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#### Introduction

Recent years have witnessed a revival of photochemical transformations, and particularly photocatalysed reactions, where transition-metal catalysts and organic photocatalysts facilitate the construction of important chemical bonds *via* mild and orthogonal reaction conditions.<sup>1-6</sup> The transformative power of modern photocatalysis and its potential impact on sustainability<sup>7</sup> are evident in the number of recent reports detailing selective and highly efficient protocols that exploit visible light, operate at ambient temperature and are devoid of stoichiometric reagents or additives.

Importantly, this has enabled the application of photochemical processes not only in academia but also in industry<sup>8</sup> to bring about the functionalisation of valuable building blocks through previously elusive methods. Key to this development is thereby not only the development of photocatalysis as a field, but also the availability of cheap and monochromatic light emitting diodes (LEDs) as attractive light sources,<sup>9</sup> and lastly the advent of new reactor technology to provide ready-to-use and standardised set-ups in batch<sup>10</sup> and continuous flow mode.<sup>11</sup>

# Discovery of a photochemical cascade process by flow-based interception of isomerising alkenes\*

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Herein we report the discovery of a new photochemical cascade process through a flow-based strategy for intercepting diradicals generated from simple alkenes. This continuous process delivers a series of unprecedented polycyclic reaction products. Exploring the scope of this novel process revealed that this approach is general and affords a variety of structurally complex reaction products in high yields (up to 81%), short reaction times (7 min) and high throughputs (up to 5.5 mmol h<sup>-1</sup>). A mechanistic rationale is presented that is supported by computations as well as isolation of key intermediates whose identity is confirmed by X-ray crystallography. The presented photochemical cascade process demonstrates the discovery of new chemical reactivity and complex chemical scaffolds by continuously generating and intercepting high-energy intermediates in a highly practical manner.

Whilst the use of blue light (*ca.* 450 nm, *ca.* 64 kcal mol<sup>-1</sup>) is commonly favoured in photoredox applications due to the tolerance of highly functionalised substrates, UV light still plays an important role in photochemistry. This can be seen in [2 + 2]photocycloaddition processes of readily available alkenes affording important cyclobutanes as well as various photochemical rearrangements.<sup>3</sup> Modern high-power LEDs emitting in the UV-A region (310–400 nm, *ca.* 71–92 kcal mol<sup>-1</sup>) are particularly interesting in view of replacing classical mediumpressure mercury lamps that do not provide monochromatic light, generate significant amounts of heat, and often require filters to block undesired wavelengths that would result in sideproduct formation.

Recent work by Noël and co-workers demonstrated the power of photocatalysed HAT processes using UV-A emitting LEDs to bring about the effective C–H functionalisation of unreactive alkanes such as methane, ethane and propane in a continuous manner.<sup>12</sup> Additional studies by Booker-Milburn,<sup>13</sup> Cochran<sup>14</sup> and others<sup>15</sup> clearly showcase the value of flow-based photochemistry exploiting uniform and selective irradiation processes to generate diverse targets in a scalable and effective manner.

Our own laboratory recently exploited a high-power LED lamp (tuneable 50–100 W) emitting at 365 nm to realise the clean generation of a set of quinolines<sup>16</sup> (Scheme 1, top). In this application the direct irradiation of chalcones triggered isomerisation of the alkene and subsequent cyclocondensation. The replacement of a traditional medium-pressure mercury lamp with this high-power UV-A LED thereby minimised sideproduct formation, which increased reaction yields by 10–25% and furthermore enabled the telescoped hydrogenation of the quinolines to furnish a set of tetrahydroquinolines including the antimalarial alkaloid galipinine.

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Scheme 1 Prior work and approach to the new photocascade process reported herein.

#### Results and discussion

Taking inspiration from this quinoline forming photo-process we embarked on a study exploiting analogous chalcones<sup>17</sup> bearing an alkyne group in place of the previous amine (Scheme 1, bottom). As the quinoline forming process involved the breaking the C–C  $\pi$ -bond and the isomerisation of the alkene prior to cyclocondensation, we proposed that such putative diradical intermediates (e.g. 6) could be trapped by an adjacent alkyne moiety to facilitate the formation of a new C-C bond as well as a new ring system. By deviating from this simple alkene isomerisation path, we anticipated to open new avenues towards creating structural complexity whilst discovering new chemical reactivities. To test this hypothesis, the required scaffold (4) was readily prepared from 2-iodo benzaldehydes (5) via a Sonogashira cross-coupling followed by an aldol condensation process under alkaline conditions (see ESI† for full details). This straightforward route created a small set of substrates providing variation on the chalcone scaffold including three points of diversity (R<sub>1-3</sub>, Scheme 1, bottom).

To study the fate of these entities under standardised photochemical conditions a Vapourtec E-series flow reactor (10 mL FEP coil, 1/16" i.d.) was exploited in combination with a high-power LED lamp emitting light at a wavelength of 365 nm (see ESI<sup>+</sup> for full details). This reactor set-up was advantageous as the LED power can be regulated between 50-100 W, while the flow processing allows for short and uniform light pathlength, narrow residence time distribution and thus excellent reproducibility. Initial efforts exploited piperonalderived substrate 4a possessing a maximum absorbance ( $\lambda_{max}$ ) at 375 nm which closely matched the emission of the LED light source. Using the described set-up, a solution of 4a (50 mM in MeCN) was processed through the LED reactor set-up operating at 100 W with a residence time of 20 minutes. Pleasingly, TLC analysis indicated full conversion of 4a and formation of a major new product. Optimisation of this flow process quickly

indicated that clean formation of this new product can be achieved using only 70 W lamp power, in combination with a shorter residence time of 7 minutes at a concentration of 80 mM while regulating the reactor temperature to 25-30 °C (Table 1). These findings indicate that shorter residence times minimise over-irradiation (*i.e.* secondary photo-reactions) through improved spatiotemporal control in the flow reactor, which renders the desired product in higher yields.

This new product was subsequently purified by silica gel chromatography rendering a beige solid that was then studied by various spectroscopic techniques to establish its chemical structure. <sup>1</sup>H-NMR spectroscopy highlighted that a single entity had been obtained whereby the presence of diastereotopic methylene protons indicated a chiral, albeit racemic structure. Most notably, a set of four resonances was observed up-field of the methylendioxybenzene unit (5.8–6.8 ppm, see Fig. 1).

These four resonances integrated for one proton each and were part of two separate alkenes. Furthermore, a doublet with a coupling constant of 2.3 Hz was observed at 3.6 ppm (1 H), as well as an AB quartet at 3.5 ppm (2  $\times$  1 H). This unexpected finding, together with the absence of the enone protons, pointed towards a set of secondary transformations that involved not only the alkene and alkyne moieties, but also the phenyl ring adjacent to the carbonyl in 4a. Analysis of <sup>13</sup>C-NMR data along with various 2D NMR data corroborated this analysis suggesting that an unexpected and likely unprecedented photochemical cascade had taken place that transformed a benzene ring into two separate alkene moieties. Highresolution mass spectrometry confirmed a formula of the base peak of  $C_{22}H_{21}O_3$  (M + H<sup>+</sup>) which is identical to that of the substrate. UV-Vis spectroscopy revealed a diminished maximum absorbance of 348 nm indicating a less conjugated system compared to the substrate ( $\Delta \lambda_{max} = 27$  nm). While IR spectroscopy additionally indicated the presence of functionalities consistent with a conjugated ketone (1674  $\text{cm}^{-1}$ ) as well as alkenes and alkanes, it was not possible to unambiguously establish the connectivity of this new structure.

To resolve this situation, the synthesis of this new photoproduct was scaled in flow mode to provide larger quantities for crystallisation and subsequent X-ray diffraction analysis.





Fig. 1 Partial  ${}^{1}$ H-NMR spectrum of the photoproduct (**7a**, CDCl<sub>3</sub>, 500 MHz).

Gratifyingly, operating the flow process under the previously established conditions for 1 h reproduced the outcome of the initial experiments and rendered a total of 1.85 g of this product after purification. This not only equates to a high productivity of  $5.5 \text{ mmol h}^{-1}$ , but moreover demonstrates the straightforward scalability of continuous photochemical processes.<sup>18</sup> Attempts to crystallise this material were soon met with success and led to identifying the molecular structure based on single crystal X-ray diffraction. Fig. 2 shows this structure,<sup>19</sup> which confirmed the prior spectroscopic data and highlighted the formation of a new fused pentacyclic ring system (excluding the methylenedioxy ring).

The structure of this unique architecture is characterised by a cyclopentenone with fused cyclopentene and cyclobutene rings that are joint by a methine carbon and two adjacent quaternary carbon atoms (C6 and C7, Fig. 2). The cyclobutene ring and butyl chain display a *cis* relationship imparting a distinct three-dimensional shape to this structure. It is noted that this tricyclic subunit of **7a** occurs in the natural product tricycloclavulone<sup>20</sup> (**8**, Fig. 3), though published studies describing the construction of this scaffold unsurprisingly require significantly more steps compared to our new photochemical cascade.

Having demonstrated proof of principle for this new photochemical cascade we next wished to study the scope of this process. To this end we modified the scaffold of substrate 4 incorporating different appendages on both aryl moieties and



Fig. 2 Molecular structure of photo-cascade product 7a (CCDC-2072295).



Fig. 3 Structural comparison between 7a and tricycloclavulone 8.

the alkyne. Pleasingly the reaction tolerated different groups on the initial 2-iodo benzaldehyde unit, including 3,4-(methylene) dioxy, 3,4-dimethoxy and even unsubstituted systems ( $R_1 = H$ , Fig. 4). A distinct blue-shift of the maximum absorbance  $(\lambda_{max})$ for substrates 4h-j to ca. 310 nm was observed, however a shoulder reaching as far as 360 nm appears to provide sufficient overlap with light emitted from the UV-LED. It was furthermore established that variations on the alkyne including phenyl and alkyl groups are well tolerated. However, placing a cyclopropyl ring next to the alkyne did not provide the desired product (7g), which is consistent with the involvement of nearby radicals.<sup>21</sup> Lastly, introduction of substituents on the benzoyl moiety is possible providing the desired products (7d, 7j) with three contiguous quaternary centres. In all these cases the desired products were isolated in high chemical yields. It was furthermore established that the cascade products possess significantly lowered  $\lambda_{max}$  values compared to the substrates (300-345 nm) which is consistent with the apparent loss of conjugation.

The next part of our study concerned probing the mechanism by which these unique photocascade products are generated. Specifically, this entailed the fate of the aryl ring that is transformed into the unusual bicyclo[3.2.0]hepta-2,6-diene scaffold. As we initially hypothesised that this ring system may have arisen from a cycloheptatriene by a photochemical  $4\pi$ -electrocyclisation process, we attempted to experimentally



Fig. 4 Structures and  $\lambda_{max}$  values of photo-cascade products 7a-j.

find proof for this proposal. We therefore wished to again exploit the benefits of continuous flow processing allowing for fine tuning of reaction conditions based on superb spatiotemporal control.<sup>22</sup>

Therefore, the flow process was repeated for the photocascade of **4a**, however, utilising conditions that would lead to incomplete conversion of the substrate. An experiment was thus run in which a more concentrated solution of **4a** (100 mM MeCN) was processed with only 5 minute residence time at 50 W lamp power (Scheme 2).

Analysing the <sup>1</sup>H-NMR spectrum of the crude product, we were pleased to identify a new species (**13a**, *ca*. 65%) besides the previous photo-cascade product (**7a**, *ca*. 35%). Purification by chromatography rendered this new product as a yellow solid ( $\lambda_{max} = 383$  nm) in 50% isolated yield. HR-MS analysis confirmed that it was an isomer of **7a** (molecular formula:  $C_{22}H_{20}O_3$ ). In addition, the <sup>1</sup>H-NMR spectrum of this compound clearly showed that the resonance for the methine proton at 3.6 ppm was absent, while a total of five olefinic resonances were visible (5.9–7.1 ppm; Fig. 5). COSY and NOESY experiments subsequently revealed that these five protons are part of the same spin system, which together with all other data suggested this species to be the cycloheptatriene intermediate **13a**.

The molecular structure of **13a** was subsequently secured through single crystal X-ray diffraction analysis<sup>19</sup> and confirmed both our initial hypothesis and structural assignment.

Considerable amounts of **13a** (*ca.* 250 mg) were soon prepared allowing for a final test to prove its role as a late-stage intermediate. Pleasingly, subjecting a solution of **13a** (80 mM MeCN) to the regular photoreaction conditions (70 W, 7 minutes residence time, Scheme 3) afforded **7a** as sole product in 97% purity (by <sup>1</sup>H-NMR).

To shed some light on the mechanistic pathway operational for this photo-cascade process we furthermore performed DFT calculations as implemented in Gaussian 16.<sup>23</sup> Specifically, calculations at the M062X<sup>24</sup>/6-31+G(d,p)<sup>25</sup> level of theory in a solvent model SMD (acetonitrile) at 298 K were used and subsequently refined for single point energies using M062X/6-311+G(d,p)<sup>25</sup> (acetonitrile). These calculations suggest a first step in which the alkene  $\pi$ -bond initially undergoes homolysis to a diradical species (**int2**) that is intercepted by the alkyne through a 5-*exo*-dig cyclisation step (Fig. 6).<sup>26</sup> The resulting intermediate (**int3**) undergoes H-atom translocation to render a fully substituted cyclic enone with an adjacent carbene (**int4**).<sup>27</sup> This carbene species then adds across the adjacent benzene ring to render a norcaradiene ring system.<sup>28</sup> These



Scheme 2 Flow synthesis of 13a and its conversion into 7a.



Fig. 5 Molecular structure (CCDC-2072296) and partial <sup>1</sup>H-NMR spectrum of cycloheptatriene **13a** (CDCl<sub>3</sub>, 500 MHz).

calculations show the possibility of both a concerted (in the singlet state) and stepwise (*via* switch to triplet intermediate) pathway for this process. Subsequent  $6\pi$ -electrocyclisation occurs in a conrotatory sense under photochemical conditions to furnish cycloheptatriene **13a**, which upon disrotatory  $4\pi$ -electrocyclisation yields the scaffold of **7a**.

As depicted in Fig. 6, the cycloheptatriene species (**int6**) is thermodynamically more stable than the final cascade product, which is in good agreement with observations that solutions of this material (*e.g.* **13a**) are stable for several weeks under laboratory conditions. However, the high-power UV-LED lamp provides enough energy to overcome this barrier towards the final products. Diminished absorbance of the photoproduct **7a** is the likely cause for the observed irreversibility of the final step in this cascade rendering the mechanism as shown in Scheme 3.

Armed with this mechanistic understanding we embarked on a final extension to probe the effect of further substituents in the *para* position of the aryl enone fragment. Both electron



Scheme 3 Mechanistic rationale for continuous photocascade process.



Fig. 6 Free-energy profile for the studied reaction of substrate 4a; S and T refer to singlet and triplet states, respectively.



**Fig. 7** Correlation between energy barrier and product formation (<sup>a</sup> yield for butyl side chain; <sup>b</sup> attainable in 50% yield, see Scheme 2).

donating (R = OMe) and electron-withdrawing (R = F, CN,  $CF_3$ ) groups were chosen to complement the previously studied cases (R = H, Me) where the substrates uniformly featured the piperonyl moiety as well as a propyl chain introduced with the alkyne. Using the *para*-fluoro substrate quickly established the

formation of the anticipated polycyclic scaffold 7 (Fig. 7) in good yield. However, when trialling either the methoxy derivative, or the electron-deficient nitrile and trifluoromethyl variants, the corresponding cycloheptatriene products 13 were obtained as principal products. These materials were found to absorb light in the UV-A region which may lead to secondary photoreactions and likely accounts for the modest yields (see ESI<sup>†</sup> for details). The results imply that the electronic effect of the para substituent is not decisive in determining the reaction outcome. To understand these results, we turned again to computations at the same level of theory as before which highlighted the interplay between the energy barrier  $(13 \rightarrow 7)$  as well as the relative stability of the ultimate product with regards to the cycloheptatriene species. In the case of the *p*-methoxy system (13l), a relatively low energy barrier together with the reversibility of this reaction favours the more stable 7-membered ring as the product. On the other hand, the energy barrier for both the trifluoromethyl and nitrile system is excessively high thus precluding the final step to occur efficiently. Conversely, in case of the *para*-substituent being a proton or methyl group (*e.g.* 7a, 7d) the energy barrier is surmountable under the reaction conditions whilst a significantly lower absorbance of the product impedes the reverse reaction.

#### Conclusions

In summary, the discovery of a novel photochemical cyclisation cascade is reported rendering complex three-dimensional structures in good to excellent yields. Flow processing in combination with a tunable high-power LED light source emitting at 365 nm was exploited to intercept *in situ* generated diradicals and rapidly generate a small selection of photoproducts with throughputs up to 5.55 mmol  $h^{-1}$ . Besides, modifications to the optimised conditions enabled the

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generation and spectroscopic characterisation of an advanced cycloheptatriene intermediate that supports the computed mechanism. Continuous flow synthesis facilitated the discovery of this new cascade process, as well as the efficient and scalable production of additional members of this unprecedented class of compounds. Further efforts towards expanding the scope and applicability of this unique light-driven cascade process are underway in our laboratories.

# Data availability

Data available in the SI includes spectroscopic and computational data.

### Author contributions

The work was conceptualised by MB and MDF. Experimentation was performed by MDF and MB. Computations were performed by CT and GSS. X-ray analysis was undertaken by ASB. The first draft of the manuscript was prepared by MB and the final draft was edited by all the authors.

# Conflicts of interest

There are no conflicts to declare.

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