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Precise graphitic nitrogen-incorporation by electrochemical oxidation

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Graphitic nitrogen (graphitic-N) plays a very important role in energy conversion and environmental protection. Although various synthesis methods have been developed, complex devices and harsh conditions are often needed, causing difficulty in flexible regulation. Electrochemical approaches are attracting increasing attention due to their mild reaction conditions, controllability, and environmental compatibility. However, precisely incorporating graphitic-N remains a significant challenge. In this study, a synthesis strategy is designed that creates carbon single vacancies *via* electrochemical oxidation and then incorporates N radicals to construct graphitic-N. Graphite paper doped with exclusively graphitic-N was achieved by using ammonium ions as the nitrogen source. By integrating multiple *operando* electrochemical characterization techniques and density functional theory calculations, the crucial regulation parameters were clarified, and the proposed doping mechanism was validated. Hydroxyl radicals generated from electrochemical water dissociation performed three functions: involving the formation of (i) carbon single vacancies and (ii) adjacent oxygen-containing functional groups, and (iii) activation of ammonium ions into N radicals. Ketones exhibited better thermodynamic behavior than hydroxyl species when assembling N radicals into carbon single vacancies. The findings offer both experimental and theoretical foundations for a deeper understanding of the structure–property relationships of graphitic-N and broaden the application prospects of graphitic-N-doped materials.

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Introduction

Graphitic-N denotes nitrogen atoms replacing sp^2 -hybridized carbon atoms in the hexagonal lattice of graphene. This involves typical in-plane nitrogen doping, where the radius of the nitrogen (N) atom (0.74 Å) is close to that of a carbon (C) atom (0.77 Å), thus preserving the coplanarity and conjugated structure of the carbon framework.^{1–3} Owing to its higher electronegativity (3.04) relative to carbon (2.55), nitrogen doping—particularly in the form of graphitic-N—alters the local electronic environment within the carbon matrix. Graphitic-N, which acts as an electron-donating species, induces an upward shift of the Fermi level.^{4–6} This electronic modulation enhances interfacial interactions governed by Lewis acid–base

sites and π – π stacking, thereby improving the material's reactivity in surface-related processes.⁷ For example, graphitic-N demonstrates a specific adsorption location for peroxide by forming a metastable intermediate with the carbon plane.⁸ It paves the way for the development of non-radical advanced oxide processes based on direct electron transfer, exhibiting stronger selectivity for electron-rich emerging contaminants.^{9–11} In addition, graphitic-N doped carbon cathodes display metal-like catalytic activity for the four-electron oxygen reduction reaction (ORR), and offer superior resistance to methanol crossover.^{12,13} This shows that carbon materials are promising alternatives for non-metal cathodes in fuel cells.

A variety of physico-chemical methods for graphitic-N synthesis have been reported, including chemical vapor deposition,^{14,15} thermal methods,^{10,11,16} and plasma methods.¹⁷ However, these approaches typically require harsh reaction conditions and complex devices, involving high temperatures, high vacuum, toxic or explosive reagents, and extended processing time. In contrast, electrochemical methods are receiving growing attention, relying on their outstanding advantages in terms of mild reaction conditions, strong controllability and greater potential for industrial production. Common electrochemical techniques such as chronopotentiometry, voltammetry, and chronoamperometry have been adopted to dope nitrogen atoms into graphite-based

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materials under normal temperature and pressure. Multiple readily available nitrogen sources, including inorganic $\text{NH}_4^+/\text{NO}_3^-$ ^{18–23} and organic compounds (e.g., ionic liquids,²⁴ glycine,²⁵ melamine²⁶), have been employed in those studies.

Although nitrogen doping has been successfully realized in electrochemical systems, mixed co-doping of pyridinic, pyrrolic, and graphitic nitrogen was generally obtained. Graphitic-N is a typical n-type dopant, which can effectively improve the carrier density and conductivity of carbon materials. Therefore, it represents a favorable structure modification for the graphite electrodes used in electrochemical energy-storage applications. However, pyridinic-N and pyrrolic-N act as p-type dopants, which tend to induce carrier scattering and increase resistance, thereby hindering conductivity-oriented device performance. Therefore, this work aims to incorporate graphitic-N into the carbon framework precisely. In addition, in the current research, the synergistic interaction between the electrolysis mode and nitrogen species is still unclear. In particular, real time control over the site-specific evolution mechanisms involved in the graphitic-N formation remains a challenge, thereby limiting the tunability of the graphitic-N content.

Given the apparent structure of graphitic-N, *i.e.* a nitrogen atom substituting for one $\text{sp}^2\text{-C}$ atom, we designed a controllable synthesis strategy consisting of vacancy-making and nitrogen-filling. Specifically, carbon single vacancies (SVs) are first created on a graphite topological plane *via* electrochemical oxidation. Then, N radicals are produced by electrochemically activating the nitrogen-containing electrolyte to occupy these SVs. The inorganic ammonium cation is a typical nitrogen source with advantages including low toxicity and good shelf stability, compared with organic amine cations and nitrate anions. In addition to our previous research on electrochemical ammonia oxidation,^{27,28} in this work, the electrooxidation of $(\text{NH}_4)_2\text{SO}_4$ was adopted to obtain N radicals. Crucial electrochemical parameters, which enable modulation of the concentrations of SVs and N radicals, were systematically examined. Advanced *operando* characterization techniques—including surface-enhanced Raman spectroscopy (SERS), electron paramagnetic resonance (EPR), differential electrochemical mass spectrometry (DEMS), and density functional theory (DFT) calculations—were utilized for comparisons. The results confirmed the feasibility of our strategy, and a sample with 100% relative graphitic-N doping was achieved. The doping mechanism and tuning approach were expounded in detail. This work provides a platform for systematically investigating the structure–property relationships between graphitic-N content and material performance, offering theoretical insights for optimizing catalysts, electrode materials, and sensors.

Compared with traditional graphitic-N doping techniques, such as CVD, high-temperature annealing, and plasma treatment, our electrochemical strategy—consisting of vacancy-making and nitrogen-filling—exhibits notable advantages. This method operates under mild conditions (room temperature and pressure) and requires simple instrumentation and nontoxic reagents. By first generating carbon vacancies and subsequently introducing nitrogen radicals for selective filling,

nitrogen atoms can directly substitute $\text{sp}^2\text{-C}$ sites, enabling spatially localized and configuration-oriented doping. This effectively suppresses the co-formation of pyridinic-N and pyrrolic-N, allowing the generation of graphitic-N with a much higher phase purity than conventional approaches. In addition, doping parameters, including potential, current density, pulse design, and electrolyte composition, can be tuned in real time, allowing programmable modulation of vacancy density and nitrogen radical concentration. This offers precise control over both doping level and nitrogen configuration. Furthermore, this electrochemical environment is compatible with *operando* characterization (e.g. SERS, EPR, DEMS, and DFT), enabling direct tracking of intermediate species and bonding evolution. Such capability strengthens mechanistic understanding and establishes a clearer structure–function correlation. Owing to its universality toward diverse carbon frameworks and scalable process conditions, this strategy shows strong potential for industrial application. In summary, the method provides a precise, green, and scalable route for high-graphitic-N incorporation, offering advantages in both incorporation precision and mechanistic controllability.

Results and discussion

Synthesis and characterization

A strip of graphite paper (1.5 cm × 0.8 cm × 0.02 mm) was used as the working electrode (Fig. 1a), and platinum wire and Ag/AgCl were used as the counter electrode and reference electrode, respectively, to construct a three-electrode cell. In the vacancy-making step, concentrated sulfuric acid (50 wt%) was selected as the electrolyte, and a constant potential of 1.8 V was applied. The oxidation period should be less than 30 min, or else the skeleton of the graphite paper will be damaged and broken into small fragments. Owing to the SO_4^{2-} intercalation as well as violent H_2O dissociation, lots of reactive oxidative radicals immediately grow on the outer and inner surfaces of the graphite layers.^{29,30} They may contribute to the rapid formation of oxygen-containing functional groups (e.g. hydroxyl –OH, carbonyl –C=O, and carboxyl –COOH) on the plane and edges and even the expected SV.^{31–33} The sample obtained in this step is labeled as electrochemical graphene oxide (EGO) (Fig. 1b). In the nitrogen-filling step, the electrolyte was replaced by 1.25 M $(\text{NH}_4)_2\text{SO}_4$ aqueous solution, and an oxidation potential lower than 1.8 V was applied for 30 min. It is assumed that the nitrogen atom in the NH_4^+ can be dehydrogenated and precisely filled into the SV as graphitic-N, producing g-ENGO, as shown in Fig. 1c.

By comparing the optical photos, it can be seen that the shape of the graphite paper remained unchanged, suggesting its convenience when working as an electrode. However, the photos exhibit significant differences in the surface roughness. According to the atomic force microscopy (AFM) measurements (Fig. 1d), the graphite surface was relatively smooth with a roughness (Ra) of ≈ 0 nm. After electrochemical oxidation, a few “stalactite”-like protrusions appeared on the surface of the EGO, corresponding to an increased roughness (Ra) of ≈ 300 nm. When the nitrogen atoms were assembled in the g-



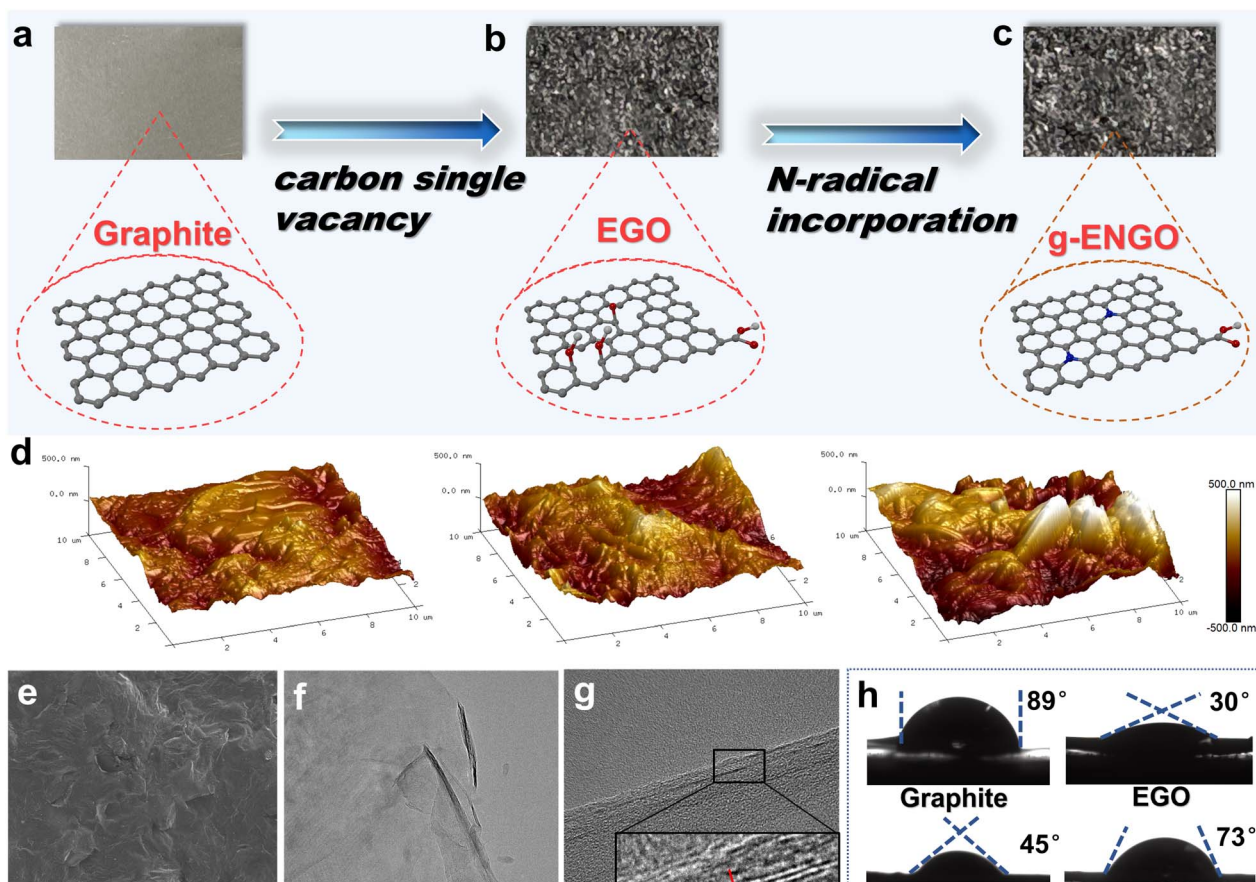


Fig. 1 Catalyst preparation process: optical images and structural model of (a) graphite, (b) EGO, and (c) g-ENGO. Atom representations are O (red), N (blue), and C (gray). (d) AFM morphological analysis of graphite, EGO, and g-ENGO. (e) SEM images, and (f and g) TEM images of g-ENGO. (h) Contact angles of graphite, EGO, g-ENGO, and g-ENGO.

ENGO, these “stalactites” transformed into “ridge”-like distributions, and the roughness (R_a) increased to ≈ 500 nm. The surface morphology of the g-ENGO was further observed using scanning electron microscopy (SEM) and transmission electron microscopy (TEM) (Fig. S2). From the SEM images with a magnification of 100 times, obvious wrinkles can be observed, which probably result from the oxygen-containing functional group generation and oxygen bubble evolution (Fig. 1e). Under the TEM view with a magnification of 63 000 times (Fig. 1f), g-ENGO looks like a smooth gauze, which is the typical morphology of a two-dimensional nanocarbon material. Because of the SO_4^{2-} intercalation, its lattice fringe spacing (0.46 nm) (Fig. 1g) is larger than the normal spacing of few-layer graphene (0.34 nm). A similar phenomenon can also be seen for the graphite paper with a thickness of 1 mm (Fig. S3).

As the surface characteristics of graphite were significantly altered after oxidation and N-doping, the hydrophilicity was correspondingly changed. According to the contact angle testing (Fig. 1h), graphite paper demonstrates the poorest hydrophilicity (contact angle: 89°) because its non-polar surface characteristics weakly interact with polar water molecules. When a substantial number of oxygen-containing functional groups were introduced in the EGO, its surface polarity was significantly enhanced, exhibiting a great enhancement in

hydrophilicity with a contact angle of 30° . Meanwhile, the hydrophobicity of the g-ENGO exhibited a slight recovery (contact angle: 45°), suggesting that the graphitic-N assembly may consume adjacent oxygen-containing groups and partially restore the non-polar sp^2 conjugated structure. The electrochemical reduction of g-ENGO was performed to further remove some oxygen-containing functional groups (g-ERNGO). As expected, the hydrophobicity increased, with a contact angle of 73° . These results support the concept that the hydrophilicity of g-ENGO can be flexibly adjusted *in situ* by using the electrochemical method, which usually plays a key role in an aqueous chemical reaction.

Precise regulation of graphitic nitrogen

In the proposed hypothesis, forming SVs is the prerequisite for incorporating graphitic-N, and a series of characterization techniques was adopted to verify their existence. EPR is a visual and quantitative method for determining the concentration of SVs in EGO (Fig. S4). It can be observed from Fig. 2a that the characteristic peak pattern of the SV emerges at a g factor of 2.003, and the content gradually increases with the oxidation time.³⁴ In particular, in the first 10 min, the SV content increased from 0 to 1.493, indicating relatively fast oxidation kinetics. As the SV theoretically stems from carbon emission,



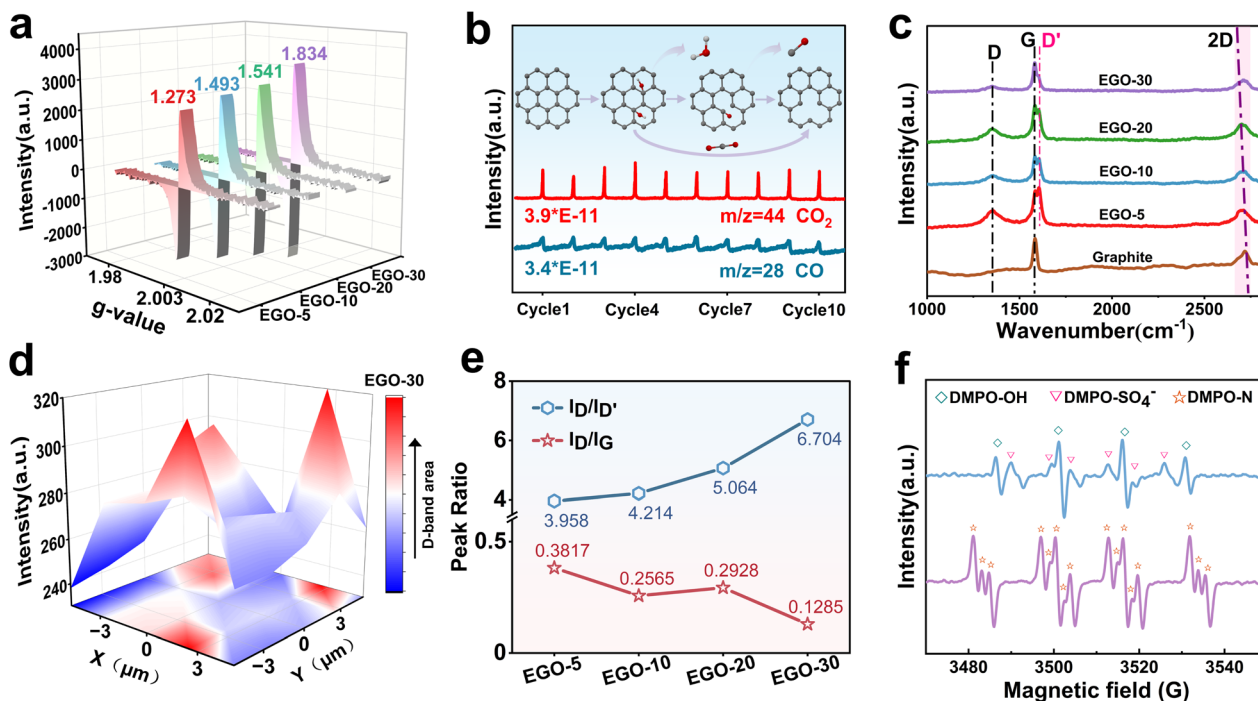


Fig. 2 (a) EPR spectra of EGO. (b) *Operando* DEMS results for EGO. Atom representations are O (red), C (gray), and H (white). (c) Raman spectra. (d) D-band Raman mapping for EGO. (e) $I_D/I_{D'}$ and I_D/I_G change curves. (f) *Operando* EPR spectra of DMPO-OH, DMPO-SO₄⁻, and DMPO-N during electrochemical processes.

operando DEMS in linear sweep voltammetry (LSV) mode for ten cycles was carried out to monitor and identify the gaseous carbon molecules being released from the graphite planes. As shown in Fig. 2b, the signals with specific mass charge ratios (m/z) of 28 (CO) and 44 (CO₂) appeared repeatedly and stably in each cycle. In addition, the deconvoluted C1s and O1s XPS spectra display three types of oxygen-containing functional groups: hydroxyl (-OH), carbonyl (-C=O) and carboxyl groups (-COOH) (Fig. S5b and 5c). Referring to the results described above, we predict that the formation of SVs is accompanied by the detachment of -OH and -C=O, which is due to continuous electrochemical oxidation at 1.8 V. The C/O ratios and high-resolution O1s spectra obtained from XPS measurements (Fig. S5a) for EGO-5, EGO-10, EGO-20, and EGO-30 provide direct evidence for the prediction. As the electrochemical oxidation proceeds, the C/O ratios fluctuate up and down, and do not exhibit the linear-like decreasing trend observed in the electrochemical exfoliation of GO.³⁰

Raman spectroscopy analysis (Fig. 2c) was further performed to characterize the carbon structure of the EGO. The overall I_D/I_G ratio shows a downward trend, indicating that as the oxidation time was extended, sp³ carbon decreased and SVs increased. Compared to the graphite, the 2D band (2720 cm⁻¹) of the EGO undergoes a red shift, which also suggests the transformation of in-plane sp² carbon to distorted sp³ carbon and the development of defects.³⁵ In addition to the typical D-band (1350 cm⁻¹) and G band (1580 cm⁻¹) peaks, the characteristic peak for the D' band was observed at 1611 cm⁻¹ for EGO. The intensity of the D' band, originating from a double resonance Raman feature induced by disorder and defects, has been

reported to be a measure for distinguishing boundary-like defects ($I_D/I_{D'} = \sim 3.5$), vacancy-type defects ($I_D/I_{D'} = \sim 7$), and sp³-type defects ($I_D/I_{D'} = \sim 13$) by comparing with the intensity of the D-band. Herein, the $I_D/I_{D'}$ ratio increased with the oxidation duration from 3.958 (EGO-5) to 6.704 (EGO-30), indicating that the boundary-like defects gradually transformed into vacancy-type defects (Fig. 2e).³⁶ Meanwhile, 3D Raman mapping of the D peak from the EGO-30 within a square area of 10 μm × 10 μm displays uniform distribution of the vacancies on the graphite plane (Fig. 2d).

Based on these results, it can be confirmed that SVs were successfully produced in the EGO, and their concentration can be adjusted by controlling the oxidation time.

Electrochemically activating NH₄⁺ to N radicals is the second step to form graphitic-N. The previous studies have speculated that hydroxyl radicals can attack ammonium ions to oxidize them into N radicals in an electrochemical system.^{37,38} In order to explore this using experiments, *operando* electrochemical EPR measurement was performed in LSV mode, and 5,5-dimethyl-1-pyrroline-N-oxide (DMPO) was used as the trapping agent. As shown in Fig. 2f, EPR signals from DMPO-OH, DMPO-SO₄⁻, and DMPO-N spin adducts all appeared, indicating the formation of hydroxyl radicals, sulfate radicals, and N radicals (*NH₂ and/or *NH).^{39,40}

Precisely doping graphitic-N with a controllable concentration is an anticipated advantage of the electrochemical method. Based on the prepared EGO at 1.8 V with different oxidation times (0–30 min), a series of g-ENGO samples was prepared under a constant 1.6 V. From the deconvoluted N1s XPS spectrum (Fig. 3a and S7), the doped N states are composed of



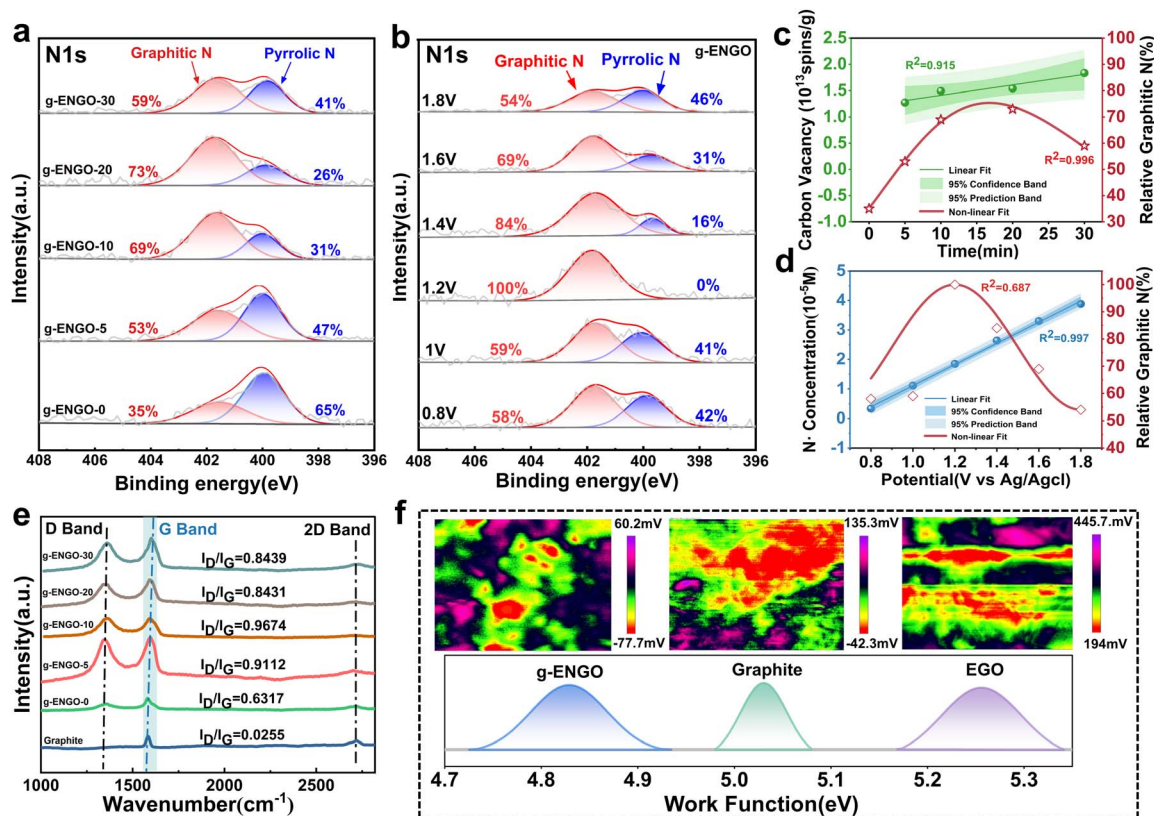


Fig. 3 (a) High-resolution N1s XPS of g-ENGO-*x* (*x* represents different oxidation times). (b) High-resolution N1s XPS of g-ENGO (at different doping potentials). (c) The relationship between carbon vacancy concentration and relative graphitic-N content as a function of oxidation time. (d) The relationship between nitrogen radical concentration and relative graphitic-N content as a function of voltage. (e) Raman spectra. (f) Plots of work functions and potential maps of graphite, EGO, and g-ENGO samples derived from KPFM results (inset).

graphitic-N (401.6 eV) and pyrrolic-N (399.9 eV), with no pyridinic-N detected (398.8 eV).¹⁶ This represents a significant difference compared to the chemical methods. By exploring the relationship between the carbon vacancy content and the relative graphitic-N content (Fig. 3c), it was found that as the carbon vacancy content increases, the relative graphitic-N content exhibits a parabolic trend. The highest relative graphitic-N content of 73% was obtained for g-ENGO-20. At this point, the absolute graphitic-N content reaches 1.581% (Fig. S9). This is because an excessive oxidation duration will transform a carbon single vacancy into a double-vacancy or even hole defects, preventing nitrogen atoms from tridentate coordination.³² Clearly, the presence of SVs is an essential condition for the formation of graphitic-N, which perfectly validates our hypothesis. Controlling the oxidation time of the EGO is one of the effective methods to adjust the graphitic-N content. The pioneers of this work believed that pyrrolic-N is formed *via* the nucleophilic attack of N-containing species on the carbonyl functional groups.⁴¹ Therefore, the lowest pyrrolic content of 26% in g-ENGO-20 can be attributed to its lowest amount of carboxyl functional groups (~0%) (Fig. S5). It can also be noted that even the graphite paper was doped by electrochemical oxidation without pre-oxidation (g-ENGO-0), and graphitic-N still emerged, which confirms the effect of water dissociation on SV generation. The negative results obtained from the methyl-

added system also indicate the important effect of water dissociation (Fig. S11 and Table S3).

We predict that anchoring a nitrogen atom into the carbon *via* the N radical is another crucial process for constructing graphitic-N. When the distribution of the SVs is determined, the N radical production is probably an adjustable parameter to control the concentration of graphitic-N in the g-ENGO. As the oxidation potential has a significant influence on this process (Fig. S6), a series of doping potentials (0.8 V to 1.8 V) for the g-ENGO was examined with EGO-30 as the basic sample (Fig. 3b and S8). Similar to the influence of the oxidation time during EGO synthesis, the relative graphitic-N content also exhibited a parabolic trend with the doping potential (Fig. 3d and S10). Specifically, 100% relative content of graphitic-N was obtained under 1.2 V, indicating that N radicals play a key role in the formation of graphitic-N. A quantitative comparison with prior work, including synthesis parameters, total nitrogen content, and nitrogen species distribution, is provided in Table S1. The advantage of our electrochemical method for the precise incorporation of graphitic-N can be clearly understood. According to the assumption from the contact angle test that oxygen-containing functional groups may be involved in graphitic-N formation, we conducted a nitrogen-doping experiment by pyrolyzing SV-contained graphite and melamine at 900 °C (Fig. S12). Due to the lack of oxygen-containing functional groups on the graphite plane, no nitrogen atoms were



doped in. This suggests that oxygen-containing functional groups are the third essential participants for graphitic-N assembly, especially the adjacent ones at the SVs.

The incorporation of graphitic-N on the graphite plane was further validated *via* Raman spectroscopy (Fig. 3e) and Kelvin probe force microscopy (KPFM) using the series of g-ENGO adjusted by oxidation time (Fig. 3f). Compared with the original graphite, the I_D/I_G ratio of the g-ENGO increased significantly, confirming the gradual formation of defect structures due to nitrogen doping. In addition, the G band (1580 cm^{-1}) of the g-ENGO exhibits a blue shift, which is a typical demonstration of nitrogen doping in graphite.⁴² Meanwhile, KPFM work function measurements show opposite regulatory effects of vacancy doping and graphitic-N doping on the Fermi level of graphene. Using the work function ($\Phi \approx 5.03\text{ eV}$) of the graphite as a reference, EGO gives a higher work function of $\sim 5.26\text{ eV}$, indicating a downward shift in the Fermi level. In contrast to that, g-ENGO-1.2 gives a lower work function of $\sim 4.82\text{ eV}$, indicating an upward shift in the Fermi level.⁴³

Synthesis mechanism

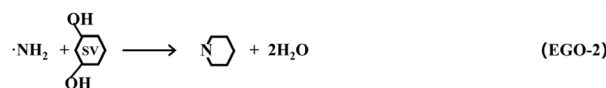
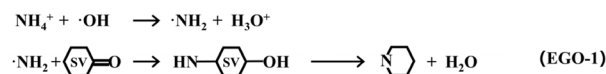
Based on the experimental results, the proposed mechanism regarding nitrogen transformation from ammonia ions to N radicals was further verified by SERS (Fig. 4a and S13). Detailed experimental procedures are provided in the SI. By performing LSV over a potential range of 0.8 V to 1.95 V (*vs.* Ag/AgCl), we successfully detected the stretching vibration modes of $^*\text{NH}_2$ (410 cm^{-1}), $^*\text{NH}$ (662 cm^{-1}), and $^*\text{NNH}$ (585 cm^{-1}), which are key intermediates needed in the graphitic-N doping process.^{27,28} Additionally, the stretching vibration peak of the active oxygen radical ($^*\text{O}$) was consistently observed at 516 cm^{-1} . Within the voltage range of 0.8 V to 1.7 V, the strong intensities of the $^*\text{NH}_2$ and $^*\text{O}$ peaks provide strong evidence that the primary N radicals are $^*\text{NH}_2$, and they are achieved depending on the attack of the hydroxyl radicals on the ammonium ions. When the potential continually increases to 1.9 V, the intensity of the $^*\text{O}$ peak decreases and the $^*\text{NH}_2$ peak disappears, indicating that excessive potential will cause vigorous hydrolysis, resulting in the undesirable production of oxygen. Peaks corresponding to $^*\text{NH}$ are also observed in the 0.8 V to 1.7 V range, and their intensities continuously increase with the potential. The similar trends for $^*\text{NH}_2$ and $^*\text{NH}$ indicate that the adsorbed $^*\text{NH}_2$ undergoes a dehydrogenation reaction on the EGO surface to form $^*\text{NH}$. Additionally, the $^*\text{NNH}$ peak becomes prominent in the range of 1.4 V to 1.7 V, but nearly disappears when the potential continues to rise. It indicates that a significant portion of the adsorbed $^*\text{NH}$ undergoes N–N coupling reactions to form N_2 .²⁸

DFT calculations were performed to clarify the reaction mechanism from EGO to g-ENGO from a thermodynamic perspective (Fig. 4b and c). Given the experimental results above, we constructed two EGO models: a single carbon vacancy on a finite graphene flake with one carbonyl ($-\text{C}=\text{O}$) group (EGO-1) or adjacent two hydroxyl ($-\text{OH}$) groups (EGO-2) (Fig. S14). The three unsaturated carbon atoms surrounding the vacancy were labeled as C_1 , C_2 , and C_3 (Fig. 4b and S15).

Route-a was exhibited for graphitic-N doping in EGO-1, with localized amplification of a-INT1. The carbonyl group is bonded to C_1 , and an external $-\text{NH}_2$ group adsorbs onto the carbonyl site with a binding energy of $3.03\text{ kcal mol}^{-1}$. Subsequently, the nitrogen atom bonds with C_2 , followed by the transfer of one hydrogen atom from $-\text{NH}_2$ to the oxygen atom, forming an $-\text{OH}$ group. The nitrogen then bonds with C_3 , resulting in two new C–N bonds on the carbon surface. Finally, the nitrogen forms a bond with C_1 , undergoing a second N–H dissociation (ab-INT2). The released hydrogen atom attacks the oxygen atom, forming an H_2O molecule and simultaneously completing the formation of graphitic-N (abc-INT2). Route-b and route-c were both calculated for graphitic-N doping in EGO-2 (bc-INT1). The two hydroxyl groups are bonded to C_1 and C_3 , and the $-\text{NH}_2$ group adsorbs onto them with a binding energy of $3.39\text{ kcal mol}^{-1}$. The nitrogen atom sequentially bonds with C_2 and C_1 , forming two C–N bonds and two $-\text{OH}$ groups. The route bifurcates based on which oxygen atom is attacked first by the hydrogen in $-\text{NH}_2$. In route-b, the nitrogen bonds with C_3 , and the hydrogen attacks the oxygen on C_3 to form an H_2O molecule. A second N–H cleavage then occurs, with hydrogen attacking the oxygen on C_1 , generating a second H_2O molecule and resulting in graphitic-N formation (abc-INT2). In route-c, the hydrogen atom first attacks the oxygen on C_1 , forming H_2O , followed by nitrogen bonding with C_3 . A second N–H cleavage yields another H_2O molecule through attack on the oxygen at C_3 , again resulting in graphitic-N.

As shown in Fig. 5c, in all three routes (a, b, and c), the attachment of the $-\text{NH}_2$ group to the carbon atom (a-TS1 and bc-TS1) is the rate-determining step, with activation energies of $18.08\text{ kcal mol}^{-1}$ for route-a and $20.94\text{ kcal mol}^{-1}$ for routes b and c. These values suggest that graphitic-N doping is kinetically accessible in all three cases. The calculated reaction energies are $-149.68\text{ kcal mol}^{-1}$ for route-a and $-179.22\text{ kcal mol}^{-1}$ for routes b and c, confirming that the graphitic-N deriving from the SV, $-\text{NH}_2$, and adjacent oxygen-containing functional groups is thermodynamically favorable under electrochemical conditions.

Based on the experimental research and DFT calculations mentioned above, schematic equations to elucidate the mechanism for NH_4^+ ion conversion into graphitic nitrogen under our electrochemical conditions are provided as follows.



g-ENGO applications

In our previous research, the experiments and DFT calculations clearly showed that peroxydisulfate (PDS) can preferentially adsorb on graphitic-N (-2.46 eV on graphitic-N, -1.30 eV on pyridinic-N, and -1.34 eV on N vacancies).³² This means that electron transfer can occur from graphitic-N to the PDS without



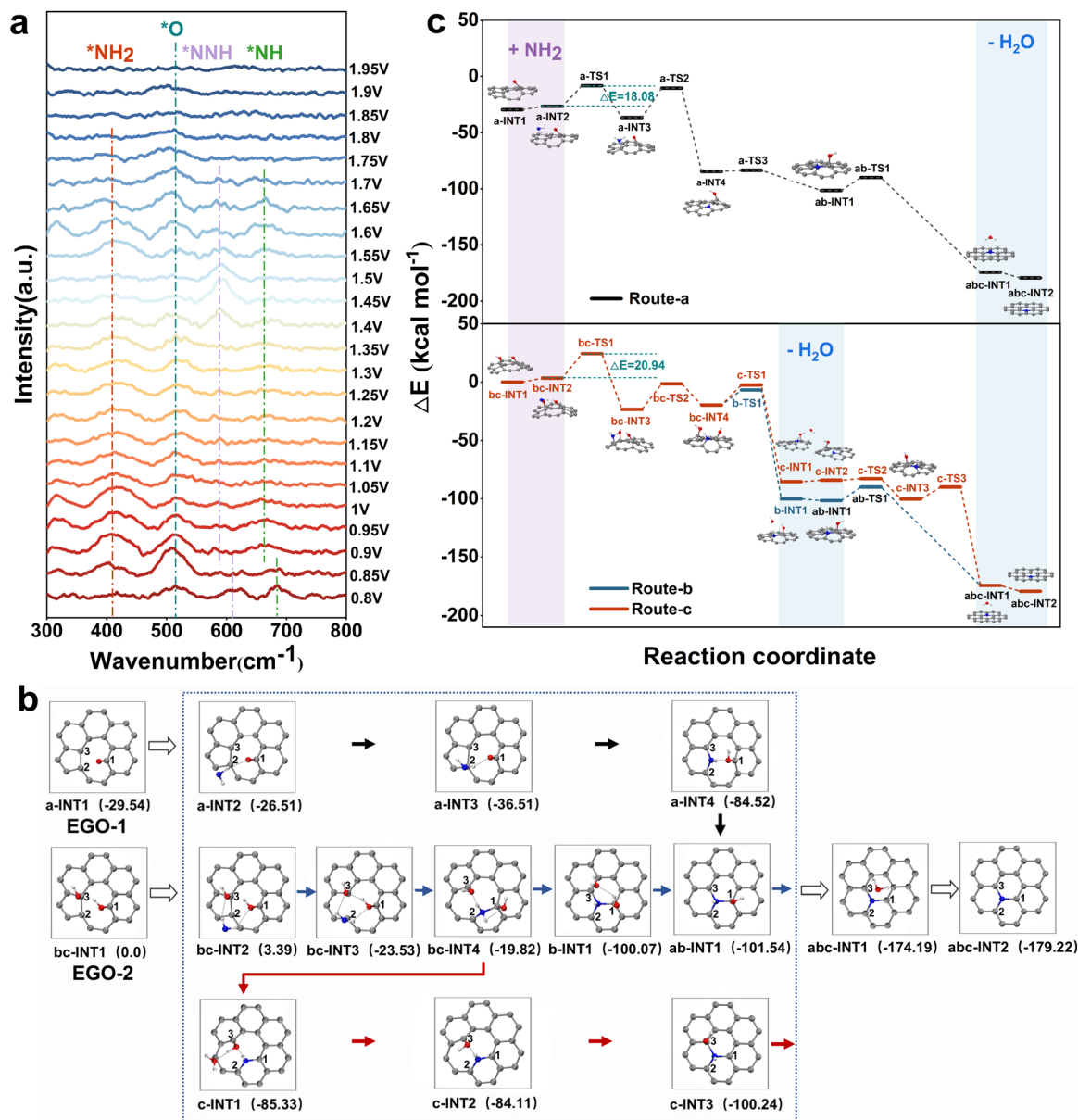


Fig. 4 (a) SERS spectra of ammonia dehydrogenation in an aqueous solution of 1.25 M $(\text{NH}_4)_2\text{SO}_4$. (b) Fragmental structures for the stationary points involved in route-a, route-b, and route-c. Atom representations are O (red), N (blue), C (gray), and H (white). Other carbon atoms are omitted for clarity. The distances are represented in units of angstroms. Relative energies (in parentheses) are given in kcal mol^{-1} . (c) Reaction energy distribution of ammonia-mediated a-INT1 (route-a in black) and bc-INT1 (route-b in blue and route-c in red) doping reactions.

any external energy input. This can be attributed to the increased Fermi level of the carbon material by the graphitic-N doping (proved by KPFM here). Therefore, we performed open-circuit potential (OCP) measurement for detecting the electron transfer from graphitic-N to PDS as a reference to evaluate the performance of the g-ENGO obtained using different doping potentials (Fig. 5a). To avoid the interference of the oxygen-containing functional groups, the g-ENGO was reduced at -1 V for 30 min (Fig. S16). The initial OCP of g-ERNGO remained stable, and upon injection of 1 mM PDS at 300 s, the OCP of the g-ERNGO immediately increased by 0.2 V to 0.5 V. Notably, the amplification of the OCP was positively

correlated with the relative content of graphitic-N. This result gives a reliable prediction that g-ERNGO can be used as an electrochemical sensor for PDS.

OER performance was also examined as a demonstration of the graphitic-N doping. LSV tests (Fig. 5b) show that the initial overpotentials of g-ERNGO-1.6 V, 1.4 V, 1.2 V, 1 V, and 0.8 V are 1.39 V, 1.41 V, 1.45 V, 1.51 V, and 1.63 V, respectively, significantly lower than that of graphite (1.84 V relative to RHE). In particular, the increase of the absolute graphitic-N content has an obvious role in negatively shifting the overpotential of the OER. Good stability of the g-ERNGO was demonstrated through



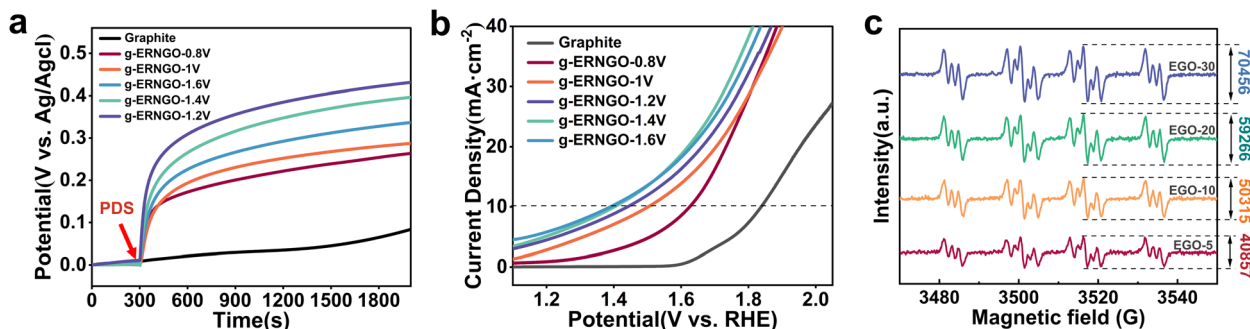


Fig. 5 (a) Open-circuit potential curves with the addition of 1 mM PDS on g-ENGO-X (X represents the doping potential). (b) LSV curves of g-ENGO-X in 1 M KOH solution, showing the OER catalytic activity of the electrode. (c) *Operando* EPR spectra of DMPO-N during g-ENGO preparation.

consecutive cyclic voltammetry (CV) scans for 50 cycles, indicating its application potential in practice (Fig. S18).

As the active oxide radicals derived from OER are crucial for N radical generation during g-ENGO synthesis, we designed a masking experiment by using the *operando* EPR measurement (Fig. 5c) to verify the active role of graphitic-N in the OER, in which EGO-5, EGO-10, EGO-20, and EGO-30 were used as the working electrode, and graphitic-N doping was performed by oxidizing them for 10 min in 1.25 M $(\text{NH}_4)_2\text{SO}_4$. At the same time, the scavenger DMPO is added to the scavenging system to capture N radicals that are not used for doping. Theoretically, the peak intensity of the N radicals should decrease, resulting from the capture by SVs in the graphite. However, as the content of SVs gradually increased from EGO-5 to EGO-30 (Fig. 2a), the concentration of the captured N radicals also exhibited an increasing trend. The reason for this may be that the graphitic-N generated during doping reduces the energy barrier for hydrolysis to hydroxyl groups, thereby promoting the accelerated production of N radicals. This is consistent with our predicted conclusion.

Conclusions

In this work, we proposed a controllable and easy-to-operate electrochemical method enabling the selective synthesis of graphitic-N at ambient temperature and pressure. The sequence process, including vacancy engineering and N radical assembly, allows precise regulation of the graphitic-N content, and 100% relative content of graphitic-N was achieved. By integrating advanced characterization and DFT calculation, we elucidated the formation mechanism of graphitic-N and identified the crucial parameters for its precise regulation. The SVs, N radicals, and adjacent oxygen-containing functional groups are the three essential participants in graphitic-N doping, which significantly depend on electrochemical water dissociation to hydroxyl radicals. This study provides a flexible framework for the systematic preparation of nitrogen-doped carbon materials and paves the way for their application in electrochemical energy conversion (*e.g.* fuel cells and metal-air batteries), environmental remediation (*e.g.* advanced oxidation processes and pollutant adsorption), electrochemical sensing, *etc.*,

thereby contributing to the development of sustainable, green, and low-carbon technologies.

Author contributions

Conceptualization: Leilei Xu, Heng Dong; investigation: Leilei Xu, Zhibo Zhang, Hong Zhou, Ziqi Wen; supervision: Heng Dong, Wei Xie; writing – original draft: Leilei Xu; writing – review & editing: Heng Dong, Wei Xie, Yuxuan Liu.

Conflicts of interest

The authors declare no competing interests.

Data availability

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information: experimental details (materials and characterization); synthesis methods; TEM images, XRD pattern, XPS analysis, results of EPR, construction of models and computation details. See DOI: <https://doi.org/10.1039/d5sc08053c>.

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