**Ag loaded B-doped-g C3N4 nanosheet with efficient properties for photocatalysis**

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**Abstract:**

Three material engineering strategies in the form of doping (Boron-doping), nanostructuring (nanosheet (NS) formation) and decorating with plasmonic nanoparticles (loading with Ag metal), were integrated to improve the photocatalytic activity of graphitic carbon nitride (gC3N4). Concentrations of B-doping and Ag-loading were optimized to maximize the catalytic performance in the final nanocomposite of Ag-loaded B-doped gC3N4 NS. Combined effect of all three strategies successfully produced over 5 times higher rate towards degradation of organic dye pollutant when compared to unmodified bulk gC3N4. Detailed characterization results revealed that incorporation of B in gC3N4 matrix reduces the band gap to increase the visible light absorption, while specific surface area is significantly enhanced upon formation of NS. Decoration of Ag nanoparticles (NPs) on B-doped gC3N4 NS assists in fast transfer of photogenerated electrons from gC3N4 to Ag NPs owing to the interfacial electric field across the junctions and thus reduces the recombination process. Investigations on individual strategies revealed that decoration of Ag NPs to induce better charge separation, is the most effective route for enhancing the photocatalytic activity.

**Keywords:** Graphitic carbon nitride, B doping, Ag nanoparticle, plasmonic, g-C3N4 nanosheet, recombination rate.

**1. Introduction:**

Rapid industrial growth has resulted in huge amount of waste water containing various organic and inorganic pollutants. Advanced oxidation processes (AOPs) such as photo Fenton [Pouran, et al., 2015; Pouran, et al., 2016; Pouran, et al., 2017, Pouran, et al.2018] and photocatalysis [Khaki et al., 2018; Khaki et al., 2018; Khaki et al., 2017] have proven to be highly effective methods for mineralization of these organic pollutants, among which photocatalysis has gained special interest owing to its simplicity of implementation. Photocatalysis involves three major steps 1) absorption of photons with energy greater than the band gap of the semiconductor, 2) excitation of electrons to the conduction band leaving behind holes in the valance band and 3) charge carrier transfer process, in which electrons located in conduction band are trapped by O2 molecule adsorbed on photocatalyst to produce superoxide anion radicals. Meanwhile, holes accumulated in valence band combines with water molecule to give rise to hydroxyl radicals (OH**.**). These highly active radicals readily react with organic pollutant and decompose them into CO2 and H2O.

Several semiconductor materials such as TiO2, ZnO, WO3, etc. have been extensively studied for photocatalytic degradation of organic pollutants [Khaki et al., 2018; Khaki et al., 2018; Khaki et al., 2017, Sakthivel et al. 2002]. However, wide band gap and inappropriate band edge position limits their efficiency to a great extent. Recently, graphitic carbon nitride (g-C3N4) has gained enormous attention due to their wide application in various fields, such as decontamination of water [Gao et al., 2018; Caux et al. 2017], H2 production via photocatalytic water splitting [Chen et al., 2018; Li et al., 2016] and electrolysis [Xu et al., 2016], etc., owing to its moderate band gap and excellent chemical stability. Nevertheless, photocatalytic efficiency of gC3N4 is limited due to low surface area, narrow visible light absorption and high recombination rate [Tonda et al., 2014; Hu et al., 2014]. Several strategies have been employed to overcome these limitations since its first application for environmental issues. Stacked layered structure of gC3N4 leads to low surface area, thus various chemical and physical methods have been employed to break the weak Vander Waals forces between stacked layers of gC3N4. Exfoliation of gC3N4 into nanosheets (NSs) not only provides high surface area associated with large number of active sites, but is also advantageous in terms of having shorter diffusion length with low recombination rate probability compared to bulk gC3N4 [Niu et al., 2012]. Gao et al synthesized gC3N4 NS by rapid cooling treatment using liquid N2 and demonstrated significant improvement in specific surface area (142.84 m2/g) compared to bulk gC3N4 (15.67 m2/g) and suppressed recombination rate of charge carriers. A group of researchers successfully synthesized monolayered gC3N4 NS using simple mixed-solvent method with superior photodegradation ability under visible light irradiation towards Rhodamine B (Rh B) [Lin et al., 2012]. The effective improvement in photoactivity was achieved owing to the favourable band structure, high surface area, prolonged lifetime of photogenerated charge carriers, high number of exposed active sites and low surface defects. Recently, easy and facile one–step approach involving acidification of melamine was proposed to prepare gC3N4 NSs, which revealed 3-fold increase in photocatalytic degradation rate under visible light illumination compared to gC3N4 [Zhang et al., 2014]. In addition to that, morphological control through post acidification, high temperature annealing and sonication in organic solvents etc. have also been reported as an efficient routes to improve the surface area of pristine gC3N4 [She et al., 2016; Kang et al., 2018]. Numerous efforts have been taken to synthesize different structures by modifying the surface texture of gC3N4 to obtain optimal photocatalyst. Chen and his group [Chen et al., 2014] reported great improvement in photodegradation ability of gC3N4/Ag/TiO2 microspheres towards methyl orange and phenol due to efficient electron transfer process. Triple-shell gC3N4 nano-capsules showed superior photodegradation performance compared to single and double-shell gC3N4 nano-capsule towards Rh B dye owing to improved visible light absorption and charge separation [Tong et al., 2017]. A novel hybrid architecture of mesoporous gC3N4 with BiPO4 nanorods displayed significant improvement in photocatalytic degradation of methyl orange dye due to ultrafast migration of charge carriers at the interface [Li et al., 2014].

The strategy of doping with metals [Deng et al., 2016; Sun et al, 2015] and non-metals [Zhu et al., 2015; Sagra et al., 2016] has also proven to be an excellent method for band gap engineering in order to extend light absorption ability in visible region. It was observed that anion doping induces conduction band tuning through hybridization between p orbital of doped element and C atom present in gC3N4 matrix. On the other hand, cation doping modifies valance band edge by forming discrete band of 3d orbitals of transition element. It was found that boron doping lowers valence band position of resulting material and hence increases the potential oxidation strength of photogenerated holes as compared to gC3N4 [Sagara et al., 2016]. Yan et al. described that boron doping promotes photodegradation of pollutants by improving the dye adsorption and light absorption ability. Superior photocatalytic removal of was reported for B-doped gC3N4 due to band gap narrowing [Lu et al., 2016]. High concentration of boron doping leads to formation of ternary compound boron carbon nitride (BCN), possessing properties of boron nitride and gC3N4 [Sivaprakash et al., 2018]. Depending upon the concentration of B, C and N, properties such as electron affinity, bandgap (0-5.5 eV) and effective barrier potential varies [Sivaprakash et al., 2018]. Hu et al. demonstrated significant improvement in photocatalytic performance of Fe doped bulk gC3N4 compared to gC3N4 for both degradation of organic pollutant and water splitting process, however low surface area of Fe doped gC3N4 limits its photocatalytic performance. It has been established that addition of Fe and P codopant in gC3N4 photocatalyst results in enhanced surface area, extended visible light absorption and narrowing of band gap in addition to decreased electron-hole pair recombination rate, thus exhibiting much higher Rh B photodegradation rate [Hu et al., 2014].

The issue of high recombination rate in gC3N4 can be suppressed by noble metal loading on gC3N4 surface, or coupling it with other semiconductors such as TiO2, graphene oxide, carbon nanodots etc [Li et al., 2017; Xiang et al., 2011; Dadigala et al., 2017]. Bu et.al. [Bu et al., 2014] prepared Ag modified gC3N4 which displayed significant improvement in electron hole pair separation with prolonged lifetime. Ge and his [Ge et al., 2011] group synthesized Ag loaded gC3N4 and reported substantial enhancement in photocatalytic performance which was attributed to enhanced visible light absorption and efficient migration of photogenerated charge carriers. Ag loaded gC3N4 was also prepared by Zhang et al. [Zhang et al., 2016] by photo-deposition method under ambient condition which showed improved photodegradation performance than pure gC3N4 as a result of close contact between Ag and gC3N4 and localized surface plasmon resonance (SPR) of Ag. Cheng et al. [Chang et al., 2013] found that Pd decorated gC3N4 facilitates efficient separation of photoelectrons and holes through electron trapping on Pd0 atom.

There are very few reports that involves combining two of the above mentioned three strategies to modify gC3N4 and improve its photocatalytic performance. Li et al. [Li et al., 2017] synthesized Ti3+-TiO2 decorated exfoliated oxygen doped gC3N4 and showed remarkable increase in photocatalytic activity due to combined effect of efficient photogenerated electron hole pair separation and decreased charge carrier resistance at the interface. B-doped gC3N4 in form of nanosheets showed 12 times better photocatalytic performance for H2 evolution as compared to pristine gC3N4 [Thaweesak et al., 2017], due to the synergistic effect of enhanced visible light absorption and higher surface area. Dadigala et al. [Dadigala et al., 2017] synthesized gC3N4 NS and decorated it with carbon dots as well as Ag nanoparticles to provide improved separation of photogenerated carriers and enhanced visible light absorption.

To the best of our knowledge, there is scarcity of report combining all the above-mentioned strategies to improve the photocatalytic properties of gC3N4. The present work aims not only to extend visible light absorption but also to enhance electron-hole pair separation and achieve higher active surface area. Therefore in this work, boron doped gC3N4 was first synthesized to narrow the band gap. Subsequently, Ag NPs were decorated over B doped gC3N4 to reduce electron hole pair recombination rate. In order to achieve higher active surface area with more number of exposed active sites, bulk gC3N4 was exfoliated to obtain nanosheet form. Thus, the present report gives a comprehensive study of combining all the above-mentioned strategies and its effect to improve the performance of gC3N4 towards degradation of industrial dyes. This form of gC3N4 showed over 5 times superior photodegradation performance compared to bulk gC3N4 towards Rh B under light illumination. Role of each of the strategies in enhancing photocatalytic activity of Ag NPs loaded B-doped gC3N4 NS compared to bulk gC3N4 is explained using various characterization techniques.

**2. Experimental methods:**

**2.1 Materials**

Melamine (pure) (C3H6N6, ≥ 99%) was purchased from Loba chemie laboratory reagents and fine chemicals. Boric acid (H3BO3, 99.5%) and Silver nitrate (extrapure) (AgNO3, 99.8%) were purchased from S.D. Fine-Chem Limited. Rh B (AR) was obtained from central drug house private limited. Nitric acid (HNO3, 65 wt%) was obtained from Merck life science private limited. Ethanol (99.9 %) was purchased from Changshu Hongsheng fine chemicals. All the above mentioned reagents used in the present work were of analytically pure grade and were used without any further purification.

**2.2 Synthesis of photocatalyst materials:**

***2.2.1 Bulk g-C3N4 & B-doped-gC3N4:***

B-doped-gC3N4 was synthesized by thermal polymerization of melamine at high temperature in a programmable furnace. In brief, required amount of boric acid (H3BO3) was dissolved in 50 ml of ethanol followed by addition of 4gm of melamine powder and dispersed through ultrasonic treatment. The quantity of H3BO3 was selected to obtain 4, 6, 8 and 10 at.% of Boron in final gC3N4 powder. The mixture was stirred for 15 minutes at room temperature and then dried at 80oC. This dried mixture was transferred to a ceramic crucible with cover lid and calcinated in air at 650oC for 4h with a ramp rate of 3 ̊C/min. After calcination, a yellowish orange compound of B-doped-gC3N4 was obtained, which was grounded into fine powder. Pure gC3N4 powder was also prepared by using the same procedure without the addition of boron precursor.

***2.2.2 gC3N4 & B-doped-gC3N4 NS:***

gC3N4 and B-doped-gC3N4 NS were prepared by liquid exfoliation of the as-prepared powders in diluted HNO3. In detail, 500 mg of as synthesized powder was dispersed in 30 ml HNO3 (65 wt%) and stirred for 7h at room temperature to weaken the bonds between interlayers. This solution was diluted, using 200 ml double distilled water (DDW) and ultrasonicated for 16 hours to exfoliate the gC3N4 layers. The suspension was separated from residual solution and washed with DDW and ethanol several times to remove any residual acid. Finally, the mixture was dried at 80 oC to obtain NSs of gC3N4 & B-doped-gC3N4. The obtained powder was light yellowish orange in colour.

***2.2.3 Ag loaded B-gC3N4:***

100 mg of B-doped-gC3N4 (bulk or nanosheet) and appropriate amount (5, 10 & 15 mg) of silver nitrate (AgNO3) was dispersed in 25 ml of ethanol and stirred for 5 h at room temperature. Finally, the suspension was dried at 80 oC to obtain Ag metal loaded B-gC3N4.

**2.3 Characterization of samples:**

To identify the different phases and crystalline structure in the samples, X-ray diffraction (XRD) measurements were performed on a XRD-6100 (Shimadzu) instrument with a Cu Kα source (Applied voltage & current values of 30 kV & 30 mA). The morphology of the samples were investigated by JEOL (2100F Cs corrected) Field emission gun based transmission electron microscope (FEG-TEM) using an accelerating voltage of 200 kV. Brunauer-Emmett-Teller (BET) specific surface area of the catalyst was measured by nitrogen (N2) adsorption-desorption isotherms at -196 oC on a BET surface area system (Smart SORB 93) after degassing all samples at 150 oC for 3h. The optical absorption spectra of the samples were measured using a Cary 500 UV-vis-NIR spectrometer in diffuse reflectance mode in the range of 200-800 nm.   
Photoluminescence (PL) studies were carried out using fluorescence spectrophotometer (Varian, Cary Eclipse). The emission spectra was collected at an excitation wavelength of 385nm. The average lifetime of charge carriers of the samples was examined by Time resolved photoluminescence (TRPL) spectroscopy (ISS chronos BH model). To acquire PL and TRPL spectra, 2 mg powder catalyst was dispersed in 10ml of DDW and transferred into a 1cm x 1cm cuvette for measurement. The chemical state and surface composition of the samples was studied by X-ray photoelectron spectroscopy (XPS) (PHI 5000 Versaprobe scanning ESCA microprobe) equipped with a monochromatic Al Kα (1486.6 eV) X-ray source and a hemispherical analyzer. Appropriate electrical charge compensation was employed to perform the analysis and binding energy was referenced to the C 1s peak.

**2.4 Photocatalytic performance:**

The photocatalytic performance of all the samples were carried out in a vertical photoreactor set up using 150W Xenon lamp (Hamamatsu) as a source of light. Typically, to study photocatalytic activity, 10 mg of photocatalyst sample was dispersed in 50 ml aqueous suspension solution of Rh B (10 mg/ L) via ultrasonic treatment for 5 mins. Prior to photocatalytic degradation test, the suspension was kept under constant stirring for 30 mins in dark in order to achieve adsorption- desorption equilibrium between photocatalyst and dye. During light irradiation, after each successive time interval, 2 ml of suspension was drawn and centrifuged to separate supernatant from photocatalyst powder. The concentration of dye in residual solution was evaluated by recording intensity of the absorption band of Rh B at 547 nm using UV-Vis spectrophotometer (Implen). Throughout the experiment, the pH of the solution was maintained to neutral and activity was performed in ambient condition. For the complete duration of the experiment, the distance between surface of suspension solution and source of light was kept constant and equal to 47cm. In order to discard the possibility of self-degradation of dye under light illumination, blank test of Rh B suspension solution was performed in the absence of photocatalyst powder. All the tests were repeated at least twice to confirm the obtained results.

**3. Results and Discussion:**

Three material engineering strategies were integrated in order to solve the most basic problems of gC3N4 photocatalyst i.e. high recombination rate of photogenerated charges, absorption of a narrow region of the visible light and low surface area with limited number of active sites. Firstly, boron was doped in gC3N4 which is expected to replace C in the matrix; secondly, this B-doped-gC3N4 is exfoliated to form NSs and lastly decorate this doped-NS with plasmonic materials in the form of Ag NPs. However, before combining all three strategies, each step is optimized before moving to the next. It is also important to investigate the effect of each strategies in the nanocomposite, thus the combinations of two strategies were also studied before preparing the final composite.

gC3N4 was first mono doped with four different concentrations of boron (4, 6, 8 & 10 at.%) in order to select the optimum concentration for preparing nanocomposite. The range of doping was chosen such that boron is doped in the matrix of gC3N4 rather than forming segregated boron species over gC3N4 sheets. The optimization of B concentration was done on the basis of the photocatalytic performance for degradation of Rh B dye. By measuring the relative intensity of the peak at ~547nm with respect to time, the decrease in Rh B dye concentration was estimated. Self-degradation of Rh B dye was found to be negligible under the light irradiation in the absence of photocatalyst. Among the different concentrations, boron doping of 8 at.% shows the best photocatalytic activity (Fig. S1) with 1.8 times higher rate of degradation than undoped gC3N4. This optimized B-doped gC3N4 was later converted into ‘single layer’ or ‘few layers based NS’ using acid treatment followed by exfoliation under ultrasonication for long duration of time (16 hr). Different amount of Ag metal NPs (3, 6 or 9 wt.%) were loaded over B-doped gC3N4 NS and it was discovered that for each loading amount, the photocatalytic activity increases as compared to B-doped gC3N4 NS and undoped gC3N4 NS (Fig. S2). However, no major variation was observed within all three loading concentrations and thus a minimum loading of 5 wt.% was selected as the optimized loading. Further studies were carried out on samples namely, bulk g-C3N4, B-doped gC3N4 bulk, undoped gC3N4 NS, B-doped gC3N4 NS, Ag NPs loaded over B-gC3N4 bulk and Ag NPs loaded over B-gC3N4 NS which are henceforth recognized as bulk gC3N4, B-bulk-gC3N4, NS-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 respectively.

Structural modification of gC3N4 was studied by XRD and reported in Fig.1. Two characteristic peaks with a strong intensity at 27.6o and a weak signal at 12.9o corresponding to (002) and (100) planes of gC3N4 (JCPDS-87-1526) were observed for all six samples. The peak at 27.6°corresponds to a characteristic interlayer stacking reflection and the one at 12.9° represents the in plane structural packing motif of the heptazine units. As compared to bulk-gC3N4, the peak intensity of the main (002) peak is reduced in case of NS-gC3N4,suggesting a decrease in the number of layers in the stacking. In all B-doped gC3N4 samples, the same peak is slightly shifted to higher angle by 0.2o, thus indicating that B is incorporated in the gC3N4 lattice causing a structural variation.

The optical properties were investigated by using UV-vis spectroscopy in diffuse reflectance mode (Fig. 2) and the corresponding band gap was estimated using Tauc plot. The band gap of bulk-gC3N4 was determined to be about 2.77eV which is in close proximity to other reports on gC3N4 [Zou et al., 2018] and its theoretically reported value of 2.7eV [Kaining et al., 2016]. Upon B-doping, the band edge shifts marginally towards the longer wavelength with a weak tail in the visible region. This feature again confirms the presence of B in gC3N4 lattice where B, by replacing C, forms localized electronic states within the band gap [Kaininf et al., 2016] and contributes in enhancing the visible light absorption. The band edge is slightly blue shifted upon formation of NS (NS-gC3N4 and B-NS-gC3N4), which is expected due to the quantum confinement effect in nanostructures. When Ag NPs are decorated over NS and bulk B-doped-gC3N4, the band edge is red shifted mainly by the contribution of the surface plasmonic band of Ag NPs in the range of 420-520 nm. This shows that both B and Ag are beneficial for gC3N4 in absorbing visible light which further assists in photocatalytic reaction.

XPS spectra of C1s, N1s, Ag 3d and B1s levels for all the six samples are displayed in Fig.3. Bulk-gC3N4 showed typical three peaks in C1s band at 284.6 eV, 288.2 eV, and 286.2 eV attributed to adventitious carbon bond (C-C), carbon bonded to nitrogen in sp2 bonds (C-N-C) and trace amount of C-O bond respectively. As samples were calcinated in air, small amount of C-O is formed on the surface of the catalysts. Same set of peaks are present for all other samples irrespective of the type of strategies used to modify gC3N4. In case of N 1s level, four peaks due to, C-N=C at 398.7 eV, N-(C)3 at 399.7 eV, C-H-N at 401.1eV, and N-O at 404.2eV are present in all samples. The XPS peak of surface boron is distinctly visible only for the gC3N4 samples in the form of NS (B-NS-gC3N4 and Ag-B-NS-gC3N4) (Fig. 3). For B-doped bulk-gC3N4, the signal is very weak and cannot be distinguished evidently from the background. Single peak at 192.0 eV is detected in B 1s spectrum of B-NS-gC3N4 and Ag-B-NS-gC3N4 samples, which is attributed to C-NB group for B-doped-gC3N4. The spectrum of Ag 3d band (Fig. 3) is composed of two peaks at 368.5 and 374.5 eV with a spin energy separation of 6.0 eV. Both these peaks are standardized binding energy (BE) of Ag 3d5/2 and Ag 3d3/2 levels of metallic silver. The XPS results indicate that the chemical structure of gC3N4 remains intact upon doping with B and Ag loading. Boron is incorporated in the gC3N4 matrix by replacing C and forms B-N bonds which are exposed on the surface only upon the formation of NS. On the contrary, metallic Ag NPs are decorated over the gC3N4 bulk as well as on NS.

TEM images of bulk-gC3N4, NS-gC3N4, B-NS-gC3N4 and Ag-B-NS-gC3N4 are shown in Fig. 4. Morphology of B-bulk-gC3N4 and Ag-B-bulk-gC3N4 was similar to bulk-gC3N4, thus was not shown here. Sheet-like morphology is clearly visible for bulk-gC3N4 (Fig. 4a), with length of a few microns (1-2 µm), thus showcasing the graphitic nature. Upon formation of NS (Fig. 4b), the length reduces in the range of 100-400 nm which is further reduced to below 100 nm for NS of B-doped-gC3N4 (Fig. 4c). No major variation in the average length of NS is observed upon Ag loading (Fig. 4d). On comparing the HRTEM images of B-NS-gC3N4 (Fig. 4e) and Ag-B-Ns-gC3N4 (Fig. 4f), dark spot of diameter 5-10 nm is observed in latter, due to the presence of Ag NPs decorated on gC3N4 NS. For these NPs, an interplanar spacing of 0.24 nm was measured from the lattice fringes, which is attributed to (111) planes of metallic Ag. Morphological analysis by TEM clearly confirms the formation of NS which is decorated with Ag NPs in Ag-B-NS-gC3N4.

To verify the effect of nanosheet formation on the overall surface area of gC3N4, BET surface area was measured and summarised in Table1. In comparison to bulk samples (bulk-gC3N4, B-bulk-gC3N4 and Ag-B-bulk-gC3N4), the surface area is drastically improved for same samples in the form of NS (NS-gC3N4, B-NS-gC3N4 and Ag-B-NS-gC3N4). The highest surface area is recorded for B-NS-gC3N4 (57.3 m2/g), which decreases by a small amount (49.6 m2/g) upon loading with Ag NPs for Ag-B-NS-gC3N4 sample. It was also noticed that addition of B and Ag in bulk-gC3N4 disrupts the interlayer to increase the surface area by a small amount for Ag-B-bulk-gC3N4.

The effect of B and Ag on recombination of photogenerated charges can be qualitatively studied by measuring the emission radiation generated by the recombination process. PL spectra of all six samples were recorded at an excitation wavelength of 385 nm and is shown in Fig. 5. The intensity of the peak centred at 465 nm, due to band edge emission, is lower for NS-gC3N4 when compared to its bulk counterpart. This is mainly due to the shorter path length required for the photogenerated charges in case of NS. Upon boron doping, the intensity further reduces, thus indicating that boron doped in gC3N4 lattice can act as the trapping site for charges, resulting in reduced recombination process. However, lowest intensity of the PL peak is detected after loading with Ag NPs in both the cases of bulk and NS of B-doped-gC3N4. Thus, Ag in form of plasmonic NPs are highly effective in decreasing the recombination of photogenerated electron and hole pairs, owing to the electronic levels of Ag and gC3N4. Fermi level of Ag lies in between conduction band (CB) and valence band (VB) of gC3N4, which leads to migration of electrons from gC3N4 to Ag when junction between Ag and gC3N4 is formed. This electron transfer leads to band bending at the CB which generates electric field across junction. During light irradiation, the photogenerated electrons are unidirectionally transferred from CB of gC3N4 to Ag NPs under the influence of interfacial electric field resulting in better charge separation. To investigate the lifetime of charge carriers, time resolved photoluminescence (TRPL) spectra were acquired for all the samples. The decay curve was fitted using three components to obtain average lifetime of charge carriers. The average lifetime of bulk gC3N4 (1.45 ns) is increased to 2.05 ns when all three strategies are integrated in Ag-B-NS-gC3N4 sample, thus suggesting that both Ag and B effectively delay the recombination processes, with Ag being more efficient than B.

The photocatalytic activity of all the samples for degradation of Rh B dye was tested and presented in Fig. 6. Except for bulk gC3N4, all other modified samples were able to complete the degradation within the maximum time limit of 180 min, thus showing improved activity than bulk. Duration of 180, 160, 120, 80 and 60 mins were required by B-bulk-gC3N4, NS-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 to complete the reaction. The highest photocatalytic performance was obtained when all the three strategies of B-doping, NS formation and Ag loading were combined in Ag-B-NS-gC3N4 photocatalyst to produce 5.2 times higher reaction rate than bulk-gC3N4. When single strategy was adopted to modify gC3N4, for the case of B-doped-gC3N4 and NS-gC3N4, the activity is improved by 1.8 and 2 times, respectively, as compared to bulk gC3N4. This result indicates that NS formation is more efficient in improving the photocatalytic activity than B-doping. When two strategies were combined in the form of B-NS-gC3N4 and Ag-B-bulk-gC3N4 samples, the reaction rate increases by 2.6 and 4.2 times, respectively. This indicates that loading Ag is even more effective strategy in enhancing the photocatalytic activity.

Photocatalytic reactions are mainly driven by three factors - surface area, charge separation and visible light absorption capability of photocatalyst material. Thus, adopting a single strategy can improve only one of these factors, such as B-doped-gC3N4 and NS-gC3N4 increases the visible light absorption and surface area (as confirmed by UV-vis-spectroscopy and BET surface area), respectively. When both of these strategies are combined together in B-NS-gC3N4, surface area as well as visible light absorption increases, thus leading to higher activity as compared to individual strategy. As explained earlier, Ag NPs are very efficient in improving the charge separation by creating interfacial electric field across the junction with gC3N4 to cause fast transfer of photogenerated electrons from CB of gC3N4 to Ag NPs, avoiding recombination process. At the same time, plasmonic Ag NPs are also capable of creating intense SPR-induced electromagnetic field to increase the rate of electron-hole formation on the surface of gC3N4 in vicinity of Ag NPs. This mechanism assists in further improvement of the photocatalytic activity. Therefore, when Ag is loaded on B-NS-gC3N4, all three strategies create a synergic effect to promote the necessary features resulting in remarkable photocatalytic activity. The effect of Ag is also noticeable when loaded on B-bulk-gC3N4 in Ag-B-bulk-gC3N4 showing second best activity among all combinations. The important point to be noticed from above results is that Ag loading turns out to be the most effective strategy in increasing the activity and hence charge recombination seems to be the rate limiting process for photocatalytic reaction.

The mechanism involved in Ag-B-NS-gC3N4 for degradation of pollutant is schematically presented in Fig 7. Briefly, upon light irradiation Ag-B-NS-gC3N4, creates electron and hole pairs in gC3N4. The photogenerated electrons are immediately transferred from CB of gC3N4 to Ag NPs due to the electric field created at their interface, whereas photogenerated holes are left in the VB of gC3N4. The electrons on the Ag NPs react with adsorbed oxygen on the surface to form superoxide radicals (O2**.-**), while holes on gC3N4 readily react with surface bonded OH- to form hydroxyl radical species (OH.). Both these species are highly oxidizing species and are very active for the degradation process. In this mechanism, the present form of photocatalyst is very useful mainly considering the role of each engineering strategies involved. The B-doping is able to improve visible light absorption as confirmed by UV-vis spectra, thus producing large number of electron and hole pairs. It is also observed in PL spectra that B-doping decreases the recombination process by small amount. High surface area in case of nanosheet provides large number of active sites to promote photocatalytic reaction. On the other hand, Ag seems to be very proficient in separating electron and hole pair by generated electric field across the metal-semiconductor junction. Due to SPR effect, Ag NPs absorbs visible light to create extra electrons to produce higher number of superoxide radicals. The electromagnetic field generated by SPR effect of plasmonic Ag NPs is also able to create more electron-hole pairs across gC3N4 surface. All these stimulating effects generated in Ag-B-NS-gC3N4 assist the photo-degradation reactions in an outstanding manner.

**4. Conclusion:**

In summary, combination of three strategies has been employed to synthesize efficient Ag-B-NS-gC3N4 photocatalyst with enhanced visible light absorption, surface area, and electron hole pair separation. Thorough characterization studies show that B-doping results in superior photocatalytic performance with better visible light absorption capability. Subsequent acid treatment of B-gC3N4 to synthesize nanosheets resulted in higher exposed specific surface area with large number of active sites for organic pollutant adsorption during photocatalysis reaction. On the other hand, successive Ag loading has proven to be a highly effective strategy to facilitate efficient charge carrier transport and prolong charge carrier lifetime as a consequence of better electron-hole pair separation at the interface of gC3N4 and Ag NPs due to the interfacial electric field. On comparison of photocatalytic activity, it was found that B-gC3N4, B-NS-gC3N4 and Ag-B-NS-gC3N4 show 1.8, 2.6 and 5.2 times higher photocatalytic performance compared to pure gC3N4, respectively. This implies that combined effect of all the three strategies has great influence on enhancing the degradation ability of resultant photocatalyst with highest contribution from loading of Ag NPs.

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**Figure caption:**

**Figure 1:** XRD pattern of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.

**Figure 2:** UV-vis absorption spectra in diffuse reflectance mode of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.

**Figure 3:**XPS spectra of C1s and N1s of (a)bulk-gC3N4, (b) NS-gC3N4, (c) B-bulk-gC3N4, (d) B-NS-gC3N4, (e) Ag-B-NS-gC3N4 and (f) Ag-B-bulk-gC3N4 samples, Ag 3d of (a) Ag-B-NS-gC3N4 and (b) Ag-B-bulk-gC3N4 and B 1s of (a) Ag-B-NS-gC3N4 and (b) B-NS-gC3N4.

**Figure 4**: TEM images of (a) bulk-gC3N4, (b) NS-gC3N4, (c) B-NS-gC3N4 and (d) Ag-B-NS-gC3N4 and HRTEM images of (e) B-NS-gC3N4 and (f) Ag-B-NS-gC3N4.

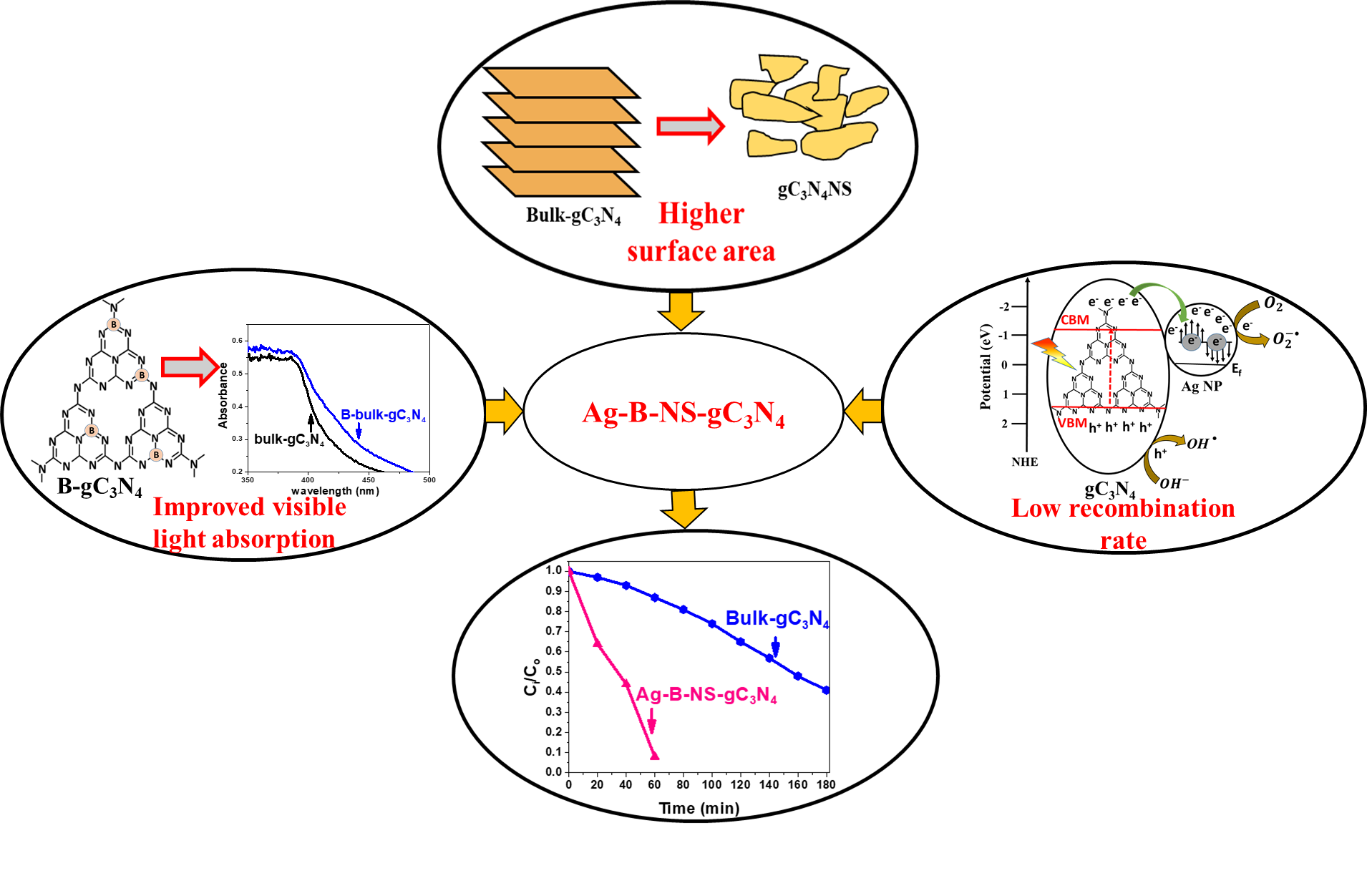
**Figure 5:** Photoluminescence emission spectra of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.

**Figure 6:** Comparison of photocatalytic degradation of Rh B under light irradiation in presence of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.

**Figure 7:** Mechanism involving charge separation in Ag loaded B-NS-gC3N4 during photodegradation of Rh B under light illumination.

**Table 1:** BET surface area and average lifetime of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.

**Graphical Abstract**

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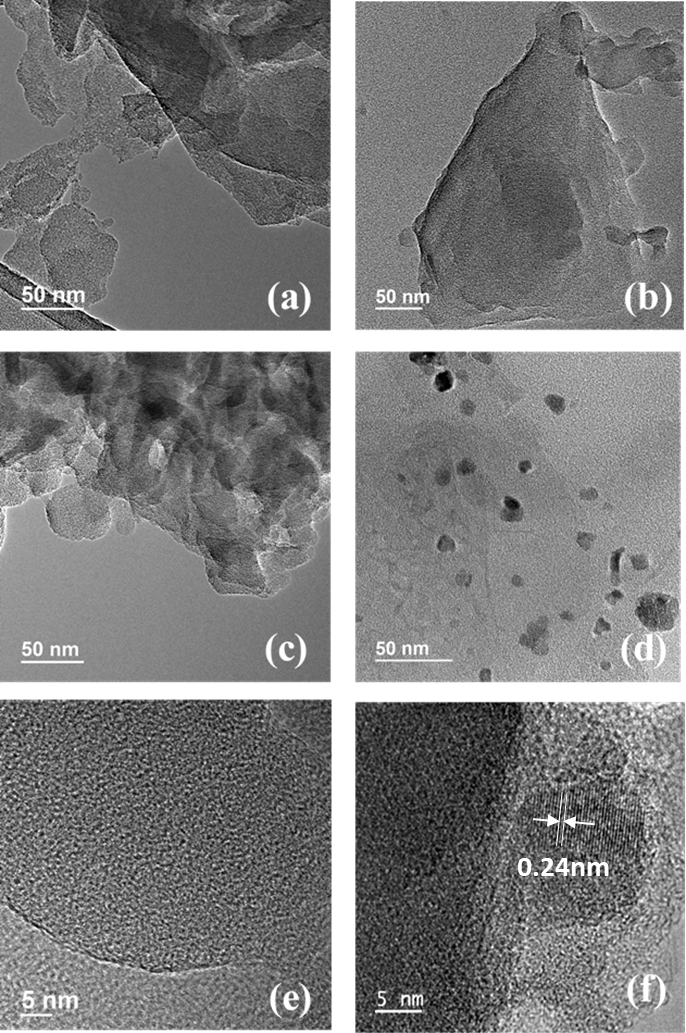
**Fig. 1:** XRD pattern of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.



**Fig. 2:** UV-vis absorption spectra in diffuse reflectance mode of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.



**Fig. 3:** XPS spectra of C1s and N1s of (a) bulk-gC3N4, (b) NS-gC3N4, (c) B-bulk-gC3N4, (d) B-NS-gC3N4, (e) Ag-B-NS-gC3N4 and (f) Ag-B-bulk-gC3N4 samples, Ag 3d of (a) Ag-B-NS-gC3N4 and (b) Ag-B-bulk-gC3N4 and B 1s of (a) Ag-B-NS-gC3N4 and (b) B-NS-gC3N4.



**Fig. 4:** TEM images of (a) bulk-gC3N4, (b) NS-gC3N4, (c) B-NS-gC3N4 and (d) Ag-B-NS-gC3N4 and HRTEM images of (e) B-NS-gC3N4 and (f) Ag-B-NS-gC3N4.

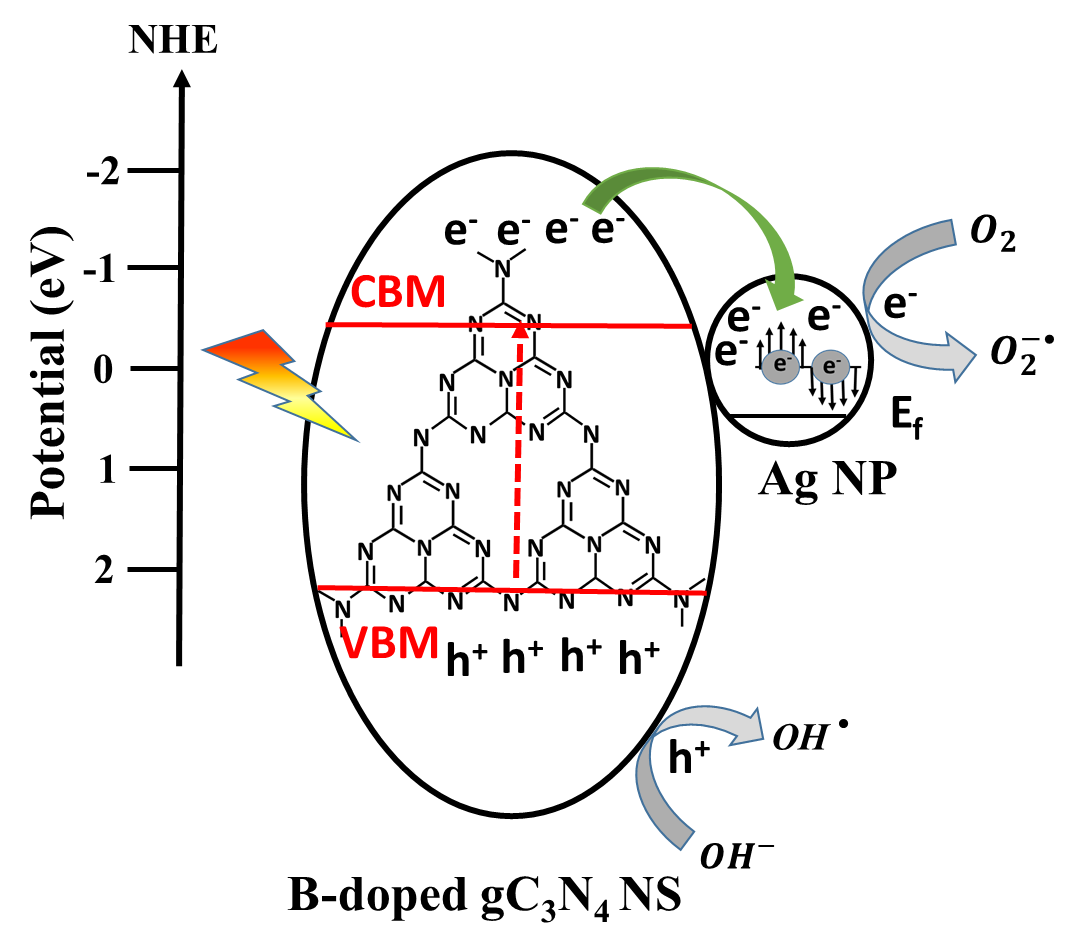


**Fig. 5:** Photoluminescence emission spectra of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.

.



**Fig. 6:** Comparison of photocatalytic degradation of Rh B under light irradiation in presence of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.



**Fig. 7:** Mechanism involving charge separation in Ag loaded B-NS-gC3N4 during photodegradation of RhB under light illumination.

|  |  |
| --- | --- |
| **Samples name** | **Surface area**  **(sq. m/gm)** |
| bulk-gC3N4 | 29.07 0.29 |
| NS-gC3N4 | 49.47 0.74 |
| B-bulk-gC3N4 | 34.18 0.48 |
| B-NS-gC3N4 | 57.27 0.91 |
| Ag-B-NS-gC3N4 | 49.64 0.64 |
| Ag-B-bulk-gC3N4 | 38.91 0.42 |

**Table 1:** BET surface area of bulk-gC3N4, NS-gC3N4, B-bulk-gC3N4, B-NS-gC3N4, Ag-B-bulk-gC3N4 and Ag-B-NS-gC3N4 samples.

**Supporting Information**

**For**

**Ag loaded B-doped-g C3N4 nanosheet with efficient properties for photocatalysis**

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**Fig. S1**: Comparison of photocatalytic degradation of Rh B under light irradiation in presence of Bulk-gC3N4, B4%-Bulk-gC3N4, B6%-Bulk-gC3N4 B8%-Bulk-gC3N4 B10%-Bulk-gC3N4.



**Fig. S2**: Comparison of photocatalytic degradation of Rh B under light irradiation in presence of B-NS-gC3N4, 5%Ag-B-NS-gC3N4, 10%Ag-B-NS-gC3N4 and 15%Ag-B-NS-gC3N4.