Biochar Aerogel Decorated with Thiophene S Manipulated 5-membered Rings Boosts Nitrogen Fixation

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Abstract

Algae, which contains most of the marine biogenic sulfur (S), is the main producer of dimethylsulfide (DMS). The oxidation products of DMS (methanesulfonic acid (MSA) and non-sea salt sulfates (nss-SO42-)) are important contributors to acid rain acidity (~ 40%) and have great influence on global climate change and acid rain formation. The biogenic S of red algae mainly exists in carrageenan molecules, the polysaccharide extract of red algae. In this work, the carrageenan biogenic S was fixed into biochar aerogel (SCA) synchronically by pyrolysis with the assistance of the unique hydrogel property of carrageenan. During this process, 89% of C and 96% of marine biogenic S are preserved in SCA, efficiently preventing the loss of C and S element. More importantly, the thiophene S structure in SCA framework derived from the marine biogenic S could endow the carbon aerogel superior electrocatalysis activity by regulating the charge density of adjacent C atoms. For instance, thiophene S structure can promote the cleavage and protonation of N2 molecules and efficiently lower the energy barrier of the promote the protonation of N2, thus greatly enhance the activity of electrocatalytic NRR. This study provided a promising avenue for the synchronization fixation of marine biogenic S and C to reduce the the formation of acid rain and carbon dioxide emissions. It also developed a green and sustainable way to synthesize high-performance biochar materials for advanced energy conversion.

Main Text

Sulfur (S) cycle is a process of transformation and reciprocation of the S elements in the ecosystem and environment (Supplementary Figure 1), wherein the acid rain is a big problem due to its great harm to ecological environment, agriculture and even human beings. Algae, which contains most of the marine biogenic S, is the main producer of dimethylsulfonio-propionate (DMSP) and contributes ~ 40% of the acidity of acid rain. Besides, the decay of algae will cause large amount of S and C elements to be lost in the atmosphere to form toxic substances or greenhouse gases. Therefore, it is urgent and meaningful to explore effective strategies to prevent the loss of S and C elements from the perspective of sustainable and environmental protection. In red algae, most of the biogenic S is found in polymers called carrageenan. Carrageenan contains a large number of sulfonic groups and can be easily treated to obtain porous biochar aerogel structure. Surprisingly, carrageenan has the properties of temperature-sensitive gel, and its molecular chain will show "spiral" transformation during the cooling process. The carrageenan with "spiral" molecular chains has potential to generate three-dimensional (3D) hierarchical porous biochar aerogels. Recently, biochar materials and their applications in emerging energy storage and conversion have become an attractive topic in the field of sustainable development. The utilization of biochar materials could play the role of carbon sequestration and carbon reduction, and greatly reduce the emission of CO2, which is an effective way to mitigate global warming. In addition, the introduction of S into biochar can endow the carbon aerogel superior electrocatalysis activity by regulating the charge density of adjacent C atoms, and thus improve the added value of biochar materials. Therefore, large-scale utilization of carrageenan to synthesize biochar can not only play an
Important role in fixing S and C, but also provide an excellent electrocatalyst for advanced energy conversion.

Ammonia (NH₃) is one of the most important fundamental chemicals for humans. Since Davy et al. discovered the electrochemical N₂ reduction reaction (NRR), it has become a promising alternative to traditional industrial ammonia synthesis (Haber-Bosch) approach for the synthesis of NH₃.\textsuperscript{14,15,16} Compared to traditional Haber-Bosch process, the electrocatalytic process using the sustainable protons in water to replace the hydrogen gas used in industry, and convert N₂ into NH₃ by controlling the potential, which greatly reduces energy consumption and carbon dioxide emissions, and promote the sustainable development of human society.\textsuperscript{17,18} However, the chemical superstability of N₂ molecules make the realization of electrocatalytic NRR difficult, which requires the search of catalysts with special active sites to accelerate ammonia synthesis.\textsuperscript{19} The designed NRR electrocatalysts should exhibit excellent apparent activity to efficiently promote the diffusion of solvated gas (N₂) and electrolytes, and exhibit outstanding intrinsic activity to improve the catalytic ability of active sites, thus accelerating the cleavage and protonation of N₂ to enhance the activity of NRR.\textsuperscript{19}

Herein, we fabricated biochar aerogel electrocatalysts decorated with thiophene S (SCA) for electrochemical NRR by directly pyrolyzing carrageenan. During this process, 89% carbon was retained and converted to biochar structure. Meanwhile, 96% marine biogenic S was cured to thiophene S and oxidized S species into the biochar, efficiently prevents the loss of S element. More importantly, the thiophene S structure in the SCA achieves excellent electrochemical N₂ reduction reaction (NRR) performance with a highest NH₃ yield rate of 36.69 μg h⁻¹ mg⁻¹\textsubscript{cat} and Faradaic efficiency (FE) of 8.72% in acid solution under ambient conditions. Density functional theory (DFT) calculation results further confirm that thiophene S structure can effectively regulate the charge density of adjacent C atoms to facilitate the activation of N₂ molecules. The preferable reaction pathway for NRR is *N₂ → *NNH → *NHNH → *NHNH₂ → *NH₂NH₂ → *NH₂ → *NH₃.

As illustrated in Figure 1a, the S element in carrageenan was fixed in the SCA framework and converted to thiophene S and oxidized S species by using a simple freeze-drying and pyrolysis process. Specifically, the hot aqueous carrageenan solution converted to transparent hydrogel after cooling down at a temperature lower than 80 °C, since the random coiled carrageenan macromolecules were cross-linked to form “double-helix” structure.\textsuperscript{20} After subsequent freeze-drying, the carrageenan hydrogel converted to highly porous aerogel. Finally, the carrageenan aerogel was pyrolyzed at T °C (600, 700, and 800) under Ar atmosphere to obtain ultralight S-doped carbon aerogels (SCA-T). Besides, the inductively coupled plasma atomic emission spectroscopy (ICP-AES) also was performed to prove that no common metals existed in SCA electrocatalyst (Supplementary Table 1). The as-synthesized samples were characterized by using scanning electron microscopy (SEM) and transmission electron microscopy (TEM). Obviously, as displayed in the SEM (Figure 1b–c) and TEM image (Figure 1d), the SCA is a 3D self-supporting porous network structure. A large amount of macropores, mesopores (25–50 nm) and
micropores can be observed on the surface of SCA. The mesopores and micropores can provide large electrode/electrolyte interface to accelerate electron transfer.\textsuperscript{21,22} EDS mapping spectrum indicates that C, S and O elements are uniformly dispersed in the whole SCA (Figure 1e).

Figure 2a displays the X-ray diffraction (XRD) patterns of SCA−T. The two diffraction peaks (23.0 and 43.5°) are assigned to the (002) and (101) diffraction plane of graphitic structure in SCA matrix. The N\textsubscript{2} adsorption–desorption isotherms and pore size distribution of SCA−T are shown in Figure 2b. The adsorption occurred at the relative pressure (\(P / P_0 < 0.02\)) indicates the existence of micropores in the samples, which is consistent with the results show in the TEM above. The H4–type hysteresis loop originates from the presence of mesopores.\textsuperscript{23} The SCA−700 owns the highest Brunauer–Emmett–Teller (BET) surface area (1668.9 m\textsuperscript{2} g\textsuperscript{−1}). From the pore size distribution, the pore diameters are located at ~1.0 nm (micropores) and ~ 2.3 nm (small mesopores) by the desorption branch of isotherm calculation. The appearance of micropores and mesopores in SCA−T is due to the decomposition of sulfur and oxygen groups (−SO\textsubscript{4}H, −COOH, and −OH) in carrageenan macromolecules. The high BET surface area combined with hierarchical porous structure could efficiently promote the diffusion of solvated gas (N\textsubscript{2}) and electrolytes, thus accelerating the protonation of N\textsubscript{2} to enhance the activity of NRR.\textsuperscript{18} The thermogravimetric analysis (TG/DTG) and the corresponding mass spectra (TG-MS) were been carried out to reveal the pyrolysis process of carrageenan aerogel (Figure 2d). The DTG curve shows three more distinct stages of heat gains at 207 °C, 310 °C and 726 °C during the whole pyrolysis process. Meanwhile, the first, second and third major weight loss steps could be found in the TG curve, that occur in the temperature range of 190−221 °C, 227−380 °C and 672−741 °C, respectively (Figure 2c). As shown in the Figure 2d, the corresponding TG-MS curves display the volatiles from aerogel pyrolysis are CO\textsubscript{2} (m/z: 44), H\textsubscript{2}O (m/z: 18), SO\textsubscript{2} (m/z: 64) and CO (m/z: 28). The sharp peaks could be found at the temperature of 194 °C from the Pink and purple curves (H\textsubscript{2}O and CO\textsubscript{2}), respectively, which corresponds to the first rapid weight loss step (190−221 °C) of the TG curve. Besides, the peaks at 296 °C in pink and orange curves (CO\textsubscript{2} and SO\textsubscript{2}), and the peaks at 700 °C in pink and olive curves (CO\textsubscript{2} and CO) could be ascribed to the second (227−380 °C) and third (672−741 °C) major decomposition steps, respectively, corresponding to the TG. Thus, during the pyrolysis process, the oxygen-containing groups (−SO\textsubscript{4}H, C−O−C and −OH et al.) in carrageenan molecular are converted to H\textsubscript{2}O, CO\textsubscript{2}, SO\textsubscript{2} and CO by the TG and TG-MS analysis. TG–Fourier transform infrared spectrisopy spectra (TG–FTIR) was carried out to further confirm the volatiles components in the thermal decomposition of carrageenan aerogel.

As shown in the 3D TG–FTIR spectrum, the peaks at wavenumbers 3851−3253 cm\textsuperscript{−1} indicated the existence of H\textsubscript{2}O by the stretching vibration of O−H bonds.\textsuperscript{24} The adsorption bands at 2400−2240 cm\textsuperscript{−1} were assigned to CO\textsubscript{2} and CO. The peaks at wavenumber 1700−1600 cm\textsuperscript{−1} and 1600−1520 cm\textsuperscript{−1} were the stretching vibration of C=O bonds and C=C bonds, respectively, which confirms the existence of some carbonyl compounds (such as ketones, alkane, and acids).\textsuperscript{25} The peaks at wavenumbers 900-1131 cm\textsuperscript{−1} representing the S=O bonds stretching vibration, which indicated the co-existence of oxidized S and sulfonate groups. In addition, at the wave number of 1200-1354 cm\textsuperscript{−1}, no peak was found in the original
carrageenan, but with the increase of pyrolysis temperature, the C-S-C vibration peaks representing the thiophene-S structure began to appear.\textsuperscript{26} The X-ray photoelectron spectroscopy (XPS) measurement of SCA-700 was carried out to probe the chemical environments of C, S and O elements (Figure 2f, Supplementary Figure 2). As shown in high-resolution S 2p spectra, the thiophene S (S 2p\textsubscript{1/2} 163.7 and S 2p\textsubscript{3/2} 165.1 eV) was determined to be the dominant S species.\textsuperscript{27} The residual S located at 168.4 and 169.2 eV are attributed to the −C−SO\textsubscript{x} species.\textsuperscript{28,29} Furthermore, the S K-edge X-ray absorption structure (XAS) spectra further confirmed the existence of thiophene S structure. As shown in the Figure 2g, the two major absorption bands were observed in the XAS of SCA-700 spectra, representing thiophene S (2471.1 eV) and oxidized-S (2481.1 eV), respectively.\textsuperscript{30} However, in the carrageenan XAS spectra, only sulfonate (2479.8 eV) and oxidized-S (2481.1 eV) are visible, which indicates that the sulfonate and oxidized-S in carrageenan were transformed or removed into thiophene S and oxidized-S after the fixation process.\textsuperscript{31} The XPS and XAS results demonstrated that the contained S in carrageenan was successfully fixed in a carbon framework and converted to thiophene S and oxidized S (Figure 2h).

For NRR tests, we used the carbon paper coated with SCA−T as working electrode. The NH\textsubscript{3} yield rates and FEs were obtained by spectrophotometry with indophenol blue method (Supplementary Figure 3, Supplementary Figure 4).\textsuperscript{32} Figure 3a shows the NRR performance of SCA−T in 0.1 M HCl electrolyte. Apparently, the sample of SCA−700 exhibits much higher NH\textsubscript{3} yield rates than SCA−600 and SCA−800, attributing to its higher specific surface area. Then, we measured the linear sweep voltammograms (LSVs) in Ar and N\textsubscript{2}−saturated 0.1 M HCl electrolyte to further appraise the NRR electrocatalytic activity of most active SCA−700 (Figure 3b). Apparently, the current density under N\textsubscript{2} is higher than that under Ar, indicating the occurrence of electrocatalytic N\textsubscript{2} reduction.\textsuperscript{33} The chronoamperometry curves at different working potentials implying that as−prepared SCA−700 can ensure the stability for reaction process (Figure 3c).\textsuperscript{34} By calculation, Figure 3d displayed the corresponding NH\textsubscript{3} yield rates and FEs of SCA−700 under different potentials. The N\textsubscript{2} can be reduced to NH\textsubscript{3} at the potential of −0.1 V to −0.5 V (vs. RHE). Besides, both NH\textsubscript{3} yield rate and FE improve as the potential is negatively shifted, and achieve the highest value at the potential of −0.3 V. The highest NH\textsubscript{3} yield rate and FE are 36.69 µg h\textsuperscript{−1} mg\textsuperscript{−1} cat and 8.72\%, respectively. Obviously, the NH\textsubscript{3} yield rate of SCA−700 is higher than the most of the previously reported metal-free electrocatalysts (Figure 3e and Supplementary Table 2). In addition, to further confirm that the detected NH\textsubscript{3} was generated \textit{via} the N\textsubscript{2} fixation process over SCA−700, we also performed electrolysis in Ar−saturated solution and pure carbon paper in N\textsubscript{2}−saturated solution at −0.3 V for 2 h. The corresponding UV−Vis absorption spectra show no detectable NH\textsubscript{3} generated in either condition (Supplementary Figure 5). Furthermore, we tested the \textsuperscript{1}H NMR analysis of the electrolyte fed by \textsuperscript{15}N\textsubscript{2} and \textsuperscript{14}N\textsubscript{2} after electrolytic reaction at -0.3 V to further determine the N source of produced NH\textsubscript{3}. As shown in Figure 3f, only the two peaks of \textsuperscript{15}NH\textsubscript{4}\textsuperscript{+} were found in the \textsuperscript{15}N saturated electrolyte, indicating that N\textsubscript{2} is the only source of produced NH\textsubscript{3} by SCA-700. Notably, the N\textsubscript{2}H\textsubscript{4} (by−product) could be detected by spectrophotometry with the method of Watt and Chrisp.\textsuperscript{35} The detecting calibration curve of N\textsubscript{2}H\textsubscript{4} is
shown in Supplementary Figure 6. We failed to detect the production of N₂H₄ in N₂−saturated electrolyte (after charging at -0.3 V for 2 h), indicating that SCA−700 is an electrocatalyst with excellent selectivity for NRR (Supplementary Figure 7). Then, we performed cycling and chronoamperometric tests of SCA−700 in 0.1 M HCl media to study the stability of catalyst for practical. As shown in the Figure 3g, no obvious decline in NH₃ yield rate and FE of SCA−700 after 5 times cycling tests. And after the NRR test lasted for 10 h, the SCA−700 shows a relatively stable current density at −0.3 V (Supplementary Figure 8).

In order to further emphasize the significance of the fixed S unit in N₂ fixation, we annealed the SCA−700 at 1100 °C to remove S atoms in SCA, fabricating SCA−700−1100. The XPS of SCA−700−1100 was displayed in Supplementary Figure 9. Apparently, only very weak peaks are observed in S 2p spectra of SCA−700−1100, indicating that most of S atoms are removed after the pyrolysis at 1100 °C. The atomic percentage of S content is largely reduced from 4.64% for SCA−700 to 0.15% for SCA−700−1100 (Supplementary Table 3). As shown in Figure 3h, the SCA−700−1100 only delivers a yield rate of 8.2 μg h⁻¹ mg⁻¹ cat, much lower than the yield rate of SCA−700. The result indicates that fixed S unit is most likely the origin of the boosted NRR performance in SCA−700. In addition, as shown in electrochemical impedance spectroscopy (EIS) (Supplementary Figure 10), the charge transfer resistance (Rₓ) value of SCA−700 is lower than SCA−700−1100, indicating that the fixed S can improve the conductivity and fast the charge transport ability. Given that HER is the primary competitive reaction of NRR, we tested the HER performances of SCA−700 and SCA−700−1100 in acid solution as well. Obviously, as shown in Figure 3i, the SCA−700 shows lower HER catalytic activity than SCA−700−1100, which indicated that the fixed S unit is the key to inhibit HER activity and promote NRR activity. However, the prepared SCA-700 contains two types of S units, thiophene S and S oxides, which is difficult to verify experimentally which of the two S units is the main active species for catalytic N₂ reduction. Therefore, we compared the Gibbs free energy of N₂ protonation (∆G*ₐₙ₂H) at different sites of the two S units to determine the best active unit for NRR by DFT calculations. Apparently, compared with thiophene S, the oxidized S unit exhibits very high ∆G*ₐₙ₂H in the N₂ protonation process, and is ineffective on electrocatalytic N₂ fixation (Supplementary Figure 11 and Supplementary Table 4). Based on the above results, the thiophene S unit is the main active site for N₂ fixation in SCA-700 electrocatalysts.

To reveal the role of thiophene S structure of SCA on NRR, we studied the electronic structure of SCA and the Gibbs free energy diagram of NRR on SCA by the DFT calculations. As shown in Figure 4a, we constructed computational calculations on two types of models (pure carbon substrate (G) and thiophene 5-membered ring on carbon substrate (S−G−n, n represents the position of active sites in Figure 4a). Firstly, the charge density difference is calculated to study the electron transfer behavior between N₂ and S−G or G. As shown in Figure 4b, more electron transfer between S−G and N₂ is observed than that between G and N₂. Figure 4c compares the density of states (DOS) for G and S−G. The DOS of S−G in Fermi level is stronger than that of G, which indicates that the S−G has better conductivity than G. Gibbs free energy of N₂ protonation (∆G*ₐₙ₂H) was calculated to highlight the effect of thiophene S for N₂ fixation (Figure 4d). Apparently, the S−G−2 exhibits much lower ∆G*ₐₙ₂H (1.12 eV) than that of G.
(2.59 eV), indicating that thiophene S can reduce the energy barrier of N₂ protonation and enhance the NRR activity by regulating the charge density of adjacent C atoms (Supplementary Table 4). It is known that the N₂ molecules can be absorbed in a vertical or horizontal way, and the adding H atoms can perform hydrogenation to the catalyst via a distal or alternating mechanism for NH₃ synthesis. As shown in Figure 4e and Supplementary Figure 12, we performed three possible reaction pathways for NH₃ synthesis on S−G. By contrast, the preferable pathway for NRR on S−G is *N₂ → *NNH → *NHNH → *NHNH₂ → *NH₂NH₂ → *NH₂ → *NH₃ (vertical alternating pathway). The rate-determining step (RDS) of vertical alternating pathway is the protonation of pre-absorbed N₂ (*N₂ → *NNH). The energy barrier (1.12 eV) of RDS in vertical alternating pathway is much lower than that in vertical distal pathway (1.68 eV) and horizon alternating pathway (2.62 eV). The thiophene S unit is critical factor to accelerate N₂ dissociation and protonation on carbon substrate. Therefore, the lowest energy barrier on S−G predicts that the SCA is an ideal electrocatalyst for catalyzing NRR.

In summary, large-scale exploitation and utilization of algae could effectively fix S and C, thus preventing the emission of sulfur dioxide and carbon dioxide. In this work, we synchronously fix the carrageenan biogenic S into the SCA by using pyrolysis with the assistance of the unique hydrogel property of carrageenan. It not only perfectly completes the role of C and S fixation, but also the generated SCA exhibits excellent NRR activity and selectivity. Under ambient conditions, the yield rate of NH₃ and FE can reach 36.69 μg h⁻¹ mg⁻¹ cat and 8.72%, respectively, which exceeds mostly reported metal-free catalysis. DFT calculation results suggested that the thiophene S structure could reduce the energy barrier of N₂ protonation by regulating the electronic structure of adjacent C atoms. The preferable reaction pathway for NRR was *N₂ → *NNH → *NHNH → *NHNH₂ → *NH₂NH₂ → *NH₂ → *NH₃. This study opens up a new avenue for the synchronization fixation of marine biogenic S and C, and explores a sustainable strategy to synthesize high-performance biochar materials for electrochemical.

Declarations

Supplementary Information is available in the online version of the paper.

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**Figures**
Figure 1

a, Schematic illustration on the process of marine biogenic S fixation. b–c, FESEM images of SCA-700. d–e, TEM image and the corresponding EDS mappings for C, S, and O elements of SCA-700.
Figure 2

Characterization of SCA. a, XRD patterns of SCA−T. b, Nitrogen adsorption−desorption isotherm for SCA−T (inset: Pore size distributions of SCA−T). c, TGA and DTG curves of carrageenan aerogel in N2 atmosphere (heating rate 5 °C/min). d, The analysis of outlet gases from the decomposition of carrageenan aerogel TG-MS: H2O (pink), CO (olive), CO2 (purple) and SO2 (orange). e, 3D TG-FTIR spectra of carrageenan aerogel. f, S 2p XPS spectra of SCA−700. g, S K-edge XAS spectra of carrageenan and SCA-700. h, The schematic diagram of the process of the S fixation.
Figure 3

a, NH3 yield of SCA−T in 0.1 M HCl electrolyte. b, LSV curves of SCA−700 in Ar−saturated and in N2−saturated electrolyte. c, The corresponding chronoamperometry curves of SCA−700 at different potentials in 0.1 M HCl solution. d, NH3 yield and FEs of SCA−700 under different potentials. e, Comparison of the NH3 yield rate among SCA-700 and available reported metal-free catalysts. f, 1H NMR analysis of the electrolyte fed by 15N2 and 14N2 after electrolytic reaction at -0.3 V. g, Cycling test of SCA−700 at -0.3 V. h, NH3 yield and i, HER performances of SCA−700 and SCA−700−1100.
Figure 4

a, Schematic of computational models. Brown and yellow represent C and S atoms, respectively. b, The images of charge density difference at G and S−G. The yellow and cyan color represents the charge accumulation and depletion, respectively. c, DOS of G and S−G. d, Calculated free energy diagram of *N2H for G and S−G−2. e, Free energy diagram of N2 reduction on S−G.
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