Heterogeneous photoredox flow chemistry for the scalable organosynthesis of fine chemicals

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Large-scale photochemical synthesis of high value chemicals under mild conditions is an ideal method of green chemical production. However, a scalable photocatalytic process has been barely reported due to the costly preparation, low stability of photosensitizers and critical reaction conditions required for classical photocatalysts. Here, we report the merging of flow chemistry with heterogeneous photoredox catalysis for the facile production of high value compounds in a continuous flow reactor with visible light at room temperature in air. In the flow reactor system, polymeric carbon nitrides, which are cheap, sustainable and stable heterogeneous photocatalysts, are immobilized onto glass beads and fibers, demonstrating a highly flexible construction possibility for devices of the photocatalytic materials. As an example of the production of high value chemicals, important chemical structures such as cyclobutanes, which are basic building blocks for many pharmaceutical compounds, like magnosalin, are synthesized in flow with high catalytic efficiency and stability.
Photocatalytic synthesis of high valuable chemicals under mild reaction conditions is a green and sustainable manner for organic transformations, which has been actively pursued by chemists. Many valuable natural compounds and pharmaceutical products have been synthesized in laboratory using photochemistry protocols. However, when scaling up these laboratory-developed photocatalytic reactions in dimension-enlarged reactors for scalable productions, several associated issues came up. One is the decrease in the irradiation efficiency. In principle, the light cannot penetrate deeply into the reaction mixtures with low surface-area-to-volume ratios in traditional batch reactors, leading to the decrease of productivity. In addition, a poor selectivity toward target products is often obtained when prolonging the reaction time to improve the conversion. The development and application of flow reactors is therefore recommended to conduct photocatalytic reactions, efficiently.

Continuous-flow chemistry with inherent advantages of efficient mass- and heat transfer and easy scale-up production has been intensively used for decades in the chemical industry. The special module (e.g., narrow channel) of continuous-flow microreactor enables the uniform irradiation of the entire reaction mixtures, thus ensuring the maximum utilization of irradiation energy to drive photochemical reactions and avoiding the over-irradiation to induce the formation of byproducts. Some homogeneous continuous-flow photochemical systems have been developed for organic synthesis. Even though significant progress has been achieved in these homogenous flow systems, the issues inherited from homogenous photocatalysts still remain, in particular, the poor catalyst stability and the complicated process for catalyst separation and product purification. On this point, the use of heterogeneous photocatalyst in flow reactors ought to be a promising solution because of their facile separation and cost-effective feature. Nevertheless, the development of heterogeneous flow photocatalysis has been greatly hampered by the solid and insoluble nature of heterogeneous photocatalysts, which could induce the blockage of flow systems and the shielding of light absorption. Therefore, a new design of a photoredox flow system is highly demanded, which involves suitable solid photocatalysts, reactors, and their immobilizations, as well as device fabrications in a harmonic way.

Metal-free polymeric carbon nitride (PCN) photocatalyst has attracted extensive attentions owing to its outstanding properties such as tunable redox potentials, high photo- and thermal-stability, facile preparation, and easy processibility in various forms. In fact, PCN has been frequently employed as a sustainable heterogeneous photocatalyst for various photoredox reactions including water splitting, CO₂ reduction, and organic transformations. The ease engineering of PCN organic photocatalyst in band structure and surface functionality makes it particularly promising for various organic transformations. For instance, the oxidations of primary benzylic amines, alcohols, and sulfides were reported under O₂ atmosphere by PCN with a high conversion and selectivity. The sp² C-H bond on phenyl ring could be activated by a PCN/Fe hybrid material. C-C bond formation such as Suzuki coupling reaction has been achieved with palladium-loaded PCN. Photocatalytic Diels–Alder reaction between electron-rich olefins and dienes was also achieved by PCN, and very recently PCN-based organic photocatalysis has been further expanded to bifunctionalize arenes and heteroarenes. Nonetheless, most of the above PCN photoredox catalysis systems mainly focus on milligram scale organic synthesis, and the scalable trial such as reaction in a large-scale batch using PCN as photocatalysts has been rarely reported. For continuous-flow reactor, carbon nitride solids were adhered on glass beads using abundant silica gel as the binder or transformed to water gel using large amount of cross-linker. Therefore, it is highly desired to develop an alternative strategy for carbon nitride-based continuous-flow reaction system in the field of photocatalytic organic synthesis.

Herein, we merge flow chemistry with heterogeneous PCN photocatalysis for the production of cyclobutanes through 2 + 2 cycloadditions with visible light. Cyclobutanes are found in many valuable natural compounds and dienes were also achieved by PCN, and very recently PCN to UCN, also illustrating the decrease of the thickness of these CNs. The physical and optical properties of the three PCNs including the Brunauer–Emmett–Teller (BET) surface areas, pore sizes, band gap, and conduction band (CB)/valance band (VB) position of these three CNs were listed in Table 1. The BET surface of MCN and TCN were both ~10 m² g⁻¹, whereas the BET surface of UCN was larger with ~52 m² g⁻¹. The average

**Results**

**Structural characterization of PCN photocatalysts.** First, three different PCN materials were obtained by simply heating cheap and commercially available precursors such as urea, thiourea, and melamine, which were labeled as UCN, TCN, and MCN, respectively. Details on preparation procedures and characterizations are described in the Supplementary Information. The transmission electron spectroscopy (TEM) images showed different morphologies for the three PCN materials (Fig. 1a-c). UCN possessed a layered structure with a thickness of 10 nm–20 nm, whereas TCN and MCN showed a rather denser morphology with a thickness between ~300 nm and ~600 nm measured by atomic force microscope (AFM) displayed in Supplementary Fig. 2. The functionalized glass beads or fibers can be easily constructed in a continuous-flow photoreactor with high light penetration, and cyclobutanes can be synthesized in gram scale in the continuous-flow photoreactor with a high yield of 81%. The success of this case is expected to open a new avenue for the application of heterogeneous photocatalysis in the green synthesis of added-value fine chemicals in flow.
absorption pore width was obtained by the automatic system calculation from Barret–Joyner–Halenda (BJH) equation, which was showed in Supplementary Fig. 3. From the UV–vis diffuse reflection spectroscopy, optical band gaps in a range from 2.65 to 2.75 eV could be derived for three PCNs (Supplementary Fig. 4). The conduction band (CB) positions could be determined by the electrochemical Mott–Schottky plots (Supplementary Fig. 4c–e), and VB positions were calculated from above two values. All PCNs illuminated light activity properties as demonstrated in photocurrent measurement with clear response for the light-on and light-off events (Supplementary Fig. 5). The highest intensity indicated the most efficient light-induced electronic conductivity, as well as charge separation and charge transfer in UCN among PCNs46, which could be further illustrated by electrochemical impedance spectrum (EIS) in Supplementary Fig. 6.

**Photocatalytic performance on milligram scale.** To test the photocatalytic efficiency of the PCN materials, visible light-induced [2 + 2] cycloaddition between trans-anethole and styrene was chosen as a model reaction. The monitoring experiments indicated that three PCN photocatalysts exhibited the same reaction character of second kinetic order for catalyzing [2 + 2] cycloaddition reaction under the irradiation of a white LED lamp (0.1 W/cm²) (Supplementary Fig. 7). However, reaction conversion differed after 8 h with 99%, 83%, and 35% for UCN, TCN, and MCN, respectively (Supplementary Fig. 8), which was in accordance with the results of photocurrent and EIS test. The highest photocatalytic efficiency for UCN probably resulted from reduced exciton-binding energy in its two dimensional structure, which likely promoted the light-induced charge transfer in CN sheets and thereby accelerated the electron transfer between the substrate and photocatalyst.

A series of control experiments with UCN photocatalyst were performed as listed in Supplementary Table 1 to optimize the reaction condition and reveal the possible interaction between the substrate and photocatalyst. Trace conversion was determined in darkness or without UCN, indicating the essential role and light-driven nature of the UCN photocatalyst (Supplementary Table 1, entry 4–5). Nitromethane was found to be the best reaction medium among tested solvents, while no product was obtained in DMSO and toluene (Supplementary Table 1, entry 6–10). In addition, molecular oxygen could accelerate the cycloaddition reaction, as an obvious decrease was found in nitrogen atmosphere (Supplementary Table 1, entry 11). Furthermore, specific electron and hole scavengers were added into the reaction to analyze the specific roles of photo-generated electron/hole pair. With the addition of KI, trace conversion was detected (Supplementary Table 1, entry 12). Although K₂S₂O₈ was added, both conversion and selectivity decreased markedly (Supplementary Table 1, entry 13).

Based on the observations and previous reports47–51, we proposed the reaction mechanism with the oxidation of trans-anethole by the photo-generated hole as the initiation step (Supplementary Fig. 9). As the VB of UCN lay at 1.36 V vs. SCE, it was sufficient to oxidize trans-anethole (E°ox = 1.17 V vs. SCE)52. Then, the generated cationic radical underwent the [2 + 2] cycloaddition with another alkene partner to form a cyclic intermediate, which afterwards got

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**Table 1 The physical and optical properties of three CNs.**

| Photocatalyst | BET surface (m² g⁻¹) | Pore diameter (nm) | Band gap (eV) | CB/VB position (V vs NHE) | Conversion in 8 h (%) |
|--------------|-----------------------|-------------------|---------------|---------------------------|----------------------|
| MCN          | 10                    | 32.9              | 2.65          | −1.29/1.36                | ~35%                 |
| TCN          | 10                    | 33.1              | 2.67          | −1.30/1.37                | ~83%                 |
| UCN          | 53                    | 21.9              | 2.74          | −1.36/1.38                | >99%                 |

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**Fig. 1 The morphological and structural characterization of PCNs.** a–c TEM images of MCN, TCN, and UCN, respectively. d XRD pattern. e FT-IR spectrum. f Solid-state ¹³C MS-MAS NMR.
an electron from the photocatalyst UCN, the superoxide radical or the reactive neutral trans-anethole to form the final cyclobutane product. As shown in Supplementary Fig. 10, oxygen could quench the fluorescence of UCN suspension with high efficiency for several cycles, indicating that electron transferred from photocatalyst to oxygen. The electron spin resonance (EPR)-trapping experiment using 5,5-dimethyl-1-pyrroline N-oxide (DMPO) as active oxygen trapping agent revealed typical EPR patterns for superoxide oxygen. The electron spin resonance (EPR)-trapping experiment showed the characteristic patterns of the obtained PBN-radical, which indicated clearly the formation of the cationic radical of anethole (Supplementary Fig. 11a). In addition, with N-tert-butyl-α-phenylnitrone (PBN) as radical trapping agent in EPR trapping experiment showed the characteristic patterns of the obtained PBN-radical, which indicated clearly the formation of the cationic radical of anethole (Supplementary Fig. 11b). The region- and stereo-selectivity was attributed to the sterically favored bond formation at β-position of the double bond than that at α position.

To further understand the adsorption patterns between anethole derivatives and UCN, the possible interaction models were calculated. As shown in Fig. 2, one π–π interaction is formed in all three systems. The π–π interaction between the electron-rich PCN and aromatic molecules is beneficial for activation of the molecules by promoting further charge transfer. The adsorption energies of anethole derivatives on UCN are \( \approx -0.8 \) eV, which indicates that the main contribution of the adsorption is from the π–π interactions, while the –OCH\(_3\) group has an ignorable effect.

Photocatalytic performance on gram scale. As an important example of potential industrial application, magnosalin was prepared in a gram-scale batch condition with UCN photocatalyst. Starting with 4 g α-asarone (the moles of substrate increased by factor of 80, 0.1 M), CH\(_3\)NO\(_2\) (200 mL), and 200 mg UCN (Supplementary Fig. 12), the conversion of α-asarone reached 79%, but the selectivity to the product, magnosalin, was 63% (with an isolated yield of 48%) (Table 2, entry 1). The decreased yield was mainly attributed to more oxide side-products formed because of the reaction heat from the amplification system. In order to demonstrate the scalability of our reaction condition and the possibility of carbon nitride as a highly efficient photocatalyst for large-scale preparation of pharmaceutical product, a new type of heterogeneous photoredox flow system was designed and constructed.

With the optimized reaction condition on hand, a photochemical continuous-flow setup was performed in order to simulate the general process of industrial production (Fig. 3a, b). Here, the photocatalyst UCN was immobilized into a photoreactor \((d = 0.7 \text{ cm, } l = 7 \text{ cm})\) to construct a heterogeneous flow reaction system. Originally, commercially available glass fiber was chosen as the support to immobilize the photocatalyst UCN. Precisely, a UCN-coated glass fiber could be obtained via direct thermal condensation of urea on the surface of fresh fiber (detailed information in Methods). As shown in Supplementary Fig. 13a, b, the fresh glass fiber demonstrated a smooth and glassy surface, the surface became rough and flake-decorated after coating with UCN photocatalyst. Strong blue emission could be clearly observed in fluorescence microscope images (Supplementary Fig. 13c, d), indicating that glass fiber was successfully covered with UCN photocatalyst. The average thickness of the UCN layer was \( \approx 20 \text{ nm} \) (Supplementary Fig. 13b) and the amount was estimated to be 1.2 wt.% after calcination. Subsequently, the UCN-coating glass fiber was filled into photoreactor. Herein, the cycloaddition

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**Table 2 Screening and control experiment of gram-scale [2 + 2] dimerization of α-asarone catalyzed by UCN under white light.**

| Entry | Light | Reactor | Reaction condition variations | Yield |
|-------|-------|---------|------------------------------|-------|
| 1     | +     | Batch   | UCN powders<sup>a</sup> | 48%   |
| 2     | +     | Flow    | UCN@glass fibers<sup>d</sup> | 70%   |
| 3     | +     | Flow    | UCN@glass beads            | 81%   |
| 4     | +     | Flow    | In dark                      | Trace |
| 5     | +     | Flow    | In acetonitrile              | 2%<sup>e</sup> |
| 6     | +     | Batch   | UCN@glass beads             | 53%   |
| 7<sup>f</sup> | +  | Flow    | UCN@glass beads             | 87%   |

<sup>a</sup> Standard reaction conditions of flow system: (α-asarone) – 0.167 M, V\(_{CH_3NO_2}\) – 60 mL, white LED lamp (0.1 W/cm\(^2\)), room temperature, air, irradiation time 48 h.

<sup>b</sup> Isolated yield.

<sup>c</sup> The amount of UCN is 200 mg.

<sup>d</sup> The amount of UCN coated on glass fibers is 80 mg.

<sup>e</sup> Conversion from GC-MS.

<sup>f</sup> Using a photoreactor assembled from six paralleled glass tubes with irradiation time 8 h.

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Fig. 2 The DFT calculation of absorption model. Optimized structures and adsorption energies of a anethole. b Methyl isoeugenol. c α-asarone on the surface of UCN.
of α-asarone was employed in flow because the generated product magnosalin is an important anti-inflammatory agent. As expected, a high reaction conversion of 79% for α-asarone and yield of 70% for magnosalin was achieved with almost quantitative selectivity (Table 2, Entry 2, and Supplementary Fig. 14a), which indicated a high catalytic efficiency of UCN photocatalyst and feasibility of creating a heterogeneous flow photochemistry. Unfortunately, a decreased conversion was found during the repeating experiments (Supplementary Fig. 14a). The possible reason lay in the separation and loss of UCN from glass fiber due to the high pressure and speed during the flow process, which was also demonstrated by the decrease of absorption for the UCN-coating glass fibers after 50 h and 200 h, respectively (Supplementary Fig. 13e). In order to enhance the binding between support and photocatalyst UCN, a revised coating method needed to be designed.

Afterwards, commercially available glass beads with diameter of 1 mm were employed as another support for UCN immobilization through a modified approach. With details described in methods (see in Fig. 3d), the glass beads were successively etched with piranha solution, decorated with silylation reagent (3-aminopropyltriethoxysilane, APTES) and coated with UCN via thermal condensation of urea, thus producing the final UCN-coated glass beads (showed in Fig. 3c). The introduction of APTES could function as a bridge for connecting glass bead and UCN (Supplementary Fig. 15). With this modified method, the obtained UCN@glass beads showed obvious yellow and rough surface (Fig. 3c). In addition, the glass beads displayed a strong blue fluorescence, indicating that the surface was completely covered with UCN photocatalyst (Fig. 4a, b). The thickness of the UCN flake was 5–10 nm from SEM (Fig. 4c) and the amount was calculated to be ~1 wt. % from calcination test. The element mapping on the deliberately damaged surface demonstrated a clear separation between support and photocatalyst, offering a direct proof for the successfully coating of UCN on the surface of glass beads (Fig. 4d–g). Chemical structure and morphology of UCN coated on the surface of glass beads were further demonstrated by XRD (Supplementary Fig. 16a), FT-IR (Supplementary Fig. 16b), XPS (Supplementary Fig. 17), and SEM (Supplementary Fig. 18). In addition, the MCN and TCN on the surface of glass beads were characteristic by PL (Supplementary Fig. 19), FT-IR (Supplementary Fig. 20) and XPS (Supplementary Fig. 21) to show the universality of this method. At last, UCN-coated glass beads exhibited a highly flexible processing possibility in various shapes, as demonstrated in Supplementary Fig. 22.

The obtained UCN-coated glass beads were then employed to test the photocatalytic efficiency and stability in continuous-flow condition. In all, 8 g UCN-coated glass beads were filled in the flow photoreactor with α-asarone concentration being 0.167 M in 60 mL CH₂NO₂ and a flow speed of 0.5 mL min⁻¹. As shown in Table 2 (Entry 3), a comparable catalytic efficiency was achieved. The reaction conversion reached 61% after 24 h and further increased to 89% after 48 h with a final isolated yield of 81%, corresponding to 1.6 g magnosalin. No conversion was determined in darkness and trace conversion was detected in other reaction medium such as acetonitrile (Table 2, Entry 4–5). Moreover, an obviously improved stability was found with UCN-coated glass beads. In repeating experiments, the reaction conversion stabilized in five cycles without obvious decrease (Supplementary Fig. 14b). Therefore, the turn over number for the reaction was calculated with a value of 106 based on the moles of UCN on the surface of glass beads for total five cycles. This improvement is mainly attributed to the strong binding force between the glass beads and UCN photocatalyst in the modified approach, which offers a miniature for continuous industrial pharmaceutical compound, magnosalin, production. The flow reactor delivered a much higher reaction yield than that in batch
Table 2, Entry 6), indicative of a great advantage in the flow reaction system. AQY measurement of both batch and flow system under a 420 nm monochromatic light (Supplementary Fig. 23) was compared with further demonstrate the advantage of the continuous-flow system for scale-up (description in detail shown in Supplementary Discussion and Supplementary Figs. 24 and 25). The photocatalytic efficiency can be further improved when using a photoreactor assembled from six paralleled glass tubes (Supplementary Fig. 26) to enlarge the irradiation surface area (Table 2, Entry 7 vs. Entry 2).

At last, to demonstrate the general accessibility of the [2 + 2] cycloaddition reaction with UCN photocatalyst, the unsymmetrical [2 + 2] cycloaddition reaction was tested with UCN-coated glass beads in continuous-flow photoreactor under the visible light irradiation. As showed in Fig. 5, various styrene derivatives with both electron-donating groups such as methyl (b–d) and electron-withdrawing groups such as ester (i) and halides (e–j) were accessible for the reaction with remarkable reaction yield. Although some side-reactions such as dimerization and oxidation of anethole may present, the crossed-cycloaddition with styrene derivatives always took place as the major pathway, with a considerable amount of unsymmetrical cyclobutane products being achieved. In addition, the position of the functional group had slight influence on the reaction yield. Both methyl-substituted (b–d) and bromine-substituted (e–g) styrene on ortho-, meta-, or para- positions demonstrated generally comparable reaction yield, respectively. Furthermore, anethole derivatives, such as methyl isoeugenol (k), methoxycinnamyl alcohol (l), and asarone (m) were also tolerated with moderate reaction yield. The slight lower conversion for methyl isoeugenol and asarone starting compound may be resulted from the over-oxidation and decomposition of electron-rich cycloadduct intermediates. However, trace products were determined even after a longer reaction time (48 h) when p-methoxy cinnamaldehyde (n) and p-methoxy cinnamic acid (o) were employed as reactive substrates, probably due the high oxidation potentials of the starting compounds. Although many reports have been conducted about cyclobutanes preparation, direct synthesis of pharmaceutical product, such as magnosalin, as far as we know, is still rare. The UCN-catalyzed [2 + 2] cycloaddition in continuous-flow reaction system offers a green approach for drugs production in a mild condition.

Discussion

The most-challenging feature of using photocatalysts in heterogeneous reaction lies in the terms of the light absorption, design of photocatalytic devices and a balance between activity and stability during the amplification reaction. We have demonstrated the use of carbon nitride as cheap, processable, and stable photocatalyst in scalable and continuous-flow system for cyclobutanes production, especially for magnosalin preparation in gram scale. Both commercial available glass fiber and glass beads have been employed as translucent support for the carbon nitride photocatalyst. The flow photoreactor with UCN-coated glass beads demonstrates the highest reaction yields. Unsymmetrical [2 + 2] cycloaddition with extended styrene derivatives have been conducted with UCN photocatalyst with considerable conversion. The flow photochemical system is compatible for crossed [2 + 2] cycloaddition reaction with corresponding cyclobutanes being achieved with a remarkable yield. We believe that this study can demonstrate an important step toward the utilization of carbon nitride for photo-catalyzed industrial production of fine chemicals under mild and sustainable reaction conditions.
Synthesis of UCN-coating glass fibers. The glass fibers were first cleaned in piranha solution (H2O2:H2SO4 = 1:3) to remove the surficial contaminants. Then, the glass fibers were immersed into a saturated urea solution (10 g/mL), followed by reflux at 60 °C for 12 h. After water evaporation at 80 °C for overnight, the crude mixture was heated in a muffle furnace with the same condition as that of UCN synthesis. A UCN-coated glass fiber could be obtained after directly thermal condensation polymerization on the surface of glass fiber.

Synthesis of UCN-coating glass beads. As shown in Fig. 3d, commercially available glass beads were first cleaned and etched with piranha solution. Then, silylation reaction was conducted in order to introduce an amine function group. Precisely, a 20 mL anhydrous toluene was added with acid etched glass beads and silylation reaction was conducted in order to introduce an amine function group. The heating program as that of UCN synthesis to produce UCN-coated glass beads. The charged with UCN (2.4 mg), a glass tube was sealed with a glass vial and used as photoreactor in oven at 60 °C for overnight. The amine functionalized glass beads were mixed with urea with a mass ratio of 1:1, and then heated in a muffle furnace using the same heating program as that of UCN synthesis to produce UCN-coated glass beads.

Photosynthesis of cyclobutanes in batch condition. A flame-dried 40 ml vial was charged with UCN (2.4 mg), E-anethole (50.0 mg, 0.24 mmol), and CH3NO2 (2.4 ml). Then, 10 equivalent of styrene derivatives partner was added into the reaction mixture. Afterwards, the vial was placed under the irradiation of a white LED lamp (0.1 W/cm²) in air condition. The conversion of was determined by GC-MS measurement based on the consumption of anethole derivatives. After the reaction was finished, the mixture was poured into a separatory funnel and extracted with a mixture of 60 ml Et2O and water (v/v, 1:1). A crude product was received after collecting organic layer, drying with anhydrous MgSO4, and concentrating with rotary evaporator. Finally, the pure cycloaddition products could be obtained after further purification with flash-column chromatography.

Data availability
All data supporting the findings of this study are available within the article as well as the Supplementary Information file, or available from the corresponding authors on reasonable request.

Received: 13 July 2019; Accepted: 11 February 2020; Published online: 06 March 2020

References
1. Scholes, G., Fleming, G., Olaya-Castro, A. & van Grondelle, R. Lessons from nature about solar light harvesting. Nat. Chem. 3, 763–774 (2011).
2. Demmig-Adams, B., Stewart, J., Burch, T. & Adams, W. Insights from placing photosynthetic light harvesting into context. J. Phys. Chem. Lett. 5, 2880–2889 (2014).
3. Scholes, G., Mirkovic, T., Turner, D., Fassioli, F. & Buchleitner, A. Solar light harvesting by energy transfer: from ecology to coherence. Energy Environ. Sci. 5, 9374–9393 (2012).
4. Nicholls, P., Leoniotti, D. & Bisemben, C. Applications of Visible light photocatalysis to the synthesis of natural products and related compounds. Nat. Prod. Rep. 33, 1248–1254 (2016).
5. Douglas, J., Sevin, J. & Stephenson, J. Visible light photocatalysis: applications and new disconnections in the synthesis of pharmaceutical agents. Org. Process Res. Dev. 20, 1134–1147 (2016).
6. Cambé, D., Bottecchia, C., Straathof, J., Hessle, V. & Noël, T. Applications of continuous-flow photochemistry in organic synthesis, material science, and water treatment. Chem. Rev. 116, 10276–10341 (2016).
7. Porta, R., Benaglia, M. & Puglisi, A. Flow chemistry: recent developments in the synthesis of pharmaceutical products. Org. Process Res. Dev. 20, 2–25 (2016).
8. Tucker, J., Zhang, Y., Jamison, T. & Stephenson, C. Visible-light photoxidation catalysis in flow. Angew. Chem. Int. Ed. 51, 4144–4147 (2012).
9. Wang, X. et al. A metal-free photocatalyst for hydrogen production from water under visible light. Nat. Mater. 8, 76–80 (2009).
10. Wang, X., Lee, C. & Antonietti, M. Polymeric graphic carbon nitride as a heterogeneous organocatalyst: from photobiology to multipurpose catalysis to sustainable chemistry. Angew. Chem. Int. Ed. 51, 68–89 (2012).
11. Ong, W. J., Tan, L., Ng, Y. H., Yong, S. T. & Chai, S. P. Graphitic carbon nitride (g-C3N4)-based photocatalysts for artificial photosynthesis and environmental remediation: are we a step closer to achieving sustainability? Chem. Rev. 116, 7359–7382 (2016).
12. Hou, Y. et al. Layered nanojunctions for hydrogen-evolution catalysis. Angew. Chem. Int. Ed. 52, 3621–3625 (2013).
13. Zhang, J. et al. Photocatalytic oxidation of water by polyaniline with nanostructures made of a supramolecular agent. J. Phys. Chem. Lett. 3, 443–446 (2012).
14. Zhai, G., Lan, Z., Lin, L., Lin, S. & Wang, X. Overall water splitting by Pt/C3N4 photocatalysts without using sacrificial agents. Chem. Sci. 7, 3062–3066 (2016).
15. Martin, D. J., Beardon, P., Moniz, S. & Tang, J. Visible light-driven pure water splitting by a nature-inspired organic semiconductor-based system. J. Am. Chem. Soc. 136, 12568–12571 (2014).
16. Lin, J., Pan, Z. & Wang, X. Photocatalytic reduction of CO2 by g-C3N4 photocatalysts. ACS Sustain. Chem. Eng. 2, 353–358 (2014).
17. Zhao, G. et al. Co-porphyrin/carbon nitride hybrids for improved photocatalytic CO2 reduction under visible light. Appl. Catal. B: Environ. 200, 141–149 (2017).
18. Savateev, A. & Antonietti, M. Heterogeneous organocatalysis for photoredox catalysis. ACS Catal. 8, 9790–9808 (2018).
19. Savateev, A., Ghosh, I., König, B. & Antonietti, M. Photoredox catalytic organic transformations using heterogeneous carbon nitrides. Angew. Chem. Int. Ed. 57, 15936–15947 (2018).
20. Su, F. et al. Aerobic oxidative coupling of amines by carbon nitride photocatalysis with visible light. Angew. Chem. Int. Ed. 50, 657–660 (2011).
21. Su, F. et al. mp-C3N4-catalyzed selective oxidation of alcohols using O2 and visible light. J. Am. Chem. Soc. 132, 16299–16301 (2010).
22. Zhang, P., Wang, Y., Li, H. & Antonietti, M. Metal-free oxidation of sulfides by carbon nitride with visible light illumination at room temperature. Green Chem. 14, 1904–1908 (2012).
23. Chen, X., Zhan, J., Fu, X., Antonietti, M. & Wang, X. Fe-g-C3N4-catalyzed oxidation of benzene to phenol using hydrogen peroxide and visible light. J. Am. Chem. Soc. 11658–11659 (2009).
24. Li, X., Baar, M., Blechert, S. & Antonietti, M. Facilitating room-temperature Suzuki coupling reaction with light: Mott-Schottky photocatalyst for C-C coupling. Sci. Rep. 3, 1743 (2013).
25. Zhao, Y. & Antonietti, M. Visible-light-irradiated graphitic carbon nitride photocatalyst photocatalyzed diels–alder reactions with dioxygen as sustainable mediator for photoinduced electrons. Angew. Chem. Int. Ed. 56, 9336–9340 (2017).
26. Ghosh, I. et al. Organic semiconductor photocatalyst can bifunctionalize amines and heteroaromatics. Science 365, 360–366 (2019).
27. Dai, Y. et al. Light-tuned selective photosynthesis of azo- and azoxy-aromatics using graphic C3N4, Nat. Commun. 9, 60 (2018).
28. Woznica, M., Chaoui, N., Taabache, S. & Blechert, S. THF: an efficient electron donor in continuous flow radical cyclization photocatalyzed by graphitic carbon nitride. Chem. Eur. J. 20, 14624–14628 (2014).
29. Cai, Y. et al. Heterogeneous visible-light photocatalysis catalysis with graphic carbon nitride for α-aminonitriles radical addition, alkylation, and heteroaclylations. ACS Catal. 8, 9471–9476 (2018).
30. Pieber, B., Shalom, M., Antonietti, M., Seeberger, H. P. & Gilmore, K. Continuous heterogeneous photocatalysis in serial micro-batch reactors. Angew. Chem. Int. Ed. 57, 9976–9979 (2018).
31. Dembitsky, V. M. Bioactive cyclobutane-containing alkaloids. J. Nat. Med. 62, 1–33 (2008).
32. Yoon, T. P. Visible light photocatalysis: the development of photocatalytic radical ion cyclizations. ACS Catal. 3, 895–902 (2013).
33. Xu, Y., Conner, M. L. & Brown, M. K. Cyclobutane and cyclobutene synthesis: catalytic enantioselective [2 + 2] cyclizations. Angew. Chem. Int. Ed. 54, 11918–11928 (2015).
34. Boxer, M. B. & Yamamoto, H. Remarkable tris(trimethylyl)silyl group for diastereoselective [2 + 2] cyclizations. Org. Lett. 7, 3127–3129 (2005).
35. Ishihara, K. & Nakano, K. Enantioselective [2 + 2] cycloaddition of unactivated alkenes with α-acetoxyacetals catalyzed by chiral organoammonium salts. J. Am. Chem. Soc. 129, 8930–8931 (2007).

36. Shibata, T., Takami, K. & Kawachi, A. Rh-catalyzed enantioselective [2 + 2] cycloaddition of allylnyl esters and norbornene derivatives. Org. Lett. 8, 1343–1345 (2006).

37. Blum, T., Miller, Z. D., Bates, D., Guzei, I. & Yoon, T. P. Enantioselective photochemistry through Lewis acid-catalyzed triplet energy transfer. Science 354, 1391–1395 (2016).

38. Ishchay, M., Zhan, L. & Yoon, T. P. [2 + 2] photocycloaddition of alkynyl esters and norbornene derivatives. Org. Lett. 8, 963–966 (2006).

39. Daub, M. et al. Enantioselective [2 + 2] cycloadditions of enamine substrates: generalizing Lewis acid catalysis of triplet energy transfer. J. Am. Chem. Soc. 141, 9543–9547 (2019).

40. Hu, N. et al. Catalytic asymmetric dearylation by visible-light-activated [2 + 2] photocycloaddition. Angew. Chem. Int. Ed. 57, 6242–6246 (2018).

41. Zhou, Q., Zou, Y., Lu, L. & Xiao, W. Visible-light-induced organic photochemical reactions through energy-transfer pathways. Angew. Chem. Int. Ed. 58, 1586–1604 (2019).

42. Poplata, S., Tröster, A., Zou, Y. & Bach, T. Recent advances in the synthesis of cyclobutanes by olefin [2 + 2] photocycloaddition reactions. Chem. Rev. 116, 9748–9815 (2016).

43. Okada, Y., Maeta, N., Nakayama, K. & Kamiya, H. TiO₂ photocatalysis in aqueous solution. J. Am. Chem. Soc. 129, 6202–6203 (2007).

44. Shibata, T., Takami, K. & Kawachi, A. Rh-catalyzed enantioselective [2 + 2] cycloaddition of allylnyl esters and norbornene derivatives. Org. Lett. 8, 1343–1345 (2006).

45. Blum, T., Miller, Z. D., Bates, D., Guzei, I. & Yoon, T. P. Enantioselective photochemistry through Lewis acid-catalyzed triplet energy transfer. Science 354, 1391–1395 (2016).

46. Ishihara, K. & Nakano, K. Enantioselective [2 + 2] cycloaddition of unactivated alkenes with α-acetoxyacetals catalyzed by chiral organoammonium salts. J. Am. Chem. Soc. 129, 8930–8931 (2007).

47. Ishchay, M., Zhan, L. & Yoon, T. P. [2 + 2] photocycloaddition of alkynyl esters and norbornene derivatives. Org. Lett. 8, 963–966 (2006).

48. Daub, M. et al. Enantioselective [2 + 2] cycloadditions of enamine substrates: generalizing Lewis acid catalysis of triplet energy transfer. J. Am. Chem. Soc. 141, 9543–9547 (2019).

49. Hu, N. et al. Catalytic asymmetric dearylation by visible-light-activated [2 + 2] photocycloaddition. Angew. Chem. Int. Ed. 57, 6242–6246 (2018).

50. Zhou, Q., Zou, Y., Lu, L. & Xiao, W. Visible-light-induced organic photochemical reactions through energy-transfer pathways. Angew. Chem. Int. Ed. 58, 1586–1604 (2019).

51. Poplata, S., Tröster, A., Zou, Y. & Bach, T. Recent advances in the synthesis of cyclobutanes by olefin [2 + 2] photocycloaddition reactions. Chem. Rev. 116, 9748–9815 (2016).

52. Okada, Y., Maeta, N., Nakayama, K. & Kamiya, H. TiO₂ photocatalysis in aqueous solution. J. Am. Chem. Soc. 129, 6202–6203 (2007).

53. Shibata, T., Takami, K. & Kawachi, A. Rh-catalyzed enantioselective [2 + 2] cycloaddition of allylnyl esters and norbornene derivatives. Org. Lett. 8, 1343–1345 (2006).

54. Blum, T., Miller, Z. D., Bates, D., Guzei, I. & Yoon, T. P. Enantioselective photochemistry through Lewis acid-catalyzed triplet energy transfer. Science 354, 1391–1395 (2016).

55. Ishihara, K. & Nakano, K. Enantioselective [2 + 2] cycloaddition of unactivated alkenes with α-acetoxyacetals catalyzed by chiral organoammonium salts. J. Am. Chem. Soc. 129, 8930–8931 (2007).

56. Shibata, T., Takami, K. & Kawachi, A. Rh-catalyzed enantioselective [2 + 2] cycloaddition of allylnyl esters and norbornene derivatives. Org. Lett. 8, 1343–1345 (2006).

57. Blum, T., Miller, Z. D., Bates, D., Guzei, I. & Yoon, T. P. Enantioselective photochemistry through Lewis acid-catalyzed triplet energy transfer. Science 354, 1391–1395 (2016).

58. Ishihara, K. & Nakano, K. Enantioselective [2 + 2] cycloaddition of unactivated alkenes with α-acetoxyacetals catalyzed by chiral organoammonium salts. J. Am. Chem. Soc. 129, 8930–8931 (2007).