

Monolithic 3D Cross-Linked Polymeric Graphene Materials and the Likes: Preparation and Their Redox Catalytic Applications

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ABSTRACT: While there have been tremendous studies about graphene and its applications in the past decade, so far the proposed huge potential of this material has not been materialized. One of the prerequisites to overcome these challenges is maintaining the nature and intrinsic properties of individual graphene sheets while in the state of bulk material. Thus, in this Perspective contribution, the fabrication/synthesis of the monolithic polymeric and three-dimensional (3D) cross-linked bulk materials (3DGraphene) with (doped) 2D graphene sheets as the building block will first be briefly summarized. Then, the second part will cover the redox catalytic application of these bulk materials including doped 3DGraphene, the graphene-like material polymeric C₃N₄ and their hybrid materials. These will include mainly oxygen reduction reactions (ORR), hydrogen evolution reactions (HER), and CO₂ reduction for their latest development. Finally, challenges and outlook related to the design of 3D crosslinked graphene based materials and their catalytic applications will be briefly discussed.

■ INTRODUCTION

Currently, the most challenging issue for truly large scale applications of graphene is preparing this material in bulk state, while simultaneously maintaining its unique two-dimensional (2D) structure, thus keeping its many excellent properties 1,2 as it has been well-known that perfect 2D graphene cannot exist in the free state.³ While this appears obvious and straightforward, it has yet to be fully realized and has been hampered mainly by the strong and intrinsic intention of restacking of graphene sheets due to strong $\pi - \pi$ interactions between the sheets.⁴ In fact, most, if not all, reported graphene bulk materials so far have such issues,⁵ and many of the wonderful properties of graphene have only been partially utilized in the bulk state.^{6,7} Clearly, these reported materials only show incremental/partial utilization of graphene's properties and miss the huge potential for some truly killer application widely expected.⁸ In this content, the bulk state and monolithic materials with some types of three-dimensional (3D) cross-linked graphene (3DGraphene) structures become the choice, where the graphene sheets must be separated in the direction (*c*-axis, out of plane) perpendicular to the graphene basal plane, and better further locked to overcome the intrinsic $\pi - \pi$ interaction forces and achieve macroscopic structural stability.^{3,9,10}

One effective strategy, but with some sacrifice, to achieve such 3DGraphene materials is to engineer a graphene material in which individual graphene sheets are bonded together at the edges with C-X bonds ($X_{i} = C_{i}$, O_{i} , S_{i} , etc., and even some molecules).^{4,11} In the proposed 3DGraphene materials to be covered in this work, graphene sheets are kept separated in the *c*-axis because of the chemical linkages at the edges and are also locked to have bulk structural integrity stability for the materials.^{12,13} Thus, in principle these bulk materials should exhibit intrinsic graphene properties at macroscopic scale. The combination of graphene properties and the material's structural stability should ensure their exciting in electronics, optical, catalytic, mechanical devices (units) and other applications.¹⁴⁻¹⁷

On the other side, among many proposed possible applications of graphene materials, one of the focuses has been on using this carbon-only material for catalytic applications.¹⁸⁻²⁰ One of the driving forces in this push is due to the issues of current state-of-art catalysts used in industry. One of such issues is that many of the most popular and efficient industrial catalysts are noble metal based materials, resulting in high costs and other issues.^{21,22} Thus, recently an increasing thrust has been emerging to develop metal-free or even carbon-only catalysts with the advantages of different morphologies, doping, and an easily tunable molecular structure as a cost-effective alternative for various photo/electro-catalytic reactions. As a typical 2D nanocarbon material, graphene based materials have attracted much attention due to its many unique and excellent properties, including its large surface area, efficient light absorption, large π conjugation system and high charge mobility, and chemical stability, all beneficial for high catalytic activity.^{23,24} As has been mentioned above, 3DGraphene materials, including heteroatom doped ones, could utilize better the advantages of graphene's many intrinsic properties at bulk state and thus are expected to demonstrate superior catalytic activity.^{25–27}

Thus, in this Perspective, we will first briefly summarize the synthesis and fabrication of these chemically cross-linked 3DGraphene materials with/out doping in bulk state, as this is the starting point for any further studies and serious applications. This include two main parts: (1) template directed methods, including chemical vapor deposition (CVD) growth and solution based template method, (2) the wet chemistry method from graphene oxide (GO) through cross coupling in solution.



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Currently, probably due to the extreme importance of redox catalytic reactions and also the matching features of graphene materials, the most studied and also important catalytic reactions using these 3D cross-linked graphene materials are oxygen reduction reactions (ORR), hydrogen evolution reactions (HER), and CO₂ reduction. Thus, these types of catalytic reactions with 3DGraphene materials will be discussed, where at least some intrinsic properties of individual graphene sheets have been utilized, particularly at the bulk state. To be complete and also for comparison, normal crosslinked 3DGraphene materials with/out doping, graphene like polymeric C₃N₄ and their hybrids will be covered. Final notes will be given for the outlook and challenges in terms of both the preparation and catalytic applications for these materials, where much more attention is required to further control the morphology, active sites, and better understand the catalytic mechanism to prompt the truly possible application of these wonderful materials.

■ FABRICATION AND PREPARATION OF 3DGRAPHENE

Even before and also after the emerging of graphene studies, there have been many theoretical proposals and studies about such 3DGraphene polymeric structures.^{28–30} Importantly, in addition to the unique electronic and mechanical properties derived from graphene,^{31,32} most of these proposed graphene based networks, including that with sp^2-sp^3 hybridized structures, are thought to be highly stable. While the ordered (crystalline) 3DGraphene materials have not been proved/ prepared completely, some materials with the proposed network structures but in the amorphous form have been reported, and indeed they demonstrate many excellent and unique properties.^{12,16,33–35}

Regarding to the synthesis/fabrication, there are mainly two types of approaches to achieve such 3DGraphene materials, as shown in Scheme 1. One is to use conventional CVD growth

Scheme 1. Schematic of Two Typical Synthetic Methods of $3DGraphene Materials^{a}$



 $a^{\prime}(I)$ The template-based method, including CVD growth or solution assembly protocol. (II) The template-free method using the wet chemistry approach by cross-linking of GO sheets.

or solution assembly with some types of 3D network templates, where the graphene network grows directly on such 3D templates. The other one is based on the wet chemistry approach without a template, using graphene oxide (GO) sheets directly and then cross-linking them together either with themselves or some linking agents. These two approaches will be summarized and discussed separately in the following sections.

CVD GROWTH OR SOLUTION ASSEMBLY ON 3D TEMPLATE FOR 3DGRAPHENE MATERIALS

While almost all CVD approaches use some types of 3D templates, there is at least one exception coming from the excellent work of the Krainyukova group.³⁶ Using a modified arc approach (but without arc discharge) where only carbon sublimation in vacuum is allowed, a very stable honeycomb carbon allotrope was obtained. High resolution TEM and other analysis results indicate that this material consists of both periodic and random Y-junction type repeating units with a honeycomb structure built exclusively from sp²-bonded carbon atoms as shown in Figure 1, and is considered as a truly



Figure 1. Proposed periodic and random honeycomb carbon structures prepared by a modified arc method from carbon rod directly. Reproduced with permission from ref 36. Copyright 2016, American Physical Society.

3DGraphene. This foam material, with an estimated density of \sim 1.48 g/cm³, demonstrates high levels of physical absorption of various gases unattainable in other carbon forms such as fullerenes or nanotubes.

Using various prefabricated/defined 3D templates and CVD growth on such 3D templates, graphene foams (GF) have been fabricated by different groups. The first and probably most important work comes from Cheng's group in 2011,³⁴ where a monolith of a continuous and interconnected foam-like graphene 3D network was obtained using the predefined 3D Ni foam template by a direct CVD growth with CH₄ as the carbon source (Figure 2A). After the growth, Ni was etched off and a free-standing GF was obtained which inherits the interconnected 3D scaffold structure of the nickel foam template. All the graphene sheets in the GF are directly bonded each other, resulting in an interconnected and monolithic flexible network of graphene. The free-standing GF is extremely light ($\sim 2-3 \text{ mg/cm}^3$) and flexible, along with a very high specific surface area (up to $850 \text{ m}^2/\text{g}$) and electrical conductivity. Similar work was soon reported afterward by Ruoff's group but with a different etchant.³⁷ With the input of N, B precursors to the carbon source and following a similar approach, N,B-codoped 3D GFs could also be made in the same way.³⁸ In 2012, Polsky's group started with an amorphous carbon structure prepared by lithographically predefined 3D pyrolyzed photoresist films, and after conformably sputtering Ni on such predefined 3D carbon structure then followed by annealing and acidic etching off the nickel, a well-defined 3D porous and multilayered graphene network could be prepared.³⁹ Cyclic voltammograms using this material as the electrode indicate that this material exhibits more favorable electron transfer kinetics than conventional glassy carbon. Yoon et al. in 2013 reported an approach to synthesize CVD-grown 3DGraphene nanonetworks (3D-GNs) that can be mass produced with large-area coverage for any arbitrary shape and on any electronic device-compatible



Figure 2. (A) Synthesis of a GF and integration with PDMS. All the scale bars are 500 μ m. Reproduced with permission from ref 34. Copyright 2011, Nature Publishing Group. (B) Schematic illustration of the fabrication process for a 3D-GN. Reproduced with permission from ref 40. Copyright 2013, Nature Publishing Group.

substrate, such as Al_2O_3 , Si, GaN, or quartz.⁴⁰ The procedure (Figure 2B) consists of four steps, with an annealing step of a PVA/iron precursor infiltrated into 3D-assembled-colloidal silica under a hydrogen environment. This reduces iron ions and generates few-layer graphene by precipitation of carbon on the iron surface and allowing the 3D-GN to be grown. Afterward, the iron/SiO₂ could be removed with HF/HCl solution, leaving few-layer 3D-GN structures on the SiO₂/Si substrate. The produced 3D-GN exhibits a high conductivity of 52 S/cm and high surface area of 1025 m²/g.

In 2013, Shen's group started with a macroporous acrylictype resin as the carbon precursor with infiltrated Ni catalyst precursor and KOH activator, followed by heating treatment at Perspective

850 °C and HCl solution washing, a 3D hierarchical porous graphene-like (3D HPG) material was prepared (Figure 3A).⁴¹ Structural and spectrum analysis confirms that the material consists of an interconnected 3D porous graphene network, arising from the thin walls of a few layers of graphene sheets. The material exhibits high surface area up to $1810 \text{ m}^2/\text{g}$, total pore volume of 1.22 cm³/g and high supercapacitor performance when used as the active materials. Using a H-type zeolite as the template and a direct CVD growth followed by HF washing of the zeolite (Figure 3B),³⁵ a cross-linked-fullerene-like framework, containing carbon hexagons and pentagons with all sp² carbon, was obtained, and the building unit of the carbon framework in each supercage of the zeolite Y has an average composition of $C_{63}H_{4,9}O_{1,2}$, very close to that of fullerene C60 cross-linked-fullerene-like framework Schwarzite carbon structure.³⁰ It also exhibits a typical semimetal band structure with zero bandgap.

Recently, a very interesting and unique approach was developed by Chen's group to prepare free-standing and high quality 3DGraphene networks.³³ The procedure started by chemically echoing out Mn in a Ni₃₀Mn₇₀ alloy to first generate a nanoporous Ni foam film, followed by a CVD process under a mixed atmosphere of H₂, Ar, and benzene. The free-standing complex 3D network comprised of interconnected graphene was then obtained by chemically removing the nanoporous Ni substrate in HCl solution. Importantly, the obtained freestanding film retains many intrinsic properties of individual graphene sheets, including the basic electronic properties of graphene, indicating the high quality of the graphene units in the structure and its potential for the practical application of graphene in 3D devices in the future.³³ In 2016, Wu and coworkers developed an ambient-pressure CVD method followed by postdoping with a solid nitrogen rich precursor graphitic- C_3N_4 to prepare a N-doped 3DGraphene foam.⁴² The 3DGraphene foam incorporated with nitrogen defects as a metal-free catalyst for CO₂ reduction shows excellent activities, and this will be discussed more in next section.

Besides CVD growth, solution templates have also be widely used to construct 3DGraphene materials through the assembly of GO sheets onto 3D templates followed by chemical reduction process. Several typical templates, such as silica,⁴³



Figure 3. (A) Synthetic steps of 3D HPG. Reproduced with permission from ref 41. Copyright 2013, Wiley-VCH. (B) Synthetic scheme for conventional ZTC consisting of a cross-linked-buckybowl-like framework. Reproduced with permission from ref 35. Copyright 2013, Elsevier.

polystyrene colloidal particles,⁴⁴ nickel foam,⁴⁵ alumina fiber blanket (AFB)⁴⁶ and even ice,⁴⁷ have been used. Note that these studies and development have been comprehensively reviewed.^{27,48}

CVD methods, along with the solution template based method, are capable of fabricating these materials with highly defined structures. This might offer more ability to control the structures including the morphology of these materials required to optimize their catalytic activity.

WET CHEMISTRY APPROACHES FOR 3DGRAPHENE MATERIALS

Another main approach to synthesize 3D cross-linked graphene materials is to use the wet chemistry method with graphene oxide as the starting material. The rich functional groups, such as OH, COOH, C==O, and even C—H at the edge on the graphene sheets, can be used to cross-link directly them together using rich organic chemistry via ester, ether or even C—C bond. Also, with rich organic chemistry, many bifunctional organic/polymeric molecules can also be used to bridge these graphene sheets together. Because such 3DGraphene materials cross-linked by other molecules have been summarized in many works,^{5,49} and these materials in general exhibit a combination of graphene and other components, this type of materials will not be covered in general in this Perspective.

For the preparation of additive-free cross-linked 3DGraphene, the earliest report should come from Shi's group, who in 2010 reported a highly electrically conducting and mechanically strong self-assembled graphene hydrogel (SGH) consisting of cross-linked graphene sheets using a hydrothermal process (Figure 4A).⁵⁰ Initially, the holding force



Figure 4. (A) Proposed formation mechanism for 3DGraphene. (Right) Typical SEM image of its interior microstructures. Reproduced with permission from ref 50. Copyright 2010, American Chemical Sociey. (B) Synthesis and structure of graphene sponges. Reproduced with permission from ref 16. Copyright 2015, Nature Publishing Group.

between the sheets for the increased stability and mechanical strength was proposed to be the $\pi-\pi$ stacking interaction between the graphene sheets. However, it is postulated that covalent cross-bonding between the sheets at the edge should be generated in such hydrothermal treatment.¹⁶ With such a highly interconnected 3D network structure, the hydrogel exhibits both remarkable electrical conductivity and mechanical robustness and thus renders it an excellent material for

flexible energy storage device and other applications.⁵¹ Additionally, the porous microstructures and mechanical properties of the generated graphene sponges could be adjusted and controlled somehow by changing the parameters of the process.⁵²

Later on, this method has been used widely and further developed. For example, when proper size of the graphene (oxide) units is used and processed in organic solvents, such as ethanol or acetone, the cross-linking between the sheets can be controlled better. Thus, a solvothermal method has been developed, and using such a method, a unique and highly elastic graphene-only 3D cross-linked graphene has been obtained (Figure 4B).^{12,16} Intensive and various structural and morphology analyses prove that a monolithic 3D chemically linked network of graphene sheets is indeed formed in such spongy material. The cross-linking is mainly due to the covalent cross-linking of the functional groups such as OH, COOH, and epoxy groups mainly located on the GO sheets edges, which generates such a monolithic 3D chemically linked graphene network. This monolithic and cross-linked graphene homogeneous material, additive-free with only graphene units, indeed exhibits many interesting and unique properties at macroscale, including zero Poisson's ratios and super elasticity in all directions and under large strain,¹⁶ and excellent organic liquid absorbing capability.⁵³

Linking the 2D graphene sheets using other molecules is another strategy to build the 3DGraphene network. One example came from Sudeep et al., where an ordered and stacked macroscopic 3D cross-linked nanoporous architecture via covalently interconnecting graphene sheets using organic molecule GAD has been reported (Figure 5).⁵⁴ The prepared 3D architectures with high hierarchical ordered morphology of uniform porosity and 3D conductive graphene scaffolds have demonstrated potential applications such as for gas storage and electrochemical electrodes.

Another interesting wet chemical approach but using a 3D template was demonstrated by Chen et al.,⁵⁵ who constructed 3DGraphene porous material (ERGO) in situ on the surface of an electrode by electrochemically reducing a concentrated graphene oxide dispersion based on the principle of graphene 3D self-assembly. Moreover, the electrochemical reduction of GO can be carried out on different types of electrode, including graphene paper, foamed nickel and platinum foil. The electrochemical reduction of GO on the graphene paper leads to the formation of a flexible graphene-based electrode in which ERGO layer tightly adheres to the substrate of graphene paper, a promising electrode for many electrochemical applications.

REDOX CATALYTIC APPLICATIONS OF 3DGRAPHENE MATERIALS

One of the most important types of catalytic reactions are the redox reactions, which all involve electron transfer. Due to the rich and large π conjugation in graphene sheets and the similarity of its building unit (benzene ring) with aromatic or conjugated compounds, it is intuitive to believe that graphene materials must have strong interactions with such substrates and can prompt electron transfer for catalytic redox reactions. With the structural stability of 3DGraphene, the large 3D network would offer extra benefits for catalytic activity due to the excellent charge mobility for electron transfer. Furthermore, heteroatoms such as N, P, B, S etc. could be incorporated into graphene sheets, offering more possible



Figure 5. Schematic of the poly-GO synthesis process from initial GO powder. (a) Photograph showing the synthesized GO powders (left). (b) The linkage of glutaraldehyde (GAD, yellow) with two different –OH groups (blue). Two different mechanisms for large-scale GO cross-linking via pure glutaraldehyde (top) and via glutaraldehyde-resorcinol (bottom). (c) The 3D networked structure of poly-GO formed via two different mechanisms. Reproduced with permission from ref 54. Copyright 2013, American Chemical Society.

active sites.⁵⁶ Together with the advantages as a 3D template to host other active materials for cocatalysts, 3DGraphene materials would therefore be promising materials for many catalytic redox reactions.

In this content, we will focus on the recent development of these materials as metal-free catalysts for the currently most studied catalytic reactions, ORR and HER, which play a vital role in the development of renewable energy sources. Because the major 3DGraphene materials used for these catalytic reactions come with some types of doped atoms, the following section will cover these doped materials first. Also, due to the extreme importance of CO_2 reduction which also belongs to redox reaction, though it has been not widely studied, it will be covered briefly in the last part of this section.

1. Oxygen Reducing Reaction. The oxygen reducing reaction, a crucial reaction occurring on the cathode to enable the fuel cell to work efficiently, plays an important role in the energy conversion devices of fuel cells and metal-air batteries.^{57–59} Along with the rapid development of carbon nanomaterials, various carbon based and metal-free electrocatalysts have been exploited, exhibiting electrocatalytic activities comparable or even better than that of the noble metal catalysts and with better stability and fuel tolerance.⁶⁰ Some recent ORR studies using 3DGraphene based materials are summarized and compared in Table 1.

Among these recently studied metal-free catalysts, materials with 3D cross-linked networks from doped graphene sheets have been widely studied as metal-free electrocatalyst for ORR due to the synergic effect combing the intrinsic nature of graphene, the unique structure and characteristics of 3DGraphene, and the extra effect from doped atoms. These include catalysts with one-type heteroatom or multitypes of doped atoms, including nitrogen, phosphorus, boron, and sulfur atoms.

In 2011, Huang's group prepared a 3D nitrogen doped carbon nanotubes/graphene (NCNTs/G) composite by pyrolysis of pyridine over a graphene sheet supported Ni catalyst.⁶¹ Tangled NCNTs with lengths of several hundred nanometers are sparsely, but tightly distributed on graphene sheets, forming quasi-aligned NCNT arrays. With 6.6 at. % N content, the NCNTs/G shows a higher activity and selectivity to ORR in alkaline electrolyte. In 2012, Qu's group fabricated an ultralight and nitrogen doped 3D cross-linked graphene framework (GF) through a hydrothermal reaction of GO aqueous suspension and pyrrole followed by a high temper-

ature annealing process, and this material consists of mainly the network of only few graphene layers, and has an ultralow density of 2.1 \pm 0.3 mg/cm^{3.62} Besides the remarkable adsorption capacity and excellent capacitor performance, the macroscopic monolithic material with a tunable hierarchical morphology and high specific surface area shows extremely high activities for electrocatalytic ORR. A slightly lower onset potential of ORR on GF (ca. -0.18 V) and notably higher diffusion current density over a large potential range (-0.4 to)-0.8 V) than that of Pt/C electrodes, as well as excellent tolerance of the GF electrode to the crossover effect were observed. The excellent performance is attributed to the unique 3D pore rich structure with maximum access to the nitrogen sites within highly exposed graphene sheets and multidimensional electron transport pathways. In 2015, nitrogen doped 3D macroporous graphene (N-MGPPy) and porous graphene foams (PNGFs) were prepared by electrochemically and hard templating approaches, respectively.^{63,0} Both of them exhibit excellent catalytic activities for ORR owing to the 3D porous structures and the nitrogen active sites of the electrode materials.

Compared with single-atom doped cases, the biatom or codoped catalysts can deliver better catalytic activity due to the synergetic effect of different dopants. One excellent work came from the Dai group's systematic investigation for a threedimensional nitrogen and phosphorus codoped mesoporous carbon foams (NPMC) used for Zn-air battery. The material was fabricated using a scalable, one-step process involving the pyrolysis of a polyaniline aerogel synthesized in the presence of phytic acid (Figure 6a-d), and exhibits outstanding catalytic activities for both oxygen reduction and oxygen evolution reactions.⁶⁵ The electrochemical performance of the N,Pdoped carbon foam as an air electrode for both primary and rechargeable Zn-air batteries was also investigated. Typically, an open-circuit potential of 1.48 V, a specific capacity of 735 mAh/g_{Zn} (corresponding to an energy density of 835 Wh/ kg_{7n}), a peak power density of 55 mW/cm², and stable operation for 240 h with mechanical recharging in primary batteries were obtained in an aqueous KOH electrolyte. A three-electrode rechargeable battery built with two NPMC metal-free air electrodes to have ORR and OER separately also demonstrated good stability (600 cycles for 100 h of operation). The carbon foam electrode material for both OER and ORR in a three-electrode configuration shows remarkable activity with a positive onset potential of 0.94 V

Table 1. Sum Performance :	mary and Con and the Activit	parison of Some Recent Reported Typical 3DGraphene Based Materials for ORR y Origin	in Terms of Their Structural Characteristic, Cata
3DGraphene based material	Structural char- acteristics	Catalytic performance	Proposed main (high) activity origin
3D NCNTs/G	N doping	$E_{\rm owert}\sim-0.191$ V vs SCE in alkaline electrolyte, slightly lower than that of the Pt/C; diffusion current close to that of the Pt/C	Nitrogen doping
3D GF	N doping	$E_{onset} \sim -0.18$ V vs SCE in alkaline electrolyte, slightly lower than that of the Pt/C; higher diffusion current density than that of Pt/C over a large potential range from -0.4 to -0.8 V	3D pore rich structure with maximum access to the N sites within exposed graphene sheets and multidimensional electron transport pathways
PNGFs	N doping	ORR activity and long-term stability in both alkaline and acidic solutions; E_{usset} of 1.02 V vs RHE in alkaline solution (diffusion current density of $c_2 = 7 \mod 20$ and 0.83 V in acidic modu	N doping, large specific surface area and diversified porosity

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ref.	61	62	63	64	65	58	38	66	67	68	69	20	71	72	73	74	75
Proposed main (high) activity origin	Nitrogen doping	3D pore rich structure with maximum access to the N sites within highly exposed graphene sheets and multidimensional electron transport pathways	N doping, large specific surface area and diversified porosity	High SSA and conductivity of graphene network; high mass transport rate owing to the 3D porous structure	Synergistic effects of N, P codoping and graphene edge effects for the catalytic activities toward both OER and ORR	Synergistic effects of N _i P codoping; 3D porous graphitic network for the electron and electrolyte transports and the stabilities	Doped N and B atoms act as active sites; interaction between adjacent N and B atoms reduces the bandgap energy; also the increased electroactive surface area	Large SSA, 3D interconnected framework; macro/mesoporosity structures ensuring accessible porosity and effective mass transport; high nitrogen and boron content, dominant active pyridinic and graphitic nitrogen and BC3 species; synergistic effect of N and B	N and S codoping	Edge effect and heteroatom synergistic effect	High amount (74.8 at. %) of pyridinic N and graphitic N; high amount (79.8 at. %) of active thiophene-S; high specific surface (391.9 m ² /g) area and multiporous structure	Synergistic coupling of N and S resulting in more exposed "active sites" in graphene frame	Increased SSA, hierarchical porous structure, and $\mathrm{sp}^3\text{-amide}$ linkage between rGO layers	Well interconnected and porous; high SSA and fast electron transfer	$g\text{-}C_3N_4$ nanosheets as active centers; rGO sheets as "highways" for electron transport; an efficient 3D electron transport network	Synergistic effect of the carbon network and B—N, resulting into active centers for ORR kinetics	Synergistic effect between broken C=O bonds and the unique porous structure of 3DGraphene
Catalytic performance	$E_{\rm overt} \sim -0.191$ V vs SCE in alkaline electrolyte, slightly lower than that of the Pt/C; diffusion current close to that of the Pt/C	$E_{\rm onset} \sim -0.18$ V vs SCE in alkaline electrolyte, slightly lower than that of the Pt/C; higher diffusion current density than that of Pt/C over a large potential range from -0.4 to -0.8 V	ORR activity and long-term stability in both alkaline and acidic solutions; E_{onset} of 1.02 V vs RHE in alkaline solution (diffusion current density of ca. 7 mA/cm ²) and 0.83 V in acidic media	$E_{\rm ouset}$ of -0.132 V vs Ag/AgCl in alkaline electrolyte; a current density value of 5.56 mA/cm ²	High activities toward both ORR and OER in alkaline electrolyte. For ORR, $E_{\rm onset}$ of 0.94 V vs RHE and a half- wave potential of 0.85 V in alkaline solution; For Zn–air battery based on NPMC, an open-circuit potential of 1.48 V, a specific capacity of 735 mAh/ $g_{\rm Zn}$ (corresponding to an energy density of 835 Wh/ $kg_{\rm Zn}$), a peak power density of 55 mW/cm ² , and stable operation for 240 h	High activities toward both HER and ORR. For ORR, a comparable E_{onset} and kinetic current density to that of Pt/C. For Zn-air battery, a high peak power density of 310 W/g and an excellent durability without potential drop over 90 h	E_{onset} of ca. -0.16 V vs SCE in alkaline electrolyte; a current density of 7.8 mA/cm ² at -0.03 V; excellent stability with 89% current retention after 20000 s	E_{onset} of -0.07 V vs SCE in alkaline electrolyte; a higher diffusion current density	A current density of 3.90 mA/cm ² in alkaline electrolyte; excellent stability with current density retention of 85.2% over 20000 s	A current density of about 0.17 mA/cm² at a voltage of 0.7 V in alkaline electrolyte	$E_{\rm onset}$ of 0.895 V vs RHE in alkaline electrolyte; diffusion-limiting current of 5.23 mA/cm ² at 0.2 V	$E_{ m onset}$ of $-0.1~ m V$ vs Ag/AgCl in alkaline electrolyte; kinetic current density of 26.1 mA/cm ²	E_{oust} of -0.1 V vs Ag/AgCl in alkaline electrolyte; higher limiting diffusion current density than that of Pt/C electrode at large potential range	An increased peak current compared with the GNR modified electrodes; a slight change in the onset potential	$E_{\rm overt}$ of -0.21 V vs Ag/AgCl in alkaline electrolyte; a wave current enhanced by 1.6 and 2.3 times compared with g- C_3N_4 and rGO; superior durability over Pt/C catalyst	E_{ousci} of 0.798 V vs RHE in alkaline conditions; a current density of 3 mA/cm ² ; higher stability than commercial Pt/C catalysts even after 10000 cycles	$E_{ m onset}$ close to that of Pt/C electrode; higher stability than that of Pt/C electrode
acteristics	N doping	N doping	N doping	N doping	N and P codop- ing	N and P codop- ing	N and B co- doping	N and B co- doping	N and S codop- ing	N and S codop- ing	N and S codop- ing	N and S codop. ing	cross-linking via covalent amide	cross-linking via glutaraldehyde	cross-linking of g - C_3N_4 and GO	wrapping of BN sheets into graphene net- work	without any other foreign atoms or doped agents
based material	3D NCNTs/G	3D GF	PNGFs	3D nN-MGPPy	3D NPMC	3D MPSA/GO	3D GF	3D GF	3D N/S-GFs	3D GF	NS-3DrGO	3D NSGAs	3D GF	3D GNRs	3D g-C ₃ N ₄ /rGO	3D rGO/BN	3DG



Figure 6. Preparation of N,P-codoped porous carbon (NPMC) electrocatalysts of (a) Schematic illustration of the preparation process for NPMC foams. (b,c) SEM images of PANi aerogel (b) and NPMC-1000 (c). Inset in panel c: Digital photo images of PANi aerogel before (left) and after (right) pyrolysis at 1,000 °C. (d) High-resolution TEM image of NPMC-1000. Electrocatalytic activity for ORR and OER of (e) Linear scan voltammogram (LSV) curves for NPMC samples prepared under different temperatures and commercial Pt/C catalyst at an RDE (1,600 r.p.m.) in O₂-saturated 0.1 M KOH solution. Scan rate, 5 mV/s, and (f) LSV curves of NPMC-1000, NPMC-1100, RuO₂ and commercial Pt/C catalyst on an RDE (1,600 r.p.m.) in 0.1 M KOH (scan rate, 5 mV/s). Reproduced with permission from ref 65. Copyright 2015, Nature Publishing Group.

versus reversible hydrogen electrode (RHE) and a half-wave potential of 0.85 V versus RHE, which are comparable to those of Pt/C (Figure 6e,f). Density functional theory (DFT) calculations, together with experimental results, indicate that the excellent catalytic activity came from the synergistic effect of the N,P codoping and highly porous graphene network and the main active sites are located at the doped edge positions of graphene network. Later on, they prepared another N,Pcodoped 3D porous carbon network (MPSA/GO) used as the catalysts for ORR/HER through a template free approach by pyrolysis of a supermolecular aggregate of self-assembled melamine, phytic acid, and graphene oxide. Zn–air batteries based on this MPSA/GO air electrode also show a high peak power density (310 W/g) and excellent durability.⁵⁸

Besides N,P-codoped materials, N,B-codoped 3DGraphene has also attracted a lot of attention. In 2013, a 3DGraphene foam codoped with nitrogen (4.5 atom %) and boron (3 atom %) (BN-GFs) was prepared by a CVD method using melamine diborate as the precursor,³⁸ and it shows significantly better electrocatalytic activity toward ORR than the undoped and B or N monoheteroatom doped foams, even slightly better than

the compared Pt-C/GC electrode in 0.1 M KOH aqueous solution, including higher reduction peak current, more positive onset potential and higher stability. The reaction kinetics studies indicate that the number of electrons transferred per oxygen molecule is 3.4-3.8 at the potential range from -0.2 to -0.5 V. The excellent catalytic activities were contributed to the following reasons: (1) both the isolated N and B atom sites can act as active sites for ORR through charge transfer with neighboring C atoms; (2) interaction between adjacent N and B atoms could reduce the bandgap energy to further facilitate the ORR performance of the BN-GF electrode; (3) B and N codoping could also enhance the electroactive surface area significantly. Then in 2015, Xu et al. also demonstrated similar results, where based on the synergistic coupling of the B and N dopants within the graphene domains, the dual 3D N/B-doped graphene aerogel (foam) (N,B-GA) prepared by a hydrothermal reaction and a pyrolysis procedure showed superior catalytic activity for the ORR than that of the single N- or B-doped materials.⁶⁶ Nitrogen and sulfur could also act together as the codoped heteroatoms to 3DGraphene for catalytic purpose materials. In



Figure 7. (A) Crystal and electronic structure of graphitic carbon nitride $(g-C_3N_4)$ (a) Schematic diagram of a perfect graphitic carbon nitride sheet constructed from melem units. (b) DFT band structure for polymeric melon calculated along the chain ($\Gamma-X$ direction) and perpendicular to the chain ($Y-\Gamma$ direction). Reproduced with permission from ref 77. Copyright 2009, Nature Publishing Group. (B) Characterization of the physical structure and photocatalytic water-splitting performance of the CDots- C_3N_4 composite catalyst. (a) A TEM image and (b) magnified TEM image of the CDots- C_3N_4 region of panel a marked in red. (c) Typical time course of H_2 and O_2 production from water under visible light irradiation. Reproduced with permission from ref 81. Copyright 2015, American Association for the Advancement of Science. (C) The performance of catalyst. (a) Photocatalytic activity for H_2 production, (b) DRS spectrum, (c) PL emission spectra, and (d) EIS plot and photocurrent–time dependence of PCNM (solid line) and the g-CN powder (dotted line). Reproduced with permission from ref 82. Copyright 2015, Wiley-VCH.

2013, Feng's group reported nitrogen/sulfur-codoped 3DGraphene frameworks (N/S-GFs) prepared by a one-pot hydro-thermal approach employing graphene oxide and ammonium thiocyanate as the precursors, and achieved an excellent catalytic behavior for ORR in alkaline condition.⁶⁷ Later,

similar works have been reported also by several other groups. $^{68-70}$

Besides doped 3DGraphene materials, some 3DGraphene prepared by cross-linking with organic small molecules have been also investigated for catalytic purposes. For example, Kim's group reported a novel 3DGraphene material prepared through covalent amidation reaction using diamine compounds and has studied them as a metal-free electrocatalyst for ORR in alkaline media.⁷¹ It has been proposed that the linking compounds used as a junction between functionalized rGO layers improve the electrical conductivity and impart electrocatalytic activity for ORR due to the facilitated interlayer charge transfer. Investigation of the ORR mechanism reveals that higher pyridinic-like N content and π -electron interactionfree interlayer charge transfer contributed to their higher catalytic activities and stability than that of the compared materials prepared from nonfunctionalized rGO. Similarly, using glutaraldehyde as the cross-linking agent, a 3D crosslinked graphene based electrocatalytic catalyst with high activity was assembled by individual 2D graphene nanoribbons.⁷² With ultrathin g-C₃N₄ nanosheets and hexagonal boron nitride as additional agents respectively, 3DGraphene based composites have been synthesized and delivered outstanding catalytic activities for ORR owing to the synergistic effects of the graphene network and the additional agents resulting into active centers.^{73,74}

In the works discussed above, where doped 3DGraphene or 3DGraphene hybrids with other additional components have been studied for their catalytic activity, the active sites of these catalysts come mainly from the foreign atoms (sites) doped on the intrinsic atoms (sites) of graphene. As has been noted and discussed above, when other agents are introduced to graphene, its intrinsic properties will be largely impaired and cannot be fully utilized. So, it would be highly interesting to study the catalytic activity of pristine 3DGraphene without any doped or hybrid agents.

Based on this, Zhang et al. have studied thermally treated 3DGraphene prepared directly by heating GO without any other foreign atoms or doped agents as a highly efficient metalfree electrocatalyst toward ORR.75 Interestingly, they pointed out that the breaking up of the initial C=O bonds in the 3DGraphene materials during the high temperature treatment (600 °C) would leave vacant sites at the original C=O positions. The synergetic effect between broken C=O bonds on 3DGraphene and its unique porous structure generated competitive catalytic activity, higher selectivity and better stability toward ORR compared with that of commercial Pt/C. While it has not been fully studied, it is postulated that the observed catalytic activity should come mainly from the defect (vacant) sites generated by the decomposition of some functional groups including C=O as the authors indicated. Also, the unique porous structure of 3DGraphene materials are able to trap oxygen molecules more efficiently, contributing to a decrease in the diffusion resistance and enhancing electrolyte-electrode accessibility for fast mass transport. Note that the catalytic studies for pristine 3DGraphene are very limited so far and the measured catalytic activity is still lower than some doped 3DGraphene.^{38,64} Thus, many more studies for this type of graphene material are needed to fully capitalize their catalytic potential, including controlling and optimizing defect and vacant sites, which is believed to be the main source of the catalytic activity.

2. Hydrogen Evolution Reaction. The hydrogen evolution reaction is a vital catalytic reaction that could be conducted by either photo- or electro-catalysis, and both approaches have been intensively studied in the past few decades using various carbonaceous (metal-free) materials, including recently with 3D cross-linked bulk graphene based material and the semiconductor but graphene-like polymeric

3D cross-linked g-C₃N₄. The efficient light absorption, tunable and no-zero but tunable band gap and its 3D conjugation network makes these materials particularly attractive for photocatalytic HER. Thus, in this section photocatalytic HER will be covered first, followed by electrochemical HER using these materials.

2.1. Photocatalysis. The intrinsic zero bandgap of pristine graphene has been limiting its direct application as a photocatalyst. Thus, the most commonly reported graphene based photocatalysts come from either heteroatom (e.g., N, S, B, P) doped graphene, chemically modified graphene materials (e.g., GO and reduced GO) or hybrid/composites where graphene is used as an electron acceptor or conductor and is combined with other semiconductors.⁷⁶ Among the metal-free 3D cross-linked carbon based catalysts, the graphene-like polymeric material g-C₃N₄ is probably the most studied. Thus, we will cover this material first.

Early in 2009, Wang and colleagues reported the catalytic hydrogen production activity from water under visible-light irradiation using old polymeric carbon nitrides (g-C₃N₄) for the first time as a metal-free photocatalyst.77 With their impressive optical properties, electronic structures as well as a 2.7 eV bandgap, the polymeric material exhibits an intrinsic semiconductor-like absorption with energetically possibility for water splitting (Figure 7A). The $g-C_3N_4$ achieved steady H_2 production with a rate of 0.1–0.4 μ mol/h from water and triethanolamine as a sacrificial electron donor on light illumination (>420 nm) in the absence of noble metal catalysts. Comparing the activity results of different materials prepared under different temperatures, they pointed out that some defect structures are crucial, which might facilitate the charge localization at specific surface termination sites where they can be transferred to the water molecules.

Although intensive studies of polymeric g-C₃N₄ as a metalfree catalyst for water splitting have been reported following this pioneering work,^{78,79} the reactions still suffer from a low hydrogen production and quantum efficiency (QE) and also require a sacrificial reagent.⁸⁰ Until 2015, Lee's group constructed carbon a dots/g-C₃N₄ composite (CDots-C₃N₄) photocatalyst for water splitting under visible light.⁸¹ The lowcost and environmentally benign materials can achieve QEs of 16% to split water into H_2 and O_2 at a wavelength of 420 nm, and also demonstrated a robust stability in 200 runs of recycling use over 200 days. A 2.0% of overall solar energy conversion efficiency, at least 1 order of magnitude larger than previous reports for any stable water splitting photocatalyst, was achieved (Figure 7B). Different from the one-step fourelectron mechanism of conventional photocatalysis, the impressive performance of this composite catalyst is believed to have a stepwise two-electron/two-electron two-step pathway, where H_2O oxidation to H_2O_2 is the first and rate limiting step (photocatalysis), and for the second and fast step (chemical catalysis) CDots are responsible for decomposing H_2O_2 to O_2 and finishing the entire catalytic cycle.

Following the exciting development of the C_3N_4 catalyst above, a macroscopic 3D porous $g-C_3N_4$ monolith (PCNM) consisted of 2D porous C_3N_4 nanosheets was synthesized by Yang's group through one-step thermal polymerization of urea inside the framework of melamine sponge (MS).⁸² With a large specific surface area (SSA), multiple junctions and open pores, the unique 3D interconnected network of PCNM exhibits significantly improved photocatalytic performance for hydrogen evolution under visible light compared to the powder

Perspective



Figure 8. Photocatalytic hydrogen evolution activities from water splitting. (a) Production rate over a series of 3DG samples synthesized at various annealing temperatures. (b) Pathways of electron transfer and mechanism of photocatalytic hydrogen production catalyzed by 3DG. Reproduced with permission from ref 83. Copyright 2017, Springer Nature.

counterpart catalyst (Figure 7C-a). The significantly enhanced catalytic activity compared to that of the powder $g-C_3N_4$ is postulated due to the following reasons: (1) larger SSA; (2) more efficient light absorption light (Figure 7C-b); (3) better separation of photogenerated charge carriers (Figure 7C-c) and faster interface charge transport (Figure 7C-d).

While it is rare, additive/doping free 3DGraphene has also been studied for its photocatalytic HER. We have exploited such a monolithic homogeneous and cross-linked 3DGraphene material for photocatalytic HER which followed a very unique mechanism.⁸³ The combination of its unique band structure of graphene and macroscopic morphology of this material ensures an efficient light absorption of graphene and an easily achievable reverse saturation state under light illumination.¹² All these together makes it capable of ejecting efficiently hot electrons under visible illumination and thus capable for wide redox chemical conversion. Thus, for the first time, a carbononly 3DGraphene catalyst for efficient water splitting to generate H₂ has been achieved under visible light. This material exhibits a remarkable hydrogen production rate of 270 μ mol/h/g_{cat} under full-spectrum light (Figure 8) following a light-induced ejected hot-electron relaxing mechanism.⁸³

2.2. Electrocatalysis. For electrocatalysis HER, the highly conductive cross-linked network existing in 3DGraphene is expected to facilitate exciton charge separation and also charge transportation during the process of HER. Furthermore, the rich macropores in these 3D structures should provide a facilitated electrolyte accessing to the inner structure of the catalysts and improve overall the HER kinetics.²⁷

By rationally engineering the macroscopic architecture of graphene and its chemical/defective structures, Tian et al. designed a 3D N-doped and plasma-etched graphene (3DNG-P) and used it as a HER catalyst.⁸⁴ Combined the merits of freestanding 3D porous architecture, high-level N-doping and plasma-induced enriched defects, the prepared 3DNG-P leads to an excellent electrocatalytic performance for HER with both highly enhanced activity with a low overpotential of 128 mV at 10 mA/cm^2 and stability over a wide pH range. The authors proposed that the enhanced catalytic performance is attributed to the following reasons: (1) the 3D architecture affords efficient electron/ion-transport pathways to lower the HER polarization resistance; (2) both N dopants and plasmainduced defects offer more active centers for HER; and (3) the coupling of N dopants with graphene defects produces a synergistic effect in further enhancing the HER activity of the active centers.

Combing doped graphene and carbon nitride, a metal-free electrocatalyst ($CN_x@N$ -RGO) has been prepared by annealing GO-soaked melamine foam, which produced a three-dimensional nanostructure consisting of nitrogen-doped graphene (N-RGO) layers distributed on the interconnected arms of carbon nitride (CN_x) tetrapods (Figure 9).⁸⁵ As an



Figure 9. (a) Pictorial representation illustrating the preparation of the $CN_x@N$ -RGO catalyst using melamine foam and GO. (b,c) SEM images representing an interconnected 3D network of $CN_x@N$ -RGO at different magnifications. (d) Tetrapod structure of $CN_x@N$ RGO; (e) single arm of the tetrapod of $CN_x@N$ -RGO, showing the wrapping of the arm by the N-RGO sheet. Reproduced with permission from ref 85. Copyright 2017, Wiley-VCH.

electrocatalyst for HER, it shows rather low overpotential of 193 mV to reach a high current density of 10 mA/cm² and low Tafel slope value of 54 mV/dec, following a Volmer–Heyrovsky mechanism. The study shows that the structural modifications and spatial rearrangement of N-RGO on CN_x provides a high density of in-plane nitrogen coordinated active centers, whereas the 3D macroporous morphology assists efficient reactant/product distribution. Furthermore, DFT calculations show that the synergy between CN_x and N-RGO facilitates good electrical coupling between the two moieties and provides optimal binding to H⁺ ions on the catalyst, which in turn results in efficient reduction of hydrogen ions.



Figure 10. Morphology of (a) A SEM image showing a representative graphene foam with an open frame structure. The electrocatalytic activities of N-doped graphene foam of (b) Faradaic efficiency of CO versus potential and (c) Maximum Faradaic efficiency of CO and its corresponding potential versus N functionality content. DFT modeling of electrocatalysis of CO_2 on NG of (d) Free energy diagram of electrochemical reduction of CO_2 to CO on NG and (e) schematic of N configuration and CO_2 reduction pathway. Reproduced with permission from ref 42. Copyright 2016, American Chemical Society.

3. CO₂ Reduction. As a visible-light responsive photocatalyst for CO₂ reduction, Tong et al. designed a 3D porous g-C₃N₄/graphene oxide aerogel (CNGA) by a hydrothermal coassembly of 2D g-C₃N₄ and GO nanosheets, in which g-C₃N₄ acts as an efficient photocatalyst, and GO supports the 3D framework and also promotes the electron transfer simultaneously.⁸⁶ Excellent photocatalytic performance with a high yield of CO of 23 mmol/g (within 6 h) from CO_2 reduction was achieved. The mechanism studies indicate that the enhanced photocatalytic activity of CNGA is due to the synergetic effect of both graphene and C_3N_4 moieties: (1) the photogenerated electrons from g-C₃N₄ can be transferred to graphene network due to their large coherent interface, thus the radiative recombination of electron-hole pairs was significantly inhibited; (2) the 3D porous aerogel structure of CNGA is beneficial to the visible-light absorption and promoting more electron-hole pairs; (3) the high adsorption capacity and rapid mass transport due to the 3D porous framework are kinetically favorable for the surface chemical reaction.

In 2015, Ajayan's group prepared a N-doped 3DGraphene foam through ambient-pressure CVD method followed by postdoping with a solid nitrogen rich precursor graphitic- C_3N_4 and studied this material for CO₂ reduction.⁴² The 3DGraphene foam incorporated with nitrogen defects as a metal-free catalyst for CO₂ reduction shows a negligible onset overpotential (-0.19 V) for CO₂ reduction to CO formation, and exhibits superior activity over Au and Ag, achieving similar maximum Faradaic efficiency for CO production (~85%) at a lower overpotential (-0.47 V) and better stability for at least 5 h. Furthermore, both theoretical calculations and experimental results suggest pyridinic-N is the most active site for CO₂ reduction. The structure and catalytic performance is shown in Figure 10.

CHALLENGE AND OUTLOOK

For the synthesis part, bulk and 3D cross-linked graphene materials and the likes such as $g-C_3N_4$ with monolithic 3D cross-linked networks of the 2D units, including the doped and hybrid ones, are expected to offer a much better position to achieve the full potential of these 2D materials at the bulk state, and thus deserve much more attention and systematic studies. So far, all these prepared materials have neither

uniform building units nor ordered structure. Thus, the truly challenging and extremely important task is to have some ordered 3D cross-linked structure with uniform graphene sheets as the building unit. These studies might also offer some exciting opportunities to win the war to fully realize the potential of graphene and bring the truly killer application of graphene to table.

Among all the proposed applications, owing to the combination of the excellent intrinsic properties of graphene and the 3D porous structures, 3D cross-linked graphene materials, including the doped and hybrid ones with $g-C_3N_4$, have demonstrated exciting catalytic potential compared to traditional metal based catalysts. The high or superior catalytic activity arise generally from the synergetic effect of the combination of several characteristics of these 3DGraphene or the like materials, including surface functional groups, doped heteroatoms and vacancies as the active sites, the collective properties of individual graphene sheet, strong adsorption and preabsorption of reactants, the large 3D and porous crosslinked conjugation network for efficient charge separation and transportation, high chemical and thermal stability, and etc. To achieve better/higher activity, the control of doping and vacant sites and their distribution and band structure is probably the first issue to address. Note since most of such materials so far do not have an ordered structure, and their exact structure is still unresolved, even the reproducibility of the preparation of these materials and their catalytic activity is another big issue. This again prompts the challenging task to have some ordered or crystalline 3D structure with graphene sheets as the building unit as mentioned above. Overcoming such a challenge will not only address many issues mentioned above for catalytic applications but should also have important implications for other graphene related applications. Though there have been many studies so far, the identification of the exact active location and nature of the catalytic active centers, and mechanism of catalysis are far from clear. All these issues require much more systematic studies before we can possibly utilize the important properties of 3DGraphene or the like materials for the extreme important catalytic application.

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Notes

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