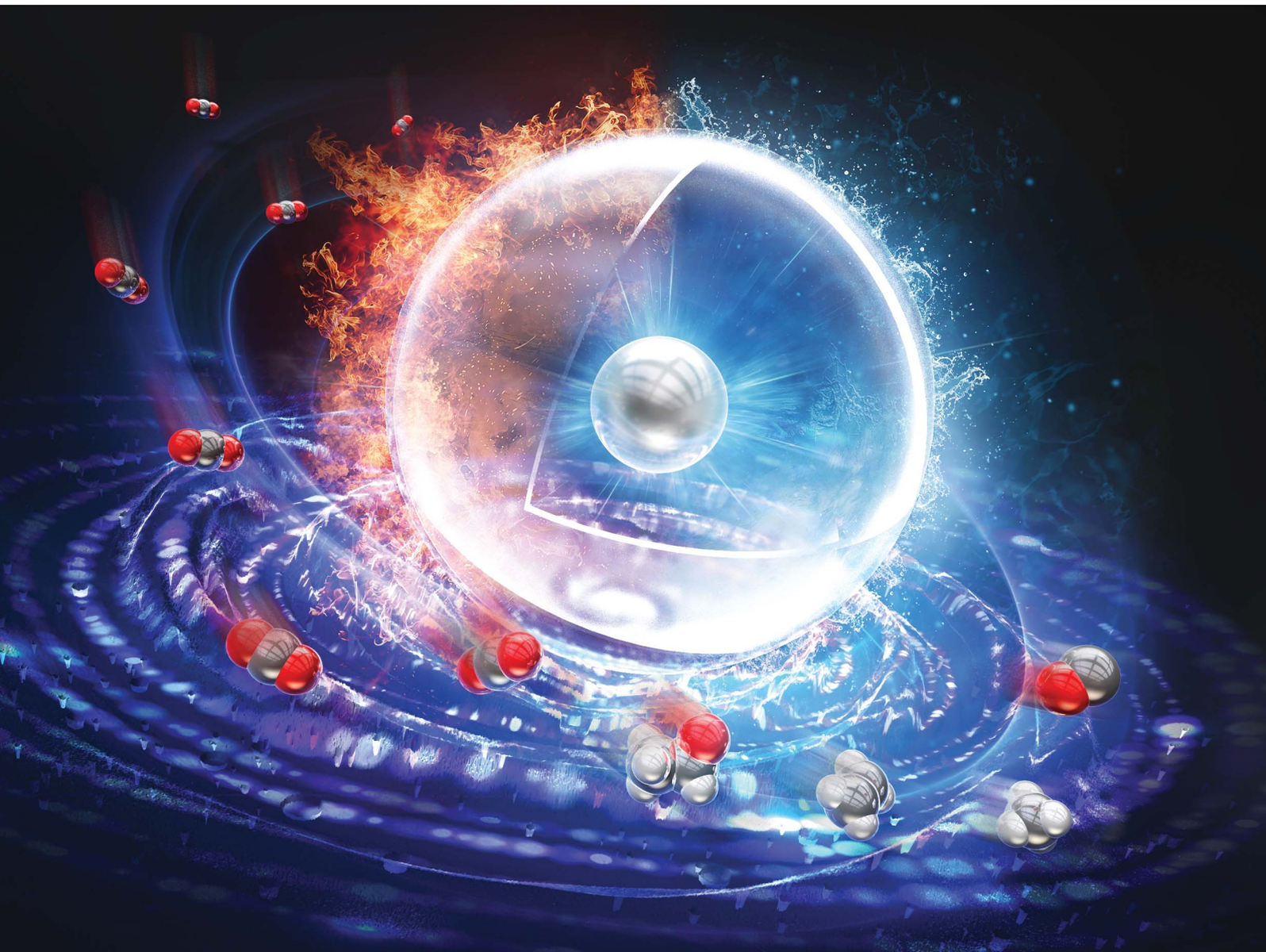


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REVIEW ARTICLE

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Hollow carbon-based materials for electrocatalytic
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Hollow carbon-based materials for electrocatalytic and thermocatalytic CO₂ conversion

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Electrocatalytic and thermocatalytic CO₂ conversions provide promising routes to realize global carbon neutrality, and the development of corresponding advanced catalysts is important but challenging. Hollow-structured carbon (HSC) materials with striking features, including unique cavity structure, good permeability, large surface area, and readily functionalizable surface, are flexible platforms for designing high-performance catalysts. In this review, the topics range from the accurate design of HSC materials to specific electrocatalytic and thermocatalytic CO₂ conversion applications, aiming to address the drawbacks of conventional catalysts, such as sluggish reaction kinetics, inadequate selectivity, and poor stability. Firstly, the synthetic methods of HSC, including the hard template route, soft template approach, and self-template strategy are summarized, with an evaluation of their characteristics and applicability. Subsequently, the functionalization strategies (nonmetal doping, metal single-atom anchoring, and metal nanoparticle modification) for HSC are comprehensively discussed. Lastly, the recent achievements of intriguing HSC-based materials in electrocatalytic and thermocatalytic CO₂ conversion applications are presented, with a particular focus on revealing the relationship between catalyst structure and activity. We anticipate that the review can provide some ideas for designing highly active and durable catalytic systems for CO₂ valorization and beyond.

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1. Introduction

The rapid advancement of industrialization has led to high fossil-fuel consumption, causing an ever-increasing atmospheric CO₂ concentration. Excessive CO₂ emissions are the main contributors to global warming and the resulting abnormal climatic variation, glacial ablation, ecosystem destruction, etc.,^{1,2} which seriously threaten the shared future of mankind. In the past decades, many efforts have been devoted



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using porous materials and their applications to green chemical reactions and the valorization of CO₂.



to controlling the CO₂ concentration worldwide (e.g., adoptions of the Kyoto Protocol and the Paris Agreement). The objective of the Paris Agreement is to keep the global average temperature below 2 °C above the pre-industrial mean, with a temperature increase of 1.5 °C limit.^{3,4} Nevertheless, the current trend of anthropogenic CO₂ growth makes achieving this goal an urgent task for a sustainable community and economic society.⁵

Catalytic CO₂ reduction conversions (e.g., photocatalysis, thermocatalysis, and electrocatalysis) to produce high-value-added fuels and feedstocks, offer a promising means of realizing carbon neutrality and potentially help to mitigate the global warming issue.^{6–9} In comparison to photocatalysis, thermal catalytic and electrocatalytic CO₂ conversions are easier to scale up, due to the appropriate reaction kinetics and lack of dependence on the availability and cost of high-intensity light sources. Herein, our review mainly focuses on these two reactions.

Electrochemical CO₂ reduction reaction (eCO₂RR), powered by electrical energy, refers to the process of using electrochemical cells to convert CO₂ to valuable compounds, such as carbon monoxide (CO), methane (CH₄), methanol (CH₃OH), ethylene (C₂H₄), etc. A typical electrocatalytic CO₂ reduction reaction involves four main steps:^{10,11} (1) diffusion of CO₂ to the electrolyte/cathode interface; (2) CO₂ molecules are adsorbed on the cathode surface; (3) when an external bias is applied, the oxygen evolution takes place at the anode due to the water oxidation, and the generated electrons are then transferred to the cathode through an external circuit, where they can participate in the CO₂ reduction processes; (4) desorption of final products from the cathode. These steps jointly determine the overall efficiency of eCO₂RR. Recently, eCO₂RR has become the research hotspot in CO₂ utilization owing to its multifarious advantages, including mild operation conditions (room temperature and atmospheric pressure), flexible reactor platforms, and relatively high conversion efficiency.^{12–16} Integrating renewable energy with eCO₂RR, simultaneously achieving a net CO₂ reduction and chemical raw material supply is possible. Despite eCO₂RR possesses numerous advantages, its pragmatic commercialization is still confronting great challenges. Firstly,

as a chemically inert and thermodynamically stable molecule, CO₂ is difficult to be activated because of the stable C=O bond (806 kJ mol⁻¹);^{17,18} thus, a high potential is inevitably needed to drive the eCO₂RR. Secondly, most eCO₂RR can yield multiple products, which makes the direct utilization of products difficult, requiring an extra target product separation process. Third, noble metal-based catalysts (Au, Ag) usually deliver high electroactivity, but are less desirable considering their cost-efficiency and scarcity. For these reasons, efficient electrocatalysts with the characteristics of highly exposed active sites, good conductivity, abundant pore structure for target molecule transfer, and high earth-abundance are urgently needed.^{19,20}

Thermocatalytic CO₂ conversion, mainly including CO₂ hydrogenation and CO₂ dry reforming of methane (DRM), is a feasible process to alleviate the large CO₂ emissions from the viewpoint of cost-efficiency.²¹ Among them, DRM is a highly endothermic reaction with the equation CH₄ + CO₂ → 2CO + 2H₂, ΔH° = 247 kJ mol⁻¹.²² The large energy consumption and the deactivation of catalysts caused by active metal sintering or coke deposition during the DRM process seriously hindered its applicability. Compared to DRM, thermal hydrogenation of CO₂ shows a lower thermodynamic limitation, which involves the reaction of CO₂ and H₂ and can afford high levels of variety and tailorability for the production of desirable hydrocarbons and oxygenates.^{23,24} Conventionally, the catalytic performance and product selectivity highly depend on the catalyst. Accordingly, the development of CO₂ hydrogenation catalysts with high CO₂ adsorption and activation ability, robust stability, and good selectivity, is of great practical relevance.

In nature, cells with hollow structures are the fundamental unit of an organism, and this unique structure endows them with distinctive properties that allow for the adjustment of adsorption, separation, and exchange for substrates required to maintain vital life activities.^{25,26} Inspired by nature, many encouraging hollow-structure nanoreactors have been developed for energy storage,^{27–31} biomass conversion,^{32,33} and environmental catalysis applications.^{34,35} In recent years, hollow-structured carbon (HSC) and its derived nanomaterials have attracted considerable attention in the fields of thermocatalysis and electrocatalysis for CO₂ conversion due to their advantageous features. Firstly, guest molecules and catalytically active species can be encapsulated in the cavity; by limiting the moving space of the inclusions, the leaching and aggregation of catalytically active species can be suppressed during the reaction, leading to the excellent durability and reusability of the catalyst.^{36,37} Secondly, adjustable pore structure and size of the carbon shell can regulate the mass transport and reaction characteristics, and further alter the catalytic selectivity.³⁸ Thirdly, the carbon shell provides a platform to anchor single atoms,^{39,40} load metal particles,^{41,42} and couple with multifunctional polymers,⁴³ which shows vast potential in variable catalysis applications. The rational design of nano-cavity or channel of HSC material can confine the metal nanoparticles (NPs) or cluster in a limited volume space, offering nanoconfinement effects to improve catalytic CO₂ reduction performance in terms of activity and selectivity. The positive influences of nanoconfinement mainly include (1) increasing



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reactant molecule concentration in the confined space and enriching the surface coverage of adsorbed species on the active sites,^{44,45} (2) regulating the local pH value to hinder the competitive hydrogen evolution reaction during the eCO₂RR process,⁴⁶ and (3) tailoring the intrinsic electronic structures and optimizing the intermediate adsorption energy of the catalytic sites.⁴⁷ For instance, Zheng *et al.* reported that the confinement effect of N-doped carbon nanotubes (NCNT) enabled the attenuation of the binding strengths between Ni NPs and *CO intermediates; therefore, the NCNT-confined Ni NPs sample (Ni@NCNT) showed enhanced CO selectivity in eCO₂RR in comparison with the Ni NPs-supported sample (Ni-NCNT).⁴⁸ Pan *et al.* proposed that, in the hollow mesoporous carbon spheres (HMCS) confining Cu clusters electrocatalyst system, the nano-confinement effect of HMCS contributed to the facilitation of C–C bond coupling to produce C₂ products in eCO₂RR.³⁷ In our previous works, a series of novel hollow carbon nanocatalysts encapsulating alloys have been developed, which exhibit excellent thermocatalytic activity and stability for CO₂ hydrogenation to produce formate owing to the nano-confinement effects provided by the hollow carbon sphere.^{49–51} Apart from CO₂ thermal conversion, our recent research demonstrated that HSC with superior conductivity and abundant pore structure can also serve as an effective support for syngas electrosynthesis from CO₂.⁵² In our opinion, HSC-based catalysts are promising materials for CO₂ conversion, and they will continue to be hotspots in the thermocatalytic and electrocatalytic fields.

In the past few years, several valuable review articles related to the advancement and current status of hollow nanoreactors have been published. For instance, the review from Das *et al.* captured the advancements in catalytic CO₂ conversion, with a focus on novel core–shell catalyst development, activity, and selectivity improvement.¹⁰ Kuang *et al.* reviewed the recent advances of metal@hollow carbon sphere catalysts in the fields of sustainable biomass and CO₂ conversion.²⁵ Li and co-workers reviewed the recent achievements of hollow carbon nanocages in the fields of energy storage and electrocatalysis.⁵³ Yu *et al.* reviewed the theoretical development of the hollow nanoreactor-based system, emphasizing the opportunities for the study of molecular kinetic behaviors in the hollow nanoreactors for achieving controllable catalysis.⁵⁴

However, most of the reviews of HSC materials mainly focus on the energy storage field and biomass conversion, with a small amount concentrating on thermo- and electrocatalysis towards CO₂ conversion. Besides, a comprehensible summary of the structure–performance relationship of hollow carbon-based materials for CO₂ conversion remains to be refined. More importantly, the review content related to the popularly studied single-atom anchored on hollow carbon-based materials is relatively scarce. Considering the rapid development of studies on HSC-based catalysts in thermocatalytic and electrocatalytic CO₂ conversion, a timely cutting-edge review on this topic is of imperative need.

This review aims to offer a critical overview of the latest progress of HSC-based materials in thermocatalytic and electrocatalytic CO₂ conversion fields, with a discussion on the linkage

of catalyst structure and activity (Fig. 1). The synthesis methods and functionalization strategies of HSC are also introduced. In the last section, an outlook for future research direction is proposed, targeting the current challenges.

2. Synthesis and functionalization of hollow-structured carbon materials

Generally, the HSC materials can be classified as hollow carbon spheres, hollow carbon nanotubes, hollow carbon polyhedrons, *etc.* motivated by the potentially utilitarian value for catalysis and energy storage, many efforts have been made toward the controllable synthesis and accurate design of hollow carbon with well-defined structures.^{55–57} In this section, the synthetic methods and functionalization strategies are discussed.

2.1 Synthesis methods of HSC

2.1.1 Hard template synthesis. Hard template synthesis is by far the most commonly used way to prepare HSC.⁵⁸ As shown in Fig. 2a, a complete process of hard template technology involves four major steps: (1) preparation of the hard template, (2) coating template core with carbon shell precursors, (3) carbonization to obtain template@carbon under calcination treatment, and (4) template removal. The HSC synthesized by this method shows a highly regulated appearance, and the shape and size of its cavity can be tuned by choosing the suitable template material; usually, the morphology of the final product can be well predicted. Furthermore, the choice of carbon source determines the composition and physico-

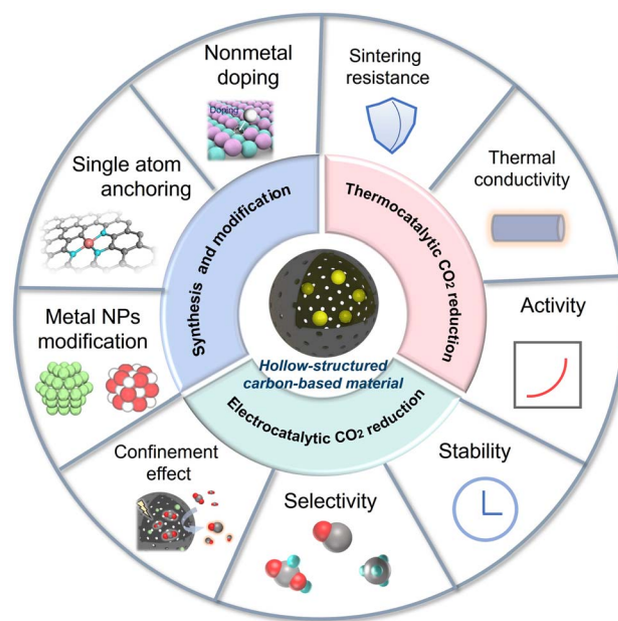


Fig. 1 Overview of the topics covered in this review. This review presents the syntheses, functionalization strategies of hollow-structured carbon (HSC) based materials, as well as their fascinating applications in electrocatalytic and thermocatalytic CO₂ conversion.



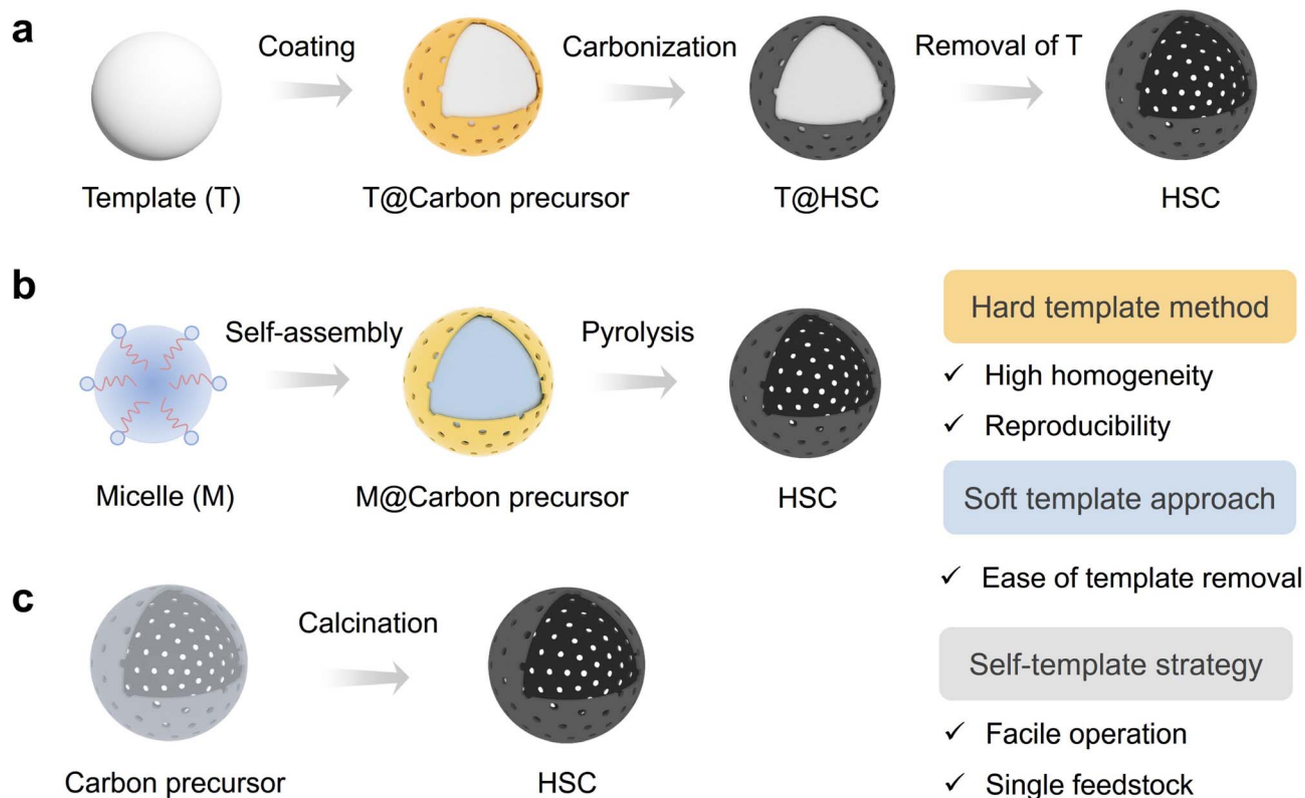


Fig. 2 Synthetic methods for hollow structured carbon (HSC). (a) Hard template method, (b) soft template approach, and (c) self-template strategy.

chemical properties of the carbon shell to some extent. A summary of recent representative works is presented in Table 1.

SiO₂ spheres are often used as hard templates to fabricate the HSC material due to the advantages of high stability, good uniformity, and ease of preparation. According to the study of

Zhang *et al.*, mesoporous carbon hollow spheres (MCHS) were prepared using SiO₂ core particle and resorcinol–formaldehyde (RF) resin as a hard template and a carbon precursor, respectively (Fig. 3a).⁵⁹ As displayed in Fig. 3b and c, hollow cavity spaces were clearly seen in the spherical structure of MCHS.

Table 1 Parameter comparison of hard templating synthesis for HSC

Template	Carbon precursor	Template removal reagent	Hollow carbon type	Application	Ref.
SiO ₂	Resorcinol and formaldehyde	HF	Mesoporous hollow carbon sphere	Supercapacitor	59
PMMA	Resorcinol and formaldehyde	Calcination	Hollow carbon sphere	Battery	64
PMMA	Dopamine	Calcination	Hollow carbon sphere	Electrocatalysis	65
SiO ₂	Aniline	HF	Hollow carbon sphere	Supercapacitor	66
CaCO ₃	Dopamine	HCl	Mesoporous hollow carbon sphere	Battery	67
SiO ₂	Resorcinol and formaldehyde	NaOH	Mesoporous hollow carbon sphere	Microwave absorption	68
SiO ₂	Dopamine	NaOH	Mesoporous hollow carbon sphere	Electrocatalysis	69
SiO ₂	Resorcinol and formaldehyde	NaOH	Hollow carbon sphere	Thermal catalysis	51
CuO ₂	3-Aminophenol and formaldehyde	HCl, H ₂ O ₂	Hollow carbon sphere	Supercapacitor	70
Bi ₂ S ₃	L-Cysteine and resorcinol	Calcination	Nanotube	Electrocatalysis	71
SiO ₂	Resorcinol and formaldehyde	HF	Hollow carbon sphere	Supercapacitor	72
Urchin-like nickel	Urea	HCl	Urchin-like hollow carbon	Electrocatalysis	60
SiO ₂	Pyrrole	HF	Hollow carbon sphere	Battery	73
SiO ₂	Resorcinol and formaldehyde	HF	Hollow carbon sphere	Battery	74
SiO ₂	Resorcinol, formaldehyde and melamine	KOH	Hollow carbon sphere	Electrocatalysis	75
SiO ₂	Resorcinol and formaldehyde	HF	Hollow carbon sphere	Oil emulsification	76
SiO ₂	Dopamine	NaOH	Hollow carbon sphere	Microwave absorption	77
SiO ₂	Resorcinol and formaldehyde	NaOH	Hollow carbon sphere	Electrocatalysis	52



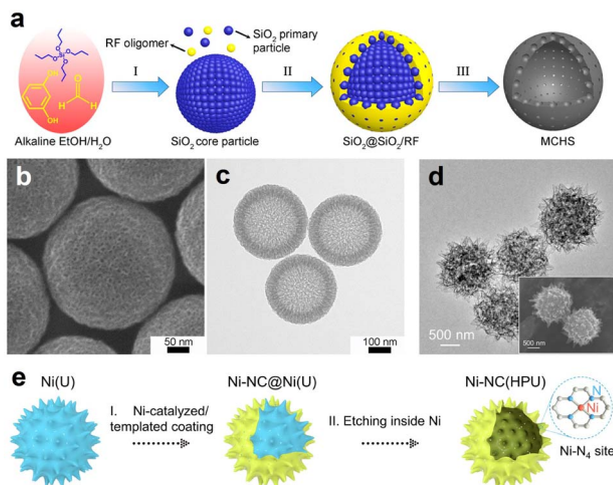


Fig. 3 (a) Schematic illustration of the synthesis procedure for mesoporous carbon hollow spheres (MCHS). (b) SEM and (c) TEM images of MCHS. Reproduced with permission from ref. 59. Copyright 2016, American Chemical Society. (d) TEM image, inset of (d) SEM image, and (e) preparation for single-Ni atoms anchored on hollow porous urchin-like N-doped carbon (Ni-NC(HPU)). Reproduced with permission from ref. 60. Copyright 2022, Wiley-VCH.

Notably, the pore size of carbon spheres can be manipulated by adjusting the synthesis conditions, such as the ratios of SiO₂ precursors (TEOS/TPOS) or solvents (ethanol/water). Taking the TEOS/TPOS ratio as an example, as the proportion of TPOS increases, the pore size of MCHS enlarges. In contrast to TEOS, the polymerization and condensation speeds of TPOS are slower. Consequently, larger silica aggregates are easily formed in the case of high-fraction TPOS, which contributes to the large mesopore formation in the following interaction process with RF oligomers. Therefore, the optimized preparation parameter is of great significance to obtain the desired HSC material. In addition to SiO₂, a metallic hard template is also available for the HSC preparation. Recent work from Li *et al.*, for instance, provided a strategy to prepare a N-doped urchin-like hollow carbon loaded with single-Ni atoms using solid urchin-like Ni and urea (providing C, N sources) as raw materials (Fig. 3d and e), where the Ni not only served as a hard template to assist the creation of hollow structure, but also offered the Ni source for the formation of Ni single sites.⁶⁰ Actually, the aforementioned method of integrating the metallic hard template with appropriate carbon sources may offer a new perspective for developing advanced nano-structured single-atom carbon-based catalysts.

2.1.2 Soft template method. Even though the hard template method offers numerous benefits, some drawbacks still exist. For example, the synthesis process of this approach is laborious; and the removal of the template involves the use of hazardous strong acids or bases. These issues stimulate the material researcher to exploit more facile and efficient syntheses of HSC. As illustrated in Fig. 2b, the soft templating method involves the utilization of thermally decomposable components (*e.g.*, micelles, emulsion droplets, or vesicles) as templates, wherein a carbon precursor is used to encapsulate

these templates, obtaining a core-shell structured composite intermediate.^{61,62} During the subsequent thermal treatment, the internal template decomposes while the external shell undergoes thermal fixation and carbonization. Eventually, a hollow carbonaceous material can be produced.

The soft templating approach for preparing HSC has gained considerable interest (Table 2), mainly due to the facile template fabrication and removal processes. In 2014, Wang *et al.* proposed a strategy to synthesize HSC using the mixed micelle converted from the P123 (EO₂₀-PO₇₀-EO₂₀) and sodium oleate as a soft template;⁷⁹ following this method, 2,4-dihydroxybenzoic acid and formaldehyde were served as carbon sources. Their results demonstrated that the diameter and shell thickness of HSC are highly related to the synthesis parameters, such as reaction temperature and the usage of polymer precursors.

The preparation of HSC with a single opening on the shell is an attractive method for increasing the diffusivity and accessibility of the guest species to the interior surface, while preserving the porous structure. Therefore, based on the traditional soft templating technique, Yu and co-workers developed a single-hole hollow carbon sphere through a poly(ethylene glycol) (PEG)-assisted emulsion-templating method, where PEG molecule functioned as a reverse demulsifier to manipulate the structure of HSC (Fig. 4a).⁶³ As revealed in Fig. 4b–d, hollow structured carbon spheres with a single hole can be observed, and the sizes of the hole and sphere are determined to be ~38 nm and ~138 nm, respectively. Additionally, by adjusting the molecular weight of PEG molecules, a closed-shell hollow carbon sphere, or bowl-like carbon can also be obtained. This work provides a new inspiration for the subtle preparation of HSC materials.

2.1.3 Self-template method. Different from the hard/soft template method, self-templating synthesis only involves the direct carbonization of the carbon-containing precursors to produce HSC (Fig. 2c). The emergence of the self-templated technique has reinvigorated the family of HSC synthetic approach (Table 2), which avoids the reliance on external templates and shows great potential for scalable fabrication.

Zhang *et al.* reported the fabrication of hollow carbon nanobubble with a shell thickness of only ~10 nm by simply calcining hollow ZIF-8 in a N₂ atmosphere (Fig. 5a).⁸⁵ It was proposed that, the distinctive hollow structured carbon featuring with ultra-thin shell not only serves to shorten the distance of ion diffusion distance but also provides a more accessible surface area for ions. Consequently, hollow carbon nanobubbles showed enhanced ion storage performance, as compared to the solid carbon nanoparticles. This synthesis concept is expected to stimulate the exploitation of more multifunctional hollow carbon materials derived from MOFs. In a separate study, Zheng *et al.* synthesized N, S co-doped hollow mesoporous carbon spheres (N/S-HMCS) using sulfur-bridged covalent triazine frameworks (S-CTF) sphere as a precursor (Fig. 5b).⁸⁹ The key point of this synthesis method is to control the pyrolysis temperature to regulate the polymerization degree of the inner and outer layers of S-CTF, which triggers the formation of the inner cavity. During the pyrolysis process, the



Table 2 Summary of the soft-templating and self-templating syntheses for HSC

Preparation method	Carbon precursor	Template	Morphology	Application	Ref.
Soft template	Glucose	Latex	Hollow carbon sphere	Battery	78
	2,4-Dihydroxybenzoic acid and formaldehyde	P123 and sodium oleate	Hollow carbon sphere	Hydrogenolysis of 5-hydroxymethylfurfural	79
	Ribose	Oleic acid emulsion	Flasklike hollow carbon	Supercapacitor	80
	Glucose	Oleic acid emulsion	Hollow carbon sphere	Biochemistry and supercapacitor	81
	Aniline and pyrrole	F127	Hollow carbon sphere	Sodium storage devices	82
	2,4-Dihydroxybenzoic acid and hexamethylenetetramine	P123 and sodium oleate	Single-hole hollow carbon sphere	Liquid-phase adsorption	63
Self-template	Melamine and formaldehyde resin	—	Hollow carbon microspheres	Supercapacitor	83
	Poly(amic acid)	—	Hollow carbon microspheres	Supercapacitor	84
	Monocrystalline zeolitic imidazolate framework (ZIF-8) nanobubble	—	Hollow carbon nanobubble	Battery	85
	Deep eutectic solvent (urea, 2,5-dihydroxy-1,4-benzoquinone and ZnCl ₂)	—	Hollow carbon nanorod	Supercapacitor	86
	Poly(hexachlorocyclotriphosphazene-tannic acid-4,4'-sulfonyldiphenol) hollow nanosphere	—	Hollow carbon sphere	Electrocatalysis	87
	Polydopamine vesicle	—	Hollow carbon cage	Cathode for Li-S batteries	88
	Sulfur bridge covalent triazine framework sphere	—	S, N-doped hollow carbon sphere	Electrocatalysis	89
	Tetrahydrofuran-treated resorcinol-formaldehyde (RF) polymer sphere	—	Concave hollow carbon sphere	Anode material for Na/K-ion battery	90

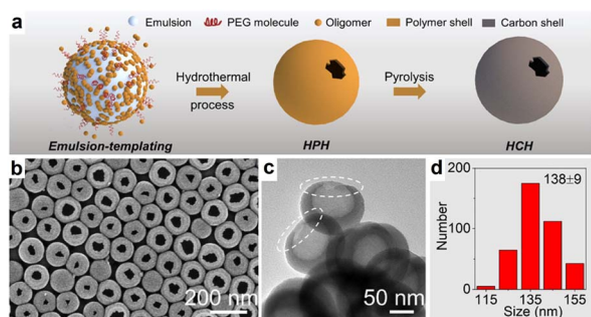


Fig. 4 (a) Schematic diagram of the synthesis for single-hole hollow carbon spheres (HCH). (b) SEM and (c) TEM images, and (d) diameter distribution of HCH. Reproduced with permission from ref. 63. Copyright 2022, American Chemical Society.

volatile substance generated from the thermal decomposition of the central oligomer with a lower polymerization degree gradually migrated to the outside to reduce the surface energy and maintain the structural stability. With the temperature increase from 500 °C to 900 °C, the internal cavity size gradually became larger and the specific surface area increased from 12 m² g⁻¹ (N/S-HMCS500) to 331 m² g⁻¹ (N/S-HMCS900). This study proposes a promising route toward the controllable synthesis of hollow carbon material with tailored heteroatom doping.

In general, the selection of an appropriate synthesis method for HSC should meet the demand of the intended application scenario. The hard template method is more recommended for applications requiring high product homogeneity and experimental reproducibility. Nevertheless, this approach still has

some limitations, including the complex and time-consuming procedures and the use of corrosive reagents for template removal. In contrast, the soft template method offers a relatively easy template removal process; however, challenges arise concerning the maintenance of structure rigidity throughout

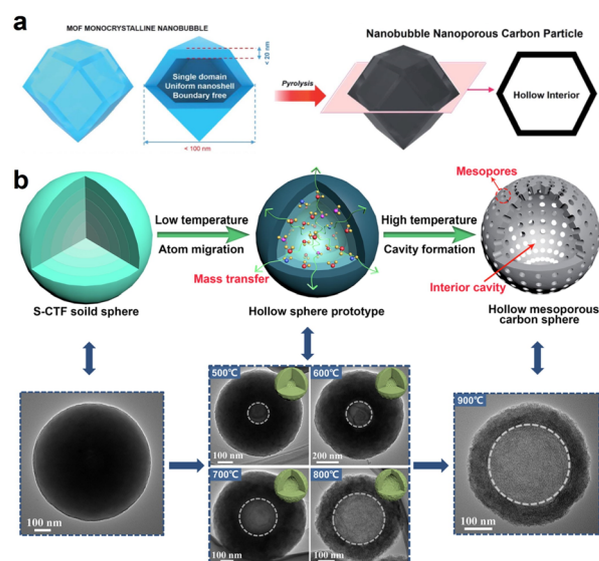


Fig. 5 (a) Preparation of nanobubble carbon via the pyrolysis of MOF nanobubbles. Reproduced with permission from ref. 85. Copyright 2017, Royal Society of Chemistry. (b) The fabrication process of N, S co-doped hollow mesoporous carbon sphere (N/S-HMCS) and the corresponding TEM images. S-CTF represents the sulfur-bridged covalent triazine frameworks. Reproduced with permission from ref. 89. Copyright 2021, Elsevier.



subsequent applications. Although the template method shows some advantages in the aspect of controlling the particle size uniformity and dispersion of HSC, the self-templated method is still regarded as a promising route for achieving scalable production due to its facile operation and utilization of only a single feedstock.

2.2 Functionalization strategies

2.2.1 Nonmetal doping. External nonmetal heteroatoms (nitrogen (N), phosphorus (P), sulfur (S), fluorine (F), *etc.*) doping, which has long been well-established in the study of carbon materials, provides a feasible modification route to regulate the charge redistribution, CO₂ affinity, and electrical conductivity.^{91–94} Thus far, the most well-established strategy is the introduction of N with high electronegativity (3.04) into the carbon framework (carbon electronegativity: 2.55), and it can change the asymmetric electron spin density and improve the intrinsic activity.^{95–97} For example, Wu *et al.* synthesized a high-efficiency nitrogen-doped carbon nanotube catalyst (NCNT) with the aid of a liquid chemical vapor deposition (CVD) method.⁹⁸ It was found that the electrochemical CO₂-to-CO conversion performance of NCNT even surpassed those of noble metals (Au and Ag), achieving a similar CO selectivity at a lower overpotential. In sharp contrast to NCNT, the pristine CNT exhibited poor activity and selectivity. As evidenced by the experimental characterization and DFT theoretical calculations, the features induced by N doping, including high conductivity, efficient catalytic sites, and a low energy barrier for CO₂ activation, benefits CO₂ electroreduction process. In another example, Wang *et al.* integrated NCNT and N-doped porous carbon to a composite membrane, and it was demonstrated to serve as an efficient metal-free catalyst for electrochemical CO₂ reduction to produce formate;⁹⁹ the results highlighted the advantage of the nitrogen doping effect in the improvement of electrochemical stability and anti-oxidizability.

In addition to N, F doping is identified as an efficient way to tailor the electronic structure of carbon catalysts and enhance eCO₂RR activity by forming a local positive charge area, and the local positive charge area can elevate COOH* (an important intermediate) adsorption and suppress the side-competing hydrogen evolution reaction (HER).^{100,101} To further increase the surface area, Ni *et al.* developed a F-doped cage-like carbon electrocatalyst, which exhibits excellent CO product selectivity at high overpotential.¹⁰² The results demonstrated that hollow morphology with abundant mesopores and micropores can facilitate the capture and diffusion of CO₂ molecules. Additionally, a strong electric field formed at the edge site of the opening pores on the F-doped carbon shell can increase the partial concentration of electrolyte cations (K⁺), thus lowering the eCO₂RR thermodynamic energy barrier and boosting the activation of CO₂.

Besides single-element doping, multiple-element doping has recently been proven to be effective in synergistically improving the eCO₂RR performance of HSC materials. Li *et al.* developed a N, S dual co-doped hollow carbon sphere, which showed a high CO faradaic efficiency (FE_{CO}) of 93% at a low potential of

−0.6 V (*vs.* RHE), surpassing that of single-N-doped counterpart.¹⁰³ The addition of S doping provided more active sites, and decreased the Gibbs free energy for the formation of COOH* intermediate, thus promoting the eCO₂RR activity. Furthermore, N, B co-doped hollow carbon spheres reported by Cheng *et al.* also exhibited a high FE_{CO} of 95.1% at a low overpotential of 310 mV.¹⁰⁴ It was suggested that doped B sites enhance CO₂ adsorption, while N sites promote the hydrogenation of *CO₂ to form COOH*.

Although encouraging progress has been made in developing non-metal-doped carbon materials, some issues still need to be solved. Taking N-doping carbon as an example. The identification of actual active sites in N-doped carbon catalyst is still controversial,¹⁰⁵ because, during the doping process, different kinds of nitrogen configurations, including pyridinic-N, pyrrolic-N, and graphitic-N, will be introduced to the carbon framework, resulting in structural inhomogeneity.¹⁰⁶ It is difficult to precisely control the N configuration. Furthermore, interactions between different types of nitrogen potentially exist, which may affect the CO₂ catalytic reaction process.⁹⁶ The combination of advanced *in situ* characterization techniques and DFT simulation is needed to further clarify the structure–activity relationship and understand the relevant reaction mechanism.

2.2.2 Metal single-atom anchoring. Since the emergence of single-atom catalysts (SACs) concept in 2010, SAC has received interdisciplinary attention and become the frontier hotspot in catalysis field due to its exceptional catalytic properties.¹⁰⁷ The atomic scale metal sites endow the catalyst with many attractive advantages in catalytic CO₂ reduction: (1) high atom utilization efficiency allows to lower the cost, especially in the case of noble metal.^{108,109} (2) Excellent activity and selectivity can be achieved by regulating the single atom coordination environment and electronic structure.^{110,111} (3) Well-defined active sites provide ease for the catalytic mechanism study.¹¹²

Normally, the isolated metal atoms with high surface energy are unstable and easy to migrate and coalesce.¹¹³ Integrating single-atom active sites with the aforementioned heteroatoms-doping HSC material prevents the aggregation of single atoms and endows HSC considerable catalytic activity simultaneously. For example, Pan *et al.* developed a highly dispersed Co–N₅ site anchored on a hollow carbon sphere (Fig. 6a and b), and it showed high-efficiency CO₂ electroreduction reactivity with FE_{CO} reaching 99.2% and 99.4% at −0.73 and −0.79 V *vs.* RHE, respectively (Fig. 6c).¹¹⁴ In contrast, cobalt phthalocyanine (CoPc, Co–N coordination number is 4) showed relatively low FE_{CO}, implying the positive roles of Co–N₅ sites. As verified by the DFT calculation results (Fig. 6d), the excellent eCO₂RR activity is associated with the existence of single-atom Co–N₅ sites and the resulting enhanced CO₂ activation, rapid formation of COOH*, and improved CO desorption. Similarly, by loading single-atom Ni–N_x sites on the well-regulated carbon structure of HSC-based catalysts, desired electrochemical eCO₂RR performance can be achieved.^{115–117} Furthermore, atomically dispersed Ru(III) supported on a N-doped carbon sphere also showed satisfactory activity and stability in continuous CO₂ hydrogenation test, owing to the strong



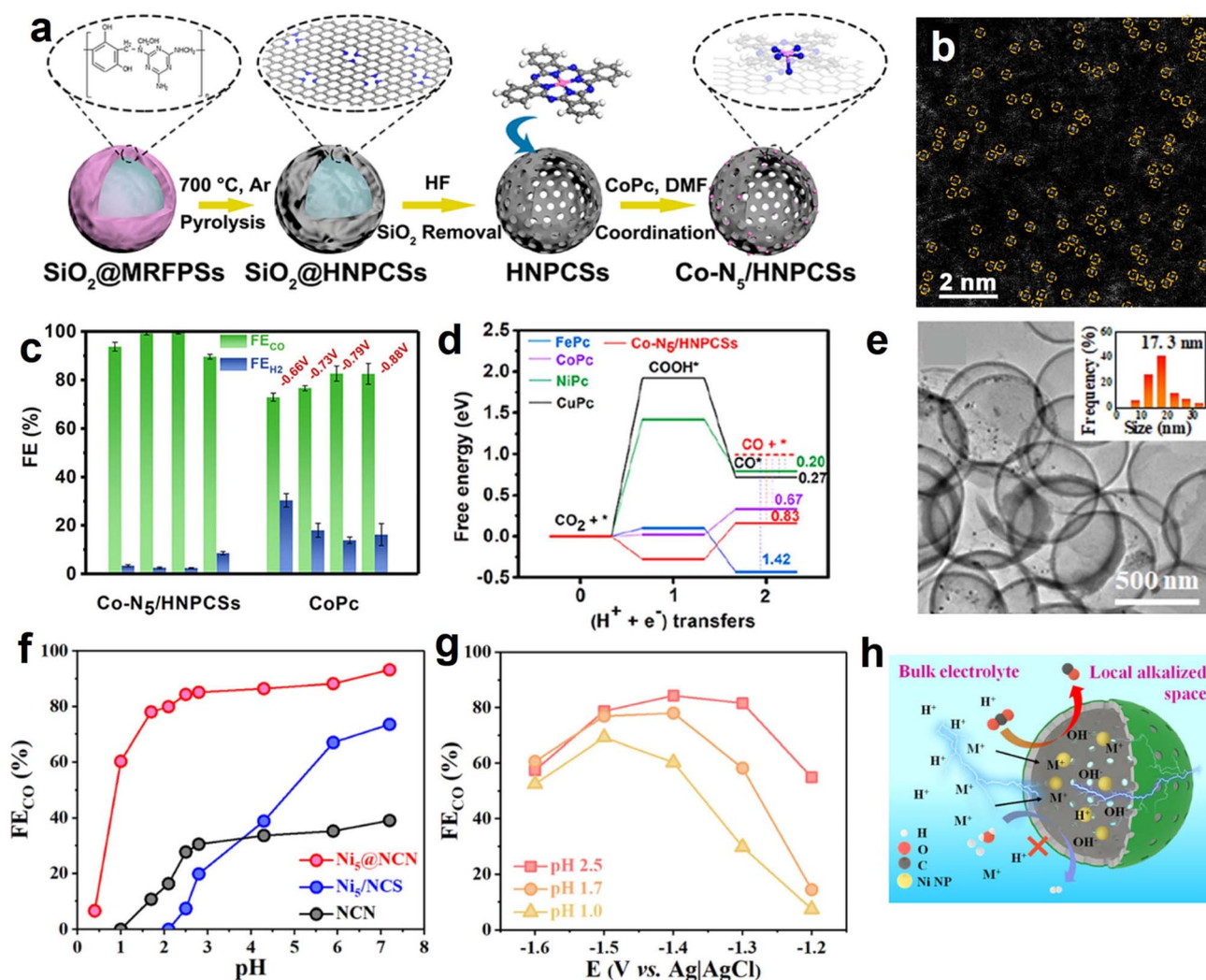


Fig. 6 (a) Synthesis procedures and (b) magnified HAADF-STEM image of hollow N-doped porous carbon spheres with single-atom Co-N₅ sites (Co-N₅/HNPCSS). (c) Faradaic efficiencies for CO and H₂ production at the given potentials over Co-N₅/HNPCSS and CoPc. (d) Calculated eCO₂RR free energy profiles of Co-N₅/HNPCSS catalyst and MPc (M = Co, Fe, Ni, Cu). Reproduced with permission from ref. 114. Copyright 2018, American Chemical Society. (e) TEM image of Ni NPs@N-doped carbon nanocages (Ni@NCN). The inset shows the metal NPs size distribution. (f) FE_{CO} and (g) current density of the best-performing Ni@NCN sample in acidic electrolyte condition. (h) Schematic diagram of the pH environment and ions transportation around the Ni@NCN catalyst. Reproduced with permission from ref. 46. Copyright 2022, American Chemical Society.

binding strength between Ru(III) species and pyridinic N/pyrrolic N site.¹¹⁸

As mentioned above, single-atom modification plays a significant role in improving the eCO₂RR performance of HSC-based materials. However, some important aspects still need to be further investigated: (1) the coordination environment has great impact on the electronic structure of SACs, so unveiling the relationship between the coordination number or connecting heteroatom type of SACs and eCO₂RR activity is important; (2) at present, most of the SACs syntheses are still at the laboratory stage; a universal approach for large-scale SACs production is required to meet the demands of practical applications;¹¹⁹ (3) some studies suggest that single-atom site configuration only shows high reactivity in the primary reaction process and is inadequate for the subsequent catalytic reaction

steps involving multiple electron transfer and intermediate conversion, accordingly, developing novel catalysts with dual atomic sites may overcome this limitation.^{120,121}

2.2.3 Metal nanoparticles modification. Metal nanoparticles (NPs) modified HSC material as active and reusable catalysts have attracted wide interest in the field of catalytic CO₂ reduction, in which metal NPs act as the active sites. The catalytic activity and selectivity can be tuned by the adjustment of their morphology,¹²² crystal facet,¹²³ and composition,¹²⁴ etc. For HSC supports, they can provide a unique space-confined structure to stabilize metal NPs; and they serve to prevent the agglomeration and leaching of the metal NPs, while allowing the diffusion of the reactant molecules to the active sites (metal NPs).

In view of the fact that HCS coupling with metal NPs can build a high-efficiency nanoreactor, Liu *et al.* designed a hollow



N-doped carbon nanocage catalyst with Ni nanoparticles being encapsulated within the cavity (Ni@NCN) (Fig. 6e).⁴⁶ Benefiting from nanoconfinement, the optimized sample (Ni₅@NCS) showed excellent eCO₂RR activity, achieving a high FE_{CO} of 93.2% in neutral media. Furthermore, Ni₅@NCS likewise exhibited decent performance in acid conditions, markedly exceeding the control samples of non-hollow-structured Ni/N-doped carbon sphere (Ni-NCS) and Ni-free N-doped carbon nanocage (NCN) (Fig. 6f). It is noteworthy that the FE_{CO} of Ni₅@NCS can achieve 84.3% even in the electrolyte of pH 2.5 at -1.4 V vs. Ag/AgCl (Fig. 6g). The superior activity is ascribed to the local pH-regulator effects of hollow carbon nanoreactor by concentrating the OH⁻ ions within the cavity (Fig. 6h). Specifically, the consumption rate of the H⁺ in the hollow carbon nanocage is faster than the replenishment rate of H⁺ because of the proton transport limitation, leading to a local alkaline environment and suppressing the competing hydrogen evolution reaction (HER). Recently, the effectiveness of metal NPs and HSC composite in electrochemical CO₂ reduction for CO production has also been demonstrated on the Ni NPs@N-doped nanotube systems.^{125,126} Previous studies have shown

that alloy NPs show superior performance over monometallic NPs in CO₂ activation and formate formation.^{36,127} Motivated by this, our group developed a PdCu alloy NPs confined within hollow carbon sphere catalyst (PdCu-N@HCS) (Fig. 7a and b), which showed an excellent thermal-catalytic CO₂ hydrogenation activity, delivering the highest turnover number hydrogenation activity, delivering the highest turnover number (TON) of 1432 at 100 °C for 24 h compared to single-metal-component samples (Fig. 7c);⁴⁹ DFT calculation result demonstrated that the alloying of Pd with Cu effectively reduces the energy barrier of the H₂ dissociation, and is capable of facilitating the HCO₃⁻ transformation.

In conclusion, metal NPs coupled with HSC showed satisfactory catalytic activity in some electrocatalysis and thermal catalysis applications. However, the activity improvement may still profit from the consideration of the following aspects. Incorporating HSC with novel high-entropy alloys (HEAs), a new class of multi-component alloys that possess numerous advantages (*e.g.*, flexible component adjustability, high oxidation resistance and excellent corrosion resistance), is expected to design high-efficiency catalysts. Recently,

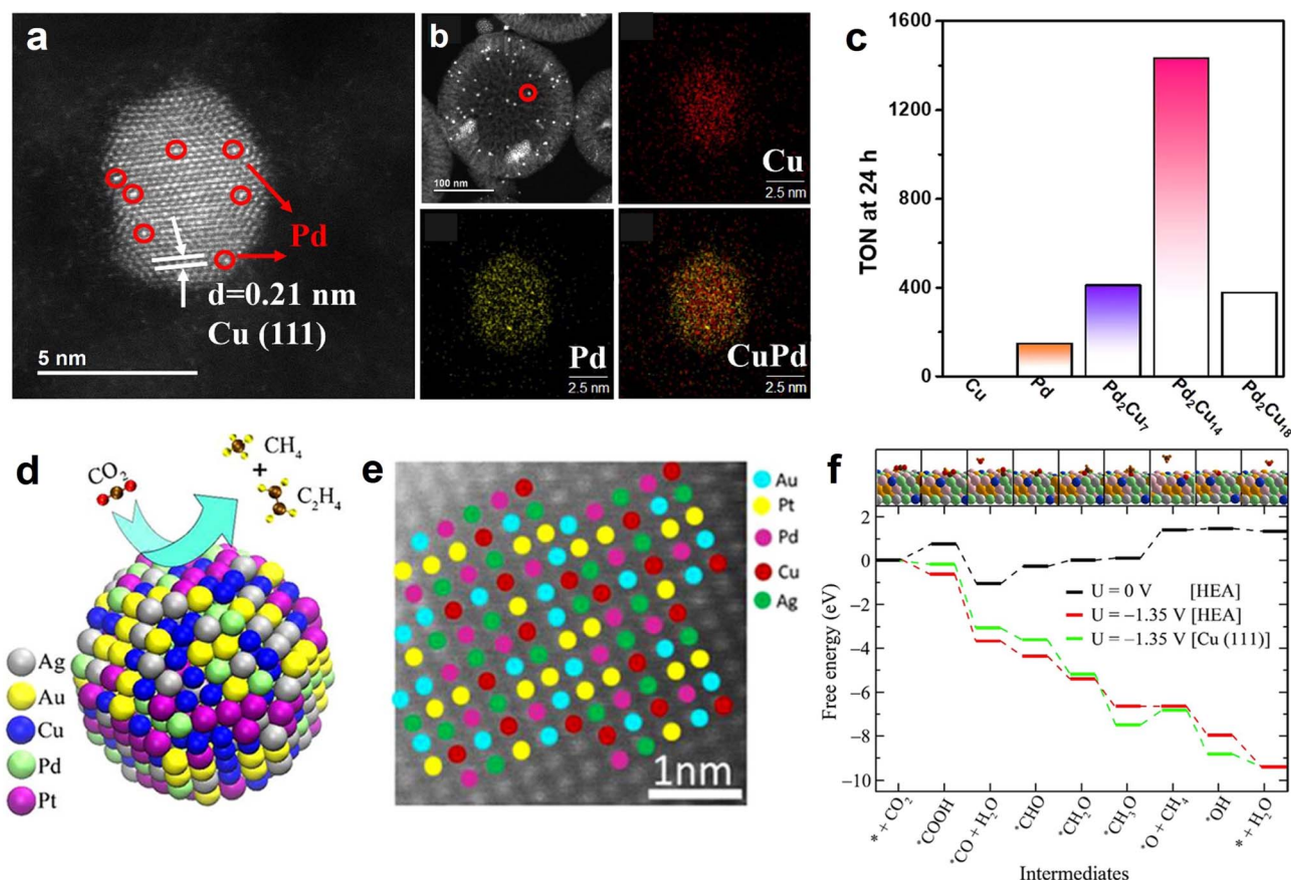


Fig. 7 (a) Magnified HAADF-STEM image of CuPd alloy NPs and (b) STEM image and the corresponding EDS elemental maps of Cu, Pd and CuPd of PdCu-N@MHCS sample. (c) Effect of the Pd/Cu molar ratio on Pd_xCu_y-N@MHCS for the hydrogenation of CO₂ to formate. Reproduced with permission from ref. 49. Copyright 2021, American Chemical Society. (d) Illustration of electrochemical CO₂ conversion and (e) HR-STEM image of HEA alloy NPs (AuAgPtPdCu). (f) Comparison of eCO₂RR free energy on AuAgPtPdCu HEA and Cu (111). The Pt, Pd, Ag, Au, Cu, C, O, and H atoms are shown in gray, green, pink, yellow, blue, brown, red, and orange, respectively. Reproduced with permission from ref. 128. Copyright 2020, American Chemical Society.



Nellaiappan *et al.*, delicately fabricated a HEA catalyst (AuAgPtPdCu) (Fig. 7d and e), which can be used for efficient electrochemical CO₂ conversion, with the FE of 67.5% for gaseous hydrocarbons (CH₄, C₂H₄) at -0.3 V vs. RHE;¹²⁸ the free energy diagram of eCO₂RR results showed that, at a given potential of -1.35 V, the reaction profiles of HEA and Cu (111) are downhill in general. Notably, at the *OCH₃ to *O transition step, the energy barrier on Cu (111) surface is obviously larger than that of HEA (Fig. 7f), suggesting that the thermodynamic favorability of eCO₂RR on the HEA system is higher than on pristine Cu. This work demonstrates that HEA can effectively alter the reaction energy barrier of the intermediates, and accelerate the CO₂ multi-electron reduction process. In another study, Pedersen *et al.* corroborated the feasibility that HEA material with optimized components and content can be employed as an efficient eCO₂RR catalyst, by using DFT calculation and machine learning.¹²⁹ In addition, the introduction of aminopolymers into metal NPs@HSC catalyst system is also worth studying, because aminopolymers contain plenty of amine groups with basicity that can promote CO₂ capture ability.¹³⁰ Another merit of aminopolymer is the anchoring of metal ions prior to the metal nucleation, thus facilitating the dispersion of metal NPs.¹³¹ Our previous works have proved the positive effects of aminopolymer in the nanoconfined reactor for CO₂ electroreduction and thermo-catalytic conversion.^{52,132} The above-mentioned directions may provide more possibilities for designing high-performance metal NPs@HSC-based catalysts.

3. Electrocatalytic CO₂ conversion of hollow carbon-based materials

Electrochemical CO₂ conversion to produce fuels and chemicals is considered as an ideal option for responding to the increasing energy supply requirement and environmental concerns.^{135,136} Typically, a CO₂ electroreduction cell comprises two electrodes: cathode (for eCO₂RR) and anode (for water oxidation). By applying an external bias, the eCO₂RR occurs at the three-phase interface of the cathode, electrolyte, and the adsorbed CO₂, with value-added products (CO, HCOO⁻, C₂H₄, *etc.*) being obtained. The commonly used reactors for eCO₂RR can be categorized into three types: (1) H-type cell (Fig. 8a), (2) liquid-flow cell (Fig. 8b), and (3) membrane electrode assembly (MEA) cell (Fig. 8c). The H-type cell involves the use of aqueous electrolytes, wherein the solubility of CO₂ is relatively low (approximately 34 mM). Therefore, CO₂ electrolysis on the H-type cell is susceptible to the influence of mass transport limitations, leading to low current densities (<100 mA cm⁻²).^{110,133} In the liquid-flow cell, an ion exchange membrane is used to separate the cathode and anode to prevent the CO₂ reduction products in the cathode from transferring to the anode and undergoing re-oxidation. Additionally, the anolyte and catholyte are continuously cycled. Different from the H-type cell, eCO₂RR occurs at the gas diffusion electrode (GDE), overcoming the solubility and mass transport limitations of CO₂ in aqueous electrolytes and enabling a high current density to be achieved (>100 mA cm⁻²).¹³⁴ In contrast to H-cell and liquid-flow cells,

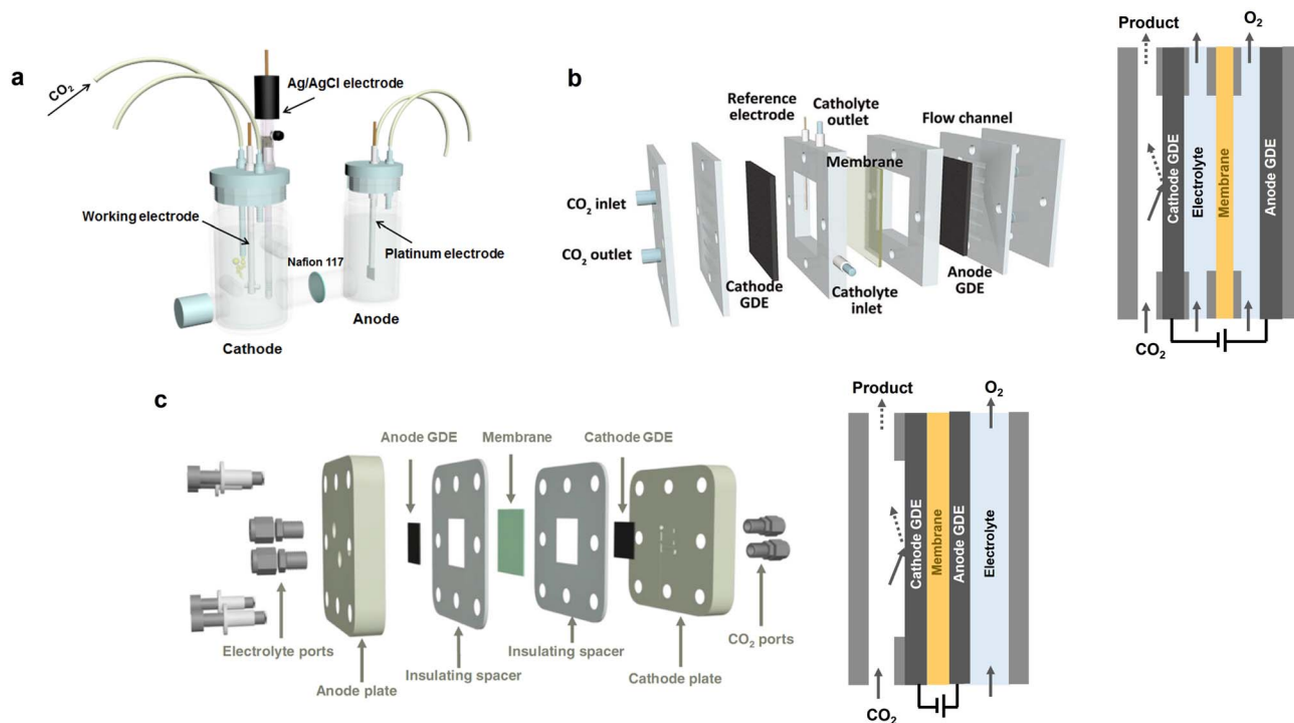


Fig. 8 Schematic illustrations of the eCO₂RR reactors. (a) H-cell. Reproduced with permission from ref. 133. Copyright 2018, Elsevier. (b) Liquid-flow cell. Reproduced with permission from ref. 134. Copyright 2021, Wiley-VCH. (c) MEA cell. Reproduced with permission from ref. 110. Copyright 2023, Springer Nature.



Table 3 Performance comparison of HSC-based catalysts for electrochemical CO₂ conversion

Catalyst	Faradaic efficiency (%)	Partial current density (mA cm ⁻²)	Potential (V vs. RHE)	Electrolyte	Reactor type	Ref.
Co-N ₅ /HNPCSS	CO, 99.4	~4.5	-0.73	0.2 M NaHCO ₃	H-type cell	114
Ni/N-doped carbon sphere	CO, ~80	~4	-0.78	0.5 M KHCO ₃	H-type cell	116
F-doped cage-like carbon	CO, 88.3	~32	-1.0	0.5 M KHCO ₃	H-type cell	102
Ni SAs/N-doped carbon sphere	CO, 95.1	~7.6	-0.8	0.5 M KHCO ₃	H-type cell	155
Ni-N ₄ /carbon sphere	CO, 95%	10.5	-1.0	0.5 M KHCO ₃	H-type cell	115
Ni/N-HCS	CO, 97%	10	-0.8	0.5 M KHCO ₃	H-type cell	117
N, B co-doped HCS	CO, 95.1	~1	-0.42	1.0 M KHCO ₃	H-type cell	104
Ni/N doped hollow carbon plates	CO, ~100	18.2	-1.0	0.5 M KHCO ₃	H-type cell	156
Ni-N ₃ /NHCSs	CO, 98.57	~14.2	-0.87	0.5 M KHCO ₃	H-type cell	146
Ni NPs/N doped carbon cage	CO, 93.2	17.1	-0.8	0.5 M KHCO ₃	H-type cell	46
Ni-N doped HGS-Micro/Meso	CO, 98.3	16.2	-1.0	0.5 M KHCO ₃	H-type cell	38
Ni SAs/N-doped urchin-like carbon	CO, 91	24.7	-0.8	0.5 M KHCO ₃	H-type cell	60
Hollow carbon with CoN ₄ O and ZnN ₄ dual atomic sites	CO, 92.6	15.57	-1.0	0.5 M KHCO ₃	H-type cell	157
Ni SAs/N-doped carbon sphere	CO, 94.91	15.35	-0.8	0.5 M NaHCO ₃	H-type cell	147
CO, 98.41	100	—	—	1 M KHCO ₃	Flow cell	
SnO ₂ /carbon sphere	Formate, 54.2	Formate, 3.7	-0.9	0.1 M KHCO ₃	H-type cell	158
	CO, 21.8	CO, 1.4				
Bi NRs@NCNT	Formate, 90.9	5.9	-0.9	0.1 M KHCO ₃	H-type cell	71
Cu/HCS	Formate, 82.4	26	-0.81	0.5 M KHCO ₃	H-type cell	41
SnS _{2-x} /NHCS	Formate, >80	~30	-1.2	0.5 M KHCO ₃	H-type cell	159
Cu cluster/HMCS	C ₂₊ products (C ₂ H ₅ OH and C ₂ H ₄), 88.7	~240	-1.0	1.0 M KOH	Flow cell	37
Cu ₂ O/NCNT	C ₂₊ products (C ₂ H ₅ OH and C ₂ H ₄), 77.61	16.79	-1.1	0.1 M KHCO ₃	H-type cell	160

the anode and cathode in MEA cell are directly compressed with the ion exchange membrane, forming a sandwich structure (Fig. 8c). This zero-gap structure significantly reduces the distance between the anode and cathode, thus minimizing ohmic resistance and markedly enhancing energy efficiency and stability.¹³⁷ Generally, the H-type cell is suitable for the fundamental study of eCO₂RR as well as the screening of efficient catalysts, due to the ease of operation and the rapid testing. Among the above-mentioned eCO₂RR cells, the MEA cell holds the most promising industrial potential for eCO₂RR, as it shows better energy utilization efficiency and stability. The development of more high-performance membranes will benefit the further improvement of the efficiency for MEA cells.

The choice of electrolytes is important to increase the activity and selectivity of electrochemical CO₂ reduction.^{138,139} Taking the widely used alkali metal bicarbonate electrolytes as examples, the size of alkali metal cation is able to affect the hydrolysis constants of hydrated cations, thereby altering the local pH value and CO₂ concentration around the electrode; eventually, it will cause a change of the selectivity toward C₁ products on Ag and C₂ products on Cu.¹⁴⁰ Additionally, the selectivity of eCO₂RR is also associated with the concentration of cations. For instance, Marcandalli *et al.* reported that at low potentials (>−0.4 V vs. SHE), a high concentration of Na⁺ cations is beneficial for enhancing the FE_{CO}, contrasting with the situation at more negative potentials (<−0.4 V vs. SHE), where a low concentration of Na⁺ cations is more favorable for suppressing hydrogen evolution and improving the FE_{CO}.¹⁴¹ In general, the electrolyte type can serve as a crucial design parameter, enabling the tailoring of the electrocatalyst–electrolyte interfacial environment to obtain desired products in the reduction of CO₂.

eCO₂RR involving multiple proton-coupled electron transfer processes usually requires overcoming high thermodynamic and kinetic barriers.^{142,143} The development of reliable electrocatalysts is of great significance to regulate the charge transfer and reaction intermediate adsorption/activation, overcoming the aforementioned obstacles, and achieving high catalytic activity and selectivity. As a result of recent research efforts, a series of high-performance HSC-based electrocatalysts have emerged (Table 3), which have creatively exploited the structural advantages of HSC, broadening the understanding of the mechanism behind eCO₂RR. In this section, the discussion mainly focuses on product selectivity with the aim of providing insights into the structure–activity relationships of HSC-based materials.

3.1 Selective production of CO or syngas (CO and H₂)

Atomically dispersed metal–nitrogen sites (M–N_x) modified carbon materials are widely studied in electrocatalysis, especially electrocatalytic CO₂ reduction, due to the well-tuned electronic characteristic and appropriate binding strength for the reaction intermediate. The merits of abundantly accessible sites, unique cavity space, and good permeability enable HSC materials to be appealing supports for anchoring the atomically dispersed M–N_x sites.

Notably, a majority of the recent studies focus on HSC-based catalysts with Ni–N_x sites and their electrochemical CO₂-to-CO conversion applications (Table 3). Herein, we use this type of material as the main line to introduce their CO₂-to-CO applications. To begin with, Xiong *et al.* developed a single-atom hollow carbon sphere catalyst containing Ni–N₄ atomic sites for eCO₂RR to CO.¹⁴⁵ The best-performing sample (Ni-HMCS-3-800) delivered an extremely high turnover frequency value of 15 608 h^{−1}, and the FE_{CO} above 95% at a wide potential range (−0.7 to −1.1 V vs. RHE) (Fig. 9a and b). It was demonstrated that the remarkable activity originated from the improved CO₂ adsorption and activation capacity, which are associated with the optimization of the geometrical structures of carbon support, including the shell thickness and compactness. Besides, this work also highlighted the importance of pore structure in eCO₂RR; the appropriate mesopore size distribution (~7–10 nm) endows Ni-HMCS-3-800 with enhanced diffusion and overflow of the reactants. As a result, Ni-HMCS-3-800 showed higher catalytic activity in comparison to the Ni-HMCS-3-700 sample with a smaller mesopore size (~3.1 nm).

Recently, the effects of the coordination environment of the central metal Ni–N_x on eCO₂RR activity have received considerable attention.¹⁴⁴ As documented in the literature, the energy barrier for *COOH intermediate formation on the Ni–N₃ site is lower than that of the Ni–N₄ site, suggesting coordinatively unsaturated Ni–N₃ may be favorable for eCO₂RR.¹⁴⁵ Accordingly, Gong *et al.* synthesized Ni–N₃ units embedded in nitrogen-doped hollow carbon spheres (Ni-NC/NHCSSs) by a hard template method using NiPc (Ni source) and dopamine (carbon and nitrogen sources) as precursors (Fig. 9c).¹⁴⁶ The optimized catalyst (Ni-NC/NHCSSs-600) exhibited a maximal FE_{CO} of 98.57% at −0.87 V vs. RHE, which is significantly larger than the value of its counterpart NiPc@NHCSSs (FE_{CO} = 19.61%). Additionally, in sharp contrast to NiPc@NHCSSs (stability < 2000 s), Ni-NC/NHCSSs-600 showed a long-term stability of 14 h with a negligible current attenuation. In a separate study, the researchers of the same group rationally developed a novel catalyst consisting of Ni–C₃N₁ moiety confined in hollow carbon reactors (I-Ni SA/NHCRs), which was reported to achieve a high FE_{CO} of 94.91% at −0.80 V in H-cell, and showing higher CO partial current density (−15.35 mA cm^{−2}) than that with outer Ni–C₂N₂ moiety (O–Ni SA/NHCRs, −12.06 mA cm^{−2}), or without hollow structure (Ni-NC, −5.39 mA cm^{−2}).¹⁴⁷ More importantly, a high FE_{CO} of up to 98.41% was achieved for I-Ni SA/NHCRs at 100 mA cm^{−2} in a flow cell with 1 M KHCO₃ electrolyte, implying a huge potential for practical application. It was proposed, based on COMSOL multiphysics finite element simulations, that the diffusion time of CO₂ molecules in hollow nano-reactor structure (model II, I-Ni SA/NHCRs-based electrode) (CO₂ diffusion time, ~1.25 s) is much faster than that of solid sphere structure (model I, Ni-NC-based electrode) (CO₂ diffusion time, >2.0 s) (Fig. 9d), as a result of the intrinsically structural virtue; in the meantime, the bottom and average concentration of CO₂ in the above two models was dynamically monitored *via* the simulation probe (Fig. 9e and f), and the results fully confirmed the facilitated CO₂ diffusion rate in hollow structural carbon. Obviously, it is conducive to the accessibility of isolated Ni



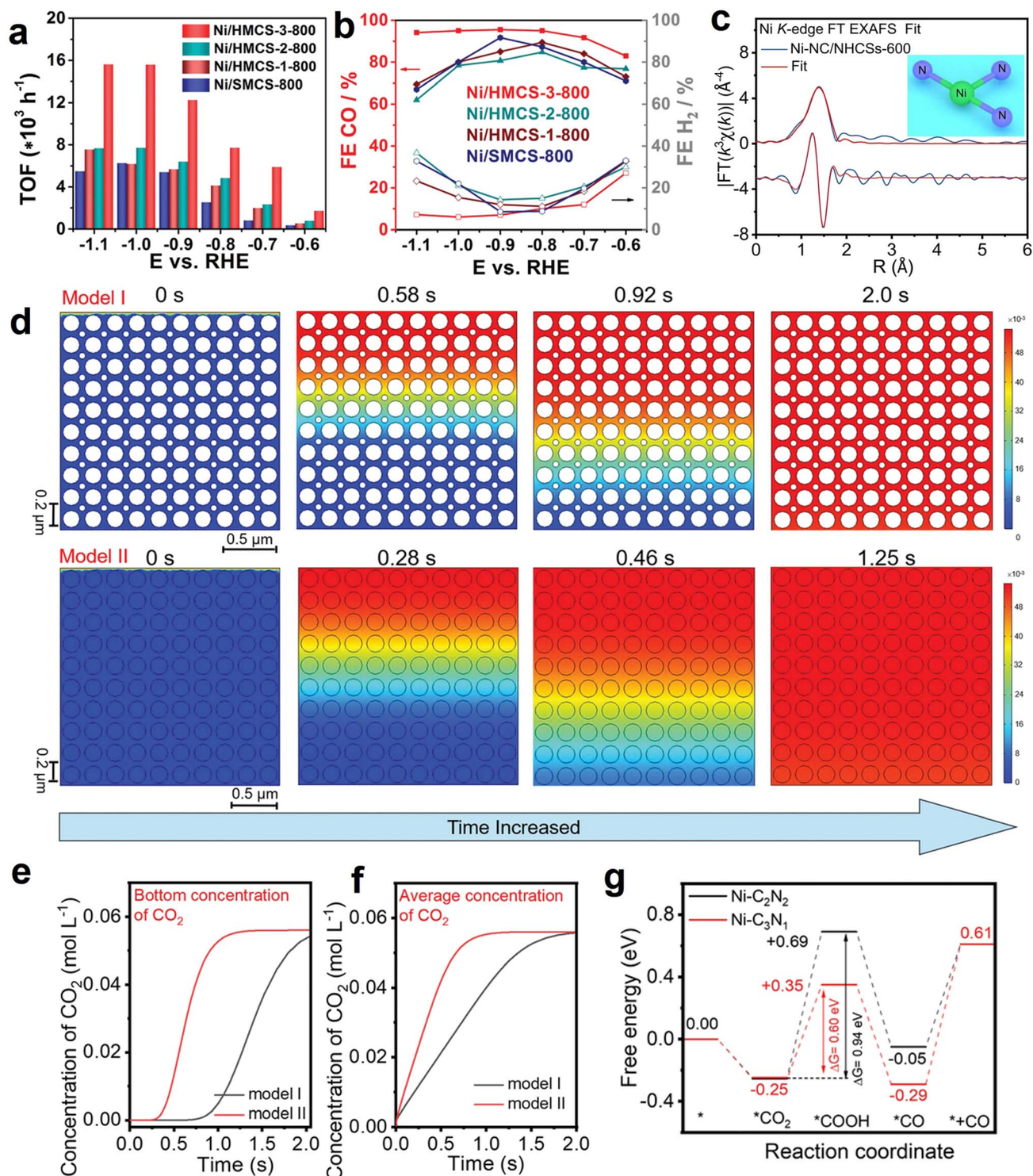


Fig. 9 (a) Turnover frequency (TOF) and (b) faradaic efficiencies of CO and H₂ of as-synthesized samples at given potentials. Reproduced with permission from ref. 115. Copyright 2020, Wiley-VCH. (c) Ni K-edge fitting curve of Ni-NC/NHCSs-600 in *R* space. The upper and lower traces represent the magnitude and imaginary part, respectively. The inset shows the model of Ni-N₃ moiety. Reproduced with permission from ref. 146. Copyright 2022, Elsevier. (d) COMSOL multiphysics finite element simulation results of CO₂ diffusion time in 0.5 M NaHCO₃ aqueous solution for solid sphere structure (model I, Ni-NC-based electrode) and hollow nano-reactor structure (model II, I-Ni SA/NHCRs-based electrode). (e) Bottom and (f) average concentration of CO₂ for model I and model II via probe monitoring. (g) Calculated eCO₂RR free energy profiles of Ni-C₃N₁ and Ni-C₂N₂ moiety. Reproduced with permission from ref. 147. Copyright 2023, Wiley-VCH.



active sites in I-Ni SA/NHCs to CO₂ molecules. The comparison of calculated free-energy diagrams for eCO₂RR of Ni-C₃N₁ and Ni-C₂N₂ configurations is shown in Fig. 9g, and the energy barrier for the protonation of *CO₂ to *COOH over Ni-C₃N₁ moiety (0.60 eV) is lower than the value of Ni-C₂N₂ structure (0.94 eV), implying that the formation of the key *COOH intermediate is more thermodynamically favorable on Ni-C₃N₁ configuration. These works provide inspiration for designing efficient HSC materials with tailored coordination of single atoms.

Despite most of the current studies being dedicated to improving CO selectivity and suppressing the competitive hydrogen evolution reaction from CO₂ reduction, the production of the mixture of CO and H₂ (syngas) is of great applied relevance. The syngas with specific ratios can serve as the feedstock for the synthetic chemical industry (*e.g.*, Fischer-Tropsch synthesis), especially in the range of 0.25–3.33 CO/H₂.¹⁴⁸ HSC-based materials are reported to function as feasible catalysts for controllable syngas production. For example, Song *et al.*, reported a nitrogen and cobalt co-doped hollow carbon catalyst (Co-HNC), with the Co-C₂N₂ moieties acting as catalytic sites for CO₂ reduction and the N functional groups (*i.e.*, pyridinic and graphitic N) serving as HER sites.¹⁴⁹ The obtained Co-HNC sample showed a faradaic efficiency of nearly 100% and a production rate of ~425 mmol g⁻¹ h⁻¹ for syngas evolution at -1.0 V *vs.* RHE. Notably, the syngas ratio is close to the desired 1/2 (CO/H₂) that is appropriate for downstream methanol production.¹⁵⁰ The hollow structure and sponge-like thin shell were claimed to be helpful for accelerating the mass transport and maximizing the catalytically active area, rendering the catalyst an excellent electrocatalytic performance. Additionally, in our previous work, a tunable syngas production

(CO/H₂ ratio, 1.09 to 2.54) could be achieved from eCO₂RR over a wide electrochemical window from -0.7 to -1.1 V *vs.* RHE, using an aminopolymer-modified hollow carbon sphere incorporating Ag NPs catalyst (Ag-P@HCS).⁵² Detailed characterizations suggested that the introduction of aminopolymers led to the finely increased dispersion of Ag NPs and improved CO₂ affinity of Ag-P@HCS, thus optimizing its eCO₂RR performance. The work emphasized the positive effects of aminopolymers in HSC-based eCO₂RR systems.

3.2 Selective production of formate

As an important liquid-phase product of eCO₂RR, formate has extensive applications in the pharmaceutical industry, animal feed, and hydrogen energy transportation.¹⁵¹ Nowadays, indium (In), copper (Cu), and tin (Sn)-containing catalysts are widely studied in eCO₂RR for CO₂-to-formate production,^{152–154} owing to the favorable *OCHO formation and suitable binding intensity with *HCOOH; but some limitations towards pragmatic application still exist, such as unsatisfied selectivity, moderate stability, and large overpotential. In fact, except for the shape and constituent control of the metal/oxides NPs, the electrocatalytic performance is also determined by the electronic and morphologic characteristics of carbon substrates to some extent. Encouragingly, some interesting works have been reported on constructing metal-carbon hollow nano-structured hybrids with space-confinement effects and abundant exposed active sites to address the problems and achieve desirable electro-reactivity. As exemplified by the study conducted by Zhang *et al.*, the pipet-like bismuth (Bi) nanorods semifilled in N-doped carbon nanotubes (Bi-NRs@NCNTs) catalyst (Fig. 10a) delivered a highly efficient CO₂-to-formate electro-conversion,

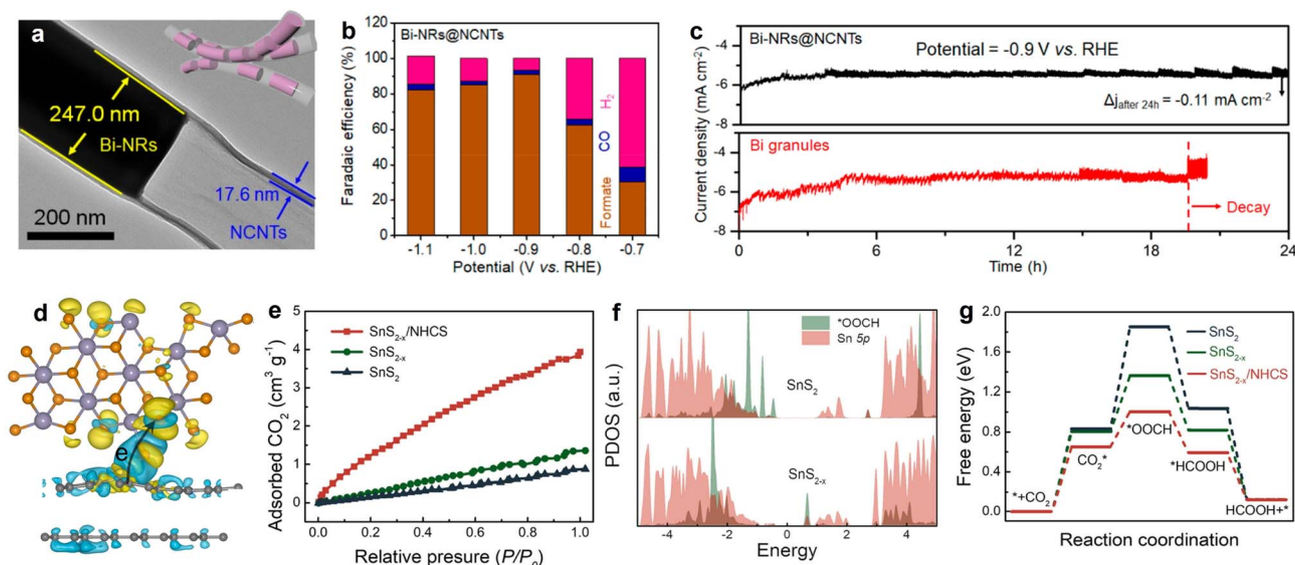


Fig. 10 (a) HRTEM image and morphology illustration (inset) of Bi-NRs@NCNTs. (b) Faradaic efficiencies of the eCO₂RR products of Bi-NRs@NCNTs. (c) Stability comparison of Bi-NRs@NCNTs and Bi granules. Reproduced with permission from ref. 71. Copyright 2021, American Chemical Society. (d) Charge density difference diagram SnS_{2-x}/N-doped carbon. (e) CO₂ adsorption isotherms. (f) Partial density of states (PDOS) of *OOCH and Sn 5p orbital for SnS₂ and SnS_{2-x}. (g) Calculated CO₂RR free energy profiles. Reproduced with permission from ref. 159. Copyright 2022, Elsevier.



showing a high selectivity for formate production with the peak faradaic efficiency value of 90.9% at -0.9 V vs. RHE (Fig. 10b).⁷¹ Different from Bi granules, no obvious current density attenuation was observed on Bi-NRs@NCNTs throughout the 24 h electrolysis (Fig. 10c), implying its structural superiority. Specifically, the spatial restriction function of NCNTs efficiently prevents the self-aggregation and oxidation of metallic Bi nanorods, ensuring good durability during long-term testing. Meanwhile, the results demonstrated that the synergy of nanocapillary and nanoconfinement effects induced by NCNTs contributed to the accelerated mass transfer, improved CO₂ adsorption, and concentrated CO₂ around the active sites. This work not only highlights the functions of HSC material in CO₂ valorization, but also provides a promising design philosophy on HSC-based material for energy conversion and beyond.

In another example, Du *et al.* developed a hollow nanostructured hybrid electrocatalyst consisting of Cu, Cu₂O, and nitrogen-doped carbon sphere (HCS/Cu) for highly selective formate production from eCO₂RR.⁴¹ The best-performing sample HCS/Cu-0.12 showed 82.4% FE_{formate} with a current density of 26 mA cm⁻² at -0.81 V vs. RHE, in which the optimized Cu/C heterogenous interfaces could be responsible for the enhanced electron transport and the resulting excellent activity.

Very recently, nitrogen-doped hollow carbon spheres (NHCS) were employed to couple with tin disulfide containing sulfur vacancies to construct a catalyst (SnS_{2-x}/NHCS) with improved CO₂ activation ability and excellent conductivity.¹⁵⁹ DFT simulation analysis confirmed that a migration channel of electrons was built with the direction from NHCS to SnS_{2-x} (Fig. 10d). Benefited from the synergy of S defects and the integration with NHCS, SnS_{2-x}/NHCS exhibited the highest uptake of CO₂ (Fig. 10e). Moreover, simulations of projected density of states (PDOS) for *OOCH and Sn 5p on SnS_{2-x} show a larger overlapping area than on its SnS₂ counterpart, indicating the stronger interaction between *OOCH and SnS_{2-x} (Fig. 10f). In addition, significantly decreased energy barriers for *CO₂ formation and *OOCH generation can be observed on SnS_{2-x}/NHCS compared with those of SnS₂ and SnS_{2-x}, signifying the promoted CO₂ activation and conversion capacities (Fig. 10g). As a consequence, the partial current density and faradaic efficiency of SnS_{2-x}/NHCS for formate production increased to 3.4 and 1.2 times, respectively, in comparison to the value of pristine SnS₂. In general, the work disclosed the fundamental correlation between catalyst structure and performance, which may trigger more consideration of the material design by coupling defect engineering and HSC material with matched energy-level structure.

3.3 Selective production of multicarbon products

Of the prevalent eCO₂RR HSC-based electrocatalysts, most mainly produce C1 products (CO or HCOO⁻), as presented in Table 3. The production of energy-dense and valuable multicarbon (C₂₊) oxygenates and hydrocarbons, such as ethylene (C₂H₄) and ethanol (C₂H₅OH), is of great value for industrial manufacture, but the relevant reports are relatively rare; the

primary obstacle lies in the difficulty of coupling C–C bonds.¹⁶¹

Cu-based catalysts are of particular interest in the field of eCO₂RR and are capable of converting CO₂ to a series of C₁/C₂ products. However, suffering from product complexity and unsatisfactory activity, more efficient Cu-based catalysts are required. Considering the structural features of HSC materials for the carbonaceous intermediates, Pan *et al.* designed hollow mesoporous carbon spheres confining Cu cluster catalyst (Cu/HMCS) for improving the C₂ selectivity.³⁷ *In situ* Fourier transform infrared spectroscopy (FTIR) analysis revealed that, under the aid of nanoconfinement provided by HMCS, with suitable Cu cluster loading, the conversion of *CO to *CHO could be facilitated; additionally, the nanocavity protection effects of HMCS allowed to boost the C–C bond coupling to generate C₂ products. As a result, the sample (Cu/HMCS-20%) with optimized conditions exhibits a C₂ (C₂H₄ and C₂H₅OH) faradaic efficiency of 88.7% at -1.0 V vs. RHE. This work provides insights for designing robust catalysts with high C₂ selectivity.

In addition to hollow carbon spheres, carbon nanotubes (CNTs) are also commonly used in electrocatalysis; they can not only offer excellent conductivity, but also prevent the agglomeration of catalytic components.¹⁶² Liu *et al.* investigated the surface modification of N-doped CNTs on Cu₂O NPs for enhanced CO₂-to-C₂ electro-conversion.¹⁶⁰ The Cu₂O NPs-NCNT sample showed a maximal C₂ (C₂H₄ and C₂H₅OH) faradaic efficiency of 77.61% at -1.1 V vs. RHE, suppressing the value of the NCNT-free sample (Cu₂O NPs, FEC₂ 38.15%); noteworthy, the C₂ partial current density of Cu₂O NPs-NCNT (16.79 mA cm⁻²) is 7.30 folds higher than that of Cu₂O NPs (2.30 mA cm⁻²) at the same applied potential. It is proposed that NCNT is helpful for catalyzing the conversion of CO₂ to CO, which subsequently enriches the CO concentration at the heterogeneous interface of NCNT and Cu₂O NPs and is conducive to C–C coupling to generate C₂ products.

Recently, some strategies (*e.g.*, alloying and surface doping) have been demonstrated to enhance the FE towards C₂₊ production of Cu materials. For instance, Chen *et al.* reported Cu–Ag tandem alloy catalysts, and the results illustrated that the CO-enriched environment introduced by Ag facilitated the subsequent C–C coupling processes occurring at the Cu sites, thereby enhancing the formation of C₂₊.¹⁶³ Additionally, Wu *et al.* found that employing molecular doping strategies, including the introduction of aromatic heterocycles (*e.g.*, thiazole and triazole derivatives) to CuAg alloy, can orient the reaction route towards C₂₊ production, achieving a high FE of C₂₊ ($\approx 80\%$); detailed analyses demonstrated that the functional groups with electron-withdrawing feature are capable of modulating the electronic structure of Cu atoms and thereby enhancing the catalytic reaction rate.¹⁶⁴ In general, incorporating modern strategies such as alloying and molecular doping into the design of copper-containing HSC-based catalysts, alongside the optimization of component interactions, holds the potential to further enhance the efficiency of catalyzing C₂₊ production and accelerate the pace of approaching practical application.



4. Thermocatalytic CO₂ conversion of hollow carbon-based materials

Thermocatalytic CO₂ conversion to produce commodity chemicals is a promising way to alleviate the dependence on fossil resources; it can be classified as hydrogenation and reforming routes loosely. For the reforming reaction, CO₂ serves as an oxidant for catalytically reforming methane to generate syngas (CO and H₂) under high temperature condition (600–1000 °C).¹⁰ Nevertheless, the extremely high temperature of the reforming reaction may lead to the decomposition of the carbon material; and there are few researches of HSC-related material reported in this field. As for CO₂ hydrogenation, it refers to the conversion of CO₂ with H₂ to formic acid, methane and olefins, *etc.*, under a relatively mild ambient condition. CO₂ hydrogenation is in line with the concept of green chemistry, and holds important industrial application prospects, therefore, many efficient HSC-based materials were designed for this reaction (Table 4). In this section, the recent applications of HSC-based materials in relatively popular CO₂ hydrogenation applications will be summarized and discussed.

4.1 CO₂ hydrogenation to formate

Formate synthesis from CO₂ hydrogenation is regarded as one of the most sustainable and economically feasible for utilization of CO₂. However, designing efficient catalysts with high stability for formate production from CO₂ hydrogenation is challenging because of the difficulty of CO₂ activation and harsh reaction conditions. In this regard, our group has developed a series of hollow carbon sphere-based materials *via* rationally engineering the catalyst component/structure, and achieved desirable performance. Compared to the conventional supported material, encapsulation effects of the cavity in hollow carbon spheres efficiently prevent the leaching of some functionalized modifiers (*e.g.*, aminopolymer) and the active metals, resulting in improved robustness and formate conversion efficiency. To begin with, a nanostructured catalyst consisting of sub-3-nm-sized PdAg NPs with high dispersity and aminopolymers encapsulated in mesoporous hollow carbon spheres (PdAg-P@MHCS) was synthesized (Fig. 11a–e).⁵¹ MHCS provided immobilization spaces for aminopolymers; and the amine groups of polymers promoted the metal ions adsorption, enabling the selective formation of PdAg NPs within P@MHCS. As a result, the turnover number (TON) of optimized PdAg-P@MHCS for formate production reached 2680 at 100 °C for 24 h. With the combination of experimental characterization and kinetics analysis, the possible reaction mechanism was proposed, as illustrated in Fig. 11f. Firstly, H₂ dissociation occurs and further forms Pd-hydride species, which are active in the hydrogenation process (step 1). Then, the bicarbonate (HCO₃[−]) adsorbs onto Ag atoms that are in an electron-deficient state after alloying with Pd (step 2); meanwhile, CO₂ dissolution in the alkaline reaction solution continuously provides HCO₃[−] (step 3). The dissociative formate is formed under the attack of active H to HCO₃[−] adsorbed on Ag (step 4). This work demonstrated a new

design strategy for developing highly efficient CO₂ hydrogenation catalysts for formate production.

To alleviate the use of hazardous reagents (*i.e.*, carbon tetrachloride) and the complex operation procedures in the aforementioned work, we further exploited a novel and facile synthesis method for monodispersed N-doped microporous hollow carbon spheres encapsulating PdAg alloy catalyst *via* an ethylenediamine-assisted pyrolysis.³⁶ And it showed competitive CO₂ hydrogenation activity with the TON value of 2750 at 100 °C for 24 h.

In addition to alloy, the MHCS was also employed to confine a composite of mononuclear Ru³⁺ and N, P-containing porous organic polymers (Ru³⁺-POPs) in another study of ours (Fig. 11g).⁵⁰ The Ru³⁺-POPs@MHCS catalyst with optimized content of Ru³⁺-POPs exhibited good activity for CO₂ hydrogenation to formate (TON = 1228 at 100 °C for 24 h), surpassing that of Ru³⁺-POPs (without MHCS) sample (Fig. 11h). The superior performance originated from the promoted adsorption abilities of CO₂ and H₂ induced by MHCS, as evidenced by the kinetics analysis (Fig. 11i and j). Furthermore, the Arrhenius curve displayed that the activation energy of hydrogenated CO₂-to-formate conversion decreased with Ru³⁺-POPs being encapsulated in MHCS (Fig. 11k). Based on the above discussion and reasoning, we demonstrate here that hollow carbon spheres benefit the overall performance of formate production from CO₂ hydrogenation. The optimization of nanoconfined space is still worthy of further study, and our relevant researches are in progress.

4.2 CO₂ methanation

Methane synthesis from CO₂ hydrogenation (CO₂ + 4H₂ → CH₄ + 2H₂O, ΔH° = −165 kJ mol^{−1}), known as the Sabatier reaction, is an exothermic process that is more thermodynamically favorable than many other hydrogenation routes.^{21,171} However, it still suffers from significant kinetic limitations due to the intrinsic inertness of linear CO₂ and the involvement of an eight-electron transfer process.^{165,172,173} Several metal-containing catalysts have been demonstrated as effective candidates for this reaction, typically based on Ru, Rh, Pd, and Ni NPs on supported materials, with Ni-based catalysts being the most popular one because of the low cost and rich reserves.^{174–176} Nevertheless, the high temperature in the catalytic reaction may give rise to issues of stability and sintering in Ni-based catalysts. The incorporation of hollow structured carbon enables the encapsulation or embedding of metal sites inside its structure, which can effectively alleviate the problems of coke formation and metal sintering, and ultimately improve the activity, stability, and selectivity for methane synthesis from CO₂ hydrogenation.^{42,177} For instance, Lin *et al.*, presented a MOF-derived hollow carbon nanostructured catalyst encapsulating dispersed Ni nanoparticles (Ni@C) which affords better CO₂ methanation performance than Ni/C (control sample, Ni NPs on graphite carbon) owing to the unique carbon-confined configuration.¹⁶⁶ Additionally, no obvious deactivation of Ni@C catalyst was observed in the long-term stability test at 250 °C for 24 h; in contrast, Ni/C without carbon shell protection displayed a 50% decline in CH₄ yield by 24 h.



Table 4 Summary of HSC-based catalysts for thermocatalytic CO₂ conversion

Catalyst	Target product	Reaction condition	Catalytic performance	Ref.
NiCe/CNT	CH ₄	0.1 g catalyst, 350 °C, atmospheric pressure, H ₂ /CO ₂ = 4 : 1 (flow rate), continuous fixed-bed, gas hourly space velocity (GHSV) = 30 L g ⁻¹ h ⁻¹	CO ₂ conversion rate: 83.8% Selectivity: 99.8%	165
Ni NPs/N-CNT	CH ₄	0.2 g catalyst, 360 °C, atmospheric pressure, H ₂ /CO ₂ = 6 : 1, continuous fixed-bed, GHSV = 120 L g ⁻¹ h ⁻¹	CO ₂ conversion rate: 92% Selectivity: 98%	45
Ni NPs@carbon hollow sphere	CH ₄	0.1 g catalyst, 325 °C, atmospheric pressure, H ₂ /CO ₂ = 4 : 1, fixed-bed flow reactor, GHSV = 33 L g ⁻¹ h ⁻¹	CO ₂ conversion rate: ~100% Selectivity: 99.9%	166
Ni NPs/NCNT	CH ₄	0.04 g catalyst, 340 °C, atmospheric pressure, H ₂ /CO ₂ /Ar = 4 : 1 : 0.6, fixed-bed flow reactor, GHSV = 50 L g ⁻¹ h ⁻¹	CO ₂ conversion rate: ~50% Selectivity: 95%	42
Ni NPs/NCNT	CH ₄	0.1 g catalyst, 400 °C, atmospheric pressure, H ₂ /CO ₂ /He = 4 : 1 : 5, weight hourly space velocity (WHSV) = 60 L g ⁻¹ h ⁻¹	CO ₂ conversion rate: 81.2% Selectivity: 99.2%	162
PdAg-aminopolymer@MHCS	Formate	100 °C, 2 MPa (1 MPa H ₂ and 1 MPa CO ₂), 15 mL 1.0 M NaHCO ₃ aqueous reaction solution, closed reactor, 24 h	Turnover number (TON): 2680 Turnover frequency (TOF): 112 h ⁻¹	51
PdAg@NHCS	Formate	100 °C, 2 MPa (1 MPa H ₂ and 1 MPa CO ₂), 15 mL 1.0 M NaHCO ₃ aqueous solution, closed reactor, 24 h	TON: 2750 TOF: 115 h ⁻¹	36
PdCu@MHCS	Formate	100 °C, 2 MPa (1 MPa H ₂ and 1 MPa CO ₂), 15 mL 1.0 M NaHCO ₃ aqueous solution, closed reactor, 24 h	TON: 1432 TOF: 60 h ⁻¹	49
Ru ³⁺ -N,P-containing porous organic polymers@MHCS	Formate	120 °C, 4 MPa (2 MPa H ₂ and 2 MPa CO ₂), 7.8 mL 1.0 M NaHCO ₃ aqueous solution, closed reactor, 24 h	TON: 1228 TOF: 51 h ⁻¹	50
Ru/N-MHCS	Formate	120 °C, 8 MPa (4 MPa H ₂ and 4 MPa CO ₂), 40 mL 3 M aqueous Et ₃ N solution, closed reactor, 2 h	TON: 7550 TOF: 3775 h ⁻¹	118
Fe@CNTs	C ₂₊ products	0.4 g catalyst, 370 °C, atmospheric pressure, H ₂ /CO ₂ = 3 : 1 (flow rate), fixed-bed reactor	CO selectivity: 45.1% CH ₄ selectivity: 29.3% C ₂ -C ₄ selectivity: 24.3% C ₅₊ selectivity: 1.3%	167
FeZn-NC (Fe@C matrix)	C ₂₊ products	0.5 g catalyst, 320 °C, 3 MPa, H ₂ /CO ₂ = 3 : 1 (molar ratio), fixed-bed reactor, space velocity = 7200 mL g ⁻¹ h ⁻¹ , 5 h	CO selectivity: 19.9 C mol% CH ₄ selectivity: 16.6 C mol% Long-chain hydrocarbons (C ₅₊) selectivity: 27.4 C mol% Light olefins (C ₂ -C ₄) selectivity: 30.0 C mol% CO selectivity: 21.2 C mol% C ₅₊ selectivity: 22.1 C mol% C ₂ -C ₄ selectivity: 32.0 C mol% Olefins selectivity = 62.3% (C ₅₊ selectivity = 39.8%, C ₂ -C ₄ selectivity = 22.5%)	168
FeZnK-NC (Fe inserted in carbon shell)	C ₂₊ products	0.2 g catalyst, 340 °C, 2 MPa, H ₂ /CO ₂ /N ₂ = 72 : 24 : 4 (volume ratio), fixed-bed reactor, GHSV = 9000 mL g ⁻¹ h ⁻¹ , 24 h	FTY (iron-time yield) for heavy olefins = 26.5 μmol _{CO₂} g _{Fe} ⁻¹ s ⁻¹ Olefins selectivity = 52.5% (C ₅₊ selectivity = 21.8%, C ₂ -C ₄ selectivity = 30.7%)	169
FeK/SWNT	C ₂₊ products	0.2 g catalyst, 340 °C, 2 MPa, H ₂ /CO ₂ /N ₂ = 72 : 24 : 4 (volume ratio), fixed-bed reactor, GHSV = 9000 mL g ⁻¹ h ⁻¹ , 24 h	FTY (iron-time yield) for heavy olefins = 10.2 μmol _{CO₂} g _{Fe} ⁻¹ s ⁻¹ CO selectivity = 18 mol% CH ₄ selectivity = 35 mol% C ₂₋₄ selectivity = 35 mol% C ₅₊ selectivity = 12 mol% Iron-time yield = 12.2 mol _{CO₂} g _{Fe} ⁻¹ s ⁻¹	170
FeK/MWNT	C ₂₊ products	0.5 g catalyst, 350 °C, 8.5 MPa, H ₂ /CO ₂ = 2 : 1, fixed-bed reactor, GHSV = 14.4 L g ⁻¹ h ⁻¹		

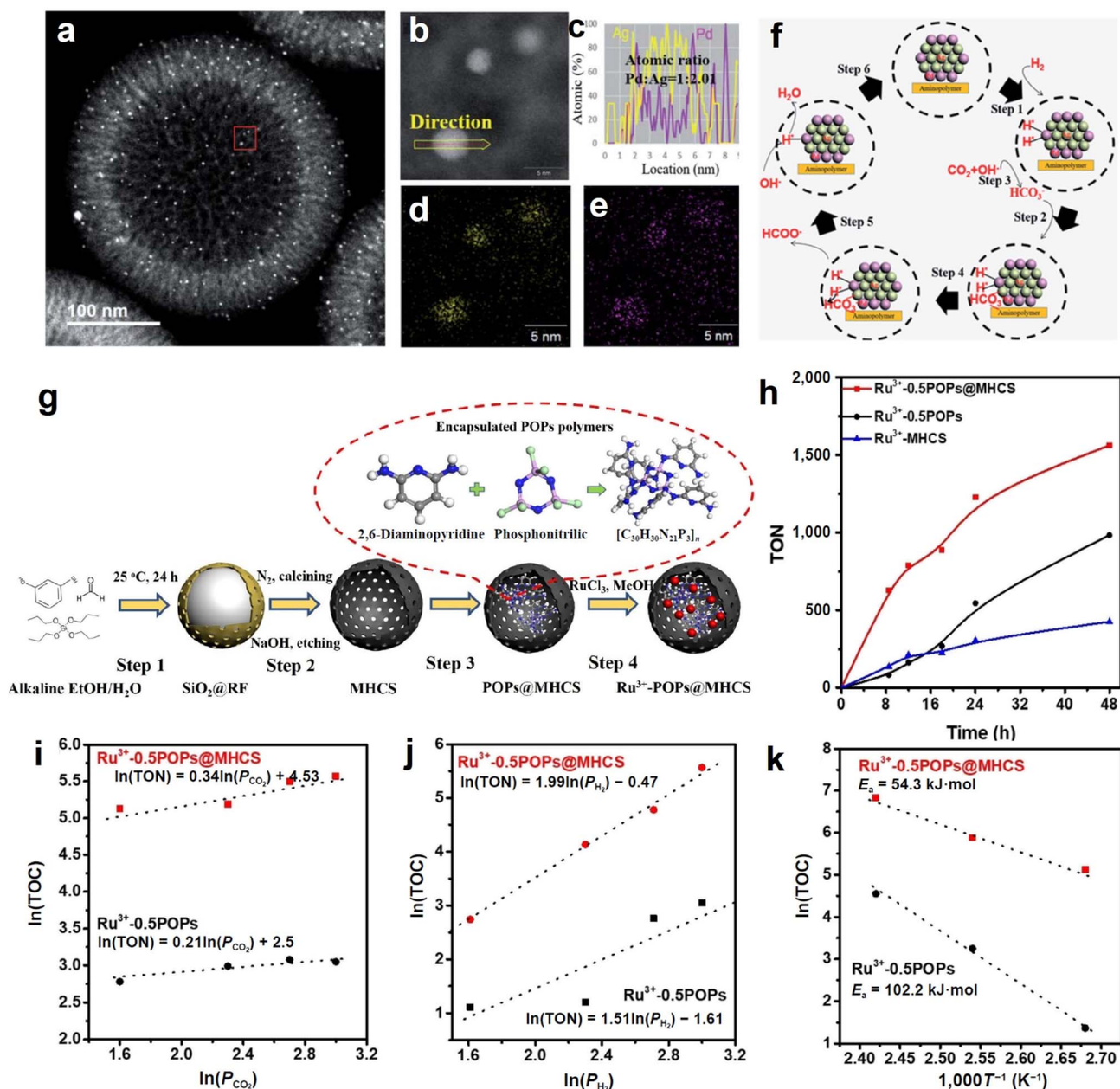


Fig. 11 (a) HAADF-STEM image of PdAg-P@MHCS. (b) Magnified image of the red frame zone in (a). (c) Elemental line scan across the PdAg NPs in (b). (d and e) Elemental maps of (d) Ag and (e) Pd. (f) Hydrogenated CO₂-to-formate conversion pathway over PdAg-P@MHCS catalyst. Reproduced with permission from ref. 51. Copyright 2020, Royal Society of Chemistry. (g) Synthesis of Ru³⁺-POPs@MHCS catalyst. (h) Reaction time profile of hydrogenated CO₂-to-formate conversion over the as-prepared samples. (i–k) Double logarithm plots of TON and the partial pressure of (i) CO₂ and (j) H₂ in the hydrogenation process, and (k) Arrhenius plots over optimized Ru³⁺-0.5POPs@MHCS and Ru³⁺-0.5POPs (MHCS-free) samples. Reproduced with permission from ref. 50. Copyright 2021, Tsinghua University Press and Springer-Verlag GmbH Germany, part of Springer Nature.

Generally, the excessive local heat generated during CO₂ methanation may result in the formation of “hot spots” that affect the stability duration and methane selectivity. Introducing an appropriate carbon material with good thermal conductivity can benefit the uniform dispersion of the reaction heat and ensure the methanation process. For example, Wang *et al.* reported that Ni NPs decorated N-doped carbon nanotubes (N-CNT) catalyst (Ni/N-CNT) (Fig. 12a and b) exhibited high

methane selectivity of 98% and CO₂ conversion rate of 92%, even at a high gas hourly space velocity (GHSV) of 120 000 mL g⁻¹ h⁻¹ with a H₂-to-CO₂ v/v ratio of 6 (Fig. 12c).⁴⁵ The existence of N-CNT with unique characteristics contributed to the superior performance because (1) the excellent thermal conductivity limited the generation of local “hot spots”, (2) the nanotube configuration optimized the diffusion pathway of the reactants, and (3) the alkalized surface of N-CNT is favorable for



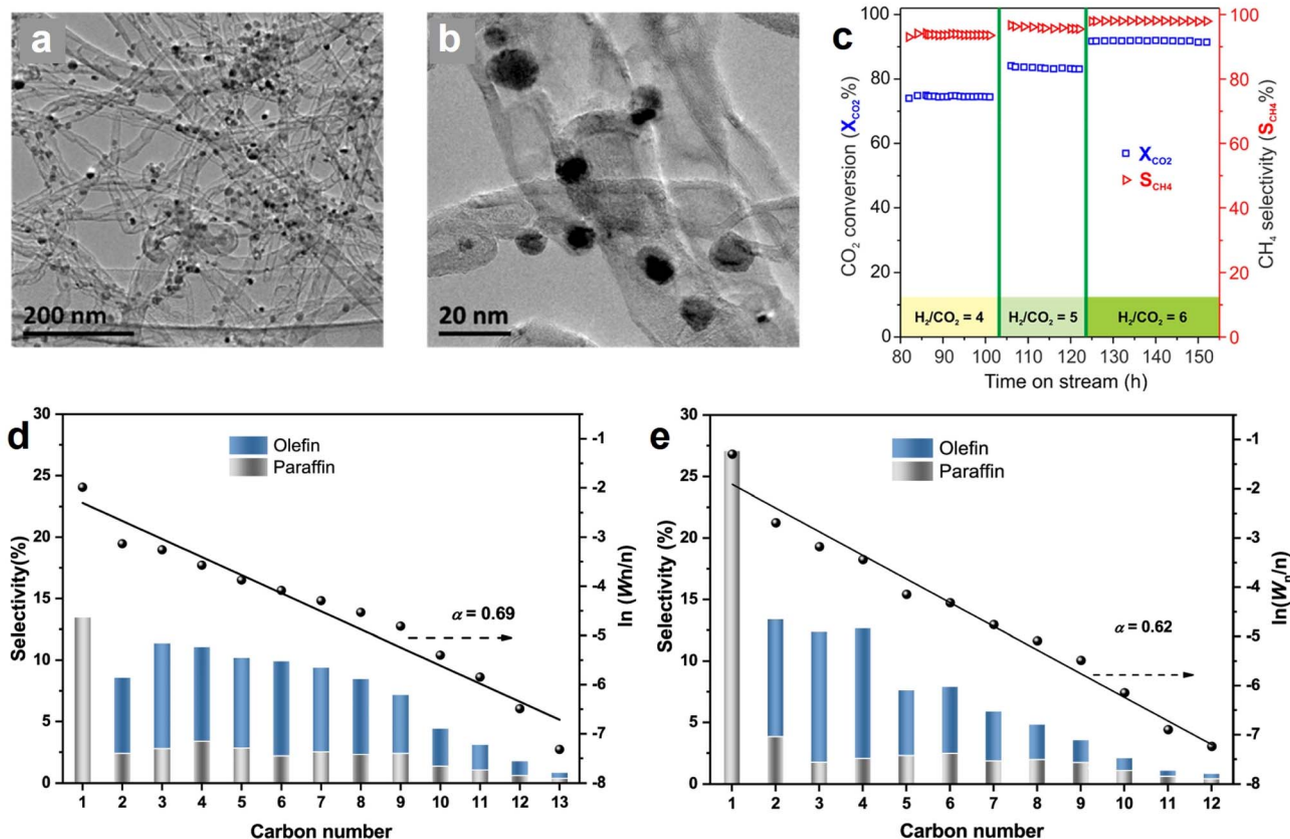


Fig. 12 (a and b) TEM images of Ni/N-CNT catalyst in different magnifications. (c) Influence of H₂/CO₂ ratio on the CO₂ methanation activity and selectivity Ni/N-CNT catalyst at a GHSV of 120 000 mL g⁻¹ h⁻¹. Reproduced with permission from ref. 45. Copyright 2018, American Chemical Society. Hydrocarbon production distributions over the (d) FeK/SWNTs and (e) FeK/MWNTs catalysts after CO₂ hydrogenation test for 24 h. Reaction conditions: 0.2 g of catalyst, $T = 613$ K, $P = 2.0$ MPa, H₂/CO₂ = 3, GHSV = 9000 mL g_{cat}⁻¹ h⁻¹. Anderson-Schulz-Flory (ASF) plots and α -values (chain propagation probability) are also presented; W_n represents the weight fraction of a hydrocarbon with n carbon atoms. Reproduced with permission from ref. 169. Copyright 2021, American Chemical Society.

concentrating CO₂ in the proximity of catalytically active Ni sites. Recently, another study disclosed the importance of N-doping catalyst towards CO₂ methanation, and the results demonstrated that N-doping enabled the formation of highly dispersive Ni sites that were responsible for the enhanced performance.¹⁶² Inspired by these intriguing studies, in our opinion, it is deemed to be of great significance to conduct more comprehensive investigations of the metal-support interaction, namely, the correlation between the intrinsic characteristics of hollow carbon internal/external surfaces and the active metals.

4.3 CO₂ hydrogenation to multicarbon products

Among the potential products from CO₂ hydrogenation, C₂₊ hydrocarbon products such as olefins and paraffins are of particular interest due to their large energy density and industrial relevance.^{178,179} CO₂ hydrogenation to hydrocarbons commonly involves reverse water-gas shift (RWGS) reaction and subsequent Fischer-Tropsch synthesis (FTS); the tandem of these two processes is thermodynamically feasible, but requires an effective catalyst to drive.¹⁸⁰ The carbon-supported Fe-based catalysts receive considerable attention because they show good activity in both RWGS and FTS.^{168,181} For the potential hollow

carbon support candidates, the CNTs with tubular morphology is the most popular one, owing to their good adhesion to metal NPs, high mechanical strength, good thermal conductivity, and ease of surface functionalization.^{182,183}

In 2013, Fe NPs@CNT catalyst was proved to be active for CO₂ hydrogenation to produce hydrocarbons by O'Byrne *et al.*, but the C₂₊ products selectivity and iron time yield (FTY) were relatively low.¹⁶⁷ Encouragingly, an iron-potassium catalyst supported on single-wall carbon nanotubes (FeK/SWNT) reported by Wang *et al.* delivered an extremely competitive CO₂ hydrogenation performance, with the olefins selectivity achieving 62.3%, where the heavy olefins (C₅₊) selectivity and light olefins (C₂-C₄) were 39.8% and 22.5%, respectively.¹⁶⁹ Fig. 12d and e show the comparison of CO₂ hydrogenation results after 24 h measurement over FeK/SWNT and FeK/MWNT (control sample with multi-walled carbon nanotubes). According to the distribution of the hydrocarbons, both of the samples show higher selectivity for olefins instead of paraffins. Notably, FeK/SWNTs favor the formation of heavy olefins, whereas the production of light olefins is preferred on FeK/MWNT (see Table 4 Row 14 and 15 for more details), as reflected by the larger α -values (chain propagation probability)



of FeK/SWNTs (0.69) as compared to that of FeK/MWNT (0.62). Detailed characterizations demonstrated that the excellent catalytic activity of FeK/SWNTs catalyst was associated with the presence of SWNTs with large curvature, which manipulated the electron distribution¹⁸⁴ and accelerated the dissociation of CO on Hägg carbide, eventually boosting CO₂ hydrogenation to heavy olefins. In extension to the conventional CNTs, the dense CNT materials are suitable for specific catalytic applications. Chernyak *et al.* prepared a new type of densely packed CNTs with embedding Fe NPs catalyst (Fe/CNT) *via* a spark plasma sintering method.¹⁷⁰ Under supercritical conditions (350 °C, 8.5 MPa), the catalyst exhibited high specific activities at low H₂/CO₂ ratios (1 or 2), with the iron-time yields of 5.4–12.2 mol_{CO₂} g_{Fe}⁻¹ s⁻¹ and C₂₊ products selectivity of 40–50 mol%, in CO₂ hydrogenation without pretreatment. This work highlighted the importance of CNT support with increased density for stabilizing the Fe NPs, which contributed to the strong metal-support interaction and CO intermediate activation.

Apart from the catalyst modification, the rational design of thermocatalytic reactors for CO₂ reduction also contributes to enhancing catalytic activity and selectivity. It is recommended to establish an appropriate dynamic mathematical model to comprehend heat conduction and mass transfer during the reaction process. Furthermore, computational fluid dynamics analysis can be employed to investigate parameters such as fluid flow and pressure drop. The integration of dynamic mathematical models, computational simulation tools, and experimental validation can guide the development of efficient thermocatalytic CO₂ reduction systems.

5. Conclusion and outlook

Herein, the recent progress of HSC-based materials in the fields of electrocatalytic and thermocatalytic CO₂ utilization has been reviewed. The synthetic methods of HSC were first introduced. Then, a comprehensive overview of functionalization strategies for HSC was provided, including nonmetal doping, metal single atom anchoring, and metal nanoparticle modification; and they were found to offer numerous benefits, such as optimization of electronic structure, superior CO₂ adsorption affinity, and improved reaction intermediate activation, allowing for the efficient enhancement of overall catalytic CO₂ conversion behavior. Finally, the discussion of HSC-based materials towards electrocatalytic and thermocatalytic CO₂ conversion applications was presented, based on the CO₂ reduction product category. While encouraging progress has been witnessed in these fields, some key issues still exist that merit further study.

(1) The synthesis of HSC materials with good rigidity, homogeneous morphology, and favorable reproducibility is mostly dependent on the utilization of the hard template method. However, this approach involves the use of highly corrosive agents to remove the template, potentially leading to adverse environmental consequences. Hence, the development of a green and environmentally friendly synthesis for high-quality HSC is urgently needed.

(2) The regulation of the pore structure of HSC is an important issue, as the porosity can affect the mass transport and the number of active sites, both of which have large influences on the activity and product selectivity. As exemplified by the work from Cao *et al.*, in Ni–N doped hollow carbon sphere catalyst system used for eCO₂RR, the mesopores on carbon shell were found to be favorable for the CO₂ diffusion; while the micropores with confinement effect were conducive to the formation of Ni–N active sites during the synthesis process. By balancing the content of micropore and mesopore, the CO₂-to-CO reaction rate can be efficiently improved.³⁸ In thermocatalytic CO₂ hydrogenation, the carbon-supported Fe–Co–K oxide catalyst using carbon support containing both micropores and mesopores shows higher olefin selectivity, as compared to that of the analogue with only micropores; the results suggest that the mesopores with sufficient accommodation space enable control over the intimate contact of K and Fe_xO_y species, making it possible to realize the boosted olefin production.¹⁸⁵

(3) Most of the works are rather ambiguous in describing the contribution of confinement effects of HSC to the enhanced catalytic performance, which should be clarified. Here are some good examples. In CO₂ electroreduction, the confinement effects of hollow carbon spheres contribute to the pH modulation of the internal cavity, which inhibits the competitive hydrogen evolution reaction by controlling the rate of H⁺ replenishment and creating a localized alkaline environment in the proximity of the catalytically active sites.⁴⁶ In thermocatalytic CO₂ reduction, in addition to improving the dispersion and stability of metal nanoparticles, HSC contributes to concentrating CO₂ around the catalytically active sites and thus accelerating the reaction rate.⁴⁵ In our opinion, more research efforts should be placed on emphasizing and clarifying the relationship between the confinement structure, active site, and substrate molecule, which allows to better utilize the positive effects and develop more advanced catalysts.

(4) Currently, the research on single-atom-based HSC materials is mainly focused on CO₂ electroreduction, and only few cases in thermocatalytic CO₂ conversion because the stabilization of single atoms with high surface free energy is a major challenge under the harsh conditions of thermocatalysis. Notably, in some specific situations, single atoms exhibit surprising stability, even better than metal nanoparticles. For example, the transformation of metal nanoparticles (Ru, Rh, and Pd) can be achieved by phosphine (PH₃)-assisted pyrolysis to form single-atom materials, in which single-atom species are firmly fixed on thermodynamically stable sites, which help inhibit the agglomeration and leakage of single-atom species.¹⁸⁶ Additionally, there are some studies reported the thermal atomic capture method allows the conversion of noble metal Pd, Pt, and Au nanoparticles to thermally stable single atoms.^{187,188} As a result, the stabilized single atoms showed better catalytic performance than nanoparticles; these works may offer some inspiration for the development of stable single-atom-based HSC materials for thermocatalytic CO₂ conversion.

(5) Furthermore, the types of single-atom-related HSC catalysts are limited, with the majority of them focusing only on Ni or Co single atoms in eCO₂RR for CO production (Table 3), and



it is important to further exploit high-performance catalysts. In this regard, the following aspects are worthy of further consideration. To begin with, novel active metal single atoms are suggested to be introduced into the matrix of HSC material. For instance, the fluxional property of gallium (Ga) has been proven to be utilized to efficiently enhance eCO₂RR behavior;¹⁸⁹ the analogous unique characteristics of more single atoms need to be explored. Secondly, studies on the coordination environment of the central metal atom should be conducted, because the coordinated non-metal atoms can affect the electronic structure of the central metal atom, resulting in the change of the adsorption/activation ability towards reactant molecules. Thirdly, rationally designing dual single-atom sites with different functions is helpful in realizing some complex tandem reactions and in achieving the controllable production of specific products.

(6) HSC materials possess unique cavities and abundant pore structures, which provide a flexible platform for designing multifunctional catalysts. Integrating with other novel materials, the hybrid catalyst suitable for various catalytic application scenarios can be expected. As an example, the novel high-entropy alloys are active for both eCO₂RR¹²⁸ and thermocatalytic CO₂ hydrogenation.¹⁹⁰ Combining similar materials with HSC carbon materials to develop novel catalysts is also a research direction of interest.

(7) During the processes of electro-/thermocatalytic CO₂ conversion, the intrinsic structure of the HSC materials may be changed under harsh conditions, such as large applied potential or high temperature. In this regard, *ex situ* characterizations may not be able to accurately characterize the catalysts; therefore, *in situ* measurements, such as *in situ* XAFS, Raman, XRD, and DRIFTS, should be employed to dynamically assess the structural change, which enables the guidance of a reasonable HSC material design. Theoretical simulation is a powerful tool to uncover catalytic mechanisms. Except for the commonly used DFT theoretical simulation, finite element modeling^{191,192} and molecular dynamics^{193,194} simulations are recommended to understand the mass transport behavior during the catalytic CO₂ reduction reaction.

(8) The continuous exploration of the unique properties of HSC materials has pushed their application boundaries. For instance, a recent study revealed that HSC played an important role in enhancing photocatalytic CO₂ reduction performance in a ternary HSC/CdS@ZnIn₂S₄ photocatalyst by offering multiple benefits, including improved light harvesting, increased CO₂ adsorption, protection of CdS against photocorrosion, *etc.*¹⁹⁵ In another work, HSC was demonstrated to be effective in boosting the separation of photoexcited electron-hole pairs and providing more catalytically active sites in the g-C₃N₄/Bi₂O₃@-HSC photocatalyst system; as a result, enhanced photocatalytic antibiotic degradation can be achieved.¹⁹⁶ In fact, the carbon shell of HSC material can absorb and convert light to heat under irradiation, resulting in the formation of a local hotspot, which is helpful for boosting the reaction kinetics.¹⁹⁷ As there is a lack of relevant reports, further study on the photo-thermal features of HSC and the development of HSC-based photothermal catalytic reactors will be of great significance.

The unprecedented development of HSC-based materials in recent years provides infinite possibilities for sustainable CO₂ utilization. We expect that this timely review will bring fellow researchers some inspiration to design more advanced HSC-based materials.

Author contributions

All of the authors contributed to the manuscript preparation. K. L. and Y. K. conceived the outline of the manuscript. K. L. wrote the original draft of the manuscript. Y. K. and H. Y. discussed and helped revise the manuscript.

Conflicts of interest

There are no conflicts to declare.

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