Simulation of nitrogen transformation in pressurized oxy-fuel combustion of pulverized coal

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Chemical kinetic modeling was applied to simulate N transformation in the pressurized oxy-fuel combustion process of pulverized coal. Modeling accuracy was validated by experimental data at different operation pressures. The key reaction paths from fuel-N to different N products were revealed by analyzing the rate of production. NO formation was synergistically affected by six elementary reactions, in which NCO and other intermediate species were involved. The reactions among N, NH, NH₂, and NO were the key paths of N₂ formation. After pressurizing the combustion system, NO and N₂ contents decreased and increased, respectively. High operation pressure inhibited the diffusion of NO from the internal to the external part of char. This condition prolonged the residence time of NO inside the char, triggered a typical heterogeneous reaction between gaseous NO and unburned char, and reduced the conversion from fuel-N to NO. Moreover, modeling was performed to predict NOₓ emission in pressurized oxy-fuel combustion as a function of various operating parameters, including temperature and excess air and recycling ratios. This study may provide guidance for reducing NOₓ emissions and improving combustion efficiency in oxy-fuel combustion, and it can serve as a reference for industrial applications that involve pulverized coal combustion.

1 Introduction

Greenhouse gases emitted from coal-fired power plants, industrial boilers, and other energy utilization equipment are preferentially enrolled in the blacklist for emission reduction due to the pressure posed by increasing environmental awareness.¹−³ Researchers worldwide are developing new combustion techniques for energy conservation and emission reduction. Oxy-fuel combustion is one of the most promising approaches due to its low CO₂, NOₓ, and SO₂ emissions and high combustion efficiency of pulverized coal.⁴−⁶

In oxy-fuel combustion, a blend of nearly pure oxygen is used as an oxidant instead of air. This usage results in a flue gas that consists mainly of CO₂ and H₂O and allows the occurrence of a simple gas purification process before CO₂ capture. Oxy-fuel combustion always involves flue gas recirculation (FGR) in practice. The formation mechanism of air pollutants is remarkably different from that of air-firing combustion due to the influence of a low flame temperature and CO₂-rich atmosphere. The transformation of N from fuel-N to NOₓ has elicited widespread attention.⁷−⁹ Several studies have reported that NOₓ emissions in oxy-fuel combustion with the FGR process are inferior to those under air-firing conditions because the combustion process has lower temperatures and higher char concentrations than air-fired combustion.¹⁰−¹¹

The formation mechanism of NOₓ during oxy-fuel combustion has been investigated mainly under atmospheric pressure conditions, and only a few studies have reported N transformation under pressurized oxy-fuel combustion. Lasek et al.¹⁴ set up a pressurized oxy-fuel combustion process and observed reduced NO emission in air-fired and oxy-fuel pressurized combustion. The authors concluded that the positive influence of pressure on NO reduction is the creation of preferential conditions for the NO–char reaction. Lei et al.¹⁵ investigated the release behavior of NOₓ during pressurized oxy-fuel combustion of Datong bituminous coal and discovered that NOₓ emission declines at 3 MPa due to the accelerated decomposition rate of NOₓ at high combustion temperatures. However, the characterizations applied in these studies limited the demonstrations, and the elementary reactions involving the combustion process were unclear. Therefore, the N transformation mechanism during pressurized oxy-fuel combustion still requires clarification. Moreover, modeling studies on NOₓ emissions in oxy-fuel combustion focused on gaseous fuels.¹²,¹³ Modeling of pulverized coal relied on computational fluid dynamics codes with simplified schemes of nitrogen transformation and exhibited limited accuracy when compared with experimental results.¹⁷

In the present study, a chemical engineering model was applied to simulate NOₓ formation and consumption during pressurized oxy-fuel combustion. This work aims to predict NOₓ...
emission as a function of various operating parameters, including operation pressure, temperature, and excess air and recycling ratios. Rate of production (ROP) and sensitivity analyses were performed to determine the key reaction paths from fuel-N to different N products. The analysis results were compared with and validated against experimental data. This study may provide guidance for reducing pollutant emissions and improving oxy-fuel combustion efficiency, and it can serve as a reference for industrial applications that involve pulverized coal combustion.

2 Experimental and modeling

2.1 Modeling

Simulations of the oxy-fuel combustion of pulverized coal were performed using the plug flow reactor model of CHEMKIN-PRO. Mixing, pyrolysis, combustion, and gas–solid reactions were represented by expressions that can be handled in CHEMKIN. The physical model was based on conservation equations for mass, energy, and species in a specified system. The influence of thermal diffusion was considered in the simulations, and the multicomponent method was used to calculate the transport parameters. The initial conditions, including initial temperature, operation pressure, flow rate, and reactant fractions, were inputted for computation. The coal sample used in this study was a type of bituminous coal, and the pyrolysis process was conducted in oxy-fuel (O₂/CO₂) atmosphere at a high heating rate. According to previous research, pyrolysis of bituminous coal at a high heating rate produces HCN, especially in oxy-fuel atmosphere. Therefore, HCN was considered as the main pyrolysis product of fuel-N in this study.

The thermodynamic and kinetic parameters modified by Hashemi et al. were applied in this work. The detailed elementary reactions used were proposed by the Lawrence Livermore National Laboratory, and these reactions have been widely tested and verified by several studies on pulverized coal combustion.

2.2 Materials

The Shenhua coal utilized in this study was purchased from the largest coal producer (Shenhua Group Corporation Ltd.) in China. The proximate and ultimate analyses that were conducted are presented in Table 1. Coal samples were pulverized into 60–100 meshes by using a coal mill before the combustion experiments. Then, the coal powders were dried at 105 °C for 12 h to remove moisture.

The major pyrolysis products of pulverized coal consisted of conventional light volatiles (H₂O, CO₂, CH₄, CO, etc.) and char. HCN was the main pyrolysis product of fuel-N. In this modeling, we assumed that char is largely converted into soot and C₂H₂ according to the C/H ratio, as reported in similar studies. The molar volatile composition of the present coal was estimated in accordance with Mackrory correlations, which are illustrated in Table 2.

2.3 Experimental

The experimental apparatus and procedure are illustrated in Fig. 1. The pressurized oxy-fuel combustion process was implemented in a fixed-bed reactor. The reactor tube was made of nickel-base alloy steel to ensure device stability under high operation pressures and temperatures. The lengths of the furnace and tube were 660 and 1000 mm, respectively. The inner and external diameters of the tube were 50 and 85 mm, respectively. Furthermore, the furnace consisted of three heating zones to stabilize the temperature of the combustion zone. The combustion zone was in the middle of the furnace, and the two ends of the furnace worked as heat preservation areas. The temperature of the furnace bed was monitored automatically by thermocouples. The monitoring point was set inside the steel tube where the coal samples were placed to ensure accurate temperature control.

Initially, CO₂ and O₂ with 99.99% purity were used as the carrier gas and oxidant in oxy-fuel combustion, respectively. The O₂/CO₂ volume ratio was set to 21 : 79 in the carrier gas, and the total feed flow rate was maintained at 1000 mlN min⁻¹ with a mass flow controller. The operation pressure of the system was monitored by two pressure sensors located at the inlet and outlet of the steel tube. During each test, the reactor was heated to the corresponding temperature and fed with O₂/CO₂ for 30 min. Thereafter, 1.0 g of coal sample was uniformly spread in a quartz boat (length of 200 mm and width of 35 mm) and rapidly pushed into the center of the steel tube. The combustion experiments were conducted at a temperature of 900 °C for 10 min. A Fourier transform infrared (FTIR) gas analyzer, Gasmet DX4000 (Finland), was utilized to measure NO and CO₂ concentrations. The FTIR instrument was calibrated by selecting the appropriate positions of characteristic peaks before the experiments to ensure highly accurate species concentrations. During combustion, the gaseous species were collected into a gas bag for subsequent analysis.

Table 1: Proximate and ultimate analyses of the present coal

| Proximate analysis (%) | Ultimate analysis (%) |
|------------------------|-----------------------|
| M₂₀ | A₂₀ | V₂₀ | FC₂₀ | Q₂₀ | C₂₀ | H₂₀ | N₂₀ | S₂₀ | O₂₀ |
| 1.66 | 10.18 | 30.83 | 57.33 | 283.27 | 73.57 | 4.23 | 0.9 | 0.53 | 8.93 |
combustion, the gas bag was transferred and connected to the FTIR instrument. The pipeline that connected the gas bag and FTIR instrument was maintained at 180 °C to prevent the gaseous species from condensing. The measurement frequency of the FTIR instrument was set to 3 s. The species concentration of each sampling point was measured by the FTIR instrument three times and averaged to ensure high accuracy. Additionally, a gas chromatograph (Agilent 7890A, USA) equipped with a thermal conductivity detector (TCD) was utilized to measure the N₂ concentration. Each experiment was conducted three times and averaged to ensure that the relative standard deviations were below 3%.

3 Results and discussion

3.1 Model validation

Modeling accuracy was validated by comparing the molar fraction of the major combustion products at different operation pressures in the numerical simulation and experimental results. The experiments and modeling were conducted at a temperature of 900 °C. As shown in Fig. 2, the NO and N₂ contents in the experimental data and modeling predictions decreased and increased, respectively, as the operation pressure increased. However, the modeling results of NO content were slightly underestimated compared with those of the experimental data. This result can be ascribed to the temperature variation during the combustion process. Although the temperatures in the modeling and experiment part were set to be similar (900 °C), the actual temperature in the experimental part might have slightly exceeded 900 °C due to the exothermic reaction of combustion. Therefore, the oxidation of fuel-N into NO accelerated, resulting in high values of NO content in the experimental results. Additionally, the concentration of CO₂ emitted from the flue gas was nearly constant in the entire pressure range, implying that the majority of volatiles were completely combusted in pressurized oxy-fuel combustion. As demonstrated in Fig. 2, the simulation results are in good agreement with the experimental data. Therefore, the reaction mechanism and calculation method applied in this work are reasonable and convincing.

3.2 Nitrogen products and free radicals

Fuel-N is the only source of final nitrogen products in oxy-fuel combustion. In the current modeling of pulverized coal, HCN was added as fuel-N. The modeling results showed that NO, NO₂, N₂O, and N₂ were the primary N products. The molar fractions of the four products varied along the homothermal length of the reactor, as illustrated in Fig. 3. The coal sample was rapidly pushed into the center of the steel tube after the reactor was preheated, as indicated in Section 2.3. Thus, the start point of the distance in Fig. 3 is the center of the steel tube.

As displayed in Fig. 3a, the molar fraction of NO increased sharply to 1.16 × 10⁻³ at the initial stage of oxy-fuel combustion then remained almost unchanged until the end of the combustion. The molar fraction of N₂ increased to 1.08 × 10⁻⁴, indicating that N₂ was the second most abundant product. Two other conventional N compounds, namely, NO₂ and N₂O, were observed among the products. The molar fractions of NO₂ and...
N\textsubscript{2}O were 3.43 \times 10^{-6} and 1.18 \times 10^{-5}, respectively. After the initial stage of oxy-fuel combustion, the molar fractions of the four N products remained constant, indicating that the combustion system was stable.

However, the yields of the four N products differed remarkably after introducing operation pressure (1.1 MPa) into this combustion system. As indicated in Fig. 3b, the molar fraction of NO decreased from 1.16 \times 10^{-3} to 7.78 \times 10^{-4} during atmospheric and pressurized oxy-fuel combustion. Meanwhile, the molar fraction of N\textsubscript{2} increased gradually and stabilized at 2.93 \times 10^{-5}. Notably, the yield of NO declined by 30%, whereas that of N\textsubscript{2} increased by nearly three times compared with atmospheric pressure. The NO\textsubscript{2} and N\textsubscript{2}O contents were 1.78 \times 10^{-5} and 1.33 \times 10^{-5}, respectively, suggesting that the production of NO\textsubscript{2} was promoted and slightly exceeded that of N\textsubscript{2}O.

The molar fraction changes of H, O, and OH radicals in the combustion process are also shown in Fig. 4. The maximum molar fractions of the three radicals (Fig. 4) are much higher than those of the N products in atmospheric and pressurized oxy-fuel combustion (Fig. 3). This result implies that H, O, and OH radicals are vital to the ignition process at the initial stage of combustion. Studies\textsuperscript{28,29} have proven that in the ignition process of fuel combustion, the chain branch reaction R1: H + O\textsubscript{2} \rightarrow O + OH (Table 3) plays the leading role in promoting combustion. Additionally, all three radicals' molar fractions in the initial stage of pressurized oxy-fuel combustion were superior to those in atmospheric oxy-fuel combustion. This result indicates that the reaction rate is much faster in pressurized oxy-fuel combustion than in atmospheric oxy-fuel combustion. This result can be explained by classic chemical reaction collision theory.\textsuperscript{30} The quantity of activated molecules in the reaction system increases as the pressure increases, thereby increasing the number of effective collisions and improving the rate of chain branch reaction R1. Furthermore, R1 consumes one H radical and generates two radicals (O and OH) simultaneously.

### Table 3  Major elementary reactions involving nitrogen transformation

| No.\textsuperscript{a} | Chemical reaction equation | No.\textsuperscript{a} | Chemical reaction equation |
|------------------------|---------------------------|------------------------|---------------------------|
| R1            | H + O\textsubscript{2} ⇌ O + OH | R442      | NH + NO ⇌ NNH + O |
| R351          | HNO + H ⇌ NO + H\textsubscript{2} | R514      | HCN + M ⇌ HNC + M |
| R361          | NO + HO\textsubscript{2} ⇌ NO\textsubscript{2} + OH | R516      | HCN + O ⇌ NCO + H |
| R391          | N\textsubscript{2}O + H ⇌ N\textsubscript{2} + OH | R521      | HCN + OH ⇌ HNCO + H |
| R417          | NH\textsubscript{2} + NO ⇌ N\textsubscript{2} + H\textsubscript{2}O | R526      | HNC + O ⇌ NH + CO |
| R418          | NH\textsubscript{2} + NO ⇌ NNH + OH | R527      | HNC + OH ⇌ HNCO + H |
| R422          | NH + H ⇌ N + H\textsubscript{2} | R537      | CN + NO\textsubscript{2} ⇌ N\textsubscript{2} + CO\textsubscript{2} |
| R423          | NH + O ⇌ NO + H | R544      | HNCO + H ⇌ NH\textsubscript{2} + CO |
| R430          | NH + NO ⇌ N\textsubscript{2}O + H | R546      | HNCO + O ⇌ NCO + OH |
| R432          | NH + NO ⇌ N\textsubscript{2} + OH | R547      | HNCO + O ⇌ NH + CO\textsubscript{2} |
| R434          | NH + NO\textsubscript{2} ⇌ N\textsubscript{2}O + OH | R548      | HNCO + O ⇌ HNO + CO |
| R435          | N + OH ⇌ NO + H\textsubscript{2} | R571      | NCO + O ⇌ NO + CO |
| R436          | N + O\textsubscript{2} ⇌ NO + O | R697      | C + NO ⇌ CN + O |
| R437          | N + NO ⇌ N\textsubscript{2} + O | R698      | C + NO ⇌ CO + N |
| R438          | NNH ⇌ N\textsubscript{2} + H | \ |

\textsuperscript{a} The sequence numbers (no.) of elementary reactions are derived from the reaction mechanism database.\textsuperscript{17}
Thus, the increase in pressure results in an exponential increase in free radical concentration and promotes the establishment of a radical pool. This radical pool then triggers numerous elementary reactions, sustains the entire combustion process, involves N conversion, and consumes fuel to the end.

3.3 ROP analysis of NO

ROP analysis of the modeling calculations was performed to study the transformation of N and obtain an improved understanding of N transformation in pressurized oxy-fuel combustion. The major elementary reactions involving N transformation are presented in Table 3. Positive and negative ROP values indicate the rates of formation and consumption of NO, respectively.

In atmospheric oxy-fuel combustion, the six major elementary reactions involving NO production are R423, R435, R436, R571, R697, and R698, as shown in Fig. 5a. The first four elementary reactions (R423, R435, R436, and R571) participated in the entire reaction process and displayed the highest reaction rate (with a maximum reaction rate of $10^{-3}$ mol cm$^{-3}$ s$^{-1}$) of NO production, implying that NCO might be the prior intermediate product with regard to NO formation. R423, R435, and R436 also exhibited remarkable acceleration in NO formation, verifying that NH and N acted as the two other chief intermediate products. On the contrary, a fraction of generated NO was consumed mainly through R697 and R698. This result can be ascribed to the reduction atmosphere at the initial stage of combustion. The generated NO diffused and transferred to the vicinity of unburned char then triggered a typical heterogeneous reaction between gaseous NO and unburned char. After adsorption on the surface of char, the NO molecules combined with the active species on the carbon surface and were reduced into low-valence species thereafter. However, the maximum reaction rates of R697 and R698 were relatively lower than those of R423, R435, R436, and R571. This result implies that considerable quantities of NO existed in the flue gas, which is in accordance with the results in Fig. 3a.

After pressurizing the combustion process, the six major elementary reactions that involved NO production and consumption were unchanged, as illustrated in Fig. 5b. Notably, the reaction rates of these six elementary reactions were improved greatly. With regard to the main NO production reaction (R571), the maximum reaction rate increased to $3.82 \times 10^{-2}$ mol cm$^{-3}$ s$^{-1}$. The maximum reaction rates of the main NO consumption reactions (R697 and R698) were also significantly enhanced. Furthermore, the maximum reaction rates of R697 and R698 increased to the same magnitude order as R571, which is slightly different from the situation in atmospheric combustion. Thus, we infer that the NO content in the flue gas decreased distinctly, whereas the low-valence N species ($N_2$) increased, as verified by the results in Fig. 3b.

The rapid increase in reaction rate under the pressurized condition can be explained by chemical reaction rate (1), Arrhenius (2), and Clapeyron (3) equations.

\[
    r = k[HCN]^{M}[O_2]^N, \quad (1)
\]

\[
    k = A \exp \left(\frac{-E_a}{RT}\right), \quad (2)
\]

\[
    PV = nRT, \quad (3)
\]

where $r$ is the chemical reaction rate in mol L$^{-1}$ s$^{-1}$, $[x]$ is the concentration of reactant $x$ in mol L$^{-1}$, $M$ and $N$ are the orders of the reaction, $k$ is the reaction rate constant in s$^{-1}$, $A$ is the pre-exponential factor in s$^{-1}$, $E_a$ is the activation energy in J mol$^{-1}$, $R$ is the ideal gas constant (8.314 J K$^{-1}$ mol$^{-1}$), $T$ is temperature in K, $P$ is pressure in Pa, $V$ is the gas volume in L, and $n$ is the molar amount in mol.

We assumed that the gaseous components in this reactor are ideal. In this modeling, the reaction rate constant ($k$) was unchanged because the temperature ($T$) was constant according to eqn (2). As the pressure ($P$) increased, the molar amount ($n$) and/or concentration $[x]$ of the reactants increased because the gas volume ($V$) was constant according to eqn (3). Thereafter, the reaction rate ($r$) augments exponentially with the increase of...
[x] according to eqn (1). This conclusion has been proven by many studies and is in accordance with our modeling results.13–35

In summary, after pressurizing the combustion system, the promotion of NO consumption rate exceeded that of NO production rate. The decrease in emitted NO content in pressurized oxy-fuel combustion synergistically resulted from the six major elementary reactions, namely, R423, R435, R436, R571, R697, and R698.

3.4 Key reaction paths

The key reaction paths involving N transformation in oxy-fuel combustion were described in accordance with the ROP and sensitivity analyses in this modeling, as demonstrated in Fig. 6. The reaction paths were complicated processes that involved multiple elementary reactions toward multiple potential directions. Nevertheless, fuel-N (HCN) generally reacted initially with H, O, and OH radicals to generate NCO, HNCO, and HNC. These three species are pivotal products to start HCN conversion, not only to NO, but also to N2.

During the entire oxidation process from HCN to NO, the O radical acted as the main oxidant. Initially, HCN was converted to NCO and HNCO through R516 and R521, followed by the oxidation of HNCO to NCO via R546. Plenty of NCO was formed and accumulated by these three reaction paths. According to Glarborg,36 the subsequent reaction of NCO largely determines the fate of N atoms in the NO formation process, which is in good agreement with our modeling results. As the major precursor of NO, the NCO generated from R516, R521, and R546 was further oxidized by the O radical through R571. However, a small fraction of the generated HNCO from R521 was converted to NH, N, and HNO through R547, R422, and R548, respectively. The generated NH, N, and HNO were further oxidized to NO through R423, R435, and R351. In this manner, the migration and conversion of N atoms from HCN to NO was completed. Thereafter, the conversion from NO to NO2 and N2O was triggered through R361, R430, and R434. According to these reaction paths, the free radical pool played a crucial role throughout the entire oxidation process involving NOx formation.

Fig. 6 also illustrates the conversion from HCN to N2. Initially, HCN was converted to HNCO and HNC through R521 and R514. The generated HNC was partially converted into HNCO through R527. Then, conversion from HNCO and HNC to N, NH, and H2 was realized through R547, R544, R526, and R422. Notably, the subsequent reactions between N/NH/NH2 produced from the abovementioned reactions and the NO generated in the aforementioned paths were the key step of N2 formation in the process.37,38 As shown in Fig. 6, N, NH, and H2 reacted with NO through R437, R432, and R417. Another intermediate product, NNH, which participates in the N transformation of most oxy-fuel combustion,39 was also evident. NNH was produced through R442 and R418. As an unstable species, NNH dissociated automatically into N2 through R442 and R418. However, a small fraction of NO2 and N2O generated in the aforementioned paths was reduced into N2 through R537 and R391. Herein, the migration and conversion of N atoms from HCN to N2 were demonstrated.

In summary, H, O, and OH radicals were vital to sustaining and accelerating the entire reaction process. NCO acted as the major precursor to form NO. NO could be further converted into NO2 and N2O. Furthermore, the reactions among N, NH, NH2, and NO were the key steps of N2 formation. NO2 and N2O could also be converted into N2 through particular paths. Generally, the migration and conversion of N atoms in oxy-fuel combustion are complicated with regard to the formation and consumption of N species.

3.5 Effect of operation pressure

Fig. 7 demonstrates the trend of N products as a function of operation pressure in pressurized oxy-fuel combustion. The NO molar fraction displayed a significant reduction when the operation pressure increased from 0.1 MPa to 2.1 MPa. On the contrary, the molar fraction of N2 showed an opposite trend as the pressure increased. In this modeling, oxy-fuel combustion was conducted under O2/CO2 atmosphere with a temperature of less than 1000 °C. Thus, almost all of the NO can be considered to have been produced from fuel-N.

The increasing operation pressure exerted two effects on NO emission. First, the concentration [x] of reactants increased with the increase in total pressure, leading to the acceleration of the oxidation reaction rate from fuel-N to NO, as discussed in Section 3.3. Therefore, the molar fraction of NO in the flue gas increased accordingly. Second, the partial pressure and concentration of CO2 also increased with the increase in total pressure, resulting in the decrease of NO content in the flue gas. In this manner, the migration and conversion of N atoms from HCN to NO was completed.
The high CO partial pressure resulted in an elevated concentration of CO through the following reactions.

\[ \text{CO}_2 + \text{C} \rightarrow 2\text{CO} \]  
\[ 2\text{CO}_2 \rightarrow 2\text{CO} + \text{O}_2 \]  

Thereafter, the generated CO promoted the reduction of NO through the following reaction.

\[ 2\text{NO} + 2\text{CO} \rightarrow \text{N}_2 + 2\text{CO}_2 \]  

The high pressure inhibited the diffusion of NO from the internal to the external part of char, thereby prolonging the residence time of NO inside the char. As discussed in Section 3.3, the high NO concentration triggered a typical heterogeneous reduction with unburned char through the following reaction.

\[ 2\text{NO} + 2\text{C} \rightarrow \text{N}_2 + 2\text{CO} \]  

According to these demonstrations, the increase in total pressure was beneficial to NO reduction and N\textsubscript{2} formation. The molar fraction of NO\textsubscript{2} also increased slightly with the increase in pressure. This condition can be ascribed to the promotion of the following reaction.

\[ 2\text{NO} + \text{O}_2 \rightarrow 2\text{NO}_2 \]  

However, this reaction was limited because the pressure and residence time were insufficient to realize complete oxidation to NO\textsubscript{2}.  

In summary, operation pressure exerted a joint effect on NO formation and reduction. The molar fraction of NO decreased with the increase in total pressure, whereas that of N\textsubscript{2} increased.

### 3.6 Effect of temperature

Temperature is an important factor that influences the conversion of combustion products, especially N oxides. Improving the temperature of a combustion system exerts complex effects on N evolution. The oxidation of N into NO is accelerated, and the reduction of N from high valence to low valence is facilitated. Thus, the NO molar fraction in flue gas is an integrated result of multiple reactions.

As shown in Fig. 8, the final molar fraction of NO emitted from the reactor increased with the increase in temperature, whereas N\textsubscript{2} and NO\textsubscript{2} displayed the opposite trend. Generally, not all of N atoms can be deeply oxidized to high-valence N (NO and NO\textsubscript{2}) at a low-temperature range due to the relatively low reaction rate and reactivity. Moreover, reductive N species, such as HCN, NH\textsubscript{3}, and HNCO, that are not timely oxidized certainly react with NO to generate a considerable amount of N\textsubscript{2} through a similar path of the SNCR process. Therefore, the oxidation products consist of medium-valence N (N\textsubscript{2}) and high-valence N (NO and NO\textsubscript{2}) at a low-temperature range.

After the temperature range moved to a high value, the reaction rate and reactivity of the oxidant were activated, and many nitrogen species were oxidized to a high valence. Meanwhile, a large amount of NO was retained because the reductive N species were rapidly consumed. Therefore, the molar fraction of NO increased with increasing temperature, whereas a reverse trend was obtained for N\textsubscript{2} conversion, which is consistent with the results of previous studies. In a similar combustion, the amount of fuel-N converted to NO\textsubscript{2}, N\textsubscript{2}O, and NH\textsubscript{3} is relatively small, with only NO being formed in large amounts, which is in accordance with this study. In summary, the formation of NO is thermodynamically favored as the temperature increases.

### 3.7 Effect of excess air ratio

Excess air ratio is another crucial factor that affects pulverized coal combustion and N conversion. The conversion ratio from fuel-N to NO (denoted as CR\textsubscript{NO}) was calculated with the following equation to evaluate the conversion degree of fuel-N.

\[ \text{CR}_{\text{NO}} = \frac{\epsilon_{\text{NO}}}{\epsilon_{\text{fuel-N}}} \times 100\% \]  

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where CRNO is the conversion ratio from fuel-N to NO (%), cNO is the concentration of gaseous NO, and cfuel-N is the concentration of N in the estimated volatile composition illustrated in Table 2. The molar fractions of main N products and CRNO as a function of the excess air ratio in pressurized oxy-fuel combustion are presented in Fig. 9.

As shown in the figure, CRNO increased gradually with the increase in the excess air ratio. When the excess air ratio exceeded 1.0, the concentration of inlet oxygen was higher than the required molar ratio for complete combustion of N and C species. In this case, the reductive species, such as HCN, NH3, and HNCO, were rapidly consumed and adequately oxidized into high-valence N (NO and NO2). Meanwhile, the incomplete combustion carbon species, such as CO and unburned char, were reduced due to the high oxygen content, thus decreasing the chance of CO/NO and char/NO reduction. In short, the reduction reactions in the combustion system were greatly suppressed, leading to the high CRNO in the excess air ratio range over 1.0.

However, the N conversion ratio to NO decreased substantially as the excess air ratio declined. A 17% decrease in CRNO was observed as the excess air ratio decreased from 1.0 to 0.8, but the N2 molar fraction increased by 7%. The formation of NO was inhibited under the low excess air ratio condition, and the generation of N2 was promoted. In other words, appropriately reducing the excess air ratio is conducive to lowering the NO emission.

### 3.8 Effect of recycling ratio

In practice, oxy-fuel combustion always involves the FGR process. As discussed in Section 3.1, the flue gas of oxy-fuel combustion mainly consists of CO2 (approximately 80%, as shown in Fig. 2) and H2O (nearly 20%, not shown here). Therefore, a part of the flue gas is recycled into the intake system to replace proportional inlet CO2 after simple drying and filtration. Accordingly, the cost of CO2 capture for oxy-fuel combustion is reduced, and the net power generation efficiency of the system is improved. A series of modeling with different recycling ratios ranging from 0 to 0.8 was conducted to obtain insights into the actual N evolution process during pressurized oxy-fuel combustion.

Fig. 10 presents the effect of recycling ratio on the main N products. The NO molar fraction in the flue gas generally increased as the recycling ratio increased. In the experiments with FGR, a high concentration of NO was recycled along with CO2 into the inlet mixture, thereby improving the NO molar fraction in the final flue gas. Thus, the higher the recycling ratio is, the more concentrated the emitted NO is.

Additionally, the conversion ratio from fuel-N to NO (CRNO) decreased with the increase in recycling ratio. This result can be ascribed to the reaction between reductive N species (HCN, NH3, and HNCO) and recycled NO. Large amounts of reductive N species were rapidly consumed by recycled NO as the recycling ratio increased. These N species are vital precursors to form NO, as discussed in Section 3.4. Thus, the oxidation paths to form NO were inhibited, leading to a continuous reduction of CRNO as the recycling ratio increased. Moreover, the reactions between reductive N species and recycled NO produced abundant N2, resulting in an evident increase in N2 molar fraction in the flue gas, as illustrated in Fig. 10. These results are in accordance with those reported by Yang in a similar investigation on the coal reburning process.29 The authors concluded that an increase in NO concentration in inlet gas might lead to a decrease in HCN and an increase in N2 in the exhausted gas. In this study, fuel-derived HCN accounted for 10% of the NO reduction in the flue gas at the initial stage of combustion. However, the detailed mechanism still needs to be explored.

Notably, the conversion ratio from fuel-N to NO (CRNO) displayed an opposite trend of final NO concentration. However, no contradiction was noted. A part of the flue gas was separated to be recycled and serve as inlet CO2 after involving FGR, and the rest was emitted to the vent. The FGR process decreased the flow rate of the final emitted flue gas. With regard to combustion with FGR, although the NO concentration in the flue gas was high, the final NO emission load was lower than the non-recycle runs.
4 Conclusions

In this study, chemical kinetic modeling was applied to simulate N transformation in the pressurized oxy-fuel combustion process of pulverized coal. Modeling accuracy was validated by experimental data at different operation pressures, and an ROP analysis was performed to determine the key reaction paths from fuel-N to different nitrogen products. NO emission was synergistically affected by six elementary reactions, in which NCO and other intermediate species were involved. The reactions among N, NH, NH2, and NO were the key paths of N2 formation. NO and N2 contents decreased and coal combustion.

This condition prolonged the residence time of NO inside the combustor system, respectively. Accordingly, the high operation pressure inhibited the diffusion of NO from the internal to the external part of char. This condition prolonged the residence time of NO inside the char, triggered a typical heterogeneous reaction between gaseous NO and unburned char, and reduced the conversion from fuel-N to NO. Moreover, modeling was performed to predict NOx emission in pressurized oxy-fuel combustion as a function of various operating parameters, including temperature and excess air and recycling ratio. This study may provide guidance for reducing pollutant emissions and improving combustion efficiency in oxy-fuel combustion, and it can serve as a reference for industrial applications involving pressurized coal combustion.

Conflicts of interest

The authors declare no conflicts of interest.

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References

1 M. B. Toftegaard, J. Brix, P. A. Jensen, P. Glarborg and A. D. Jensen, Prog. Energy Combust. Sci., 2010, 36, 581–625.
2 S. D. Kenarsari, D. Yang, G. Jiang, S. Zhang, J. Wang, A. G. Russell, Q. Wei and M. Fan, RSC Adv., 2013, 3, 22739–22773.
3 Y. Lu, Y. Wang, Y. Zhao, Z. Wei, Y. Li, W. Hao and Y. Zhang, RSC Adv., 2017, 7, 51036–51045.
4 Q. Shen, Y. Zheng, C. Luo, N. Ding, C.-G. Zheng and M. Thern, RSC Adv., 2015, 5, 39785–39790.
5 M. C. Stern and T. A. Hatton, RSC Adv., 2014, 4, 5906–5914.
6 J. Sun, Z. Yi, X. Zhao, Y. Zhou and C. Gao, RSC Adv., 2017, 7, 14678–14687.
7 Z. Sun, S. Su, J. Xu, K. Xu, S. Hu, Y. Wang, L. Jiang, N. Si, Y. Zhou and S. S. A. Syed-Hassan, Energy Fuels, 2017, 31, 8392–8399.
8 L. Álvarez, J. Riaza, M. V. Gil, C. Pevida, J. J. Pis and F. Rubiera, Greenhouse Gases: Sci. Technol., 2011, 1, 180–190.
9 M. C. Stewart, R. T. Symonds, V. Manovic, A. Macchi and E. J. Anthony, Fuel, 2012, 92, 107–115.
10 H. I. Matheka, B. Oboirien, A. Engelbrecht, B. North and K. Premlal, Energy Fuels, 2016, 30, 6756–6763.
11 L. Duan, C. Zhao, W. Zhou, C. Qu and X. Chen, Fuel Process. Technol., 2011, 92, 379–384.
12 K. Andersson, F. Normann, F. Johnsson and B. Leckner, Ind. Eng. Chem. Res., 2008, 47, 1835–1845.
13 Y. Tan, E. Croiset, M. A. Douglas and K. V. Thambimuthu, Fuel, 2006, 85, 507–512.
14 J. A. Lasek, M. Janusz, J. Zuwala, K. Glöd and A. Iluk, Energy, 2013, 62, 105–112.
15 M. Lei, X. Huang, C. Wang, W. Yan and S. Wang, J. Therm. Anal. Calorim., 2016, 126, 1067–1075.
16 T. Mendiara and P. Glarborg, Energy Fuels, 2009, 23, 3565–3572.
17 H. Hashemi, S. Hansen, M. B. Toftegaard, K. H. Pedersen, A. D. Jensen, K. Dam-Johansen and P. Glarborg, Energy Fuels, 2011, 25, 4280–4289.
18 P. Glarborg, A. Jensen and J. E. Johnsson, Prog. Energy Combust. Sci., 2003, 29, 89–113.
19 J. Leppälähti, Fuel, 1995, 74, 1363–1368.
20 Y. Duan, L. Duan, E. J. Anthony and C. Zhao, Fuel, 2017, 189, 98–106.
21 M. Østberg, P. Glarborg, A. Jensen, J. E. Johnsson, L. S. Pedersen and K. Dam-Johansen, Symposium (International) on Combustion, 1998, 27, 3027–3035.
22 M. U. Alzueta, R. Bilbao, A. Millera, P. Glarborg, M. Østberg and K. Dam-Johansen, Energy Fuels, 1998, 12, 329–338.
23 P. Glarborg, P. G. Kristensen, K. Dam-Johansen, M. Alzueta, A. Millera and R. Bilboa, Energy Fuels, 2000, 14, 828–838.
24 E. Vilas, U. Skifter, A. D. Jensen, C. Lópex, J. Maier and P. Glarborg, Energy Fuels, 2004, 18, 1442–1450.
25 S. Niksa and S. Cho, Energy Fuels, 1996, 10, 463–473.
26 J. C. Chen and S. Niksa, Energy Fuels, 1992, 6, 254–264.
27 A. J. Mackrory, Ph.D. thesis, Brigham Young University, 2008.
28 J. Huang and W. Bushe, Combust. Flame, 2006, 144, 74–88.
29 P. Zhang, J. Ran, C. Qin, X. Du, J. Niu and L. Yang, Energy Fuels, 2018, 32, 3900–3907.
30 M. S. Child, Molecular collision theory, Courier Corporation, 1996.
31 W. Shangguan, Y. Teraoka and S. Kagawa, Appl. Catal., B, 1997, 12, 237–247.
32 M. Higashi, S. Uchida, N. Suzuki and K. Fujii, IEEE Trans. Plasma Sci., 1992, 20, 1–12.
33 T. F. Wall, G.-s. Liu, H.-w. Wu, D. G. Roberts, K. E. Benfell, S. Gupta, J. A. Lucas and D. J. Harris, Prog. Energy Combust. Sci., 2002, 28, 405–433.
34 D. Roberts and D. Harris, Energy Fuels, 2000, 14, 483–489.
35 R.-L. Du, K. Wu, D.-A. Xu, C.-Y. Chao, L. Zhang and X.-D. Du, Fuel Process. Technol., 2016, 148, 295–301.
36 P. Glarborg and P. Marshall, Energy Fuels, 2017, 31, 2156–2163.
37 Y. He, X. Zheng, J. Luo, H. Zheng, C. Zou, G. Luo and C. Zheng, Energy Fuels, 2017, 31, 10093–10100.
38 H. Xiao, Z. Wang, A. Valera-Medina and P. J. Bowen, J. Therm. Sci., 2018, 27, 270–276.
39 S. J. Klippenstein, L. B. Harding, P. Glarborg and J. A. Miller, *Combust. Flame*, 2011, **158**, 774–789.
40 J. A. Lasek, K. Głod, M. Janusz, K. Kazalski and J. Zuwala, *Energy Fuels*, 2012, **26**, 6492–6500.
41 K. Okazaki and T. Ando, *Energy*, 1997, **22**, 207–215.
42 Y. Hu, N. Kobayashi and M. Hasatani, *Energy Convers. Manage.*, 2003, **44**, 2331–2340.
43 F. Normann, K. Andersson, B. Leckner and F. Johnsson, *Prog. Energy Combust. Sci.*, 2009, **35**, 385–397.
44 Y. Hu, N. Kobayashi and M. Hasatani, *Fuel*, 2001, **80**, 1851–1855.
45 F. Winter, C. Wartha and H. Hofbauer, *Bioresour. Technol.*, 1999, **70**, 39–49.
46 L. Duan, Y. Duan, C. Zhao and E. J. Anthony, *Fuel*, 2015, **150**, 8–13.
47 F. Kapteijn, J. Rodriguez-Mirasol and J. A. Moulijn, *Appl. Catal., B*, 1996, **9**, 25–64.
48 A. Hayhurst and A. Lawrence, *Combust. Flame*, 1996, **105**, 341–357.
49 C. J. Tullin, S. Goel, A. Morihara, A. F. Sarofim and J. M. Beer, *Energy Fuels*, 1993, **7**, 796–802.
50 Y. B. Yang, E. Hampartsoumian and B. M. Gibbs, *Symposium (International) on Combustion*, 1998, **27**, 3009–3017.