Nitrogen Migration during Pyrolysis of Raw and Acid Leached Maize Straw

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1. Introduction

Biomass energy is a form of renewable energy derived directly or indirectly from plant photosynthesis. Solid biomass fuel has been gaining popularity as a way to reduce reliance on fossil fuels while also dealing with climate change [1]. However, biomass fuel combustion is reported to result in high NOₓ (NO, NO₂, N₂O, etc.) emissions [2]. The NOₓ produced from fuel combustion can be divided into thermal-NOₓ, fuel-NOₓ, and prompt-NOₓ according to their origins. The formation mechanism of prompt-NOₓ is...
complicated and its amount is relatively small. Thermal-NOx is primarily formed by the reaction of nitrogen and oxygen at high temperatures (>1300 °C) [3]. While the furnace temperature of household biomass stove is mostly lower than 700 °C (refer to previous studies [4,5]), fuel-NOx accounted for most of all NOx [6,7]. Fuel-N can be converted into various nitrogen-containing functional groups during the pyrolysis process, and these compounds can react with oxygen to produce various NOx species [8]. For NOx regulation, it is critical to have a thorough understanding of the transformation and migration behavior of fuel-N during biomass pyrolysis.

Both biomass characteristics (species, contents of nitrogen, volatile matter and ash content, etc.) and pyrolysis operation conditions (temperature, heating rate, air supply, etc.) influence the pyrolysis performance of the biomass. Researchers investigated the migration course of nitrogen elements in raw biomass, proteins, and N-containing compounds (formed by protein mixed with hemicellulose, cellulose, and lignin) under various operating conditions [9–11]. With the increase of temperature, NOx precursors are primarily formed by the following three ways: the pyrolysis of unstable fuel-N, secondary decomposition of char, and tar products [12]. The main NOx gaseous precursors are NH3, HCN, and HNCO, with HNCO accounting for little and being easily transferred to the other two [13,14]. The dominant N-species are determined by the biomass type and origin. In some biomass, such as sewage sludge, nitrogen mainly exists in the form of heterocyclic-N, HCN can be found to be the main NOx precursor during pyrolysis [8]. However, the majority of nitrogen in plant residues is found in proteins and free amino acids, with only a small amount in the form of nucleic acid, chlorophyll, enzymes, vitamins, alkaloids, and inorganic nitrogen [15], NH3 will become the dominant nitrogen-containing precursor [16].

Agricultural and woody residues are the two main materials of solid biofuels. According to the literature, agricultural residues (straw fuels) may have 5–20 times higher ash content than those of woody biomass, which could contribute to inefficient combustion [17]. Besides, alkali and alkaline earth metallic species (AAEMs) in ash (K, Ca, Na, Mg, etc.) are thought to influence the devolatilization and combustion process [18,19]. Acid-leaching treatment can reduce the content of ash and AAEMs, effectively improve the combustion performance and reduce ash melting, slagging, and corrosion [20–23], but the impact of AAEMs on the NOx precursor generations of biomass pyrolysis remains unclear. Deep research on the effects of de-ashing and demineralization by acid-leaching on nitrogen transfer and conversion would help the clean utilization of straw biomass fuel.

Therefore, nitrogen migration behaviors during pyrolysis of raw and acid-leached maize straw were investigated in this report. To reduce the ash and AAEMs contents in maize straw, pre-treatment of acid-leaching with CH3COOH was introduced. The thermal gravimetric analysis (TGA) and pyrolysis experiments at various temperatures were performed to evaluate the decomposition characteristics of the two biomass samples. X-ray photoelectron spectroscopy (XPS) was used to investigate the distribution and migration of nitrogen functional groups in biomass and corresponding biochar, and the effect of acid-leaching on NOx precursors generation was further analyzed.

2. Materials and Methods

2.1. Materials and Pretreatment Process

In this study, raw maize straw (RMS) was obtained from Shangzhuang experimental station of China Agricultural University, Beijing, China. The samples were crushed and sieved through a 100 mesh sieve, then dried in an oven at 105 °C for 12 h.

Various organic and inorganic acids have been used for acid treatment in previous researches. Strong acids, such as sulfuric acid and nitric acid were reported to enhance dehydration reactions in biomass. Hydrochloric acid is also a commonly leaching solution, but chloride ions will affect the subsequent HCN concentration measurement. To effectively remove AAEMs, while reducing the damage to the biomass structure and not introducing other ions [24], CH3COOH solution was adopted in this study. The solution was prepared by adding 30 mL of glacial acetic acid (1.05 g/mL, 99.5%) into 970 mL of distilled water
and mix thoroughly, which was converted to a mass concentration of 3.13%. Acid-leached maize straw (AMS) was pretreated by soaking 10 g of raw maize straw samples in 200 mL acid solution for 2 h at room temperature while stirring. After removing the acid liquor, the acid-leached biomass was washed with deionized water to neutral pH and then dried at 105 °C for 12 h.

Ultimate analysis of the samples was performed by Vario EL cube (Elementar, Langenselbold, Germany), and oxygen content was detected by difference (Table 1). Proximate analysis of the biomass fuels was carried following the existing Chinese standards [25,26]. The ash content for the biomass sample was reduced from 12.69 wt% to 8.67 wt% after acid treatment. The main residual material in the ash should be silica, since more than half of the ash of the maize straw consists of SiO2 [27], and it is hardly removed by acid leaching. The contents of cellulose, hemicellulose, and lignin in biomass were determined using the methods mentioned by Guo et al. [28]. To better understand the effect of the pre-treatment, inductively coupled plasma with optical emission spectroscopy (ICP-OES) (ICPOES730, Agilent, Santa Clara, CA, USA) was performed on RMS and AMS to provide selected elemental compositions of the fuel ashes (Table 1). It should be noted that the acid leaching significantly decreased the content of K, Na, and Mg, which were abundantly found in biomass ash.

Table 1. Basic characteristics of the two biomass samples.

| Characteristic | Biomass Sample | Removal Ratio |
|---------------|----------------|---------------|
|               | RMS            | AMS           |
| Ultimate analysis wt%, dry basis | C 41.83 | 44.69 |
|               | H 5.08 | 6.18 |
|               | S 0.58 | 0.68 |
|               | N 1.39 | 1.13 |
|               | O 1 51.13 | 47.33 |
| Proximate analysis wt%, dry basis | Volatile matter 68.68 | 75.70 |
|               | Fixed carbon 16.94 | 13.01 |
|               | Ash 12.69 | 8.67 |
| Composition% | Hemicellulose 31.11 | 36.49 |
|               | Cellulose 29.03 | 34.47 |
|               | Lignin 3.72 | 3.62 |
| Inorganic elements mg/kg | K 21,245.4 | 92.1 | 99.57 |
|               | Na 1474.9 | 61.5 | 95.83 |
|               | Mg 2171.9 | 91.7 | 95.78 |
|               | P 2254.1 | 364.4 | 83.84 |
|               | Fe 379.1 | 203.3 | 46.38 |
|               | Ca 3979.7 | 2444.4 | 38.58 |

1 By difference. 2 By ICP-OES analysis.

2.2. Pyrolysis System and Experimental Procedure

The slow pyrolysis system is shown in Figure 1. A gas supply, gas flowmeter, tubular resistance furnace (OTF-1200X, Hefei, China), condensation unit, filter unit, and N-containing gas collection unit were all included in the system. At the start of each experiment, a 5.00 ± 0.02 g sample was mounted in the center of the furnace tube (constant temperature zone length: 150 mm) in a quartz boat (length: 80 mm). The system was purged for 40 min (400 mL/min) with high purity Ar (99.999%). Subsequently, the flow rate was adjusted to 200 mL/min, and the reactor was heated to the desired temperature (350 °C, 450 °C, 550 °C, or 650 °C) with a heating rate of 10 °C/min and then kept steady for 30 min to ensure the completion of pyrolysis. The outlet of the tubular quartz reactor was connected with the condensing device placed in the ice-water mixture. After that, the pyrolysis gas was filtered by absorbent cotton and entered the nitrogen-containing gas collection bottle containing 200 mL absorbent solution. The two kinds of NOx precursors, NH3 and
HCN, were absorbed by 5 g/L HBO$_3$ and 8 g/L NaOH solutions, respectively [29], and converted to the corresponding ions (NH$_3$→NH$_4^+$, HCN→CN$^-$). To prevent interference, NH$_3$ and HCN collection experiments were carried out separately. Each experiment was duplicated under the same condition and the average value was adopted.

2.3. TG Analysis

The TG analysis of raw and acid-leached maize straw was performed by a thermal gravimetric analyzer (TGA 2, Mettler-Toledo, Greifensee, Switzerland) in the Ar atmosphere (50 mL/min). In each experiment, 5.00±0.05 mg of the straw sample was heated from 30°C to 900°C at a constant heating rate of 10°C/min. The experiments were performed twice to test the repeatability.

2.4. XPS Analysis

The nitrogen functional groups in the biomass and corresponding biochar were characterized using XPS analysis. The sample was uniformly glued to the conductive adhesive tape and degassed for 12 h. An X-ray photoelectron spectrometer (ESCALAB 250Xi, Thermo Fisher Scientific, Waltham, MA, USA) with a monochromatic Al K$_{\alpha}$ X-ray source (150 W, hv = 1486.6 eV) was used, the following were the operating conditions: a spot size of 650 µm, a voltage of 14.8 kV, a current of 1.6 A, a constant analyzer pass energy mode (100 eV for survey scans and 20 eV for narrow scans) and a pressure less than 10$^{-10}$ mbar. The XPS results were evaluated by Thermo Avantage software. The Shirley-type background was subtracted and all spectra were calibrated based on the principal C1s peak at 285.0 eV [30–33]. Thereafter, the N1s signal was curve-resolved using peaks with a 70% Gaussian and 30% Lorentzian line shape and the FWHM of 1.4 eV [32]. Peaks at 398.8±0.2 eV, 399.9±0.2 eV, 400.4±0.2 eV, 401.4±0.2 eV and 402–405 eV corresponding to the energy positions were considered as N functional groups of pyridine-N (N-6), amine-N/amide-N/protein-N (N-A), pyrrole-N (N-5), quaternary-N/inorganic-N (N-Q/N-IN) and N-oxide (N-X), respectively [12,34–36].

2.5. Scanning Electron Microscopy (SEM)

The sample powers were fixed on the conductive adhesive tape and coated with a gold film. The morphology images of two biomass and corresponding char products were performed using an SEM device (SU3500, HITACHI, Tokyo, Japan, an accelerating voltage of 10 kV).

2.6. Calculations

The difference between the initial and final weight of containers was used to calculate char and tar production. According to the forms of pyrolysis product, nitrogen
compounds were classified into char-N, gas-N (NH$_3$-N, HCN-N) and rest-N (tar-N and other N-containing products). The following equations were used to measure the yields of char-N and nitrogen functional groups (group-N) in char products [36]:

$$Y_{\text{char-N}} = \frac{m_{\text{char}} \times w_{\text{char-N}}}{m_{\text{biomass}} \times w_{\text{N}}} \times 100\%$$ (1)

$$Y_{\text{group-N}} = \frac{\sum A_{\text{group-N}}}{\sum A_{\text{group-N}}} \times Y_{\text{char-N}}$$ (2)

where $Y_{\text{char-N}}$ and $Y_{\text{group-N}}$ mean the yields of char-N and group-N in char products, respectively (wt%), $m_{\text{biomass}}$ and $m_{\text{char}}$ mean the mass of biomass and corresponded char products, respectively (g), $w_{\text{N}}$ and $w_{\text{char-N}}$ mean the N contents from biomass and char products, respectively (wt%), and $A_{\text{group-N}}$ means the peak area of each nitrogen functional group in char according to the XPS analysis.

Based on relevant Chinese standards, the absorption solutions of NO$_x$ gaseous precursors were analyzed using a UV2800 ultraviolet spectrophotometer (UNICO, Princeton, NJ, USA) [37,38]. Yields of these gas-N were calculated by Equation (3):

$$Y_{\text{gas-N}} = \frac{c_n \times V}{m_{\text{biomass}} \times w_{\text{N}}} \times 100\%$$ (3)

where $Y_{\text{gas-N}}$ means the yields of NH$_3$-N or HCN-N (wt%), $c_n$ is the concentrations of NH$_3$-N or HCN-N in the absorption solutions (g/mL), and V represents the volume of absorption solution (200 mL).

3. Results and Discussion

3.1. Characteristics of Raw/Acid Leached Maize Straw

TG and derivative thermogravimetric (DTG) curves of RMA and AMS samples are shown in Figure 2. Similarly, both two biomass samples were found in three stages of mass loss. According to relevant researches, these three stages were summarized as water evaporation (below 200 °C), rapid decomposition of hemicellulose and cellulose (around 200–400 °C), and slow pyrolysis of lignin and other residual materials (above 400 °C) [39–41]. By comparison with the TG curve of RMS (Figure 2a), the second pyrolysis stage of AMS began later (at around 300 °C). This hysteresis was due to the fact that AAEMs in the ash could promote the decomposition of cellulose and hemicellulose under low temperature [42]. A shoulder peak at 305.67 °C was observed during the pyrolysis of acid-leached maize straw (Figure 2a). It is understood that the peak at around 300 °C originated from hemicellulose degradation [43]. RMS had a maximum degradation rate of 8.43%/min at 322.33 °C, which was lower than AMS’s (10.60%/min at 356.83 °C) (Figure 2b). As the temperature increased to about 350 °C, the AMS samples continued to release volatiles at a rapid rate, while the weightlessness rate of RMS slowed. These results are consistent with those reported in other studies [42,44,45]. According to the description of Zareihasanregheshlaghi et al. [23], the increased concentration of metal impurities on the surface of straw biomass ash could be found during the conversion, which could promote the formation of agglomeration and char, thereby hindering the following biomass decomposition. Also, acid-leaching treatment could increase the porosity of biomass and promote the conversion of cellulose and hemicellulose into sugars, leading to a large release of volatiles [20].
The functional group characteristics of specific elements in different samples were investigated using XPS analysis. The predominant peak was observed in both biomass samples at 399.9 ± 0.2 eV, which can be verified as N-A (Figure 2c,d) [12]. Maize can absorb nitrogen elements from soil and chemical fertilizer for growth, the fixed nitrogen in biomass mainly existing in the forms of amine-N/amide-N/protein-N [46]. Some nitrogen in biomass exists in inorganic nitrogen such as NH$_4^+$-N, while quaternary-N groups are
usually observed after heating treatment [47–49], thus the small peaks at 401.88 eV of RMS and 401.39 eV of AMS should be defined as N-IN in this study (Figure 2c,d) [36]. Moreover, the RMS spectrum contained the N-6 peak at 398.93 eV, which was missing in the AMS spectrum. These losses may be due to the high solubility of pyridine nitrogen and its reaction with acids during the pre-treatment. Moreover, a lot of floccules and small particles were observed in raw maize straw compared to acid leached straw (Figure 2e,f), which mainly belong to various salty, inorganic constituents in ash.

3.2. Distribution of Pyrolysis Products

In this study, solid products were represented by biochar, liquid products consisted of tar and condensed water, and the remaining non-condensable gases were referred to as gaseous products. According to the TG analysis reported above, the moisture in biomass was extracted before 200 °C, so the change of liquid products from 350 °C to 650 °C could be attributed to different tar productions. Under different temperatures, similar change trends in the pyrolysis product yields were obtained for the two biomass samples (Figure 3). Biochar was the predominant product during biomass samples pyrolysis at 350 °C, accounting for 48.55% for RMS and 44.81% for AMS. When the temperature increased to 450 °C, the char proportions decreased quickly for RMS (39.06%) and AMS (35.66%) due to the devolatilization. Gaseous product yields increased as the temperature increased from 350 °C to 450 °C, while liquid product yields increased from 450 °C to 550 °C and peaked at 550 °C. The highest yields of liquid products were 42.33% and 46.92% for RMS and AMS, respectively. During pyrolysis at low temperatures, the unstable chemical bonds of biomass break quickly and primarily produce non-condensable gases, while the ring cracking reactions in lignin and char were much stronger at higher temperatures and a large amount of tar are generated [50]. More surface folds and structural cracking were observed in SEM images from high-temperature pyrolysis (Figure 4). As the reaction temperature increased from 550 °C to 650 °C, the liquid product yields fell rapidly, which were attributed to the secondary decomposition of tar products. Moreover, higher solid product yields and lower liquid product yields were observed in RMS groups than that in AMS groups at the same pyrolysis temperature. According to Figure 4, ash agglomeration and sintering of particles resulted in larger ash particles coating the surface of char products from RMS. The acid-leaching pre-treatment could increase the porosity of the biomass and promoted the removal of volatiles [20].

Figure 3. Pyrolysis products from RMS (a) and AMS (b) under different pyrolysis temperatures.
Figure 4. SEM images of the char products: (a) RMS$_{350}$; (b) RMS$_{450}$; (c) RMS$_{550}$; (d) RMS$_{650}$; (e) AMS$_{350}$; (f) AMS$_{450}$; (g) AMS$_{550}$; (h) AMS$_{650}$.
3.3. Distribution of Gaseous NO\textsubscript{x} Precursors

As shown in Figure 5, NH\textsubscript{3} was the dominant NO\textsubscript{x} gaseous precursors during pyrolysis of the two straw samples. According to the literature, when the fuel-N mostly existed as aromatic ring-N, the dominant gaseous NO\textsubscript{x} precursors was HCN. When it existed in amine-N groups, NH\textsubscript{3} would become the key intermediate product. The rapid increase of NH\textsubscript{3} yield from 350 °C to 450 °C was primarily due to the decomposition of N-A, while the decomposition of char and tar could also produce NH\textsubscript{3} at the higher temperature. For HCN-N yield, it remained at a low level in the low-temperature pyrolysis for RMS and AMS, while a large increase was achieved at 650 °C. This meant that more heterocyclic-N was broken and HCN was released as the temperature rose.

![Figure 5. Yields of NH\textsubscript{3}-N (a) and HCN-N (b) from RMS and AMS under different pyrolysis temperatures.](image-url)

In the literature, it is reported that the removal of AAEM species from coal fuel before pyrolysis will reduce nitrogen conversion to NO\textsubscript{x} gaseous precursors, but the situation is more complicated for biomass fuels [51]. Different amino acid structures have different nitrogen release characteristics during the chemical chain cracking. Potassium, for example, has been found to promote the NH\textsubscript{3}-N production during aspartic acid pyrolysis, while inhibiting the HCN-N conversion from phenylalanine thermal decomposition [6]. By comparison with RMS, AMS released more NH\textsubscript{3}-N at each pyrolysis experiment, while obtained much higher HCN-N yields in high-temperature pyrolysis. The highest NH\textsubscript{3}-N and HCN-N yields were 3.17% ± 0.14% and 0.90% ± 0.07%, respectively, obtained from AMS pyrolysis at 650 °C. It’s clear that the gas-N proportion was small for the total fuel-N. In the following part, we investigated the characteristics of char-N at various temperatures to further quantitatively confirm the nitrogen conversion from the two biomass samples pyrolysis.

3.4. Nitrogen Migration in Char-N during Biomass Pyrolysis

During pyrolysis, C, H, O, N, and S elements can be removed in various compound forms. However, since decarburization and denitrification take far longer than other reactions like dehydrogenation and deoxygenation, the increased carbon and nitrogen contents could be observed in biochar compared to the biomass sample (Table 2).

![Table 2. Ultimate analysis (wt%, dry basis) of char products from raw/acid-leached maize straw.](table-url)
Table 2. Cont.

| Sample  | C     | H     | S     | O \(^1\) | N     |
|---------|-------|-------|-------|----------|-------|
| AMS\(_{350}\) | 58.56 ± 0.16 | 4.40 ± 0.03 | 0.65 ± 0.06 | 34.49 ± 0.29 | 1.91 ± 0.04 |
| AMS\(_{450}\) | 61.42 ± 0.14 | 3.87 ± 0.04 | 0.93 ± 0.04 | 31.78 ± 0.27 | 2.01 ± 0.04 |
| AMS\(_{550}\) | 63.93 ± 0.20 | 2.98 ± 0.08 | 0.64 ± 0.06 | 30.37 ± 0.38 | 2.09 ± 0.04 |
| AMS\(_{650}\) | 65.52 ± 0.20 | 2.36 ± 0.04 | 0.53 ± 0.05 | 29.83 ± 0.35 | 1.77 ± 0.06 |

\(^1\) By difference.

The XPS spectra of biochar obtained at different pyrolysis temperatures are shown in Figure 6 and yields of nitrogen functional groups in the solid phase are shown in Figure 7. The biochar obtained from low-temperature pyrolysis kept some residual N-A groups and N-IN groups, which eventually vanish at 450 °C and higher temperatures. They were considered as the main source of NH\(_3\)-N in low-temperature pyrolysis [49]. N-6 and N-5 were mainly produced from N-A through direct cyclization, dimerization, and other reactions [52, 53]. During the pyrolysis process, N-5 and N-6 dominated in nitrogen functional groups from solid products and accounted for more than 50% of biochar-N. The yields of N-6 and N-5 increased with the temperature in the range from 350 °C to 450 °C, but decreased at higher temperatures due to the secondary decomposition (Figure 7). The ring-opening reaction of N-5 was the important way of HCN formation [54]. The binding energies of N-IN and N-Q groups were very close (401.4 ± 0.2 eV). Considering the thermal stability, N-IN could easily decompose when heated, while N-Q was primarily formed by the cyclo-condensation and hydrogenation reaction of N-6 at high temperature [55]. Previous studies have reported that N-6 and N-Q conversions play an important role in NH\(_3\)-N generation at high pyrolysis temperature [36, 56]. N-X stands for various nitrogen oxide functional groups, which can be observed in char products under each temperature. Zhan et al. [29] suggested that it could be produced by the combination of N-6 and oxygen functional groups.

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Figure 6. Cont.
Figure 6. XPS analyses of the char products at various temperatures: (a) RMS$_{350}$; (b) RMS$_{450}$; (c) RMS$_{550}$; (d) RMS$_{650}$; (e) AMS$_{350}$; (f) AMS$_{450}$; (g) AMS$_{550}$; (h) AMS$_{650}$.

Figure 7. Yields of nitrogen functional groups in char products from RMS (a) and AMS (b) under different pyrolysis temperatures.

For nitrogen conversion in RMS pyrolysis, N-A yield dramatically dropped from 80.00 wt% to 20.82 wt% at 350 °C, while the corresponding gaseous-N (NH$_3$-N and HCN-N) only accounted for 1.06 wt%. It meant that in addition to the conversion of N-A to gaseous-N and N-5 and N-6 in char-N, a large part of N-A flowed into tar products. Moreover, by comparing the pyrolysis performance of RMS and AMS, delayed removals of N-IN and N-A were found in AMS pyrolysis with the same temperature. As the temperature increased from 350 °C to 450 °C, N-IN was totally converted, and N-A yield showed a decrease from 34.57 wt% to 10.94 wt%, which were responsible for the significant increase in NH$_3$-N yields in AMS pyrolysis (Figure 5a).

Based on the characteristic analysis of the pyrolysis products and related literature [12,36,57], nitrogen migration pathways during raw and acid-leached maize straw
pyrolysis were illustrated in Figure 8. Thermal decomposition of N-A and N-IN was the main source of NH₃-N at low temperature, and the conversions of N-5 (char-N) and heterocyclic-N (tar-N) contributed to HCN-N yield at high temperature.

![Figure 8](image)

**Figure 8.** Possible fuel-N migration pathways during pyrolysis of RMA and AMS.

### 3.5. Nitrogen Distribution in Pyrolysis Products of Raw and Acid-Leached Maize Straw

In general, the distribution of nitrogen in each phase product from RMS and AMS were shown in Table 3. The yield of rest-N was detected by difference, of which tar-N accounted for more than 90%. During low temperature (350 °C, 450 °C) pyrolysis processes, more than half of the biomass-N was deposited in biochar products. As pyrolysis temperature increased, it can be seen that nitrogen elements gradually flowed from solid products to gas and tar. Among the three-phase products, gas-N took a small part of the total, and the majority of nitrogen was combined in the tar products during high-temperature pyrolysis. The proportion of gaseous NOₓ precursors of sludge and some industrial waste biomass can be higher than that of straw biomass [49]. Zhan et al. [49] reported the gas-N yield (<8.5 wt%) and variation trend from straw biomass pyrolysis between 200 °C and 500 °C, which was close to the results in this study.

| Sample | Char-N | Gas-N | Tar-N and Others ¹ |
|--------|--------|-------|--------------------|
| RMS₃₅₀ | 67.80  | 1.06  | 31.14              |
| RMS₄₅₀ | 51.87  | 2.04  | 46.09              |
| RMS₅₅₀ | 45.23  | 2.28  | 52.49              |
| RMS₆₅₀ | 34.98  | 3.18  | 61.84              |
| AMS₃₅₀ | 76.39  | 1.76  | 21.85              |
| AMS₄₅₀ | 63.32  | 3.03  | 33.65              |
| AMS₅₅₀ | 61.53  | 2.96  | 35.51              |
| AMS₆₅₀ | 48.13  | 4.07  | 47.80              |

¹ By difference.

The higher tar-N yield was obtained from RMS than AMS with the same temperature, AAEMs in the ash may promote the generation of tar-N in raw straw. N-rich tar products can be used in the production of high value-added chemicals, which is meaningful for
industrial production. However, the complex nitrogen bonds in tar were labile and would cause severe NO\textsubscript{x} emissions during fuel combustion. The slow nitrogen release from a stable char structure would support nitrogen control of straw fuel combustion. Although a slight increase of gas-N yield was found in AMS, the char-N yield was much higher than that of RMS. The char-N yield (76.39 wt\%) was peaked at 350 °C from AMS pyrolysis, which was higher than that of other studies \cite{34,49,58}. These results can provide some guidance for straw fuel modification and its low NO\textsubscript{x} combustion.

4. Conclusions

The pyrolysis performance and nitrogen migration pathways of raw and acid leached straw biomass at various temperatures were investigated in this paper. The acid-leaching pretreatment effectively reduced the AAEMs in the ash of the straw. Thermogravimetric analysis showed that AMS released more volatiles than RMS during pyrolysis, but the maximum degradation rate of the former moved to a higher temperature than that of the latter. NH\textsubscript{3} dominated in gaseous NO\textsubscript{x} precursor from slow pyrolysis of two straw biomass. The conversions of unstable N-IN and N-A groups in straw contributed to the NH\textsubscript{3} generation at low pyrolysis temperatures (350 °C and 450 °C), while more than half of the HCN was formed from the secondary reactions of N-5 in char and other heterocyclic-N in tar products. Particularly, higher char-N and gas-N yields were observed from AMS than RMS with the same temperature. The highest char-N yield of 76.39 wt\% was obtained from AMS pyrolysis at 350 °C. By operating low pyrolysis temperature and acid-leaching treatment, the fuel nitrogen can be effectively stored in the stable char-N structure, rather than gaseous NO\textsubscript{x} precursors or labile tar-N, which could provide support for NO\textsubscript{x} emissions control from straw fuel combustion.

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