Isotopic labelling studies for a gold-catalysed skeletal rearrangement of alkynyl aziridines

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Abstract
Isotopic labelling studies were performed to probe a proposed 1,2-aryl shift in the gold-catalysed cycloisomerisation of alkynyl aziridines into 2,4-disubstituted pyrroles. Two isotopomers of the expected skeletal rearrangement product were identified using $^{13}$C-labelling and led to a revised mechanism featuring two distinct skeletal rearrangements. The mechanistic proposal has been rationalised against the reaction of a range of $^{13}$C- and deuterium-labelled substrates.

Introduction
Gold-catalysed cycloisomerisation reactions have emerged as powerful methods to construct a diverse array of hetero- and carbocyclic motifs under generally mild reaction conditions using straightforward preparative procedures [1-9]. Many such processes also incorporate a rearrangement of the hydrocarbon skeleton present in the starting material, which can lead to unexpected and synthetically interesting outcomes [10,11].

One of the most common fundamental steps in these skeletal rearrangements involves C–C bond fission through 1,2-migration. This step is triggered by the generation of carbocationic character, which is generally stabilised in some form, either by an adjacent gold atom or through extended delocalisation. While 1,2-alkyl migrations are well-established in gold-catalysed heterocyclic synthesis [12,13], 1,2-aryl shifts are much less common [14-17].

In relation to these processes, we recently reported the gold-catalysed reactions of aryl substituted alkynyl aziridines, such as 1. Careful choice of reaction conditions resulted in selective cycloisomerisation to produce either the 2,5-disubstituted pyrrole 2 or the skeletally rearranged 2,4-disubstituted pyrrole 3 (Scheme 1) [18].

The nature of both the solvent, and the counter ion to the cationic gold catalyst, proved crucial to the reaction outcome:
Scheme 1: Gold-catalysed cycloisomerisations of aryl–alkynyl aziridine to pyrroles.

Scheme 2: Working mechanism to rationalise the formation of two regiosomeric pyrroles in the gold catalysed cycloisomerisation of alkynyl aziridines.

Path I follows proton elimination (B→C) and subsequent protodemetallation to give 2,5-disubstituted pyrrole D, in keeping with a previously reported furan synthesis [24,25]. The 2,4-disubstituted skeletal rearrangement product G was rationalised through Path II, where a 1,2-aryl shift (B→E) precedes proton elimination (E→F) and protodemetallation. The observed solvent and counter ion effects could be rationalised by considering that the 1,2-aryl shift at B is disfavoured relative to proton elimination as its cationic character is diminished, either by solvent stabilisation or the formation of a closer contact ion pair, and/or by more basic solvents/counter ions assisting proton elimination [18,26,27].

However, for the proposed skeletal rearrangement mechanism to be valid, gold bound carbocation B must be prone to undergo a relatively rare 1,2-aryl shift in preference to either proton elimination or a 1,2-hydride shift. We therefore sought to validate this proposal experimentally by isotopic labelling.

Results and Discussion
A deuterium labelled precursor
Alkynyl aziridine 4 was selected as the initial probe for the mechanism which requires fission of three bonds: The propargylic C–N bond in ring expansion; the aryl–aziridinyl C–C bond in the 1,2 shift and the propargylic C–H bond for aromatisation (Scheme 3). The positioning of a deuterium atom at the benzylic carbon in 4 enables the carbons of the aziridine ring to be distinguished and labels the carbon from which the aryl shift would occur. The tolyl and butyl groups were selected as substituents in place of other aromatic units to aid analysis by simplifying the key aromatic region in the resulting NMR spectra.
Deuterated aziridine 4 was prepared in four steps from ester 5 (Scheme 4). Reduction using lithium aluminium deuteride led to the insertion of the label at the benzylic position. Oxidation to the aldehyde followed by condensation with tosylamide afforded the deuterated imine 6. Aziridination using the sulphonium ylide generated in situ from 7 proceeded smoothly to afford the desired cycloisomerisation precursor 4 [28]. The aziridines were formed as a mixture of diastereomers, with the cis diastereomer predominating, and were employed as such in the cycloisomerisation reactions.

Deuterated aziridine 4 was reacted under the standard conditions for skeletal rearrangement (5 mol % of Ph3PAuOTf generated in situ in CH2Cl2), to give a mixture of 2,4- and 2,5-pyrrole isomers 8c and 9 alongside the main product, 3-deutero-2,4-disubstituted pyrrole 8b (Scheme 5). Isomer 8a, which was predicted as the major product under the proposed mechanism (Scheme 2), could not be indentified from the reaction mixture. Though H/D exchange in pyrroles has been noted in gold-catalysed reactions [13], control reactions had shown that no H/D exchange occurred in this specific system [18,19]. There was also no interconversion between 2,4- and 2,5-isomers under the same reaction conditions [18]. Surprisingly, despite several repetitions and the use of different batches of starting materials, the reaction of 4 was always less clean than its non-deuterated analogue, even when the substrates were tested side-by-side under identical conditions.

Despite this unexplained difference, the proposed mechanism does not account for the formation of 2,4-disubstituted pyrrole without deuterium incorporation at the C-5 position. More information on whether a 1,2-aryl shift occurs was therefore sought using a non-labile 13C label at the aziridine carbon.

13C labelling studies
13C-Enriched benzoic acid was converted into imine 10 and subsequently coupled with 7 to give the aziridine 11, in four steps, as a 1:5 13C:12C mixture (Scheme 6). Two further 13C-
enriched alkynyl aziridines, 14 and 15, with different substituents on the alkyne, were later prepared using the sulfoxonium salts 12 and 13.

On subjecting 11 to the conditions for skeletal rearrangement (Scheme 7), three enhanced resonances were seen in the $^{13}$C NMR spectra of the pyrrole products (Supporting Information File 1). One of the enhanced resonances was associated with the minor 2,5-disubstituted isomer 16 showing $^{13}$C enrichment at the expected C-2 position (Scheme 2, Path I). This result was confirmed by carrying out the control cycloisomerisation of 11 under the conditions known to give solely 2,5-disubstituted products (Scheme 8) [19]. All other resonances were associated with the 2,4-disubstituted pyrrole 17 and the remaining enhancements were therefore assigned to the unexpected formation of two separate isotopomers of 17. $^{13}$C-enrichment at C-5 (117.7 ppm) corresponds to the isotopomer 17a anticipated from the proposed 1,2-aryl shift mechanism. Isotopomer 17b was identified by enrichment at the quaternary C-4 position (127.0 ppm). HMBC experiments confirmed this assignment showing clear $^{3}$J coupling between the C-4 $^{13}$C-enriched quaternary centre and the protons of the phenyl ring.

Cycloisomerisation of the alkynyl aziridine 14, which bears a phenyl substituent on the alkyne, resulted in a different outcome under the same reaction conditions: Only a single isotopomer of the 2,4-disubstituted pyrrole 19 was formed with $^{13}$C-enrichment at the C-5 position, alongside the 2,5-disubstituted isomer 18 (Scheme 9).

The observed inconsistency between the reactions of 11 and 14 was investigated using alkyl substituted alkynyl aziridine 15. Cycloisomerisation of 15 once again afforded a mixture of isotopomers of the 2,4-disubstituted pyrrole 21a/b alongside 2,5-disubstituted pyrrole 20 (Scheme 10).

The formation of the isotopomers 17a, 19 and 21a with $^{13}$C-enrichment at C-5 confirms C–C bond fission between the aryl group and the aziridine during the reaction. This outcome is consistent with an operative 1,2-aryl shift in the skeletal
rearrangement of alkynyl aziridines to form 2,4-disubstituted pyroles (Scheme 2, Path II).

However, the initial use of the alkyl substituted aziridine 11 rather than the aryl-substituted aziridine 14 was fortuitous. A second accessible pathway must exist to account for the formation of the 2,4-disubstituted pyrrole isotopomers 17b, and 21b by an alternate skeletal rearrangement.

The mechanistic proposal for the skeletal rearrangement was therefore reappraised to explain these results (Scheme 11). Alongside the ring expansion of metal coordinated alkynyl aziridines to the 5-membered cyclic cation B, which leads to isotopomer G (Path II) and 2,5-disubstituted pyrroles D (Path I), a competing regioisomeric ring-expansion could afford 4-membered ring intermediate I. Evolution of I by a ring-expanding 1,2-shift of the metal stabilised vinyl unit onto the benzylic cation leads to a new 5-membered cyclic intermediate J [29]. Deprotonation and protodemetalation of J results in a 2,4-disubstituted isomer G’ where the $^{13}$C-enriched carbon maintains its connectivity to the aryl unit, and C–C bond fission occurs between the aziridine and the alkyne fragment.

The observed effects of the alkyne substituent can be rationalised if the pathway is dependent on the relative stabilisation of the developing positive charge at either carbon of the aziridine by the alkynyl aziridine intermediate H (Scheme 11). The formation of either benzylic cation I (Path III) or the enamide stabilised cation B (Path II), under a cationic ring-expansion manifold, is apparently closely balanced: Products from both pathways are observed for alkyl substituted systems such as 11 and 15. However, when the alkyne substituent is a phenyl group Path II is favoured over Path III due to the additional stabilisation of B by delocalisation through an extended π-system.
The initial report showed how the ratio of 2,5- to 2,4-disubstituted pyrroles varied with the electronic properties of the aromatic unit substituent directly attached to the aziridine [18]. While the trend was consistent with that expected for a 1,2-aryl shift, the more complex mechanism also accounts for the observed outcomes. An increasingly electron-rich aryl group will favour both formation of benzylic cation I and 1,2-aryl migration from B, leading to the same products from each route. Electron-deficient aromatics would have a destabilising effect on the formation of I and also be less prone to undergo 1,2-aryl shift from B thus favouring Path I and formation of the 2,5-disubstituted pyrroles.

Two further $^{13}$C labelled compounds were prepared to test this rationalisation by predictably affecting the relative stability of just one of the key intermediates: An electron-deficient aromatic substituent to the alkyne (R = $p$-C$_6$H$_4$CF$_3$) should disfavour Path II by destabilising B over I, whereas an electron-rich aromatic substituent on the alkyne (R = $p$-C$_6$H$_4$OMe) should favour Path II by further stabilising B.

The substrates were prepared by Sonogashira reaction between propargyl alcohol and the appropriately substituted iodobenzene substrate 22, 23 (Scheme 12). Functional group interconversion of the alcohols by bromination and substitution results in the sulfonium salts 28, 29 which were reacted with $^{13}$C labelled imine 10 to yield the desired alkynyl aziridines 30 and 31.

The anticipated outcomes of the catalysis were borne out under the standard conditions (Scheme 13). Two isotopomers of $^{13}$C enriched 2,4-disubstituted pyrrole 32 were produced from the
In keeping with the previous observation that strongly electron-rich aromatic groups were poor substituents for this cycloisomerisation [19], only a low yield of the pyrrole products was obtained on reaction of 31. However, this was sufficient to determine that Path II predominated over Path III, with no significant isotopeomer identified from intermediate I. Isomer 34 was identified alongside a significant quantity of the 2,5-disubstituted pyrrole 33. While solvents and/or counter ions that stabilise positive charge favour the formation of the 2,5-disubstituted pyroles, likewise, reducing the localised cationic charge in the 5-ring intermediate B by conjugation appears to have the same effect.

Re-evaluation of the cycloisomerisation of the deuterium-labelled alkynyl aziridine 4 (Scheme 4) shows the results to be consistent with both the modified mechanistic scenario and the trends shown in the 13C-labelling study. Relative to all the other substrates tested, excepting perhaps 30, substrate 4 would be the most likely to favour Path III over Path II (Scheme 11). Whilst the butyl substituent does not lend any additional stabilisation to intermediate B, the tolyl group stabilises the benzyl cation I better than a phenyl group. Significant deuterium incorporation at C-5 in the product would therefore not be expected. Following Path III, the deuterium labelled carbon ends-up at C-4 (Scheme 14, I-4→J-4). Aromatisation results in C–D bond cleavage with deuterodemetallation of K-4 affording the observed major product 8b (Scheme 5). The intermolecular deuteron transfer allows for H/D exchange in the reaction media and hence the partial incorporation of a proton at C-3.

Conclusion

There are two operative skeletal rearrangement pathways for the formation of 2,4-disubstituted pyrroles by gold-catalysed cycloisomerisation of aryl-substituted N-tosyl alkynyl aziridines. The two competing pathways coincidentally give rise to the same isomer but with different absolute connectivities, as determined by the formation of two 13C isotopomeric products.

A modified mechanistic scheme accounts for all of the results of the labelling studies. On activation by a cationic gold source and in the absence of coordinating solvents and/or counter ions, regioisomeric ring expansion of the alkynyl aziridine can lead to two different cationic cyclic intermediates, featuring either a 4-membered or a 5-membered ring. The regioselectivity, and hence the isotopomeric reaction selectivity, is controlled by the relative ability of the substituents to stabilise the respective cations in favour of a particular pathway. The observed skeletal rearrangements are consistent with either a 1,2-aryl shift or a 1,2-(metal-stabilised)-vinyl shift in the 5- or 4-membered cyclic intermediates, respectively. These studies highlight some of the complexities associated with gold-catalysed cycloisomerisation reactions and the continued relevance of labelling studies in this field.

Supporting Information

Supporting Information contains full experimental details for the preparation of the cyclisation precursors and their subsequent reactions. NMR spectra for the labelling studies are provided.

Supporting Information File 1

Full experimental details.

[http://www.beilstein-journals.org/bjoc/content/supplementary/1860-5397-7-96-S1.pdf]

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