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1,3- and 1,4-Benzdiyne equivalents for regioselective synthesis of polycyclic heterocycles†

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We have devised a novel 1,3-benzdiyne equivalent, capable of quadruple functionalization by sequential benzyne generation and reaction with arynophiles. The key features of this method include the chemoselective generation of two triple bonds in a single benzene ring under fluoride-mediated mild conditions, and the regiocontrol of each benzyne reaction by the substituent next to the triple bond. This method produced various benzo-fused heteroaromatic compounds *via* reactions with arynophiles, such as furans, azides, and diazo compounds. A validation of the method is given in the convergent synthesis of the antipsychotic drug risperidone. A similar strategy has also been applied to a 1,4-benzdiyne equivalent to construct linearly benzo-fused heteroaromatics.

Introduction

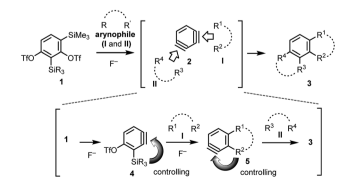
The reactions of benzynes with arynophiles are widely utilized for introducing substituents to adjacent carbons of benzene rings. The direct installation of fused rings onto benzenes is an advantage specific to the benzyne reaction and is not possible through other methods. Furthermore, a variety of new arynophiles have been recently reported, enriching the diversity of the method.²

The reactions of benzdiynes, possessing two triple bonds in a single benzene ring, and two arynophiles, would provide a few-step synthesis for the convergent preparation of multifused benzenes. However, benzdiynes are observed only under gas-phase conditions due to their extreme instability,³ and it would be impossible to react one with two different arynophiles for the synthesis of unsymmetrically fused benzene rings.

An alternative approach is to use benzdiyne equivalents, where two benzynes are generated sequentially in one pot to provide substituted acenes and polycyclic aromatic compounds. If we could control the regiochemistry of consecutive benzyne reactions, starting from benzdiyne equivalents, each with different arynophiles, we could produce a wide range of multiring fused unsymmetrical aromatic compounds. However, only a limited number of such reactions have been reported, and most of them require several steps for functional group transformations to generate the second benzyne. Therefore, the development of more sophisticated benzdiyne equivalents is needed to facilitate two-step sequential benzyne reactions. Crucial factors in the design of these benzdiyne equivalents

include suitable functional groups which enable the generation of the second benzyne without further transformations, and a way to control the regiochemistry of each benzyne reaction. The work of Suzuki *et al.* involving their original 1,4-benzdiyne equivalent meets these requirements, which uses *n*-butyllithum to generate the benzynes. ^{6b} Very recently, Peña *et al.* have demonstrated that triple bonds were sequentially generated twice under fluoride-mediated mild conditions from 1,4-benzdiyne equivalents and reacted with two different arynophiles in both stepwise and one-pot manners. ⁷ In contrast, there have been no reports of a suitable 1,3-benzdiyne equivalent. ^{5,8,9}

We have attempted sequential benzyne reactions starting from 1,3-benzdiyne equivalents 1, with various arynophiles (Scheme 1). This method was designed to afford unsymmetrically substituted polycyclic aromatic compounds 3, possessing consecutive fused-rings, as are often seen in material and pharmaceutical science. While compounds like 3 have been



Scheme 1 Design of benzdiyne equivalent 1 that can sequentially generate two triple bonds in a single benzene ring and control the regiochemistry of two benzyne reactions by the substituents next to each triple bond.

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mainly synthesized via linear, multi-step routes, our approach is convergent and rapid, proceeding by the combination of $\mathbf{1}$ and two different arynophiles (\mathbf{I} and \mathbf{II}), and allows a rational design for the production of a library of compounds. We were particularly interested in its application to the synthesis of benzofused heterocycles for medicinal chemistry. Therefore, we planned reactions using heteroatomic 1,3-dipoles, such as azides, nitrones, diazo compounds, and nitrile oxides, as the arynophiles.

We aimed to develop a synthetic methodology in which (1) two benzynes (4 and 5) are chemoselectively generated in a stepwise manner without any additional functionalization steps, (2) each benzyne is generated under mild conditions using a fluoride, and (3) the two cycloaddition reactions of 4 and 5 with I and II proceed in a highly regioselective manner (Scheme 1). In this article, we report the preparation of a new 1,3-benzdiyne equivalent 1b [SiR₃ = Si(t-Bu)Me₂], and a method for the preparation of unsymmetrical, angular, and multi-ring fused heterocyclic compounds 3, which satisfies the above criteria. One significant advantage of this method is the high regioselectivity of both benzyne reactions, in which the first step is controlled by the traceless directing group, R₃Si (ref. 11) of 4, and the second step by the cyclic systems¹² of 5.

Results and discussion

We synthesized two 1,3-benzdiyne equivalents, 1a and 1b, which were treated with CsF in the presence of 2,4-dimethylfuran 6a. The reaction of 1a with 6a produced the undesired cycloaddition product 8 via benzyne 7 (Scheme 2). However, the reaction of 1b afforded the desired cycloaddition product 10a (78% yield) through the Diels-Alder reaction of the expected benzyne 4a with 6a. An important observation is that the double cycloaddition product 3a was not detected by GC analysis of the crude reaction mixture after 30 min (see ESI†). This may be due to the lower reactivity of the Me₂(t-Bu)Si group, even in the presence of excess CsF and 6a. The generation of the second benzyne 5a was achieved after long reaction time (19 h) under the same reaction conditions using CsF to give 3a in 90% isolated yield.

We attempted to synthesize compounds 3b-f through stepwise benzyne cycloaddition reactions from 1b (Table 1). All

reactions of 3-silylbenzyne 4a with arynophiles 6b-e provided cycloaddition products 10 with good regioselectivities due to synergetic effect of the neighboring Me2(t-Bu)Si (ref. 11f) and the distant triflyloxy (TfO) groups.13 Among them, unexpected proximal regioselectivity (proximal-10d : distal-10d = 78 : 22) was observed in the reaction between 3-silylbenzyne 4a and nitrone 6d (entry 4-1), which was opposite to the previously reported reactions of 3-silylbenzynes with nitrones (for structural determinations, see ESI†).11f-h This result is probably due to the inductively electron-withdrawing effect of the TfO group at C4.13a The reaction of 4a with sydnone 6e to give distal-2Hindazole 10e selectively (entry 5-1) is particularly noteworthy, as the reactions of unsymmetrical benzynes such as 3-methoxybenzyne with sydnones have been reported to afford mixtures of regioisomers in 1:1 ratio. 2f The reaction of benzynes 5 with arynophiles 6b and 6f-g provided polycyclic compounds 3b-f. This is the first report of the generation and reaction of 4,5benzotriazolyne **5b** (entry 2-2), 6,7-benzisoxazolyne **5c** (entry 3-2), 4,5-benzisoxazolinyne 5d (entry 4-2), and 6,7-2H-indazolyne 5e (entry 5-2). The regioselectivity of these reactions is higher than that of sterically similar 4,5-indolyne (see preliminary theoretical discussion of these regionelectivities in ESI†). 12a,b The reactions of 5c with 6b and 5e with 6g provided distal-3d and proximal-3f exclusively (entries 3-2 and 5-2).

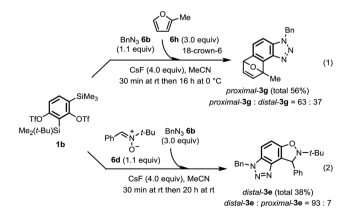
Next, one-pot sequential benzyne cycloadditions from **1b** were demonstrated for the synthesis of angular tricyclic heterocycles **3** without isolating **10** (Scheme 3). After a mixture of **1b** (1.0 equiv.), benzyl azide **6b** (1.1 equiv.) and CsF (4.0 equiv.) in MeCN was stirred at room temperature for 30 min, 2-methylfuran **6h** (3.0 equiv.) and 18-crown-6 (4.0 equiv.) were added and then the reaction mixture was stirred for 16 h at 0 °C (Scheme 3-1). Gratifyingly, *proximal-3g* was obtained as a main product (*proximal-3g*: *distal-3g* = 63:37, total 56%). The tricyclic compound, *distal-3e* was also synthesized as the predominant product (*distal-3e*: *proximal-3e* = 93:7, total 38%) by a similar one-pot combination of arynophiles, nitrone **6d** and **6b** (Scheme 3-2). The yield and regioselectivity of these products were comparable to those obtained by the stepwise method (Table 1, entries 4-1 and 4-2).

We applied these findings to the convergent synthesis of the antipsychotic drug risperidone **14** (Scheme 4). The silylbenzyne **4a** and a nitrile oxide **6i** (ref. 2*e*) were simultaneously generated

Scheme 2 Sequential benzyne generation from benzdiyne equivalents 1a and 1b followed by Diels-Alder reaction with furan 6a.

Table 1 Reactions of 1b with two different arynophiles 6 for the synthesis of angular polycycles 3'

^a Conditions: **1b** or **10** (1.0 equiv.), arynophile (3.0 equiv.), CsF (3.0 equiv.) in MeCN at rt. ^b The ratio of major and minor products was determined by ¹H NMR. ^c Total isolated yield of *distal*-**10** (or *distal*-3) and its regioisomer *proximal*-**10** (or *proximal*-3). Mes = C₆H₂-2,4,6-Me₃.



Scheme 3 One-pot synthesis of unsymmetrical angular heterocycles 3g and 3e.

from a mixture of **1b** and a chloro-oxime **11** and then reacted *in situ* to form *distal***-10f** as a single regioisomer. The next reaction

of 6,7-benzisoxazolyne 5**f**, generated by BnMe₃NF (ref. 14 and 15) and a fluoride 6**j**, provided 3**h** with excellent regioselectivity. Finally, the synthesis was completed by the *N*-deprotection of 3**h** to give 12 and the alkylation with 13.¹⁶ This result suggests that 1**b** should be useful tool for the expeditious divergent synthesis of a wide variety of biologically active compounds and their derivatives by choosing different arynophiles 6 once 1**b** become easily available (for the first synthesis of 1**b**, see ESI†).

We also report the synthesis of linearly fused, unsymmetrical polycyclic aromatics **19** using **15** (ref. 17–19) as a 1,4-benzdiyne equivalent (Table 2). The first benzyne generation proceeds using CsF at room temperature in MeCN for a short time, under which conditions, generation of the second benzyne does not occur (see ESI†). The mono-cycloaddition products **17**, obtained as a mixture of two regioisomers, were subjected to the second reaction with arynophiles **II**, without separation of the regioisomers, ²⁰ to afford the multicyclic compounds **19**. Due to the dual effect of the TfO group^{13a} and Me₃Si group^{11e} of **16**, the all first benzyne reactions proceeded in a regioselective manner

Scheme 4 Application of 1.3-benzdivne equivalent 1b to the synthesis of risperidone 14

Table 2 Reactions of a 1,4-benzdiyne equivalent 15 with two different arynophiles 6 for the synthesis of linear polycycles 19^{a}

TfO Me ₃ Si ¹	SiMe ₃	R2 6 arynophile I CsF, MeCN, rt 15–45 min		TfO Me ₃ Si	R ¹ .	R ³ 6 arynophile II CsF, MeCN, rt 3–11 h	$\begin{bmatrix} \begin{pmatrix} R^3 & \mathbf{R}^1 \\ \mathbf{R}^4 & \mathbf{R}^2 \end{pmatrix} \end{bmatrix}$	$\begin{array}{c} \longrightarrow & \left(\begin{array}{c} R^3 \\ \\ R^4 \end{array} \right) \end{array}$
	Product 17			.			Product 19	
Entry	Arynophile I	Major isomer	Regioselectivity ^b Yield ^c	Entry	Arynophile II	2nd benzyne 18	Major isomer	Regioselectivity ^b Yield ^c
1-1	6b	Me ₃ Si N N N N N N N N Proximal-17a Bn	85 : 15 61%	1-2	MeN NMe	N N Bn	MeN N N N N N N N N N N N N N N N N N N	N 75:25 71% (74:26) ^d (37%) ^d 3n
2-1	6d	TfO Ph Me ₃ Si O N—t-Bi	u 73:27 71%	2-2	6g	Ph N- <i>t</i> -Bu	N N	Ph (N— <i>t-</i> Bu 70 : 30 57%
3-1	EtO N ₂	Me ₃ Si N N N N N N N N N N N N N N N N N N N	76 : 24 66% it	3-2	MeO OMe	H N N 18c CO ₂ Et	MeO HNNN Notes and MeO HNNN Notes and MeO HNNNN Notes and MeO HNNNNN Notes and MeO HNNNNNN Notes and MeO HNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN	76 : 24 83% ^[e] ₂ Et

^a Conditions: **15** or **17** (1.0 equiv.), arynophile **6** (3.0 equiv.), CsF (3.0 equiv.) in MeCN at rt. ^b A ratio of major and minor products was determined by ¹H NMR. ^c Total isolated yield of *distal-***17** (or *distal-***19**) and its regioisomer *proximal-***17** (or *proximal-***19**). ^d One-pot reaction conditions: **15** (1.0 equiv.), 6b (1.1 equiv.), 6l (3.0 equiv.), CsF (4.0 equiv.) in MeCN (0.1 M) at rt for 30 min and then at rt for 14 h. The yield was calculated based on 15. ^e Isolated as a corresponding ketone 19c' after the hydrolysis of acetal 19c.

beyond expectation (Table 2, entries 1-1, 2-1 and 3-1), although these selectivities were only a little lower than those of the 1,3benzdiyne equivalent 1b (see, Table 1). Interestingly, the second benzyne reactions also regioselectively provided cycloaddition products 19 probably because of the inductive effect of heteroatoms such as nitrogen and oxygen constructing heterocycles (entries 2-2 and 3-2). These results provide useful information for regioselectivity control of benzyne cycloadditions from distant positions. Importantly, the one-pot synthesis of a linear tricyclic compound 19a from 15, 6b and 6l was also successfully achieved (see entry 1-2).

Conclusions

In conclusion, we have developed a novel synthetic route to multi-ring fused heterocycles by the combination of benzdiyne equivalents and arynophiles. In this study, the newly generated azole-fused benzynes were found to exhibit higher regioselectivities than those of sterically similar 4,5-indolyne. 12b This method has facilitated the convergent synthesis of the antipsychotic risperidone. Therefore, we believe that this synthetic methodology will be invaluable to drug discovery. Work is ongoing into easier, scalable synthetic methods for these benzdiyne equivalents,19 analysis of the origin of the regioselectivity (see ESI†), and applications to medicinal chemistry.

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- two equivalents of a single arynophile.^{4,18} Peña *et al.* have recently used **15** as a quasi-step-by-step benzdiyne equivalent for the synthesis of polyaromatic hydrocarbon (PAH), in which the second benzyne was not generated because of the precipitation of the mono-cycloaddition product^{7a-c}.
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