# Energy & Environmental Science

## COMMUNICATION

## **RSC**Publishing

View Article Online View Journal | View Issue

**Cite this:** *Energy Environ. Sci.*, 2013, **6**, 3595

Received 1st September 2013 Accepted 27th September 2013

DOI: 10.1039/c3ee42951b

#### www.rsc.org/ees

Photocatalytic oxygen evolution using BaNbO<sub>2</sub>N modified with cobalt oxide under photoexcitation up to 740 nm<sup>+</sup>

Takashi Hisatomi,<sup>a</sup> Chisato Katayama,<sup>bc</sup> Yosuke Moriya,<sup>a</sup> Tsutomu Minegishi,<sup>a</sup> Masao Katayama,<sup>a</sup> Hiroshi Nishiyama,<sup>d</sup> Taro Yamada<sup>d</sup> and Kazunari Domen<sup>\*a</sup>

BaNbO<sub>2</sub>N was activated for photocatalytic sacrificial water oxidation and reduction by modifying the starting material for nitridation and loading appropriate cocatalysts. Addition of BaCO<sub>3</sub> to the Ba<sub>5</sub>Nb<sub>4</sub>O<sub>15</sub> precursor improved the crystallinity and uniformity of BaNbO<sub>2</sub>N as a nitridation product, leading to higher oxygen evolution activity. BaNbO<sub>2</sub>N generated oxygen from an aqueous AgNO<sub>3</sub> solution under illumination up to 740 nm.

Water splitting on semiconductor photocatalysts has received much attention for large-scale production of renewable hydrogen from solar energy and water.1-8 The hydrogen thus produced is storable and can be efficiently converted into electricity or heat at any desired time using fuel cells. Moreover, hydrogen can be employed to produce feedstock in the chemical industry.<sup>2</sup> Since visible and infrared rays account for the greater portion of solar energy, it is necessary to extend the wavelength region available for photocatalytic water splitting by developing narrow band gap semiconductors. In fact, photocatalytic materials active for water splitting under irradiation up to wavelengths of 600 nm or even longer are needed to achieve a high solar energy conversion efficiency at a reasonable quantum efficiency.<sup>2</sup> However, photocatalysis is triggered by photoexcited electrons and holes, which relax to the bottom of the conduction band and the top of the valence band, respectively. Thus, it

#### Broader context

Water splitting on semiconductor photocatalysts has received much attention for the large-scale production of renewable hydrogen from solar energy and water. Photocatalysts should be active for water splitting under irradiation up to 600 nm or even longer wavelengths in order to achieve a high solar energy conversion efficiency at a reasonable quantum efficiency. In this regard, BaNbO2N is a promising candidate because of its small band gap energy of 1.7 eV, allowing light absorption up to 740 nm. The present study demonstrates that BaNbO2N can be activated for photocatalytic water oxidation and reduction in the presence of sacrificial reagents by modifying the starting material for nitridation and loading appropriate cocatalysts. BaNbO2N generates oxygen from an aqueous AgNO3 solution under illumination up to 740 nm, the longest wavelength ever reported for (oxy)nitride photocatalysts. In addition, BaNbO2N has a conduction band edge that is more negative than the hydrogen evolution potential. This means that BaNbO2N electrodes could drive photoelectrochemical water splitting with a smaller applied voltage than other visible-light-driven oxide photoanodes. The promising semiconducting properties of BaNbO2N in solar water splitting and the critical roles of the cocatalysts in boosting the photocatalytic activity will be key to achieving practical solar fuel production.

is more challenging to drive photocatalytic water splitting as the band gap energy decreases.

Perovskite-type and relevant semiconductors modified with appropriate cocatalysts have produced a variety of benchmarks in the field of photocatalytic water splitting. La-doped NaTaO<sub>3</sub> shows excellent quantum efficiencies for overall water splitting under UV light (56% at 270 nm)<sup>9</sup> and Rh-doped SrTiO<sub>3</sub> is used as a hydrogen evolution photocatalyst in *Z*-scheme water splitting under visible light.<sup>10</sup> Non-oxide perovskite-type materials also exhibit promising activity in water splitting. Porous singlecrystalline LaTiO<sub>2</sub>N modified with CoO<sub>x</sub> as an oxygen evolution cocatalyst exhibited a quantum efficiency of 27% at 440 nm for an oxygen evolution reaction in the presence of Ag<sup>+</sup> cations as sacrificial electron acceptors.<sup>11</sup> BaTaO<sub>2</sub>N and its solid solutions with BaZrO<sub>3</sub> have attracted growing interest owing to its response to long-wavelength photons up to 660 nm and the ability to generate both hydrogen and oxygen in the presence of

<sup>&</sup>lt;sup>a</sup>Department of Chemical System Engineering, School of Engineering, The University of Tokyo, 7-3-1 Hongo, Bunkyo-ku, 113-8656 Tokyo, Japan. E-mail: domen@chemsys.t. u-tokyo.ac.jp; Fax: +81 3 5841 8838; Tel: +81 3 5841 1148

<sup>&</sup>lt;sup>b</sup>Japan Technological Research Association of Artificial Photosynthetic Chemical Process (ARPChem), 5-1-5 Kashiwanoha, Kashiwa-shi, 277-8589 Chiba, Japan

<sup>&</sup>lt;sup>c</sup>FUJIFILM Corporation, 577, Ushijima, Kaisei-Machi, Ashigarakami-gun, 258-8577 Kanagawa, Japan

<sup>&</sup>lt;sup>d</sup>Department of Chemical System Engineering, School of Engineering, The University of Tokyo, 5-1-5 Kashiwanoha, Kashiwa-shi, 277-8589 Chiba, Japan

<sup>†</sup> Electronic supplementary information (ESI) available: Detailed experimental, SEM images, XRD patterns, DRS, hydrogen evolution reaction, and *J*–*E* curves of the samples. See DOI: 10.1039/c3ee42951b

sacrificial reagents.<sup>12</sup> In photoelectrochemical applications, in which necessary electrodes can be readily fabricated *via* electrophoretic deposition<sup>13</sup> and particle transfer<sup>14</sup> using photocatalyst powder, photoanodes of SrNbO<sub>2</sub>N decorated with IrO<sub>2</sub> can utilize photons up to 700 nm for water oxidation.<sup>13</sup> The various physical and semiconducting properties of perovskitetype materials, a consequence of their adjustable compositions and structural stability, allow flexible synthetic routes and establish the potential of perovskite-type photocatalysts for use in versatile approaches to solar energy conversion. Spectroscopic and computational studies are also active to understand band structures of perovskite-type photocatalysts.<sup>15-17</sup>

Nb-based perovskite-type oxynitrides have smaller band gap energies than the corresponding Ta-based oxynitrides because of the higher electronegativity of Nb.<sup>18</sup> BaNbO<sub>2</sub>N is of particular interest for solar energy conversion because of its narrow band gap of 1.7 eV although it was found to be inactive for both photocatalytic oxygen and hydrogen evolution reactions.<sup>19</sup> In the present study, however, we successfully activated BaNbO<sub>2</sub>N for both of these reactions by modifying the preparation conditions of the starting material for nitridation and loading suitable cocatalysts such as  $CoO_x$  (ref. 11 and 20) and Pt. Here, we report the effect of the preparation conditions for BaNbO<sub>2</sub>N on its physical properties and photocatalytic activity for O<sub>2</sub> evolution.

BaNbO<sub>2</sub>N was prepared by nitriding Ba<sub>5</sub>Nb<sub>4</sub>O<sub>15</sub> under an NH<sub>3</sub> flow. Ba<sub>5</sub>Nb<sub>4</sub>O<sub>15</sub> was prepared by calcining a mixture of BaCO<sub>3</sub> and Nb<sub>2</sub>O<sub>5</sub>, with NaCl added as a flux at 1173 K unless otherwise noted. After a slow cool-down to room temperature, the oxide sample was rinsed with copious amounts of distilled water to eliminate the NaCl flux. BaCO3 was added to Ba5Nb4O15 before nitridation to suppress the possible generation of niobium oxynitride.19 The precursor was nitrided at 1202 K for 15 h under an NH<sub>3</sub> flow of 200 mL min<sup>-1</sup>. Excessive Ba species generated during the nitridation were dissolved by agua regia. Photocatalytic reactions were carried out using a closed-circulation system. BaNbO2N was modified with nanoparticles of  $CoO_x$  and Pt, which served as cocatalysts for oxygen and hydrogen evolution, respectively. Aqueous solutions of AgNO<sub>3</sub> and CH<sub>3</sub>OH were used for sacrificial oxygen and hydrogen evolution reactions, respectively. The reaction vessel was evacuated to remove air before irradiation. The light source used was a 300 W Xe lamp equipped with cutoff filters. Evolved gases were analysed by gas chromatography. Other experimental details are described in the ESI.†

Fig. 1A shows X-ray diffraction (XRD) patterns of samples at different preparation stages.  $Ba_5Nb_4O_{15}$  was obtained by calcining the starting materials at 1173 K with a NaCl flux while unreacted  $BaCO_3$  remained as a by-product in the absence of the flux. A mixture of  $Ba_5Nb_4O_{15}$  and  $BaCO_3$  at a Ba/Nb ratio of 2.00 was converted into  $BaNbO_2N$  and by-products originating from excessive Ba-species. XRD-pure  $BaNbO_2N$  was obtained by dissolving the by-products by treatment with aqua regia. On the other hand,  $Ba_5Nb_4O_{15}$  calcined at 1423 K remained partly unreacted when nitrided under the same conditions (data not shown), presumably because of the larger particle size of the oxide. Scanning electron microscopy (SEM) observation revealed



**Fig. 1** (A) XRD patterns and (B) UV-Vis DRS for (a)  $Ba_5Nb_4O_{15}$  prepared by the flux method, (b)  $BaNbO_2N$  nitrided from a mixture of  $Ba_5Nb_4O_{15}$  and  $BaCO_3$  with a Ba/Nb ratio of 2, and (c)  $BaNbO_2N$  after treatment with aqua regia.

that particles of the nitridation product were huge aggregates, several micrometres in size, and consisted of partly fused grains (Fig. S1<sup>†</sup>). After the aqua regia treatment, most of the particles became porous. This is presumably because the excessive Ba species between BaNbO2N domains, which were compressed during the crystallographic conversion from  $Ba_5Nb_4O_{15}$  to BaNbO<sub>2</sub>N, were dissolved. XRD-pure BaNbO<sub>2</sub>N was also obtained at different Ba/Nb ratios between 1.25 and 2.00 after treatment with aqua regia (Fig. S2<sup>+</sup>). Note that NbO<sub>x</sub>N<sub>y</sub> was easily produced as a by-product in our previous work particularly when the Ba/Nb ratio was lower than 2.00.19 The full width at half maximum of the (110) peak at 30.5-30.6° decreased from 0.246 to 0.213, 0.210, and 0.207 as the Ba/Nb ratio increased from 1.25 to 1.50, 1.75, and 2.00, respectively. This suggests that higher Ba/Nb ratios in the precursor oxide could improve the crystallinity and uniformity of the nitridation products.

In the diffuse reflectance spectra (DRS) of the nitridation products, the onset of light absorption characteristics of the band gap excitation of BaNbO2N was observed at around 740 nm (Fig. 1B). Additional light absorption was observed in the longer wavelength region, which could be attributed to defects such as reduced B-site cations (Nb) and anion vacancies.18,19,21 The background absorption was weakened by the aqua regia treatment, suggesting that defect species on the surface were eliminated to some extent. However, considerable background absorption remained even after the acid treatment. This indicated that the bulk of BaNbO<sub>2</sub>N also held reduced Nb species and anion vacancies. Since no diffraction peaks other than those for the perovskite-type material were observed, such defect phases could be expressed nominally as solid solutions of BaNbO<sub>2</sub>N with BaNbO<sub>3</sub> containing Nb<sup>4+</sup> states and/or  $BaNbO_{2-\delta 1}N_{1-\delta 2}$  with anion deficiencies if crystalline. In addition, amorphous defect by-products could contribute to the background absorption although their impact could not be determined unambiguously. The addition of BaCO<sub>3</sub> to Ba<sub>5</sub>Nb<sub>4</sub>O<sub>15</sub> was expected to weaken the light absorption due to defect species, similar to the case of LaTiO2N nitrided from La<sub>2</sub>Ti<sub>2</sub>O<sub>7</sub> with La deficiency.<sup>22</sup> In reality, however, the intensity of the background absorption increased by the addition of BaCO<sub>3</sub> to Ba<sub>5</sub>Nb<sub>4</sub>O<sub>15</sub> (Fig. S3<sup>†</sup>). Moreover, the nitridation of Ba<sub>5</sub>Nb<sub>4</sub>O<sub>15</sub> obtained at higher temperatures than 1173 K tended to result in stronger background light absorption. The generation of defect species was not controllable solely by the Ba/Nb ratio during nitridation, presumably because it was too sensitive to the degree of nitridation from Ba<sub>5</sub>Nb<sub>4</sub>O<sub>15</sub> to BaNbO<sub>2</sub>N, which was in turn governed by both the nitridation conditions and the nature of the precursor oxides.

Fig. 2 shows reaction time courses for photocatalytic oxygen evolution using BaNbO<sub>2</sub>N loaded with the  $CoO_x$  cocatalyst.



The oxygen evolution rates were increased by adding  $BaCO_3$ to  $Ba_5Nb_4O_{15}$  as a starting material for nitridation. The XRD patterns of the samples indicated that the crystallinity and



**Fig. 2** Reaction time courses for oxygen evolution under visible light ( $\lambda > 410$  nm) using 2 wt% CoO<sub>x</sub>-loaded BaNbO<sub>2</sub>N nitrided at a Ba/Nb ratio of (a) 1.25, (b) 1.50, (c) 1.75, and (d) 2.00. Photocatalyst, 0.20 g; solution, 200 mL of 50 mM AgNO<sub>3</sub> aqueous solution containing 0.20 g of La<sub>2</sub>O<sub>3</sub>; and light source, 300 W Xe lamp.



**Fig. 3** Reaction time course for hydrogen evolution under visible light ( $\lambda$  > 410 nm) using 1 wt% Pt-loaded BaNbO<sub>2</sub>N nitrided at a Ba/Nb ratio of 1.75. Photocatalyst, 0.20 g; solution, 200 mL of 80 vol% aqueous CH<sub>3</sub>OH solution; and light source, 300 W Xe lamp.

uniformity of the nitridation products were improved by the addition of BaCO<sub>3</sub> to the precursor. This would improve oxygen evolution rates, because higher crystallinity and uniformity are thought to be favourable for higher photocatalytic activity owing to the lower probability of carrier trapping. Similar results were reported for LaTiO<sub>2</sub>N (ref. 23) and Ta<sub>3</sub>N<sub>5</sub>,<sup>24</sup> where the photocatalyst samples with sharper XRD peaks exhibited higher oxygen evolution activity. On the other hand, the background absorption due to defect species was progressively enhanced. The influence of defect species on photocatalytic oxygen evolution in the presence of sacrificial reagents is still controversial. In the present study, they likely did not contribute to charge trapping and recombination significantly. The present results demonstrate that adding an appropriate amount of BaCO<sub>3</sub> to the precursor oxide could modify the structural properties and improve the photocatalytic activity of BaNbO<sub>2</sub>N. More comprehensive synthesis and analysis of BaNbO2N samples are currently underway.

Fig. 4 shows the dependence of the initial oxygen evolution rate on the cutoff wavelength of the incident light, where BaNbO<sub>2</sub>N nitrided at a Ba/Nb ratio of 1.5 and modified with 2 wt% cobalt oxide was employed. The DRS trace of the sample is also shown for comparison. Photocatalytic oxygen evolution from the AgNO<sub>3</sub> solution was observed up to a cutoff wavelength of 740 nm (0.2 µmol h<sup>-1</sup>) although the oxygen evolution rate decreased as the cutoff wavelength increased. The apparent quantum efficiency of the oxygen evolution reaction was measured to be 0.04% at 640 ± 30 nm. The onset of light absorption reasonably matched the longest wavelength available for photocatalytic oxygen evolution. On the other hand, continuous light absorption by defect species did not contribute to photocatalysis. Thus, it was concluded that water oxidation



**Fig. 4** Dependence of the initial oxygen evolution rate of 2 wt%  $CoO_x$ -loaded BaNbO<sub>2</sub>N on the cutoff wavelength of the incident light. The DRS trace of the BaNbO<sub>2</sub>N used is also shown. Photocatalyst, 0.10 g; solution, 200 mL of 50 mM aqueous AgNO<sub>3</sub> solution containing 0.20 g of La<sub>2</sub>O<sub>3</sub>; and light source, 300 W Xe lamp.

proceeded *via* band gap transitions for the BaNbO<sub>2</sub>N photocatalyst.

## Conclusion

We demonstrated activation of BaNbO<sub>2</sub>N by loading a CoO<sub>x</sub> cocatalyst for photocatalytic O2 evolution under visible light. The apparent quantum efficiency of the reaction using  $CoO_x$ loaded BaNbO<sub>2</sub>N was lower than the value for the porous singlecrystalline LaTiO<sub>2</sub>N,<sup>11</sup> likely because of the smaller band gap energy of BaNbO<sub>2</sub>N (smaller by 0.4 eV) in addition to insufficient optimization of the preparation conditions for the material. Nevertheless, BaNbO2N can utilize photons up to 740 nm for photocatalytic  $O_2$  evolution. This wavelength is the longest ever reported for an (oxy)nitride photocatalyst. Managing longer wavelength photons could help overcome the barrier against effective production of solar fuel via water splitting.2 Moreover, BaNbO<sub>2</sub>N had a band gap straddling the hydrogen and oxygen evolution potentials. This means that overall water splitting and unassisted photoelectrochemical water splitting could be achieved under illumination of red light by improving the semiconducting quality of the BaNbO2N photocatalyst. These findings motivate further studies on refinement of the nitridation and cocatalyst loading conditions,11,20 modifications of oxide precursors,<sup>23,24</sup> and doping<sup>25</sup> to improve the photocatalytic and photoelectrochemical activity of BaNbO2N.

## Acknowledgements

This work was supported by the Artificial Photosynthesis Project of the Ministry of Economy, Trade and Industry (METI) of Japan, a Grant-in-Aid for Specially Promoted Research (no. 23000009) and for Young Scientists (B) (no. 25810112) of the Japan Society for the Promotion of Science (JSPS). JSPS also contributed through the Funding Program for World-Leading Innovative R&D on Science and Technology (FIRST), initiated by the Council for Science and Technology Policy (CSTP).

## Notes and references

- 1 A. Kudo and Y. Miseki, Chem. Soc. Rev., 2009, 38, 253-278.
- 2 K. Maeda and K. Domen, J. Phys. Chem. Lett., 2010, 1, 2655–2661.
- 3 X. Chen, S. Shen, L. Guo and S. S. Mao, *Chem. Rev.*, 2010, **110**, 6503–6570.
- 4 R. Abe, J. Photochem. Photobiol., C, 2010, 11, 179-209.
- 5 K. Maeda, J. Photochem. Photobiol., C, 2011, 12, 237-268.
- 6 T. Hisatomi, T. Minegishi and K. Domen, *Bull. Chem. Soc. Jpn.*, 2012, **85**, 647–655.
- 7 J. Yang, D. Wang, H. Han and C. Li, *Acc. Chem. Res.*, 2013, **46**, 1900–1909.
- 8 F. E. Osterloh, Chem. Soc. Rev., 2013, 42, 2294-2320.
- 9 H. Kato, K. Asakura and A. Kudo, *J. Am. Chem. Soc.*, 2003, **125**, 3082–3089.
- 10 Y. Sasaki, A. Iwase, H. Kato and A. Kudo, *J. Catal.*, 2008, **259**, 133–137.

- 11 F. Zhang, A. Yamakata, K. Maeda, Y. Moriya, T. Takata, J. Kubota, K. Teshima, S. Oishi and K. Domen, J. Am. Chem. Soc., 2012, 134, 8348–8351.
- 12 K. Maeda and K. Domen, Angew. Chem., Int. Ed., 2012, 51, 9865–9869.
- 13 K. Maeda, M. Higashi, B. Siritanaratkul, R. Abe and K. Domen, J. Am. Chem. Soc., 2011, 133, 12334– 12337.
- 14 T. Minegishi, N. Nishimura, J. Kubota and K. Domen, *Chem. Sci.*, 2013, **4**, 1120–1124.
- 15 I. E. Castelli, D. D. Landis, K. S. Thygesen, S. Dahl, I. Chorkendorff, T. F. Jaramillo and K. W. Jacobsen, *Energy Environ. Sci.*, 2012, 5, 9034–9043.
- 16 S. Balaz, S. H. Porter, P. M. Woodward and L. J. Brillson, *Chem. Mater.*, 2013, **25**, 3337–3343.
- 17 P. Reunchan, S. Ouyang, N. Umezawa, H. Xu, Y. Zhang and J. Ye, *J. Mater. Chem. A*, 2013, **1**, 4221–4227.

- 18 Y.-I. Kim, P. M. Woodward, K. Z. Baba-Kishi and C. W. Tai, *Chem. Mater.*, 2004, **16**, 1267–1276.
- 19 B. Siritanaratkul, K. Maeda, T. Hisatomi and K. Domen, *ChemSusChem*, 2011, 4, 74–78.
- 20 S. S. K. Ma, K. Maeda, T. Hisatomi, M. Tabata, A. Kudo and K. Domen, *Chem.-Eur. J.*, 2013, **19**, 7480–7486.
- 21 M. Miyauchi, M. Takashio and H. Tobimatsu, *Langmuir*, 2003, **20**, 232–236.
- 22 T. Moriga, K. Ikeuchi, R. Mashima, D. Aoki and K. Murai, *J. Ceram. Soc. Jpn.*, 2007, **115**, 637–639.
- 23 A. E. Maegli, T. Hisatomi, E. H. Otal, S. Yoon, S. Pokrant, M. Grätzel and A. Weidenkaff, J. Mater. Chem., 2012, 22, 17906–17913.
- 24 S. S. K. Ma, T. Hisatomi, K. Maeda, Y. Moriya and K. Domen, *J. Am. Chem. Soc.*, 2012, **134**, 19993–19996.
- 25 T. Takata and K. Domen, J. Phys. Chem. C, 2009, 113, 19386– 19388.

This journal is © The Royal Society of Chemistry 2013