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The study on continuous denitrification, desulfurization of pyrolusite/activated coke hybrid catalyst†

Lin Yang,^{ID}^a Lu Yao,^{ab} Wenju Jiang,^{*ab} Xia Jiang^{ab} and Jianjun Li^{ab}

In this study, the industrial application of pyrolusite/activated coke hybrid catalysts (ACP) for continuous SO₂ and NO removal was simulated and studied systematically. The obtained results show that the denitrification process of ACP before SO₂ removal had almost no influence on its desulfurization activity even though the surface functional groups showed variation. The denitrification property of ACP, however, significantly enhanced after desulfurization and regeneration. The NO removal efficiency of ACP increased from 74.2% to 100% and remained relatively stable, and the N₂ selectivity was higher than 90.2%. The formed metal sulfate (MnSO₄ and Fe₂(SO₄)₃), showed the catalytic activity, which synergistically worked with surface functional groups to improve the selected catalytic reduction (SCR) of NO. However, large quantity of the formed metal sulphates restricted the mass transfer on the surface of ACP, which caused the reduction of desulfurization performance and the hypothesis of NO conversion of ACP-Rn.

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1. Introduction

With the significant efforts contributed from the national and local government for air pollution control in recent years, the sulphur oxides (SO_x) and nitric oxides (NO_x) emission, which primarily caused the acid rain, photochemical smog and haze pollution,^{1–3} has been effectively controlled and the environmental quality greatly improved across China. However, a further control on the emission of such compounds is needed because of the implementation of new Ambient Air Quality Standards (GB 3095-2012), which reported a more strict emission limitation on almost all pollutants. Therefore, improvements in desulfurization and denitrification of the flue gas (FGD), the most common purification method of the sintering exhaust, are still an important challenge for air quality improvement.

After years of survey and investigation, significant amount of funds and effort have been devoted to develop new technologies and materials in order to control the emission of SO₂ and NO_x.

Activated coke (AC), a high efficiency and low cost porous carbon material, has obtained increasing attention in recent decades because it can be control the emission of SO₂, NO_x,^{4–6} as well as the heavy metal/toxic waste.^{7,8} However, the issue of deactivation of catalytic activity of AC due to formation of (NH₄)₂SO₄/NH₄HSO₄ during the simultaneous removal of SO₂ and NO_x has not been dealt effectively,⁶ which limits its applicability significantly because of the decreased activity and increased operation cost.⁹ Therefore, the reactor designers usually divide the moving bed reactor into two parts (upper and bottom section) to ensure the high efficiency of purification. The bottom section is used to remove SO₂, heavy metals and toxins, while the upper section is used for denitrification. The flue gas, primarily containing SO₂, NO_x, and other toxic gases, is introduced into the system from the bottom to first remove SO₂, toxic gases and most of the steam, and then completely remove NO_x by passing NH₃ or CO in the upper section.¹⁰ The direction of AC flow is kept opposite to that of the flue gas; AC enters the top of reactor and first removes NO, and then descends to the bottom section to remove SO₂, air toxins and steam. After NO and SO₂ removal, the SO₂/toxics-filled AC is discharged and sent to the regenerator, and then back to the purification plant for reuse of AC after regeneration. Unfortunately, almost all of the previous research were based on only the desulfurization and denitrification separately¹¹ or simultaneously by AC,^{12,13} nobody had studied the continuous desulfurization and denitrification of AC, which is essential for industrial applications. Tsuji *et al.* had reported that the NO removal efficiency increased from 45.0 to 80.0% when AC was regenerated after NO and SO₂ removal,¹⁴ but they only described the variation of NO removal activity, and

^aCollege of Architecture and Environment, Sichuan University, Chengdu 610065, P. R. China. E-mail: wenjujiang@scu.edu.cn; xjiang@scu.edu.cn

^bNational Engineering Research Center for Flue Gas Desulfurization, Chengdu 610065, P. R. China

† Electronic supplementary information (ESI) available: Fig. S1 the diagram of the denitrification and desulfurization system. Fig. S2 the N₂ adsorption–desorption curves of the ACP and ACP-Rn. Fig. S3 the wide-scan XPS spectrums of the ACP-Rn. Fig. S4 the C 1s curves deconvolution of the ACP and ACP-Rn before and after NO removal. Fig. S5 the Mn 2p and Fe 2p curves deconvolution of the ACP and ACP-Rn before and after denitrification. Fig. S6 the NO removal curves of the ACP-400-Rn. See DOI: 10.1039/c7ra11720e



there was no mention of why the NO removal efficiency improved and its mechanism. It was reported that the physicochemical properties of AC constantly change with continuous usage, it is important to point out what the reciprocities of these two process and how it works.

Moreover, researchers have also made quantity of efforts to improve activity and reduce preparation cost of AC in order to make it more suitable for industrial application.¹⁵ Among these technologies, surface modification by loading transition metals attracted great attention.^{6,16} Natural minerals, rich in transition metals, showed great prospect due to its relatively low cost and the simple preparation procedure when it used as precursor,¹⁷ and the catalytic activity and stability of some kinds of natural minerals for SO₂ and NO removal also have been proved.^{18,19} However, it is noteworthy that the formation of metal sulphates during the desulfurization process on the surface of modified AC is unavoidable for almost all the transition metals, and they are difficult to be decomposed under commonly used regeneration temperatures (≤ 500 °C).²⁰ In our previous research, we had discussed the effects of MnSO₄ on SO₂ removal over MnO₂ modified AC by lending method (*i.e.* Mn₄/AC).²¹ However, what the effects of the formed metal sulphates on NO removal of such hybrid carbon-based catalyst are still unknown. Therefore, it has as great theoretical significance and practical value (in terms of industrial applications) to study the continuous application of AC for denitrification and desulfurization.

In this study, pyrolusite/activated coke catalyst (ACP) was prepared by the blending modification of fresh AC. The continuous denitrification and desulfurization property of prepared sample was investigated on a lab-scale to evaluate its potential for industrial applications. The structure properties and surface chemistry were characterized by BET, XPS, XRD and FTIR. The metal sulphate impregnated AC (AC-IM) and dilute sulphuric acid treated AC (AC-DSC) were prepared to investigate the functions of metal sulphate and surface functional groups formed on regenerated ACP after NO and SO₂ removal, which are helpful to point out the reciprocities and the work mechanism between these two processes.

2. Materials and methods

2.1 Preparation of activated coke

In this study, two types of coal as raw carbon materials, coal tar as binder and pyrolusite as catalyst precursor were used to prepare the pyrolusite-doped AC (ACP); the approximate elemental analysis of the two types of coal and coal tar are listed in Table 1. Pyrolusite used in this study majorly consists of Mn

(39.40%), Fe (6.27%), Si (4.10%), Ca (1.30%), Ni (0.07%), Cu (0.06%) and Co (0.06%). The ACP preparation is almost similar to our previous study;¹⁹ the difference is that the blending ratio of pyrolusite is 10.0 wt%. The metal sulphate impregnated AC (AC-IM) was prepared by equivalent-volume impregnation method. Initially, 3.0 wt% of MnSO₄ (AR grade) and 0.6 wt% of Fe₂(SO₄)₃ (AR grade) based on the weight ratio of AC sample were simultaneously dissolved in deionized water. AC was soaked in sulphate solution for 24 h to ensure uniformity of mixture. Following this, the mixture was first dried in a water bath at 60 °C for 2 h to release most of the water, and then placed in an oven at 105 °C overnight. The dilute sulphur acid treated AC (AC-DSC) was also prepared by equivalent volume impregnation method. Fresh AC was immersed in dilute sulphuric acid over 24 h to obtain 10% loading ratio of sulphur acid, and then dried in an oven at 105 °C overnight. Following this, the sulphur acid impregnated AC was calcined at 500 °C for 60 min in N₂ atmosphere to simulate the regeneration of AC after SO₂ removal.

2.2 Characterization of activated coke

Nitrogen adsorption capacity of ACs was measured using ASAP 2460 analyser (Micromeritics, USA) at 77 K. The samples were degassed at 523 K for 8 h before analysis. The specific surface areas (SSA) were calculated using Brunauer–Emmett–Teller equation.²² The total pore volume was estimated with respect to the liquid volume of N₂ adsorbed at a relative pressure of 0.995. The micropore (V_{mic}) volume and micropore surface area (MSA) were obtained by *t*-plot theory. The mesopore volume (V_{mes}) was determined by Barret–Joyner–Halenda (BJH, adsorption) model. X-ray photoelectron spectroscopy (XPS) characterization of the surface was carried out on a XSAM-800 spectrometer (KRATOS, UK) with Al-K α radiation (1486.60 eV) under ultra-high vacuum; the source was operated at 12 kV and 15 mA. Energy calibration was carried out by recording the core level spectra of Au 4f_{7/2} (84.00 eV) and Ag 3d_{5/2} (368.30 eV). High-resolution scans (0.10 eV) were recorded over the range of 278.00–294.00 eV (C 1s), 630.00–665.00 eV (Mn 2p) and 700.00–738.00 eV (Fe 2p). The crystal structure of activated coke was determined by X-ray diffraction (XRD) using an X-Pert PRO MPD diffractometer (Panalytical, NL) employing CuK α radiation at 30 kV and 20 mA and step-scanning over 2θ range of 10.00–80.00°. The crystalline phases were identified by comparison with the reference data from the International Center for Diffraction Data (JCPD). The Fourier-Transform Infrared Spectroscopy (FTIR) characterizations were carried out in the 4000–400 cm⁻¹ region with a spectrometer (FTIR 6700, Nicolet, USA) at a resolution of 2 cm⁻¹.

2.3 Desulfurization, denitrification and regeneration

The denitrification and desulfurization activity tests were carried out on a lab-scale fixed-bed reactor system (Fig. S1†). The conditions and operation of SO₂ removal test were same as our previous study,¹⁹ the simulated flue gas contains 3000 ppm of SO₂, 8.0% of O₂, 10.0% of steam and N₂ as balance gas. The NO removal test of the activated coke was carried out under

Table 1 Main ingredients of raw materials and main binder (%)

Samples	Proximate analysis				Elemental analysis			
	M _{ad}	A _d	V _d	FC _d	C _{ad}	H _{ad}	N _{ad}	S _{ad}
Bituminous coal	0.85	11.83	34.11	54.06	73.78	9.69	2.29	1.76
1/3 coking coal	1.32	2.74	35.55	61.71	74.23	4.83	2.08	0.47



150 °C with the space velocity of 1000 h⁻¹. The simulated flue gas flow contains 500 ppm (670 mg m⁻³) of NO, 550 ppm (417 mg m⁻³) of NH₃, 5.0% of O₂ and N₂ as the balance gas. Before the catalytic reduction of NO with NH₃, all samples were saturated with adsorbed NO at the same conditions of denitrification in order to eliminate the adsorption effect. The selective catalytic reduction (SCR)-based NO removal was initiated when the inlet and outlet concentrations of NO were equal. The inlet and outlet SO₂ and NO were monitored online by Testo 350, and the 8 h NO removal efficiency was defined as the activity index. For N₂ selectivity calculation, the inlet and outlet gas composition were analysed by FT-IR analysis (Antaris IGS-6700, Nicolet Thermo Fisher Scientific). The NO and SO₂ removal were simulated by static experiments in order to get uniform samples. The regeneration of SO₂ captured ACP was carried out at 500 °C for 60 min in N₂ atmosphere (100 mL min⁻¹), and the heating rate of regeneration was 5 °C min⁻¹. The regenerated catalyst-sorbent was labelled as ACP-Rn, and its name was updated as ACP-Rn-NO after denitrification.

3. Results and discussion

3.1 The continuous denitrification and desulfurization performance

The NO removal efficiency of fresh and regenerated ACP (ACP and ACP-Rn) are shown in Fig. 1a. It can be observed that the NO conversion of the fresh ACP is 74.2%, which significantly improved after continuous NO and SO₂ removal and regeneration, and the improvement showed a linear increase with the increase of regeneration cycles. The NO removal efficiency increased to 93.8% after one regeneration cycle (ACP-R1), which is 19.6% higher than that of ACP. After three times of regeneration (ACP-R3), the NO removal efficiency was higher than 99.5% and remained relatively stable with further reuse. This result is clearly consistent with the previous studies that the NO removal activity was enhanced when AC was regenerated after SO₂ removal.^{14,23} The N₂O concentration in the outlet gas remained stable in the range from 30 to 35 ppm (53–62

mg m⁻³), and the corresponding N₂ selectivity was higher than 90.2%. It was reported that the coexistence of Mn and Fe can accelerate the electron transfer and promote NH₃ activation, which is favourable for the reaction between the NO and NH₃.²⁴ However, it is interesting that the ACP-Rn needs a longer working time to reach a stable NO removal activity than that of fresh ACP, and showed a strong dependence on the number of regeneration cycles. This hysteresis of NO reduction in the previous stage of denitrification could be caused by the metal sulphate generated during the desulfurization process, but this is just our deduction. The real reason and mechanism of the hysteresis is still unclear.

The desulfurization breakthrough curves of ACP and ACP-Rn after NO removal are shown in Fig. 1b. The breakthrough working time of ACP-Rn-NO linearly reduced after each regeneration cycle. The ACP-Rn, however, still had a relatively high activity for SO₂ removal after five times of regeneration; the 100% SO₂ removal time of ACP-R5 was still longer than 8 h. The sulphur capacity of ACP-R5 was 71 mg g⁻¹; which is significantly higher than that for many non-regenerated AC samples reported earlier.^{25,26} The same variation was reported by Guo *et al.*; the group found that the desulfurization performance of the used catalyst-sorbent linearly decreased, while its NO removal efficiency clearly improved.²⁷ The excellent NO removal activity and improved desulfurization performance illustrate that the used pyrolusite-modified AC (ACP) synthesized by the blending method is a good low temperature desulfurization and denitrification catalyst.

3.2 Physicochemical analysis of the regenerated ACP

3.2.1 Specific surface area and porosity. The N₂ adsorption-desorption of the used ACs were investigated to evaluate the influence of continuous denitrification, desulfurization and regeneration on their structure properties; the N₂ adsorption-desorption curves are depicted in Fig. S2.† Table 2 lists the calculation based on the analysis. The surface area (SSA) of AC is 479 m² g⁻¹, which decreased to 454 m² g⁻¹ after the first cycle of application and then remained stable after the second

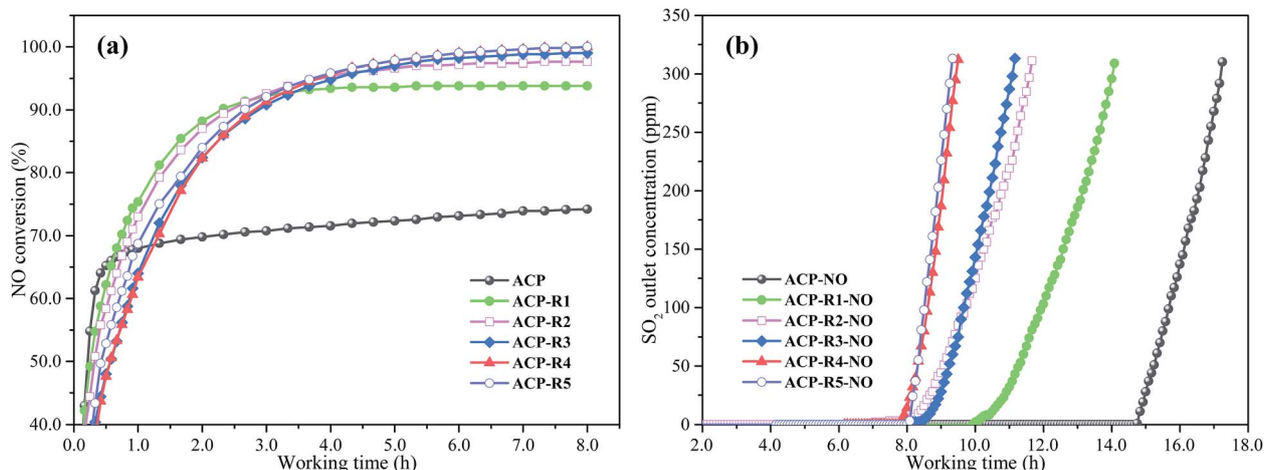


Fig. 1 NO conversion (a) and desulfurization breakthrough curves (b) of ACP-Rn.



Table 2 Textual properties of activated coke samples^a

Samples	SSA (m ² g ⁻¹)	MSA (m ² g ⁻¹)	V _{tot} (cm ³ g ⁻¹)	V _{mic} (cm ³ g ⁻¹)	V _{mes} (cm ³ g ⁻¹)	R _{mean} (nm)
ACP	479	385	0.277	0.186	0.091	2.31
ACP-R1	454	362	0.257	0.179	0.079	2.37
ACP-R2	452	377	0.275	0.186	0.088	2.48
ACP-R3	470	393	0.287	0.194	0.093	2.49
ACP-R4	477	397	0.289	0.196	0.093	2.38
ACP-R5	506	415	0.294	0.205	0.089	2.37
ACP-R5W	590	490	0.327	0.234	0.093	2.25
AC	458	379	0.267	0.187	0.080	2.15
AC-IM	431	346	0.245	0.170	0.075	2.27
AC-DSC	502	413	0.279	0.205	0.074	2.11

^a SSA is specific surface area, MSA is micropore surface area, V_{tot} is total pore volume, V_{mic} is micropore volume, V_{mes} is mesopore volume and R_{mean} is average pore diameter.

regeneration cycle (452 m² g⁻¹ of ACP-R2). After that, the SSA linearly increases with the increased reusing cycles and completely recovered after the fourth regeneration cycle (ACP-R4) with SSA at 477 m² g⁻¹. The total pore volume (V_{tot}) and micropore volume (V_{mic}) showed the same change trend with SSA and recovered after two cycles of regeneration. In addition, it should be noted that there was almost no change in V_{mes}, which maintained at around 0.090 cm³ g⁻¹ except for ACP-R1 (0.079 m³ g⁻¹). The reduction of SSA, V_{tot} and V_{mic} in the first few cycles is due to the formation of metal sulphates during the SO₂ removal; the removed SO₂ primarily transformed to form sulphate, and they could not decomposed at the regeneration temperature.²¹ With the increasing number of regeneration cycles, there is a higher ratio of removed SO₂ stored in AC as sulphur acid. The sulphuric acid, reacting with carbon, played the role of further activation of ACP and improved its porosity. The variation of V_{mes}, shown in Table 2, suggests that the regeneration of ACP at 500 °C primarily created the micropore structure, and there was no clear pore diameter expansion because the mesopore volume was not clearly changed in the first few regeneration cycles. The relative stable average pore diameter (R_{mean}) was a good testimony.

Our previous study showed that the SSA of pyrolusite-doped AC recovered quickly and then continuously increased for more than 10 reusing cycles.¹⁹ The slower recovery rate in this study is attributed to the higher blending ratio of pyrolusite; the 10.0 wt% blending ratio of pyrolusite is 2.5 times higher than the sample used in our previous study (4.0 wt%). This indicates that there was higher metal sulphate formation when the ACP was used for SO₂ removal. To confirm this deduction, ACP-R5 was washed using deionized water (labelled as ACP-R5W). The BET analysis shows that the SSA and V_{tot} of ACP-R5W (590 m² g⁻¹ and 0.327 cm³ g⁻¹) were dramatically higher than that of ACP-R5 (506 m² g⁻¹ and 0.294 cm³ g⁻¹). The V_{mes} of ACP-R5W (0.093 cm³ g⁻¹) conformed with our previous result, indicating that the reaction between the sulphur acid and carbon at 500 °C only formed some content of micropore structure, and there was no clear pore diameter expansion during the regeneration. In addition, increased micropore volume caused the reduction of R_{mean} of ACP-R5W (2.25 nm).

Corroboration of the NO removal activity of regenerated ACP and their structure properties suggests that the SSA and pore structure were not the limiting factors for NO removal because the NO conversion of ACP-R1 and ACP-R2 were significantly higher than that of ACP although their structure properties are worse. The variation of the desulfurization and denitrification could be primarily controlled by their surface chemical properties, such as modification of the functional groups and metals, which would be discussed in the following section.

3.2.2 Surface functional groups. The XPS characterization was carried out to analyse the surface functional groups variation of ACP after continuously used for denitrification and desulfurization; the wide-scanning spectrums of ACP and ACP-Rn are exhibited in Fig. S3.† For the fresh ACP, only four clear peaks attributed to the Mn, Fe, O and C were detected, while there was a new peak, which is attributed to sulphur (S 2p), detected on the surface of ACP-Rn. Furthermore, the low intensity new peak corresponding to N 1s also appeared after three regeneration cycles. The S 2p peak is attributed to the formed metal sulphates after flue gas desulfurization; many previous studies had discussed about this.²⁰ The presence of N 1s could be due to the adsorbed NH₃ and NO, particularly the NH₃ species, which could react with H₂SO₄ to form (NH₄)₂SO₄/NH₄HSO₄, and then aminate the coke surface during the high temperature regeneration.²⁸ No N-containing functional groups on the surface of ACP-R1 and ACP-R2 were detected, which indicates that the adsorbed NO and NH₃ participated in SCR NO removal effectively. However, with the increase of regeneration number, the textual properties of activated coke improved (Table 2), which are more favourable for NO and NH₃ adsorption. In addition, there was some amount of adsorbed NO and NH₃ available for the reaction following desulfurization.

To evaluate the variation of surface oxygen-containing functional groups, the binding energy (BE) and relative content (RCs) of ACP and ACP-Rn corresponding to the C 1s spectra (Fig. S4†) are calculated. As summarized in Table 3, the RCs of C–C bond on the surface of catalyst-sorbent reduced gradually from 1st to 4th regeneration cycle, and then showed a slight increment with further reuse. However, the content of C=O and π–π* transition peak was higher than that of ACP even though ACP-R5 showed a reduction compared with ACP-



Table 3 Binding energy (BE) and relative content (RCs) of C 1s for activated coke

Samples	C-C		C-O		C=O		$\pi-\pi^*$	
	BE (eV)	RC (%)	BE (eV)	RC (%)	BE (eV)	RC (%)	RC (%)	BE (eV)
ACP	284.63	70.34	285.90	19.09	287.61	6.18	289.97	4.39
ACP-NO	284.64	68.56	285.73	19.75	28.15	6.24	289.47	5.45
ACP-R1	284.65	69.95	285.80	18.02	287.32	7.08	289.49	4.96
ACP-R1-NO	284.65	67.87	285.79	19.59	287.14	7.10	289.16	5.43
ACP-R2	284.63	67.08	285.93	19.06	287.39	8.84	289.40	5.03
ACP-R2-NO	284.60	64.57	285.98	20.03	287.70	9.26	289.76	6.14
ACP-R3	284.67	65.45	285.87	19.87	287.35	9.19	289.23	5.49
ACP-R3-NO	284.60	63.98	285.74	21.03	287.32	9.00	289.52	6.00
ACP-R4	284.64	65.16	286.03	18.81	287.51	9.56	289.57	6.47
ACP-R4-NO	284.67	63.74	286.03	20.10	287.36	9.27	289.37	6.89
ACP-R5	284.63	69.14	285.94	18.83	287.59	6.24	289.16	5.79
ACP-R5-NO	284.66	67.28	285.92	20.76	287.42	6.02	289.28	5.95

R4. The RCs of the C-O group initially increased, and then reduced with the number of regeneration cycles; ACP-R3 had the highest content of C-O at 20.87%. In general, the RCs of oxygen-containing functional groups of ACP-Rn were higher than those of ACP, which is primarily attributed to the high temperature carbon acid reaction, *i.e.*, $C + H_2SO_4 \rightarrow CO + CO_2 + SO_2 + H_2O$ and the pyrolysis of oxygen containing macromolecules during the regeneration.^{29,30} On comparing ACP/ACP-Rn before and after NO removal, it was found that the RCs of C-C showed a certain reduction when the ACP/ACP-Rn was used for NO removal, while the RCs of C-O, C=O and $\pi-\pi^*$ transition peak mildly increased alternatively. This could be attributed to the adsorbed oxygen, which transformed to radical oxygen and expressed as the increment of oxygen-functional groups. It was reported that the catalytic activity for NO_x and SO₂ removal of AC-based catalyst could be improved with the increase in the surface oxygen-containing functional groups,^{28,31} and it also proved to be the controlling factor for the SO₂ removal of the pyrolusite-doped AC.¹⁹ The linearly reduced

desulfurization performance of ACP-Rn was primarily due to its large content of metal sulphate dispersed on the surface, which limited the catalytic oxidation of SO₂ on the surface of catalyst-sorbent.²¹

3.2.3 Surface metal chemistry. The X-ray powder diffraction (XRPD) of ACP and ACP-Rn was applied to clarify the metal chemistry. As shown in Fig. 2, the detected SiO₂ on the surface of all samples is due to the two types of coals and pyrolusite as raw materials, which contain certain amount of SiO₂.¹¹ The peaks corresponding to crystalline MnO (JCPDs no. 75-0257/04-0326) and Fe⁰ (JCPD 87-0722) on the surface of ACP imply that manganese and iron oxides in the pyrolusite were reduced in the strong reduction atmosphere under the high activation temperature (900 °C).²¹ After desulfurization and denitrification, however, MnO and Fe⁰ diffraction peaks weakened or even disappeared (ACP-Rn). There were some new peaks corresponding to MnSO₄·H₂O ($2\theta = 18.22^\circ$, JCPD 07-0230), MnSO₄ ($2\theta = 25.32^\circ$ (ref. 20)), MnO₂ ($2\theta = 45.26^\circ$, JCPD 42-1169) and Mn₃O₄ ($2\theta = 42.94^\circ, 62.32^\circ$, JCPD 13-0162) that were detected. In addition, two small peaks at $2\theta = 38.82^\circ, 50.29^\circ$ corresponding to FeS were also detected. It is known that the coexistence of Mn⁴⁺ and Fe²⁺ could accelerate the electron transfer, which promote the NH₃ activation on the surface of ACP and further accelerate NO removal.²⁴ The detected sulphide could be due to reactions between iron oxide and SO₂ during the thermal regeneration³² under a strong reducing atmosphere along with transition metal catalysts.

The chemical properties of Mn and Fe were further confirmed by XPS analysis, and high-resolution Mn 2p and Fe 2p spectra as depicted in Fig. S5.† The metal chemistry of ACP-Rn changed slightly with continuous application. The peaks of ACP-Rn at around 642.00–642.50 eV and 710.50–711.50 eV were consistent with those of Mn²⁺ of MnSO₄ (ref. 20) and Fe²⁺ of FeSO₄,³³ respectively. The unclear binding energy shift of Mn 2p and Fe 2p of ACP-Rn is attributed to the presence of large quantity of metal sulphates, which covered the new exposed metal oxides after regeneration. Fig. 3 shows the high-resolution spectrum of sulphur in ACP-Rn to corroborate the metal chemistry. ACP-Rn showed doublet peaks in the BE range

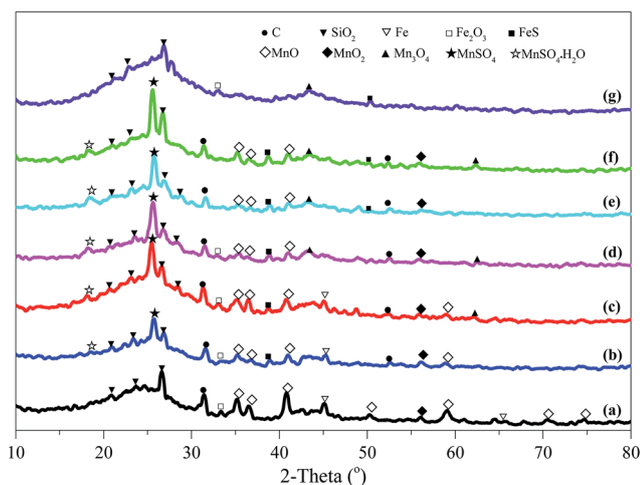


Fig. 2 XRD patterns of cyclically regenerated activated coke. (a) ACP, (b) ACP-R1, (c) ACP-R2, (d) ACP-R3, (e) ACP-R4, (f) ACP-R5, (g) ACP-R5W.



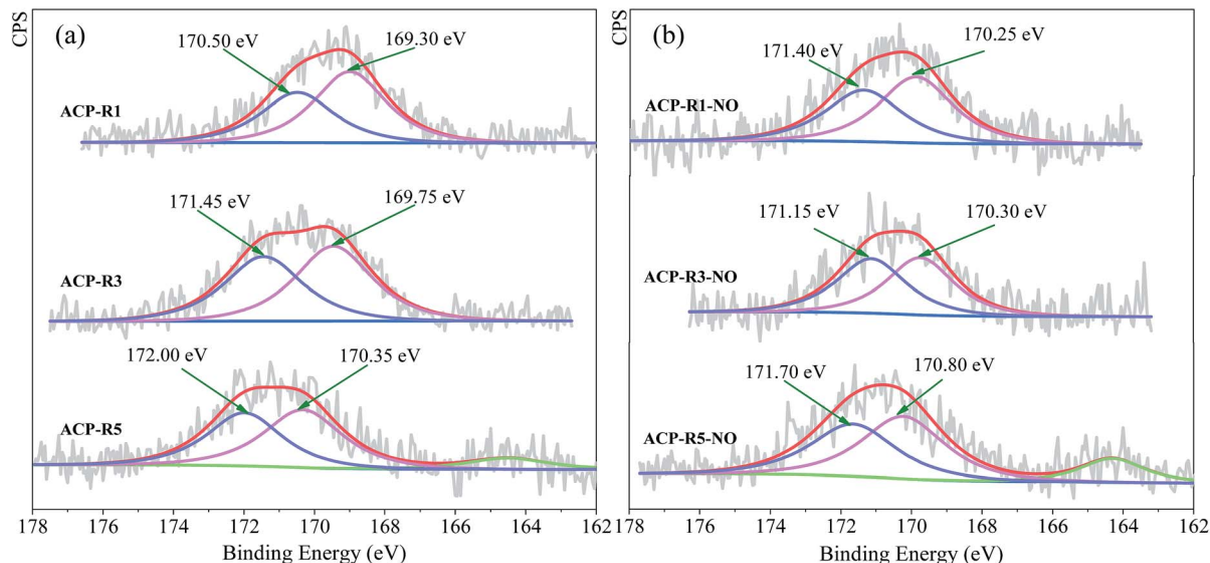


Fig. 3 Deconvoluted S 2p peaks corresponding to ACP-Rn (a) and ACP-Rn post NO removal (b).

of 169.30–172.00 eV, which is normally associated with those of SO_4^{2-} .³⁴ The relative higher binding energy was due to the large amount of metal sulphate;³⁵ the increase in binding energy with the increase in number of regeneration cycles, 170.50 of ACP-R1, 171.45 of ACP-R3 and 172.00 eV of ACP-R5 (Fig. 4a), is good evidence to affirm the above hypothesis.

The comparison of ACP-Rn before and after denitrification suggests that the right B.E. peak shifted to a higher binding energy, while the that on the left reduced slightly after the denitrification except for ACP-R1-NO (Fig. 4b). This could be due to the NO, NH_3 and O_2 adsorption during the NO removal, but this is just a hypothesis and needs a further discussion. In addition, there was no XPS peak corresponding to S^{2-} (Fig. 3), which could be due to its low content and uniform distribution; also the large content of sulphate covered the surface completely. However, a new peak for ACP-R5 and ACP-R5-NO at 164.55 and 164.30 eV, respectively, shows the presence of elemental sulphur.³⁶ Klinzing *et al.*

had reported that carbon could react with SO_2 to form the elemental sulphur ($\text{C} + \text{SO}_2 \rightarrow \text{S}_2 + \text{CO}_2$).³⁷ Therefore, whether the metals accelerate this reaction needs to be studied further.

3.3 The reciprocities between denitrification and desulfurization process

3.3.1 The influence of NO removal on desulfurization. Sulphur removal capacity of the ACP-Rn before and after NO removal, and before desulfurization is shown in Fig. 4. The desulfurization performance of ACP-Rn after NO removal gradually reduced with increasing number of regeneration cycles, while the sulphur removal capacity reduced from 138.5 to 71.6 mg g^{-1} after five using cycles, which is only 58.2% of the sulphur removal capacity of ACP. The sulphur removal capacity of ACP (without denitrification, ACP-Rn-N) showed the similar change; its sulphur capacity reduced from 138.5 to 72.9 mg g^{-1} . The maximum difference was obtained after the third using cycle, but it was only 6.3 mg g^{-1} , which could be attributed to the experimental error. The ignorable difference of sulphur removal capacity between ACP-Rn and ACP-Rn-N illustrates that NO removal only had a slight effect on its SO_2 removal activity; this finding is well consistent with the previous report.³⁸

3.3.2 The influence of desulfurization-regeneration on NO removal. To evaluate the effect of the formed metal sulphates during SO_2 removal after NO reduction, MnSO_4 and $\text{Fe}_2(\text{SO}_4)_3$ co-impregnated AC (AC-IM) was prepared to simulate the regenerated ACP, but without any variation of surface functional groups and metals. As summarized in Table 2, the impregnation of sulphates caused a great reduction of SSA (from 458 to 431 $\text{m}^2 \text{g}^{-1}$) and pore volume (from 0.267 to 0.245 $\text{m}^3 \text{g}^{-1}$) of the fresh AC, which would limit the NO removal activity of AC, in theory, because of poor molecular transfer. The AC-IM, however, showed a great efficiency of denitrification (Fig. 5a), whose NO removal efficiency increased to 60.5%,

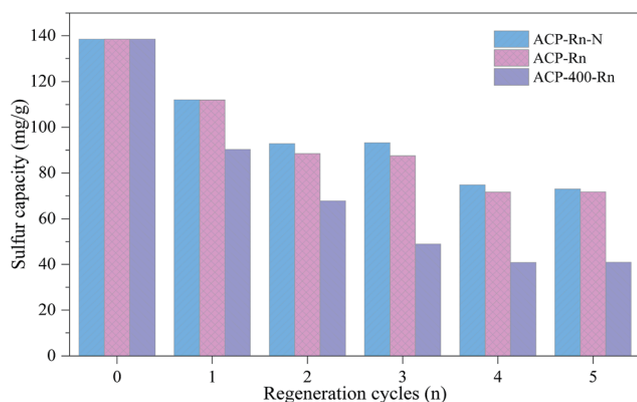


Fig. 4 Sulphur removal capacity of ACP-400-Rn and ACP-Rn before and after denitrification, and before desulfurization.



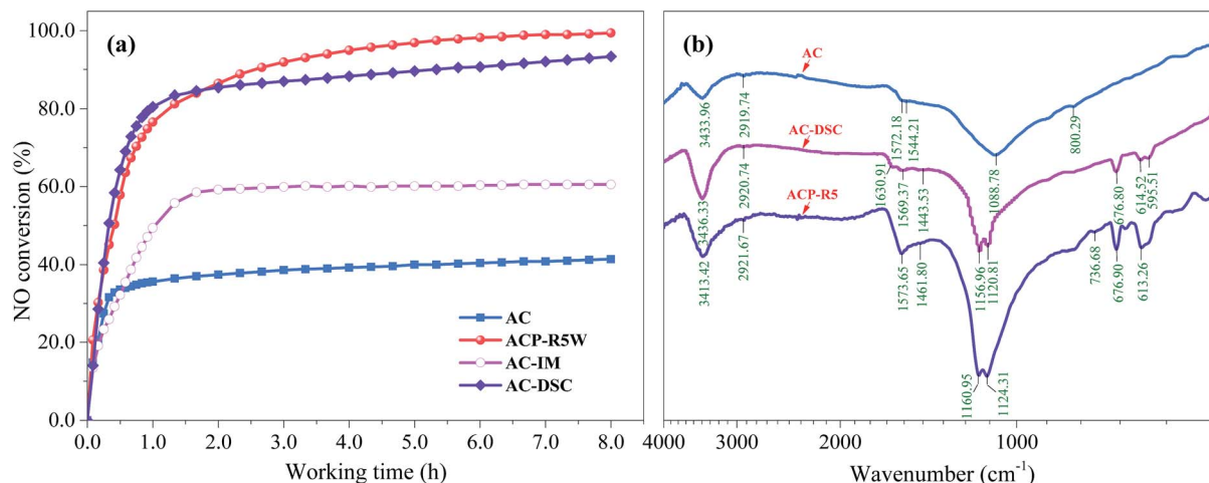


Fig. 5 NO removal activity (a) and FTIR spectra (b) of treated AC, AC-DSC and AC-R5.

which is 19.1% higher than fresh AC. Such improvement in efficiency reveals that the metal sulphate on the surface of AC enhanced the catalytic reduction of NO. Furthermore, the NO removal curve showed a similar hysteresis for ACP-Rn, which confirmed the coverage effect of metal sulphate.

To further discuss the variation of NO removal activity of AC-Rn, ACP-R5W was also applied for NO removal. Fig. 5a shows that the NO removal efficiency of ACP-R5W was 99.4%, which was only slightly different compared with ACP-R5; also, the activity hysteresis inevitably weakened. Due to the trace metals on the surface of ACP-R5W (Fig. 2g), the dilute sulphuric acid treated AC (AC-DSC) was prepared to simulate the surface chemistry of regenerated AC. The NO removal efficiency of AC-DSC was 93.4% (Fig. 5a), which is 53.0% and 32.9% higher than that of fresh AC (40.4%) and AC-IM (60.5%), respectively. These results indicate that the improvement of NO removal activity of regenerated ACP was due to the synergistic effect of surface chemistry and metal sulphates, but the surface chemistry played a prominent role for NO reduction than that of metal salts.

It has been reported in literature that sulphur-containing species promote the NO conversion of AC effectively, which could reduce the energy barrier of the catalytic oxidation of SO₂.³⁹ Therefore, the FTIR characterization of AC, AC-DSC and AC-R5 were carried out and the resultant spectra are depicted in Fig. 5b. It could be observed that there were two new peaks of AC-DSC and ACP-R5 detected at the wavenumber range from 600.00 to 680.00 cm⁻¹. The peaks at 614.52 and 613.26 cm⁻¹ of AC-DSC and AC-R5 attributed to the symmetric and asymmetric stretching vibration of S–O or S=O in sulphate ion,⁴⁰ while the peaks located at 676.80 cm⁻¹ of AC-DSC and 676.90 cm⁻¹ of ACP-R5 attributed to the O=C=O bond.⁴¹ Furthermore, the peaks around 1570.00 cm⁻¹ of the AC-DSC and ACP-R5, attributed to the asymmetric stretching vibration of C=O, had higher intensity.²⁸ Such variation of surface chemistry could partially explain the improvement of NO removal by regenerated ACP.

3.3.3 The effect of regeneration temperature. Although the lower regeneration temperature (<500 °C) could not affect the

chemistry of metal sulphates, the functional groups would be greatly varied.⁴² Therefore, a new batch ACP was regenerated at 400 °C (labelled as ACP-400-Rn) after NO and SO₂ removal and was thoroughly analysed to further investigate the influence of surface functional groups. As is shown in Fig. 6, the NO removal capacity of ACP-400-Rn increased with the increase in the number of regeneration cycles, but the increment was lower than that of ACP-Rn, particularly in the first and second reusing cycle, and the stable time of NO removal incontestably became longer (Fig. S6†). The NO removal efficiency of ACP-400-R1 and ACP-400-R2 were 77.8% and 85.4%, which are drastically lower than those of ACP-R1 (93.8%) and AC-PR2 (97.6%). The ACP-400-R4 had the highest NO removal efficiency (98.6%). However, the key difference is the slight reduction of NO removal efficiency of ACP-400-R5 compared with ACP-400-R4. This indicates that the activity of regenerated ACP cannot hold with a lower regeneration temperature. The sulphur capacity of ACP-400-Rn was lower than ACP-Rn. The ACP-400-R1 showed the largest reduction; the sulphur removal capacity decreased by 47.7 mg g⁻¹ to 90.8 mg g⁻¹, and the difference

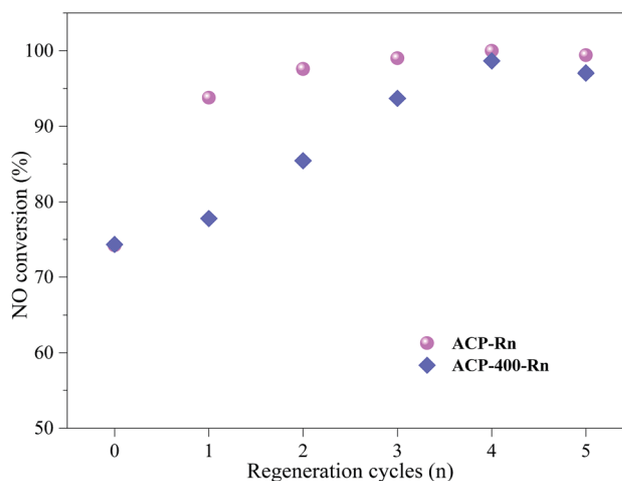


Fig. 6 8 h NO conversion of ACP-Rn and ACP-400-Rn.



became greater with the number of regeneration cycles. These differences reveal that ACP prepared by blending method needs a relatively higher regeneration temperature (500 °C) in order to have a longer life time and high efficiency of reuse.

4. Conclusion

Continuous denitrification, desulfurization and regeneration of ACP showed that the NO removal application before desulfurization had almost no influence on the SO₂ removal of sample even though the surface functional groups of ACP showed a variation after NO removal. The denitrification property of ACP, however, was significantly improved after the followed desulfurization and regeneration. The NO removal efficiency of ACP-Rn increased from 74.2% to almost 100% after three cycles of regeneration, and then remained relatively stable. The N₂ selectivities were higher than 90.2% for all samples. Metal sulphates (MnSO₄ and Fe₂(SO₄)₃), showed catalytic activity for SCR NO removal. The synergistic effect of metal sulphates and surface functional groups improved the SCR NO removal of ACP even though the later one was proved to be more important. However, large quantity of metal sulphate restricted the migration of reactants molecular (*i.e.* NO, NH₃, SO₂ and O₂), leading to the reduction in desulfurization activity and the hypothesis of NO conversion at beginning.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- I. Mochida, Y. Korai, M. Shirahama, S. Kawano, T. Hada, Y. Seo, M. Yoshikawa and A. Yasutake, *Carbon*, 2000, **38**, 227–239.
- M. Tao, L. Chen, L. Su and J. Tao, *J. Geophys. Res.: Space Phys.*, 2012, **117**, 48–50.
- W. Li, S. Zhou, X. Wang, Z. Xu, C. Yuan, Y. Yu, Q. Zhang and W. Wang, *J. Geophys. Res.: Space Phys.*, 2011, **116**, D09301.
- S. Ding, Y. Li, T. Zhu and Y. Guo, *J. Environ. Sci.*, 2015, **34**, 37–43.
- C. Lu and M. Wey, *Fuel Process. Technol.*, 2007, **88**, 557–567.
- Y. Wang, Z. Huang, Z. Liu and Q. Liu, *Carbon*, 2004, **42**, 445–448.
- Y. Lee, H. Kim, J. Park, B. Choi, D. Choi and J. Park, *Carbon*, 2003, **41**, 1881–1888.
- P. Davini, *Carbon*, 2001, **39**, 2173–2179.
- Y. Zhao, R. Hao, T. Wang and C. Yang, *Chem. Eng. J.*, 2015, **273**, 55–65.
- G. O. David, K. Tsuji and I. Shiraishi, *Fuel Process. Technol.*, 2000, **65–66**, 393–405.
- X. Gao, S. Liu, Y. Zhang, Z. Luo and K. Cen, *J. Hazard. Mater.*, 2011, **188**, 58–66.
- Y. Zhao, Y. Han and C. Chen, *Ind. Eng. Chem. Res.*, 2012, **51**, 480–486.
- Y. Guo, Z. Liu, J. Wang, X. Xing and Z. Lei, *J. Fuel Chem. Technol.*, 2008, **36**, 36–41.
- K. Tsuji and I. Shiraishi, *Fuel*, 1997, **76**, 549–553.
- C.-M. Yang and K. Kaneko, *J. Colloid Interface Sci.*, 2002, **255**, 236–240.
- L. Yang, T. Huang, X. Jiang, J. Li and W. Jiang, *RSC Adv.*, 2016, **6**, 55135–55143.
- L. Yang, X. Jiang, T. Huang and W. Jiang, *Environ. Technol.*, 2015, **36**, 2847–2854.
- L. Yang, W. Jiang, L. Yao, X. Jiang and J. Li, *J. Chem. Technol. Biotechnol.*, 2017, DOI: 10.1002/jctb.5418.
- L. Yang, X. Jiang, W. Jiang, P. Wang and Y. Jin, *Energy Fuels*, 2017, **31**, 4556–4564.
- J. Guo, Y. Qu, S. Shu, X. Wang, H. Yin and Y. Chu, *New J. Chem.*, 2015, **39**, 5997–6015.
- L. Yang, X. Jiang, Z.-S. Yang and W.-J. Jiang, *Ind. Eng. Chem. Res.*, 2015, **54**, 1689–1696.
- H. Guedidi, L. Reinert, J. L  v  que, Y. Soneda, N. Bellakhal and L. Duclaux, *Carbon*, 2013, **54**, 432–443.
- A. Boyano, M. E. G  lvez, R. Moliner and M. J. L  zaro, *Fuel*, 2008, **87**, 2058–2068.
- S. Yang, F. Qi, S. Xiong, H. Dang, Y. Liao, P. K. Wong and J. Li, *Appl. Catal., B*, 2016, **181**, 570–580.
- J. Li, N. Kobayashi and Y. Hu, *J. Environ. Eng.*, 2007, **2**, 740–751.
- K. Jastrz  b, *Fuel Process. Technol.*, 2012, **101**, 16–22.
- Y. Guo, Z. Liu, Q. Liu and Z. Huang, *Catal. Today*, 2008, **131**, 322–329.
- K. Tsuji and I. Shiraishi, *Fuel*, 1997, **76**, 555–560.
- K. Jastrz  b, *Fuel Process. Technol.*, 2012, **104**, 371–377.
- Z. Yan, L. Liu, Y. Zhang, J. Liang, J. Wang, Z. Zhang and X. Wang, *Energy Fuels*, 2013, **27**, 3080–3089.
- C. Moreno, F. J. Maldonado and C. A. F. P  rez, *Langmuir*, 2003, **19**, 5650–5655.
- S. Mitsubishi, T. Seto, S. Mitsuoka and K. Inoue, European Patent, EP0377483, 1990.
- M. Descostes, F. Mercier, N. Thromat, C. Beaucaire and M. Gautier-Soyer, *Appl. Surf. Sci.*, 2000, **165**, 288–302.
- V. Hayez, A. Franquet, A. Hubin and H. Terryn, *Surf. Interface Anal.*, 2004, **36**, 876–879.
- C. Sun, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2004, **69**, 1129–1133.
- E. Morris, C. Jia and K. Morita, *Ind. Eng. Chem. Res.*, 2010, **49**, 12709–12717.
- G. Klinzing and R. Walker, *Fuel*, 1984, **63**, 1450–1454.
- J. M. Rosas, R. Ruiz-Rosas, J. Rodr  guez-Mirasol and T. Cordero, *Chem. Eng. J.*, 2017, **307**, 707–721.
- Z. Zhu, Z. Liu, H. Niu and S. Liu, *J. Catal.*, 1999, **187**, 245–248.
- K. Saravanan, B. Tyagi and H. C. Bajaj, *Catal. Sci. Technol.*, 2012, **2**, 2512–2520.
- P. Rowe, S. Neshyba and V. Walden, *Opt. Express*, 2011, **19**, 5451–5463.
- Y. Guo, Z. Liu, Y. Li and Q. Liu, *J. Fuel Chem. Technol.*, 2007, **35**, 344–348.

