Al Sobhi, S., Bion, N., Hargreaves, J. S.J., Hector, A. L., Laassiri, S., Levason, W., Lodge, A. W., McFarlane, A. R. and Ritter, C. (2019) The reactivity of lattice nitrogen within the Ni2Mo3N and NiCoMo3N phases. Materials Research Bulletin, 118, 110519.

There may be differences between this version and the published version. You are advised to consult the publisher’s version if you wish to cite from it.

http://eprints.gla.ac.uk/189059/

Deposited on: 29 October 2019
Citrate-gel preparation and ammonia synthesis activity of compounds in the quaternary (Ni,M)$_2$Mo$_3$N (M = Cu or Fe) systems

Samia Al Sobhi,$^a$ Justin S. J. Hargreaves,$^{a, *}$ Andrew L. Hector$^{a,*}$ and Said Laassiri$^b$

$^a$ School of Chemistry, University of Southampton, Highfield, Southampton SO17 1BJ, United Kingdom

$^b$ WestCHEM, School of Chemistry, University of Glasgow, Joseph Black Building, Glasgow G12 8QQ, United Kingdom

Single phase solid solutions based on copper and iron substitution into the nickel sites of Ni$_2$Mo$_3$N are produced using a citrate gel approach followed by ammonolysis. These metal nitrides show good ammonia synthesis activity at 500 °C and ambient pressure. The activity fell with copper content and increased with iron content, although these changes appear to be related to changing surface areas of the catalysts. The oxide intermediates from the citrate gel process cleanly converted to the metal nitride under the ammonia synthesis gas stream (75% H$_2$, 25% N$_2$) providing more active catalysts than those pre-nitrided by ammonolysis.

Introduction

The Haber-Bosch process directly sustains a major fraction of the world’s population by providing a route from N$_2$ to fertilisers, and demand could rise if ammonia is employed significantly as a carbon-free fuel. However, Haber-Bosch consumes 1-2% of the world’s energy supply and results in significant carbon emissions in necessarily centralised (and hence poorly suited to renewable energy use) environments. $^7$ Hence the search for catalysts that can operate effectively under lower pressure and temperature environments, or with less pure feedstocks, continues to be important even after many decades of operation of the Haber-Bosch process.

Interest in metal nitrides as heterogeneous catalysts has focussed on hydrotreating reactions such as heteroatom removal and hydrogenations, often using high surface area molybdenum nitride derived from temperature programmed ammonolysis reactions of oxides or other molybdenum compounds. $^2-^4$ However, the high ammonia synthesis activity of some ternary molybdenum nitrides has also attracted significant work on these compounds. Notably Co$_3$Mo$_3$N can outperform commercial iron-based Haber-Bosch catalysts $^5,^6$ with its high activity linked to the possible operation of a Mars-van Krevelen type process. $^7,^8$ We previously reported quaternary compounds in the (Ni,Co)$_2$Mo$_3$N system with good activity for ammonia synthesis. $^9,^{10}$ The principle of electronic tuning through substitution of one of the atoms is used extensively in catalysis, and could be useful in optimising activity of ternary nitrides.

The filled β-Mn structure adopted by Ni$_2$Mo$_3$N contains distorted, corner sharing NMo$_6$ octahedra interpenetrated with a (10,3)-network of nickel atoms (Fig. 1). $^{11,12}$ Complex metal nitrides are often
made via the high temperature ammonolysis of an oxide with the desired metal atom ratio, for example CoMoO₄ produces Co₃Mo₃N. Attempts at this approach in the Ni₂Mo₃N system have led to metal contamination and a much reduced ammonia synthesis activity. The use of a citrate gel to produce an intimate mix of metal oxides that forms the filled β-Mn phase on ammonolysis was successful in producing active Ni₂Mo₃N and (Ni,Co)₂Mo₃N catalysts. Copper substitution of the nickel sites in Ni₂Mo₃N has not previously been reported, but the citrate gel process requires merely a water soluble copper source and was considered likely to produce phases in this space. We show herein that (Ni,Cu)₂Mo₃N phases do exist up to a solubility limit, and determine the effect of copper substitution on ammonia synthesis activity.

![Fig. 1 Unit cell of Ni₂Mo₃N viewed along [100] showing the corner linked NMo₆ octahedra and the interpenetrating (10,3)-network of nickel atoms. Nickel atoms are shaded pink, molybdenum blue and nitrogen green. Diagram prepared using Vesta and structure data from Prior and Battle.](image)

**Experimental**

Nanocrystalline samples of Ni₂ₓMₓMo₃N (M = Cu or Fe) were prepared by a citrate gel route followed by ammonolysis. Stoichiometric amounts of (NH₄)₃Mo₇O₂₄·4H₂O (2 g, Sigma Aldrich, 99.9 %), Ni(NO₃)₂·6H₂O (Sigma Aldrich, 99.9 %) and either Cu(NO₃)₂·4H₂O (Sigma Aldrich, 99%) or Fe(NO₃)₂·6H₂O (supplier, purity) were dissolved in 60 cm³ aqueous HNO₃ (2.6 mol dm⁻³, prepared from Fisher Scientific 70% HNO₃). Citric acid monohydrate (7.935 g, Sigma Aldrich, ACS reagent, ≥99%) was then added to the green solution. Actual quantities are given in ESI, Table S1. The green solution was evaporated in a sand bath at 70 °C to ~10 cm³ of viscous green gel (ESI, Fig. S1), then heated in an ashing furnace (60 °C min⁻¹ heating rate) at 500 °C for 2 h. The resulting grey foam was ground to a powder then heated in flowing ammonia (BOC anhydrous grade, further dried with molecular sieves) at 5 °C min⁻¹ to 900 °C and maintained for 12 h. Once cool, the furnace tube was flushed with N₂ for 30 mins then air was allowed to diffuse in slowly to passivate sample surfaces before the black crystalline product was recovered. Caution: a sand bath was used because citrate gels can occasionally ignite; the unpassivated metal nitrides are highly pyrophoric.
The powder X-ray diffraction (PXRD) patterns of the samples were collected using a Bruker D2 Phaser X-ray diffractometer with Cu-Kα radiation. The Rigaku PDXL2 package and the PDF database were used for pattern matching. Rietveld refinement was carried out using GSAS and structure models from ICSD. Scanning electron microscopy (SEM) was conducted using a Philips XL-30 ESEM (20 kV accelerating voltage) with a ThermoFisher Ultradyne energy dispersive spectroscopy (EDS) detector and Noran System 7 data processing. Transmission electron microscopy (TEM) was performed on Hitachi H7000 with accelerating voltage of 75 kV with samples prepared by ultrasound dispersal in propan-2-ol and deposition on carbon-coated copper grids. Thermogravimetric analysis (TGA) used a Netzsch TG 209 F1 Libra instrument with a heating rate of 10 °C min⁻¹ under 40 ml min⁻¹ of 50% oxygen and 50% argon. The surface areas of the samples were calculated by applying the Brunauer, Emmett and Teller (BET) method to nitrogen physisorption isotherms measured at 77 K using a Micromeritics Tristar II and samples previously degassed overnight at 120 °C.

The catalytic performance of the obtained catalysts in ammonia synthesis was evaluated by a method similar to that described previously. The catalyst (0.15 g) was loaded in the centre of a 4.2 mm internal diameter silica tube and held in a plug against a sinter using silica wool, then placed in a tube furnace. 13 ml min⁻¹ 75% Hydrogen/ 25% Nitrogen gas mixture (BOC, H₂ 99.998 %, N₂ 99.995 %) was passed through the tube and it was heated to 700 °C for 2 hours to activate the catalyst followed by lower temperature steps to test ammonia production. On reaching the reaction temperature, the vent gas from the reactor was bubbled through a dilute solution of sulfuric acid (0.001 mol dm⁻³, 100 ml) at ambient temperature and the rate of ammonia production was calculated from the conductivity decrease of the sulfuric acid solution with time (Orion Star A212 Portable Conductivity Meter). A linear decrease in conductivity versus time was observed during the measurement using H₂/N₂ as the reactant gas. Detailed calculations of ammonia production rates is given in ESI (Note S1).

Results and Discussion

The advantage of citrate gel synthesis to produce solid solutions in the (Ni,M)_2Mo₃N (M = Cu or Fe) system is that the 3d metals are mixed in solution, so could be well distributed in the products. A range of compositions was made starting with Ni₃Mo₄N and replacing nickel with copper or iron. Previously in the (Ni,Co)₂Mo₃N system we fired samples at 700 °C and found that, in addition to the filled β-manganese type phases, they contained a trace of a rocksalt-type impurity that resembled γ-Mo₂N. The filled β-manganese structure has been discussed in more detail elsewhere. Firing Ni₃Mo₃N samples at temperatures between 600 and 1000 °C the samples produced below 800 °C were found to contain the rocksalt-type phase (ESI, Fig. S2), but at 900 or 1000 °C the phase pure filled β-manganese material was produced. The lattice parameters increased slightly with temperature (ESI, Fig. S3). Hence the (Ni,M)₂Mo₃N (M = Cu or Fe) used herein were produced at 900 °C.

Figure 2 shows PXRD patterns for a series of (Ni,Cu)₂Mo₃N compositions produced at 900 °C. Compositions from Ni₃Mo₃N to Ni₁.₇Cu₀.₃Mo₃N were identified as the single phase filled β-manganese structure (space group P4₃₃2). However, as the copper content was increased further, γ-Mo₂N and copper metal appeared in the patterns and grew in intensity. The lattice parameter variation obtained from Rietveld refinement of the diffraction data (ESI, Figs. S4-S12) is shown in Fig. 2. The lattice parameters show a Vegard trend in the single phase compositions, suggesting a good
distribution of copper in the solid solutions. The values increase with copper content as expected from the larger ionic radius of Cu$^{+2}$ (0.73 Å) compared with Ni$^{+2}$ (0.69 Å). The lattice parameter of the filled β-manganese phase is then constant for overall compositions from Ni$_{1.7}$Cu$_{0.3}$Mo$_3$N to Ni$_{1.25}$Cu$_{0.75}$Mo$_3$N. Hence no further copper had been introduced into the Ni$_2$Mo$_3$N structure and the limit of solubility of Cu in Ni$_2$Mo$_3$N is around the Ni$_{1.7}$Cu$_{0.3}$Mo$_3$N composition.

Fig. 2 Left: Powder XRD patterns for Ni$_{2-x}$Cu$_x$Mo$_3$N (overall composition $x = 0$ to 0.75) synthesized by firing the dry powder at 900 °C for 12 hours under NH$_3$ gas, where $x = 0.0$ (a), 0.1 (b), 0.2 (c), 0.3 (d), 0.4 (e), 0.5 (f), 0.6 (g), 0.7 (h) or 0.75 (i). Asterisks indicate the Mo$_2$N reflections and black circles show reflections of Cu metal. Right: The lattice parameter of the filled β-manganese phase versus the concentration of Cu in Ni$_{2-x}$Cu$_x$Mo$_3$N, with lines added as a guide to the eye.

EDX showed that the Ni:Mo ratio of the Ni$_2$Mo$_3$N sample to be 36.4:63.6 (expected 40:60 for Ni$_2$Mo$_3$N). As copper was substituted in the copper contents were measured a little below the nominal compositions, but with an increasing trend (ESI, Table S2). Note there is some overlap between the nickel and copper EDX peaks so there is a significant uncertainty in the analysis of these small amounts of copper. The composition was also tested by TGA (Fig. 3), with the samples heated in an oxygen-rich gas mixture to convert it to oxides as shown in Equation 1. The mass was unchanged up to ~330 °C, above which the oxidation of the material was observed with a resulting mass increase. This occurred in three distinct steps, with an overall mass increase of 38.30 % observed by 550 °C for Ni$_2$Mo$_3$N. No further mass increases were observed above 650 °C, but mass was lost sharply above 750 °C. This can be attributed to the sublimation of MoO$_3$. The mass gains for the Cu-doped compositions (ESI, Table S3) were consistent with expected values. As more Cu is introduced to the lattice the oxidation temperature decreases (Fig. 3).
Fig. 3 TGA curves (oxygen-rich environment) showing mass gain % of the Ni$_2$Mo$_3$N, Ni$_{1.7}$Cu$_{0.3}$Mo$_3$N, Ni$_{1.8}$Cu$_{0.2}$Mo$_3$N and Ni$_{1.9}$Cu$_{0.1}$Mo$_3$N samples prepared at 900 °C with heating rate of 5 °C/min under ammonia.

\[
\text{Ni}_{1.8}\text{Cu}_{0.2}\text{Mo}_3\text{N} + \frac{11}{2} \text{O}_2 \rightarrow 1.8\text{NiO} + 0.2\text{CuO} + 3\text{MoO}_3 + \frac{1}{2} \text{N}_2 \quad \text{Equation 1}
\]

Not only the crystalline structure and the chemical composition are important to understand the catalytic properties, the morphology of the catalyst can also be significant. Fig. 4 shows typical images of a Ni$_{1.7}$Cu$_{0.3}$Mo$_3$N sample obtained at 900 °C. An open structure of interconnected, rounded nanoparticles with sizes around 200 nm were found, which is in a good correlation with the average crystallite size of 140 nm calculated using an adaptation of the Scherrer equation incorporated into the Rietveld fitting package.\textsuperscript{15}

Fig. 4 TEM (left) and SEM (right) images of Ni$_{1.7}$Cu$_{0.3}$Mo$_3$N obtained from the citrate-gel derived metal oxide precursors heated in ammonia at 900 °C for 12 hours.

The ammonia synthesis rates measured for the (Ni,Cu)$_2$Mo$_3$N samples are summarised in Table 1, alongside the BET surface areas and the nitrogen contents measured from combustion analysis (expected wt. % N is 3.3%). It is noteworthy that the rate of ammonia synthesis reduces systematically with copper doping. The need to produce these samples at a relatively high
temperature of 900 °C has resulted in quite low surface areas, and the surface area is reducing with increasing copper content. This can be attributed partly to crystallite growth and partly to agglomeration, both occurring through sintering at this high reaction temperature, but the surface areas are fairly typical for this type of material. However, the surface area of Mo based nitride materials can be strongly influenced by synthesis factors such as space velocities of NH₃ and temperature ramp rates, and achieving higher surface areas should be a future priority.

Table 1 BET surface areas, elemental analysis, ammonia synthesis rate and specific activity of (Ni,Cu)₂Mo₃N

| Catalyst   | Surface area (m² g⁻¹) | Crystallite size (nm) | Nitrogen content (wt.%) | Ammonia synthesis rate at 500 °C (µmol h⁻¹ g⁻¹) | Specific activity at 500 °C (µmol h⁻¹ m⁻²) |
|------------|-----------------------|-----------------------|-------------------------|-----------------------------------------------|------------------------------------------|
| Ni₂Mo₃N    | 2.2                   | 113(2)                | 3.4                     | 272(5)                                        | 124(2)                                   |
| Ni₁₃Cu₀₁Mo₃N | 2.1                   | 119(3)                | 3.6                     | 252(2)                                        | 120(1)                                   |
| Ni₁₈Cu₀₂Mo₃N | 1.8                   | 126(5)                | 3.6                     | 231(2)                                        | 128(1)                                   |
| Ni₂₇Cu₀₃Mo₃N | 1.6                   | 142(11)               | 3.6                     | 179(6)                                        | 112(4)                                   |

a Conductivity profiles used to calculate these rates are shown in ESI (Fig. S13), b calculated values are 3.3%.

Substituting iron into the Ni₂Mo₃N, the synthesis proceeded in the same way as for copper except that a brown colour developed as the sol was concentrated (ESI, Fig. S1). After firing in ammonia at 900 °C the PXRD showed phase pure filled β-manganese-structured samples with iron contents up to NiFeMo₃N (Fig. 5). An increasing linear trend in lattice parameter with iron content indicates the formation of well distributed solid solutions and is consistent with the larger ionic radius of Fe²⁺ (0.78 Å) compared with Ni²⁺ (0.69 Å). Increasing the iron content further molybdenum metal and Fe₂N impurities were observed, although in contrast with the (Ni,Cu)₂Mo₃N solid solutions the lattice parameter of the (Ni,Fe)₂Mo₃N continued to increase, suggesting that there is no strong barrier to forming solid solutions in this more iron-rich region of the phase space. Based on the PXRD data the crystallite size of (Ni,Fe)₂Mo₃N decreases with Fe content, and the BET surface areas increased with iron content (Table 2). The nitrogen content of the samples is approximately consistent with the calculated values. The TGA data during oxidation (ESI, Fig. S14) showed the expected mass gains, with thermal stability decreasing with concentration of the second metal, similarly to the (Ni,Cu)₂Mo₃N system. Electron microscopy (ESI, Fig. S15) showed similar aggregates of nanoparticles to those of (Ni,Cu)₂Mo₃N.
Fig. 5 Left: magnified PXRD patterns of Ni$_{2-x}$Fe$_x$Mo$_3$N, with asterisks indicating Mo metal reflections and black circles the reflections of Fe$_2$N. A wider 2θ range is shown in ESI (Fig. S1). Right: Lattice parameters obtained by Rietveld refinement (ESI, Figs. S17-S24 and Table S6) versus the concentration of Fe in the (Ni,Fe)$_2$Mo$_3$N systems.

Table 2 BET surface areas, elemental analysis, ammonia synthesis rate and specific activity of (Ni,Fe)$_2$Mo$_3$N.

| Catalyst      | Surface area (m$^2$ g$^{-1}$) | Crystallite size (nm) | Nitrogen content (wt.%)$^a$ | Ammonia synthesis rate at 500 °C (µmol h$^{-1}$ g$^{-1}$)$^b$ | Specific activity at 500 °C (µmol h$^{-1}$ m$^{-2}$) |
|---------------|-------------------------------|-----------------------|-----------------------------|------------------------------------------------------------|-----------------------------------------------------|
| Ni$_2$Mo$_3$N | 2.2                           | 113(2)                | 3.4                         | 272(5)                                                     | 124(2)                                               |
| Ni$_{1.8}$Fe$_{0.2}$Mo$_3$N | 2.4                           | 142(12)               | 3.5                         | 311(2)                                                     | 130(1)                                               |
| Ni$_{1.5}$Fe$_{0.5}$Mo$_3$N | 2.5                           | 119(10)               | 3.5                         | 322(2)                                                     | 129(1)                                               |
| Ni$_{1.2}$Fe$_{0.8}$Mo$_3$N | 2.7                           | 114(1)                | 3.6                         | 337(4)                                                     | 125(1)                                               |
| Ni$_{1.1}$Fe$_{0.9}$Mo$_3$N | 2.8                           | 103(2)                | 3.6                         | 354(4)                                                     | 126(2)                                               |

$^a$calculated values are 3.3-3.4%; $^b$Conductivity profiles used to calculate these rates are shown in ESI (Fig. S25).

The ammonia synthesis rates of the (Ni,Fe)$_2$Mo$_3$N catalysts increase with iron addition throughout the range of compositions where a single phase is formed, from 272 µmol h$^{-1}$ g$^{-1}$ with Ni$_2$Mo$_3$N to 354 µmol h$^{-1}$ g$^{-1}$ with Ni$_{1.1}$Fe$_{0.9}$Mo$_3$N (Table 2). However, this increase appears to be linked to the increasing surface area with temperature, as the activities normalised by surface area have similar values.

Molybdenum nitrides have been reported as active catalysts for ammonia synthesis at 400 °C, and their catalytic performance were significantly improved by the addition of a second transition metal such as Ni, Co or Fe. Kojima and Aika showed that cobalt molybdenum nitride promoted with Cs can synthesise ammonia at double the efficiency of commercial catalysts, with reported activities of 652 µmol g$^{-1}$ h$^{-1}$ and 986 µmol g$^{-1}$ h$^{-1}$. The high activity of the Co$_3$Mo$_3$N has been linked to the fact that CoMo alloy has an average nitrogen adsorption energy close to the optimal nitrogen binding energy for catalytic performance, and to lattice nitrogen activity. Activities for Ni$_2$Mo$_3$N and doped analogues are typically lower despite the structural and electronic similarities.
between these two systems, and that may be linked at least partly to the lack of lattice nitrogen mobility exhibited by this compound. Activities observed here are significantly higher than those observed in many other studies of this material and were seen to increase with iron doping, although our activities were measured at 500 °C due to a relatively low activity observed at 400 °C (ESI, Fig. S26). Comparison of PXRD patterns collected before and after catalysis (ESI, Figs. S27 and S28) shows no appreciable change to the structures throughout the series of samples. The nitrogen contents of the catalysts post-catalysis (Tables 1 and 2) also remain consistent with the stoichiometric compositions. In addition to the linear conductivity changes observed in the catalysis, this suggests that the catalysts are stable under the reaction conditions used.

It has previously been noted that Ni$_2$Mo$_3$N can be synthesised from oxides using hydrogen-nitrogen mix, a key advantage compared with Co$_3$Mo$_3$N as it means the ammonia synthesis catalyst can be produced in situ, not requiring an ammonia firing step before use. The citrate gel process led to a mixture of α-NiMoO$_4$ (JCPDS card no. 33-0948) and MoO$_3$ (JCPDS card no. 35-0609), which is consistent with the previously published observations of Alconchel et al. (Fig. 6). These were also tried in catalysis under the same conditions as the metal nitrides described above, again resulting in a steady rate of ammonia production. Interestingly these materials exhibited significantly higher mass-normalised ammonia synthesis rates (Table 3). The surface areas were found to be higher than in the pre-formed nitrides both before and after catalysis (Table 3), and post-catalysis the PXRD patterns and compositions were very similar to those of the pre-formed nitrides (Fig. 6). We infer from this that the transformation to the filled β-manganese type nitride phases proceeds smoothly under the catalyst activation conditions (700 °C under 75% H$_2$/25% N$_2$ yielding a higher surface areas that is advantageous in the catalytic process. Use of ammonia in the synthesis at this temperature yields a rocksalt-type impurity. The activity has not scaled linearly with surface area, and these catalysts have lower area-normalised activities than the pre-formed nitrides (Tables 1-3), but nonetheless the increase supports the idea that increasing surface area could lead to higher overall activity levels in these materials.
Fig. 6 PXRD patterns of catalysts after (a) and before (b) use in ammonia synthesis. The precursor compositions were Ni$_2$Mo$_3$O$_x$ (top), Ni$_{1.7}$Cu$_{0.3}$Mo$_3$O$_x$ (centre) and Ni$_{1.2}$Fe$_{0.8}$Mo$_3$O$_x$ (bottom).
Table 3 Surface areas, composition post-catalysis, ammonia synthesis rate and specific activity of metal oxide catalysts used directly in ammonia synthesis.

| Catalyst          | Surface area (m² g⁻¹) | Nitrogen content (wt %) post-catalysis | Ammonia synthesis rate at 500 °C (µmol g⁻¹ h⁻¹)ᵇ | Specific activity at 500 °C (µmol h⁻¹ m⁻²)ᶜ |
|-------------------|------------------------|----------------------------------------|--------------------------------------------------|---------------------------------------------|
| Ni₂Mo₃Oₓ          | 10                     | 3.1                                    | 466(18)                                          | 78(3)                                       |
| Ni₁.7Cu₁.3Mo₃Oₓ   | 6.6                    | 3.3                                    | 368(9)                                           | 77(2)                                       |
| Ni₁.2Fe₀.8Mo₃Oₓ   | 6.2                    | 3.2                                    | 426(4)                                           | 82(1)                                       |

ᵃ calculated values are 3.3-3.4%;ᵇ Conductivity profiles used to calculate these rates are shown in ESI (Fig. S29);ᶜ Calculation based on the surface area of the nitride phase post-catalysis.

Conclusions

Filled β-manganese type materials with composition Ni₂Mo₃N and with some Ni sites replaced by Cu or Fe were prepared by a citrate gel route. In both cases there was a limit to the amount of the second metal that could be substituted under these conditions, in the copper case that seemed to be due to a solubility limit of copper in the host phase. Iron doping increased the activity of the phase as a catalyst for ammonia synthesis at 500 °C and copper decreased that activity, but in both cases surface area changes rather than any electronic differences appear to be the main contributor to the observed change. The oxide intermediates could be converted to the nitride phases during the catalyst activation, avoiding the need to fire in ammonia. The in situ transformation of the oxides led to higher catalyst surface areas and hence to higher activity.

Acknowledgements

The authors thank the government of Saudi Arabia for a scholarship to SAS and EPSRC for support of SL under EP/L02537X/1.

References

1. Y. Nishibayashi, Inorg. Chem., 2015, 54, 9234–9247.
2. J. S. J. Hargreaves, Coord. Chem. Rev., 2013, 257, 2015–2031.
3. M. Nagai, Appl. Catal. A Gen., 2007, 322, 178–190.
4. E. Furimsky, Appl. Catal. A Gen., 2003, 240, 1–28.
5. R. Kojima and K. I. Aika, Appl. Catal. A Gen., 2001, 218, 121–128.
6. C. J. H. Jacobsen, Chem. Commun., 2000, 1057–1058.
7. C. D. Zeinalipour-Yazdi, J. S. J. Hargreaves and C. R. A. Catlow, J. Phys. Chem. C, 2015, 119, 28368–28376.
8. S. M. Hunter, D. H. Gregory, J. S. J. Hargreaves, M. Richard, D. Duprez and N. Bion, ACS Catal., 2013, 3, 1719–1725.
9. N. Bion, F. Can, J. Cook, J. S. J. Hargreaves, A. L. Hector, W. Leveson, A. R. McFarlane, M. Richard and K. Sardar, Appl. Catal. A Gen., 2015, 504, 44–50.
S. Al Sobhi, N. Bion, J. S. J. Hargreaves, A. L. Hector, S. Laassiri, W. Levason, A. W. Lodge, A. R. Mcfarlane and C. Ritter, Mater. Res. Bull., 2019, 118, 110519 (5 pages).

P. S. Herle, M. S. Hegde, K. Sooryanarayana, T. N. Guru Row and G. N. Subbanna, Inorg. Chem., 1998, 37, 4128–4130.

T. J. Prior and P. D. Battle, J. Solid State Chem., 2003, 172, 138–147.

J. S. J. Hargreaves and D. Mckay, J. Mol. Catal. A Chem., 2009, 305, 125–129.

PDF-2 (Powder Diffrr. File), 2012 release, Int. Cent. Diffrr. Data, Swart. PA.

A. C. Larson and R. B. Von Dreele, Gen. Struct. Anal. Syst. (GSAS), The Regents Univ. Calif., 1985, 748, LAUR 86-748.

B. H. Toby, J. Appl. Crystallogr., 2001, 34, 210–213.

Inorg. Cryst. Struct. Database (ICSD, Fiz Karlsruhe) accessed via EPSRC-funded Natl. Chem. Database Serv. hosted by R. Soc. Chem.

S. Brunauer, P. H. Emmett and E. Teller, J. Am. Chem. Soc., 1938, 60, 309–319.

D. McKay, PhD thesis, University of Glasgow, 2008.

R. D. Shannon, Acta Crystallogr. Sect. A, 1976, 32, 751–767.

S. Alconchel, F. Sapiña, D. Beltrán and A. Beltrán, J. Mater. Chem., 1999, 9, 749–755.

A. Srifa, K. Okura, T. Okanishi, H. Muroyama, T. Matsui and K. Eguchi, 2016, 7495–7504.

L. Volpe and M. Boudart, J. Solid State Chem., 1985, 59, 332–347.

J. G. Choi, J. R. Brenner, C. W. Colling, B. G. Demczyk, J. L. Dunning and L. T. Thompson, Catal. Today, 1992, 15, 201–222.

D. W. Kim, D. K. Lee and S. K. Ihm, Korean J. Chem. Eng., 2002, 19, 587–592.

J. G. Choi, R. L. Curl and L. T. Thompson, J. Catal., 1994, 146, 218–227.

L. Volpe and M. Boudart, J. Phys. Chem., 1986, 90, 4874–4877.

S. T. Oyama, Catal. Today, 1992, 15, 179–200.

D. Mckay, D. H. Gregory, J. S. J. Hargreaves, S. M. Hunter and X. Sun, Chem. Commun., 2007, 7, 3051–3053.

D. Mckay, J. S. J. Hargreaves, J. L. Rico, J. L. Rivera and X. L. Sun, J. Solid State Chem., 2008, 181, 325–333.

I. Alshibane, A. Daisley, J. S. J. Hargreaves, A. L. Hector, S. Laassiri, J. L. Rico and R. I. Smith, ACS Sustain. Chem. Eng., 2017, 5, 9214–9222.

R. Kojima and K. I. Aika, Appl. Catal. A Gen., 2001, 219, 141–147.
33 R. Kojima and K. Aika, *Appl. Catal. A Gen.*, 2001, **215**, 149–160.

34 C. J. H. Jacobsen, S. Dahl, B. G. S. Clausen, S. Bahn, A. Logadottir and J. K. Nørskov, *J. Am. Chem. Soc.*, 2001, **123**, 8404–8405.

35 A. Boisen, S. Dahl and C. J. H. Jacobsen, *J. Catal.*, 2002, **186**, 180–186.

36 A. Daisley and J. S. J. Hargreaves, *J. Energy Chem.*, 2019, **39**, 170–175.