This is an open access article published under a Creative Commons Attribution (CC-BY) License, which permits unrestricted use, distribution and reproduction in any medium, provided the author and source are cited.

> ACS AUTHORCHOICE



pubs.acs.org/JACS



Bismuth-Catalyzed Oxidative Coupling of Arylboronic Acids with Triflate and Nonaflate Salts

Oriol Planas, Vytautas Peciukenas, and Josep Cornella*



ABSTRACT: Herein we present a Bi-catalyzed cross-coupling of arylboronic acids with perfluoroalkyl sulfonate salts based on a Bi(III)/Bi(V) redox cycle. An electron-deficient sulfone ligand proved to be key for the successful implementation of this protocol, which allows the unusual construction of $C(sp^2)$ –O bonds using commercially available NaOTf and KONf as coupling partners. Preliminary mechanistic studies as well as theoretical investigations reveal the intermediacy of a highly electrophilic Bi(V) species, which rapidly eliminates phenyl triflate.

unctional groups such as the trifluoromethanesulfonate (triflate, OTf) or the nonafluorobutanesulfonate (nonaflate, ONf) are highly useful moieties when present in organic compounds, especially when attached to a carbon atom (C-OTf or C-ONf).¹ Indeed, $C(sp^2)$ -OTf and $C(sp^2)$ -ONf have been utilized as surrogates of aryl halides (aka pseudohalides) due to their ability to heavily polarize the C-O bond, facilitating oxidative addition by d-block metals. This strategy has been largely exploited with a myriad of combinations of both transition metals and coupling partners, thus placing aryl triflates and nonaflates as routine electrophiles in this large arena.³ From the organometallic standpoint, OTf anions have also many attractive features. The coordinating properties of triflate anions have been a matter of intense debate in the recent literature.⁴ However, it is evident that differently than Ar-Cl, oxidative addition complexes of Ar-OTf would result in a remarkably weaker interaction of the OTf anion and the metal center in solution. Furthermore, in polar and coordinating solvents the OTf anion is generally relegated to the outer sphere, leaving a vacant coordination site (Figure 1A), which has been exploited for a variety of organometallic and coordination purposes.⁵ Yet, the great attributes of OTf anions-highly electronegative, poor nucleophiles and labile ligands-inherently situates them as one of the foulest anions to undergo C-O reductive elimination.

Many examples with high-valent transition metals have been reported to accommodate OTf anions in the primary coordination sphere.⁷ However, reductive elimination primarily occurred at other anionic ligand sites and the M–OTf bond remained unaltered.⁸ During the synthesis of trisubstituted olefins, Gaunt and co-workers suggested that $C(sp^2)$ –OTf bonds could be formed through an unusual reductive elimination from a Cu(III) center,⁹ although further evidence was not provided. Indeed, examples of well-defined transition metal complexes that forge $C(sp^2)$ –OTf bonds still remain elusive. Notwithstanding, the development of a catalytic protocol which enables the formation of Ar–OTf from the corresponding organometallic reagent (Ar–M) and a com-



Figure 1. (A) OTf anions as ligands in transition metal chemistry. (B) Catalytic Ar–OTf formation through a Bi(III)/Bi(V) redox system.

mercially available triflate salt (MOTf) would be highly desirable from both the synthetic and fundamental point of view.

Our group has recently started a program to study the catalytic redox properties of bismuth (Bi) complexes, 10 to facilitate transformations beyond the reactivity of transition

 Received:
 May 15, 2020

 Published:
 June 14, 2020



metals.¹¹ Hence, based on the known oxophilicity of Bi complexes¹² and their ability to bind triflate,¹³ we envisaged that an oxidative protocol based on the redox couple Bi(III)/ Bi(V) could fulfill this synthetic challenge. Indeed, a decade ago Mukaiyama and co-workers demonstrated that C-OTf bonds could be forged from Bi(V) compounds and HOTf, albeit in low yields.¹⁴ Inspired by this early precedent, herein we report on a catalytic oxidative coupling between arylboronic acids and triflate salts to furnish Ar-OTf species (Figure 1B). A rationally designed Bi complex bearing an electron-deficient diarylsulfone ligand unlocks a catalytic redox process and enables the use of triflate (OTf) and nonaflate (ONf) salts as coupling partners. Preliminary mechanistic investigations and theoretical analysis revealed that the $C(sp^2)-O$ bond formation is extremely fast from Bi(V) and is suggested to proceed through a five-membered transition state.

We started our investigations by optimizing the coupling of phenylboronic acid (1a) with NaOTf to generate phenyltriflate (2a) (Table 1). Based on our previous studies on Bi-catalyzed



^aReactions performed at 0.025 mmol of 1a. Yields determined by ¹⁹F NMR using 1-fluoro-4-nitrobenzene as internal standard. ^bIsolated yield of pure material of a reaction performed at 0.3 mmol of 1a.

fluorination, bismines featuring a diarylsulfone backbone (4ac) were selected as catalysts,¹⁵ together with N-fluoro-2,6dichloropyiridinium tetrafluoroborate ([Cl₂pyrF]BF₄) as oxidant. To promote transmetalation, we selected K₂CO₃, as it has been recently demonstrated to be an excellent base for this purpose.¹⁶ In our initial attempts, the unsubstituted bismine catalyst (4a) provided no reactivity toward 2a (entry 1). However, when a CF₃ group was introduced in meta-position to the Bi (4b), an encouraging 11% of 2a was obtained; interestingly, the formation of protodeboronation byproduct 3 was largely suppressed (entry 2). In line with these results, when two CF₃ are introduced in the backbone of the sulfone (4c) the reactivity toward 2a increased to 32%, while the formation of 3 was still largely reduced (entry 3). When K₂CO₃ is replaced by NaF, a reversed trend in the product distribution is observed, substantially favoring undesired 3 (entry 4). Surprisingly, addition of 4 Å molecular sieves (MS) boosted the formation of 2a to 54% yield, while formation of 3 was still minimized (entry 5). Remarkably, when K₂CO₃ was

pubs.acs.org/JACS

replaced by the weaker Na_3PO_4 , nearly quantitative formation of **2a** was achieved (entry 6). The use of 5 Å MS proved crucial to completely suppress the formation of **3**, thus obtaining the desired **2a** in >95% yield (90% isolated) (entry 7). Unfortunately, lower catalyst loadings resulted in poor yields (entry 8).

With the optimal conditions in hand, the scope of the Bicatalyzed $C(sp^2)$ -OTf bond reaction was investigated using a variety of arylboronic acid derivatives (Table 2). The

Table 2. Scope of the Bi-Catalyzed Oxidative Coupling of Arylboronic Acids and Sodium Triflate a



^{*a*}Reaction conditions: 1 (0.3 mmol), NaOTf (0.33 mmol), 4c (0.03 mmol), [Cl₂pyrF]BF₄ (0.33 mmol), Na₃PO₄ (0.6 mmol), and 5 Å MS (120 mg) in CHCl₃ at 60 °C for 16 h. Yields of isolated pure material. ^{*b*}Reaction performed at 90 °C with 2.0 equiv of NaF as base. ^{*c*}Yields determined by ¹⁹F NMR using 1-fluoro-4-nitrobenzene as internal standard. ^{*d*}Reactions performed at 0.025 mmol of the corresponding arylboronic acids. ^{*e*}Reaction performed at 90 °C with 4.0 equiv of Na₃PO₄ as base.

methodology boded well with Me groups in both *para*- (2b) and *ortho*-positions (2c). Remarkably, when the steric encumbrance at the *ortho*-position was further increased, excellent yields of the corresponding triflate were obtained (2d and 2e). Furthermore, the presence of alkyl moieties in other positions of the aryl ring did not affect the reactivity (2f and 2g). The protocol accommodates various functional groups, including ethers (2h and 2i) and halogens (2j, 2k, and 2l), albeit in moderate yields. Arylboronic acids substituted with a trimethylsilyl group (TMS), Ph, or an ester at the *para*position afforded good to excellent yields of the corresponding

aryl triflates (2m-2o). Arylboronic acids bearing unsaturated moieties boded well in this methodology, as exemplified by the presence of alkynyl (2p) and vinyl (2q) groups. In spite of the large variety of arylboronic acids amenable for this transformation, moderate yields were obtained in the presence of certain functionalities. Due to the high reactivity toward oxidation with $[Cl_2pyrF]BF_4$, fluorene derivative 2r was obtained in 38% yield.¹⁵ Substrates bearing strong electronwithdrawing groups such as CF_3 (2s) and reactive carbonyl functionalities at the *para*-position (2t and 2u) also struggled to undergo C–O bond formation, demonstrating some limitations to the scope of this reaction.

Having established a protocol for the successful coupling of NaOTf, we turned our attention to the use of less nucleophilic nonaflate salts as coupling partners. A brief re-examination of the reaction parameters revealed bismine nonaflate 4d as the catalyst of choice to couple arylboronic acids with commercially available KONf.¹⁵ With the optimized conditions shown in Table 3, Ph–ONf (5a) was isolated in a satisfactory

Table 3. Scope of the Bi-Catalyzed Oxidative Coupling of Arylboronic Acids and Potassium Nonaflate^a



^aReaction conditions: 1 (0.3 mmol), KONf (0.33 mmol), 4d (0.03 mmol), $[Cl_2pyrF]BF_4$ (0.33 mmol), Na_3PO_4 (1.2 mmol) and 5 Å MS (120 mg) in CHCl₃ at 60 °C for 16 h. Yields of isolated pure material.

97% yield. The various arylboronic acids scrutinized in the nonaflate synthesis revealed comparable reactivity to NaOTf. Aryl nonaflates containing *ortho*-substituents such as Br (**5b**) and Me (**5c**) were obtained in excellent yields. Furthermore, a TMS moiety can also be accommodated to the protocol (**5d**) as well as unsaturated alkynyl functionalities (**5e**).

The unprecedented catalytic C-OTf and C-ONf bond forming reaction using 4c and 4d led us to explore the operative mechanism governing this transformation. First, we interrogated the transmetalation step between 1a and 4c (Figure 2A). When the reaction was performed in the presence of Na₃PO₄ and 5 Å MS, transmetalation occurred efficiently and 6 was obtained in 86% yield (entry 1). In the absence of base, 6 was also obtained in slightly lower yields (62%, entry 2). In sharp contrast, when the reaction was performed without MS (entry 3), formation of 6 was dramatically reduced (21%). In the absence of both MS and Na₃PO₄, 6 was not detected. These results demonstrate the importance of molecular sieves in this transformation, not only as a dehydrating agent¹⁷ but also as a potential heterogeneous Brønsted base,¹⁸ promoting transmetalation to the Bi(III) center. At this point, the oxidation-reductive elimination sequence from phenylbismine 6 was studied utilizing different oxidants and triflate sources (Figure 2B, top). After oxidizing **6** with XeF_2 to the high-valent Bi(V) difluoride species, ^{10b,15} TMSOTf was added, resulting in a rapid color change from pale to dark yellow. Analysis of the



Figure 2. (A) Study of the transmetalation step: influence of the molecular sieves and the base. (B) Stoichiometric sequence of oxidative addition-reductive elimination. (C) Theoretical analysis of the C-O bond forming step. ^{*a*}Yields determined by ¹H NMR using 1,3,5-trimethoxybenzene as internal standard. ^{*b*}Yields determined by ¹⁹F NMR using 1-fluoro-4-nitrobenzen as internal standard.

reaction crude revealed quantitative formation of **2a**. This result points to the formation of a highly electrophilic Bi(V) intermediate (7a) bearing an OTf moiety, as a consequence of fluoride abstraction by TMSOTf. Indeed, when TMSOTf was added at -41 °C, intermediate 7a could be detected by HRMS (Figure 2B). Furthermore, using [Cl₂pyrF]BF₄ as an oxidant together with NaOTf similar yields for **2a** were obtained (92%, Figure 2B, bottom). It is important to mention that only trace amounts of fluorobenzene were detected, which shows the preferential formation of C-OTf over C-F bonds (*vide infra*).¹⁹ Related intermediates have been previously postulated by Mukaiyama, in the C(sp²)-OTs coupling from Bi(V) intermediates.¹⁴ Based on these experimental results, preliminary theoretical studies were performed to investigate a putative reductive elimination from 7, bearing both a F (7a) or an OTf (7b) as counterions.¹⁵ As shown in Figure 2C, two

possible scenarios were postulated. On one hand, reductive elimination can occur through a three-membered transition state (Figure 2C, pathway a), reminiscent of concerted reductive eliminations performed by d-block elements. Alternatively, reductive elimination could also occur via a five-membered transition state (Figure 2C, pathway b), where two oxygens of the OTf are involved. This latter hypothesis has been previously invoked to explain the selectivity of Bimediated couplings such as α -arylation of phenols²⁰ and Narylation of pyridones,²¹ among other transformations.²² In accordance with these previous reports, our theoretical analysis predicts that the five-membered TS2 is slightly favored over TS1, pointing toward TS2 as the preferable pathway for the C-O bond forming event. NBO analysis on the Bi center also provided additional information about this process.²³ In the case of 7a, the NBO charge on the Bi decreases from 2.17 to 1.84 in TS1 and 1.88 in TS2 and is further reduced to 1.50 in 8. The same trend is observed from 7b.¹⁵ This progressive change in charge at the metal center has been previously observed in high-valent Cu cross-couplings,²⁴ suggesting a concerted reductive elimination through the metal.

Taken these results together, the reaction is proposed to follow the catalytic cycle depicted in Figure 3. Initially, bismine



Figure 3. Postulated mechanism for the Bi-catalyzed oxidative coupling of arylboronic acids and triflate salts.

A undergoes transmetalation (TM) with the corresponding arylboronic acid, thus forming aryl bismine **B**. Subsequently, **B** undergoes formal oxidative addition (OA) with $[Cl_2pyrF]BF_4$, furnishing the proposed high-valent Bi(V) intermediate **C**. Reductive elimination (RE) from **C** delivers the desired aryl triflate with concomitant regeneration of **A**. Due to the structural similarities between OTf and ONf, we believe that a similar mechanism is operating for the coupling of the latter.

In summary, an unprecedented oxidative coupling of arylboronic acids with triflate and nonaflate salts has been developed exploiting the reactivity of the Bi(III)/Bi(V) redox couple. A highly electron-withdrawing diarylsulfone ligand unlocked a catalytic process which proceeds under mild conditions and accommodates various functional groups. The results presented in this study unveil bismuth redox catalysis as a promising tool to perform transformations beyond the scope of transition metals, while mimicking their fundamental organometallic steps.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c05343.

Experimental procedures, analytical data (¹H, ¹⁹F, ¹¹B, and ¹³C NMR, HRMS) for all new compounds, computational results, including Tables (S1–S9) and Figures (S1–S20) (PDF)

AUTHOR INFORMATION

Corresponding Author

Josep Cornella – Max-Planck-Institut für Kohlenforschung, Mülheim an der Ruhr 45470, Germany; o orcid.org/0000-0003-4152-7098; Email: cornella@kofo.mpg.de

Authors

- Oriol Planas Max-Planck-Institut für Kohlenforschung, Mülheim an der Ruhr 45470, Germany; is orcid.org/0000-0003-2038-2678
- Vytautas Peciukenas Max-Planck-Institut für Kohlenforschung, Mülheim an der Ruhr 45470, Germany; orcid.org/0000-0001-7372-9650

Complete contact information is available at: https://pubs.acs.org/10.1021/jacs.0c05343

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

Financial support for this work was provided by Max-Planck-Gesellschaft, Max-Planck-Institut für Kohlenforschung, Fonds der Chemischen Industrie (FCI-VCI). This project has received funding from European Union's Horizon 2020 research and innovation programme under Agreements Nos. 850496 (ERC Starting Grant, J.C.) and 833361 (Marie Skłodowska Curie Fellowship, O.P.). We thank Prof. Dr. A. Fürstner for insightful discussions and generous support. We also thank Dr. Kalishankar Bhattacharyya and Dr. Dimitrios Pantazis for insightful suggestions and support in the computational studies, as well as the analytical department at the MPI-Kohlenforschung for support in the characterization of compounds.

REFERENCES

(1) (a) Hansen, R. L. Perfluoroalkanesulfonate Esters as Alkylating Agents. J. Org. Chem. **1965**, 30, 4322–4324. (b) Stang, P. J.; Hanack, M.; Subramanian, L. R. Perfluoroalkanesulfonic Esters: Methods of Preparation and Applications in Organic Chemistry. Synthesis **1982**, 1982, 85–126. (c) Ritter, K. Synthetic Transformations of Vinyl and Aryl Triflates. Synthesis **1993**, 1993, 735–762. (d) Högermeier, J.; Reissig, H.-U. Nine Times Fluoride can be Good for your Syntheses. Not just Cheaper: Nonafluorobutanesulfonates as Intermediates for Transition Metal-Catalyzed Reactions. Adv. Synth. Catal. **2009**, 351 (17), 2747–2763.

(2) (a) Zeni, G.; Larock, R. C. Synthesis of Heterocycles via Palladium-Catalyzed Oxidative Addition. *Chem. Rev.* **2006**, *106*, 4644–4680. (b) Rosen, B. M.; Quasdorf, K. W.; Wilson, D. A.; Zhang, N.; Resmerita, A.-M.; Garg, N. K.; Percec, V. Nickel-Catalyzed Cross-Couplings Involving Carbon–Oxygen Bonds. *Chem. Rev.* **2011**, *111*, 1346–1416. (c) Bisz, E.; Szostak, M. Iron-Catalyzed C–O Bond Activation: Opportunity for Sustainable Catalysis. *ChemSusChem* **2017**, *10*, 3964–3981.

(3) For representative transition-metal catalyzed C–OTf and C– ONf functionalization reactions, see: (a) Echavarren, A. M.; Stille, J.

K. Palladium-catalyzed coupling of aryl triflates with organostannanes. J. Am. Chem. Soc. 1987, 109, 5478-5486. (b) Ohe, T.; Miyaura, N.; Suzuki, A. Palladium-catalyzed cross-coupling reaction of organoboron compounds with organic triflates. J. Org. Chem. 1993, 58, 2201-2208. (c) Wolfe, J. P.; Buchwald, S. L. Palladium-Catalyzed Amination of Aryl Triflates. J. Org. Chem. 1997, 62, 1264-1267. (d) Rottländer, M.; Knochel, P. Palladium-Catalyzed Cross-Coupling Reactions with Aryl Nonaflates: A Practical Alternative to Aryl Triflates. J. Org. Chem. 1998, 63, 203-208. (e) Anderson, K. W.; Mendez-Perez, M.; Priego, J.; Buchwald, S. L. Palladium-Catalyzed Amination of Aryl Nonaflates. J. Org. Chem. 2003, 68, 9563-9573. (f) Lee, D.-Y.; Hartwig, J. F. Zinc Trimethylsilylamide as a Mild Ammonia Equivalent and Base for the Amination of Aryl Halides and Triflates. Org. Lett. 2005, 7, 1169-1172. (g) Gooßen, L. J.; Rodríguez, N.; Linder, C. Decarboxylative Biaryl Synthesis from Aromatic Carboxylates and Aryl Triflates. J. Am. Chem. Soc. 2008, 130, 15248-15249. (h) Uemura, M.; Yorimitsu, H.; Oshima, K. Cp*Li as a base: application to palladium-catalyzed cross-coupling reaction of aryl-X or alkenyl-X (X = I, Br, OTf, ONf) with terminal acetylenes. Tetrahedron 2008, 64, 1829-1833. (i) Watson, D. A.; Su, M.; Teverovskiy, G.; Zhang, Y.; García-Fortanet, J.; Kinzel, T.; Buchwald, S. L. Formation of ArF from LPdAr(F): Catalytic Conversion of Aryl Triflates to Aryl Fluorides. Science 2009, 325, 1661–1664. (j) Gooßen, L. J.; Linder, C.; Rodríguez, N.; Lange, P. P. Biaryl and Aryl Ketone Synthesis via Pd-Catalyzed Decarboxylative Coupling of Carboxylate Salts with Aryl Triflates. Chem. - Eur. J. 2009, 15, 9336-9349. (k) Shekhar, S.; Dunn, T. B.; Kotecki, B. J.; Montavon, D. K.; Cullen, S. C. A General Method for Palladium-Catalyzed Reactions of Primary Sulfonamides with Aryl Nonaflates. J. Org. Chem. 2011, 76, 4552-4563. (1) Si, T.; Li, B.; Xiong, W.; Xu, B.; Tang, W. Efficient cross-coupling of aryl/alkenyl triflates with acyclic secondary alkylboronic acids. Org. Biomol. Chem. 2017, 15, 9903-9909.

(4) Krossing, I.; Raabe, I. Noncoordinating Anions—Fact or Fiction? A Survey of Likely Candidates. *Angew. Chem., Int. Ed.* **2004**, 43, 2066–2090.

(5) (a) Hartwig, J. F. Organotransition metal chemistry: From bonding to catalysis; University Science Books: Mill Valley, CA, 2010.
(b) Maleckis, A.; Sanford, M. S. Facial Tridentate Ligands for Stabilizing Palladium(IV) Complexes. Organometallics 2011, 30, 6617–6627. (c) Racowski, J. M.; Gary, J. B.; Sanford, M. S. Carbon(sp³)–Fluorine Bond-Forming Reductive Elimination from Palladium(IV) Complexes. Angew. Chem., Int. Ed. 2012, 51, 3414–3417.

(6) Dhakal, B.; Bohé, L.; Crich, D. Trifluoromethanesulfonate Anion as Nucleophile in Organic Chemistry. *J. Org. Chem.* **2017**, *82*, 9263– 9269.

(7) (a) Lawrance, G. A. Coordinated trifluoromethanesulfonate and fluorosulfate. *Chem. Rev.* 1986, 86, 17–33. (b) Beck, W.; Suenkel, K. Metal complexes of weakly coordinating anions. Precursors of strong cationic organometallic Lewis acids. *Chem. Rev.* 1988, 88, 1405–1421.
(c) Ball, N. D.; Kampf, J. W.; Sanford, M. S. Aryl–CF₃ Bond-Forming Reductive Elimination from Palladium(IV). *J. Am. Chem. Soc.* 2010, 132, 2878–2879. (d) Lyons, T. W.; Sanford, M. S. Palladium-Catalyzed Ligand-Directed C–H Functionalization Reactions. *Chem. Rev.* 2010, 110, 1147–1169. (e) Canty, A. J. *Higher Oxidation State Organopalladium and Platinum Chemistry*; Springer-Verlag: Berlin, Heidelberg, Germany, 2011.

(8) (a) Bour, J. R.; Camasso, N. M.; Sanford, M. S. Oxidation of Ni(II) to Ni(IV) with Aryl Electrophiles Enables Ni-Mediated Aryl– CF₃ Coupling. J. Am. Chem. Soc. **2015**, 137, 8034–8037. (b) Camasso, N. M.; Sanford, M. S. Design, synthesis, and carbon-heteroatom coupling reactions of organometallic nickel(IV) complexes. *Science* **2015**, 347, 1218–1220. (c) Canty, A. J.; Ariafard, A.; Camasso, N. M.; Higgs, A. T.; Yates, B. F.; Sanford, M. S. Computational study of $C(sp^3)$ –O bond formation at a Pd^{IV} centre. *Dalton Trans* **2017**, *46*, 3742–3748. (d) Nebra, N. High-Valent Ni^{III} and Ni^{IV} Species Relevant to C–C and C–Heteroatom Cross-Coupling Reactions: State of the Art. *Molecules* **2020**, *25*, 1141. (9) Suero, M. G.; Bayle, E. D.; Collins, B. S. L.; Gaunt, M. J. Copper-Catalyzed Electrophilic Carbofunctionalization of Alkynes to Highly Functionalized Tetrasubstituted Alkenes. *J. Am. Chem. Soc.* **2013**, *135*, 5332–5335.

(10) (a) Wang, F.; Planas, O.; Cornella, J. Bi(I)-Catalyzed Transfer-Hydrogenation with Ammonia-Borane. J. Am. Chem. Soc. 2019, 141, 4235–4240. (b) Planas, O.; Wang, F.; Leutzsch, M.; Cornella, J. Fluorination of arylboronic esters enabled by bismuth redox catalysis. Science 2020, 367, 313–317.

(11) (a) Power, P. P. Main-group elements as transition metals. Nature 2010, 463, 171–177. (b) Weetman, C.; Inoue, S. The Road Travelled: After Main-Group Elements as Transition Metals. ChemCatChem 2018, 10, 4213–4228. (c) Melen, R. L. Frontiers in molecular p-block chemistry: From structure to reactivity. Science 2019, 363, 479–484. (d) Janssen-Müller, D.; Oestreich, M. Transition-Metal-Like Catalysis with a Main-Group Element: Bismuth-Catalyzed C-F Coupling of Aryl Boronic Esters. Angew. Chem., Int. Ed. 2020, 59, 8328–8330.

(12) (a) Stewart, C. A.; Calabrese, J. C.; Arduengo, A. J. Synthesis and structure of the first 20-Bi-9 system: a discrete nine-coordinate 20-electron bismuth. J. Am. Chem. Soc. 1985, 107, 3397-3398.
(b) Boyer, B.; Keramane, E. M.; Montero, J.-L.; Roque, J.-P. BiCl₃: An Efficient Agent for Selective Chlorination of Alcohols or for Halogen Exchange Reaction. Synth. Commun. 1998, 28, 1737-1741.
(c) Matano, Y.; Nomura, H. Dimeric Triarylbismuthane Oxide: A Novel Efficient Oxidant for the Conversion of Alcohols to Carbonyl Compounds. J. Am. Chem. Soc. 2001, 123, 6443-6444. (d) Yin, S.-F.; Maruyama, J.; Yamashita, T.; Shimada, S. Efficient Fixation of Carbon Dioxide by Hypervalent Organobismuth Oxide, Hydroxide, and Alkoxide. Angew. Chem., Int. Ed. 2008, 47, 6590-6593. (e) Raţ, C. I.; Silvestru, C.; Breunig, H. J. Hypervalent organoantimony and -bismuth compounds with pendant arm ligands. Coord. Chem. Rev. 2013, 257, 818-879.

(13) For selected recent examples of Bi-OTf species, see: (a) Tschersich, C.; Hoof, S.; Frank, N.; Herwig, C.; Limberg, C. The Effect of Substituents at Lewis Acidic Bismuth(III) Centers on Its Propensity to Bind a Noble Metal Donor. Inorg. Chem. 2016, 55, 1837-1842. (b) Kannan, R.; Kumar, S.; Andrews, A. P.; Jemmis, E. D.; Venugopal, A. Consequence of Ligand Bite Angle on Bismuth Lewis Acidity. Inorg. Chem. 2017, 56, 9391-9395. (c) Ritschel, B.; Poater, J.; Dengel, H.; Bickelhaupt, F. M.; Lichtenberg, C. Double CH Activation of a Masked Cationic Bismuth Amide. Angew. Chem., Int. Ed. 2018, 57, 3825-3829. (d) Ramler, J.; Poater, J.; Hirsch, F.; Ritschel, B.; Fischer, I.; Bickelhaupt, F. M.; Lichtenberg, C. Carbon monoxide insertion at a heavy p-block element: unprecedented formation of a cationic bismuth carbamoyl. Chem, Sci. 2019, 10, 4169-4176. (e) Ramler, J.; Hofmann, K.; Lichtenberg, C. Neutral and Cationic Bismuth Compounds: Structure, Heteroaromaticity, and Lewis Acidity of Bismepines. Inorg. Chem. 2020, 59, 3367-3376.

(14) (a) Imachi, S.; Mukaiyama, T. Oxidative Coupling of Carbonyl Compounds by Using Pentavalent Biphenyl-2,2'-ylenebismuth Reagents. *Chem. Lett.* 2007, 36, 718–719. (b) Sakurai, N.; Mukaiyama, T. A New Preparative Method of Aryl Sulfonate Esters by Using Cyclic Organobismuth Reagents. *Heterocycles* 2007, 74, 771–790.

(15) See Supporting Information for further details.

(16) Jurrat, M.; Maggi, L.; Lewis, W.; Ball, L. T. Modular bismacycles for the selective C–H arylation of phenols and naphthols. *Nat. Chem.* **2020**, *12*, 260–269.

(17) West, M. J.; Fyfe, J. W. B.; Vantourout, J. C.; Watson, A. J. B. Mechanistic Development and Recent Applications of the Chan–Lam Amination. *Chem. Rev.* **2019**, *119*, 12491–12523.

(18) Steinhoff, B. A.; King, A. E.; Stahl, S. S. Unexpected Roles of Molecular Sieves in Palladium-Catalyzed Aerobic Alcohol Oxidation. *J. Org. Chem.* **2006**, *71*, 1861–1868.

(19) Furuya, T.; Kamlet, A.; Ritter, T. Catalysis for fluorination and trifluoromethylation. *Nature* **2011**, 473, 470–477.

(20) Barton, D. H. R.; Bhatnagar, N. Y.; Finet, J.-P.; Motherwell, W. B. Pentavalent organobismuth reagents. Part VI. Comparative

migratory aptitudes of aryl groups in the arylation of phenols and enols by pentavalent bismuth reagents. *Tetrahedron* **1986**, *42*, 3111–3122.

(21) Ikegai, K.; Mukaiyama, T. Synthesis of N-Aryl Pyridin-2-ones via Ligand Coupling Reactions Using Pentavalent Organobismuth Reagents. *Chem. Lett.* **2005**, *34*, 1496–1497.

(22) (a) Finet, J.-P. Ligand Coupling Reactions with Heteroatomic Compounds; Elsevier: Oxford, 1998. (b) Finet, J.-P. Arylation reactions with organobismuth reagents. Chem. Rev. **1989**, 89, 1487–1501.

(23) (a) Reed, A. E.; Curtiss, L. A.; Weinhold, F. Intermolecular interactions from a natural bond orbital, donor-acceptor viewpoint. *Chem. Rev.* **1988**, *88*, 899–926. (b) Weinhold, F.; Landis, C. R. Valency and Bonding: A Natural Bond Orbital Donor-Acceptor Perspective; Cambridge University Press: Cambridge, United Kingdom, 2005.

(24) Paeth, M.; Tyndall, S. B.; Chen, L.-Y.; Hong, J.-C.; Carson, W. P.; Liu, X.; Sun, X.; Liu, J.; Yang, K.; Hale, E. M.; Tierney, D. L.; Liu, B.; Cao, Z.; Cheng, M.-J.; Goddard, W. A.; Liu, W. Csp³–Csp³ Bond-Forming Reductive Elimination from Well-Defined Copper(III) Complexes. J. Am. Chem. Soc. **2019**, *141*, 3153–3159.