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Flow chemistry as a discovery tool to access sp^2 – sp^3 cross-coupling reactions *via* diazo compounds†Duc N. Tran,^a Claudio Battilocchio,^a Shing-Bong Lou,^a Joel M. Hawkins^b
and Steven V. Ley^{*a}

The work takes advantage of an important feature of flow chemistry, whereby the generation of a transient species (or reactive intermediate) can be followed by a transfer step into another chemical environment, before the intermediate is reacted with a coupling partner. This concept is successfully applied to achieve a room temperature sp^2 – sp^3 cross coupling of boronic acids with diazo compounds, these latter species being generated from hydrazones under flow conditions using MnO_2 as the oxidant.

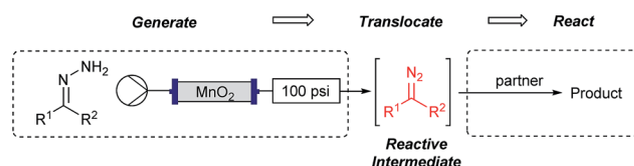
Introduction

Chemical processes define our modern world in terms of providing society's functional materials.¹ Many of these processes involve complex reaction pathways and proceed *via* transient reactive intermediates. The lifetime of these reactive species can vary enormously and their selectivity can often be difficult to exploit. Their high level of reactivity, when generated in batch mode, can be problematic because of incompatibilities with their chemical environment or functional groups, or because of associated exothermic properties. A case in point relates to diazo compounds which are valuable reactive intermediates.² For example, isolated stabilised diazo compounds, bearing one or more electron withdrawing groups, have been extensively investigated in a wide range of applications such as dipolar cycloadditions,³ carbonyl homologation⁴ and cyclopropanation.⁵ However, their unstabilised alkyl and aryl diazo counterparts are much more difficult to harness.^{2,6} Their chemical preparation is limited due to the often tedious, corrosive or toxic synthesis protocols.^{6a,7} *In situ* generation of diazo compounds therefore represents an interesting opportunity for synthesis,⁸ although any new procedure must be compatible with both the generation of the diazo compound and with its subsequent chemical transformation.

Manganese dioxide (MnO_2) has been used previously in batch mode to generate diazo compounds from hydrazones, although with low efficiency and with very specific substrates.⁹ Following on from the advantages that accrue when using MnO_2

under continuous flow reactions,¹⁰ we believed that we could devise a system capable of delivering reactive *unstable* diazo compounds on demand. Also, with the aim of better exploiting the potential of the flow chemistry,^{11,12} we anticipated that these flow-generated transient species could be rapidly transferred into another chemical environment, in order to react the aforementioned species with a suitable partner, hence avoiding interference by spent reagents (Scheme 1). This generation/translocation/reaction sequence should immediately provide access to new chemical space for the discovery of new reactions for these reactive intermediates.

Metal free sp^2 – sp^3 cross-coupling reactions have recently become a new focus of the chemical community.¹³ In 2009, Barluenga *et al.* reported an efficient coupling between tosylhydrazones and boronic acids.^{13a} Further studies with saturated heterocyclic sulfonylhydrazones by Nakagawa,^{13c} Kirschning,^{13e} and ourselves,^{13f} demonstrated a wider scope for the reaction (Scheme 2). Recently, Wang and co-workers reported an efficient synthesis of α -aryl esters and nitriles *via* a diazotization method for the generation of *stabilised* diazo compounds.^{13g} However, the relatively harsh reaction conditions (up to 110 °C), large amount of inorganic additives (bases, $NaNO_2$, NH_4Cl) and low atom economy represent potential drawbacks of these transformations. We envisaged that we could access new reactivities of *unstabilised* diazo compounds, in particular their coupling with boronic acids at room temperature.



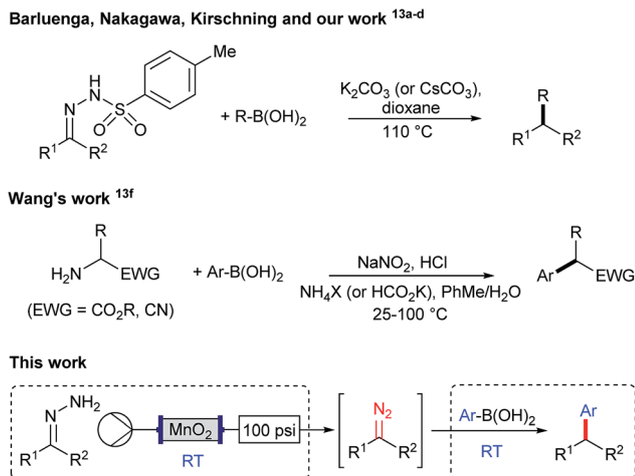
Scheme 1 Flowing scheme for the generation, translocation and reaction of the diazo species.

^aInnovative Technology Centre, Department of Chemistry University of Cambridge, Lensfield Road, Cambridge, CB2 1EW, UK. E-mail: svl1000@cam.ac.uk

^bPfizer Worldwide Research and Development, Eastern Point Road, Groton, CT 06340, USA

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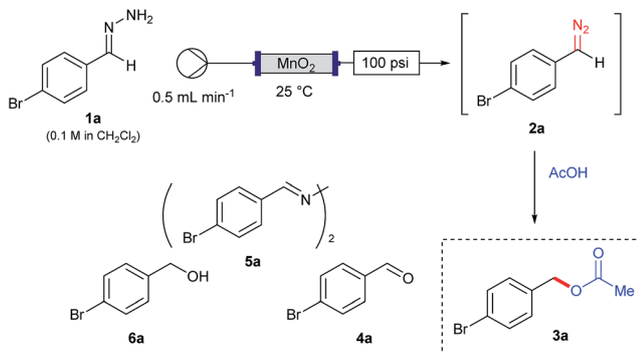


Scheme 2 Selected metal-free sp^2 - sp^3 cross-coupling reactions.

Results and discussion

Generation of diazo compound

Initial investigations focused on 4-bromophenyl hydrazine (**1a**) as model substrate for the generation of the diazo compound **2a**. To evaluate the generation of the transient diazo species, the material processed was quenched with acetic acid to yield the corresponding acetic acid ester **3a** (Scheme 3).⁶ Amorphous MnO_2 packed into the flow cartridge gave very poor conversion to **2a** with mainly starting material being recovered. By contrast, a column packed with *activated* MnO_2 gave full conversion to various products but with aldehyde **4a** being the main product observed. This result was also consistent with batch studies where aldehyde or azine by-products were generally observed at *ca.* 30% yield, even under carefully controlled reaction conditions.^{9b,c,f} However, conditioning of the packed column with a small amount of sacrificial hydrazine¹⁴ effectively reduced the formation of aldehyde and increased the production of diazo compound **2a** as determined *via* the formation of ester **3a**. Furthermore, addition of a base (Hünig's base, 2 equiv.) to the feedstock solution provided a reliable system with greatly reduced amounts of by-products generated.¹⁵



Scheme 3 Flow generation and quenching of 1-bromo-4-(diazo-methyl)benzene.

Under the optimised conditions, a 0.1 M solution of hydrazine **1** was passed through a conditioned MnO_2 column (0.86 g, activated MnO_2) at 0.5 mL min^{-1} with 2 equiv. of Hünig's base in CH_2Cl_2 . In-line IR data (collected using an in-line infrared spectrometer) suggested that a typical load of activated MnO_2 (10 mmol) could cleanly generate 2 mmol of diazo compound over 40 minutes. The generation of diazo compound, which proved to be highly practical as the whole process was carried out at room temperature, was followed by an esterification step of the transient species produced.

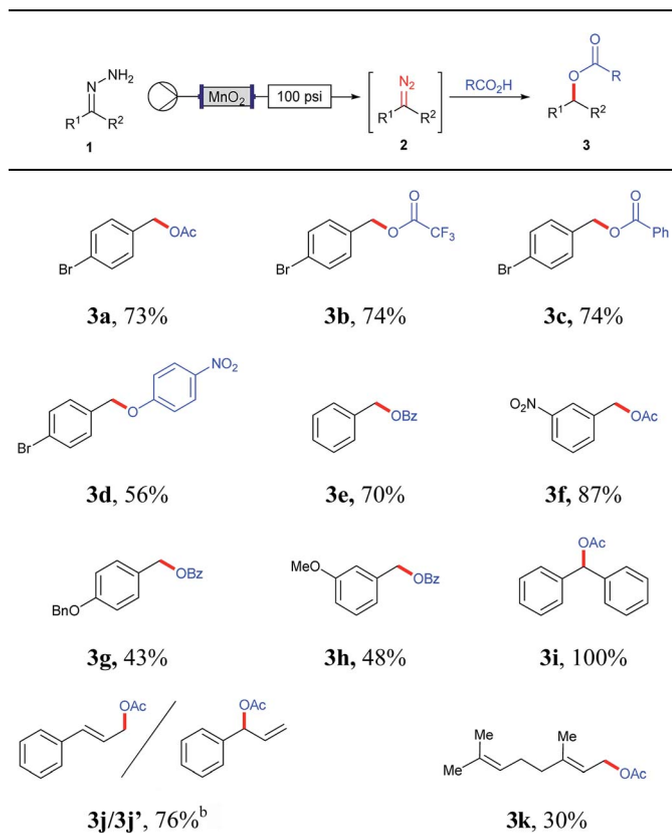
To explore the general application of this process, different hydrazone derivatives were examined (Table 1). The generated diazo compound was quenched with different carboxylic acids (**3a-3c**) and one phenol substrate (**3d**). The reaction was compatible with both electron poor and electron rich aromatic hydrazones. Electron poor aryl diazo compound gave in general better yields as they are more stabilised. Investigation of the reactivity of a vinylic diazo compound revealed the formation of two regioisomers (**3j** and **3j'**) after reaction with excess AcOH. However, when geranyl hydrazone was processed *via* our protocol only one isomer was obtained, though only in moderate yield (**3k**). On the other hand, diphenyl ketone hydrazone gave ester **3i** in excellent yield.

Coupling reaction with boronic acids

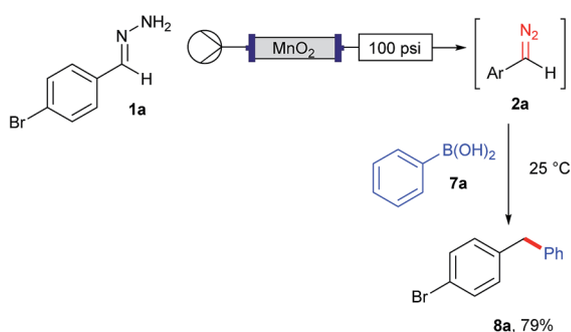
Diazo compounds are commonly proposed as intermediates for several useful transformations, in particular carbon-carbon sp^2 - sp^3 coupling between tosylhydrazone and boronic acids.¹³ High temperature and an inorganic base are typically required as the generation of the diazo compound is reported to be the limiting step of these reactions. We speculated that with our new protocol the chemical reactivity of diazo compounds could be studied separately without interference from the generation step. Pleasingly, after its generation, the stream of the diazo compound **2a** reacted quickly at room temperature with boronic acid **7a** in the first attempt, giving the coupled product **8a** in very good yield (Scheme 4). This extremely mild and fast reaction conditions demonstrated the high chemical reactivity of diazo compounds and highlighted the advantages of the flow procedure for these reactive intermediates where the equivalent batch mode reaction was found to be unsuccessful.

Different coupling combinations were investigated using the flow procedure for the generation of clean diazo compounds (Table 2). With much milder reaction conditions compared to the original Barluenga coupling, the electronic properties exerted a strong influence on the reaction time. For electron rich or neutral boronic acids the transformation was complete within 1 hour (compounds **8a-c**). By contrast, electron poor aryl boronic acids required longer reaction time or harsher reaction conditions (**8d-e**). Similar trends were observed with other diazo compounds. Electron poor (more stabilised) diazo compounds required, in general, longer reaction times when reacted with the same boronic acid (**8j** vs. **8b**). Interestingly, vinyl diazo compounds are also compatible with this protocol giving the desired products in good yields (**8n-q**). Notably, in these cases a mixture of products was obtained resulting from a



Table 1 Titration of generated diazo compound to the corresponding esters^a

^a Isolated yield over 3 steps based on aldehyde precursor. ^b Combined yield (r.s. = 1 : 1).

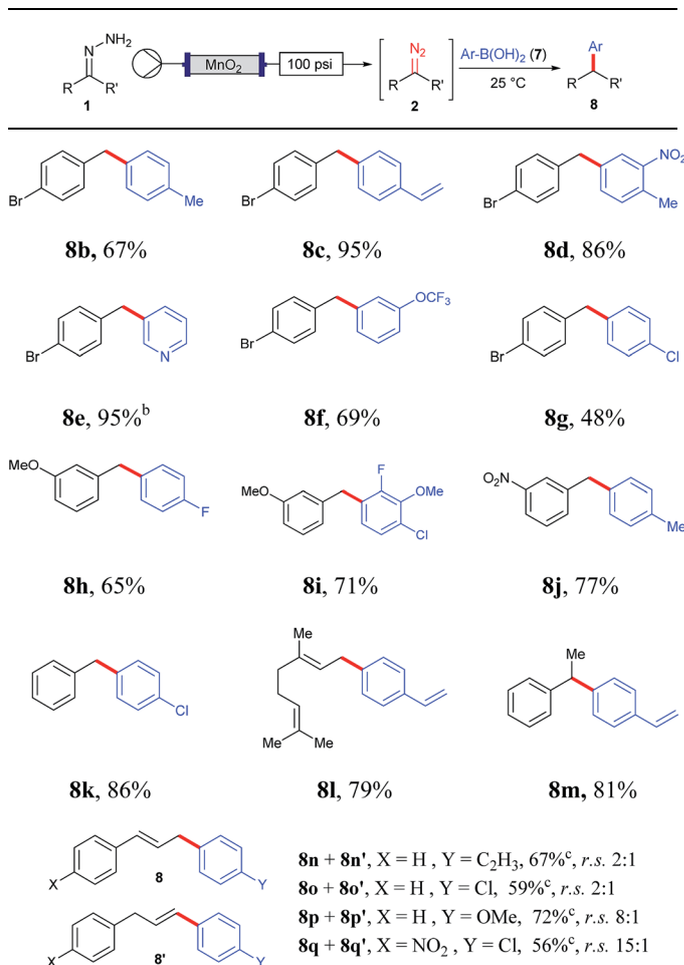
**Scheme 4** Flow generation, translocation and reaction of **2a** with **7a**.

1,3-borotropic rearrangement and the regioselectivity of the products mixtures was strongly influenced by the electronic properties of the two aromatic rings, ranging from 2 : 1 (**8n**) to 15 : 1 (**8q**). Different functionalities are tolerated such as halogens, nitro groups and ethers. Heterocycles such as pyridine (**8e**) are also compatible with this protocol. The highly reactive geranyl diazo gave the geranylated product (**8l**) in good yield, based on the titration of the corresponding diazo compound. This process is remarkable as it provides a complementary

method for geranylation, where the normal Friedel–Crafts alkylation fails. Additionally, we were able to access sp^2 – sp^3 coupling transformations for aryl–alkyl diazomethane derivatives in quite good yield (**8m**).

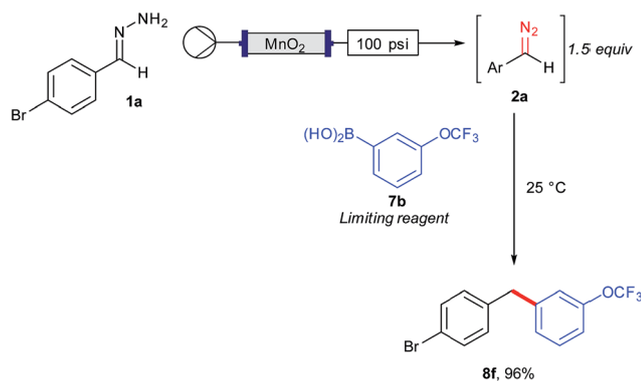
Since the boronic acids are often the more functionalised of the two starting materials, the flow-generated diazo partner was used in excess to fully convert the boronic acid **7b** to the desired coupling product (Scheme 5). Pleasingly, the diaryl compound **8f** was obtained in 96% isolated yield using only a slight excess of diazo compound (1.5 equiv.). In this case, the excess diazo was subsequently quenched *via* an acidic aqueous work-up protocol. The main by-products detected were both the corresponding azine and alcohol.

The strong dependence of the reaction kinetics on the electronic properties of both starting materials reveals some insight about the reaction mechanism. It is generally assumed that after nucleophilic attack of the diazo compound to the boronic acid, a first ate complex **9** is formed. This intermediate rapidly undergoes a 1,2-carbon migration forming the boronic species

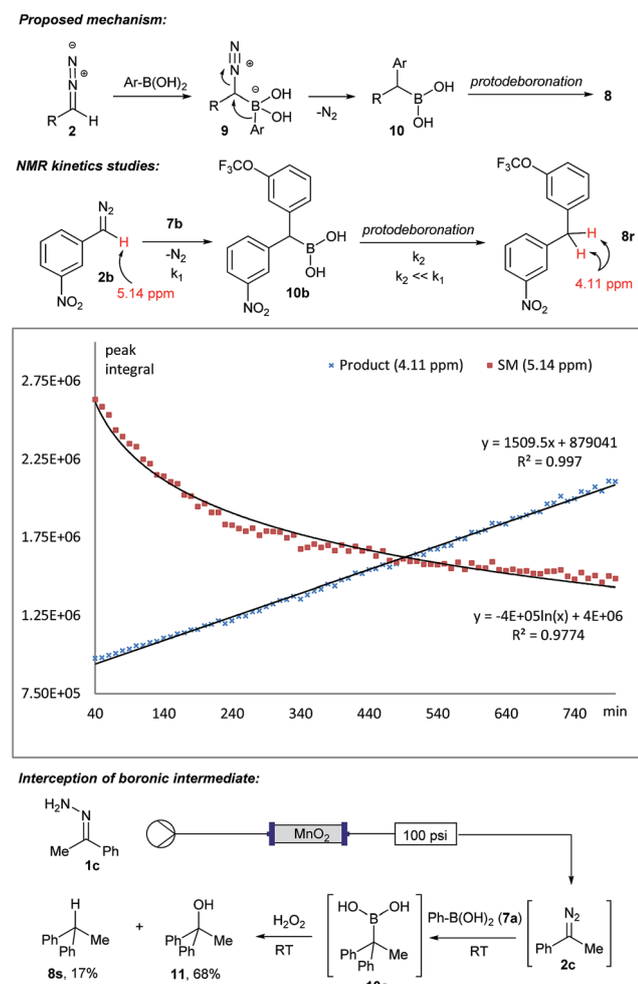
Table 2 Coupling products prepared by reaction of diazo compounds with boronic acids^a

^a Isolated yield based on titration of diazo compound with acetic acid. ^b Microwave 100 °C in 15 min. ^c Combined yields.



Scheme 5 Boronic acid **7b** used as limiting reagent.

10 which isomerises to the most stable isomer prior to a protodeboronation to give the coupling product **8** (Scheme 6). Although this mechanism is assumed to describe the actual chemical combinations of the intermediates involved, no report has either identified the kinetics of this kind of transformation or the nature of the species involved. The reasons behind this

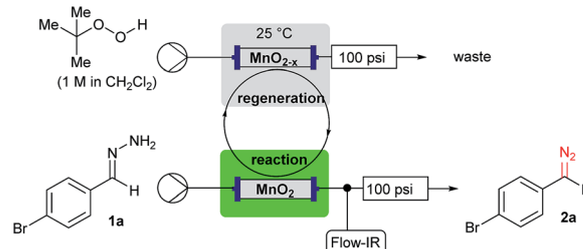
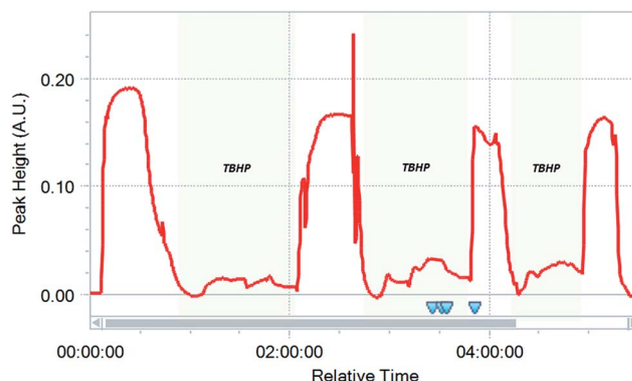


Scheme 6 Mechanistic studies of coupling reaction between diazo compound and boronic acid.

are linked to the harsh conditions employed to generate the reactive intermediate.¹³ We started by investigating the kinetics involved in the sequence of events. Initial NMR studies focused on a model reaction between diazo compound **2b** and boronic acid **7b** and the analysis showed a two-step mechanism (Scheme 6), where the first step happens at a much faster rate than the second one. Using the data collected over a large series of dynamic experiments, we were able to define that the reaction is first order with respect to the diazo compound.

Remarkably, using our mild flow protocol we were also able for the first time to intercept the secondary boronic intermediate **10**, proving the presence of this intermediate prior to protodeboronation. Indeed, we envisaged that the generation of a relatively stable boronic intermediate **10** would reduce the rate of protodeboronation and this new reactive intermediate could be intercepted *via* an alternative reaction pathway. This idea was realised with the use of acetophenone hydrazone (**1b**) as starting material, which was passed through the oxidant column, packed with activated MnO₂, to form the corresponding diazo compound (as detected by in-line IR); the transient diazo species was translocated to react with boronic acid **7a** and then further reacted with a solution of hydrogen peroxide. The analysis of the material revealed the prevalence of alcohol **11** (68%) within the mixture, corroborating the presence of a boronic species **10c** as an intermediate.

To increase the efficiency of the new protocol, the regeneration and re-use of MnO₂ column was briefly investigated. The spent manganese oxide could be oxidised by passage of a

Scheme 7 Flow regeneration of MnO₂ reagent using TBHP.Fig. 1 IR data for the four regeneration cycles of oxidation for a single MnO₂ load (0.86 g).

solution of *tert*-butyl hydroperoxide (TBHP, 1 M in CH₂Cl₂) (Scheme 7). Interestingly, the regenerated column produced diazo compound without the need for a pre-conditioning phase. The heterogeneous oxidant was recycled over three different cycles with just a slight decrease in reactivity after each recycle as detected by in-line IR (Fig. 1).

Moreover, in a preliminary attempt with tube-in-tube technology,¹⁶ oxygen gas also proved to be a useful re-oxidant for a low cost regeneration of MnO₂ but with lower efficiency.¹⁷

Conclusion

In summary, we have reported a practical flow generation of *unstable* diazo compounds as reactive intermediate using the cheap, recyclable and non-toxic oxidant, MnO₂. The diazo flow stream can be accurately titrated or followed by in-line IR. Using our protocol, we were able to deliver sp²-sp³ coupling reactions of the freshly generated diazo species with boronic acids, under extremely mild and simple reaction conditions providing alkylated products in good to excellent yield. The mechanistic studies demonstrated for the first time the presence of a specific boronic intermediate. These results demonstrate that flow generation/translocation/reaction of transient species opens up opportunities for the exploitation and study of new chemical reactivities. Further studies in this area are underway in our laboratories.

Acknowledgements

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