Reaction **Chemistry & Engineering**



COMMUNICATION

View Article Online



Cite this: React. Chem. Eng., 2024,

Received 6th November 2023, Accepted 17th January 2024

DOI: 10.1039/d3re00593c

rsc.li/reaction-engineering

Improving the energy yield of plasma-based NO_x synthesis with in situ adsorption†

Plasma-based NO_x synthesis from air is a promising option to electrify nitrogen fixation. However, the energy efficiency of direct plasma-based NO_x synthesis in a plasma reactor is severely limited by NO_x decomposition in the plasma phase. In situ NO_x adsorption on MgO improves the NO_x energy yield in a dielectric barrier discharge (DBD) plasma reactor by a factor of 15.

Plasma-based nitrogen fixation via NOx synthesis from air was commercialized in the early 20th century. Kristian Birkeland and Samuel Eyde commercialized the first industrial nitrogen fixation process, the Birkeland-Eyde process, based on an electric arc plasma reactor.^{1,2} This process was operational in Norway and Canada.^{3,4} However, the process was eventually outcompeted by the Haber-Bosch process, producing ammonia (NH₃) from nitrogen (N₂) and hydrogen (H₂) derived from steam reforming of hydrocarbons or from water electrolysis.

Plasma-based nitrogen fixation has re-emerged in scientific literature as an option to electrify N2 fixation. 5-9 Plasma-based processes have the advantage that the load can be varied quickly, in contrast to thermal processes, so that rapid changes in supply of renewable electricity can be handled. 10 Also, absence of the need for intensive energy integration makes local production at relatively small scale feasible. In particular, plasma-N₂ fixation in the form of NO_x shows promise.5,6

So far, plasma-based NO_x synthesis in warm plasma reactors such as gliding arc (GA) and microwave (MW) plasma reactors, shows the best performance, 5,6 with NO_x formation at an energy cost down to 0.42 MJ per mol NOx, 11 which is competitive with the renewable Haber-Bosch process (using electrolysis to produce H2) combined with the Ostwald process (NH3 oxidation) in terms of energy cost (0.6 MJ per mol HNO₃).6 However, such low energy consumptions for plasma reactors are only achieved when operating at low NO_x concentrations of typically a few hundred ppm, 11 which is not practical for NO_x absorption in water in an industrial process. ¹² Concentrations in the order of 5 mol% NO_x are required for efficient processing. For those conditions, the energy consumption for NO_x formation in plasma reactors is at least in the order of 2 MJ per mol NO_x.6 Furthermore, the high temperatures in warm plasmas in the order of 10³ K cause thermal NO_x decomposition¹³ and rapid quenching is required to minimize thermal NO_x decomposition.

This work introduces a novel concept for energy-efficient NO_x formation in a dielectric barrier discharge (DBD) reactor via in situ NO_x removal using a solid MgO sorbent. A DBD reactor is a non-thermal plasma reactor operating near room temperature.¹⁴ Only a few studies have been done using DBD reactors for NO_x synthesis, $^{13,15-18}$ reporting relatively high energy consumption compared to warm plasma reactors. Nevertheless, a DBD reactor has been used in this work for two reasons. Firstly, thermal NO_x decomposition is negligible at mild temperature, and secondly, in situ NOx removal via adsorption on a solid sorbent is easily achievable. Product molecules adsorbed on microporous materials are protected against plasma induced decomposition in a DBD reactor, as we have demonstrated earlier for the case of ammonia synthesis in a DBD reactor. 19 Plasma cannot develop in submicron pores while plasma activated species are too shortlived to diffuse into these pores; therefore adsorbed molecules are protected against plasma-decomposition.8 Suppressing NO_x decomposition is expected to improve both energy efficiency and the single pass conversion in a DBD

^a Catalytic Processes & Materials, MESA+ Institute for Nanotechnology, University of Twente, P.O. Box 217, 7500 AE Enschede, The Netherlands. E-mail: l.lefferts@utwente.nl

^b Ammonia Energy Association, 77 Sands Street, 6th Floor, Brooklyn, NY 11201,

^c Koolen Industries, Europalaan 202, 7559 SC Hengelo, The Netherlands

[†] Electronic supplementary information (ESI) available: Experimental procedures, including materials and preparation, material characterization, plasma characterization, NOx synthesis and adsorption experiments. Results & discussion, including material characterization, plasma characterization, thermal NO_x TPD study with MgO, and energy cost for MgO regeneration. See DOI: https://doi.org/10.1039/d3re00593c

reactor. As earth-alkali metal oxides are known to sorb NO_x, ²⁰ we have selected MgO to demonstrate this concept. We will show that both energy efficiency as well as NOx concentrations in the product stream are drastically improved.

High surface area MgO (294 m² g⁻¹) with a particle size between 250 and 300 µm was prepared and characterized as described in ESI† section S1. The experiments were performed using a packed bed DBD reactor with MgO particles, and the NO_x concentration was determined with a mass spectrometer (MS), as described in detail in section S2.1.†

Fig. 1 shows the NO_x concentration in the outlet as a function of time on igniting the plasma, operating with an O₂: N₂ ratio of 1:1, flow rate of 20 mL min⁻¹, plasma power of 6.4 W, implying a SEI (specific energy input) of 19.2 kJ L^{-1} . The corresponding Lissajous plot is shown in Fig. S3 (see ESI†). It is observed that the NO_x outlet concentration is about 0.1 mol% during the first 4 min and increases to about 0.5 mol% afterwards. The change in NO_x outlet concentration with time in Fig. 1 is attributed to adsorption of NO_x on the MgO sorbent, followed by breakthrough of the NO_x in the outlet.

The result shown in Fig. 2 confirms that NO_x is indeed adsorbed in situ. After exposure to the plasma, the reactor was heated (25 °C min⁻¹) in N₂ (10 mL min⁻¹), inducing desorption of NO_x in absence of plasma. The concentration of NO2 is much higher than NO, but we cannot rule out NO formation, as explained in ESI 3.3.† The NO_x concentration during desorption is up to 4%, one order of magnitude higher than the NO_x concentration during steady-state plasma operation. The amount adsorbed NO_x is 0.05 mol per mol MgO.

Fig. 3 compares the TPD (temperature programmed desorption) result after NO_x synthesis by plasma illumination during different times with TPD after adsorption of NO2 at room temperature. The amount of NOx formed during the plasma experiment obviously varies with plasma time. Fig. 4 shows that the amount of NO_x detected with TPD is already

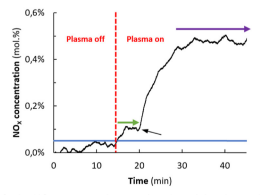


Fig. 1 Outlet NO_x concentration as a function of the plasma duration. Oven temperature set point, 25 °C; total flow rate, 20 mL min⁻¹; O₂: $N_2 = 1:1$; plasma power, 6.4 W (SEI = 19.2 kJ L⁻¹); MgO loading, 300 mg (250–300 $\mu\text{m}).$ The blue line represents the MS sensitivity limit. The green arrow line indicates in situ NOx removal with MgO. The black arrow indicates breakthrough of NO_x due to sorbent saturation. The purple arrow indicates steady-state operation.

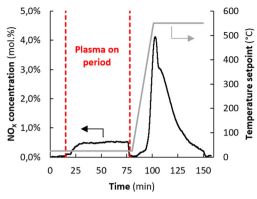


Fig. 2 Typical result of plasma-based NO_x synthesis with in situ adsorption. Reactor temperature (left axis) and outlet NOx MS signal (right axis) as a function of time. Oven temperature set point during plasma, 25 °C; total flow rate, 20 mL min^{-1} ; $O_2: N_2 = 1:1$; plasma power, 6.4 W (SEI = 19.2 kJ L^{-1}); MgO loading, 300 mg (250–300 μ m); plasma duration, 63 min; heating rate after plasma-based NO_x synthesis, 25 °C min⁻¹; pure N₂, 10 mL min⁻¹. Samples are thermally pre-treated at 550 °C in N2.

saturated after typically 5 minutes of plasma exposure. The amount is slightly lower than after thermal NO2 adsorption in absence of plasma (0.055 mol-NO2 per mol-MgO), in the same order of magnitude as reported by Duong et al.21 for MgO with a slightly lower surface area of 126 m² g⁻¹. The lower amount observed during NO_x synthesis with plasma is likely attributed to mild heating induced by the plasma, causing weakly adsorbed NO2 to desorb. A similar effect was previously reported for in situ NH3 removal by a zeolite for plasma-based NH₃ synthesis. 19 The effective NH₃ capacity on zeolite 4A decreased by 60% due to heating effects. 19

The significance of the *in situ* adsorption of NO_x from the plasma zone is evident from the reduction in energy consumption versus steady-state operation. Steady-state NO_x synthesis in this work results in an energy consumption of 93 MJ per mol NOx, in line with literature values reported in the

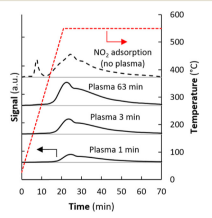
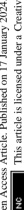
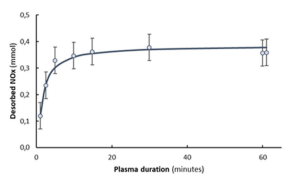


Fig. 3 TPD curve results after plasma-based NO_x synthesis $(O_2: N_2 =$ 1:1, plasma duration varied: 1 min, 3 min & 63 min), flow rate 20 mL min $^{-1}$ compared to TPD curve after adsorption of NO2 at room temperature (2 vol% NO2 in 10 vol% O2 and 88 vol% N2 balance gas during adsorption). TPD in pure N₂, 10 mL min⁻¹ and heating rate 25 °C min⁻¹.





Communication

Fig. 4 The effect of plasma duration for plasma-based NO_x synthesis with in situ adsorption on the amount of adsorbed NO_x on 300 mg MgO.

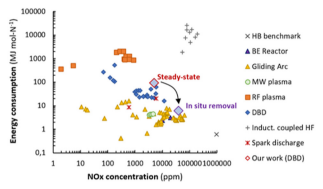


Fig. 5 Comparison of energy consumption for NO_x production in various plasma reactors. Original figure reproduced from ref. 6, and data from this work added. Steady-state: red lined diamond. In situ removal: purple lined diamond.

range 16-540 MJ per mol NO_x for DBD reactors. 13,15-18 In case of in situ product removal during 5 minutes, the energy consumption for the plasma reactor and NO_x product desorption decreases to 6.0 MJ per mol NOx as calculated in section S3.2† including the energy cost of the desorption step. Effectively, the energy consumption efficiency is improved by a factor 15. As shown in Fig. 5, in situ adsorption of NO_x in a DBD reactor results in the best performance for a non-thermal plasma-reactor, by both decreasing the energy consumption as well as increasing the NO_x concentration in the product stream. Importantly, the NO_X concentration raised well above 1%, which is considered as the minimum concentration required for nitric acid production by adsorption in water in a commercially viable process.²²

this Concluding, work demonstrates low temperature DBD plasma reactors can produce NOx with similar low energy consumptions obtained with warm plasma reactors like gliding arc reactors and microwave reactors (Fig. 5), by integrating plasma conversion and product separation. Also, the NO_x concentration in the product stream is significantly increased. This study demonstrates the concept, but optimization of the (earth-) alkali material as well of the morphology of the adsorbent is likely to result in further improvement.²⁰ Furthermore, addition of a catalyst, 23 optimization of the dielectric

properties of the adsorbent, 18 and optimization of the plasma properties could be considered.^{24,25}

In situ product removal is relevant for the wider scientific community working on plasma(-catalytic) conversion. We demonstrate that a significant improvement for NO_x synthesis and it is likely that the concept would also applicable for plasma-based CO₂ dissociation and CH₄ conversion.²⁶

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This project is co-financed by TKI-Energie from Toeslag voor Topconsortia voor Kennis en Innovatie (TKI) from the Ministry of Economic Affairs and Climate Policy, The Netherlands. The authors acknowledge B. Geerdink for technical support, K. Altena-Schildkamp for N₂ chemisorption experiments and T. Lubbers for XRF analysis. The authors acknowledge K. Van't Veer and A. M. B. Bogaerts from the University of Antwerp for discussions regarding plasma activation of N2 in DBD reactors.

Notes and references

- 1 K. Birkeland, Trans. Faraday Soc., 1906, 2, 98-116.
- 2 S. Evde, J. Ind. Eng. Chem., 1912, 4, 771-774.
- 3 A. S. Travis, Nitrogen Capture: The Growth of an International Industry (1900-1940), Springer International Publishing, 2018.
- K. H. R. Rouwenhorst, A. S. Travis and L. Lefferts, Sustainable Chem., 2022, 3, 149-171.
- 5 B. S. Patil, Q. Wang, V. Hessel and J. Lang, Catal. Today, 2015, 256, 49-66.
- 6 K. H. R. Rouwenhorst, F. Jardali, A. Bogaerts and L. Lefferts, Energy Environ. Sci., 2021, 14, 2520-2534.
- 7 N. Cherkasov, A. O. Ibhadon and P. Fitzpatrick, Chem. Eng. Process., 2015, 90, 24-33.
- 8 A. Bogaerts, X. Tu, J. C. Whitehead, G. Centi, L. Lefferts, O. Guaitella, F. Azolina-Jury, H.-H. Kim, A. B. Murphy, W. F. Schneider, T. Nozaki, J. C. Hicks, A. Rousseau, F. Thevenet, A. Khacef and M. Carreon, J. Phys. D: Appl. Phys., 2020, 53,
- 9 K. H. R. Rouwenhorst, F. Jardali, A. Bogaerts and L. Lefferts, Energy Environ. Sci., 2023, 16, 6170-6173.
- 10 A. Bogaerts and E. C. Neyts, ACS Energy Lett., 2018, 3, 1013-1027.
- 11 E. Vervloessem, Y. Gorbanev, A. Nikiforov, N. De Geyter and A. Bogaerts, Green Chem.,2022, 916-929.
- 12 W. A. Dekker, E. Snoeck and H. Kramers, Chem. Eng. Sci., 1959, 11, 61-71.
- 13 X. Pei, D. Gidon, Y. J. Yang, Z. Xiong and D. B. Graves, Chem. Eng. J., 2019, 362, 217-228.
- 14 U. Kogelschatz, Plasma Chem. Plasma Process., 2003, 23,

15 B. S. Patil, N. Cherkasov, J. Lang, A. O. Ibhadon, V. Hessel and Q. Wang, Appl. Catal., B, 2016, 194, 123-133.

Reaction Chemistry & Engineering

- 16 Q. Sun, A. Zhu, X. Yang, J. Niu and Y. Xu, Chem. Commun., 2003, 1418-1419.
- 17 A. A. Abdelaziz and H.-H. Kim, J. Phys. D: Appl. Phys., 2020, 53, 114001-114018.
- 18 Y. Ma, Y. Wang, J. Harding and X. Tu, Plasma Sources Sci. Technol., 2021, 30, 105002-105013.
- 19 K. H. R. Rouwenhorst, S. Mani and L. Lefferts, ACS Sustainable Chem. Eng., 2022, 10, 1994-2000.
- C. Verrier, J. H. Kwak, D. H. Kim, C. H. F. Peden and J. Szanyi, Catal. Today, 2008, 136, 121-127.
- 21 T. H. Y. Duong, T. N. Nguyen, H. T. Oanh, T. A. Dang Thi, L. N. T. Giang, H. T. Phuong, N. T. Anh, B. M. Nguyen, V. T.

- Quang, G. T. Le and T. Van Nguyen, J. Chem., 2019, 2019, DOI: 10.1155/2019/4376429.
- 22 K. H. R. Rouwenhorst and L. Lefferts, Catalysts, 2020, 10(9), DOI: 10.3390/catal10090999.
- 23 A. ul R. Salman, B. C. Enger, X. Auvray, R. Lødeng, M. Menon, D. Waller and M. Rønning, Appl. Catal., A, 2018, 564, 142-146.
- 24 H.-H. Kim, Y. Teramoto, A. Ogata, H. Takagi and T. Nanba, Plasma Processes Polym., 2017, 14, 1-9.
- 25 P. Peng, P. Chen, M. Addy, Y. Cheng, E. Anderson, N. Zhou, C. Schiappacasse, Y. Zhang, D. Chen, R. Hatzenbeller and Y. Liu, ACS Sustainable Chem. Eng., 2019, 7, 100-104.
- 26 K. H. R. Rouwenhorst and L. Lefferts, Plasma Processes Polym., 2023, 1-8.