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FLUORESCENCE QUENCHING ON MESOPOROUS CARBON NITRIDE BY PHENOL AND ANILINE

(Pelindapan Pendarfluor pada Karbon Nitrida Mesoliang oleh Fenol dan Anilina)

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In this study, fluorescence q 12 ching on mesoporous carbon nitride (MCN) was examined in the presence of phenol and aniline.

MCN was synthesized by thermal polymerization of cyanamide using nanocolloidal silica (7 nm) as a hard template and characterized by X-ray diffractometer (XRD), Fourier transform infrared (FTIR), transmission electron microscope (TEM), specifically specif

Keywords: mesoporous carbon nitride, phenol, aniline, fluorescence quenching, adsorption

Abstrak

Dalam kajian ini, pelindapan pendarfluor pada karbon nitrida mesoliang (MCN) telah diperiksa dengan kehadiran fenol dan anilina. MCN telah disintesis secara pempolimeran haba sianamida menggunakan silika nanokoloid (7 nm) sebagai templat keras dan telah dicirikan dengan pembelauan sinar-X (XRD), inframerah transformasi Fourier (FTIR), mikroskop elektron transmisi (TEM), luas permukaan, taburan saiz liang, dan spektroskopi pendafluor. MCN menunjukkan dua panjang gelombang pengujaan pada 275 dan 370 nm oleh kerana kehadiran masing — masing N=C dan N-C sebagai kumpulan terminal. Kedua-dua panjang gelombang pengujaan memberikan hanya satu puncak pelepasan pada kira-kira 460 nm.Sama ada dengan kehadiran fenol atau anilina, keamatan pemancaran daripada MCN disahkan mengalami pelindapan dalam fungsi linear terhadap kepekatan fenol atau anilina sebagai molekul pelindap. Kumpulan terminal N-C didapati mempunyai interaksi lebih baik sedikit untuk fenol dan anilina berbanding dengan kumpulan N=C. Oleh kerana keamatan pemancaran MCN didapati lebih terlindap oleh molekul fenol berbanding dengan anilina, adalah dicadangkan bahawa MCN berinteraksi lebih kuat dengan fenol dibandingkan dengan anilina. Keputusan ini juga disokong oleh data penjerapan, dengan MCN memberikan penjerapan yang lebih tinggi terhadap fenol dibandingkan dengan anilina.

Kata kunci: karbon nitrida mesoliang, fenol, anilina, pelindapan pendarfluor, penjerapan

Introduction

Carbon nitrides (CN) are carbon materials in which some carbon atoms in the graphite-like structure are replaced by some nitrogen atoms in a regular manner. Generally, CN stands for a large family of related compounds (C_xN_y) [1 – 3]. Even though the CN has been reported since the 1990s, this material still receives considerable attention because of its versatility until nowadays. The CN is used in various applications while the material is just composed of C, N and H atoms [4]. Other factors that make CN being well explored is because of the abundance of these atoms on earth, non-toxic and easy to be synthesized into a desired shape where the pore of carbon nitride is tunable to form mesoporous carbon nitride (MCN) [1, 5, 6]. MCN has a larger specific surface area than the CN and it facilitated better performance for 33 ous applications [7 – 12]. Moreover, CN has many advantageous properties that make it an interesting material, such as high thermal and chemical stability, and great optical properties [13 – 15].

Recently, application of CN and MCN as fluorescence sensor has been a significant interest. Owing to the presence of fluorescent active sites (tri-s-triazine units), CN and MCN are potential materials to be used as an optical sensor. Despite of the fact that these materials have a strong fluorescence pro 28 y, less attention was paid on its sensing ability until there was a report showing the utilities of cubic MCN as all-in-one chemosensor for sensing of trace amount of metal ions in aqueous solution [16]. Since the cubic MCN has the highest sensitivity and selectivity towards copper ions, further modification was also carried out by the research group on the cubic MCN incorporating copper ions for sensing of cyanide ions, which are very poisonous and used abundantly in metal plating industry [17]. CN has been also reported as a 47 ficient fluorosensor for detection of copper ions. It was suggested that the fluorescence quenching was due to strong coordination of the Lewis basic sites on the CN with the Cu ions [18]. Besides metal and cyanide ions, recent studies showed that nitrogen-containing compounds can be also detected by fluorescence quenching. It was reported that MCN was able to be used as a fluorescence sensor for *N*-nitrosopyrrolidone (NPYR) [7]. The surface interaction was proposed to involve electrostatic interactions between the electropositive N and 46 tropositive N at the NPYR molecules and terminal N-C groups of the MCN, respectively. CN was also reported as a fluorescence chemical sensor for nitrate ions [19].

In this study, fluorescence quenching on M8N was investigated in the presence of aromatic compounds, which are phenol and aniline. Phenol is considered as one of the organic pollutants that can be easily found in industrial wastewater [20 – 22]. The wastewater containing phenol may cause the contamination of groundwater and it will give bad effects for ecosystems and also for human. Meanwhile, aniline is also recognized as one of the high priority pollutants that are released from several industries, such as various manufacturers of dyes, and rubber, which caused wide distribution of aniline in the environment [23 – 25]. As an adsorbent, MCN possesses well-ordered porosity, high surface area, and basic groups (NH and NH₂) in the structures, which are the important criteria for the removal of organic pollutants. Due to the nitrogen rich structure of MCN with basicity property, it showed remarkable adsorptions towards one of the acidic persistent organic pollutants, i.e. phenol [26]. On the other hand, the MCN was also reported showing superior capability to adsorb aniline than the CN and MCM-41 [8, 9]. Therefore, based on these reported studies, the fluorescence quenching on MCN by phenol and aniline is feasible. This study demonstrated that MCN can be also used as a fluorescence sensor for the aromatic compounds.

Materials and Methods

Preparation of MCN

The MCN was prepared by a thermal polymerization of cyanamide as the precursor, in the similar way to the method reported in other literatures [7 – 9]. The cyanamider as mixed with the colloidal silica as a hard template 7 nm (mass ratio was 1:1), followed by stirring and heating at 363 K 2 til a white powder was formed. The obtained 44 te powder was ground and then heated at a temperature of 823 K for 4 hours. In order to obtain the MCN, the silica template was removed by treating the MCN powder with the NH₄HF₂ solution (4 M, 400 mL) under vigorous stirring and also with ethanol and distilled water. This washing step was repeated for three times. After washing and filtration, the powder was dried at 343 K overnight in an oven.

Characterizations

The structure of the MCN was determine 5 by an X-ray diffractimeter (XRD) on a Bruker D8 Advance Diffractometer at ambient temperature, with a scan rate of 0.05 s⁻¹ using Cu-K α radiation where λ equals to

1.5406 Å at 40 kV and 40 mA. To determine the structural information and functional groups of the MCN, a Fourier transform in 26 ed spectroscopy (FTIR) was employed using a FTIR Nicolet iS50 Thermo Scientific. For pellet preparation, a small amount of the MCN was mixed with potassium bromide (KBr) and put under a pellet press. To obtain a better background correction 21 trogen gas was purged into the system before the analysis to remove all humidity. Morphology of the MCN was measured using a transmission electron microscope (TEM) on a JEOL JEM-2100. The MC7 powder was first dispersed well in acetone, and then put onto the copper grid to be observed with the TEM. The surface area of the MCN was determined by using the Brunauer-Emmet-Teller (BET) equation, while the pore size of the MCN was determined using the Barret-Joyner-Halenda (BJH) from adsorption-desorption isotherm at 77 K obtained on a Quanta chrome Autosorb-1 instrument. The fluorescence properties were measured using a JASCO Spectrofluorometer FP-8500 to obtain the excitation and emission spectra. The excitation

wavelengths used to monitor the emission spectra were 275 and 370 nm, while the emission wavelength used to

monitor the excitation spectra was 460 nm. The analysis was carried out at room temperature.

Fluorescence quencl 42g

The quenching tests were carried out with various concentrations of phenol and aniline on the MCN (50 mg) at room temperature. The excitation wavelengths at 275 and 370 nm of the MCN were determined by observing the emission spectrum at 460 nm. Meanwhile, the emission wavelength was determined by observing excitation spectra at both excitation wavelengths. From various amounts of phenol solutions (300 – 600 ppm), 20 µL of the solution was taken and then intraduced onto the MCN for the quenching test for phenol. The changes in the emission spectra of the MCN were then measured at excitation wavelengths of 275 and 370 nm. As for the quenching test for aniline, the same steps were followed using various concentrations of aniline solution (200 –700 ppm).

Adsorption tests

The adsorption process of phenol or aniline onto the MCN was conducted under dark condition using various concentrations of phenol (60 – 400 ppm) and aniline (100 – 500 ppm) at room temperature. Different set 40 of concentrations were selected due to the different behaviors of the observable adsorption. A magnetic bar was put in a 30 mL beaker and 10 mL of phenol or aniline solution was poured into the beaker together with the MCN (50 mg). Inside a closed box, the beaker containing the mixture was placed on the st 20 g plate and stirred for 6 hours. After filtration and separation from the MCN, the solution was analysed using gas chromatography with a flame ionization detector (GC-FID) on an Agilent 7820AGC-FID. The adsorbed amount was determined by subtracting the initial phenol or aniline with the amounts of remaining phenol or aniline after the adsorption test.

Results and Discussion

Proper 39 of MCN

Shown in Figure 1(a) is the XRD puttern of the prepared MCN, which gave two diffraction peaks at 2θ of 13.1 and 27.4°. The 16 ak at 2θ of 13.1° was corresponding to the distance of 0.66 nm, which indicated the in-plana 1 epeating units and structural packing motif, such as hole-to-hole distance of the nitride 125 s, while the intense diffraction peak at the 2θ of 27.4° could be indexed as (002) for graphitic mater 38 giving an interlayer distance of 0.325 nm for the graphite-like stacking of the conjugated aromatic CN layers. This result is in good agreement with values reported elsewhere [4, 7 – 11, 13, 14]. Therefore, it can be concluded that the MCN sample was successfully synthesized.

The functional groups present on the MCN were investigated by FTIR spectroscopy. As shown in Figure 1(b), the absorption peaks of the MCN were 24 servable at the region of 809, 1200 − 1700 and 3000 − 3700 cm⁻¹. The bands at 809, 1252, 1319, 14218 and 1646 cm⁻¹ could be attributed to the vibration modes of heterocyclic tri-s-triazine ring (C₆N₇) units, while the broad peak at 3163 cm⁻¹ could be a 23 ned to two overlapping bands of stretching modes of N-H and O-H groups [9, 11, 27]. The peak at 2177 cm⁻¹ could be attributed to the presence of C≡N or N=C=N group, which was reported in MCN prepared via thermal polymerization reaction [27]. These FTIR spectra supported that the MCN sample was successfully prepared.

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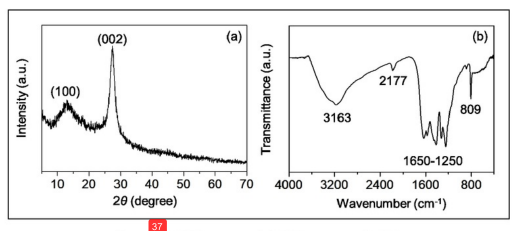


Figure 1. (a) XRD pattern and (b) FTIR spectrum of MCN

The morphology of the prepared MCN was observed by TEM 36 shown in Figure 2. It can be seen that the MCN has uneven layers due to the graphitic structures of the MCN. Figure 3(a) shows the nitrogen adsorption-desorption isotherms and Figure 3(b) shows the Barrett-Joyner-Halenda (BJH) pore size 50 tribution of the MCN. The results suggested that the MCN has type IV isotherm for mesopore 19 material with hysteresis loop at relative pressure of 0.5 – 1 due to the capillary 49 densation in pores. The BET specific surface area of the MCl 35 as determined to be 287 m² g⁻¹. Meanwhile, for 32 H pore size distribution, the MCN showed a narrow and sharp pore size of 7 nm. This pore size reflected the size of the silica template that was used for the preparation of the MCN.

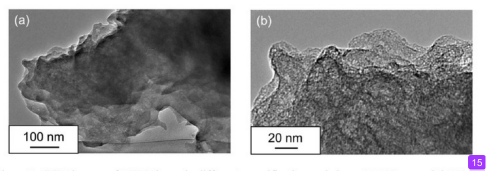


Figure 2. TEM images of MCN shown in different magnification scale bars, (a) 100 nm and (b) 20 nm

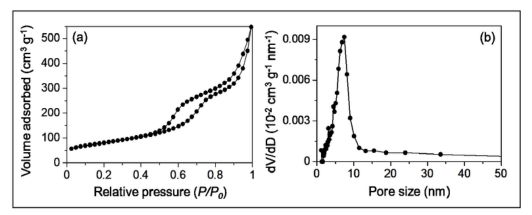


Figure 3. (a) Nitrogen adsorption-desorption isotherm and (b) BJH pore size distribution of MCN

Figure 4(a) disp 3 ys the excitation spectrum of the MCN. It was clear that the M10 I exhibited two excitation peaks at 275 and 372 nm. The excitation peak at 275 nm can be attributed to the π to π^* electronic transition of the N=C 10 ups in the aromatic 1,3,5-triazine ring of MCN. Meanwhile, the other excitation peak at 370 11 can be attributed to n to π^* electronic transition of terminal N-C group involving lone pairs of nitrogen atoms. As can be seen from Figure 4(b), the MCN gave one emission peak at a wavelength of 460 nm when the MCN was excited either at 275 or 370 nm. These results agree well with previous reports [7, 9, 19].

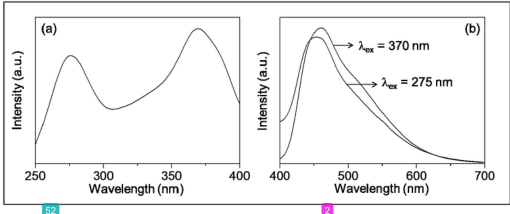


Figure 4. (a) 2 citation and (b) emission spectra of MCN that were monitored at emission wavelength of 460 nm and excitation wavelengths of 275 and 370 nm

Fluorescence quenching

Quenching tests were conducted on the MCN by the addition of various concentrations of phenol or aniline in order to investigate the sites where the interactions occurred either at N=C groups or terminal N-C groups. The quenching tests were determined by observing the changes in the emission intensity of the MCN sample before and after the addition of phenol or aniline. Since one emission spec 31 at around 460 nm was observed for both excitations at 275 and 370 nm, the quenching tests w 9 carried out by monitoring the intensity of the emission peak at 460 nm excited at both excitation wavelengths. As illustrated in Figure 5(a) and (b), the emission intensities of the MCN

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excited at both excitation wavelengths of 275 and 370 nm (N=C and terminal N-C emission sites), respectively decreased with the 6 dition of the phenol. This result showed that phenol acted as a quenching agent to deactivate the emission sites. When the added phenol concentration increased, the emission intensity was further decreased. The decrease in the emission intensity would be due to the interactions between the emission sites of MCN at an emission wavelength of 460 nm and the added phenol.

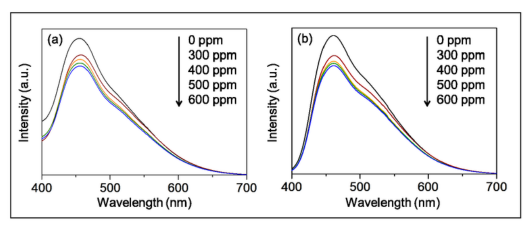


Figure 5. Fluorescence quenching on MCN in the absence and presence of phenol (0-600 ppm) when excited at (a) N=C and (b) terminal N-C groups

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In order to have a better understanding about the interestion of MCN with phenol, the quenching results were plotted according to the Stern-Volmer plot following the equation 1:

$$Io/I = K_{SV}[Q] + 1 \tag{1}$$

where Io and I are fluorescence intensities observed in the absence and presence of phenol, Q is the concentration of phenol, and K_{SV} is the Stern-Volmer quenching constant.

Based on the Stern-Volmer equation, the relative emission intensity would be expressed as the function of phenol concentra 2 n when the same amount of MCN and the constant illumination intensity are taken in account. Figure 6 shows the Stern-Volmer plots of the relative emission intensity of MCN with various concentrations of phenol. In the range of 300 – 600 ppm, the relative emission intensity excited at both emission sites gave a linear Stern-Volmer plot with the concentrations of phenol. The linear Stern-Volmer plot indicated that the MCN could detect phenol using 23 fluorescence spectroscopy. Meanwhile, the quenching efficiency of phenol on the emission site of MCN could be obtained from the slope of the linear plot, which was matched to the quenching rate constant (K_{SV} value). It was shown that the K_{SV} values for the emission sites monitored at excitation wavelengths of 275 and 370 nm were 4×10^{-4} and 5×10^{-4} ppm⁻¹, respectively. These results suggested that the emission sites of MCN excited at 275 and 370 nm have almost equal interactions with phenol, which the terminal N-C groups gave slightly better interaction than the N=C groups.

Figure 7(a) and (b) show the changes on the emission intensities excited at 275 and 375 nm, respectively in the presence of aniline. Similar to the phenol addition, the addition of aniline quenched the intensity of the MCN. Aniline also acted as a quenching agent that deactivated the emission sees of MCN. When the concentration of aniline increased, the emission intensity was also further reduced. The decrease in the emission intensity of the MCN suggested that there were interactions between the N=C and terminal N-C emission sites and the aniline. As previously mentioned, the MCN has two excitation sites at 275 and 370 nm, which belong to the N=C groups

 $(\pi - \pi^*)$ in the aromatic 1,3,5-triazine and the terminal N-C groups $(n - \pi^*)$, respectively. In order to investigate which sites were favoured to the aniline, the Stern-Volmer plot was performed.

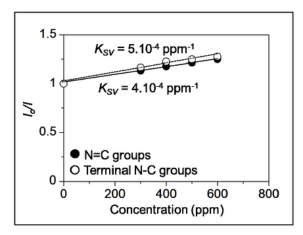


Figure 6. Stern Volmer plots of phenol quenching on MCN at N=C and terminal N-C groups

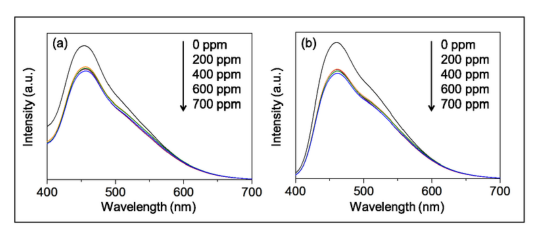


Figure 7. Fluorescence quenching on MCN in the absence and presence of aniline (0 – 700 ppm) when excited at (a) N=C and (b) terminal N-C groups

Shown in Figure 8 is the linear Stern-Volmer plots between the relative emission intensity of MCN with various concentrations of aniline in the range of 200 –700 ppm. Both emission sites, the N=C and terminal N-C groups showed the stern-Volmer plots with the addition of aniline. K_{SV} values obtained were 3×10^{-4} and 4×10^{-4} ppm⁻¹ for the N=C and terminal N-C groups, respectively. The slightly higher K_{SV} value of the terminal N-C group suggested that similar to the results when using phenol as the quencher agent, aniline also favored the terminal N-C groups. Based on the fluorescence quenching tests, it was demonstrated that the MCN showed good interactions with phenol and aniline.

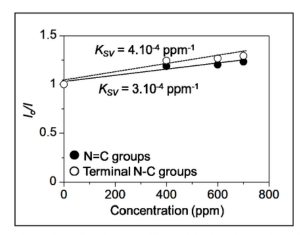


Figure 8. Stern Volmer plots of aniline quenching on MCN at N=C and terminal N-C groups

Adsorption tests

Adsorption capability of the M1N towards phenol or aniline was examined by using different concentrations. The adsorption test for phenol was carried out at room temperature with the concentration of phenol was varied from 60 to 400 ppm. Figure 9 presents the results of the adsorption test of phenol on the MCN. The amount of adsorbed phenol increased with the increase of the phenol concentration when it was below 100 ppm. Further increase of the phenol concentration at more than 100 ppm did not affect much the amount of adsorbed phenol. Figure 9 also shows the results of aniline adsorption on the MCN with various concentrations from 100 to 500 ppm. Separated experiments confirmed that no aniline adsorption occurred when the aniline concentration was lower than 100 ppm. At concentration of 100 ppm and above, the MCN showed an increased amount of adsorption with the increase of aniline concentration up to 500 ppm. Compared to the adsorption of phenol, higher amount of aniline was required for MCN to act as an adsorbent.

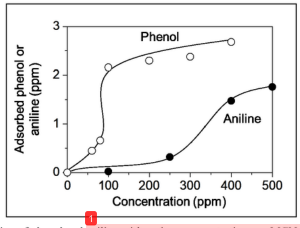


Figure 9. Adsorption of phenol and aniline with various concentrations on MCN at room temperature

From the adsorption test analyses, it can be proposed that phenol and aniline were successfully adsorbed on the MCN. However, under the same concentration for instance at 100 ppm, the MCN gave better adsorption towards phenol than the aniline, which gave adsorbed amounts of 2.16 and 0.02 ppm, respectively. The higher adsorbed

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amount of phenol than aniline might be due to the fact that phenol is a weak acid, while aniline is a weak base. Therefore, basicity of the MCN would favor phenol rather than aniline. It can be proposed that phenol would give stronger interaction to the MCN, which in ture gave higher K_{SV} value than the aniline. As has been reported elsewhere [7], the presence of electronegative N at the terminal N-C groups of the MCN provided the possibility for the formation of electrostatic interactions towards the analyte moles less. As phenol is a weak acid, the acid-base electrostatic interactions would occur between the electronegative N at the terminal N-C groups of the MCN and the electropositive p sholic proton. As for aniline, the electrostatic interactions would occur between the electronegative N at the terminal N-C groups of the MCN and the electropositive H at the NH₂ group of the aniline, which was less electropositive than the phenolic proton.

Conclusion

Characterizations by XRD, FTIR, TEM, surface a 12 and pore size analyzer, and fluorescence spectroscopy confirmed that the MCN was prepared successfully by thermal polymerization of cyanamide using nanocolloidal silica as the hard template. The MCN was shown to have good interactions with phenol and aniline, where its emission intensity was quenched almost linearly with the addition of these aromatic compounds. It was obtained that the terminal N-C groups gave slightly higher quenching efficiency than the N=C groups for both phenol and aniline. Phenol gave better interactions towards the emission sites of the MCN, owing to the stronger adsorption of phenol on to the MCN than the aniline caused by the acid-base electrostatic attractive force.

1 Acknowledgement

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References

- Vinu, A. (2008). Two-dimensional hexagonally-ordered mesoporous carbon nitrides with tunable pore diameter, surface area and nitrogen content. Advanced Functional Materials, 18(5): 816 827.
- 2. Teter, D. M. and Hemley R. J. (1996). Low-compressibility carbon nitride. Science, 271(5245): 53 55.
- Corkill, J. L. and Cohen, M. L. (1993). Calculated quasiparticle band gap of β-C₃N₄. Physical Review B, 48(23): 17622 – 17624.
- 4. Goettmann, F., Fischer, A., Antonietti, M. and Thomas, A. (2006). Chemical synthesis of mesoporous carbon nitrides using hard templates and their use as a metal-free catalyst for Friedel-Craft reaction of benzene. Angewandte Chemie International Edition, 45(27): 4467 – 4471.
- Gillan, E. G. (2000). Synthesis of nitrogen-rich carbon nitride networks from an energetic molecular azide precursor. Chemistry of Materials, 12(12): 3906 – 3912.
- Komatsu, T. (2001). Attempted chemical synthesis of graphite-like carbon nitride. *Journal of Materials Chemistry*, 11: 799 801.
- Sam, M. S., Lintang, H. O., Sanagi, M. M., Lee, S. L. and Yuliati, L. (2014). Mesoporous carbon nitride for adsorption and fluorescence sensor of N-nitrosopyrrolidine. Spectrochimica Acta Part A: Molecular and Biomolecular Spectroscopy, 124: 357 – 364.
- 8. Sam, M. S., Lintang, H. O., Sanagi, M. M., Lee, S. L. and Yuliati, L. (2014). Adsorption of aniline using novel mesoporous carbon nitride. *Advanced Materials Research*, 95: 135–139.
- Sam, M. S., Tiong, P., Lintang, H. O., Sanagi, M. M., Lee, S. L. and Yuliati, L. (2015). Mesoporous carbon nitride as a metal-free catalyst for the removal of aniline. RSC Advances, 5: 44578 – 44586.
- Lee, S.C., Lintang, H.O. and Yuliati, L. (2012). A urea precursor to synthesize carbon nitride with meso porosity for enhanced activity in the photocatalytic removal of phenol. *Chemistry An Asian Journal*, 7(9): 2139 2144.
- Cui, Y., Zhang, J., Zhang, G., Huang, J., Liu, P., Antonietti, M. and Wang X. (2011). Synthesis of bulk and nanoporous carbon nitride polymers from ammonium thiocyanate for photocatalytic hydrogen evolution. *Journal of Materials Chemistry*, 21: 13032 – 13039.
- Wang, X., Maeda, K., Chen, X., Takanabe, K., Domen, K., Hou, Y., Fu, X. and Antonietti, M. (2009). Polymer semiconductors for artificial photosynthesis: hydrogen evolution by mesoporous graphitic carbon nitride with visible light. *Journal of the American Chemical Society*, 131(5): 1680 – 1681.

- Wang, Y., Wang, X. and Antonietti, M. (2012). Polymeric graphitic carbon nitrides as a heterogeneous organocatalyst: from photochemistry to multipurpose catalysis to sustainable chemistry. *Angewandte Chemie Inter*national Edition, 51(1): 68 – 89.
- Thomas, A., Fischer, A., Goettmann, F., Antonietti, M., Müller, J.-O., Schlögl, R. and Carlsson, J. M. (2008).
 Graphitic carbon nitride materials: Variation of structure and morphology and their use as metal-free catalyst. *Journal of Materials Chemistry*, 18(41): 4893 – 4908.
- Zhang, Y., Mori, T. and Ye, J. (2012). Polymeric carbon nitrides: Semiconducting properties and emerging applications in photocatalysis and photoelectrochemical energy conversion. Science of Advanced Materials, 4(2): 282 – 291.
- Lee, E. Z., Jun, Y. S., Hong, W. H., Thomas, A. and Jin, M. M. (2010). Cubic mesoporous graphitic carbon(IV) nitride: an all-in-one chemosensor for selective optical sensing of metal ions. *Angewandte Chemie International Edition*, 49 (50): 9706 9710.
- Lee, E. Z., Lee, S. U., Heo, N. S., Stucky, G. D., Jun, Y. S. and Hong, W. H. (2012). A fluorescent sensor for selective detection of cyanide using mesoporous graphitic carbon(IV) nitride. *Chemical Communications*, 48(33): 3942 – 3944.
- Tian, J., Liu, Q., Asiri, A. M., Al Youbi, A. O. and Sun, X. (2013). Ultrathin graphitic carbon nitride nanosheet: a highly efficient fluorosensor for rapid, ultrasensitive detection of Cu²⁺. Analytical Chemistry, 85(11): 5595 – 5599.
- 19. Alim, N. S., Lintang, H. O. and Yuliati, L. (2015). Fabricated metal-free carbon nitride characterizations for fluorescence chemical sensor of nitrate ions. *Jurnal Teknologi*, 76(13): 1 6.
- Dabrowski, A., Podkościelny, P., Hubicki, Z. and Barczak, M. (2005). Adsorption of phenolic compounds by activated carbon - A critical review. *Chemosphere*, 58(8): 1049 – 1070.
- 21. Busca, G., Berardinelli, S., Resini, C. and Arrighi, L. (2008). Technologies for the removal of phenol from fluid streams: A short review of recent developments. *Journal of Hazardous Materials*, 160(2-3): 265 288.
- Ahmed, S., Rasul, M. G., Martens, W. N., Brown, R. and Hashib, M.A. (2010). Heterogeneous photocatalytic degradation of phenols in wastewater: a review on current status and developments. *Desalination*, 261(1-2): 3 18
- Soares, B. G., Amorim, G. S., Souza Jr, F. G., Oliveira M. G. and da Silva, J. E. P. (2006). The in situ polymerization of aniline in nitrile rubber. Synthetic Metals, 156 (2-4): 91 98.
- Zhao, R., Tan, C., Xie, Y., Gao, C., Liu, H. and Jiang, Y. (2011). One step synthesis of azo compounds from nitroaromatics and anilines, *Tetrahedron Letter*, 52(29): 3805 – 3809.
- Puig, A., Ormad, P., Roche, P., Sarasa, J., Gimeno, E. and Ovelleiro, J. L. (1996). Wastewater from the manufacture of rubber vulcanization accelerators: Characterization, downstream monitoring and chemical treatment.
 Journal of Chromatography A, 733 (1–2): 511 522.
- Haque, E., Jun, J. W., Talapaneni, S. N., Vinu, A. and Jhung, S. H. (2010). Superior adsorption capacity of mesoporous carbon nitride with basic CN framework for phenol. *Journal of Materials Chemistry*, 20(48): 10801 – 10803.
- Miller, D. R., Wang, J. and Gillan, E. G. (2002). Rapid, facile synthesis of nitrogen-rich carbon nitride powders. *Journals of Materials Chemistry*, 12(8): 2463 – 2469.