

Contents lists available at [ScienceDirect](www.sciencedirect.com/science/journal/09259635)

Diamond & Related Materials

journal homepage: www.elsevier.com/locate/diamond

Hydrogen catalytic performance of hybrid $Fe₃O₄/FeS₂/g-C₃N₄$ nanocomposite structures

Majed Alshammari^{a,*}, Sultan Alhassan^a, Khulaif Alshammari^a, Turki Alotaibi^a, Alhulw H. Alshammari^a, Satam Alotibi ^b, Taha Abdel Mohaymen Taha^a, Ali Ismael ^c

^a *Physics Department, College of Science, Jouf University, P.O. Box: 2014, Sakaka, Saudi Arabia*

^b *Department of Physics, College of Science and Humanities in Al-Kharj, Prince Sattam bin Abdulaziz University, Al-Kharj 11942, Saudi Arabia*

(8480 mL/g⋅min).

^c *Physics Department, Lancaster University, Lancaster, LA1 4YB, UK*

ARTICLE INFO *Keywords:* NaBH4 $Fe₃O₄/FeS₂$ nanosheet Hydrogen production Methanolysis ABSTRACT In this work, Fe3O4/FeS2/g-C3N4 nanocomposites were developed for catalytic hydrogen generation from sodium borohydride. X-ray diffraction (XRD), Fourier transform infrared spectroscopy (FTIR), and environmental scanning electron microscopy (ESEM) were used to analyze these nanocomposites. The XRD diffraction peaks of Fe3O4 and FeS2 cubic phase showed an average crystal size of calculation of 15 and 20 nm. ESEM micrographs showed a 2D broken up sheet structure having more edge sites. The BET surface areas for S@g-C₃N₄, 1.0, 2.0, and 3.0 wt% Fe₃O₄/FeS₂ were 40, 109, 137 and 162 m^2/g , respectively. Even though Fe₃O₄/FeS₂ were incorporated into the nanosheet, the pore size was increased from 2.0 to 2.15 nm. S@g-C₃N₄ has an average band gap of 2.60 eV that decreased to 2.30, 2.21 and 2.18 eV at 1.0, 2.0 and 3.0 wt% of FeS₂. In addition, Fe₃O₄/FeS₂/g-C₃N₄ nanosheets showed an emission band at 460 nm. Moreover, the intensity of this band decreased as the content of Fe₃O₄/FeS₂ reached 3.0 wt%. The rate of hydrogen production is accelerated as the percentage of Fe₃O₄/FeS₂

1. Introduction

Recently, improving hydrogen production is a research priority in various research sectors. Production improvement includes many means, including the use of catalysts. Therefore, tailoring the catalyst during the preparation process by controlling the shape, size and surface properties is very important to increase performance [1–[3\]](#page-5-0). Nanosheets have achieved great success when used as hydrogen production catalysts. The structural, electrical, and mechanical properties of carbon nitride (g-C3N4) have drawn much attention in physicochemical studies [[4](#page-6-0),[5](#page-6-0)]. Meanwhile, $g - C_3N_4$ contains carbon and nitrogen that are abundant elements. $g-C_3N_4$ exhibits a prominent UV–Vis absorption peak in the optical wavelength of 260–320 nm. The absorption peak at around 250 nm is caused by the $\pi\text{-}\pi^\ast$ electron transfer for g-C₃N₄ with *s*-triazine rings. While the absorption band located at 320 nm was formed by the transitions of electrons at n-π* that include a lone pair of electrons on nitrogen atoms in g-C₃N₄ [[6](#page-6-0),[7](#page-6-0)]. Moreover, g-C₃N₄ has improved thermal stability for catalytic application because of high strength up to 600 °C.

However, the electroactive surface area and conductivity of $g - C_3N_4$ are both low [8–[10\]](#page-6-0). The key factors limiting effectiveness are low carrier mobility and high bulk recombination.

increased from 1.0 to 3.0 wt%. The sample 3.0 wt% Fe₃O₄/FeS₂ showed the best rate of hydrogen production

The $g - C_3N_4$ nanosheet consists of stacked layers, and for this, it needs to be modified with other nanomaterials to improve its catalytic efficiency. This goal can be achieved by grafting them with different minerals, metal oxides, or metal sulfides. This procedure, in turn, will combine the features of g-C3N4 nanosheets with these nanomaterials in the final composite. The procedure must take into account the grafting ratios and the morphology of the composite as well as the surface characteristics. As a result, electron-hole recombination is suppressed.

The semiconductor nanoparticles of iron sulfide are naturally abundant and non-toxic and thus have more interest [\[10](#page-6-0)–13]. These materials are flexible and can be prepared with different sizes and morphologies. This diverse group of complexes exists in various phases, including greigite, marcasite, pyrrhotite, pyrite and troilite. The energy gap of these materials is 0.8–0.95 eV and possess an absorption coefficient of \sim 10⁵ cm⁻¹. The stoichiometric iron sulfide (FeS₂) exhibits

* Corresponding author. *E-mail address:* malshammary@ju.edu.sa (M. Alshammari).

<https://doi.org/10.1016/j.diamond.2023.110214>

Available online 22 July 2023 0925-9635/© 2023 Published by Elsevier B.V. Received 26 April 2023; Received in revised form 22 June 2023; Accepted 15 July 2023 antiferromagnetic properties at room temperature $[14-16]$ $[14-16]$. The FeS₂ catalyst can act as an electron trap and a site for H_2 production to enhance interfacial charge transfer and separation. For catalytic applications, nanoscale $FeS₂$ with an appropriate surface charge and functionality may be of significance. The development of a precise preparation technique is therefore essential to link nickel sulfides with the g-C3N4 nanosheet. Iron oxide is a commonly used co-catalyst in photocatalytic reactions due to its ability to enhance charge separation and transfer to improve the performance of the catalyst [[17\]](#page-6-0). Combining iron oxide with carbon nitride catalyst can help to improve the efficiency and effectiveness of the catalyst for various chemical reactions. Iron oxide can act as a co-catalyst and help to promote electron transfer and improve the catalytic activity of carbon nitride. This combination can also increase the stability of the catalyst and enhance its performance in various applications such as water purification, organic synthesis, and energy conversion. Therefore, the nanocomposites of $Fe₃O₄/FeS₂/g-$ C3N4 will show high surface area including many catalytic active sites.

Efforts have been made to engineer the structure of carbon nitride by modifying with metals or metal oxides to improve its efficiency as a catalyst. Moreover, the works included other matrices such as graphene and metal organic frameworks (MOF) because of their structural advantages.

Phosphorus was used to modify the surface properties of $g-C_3N_4$ -titanium oxide composites to function as a catalyst for the extraction of hydrogen from sodium borohydride [\[18](#page-6-0)]. These composites showed a significant improvement in stimulating hydrogen production, and the value of the production rate was 14,750 mL min $^{-1}$ g $^{-1}$. Meanwhile, the activation energy of the hydrogen production process was 36 kJ mol $^{-1}$. In the previous work, we dealt directly with the preparation of NiS/g- C_3N_4 sheets using the polycondensation method [\[19](#page-6-0)]. Hydrogen production measurements were completed from the methanolysis reaction of NaBH4. The study succeeded in achieving a hydrogen production rate of 8654 mL/g⋅min. Kottaikalai Ganesan et al. [\[20](#page-6-0)] have conducted a study of phosphorylated silica as a catalyst from NaBH4 methanolysis reaction. The prepared material was incorporated into the hydrogen production process and achieved a rate of 762.4 mL min $^{-1}$ g $^{-1}$. The study completed the calculation of the activation energy for the hydrogen production process and the value was 29.92 kJ mol $^{-1}$. On the other hand, an experimental study appeared for the preparation of g- $C_3N_4/SiO_2/N$ nanocomposites as a catalyst for hydrogen production [[21\]](#page-6-0). Laboratory measurements of hydrogen production rates in the case of catalyst g-C₃N₄/SiO₂/N showed a value of $11{,}400$ mL min $^{-1}$ g $^{-1}$. The activation energy for the self-hydrolysis of NaBH₄ was 33.2 kJ mol $^{\rm -1}$. In another context, the $MoO₃/S@g-C₃N₄$ nanocomposites were prepared at different carbon nitride content [[22\]](#page-6-0). The nanocomposite of $MoO₃/10wt$ %S@g-C3N4 showed the highest rate of hydrogen production (18,421 mL/g⋅min). Chemical reduction was used to prepare highly distributed CoB alloys imbedded on MOF-74-derived graphene nanosheets [\[23](#page-6-0)]. The measurements were conducted on the catalyst in the reaction of methanolysis from NaBH4 and showed a production rate of 7937 mL \rm{min}^{-1} g $^{-1}$. The activation energy for self-hydrolysis of NaBH₄ is calculated to be 38.8 kJ mol $^{-1}$. To the best of our knowledge, there is not any research investigated the preparation of $Fe₃O₄/FeS₂/g-C₃N₄$ hybrid catalyst for hydrogen evolution from NaBH4.

In order to increase the catalytic activity of g -C₃N₄, the current work seeks to identify an acceptable precursor, morphology, exfoliation condition, and synthesis techniques for carbon nitride composites. Therefore, $Fe₃O₄/FeS₂/g-C₃N₄$ sheets were prepared by the direct decomposition/polycondensation of nickel chloride and thiourea at 550 ◦C. Various techniques have been used to clarify the structural properties of nanocomposites, such as XRD, FTIR, and ESEM. A surface area analyzer investigated surface properties such as surface area and porosity of nanosheets. Electronic configuration analysis of the prepared samples was completed by optical absorption as well as emission measurements. Moreover, these nanocomposites have been applied as catalysts in hydrogen production processes from methanolysis reaction of NaBH4. All procedures were carried out at room temperature and the amount of hydrogen was determined using the water displacement technique. Interestingly, the nanosheets achieved high hydrogen production rates, especially for the sample with a concentration of 3 wt% Fe₃O₄/FeS₂.

2. Experimental

Iron trichloride hexahydrate (FeCl3⋅6H2O, 97 %), thiourea (NH2⋅CS. NH₂, 99 %) and sodium borohydride (NaBH₄, 97 %), were obtained from Loba Chemi, Mumbai, India, and were used without further purification. While, The Sigma-Aldrich, Darmstadt, Germany provided the absolute methanol.

For the synthesis of $S@g-C_3N_4$, 10.0 g of thiourea were heated in a porcelain crucible in the muffle furnace at 550 ◦C for 120 min with a heating rate of 3.0[○]/min. After cooling to room temperature, the yellow solid mass in the crucible was ground inside an agate mortar. Thereafter, Fe₃O₄/FeS₂/g-C₃N₄ nanocomposites were prepared by grinding 10 g of thiourea and 1.0, 2.0 and 3.0 wt% of $FeCl₃.6H₂O$ for 30 min. The mixture was then poured into porcelain crucibles that were placed within a muffle furnace and secured with a lid. The polycondensation procedure was completed at 550 ◦C for 2 h with a ramping rate of 3.0 ◦C/ min. After allowing the acquired nanocomposite samples to cool, they were ground.

Structure-related variables including crystal structure, crystal size, and microstrain were examined by X-ray diffraction analyses. The Shimadzu XRD 700 device used a Cu_{ka} wavelength of 1.54056 Å to generate the XRD spectra. The diffraction pattern of the produced nanocomposite was compared to the JCPDS files in the database to determine the crystal structure. A Shimadzu FTIR spectrometer, model 100 Tracer, was used to acquire FTIR measurements. Typically, frequency ranges between 4000 and 399 cm^{-1} were optimized. The energy dispersive X-ray analysis (EDX) accessory, when used in conjunction with the sensitive characterization equipment ESEM, reveals surface morphology and identifies the elemental content of materials. Using a Thermofisher Quattro environmental scanning electron microscope (ESEM), ESEM micrographs and EDX scans were collected. The prepared samples were degassed at 77 K before recording the adsorption-desorption data on the NOVA 4200e surface area analyzer, the samples were exposed to N_2 gas at 77 K. Optical spectroscopy is based on the absorption of photons to excite the electrons from valence to conduction band. The measurements of optical absorption for the nanocomposites were conducted on a Thermo fisher Evo 201 UV–Vis spectrophotometer. The spectra of photoluminescence (PL) were completed at an excitation wavelength of 350 nm on the Cary Eclipse fluorescence spectrometer.

The prepared $Fe₃O₄/FeS₂/g-C₃N₄$ nanosheets were used as a catalyst for hydrogen production from methanolysis reaction of NaBH4. All hydrogenation tests were conducted in a glass reactor with a 250 mL capacity and a glass stopper for closing. In a typical experiment, the desired amount of catalyst (20 mg) was mixed along with the NaBH₄ (25 mg). The mixture was charged into the reactor with 10 mL methanol. Immediately, the self-hydrolysis of NaBH4 started and hydrogen gas released. The reaction was conducted at constant temperature (293 K), and the H_2 volume as a function of time was measured. The water displacement method was applied to measure the volume of H_2 gas. These experiments were repeated 3.0 times with an experimental error of \pm 5.0 %. Finally, all the hydrogen catalytic reactions were performed in the ordinary visible light.

3. Results and discussion

The crystal structure and crystallite size of materials are easily obtained from XRD analysis. [Fig. 1](#page-2-0) displays the XRD data of the fabricated Fe₃O₄/FeS₂/g-C₃N₄ nanocomposite catalysts. The spectrum of S@g-C3N4 nanosheet revealed two major peaks at 13.08◦ and 27.20◦. The minor diffraction peak at 13.08 $^{\circ}$ matches the (100) plane [\[24](#page-6-0),[25\]](#page-6-0). While

the intense peak at 27.20◦ matches the (002) plane that comes from interlayer stacking of $g - C_3N_4$ structure. Moreover, the (100) plane has been attributed toward the inter-planar configuration of aromatic systems, and the crystal plane (002) to the interlayer construction of aromatic systems [[26\]](#page-6-0). The diffraction peak intensity of the planes (100) and (002) has decreased in $Fe₃O₄/FeS₂/g-C₃N₄$ composites with different concentration in comparison to $S@g-C_3N_4$ nanosheet. Whenever compared to the S@g-C3N4 peak position of 27.2◦, at 3.0 wt% of Fe₃O₄/FeS₂ shifted to a higher angle of 27.8°. The main reason for this shift is the condensation of carbon nitride or shrinkage of the layer stacking direction [\[27](#page-6-0)]. In addition, the thickness of carbon nitride layers is reduced in accordance with the shift of peak position [\[28\]](#page-6-0).

The cubic crystal phase of $Fe₃O₄$ showed diffraction peaks at 30.36, 35.78, 37.20, 43.53, 47.52, 54.02, 57.42 and 63.01◦. The reflections of these peaks are (220), (311), (222), (400), (331), (422), (511) and (440), respectively [\[29](#page-6-0)]. The diffraction angles 33.2, 40.8, 47.50, 56.39 and 64.19 $^{\circ}$ are related to the FeS₂ reflections (200), (211), (220), (311), and (321), respectively [\[30](#page-6-0)]. The broadening of these diffraction peaks nearly the same whatever the content of $FeS₂$ increased up to 3.0 wt%. Furthermore, the Scherer equation suggests the relationship between the crystallite size (D) and the diffraction peak widening (β) as follows [31–[33\]](#page-6-0);

$$
D = \frac{0.9\lambda}{\beta cos \theta} \tag{1}
$$

For the diffraction peak (002), the values of β increased upon the increase of FeS₂ wt%. This result is indicative of the small $S@g-C_3N₄$ crystalline domains. The average crystal size determined by the Scherer equation for Fe3O4 and FeS2 to be 15 and 20 nm. Furthermore, the Fe3O4/FeS2 integration into carbon nitride revealed a shift of (002) peak that indicate the preparation of nanocomposite sheets.

FTIR profiles of Fe₃O₄/FeS₂/g-C₃N₄ nanosheets were collected and displayed in Fig. 2. Analysis of the spectra in the 3100–3300 cm⁻¹ region, shows stretching vibrations of N–H and H_2O [\[34](#page-6-0)]. The vibrations gion, shows successing vibrations of N—n and H_2O [34]. The vibrations of C—N bond occur at 2170 cm⁻¹ [\[35](#page-6-0)]. The peaks in the region 1229–1628 cm^{-1} are assigned to CN heterocycles at stretching modes [[36\]](#page-6-0). The absorption bands detected at 1010 and 1132 cm^{-1} are due to stretching vibrations of C–O and C-OH [\[37](#page-6-0)].

The spectra of all samples reveal a strong band at 804 cm^{-1} for heptazine or triazine blocks $[38]$ $[38]$. Partly, the bonds of C-N(-C)-C

generate the absorption band at 1311 $\rm cm^{-1}.$ Moreover, the C—N $\rm (sp^3)$ bond vibrations show an absorption peak at 1205 cm⁻¹ [[39\]](#page-6-0). The varied levels of interlayer $FeS₂$ may be the cause of the variations in intensities and shifts. These findings support the XRD examinations and show that $Fe₃O₄/FeS₂/g-C₃N₄$ nanocomposites were successfully synthesized at 550 ◦C.

ESEM images were used to examine the shape and structure of the fabricated Fe₃O₄/FeS₂/g-C₃N₄ nanocomposite. The collected micrographs for the powder samples were illustrated in [Fig. 3](#page-3-0). The image of sulfur/g-C₃N₄ showed a 2D flake morphology. In the case of Fe₃O₄/ $FeS_2/g-C_3N_4$ nanocomposites, sheet materials have been split up throughout the growing process, revealing additional edge sites. The 2D sheet morphology of the $Fe₃O₄/FeS₂/g-C₃N₄$ nanocomposites supports the data analysis of XRD and FTIR. Moreover, the results of ESEM micrographs indicate an increase of surface area as the ratios of FeS reaches 3.0 wt%. The EDS elemental data plotted in Fig. S1 reveal the compositional ratios of C, N, Fe, O and S.

The study of the surface properties of nanocomposites represents the most important method for determining the initial catalyst efficiency. Therefore, we have measured the adsorption-desorption data of the $FeS_2/g-C_3N_4$ nanocomposites as shown in [Fig. 4](#page-3-0). The shape of isotherms agrees with the IV type that corresponds to mesoporous structure. The sample S@g-C₃N₄ showed a surface area BET of 40 m²/g. Meanwhile, the Fe₃O₄/FeS₂/g-C₃N₄ nanosheets have a surface area of 109, 137 and 162 $\rm m^2/g$ at 1.0, 2.0 and 3.0 wt%. According to these data, the intercalation of $Fe₃O₄/FeS₂$ into g-C₃N₄ nanosheets increased the surface area and thus increased the number of active sites. Moreover, the BJH model was applied to the collected data to estimate the pore size values. In that case, the pore size for the sample $S@g-C_3N_4$ was 2.0 nm. While the pore size data for the Fe₃O₄/FeS₂/g-C₃N₄ nanosheets were 1.55, 1.77 and 2.15 nm at 1.0, 2.0 and 3.0 wt%. Indeed, the sample 3.0 wt% of $Fe₃O₄/FeS₂$ has the highest surface area and pore size. Therefore, this sample may possess the highest catalytic performance to produce hydrogen.

The optical properties, such as absorbance and the optical band gap of nanocomposites were studied as follows. The absorbance parameters were measured by UV–Vis spectroscopy, as shown in [Fig. 5a](#page-4-0). The wavelength ranges are about 200 to 1000 nm. The absorbance peak is appeared at around 320 nm corresponding to $n \rightarrow \pi^*$ electronic transitions [[40\]](#page-6-0). The peaks of absorbance are red shift due to generating

Fig. 3. ESEM images of Fe₃O₄/FeS₂/g-C₃N₄ nanosheets.

Fig. 4. N₂ isotherm data of $Fe₃O₄/FeS₂/g-C₃N₄$ nanosheets.

photo-induced electrons and holes. The absorption band tail corresponding to heterocyclic aromatics $π$ -π^{*} transitions appeared around 400 nm [[41](#page-6-0)].

The study of the optical energy gap (E_{opt}) of nanocomposites represents the most important method for determining the electronic structure of a catalyst. For that reason, the equation proposed by Tauc was applied to estimate the optical band gap values [42–[44\]](#page-6-0);

$$
ahv = k\big(hv - E_{opt}\big)^{0.5} \tag{2}
$$

The optical band gaps (E_{opt}) can be determined from [Fig. 5b](#page-4-0). The values of the *Eopt* are estimated by the straight-line intercept from the $(\alpha h\nu)^2$ against hv. The values are 2.60, 2.30, 2.21 and 2.18 eV for (S/g-C₃N₄), (1 wt% of Fe₃O₄/FeS₂/g-C₃N₄), (2 wt% of Fe₃O₄/FeS₂/g-C₃N₄), and (3 wt% of $Fe₃O₄/FeS₂/g-C₃N₄$), respectively. These values decrease as the concentration increase; possibly due to the development of energy levels [\[45](#page-6-0)].

When the materials are pumped with photon energy, many electronhole pairs generated at the excited energy states. In that case, the induced electron hole pairs remain for a lifetime and then recombine. The photoluminescence spectra of the prepared $Fe₃O₄/FeS₂/g-C₃N₄$ nanosheets are plotted in [Fig. 6](#page-4-0). From inspection of this figure, all the samples show an emission peak around 460 nm that related to the transitions of n- π ^{*} [\[46](#page-6-0)]. However, the increase of Fe₃O₄/FeS₂ percentage reduced the intensity of this peak. This decrease is explained by the delay in the recombination of the electrons and holes [\[47](#page-6-0)]. For that reason, the catalytic performance of the $Fe₃O₄/FeS₂/g-C₃N₄$ nanosheets may be improved.

The method of self-decomposition of sodium borohydride in a methanol solution occupies a great deal of interest because it has high stability and provides a low temperature. Moreover, this mechanism

Fig. 5. Graphs of (a) absorbance vs wavelength and (b) $(\alpha h\nu)^2$ vs hv for $Fe_3O_4/FeS_2/g-C_3N_4$ nanosheets.

Fig. 6. Emission spectra of $Fe₃O₄/FeS₂/g-C₃N₄$ nanosheets.

produces hydrogen gas rapidly and with high rates [[48\]](#page-6-0). Therefore, this reaction was used to test the efficiency of $Fe₃O₄/FeS₂/g-C₃N₄$ nanocomposites as a catalyst for hydrogen emission from NaBH4. Firstly, the volume of H_2 gas was recorded at various times at 293 K. Fig. 7 display the curves of hydrogen volume at different times for the $Fe₃O₄/FeS₂/g-$ C3N4 nanosheets.

Fig. 7 indicates that the rate of hydrogen production is enhanced as the percentage of $Fe₃O₄/FeS₂$ increased. This comes due to the improvement of surface area and pore size that contained more catalytic active sites. In considering the decomposition of NaBH4 in methanol solution, two ions are produced Na⁺ and BH₄ [[49\]](#page-6-0). The catalyst's surface may adsorb $BH₄⁻$ and CH₃OH molecules from the solution. A new electron is added to the hydrogen atom in its hydridic form (H[−]), while the BH₃ species is still bound to the Fe₃O₄/FeS₂/g-C₃N₄ catalyst. H₂ and CH3O[−] are generated when H[−] and CH3OH react. Moreover, boron in BH₃ interacts with CH₃O[−] to form the BH₃ (CH3O)[−] ion. Once more, the $BH₃$ (OH)⁻ ion is used to transport H₂. In each cycle, the hydrogen adsorption cycle on the catalyst is repeated until the BH₃ (CH3O)[−] ion

Fig. 7. H₂ volume vs time at 20 mg of $Fe₃O₄/FeS₂/g-C₃N₄$ catalyst and 2.5 % NaBH4.

becomes $B(CH_3O)_4$ and H_2 is released [50–[54\]](#page-6-0). The best catalyst must adsorb large amount of BH $_4^-$ at very short times. Further, the preferred catalyst should increase the volume of hydrogen gas [\[55](#page-6-0)].

From the definition of hydrogen generation rate (k), we can estimate the catalytic performance of nanocomposites as a function of H_2 volume (V), mass of the catalyst (m_{cat}) and reaction time (t) [$17,56$ $17,56$];

$$
k = \frac{V}{t \cdot m_{cat}} \tag{3}
$$

The slope of the plots in Fig. 7 were used to compute the rates of hydrogen produced (k). In this case, the obtained data for different $Fe₃O₄/FeS₂/g-C₃N₄$ nanosheets are given in [Fig. 8](#page-5-0). Consequently, the rate of hydrogen production for the sample sulfur@g-C₃N₄ was 6333 mL/g⋅min. While the values of k at 1.0, 2.0 and 3.0 wt% of $Fe₃O₄/FeS₂$ were 6024, 8464 and 8480 mL/g⋅min. Now, the sample 3.0 wt% $Fe₃O₄/$ $FeS₂$ showed the best rate of hydrogen production. This result is related to the structural and surface characteristics of this sample, as it showed a high surface area and a large pore size. Therefore, many active sites were located on the surface of the nanocomposites and induced large amounts

Fig. 8. The rates of H_2 generation at 20 mg of $Fe_3O_4/FeS_2/g-C_3N_4$ catalyst and 2.5 % NaBH4.

of hydrogen.

It is appropriate to make a comparison between the performances of $Fe₃O₄/FeS₂/g-C₃N₄$ nanocomposites as a catalyst for hydrogen production with their counterparts in the literature. In our previous work, the catalyst of 1.5 wt% NiS/g-C₃N₄ was prepared by polycondensation route and tested for hydrogen production [\[19](#page-6-0)]. In that case the rate of hydrogen generation at 293 K was 8654 mL/g⋅min that is lower than the value reported in the present work. Another study conducted by F. Wang et al. [[57\]](#page-7-0) used the electroplating technique to synthesize an active catalyst of Ru/NiO-Ni foam. The morphology of the catalyst material was bush-like. Further, this work achieved a rate of production equal to 6000 mL/g⋅min that is lower than the present finding. On the other hand, the catalytic performance of rutile $TiO₂/NH₂$ was examined ac-cording to the methanolysis hydrolysis of NaBH₄ [[56\]](#page-7-0). As a result, the hydrogen production rate in this case reaches 3525 mL/g⋅min. However, this value is still lower than the estimated value in the present study. Finally, P. Dai et al. $[58]$ $[58]$ reported the self-hydrolysis of NaBH₄ at the ZIF-67@GO catalyst. The surface morphology of the composite showed 3D polyhedrons and the rate of H_2 gas production was 3200 mL/g⋅min. According to these findings, the present 3.0 wt% $Fe₃O₄/FeS₂/g-C₃N₄$ showed high potential for catalytic production of hydrogen gas.

The efficiency of the $Fe₃O₄/FeS₂/g-C₃N₄$ catalyst was investigated for 3 cycles. The data were plotted in Fig. 9 and showed a slight decrease in efficiency from 100 to 99.97 %. This small decrease may be due to the blocking of surface-active sites for the $Fe₃O₄/FeS₂/g-C₃N₄$ catalyst after 3.0 cycles.

4. Conclusions

Nanocomposites of Fe₃O₄/FeS₂/g-C₃N₄ were developed for catalytic hydrogen generation from sodium borohydride. The XRD diffraction peaks of Fe₃O₄ and FeS₂ cubic phase showed an average crystal size of calculation of 15 and 20 nm. ESEM micrographs showed a 2D broken up sheet structure having more edge sites. The surface area BET of the

Fig. 9. The reuse experiments of $Fe₃O₄/FeS₂/g-C₃N₄$ catalysts.

prepared nanosheets increased from 40 to 162 m^2/g . While the optical energy gap decreased from 2.60 to 2.18 eV. All the samples show an emission peak around 460 nm that related to the transitions of n-π*. The rate of hydrogen production is accelerated as the percentage of Fe₃O₄/ FeS₂ increased. The sample 3.0 wt% Fe₃O₄/FeS₂ showed the best rate of hydrogen production with a value of 8480 mL/g⋅min. Accordingly, the prepared 3.0 wt% $Fe₃O₄/FeS₂/g-C₃N₄$ nanosheet showed high potential for catalytic production of hydrogen gas.

Supplementary data to this article can be found online at [https://doi.](https://doi.org/10.1016/j.diamond.2023.110214) [org/10.1016/j.diamond.2023.110214](https://doi.org/10.1016/j.diamond.2023.110214).

CRediT authorship contribution statement

All the authors have contributed in the work, and they are known with the submission.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability statement

Data will be made available on reasonable request.

Acknowledgments

The authors extend their appreciation to the Deputyship for Research & Innovation, Ministry of Education in Saudi Arabia for funding this research work through the project number 223202.

References

^[1] [R. Shwetharani, M. Sakar, C.A.N. Fernando, V. Binas, R.G. Balakrishna, Recent](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0005) [advances and strategies to tailor the energy levels, active sites and electron](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0005)

M. Alshammari et al.

[mobility in titania and its doped/composite analogues for hydrogen evolution in](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0005) [sunlight, Catal. Sci. Technol. 9 \(1\) \(2019\) 12](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0005)–46.

- [2] [L. Chen, Z. Qi, S. Zhang, J. Su, G.A. Somorjai, Catalytic hydrogen production from](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0010) [methane: a review on recent progress and prospect, Catalysts 10 \(8\) \(2020\) 858.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0010)
- [3] [N.A. Roslan, S.Z. Abidin, A. Ideris, D.V.N. Vo, A review on glycerol reforming](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0015) [processes over Ni-based catalyst for hydrogen and syngas productions, Int. J.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0015) [Hydrog. Energy 45 \(36\) \(2020\) 18466](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0015)–18489.
- [4] [M. Sohail, U. Anwar, T.A. Taha, H.I.A. Qazi, A.G. Al-Sehemi, S. Ullah, A. Hayat,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0020) Nanostructured materials based on g-C₃N₄ for enhanced photocatalytic activity and [potentials application: a review, Arab. J. Chem. 15 \(9\) \(2022\), 104070.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0020)
- [5] [A. Hayat, M. Sohail, U. Anwar, T.A. Taha, H.I.A. Qazi, Z. Ajmal, Y. Orooji,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0025) [A targeted review of current progress, challenges and future perspective of g-C3N4](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0025) [based hybrid photocatalyst toward multidimensional applications, Chem. Rec. 23](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0025) [\(1\) \(2023\), e202200143.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0025)
- [6] [H. Guo, C.G. Niu, Y.Y. Yang, C. Liang, H.Y. Niu, H.Y. Liu, N. Tang, Interfacial Co-N](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0030) [bond bridged CoB/g-C3N4 Schottky junction with modulated charge transfer](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0030) [dynamics for highly efficient photocatalytic Staphylococcus aureus inactivation,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0030) [Chem. Eng. J. 422 \(2021\), 130029](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0030).
- [7] [T. Mahvelati-Shamsabadi, H. Fattahimoghaddam, B.K. Lee, S. Bae, J. Ryu,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0035) [Synthesis of hexagonal rosettes of g-C3N4 with boosted charge transfer for the](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0035) [enhanced visible-light photocatalytic hydrogen evolution and hydrogen peroxide](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0035) [production, J. Colloid Interface Sci. 597 \(2021\) 345](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0035)–360.
- [8] [C. Wan, J. Li, S. Chen, W. Wang, K. Xu, In situ synthesis and catalytic](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0040) [decomposition mechanism of CuFe2O4/g-C3N4 nanocomposite on AP and RDX,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0040) [J. Anal. Appl. Pyrolysis 160 \(2021\), 105372](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0040).
- [9] J. Wang, X. Lian, S. Chen, H. Li, K. Xu, Effect of Bi2WO6/g-C₃N₄ composite on the [combustion and catalytic decomposition of energetic materials: an efficient catalyst](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0045) [with g-C3N4 carrier, J. Colloid Interface Sci. 610 \(2022\) 842](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0045)–853.
- [10] [C. Mutalik, I.H. Lin, D.I. Krisnawati, S. Khaerunnisa, M. Khafid, Widodo, T.R. Kuo,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0050) [Antibacterial pathways in transition metal-based nanocomposites: a mechanistic](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0050) [overview, Int. J. Nanomedicine \(2022\) 6821](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0050)–6842.
- [11] [C. Mutalik, Y.C. Hsiao, Y.H. Chang, D.I. Krisnawati, M. Alimansur, A. Jazidie, T.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0055) [R. Kuo, High uv-vis-nir light-induced antibacterial activity by heterostructured](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0055) [TiO2-FeS2 nanocomposites, Int. J. Nanomedicine \(2020\) 8911](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0055)–8920.
- [12] [T.R. Kuo, H.J. Liao, Y.T. Chen, C.Y. Wei, C.C. Chang, Y.C. Chen, D.Y. Wang,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0060) [Extended visible to near-infrared harvesting of earth-abundant FeS2](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0060)–TiO2 [heterostructures for highly active photocatalytic hydrogen evolution, Green Chem.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0060) [20 \(7\) \(2018\) 1640](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0060)–1647.
- [13] [C. Mutalik, D.I. Krisnawati, S.B. Patil, M. Khafid, D.S. Atmojo, P. Santoso, T.R. Kuo,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0065) [Phase-dependent MoS2 nanoflowers for light-driven antibacterial application, ACS](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0065) [Sustain. Chem. Eng. 9 \(23\) \(2021\) 7904](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0065)–7912.
- [14] [D.N. Ma, X.M. Li, X.Q. Wang, Y.J. Luo, Preparation of g-C3N4 nanosheets/CuO](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0070) [with enhanced catalytic activity on the thermal decomposition of ammonium](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0070) [perchlorate, Eur. J. Inorg. Chem. 2021 \(10\) \(2021\) 982](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0070)–988.
- [15] [X. Liu, R. Ma, L. Zhuang, B. Hu, J. Chen, X. Liu, X. Wang, Recent developments of](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0075) [doped g-C3N4 photocatalysts for the degradation of organic pollutants, Crit. Rev.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0075) [Environ. Sci. Technol. 51 \(8\) \(2021\) 751](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0075)–790.
- [16] [Y. Shi, L. Wang, L. Fu, C. Liu, B. Yu, F. Yang, Y. Hu, Sodium alginate-templated](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0080) [synthesis of g-C3N4/carbon spheres/Cu ternary nanohybrids for fire safety](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0080) [application, J. Colloid Interface Sci. 539 \(2019\) 1](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0080)–10.
- [17] [P. Li, D.E. Miser, S. Rabiei, R.T. Yadav, M.R. Hajaligol, The removal of carbon](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0085) [monoxide by iron oxide nanoparticles, Appl. Catal. B Environ. 43 \(2\) \(2003\)](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0085) 151–[162.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0085)
- [18] [C. Saka, Phosphorus decorated g-C3N4-TiO2 particles as efficient metal-free](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0090) [catalysts for hydrogen release by NaBH4 methanolysis, Fuel 322 \(2022\), 124196.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0090)
- [19] [A.H. Alshammari, K. Alshammari, T. Alotaibi, M. Alshammari, S. Alhassan, T.A.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0095) [M. Taha, In situ polycondensation synthesis of NiS-g-C3N4 nanocomposites for](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0095) [catalytic hydrogen generation from NaBH4, Nanomaterials 13 \(5\) \(2023\) 938.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0095)
- [20] [K. Ganesan, C. Hayagreevan, R. Rahul, A.J. Jeevagan, T. Adinaveen, D.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0100) [S. Bhuvaneshwari, M. Amalraj, Catalytic hydrolysis of sodium borohydride for](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0100) [hydrogen production using phosphorylated silica particles, Environ. Sci. Pollut.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0100) [Res. \(2022\) 1](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0100)–14.
- [21] [C. Saka, Efficient and durable H2 production from NaBH4 methanolysis using N](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0105) [doped hybrid g-C3N4-SiO2 composites with ammonia as a nitrogen source, Fuel](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0105) [324 \(2022\), 124594.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0105)
- [22] [A.H. Alshammari, M. Alshammari, S. Alhassan, K. Alshammari, T. Alotaibi, T.A.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0110) [M. Taha, MoO3/S@ g-C3N4 nanocomposite structures: synthesis, characterization,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0110) [and hydrogen catalytic performance, Nanomaterials 13 \(5\) \(2023\) 820](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0110).
- [23] [H. Zhang, Q. Wei, G. Wu, S. Qiu, Y. Zou, Y. Xia, H. Chu, Zn-MOF-74-derived](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0115) [graphene nanosheets supporting CoB alloys for promoting hydrolytic](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0115) [dehydrogenation of sodium borohydride, J. Alloys Compd. 930 \(2023\), 167486](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0115).
- [24] [T. Yan, Q. Yan, X. Wang, H. Liu, M. Li, S. Lu, M. Sun, Facile fabrication of](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0120) [heterostructured gC3N4/Bi2MoO6 microspheres with highly efficient activity](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0120) [under visible light irradiation, Dalton Trans. 44 \(4\) \(2015\) 1601](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0120)–1611.
- [25] [V. Jayaraman, C. Ayappan, G. Vattikondala, A. Mani, Preparation and](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0125) [characterization of the Cu, Fe co-doped Bi2Ti2O7/EG-g-C3N4 material for organic](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0125) [model pollutants removal under direct sun light irradiation, Mater. Res. Bull. 143](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0125) [\(2021\), 111439.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0125)
- [26] J. Li, H. Hao, Z. Zhu, Construction of g-C3N4-WO3-Bi2WO6 double Z-scheme [system with enhanced photoelectrochemical performance, Mater. Lett. 168 \(2016\)](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0130) 180–[183.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0130)
- [27] [Y. Fang, X. Fu, X. Wang, Diverse polymeric carbon nitride-based semiconductors](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0135) [for photocatalysis and variations, ACS Mater. Lett. 2 \(8\) \(2020\) 975](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0135)–980.
- [28] [Q. Liang, B. Shao, S. Tong, Z. Liu, L. Tang, Y. Liu, Z. Peng, Recent advances of](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0140) [melamine self-assembled graphitic carbon nitride-based materials: design,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0140)

[synthesis and application in energy and environment, Chem. Eng. J. 405 \(2021\),](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0140) [126951.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0140)

- [29] [S. Meng, S. Sun, Y. Liu, Y. Lu, M. Chen, Synergistic modulation of inverse spinel](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0145) [Fe3O4 by doping with chromium and nitrogen for efficient electrocatalytic water](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0145) [splitting, J. Colloid Interface Sci. 624 \(2022\) 433](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0145)–442.
- [30] [H. Choi, J.Y. Seo, Y.R. Uhm, G.M. Sun, C.S. Kim, Crystalline structure and magnetic](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0150) operties of pyrite FeS2, AIP Adv. 11 (1) (2021), 015131.
- [31] [K. Khan, Z. Batool, S. Manzoor, D. Ahmad, S. Aman, D. Alhashmialameer, M.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0155) [N. Ashiq, Fabrication of substituted Y-type hexaferrites/carbon dots composites for](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0155) [recording media and photodegradation of dye, Ceram. Int. 48 \(19\) \(2022\)](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0155) [27550](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0155)–27559.
- [32] [M.H. Mahmoud, A.M. Hassan, A.E.A.A. Said, T.A. Taha, Structural, magnetic, and](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0160) [catalytic studies of microwave-combustion/ball-mill synthesized zinc ferrite](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0160) [nanoparticles, Inorg. Chem. Commun. 144 \(2022\), 109932.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0160)
- [33] [T.A. Taha, R. Saad, M. Zayed, M. Shaban, A.M. Ahmed, Tuning the surface](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0165) [morphologies of ZnO nanofilms for enhanced sensitivity and selectivity of CO2 gas](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0165) [sensor, Appl. Phys. A 129 \(2\) \(2023\) 115](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0165).
- [34] [C. Saka, Sulphur and nitrogen-doped metal-free microalgal carbon catalysts for](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0170) [very active dehydrogenation of sodium borohydride in methanol, Int. J. Hydrog.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0170) [Energy 46 \(35\) \(2021\) 18326](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0170)–18337.
- [35] [Z.T. Wang, J.L. Xu, H. Zhou, X. Zhang, Facile synthesis of Zn\(II\)-doped gC3N4 and](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0175) [their enhanced photocatalytic activity under visible light irradiation, Rare Metals](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0175) [38 \(2019\) 459](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0175)–467.
- [36] [J. Gao, J. Wang, X. Qian, Y. Dong, H. Xu, R. Song, J. Yao, One-pot synthesis of](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0180) [copper-doped graphitic carbon nitride nanosheet by heating Cu](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0180)–melamine [supramolecular network and its enhanced visible-light-driven photocatalysis,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0180) [J. Solid State Chem. 228 \(2015\) 60](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0180)–64.
- [37] [C. Fan, J. Miao, G. Xu, J. Liu, J. Lv, Y. Wu, Graphitic carbon nitride nanosheets](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0185) [obtained by liquid stripping as efficient photocatalysts under visible light, RSC](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0185) [Adv. 7 \(59\) \(2017\) 37185](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0185)–37193.
- [38] [M.J. Bojdys, J.O. Müller, M. Antonietti, A. Thomas, Ionothermal synthesis of](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0190) [crystalline, condensed, graphitic carbon nitride, Chem Eur J 14 \(27\) \(2008\)](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0190) [8177](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0190)–8182.
- [39] [P. Xia, B. Zhu, J. Yu, S. Cao, M. Jaroniec, Ultra-thin nanosheet assemblies of](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0195) [graphitic carbon nitride for enhanced photocatalytic CO2 reduction, J. Mater.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0195) [Chem. A 5 \(7\) \(2017\) 3230](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0195)–3238.
- [40] [A.J. Wang, H. Li, H. Huang, Z.S. Qian, J.J. Feng, Fluorescent graphene-like carbon](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0200) [nitrides: synthesis, properties and applications, J. Mater. Chem. C 4 \(35\) \(2016\)](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0200) [8146](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0200)–8160.
- [41] [A.B. Jorge, D.J. Martin, M.T. Dhanoa, A.S. Rahman, N. Makwana, J. Tang, P.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0205) [F. McMillan, H2 and O2 evolution from water half-splitting reactions by graphitic](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0205) [carbon nitride materials, J. Phys. Chem. C 117 \(14\) \(2013\) 7178](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0205)–7185.
- [42] [A.H. Alshammari, M. Alshammari, K. Alshammari, N.K. Allam, T.A. Taha, PVC/](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0210) [PVP/SrTiO3 polymer blend nanocomposites as potential materials for](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0210) [optoelectronic applications, Results Phys. 44 \(2023\), 106173.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0210)
- [43] [R.M. Ahmed, T.A. Taha, F.M. Ezz-Eldin, Investigation of Sm2O3 effect on opto](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0215)[electrical parameters and dielectric properties of some fluorophosphate glasses,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0215) [J. Mater. Sci. Mater. Electron. 32 \(2021\) 28919](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0215)–28934.
- [44] [A.H. Alshammari, K. Alshammari, M. Alshammari, T.A.M. Taha, Structural and](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0220) [optical characterization of g-C3N4 nanosheet integrated PVC/PVP polymer](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0220) [nanocomposites, Polymers 15 \(4\) \(2023\) 871](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0220).
- [45] [X. Wang, J. Gong, Y. Dong, S. An, X. Zhang, J. Tian, Energy band engineering of](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0225) [hydroxyethyl group grafted on the edge of 3D g-C3N4 nanotubes for enhanced](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0225) [photocatalytic H2 production, Mater. Today Phys. 27 \(2022\), 100806](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0225).
- [46] [X. Song, W. Mao, Y. Wu, M. Wang, X. Liu, W. Zhou, P. Huo, Fabricating carbon](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0230) [nitride-based 3D/0D intramolecular donor](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0230)–acceptor catalysts for efficient [photoreduction of CO2, New J. Chem. 46 \(42\) \(2022\) 20225](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0230)–20234.
- [47] [X. Rong, F. Qiu, J. Rong, X. Zhu, J. Yan, D. Yang, Enhanced visible light](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0235) [photocatalytic activity of W-doped porous g-C3N4 and effect of H2O2, Mater. Lett.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0235) [164 \(2016\) 127](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0235)–131.
- [48] [C. Saka, A. Balbay, Influence of process parameters on enhanced hydrogen](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0240) [generation via semi-methanolysis and semi-ethanolysis reactions of sodium](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0240) [borohydride using phosphoric acid, Int. J. Hydrog. Energy 44 \(57\) \(2019\)](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0240) [30119](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0240)–30126.
- [49] [D. Xu, Y. Zhang, Q. Guo, Research progress on catalysts for hydrogen generation](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0245) [through sodium borohydride alcoholysis, Int. J. Hydrog. Energy 47 \(9\) \(2021\)](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0245) [5929](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0245)–5946.
- [50] [C. Saka, Metal-free phosphorus and boron-doped graphitic carbon nitride/zeolite](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0250) [hetero-linked particles for highly efficient green hydrogen production in methanol,](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0250) [Environ. Sci. Pollut. Res. 30 \(2023\) 43480](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0250)–43495.
- [51] [C. Saka, Metal-free hybrid composite particles with phosphorus and oxygen-doped](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0255) [graphitic carbon nitride dispersed on kaolin for catalytic activity toward efficient](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0255) [hydrogen release, Int. J. Hydrog. Energy 48 \(37\) \(2023\) 13864](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0255)–13876.
- [52] [C. Saka, Surface modification of graphitic carbon nitride nanoparticles with B, O](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0260) [and S doping/carbon vacancy for efficient dehydrogenation of sodium borohydride](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0260) [in methanol, Int. J. Hydrog. Energy 48 \(35\) \(2023\) 13123](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0260)–13138.
- [53] [C. Saka, Performance of g-C3N4 nanoparticles by EDTA modification and](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0265) [protonation for hydrogen release from sodium borohydride methanolysis, Int. J.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0265) [Hydrog. Energy 47 \(28\) \(2022\) 13654](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0265)–13663.
- [54] [S.A.K.A. Cafer, Highly active hydrogen generation from sodium borohydride](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0270) [methanolysis and ethylene glycolysis reactions using protonated chitosan-zeolite](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0270) [hybrid metal-free particles, Appl. Catal. B Environ. 325 \(2023\), 122335.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0270)
- [55] [S. Demirci, A.K. Sunol, N. Sahiner, Catalytic activity of amine functionalized](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0275) [titanium dioxide nanoparticles in methanolysis of sodium borohydride for](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0275) [hydrogen generation, Appl. Catal. B Environ. 261 \(2020\), 118242](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0275).

M. Alshammari et al.

- [56] [C. Saka, g-C3N4 particles with boron and oxygen dopants/carbon vacancies for](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0280) [efficient dehydrogenation in sodium borohydride methanolysis, Int. J. Hydrog.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0280) [Energy 47 \(44\) \(2022\) 19016](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0280)–19026.
- [57] [F. Wang, Y. Luo, Y. Zhang, Y. Wang, H. Zhu, Preparation of bush-like Ru/NiO-Ni](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0285) [foam catalyst and its performance in hydrogen production from sodium](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0285) [borohydride alcoholysis, Energy Fuel 34 \(9\) \(2020\) 11365](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0285)–11372.
- [58] [P. Dai, Y. Yao, E. Hu, D. Xu, Z. Li, C. Wang, Self-assembled ZIF-67@ graphene oxide](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0290) [as a cobalt-based catalyst precursor with enhanced catalytic activity toward](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0290) [methanolysis of sodium borohydride, Appl. Surf. Sci. 546 \(2021\), 149128.](http://refhub.elsevier.com/S0925-9635(23)00539-3/rf0290)