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Cyclic Diaryl λ3-Bromanes: A Rapid Access to Molecular Complexity via Cycloaddition Reactions / Lanzi, M.; Ali Abdine, R. A.; De Abreu, M.; Wencel-Delord, J.. - In: ORGANIC LETTERS. - ISSN 1523-7060. -23:23(2021), pp. 9047-9052. [10.1021/acs.orglett.1c03278]

Availability: This version is available at: 11381/2994018 since: 2024-08-21T10:05:47Z

Publisher: American Chemical Society

Published DOI:10.1021/acs.orglett.1c03278

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Cyclic Diaryl λ^3 -Bromanes: a Rapid Access to Molecular Complexity via Cycloaddition Reactions

Matteo Lanzi, Racha Abed Ali Abdine, ‡ Maxime De Abreu, ‡ and Joanna Wencel-Delord*

Laboratoire d'Innovation Moléculaire et Applications (UMR CNRS 7042), Université de Strasbourg/Université de Haute Alsace, ECPM, 25 rue Becquerel, 67087 Strasbourg (France) Supporting Information Placeholder



ABSTRACT: Biaryls have widespread applications in organic synthesis. However, sequentially polysubstituted biaryls are underdeveloped due to their challenging preparation. Herein, we report the synthesis of unsymmetric 2,3,2',3',4-substituted biaryls *via* pericyclic reactions of cyclic diaryl λ^3 -bromanes. The functional groups tolerance and atom economy allow access to molecular complexity in a single reaction step. Continuous flow protocol has been designed for the scale-up of the reaction, while post-functionalizations have been developed taking advantage of the residual bromide.

The direct access to molecular complexity under sustainable, simple, and mild reaction conditions is a critical challenge of modern organic chemistry. Biaryl compounds are ubiquitous motifs in many fields of chemistry, including medicinal chemistry, agrochemistry, material science, and ligand design. A straightforward synthesis of highly decorated biaryls thus remains an important challenge. ¹ In particular, dissymmetrical 2,3,2',3',4-substituted biaryls are poorly explored motifs due to the arduous synthesis via standard cross-coupling reactions which require densely decorated precursors. Albeit scarcely investigated, a general and straightforward alternative approach based on a modular functionalization via pericyclic reactions of a finely designed biaryl benzyne precursor represents an attractive solution to this synthetic issue (Scheme 1a).² However, multiple drawbacks hamper the state-of-the-art of the aryne-based approach (Scheme 1b): 1) lengthy synthetic routes for the preparation of aryne precursors; 2) the limited atom economy resulting from the aryne generation with the removal of leaving groups (-OTf and -TMS; -X and -OR; Aux-I) which translate into significant waste formation and 3) the limited functional group tolerance as the base- and fluorine-sensitive moieties are generally incompatible. ³⁻⁵ Accordingly, the implementation of the pericyclic reactions to rapidly access dissymmetrical 2,3,2',3',4'-substituted biaryls diversity requests design of original biaryl aryne precursors, activated under mild reaction conditions and in presence of a weak base.

Despite being known since the 19th century and albeit their unique reactivity, hypervalent bromane (III) reagents

have received much less attention compared to the corresponding iodine compounds. $^{\rm 6}$

The lack of general and simple synthetic routes deliver-Scheme 1: a) Common aryne precursors; b) Aryne strategy to *ortho-*, *ortho-*, *meta-* substituted biaryls, c) Cyclic diaryl λ^3 -bromanes as aryne precursors in pericyclic approach towards *ortho-*, *ortho-*, *meta-* substituted



ing these unique scaffolds has drastically hampered the development of bromane (III) chemistry, 7 and thus the rare advances in this field have been reported only recently. 8 Our group has latterly established a simple, safe, and scalable protocol for the preparation of cyclic diaryl λ^3 bromanes in good to excellent yields. 9 The superior electron-withdrawing effect and the nucleofugality of hypervalent bromane (III) compared to the iodine (III) counterpart (approximately 10⁶ times higher), confer them a complementary reactivity. ¹⁰ Encouraged by the disclosed reactivity of cyclic diaryl λ^3 -bromanes towards the construction of carbon-oxygen and carbon-nitrogen bonds, we speculate that they could represent a new family of biaryl coupling partners for pericyclic reactions (Scheme 1c). Moreover, cyclic hypervalent λ^3 -bromane products carry a residual functional bromine atom which opens additional perspectives for further post-functionalizations, while warranting excellent atom economy of the overall process. Thus unique reactivity provides a new strategy for the synthesis of valuable and unprecedented highly decorated dissymmetric 2,3,2',3',4'-substituted biarvls. helicenes ¹¹ and four consequently substituted aromatic units. ^{4e,12}

Our investigation begun by studying the reaction between the well-known cyclic diaryl λ^3 -iodanes and furan **2a** (Table 1, entry 2), but no product formation was observed neither by **GC-MS** nor ¹**HNMR** (see Supporting Information). Remarkably, while using cyclic diaryl λ^3 bromane **1a**-OTf and **2a**, in a presence of cesium carbonate and DCM, the desired product **3aa** was afforded in 92% yield after 16 hours at room temperature (entry 1), unambiguously illustrating the unique reactivity of these rea-

Table 1. Optimization of the Diels-Alder Reaction with cyclic diaryl hypervalent bromanes^a

1a-0	$\begin{array}{c} & & & \\ & & \\ & & \\ & \\ & \\ & \\ & \\ & $	3aa
entry	deviation from standard conditions	yield (%) ^b
1	none	92% ^c
2	λ^3 -I-OTf/-OMs instead of 1a -OTf	_d
3	no Cs ₂ CO ₃	_ d
4	tBuONa / K ₂ CO ₃ instead to Cs ₂ CO ₃	59% / 23%
5	CH_3CN / H_2O instead to CH_2Cl_2	69% / 13%

^a Reaction conditions: λ^3 -X-Y (0.1 mmol), furan (1.2 equiv), r.t. 16h; ^b 1HNMR yields using CH₂Br₂ as internal standard; ^c Isolated yield; ^d 5 equiv furan.

gents.

No reactivity was observed in the absence of Cs_2CO_3 (entry 3). Further investigation showed that the nature of the base holds a key role. Despite being a stronger base, *t*BuONa provided **3aa** in lower yield, while K_2CO_3 performed poorly (entry 4). Finally, the role of a chlorinated polar solvent, namely CH₂Cl₂, was confirmed (entry 5). Surprising, albeit in low yield, the reaction occurred in

water media. ¹³ With the optimal reaction conditions in hand, the scope of this cycloaddition was explored, focusing on various cyclic diaryl λ^3 -bromanes **1a-l** and aromatic cyclic Diels-Alder partners 2b-2d (Scheme 2). The symmetric bromane 1b, presenting inductive electrondonating methyl groups provided the desired product 3ba in good yield (69%). Regioselective functionalizations took place in presence of asymmetric substrates, such as 1c, 1d, and 1e, where the enhanced reactivity of one aryl moieties translated into the formation of a unique products in good yields. The introduction of the methyl group in orthoposition directs the desired transformation onto one aromatic ring, thus promoting the formation of 3fa in moderate yield (51%). Bifunctional dissymmetric λ^3 -bromanes bearing methyl ester and chlorine afforded 3ga and 3ha in 59% and 98% yields respectively. Finally, monosubstituted cyclic diaryl λ^3 -bromanes were tested. A weak electron-donating methyl group, **1***i*, unsurprisingly leads to a 1:1 mixture of products 3ia and 3ia', while the enhanced electron-withdrawing character of methyl ester 1j, chloro **1k** and trifluoromethyl **1l** enabled the preferential formation of one regioisomer with good selectivity. Moreover, substituted furans, **2b** and **2c** and N-protected pyrrole **2d** were well tolerated under our reaction conditions, affording the desired products in good yields. Moreover, the dissymmetric bromane **1h** reacted smoothly with **2d** offering the highly functionalized bromo, chloro cycloadduct 3hd in 84% yield. Targeting a rapid generation of a broad range of diverse molecules, we have challenged the λ^3 bromanes with other arynophiles. Various azides have been successfully coupled with **1a**-OTf, furnishing highly decorated bromo benzotriazole derivatives. Simple benzyl azides 2e and substituted 2f - 2i supplied the desired regioisomeric 3 and 3' in moderate to good yields, which were straightforwardly separated via column providing the pure products. The single-crystal X-ray analysis of 3af confirmed the triazole structure and the regioselectivity. Remarkably, thanks to the fluorine-free conditions, the silyl-protected phenols 3ai and 3ai' could be obtained in excellent yields, highlighting the complementarity of this protocol. The cholesterol-de derived 2j and the cyclohexyl azide 2k react smoothly under the standard reaction conditions. Diazomethane 21 (TMSCHN₂) also proved to be an excellent arynophile, yielding **3al** and **3cl** in 92% and 50% yields respectively, albeit as a mixture of two cycloadducts. In accordance with the aryne distorted and steric models, a moderate regiocontrol has been observed in the 1,3dipolar cycloaddition where the nucleophilic nitrogen reacts with the more distorted alkyne terminus position furnishing the less steric demanding product as major isomer.¹⁴ While Diels-Alder reactions involving arvne have been largely investigated, ¹⁵ the [2+2] reactions have received much less attention despite the interest of the resulting peculiar structures. ¹⁶ Rewardingly, the reaction of 1a-OTf with 2,3-dihydrofuran 2m and 2,3-dihydropyran 2n conducted to the selective formation of 3am and 3an in excellent yields, furnishing a single regioisomer with a ratio up to 14.3:1. Moreover, strained cycloalkenes, namely norbornene 20 and norbornadiene 2p, provided the desired cyclobutene products **3ao** and **3ap** with good yields. Interestingly, an elusive [2+2+2] mechanism has been observed within **1a**-OTf and **2p** yielding the uncommon Scheme 3: Scope of the reaction



3ap' in 17% yield. The acrylate 2q showed sufficient reactivity under our reaction conditions. Asymmetric λ^3 bromanes **1h**, **1c** and **1g** provided the highly decorated biaryls 3hm, 3cn and 3go in excellent yields. Finally, structurally condensed polycyclic aromatic compounds, valuable motifs in material science, were targeted. We investigated the reaction of **1a**-OTf with the β -bromostyrenes **2r**. After a minor reoptimization of the reaction conditions, a cascade [4+2] cycloaddition/base-mediated aromatization occurred smoothly, at room temperature, in dioxane yielding the endo-product **3ar** as a single regioisomer. Electronrich bromo styryls showed higher reactivity, delivering 3as and 3ay respectively in 85% and 58% yields. The single-crystal X-ray analysis of 3as confirmed the formation of the endo-product. Electron withdrawing groups, such as trifluoromethyl, impact the reaction outcome, delivering **3at** in a decreased yield.

On the other hand, halogenated aromatic rings 2u, 2v, 2w and 2x, offered synthetically useful bifunctional products in moderate to high yields. The electron-rich, protected bromo caffeic acid derivative 2z turned out to be suitable for the synthesis of highly decorated phenyl

phenanthrene **3az**, providing a functional dibromo compound. Sterically demanding substrate **2A** furnished the conformationally stable atropoisomers **3aA**. Finally, when dissymmetric λ^3 -bromane **1f** was tested, a selective functionalization occurred, yielding **3fs**. Remarkably, the excellent regioselectivity reached in all of these cases, providing the sterically hindered *endo*-regioisomers as unique product, outcompetes the literature precedents. ¹⁷





Encouraged by the high versatility observed using cyclic diaryl λ^3 -bromanes, we endeavored on developing a flow protocol for this reaction (Scheme 3). Despite clear advantageous of flow chemistry approach as scalability, safety

Scheme 4: Synthesis of functional biaryl compounds from 3aa, 3ar, 3as and 3au



and reduction of the reaction time, implementation of this process into the aryne chemistry remains rare and the chemically difficult. ¹⁸ The difficulty in our case lays in the use of insoluble Cs₂CO₃, however, attempts to replace it with organic bases failed. (see Supporting Information). Accordingly, the flow-reaction was performed in a biphasic system, which involved a water solution of Cs_2CO_3 and a solution of 1a-OTf and 2a in CH₃CN. The optimal flowreaction conditions enabled the synthesis of 516 mg of 3aa (66% yield) within 22 min of residence time at 60 °C. Interesting, the substrate **1a**-OTf is stable under these reaction conditions and the unreacted portion was recovered via precipitation with Et₂O from the reaction steam. The conspicuous reduction of the reaction time and the use of green solvents make this approach a suitable alternative to the batch protocol, especially for further reaction scale-up.

Beyond the mild reaction conditions for the arvne generation and the high functional group tolerance demonstrated, the cycloadduct products feature a bromine atom which can serve as an additional handle for the incorporation of molecular complexity (Scheme 4). Initially, the robustness of our procedures was confirmed by performing the reactions on 1 or 2 mmol scales, furnishing 3aa, 3ar, 3as and 3au in multi milligram scale. The functionalization of the 3aa through lithium-mediated bromine exchange provided the corresponding carboxylic acid 4, aldehyde 5, and the new phosphine 6 in 64%, 93% and 76% yields respectively. Moreover, a standard deoxygenation protocol generated 7 in 74% yield. Also, bromo-biphenyl phenanthrenes **3ar**, **3as** and **3au** are synthetically interesting scaffolds and the residual bromine enables access to highly adorned structures. Pdcatalyzed cross-couplings, such as Sonogashira and Suzuki reactions granted the introduction of alkyne 8 and aryl 9 moieties with useful yields. Furthermore, a Buchwald-Hartwig Pd-catalyzed couplings delivered 10 and the sterically challenging 11 diaryl amines, in good yields. A lithium exchange protocol using **3ar**, **3as** and **3au**, led to the versatile aldehydes **12**, **13** and **14**. A subsequent Wittig reaction involving the aldehyde **13** provided the stereoisomers *E*-**15** and *Z*-**15** in good yield (70% + 17% respectively), which were easily separated by flash chromatography. To further illustrate the utility of the bromine moiety, the synthesis of screw-shaped helicene was envisioned. A standard Corey-Fuchs alkyne synthesis using the aldehyde **14** furnished **17**. Then, a Pt-catalyzed cyclization afforded the desired [5]-helicene **18** in 12% overall yield.

In conclusion, we demonstrated herein the peculiar versatility, chemoselectivity and atom economy of cyclic diaryl λ^3 -bromanes as biaryl aryne precursors in cycloaddition reactions for the synthesis of highly decorated dissymmetric 2,3,2',3',4'-substituted biaryl motifs. The mild aryne generation conditions, namely carbonate base and room temperature, allowed a wide arynophiles compatibility, together with great functional group tolerance. Robustness of these aryne precursors guaranteed an efficient biphasic flow protocol for the pericyclic reaction further improving the sustainability. High molecular complexity *via* multiple C-C and C-N bonds formation was attained in a single reaction step, while subsequent functionalizations further expand access to a plethora of complex functional compounds.

ASSOCIATED CONTENT

Supporting Information

Experimental procedures, synthesis of reagents, copies of NMR spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

* wenceldelord@unistra.fr

Author Contributions

*‡*These authors contributed equally.

Notes

The authors declare no conflict of interest.

ACKNOWLEDGMENT

We thank the CNRS (Centre National de la Recherche Scientifique), the "Ministère de l'Education Nationale et de la Recherche" (France) for financial support. J.W.D., M.L. and M.D.A. are very grateful to European Commission for the ERC-Starting Grant "AlCHIMIE" N°949804. We are grateful to Dr. Lydia Karmazin, Dr. Corinne Bailly and Dr. Nathalie Gruber for X-ray diffraction analysis (Service de Radiocristallographie, CNRS-Université de Strasbourg, Strasbourg, France). Authors are also very grateful to Dr. Morgan Donnard, Dr. Armen Panossian and Dr. Frédéric R. Leroux for helpful discussions on flow-chemistry.

REFERENCES

(1) (a) Wu, P.; Nielsen, T. E.; Clausen, M. H. Small-molecule kinase inhibitors: an analysis of FDA-approved drugs. *Drug Discovery Today* **2016**, *21*, 5–10. (b) Mullard, A. 2020 FDA Drug Approvals. *Nat Rev Drug Discov* **2021**, *20*, 85–90. (c) Bhutani, P.; Joshi, G.; Raja, N.; Bachhav, N.; Rajanna, P. K.; Bhutani, H.; Paul, A. T.; Kumar, R. U.S. FDA Approved Drugs from 2015–June 2020: A Perspective. *J. Med. Chem.* **2021**, *64*, 2339–2381. (d) Jhulki, S.; Ghosh, A.; Chow, T. J.; Moorthy, J. N. Twisted Biaryl-Amines as Novel Host Materials for Green-Emissive Phosphorescent Organic Light-Emitting Diodes (PhOLEDs). *RSC Adv.* **2015**, *5*, 101169–101176. (e) Esmaeili, A. A.; Moradi, A.; Mohammadi, H. K. Regioselective Synthesis of Highly-Substituted Biaryls by Reaction of Vinyl Malononitriles with Acetylenic Esters. *Tetrahedron* **2010**, *66*, 3575–3578.

(2) (a) Standera, M.; Häfliger, R.; Gershoni-Poranne, R.; Stanger, A.; Jeschke, G.; van Beek, J. D.; Bertschi, L.; Schlüter, A. D. Evidence for Fully Conjugated Double-Stranded Cycles. Chem. Eur. J. 2011, 17, 12163-12174. (b) Diemer, V.; Garcia, J. S.; Leroux, F. R.; Colobert, F. Aryne-Mediated Fluorination: Synthesis of Fluorinated Biaryls via a Sequential Desilvlation-Halide Elimination-Fluoride Addition Process. J. Fluorine Chem. 2012, 134, 146-155. (c) Neudorff, W. D.; Schulte, N.; Lentz, D.; Schlüter, A. D. 8,9-Didehydrofluoranthenes as Building Blocks for the Synthesis of Extended Polycyclic Aromatic Hydrocarbons (PAHs). Org. Lett. 2001, 3, 3115-3118. (d) Akai, S.; Ikawa, T.; Takayanagi, S.; Morikawa, Y.; Mohri, S.; Tsubakiyama, M.; Egi, M.; Wada, Y.; Kita, Y. Synthesis of Biaryl Compounds through Three-Component Assembly: Ambidentate Effect of the Tert -Butyldimethylsilyl Group for Regioselective Diels-Alder and Hiyama Coupling Reactions. Angew. Chem. Int. Ed. 2008, 47, 7673-7676. (e) Moreira, B.; Muraca, A.; Raminelli, C. Synthesis of Silylbiaryl Triflates by Chemoselective Suzuki Reaction. Synthesis 2016, 49, 1093–1102. (f) Asgari, P.; Dakarapu, U. S.; Nguyen, H. H.; Jeon, J. Aryne Cycloaddition Reactions of Benzodioxasilines as Aryne Precursors Generated by Catalytic Reductive Ortho-C H Silylation of Phenols with Traceless Acetal Directing Groups. Tetrahedron 2017, 73, 4052-4061. (g) Wei, Y.-L.; Dauvergne, G.; Rodriguez, J.; Coquerel, Y. Enantiospecific Generation and Trapping Reactions of Aryne Atropisomers. J. Am. Chem. Soc. 2020, 142, 16921-16925. (h) Werz, D. B.; Biju, A. T. Uncovering the Neglected Similarities of Arynes and Donor-Acceptor Cyclopropanes. Angew. Chem. Int. Ed. **2020**, *59*, 3385–3398

(3) (a) Shi, J.; Li, L.; Li, Y. O -Silylaryl Triflates: A Journey of Kobayashi Aryne Precursors. *Chem. Rev.* **2021**, *121*, 3892-4022. (b) Sarmah, M.; Sharma, A.; Gogoi, P. Exploration of Kobayashi's Aryne Precursor: A Potent Reactive Platform for the Synthesis of Polycyclic Aromatic Hydrocarbons. *Org. Biomol. Chem.* **2021**, *19*, 722-737. (c) Idiris, F. I. M.; Jones, C. R. Recent Advances in Fluoride-Free Aryne Generation from Arene Precursors. *Org. Biomol. Chem.* **2017**, *15*, 9044–9056. (d) Roy, T.; Biju, A. T. Recent Advances and the second s

vances in Molecular Rearrangements Involving Aryne Intermediates. *Chem. Commun.* **2018**, *54*, 2580–2594.

(a) Sundalam, S. K.; Nilova, A.; Seidl, T. L.; Stuart, D. R. A (4) Selective C-H Deprotonation Strategy to Access Functionalized Arynes by Using Hypervalent Iodine. Angew. Chem. Int. Ed. 2016, 55, 8431-8434. (b) Chen, H.; Han, J.; Wang, L. Diels-Alder Cycloadditions of N -Arylpyrroles via Aryne Intermediates Using Diaryliodonium Salts. Beilstein J. Org. Chem. 2018, 14, 354-363. (c) Zhang, Z.; Wu, X.; Han, J.; Wu, W.; Wang, L. Direct Arylation of Tertiary Amines via Aryne Intermediates Using Diaryliodonium Salts. Tetrahedron Letters 2018, 59, 1737-1741. (d) Nilova, A.; Metze, B.: Stuart, D. R. Arvl(TMP)Iodonium Tosvlate Reagents as a Strategic Entry Point to Diverse Aryl Intermediates: Selective Access to Arynes. Org. Lett. 2021, 23, 4813-4817. (e) Nilova, A.; Sibbald, P. A.; Valente, E. J.; González-Montiel, G. A.; Richardson, H. C.; Brown, K. S.; Cheong, P. H.; Stuart, D. R. Regioselective Synthesis of 1,2,3,4-Tetrasubstituted Arenes by Vicinal Functionalization of Arynes Derived from Aryl(Mes)Iodonium Salts. Chem. Eur. J. **2021**, *27*, 7168 – 7175.

(5) (a) Heinz, B.; Djukanovic, D.; Filipponi, P.; Martin, B.; Karaghiosoff, K.; Knochel, P. Regioselective Difunctionalization of Pyridines via 3,4-Pyridynes. *Chem. Sci.* **2021**, *12*, 6143–6147. (b) Lin, W.; Sapountzis, I.; Knochel, P. Preparation of Functionalized Aryl Magnesium Reagents by the Addition of Magnesium Aryl Thiolates and Amides to Arynes. *Angew. Chem. Int. Ed.* **2005**, *44*, 4258–4261. (c) Sapountzis, I.; Lin, W.; Fischer, M.; Knochel, P. Preparation of Polyfunctional Arynes via 2-Magnesiated Diaryl Sulfonates. *Angew. Chem. Int. Ed.* **2004**, *43*, 4364–4366.

(6) (a) Wirth, T. Hypervalent Iodine Chemistry in Synthesis: Scope and New Directions. *Angew. Chem. Int. Ed.* **2005**, *44*, 3656–3665. (c) Ochiai, M. Hypervalent Aryl-, Alkynyl-, and Alkenyl- λ^3 -Bromanes. *Synlett* **2009**, *2009*, 159–173. (d) Bhojgude, S. S.; Thangaraj, M.; Suresh, E.; Biju, A. T. *Org. Lett.* **2014**, *16*, 3576-3579

(7) Ochiai, M.; Miyamoto, K.; Kaneaki, T.; Hayashi, S.; Nakanishi, W. Highly Regioselective Amination of Unactivated Alkanes by Hypervalent Sulfonylimino- λ^3 -Bromane. *Science* **2011**, *332*, 448–451.

(8) (a) Yoshida, Y.; Ishikawa, S.; Mino, T.; Sakamoto, M. Bromonium Salts: Diaryl- λ^3 -Bromanes as Halogen-Bonding Organocatalysts. *Chem. Commun.* **2021**, *57*, 2519–2522. (b) Sokolovs, I.; Mohebbati, N.; Francke, R.; Suna, E. Electrochemical Generation of Hypervalent Bromine(III) Compounds. *Angew. Chem. Int. Ed.* **2021**, *60*, 15832–15837. (c) Miyamoto, K.; Saito, M.; Tsuji, S.; Takagi, T.; Shiro, M.; Uchiyama, M.; Ochiai, M. Benchtop-Stable Hypervalent Bromine(III) Compounds: Versatile Strategy and Platform for Air- and Moisture-Stable λ^3 -Bromanes. *J. Am. Chem. Soc.* **2021**, *143*, 9327–9331.

(9) Lanzi, M.; Dherbassy, Q.; Wencel-Delord, J. Cyclic Diaryl λ^3 -Bromanes as Original Aryne Precursors. *Angew. Chem. Int. Ed.* **2021**, *60*, 14852–14857.

(10) (a) Tolstaya, T. P.; Demkina, I. I.; Grushin, V. V.; Vanchikov, A. N. Nucleophilic Substitution in Diphenylbromonium and Diphenyl- chloronium Salts. *J. Org. Chem. USSR* **1989**, *25*, 2305–2310. (b) Nakajima, M.; Miyamoto, K.; Hirano, K.; Uchiyama, M. Diaryl- λ^3 -chloranes: versatile synthesis and unique reactivity as aryl cation equivalent. *J. Am. Chem. Soc.* **2019**, *141*, 6499-6503

(11) Shen, Y.; Chen, C.-F. Helicenes: Synthesis and Applications. *Chem. Rev.* **2012**, *112*, 1463–1535.

(12) Biju, A. T. Modern Aryne Chemistry, Wiley, 2021.

(13) Yoshimura, A.; Fuchs, J. M.; Middleton, K. R.; Maskaev, A. V.; Rohde, G. T.; Saito, A.; Postnikov, P. S.; Yusubov, M. S.; Nemykin, V. N.; Zhdankin, V. V. Pseudocyclic Arylbenziodoxaboroles: Efficient Benzyne Precursors Triggered by Water at Room Temperature. *Chem. Eur. J.* **2017**, *23*, 16738–16742.

(14) (a) Medina, J. M.; Mackey, J. L.; Garg, N. K.; Houk, K. N. The Role of Aryne Distortions, Steric Effects, and Charges in Regioselectivities of Aryne Reactions. *J. Am. Chem. Soc.* **2014**, *136*, 15798–15805. (b) Bronner, S. M.; Mackey, J. L.; Houk, K. N.; Garg, N. K. Steric Effects Compete with Aryne Distortion To Control