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1. Introduction

Mercury chloride is the efficient catalyst used for acetylene hydrochlorination for the production of vinyl chloride monomers in China. However, the mercury based catalysts cause environmental problems and are toxic to human beings. Therefore, it is urgent and indispensable to explore environment friendly non-mercury catalysts to replace HgCl₂ catalysts for acetylene hydrochlorination.

It is worth mentioning that the noble metal chloride catalysts were investigated in the past decades for the reaction, with significant results achieved.¹⁻⁹ Au-based catalysts and Ru-based catalysts have attracted considerable research attention due to the superior catalytic activity for acetylene hydrochlorination. On the other hand, the noble metal catalysts are also facing serious challenges for the reason of high price, poor stability, and limited storage which hampered industrial application. Therefore, the heteroatom-doped carbon catalysts have been studied by many investigation groups.

The p-block elements of N, B, P, or S create electron-donating or electron-accepting sites on carbon materials due to the induction of a redistribution of p-electrons, which influence the adsorption properties on the carbon-based catalysts. In recent years, N-doped carbon materials have been widely studied as

Sulphur-doped activated carbon as a metal-free catalyst for acetylene hydrochlorination

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A series of sulfur-doped spherical activated carbon (SAC) catalysts were prepared with phenyl disulfide $(C_{12}H_{10}S_2)$ as a sulfur source for acetylene hydrochlorination. The S-doped catalyst exhibits preferable catalytic performance compared to that of the blank carrier with the reaction conditions of GHSV of 90 h^{-1} and at 180 °C. The catalysts were characterized by N₂ adsorption/desorption (BET), elemental analysis (EA), thermogravimetric analysis (TG), temperature-programmed desorption (TPD), Raman spectrum (Raman) and X-ray photoelectron spectroscopy (XPS). The results indicate that the presence of sulfur species is favorable to promote the ability of reactant adsorption and inhibit carbon deposition. In addition, the electronic and chemical properties of catalysts were investigated by density functional theory (DFT) simulation. It is illustrated that the introduction of sulfur species can not only change the spin density and charge density but also create more active sites on a carrier. The single sulfur doped carbon material catalysts were designed for the first time and the desirable results make it a green catalyst for the industrial application of acetylene hydrochlorination.

metal-free catalysts for acetylene hydrochlorination reaction. The group of Bao¹⁰ investigated a nanocomposite of nitrogendoped carbon derived from silicon carbide (SiC@N-C) as catalyst, which exhibited considerable performance during a 150 hour test. Li *et al.* prepared¹¹ g-C₃N₄/AC catalyst which displays excellent catalytic performance and revealed that the nitrogen atom of the g-C₃N₄/AC catalyst is the active site for the HCl. Although N-doped carbon material has superiority catalytic activity, it still cannot be compared with Au-based catalysts. The dual doping strategy was investigated to improve the N-doped carbon catalytic activity for acetylene hydrochlorination by many excellent research groups.

Dai *et al.* reported¹² that N, B dually doped graphene and the catalyst could promote HCl adsorption, which showed a catalytic activity that was little lower than Au-based catalyst. Li's group¹³ proposed the co-doped carbon catalyst by the doping element of sulfur and nitrogen, that the introduction of sulfur improved the pyridinic N content effectively in N-doped carbon material. Zhu *et al.* investigated¹⁴ the sulphur and nitrogen dual-doped carbon catalyst, the presence of sulphur enhanced the catalytic activity of catalyst for the synergistic effect between S and N.

Overall, the interaction between N and S inspired us to research the effect of only S-doped carbon materials for the reaction. Diphenyl disulfide has a relatively simple structure and two S atoms in the molecule are covalently linked. In consideration of the doping of sulfur in activated carbon, there is no obstacle for the doping process due to the presence of two benzene rings. Herein, we selected diphenyl disulfide as the sulfur source to modified activated carbon and designed sulfur-

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doped carbon catalysts, combined with the DFT simulation method to study the effect of S species on the catalyst.

2. Experimental

2.1. Material

Phenyl disulfide ($C_{12}H_{10}S_2$) was purchased from Beijing Bailingwei Technology Development Co., Ltd. Pitch-based spherical activated carbon (marked as SAC, 20–40 mesh) was purchased from Shanghai Huiheda Investment Management Co., Ltd. Ethanol (purity 99.7%) and sodium hydroxide (purity 96%) were purchased from Tianjin Guangfu Fine Chemical Research Institute. The reaction gas (HCl and C_2H_2 , purity all about 99.99%) were purchased from Tianjin Dongxiang Special Gas Co., Ltd. All of the materials were used without further purification.

2.2. Catalyst preparation

A certain amount of SAC was roasted at a heating rate of 5 °C min⁻¹ ranging from 25 °C to 800 °C in an Ar atmosphere for 120 min and then dropped down to room temperature, the final sample denoted as B-SAC. Phenyl disulfide (1 g) was dissolved in ethanol solution (20 ml) and then the solution was dropwise added into 10 g B-SAC with stirring. After that, the mixture was set aside for 12 h in room temperature, followed by evaporation at 60 °C for 12 h and desiccation at 120 °C for 12 h. The sample was calcined at particular temperature (600 °C, 700 °C, 800 °C and 900 °C, respectively) for 2 h at the heating rate of 5 °C min⁻¹ under flowing Ar to finally obtain the catalysts, denoted as 3% S/B-SAC (600), 3% S/B-SAC (700), 3% S/B-SAC (800), 3% S/B-SAC (900), respectively. On the other hand, the catalysts with different loading content of phenyl disulfide were prepared by using the same method. The obtained catalysts were denoted as 6% S/B-SAC (800), 9% S/B-SAC (800), 12% S/B-SAC (800), respectively.

2.3. Catalyst characterization

Brunauer–Emmett–Teller (BET) specific surface areas were measured by using a Quantachrome Autosorb Automated Gas Sorption System (Quantachrome Instruments, USA). The catalysts were degassed at 300 °C for 4 h and then analyzed *via* liquid nitrogen adsorption at -196 °C. Thermogravimetric analysis (TGA) of samples was investigated using a TG-DTA 2 thermal analyzer (METTLER TOLEDO, Switzerland). The sample was heated from 35 °C to 900 °C with a heating rate of 10 °C min⁻¹ under the air atmosphere.

Elemental Analysis (EA) was conducted using a Vario Micro analysis instrument. Raman spectra were measured using a DXR Raman micro-scope (DXR microscope, Thermo Fisher, USA) with the He–Ne laser emitter at wavelength of 633 nm.

Temperature-programmed desorption (TPD) was performed using an AutoChem BET TPR/TPD (Quantachrome Instruments AMI-90). The samples (150 mg) were treated under pure C_2H_2 , HCl and C_2H_3Cl gas at 180 °C for 4 h. The obtained samples were performed on the instrument over the temperature range 30–900 °C with a heating rate of 10 °C min⁻¹ and a He flow rate of 25 ml min⁻¹.

X-ray photoelectron spectroscopy (XPS) was carried out on a Thermo ESCALAB 250Xl (Thermo Fisher Scientific, USA), with the C 1s peak at 284.4 eV as the internal standard.

2.4. Catalyst tests

The catalyst performance evaluation was carried out in a fixedbed microreactor (i.d. of 10 mm). A CKW-1100 temperature controller (Chaoyang Automation Instrument Factory, Beijing, China) was used to regulate the temperature of the reaction. The reactor was purged into an appropriate amount of nitrogen to remove the air and water in the system before the reaction. Hydrogen chloride gas was passed through the heating reactor to activate the catalyst (3.0 ml) with the temperature of 150 °C and the flow rate of 25 ml min⁻¹ for 30 min. Subsequently, the HCl and C_2H_2 with a molar ratio of 1.1 were fed through the reactor vessel, producing a gas hourly space velocity (GHSV, C_2H_2) of 90 h⁻¹ under the reaction temperature of 180 °C. The exit gas mixture was passed through an absorption bottle containing NaOH solution, followed by the analysis using a Beifen 3420A gas chromatograph.

3. Results and discussion

3.1. Performance of S-doped carbon catalysts

The catalytic activities of the obtained catalysts have been evaluated under the same condition. First, the evaluation results of

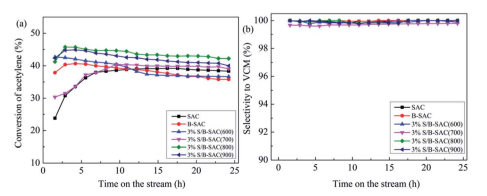


Fig. 1 Acetylene conversion (a) and selectivity to VCM (b) of 3% S/B-SAC catalysts with different calcination temperature. Reaction conditions: temperature (T) = 180 °C; (GHSV) = 90 h⁻¹; $V_{HCI}/V_{C_2H_2}$ = 1.1; 3 wt% S.

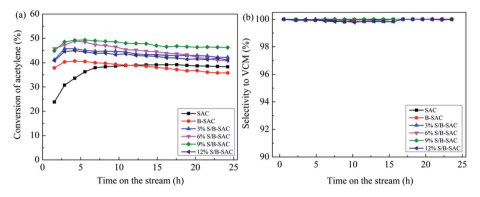


Fig. 2 Acetylene conversion (a) and selectivity to VCM (b) of S/B-SAC (800) catalysts with different S doped amount. Reaction conditions: temperature (T) = 180 °C; (GHSV) = 90 h⁻¹; $V_{HCI}/V_{C_2H_2}$ = 1.1; calcination temperature 800 °C.

Table 1	Comparison of	catalytic	performance
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Catalysts	Temperature/°C	GHSV of C_2H_2/h^{-1}	Molar ratio/C ₂ H ₂ : HCl	Acetylene conversion
NS–C–NH ₃ (ref. 12) S,N-carbon-2 (ref. 13)	220 180	35 50	1 : 1.2 1 : 1.15	80% 82.44%
9% S/B-SAC	180	90	1:1.1	50%

the 3% S-doped catalysts under the treatment condition with different calcination temperature were shown in Fig. 1(a). As the reference sample, B-SAC shows an initial acetylene conversion of 40% and decreases to 35% after 24 h reaction. Compared with B-SAC, the C2H2 conversion of S-doped catalysts was improved with different degree. 3% S/B-SAC (800) exhibits the most favorable catalytic activity with the acetylene conversion of 47%, which still maintain at 42% after 24 h. As for 3% S/B-SAC (600), 3% S/B-SAC (700) and 3% S/B-SAC (900), acetylene conversion of 36%, 38% and 39% were achieved after 24 h reaction, respectively. The selectivity to vinyl chloride over all the catalysts is higher than 99% (Fig. 1(b)). The result indicates that the catalytic activity increases until the increasing of calcination temperature at 800 °C. The 3% S/B-SAC (900) catalyst shows lower acetylene conversion than that of 3% S/B-SAC (800) catalyst. It may be due to higher calcination temperature resulting in the collapse of pore, which covering the active sites. To sum up, the optimum calcination temperature is 800 °C.

The catalytic performance of the catalysts with different sulfur contents are shown in Fig. 2(a). There is no regular variation of C_2H_2 conversion for all S-doped catalysts. Over 3% S/B-SAC, 6% S/ B-SAC, 9% S/B-SAC and 12% S/B-SAC catalysts, the acetylene conversion is respectively 45%, 48%, 50% and 44% at 5 h, whereas it decreases to 42%, 41%, 46% and 41% after 24 h. It can be seen that 9% S/B-SAC catalyst exhibits the optimal activity, indicating that the S-doped carbon catalyst can suitable for the metal-free catalysts used in the acetylene hydrochlorination reaction. For comparing the catalytic activity with that of the considerable sulphur and nitrogen dual-doped carbon catalysts, we list the detail contents in Table 1. Both of the two N and S dual-doped carbon catalysts exhibit preferable catalytic activity. The 9% S/B-SAC catalyst also shows excellent activity at relatively

Table 2 Pore structure parameters of catalysts				
$S_{ m BET} ({ m m}^2~{ m g}^{-1})$		Total pore (cm ³ g ⁻¹)	e volume	
Catalyst	Fresh	Used	Fresh	Used
SAC	1149	1077	0.661	0.627
B-SAC	830	755	0.352	0.328
3% S/B-SAC	722	679	0.321	0.304
6% S/B-SAC	659	607	0.302	0.275
9% S/B-SAC	589	556	0.273	0.256
12% S/B-SAC	524	482	0.229	0.207

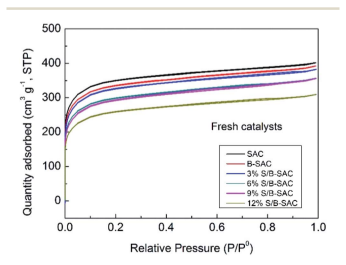


Fig. 3 The adsorption-desorption isotherm curves of fresh catalysts.

lower reaction temperature and higher GHSV condition. In addition, it is worth mentioning that the acetylene conversion only reduced by approximately 2% after 9 h, 4% after 24 h of reaction for 9% S/B-SAC catalyst. It can be seen that S-doped carbon catalyst has better stability for the reaction.

3.2. Catalyst characterization

3.2.1 Textual properties and coke deposition for the Sdoped catalysts. Nitrogen adsorption/desorption isotherms were carried out to confirm the physical and structural properties of S-doped carbon catalysts. Table 2 lists the specific surface areas and total pore volume of the fresh and used SAC, B-SAC and S/B-SAC catalysts. All the catalysts exhibit type-I adsorption isotherm due to the large number of micro-pores in the SAC surface, as shown in Fig. 3. The B-SAC displays a specific surface area of 830 m² g⁻¹ and a total pore volume of 0.352 cm³ g⁻¹, which is lower than that of SAC sample. The

Table 3 Weight loss of the fresh and used S-doped SAC catalysts

Amount of carbon deposition (%)
16
10
8.4
8.2
7.8
8.8

decline of the pore structure data over B-SAC is due to the collapse of the pore channel during the roasting process.

As listed in Table 2, the specific surface areas and total pore volumes of the fresh S-doped catalysts decrease with the increasing of sulfur doping content. The phenomenon is caused by the dilution effect, the more pores of supports are filled or

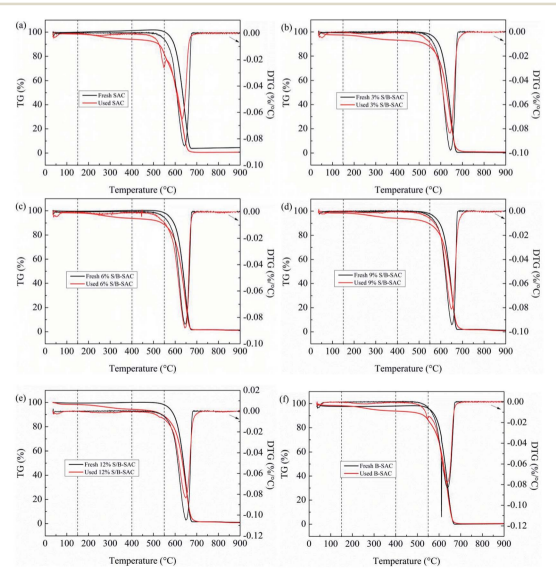


Fig. 4 TG and DTG curves of the fresh and used S-doped SAC catalysts.

blocked after the more loading active species.¹⁵ By observing the pore structure data between the fresh and used catalysts, the variation trend of the specific surface areas and the total pore volume of the catalysts are shown below. The variation trend of the specific surface areas (ΔS_{BET} %) of the catalysts are following the order: 12% S/B-SAC (8.0) > 6% S/B-SAC (7.8) > 3% S/B-SAC (5.9) > 9% S/B-SAC (5.6). The changes of the total pore volume (ΔV_{total} %) show the same trend in the following order: 12% S/B-

Table 4 Elemental analysis of catalysts				
	Bulk content (wt%)			
Sample	С	Н	Ν	S
SAC	96.30	0.964	0.27	0.16
B-SAC	96.91	0.950	0.29	0.15
3% S/B-SAC	95.80	0.865	0.28	0.40
6% S/B-SAC	96.06	0.837	0.30	0.80
9% S/B-SAC	95.57	0.796	0.33	1.29
12% S/B-SAC	94.12	0.752	0.28	1.87
Used SAC	92.64	1.077	0.28	0.12
Used B-SAC	91.72	1.061	0.28	0.11
Used 3% S/B-SAC	91.44	1.019	0.28	0.36
Used 6% S/B-SAC	91.26	0.981	0.28	0.69
Used 9% S/B-SAC	90.30	0.962	0.33	1.08
Used 12% S/B-SAC	90.62	0.950	0.27	1.65

SAC (9.6) > 6% S/B-SAC (9.2) > 3% S/B-SAC (6.2) > 9% S/B-SAC (6.0). For the results, the variation of the BET data and the activity may be due to the collapse of pore structure or carbon deposition on the catalysts during the reaction.

TGA was performed to evaluate the amount of coke deposition on the surface of the used catalysts. As seen in Fig. 4, all of the catalysts have a slight weight loss before 150 $^{\circ}$ C, due to very little water existing on the surface of the fresh and used samples. For the 3% S/B-SAC sample (Fig. 4b), the fresh catalyst

 $\label{eq:Table 5} \begin{array}{l} \mbox{Table 5} & \mbox{The relative contents and binding energies of S species in fresh catalysts} \end{array}$

Catalyst	-C-S-C (ev	V)	-C-S-C (%)
Fresh 3% S/B-SAC	164.2	165.4	75.36
Fresh 6% S/B-SAC	164.0	165.2	83.66
Fresh 9% S/B-SAC	164.0	165.2	85.19
Fresh 12% S/B-SAC	164.1	165.2	79.49
	-C-SO _r -C		$-C-SO_x-C$
	(x = 2, 3,	4) (eV)	(%)
Fresh 3% S/B-SAC	(x = 2, 3, 167.8)	4) (eV) 169.8	(%) 24.64
Fresh 3% S/B-SAC Fresh 6% S/B-SAC	(, , , ,	, , ,	
	167.8	169.8	24.64

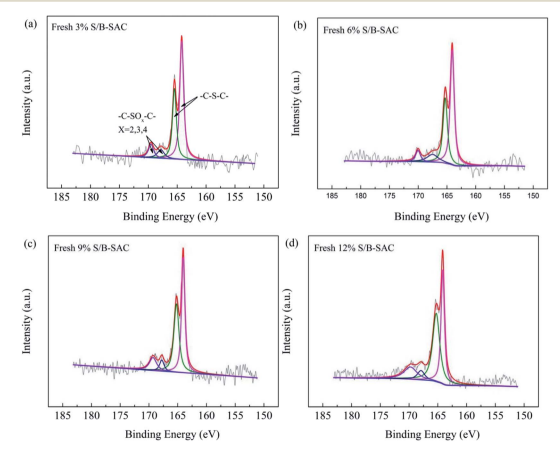


Fig. 5 High-resolution XPS spectra of S 2p for fresh S-doped catalysts.

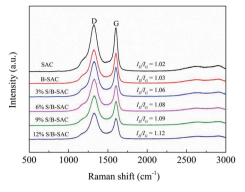


Fig. 6 Raman spectra of S-doped SAC catalysts.

shows the weight loss of 1%, whereas, there is apparent weight loss of 9.4% for the used catalyst during the temperature range of 150–500 $^{\circ}$ C. The significant change of the mass loss for the fresh and used catalyst is attributed to carbon deposition combustion. At the temperature higher than 500 $^{\circ}$ C, both of the

fresh and used 3% S/B-SAC show rapid weight loss due to the combustion of activated carbon. All the other catalysts have the same variation tendency in mass compare to the 3% S/B-SAC catalyst in the same temperature range.

In order to obtain coke deposition of every sample, the previous calculation method¹⁶ is adopted and the computed consequences are listed in Table 3. The relative amount of deposited coke is following the order: 9% S/B-SAC (7.8) < 6% S/B-SAC (8.2) < 3% S/B-SAC (8.4) < 12% S/B-SAC (8.8). Combining with the BET results, it is concluded that the doping of sulfur in the activated carbon can inhibit the formation of coke deposition and subsequently enhance the catalytic activity.

3.2.2 Determination of the active sites for S-doped carbon catalysts. Elemental analysis was performed to investigate the content variation of sulfur for fresh and used S-doped carbon catalysts. As listed in Table 4, the content of sulfur increases continuously with the increasing of sulfur precursor content for the fresh and used samples.

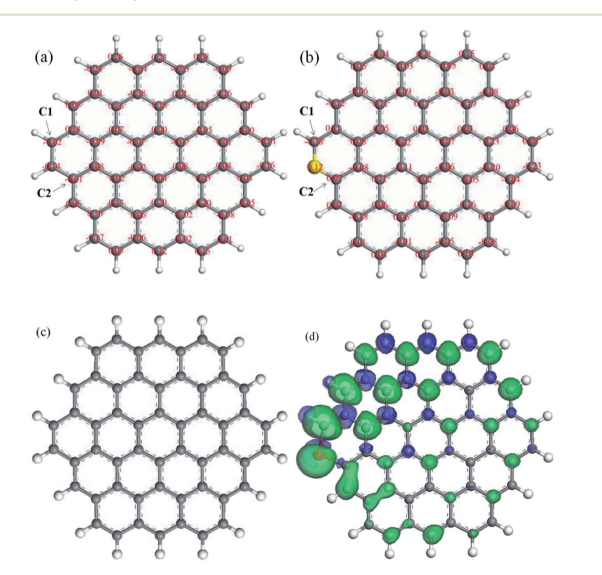


Fig. 7 On-site Bader charge (a) and spin density isosurface (c) for undoped model carbon layer; on-site Bader charge (b) and spin density isosurface (d) for doped model carbon layer.

Further, XPS was carried out to identify the chemical states and the types of sulfur active species for S-doped carbon catalysts. As shown in Fig. 5 and Table 5, the XPS S 2p was deconvoluted into two different sulfur species, the peak at 164–166 eV and 167–170 eV correspond to C–S–C and C–SO_x–C (x = 2, 3, 4), respectively.¹⁷

Table 5 shows the binding energies and the relative amounts of sulfur species clearly, for the 3% S/B-SAC catalyst, the relative amount of C–S–C species is 75.36% and C–SO_x–C species is 24.64%. It can be found that the content of C–S–C species is increased with the increasing of sulfur precursor content. The 9% S/B-SAC catalyst contains C–S–C species of 85.19% and C– SO_x–C species of 14.81%, which has the maximum content of C–S–C species. Combing with the active data of catalysts, it can be speculated that the sulfur species of C–S–C could be the catalytic active site for the reaction. For the previous study, oxidize sulfur can be converted to sulfide under the treatment condition of calcinations for carbon materials. Due to the close electroneutrality for sulfur and carbon, the C–S bonds are predominately at the edge or the defect sites and the presence of defect sites promotes the reaction.¹⁸

To further elucidate the correlations among the existence form of sulfur and catalytic activity, and ensure the catalytic active sites of the catalyst, the Raman spectroscopy was analyzed. The consequences are shown in Fig. 6, the I_D/I_G value

Zhang et al. reported¹⁹ that the spin density should be more important in determining the catalytic active sites, which can be studied by density functional theory (DFT). Therefore, the method of DFT simulation is adopted to investigate the influence of sulfur doping on electronic and chemical properties of carbon. The on-site Bader charge of undoped and doped model carbon layer are exhibited in Fig. 7(a) and (b). After the replacement of C atom to S atom on the edge of the activated carbon layer, the two C atoms, which is adjacent to S atom is named C1 and C2. The electric charge density of C1 and C2 is -0.02 and -0.01 for the undoped model carbon layer, respectively. For the doped model carbon layer, the electric charge density of C1 and C2 is -0.38 and 0.02, respectively. The doping of S on carbon can change charge density and broken the chemical inertia to a certain extent of carrier, which promotes the electron transfer between carrier surface and reactants. Fig. 7(c) and (d) show the spin density isosurface of undoped and doped model carbon layer. It can be observed clearly that the C atom adjacent to S atom on the doped model carbon layer

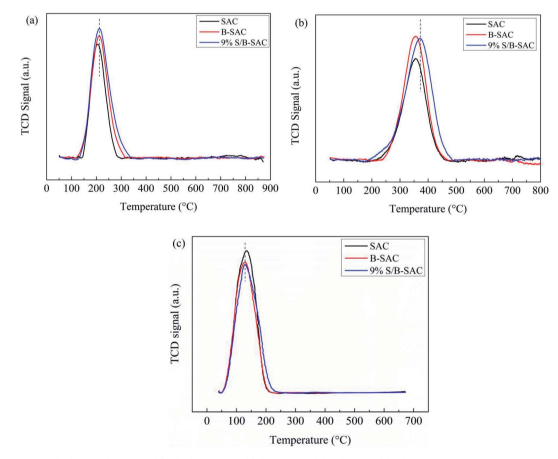


Fig. 8 TPD profiles of SAC, B-SAC and 9% S/B-SAC catalysts: (a) C₂H₂-TPD, (b) HCI-TPD, (c) VCM-TPD.

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exhibits obvious variation of spin density, which compares with that of the undoped model carbon layer. Zhang *et al.* reported¹⁹ that S-doped carbon materials would induce a larger spin and charge density to neighboring carbon and create more active sites of the ORR reaction. Combined with the DFT simulation results, we believe that S-doped carbon catalyst promotes the reaction and the C–S bond at the edge or the defect sites is the active sites for acetylene hydrochlorination.

3.2.3 Adsorption properties of S-doped catalyst. TPD experiments were measured to evaluate the adsorption properties of the reactants and product. Here, we selected three types of catalysts (SAC, B-SAC and 9% S/B-SAC) to explore the differences between undoped carbon catalyst and S-doped carbon catalyst. Fig. 8 displays the C2H2, HCl and VCM profiles on different catalysts. The individual desorption amounts are listed in Tables 6 and 7. Fig. 8(a) shows the C₂H₂-TPD profiles of the fresh SAC, B-SAC and 9% S/B-SAC catalysts. The desorption peak of C₂H₂ can be observed in the temperature range 100-350 °C in all catalysts. The fresh 9% S/B-SAC catalyst shows the maximum desorption area of acetylene comparing with the other catalysts, which suggests that the doping of sulfur on carbon enhancing the ability of acetylene adsorption. In addition, HCl-TPD profiles of the fresh SAC, B-SAC and 9% S/B-SAC catalysts are shown in Fig. 8(b). The desorption peak of HCl can be observed in the temperature range 200-500 °C in all catalysts. The binding strength (371 °C) of HCl with 9% S/B-SAC was obviously stronger than that of SAC (355 °C) and B-SAC (356 °C). The 9% S/B-SAC catalyst also exhibits the largest desorption area of HCl for all samples. The consequence of DFT shows that the doping of S creates more active sites at the edge of carbon. The broken of chemical inertia of carrier, which promotes the electron transfer between carrier and reactants. Combined with the results of HCl-TPD, the enhancement of hydrogen chloride adsorption capacity corresponds to the results of DFT. Therefore, the doping of sulfur effectively increases the adsorption strength of HCl and exhibits excellent catalytic activity.

 $\label{eq:table_$

Catalyst	Desorption area	Desorption temperature (°C)
SAC	2215.8	355
B-SAC	2595.9	356
9% S/B-SAC	3141.2	371

Table 7The desorption areas and desorption temperature of VCM forthe fresh catalysts

Catalyst	Desorption area	Desorption temperature (°C)
SAC	29 188.4	134
B-SAC	29 499.7	128
9% S/B-SAC	28 645.0	128

The detail results are listed in Table 6. It can be illustrated that the doping of sulfur on carbon enhancing the ability of hydrogen chloride adsorption. Fig. 8(c) shows the VCM-TPD profiles of the fresh SAC, B-SAC and 9% S/B-SAC catalysts. The VCM area of the 9% S/B-SAC catalyst is measured to be less than that of the other samples and the concrete results are shown in Table 7. In summary, the doping of sulfur on carbon not only enhances the adsorption of the reactants, but also promotes desorption of VCM product.

4. Conclusions

In summary, a series S-doped catalyst were prepared and applied in acetylene hydrochlorination. The 9% S/B-SAC catalyst exhibited obvious catalytic performance, compared to the blank carrier with the conditions of GHSV (C_2H_2) of 90 h⁻¹ and at 180 °C. The presence of sulfur species inhibited the formation of coke deposition and enhanced the ability of reactants adsorption. The characterizations indicated that the C-S bond at the edge or the defect sites was the active sites for the reaction. The study of DFT showed that the doping of S on carbon changed charge density and broken the chemical inertness to a certain extent of carrier, which promoted the electron transfer between carrier surface and reactants. Finally, it is the first report of single sulfur doping activated carbon to catalytic acetylene hydrochlorination. The desirable results are expected to encourage further investigation of S-doped catalysts by changing the source of sulfur or doping with other heteroatom.

Conflicts of interest

There is no conflicts to declare.

Acknowledgements

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References

- 1 G. J. Hutchings, J. Catal., 1985, 96, 292.
- 2 M. Conte, A. F. Carley, G. Attard, A. A. Herzing, C. J. Kiely and G. J. Hutchings, *J. Catal.*, 2008, **257**, 190–198.
- 3 K. Zhou, W. Wang, Z. Zhao, G. Luo, J. T. Miller, M. Wong and F. Wei, *ACS Catal.*, 2014, 4, 3112.
- 4 H. Zhang, B. Dai, W. Li, X. Wang, J. Zhang, M. Zhu and J. Gu, *J. Catal.*, 2014, **316**, 141–148.
- 5 P. Johnston, N. Carthey and G. J. Hutchings, *J. Am. Chem. Soc.*, 2015, **137**, 14548–14557.
- 6 X. Qi, W. Li, J. Gu, C. Guo and J. Zhang, *RSC Adv.*, 2016, 6, 105110–105118.
- 7 J. Zhao, S. Gu, X. Xu, T. Zhang, Y. Yu, X. Di, J. Ni, Z. Pan and X. Li, *Catal. Sci. Technol.*, 2016, **6**, 3263–3270.
- 8 S. Shang, W. Zhao, Y. Wang, X. Li, J. Zhang, Y. Han and W. Li, *ACS Catal.*, 2017, 7, 3510–3520.

- 9 Z. Zhao, M. Li, L. Zhang, L. Dai and Z. Xia, *Adv. Mater.*, 2015, 27, 6834–6840.
- 10 X. Li, X. Pan, Y. Liang, P. Ren, X. Wu, L. Sun, F. Jiao and X. Bao, *Nat. Commun.*, 2014, 5(1), 1–7.
- 11 X. Li, Y. Wang, L. Kang, M. Zhu and B. Dai, *J. Catal.*, 2014, **311**, 288–294.
- 12 B. Dai, K. Chen, Y. Wang, L. Kang and M. Zhu, *ACS Catal.*, 2015, 5, 2541–2547.
- 13 X. Dong, S. Chao, F. Wan, Q. Guan, G. Wang and W. Li, J. Catal., 2018, 359, 161–170.
- 14 J. Wang, F. Zhao, C. Zhang, L. Kang and M. Zhu, *Appl. Catal.*, *A*, 2018, **549**, 68–75.

- 15 H. Zhang, B. Dai, X. Wang, W. Li, Y. Han, J. Gu and J. Zhang, *Green Chem.*, 2013, **15**, 829.
- 16 B. Nkosi, M. D. Adams, N. J. Coville and G. J. Hutchings, *J. Catal.*, 1991, **128**(2), 378–386.
- 17 H. C. Chang, H. P. Sung and I. W. Seong, *Green Chem.*, 2011, 13, 406.
- 18 Z. Yang, Z. Yao, G. Li, G. Feng, H. Nie, Z. Liu, X. Zhou, X. Chen and S. Huang, ACS Nano, 2012, 6(1), 205–211.
- 19 L. Zhang, J. Niu, M. Li and Z. Xia, *J. Phys. Chem. C*, 2014, **118**, 3545–3553.