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Promotion effect of proton-conducting oxide BaZr$_{0.1}$Ce$_{0.7}$Y$_{0.2}$O$_{3-\delta}$ on the catalytic activity of Ni towards ammonia synthesis from hydrogen and nitrogen

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Abstract

In this report, for the first time, it has been observed that proton-conducting oxide BaZr$_{0.1}$Ce$_{0.7}$Y$_{0.2}$O$_{3-\delta}$ (BZCY) has significant promotion effect on the catalytic activity of Ni towards ammonia synthesis from hydrogen and nitrogen. Renewable hydrogen can be used for ammonia synthesis to save CO$_2$ emission. By investigating the operating parameters of the reaction the optimal conditions for this catalyst were identified. It was found that at 620 °C with a total flow rate of 200 mL min$^{-1}$ and a H$_2$/N$_2$ mol ratio of 3, an activity of approximately 250 μmol g$^{-1}$ h$^{-1}$ can be achieved. This is ten times larger than that for the unpromoted Ni catalyst under the same conditions although the stability of both catalysts in the presence of steam was not good. The specific activity of Ni supported on proton-conducting oxide BZCY is approximately 72 times higher than that of Ni supported on non-proton conductor MgO–CeO$_2$. These promotion effects were suspected to be due to the proton conducting nature of the support. Therefore it is proposed that the use of proton conducting support materials with highly active ammonia synthesis catalysts such as Ru and Fe will provide improved activity of at lower temperatures.

Introduction

Ammonia, due to its important use as a fertiliser is mass produced at approximately 150 million tons per annum [1]. That fact that ammonia fertilisers have supported approximately 27% of the world’s population over the last century confirms its importance, with the development of the high temperature Haber-Bosch process being considered one of the most important chemical processes of the last 100 years [2]. In this process, ammonia is produced from H$_2$ and N$_2$ under high temperature and high pressure. About 50% of the hydrogen produced in the world is used for ammonia production [3]. In addition, CO$_2$ emitted from the ammonia industry is equivalent to 0.77% of the world total CO$_2$ emission because the
source of hydrogen is mainly from steam-reforming of natural gas [3,4]. This green-house gas emission can be significantly reduced if the source of hydrogen can be changed to be provided through renewable sources such as hydro-electricity, wind or solar energy. Hydrogen from the electrolysis of water (hydro-electricity) was used for low-carbon ammonia production through Haber-Bosch process in Norway [5]. A Dutch company Proton Venture is developing de-centralised ammonia production technology using renewable electricity as the energy source [6]. Therefore synthesis of ammonia from renewable hydrogen could be a complementary part of hydrogen economy [3,7–11].

The Haber-Bosch process is very important in chemical industry. This led to Fritz Haber being awarded the Nobel Prize in 1918 and with Carl Bosch also receiving the award in 1931. Gerhard Ertl also won the Nobel Prize in Chemistry in 2007 for his contributions towards the understanding of the surface chemistry of iron catalyst [12]. Industrially promoted fused iron catalysts are used in the Haber-Bosch process with promoters including potassium, barium and aluminium oxides [13–18]. As well as iron, supported ruthenium has attracted a large amount of research interest as an ammonia synthesis catalyst with carbon supported ruthenium currently employed in the Kellogg advanced ammonia process (KAAP) [19]. Due to the cost of ruthenium, it is supported on a range of materials in order to be used in the ammonia synthesis process. These supports often also fill the role of promoter with very high activities recently reported for Ru/HT–C12A7:0.25 [20], Ru/Y2Si2 [21], and Ru/MgO supports. In 2013 Wang Z et al. [23] reported the use of perovskite material BaZrO3 as a catalyst support for the ammonia synthesis reaction at high pressure and intermediate temperature when ruthenium was used as the catalyst.

The reaction to produce ammonia from hydrogen is shown below [19].

\[
N_2 + 3H_2 \rightarrow 2NH_3 \quad (\Delta H^\circ = -46.1 \text{ kJ mol}^{-1})
\]  

Although this reaction is exothermic and favours high pressure and low temperature the ammonia formation rate at these conditions is extremely low even though an ammonia equilibrium concentration of close to 100% is theoretically possible. For this reason, the Haber-Bosch process is carried out at temperatures between 400 °C and 500 °C at pressures close to 200 bar.

It is commonly accepted that the rate limiting step for the ammonia synthesis reaction is the dissociative adsorption of nitrogen although the reaction could also be limited by the N binding energy due to the Sabatier principle [24]. This is shown in Eqs. (2)–(8) below:

\[
N_2 + \star \rightarrow N_2^\star \quad (2)
\]

\[
N_2^\star + \star \rightarrow 2N^\star \quad (3)
\]

This dissociated nitrogen reacts with dissociated hydrogen on the catalyst surface to form ammonia. The dissociative adsorption of hydrogen and the surface reactions are shown below where * represents the catalyst active site:

\[
H_2 + 2^\star \rightarrow 2H^\star \quad (4)
\]

\[
H^\star + N^\star \rightarrow NH^\star + ^\star \quad (5)
\]

\[
NH^\star + H^\star \rightarrow NH_2^\star + ^\star \quad (6)
\]

\[
NH_2^\star + H^\star \rightarrow NH_3^\star + ^\star \quad (7)
\]

\[
NH_3^\star \rightarrow NH_3 + ^\star \quad (8)
\]

It can be seen from this that if the dissociative adsorption of hydrogen is favoured on the catalyst surface then there will be less active sites for the rate limiting step therefore reducing activity. It has been reported that reduce dinitrogen will be feasible upon addition of protons and electrons similar to the mechanism of ammonia production by nitrogenases. This has been described in detail in an excellent review [25].

Recently, it was discovered that ammonia could be produced from hydrogen and nitrogen in an electrochemical cell [7,26–35]. This led Vasileiou E et al. [36,37] to achieve the electrochemical synthesis of ammonia using a proton conducting electrolyte. They achieved this by using BaCe0.2Zr0.7Y0.1O2.9 as the electrolyte with Ni–BaCe0.2Zr0.7Y0.1O2.9 used as the cathode and anode. When they performed the experiment with no applied voltage they discovered that ammonia was also produced through the catalysed reaction of H2 and N2 fed to the cell. Although the catalytic activity of the Ni–BaCe0.2Zr0.7Y0.1O2.9 material was noted no further investigation was done. In this paper, the effects of a similar proton conducting electrolyte on the catalytic activity towards ammonia synthesis, were investigated with the use of a Ni catalyst. BaZr0.1Ce0.7Y0.2O3−δ (BZCY) was chosen due to its excellent proton conductivity [38,39]. Ni-BZCY has been investigated as membrane materials for hydrogen separation using the proton conduction of BZCY [40,41]. By using a proton conducting support, it is proposed that the dissociated hydrogen on the catalyst active sites will be transferred to the proton conducting support, therefore freeing active sites for the dissociative adsorption of nitrogen. Through this increase in active sites, it is expected that the catalyst activity will be greatly increased. In this study, the catalytic activity of Ni and Ni supported on BZCY or MgO–CeO2 are investigated. It was found that proton-conducting oxide BZCY has promotion effects on the catalytic activity towards ammonia synthesis.

Experimental

Materials and preparation method

In order to synthesise the BaZr0.1Ce0.7Y0.2O3−δ (BZCY) perovskite catalyst support a solid state reaction was employed. Firstly stoichiometric amounts of BaCO3 (99% alfa), ZrO2 (99% alfa), CeO2 (99.5% Alfa) and Y2O3 (99.9% Alfa) were weighed and mixed using a pestle and mortar. The resulting mixture was then wet ground in isopropyl alcohol for 12 h. After drying at 80 °C the mixture was then fired at 1000 °C for 3 h with a heating and cooling rate of 5 °C min−1. After this NiO (99% Alfa) was added to the BaZr0.1Ce0.7Y0.2O3−δ...
powder with a weight ratio of 60%–40% respectively. This was further wet ground in isopropyl alcohol for 12 h. The MgO–CeO₂ support for the comparison test was prepared through a combustion synthesis in which equimolar amounts of Ce(NO₃)₃·6H₂O (99.5% alfa) and Mg(NO₃)₂·6H₂O (98% alfa) were dissolved in deionised water, citric acid (99% alfa) was then added with the mole ratio of 1:1 against total moles of metal ions. This solution was then heated on a hot plate at 200 °C until the combustion was complete with the resulting powder fired at 500 °C for 2 h. After this NiO (99% Alfa) was added to the MgO–CeO₂ powder with a weight ratio of 60%–40% respectively. This was further wet ground in isopropyl alcohol for 12 h.

Materials characterisation

The catalyst was characterised using both X-ray Diffraction (XRD) and Scanning electron microscopy (SEM). The crystal structures were determined using a Pananalytical X’Pert Pro Multi-Purpose Diffractometer (MPD) with Cu K alpha 1 radiation working at 45 kV and 40 mA. The SEM images were obtained with ZEISS SUPRA 55-VP operating at 10 kV. Thermal-gravimetry-differential scanning calorimetry (TG-DSC) analyses of pre-reduced Ni-BZCY catalysts were carried out on a NETSCH STA 449 F1 thermal analyser in flowing N₂ to 800 °C with an N₂ flow rate of 70 mL min⁻¹. FT-IR measurements were carried out on a Bruker Vertex 70V IR spectrometer. The specific surface area of both the Ni-BZCY catalyst and the Ni–MgO–CeO₂ catalyst was measured using a QUADRASORB SI surface area analyser. Both of the reduced samples were degassed at 350 °C before carrying out surface area analysis at liquid nitrogen temperature.

Catalyst activity measurement

To measure the catalytic activity 0.48 g of catalyst was loaded into an alumina reactor and was supported in the centre on glass fibre. The catalyst was then reduced at 700 °C in H₂ and N₂ with a total flow rate of 100 mL min⁻¹ and mole ratio of 9:1 H₂:N₂ for 4 h. After this the temperature, total flow rate and flow rate ratio were adjusted in order to determine the optimal conditions. H₂ and N₂ from gas cylinders were directly used without any purification process. For the stability test, the catalyst was cooling down to room temperature under the protection of mixed H₂:N₂ (3:1 m/o), then N₂ passing through room temperature water was passed through the catalyst for one hour. After this process, the gas was switched to mixed H₂ and N₂ then slowly heated to 620 °C to continue the ammonia synthesis measurement.

Dilute H₂SO₄ (0.01 M) was used to collect any produced ammonia which was then measured using ISE Thermo Scientific Orion Star A214 ammonia meter. Both hydrogen and nitrogen were used from the cylinder with no further purification.

In order to calculate the ammonia synthesis rate the following equation was used.

$$r_{\text{NH}_3} = \frac{[\text{NH}_3]}{t \times m} \times V$$  (9)

where [NH₃] is ammonia concentration in mol L⁻¹, V is volume of 0.01 M H₂SO₄ in L, t is time in hours and m is catalyst mass in g.

Results and discussion

Materials characterisation

XRD analyses
In the XRD results shown in Fig. 1, it can be seen that there are some small peaks attributed to BaCO₃ and Y-doped CeₓZr₁₋ₓO₂ present for BaZr₀.₈Y₀.₂O₃₋₄ before and after being mixed with the NiO; however, after reduction at 700 °C in H₂/N₂ mixture (90%H₂) for 4 h, these peaks are no longer present. A possible reason is that, BaCO₃ and Y-doped CeₓZr₁₋ₓO₂ were converted into amorphous phase during the reduction process thus cannot be detected by XRD. The XRD peaks for the catalyst before and after the stability test are the same although the intensity of the Ni peak has increased after the stability tested showing the possible aggregation of Ni particles whilst better crystallisation is another possible reason.

FT-IR analysis
In order to identify the BaCO₃ phase, the absorbance spectra of the catalyst were measured before and after reduction to investigate whether or not BaCO₃ and Y-doped CeₓZr₁₋ₓO₂ are converted into amorphous phases. The absorbance spectra of pure BaCO₃, ZrO₂, CeO₂ and the catalysts before and after catalysts test were measured using a Bruker Vertex 70V IR spectrometer. FT-IR was employed over the wavenumber range of 500 cm⁻¹ to 4000 cm⁻¹. Fig. 2 shows that the adsorption for BaCO₃ at around 1440 cm⁻¹ were present in both the reduced and unreduced catalyst, remaining present in an amorphous phase throughout the catalyst tests. The observed IR adsorption for BaCO₃ is consistent with that in a previous report [42]. It has been reported that a similar perovskite oxide to BZCY, BaZr₀.₈Y₀.₂O₃₋₄ does not exhibit adsorption around this wavenumber [43]. It was noticed that the adsorption peak for BaCO₃ at approximately 1250 cm⁻¹ is
weaker in reduced Ni-BZCY sample, the possible reason is
that, during the reduction process, the generated Ni particles
diffuse and coat on the surface of BaCO₃ particles thus IR ab-
sorption for BaCO₃ is reduced. The decomposition of the trace
amount of BaCO₃ is unlikely as the sample was prepared at
1000 °C which is 300 °C higher than the reducing temperature.

SEM observation
Fig. 3a and b show the SEM pictures of unreduced NiO-BZCY
catalyst. The big particles are BZCY oxide with small NiO
particles homogeneously distributed in the oxide matrix. The
secondary particle size of NiO was about 0.2 μm (Fig. 3a). After
high temperature reduction, the particle size of generated Ni
particles was about 1.5 μm Ni (Fig. 3b), due to coarsening of Ni
particles during the high temperature reduction. The particle
size of Ni did not exhibit significant increase during the long
time stability test (Fig. 3c). The increased intensity of Ni
component in XRD is more likely due to better crystallisation
during the high temperature operation. Element mapping of
reduced Ni-BZCY is shown in Fig. 4. The distribution of Ni
(Fig. 4b) and other elements is homogeneous. From Fig. 5 the
particle size of Ni in Ni-BZCY and Ni–MgO–CeO₂ was
compared, from this it was observed that a similar Ni particle
size was achieved over both catalysts despite the large dif-
ference in particle size observed between BZCY and MgO–
CeO₂. The particle size of MgO–CeO₂ prepared at 500 °C is
much smaller than that of BZCY prepared at 1000 °C. This is
consistent with the specific surface area for Ni-BZCY
(0.907 m² g⁻¹) and Ni–MgO–CeO₂ (16.940 m² g⁻¹) measured
by BET method.

TG-DSC analysis
In order to figure out the effects of moisture on the properties
of the reduced Ni-BZCY catalyst, ‘TG-DSC’ analyses were car-
rried out for both dry and wet reduced Ni-BZCY catalysts. For
the wet catalyst, reduced Ni-BZCY catalyst was exposed to
flowing air through room temperature for 1 h before the TG-
DSC measurement. The TG-DSC data for both samples are
shown below in Fig. 6(a) and (b) respectively. For the dry
catalyst, the initial weight loss below 100 °C (~0.12 wt%) is due
to the loss of absorbed water. When the wet reduced Ni-BZCY
was used, the initial weight loss continued to ~250 °C, with a
larger weight loss (~0.34 wt%) indicating BZCY can hold water
to a higher temperature. A shoulder weight gain peaked
around 450 °C was observed which is due to water uptake.
This was also observed in protonic conducting oxides [44,45].
On cooling, more water uptake (~0.18 wt%) was observed.
indicating BZCY can strongly uptake water at lower temperatures. On the DSC curves, it has been noticed that there is an endothermic shoulder peak with positive thermal effect (on right Y-axis) has been observed peaked at around 625 °C for both samples which is related to the slow desorption of water. The observed weight and thermal effects are consistent with those in previous reports [44,45].

**Catalytic activity towards ammonia synthesis**

**Effect of temperature on catalyst activity**

When a constant flow rate was kept at 120 mL min⁻¹ and H₂:N₂ were flown with a mole ratio of 3:1 the effects of changing temperature could be observed, this is shown in Fig. 7. It was observed that the activity increases up to a maximum of approximately 135 μmol g⁻¹ h⁻¹ at 620 °C before dropping.
again. At lower temperature, the catalytic activity of the Ni-BZCY catalyst is not high enough. At a higher temperature, the produced ammonia may decompose due to the thermodynamic equilibrium for the reaction shifting to the left as the reaction is exothermic, leading to lower production rate. In Fig. 6b, a weight loss at ~650°C was observed due to the loss of uptaken water. This temperature is very close to the highest catalytic activity as shown in Fig. 7. Therefore promotion effect of the BZCY could be related to the uptaken water at high temperature. This trend was also observed by Vasileiou E et al. [1] in their cell when hydrogen and nitrogen were fed to the Ni–MgO–CeO2 electrode under open circuit conditions.

The ammonia outlet concentration was plotted alongside the equilibrium concentration for the reaction in Fig. 8. Thermodynamic equilibrium data was obtained from Haber’s Nobel Lecture [46]. It can be observed that the outlet concentration reaches a maximum and starts to decrease before obtaining the outlet concentration expected at thermodynamic equilibrium. It is therefore expected that the activity over 620 °C is inhibited due to some other means. The outlet concentration of ammonia at 700 °C was still closer to the thermodynamic equilibrium than those at lower temperatures.

Effect of total flow rate on catalyst activity
The effect of total flow rate was then tested at a constant temperature of 620 °C with the results shown in Fig. 9. It can be seen that the activity increases with increasing flow rate. This increase in activity expected to be due solely to the increase in reactant gas, in order to confirm this ammonia outlet concentration was plotted against total gas flow rate.

As shown in Fig. 10, when total flow rate is plotted against ammonia outlet concentration, it rises up to a total flow rate of 120 mL min⁻¹ before levelling off. The outlet concentration was calculated using the following equation

\[
\text{Outlet NH}_3 \text{ concentration} = \left( \frac{r_{\text{NH}_3} \times m}{\text{Total molar flowrate}} \right) \times 100% \tag{10}
\]

This therefore shows that the total flow rate is independent of conversion rate over a value of 120 mL min⁻¹ in our experiments and that the activity measured at this these flow rates is solely due to catalytic activity. However, at total gas flow rates less than 120 mL min⁻¹, lower outlet ammonia concentration was observed. The possible reason is that, majority of the mixed gas passed through the edge of the glass fibre where the loading of catalyst was relatively lower thus the contact time with the catalyst was short leading to reduced ammonia formation.

Effect of feed gas ratio on catalyst activity
To determine the optimal feed ratio the gas inlet mole ratio was adjusted area when 2.6 and 3.4 (H₂/N₂) with the optimal being detected for a value of 3.2 with a rate of approximately 320 µmol g⁻¹ h⁻¹ (Fig. 11). All measurements were taken at 620 °C with a total flow rate of 200 mL/min. The reason for this deviation from the normal may be due to the proton conducting nature of the BZCY support with some of the fed H₂
being ionised and transferred to the support as $H^+$ therefore adjusting the value of $H_2$ to $N_2$ in the reactor closer to the stoichiometric value of 3. This phenomenon of a $H^+$ being transferred to the active sites was observed by Shin H et al. [47] when they used proton conducting support materials for a Cr catalyst in the degradation of ethane.

Effect of temperature on catalyst activity of Ni/MgO–CeO$_2$

In order to examine the promotion effects of the proton conducting nature of the catalyst support, a Ni catalyst supported on a non-proton conductor was tested under the same conditions. It has been reported that MgO–CeO$_2$ composite is an excellent support for Ru catalysts for ammonia synthesis [48]. In this study, for comparison, Ni supported in MgO–CeO$_2$ composite was also synthesised and the catalytic activity was also investigated. This was tested over the temperature range of $600 \degree C$ to $640 \degree C$ with a hydrogen to nitrogen mole ratio of 3 and a total flow rate of 120 mL min$^{-1}$ (Fig. 12). From this it can be seen that the maximum flow rate achieved was at $620 \degree C$ mirroring that results obtained for the BZCY support. However, the activity of this catalyst is around 4 times lower than the activity of the Ni catalyst when used with the BZCY proton conducting support (Fig. 7). However, the catalytic activity is related to the specific surface area. The specific surface area was measured to be 0.907 m$^2$ g$^{-1}$ for the Ni-BZCY catalyst and...
16.940 m² g⁻¹ for the Ni–MgO–CeO₂ catalyst. The specific surface area of Ni-BZCY is only 5.3% of that of Ni–MgO–CeO₂ but the catalytic activity to ammonia synthesis is much higher, with a specific activity of approximately 149 μmol m⁻² h⁻¹ for the Ni-BZCY catalyst compared to 2 μmol m⁻² h⁻¹ for the Ni–MgO–CeO₂ catalyst. This experiment further demonstrates that proton-conducting oxide BZCY has obvious promotion effects on ammonia synthesis.

Stability of catalytic activity in the presence of moisture

The stability of ammonia synthesis catalysts in the presence of an oxidant is a big challenge [12,19]. It has been reported that Ru–YₓSi₁₋ₓ catalyst exhibit good stability in the presence of moisture [21]. Inspired by this excellent work, we also investigated the stability of our catalysts in the same manner. The catalyst stability was investigated over 144 h at 620 °C with a H₂/N₂ mole ratio of 3 and a total flow rate of 200 mL min⁻¹. The catalyst was found to be stable over this period with no loss of activity as can be seen in Fig. 13. After this the effect of wetting the catalyst was also investigated. To perform these experiments the reactor was cooled to room temperature and wet nitrogen (100 mL min⁻¹) was bubbled through the reactor for 1 h before being heated back to 620 °C at a rate of 1 °C min⁻¹. This was repeated 5 times with the results shown in Fig. 13. It can be seen from the results that there is a drop in activity after each cycle with an overall linear drop over the 5 cycles. The activity drops to approximately a fifth of its original value after 5 cycles going from approximately 250 μmol g⁻¹ h⁻¹ to 50 μmol g⁻¹ h⁻¹ with a further drop expected on further wetting cycles. This loss of activity was suspected to be caused either due to the poisoning effect of the water on the Ni catalyst after being wetted at room temperature because slight oxidation of Ni on the surface may happen as the case for Fe-based catalysts [12]. However, upon examining the XRD patterns and SEM images of the reduced catalyst after the stability test no major changes in particle size were observed from the freshly reduced catalyst (Fig. 3b and c). However, a trace amount of NiO may still have been formed after treating the catalyst but is beyond the measurement limit for XRD. This NiO will be in situ reduced to metallic Ni at high temperature in the presence of high concentration of H₂. The oxidation and reduction cycles that the Ni catalyst undergoes in the wetted catalyst may also damage the active sites on the catalyst greatly speeding up the degradation of the catalyst that would be noticed over the catalysts lifetime [49], evidence for this was observed during the XRD which showed an increase in intensity of the Ni peak after the stability test suggesting possible better crystallisation of Ni particle leading to loss of active sites on the Ni surface. This effect of enhanced catalyst degradation may also be attributed to the heating and cooling cycles in-between each data point on the wetted catalyst stability test.

As well as the BZCY promoted catalyst pure Ni was also tested with a rate of 25.12 μmol g⁻¹ h⁻¹ observed at 620 °C with a total flow rate of 200 mL/min and a H₂/N₂ ratio of 3. This is roughly ten times lower than that for the BZCY promoted catalyst when the same weight of nickel oxide was used. This
Therefore shows the excellent promotion effects that can be achieved using the BZCY proton conducting support. When investigating materials as potential supports for ammonia synthesis catalysts the electro negativity of the support is a strong consideration [50–52]. It has been shown that the electron transferring nature of support could be attributed to increased activity [50–52]. This is mainly prominent in Ru catalyst supports where the transferred electrons change the electronic state of the Ru catalyst [53]. However, in this work, we have shown that another desirable effect of a support material may be its ability to conduct protons. This promoting ability of proton conducting supports could be explained by the ionisation of the H₂ gas fed to the reactor. It is commonly accepted that the rate limiting step for the ammonia synthesis reaction is the dissociation of the N₂ molecules on the catalyst surface [54,55] or the N-binding for Fe catalysts; however, if molecular H₂ is adsorbed on the catalyst active sites instead then the catalytic activity would suffer due the loss of active sites for the rate limiting step [56]. By using a proton conducting support it is suggested that the dissociated hydrogen on the active sites is then transferred in to the support freeing the site for the adsorption of nitrogen. The dissociated nitrogen on the active sights may then further react with the H⁺ ions in the support further increasing activity. It has been reported that the active sites for CO₂ reduction on Cu from reduced CuO is mainly on the grain boundary whilst the well crystallized BZCY has higher promotion activity that less crystallised MgO–CeO₂ support [57]. This could be related to the proton conduction of BZCY. Other possibilities such as lattice mismatch or metal/metal-oxide support interactions/adhesion cannot be ruled out, which need further investigation. Anyway, the reaction mechanisms on CO₂ reduction and N₂ reduction are very different and hard to compare. Despite these promotion effects, the reaction temperature is too high for any meaningful application of the reported catalyst; however, the excellent promotion effects of the proton conducting support has been shown with the activity increasing by an order of magnitude. It is therefore expected that when used with more conventional catalysts such as Ru, Co or Fe then high activities at lower temperatures more in line with those employed in the Haber-Bosch process can be achieved.

**Conclusion**

The use of a BZCY proton conducting support has proven to increase the activity of Ni towards ammonia synthesis by an order of magnitude. This increase in activity was observed even though no gas purification was employed. At 620 °C and a total flow rate of 200 mL min⁻¹ it was observed that an ammonia synthesis rate of approximately 250 μmol g⁻¹ h⁻¹ could be achieved which is ten times higher than that observed for unpromoted nickel under the same conditions. The activity of Ni supported on proton-conducting oxide BaZr₀.₁Ce₀.₇Y₀.₂O₃₋₈ is four times that of Ni supported on non-proton conductor MgO–CeO₂ although the specific surface area of the former is much lower, giving a specific activity of around 72 times lower than that of Ni supported on BaZr₀.₁Ce₀.₇Y₀.₂O₃₋₈. The observed activity of the promoted Ni catalysts is slightly lower than the known Fe and Ru based catalyst at 400–500 °C and a pressure of one atmosphere because Ni itself is not a good ammonia synthesis catalyst [21,58,59]. It is therefore expected that when employed with more active catalysts such as Fe, Co or Ru then an increase in activity could be achieved which we believe is tied to the proton conducting nature of the support. The relevant research are on-going in our group.

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