Electron Deficient Monomers that Optimize Nucleation and Enhance the Photocatalytic Redox Activity of Carbon Nitriles

Guigang Zhang, * Minghui Liu, Tobias Heil, Spiros Zafeiratos, Aleksandr Savateev, Markus Antonietti, and Xinchen Wang*

Abstract: Polymeric carbon nitride (PCN) is usually synthesized from nitrogen-rich monomers such as cyanamide, melamine, and urea, but is rather disordered in many cases. Now, a new allotrope of carbon nitride with internal heterostructures was obtained by co-condensation of very electron poor monomers (for example, 5-amino-tetrazole and nucleobases) in the presence of mild molten salts (for example, NaCl/KCl) to mediate the polymerization kinetics and thus modulate the local structure, charge carrier properties, and most importantly the HOMO and LUMO levels. Results reveal that the as-prepared NaK-PHI-A material shows excellent photo-redox activities because of a nanometric hetero-structure which enhances visible light absorption and promotes charge separation in the different domains.

The direct conversion of solar energy into storable chemical fuel molecules (for example, H2) via photocatalytic water splitting has been regarded as a feasible way to decrease the strong dependence on traditional fossil fuels.1-3 The key of this technique however is the development of low-cost, stable, and highly efficient photocatalysts, which are able to catalyze the desired reactions in an efficient manner. Among all the investigated photocatalysts, polymeric carbon nitride (PCN) has drawn particular interests owing to its inherent advantages including being metal-free, ease of fabrication, chemical and thermal robustness, and visible light activity.4-9 However, direct photocatalytic splitting of water by PCN is rather challenging because PCN in its pristine form features only moderate water oxidation activity.10-12

In principle, PCN is usually prepared by thermal polymerization of carbon and nitrogen containing monomers (for example, cyanamide, melamine, urea, and many more) at elevated temperatures (usually 550°C).13-15 The polymerization process is hindered at later stages by the sluggish deamination kinetics and thus generates PCN with incomplete condensation and lower local order; it is however potentially still strongly oxidizing (VB = 1.82 V, vs. RHE). The created overpotential is however not enough for coca-
talyt-free water oxidation, which is, despite its apparent simplicity, a remarkably complicated process. PCN however oxidizes many organic compounds, as described in some recent reviews.16,17 Recent progress from our group and other groups reveal that the use of suitable molten salts to mediate the polymerization significantly decreases the number of defects and improves the photocatalytic performance.16-21 Molten salts (for example, LiCl/KCl, NaCl/KCl) dissolve monomers and intermediates and give a liquid medium for polycondensation, thus improving the local order. Salt-melt synthesis provides further the opportunity to synthesize carbon nitride as new allotropes and iso-forms, many coming with higher oxidation potential. Recently, we were able to show that a more noble carbon nitride with highly crystalline structure could be obtained when electron deficient triazole/tetrazole based monomers were polycondensed in the LiCl/KCl salt melt.22-25 The as-prepared polymers bear much better oxidation activity thanks to the positively shifted HOMO (VB = 2.5 V, vs. RHE). However, here the simultaneous downshift of the LUMO (CB = −0.1 V, vs. RHE) results in a weakening of the photocatalytic reduction activity.24 Interestingly, excellent reduction ability rather than oxidation activity could be obtained when the tetrazole was polymerized in an alternative NaCl/KCl salt melt.26 It is thus exciting to search rationally for other and even more active carbon nitrides, namely more stable conjugated system with both strong oxidation power and good reduction ability for photocatalytic redox reactions, and here we use copolymerization with more electron deficient monomers in diverse molten salts which will turn out to nucleate different allotropes.

The systems are synthesized by copolymerization of electron-deficient aminotetrazole in NaCl/KCl molten salts, while we use nucleobases as natural comonomers to introduce local D-A junctions in spurious amounts (ca. 0.3 wt %). Interestingly, already that little comonomer obviously changes the local electron density, optical absorption,
The lack of the typical peak at 13\,\text{cm}^{-1} are observed, revealing the formation of the
a) Fourier filtered HR-TEM image of the selected area of
Raman spectra were conducted,
a) High-resolution N1 sX PS analysis and b) UPS analysis of
weak absorption
2019
and the strong variations of surface cyano groups
reveal the higher extension of polycondensation
bases is revealed in AC-HRTEM, where we can observe
the materials are presented in the Supporting Information,
the materials.S urvey spectra (Supporting Information, Fig-
hemation of the as-prepared heptazine-based carbon nitrides.
More information about the materials synthesis is found
in Experimental Section in the Supporting Information. In
brief, the melon-based carbon nitrides as references were
synthesized by thermal polymerization of the monomers
dicyandiamide, melamine, and urea (denoted as CND, CNM,
and CNU). NaK-PHI-based carbon nitrides were obtained by
copolymerization of 5-amino-tetrazole (1) without or with
nucleobases (2–5) in salt melts (denoted as NaK-PHI, NaK-
PHI-A, NaK-PHI-G, NaK-PHI-T, and NaK-PHI-C, respectively).
The as-synthesized poly(heptazine imide) (PHI)-based
carbon nitrides were firstly characterized by XRD, FTIR, and
XPS. As seen from the Supporting Information, Figure S1,
such carbon nitrides show only one diffraction peak located
27.4^\circ, corresponding to the graphic layer stacking distance of
the units.[26–28] The lack of the typical peak at 13^\circ, which is
related to the periodic in plane repeat of heptazine units,
and the broad diffraction peaks of PHI-based carbon nitride
already indicate a local rearrangement of the packing units,
which will become more clear in the TEM images (see below).
FTIR (Supporting Information, Figure S2) of the samples
reveals that both sample sets of carbon nitrides present
similar breathing and stretching modes of the heptazine units
at 700–850\,\text{cm}^{-1} and 1200–1600\,\text{cm}^{-1}, respectively. For the
PHI-based carbon nitrides, evident variations at 987\,\text{cm}^{-1}
and 1065\,\text{cm}^{-1} are observed, revealing the formation of the
symmetric and asymmetric variations of imide bound metal
ions (here Na/K-NC\_ groups).[22,24,29] The weak absorption
band of surface uncondensed amino groups at 3300–
3500\,\text{cm}^{-1} and the strong variations of surface cyano groups
at 2177\,\text{cm}^{-1} reveal the higher extension of polycondensation
in the presence of molten salts in comparison with the
reference without salts.[4,19,22] Raman spectra were conducted,
but both melon and NaK-PHI-A show quite similar triazine/
heptazine structures (Supporting Information, Figure S3).

XPS analysis provides more structural details of the
materials. Survey spectra (Supporting Information, Figure
S4) of the materials show that melon comprises C, N,
and O, while PHI-based carbon nitrides are composed with C,
N, O, Na, and K. High-resolution N1 s analysis of the sample
shows a clearly positive shift of the N–C\_ species, indicating
the stronger electron binding of a more stable and extended
conjugation of the PHI-based carbon nitrides, the first
indication of a more positive work function of the mater-
ial.[30,31] The successful generation of carbon nitrides with
enhanced nobility can be also proven by the positive shift of
the valence band positions, moving from 1.92\,\text{eV of melon to} 
2.14\,\text{eV of NaK-PHI-A, as shown in UPS analysis (Figure 1 b).}
This number, in comparison with the previous reported values
of PHI-K, is less positive, but it keeps the strength for a
successful photocatalytic water reduction reaction, as
certified by the slightly downshift in LUMO (from
\text{–1.4 V of melon to} \text{–1.22 V of NaK-PHI-A, Mott–Schottky plots of} 
the materials are presented in the Supporting Information,
Figure S5).
The secret of this new structure nucleated from nucleo-
bases is revealed in AC-HRTEM, where we can observe
domains with distinct locally ordered structure, different from
the weakly ordered structure of melon (Supporting Information,
Figure S6). The lamellar repeat period is unusually high
(ca. 1.65\,\text{nm}, d\_1 + d\_3), which is of the order of two heptazine
units, that is, we see a periodic superstructure, with the
electron contrast presumably created by the charge distribu-
tion within a D-A structure. In some selected parts of
the picture, we even can see that the 1.65\,\text{nm repeat is breaking
up into two layer distances of ca. 0.67 nm and ca. 0.98 nm,}
(Figure 2a). The formation of new stacking distance of
0.67\,\text{nm (smaller than the traditional packing distance of
about 0.92 nm in unmodified PHI) can indicates a tilted
stacking between neighboring heptazine units (tilt angle then
around 40^\circ), which is very similar to J-aggregates of dye
molecules. As J-aggregated usually allows a red shift of the
absorption band and higher thermodynamic stability, while
H-packing then correspond to the usual PHI-packing, we
would have a simple observation of the observed electronic
and contrast features. Electron-hole separation would then
simply occur between the different stacks, thus greatly
simplifying local charge separation and charge transport in

![Figure 1](https://example.com/figure1.png)

**Figure 1.** a) High-resolution N 1s XPS analysis and b) UPS analysis of melon, NaK-PHI, and NaK-PHI-A.

![Figure 2](https://example.com/figure2.png)

**Figure 2.** a) Fourier filtered HR-TEM image of the selected area of NaK-PHI-A; b),c) C K-edge and N-K edge EELS of b) melon and c) NaK-PHI-A.
each well aligned molecular columns (see local D-A junction as depicted in the Supporting Information, Scheme S2). We also find new optical absorption transitions beyond the intrinsic $\pi-\pi^*$ electron transitions of the standard heptazine stacks. The role of 0.3% nucleobases in this structure is still unclear to us and deserved further analysis, but we have to assume that primary condensation structures are able to nucleate such bipolar stacks, which then simply continue to grow, that is, it is the classical role of a nucleation agent.$^{[32,33]}$

Interestingly, relevant electronic details of the materials could be revealed by the C-K and N-K edges EELS of the samples (Figure 2b,c). It is clearly to observe that both samples exhibit a typical carbon nitride structure comprising of both $\pi^*$ and $\sigma^*$ transitions, while however the $\sigma^*$ transition of carbon is much sharper in the new material than that of melon, which potentially can be related to the extra C-T band also seen in optical absorption. On top of that, the distance between the carbon and the nitrogen EELS peaks is increased from 112.2 to 113.8 eV, and this positive change in the binding energy (ca. 1.6 eV) is mostly due to stronger binding of electrons also at the nitrogen position, that is, the overall electronic structure with its positive change in the binding energy indicates a lower valance band of a more stable polymorph.$^{[34]}$ This is in very good accordance with the XPS and UPS data discussed above.

Another difference of the new and the reference carbon nitrides is the surface morphology. SEM images of PHI-based carbon nitrides (Supporting Information, Figure S7) confirm the re-organization of the crystals in the presence of salt melts. Especially with the guanine containing samples we see in the surface texture well faceted primary crystals of 20–50 nm in size, thus supporting our nucleation hypothesis. These primary particles are however inter-grown to a dense bulk, with planar gains boundaries as depicted in Figure 2a. BET analysis (Supporting Information, Figure S9) of the samples illustrate that the specific surface area of NaK-PHI-A is only 11 m$^2$ g$^{-1}$, which is slightly larger than that of weakly ordered melon (for example, CND, 5 m$^2$ g$^{-1}$) but much lower than that of a porous mpg-CN (125 m$^2$ g$^{-1}$) because of the intergrowth of the primary crystals. Besides, SEM elemental mapping (Supporting Information, Figure S8) of the NaK-PHI-A sample confirm the even distribution of the C, N, Na, and K elements.

The optimized D-A heterostructured carbon nitrides also show significantly improved optical and electronic properties, when compared to the melon-based counterparts. UV/Vis spectra of both melon and PHI-based carbon nitrides in Figure 3a and the Supporting Information, Figure S10, but also sheer color reflects the obvious difference in optical absorption. The new system has a stronger transition moment as a semiconductor, but also an extra CT band partly covering the visible spectrum. The optical band gaps of the reference and the new material are calculated to be 2.65 and 2.51 eV, respectively. This absorption, when attributed to a $\sigma^*$ transition, is forbidden for perfectly symmetric and planar units$^{[35–37]}$ but might be activated by the creation of the new packing mode as described in HR-TEM. Another option is the direct optical excitation of a D-A transition. Figure 3b depicts the room-temperature photoluminescence (PL) spectra of the materials. It is easy to observe that NaK-PHI-A possesses much lower PL emission intensity than melon, indicating suppressed recombination of the radioactive charge carrier.$^{[7,8,33]}$ Time-resolved PL (Supporting Information, Figure S11) reveal that the mean radioactive lifetimes (A1) of the recombining charge carriers of melon and NaK-PHI-A were calculated to be 12.18 and 3.88 ns, respectively. It is most likely the local heterostructure in the crystalline polymer accelerates the charge transfer over the interface.$^{[36]}$

The better charge carrier transfer can also been revealed by transient photocurrent generation and the electrochemical impedance spectroscopy (Figure 3c,d). A steady and clearly enhanced photocurrent was generated for the NaK-PHI-A. Meanwhile, the smaller EIS radius of the NaK-PHI-A electrode than that of melon also suggests a faster charge carrier transfer, which is beneficial to improve the photocatalytic performance.$^{[9]}$

The photocatalytic performance of the as-prepared melon and PHI-based samples was then firstly evaluated by H$_2$ production from water in the presence of Pt as cocatalysts and triethanolamine (TEOA) as a sacrificial agent for oxidation. As shown in the Supporting Information, Figure S11, all of the PHI-based carbon nitrides exhibit much better performances (178–342 µmol h$^{-1}$) than the melon-based counterparts, even much better than the mpg-CN (94 µmol h$^{-1}$) with large surface area, indicating overall better photochemistry for the complete reaction chain. As specific BET surface area of NaK-PHI-based carbon nitrides is only about 11 m$^2$ g$^{-1}$, much lower than that of mpg-CN (125 m$^2$ g$^{-1}$), this improvement is even more remarkable for a heterogeneous photocatalyst.

Besides, the PHI-based carbon nitrides also show excellent performance for photocatalytic H$_2$ production by reforming of another, more critical sacrificial agent, that is, ethanol. From Figure 4a we can see that all melon-based carbon nitrides are inactive for photocatalytic oxidation of ethanol, as evidenced by slow H$_2$ production (lower than 5 µmol h$^{-1}$).
In conclusion, a series of PHI-based carbon nitrides were obtained by condensation of 5-amino-tetrazole in the presence of minor amounts of nucleobases. The results reveal that these materials bear an inter-columnar superstructure with different stacking motifs that can act as a D-A heterojunction. This local effect leads to enhanced optical absorption, more stable conjugated systems and improved electronic properties, which speed up the charge transfer and promote the photocatalytic activity, such as exemplifies with H₂ and CH₄ production assays. This study provides a promising way to structure photocatalytic redox active polymers by nucleation effects in a mild salt melt to modulate the local structure and charge transfer processes.

Acknowledgements
G.Z. thanks the Alexander von Humboldt Foundation for a postdoctoral fellowship. This work is financially supported by the Max Planck Institute of Colloids and Interfaces and the National Natural Science Foundation of China (2176132002 and 21425309), the National Key R&D Program of China (2018YFA0209301), and the 111 Project (D16008).

Conflict of interest
The authors declare no conflict of interest.

Keywords: carbon nitride · electron deficient monomers · packing geometry · photocatalysis · water splitting

How to cite: Angew. Chem. Int. Ed. 2019, 58, 14950–14954
Angew. Chem. 2019, 131, 15092–15096

[1] N. S. Lewis, D. G. Nocera, Proc. Natl. Acad. Sci. USA 2006, 103, 15729–15735.
[2] A. Listorti, J. Durrant, J. Barber, Nat. Mater. 2009, 8, 929–930.
[3] K. Maeda, M. Higashi, D. Lu, R. Abe, K. Domen, J. Am. Chem. Soc. 2010, 132, 8585–8586.
[4] G. Zhang, L. Lin, G. Li, Y. Zhang, A. Savateev, X. Wang, M. Antonietti, Angew. Chem. Int. Ed. 2018, 57, 9372–9376; Angew. Chem. 2018, 130, 9516–9520.
[5] W. J. Ong, L. L. Tan, Y. H. Ng, S. T. Yong, S. P. Chai, Chem. Rev. 2016, 116, 7159–7329.
[6] D. J. Martin, P. J. T. Reardon, S. I. A. Moniz, J. Tang, J. Am. Chem. Soc. 2014, 136, 12568–12571.
[7] W. J. Ong, L. L. Tan, S. P. Chai, S. T. Yong, Chem. Commun. 2015, 51, 858–861.
[8] H. Wang, X. Sun, D. Li, X. Zhang, S. Chen, W. Shao, Y. Tian, Y. Xie, J. Am. Chem. Soc. 2017, 139, 2468–2473.
[9] K. Schwinghammer, M. B. Mesch, V. Duppel, C. Ziegler, J. Seener, B. V. Lortsch, J. Am. Chem. Soc. 2014, 136, 1730–1733.
[10] G. Zhang, Z. Ran, L. Lin, S. Lin, X. Wang, Chem. Sci. 2016, 7, 3062–3066.
[11] G. Zhang, S. Zang, X. Wang, ACS Catal. 2015, 5, 941–947.
[12] M. Zhu, S. Kim, L. Mao, M. Fujitsuka, J. Zhang, X. Wang, T. Majima, J. Am. Chem. Soc. 2017, 139, 13234–13242.
[13] M. Shalom, S. Inal, C. Fettkenhauer, D. Neher, M. Antonietti, J. Am. Chem. Soc. 2013, 135, 7118–7121.

In conclusion, a series of PHI-based carbon nitrides were obtained by condensation of 5-amino-tetrazole in the presence of minor amounts of nucleobases. The results reveal that these materials bear an inter-columnar superstructure with different stacking motifs that can act as a D-A heterojunction. This local effect leads to enhanced optical absorption, more stable conjugated systems and improved electronic properties, which speed up the charge transfer and promote the photocatalytic activity, such as exemplifies with H₂ and CH₄ production assays. This study provides a promising way to structure photocatalytic redox active polymers by nucleation effects in a mild salt melt to modulate the local structure and charge transfer processes.

Acknowledgements
G.Z. thanks the Alexander von Humboldt Foundation for a postdoctoral fellowship. This work is financially supported by the Max Planck Institute of Colloids and Interfaces and the National Natural Science Foundation of China (2176132002 and 21425309), the National Key R&D Program of China (2018YFA0209301), and the 111 Project (D16008).

Conflict of interest
The authors declare no conflict of interest.

Keywords: carbon nitride · electron deficient monomers · packing geometry · photocatalysis · water splitting

How to cite: Angew. Chem. Int. Ed. 2019, 58, 14950–14954
Angew. Chem. 2019, 131, 15092–15096

[1] N. S. Lewis, D. G. Nocera, Proc. Natl. Acad. Sci. USA 2006, 103, 15729–15735.
[2] A. Listorti, J. Durrant, J. Barber, Nat. Mater. 2009, 8, 929–930.
[3] K. Maeda, M. Higashi, D. Lu, R. Abe, K. Domen, J. Am. Chem. Soc. 2010, 132, 8585–8586.
[4] G. Zhang, L. Lin, G. Li, Y. Zhang, A. Savateev, X. Wang, M. Antonietti, Angew. Chem. Int. Ed. 2018, 57, 9372–9376; Angew. Chem. 2018, 130, 9516–9520.
[5] W. J. Ong, L. L. Tan, Y. H. Ng, S. T. Yong, S. P. Chai, Chem. Rev. 2016, 116, 7159–7329.
[6] D. J. Martin, P. J. T. Reardon, S. I. A. Moniz, J. Tang, J. Am. Chem. Soc. 2014, 136, 12568–12571.
[7] W. J. Ong, L. L. Tan, S. P. Chai, S. T. Yong, Chem. Commun. 2015, 51, 858–861.
[8] H. Wang, X. Sun, D. Li, X. Zhang, S. Chen, W. Shao, Y. Tian, Y. Xie, J. Am. Chem. Soc. 2017, 139, 2468–2473.
[9] K. Schwinghammer, M. B. Mesch, V. Duppel, C. Ziegler, J. Seener, B. V. Lortsch, J. Am. Chem. Soc. 2014, 136, 1730–1733.
[10] G. Zhang, Z. Ran, L. Lin, S. Lin, X. Wang, Chem. Sci. 2016, 7, 3062–3066.
[11] G. Zhang, S. Zang, X. Wang, ACS Catal. 2015, 5, 941–947.
[12] M. Zhu, S. Kim, L. Mao, M. Fujitsuka, J. Zhang, X. Wang, T. Majima, J. Am. Chem. Soc. 2017, 139, 13234–13242.
[13] M. Shalom, S. Inal, C. Fettkenhauer, D. Neher, M. Antonietti, J. Am. Chem. Soc. 2013, 135, 7118–7121.
