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Cyclopropanation using flow-generated diazo compounds†

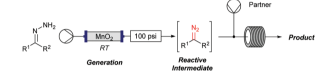
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We have devised a room temperature process for the cyclopropanation of electron-poor olefins using unstabilised diazo compounds, generated under continuous flow conditions. This protocol was applied to a wide range of different diazo species to generate functionalised cyclopropanes which are valuable 3D building blocks.

Introduction

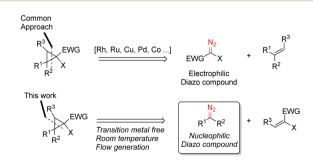
Flow chemistry is finding increasing use as an enabling technology in both industrial and academic settings, providing a number of advantages over traditional batch mode processes and thereby leading to a variety of interesting opportunities. Many reports to date have involved the adaptation of a single transformation from batch to flow modes.² From our perspective, a more important goal would be the discovery of new patterns of reactivity. Using flow chemistry methods the space/ time dynamics of a reaction profile can be adjusted along with other reaction parameters (i.e. pressure, temperature and mixing) that can be controlled accurately. New opportunities arise especially in harnessing reactive intermediates, which can be very challenging when using conventional methods. In a flow mode, it is possible to continuously generate highly reactive species free from contaminating reagents and then deliver these to a reaction zone for combination with various partners to afford a range of products.

Using this approach, we have previously established a continuous process for the production of transient diazo species on demand, as a class of interesting reactive building blocks (Scheme 1).³ Notably, this approach enabled the continuous generation of highly reactive intermediates and avoided the hazards associated with bulk handling of these species.⁴



Scheme 1 Flow generation, translocation and reaction of diazo intermediates.

The cyclopropane ring features in many natural products and often in bioactive and other functional materials due to its particular conformational constraints and properties.⁵ Accordingly, cyclopropanation of olefins using stabilised electrophilic diazo compounds has been well studied. In particular, transition-metal catalysis for the coupling of olefins and diazo compounds are amongst the most well developed methods for cyclopropane preparation. By contrast, unstabilised diazo compounds are underexplored. Interestingly, this highly energetic species often demonstrates a reversed reactivity pattern due to their amphiphilic character (Scheme 2).7 Further to our initial investigations using the flow generate/ translocate/react protocol for sp²-sp³ cross couplings, we envisaged that the diazo species could react with different electron poor olefins in order to give access to substituted cyclopropanes, under mild conditions and without the need for any transition metals in the coupling step. This method would



Scheme 2 Cyclopropanation of diazo species.

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represent an attractive alternative approach to the preparation of poly functional cyclopropanes.

Results and discussion

The study commenced with the use of the readily prepared hydrazone 1a as model substrate for the reaction.8 We soon found that in AcOEt as solvent, a consistent and clean stream of diazo compounds 2a was generated by passage through a column packed with activated manganese dioxide. As in previous studies,3 the use of Hünig's base was necessary to suppress the formation of by-products. Using this protocol, we were able to rapidly establish the cyclopropanation procedure using the olefin substrate 3a. Under optimised conditions, a solution of 1a (0.1 M in AcOEt, 2 equiv. of DIPEA) was passed through a column reactor packed with activated MnO2. The exiting stream of diazo (0.5 mL min⁻¹) was combined with a solution of olefin (3a, 0.05 M in AcOEt, 0.5 mL min⁻¹) after mixing at a T-piece and then stirred at room temperature for 2 h. With these optimised conditions, 4a was isolated in 89% yield (Scheme 3). The process was scaled up using a column of

Scheme 3 Cyclopropanation reaction via the generate/translocate/ react protocol (major isomer reported in scheme); a 0.2 mmol scale; ^b 4.25 mmol scale.

Scheme 4 Flow diazo preparation and cyclopropanation sequence using: (a) a heated reactor coil; (b) direct evaporation.

3 g MnO₂ to produce 1.14 g of cyclopropane 4a (86% yield) in one run using only 1.2 equiv. of diazo compound. Notably, the transformation was also achieved with high diastereoselectivity (10:1). No attempt was made at this stage to further increase the scale of the reaction.

With the aim of continuously processing material, we briefly investigated the flow cyclopropanation under different conditions (i.e. temperature and residence time) and rapidly found that running the reaction continuously in a 20 mL

Table 1 Cyclopropanation of different diazo species with olefin 3a^a

	CO₂Me NHAc 3a ○ 0.5 mL min ⁻¹				
N, NH ₂ MnO ₂ 100 R R ₁ R ₁ 0.5 mL min ⁻¹	$ \begin{array}{c} $	$R_1 \xrightarrow{R_1 \text{NHAc}} R_1 \xrightarrow{\text{NHAc}}$			

•		-			
Entry	Substrate	Product		Yield ^b	dr
1	N ₂	NHAc CO ₂ Me	4a	89%	10:1
2	N ₂ H Me	NHAc CO ₂ Me	4b	99%	10:1
3	N ₂ H OMe	NHAc CO ₂ Me	4c	97%	10:1
4	MeO H	NHAc CO ₂ Me	4d	89%	10:1
5	F H	ÖMe NHAc CO ₂ Me	4e	80%	8:1
6	O_2N H	NHAc CO ₂ Me	4f	79%	7:1
7	N ₂ H	NHAc CO ₂ Me	4g	59%	6:1
8	N_2	NHAc CO₂Me	4h	99%	10:1
9	N_2	NHAc CO ₂ Me	4i	39%	5:1
10	N ₂ Me	Me, NHAc CO ₂ Me	4j	55%	5:1
11	N ₂	NHAc CO ₂ Me	4k	63% ^c	5:1

 $[^]a$ 0.2 mmol scale. b Isolated yield after aqueous work-up and silica gel plug filtration. ^c Ratio of E/Z 3 : 1.

polymer (PFA) coil at 60 °C, a continuous throughput of 0.93 g h⁻¹ of material could be obtained. Notably, in addition to this we found that the combination of the two reaction streams and the direct feed into a rotary evaporator gave complete conversion to the cyclopropyl derivative 4a in situ (Scheme 4).

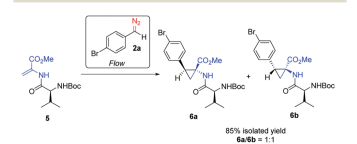
Next we wished to assess the scope of this new room temperature cyclopropanation process. Electron-withdrawing substituents on the aryl ring of diazo species (Table 1) were well tolerated, giving cyclopropanes 4a, 4e and 4f in very good yields (79-89%). Similarly, electron-donating groups also generated cyclopropanes in high yields (4b-d, 89-99%). Importantly, the steric hindrance exerted by ortho substituents seems not to affect the reaction outcome, as illustrated by comparison between 4c and 4d. This unusual feature of the protocol contrasts with metal-catalysed procedures for cyclopropanation.⁶ Heterocycles are well tolerated (4g, 4h) as is the presence of unsaturation (4i and 4k). Notably, the use of the ketone derivatives (Table 1, entry 10) gave an interesting 55% isolated yield of the product 4j, with a good level of diastereoselectivity (5:1). One remarkable aspect of the protocol is that all the products were obtained with good to excellent diastereoselectivity (Table 1).

Additionally, we were also pleased to find that increasing the functional complexity of the olefinic partner did not affect the outcome of the reaction as shown by reacting the diazo species 2a with the more elaborate structures (i.e. 5, Scheme 5) which gave high yield of the corresponding cyclopropyl peptides 6a and 6b.

We also decided to investigate the preparation of specifically challenging, unstable and difficult to access cyclopropyl compounds, in particular substituted donor-acceptor cyclopropanes.9

For example, the reaction of the diazo 2a with olefin 3c gave 7a in 55% yield as a single diastereoisomer (Scheme 6).

While this yield was lower than desired, the olefin starting material had all been consumed. Further attempts with different diazo species failed to give any isolable products. We quickly realised the chemical sensitivity of this class of cyclopropane products to silica gel chromatography and devised an alternative "work-up" involving the use of LiAlH₄ (Scheme 7) to give the corresponding diols which were readily isolated (Table 2). This 2-step transformation provides synthetically useful yields with very high diastereoselectivities.



Scheme 5 Diastereoselective cyclopropanation of dipeptide 5 to 6a/6b

Scheme 6 Diastereoselective cyclopropanation of 3b.

Scheme 7 Diastereoselective cyclopropanation of 3b, followed by reduction to diol 8.

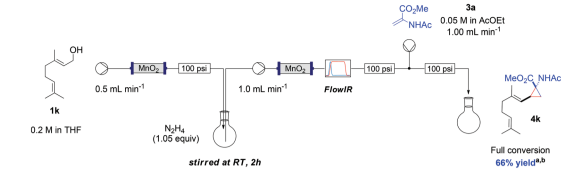
Table 2 Cyclopropanation of different diazo species with olefin 3b^a followed by reduction to diol 8

Entry	Substrate	Product		$Yield^b$	dr
1	N ₂	Me OH	8a	48%	>20:1
2	Br N ₂	Me OH	8b	55%	>20:1
3	F II	Me Me OH	8c	58%	>20:1
4	N ₂	Me OH	8d	40% ^c	>20:1

^a One-pot procedure at 0.2 mmol scale. ^b Isolated yield after aqueous work-up and silica gel plug filtration. c Ratio of E/Z 1.2 : 1.

Another valuable aspect of flow chemistry is associated with the advantages of continuously flowing material though a multistep sequence without the need to isolate intermediates. To showcase this aspect, we devised a 4-step sequence, whereby the use of a very cheap source of material (geraniol) would generate a more complex molecular product, 4k (Scheme 8).

During optimisation of this procedure, we discovered that an excess of MnO2 (7 g) was required for the complete oxidation of geraniol (delivered at 0.5 mL min⁻¹ as a 0.2 M solu-



Scheme 8 Telescoped sequence for the preparation of cyclopropane amino acid 4k. ^a Yield following concentration, measured by ¹H-NMR compared to an internal standard; ^b reaction run on 1 mmol scale.

tion in THF) to geranial (step 1).¹⁰ The output from the reagent column was collected in a reactor flask containing 1.05 equiv. of hydrazine. After stirring for 2 h at room temperature (step 2), the crude reaction mixture was pumped through a pre-activated column of MnO₂ (1.0 mL min⁻¹) to generate the diazo intermediate (step 3). The MnO₂ column served also to decompose any residual hydrazine.¹¹ In-line IR monitoring at around 2070 cm⁻¹ identified the diazo product exiting the reactor. This diazo stream was finally combined in a T-piece with a stream (1.0 mL min⁻¹) containing olefin 3a (step 4). Direct evaporation of the final reaction mixture in a rotary evaporator gave 4k in good yield.

Conclusions

In conclusion, we devised a system that could deliver, rapidly and efficiently, differently decorated cyclopropanes under flow conditions, with no need for precious metal catalysis and under remarkable mild conditions (room temperature). The process was successfully applied to a wide range of different *unstabilised nucleophilic diazo* species which were combined with electron-poor olefins. In this work, we demonstrated how flow chemistry can serve as a discovery tool for generating new chemical reactivity windows to prepare valuable products. This approach again exemplifies the opportunities that arise by using flow chemistry methods.

Acknowledgements

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