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Nitrogen-doped nanocarbon as a metal-free catalyst for CO₂ hydroboration†

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Designing heterogeneous metal-free catalysts for highly efficient transformation of CO_2 into valuable products is a dream goal and a long-standing challenge in thermal catalysis. Herein, N-doped nanocarbon catalysts, NOLCx, are synthesized and present an attractive ability to catalyse CO_2 hydroboration. N doping on the NOLCx surface mainly exists in the form of pyridinic N, pyrrolic N and graphitic N, where pyridinic N and pyrrolic N are the possible catalytic active sites. *In situ* IR was utilized to examine the catalytic mechanism and capture the reaction intermediate. These findings could promote the in-depth understanding of carbon surface chemistry and the rational design of heterogeneous metal-free nanocarbon catalysts for CO_2 thermal transformation.

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1. Introduction

The sequestration, activation and utilization of CO₂ through chemical methods has become a cutting-edge and booming

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istry and related applications in catalysis and energy-associated areas.

research area owing to increasing concerns regarding ecology crisis resulting from high atmospheric CO_2 concentration. Meanwhile, CO_2 , as an appealing C1 resource, could be utilized to produce fine chemicals, fuels, polymers and other products, wherein catalytic CO_2 reduction, including hydrogenation, hydrosilylation, hydrosilylation, and hydroboration, are valuable and widely used methods for CO_2 transformation.

Owing to the thermodynamic stability ($\Delta G_f^{\Theta} = -394.4 \text{ kJ}$ mol⁻¹) and chemical inertness inherent in non-polar CO₂ molecules, catalytic CO2 hydrogenation typically necessitates harsh reaction conditions (high temperature and high hydrogen pressures) or the use of transition metal catalysts. However, catalytic CO2 hydrosilylation or hydroboration requires significantly milder operating conditions because of the presence of relatively weak Si/B-H bonds and stronger Si/ B-O bonds and has drawn great interests as an alternative to hydrogenation. 18,23 CO2 hydroboration can produce formoxyborane, bis(boryl)acetal, and methoxyborane, along with bis (boryl)ether, which are important value-added chemicals. Transition metal catalysts, 19,20,24 alkali or alkaline earth metal catalysts, 25,26 and metal-free catalytic systems 21,22 can catalyze hydroboration reactions. Among them, metal-free catalytic systems could displace metal or metal oxide catalysts and have demonstrated significant potential in activating CO2, facilitating the creation of an environmentally friendly and sustainable society.

Much efforts have been made to construct metal-free catalysts for CO₂ activation and transformation. Frustrated Lewis pairs (FLPs), mainly composed of main group elements, are known to bind CO₂ through the interaction between unquenched Lewis acid sites/Lewis base sites with the O/C of

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capacity catalytic exhibiting for hydroboration.^{27,28} Ashley et al. used 2,2,6,6-tetramethylpiperidine (TMP) and B(C₆F₅)₃ to catalyze CO₂ reduction and obtained a unique formatoborate complex [TMPH]-[HCO2B $(C_6F_5)_3$] that could be further converted into $CH_3OB(C_6F_5)_3$ and methanol.²⁹ Fontaine et al. reported that intramolecular FLP 1-Bcat-2-PPh₂-C₆H₄ (cat = catechol) could catalyze CO₂ reduction by hydroboranes to methoxyborane. 21,30 Gao et al. found that a 6-amino-2-picoline-borane catalyst could reduce CO₂ with a borane-trimethylamine complex in an intramolecular FLP manner.22 Apart from FLPs, molecular guanidine and amidines prove to be highly efficient metal-free catalysts for CO₂ reduction with hydroboranes such as 9-borabicyclo [3.3.1]nonane (9-BBN) and catecholborane (catBH).31 Additionally, Stephan et al. reported that phosphines can serve as catalysts for the reduction of CO2 to methoxyboranes using 9-BBN as the reducing agent.³² An N-phosphinoamidinato NHC-diborene catalyst displayed catalytic CO2 hydroboration capacity with a B=B active site. 33 To date, the reported metalfree catalysts for CO2 hydroboration are mainly homogeneous, while heterogeneous metal-free catalysts have been rarely

Herein, we reveal that the nitrogen doped nanocarbon could serve as an efficient heterogeneous metal-free catalyst for CO₂ hydroboration for the first time, in contrast to the already published works where N-doped carbon materials are widely used as efficient metal-free catalysts for CO₂ electrocatalysis^{34–37} and infrequent nonreductive thermal conversion.³⁸ Successful N doping on the nanocarbon surface is confirmed by HRTEM, EDX, XPS and Raman spectroscopy. N species plays an important role in catalytic CO₂ hydroboration, where pyridinic N and pyrrolic N are the possible catalytic active sites. Finally, the catalytic mechanism is proposed based on *in situ* FTIR characterization, shedding light on the application prospect of heterogeneous metal-free catalysts in thermal activation and transformation of CO₂.

2. Experimental section

2.1 Reagent

Nanoscale

Purified ultra-dispersed nanodiamonds (UDDs) were purchased from Beijing Grish Hitech Co., Ltd. UDDs were washed with acid to remove amorphous carbon and metal impurities. 1-Butyl-3-methylimidazolium dicyanamide (BMIM·DCN, 98%) was provided by Shanghai Chengjie Chemical Co., Ltd. Chloroform-D (99.8% D, 0.03% TMS) and 1,3,5-trimethoxybenzene (98%) were provided by Adamas Chemical Co., Ltd. Toluene (99.5%), pinacolborane (HBpin, 4,4,5,5-tetramethyl-1,3,2-dioxaborolane, 97%) and 2-methoxy-4,4,5,5-tetramethyl-1,3,2-dioxaborolane were purchased from Energy Chemical Co., Ltd. Ultra-pure CO₂ gas was provided by Air Liquide.

2.2 Preparation of N-doped nanocarbons

N-doped nanocarbons were prepared from the carbonization of an IL containing the cyano group, namely, BMIM·DCN, on a

nanocarbon surface with a graphitic structure. 39,40 Onion-like carbon (OLC) was selected as the starting nanocarbon material because it has an almost perfect graphitic structure and large surface area; it was prepared by annealing UDDs at 1500 °C for 240 min under an Ar atmosphere. 41,42 The IL, BMIM·DCN was first completely dissolved in ethanol and BMIM·DCN alcoholic solutions was obtained. After that, OLC with the same weight as BMIM·DCN was put into BMIM·DCN alcoholic solutions and finely dispersed through ultrasonic treatment for 60 min. OLC impregnated with BMIM·DCN was obtained after the complete evaporation of ethanol and ground using an agate mortar. Subsequently, the resultant mixture was loaded in a quartz crucible and placed in the central zone of a tubular furnace, where residual oxygen and moisture were eliminated as much as possible through vacuuming and Ar purging three times. The temperature in the furnace was then incrementally raised from ambient conditions to a desired temperature at a heating rate of 10 °C min⁻¹ under an Ar atmosphere and maintained for 60 min. Following gradual cooling to room temperature under an Ar atmosphere, nitrogen-doped nanocarbon catalysts were obtained at 400 °C, 600 °C and 800 °C, denoted as NOLC400, NOLC600 and NOLC800, respectively.

2.2 Catalytic test

The catalytic reduction of CO₂ on N-doped carbon catalysts was conducted according to the following procedure. Initially, a precise quantity of certain catalyst was loaded into a high-pressure stainless autoclave equipped with a temperature controller and magnetic stirrer. Subsequently, 1 mL toluene and 2 mmol HBpin were introduced into the autoclave, serving as the solvent and reductant, respectively. The air within the sealed autoclave was purged three times with 4 MPa CO₂, followed by the final charging with CO₂ to 4 MPa. The autoclave was then positioned within a preheated jacket, which had attained the reaction temperature, and the moment of stirring at 800 rpm marked the initiation of the reaction. After a specific reaction time, the autoclave was removed from the heating jacket and immersed in a cold-water bath.

The gas inside the autoclave was collected in a gasbag and analyzed using a gas chromatography (GC9720 plus, Fuli, equipped with two FID detectors and one TCD detector). There were no products from CO₂ reduction in the gas phase, as shown in Fig. S3–S5.† Following the evacuation of gas, the autoclave was opened to extract the liquid product, from which the N-doped nanocarbon catalyst was separated through filtration. The resultant liquid product was collected and supplemented with 1,2,4-trimethoxybenzene as an internal standard for quantitative analysis *via* ¹H NMR (AVANCE NEO 400 MHz, Bruker).

2.4 Characterization

XPS spectra of samples was collected on the Thermo Scientific NEXSA Surface Analysis System with a monochromatized Al K α X-ray source ($h\nu=1486.6$ eV) and a power of 150 W. The surface structure of N-doped nanocarbon was studied on a laser confocal micro-Raman spectrometer (Thermo Scientific

Paper Nanoscale

Company, DXR). The instrument employs a 532 nm laser source (cobalt) and the power directed to the sample surface at full strength is set at 1 mW with an exposure time of 4 min. The microscopic morphology of N doped nanocarbon catalysts was observed through high resolution transmission electron microscopy (HRTEM, Thermo Scientific Talos F200X). EDX mapping was also conducted to study elemental distribution.

N₂ physical adsorption on NOLCx samples was carried out with the help of a physical adsorption instrument (Micromeritics 3Flex). The samples were first degassed under vacuum at 473 K for 6 h before the physical adsorption test. Nitrogen physical adsorption was then performed at 77 K at a relative pressure range of 0-1. The specific surface area of samples was calculated using the BET model, and the pore size distribution of samples was derived using the BJH model.

In situ FTIR analysis of CO2 hydroboration on N-doped nanocarbon catalysts was conducted employing a Nicolet iS50 spectrometer (Thermo Scientific) equipped with a diffuse reflection accessory comprising a heat chamber, DRIFT cell, and MCT-A detector. All spectra were obtained via 64 scans at a resolution of 4 cm⁻¹ under specific experimental conditions. In the standard procedure, the N-doped nanocarbon catalyst was initially loaded into the DRIFT cell and subsequently subjected to pretreatment at 300 °C under an Ar flow for 60 minutes, followed by cooling to 40 °C. The background spectrum of the sample was then acquired and utilized as a benchmark in subsequent experiments. Initially, HBpin was introduced into the cell via Ar passing through a bubbler containing HBpin. The initiation of argon flowing into the bubbler marked the starting point, and spectra were recorded over specific time intervals. Once the spectra were stabilized, the argon flow was switched to CO₂ bubbling through HBpin, and spectra were recorded over defined time intervals. Furthermore, to gain a more comprehensive picture of CO₂ reduction on the N-doped nanocarbon catalysts, the first introduction of CO2 into the DRIFT cell, followed by the bubbling of HBpin by CO2, was conducted, and spectra were collected at specific time intervals.

3. Results and discussion

The microscopic structure of UDDs and OLC was studied through HRTEM, and the results are shown in Fig. 1(a) and (b), indicating that OLC with an almost graphitic fullerene-like structure was obtained after the thermal annealing of UDDs. Subsequently, nitrogen doping was conducted on the OLC surface with a dicyanamide containing IL as a precursor at varying temperatures. The HRTEM micrograph of NOLC400 revealed a slightly uneven spherical structure, comprising densely arranged multilayered sp² fullerene-like shells, with a lattice spacing of approximately 0.353 nm between adjacent carbon layers (Fig. 1c). XRD spectra were collected to investigate the bulk structure, as shown in Fig. S1,† where the peak at about 25° for OLC suggests the graphitic structure of the OLC bulk phase, in line with the HRTEM pattern. There exists

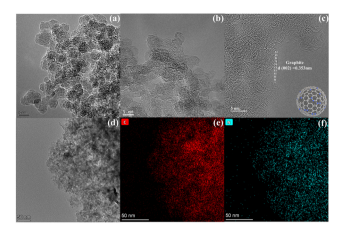


Fig. 1 (a) HRTEM of UDDs. (b) HRTEM of OLC: (c) HRTEM of NOLC400. (d) Selected area of NOLC400 for EDX mapping. (e) EDX mapping of C atoms on NOLC400. (f) EDX mapping of N atoms on NOLC400.

a peak at about 25° for NOLC400, indicating that N doping does not disturb the bulk phase of OLC. Moreover, C and N elemental EDX mappings of NOLC400 were collected to analyze the composition of N-doped nanocarbon, as depicted in Fig. 1(e) and (f). Nitrogen atoms and carbon atoms are uniformly distributed across the whole surface of NOLC400, confirming the successful doping of nitrogen atoms onto OLC. According to our previous research, 39,43,44 the graphitic structure of the nanocarbon surface can have rather strong staticassistant CH- π interactions with an interaction energy of -275.9 kJ mol⁻¹ where the heterocyclic CH of imidazolium ions serves as a proton donor and the conjugated π structure of OLC serves as a proton acceptor. IL molecules that do not have direct interactions with the graphitic surface evaporate when temperature increases. In contrast, the decomposed debris of IL molecules supported on OLC are entrapped by the graphitic surface of OLC. This phenomenon is similar to what occurs in the preparation of pristine-graphene-supported nitrogen-doped carbon with glucaminium-based ionic liquids (GILs) as a precursor, where hydrophobic and π - π stacking interactions promote the conformal in situ self-assembly of GILs on graphene. 45,46 Therefore, it is reasonable to infer that the N-doped carbon layer is coated on OLC.

X-ray photoelectron spectroscopy (XPS) is a powerful tool for scrutinizing the surface structure and nitrogen species within NOLCx. The surface compositions of OLC and NOLCx were first analyzed through XPS survey scans, as shown in Fig. S2,† and the data are summarized in Table S1.† Pristine OLC exhibits a composition of 98.3 at% of C atoms and 1.7 at% of O atoms, with the absence of N atoms. However, subsequent N doping introduces nitrogen atoms onto OLC successfully, which can be confirmed by the XPS N 1s spectra of NOLCx and EDX. The XPS N 1s spectra of three NOLCs samples were all deconvoluted into graphitic N (N1), pyrrolic N (N2), and pyridinic N (N3) species, as shown in Fig. 2(a-c). Increasing the preparation temperature resulted in a discernible variation in both the nitrogen content and nitrogen

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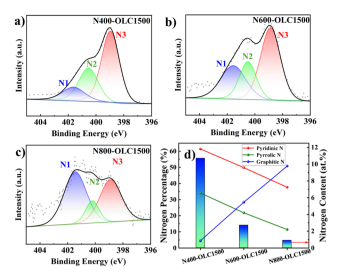


Fig. 2 (a) The deconvoluted N 1s XPS spectra of NOLC400, (b) NOLC600 and (c) NOLC800; (d) the change in the N content and percentages of different N species of NOLCx samples.

species percentage, as depicted in Fig. 2(d). A decline in the percentage of pyridinic and pyrrolic nitrogen species was accompanied with nitrogen content reduction as the preparation temperature was elevated—a phenomenon also documented in the literature on N-doped carbon materials. 47,48 While the graphitic nitrogen percentage increased concomitantly with an elevation in the preparation temperature, which could be attributed to the chemical characteristics of N covalent bonding with three carbon atoms. 49 Graphitic nitrogen tends to be generated and remain stable as temperature is elevated, suggesting the graphitization of surface carbonous

The nitrogen-doped structure on the NOLCx surface could also be understood in combination with the possible carbonization process of BMIM·DCN. According to previous research, 40 BMIM·DCN is first decomposed with the elimination of alkyl chains and then condensed via the trimerization of nitrile groups, thus forming triazine rings containing pyridinic N species. Meanwhile the imidazolium cationic ring structure remains intact and is then incorporated into carbonous debris through the nucleophilic attack of the DCN⁻ anion, thus yielding pyrrolic N species. As the preparation temperature increases, the elimination of hydrogen, nitrogen and alkyl fragments becomes more pronounced, and the restructuring of the carbonous debris occurs to fit the requirements of the locally graphitized structure. The possible carbonization process of BMIM·DCN can well explain the evolution process of nitrogen species in NOLCx.

Raman spectroscopy is a valuable tool for elucidating the structural characteristics of carbonaceous materials, simultaneously obtaining defect information. From the deconvolution of the Raman spectra of NOLCx samples, given in Fig. 3, it could be found that the NOLCx catalysts all possess the D1 band, D2 band and G band at 1350 cm⁻¹, 1500 cm⁻¹ and

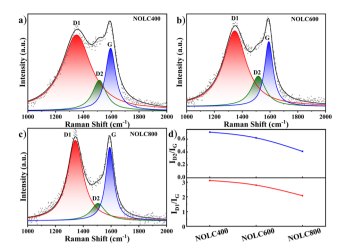


Fig. 3 (a) The deconvoluted Raman spectra of NOLC400 sample; (b) the deconvoluted Raman spectra of the NOLC600 sample; (c) the deconvoluted Raman spectra of the NOLC800 sample; (d) I_{D1}/I_{G} and I_{D2}/I_{G} $I_{\rm G}$ of the NOLCx samples.

1580 cm⁻¹, respectively. ^{39,41} The D1 band is attributed to the edge of disordered graphitic microcrystals and originates from lattice vibrations (A_{1g}), which are exclusively detectable in the presence of graphitic lattice vibrations in the vicinity of structural defects.⁵⁰ The G band (~1580 cm⁻¹) is generated by the stretching motion of all sp² atomic pairs in the carbon rings or long chains, representing the ideal graphitic structure. The D2 band represents amorphous carbon. Therefore, based on the origin of these Raman signals of carbon materials, the relative intensity ratio of the D band compared to the G band ($I_{\rm D1}/I_{\rm G}$ or $I_{\rm D2}/I_{\rm G}$) is commonly used for roughly estimating the defect type and content of nanocarbon materials in research. As illustrated in Fig. 3(d), I_{D1}/I_{G} exhibits a progressive decline with increasing preparation temperature, which could be ascribed to the decreasing N content and the graphitization of surface carbonous debris impelled by high temperature. 48 In addition, the appearance of a D2 band suggests the existence of an amorphous carbon part in the NOLCx surface; the reduction in $I_{\rm D2}/I_{\rm G}$ with increase in preparation temperature could be attributed to the volatilization or graphitization of this amorphous carbon part.

The surface area and pore size distribution of samples were studied using N2 physical adsorption. The N2 adsorption/desorption isotherms are shown in Fig. 4(a). The surface area of N-doped samples is smaller than that of OLC and increases with an increase in the preparation temperature. The three NOLCx samples have a similar pore structure, as depicted in Fig. 4(b). The pore volume of NOLCx samples also increases with the preparation temperature, whereas the mean pore size decreases (Table S2†).

After confirming successful N doping onto OLC, the catalytic activity of NOLCx materials for CO2 hydroboration was tested, and their catalytic performance is summarized in Table 1. It was found that only a negligible amount of the CO₂ reduction product (MeOBpin) was obtained when no catalysts Paper

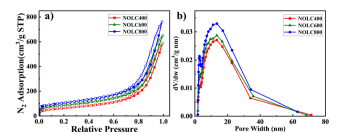


Fig. 4 (a) The N_2 adsorption/desorption isotherms and (b) pore size distribution of NOLCx samples.

Table 1 Catalytic performance of different catalysts under different reaction conditions

Catalyst	HBpin/ mmol	Temp/ °C	Time/h	MeOBpin yield mmol
No	2	100	2	0.013
OLC	2	100	2	0.021
NOLC400	0	100	2	0.000
NOLC400	2	100	2	0.525
NOLC600	2	100	2	0.391
NOLC800	2	100	2	0.123
NOLC400	2	80	2	0.264
NOLC400	2	60	2	0.049
Acridine a	2	100	2	0.023
Acridine b	2	100	2	0.018
A-OLC ^c	2	100	2	0.021
Carbazole d	2	100	2	0.028
Carbazole ^e	2	100	2	0.023
C-OLC f	2	100	2	0.028
	No OLC NOLC400 NOLC400 NOLC600 NOLC800 NOLC400 Acridine ^a Acridine ^b A-OLC ^c Carbazole ^d Carbazole ^e	Catalyst mmol No 2 OLC 2 NOLC400 0 NOLC400 2 NOLC800 2 NOLC800 2 NOLC400 2 NOLC400 2 Acridine a 2 Acridine b 2 A-OLC c 2 Carbazole d 2 Carbazole e 2	Catalyst mmol $^{\circ}$ C No 2 100 OLC 2 100 NOLC400 0 100 NOLC400 2 100 NOLC800 2 100 NOLC400 2 80 NOLC400 2 60 Acridine a 2 100 Acridine b 2 100 A-OLC c 2 100 Carbazole d 2 100 Carbazole e 2 100	Catalyst mmol °C Time/h No 2 100 2 OLC 2 100 2 NOLC400 0 100 2 NOLC400 2 100 2 NOLC800 2 100 2 NOLC400 2 80 2 NOLC400 2 60 2 Acridine a 2 100 2 Acridine b 2 100 2 A-OLC c 2 100 2 Carbazole d 2 100 2 Carbazole c 2 100 2

 $[^]a$ 10 mg acridine. b 50 mg acridine. c 10 mg A-OLC, which is 10 wt% acridine supported on OLC. d 10 mg carbazole. e 50 mg carbazole. f 10 mg C-OLC, which is 10 wt% carbazole supported on OLC.

or OLC was used in the catalytic reaction, suggesting that CO₂ reduction by borane is incapable of proceeding without an effective catalyst and OLC. Moreover, CO2 cannot be reduced in the absence of a reductant. A significant MeOBpin yield was obtained when NOLC400 was used as the catalyst. NOLC600 and NOLC800 could effectively catalyze CO2 hydroboration, although their catalytic activity was poorer than that of NOLC400. At a lower reaction temperature, the catalytic activity of NOLC400 was weakened. However, it was still better than that of OLC at a higher temperature. NOLC400 also exhibits good stability even after recycling 6 times, as shown in Fig. S6.† Compared with the reported catalysts for CO₂ hydroboration, as shown in Table S3,† NOLCx catalysts showed better catalytic performance with their feature of metal-free heterogeneous catalytic systems being the most significant advantage.

The change in CO₂ hydroboration product yield with the reaction time catalyzed by NOLCx was acquired, as depicted in Fig. 5(a). The CO₂ hydroboration product yield catalyzed by the three samples all increased with the reaction time, from which the initial reaction rate could be derived. During the whole reaction period, NOLC400 possessed better catalytic activity

than NOLC600 and NOLC800. All these catalytic activity data underlines that N doping plays an important role in the catalysis of CO₂ hydroboration by NOLCx. To explore the nature of catalytic sites, the initial reaction rate is correlated with the different N species content, as shown in Fig. 5(b-d). It is found that there exists a substantial positive correlation between pyridinic nitrogen or pyrrolic nitrogen and the initial reaction rate, whereas there is no obvious correlation between graphitic nitrogen and the initial reaction rate. Therefore, it is reasonable to posit that pyridinic nitrogen or pyrrolic nitrogen may serve as the potential active site in this catalytic process.

Furthermore, aromatic molecules containing specific N species were used to mimic the N doping structure on the carbon surface, which helped to disclose the nature of active sites of N-doped carbon materials, whose validity has been verified by Lin et al.51 Here, acridine and carbazole were utilized to model pyridinic N and pyrrolic N species, respectively, as shown in Fig. S7.† Unfortunately, acridine and carbazole showed negligible catalytic activity, where increasing their dosage five times had no positive effect on their catalytic activity. To eliminate the influence of the OLC matrix, acridine and carbazole at 10 wt% were supported on the OLC surface and utilized in the catalytic reaction, which produced negligible yields of CO2 hydroboration products. These results underscore the unique catalytic function of the N-doping structure on the nanocarbon surface, exceeding that of small aromatic molecules with isolated N species. This catalytic performance could be ascribed to the special electron-rich N-doping structure incorporated into the rather big π -conjugated carbon skeleton with electron donation from the graphitic structure to the N-doping sites on the nanocarbon surface. Their electronic structure is apparently different from

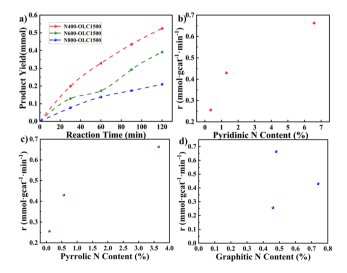


Fig. 5 (a) The change in CO₂ hydroboration product yield with the reaction time (10 mg catalyst, 2 mmol HBpin, 100 °C); (b) the correlation between the initial reaction rate and the pyridinic N content; (c) the correlation between the initial reaction rate and the pyrrolic N content; (d) the correlation between the initial reaction rate and the graphitic N content

Nanoscale Paper

that of the isolated molecules with small aromatic structures, which would significantly influence catalytic performance. 43,52 Therefore, special caution should be taken while using small aromatic molecules with a functionality to mimic the active sites on the carbon surface.

To further explain the CO₂ reduction mechanism by HBpin on NOLCx, in situ FTIR spectra were collected and analyzed. HBpin was first introduced into an in situ cell through Ar bubbling, and the IR signal of HBpin was acquired. As shown in Fig. 6(a), the IR peak at 2595 cm⁻¹ attributed to B-H stretching vibration became obvious after HBpin introduction and decreased with the subsequent introduction of CO₂, indicating the reaction of B-H with CO2. Meanwhile, the peak attributed to C=O vibration located at 1692 cm⁻¹ appeared at the beginning and was then weakened with an increase in CO2 blowing time, as depicted in Fig. 6(b), suggesting its characteristics of a reaction intermediate. It is worth noting that the IR peak position of C=O is very close to that of CO2 activated by frustrated Lewis pairs. 28,53 Peaks attributed to the B-O and C-O stretching vibrations of the reaction intermediate also appeared while B-O¹ stretching vibration from the dioxaborolane ring of the reductant HBpin was simultaneously weakened, as shown in Fig. 6(b) and Fig. S8.†

To gain a more comprehensive picture of the reaction mechanism, CO2 was first introduced into the in situ DRIFT cell, and later, HBpin was added. As shown in Fig. 7(a), a striking IR peak at 1660 cm⁻¹ appeared just after the CO₂ introduction, and its intensity became stronger with an increase in CO₂ blowing time, which is attributed to the asymmetric O-C-O stretching vibration of chemisorbed CO2 on the NOLC400 surface.^{54,55} This IR peak is apparently different from the IR peak at 1692 cm⁻¹ shown in Fig. 6(b), and the difference suggests that CO₂ is first chemically adsorbed on the NOLC400 and then transformed into the reaction intermediate O=CH-O-Bpin with the help of HBpin. The in situ IR spectra in Fig. 7(b) after the HBpin introduction through CO₂ bubbling verified the hypothesis to some extent. The IR peak at 1660 cm⁻¹ was weakened, whereas that at 1692 cm⁻¹ became apparent with an increase in the HBpin addition time, which is consistent with the trend given in Fig. 6(b). At the same time, the IR peaks corresponding to N-C=O and N-CO-O vibrations appeared and became obvious.

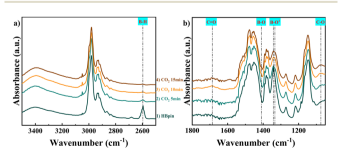


Fig. 6 (a) The in situ FTIR spectra of HBpin and CO2 on NOLC400 in the range of 3500-2500 cm⁻¹ and (b) 1800-1050 cm⁻¹

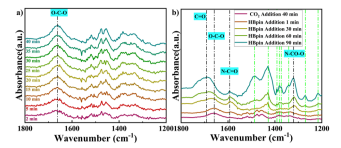


Fig. 7 (a) The in situ FTIR spectra of CO2 adsorption on NOLC400 in the range of 1800-1200 cm⁻¹; (b) the in situ FTIR spectra of CO₂ reacting with HBpin on NOLC400 (IR peaks marked with green lines result from HBpin).

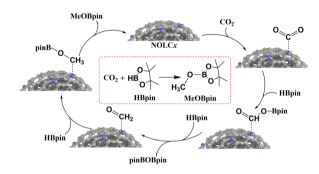


Fig. 8 The proposed CO₂ reduction mechanism catalyzed by NOLCx.

Based on the catalytic test and in situ FTIR results, a possible catalytic mechanism is proposed, as shown in Fig. 8. The chemisorption of CO₂ on the N-doping sites of NOLCx occurs first and chemisorbed CO2 is transformed into the O=CH-O-Bpin intermediate with the help of HBpin. Another HBpin molecule reacts with the intermediate, and the byproduct pinBOBpin is released. Finally, another HBpin molecule reacts with the O=CH₂ intermediate, and MeOBpin is obtained.

4. Conclusion

In this study, metal-free nanocarbon catalysts for effectively catalyzing CO2 reduction were synthesized via N doping on the OLC surface, which was confirmed by HRTEM, EDX, XPS and Raman spectroscopy. NOLCx demonstrated the novel and excellent capacity of catalyzing CO₂ hydroboration, where N-doping species and content play an important role. Pyridinic N and pyrrolic N might be the potential catalytic active sites, rather than graphitic N species. In situ IR was utilized to discover the catalytic mechanism by which chemisorbed CO2 would be transformed into the reaction intermediate O=CH-O-Bpin with the help of N doping and HBpin. To the best of our knowledge, this work presents the first report on CO₂ hydroboration catalyzed by a heterogeneous metal-free nanocarbon catalyst, thus not only expanding the catalytic

Paper Nanoscale

application scenario of nanocarbon materials but also providing a deeper understanding of carbon surface chemistry.

Author contributions

Tao Du: writing - original draft, visualization, validation, software, investigation, and formal analysis. Peng Zhang: writing original draft, visualization, and software. Guofeng Wang: investigation, formal analysis, data curation, Conceptualization. Zhen Jiao: writing - original draft, visualization, and software. Jiancheng Zhou: supervision, resources, and formal analysis. Yuxiao Ding: writing - review & editing, writing - original draft, project administration, methodology, investigation, funding acquisition, formal analysis, data curation, and conceptualization.

Data availability

The authors confirm that the data supporting the findings of this article are available within the main article and its ESI.†

Conflicts of interest

There are no conflicts to declare.

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Nanoscale Paper

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