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Selective access to dihydrophenanthridines and phenanthridinones *via* cyclisation of aryl amines onto *N*-tethered arynes†

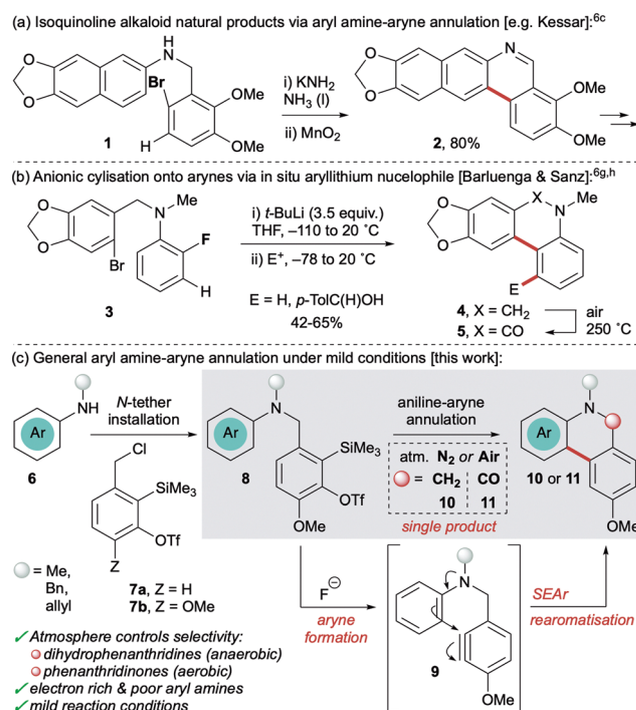
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5,6-Dihydrophenanthridines are prepared from aryl amines *via* intramolecular addition to *N*-tethered arynes under mild conditions. A new *o*-silylaryl triflate precursor was developed to increase reactivity and enable electron-rich and electron-poor aryl amines to undergo cyclisation. A complete switch in product selectivity occurs when the reaction is conducted in air, affording the corresponding phenanthridin-6(5*H*)-one as the sole product under otherwise identical reaction conditions.

Phenanthridines and phenanthridinones are privileged heterocyclic scaffolds found in a range of natural products and therapeutically active compounds.¹ They possess broad biological properties, including anticancer,^{2a} antitumour,^{2b} antiviral,^{2c} antimicrobial,^{2d} antifungal^{2e} and antimalarial^{2f} activity. In addition, a high charge mobility renders these frameworks versatile building blocks for functional materials.³ As a result, significant synthetic effort has been devoted to the synthesis of phenanthridines and their derivatives.⁴ This includes the Bischler-Napieralski reaction,^{5a} photochemical,^{5b} radical^{5c} and microwave-assisted^{5d} cyclisations, transition metal free C–H arylation,^{5e} aza-Wittig,^{5f} anionic ring-closure,^{5g} hypervalent iodine^{5h} and metal catalysed approaches.⁵ⁱ

Arynes have also been utilised in the synthesis of phenanthridine derivatives and related alkaloid natural products.^{6,7} These versatile reactive intermediates afford valuable benzenoid and heterocyclic frameworks⁸ and have experienced a recent resurgence in interest due to the advent of aryne precursors that act under mild conditions, namely the *o*-trimethylsilylaryl triflates (*o*SATs)⁹ and the hexadehydro-Diels–Alder reaction of polyalkynes.¹⁰ Cyclisation onto a pendant aryne has proven a valuable method to furnish phenanthridine derivatives, most notably in the synthesis of natural products.^{6a–e,g,h} However, these approaches all generate arynes using organometallic reagents or strong bases at low temperatures

which restricts the functional group tolerance. Methods that operate *via* electrophilic aromatic substitution (SEAr) have also been limited to electron-rich aryl nucleophiles such as **1** (Scheme 1a),^{6a–e} which although common in the natural products, precludes preparation of more diverse substrate analogues. Alternatively, Barluenga and Sanz utilised anionic cyclisation onto pendant arynes (accessed from **3**) to yield phenanthridine derivatives **4** and **5** by exploiting stronger *in situ*-formed aryllithium nucleophiles (Scheme 1b).^{6g,h} However, this strategy also required organolithium reagents (in excess) to form both the aryne and the



Scheme 1 Cyclisation onto tethered arynes for the synthesis of phenanthridines and derivatives.

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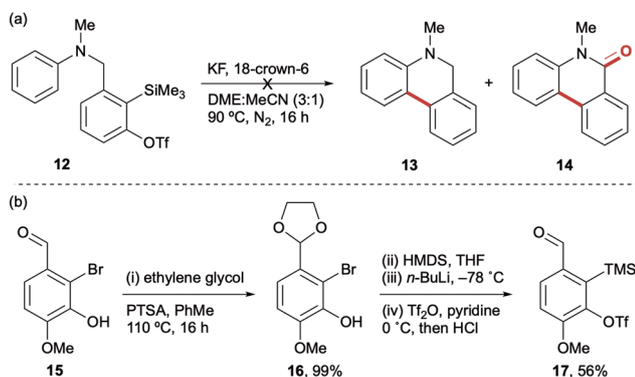
nucleophile, plus pre-functionalisation (*o*-bromination) of the pronucleophilic arene.

Given our interests in the chemistry of arynes,¹¹ we sought to develop a cyclisation approach to phenanthridine derivatives that exploited the benzylic *o*SAT precursor **7a** previously used in our group (Scheme 1c).^{11b} Inspired by the strategy of both Kessar and Stermitz in their syntheses of alkaloid natural products,^{6c,d} it was proposed that aryl amines bearing an *N*-tethered aryne precursor (**8**) would undergo SEAr, upon generation of the aryne **9**, to afford 5,6-dihydrophenanthridine derivatives **10**. Given the propensity for such frameworks to undergo oxidation there would also be potential to access the corresponding phenanthridinones **11**. Importantly, by using the *o*SAT precursor it removes the harsher reaction conditions of previous reports and should facilitate a wider range of aryl amine nucleophiles **6**, especially those containing useful functional handles. Furthermore, tethering the aryne at nitrogen enables more facile synthesis of the starting materials, compared to linking *via* C–C bonds,^{6a,b} thereby expediting the preparation of structural analogues.

To test this cyclisation hypothesis, benzylic *o*SAT precursor **7a** was tethered to *N*-methyl aniline to afford precursor **12**, which was then subjected to our standard aryne-forming conditions previously used with this tether (Scheme 2a).^{11b} Unfortunately, this afforded a mixture of compounds and no clear evidence of cyclisation to dihydrophenanthridine **13** nor phenanthridinone **14**. Intriguingly, a significant by-product was observed in the ¹H NMR spectrum of the crude reaction mixture that suggested intermolecular attack at the aryne was taking precedence over the desired intramolecular process (although we were unable to isolate a clean sample for characterisation).¹² With a goal to promote the cyclisation and suppress undesired intermolecular reactivity, a modified

aryne precursor **17** was developed (Scheme 2b). It was rationalised that a *p*-methoxy substituent in **17** should increase reactivity by inductively polarising the aryne towards nucleophilic addition *ortho* to the tether – as supported by Garg and Houk's aryne distortion model¹³ – in addition to providing increased stabilisation of the resulting aryl anion.^{8a} To this end, the novel *p*-methoxybenzyl aryne precursor **17** was prepared in four steps from commercially available 2-bromo-isovanillin **15** with no purification of the intermediates required.

With the second generation aryne tether **17** in hand, the corresponding *N*-methyl aniline precursor **18a** was then subjected to the conditions used for attempted cyclisation of **12**. Gratifyingly, the new aryne tether promoted significant intramolecular reaction, affording dihydrophenanthridine **19a** in 50% yield and the corresponding phenanthridinone **20a** in 23% yield, with a marked decrease in deleterious *N*-arylation (8% by-product) (entry 1, Table 1). Having addressed the issue of overall reactivity, attention turned to optimising the reaction conditions to avoid mixtures of cyclisation products (see Table 1 for selected optimisation experiments). Evaluation of some common *o*-SAT activators (entries 2–5) identified CsF in acetonitrile and KF/18-crown-6 in THF as promoting cyclisation to the dihydrophenanthridine **19a** in good yields, albeit with phenanthridinone and by-product still present. Encouragingly, replacing THF with DME led to an excellent yield of the desired dihydrophenanthridine **19a**, with only trace amounts of the phenanthridinone **20a** and no evidence of the competing intermolecular by-product (entries 5 & 6). Further investigations found that lowering the reaction temperature and concentration both decreased the overall yield (entries 7 and 8). Finally, increasing concentration led to more of the competing



Scheme 2 Investigating cyclisation of aryl amine onto *N*-tethered aryne, (a) unsuccessful preliminary attempt to form 5,6-dihydrophenanthridine **13** or phenanthridinone **14**. Reaction conditions: aryl amine (1.0 equiv.), KF (2.0 equiv.), 18-crown-6 (2.0 equiv.), DME : MeCN (3 : 1 by volume, 0.01 M), 90 °C, 24 h, N₂ atmosphere. (b) Synthesis of a new *p*-methoxy benzylic aryne precursor **17**. Reaction conditions: (i) ethylene glycol (5.0 equiv.), pyridinium *p*-toluenesulfonate (0.12 equiv.), PhMe, 110 °C, (ii) hexamethyl-disilazane (0.8 equiv.), THF, 66 °C, 2 h, (iii) *n*-BuLi, THF, -78 °C, 30 min, (iv) trifluoromethanesulfonic anhydride (1.3 equiv.), pyridine (3.0 equiv.), CH₂Cl₂, 0 °C, 3 h, then HCl (4 M aq.), 16 h. Yields of isolated products throughout.

Table 1 Selected optimization studies for the preparation of dihydrophenanthridines^a

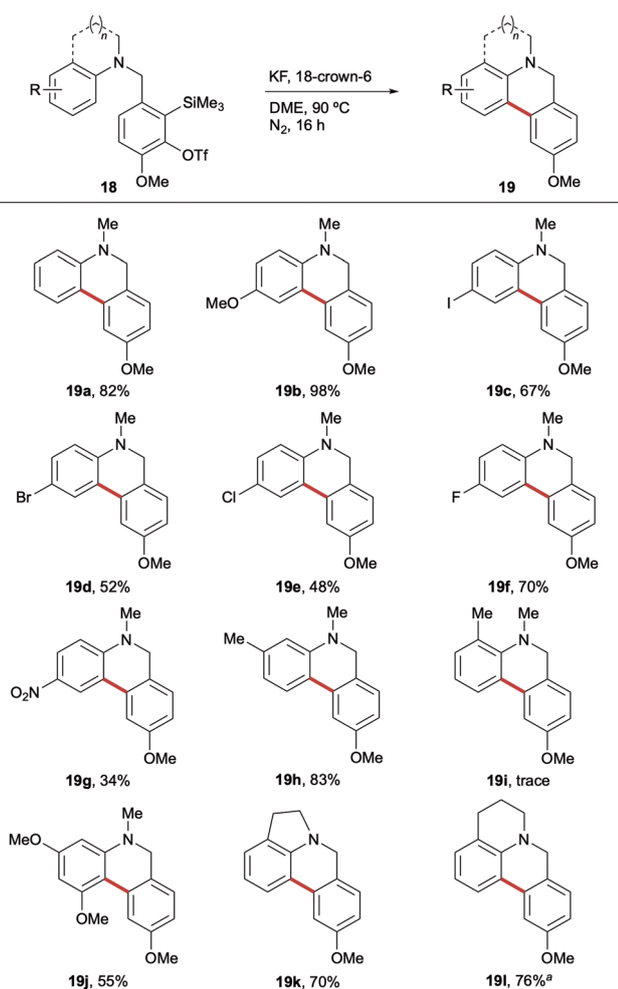
Entry	Activator	Solvent	T (°C)	Yield ^b (%)		
				19a	20a	By-product
1	KF 18-crown-6	DME:MeCN (3:1)	90	50	23	8
2	CsF	MeCN	90	69	2	7
3	CsF	PhMe:MeCN	90	64	10	11
4	TBAF ^c	THF	90	36	—	—
5	KF 18-crown-6	THF	90	69	10	5
6	KF 18-crown-6	DME	90	84(82)	2	—
7	KF 18-crown-6	DME	70	69	2	5
8	KF 18-crown-6	DME ^d	90	69	—	9
9	KF 18-crown-6	DME ^e	90	46	—	35

^a Reaction conditions: aryl amine (1.0 equiv.), activator (2.0 equiv.), additive (2.0 equiv.), solvent [0.01 M], 16 h, N₂ atmosphere. ^b ¹H NMR yield *vs.* dibromomethane internal standard, isolated yield in parentheses, all reactions proceeded to full conversion after 16 h. ^c 1.0 M in THF. ^d 0.002 M. ^e 0.05 M.



intermolecular *N*-arylation and subsequent erosion of the dihydropheanthridine yield (entry 9).

Having optimised the reaction conditions, attention turned to investigating substrate scope. Pleasingly, a range of substituted aniline derivatives were found to undergo cyclisation in generally good to excellent yields (Scheme 3). 4-Substituted anilines were evaluated first, with the electron-rich 4-methoxy derivative affording the corresponding dihydropheanthridine **19b** in 98% yield. The halogenated series showed an interesting trend, with the 4-iodo and 4-fluoro precursors undergoing efficient cyclisation to **19c** (67%) and **19f** (70%), respectively, however, the 4-bromo and 4-chloro derivatives furnished **19d** and **19e** in more moderate yields (52% and 48%). With the electron deficient 4-fluoro analogue proving highly amenable to the cyclisation, it was encouraging to observe that the 4-nitroaniline precursor also underwent the transformation to generate the corresponding electron-poor dihydropheanthridine **19g** in 34% yield. These are particularly noteworthy as

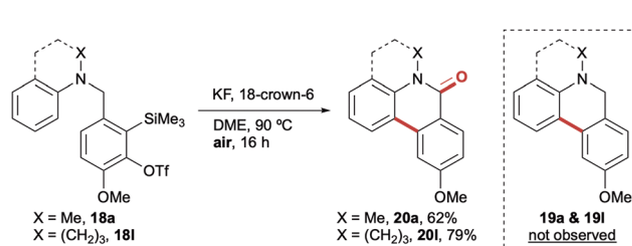


Scheme 3 Synthesis of 5,6-dihydropheanthridine derivatives **15**. Reaction conditions: aryl amine (1.0 equiv.), KF (2.0 equiv.), 18-crown-6 (2.0 equiv.), DME (0.01 M), 90 °C, 16 h, N₂ atmosphere. Yields of isolated products throughout. ^a ¹H NMR yield vs. dibromomethane internal standard, note: **19l** partially oxidised upon purification to give isolated 1:1.2 mixture of **19l** and corresponding phenanthridinone **20l**.

most reports of intramolecular SEAr with arynes rely on markedly electron rich aromatic systems. A comparison of the propensity for cyclisation of the 3-methyl (**18h**) and 2-methylaniline (**18i**) precursors revealed a key role played by sterics. 3-Methylaniline **18h** afforded the corresponding dihydropheanthridine **19h** in an excellent 83% yield, whereas only a trace of the 2-methyl analogue was observed. This suggested that **19i** experiences significant 1,5-strain in the transition state required for cyclisation; instead favouring a mixture of the *N*-arylation by-product and an intramolecular dearomative aryne Diels–Alder cycloaddition.¹⁴ Interestingly, the more conformationally rigid indoline (**18k**) and tetrahydroquinoline (**18l**) precursors underwent efficient SEAr to furnish the corresponding tetracyclic products, **19k** and **19l**, respectively. It was noted that the THQ-derived framework afforded **19l** in 76% yield by ¹H NMR spectroscopic analysis; however, during attempted purification it proved susceptible to oxidation, leading to an isolated mixture (1:1.2) of **19l** and the corresponding phenanthridinone.

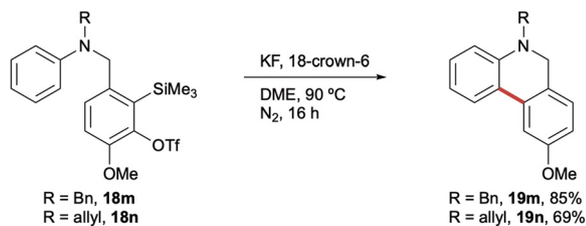
Having established the formation of dihydropheanthridines from a range of electron rich and electron deficient aryl amines, we next investigated whether the product selectivity could be reversed to instead access phenanthridinones. During the initial optimisation studies, dihydropheanthridine **19a** had been found to be susceptible to partial oxidation upon exposure to air, whilst **19l** had part-oxidised during purification. This afforded mixtures of the dihydropheanthridines and phenanthridinones; however a discrete second oxidative step was always required to effect complete conversion. With a view to accessing the phenanthridinone derivatives in a single step from the analogous aryl amine precursors **18**, the reaction was performed in the presence of a range of oxidants, with most attempts affording mixtures of the two cyclisation products. However, it was extremely pleasing to observe that when **18a** and **18l** were exposed to the previously optimised cyclisation conditions, only in air rather than under an inert atmosphere, it resulted in a complete switch of selectivity to exclusively afford the corresponding phenanthridin-6(5*H*)-ones **20a** and **20l** in 62% and 79% yields respectively (Scheme 4).¹⁵

Finally, *N*-benzyl (**18m**) and *N*-allyl (**18n**) aniline precursors were also found to be amenable to cyclisation (Scheme 5). Given the increased potential for subsequent protecting group cleavage, the comparable yields obtained for the analogous



Scheme 4 Cyclisation in air exclusively affords phenanthridinones **20a** and **20l**. Reaction conditions: aryl amine (1.0 equiv.), KF (2.0 equiv.), 18-crown-6 (2.0 equiv.), DME (0.01 M), 90 °C, 16 h, air. Yields of isolated products throughout.





Scheme 5 5,6-Dihydrophenanthridines bearing alternative *N*-protecting groups. Reaction conditions are as shown in Scheme 3. Yields of isolated products throughout.

N-benzyl (**19m**, 85%), *N*-allyl (**19n**, 69%) and *N*-Me (**19a**, 82%) dihydrophenanthridines were particularly encouraging for the synthetic utility of the method.

In conclusion, the cyclisation of aryl amines onto *N*-tethered arynes has been developed under mild reaction conditions using a novel silylaryl triflate precursor. This enabled the new biaryl linkage to be generated by SEAr using both electron rich and poor aryl amines and furnished a range of phenanthridine derivatives with handles for subsequent functionalisation. Selective access to dihydrophenanthridines or the corresponding phenanthridinones is dependent upon a simple switch between anaerobic and aerobic reaction environments.

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Conflicts of interest

There are no conflicts to declare.

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