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Continuous synthesis of pyridocarbazoles and initial photophysical and bioprobe characterization†

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Pyridocarbazoles when ligated to transition metals yield high affinity kinase inhibitors. While batch photocyclizations enable the synthesis of these heterocycles, the non-oxidative Mallory reaction only provides modest yields and difficult to purify mixtures. We demonstrate here that a flow-based Mallory cyclization provides superior results and enables observation of a clear isobestic point. The flow method allowed us to rapidly synthesize ten pyridocarbazoles and for the first time to document their interesting photophysical attributes. Preliminary characterization reveals that these molecules might be a new class of fluorescent bioprobe.

Recently a family of complexes1 exhibiting high affinities and selectivities for individual protein kinases was reported.2 These inhibitors are powerful signal transduction probes and may eventually lead to transition metal-based drugs.3 The inspiration for the GSK-3 inhibitor (2, one representative of this family of complexes)4 was the natural product staurosporine (1) in which the indolocarbazole glycine was mimicked by the novel pyrido[2,3-a]pyrrolo[3,4-c]carbazole-5,7(6H)-dione ligand. A key step in the synthesis of such pyridocarbazole ligands is a 6π-electrocyclization (eqn (1)).

[Diagram of staurosporine (1) and Meggers Group’s GSK-3 inhibitor (2)]

The success of this class of heterocycles in combination with difficulties in scaling up the photochemical cyclization

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are soluble in toluene. A solution of 3 (1 mg mL$^{-1}$) was drawn through the tubular microreactor using a syringe pump at flow rates corresponding to the residence times described in Fig. 1B. Each reaction used 2 mg of material (Fig. 4).

The UV/Vis spectra of 3 and 4 overlap exhibiting molar absorptivities of 16 000 and 12 000 M$^{-1}$ cm$^{-1}$ (320 nm), respectively, indicating that a small inner-filter effect will inhibit the rate of photocyclization. Regardless, we monitored the reaction as a function of residence time by UV/Vis (0.12 min to 10 min; Fig. 1). The spectra exhibit a well-defined isosbestic point indicating that the photocyclization proceeds from 3 to 4 via no observable intermediate (Fig. 1B), consistent with a rate-determining photoelectrocyclization followed by rapid loss of HBr (Fig. 3). Crude $^1$H-NMR spectra recorded after each residence time revealed that the reaction proceeds cleanly from 3 with only a small amount of by-product formed (5). Fig. 2 features $^1$H-NMR data for a segment of the aromatic region. Product is formed even at residence times of just 0.12 min and the reaction is near complete in 20 min. By-product to product ratio decreases from 1 : 3 to 1 : 8 from earliest to latest residence times. NMR data for the by-product are consistent with the previously described structure 5 shown in Fig. 3.

The observation that 4/5 increases with time suggests that 5 forms directly from 3 with a slower rate constant compared to the formation of 4, and that 5 can revert back to 3. Both the NMR and UV/Vis data support the model that 3 proceeds directly to product via intermediates too short-lived to be observed by these techniques. In addition, these data indicate that the desired photoelectrocyclizations occur cleanly under continuous flow conditions (see Fig. 5 and text below).

The method’s generality was demonstrated through the synthesis of 10 pyridocarbazoles (4 and 6–14). The substrates fit into two categories: high and low solubility in toluene. Substrates exhibiting low solubility yielded products that precipitated out of solution (Fig. 4). Fortunately, we observed no clogging even at flow rates as low as 330 µL min$^{-1}$. Below 300 µL min$^{-1}$, we observed variable amounts of solid emerging from the reactor as well as solid left in the reactor once the experiment was concluded. The insoluble substrates typically lacked a solubilizing group such as a silyl imide or ether. Regardless,
yields were good to excellent, such that some products showed no evidence of impurities by NMR (>95% pure). Most reactions required only a silica plug to remove the pyridinium by-product described above.

Flow methods are more attractive than batch for preparing pyridocarbazoles because the residence time is short (<20 min) relative to batch (>5 h). Comparison of the starting material, product (13) from flow and product (13) from batch irradiation shows distinct differences (Fig. 5). For the batch reactor case, significant amounts of a by-product form (arrows) with an \( R_t \) immediately above the desired product. This emerging by-product hampers purification.

Preparing the pyridocarbazoles rapidly enabled comparison of the photophysical properties. These fluorophores are interesting because they can have three orthogonal points of attachment (Fig. 6). Most exhibit a blue-green emission, but when strong \( \pi \)-donors are present, the emission is red-shifted to yellow/orange color (10). As representatives, we measured photophysical data for 11–14 at the same concentration (Fig. 7). This relative data indicates that 11/14 are 100× more fluorescent than 12/13 (see inset for fluorescent signal of 12/13). Compounds 11/14 have large Stokes shifts, 74 nm/83 nm, and ns decay times (both ca. 1.5 ns), making them similar to probes such as Cy5. While we have yet to collect quantum yield data, the ease with which these chromophores can be modified combined with their promising photophysical properties indicates that pyridocarbazoles could be very attractive for bioconjugation applications.

To demonstrate that these chromophores have potential as bioprobes, macrophage cell line RAW 264.7 was treated with compounds 7 and 8 for 20 min at 15 \( \mu \)g mL\(^{-1} \) concentration with a final DMSO content of 0.5%. The treated cells were imaged using a LSM-700 confocal microscope (Carl Zeiss AG; FITC channel) to detect the localization of the compounds. As can be seen in Fig. 8, the chromophores are easily observed and appear localized within membranes. This proof of concept illustrates that these chromophores remain fluorescent under biological conditions.

We have demonstrated that a flow photoreactor is an excellent tool to produce pyridocarbazoles. The reaction occurs with no observable intermediates. Considering that 366 nm light is unlikely to break C–Br bonds, we propose that this reaction proceeds via a rate-limiting [6\( \pi \)]-cyclization followed by rapid extrusion of HBr to yield the desired products. Comparing long and short irradiation times, we have found that long irradiation results in formation of a new unwanted product that would be challenging to remove. The rapid and clean conversions are in part due to the high photoflux provided by the small dimensions of the photoreactor. These data demonstrate that the study and synthesis of compounds via photoelectrocyclizations benefit from being run in flow and that pyridocarbazoles are exciting from the standpoint of both metal coordination and photophysics. Furthermore, we have demonstrated that these compounds exhibit interesting photophysical properties including large Stokes shifts and are bright under physiological conditions. Strong \( \pi \)-donors such as dimethylamine result in significant red shifting of the fluorescence emission, indicating that the color of these chromophores can readily be tuned.

We are now working to develop novel bioprobes based on the pyridocarbazole core.

**Experimental section**

The monobromides were synthesized according to published procedures. See ESI for experimental details. A representative photocyclization: compound 3 (33 mg, 0.062 mmol, 33 mL Toluene) was filtered through a 0.45 micro PTFA syringe filter and the solution was primed into a Vapourtec R2C+ system. The solution was pumped through a 5 mL photoreactor at 333 \( \mu \)L min\(^{-1} \) (15 min res time). The desired material was collected and concentrated to yield a dark orange solid and chromatographed over silica gel using a 3.5 Hex, 1.5 2% Et\(_3\)N/EtOAc solvent system (\( R_t = 0.21 \)). The fractions containing product were concentrated to yield 25 mg (89% yield) of an orange solid.
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Notes and references

12 Vapourtec Ltd. http://www.vapourtec.co.uk is gratefully acknowledged for providing R2C+ and R4 flow-chemistry systems.
15 All compounds matched the characterization data referenced for each compound. For those compounds lacking a citation, we provide characterization information in the ESL†.
17 The origin of the quenching observed for compounds 12 and 13 is yet to be determined. Possible explanations being considered are side chains enabling non-radiative relaxation or that the fluorine present introduces low-lying πσ* states – see: M. Z. Zgierski, T. Fujiwara and E. C. Lim, J. Chem. Phys., 2005, 122, 144312 or that the fluorescent present lowers the indole N–H ππ* facilitating photoinduced proton transfer.