Simultaneous absorption of NO and SO\textsubscript{2} by combined urea and Fe\textsuperscript{II}EDTA reaction systems

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SO\textsubscript{2} and NO emitted from coal-fired power plants have caused serious air pollution in China. In this work, a novel mixed absorbent, Fe\textsuperscript{II}EDTA/urea, was employed for simultaneous removal of SO\textsubscript{2} and NO in a packed tower, with a corresponding optimal ratio of 0.014 mol L\textsuperscript{-1} : 5%. The effects of various factors, such as mixed absorbent constitutions, reaction temperature, pH, O\textsubscript{2} concentration, as well as concentrations of SO\textsubscript{2} and NO, on simultaneous removal were investigated. The desulfurization efficiency was 95–99% in all tests, whereas denitrification was affected significantly by various conditions. NO removal efficiency decreased increasing oxygen concentration as well as increasing NO concentration. With an increase in temperature, pH, or SO\textsubscript{2} concentration, NO removal efficiency increased first and then decreased. Under optimal conditions, SO\textsubscript{2} removal efficiency was 100% and NO removal efficiency could exceed 91% within 80 min. The reaction mechanism was speculated according to relevant literature.

1. Introduction

Emission of SO\textsubscript{2} and NO accounts for 90–95% of NO\textsubscript{x} in fuel combustion from power plants and results in various environmental problems (e.g., acid rain, photochemical smog, and other pollution) to damage the environment and human health.\textsuperscript{1} Usually, wet flue gas desulfurization (WFGD) such as the limestone–lime method\textsuperscript{2} and selective catalytic reduction (SCR)\textsuperscript{3,4} processes can, respectively, realize SO\textsubscript{2} and NO removal for industrial application on a large scale. However, the combination of WFGD and SCR processes for simultaneous desulfurization and denitrification suffer from the need for huge areas of chemical plants, intricate processes, high operating costs, ammonia leakage, and spent catalysts. There is an urgent need for a more economical method for controlling NO and SO\textsubscript{2} emission.

The simultaneous absorption technologies of NO and SO\textsubscript{2} can be split into “dry” and “wet” processes. The former includes plasma degradation, electron beam radiation photocatalysis, sorbent adsorption, and microwave catalytic decomposition, and necessitates high cost and complex processes to obtain high absorption efficiency of SO\textsubscript{2} and NO.\textsuperscript{5–9} Wet processes applied to industrial furnaces are promising ways to conform to an ultra-low emission standard.

Simultaneous desulfurization and denitrification by wet processes mainly adopt oxidation absorption using KMnO\textsubscript{4},\textsuperscript{10} NaClO\textsubscript{3},\textsuperscript{11} NaClO\textsubscript{4},\textsuperscript{12} O\textsubscript{3} (ref. 13–15) Fenton reagents,\textsuperscript{16} UV/H\textsubscript{2}O\textsubscript{2},\textsuperscript{17} thermal/transition metal ions/persulfate,\textsuperscript{18} or ultrasound/Fe\textsuperscript{2+}/heat-coactivated persulfate.\textsuperscript{19} However, oxidation absorption processes are mostly at pilot exploration or laboratory stages due to their significant costs, low oxidation efficiency or secondary pollution. Therefore, trying to exploit new and more promising NO\textsubscript{x} and SO\textsubscript{2} simultaneous removal wet technologies has important theoretical and practical importance.

Reducing absorption with urea ([NH\textsubscript{2}]\textsubscript{2}CO) and complexing absorption (e.g., Fe(II)EDTA, hexamminecobalt(II)) are also employed for simultaneous removal of NO and SO\textsubscript{2}.\textsuperscript{20,21} However, NO removal through the reduction of [NH\textsubscript{2}]\textsubscript{2}CO is not satisfactory. NO removal efficiencies are usually <40% without other additives because of the low solubility of NO in solution. Complex absorption could weaken the impact of the low solubility of NO, but they are easily oxidized, and lose their complexing ability.\textsuperscript{22} Therefore, simultaneous removal of NO and SO\textsubscript{2} by reducing absorption or complexing absorption is barely satisfactory.

Previously, we combined urea and Fe(II)EDTA for NO removal. We noted a synergistic reaction between urea and Fe\textsuperscript{II}EDTA upon NO absorption in which urea could restrain Fe\textsuperscript{II}EDTA oxidation and, importantly, it could react with Fe\textsuperscript{II}EDTA–NO to generate Fe\textsuperscript{II}EDTA, N\textsubscript{2}, and CO\textsubscript{2}.\textsuperscript{23} A mixed absorbent was shown to improve NO removal efficiency significantly, and could maintain >76% for 80 min at optimal conditions in our previous study. However, simultaneous removal of SO\textsubscript{2} and NO with a mixed urea ([NH\textsubscript{2}]\textsubscript{2}CO) and Fe\textsuperscript{II}EDTA system has not been shown. It is necessary to study the effect of the interrelationship between SO\textsubscript{2} and NO on their

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removal efficiencies by mixed absorbents containing Fe$^{II}$EDTA and ([NH$_2$)$_2$CO].

We wished to investigate the simultaneous removal of SO$_2$ and NO from flue gas using a mixed urea ([NH$_2$)$_2$CO] and Fe$^{II}$EDTA system in a “packing tower”. The key influencing factors and mechanism of simultaneous removal of SO$_2$ and NO were investigated. Our results could provide crucial theoretical guidance for follow-up studies and industrial applications of this technology.

2. Experimental section

2.1 Chemicals

FeSO$_4$·7H$_2$O (99.0%), urea (99.0%), C$_{10}$H$_{14}$N$_2$O$_8$Na$_2$·7H$_2$O (99.0%), NH$_4$Fe(SO$_4$)$_2$·12H$_2$O (99.0%) and CaCl$_2$ (99.0%) were obtained from Tianjin Kermel Chemical Reagent Co., Ltd. (China). N$_2$ (99.999%), SO$_2$ (99.99%) and NO (99.99%) were provided by Guangzhou Yuejia Gas Co. (China). Other reagents were of analytical grade.

2.2. Experimental apparatus

A schematic diagram of the experimental setup is shown in Fig. 1. The system comprised three main parts: gas-simulation system, packed tower, and a gas sampling-analysis system. Absorption of NO and SO$_2$ was conducted in the packed tower of length 1000 mm and internal diameter 20 mm. The pressure in the packed tower was atmospheric. The concentration of NO and SO$_2$ was measured by a flue gas analyzer (KM950, KANE International) after drying with anhydrous calcium chloride. The resolution and accuracy of this gas analyzer for NO was 1 ppmv and ±5%, respectively.

2.3. Simultaneous removal of SO$_2$ and NO by urea and Fe$^{II}$EDTA

At first, the pressure of the mixed gas cylinder was set to 0.2 M Pa. According to the pressure and volume of the mixed gas cylinder, the volume of the desired NO and SO$_2$ concentrations was obtained. NO and SO$_2$ were poured successively into the mixed gas cylinder by adjusting the corresponding rotameter. The desired O$_2$ concentration was obtained by mixing air and N$_2$. The absorbing liquid was prepared by adding urea to Fe$^{II}$EDTA solution. The pH of the mixed solution was adjusted by NaOH solution and H$_2$SO$_4$. The experiment was started after N$_2$ flowed through the reactor to clean up the remaining air. Absorption was carried out for 80 min, and data were measured every 10 min. The absorbing-liquid volume was ~300 mL, the gas flow rate was 0.8 L min$^{-1}$, and the flow rate of the absorption liquid was 45 mL min$^{-1}$. The effect of mixed absorbent constitutions, temperature, pH, O$_2$ concentration of simulated flue gas, as well as SO$_2$ and NO concentrations on simultaneous desulfurization and denitrification were noted.

2.4. Analytical methods

The concentrations of NO and SO$_2$ were analyzed by a flue gas analyzer (KM950). NO and SO$_2$ removal efficiency were defined as

$$\eta = \frac{C_{in} - C_{out}}{C_{in}} \quad (1)$$

where $\eta$ is NO or SO$_2$ removal efficiency given as %; $C_{in}$ is the inlet concentration of NO or SO$_2$ in ppmv; $C_{out}$ is the outlet concentration of NO or SO$_2$ in ppmv.

$$\bar{\eta} = \frac{1}{T} \int_0^T \eta(t) \, dt \quad (2)$$

where $\bar{\eta}$ is the average removal efficiency of NO or SO$_2$, $T$ is the total absorption time (min), and $\eta(t)$ is the removal efficiency of NO or SO$_2$ as a function of time.

Fig. 1 Experimental apparatus for NO and SO$_2$ removal (schematic): (1) NO cylinder, (2) N$_2$ cylinder, (3) SO$_2$ cylinder, (4) air compressor, (5) mixed cylinder, (6) gas rotameter, (7) piezometer, (8) packed tower, (9) liquid storage tank, (10) pump, (11) liquid rotameter.
3. Results and discussion

3.1. Effect of mixed absorbent constitutions on simultaneous removal

Fig. 2 shows the effect of urea concentration on simultaneous removal. As the urea concentration increased from 0 wt% to 10 wt%, the SO2 removal efficiency increased from 94.5% to 99.6%. These experimental results were similar to those in the literature, and SO2 removal efficiency using urea solution was almost the same as that of other technologies, all being up to almost 100%. Hence, urea was a good absorbent for SO2. Nevertheless, NO removal efficiency was <40% in all tests, suggesting that NO removal could not proceed well without the help of a complexing agent. This is because the solubility of SO2 in water is 700-times more than that of NO (one volume of water dissolves about 39 volumes of SO2 or ~0.05 volume of NO at 20 °C).

NO could be captured readily by FeIIEDTA in solution to form stable ferrous-nitrosyl complexes FeIIEDTA–NO (eqn (3) and (4)).

\[
\text{NO(g)} \rightarrow \text{NO(aq)} \quad (3)
\]

\[
\text{NO(aq)} + \text{FeIIEDTA}^{2-} \rightarrow \text{FeIIEDTA} – \text{NO}^{2-} \quad (4)
\]

Previously, we noted the synergistic effect of (NH2)2CO and FeIIEDTA on NO removal. Urea can react with FeIIEDTA–NO to produce FeIIEDTA for continued complexing of NO, but can also restrain oxidation of FeIIEDTA to FeIIIEDTA. Hence, a combination of FeIIEDTA and urea is suitable and helpful for the removal of NO and SO2.

Before seeking an optimal concentration ratio of FeIIEDTA to urea, reagent prices were considered to ensure that industrial scale up is prohibitively expensive. According to the data from the Alibaba Internet website, the prices of industrial FeIIEDTA was ~$1891/ton ($658.5/kmol), which is approximately five-times higher than that of industrial urea ($366/ton). Then, experiments to determine the optimal concentrations of FeIIEDTA to urea were carried out (Fig. 3).

The prime costs of the five groups from Fig. 3 were identical, and the optimal concentration ratio of FeIIEDTA to urea could be reflected clearly from simultaneous removal efficiencies. As shown in Fig. 3, with the change in the mass ratio, desulfurization efficiency was almost constant at 100%, whereas denitrification efficiency increased significantly. When the concentration ratio of FeIIEDTA to urea increased from 0 mol L⁻¹:10 wt% to 0.014 mol L⁻¹: 5 wt%, the average denitrification efficiency increased from 39.4% to 85.5%, because the FeIIEDTA concentration increased. In addition, the viscosity of the absorption liquid increased because the urea concentration was too high, so the liquid diffusivity of NO decreased. As the ratio increased further, NO removal efficiency decreased because a too-low urea concentration can result in with high concentrations of FeIIEDTA–NO and FeIIIEDTA accumulating in the complex absorbent. Thus, the optimal ratio of FeIIEDTA to urea was 0.014 mol L⁻¹: 5 wt%.

3.2. Effect of temperature on simultaneous removal

3.2.1. Effect of mixed absorbent constitutions on simultaneous removal

The temperature of the absorption liquid plays a significant part in the transfer of NO or SO2 from the gas phase to the liquid phase, and is governed by the diffusion, dissolution and reaction characteristics of the species in solution. The dependence of simultaneous removal on temperature was evaluated. As shown in Fig. 4a, the absorption temperature had a slight effect on desulfurization but a significant effect on NO removal. The denitrification efficiency increased slightly when the reaction temperature was 25–30 °C but, when the reaction temperature increased from 30 °C to 50 °C, there was a decrease in average NO removal efficiency from 86.2% to 58.1%. The effect of temperature on NO removal is highly complicated. On the one
3.3 Effect of the pH on simultaneous removal

pH is a vital parameter on SO2 and NO removal using a FeIIEDTA and urea mixed solution. To investigate the effect of the initial pH of the mixed solution on simultaneous absorption, experiments were carried out by changing the pH of the absorption liquid from 5.0 to 11.0. Hence, the effect of the pH of mixed absorbents on simultaneous removal was investigated. As shown in Fig. 5a, an alkaline atmosphere was favorable for desulfurization and the average SO2 removal efficiency increased from 95.5% to 100%. Similarly, NO removal efficiency increased from 66.3% to 88.0% with increasing pH from 5.0 to 9.0, which could be because a low pH can promote further oxidation of FeIIEDTA:

$$4\text{FeIIEDTA}^{2-} + \text{O}_2 + 4\text{H}^+ \rightarrow 4\text{FeIIIEDTA}^- + 2\text{H}_2\text{O}$$  (5)

Hand, a high temperature can accelerate the movement of molecules and increase the chemical rate. As the temperature increases, these factors result in an increase in NO removal efficiency. On the other hand, a lower temperature is favorable for the formation of iron(II) nitroxyl species. Simultaneously, as the temperature increases, the solubility of NO and O2 in the aqueous solution decreased and the consequent decomposition of FeIIEDTA begins to dominate the process. In addition, a higher temperature accelerates the change of FeIIEDTA to FeIIIEDTA. As the temperature increases, these factors can cause a decrease in NO removal efficiency. Hence, the slight decrease and then rapid decrease in NO removal efficiency with an increase in temperature was a result of the interaction of many positive and negative aspects. In summary, 30 °C was the best absorption temperature. However, the temperature of flue gases emitted from industrial boilers is typically ~100 °C. Hence, before simultaneous desulfurization and denitrification by mixed absorbent FeIIEDTA/urea in industrial applications, flue gases at high temperatures can be treated by different cooling technologies such as water cooling, vaporization cooling, and spray cooling.

A sharp decrease in average NO removal efficiency from 87.4% to 60.0% was observed when the pH was between 10.0 and 11.0 because Fe2+ hydrolyzes readily and forms a precipitate in strong alkaline solution, which greatly affects NO absorption. Thus, the optimal pH of the mixed absorbent was determined to be 9.0–10.0.

3.4 Effect of O2 concentration on simultaneous removal

Industrial flue gases in China, in general, contain 3–11% (v/v) oxygen, which has a considerable effect on the oxidation of FeIIEDTA. The O2 concentration in flue gas was varied from 3% to 11% (v/v) and its effect on simultaneous absorption illustrated in Fig. 6a. SO2 removal efficiency was almost unchanged at ~99%, and the average NO removal efficiency decreased from 91.9% to 75.6% with an increase in O2 concentration from 3% to 11% (v/v). Low NO removal efficiency under high O2 concentration was due to the oxidation of FeIIEDTA to FeIIIEDTA.

3.5 Effect of SO2 and NO concentrations on simultaneous removal

In practice, the components as well as the concentrations of flue gas are varied with boiler loads and operational conditions. Thus, simultaneous-removal experiments with various concentrations of SO2 and NO were carried out. As shown in Fig. 7a, the effect of SO2 on desulfurization was dependent on its concentration. When the SO2 concentration increased from 500 ppm to 3000 ppm, the average desulfurization efficiency increased from 94.3% to 99.5% and, then, as the SO2 concentration increased further to 4000 ppm, desulfurization efficiency began to decrease. The reason for the increase in SO2 removal efficiency was that the increase in SO2 concentration enhanced mass transfer and then promoted the gas–liquid mass transfer rate, so efficiency was increased. When the SO2 concentration was excessive, despite mass transfer being promoted further, the
reaction rate became the controlling step, so SO₂ removal resulting from SO₂ accumulation decreased.

Fig. 7b reveals that NO removal efficiencies decreased with operation time, and when the SO₂ concentration increased from 500 ppm to 3000 ppm, the average efficiency of NO removal increased from 80.3% to 85.5%. Then, as the SO₂ concentration increased further to 4000 ppm, the average denitrification efficiency decreased to 82.6%. The reasons for the increase in NO removal were as follows:

$$\text{2Fe}^{2+}\text{EDTA} + \text{NO}_2^- + \text{SO}_3^{2-} \rightarrow \text{2Fe}^{2+}\text{EDTA}^{2-} + \text{N}_2\text{O}_2^{2-} + \text{SO}_4^{2-} \quad (6)$$

$$\text{2Fe}^{3+}\text{EDTA}^- + 2\text{SO}_3^{2-} \rightarrow \text{2Fe}^{2+}\text{EDTA}^{2-} + \text{S}_2\text{O}_6^{2-} \quad (7)$$

$$\text{4Fe}^{2+}\text{EDTA} - \text{NO}_2^- + 4\text{HSO}_3^- \rightarrow 4\text{Fe}^{2+}\text{EDTA}^{2-} + 2\text{H}^+ + \text{H}_2\text{N}_2\text{O}_2 \quad (8)$$

$$\text{2Fe}^{3+}\text{EDTA}^- + 2\text{HSO}_3^- \rightarrow \text{2Fe}^{2+}\text{EDTA}^{2-} + \text{S}_2\text{O}_6^{2-} + 2\text{H}^+ \quad (9)$$

Sulfite and hydrosulfite ions can restore the active component \(\text{Fe}^{3+}\text{EDTA}\) from \(\text{Fe}^{3+}\text{EDTA-NO}\), and \(\text{Fe}^{3+}\text{EDTA}\) accumulates in the mixed absorption liquid. Thus, denitrification efficiency was increased with an appropriate increase in SO₂ concentration. However, if the SO₂ concentration was >4000 ppm, a small amount of SO₄ would compete with NO for the limited complexant (FeIIEDTA) in the gas phase. Moreover, absorption competition between SO₂ and NO also occurred in urea solution, resulting in an increase in mass transfer resistance between NO and aqueous phases, from which the denitrification efficiency began to decrease.

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The effect of the NO concentration on simultaneous removal is shown in Fig. 8a. An increase in the NO concentration from 300 ppm to 500 ppm could increase SO₂ removal efficiencies because NO₂ (the oxidation product of NO) could accelerate the oxidation of sulfite to sulfate (eqn (10) and (11)).

$$2\text{NO} + \text{O}_2 + \text{HSO}_3^- + \text{H}_2\text{O} \rightarrow 3\text{H}^+ + 2\text{NO}_2^- + \text{SO}_4^{2-} \quad (10)$$
2NO + O₂ + SO₃⁻⁻ + H₂O → 2H⁺ + 2NO₂⁻ + SO₄²⁻ \quad (11)

A further increase in the NO concentration causes severe competition between SO₂ and NO for the absorbent, resulting in a decrease in SO₂ removal. The increase of the NO concentration would cause a linear decrease in denitrification efficiency, which could be interpreted as a decrease in the molar ratio of oxidant to NO.

3.6 Validation experiments

Comprehensive consideration of the process parameters and experimental data led to optimal experimental conditions for simultaneous removal using a mixed absorbent were determined: 0.014 mol L⁻¹ Fe⁺⁶EDTA + 5 wt% urea, SO₂ and NO initial concentration of ~3000 and ~300 ppm, respectively, a temperature of 30 °C, pH of 9.0, and O₂ concentration of 3%. In this case, SO₂ removal efficiency was 100% and NO removal efficiency could exceed 91% within 80 min under optimal conditions with 300 mL of absorbing liquid volume, gas flow rate of 0.8 L min⁻¹, and flow rate of the absorption liquid of 45 mL min⁻¹.

3.7 Reaction mechanism

The mechanism involved in simultaneous desulfurization and denitrification using Fe⁺⁶EDTA and urea was speculated according to our previous work and related literature.²³,²⁴,³¹ The desulfurization mechanism was relatively simple because SO₂ is highly soluble in water. If the reactions take place in the liquid phase, SO₂ can react with the ammonium carbamate generated by urea hydrolysis. The possible forms of sulfur species are SO₃⁻, SO₂⁻H₂O, HSO₃⁻, SO₃²⁻, and SO₄²⁻, of which HSO₃⁻ and SO₄²⁻ are the main forms of S⁴⁺ species and S⁶⁺ species, respectively. In the present study, the reaction between SO₃²⁻ and urea-hydrolysis products could promote SO₂ absorption. Hence, the gas–liquid reaction between SO₂ and (NH₂)₂CO solution was completely gas-film controlled. The related reactions were:
The denitrification mechanism was much more complicated. NO having empty anti-bonding π orbitals can form π bonds by overlapping the d orbit of the central ion Fe\(^{2+}\).\(^{21}\) Thus, NO could be combined readily with Fe\(^{II}\)EDTA (eqn (3) and (4)). Meanwhile, in industrial applications, because of O\(_2\) in the flue gas, Fe\(^{II}\)EDTA is oxidized readily to Fe\(^{III}\)EDTA, and activity is lost (eqn (5)). Hence, during absorption, Fe\(^{II}\)EDTA–NO and Fe\(^{III}\)EDTA are the major byproducts. Urea can solve these problems to some degree (eqn (17)). HSO\(_3^-\) and SO\(_3^{2-}\) from the absorption liquid after SO\(_2\) removal can also restore Fe\(^{II}\)EDTA from Fe\(^{III}\)EDTA–NO and Fe\(^{III}\)EDTA (eqn (6)–(9)).

\[
\begin{align*}
\text{SO}_2 + \text{H}_2\text{O} & \rightarrow \text{SO}_2\cdot\text{H}_2\text{O} \rightarrow \text{HSO}_3^- + \text{H}^+ \tag{12} \\
\text{HSO}_3^- & \rightarrow \text{SO}_3^{2-} + \text{H}^+ \tag{13} \\
\text{SO}_3^{2-} + \frac{1}{2}\text{O}_2 & \rightarrow \text{SO}_4^{2-} + \text{H}^+ \tag{14} \\
2(\text{NH}_2\text{CO} + \text{H}_2\text{O}) & \rightarrow \text{NH}_2\text{COONH}_4 \tag{15} \\
\text{SO}_3^{2-} + 2\text{H}^+ + \text{NH}_2\text{COONH}_4 + \text{O}_2 & \rightarrow (\text{NH}_4)_2\text{SO}_4 + \text{CO}_2 \tag{16} \\
\text{NO} + \frac{1}{2}\text{O}_2 & \rightarrow \text{NO}_2 \tag{18} \\
2\text{NO}_2 & \rightarrow \text{N}_2\text{O}_4 \tag{19} \\
2\text{NO}_2 + \text{H}_2\text{O} & \rightarrow \text{HNO}_2 + \text{HNO}_3 \tag{20} \\
\text{N}_2\text{O}_4 + \text{H}_2\text{O} & \rightarrow \text{HNO}_2 + \text{HNO}_3 \tag{21} \\
2\text{HNO}_2 + (\text{NH}_2\text{CO}) & \rightarrow 2\text{N}_2 + \text{CO}_2 + 3\text{H}_2\text{O} \tag{22} \\
2\text{HNO}_2 + \text{NH}_2\text{COONH}_4 & \rightarrow 2\text{N}_2 + \text{CO}_2 + 4\text{H}_2\text{O} \tag{23} \\
6\text{HNO}_3 + 5(\text{NH}_2\text{CO}) & \rightarrow 8\text{N}_2 + 5\text{CO}_2 + 13\text{H}_2\text{O} \tag{24} \\
6\text{HNO}_3 + 5\text{NH}_2\text{COONH}_4 & \rightarrow 8\text{N}_2 + 5\text{CO}_2 + 18\text{H}_2\text{O} \tag{25} \\
6\text{Fe}^{II}\text{EDTA} - \text{NO}^{2-} + 2(\text{NH}_3)_2\text{CO} & \rightarrow 6\text{Fe}^{II}\text{EDTA}^{2-} + 5\text{N}_2 + 2\text{CO}_2 + 4\text{H}_2\text{O} \tag{17}
\end{align*}
\]

Besides being absorbed by Fe\(^{II}\)EDTA, some NO is oxidized to NO\(_2\) and N\(_2\)O\(_4\), which are dissolved readily in solution, producing nitrous acid and nitric acid, respectively. Subsequently, HNO\(_2\) and HNO\(_3\) can react with urea to form N\(_2\), which can also react with ammonium carbamate to form N\(_2\).

4. Conclusion

A mixed absorbent of Fe\(^{III}\)EDTA/urea was first employed for simultaneous removal of SO\(_2\) and NO, and the best mass concentration ratio was 0.014 mol L\(^{-1}\) : 5 wt%. Desulfurization was affected slightly by different factors, and the SO\(_2\) efficiency was 95–99% in all tests, whereas denitrification was affected significantly by various conditions. NO removal efficiency declined with increasing oxygen concentration as well as the increase in NO concentration. However, with increasing...
temperature, pH, or SO2 concentration, NO removal efficiency initially increased up to the maximum value and then decreased. Under optimal conditions, SO2 removal efficiency was 100% and NO removal efficiency could exceed 91% within 80 min. These research findings suggest that the Fe^{II}EDTA and (NH2)2CO solution could be used as an efficient and low-cost absorbent for simultaneous desulfurization and denitrification.

Conflicts of interest
There are no conflicts to declare.

References
1 L. Zheng, G. Liu and C.-L. Chou, Sci. Total Environ., 2007, 384, 374–383.
2 A. Poullikkas, Energy. Technol. Policy., 2015, vol. 2, pp. 92–103.
3 Y. Jin, M. C. Veiga and C. Kennes, J. Chem. Technol. Biotechnol., 2005, 80, 483–494.
4 S. Andreoli, F. A. Deorsola, C. Galletti and R. Pirone, Chem. Eng. J., 2015, 278, 174–182.
5 L. C. Lau, K. T. Lee and A. R. Mohamed, Fuel, 2011, 90, 1811–1817.
6 Z.-s. Wei, H.-j. Niu and Y.-f. Ji, Fuel Process. Technol., 2009, 90, 324–329.
7 Y. Han, J. Zhang and Y. Zhao, Energy, 2016, 103, 725–734.
8 C.-j. Li, R. Zhao, M.-q. Peng, L. I. U. Hao, Y. U. Gang and X. I. A. Dong-sheng, J. Fuel Chem. Technol., 2015, 43, 1516–1522.
9 L. Huang and Y. Dang, Chin. J. Chem. Eng., 2011, 19, 518–522.
10 H. Chu, T.-W. Chien and S. Li, Sci. Total Environ., 2001, 275, 127–135.
11 Y. Zhao, T.-x. Guo, Z.-y. Chen and Y.-r. Du, Chem. Eng. J., 2010, 160, 42–47.
12 C. V. Raghunath and M. K. Mondal, Chem. Eng. J., 2017, 314, 537–547.
13 J. Zhang, R. Zhang, X. Chen, M. Tong, W. Kang, S. Guo, Y. Zhou and J. Lu, Ind. Eng. Chem. Res., 2014, 53, 6450–6456.
14 C. Sun, N. Zhao, H. Wang and Z. Wu, Energy Fuels, 2015, 29, 3276–3283.
15 Q. Wu, C. Sun, H. Wang, T. Wang, Y. Wang and Z. Wu, Chem. Eng. J., 2018, 341, 157–163.
16 Y. Zhao, R.-L. Hao, Q. Guo and Y.-N. Feng, Fuel Process. Technol., 2015, 137, 8–15.
17 Y. Liu, J. Zhang, C. Sheng, Y. Zhang and L. Zhao, Chem. Eng. J., 2010, 162, 1006–1011.
18 Y. G. Adewuyi, N. Y. Sakyi and M. Arif Khan, Chemosphere, 2018, 193, 1216–1225.
19 Y. Liu, Z. Liu, Y. Wang, Y. Yin, J. Pan, J. Zhang and Q. Wang, J. Hazard. Mater., 2018, 342, 326–334.
20 P. Fang, C. Cen, Z. Tang, P. Zhong, D. Chen and Z. Chen, Chem. Eng. J., 2011, 168, 52–59.
21 Y.-P. Mao, W. Bi, X.-L. Long, W.-D. Xiao, W. Li and W.-k. Yuan, Sep. Purif. Technol., 2008, 62, 183–191.
22 E. Sada, H. Kumazawa and H. Machida, Ind. Eng. Chem. Res., 1987, 26, 1468–1472.
23 F. He, X. Deng and M. Chen, Chemosphere, 2017, 168, 623–629.
24 P. Fang, C. Cen, Z. Tang, P. Zhong, D. Chen and Z. Chen, Chem. Eng. J., 2011, 168, 52–59.
25 X. Wang, Y. Zhang, X. Dong, M. Chen, Z. Shi and J. Zhou, Energy Fuels, 2013, 27, 6024–6030.
26 J. Wei, Y. Luo, P. Yu, B. Cai and H. Tan, J. Ind. Eng. Chem., 2009, 15, 16–22.
27 P. Gans, J. Chem. Soc. A, 1967, 943–946.
28 L. Wang, W. Zhao and Z. Wu, Chem. Eng. J., 2007, 132, 227–232.
29 B. Yan, J. Yang, M. Guo, G. Chen, Z. Li and S. Ma, J. Ind. Eng. Chem., 2014, 20, 2528–2534.
30 W. Li, L. Zhang, N. Liu, Y. Shi, Y. Xia, J. Zhao and M. Li, Energy Fuels, 2014, 28, 4725–4730.
31 E. Sada, H. Kumazawa and Y. Takada, Ind. Eng. Chem. Fundam., 1984, 23, 60–64.
32 R. Hao, S. Yang, B. Yuan and Y. Zhao, Fuel Process. Technol., 2017, 159, 145–152.
33 J. Enemark and R. Feltham, Coord. Chem. Rev., 1974, 13, 339–406.