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Cite this: DOI: 10.1039/c0xx00000x

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Published on 22 August 2013. Downloaded by University of Newcastle on 23/08/2013 00:53:58.

Communication

Communications Accepted Manuscrip

Palladium-catalyzed cyclization of bromoenynamides to tricyclic azacycles: Synthesis of trikentrin-like frameworks

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s Received (in XXX, XXX) Xth XXXXXXX 20XX, Accepted Xth XXXXXXXX 20XX DOI: 10.1039/b000000x

Palladium-catalyzed cascade cyclization of bromoenynamides equipped with an additional alkyne or ynamide substituent affords azatricyclic products. Using 5- to 7-membered ring 10 tethers, this chemistry offers a regiospecific route to highlyfunctionalized azacycles. Elaboration to the trikentrin B skeleton is achieved from the arylsilane cyclization products.

The trikentrin and herbindole families of natural products, isolated from the marine sponges *Trikentrion flabelliforme*^[1a] and ¹⁵ *Axinella* sp.,^[1b] display a range of bioactivities including antimicrobial, antifeedant and cytotoxic properties. The heavily-substituted tricyclic indole systems which feature in these compounds (e.g. *cis*-trikentrin B and herbindole B, Scheme 1) represent a particular synthetic challenge that has inspired a ²⁰ number of elegant solutions.¹

In recent work, we have developed a number of routes to from ynamides, including via ynamide azabicycles carbopalladation.² We have also reported a strategy to access the tricyclic 7,6,5-CDE ring cores of rubriflordilactones A and B, 25 which contain penta- and tetrasubstituted arenes respectively, through palladium-catalyzed cascade cvclization of bromoenediynes.³ We noted that the combination of these two methodologies could provide access to the tricyclic indole core of the trikentrins, via cyclization of a bromoenynamide equipped 30 with a remote alkyne $(1 \rightarrow 2$, Scheme 1). The construction of fused ring arenes in this manner⁴ has rarely been employed in synthetic endeavours,³ and in the context of ynamides offers a useful alternative to the elegant cyclotrimerization methodology pioneered by Witulski,5 which has recently been applied to the

³⁵ herbindole system.^{1c} This sequenced carbopalladation strategy offers advantages over intramolecular cyclotrimerization, where long tethers restrict the formation of larger rings, presumably due to competing intermolecular reactions.⁶ Here we describe the development of this novel ynamide chemistry,⁷ and its ⁴⁰ application to a number of azatricyclic systems, including aza-and benzazepine trikentrin analogues. Elaboration to the *bis*-desmethyl-trikentrin B framework is also described.

We first set about the synthesis of a series of bromoenynamide alkynes suitable for cyclization, through the preparation of ⁴⁵ appropriate sulfonamide and bromoalkyne precursors (Scheme 2). The bromoalkenyl sulfonamides **3a** and **3b** were prepared from alkynes **4a** and **4b** *via* bromoboration / protodeborylation (with simultaneous Boc deprotection), whilst bromoalkynes **5a**



Scheme 1 *cis*-Trikentrin B, herbindole B, and the general bromoenynamide cyclization strategy.

and **5b** were synthesized from 1,6-heptadiyne by monosilylation then bromination. Both trimethylsilyl and benzyldimethylsilyl groups were installed, which we anticipated would enable various ⁵⁵ strategies for the attachment of trikentrin-like sidechains following cyclization. These building blocks were coupled using Hsung's copper-catalyzed methodology for ynamide formation,⁸ which provided ynamides **1a-c** in moderate to good yield, albeit







 a Reaction concentration 0.017 M unless indicated otherwise. b Isolated yield. c Reaction concentration 0.16 M. d Complex mixture.

- ⁵ accompanied by a degree of desilvlation in the case of TMS-substituted diyne 5a. To further extend the cyclization methodology, we targeted a substrate featuring two ynamides, which would lead to an aza-trikentrin (pyrroloindoline) framework. The *bis*-ynamide 1d was readily prepared in four
 ¹⁰ steps from 4a by bromination / carbamate deprotection (to afford the sulfonamide 6), followed by sequential Hsung couplings firstly with 1-bromo-oct-1-yne (used in excess to minimize intermolecular homocoupling of 6), then with sulfonamide 3a.
- With a selection of substrates in hand, the cascade cyclizations ¹⁵ were investigated (Table 1). Using our previously reported conditions (10 mol% Pd(PPh₃)₄, Et₃N, 0.017 M in MeCN, 80 °C),³ we were pleased to obtain the 5,6,5-tricyclic trikentrin frameworks **2a** and **2b** in excellent yields (90%, Entries 1, 4). The catalyst loading could be lowered to 5 mol% with a slight
- ²⁰ reduction in yield (Entries 2, 5); however, by increasing the concentration (to 0.16 M), catalytic efficiency was restored, with **2a** isolated in 88% yield (Entry 3).

Cyclization to the challenging 7,6,5-tricyclic analogue of the trikentrin framework was next attempted. At higher catalyst

- ²⁵ loading and dilution, the desired tricycle 2c was obtained as a component of a complex mixture (Entry 6). However, by performing this reaction at higher concentration, 2c was formed as the sole product in excellent yield (Entry 7), a result that highlights the advantages of the sequenced carbopalladation
- ³⁰ strategy. Finally, diynamide **1d** was subjected to the range of reaction conditions (Entries 8-10). To our delight, tricycle **2d**, which represents the first example of such a pyrroloindoline framework, was isolated in high yield when reacted at the higher

concentration (70%, Entry 10).

With efficient access to azatricycles established, we aimed to demonstrate the utility of the methodology by preparing a natural product analogue - bis-desmethyl-trikentrin B 13 (see Scheme 3) – from the 5,6,5-indolines 2a or 2b. This required installation of the requisite butenyl sidechain, and conversion of the protected 40 indoline to the free indole. For the former of these tasks, we recognised the synthetic value of the silane present in 2a/b, which enables a variety of sidechain attachment strategies. We first addressed Hiyama cross-coupling of 2b, which offers a direct route to the butenvl substituent and is an attractive alternative to 45 other coupling methods (e.g. Stille, Suzuki) due to the low toxicity of silicon and its stability to multistep synthesis.⁹ To our knowledge, no Hiyama couplings between arylbenzyl dimethylsilanes and *alkenyl* halides have been reported, with only the reverse process being described (i.e. the coupling of ⁵⁰ alkenylbenzyldimethylsilanes with aryl halides).¹⁰

Standard conditions for the coupling of alkenyl benzyldimethylsilanes (TBAF, Pd₂dba₃•CHCl₃ or Pd(dba)₂),¹⁰ using either β -styrenyl iodide 7a or butenyl iodide 7b as the halide partner, afforded no cross-coupling product (Entries 1, 2). 55 As benzylsilanes are 'safety-catch' silanols, and indeed are hydrolysed to the latter on treatment with TBAF, alternative conditions for the coupling of alkenylsilanols¹¹ were also investigated, without success (Entry 3). In all of these trials, mixtures of silanol, disiloxane, and desilylated arene were 60 recovered,¹² suggesting that the aryl silanol revealed on unmasking of the benzylsilane was resistant to transmetallation. The addition of Ag₂O has been reported by Hiyama to accelerate transmetallation,¹³ and we were delighted to find that the coupling of styrenyl iodide 7a under these conditions smoothly 65 afforded the styrenyl trikentrin framework 8a (68%). Disappointingly, only desilvlated arene was returned on attempted coupling with butenyl iodide 7b, which for this study presented an insurmountable limitation.

A more classical route to install the butenyl sidechain was thus ⁷⁰ developed (Scheme 3).¹⁴ Aryltrimethylsilane **2a** was subjected to a Friedel-Crafts acylation, which proceeded with exclusive *ipso*selectivity to give ketone **9** (79%). This ketone then underwent a high-yielding reduction / dehydration sequence to deliver the

Table 2 Hiyama Cross-coupling of Arylsilane 2b.

75	N Ts	2b	SiMe ₂ Bn R THF, 22 h Conditions (See Table)	R = Ph R = Et	N Ts 8a (R = Ph) 8b (R = Et)	R
	Entry	Alkenyl Iodide	[Pd] cat. (mol %)	TBAF (equiv.)	Temp (°C)	Yield (%) ^a
	1	7a	Pd ₂ dba ₃ •CHCl ₃ (2.5) or Pd(dba) ₂ (5)	2.2	20→50	_ ^b
	2	7b	Pd ₂ dba ₃ •CHCl ₃ (2.5)	2.2	20→50	_ ^b
	3	7a	(allylPdCl) ₂ (2.5)	2.2	20→50	_ ^b
	4	7a	Pd(PPh ₃) ₄ (5), Ag ₂ O ^c	1.1	20	68
	5	7b	$Pd(PPh_{3})_{4}$ (5), $Ag_{2}O^{c}$	1.1	20	_ ^d
			4			

^{*a*} Isolated yield. ^{*b*} A mixture of silanol, disiloxane, and desilylated arene was recovered. ^{*c*} 1.1 equiv. Ag₂O. ^{*d*} Desilylated **2b** was isolated (67%).



Scheme 3. Synthesis of bis-desmethyl-trikentrin B

targeted butenyl sidechain (**8b**). Completion of the synthesis now required indoline detosylation and oxidation to reveal the indole ⁵ moiety. However, all attempts to oxidise **8b** to the corresponding sulfonyl indole were unsuccessful, leading mainly to degradation.¹⁵ Inverting this sequence of events resolved this issue; although Mg/MeOH/sonication (which is usually effective for such detosylations)^{2a,16} effected partial deprotection (<25%), ¹⁰ treatment of **8b** with sodium naphthalenide gave the deprotected indoline **11** with high efficiency. Somewhat surprisingly, **11** underwent rapid aerobic decomposition, presumably due to the indoline-enhanced reactivity of the electron-rich styrene,¹⁷ and isolation of the pure indoline proved difficult. However, we were ¹⁵ pleased to find that direct dehydrogenation of the synthesis, giving the definite of the synthesis, giving the definite of the synthesis, giving the definite of the synthesis, giving the synthesis of the s

bis-desmethyl-trikentrin **12** in good yield over the two steps. In conclusion, we have developed a facile method for the

preparation of azatricycles from bromoalkenyl ynamides. The 20 reaction enables formation of five- to seven-membered rings, and

- offers an attractive alternative to cyclotrimerization strategies. The utility of this chemistry is demonstrated by installation of the trikentrin B alkenyl sidechain in a further four steps using Friedel-Crafts *ipso*-substitution of the arylsilane cyclization
- ²⁵ products. As an alternative, we report the first example of an alkenyl iodide / arylbenzylsilane Hiyama cross-coupling, which affords a styrenyl-trikentrin analogue.

We thank the EPSRC (EP/H025839/1, CDC; EP/E055273/1, Advanced Research Fellowship to E.A.A.), and Syngenta Ltd. for ³⁰ a studentship (to R.L.G.).

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[†] Electronic Supplementary Information (ESI) available: Experimental details, characterization and copies of ¹H and ¹³C NMR spectra for novel compounds. See DOI: 10.1039/b000000x/

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