Observation of Replacement of Carbon in Benzene with Nitrogen in a Low-Temperature Plasma

Zhiping Zhang¹, Xiaoyun Gong², Sichun Zhang², Haijun Yang², Youmin Shi², Chengdui Yang², Xirong Zhang², Xinghuang Xiong³, Xiang Fang³ & Zheng Ouyang⁴

¹School of Chemistry and Chemical Engineering, Xi’an Shiyou University, Xi’an 710065 (China), ²Department of Chemistry, Tsinghua University, Beijing 100084 (China), ³National Institute of Metrology, Beijing 100013 (China), ⁴Weldon School of Biomedical Engineering, Purdue University, West Lafayette, Indiana 47907 (USA).

Selective activation of benzene has been mainly limited to the C-H activation. Simple replacement of one carbon in benzene with another atom remains unresolved due to the high dissociation energy. Herein, we demonstrate a direct breakage of the particularly strong C=C bond in benzene through ion-molecule reaction in a low-temperature plasma, in which one carbon atom was replaced by one atomic nitrogen with the formation of pyridine. The mechanism for the formation of pyridine from benzene has been proposed based on the extensive investigation with tandem mass spectrometry. The reaction pathway also works to other aromatics such as toluene and o-xylene. This finding provides a new avenue for selective conversion of aromatics into nitrogen-containing compounds.

Results

The experiment was carried out by introducing benzene vapor with air into a low-temperature plasma region (Figure S1). The reaction product was on-line monitored using a high resolution Orbitrap mass spectrometer. Besides the peaks at mass-over-charge ratio (m/z) 94.0651 (C₆H₅N⁺), 110.0600 (C₆H₅ON⁺) and 126.0551 (C₆H₅NO₂⁺), the most abundant ionic species observed was m/z 80.0494 (Figure 1A), which was different from the typical molecular ion of benzene (m/z 78.0469) as shown in Figure S2 in mass spectrometry (MS) analysis. In a previous study under a similar plasma condition but with benzene in helium, ion [M + 2]⁺ was observed and was...
identified as C6H8 based on deuterium isotopic labeling experiments. The chemical formula of the ion m/z 80.0494 observed in this study was assigned as C5H6N. This peak was the most abundant in the spectrum (Figure 1A), which suggests that this reaction was highly efficient in producing this unusual product.

To confirm the chemical structure of the product C5H6N (m/z 80) generated from benzene, MS/MS analysis was performed and the fragmentation pattern was compared with that from pyridine (analytical standard). As shown in Figure 1B and C, both cations C5H6N (m/z 80) generated from benzene through the plasma reaction and from pyridine standard sample by nano-electrospray ionization (nano-ESI) have identical fragmentation patterns, with fragments of m/z 28 (CNH2), 50 (C4H2), 51 (C4H3), 53 (C4H5) and 78 (C5NH4) observed at similar relative intensities. This result suggests the product C5H6N (m/z 80) probably has a structure of pyridine.

D6-benzene was also used for the plasma reaction and ions at m/z 85.0810 and 86.0871 were observed, corresponding to C5D5HN and C5D6N, respectively (Figures 2A, S5 and S6). The most abundant ion C5D5HN at m/z 85.0810 could be attributed to the transfer of a proton from water in air during the reaction. In its MS/MS spectrum, a similar dissociation pathway as those from benzene and pyridine was observed, with fragments of m/z 29 (CNHD), 30 (CND2), 51 (C4HD), 52 (C4D2), 53 (C4HD2), 54 (C4D3), 57 (C4HD4), 58 (C4D5), 81 (C5NHD3) and 82 (C5ND4) (Figures 2B and S6D), confirming the occurrence of conversion from benzene to pyridine.

Further convinced evidence on the production of pyridine from benzene has been obtained by gas chromatography/mass spectrometry (GC/MS) analyses. When pure benzene was used for the analysis, no pyridine peak was observed in the chromatogram (Figure 2C). However, the retention peak of pyridine appeared at 3.44 min for the collected product from the plasma reaction of
benzene (Figure 2D) by using the apparatus as shown in Figure S7. This is in good agreement with the retention behavior of pyridine standard (Figure S8).

In an effort of identifying the source of nitrogen atom involved in the replacement of the carbon from benzene, different gases, including air, pure N2, and the mixture of N2 and NO (99:1, v/v), were used for the plasma reaction. The results shown in Figure S9 clearly demonstrate that NO-containing gas more favors the generation of pyridine than air and pure N2 in this reaction. NO was then added into the reaction for tracing the nitrogen source via isotope ratio analysis. After a reaction period of 30 min, the liquid sample was analyzed using an Orbitrap mass spectrometer. As expected, the peak intensity of 15N-labeled cation (C5H6$^{15}$N) sharply increased, and the ratio between 15N-labeled cation (C5H6$^{15}$N) and C5H6$^{14}$N was 1.53 (Figure 3A). On the contrary, the peak intensity ratio was only 0.05 for the reaction using NO (Figure 3B). This provides a direct evidence that the pyridine product was generated from the reaction with exogenous nitrogen-containing species rather than contaminants in the original benzene solvent.

**Discussion**

Besides the dielectric barrier discharge apparatus (Figure S1), other reaction systems such as paper- or needle-based discharge (Figure S10) were also used to investigate the discharge reaction products of benzene at atmospheric pressure without any other auxiliary gases. It is interesting that the same reaction product patterns were observed (Data not shown). These results demonstrate that the production of C6H8N$^+$ under the discharge of benzene is a general case.

To give some insights into the formation of C6H8N$^+$ from benzene, efforts were carried out to its origin. As shown in Figure 1, besides the peak C6H8N$^+$ at m/z 80.0494, three other abundant peaks, assigned to C6H8O2N (m/z 126) and C6H8ON (m/z 110) through loss of one molecule H2O or CO or C group by collision induced dissociation (CID) as shown in Figure 4. For the ion m/z 94, no direct experimental results were found to be related with the production of m/z 80 (C5H6$^{15}$N) as shown in Figure 4C. Also, the component of m/z 94 varied from C6H8N to C6H6O with reaction time (Figures S12), and their possible formation mechanisms were suggested as Scheme S1. From Figure S12, it can be seen that the formation of C6H8N and C6H6O should be one pair of competing reactions. Further detailed investigation is needed to elucidate the source and conversion of C6H8N to C6H6O (m/z 94) as well as C6H8ON$^+$ (m/z 110) and C6H8O2N$^+$ (m/z 126).

According to the experimental results, a possible mechanism for pyridine formation from benzene is suggested in Figure 5a. The intermediate C6H8O2N$^+$ with m/z 126 (Figure 4A) is first formed through the interaction between benzene and NO, H2O in air or...
H$_2$ONO$^+$, probably derived from a three body dissociation of NO$^+$, H$_2$O in air and neutral molecules$^{26}$. The rapid loss of H$_2$O molecule$^{27,28}$ from C$_6$H$_4$NO$_2^-$ results in the production of C$_6$H$_4$NOH$^+$ (m/z 108)$^{29,30}$, followed by ring- opening and closing reactions with formation of C$_6$H$_5$N$^+$ (m/z 80). In addition to this proposed pathway (Figure 5a), other similar mechanism (Figure 5b) might simultaneously undergo due to the existence of the isomers$^{23,24}$. For instance, the peak at m/z 109 (Figure 4A) could be attributed to the loss of OH group from the isomeric species of C$_6$H$_5$NHOOH$^-$ (m/z 126), followed by formation of C$_5$NH$_6$$^+$. The suggested schemes agree well with the formation of pyridine from m/z 110 C$_6$H$_4$ON$^+$ (Figure 4B) as well as the correspondingly isotopically labeled products from D$_3$-benzene (Figures S13–S14).

The reaction pathway also works to other aromatics such as toluene (Figures S15–S21 and Schemes S2–S3) and o-xylene (Figure S22). The occurrence of conversion from toluene to 3-picoline isomers$^{23,24}$ is generally applicable to perform a direct replacement of one carbon atom in benzene related aromatics with nitrogen atom in the plasma reaction. Due to the limit of the experiment results on the reaction intermediates, there may be other processes leading to the observed products. In summary, a reaction pathway for the replacement of one carbon atom in benzene with atomic nitrogen, leading to the generation of nitrogen-heterocyclic compounds, was observed in a plasma process. We anticipate the reaction pathway described herein open a door to produce a wide variety of hetero-rings using aromatics as the starting materials.

**Methods**

**On-line monitoring the reaction product.** The isotopically labeled D$_3$-benzene and D$_3$-toluene (99.96 atom% D), as well as pyridine, 2-picoline, 3-picoline, 4-picoline, and o-xylene, were purchased from Sigma-Aldrich (St. Louis, MO). The used benzene and toluene (Beijing Chemical Works, Beijing, China) were chromatographic grade. The isotopically labeled $^{15}$NH$_4$Cl ($>$98 atom% N) was purchased from Shanghai Engineering Research Center of Stable Isotope (Shanghai, China). All chemicals were used as received. For on-line monitoring the products from the plasma reaction of benzene, D$_3$-benzene, toluene, D$_3$-toluene and o-xylene, the experiments were carried out on a plasma reaction chamber (Figure S51) in an air, nitrogen (N$_2$) or nitric oxide (NO) atmosphere. Due to the strong toxicity of NO, it was balanced with 99% nitrogen gas(N$_2$). The plasma reaction was carried out in a quartz tube (O.D.: 40 mm, I.D.: 30 mm, length: 4.0 cm) with one stainless steel electrode inside and the other one touching the outside of the tube. The gas flow rate was maintained at 0.3 L·min$^{-1}$. The plasma reaction was controlled with a custom-built 5 W alternating current (AC) power supply (2.5 kV$^{-3}$F$^{-1}$ with a frequency of 3.0 kHz).

**Sample collection.** For collecting the plasma reaction product, the apparatus as shown in Figure S7 was used. A 5 W AC power supply, as described above, was applied on a surface dielectric barrier discharge electrode for the plasma reaction. The reaction gas flow rate was maintained at 0.15 L·min$^{-1}$, and the reaction temperature was kept at the room temperature. The collected sample was used for analysis without pretreatment unless explicitly indicated otherwise.

**Mass spectrometry analysis.** A Thermo Scientific Exactive Orbitrap mass spectrometer in a high-resolution mass measurement mode was employed to assign the composition of the resulting products. A TSQ Quantum Access Max (Thermo Scientific, San Jose, CA), operated in the selected reaction monitoring (SRM) mode, was used, and the specific product ions produced by collision-induced dissociation (CID) were monitored. The Xcalibur software was used for control of the TSQ Quantum Access Max MS system and data acquisition. Argon gas (99.995% purity) was used as collision gas. The temperature for MS inlet capillary was 300°C.

**GC-MS analysis.** The GC-MS analysis was completed on a TRACE GC-DSQ (Thermo Fisher Scientific, San Jose, CA) equipped with an AB-5MS capillary column (30 m × 0.25 mm i.d. × 0.25 μm). The analysis of the collected sample was operated in a split mode (50:1). High purity helium was used as carrier gas, and the flow rate was 1.0 mL·min$^{-1}$; injector temperature: 250°C; detector temperature: 250°C; column temperature: 60°C for 2 min to 300°C at 10°C·min$^{-1}$; mass spectrometry scan mode: Full scan; Mass range: m/z 30–650; electron impact: 70 eV. Xcalibur 2.0 software (Thermo Fisher Scientific Corp., San Jose, CA) was used for instrument control and data acquisition.

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**Author contributions**

Z.Z., Z.O. and X.Z. supervised the project. Z.Z., S.Z., H.Y., X.Z., X.F. and Z.O. conceived and designed the experiments. Z.Z., X.G., C.Y., Y.S. and X.X. performed the experiments. Z.Z., Z.O. and X.Z. wrote the paper and analyzed the data. All authors discussed the results and commented on the manuscript.

**Additional information**

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