Visible Light Flow Reactor Packed with Porous Carbon Nitride for Aerobic Substrate Oxidations

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Abstract

A triphasic photocatalytic reactor employing a mesoporous carbon nitride photocatalyst and aerobic O$_2$ was assembled to operate under continuous flow conditions. This reactor design allows for facile downstream processing and reusability in multiple flow cycles. The selective aerobic oxidation of alcohols and amines was chosen to demonstrate the applicability and performance advantage of this flow approach compared to conventional batch photochemistry. This precious-metal-free photocatalytic flow system operates under benign reaction conditions (visible light, low pressure, and mild temperature) and will stimulate the exploration of other oxidative reactions in a sustainable, scalable, and affordable manner.

Keywords: photocatalysis, flow chemistry, aerobic oxidation, carbon nitride, green chemistry

Introduction

Continuous flow chemistry has grown tremendously in recent years due to its advantages over conventional batch reactors, particularly for photocatalytic processes.$^{1-5}$ Batch photoreactors tend to be comprised of a simple tube or flask, and hence suffer from a lack of design specificity towards light driven transformations. Alternatively, flow devices can be fabricated to have a high surface area-to-volume (SA-to-V) ratio, guaranteeing extensive transmission of light throughout the reaction mixture even at high chromophore concentrations.$^6$ Furthermore, this characteristic also ensures efficient thermal transfer, reducing the risk of runaway reactions from exothermic pathways.$^{7,8}$ The continuous operation also allows for accurate control of residence and irradiation time via the flow rate.$^9$ Photocatalyst immobilization within the reactor can assist with downstream processing of reaction mixtures as it avoids catalyst separation and thereby facilitates reusability and scalability.$^{10}$

Progress in the field of photoredox catalysis has mostly been confined to homogeneous systems, involving costly Ru and Ir complexes,$^{11}$ and tailor-made organic chromophores.$^{12}$ Graphitic carbon nitride, a polymeric, easily synthesizable, robust, metal-free material composed of tri-s-triazine units (Figure S1),$^{13}$ is one of the most promising heterogeneous photocatalysts, with applications in visible light
mediated organic transformations, and early-stage development in flow photochemistry.

Two main approaches for carbon nitride-based flow photochemistry have been adopted to date: (i) featuring a stationary carbon nitride phase within a packed column approach, and (ii) using a suspension in-flow, that is divided into smaller sub-units by means of an inert gas spacer (termed serial micro-batch reactors, SMBR). The former approach allows for the effective immobilization of the carbon nitride material, and the overall setup can therefore easily be re-used with minimal effort to recover the photocatalyst. However, this configuration was limited to liquid-only flow applications, and not for triphasic (gas–liquid–solid) flow chemistry. The innovative design of the SMBR made use of toroidal vortices in the segmented flow pattern to produce ‘well-stirred’ micro-suspensions containing the carbon nitride photocatalyst. This internal mixing phenomenon was therefore responsible for homogenizing the mixture and resulted in the formation of uniformly irradiated suspensions in the flow sequence. However, the separation strategy for product isolation, and recovery of the catalyst and ionic liquid was demanding (featuring filtration, liquid extraction, and solid extraction stages), and hence offered some additional complexity to the reaction protocol in comparison to the packed column design. The SMBR employed inert gas (N₂) to generate the flow pattern, and hence the study precluded the use of a reactive atmosphere.

The demand from industry for aerobic oxidation catalysts is growing. O₂ can be considered as an ideal oxidant, as it is readily abundant and usually generates water as a benign by-product. This is especially advantageous for the production of active pharmaceutical ingredients, where impurities are strongly regulated. A review of large-scale oxidation processes concluded that O₂ gas has the advantage of being a cost-effective and readily available oxidant, but the safety issues surrounding its use (mixtures of O₂ and organic solvents can potentially generate highly flammable mixtures) make it one of the least frequently used reagents. The recent revival in the development of catalytic systems for aerobic substrate oxidations is testimony to the commitment from industry to pursue the application of aerobic O₂. However, such systems are either confined to a small scale, feature catalysts containing precious metals, make use of a homogeneous photocatalyst that hinders downstream processing and separation, or are unsuitable for a continuous flow platform.
Conducting aerobic oxidations in-flow with a carbon nitride material is therefore an attractive option due to the attributes of this photocatalyst, and the potential to explore this chemistry at a more commercial level. The packed column design offers both a large interfacial area and simulates high local catalyst loading, and is in fact more applicable when heterogeneous catalysis is needed for continuous, multiphase (gas–liquid–solid) chemical transformations, such as catalytic hydrogenation reactions.

Here we present a triphasic, packed column photoreactor (PCP) system for the continuous flow oxidative photochemistry of alcohols and amines during visible light irradiation, in which a mesoporous, graphitic carbon nitride material (mpg-CNₓ) is incorporated as the stationary photocatalyst, and aerobic O₂ is used as the terminal electron acceptor and oxidant (Figure 1, Scheme 1). In addition, we demonstrate that this chemistry can be accomplished using more benign conditions compared to previously reported mpg-CNₓ-based batch photoreactors, which typically require temperatures and O₂ pressures in the range of 80–100 °C and 5–8 bar, respectively. The previously proposed photocatalytic mechanism for this reaction suggests that photoactivated mpg-CNₓ oxidizes the benzyl alcohol/amine to the corresponding aldehyde/imine and reduces O₂ to hydrogen peroxide and water (Figure S1). Aldehydes and imines are commonly used synthetic building blocks for the pharmaceutical industry.

Results and Discussion

A flow setup was constructed to generate a stable gas–liquid segmented (slug) flow pattern (Figure 1). This pattern formed downstream of a 4-port 3-way valve, where the substrate and air fronts meet and mix. The resultant laminar flow feeds into the PCP, which was comprised of a flexible, transparent fluorinated ethylene propylene (FEP) tube (l = 75 cm, i.d. = 2 mm), which was packed with the carbon nitride material (3.5 wt.% in a glass bead mixture; see Supporting Information for more details). The PCP was coiled around a reflective support, and then irradiated in a 360° fashion with a blue LED setup (14.4 W at λ = 470 nm). Typically, 2.5 mL of the substrate solution (concentration in the range of 50 – 100 mM) was first injected into the reactor, followed by 2.5 mL of pure solvent. The residence time of the starting material, τ, within the
photoactive zone was varied by changing the flow rate as the reactor volume was kept fixed at \(~1\) mL (see Supporting Information, Eq. S2).

**Figure 1.** Schematic diagram of the triphasic flow photoreactor incorporating mpg-CN\(_{x}\) within the PCP. Irradiation is carried out via 360° blue LEDs (14.4 W at 470 nm) (MFC: mass flow controller). TEM image highlighting the mesoporous morphology of the mpg-CN\(_{x}\) material and illustration of the gas–liquid flow injected into the PCP is also shown.

We observed that the morphology of the carbon nitride greatly impacts the flow performance of the PCP. In preliminary tests, both mpg-CN\(_{x}\) and conventional bulk carbon nitride (b-CN\(_{x}\)) were individually incorporated within the PCP. Transmission electron microscopy (TEM) images show the nonporous, flat morphology of b-CN\(_{x}\) (Figure S2a), which is in stark contrast to the mesoporous network of mpg-CN\(_{x}\) (Figure 1, Figure S2b). This morphological difference is also corroborated by N\(_2\) sorption measurements (Figure S3a), which demonstrate the large Brunauer–Emmett–Teller (BET) surface area of mpg-CN\(_{x}\) (187 m\(^2\) g\(^{-1}\)) in comparison to b-CN\(_{x}\) (5 m\(^2\) g\(^{-1}\)). The pore size distribution gives an average pore size of 5.5 nm for mpg-CN\(_{x}\) (Figure S3b).

The b-CN\(_{x}\) PCP could not support the movement of a gas–liquid flow, presumably due to the agglomeration and clustering of the flat b-CN\(_{x}\) particles, which resulted in a high back-pressure (~1.5 MPa) during operation. On the other hand, the porous framework of mpg-CN\(_{x}\) allowed for facile gas–liquid permeation, making it suitable for flow applications. The absence of any critical back-pressure issues permitted the use of small diameter reactors, satisfying the high SA-to-V constraint.
imposed for photoinduced reactions. The high SA-to-V ratio is necessary in order to minimize the amount of photocatalyst (and hence reactive sites) within the central regions of the reactor which, as governed by the Beer–Lambert–Bouguer law, would experience a diminished photon flux. The larger surface area of mpg-CN\textsubscript{x} compared to b-CN\textsubscript{x} gives the additional benefit of improved charge separation and interfacial electron transfer\textsuperscript{32}.

![Scheme 1](image)

**Scheme 1.** (a) Alcohol and (b) amine oxidation reactions studied under continuous flow conditions.

The photoreactor assembly allowed for substrate and solvent injection, and the air feed was continuously switched on to ensure the formation of a gas–liquid flow, and to constantly supply aerobic O\textsubscript{2} to the reactive centers. Acetonitrile (MeCN) was the most effective solvent (Figure S4), and the heat generated from the LED setup (~55 °C) increased the overall conversion yield. To facilitate downstream processing, an MeCN suppression pulse program for \textsuperscript{1}H NMR analysis was designed and implemented, to allow for direct sampling and measurement of the crude mixture without the need for reaction treatment (see Supporting Information for more details).

The system was initially studied using the photocatalyzed aerobic oxidation of benzylic alcohols\textsuperscript{30} with 1-naphthalenemethanol selected as the model compound (Scheme 1a, Figure 2). The overall *starting material conversion (Conv)*, into the resultant aldehyde and carboxylic acid, and *product selectivity (Sel)* to the corresponding aldehyde, was controlled by varying the residence time in the PCP. The overall Conv approaches 100% at \( \tau > 90 \text{ min} \), and the formation of the ‘overoxidized’ product (carboxylic acid) only becomes appreciable at \( \tau > 75 \text{ min} \) (Figure 2). The significance of this result is that a lower value of \( \tau \) could be employed to ensure high selectivity for the aldehyde; the starting material can then be separated from the reaction mixture in an in-line manner at the PCP exit and recycled back into the photoreactor setup.
The fit applied to the Conv dataset in Figure 2 is of the form: \(1 - e^{-k_{PCP} \tau}\), where \(k_{PCP}\) represents a form of ‘rate constant’ (units: \(\text{min}^{-1}\)) for the triphasic oxidation reaction within the PCP. \(k_{PCP}\) is therefore expected to vary with both an alteration in the reactor parameters (such as catalyst loading, and diameter of the packed column, amongst others), and a change in substrate. Effectively, it can be used as a design guide to compare the performance of different reactor configurations (in the presence of the same substrate), or to compare the reactivity of different substrates within the same PCP design.

The inset in Figure 2 highlights the recyclability of the photocatalytic system, which maintains a high level of activity for multiple flow cycles. To compile this plot, a 1-naphthalenemethanol solution in MeCN was injected into reactor (2.5 mL, 100 mM), followed by 2.5 mL of pure MeCN. This procedure was repeated for a total of 25 times, and the Conv metric was computed after each individual flow reaction. Relative to the initial run, there was only an approximate 10% drop in activity by the 25th flow experiment, most likely due to the sedimentation (and possible aggregation) of the mpg-CN\(\_x\) particles within the packing mixture of the PCP (Figure S5), which would effectively lead to a reduction in the active surface area of the photocatalyst.

The benzylic oxidation reaction (Scheme 1a) was compared under both batch and flow conditions and the performance was assessed by determining a number of metrics, namely the Conv, Sel, and product–substrate recovery, with the latter henceforth denoted as the mass balance (M.B.). The M.B. metric therefore provides a means to evaluate the moles of species exiting the reactor, relative to the amount injected into the system (see Supporting Information, Eq. S5). For the batch photoreactor, the reaction time was set to 90 min and the optimal mpg-CN\(\_x\) loading was established (Figure S6). A wide range of para-substituted benzyl alcohols were studied, and the PCP setup displayed a substantial improvement in Conv compared to its batch counterpart (Table 1). The enhancement of the flow setup compared to batch stems from a higher local loading of the photocatalyst in the light-responsive zone; by directing more of the substrate over the catalyst, and constantly replenishing the \(O_2\) at the reactive sites (by means of the constant airflow), a higher activity can be achieved for a given reaction. The reaction time in the batch photoreactor had to be increased to 4 h in order to obtain a Conv of similar magnitude to that obtained under continuous flow conditions with \(\tau\) set to 90 min (see Table 1, entry 1.i and 1,
respectively). As in the case of the PCP, carboxylic acid formation in-batch became more appreciable for the higher \( \text{Conv} \) value. However, a better \( \text{Sel} \) for the aldehyde product was obtained for the flow photoreactor (93% relative to 90% in-batch, Table 1 entry 1 and 1.i, respectively). This supports another general advantage of flow over batch photoreactors, in which the reaction mixture can be more easily removed from the photoactive zone, to minimize or lessen the formation of side-products resulting from secondary reactions or decomposition pathways\(^{33}\) (in this case, overoxidation of the aldehyde compound).

The external quantum efficiency (EQE) in-batch and in-flow was calculated to be 1.5\% and 1.6\% at \( \lambda = 470 \) nm, respectively (see Supporting Information, Eq. S6). Although the ascertained values are similar for the two photoreactors, it is noted that the flow system was not specifically optimized for this parameter. Further improvements in the \( \text{SA-to-V} \) ratio of the PCP, along with the positioning and intensity of the light source, can lead to enhancements in the EQE value.

Control experiments with the PCP in the absence of light did not result in alcohol conversion (Table 1, entry 1.ii). Using a biphasic setup (i.e. liquid-only flow, with the air feed switched off during operation), in which the substrate solution and solvent feeds were saturated with air prior to transferring into the syringes, resulted in a much lower \( \text{Conv} \) relative to the triphasic assembly (11\% compared to 80\%, Table 1, entry 1.iii and 1, respectively). Purging the PCP with \( \text{N}_2 \) and using \( \text{N}_2 \) purged substrate and solvent feed stocks resulted in \( \text{Conv} \) of 8\% (Table 1, entry 1.iv). Similarly, a low but significant \( \text{Conv} \) of 4\% was measured in batch under a \( \text{N}_2 \) atmosphere (Table 1, entry 1.iv). This low anaerobic background activity can be associated with the charge storage abilities of carbon nitride, whereby the oxidation and reduction steps can be decoupled, thus allowing for a small amount of substrate oxidation even in the absence of a suitable electron acceptor.\(^{34-36}\) Incomplete elimination of \( \text{O}_2 \) within the flow setup, and the trapping of some \( \text{O}_2 \) molecules within the mesoporous network of \( \text{mpg-CN}_x \),\(^{32}\) could also contribute to this non-zero \( \text{Conv} \) value under inert conditions. These results highlight the importance of the aerobic conditions and the advantage of using a gas–liquid (rather than a liquid-only) flow in which \( \text{O}_2 \) acts as both an electron acceptor and a subsequent reagent in the overall oxidation process.
Figure 2. Residence time profile with both starting material conversion and selectivity for aldehyde production (1-naphthalenemethanol concentration = 50 mM in MeCN). Inset: photoreactor re-usability with a total of 25 flow experiments in the PCP setup (1-naphthalenemethanol concentration = 100 mM in MeCN, τ = 90 min). General conditions for both plots: \( V_{air} = 0.5 \) sccm, blue LEDs (14.4 W at 470 nm), \( T = 55 \) °C.

Para-substituents with the lowest Hammett parameter (\( \sigma \)) generally achieved the highest conversions, as they are more effective in stabilizing the radical intermediate (Table 1, entries 2 and 3, versus 4 and 5; Figure S1).\(^{37}\) 4-tert-butylbenzyl alcohol displays a lower than expected Conv, which is presumably due to unfavorable steric effects (Table 1, entry 6). A higher Conv was achieved for all substrates under continuous flow compared to batch, as a consequence of the higher throughput efficiency provided by the PCP environment. The slightly higher Sel metric of the batch photoreactor is a consequence of the lower Conv. As discussed above (Figure 2), formation of the overoxidation product only becomes considerable once the concentration of the starting material decreases substantially, such that the oxidation of aldehyde begins to take place. Some variance in the M.B. between batch and flow was observed (average of 95% and 90% for entries 1 – 6 in Table 1, respectively; see Table S1 for full list of M.B. values), which is most likely due to the volatility of the aldehyde, which can escape from the flow reactor via the gas output stream. Thus, in
reality, **Conv** and **Sel** for the flow PCP system are likely higher than reported in Table 1, and system improvements are currently being made to increase the **M.B.**

### Table 1. Alcohol oxidation scope under batch and flow conditions, featuring control experiments performed in flow, based on Scheme 1a.

<table>
<thead>
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<th>Entry</th>
<th>Substrate</th>
<th>Flow</th>
<th>Batch</th>
</tr>
</thead>
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<tr>
<td></td>
<td></td>
<td><strong>Conv</strong></td>
<td><strong>Sel</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>(%)</td>
<td></td>
</tr>
<tr>
<td>1</td>
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<td>80</td>
<td>93</td>
</tr>
<tr>
<td>1.i</td>
<td><img src="https://example.com" alt="OH" /></td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>1.ii</td>
<td>No light</td>
<td>0</td>
<td>N.A.</td>
</tr>
<tr>
<td>1.iii</td>
<td>No air flow</td>
<td>11</td>
<td>100</td>
</tr>
<tr>
<td>1.iv</td>
<td>No air flow, N(_2) purged</td>
<td>8</td>
<td>100</td>
</tr>
<tr>
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<td>100</td>
<td>76</td>
</tr>
<tr>
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<td>76</td>
<td>88</td>
</tr>
<tr>
<td>4</td>
<td><img src="https://example.com" alt="OH" /></td>
<td>46</td>
<td>93</td>
</tr>
<tr>
<td>5</td>
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<td>44</td>
<td>93</td>
</tr>
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<td>6</td>
<td><img src="https://example.com" alt="OH" /></td>
<td>55</td>
<td>92</td>
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**a** General conditions: substrate concentration 75 mM in MeCN (CD\(_3\)CN in batch), blue LEDs (14.4 W at 470 nm), \(T = 55\) °C. Control flow or batch experiments detailed as sub-entries within the table. **b** Flow conditions: \(r = 90\) min, \(V_{air} = 0.5\) sccm. **c** Batch conditions: reaction time = 90 min, mp-g-CN, photocatalyst loading = 2 mg. **d** **Conv** calculated using 1,3,5-trimethoxybenzene as internal standard. **e** **Sel** for corresponding aldehyde. **f** Reaction time for batch photoreactor set to 4 h. **g** Substrate solution and solvent feeds purged with air for 15 min prior to transferring into the syringe. **h** PCP setup first purged with N\(_2\) (25 sccm) for 20 min, and then MeCN (1.5 mL, 0.028 mL min\(^{-1}\)); substrate solution and solvent feeds purged with N\(_2\) for 15 min prior to transferring into the syringe. **i** For the batch control, a N\(_2\) atmosphere was used.

Upon replacing the alcohol by a primary benzyl amine, a dibenzylc secondary (E)-imine was obtained as the major product (Scheme 1b). This resulted from the cascade condensation reaction between the primary aldimine intermediate with the
starting material in the presence of mpg-CN\textsubscript{x}.\textsuperscript{31} Preliminary batch studies revealed that the oxidation of benzylic amines proceeds at a faster rate than that of benzylic alcohols – hence a lower value for \( \tau \) was selected (60 min) for the continuous flow experiments. Analogous to the aerobic alcohol oxidations, the enhanced performance of the PCP in comparison to batch was observed (Table 2). We observe that the PCP setup does not impede the condensation step following initial amine oxidation. This result highlights the versatility of the flow system for various aerobic oxidation processes.

\textbf{Table 2.} Amine reaction scope under batch and flow conditions, featuring control experiments performed in flow, based on Scheme 1b.

<table>
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<tr>
<th>Entry</th>
<th>Substrate\textsuperscript{a}</th>
<th>\textbf{Flow\textsuperscript{b}}</th>
<th>\textbf{Batch\textsuperscript{c}}</th>
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<tr>
<td></td>
<td>\textbf{Conv\textsuperscript{d}}</td>
<td>\textbf{Sel\textsuperscript{e}}</td>
<td>\textbf{Conv\textsuperscript{d}}</td>
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<td>99</td>
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<td>N.A.</td>
</tr>
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<td>100</td>
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<td>99</td>
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<td>78</td>
</tr>
<tr>
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<td>No air flow\textsuperscript{f}</td>
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<td>88</td>
</tr>
<tr>
<td>6.ii</td>
<td>No light</td>
<td>0.8</td>
<td>93</td>
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</tbody>
</table>

\textsuperscript{a} General conditions: substrate concentration 75 mM in MeCN (CD\textsubscript{3}CN in batch), blue LEDs (14.4 W at 470 nm), \( T = 55 \) \textdegree C. Control flow experiments detailed as sub-entries within the table.

\textsuperscript{b} Flow conditions: \( \tau = 60 \) min, \( V\textsubscript{air} = 0.5 \) sccm.

\textsuperscript{c} Batch conditions: reaction time = 60 min, mpg-CN\textsubscript{x}, photocatalyst loading = 2 mg.

\textsuperscript{d} \textbf{Conv} calculated as per Table 1.

\textsuperscript{e} \textbf{Sel} is for the dibenzylic secondary (\( E \))-imine; minor impurity due to presence of primary benzyl imine.

\textsuperscript{f} Residual O\textsubscript{2} (from ambient air) still present in the liquid feeds.
However, there was a discrepancy in the M.B. metric when comparing the primary amine oxidation in the PCP setup and in-batch. The M.B. for the primary amines in-flow was on average ~70% (Table 2, entries 1 – 5; Table S2), whereas a M.B. of ~98% was obtained in batch. Flow experiments using 1,3,5-trimethoxybenzene as a tracer molecule (which was added to the amine solution and pumped simultaneously into the PCP, but was unreactive towards light activated mpg-CN₅) gave complete M.B. for the tracer compound (DMSO was used as internal standard in this case). Secondary amines faced no such issue (Table 2, entry 6) and a M.B. of 95% was achieved in-flow. This implied that a side-product from the reaction with the primary amine may have deposited within the PCP, and hence could not be accounted for during downstream liquid analysis.

High concentrations of benzylic primary amines are known to react with aerobic CO₂ to form a carbamic acid/carbamate salt mixture, which consists of a white precipitate that is insoluble in a number of organic solvents.³⁸,³⁹ We therefore propose that the mass imbalance could originate from the formation of a carbamic acid precipitate in the PCP. To verify this hypothesis, ¹³CO₂ was bubbled into a primary benzyl amine solution in MeCN (100 mM). This rapidly resulted in the formation of a white precipitate within the vial, that was found to be soluble in basic solution (TBAOH, 10 mM in MeCN). An infrared spectrum of this solid revealed the appearance of a single sharp band at 3380 cm⁻¹, indicative of the loss of a primary amine, and characteristic peaks centered at 1650 cm⁻¹ and 1620 cm⁻¹, which can be attributed to the asymmetric (C=O) stretch of the COOH moiety of the carbamic acid and the carboxylate anion, respectively (Figure S7).⁴⁰ Furthermore, a strong peak at ~158 ppm was observed for the carbamic acid carbon atom in the ¹³C NMR spectrum (Figure S8).⁴⁰ Finally, continuous flow experiments under alkaline conditions (10 mM TBAOH present in the substrate feed) increased the M.B. by 12%, as the hydroxide ions facilitate hydrolysis of the carbamic acid/carbamate salt.⁴¹ Our rational understanding of the flow reaction shown in Scheme 1b is that some degree of solvent evaporation occurs upon passing the solution through the heated zones (~55 °C) of the PCP, causing the local concentration of amine to increase at such points. Here, the low pressure of CO₂ from the compressed air feed is sufficient to react with the locally concentrated starting material to yield the carbamic acid/carbamate salt. Due to the insolubility of this mixture in MeCN, it precipitates from the solution and settles in the
PCP during operation. With the light switched off, and thus the temperature of the PCP maintained at ~22 °C rather than 55 °C, the M.B. for the primary benzyl amine increased from 68% to 82% (Supporting Information, Table S2, entry 1.i), supporting our rationale on the behavior of such compounds within the continuous flow environment. This kind of reaction environment is not experienced within the batch reactor, and hence, could explain the difference in the M.B. metric between the two systems. The PCP flow setup is therefore being investigated as a potential tool to explore other organic transformations, as it could potentially lead to a product distribution that differs from that obtained in a typical batch setup.

Conclusions

In conclusion, we have developed a triphasic photoreactor employing mpg-CN$_x$ to explore aerobic substrate oxidations and reactions in-flow under benign conditions. We investigated the impact of the morphology of the carbon nitride within the PCP under a continuous flow environment, and established that the mesoporosity plays a vital role with regards to the gas–liquid dynamics through the heterogeneous catalyst. Our mpg-CN$_x$–based continuous flow system is not only low cost and reusable for multiple flow cycles, but is also more practical and effective than a conventional batch photoreactor. This photocatalytic flow reactor is able to conduct oxidative transformations using a well-controlled air feed, which makes it more compatible with the safety restrictions surrounding the use of O$_2$ for industrial applications. Efforts are now being placed to utilize the overall visible light activated mpg-CN$_x$ PCP setup to investigate the improvement in performance of other known aerobic reactions,$^{31,42,43}$ and to probe novel organic transformations under continuous flow operation.

Associated Content

Further details on the experimental conditions (mpg-CN$_x$ fabrication, batch and flow photoreactor setup, MeCN suppression $^1$H NMR programme), extra figures (TEM, nitrogen sorption data, $^{13}$C NMR and FTIR spectra, optimization plots, images of the flow assembly), and extra tables (list of mass balance values) can be found in the Supporting Information.
Conflict of interest

There are no conflicts of interest to declare.

Acknowledgements

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References


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(X = O, NH)

\[ \text{R} \xrightarrow{\text{hv}} \text{R} \xrightarrow{\text{O}_2} \xrightarrow{\text{H}_2\text{O}} \]

\[ \text{R} \xrightarrow{\text{XH}} + \text{O}_2 \]

\[ \text{R} \xrightarrow{\text{X}} + \text{H}_2\text{O} \]