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Using sustainable energy-based electricity to synthesize $NH₃$ from $H₂O$ and $N₂$ to release $O₂$ not only contributes to making chemical fertilizer production carbon neutral, but also holds promise for the use of $NH₃$ as a fuel. $NH₃$ synthesis from water and nitrogen was conducted at around 250 $^{\circ}$ C and below 1.0 MPa by combining a molten salt electrolyte of NaOH–KOH, a Pd alloy hydrogenpermeable membrane cathode, a Ni anode, and a Ru-based catalyst on the cathode backside. The rate and current efficiency for NH₃ formation were obtained as 11 nmol s $^{-1}$ cm $^{-2}$ (38 μ mol h $^{-1}$ cm $^{-2})$ and 25%, respectively, at 30 mA cm $^{-2}$, 1.0 MPa, and 250 °C. It was confirmed that the remaining percentage from the 100% current efficiency for $NH₃$ production was attributed to the current efficiency for H_2 production. The cell voltage was as low as 1.47 V at 30 mA cm $^{-2}$ and increased to 1.95 V at 100 mA cm $^{-2}$. The potential of this electrochemical system is discussed. COMMUNICATION
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Introduction

Ammonia (NH_3) not only supports human food production as a synthetic fertilizer itself or as its raw materials, but also holds promise as a carbon-free fuel when obtained from renewable energy sources.^{1–3} At present, the most reliable methods for obtaining large amounts of renewable energy involve using photovoltaics, solar thermal, wind power, $etc.⁴$ However, in advanced countries, surplus electricity from these sources is already being generated, leading to situations where it is shut down.^{5,6} It is important to convert this electricity into chemical energy carriers and store it for use at times and places when humans need it.^{7,8} Therefore, a huge number of research studies have been conducted on electrochemical reduction of nitrogen to produce NH_3^{9-13} and electrochemical reduction of CO_2 to produce hydrocarbons.^{14,15}

However, while the electrochemical reduction of $CO₂$ to CO is approaching practical efficiency,¹⁴ the production of NH₃ via electrochemical reduction of N_2 ¹⁶ and the production of hydrocarbons or alcohols *via* electrochemical reduction of CO_2 ¹⁵ are not considered to have reached practical levels due to low current efficiency and significant overpotentials. The most promising electrochemical method for obtaining ammonia is the lithium-mediated method, but the use of metallic lithium is expected to require a very large overpotential.^{16,17}

It is possible to produce $NH₃$ from electricity by utilizing water electrolysis for hydrogen production and the conventional Haber-Bosch process.¹⁸ This approach has been attempted since the early stages of $NH₃$ synthesis.¹⁸ However, large-scale chemical plants like the Haber–Bosch process face challenges in operating under fluctuating conditions, making it difficult to integrate with highly variable sources of energy, such as solar and wind power. As a result, complex and expensive systems including hydrogen storage become necessary. If NH₃ could be directly obtained from water and nitrogen within a single electrochemical device, not only would the thermodynamic energy requirements be reduced, but also the system could be operated solely with easy start–stop controls, eliminating the need for hydrogen storage.

Our group has been addressing the synthesis of $NH₃$ from $H₂O$ and $N₂$ using phosphate-based electrolytes, a composite of CsH_2PO_4 and SiP_2O_7 , suitable for operation at temperatures between 200 and 250 $\mathrm{^{\circ}C}.^{19,20}$ In this temperature range, phosphate electrolytes are effective for synthesizing various molecules.²¹⁻²⁵ The distinguishing feature of our technology lies not in electrochemically reducing nitrogen, but in catalytically activating nitrogen, which is then hydrogenated by hydrogen obtained from water electrolysis.²¹ Our goal is not to discover a new chemical phenomenon of direct electrochemical reduction of nitrogen as fundamental research, but to find out how to produce ammonia efficiently and easily from water and nitrogen using electricity. To achieve this, it is necessary to form a catalytic layer with two distinct functionalities: one portion exposed to the gas phase without being immersed in the electrolyte, and another portion

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that performs proton reduction in contact with the electrolyte. For this purpose, we devised an electrolysis cell with a Pd–Ag alloy hydrogen-permeable membrane interfacing with the electrolyte as the cathode, while a Ru-based catalyst for $NH₃$ synthesis is placed on the backside of the Pd-Ag membrane.^{19,20} The hydrogenpermeable membrane prevents the electrolyte from permeating into the Ru-based catalyst for NH₃ synthesis, avoiding the absorption of the generated $NH₃$ into the electrolyte and contamination of introduced steam into NH3. Therefore, even unstable catalysts prone to dissolution when in contact with high-temperature electrolytes can be used reliably. Furthermore, mechanical support by the Pd–Ag alloy hydrogen-permeable membrane facilitates operation under pressure, up to 1.0 MPa. Even if spanning a two-step reaction pathway involving hydrogen formation and nitrogen hydrogenation, the production of $NH₃$ and oxygen from water and nitrogen within a single-temperature electrochemical device maintains thermodynamic favorability similar to the electrochemical reduction of nitrogen.^{19,20} In other words, the exothermic heat generated by the hydrogenation of N_2 to form NH_3 can be utilized for the endothermic process of water electrolysis. Furthermore, it offers practical advantages such as eliminating the need for hydrogen storage and simplifying device start–stop procedures. However, it has been noted that in systems operating at over 60 mA cm⁻² with this CsH₂PO₄/SiP₂O₇ electrolyte, the overpotential gradually increases, leading to degradation of the anode.²⁰ Communication

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Phosphate electrolytes have been studied with the expectation of being a practical electrolyte, given the commercialization of phosphoric acid fuel cells (PAFCs) operating at around 200 °C.^{26,27} However, large-scale water electrolysis predominantly utilizes alkaline water electrolysis (AWE) at temperatures below 100 \degree C.^{28,29} There is hope for the application of this electrolyte in $NH₃$ synthesis by raising the temperature range to between 200 and 250 \degree C.^{30–32} NaOH and KOH form eutectic salts with a melting point of around 160 \degree C, which allows them to be used as molten salt electrolytes at temperatures above 200 $^{\circ}$ C.^{33–36} With sufficient humidification, steam can undergo dissolution in the molten salt resembling a concentrated alkaline solution.³³⁻³⁶ Phosphate electrolytes are in a semi-solid state at the operating temperatures, and while the usefulness of solid electrolytes is evident in fuel cells because the gas-phase network and threephase interfaces for rapidly diffusing gaseous reactants to the electrode layer are crucial, their advantages seem not to be significant in water electrolysis. This is because the products are gaseous and are evolved as bubbles even in liquid electrolytes, and the superiority of solid electrolytes is not significant. Therefore, there is potential for liquid electrolytes such as molten salts as well. Unlike acidic ion exchange membranes used in proton exchange membrane electrolysis, alkaline electrolytes like NaOH–KOH allow for the use of non-noble metals such as Ni as electrode catalysts, which is another advantage.^{28,29} In this study, we investigated $NH₃$ synthesis using a Pd–Ag hydrogen-permeable membrane cell with an electrolyte consisting of NaOH–KOH. There were past attempts at $NH₃$ synthesis using hydroxide molten salts, but none have reliably achieved nitrogen reduction.^{37,38} Separating the functions of water electrolysis and nitrogen activation/hydrogenation, it is demonstrated that hydroxide melts can be

applied to $NH₃$ synthesis in this work. The characteristics and practicality of $NH₃$ synthesis from water and nitrogen using alkaline electrolytes were discussed.

Experimental

The sectional drawing of a hydrogen-permeable membrane electrochemical cell for $NH₃$ synthesis with an NaOH–KOH molten salt electrolyte is illustrated in Fig. 1. The main components of the electrochemical cell included a catalyst powder of Ru/Cs⁺ /C weighing 0.20 g, a Pd–Ag membrane (3 : 1 atomic ratio) with a thickness of 100 μ m, an NaOH-KOH aqueous solution with a 1 : 1 molar ratio, and Ni foam with a diameter of 10 mm and a thickness of 1 mm on a Ni disk with a diameter of 10 mm and a thickness of 5 mm. The major differences from previous phosphate electrolyte electrochemical cells are that the material of the anode vessel has been changed from Ti to polytetrafluoroethylene (PTFE), and the material of the anode has been changed from Pt to Ni metal. The gap between the end faces of the cathode and anode was set to 3 mm. After assembling the anode and cathode vessels of the electrochemical cell, 5.0 mL of NaOH–KOH solution was introduced into the cell using a syringe with a narrow tube connected to the gas tube. The NaOH–KOH solution was a mixture of 0.20 mol of NaOH, 0.20 mol of KOH, and 20 mL of $H₂O$.

The cathode of the Pd–Ag membrane had a contacting area of 3.1 cm^2 to the electrolyte, and the anode of the Ni foam disk had a geometric area of 0.79 cm^2 . In this paper, the current density was defined relative to the cathode area, taking into account the uncertainty in the practical area of the Ni foam anode at the electrolyte interface. Due to length restrictions in the communication, the synthesis details of Ru/Cs⁺/C have been provided in our previous paper and the ESI,† for this article. Additionally, methods for detecting generated $NH₃$ or H2, cell heating techniques, gas supply methods, and ways to

Fig. 1 Sectional drawing of the hydrogen-permeable membrane electrochemical cell for NH₃ synthesis with an NaOH-KOH molten salt electrolyte.

increase cell pressure can be referenced in our previous papers or the ESI† of this paper. It should be noted that the flow rate of N_2 was optimized for maximizing NH₃ production, resulting in an H_2/N_2 ratio of 0.070 against current density.^{19,20}

Results and discussion

The rates of NH₃ and H₂ formation for NH₃ synthesis from H₂O and N_2 with varying temperature are plotted in Fig. 2. The current efficiencies for the respective rates are also provided in Fig. 2. The current efficiencies for the formation of $NH₃$ and $H₂$, $\mathrm{CE_{NH_3}}$ and $\mathrm{CE_{H_2}}$, respectively, were defined as follows.

$$
CE_{NH_3} = \frac{3F r_{NH_3}}{j} \times 100\ (\%)
$$
 (1)

$$
CE_{H_2} = \frac{2Fr_{H_2}}{j} \times 100\ (\%)
$$
 (2)

where $r_{NH₂}$ and $r_{H₂}$ are the rates of formation of NH₃ and H₂, respectively, in mol s^{-1} cm⁻². *F* and *j* represent the Faraday constant of 96 485 C mol $^{-1}$ and current density in A cm $^{-2}$. The product of the ratio of the thermoneutral potential of 1.32 V to the actual cell voltage and the current efficiency is the energy conversion efficiency, which is a critically important factor in energy conversion technologies.

The current efficiency for $NH₃$ synthesis was obtained as 4.0– 4.2% at 230–250 \degree C. This efficiency was similar to our previous papers. The theoretical limit of current efficiency for $NH₃$ formation at 250 °C under this N_2 flow condition was derived as 6.4% from the thermodynamical equilibrium among $NH₃$, $H₂$, and N_2 ^{19,20} The attainment of equilibrium was estimated to be 65% at 250 °C. The fact that there is little difference in the rate of $NH₃$ formation between 230 °C and 250 °C suggests that at 250 °C, the catalytic reaction is not limiting the rate, and the rate of $NH₃$ formation is constrained by the equilibrium. The decrease in $NH₃$ formation rate below 210 \degree C is attributed to insufficient catalytic reaction rates. The current efficiency for H_2 formation exceeded 100%, and when the sum of the current efficiencies of ammonia

Fig. 2 Rates and current efficiencies of $NH₃$ and $H₂$ formation for $NH₃$ synthesis from H_2O and N_2 as a function of temperature. The current density was 10 mA $\rm cm^{-2}$, and pressure was 0.10 MPa in absolute pressure. The cathode and anode gases were 3.0 $\mathsf{cm_{STP}}^3$ $\mathsf{min^{-1}}$ of $\mathsf{N_2}$ and 10 $\mathsf{cm_{STP}}^3$ min^{-1} + 10 μ L min $^{-1}$ of Ar + H₂O_{liq}, respectively.

Fig. 3 Rates and current efficiencies of NH₃ and H₂ formation for NH₃ synthesis from H_2O and N_2 as a function of current density. The absolute pressure and temperature were 0.10 MPa and 250 °C. The cathode and anode gases were 3.0 cm_{STP}^3 min⁻¹ of N₂ per 10 mA cm⁻² and 10 cm_{STP}³ min⁻¹ + 10 μ L min⁻¹ of Ar + H₂O_{liq}, respectively.

production and hydrogen production was calculated, it was approximately 100%.

The advantage of NaOH–KOH electrolyte is its ability to maintain stability and conduct long-term experiments at a relatively high current density of 100 mA cm^{-2} . In our conventional system using phosphate electrolytes, when the current density exceeds 50 mA cm^{-2} , the cell voltage starts to rise within a few hours, leading to deterioration. The rates of $NH₃$ and H_2 formation for NH₃ synthesis from H₂O and N₂, along with their current efficiencies, are depicted in Fig. 3 as a function of current density up to 100 mA cm^{-2} . Exceeding a current density of 100 mA cm^{-2} was not possible due to limitations in the capacity of the power supply. The rates of NH3 and H2 formation increased monotonically with increasing current density. Looking at the current efficiency, between 10 and 30 mA $\rm cm^{-2}$, both the current efficiency for NH₃ formation and for $H₂$ formation remain almost constant. However, when the current density is increased beyond 50 mA cm^{-2} , it becomes evident that both the current efficiency for $NH₃$ formation and for H_2 formation decrease.

The decrease in current efficiency for $NH₃$ formation at high current densities can be understood as insufficient catalytic activity. In such cases, the current efficiency for H_2 formation must increase by the amount that the current efficiency for $NH₃$ formation decreases. The most likely cause of both the current efficiency for $NH₃$ and $H₂$ formation decreasing, resulting in a decrease in overall current efficiency, is the $O₂$ generated at the anode either being reduced at the cathode by passing through the electrolyte or the hydrogen reduced at the cathode being

unable to be absorbed by the Pd–Ag membrane and instead released as H_2 molecules in the electrolyte, subsequently oxidizing at the anode. The issue of such cross-leakage becoming problematic under high current densities was already recognized in our previous studies using phosphate electrolytes,²⁰ and it became pronounced even with NaOH–KOH electrolytes. Particularly in liquid electrolytes such as molten salts, crossleakage poses a serious problem, as the evolved gases might pass through the electrolyte as bubbles or absorbed gases between the anode and cathode. Therefore, it seems necessary to explore measures such as optimizing the electrode spacing and installing separators with semi-permeable membranes.

Next, the results of time-dependent changes in cell voltage during $NH₃$ synthesis at constant current are shown in Fig. 4A for various current densities. At 10 mA $\rm cm^{-2},$ the cell voltage was about 1.28 V, which was between the equilibrium potential (1.16 V at 250 $^{\circ}$ C) and thermoneutral potential (1.32 V at 250 $^{\circ}$ C). With increasing current densities, cell voltage was increased to 1.96 V at 100 mA $\rm cm^{-2}.$ At any current density, the cell voltage remained nearly constant over a period of 3 h, exhibiting remarkable stability. Fig. 4B shows the cell voltage changes in an operation for 3 weeks. While minor fluctuations in cell voltage were observed, there were no noticeable voltage increases indicative of degradation. The catalyst is inherently non-degrading, so consistent NH₃ production was observed at

any given point as long as water electrolysis continued at the stated current density. This was a highly favourable outcome compared to cells using phosphate electrolytes, as demonstrated in our previous papers, where an increase in cell voltage was observed within several hours at current densities exceeding 50 mA cm $^{-2}$.

The synthesis of $NH₃$ by this system is strongly limited by chemical equilibrium, and theoretically, at a pressure of 0.10 MPa, temperature of 250 °C, and the present ratio between N_2 and current density, only up to 6.4% current efficiency of $NH₃$ can be expected. To relax the constraints of equilibrium, lowering the temperature while increasing the pressure is an option; however, as shown in Fig. 2, reducing the temperature shifts the limitation from equilibrium to kinetic, resulting in a decrease in the rate of $NH₃$ formation. Therefore, the results of investigating the rate of $NH₃$ formation when increasing the total pressure are shown in Fig. 5. At 1.0 MPa and 250 $^{\circ}$ C, the theoretical limit is estimated to be 36% .^{19,20} As shown in Fig. 5, while the current efficiency for $NH₃$ formation was 4.76%, at 0.10 MPa, 10 mA cm^{-2} and 250 °C, it was increased to 30% with elevating the pressure to 1.0 MPa. For 30 mA $\rm cm^{-2}$ current density at 1.0 MPa and 250 $^{\circ}$ C, the current efficiency for NH3 formation decreased to 25% because of insufficient catalytic activity. Communication

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The results of this study show that, unsurprisingly, the current efficiency of $NH₃$ production was very similar to the values obtained in our previous research, as the catalyst layer used was the same.^{19,20} However, the significant advantage lies in the stable operation at higher current densities and the absence of observations such as electrode degradation. An important point not to overlook is that concentrated alkaline electrolytes, being strong alkalis, do not absorb $NH₃$. In previous electrochemical cells using phosphate-based electrolytes, the use of a Pd–Ag membrane was imperative as it prevented the absorption of $NH₃$ by the protonic electrolyte and the subsequent neutralization of the electrolyte. $19,20$ If the electrolyte does not absorb $NH₃$, it might be possible to use hydrophobic filters instead of Pd–Ag membranes, as in the synthesis

Fig. 4 Time courses of cell voltage for $NH₃$ synthesis from $H₂O$ and $N₂$ at various current densities and 250 °C. A summary of short-term current fluctuations (A) and the results obtained over a period of three weeks of operation (B). Note that (A) and (B) represent the experimental results from different cells that were reassembled, so the results do not perfectly match.

Fig. 5 Current efficiency of NH₃ and H₂ formation for NH₃ synthesis from H_2O and N_2 at 250 °C with varying absolute pressure and current density. The rates of $NH₃$ formation are shown in this figure. The cathode and anode gases were 3.0 cm_{STP}^3 min⁻¹ of N₂ per 10 mA cm⁻² and 10 cm_{STP}³ min⁻¹ + 10 µL min⁻¹ of Ar + H₂O_{liq}, respectively.

of methane from $CO₂$ and water using acidic phosphate electrolytes in this electrochemical system.²² Alkaline water electrolysis conducted in aqueous solutions below 100 °C is considered crucial for hydrogen production in a hydrogen energy society. By increasing the operating temperature to around 250 \degree C, it is conceivable to synthesize NH₃ and produce various other fuel substances and useful materials.

Conclusions

With KOH–NaOH eutectic molten salt electrolytes, an electrochemical-catalytic cell for $NH₃$ synthesis from nitrogen and water was studied at 250 $^{\circ}$ C and 1.0 MPa. The current efficiency for NH_3 formation was achieved to 30%, and the residue was used for H_2 formation. The cell voltage was extremely stable up to 100 mA $\rm cm^{-2},$ and no corrosion of the electrodes or components was observed, suggesting that this study shows significant potential for obtaining ammonia from water and nitrogen using electricity. Energy Advances

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Author contributions

Sagara primarily contributed to the investigation, which included experimental work, analysis of the obtained data, and paper writing. Hayashi, Hirata, and Nagaishi were primarily involved in practical experimental works and data analysis. Kubota played multifaceted roles including supervision in this study.

Conflicts of interest

There are no conflicts to declare.

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