Lewis acid–base 1,2-addition reactions: synthesis of pyrylium borates from en-ynoate precursors†

Lewis C. Wilkins, a Hugh B. Hamilton, a Benson M. Kariuki, a A. Stephen K. Hashmi, b,c Max M. Hansmannb and Rebecca L. Melen*a

Treatment of methyl (Z)-2-alken-4-ynoates with the strong Lewis acid tris(pentafluorophenyl) borane, B(C6F5)3, yield substituted zwitterionic pyrylium borate species via an intramolecular 6-endo-dig cyclisation reaction.

Some of the first examples of pyrylium salts were reported over 100 years ago through the reaction between acetophenone and sulphuric acid to yield the 2,4,6-triphenyl pyrylium cation.1 While the significance of this compound was not fully appreciated at the time, it has since spawned intense interest in various areas such as a basis for photo-sensitisers,2 redox mediators3 and photo-induced free radical polymerisations.4 Due to these interesting physical properties, pyrylium salts play a fundamental role in materials chemistry.5 Alongside this prominence in materials chemistry, pyrylium compounds are extremely important intermediates in synthetic chemistry and are vital in the generation of complex cyclic compounds with various functionality such as extended macrocyclic arenes6 inter alia.7 In particular, these electron deficient aromatic compounds have significance in medicinal and pharmaceutical chemistry for the synthesis of many biologically active drugs and bio-imaging compounds.8

Much of the reason why these pyrylium systems are useful in such applications arise from the 6-membered ring system similar to phenyl and pyridyl derivatives, albeit with reduced aromaticity compared to benzene as a result of the incorporation of the more electronegative oxygen atom.9 The perturbation to the aromaticity caused by the incorporation of an oxyacetylene renders these pyrylium systems susceptible to attack by nucleophiles at the 2-, 4- or 6-positions. This reactivity is of necessity in the formation of aryl-substituted phosphinines,10 pyridinium salts11 and substituted furans.12

The formation of pyrylium salts as intermediates using π-Lewis acidic transition metals, such as gold, has been shown to be a reliable synthetic methodology through π-activation of the alkyne unit of enynone and enynal reagents.13 This proceeds through a trans-oxy-auration step across the alkyne moiety via a 6-endo-dig cyclisation mechanism. This intermediate can then undergo many transformations such as acting as the diene component in [4 + 2] intermolecular cycloaddition reactions with alkenes to yield acylphosphine derivatives.14 Our previous studies have shown that the reactions of B(C6F5)3 with organic substrates containing π-bonds together with nucleophilic ketones,15 esters,16 or amides17 result in a range of carboboration and 5-exo-dig 1,2-addition reactions.18 The work discussed here extends our research to en-ynoate substrates in 6-endo-dig cyclisation reactions to generate pyrylium borate species.

The reaction between the strong Lewis acid B(C6F5)3 and functionalised methyl (Z)-2-alken-4-ynoates (1a and 1b) proceeded rapidly at room temperature with an immediate colour change to a dark orange-red colour associated with the formation of the 2-methoxy pyrylium borate products (2a and 2b, Scheme 1). In situ multinuclear NMR studies showed almost

Scheme 1 Proposed mechanistic pathway of the reaction between B(C6F5)3 and methyl en-ynoates.
quantitative conversion in as little as 1 h, although isolated yields were more modest at 65% [2a] and 64% [2b].

The $^1$H NMR spectra were diagnostic revealing a prominent resonance of the vinylic proton $\beta$ to the carbonyl at C(4) at $\delta = 8.65$ and 8.47 ppm for 2a and 2b respectively (cf. $\delta = 6.37$ ppm (1a) and 6.15 ppm (1b)). In addition, the bulky nature of the three C$_8$F$_5$ groups at boron would appear to restrict motion of the adjacent C(6) substituent, leading to two independent proton environments of the CH$_2$ group in the nBu functionality of 2b reflected in the observation of two distinct broad singlets in a 1 : 1 ratio at $\delta = 1.43$ and 0.64 ppm. Similarly, the constrained geometry is reflected in separate chemical environments for the F atoms of the C$_8$F$_5$ groups, evident in the $^{19}$F NMR spectra of 2b (Fig. 1, bottom) which reveals a total of fifteen resonances for the ortho ($\delta = -128.2$ to $-135.0$ ppm (6F)), para ($\delta = -158.9$ to $-159.9$ ppm (3F)) and meta ($\delta = -162.8$ to $-165.3$ ppm (6F)) fluorine atoms. Heating an NMR sample of 2b in $d_6$-toluene to 80 °C showed no coalescence of the proton environments of the methylene group attached to C(6) in the $^1$H NMR spectrum. In addition, the $^{19}$F NMR showed no discernible coalescence of the fluorine environments on the C$_8$F$_5$ rings suggesting a considerable energy barrier to rotation of the pyrylium cation with respect to the B(C$_6$F$_5$)$_3$ group. The $^{19}$F NMR of 2a (Fig. 1, top) exhibits similar behaviour, although the spectra in this case are somewhat simpler with just eight peaks for the ortho ($\delta = -128.6$ (1F), $-130.3$ (4F) and $-131.1$ ppm (1F)), para ($\delta = -159.2$ (1F) and $-160.0$ ppm (2F)) and meta ($\delta = -163.5$ (1F), $-164.7$ (1F) and $-165.4$ ppm (4F)) fluorine atoms. $^{11}$B NMR spectra lead to sharp resonances at $\delta = -14.4$ and $-14.5$ ppm for 2a and 2b which has become synonymous with these four coordinate vinyl borate complexes.$^{17,18}$

Examination of the solid-state single crystal structures of 2 unambiguously confirmed the product of the cyclisation process as the aforementioned pyrylium-borate species (Fig. 2). 2a and 2b both crystallised in the monoclinic $P2_1/n$ space group with two and one molecule in the asymmetric unit respectively. The phenyl substituent attached to the ring in 2a rotates out of the plane of the pyrylium ring by 66.1(2)$^\circ$ and 58.4(2)$^\circ$, this loss of planarity also prevents an extended aromatic network with the pyrylium fragment. This is in contrast to that observed previously for dioxolium compounds which contain a cationic 5-membered [C$_2$O$_2$C]$^+$ ring.$^{16a}$ In these cases, when the C' carbon is substituted by an aromatic ring then the two rings lie in the same plane offering the possibility for stabilisation of the positive charge through delocalisation.$^{16a}$ Regarding compound 2a, the rotation of the phenyl group out of the plane of the pyrylium ring arises chiefly through reduction of steric congestion. This can be clearly observed in the space-filling models (Fig. 3) of both 2a and 2b. Additionally the reaction of B(C$_6$F$_5$)$_3$ with 1c gave inconclusive results presumably due to the steric congestion that would
arise from the neighbouring borate and tBu groups. In 2a the rotation of the C₆H₅ ring out of the plane of the pyrylium ring allows it to interact with one of the C₆F₃ rings through a displaced π-stacking interaction with a centroid-centroid distance of 3.6421(10) Å leading to an essentially eclipsed geometry in which the C(5)C(6)Ph plane deviates out of the C(5)BC₆F₅ plane by 0.11(14)°. 2b however, adopts a more conventional staggered geometry whereby the C(6)Ph-butyl bond is out of C(5)BC₆F₅ plane by 49.9(2)° which is consistent with the higher symmetry observed for 2a with respect to 2b as detected via NMR spectroscopy. The structures of both 2a and 2b show similar metric parameters for the pyrylium ring. In both cases the C₅O heterocycle is essentially planar verified by similar metric parameters for the pyrylium ring. In both cases the C–O bond lengths (1.372(3)–1.377Å) are intermediate between a formal double and single bond. The C–C bond lengths (Table 1) (1.384 Å) are more typical for C–C single bond (1.33 Å) and are more intermediate between those for a typical C–C double bond (1.316 Å) and are more intermediate for C–C bonds found in more delocalised systems (1.38 Å in benzene).19

The C–O bond lengths associated with the O(1)C(2)O(7) fragment are almost identical within error (1.303(2)–1.316(2) Å) and are intermediate between a formal double and single carbon-oxygen bond (ca. 1.23 and 1.42 Å respectively).19 The nature of the bonding in 2 has been investigated by way of DFT calculations carried out at the B3LYP/6-311G* level (see ESI†). The calculated structure was in good agreement with that observed experimentally with similar geometric parameters for the cationic C₅O ring (Table 1) with C–C bond lengths intermediate between single and double bonds characteristic of a delocalised structure. An NBO analysis clearly reveals strong delocalisation around the carbon-containing part of the cationic pyrylium ring but a more localised polar structure in the vicinity of the oxygen atoms. The Wiberg bond

### Table 1 Experimental and calculated bond lengths for 2a and 2b

<table>
<thead>
<tr>
<th>Bond</th>
<th>2a</th>
<th>2b</th>
</tr>
</thead>
<tbody>
<tr>
<td>C–C</td>
<td>1.371(3)</td>
<td>1.388(2)</td>
</tr>
<tr>
<td>C–C</td>
<td>1.362(3)</td>
<td>1.360(2)</td>
</tr>
<tr>
<td>C–C</td>
<td>1.430(2)</td>
<td>1.435(2)</td>
</tr>
<tr>
<td>C–C</td>
<td>1.361(2)</td>
<td>1.365(2)</td>
</tr>
<tr>
<td>C–O</td>
<td>1.316(2)</td>
<td>1.304(2)</td>
</tr>
<tr>
<td>C–O</td>
<td>1.390(2)</td>
<td>1.397(2)</td>
</tr>
<tr>
<td>C–O</td>
<td>1.306(2)</td>
<td>1.303(2)</td>
</tr>
<tr>
<td>C–B</td>
<td>1.665(2)</td>
<td>1.650(2)</td>
</tr>
</tbody>
</table>

Fig. 4 (left) Wiberg bond indices for 2a and (right) NBO charges for the core of the pyrylium cation 2a and 2b.

Fig. 5 Aromaticity of compounds 2 relative to benzene using nucleus independent chemical shift calculations.
in these systems we can isolate the vinyl borate species as a stable product in the reaction, this is in contrast to that seen in transition metal catalysed systems as the analogous vinyl gold intermediates have been seldom observed or isolated until recent years as the deauration step is typically rapid.23 Further work will probe the breadth of substrate scope and subsequent reactivity of the pyrylium borate products, all of which are ongoing projects in our group. The hydrolysis and/or proto-deboronation of these products will be investigated in the synthesis of pyrones. Indeed, isocoumarins and α-pyrones are key structural units in natural products and many biologically active pharmaceuticals e.g. in HIV protease inhibitors.24 Such systems have been synthesised by similar electrophilic cyclisation reactions using other main group Lewis acids (e.g. using iodine or selenium electrophiles).25 Unlike the cyclisations described above, and those performed by transition metals such as gold, the ability for this transformation to trap the pyrylium species en route to the pyrone product also permits these products to be used for further transformations including reactions with nucleophiles, cycloaddition reactions or cross-coupling of the boronate with organo-halides. Our ongoing studies in this area will permit us to develop these types of reagents in the synthesis of useful precursors in a variety of areas of synthetic, biological and materials chemistry.

Acknowledgements

H. B. H. would like to thank the EPSRC-funded Bath/ Bristol/Cardiff Catalysis Centre for Doctoral Training (EP/L016443/1). M. M. H. is grateful to the Fonds der Chemischen Industrie for a Chemiefonds scholarship and the Studienstiftung des deutschen Volkes. We thank W. R. Grace for the kind donation of B(C₆F₅)₃. We also thank Dr James Platts for his assistance in NICS calculations as well as Dr Joel Loveridge for his support with NMR spectroscopy.

Notes and references