CHEMISTRY

Templated deprotonative metalation of polyaryl systems: Facile access to simple, previously inaccessible multi-iodoarenes

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The development of new methodologies to affect non–ortho–functionalization of arenes has emerged as a globally important arena for research, which is key to both fundamental studies and applied technologies. A range of simple arene feedstocks (namely, biphenyl, meta-terphenyl, para-terphenyl, 1,3,5-triphenylbenzene, and biphenylene) is transformed to hitherto unobtainable multi-iodoarenes via an s-block metal sodium magnesiate templated deprotonative approach. These iodoarenes have the potential to be used in a whole host of high-impact transformations, as precursors to key materials in the pharmaceutical, molecular electronic, and nanomaterials industries. To prove the concept, we transformed biphenyl to 3,5-bis(N-carbazolyl)-1,1′-biphenyl, a novel isomer of 4,4′-bis(N-carbazolyl)-1,1′-biphenyl (CPB), a compound which is currently widely used as a host material for organic light-emitting diodes.

INTRODUCTION

Until 1938, organometallic reagents were considered as exotic rarities, commanding little application in synthetic chemistry (1). However, the discoveries by Gilman and Bebb (2) and Wittig et al. (3) that alkyllithium reagents could be used to selectively metatalate anisole (and other heterosubstituted arenes) at an ortho position opened the doors to the first major utilization of organometallics in synthesis. This discovery was later coined directed ortho–metalation (DoM) (4), and from this juncture, organometallic reagents were seen as “indispensable tools in chemical synthesis” (1). DoM can be used to install new functionalities on arenes, thus allowing the construction of new molecules (5–7), and relies on converting a relatively inert C–H bond in an organic molecule into a more reactive and versatile C–metal bond by treating it with a metalating reagent (normally an organolithium reagent). The beauty of the DoM reaction is it efficiently and selectively metalates the C–H bond adjacent to a directing group (DG), which promotes reaction due to its dual coordination/inductive roles. Since the pioneering works of Gilman and Bebb (2) and Wittig et al. (3), the DoM concept has been advanced using monometallic (8–12) and bi-metallic (13–19) systems.

In his influential text, Schlosser stated, “deviations from the ortho-rule do exist but are scarce” (1). Enabling activation of arene C–H bonds at positions other than ortho has become a high interest and impact area of research, with the goal of allowing easy access to key pharmaceuticals, agrochemicals, and novel materials. In the main, new methodologies involve transition metal C–H activation chemistry (20–28). In 2014, we exploited s-block organometallic chemistry and reported directed ortho–meta– and meta–meta–dimetalations using a sodium magnesiate reagent [Na4Mg2(TMPS)2(Bu)] (TMP is 2,2,6,6-tetramethylpiperidide) as an alkyl base (29). It functions in a templated manner, allowing double metalations to occur, and has been shown to exhibit regioselective 2,5-double alkyl basicity toward a range of substituted arenes, including amides and carbamates and 3,5-double basicity toward anilines, culminating in the isolation of a group of heterometallic inverse crown molecules (30). Here, we use the template metalation strategy to functionalize unactivated multiaryl compounds (non-DoM substrates) that lack acidity due to the absence of functional groups, regioselectively at meta sites. The biphenyl motif (Fig. 1A) is an important building block in the pharmaceutical, agrochemical, and polymer/materials sciences. It is present in telmisartan, valsartan, and oriatavancin and is a key component of molecules for organic light-emitting devices, such as N,N′-di(1-naphthyl)-N,N′-diphenyl-(1,1′-biphenyl)-4,4′-diamine (NPD) and 4,4′-bis(N-carbazolyl)-1,1′-biphenyl (CPB). Biphynyl-based hydrocarbons, such as terphenyls and 1,3,5-triphenylbenzene (tpb) (Fig. 1B), are also cores for molecules used in the devices and for the synthesis of nanomaterials, such as graphene/graphene nanoribbons (31). Novel methods to improve the preparation of functionalized multiarenes for these applications are continuously being sought. Reflecting the importance of developing new methods to access novel polyaryl systems, Suzuki et al. (32) have recently reported the elegant synthesis of a new class of “multitalented” hexaarylbenzenes (33) that contain five or six different arene substituents on a single benzene molecule.

RESULTS AND DISCUSSION

We targeted the templated metalation of simple polyaryl compounds (namely, biphenyl, para-terphenyl, meta-terphenyl, tpb, and antiaromatic biphenylene) using I and subsequently transformed the organometallic intermediates into novel versatile organic precursors (34). Because of their lack of DGs (hence, the absence of docking site and strong acidifying group), the substrates studied are difficult to metalate using conventional s-block organometallics, unless a Lewis basic functionality activates them. Indings that this work may be successful stemmed from the fact that we had previously shown that I can be used to doubly metatalate naphthalene at the 1 and 4 positions regioselectively, which can be subsequently converted to 1,4-diodonaphthalene (35).

Substituted biphenyls are often prepared by coupling reactions (36). In terms of metalation chemistry, both rings of biphenyl can be singly deprotonated using two equivalents of n-butyllithium/TMEDA (N,N,N′,N′-tetramethylethlenediamine) (37). Synthetically useful yields of monolithiobiphenyls can be obtained using lithium-halogen exchange reactions (38, 39). Introducing a wholly new situation,
by treating biphenyl with a methylcyclohexane solution of 1 in a 1:1 molar ratio, it is possible to regioselectively dimagnesiate biphenyl at the 3 and 5 positions of a single phenyl ring, as evidenced by nuclear magnetic resonance (NMR) spectroscopic analysis of the reaction mixture.

\[ \text{[Na}_4\text{Mg}_2(\text{TMP})_6(3,5\text{-biphenyl-di-ide})] \]

was isolated in 70% yield and characterized by x-ray crystallography (Fig. 2A). When two or more equivalents of 1 were used, 2 was again observed, presumably because of the high steric protection afforded to the second phenyl ring, disfavoring further reactivity. When biphenyl is treated with one or two equivalents of the monometallic reagents NaTMP or TMPMg\text{Bu}, intact unreacted biphenyl was obtained (Supplementary Materials). The key bond parameters of 2 (Supplementary Materials) resemble those of other comparable inverse crown molecules (29,40). When 2 is reacted with an excess of iodine/tetrahydrofuran (THF) solution, 3,5-diiodobiphenyl 3 can be prepared in 64% isolated yield (Fig. 3, i).

The next arene molecules studied were meta- and para-terphenyl. Terphenylide anions—prepared via lithium-halogen exchange reactions of 1-halo-2,5-diphenylbenzenes and n-butyllithium—are useful ligands in coordination chemistry because they provide a pocket, which protects the metal from forming oligomers despite occupying only one coordination site. Power exploited this special chemistry using a bulky meta-terphenyl, 2,6-bis-[(2,6-diiisopropyl)phenyl]phenyl to isolate the first quintuple bond between two metals, namely, two Cr(I) cations (41). It appears that the parent terphenyls have not thus far been deprotonated using any base. Focusing on para-terphenyl, when treated with an equivalent of 1 in methylcyclohexane solution, NMR spectroscopic analysis showed dimetalation of the arene at 3,5 positions of a single phenyl ring, likely producing [Na\text{Mg}_2(\text{TMP})_6(3,5\text{-para-terphenyl-di-ide})] 4 (Supplementary Materials). Because of its high solubility, 4 was isolated in low to moderate yield (26%). In situ mixture was reacted with iodine/THF solution and successfully converted to 3,5-diiodo-para-terphenyl 5 in 59% yield (Fig. 3, ii). As well as characterization by NMR spectroscopy, good quality crystals of 5 were successfully grown from an ethanol solution and were characterized by x-ray crystallography (Supplementary Materials). Increasing the base/substrate ratio (to 2:1) allowed further metalation of the para-terphenyl ligand to occur presumably because of the greater distance between the two “outer” phenyl rings (when compared with biphenyl), as evidenced by NMR spectroscopy. Crystals were obtained, which after x-ray analysis showed fourfold metalation of para-terphenyl culminating in the isolation of \([\text{[Na}_4\text{Mg}_2(\text{TMP})_6(3,3″,5,5″\text{-para-terphenyl-tetra-ide})]} \) 6 in 69% yield (Fig. 2B). The outer rings of the para-terphenyl have each been doubly metalated, and each affected “arene dianion” is hosted by separate inverse crown rings. The key metrics of the inverse crown rings in 6 are essentially identical to those in the benzyl derivative 2. Complex 6 represents the first example of a molecule, which contains two inverse crowns within its molecular structure. Previously, it has been shown that metallocenes can also be tetradeprotonated; however, these tetranions are accommodated within a single 16-membered inverse crown ring (42,43). When isolated 6 is reacted with iodine/THF, NMR spectroscopy reveals that it is transformed to 3,3″,5,5″-tetraiodo-para-terphenyl 7 in 51% yield (Fig. 3, ii). When an in situ mixture of 6 is reacted with excess iodine/THF to prepare 7, a slightly lower yield (44%) was obtained.

The chemistry observed with meta-terphenyl (to produce 8 to 11) essentially mirrors that of 4 to 7, that is, di- and tetrametalated.

Fig. 1. Prevalence of the biphenyl skeleton in pharmaceutical compounds and organic light-emitting diode materials. (A) Biphenyl and the structural prevalence of its framework in pharmaceutical agents and host materials for organic light-emitting diodes (OLED). (B) Biphenyl-based polyaryl hydrocarbons.
compounds \([\{\text{Na}_4\text{Mg}_2(\text{TMP})_6\}_2(3,5,\text{-meta-terphenyl-di-ide})\}\text{8} \) and \([\{\text{Na}_4\text{Mg}_2(\text{TMP})_6\}_2(3,3',5,5'\text{-meta-terphenyl-tetra-ide})\}\text{10}\), respectively, were obtained and characterized by NMR spectroscopy, as well as their respective di- and tetraiodo derivatives \(3,5\text{-diiodo-meta-terphenyl 9}\) (54%) and \(3,3',5,5'\text{-tetraiodo-meta-terphenyl 11}\) (40%).

Turning to tpb, it has emerged in recent years as an important ligand, formally allowing stabilization of Ca(I) cations (44). C–H metalation of tpb is rare, and complexes derived from it generally emerge from the use of a pre-halo-substituted (normally at the central \(C_6\) ring) reagent. One example of where a genuine C–H metalation is observed is when 1-bromo-2,4,6-triphenylbenzene is reacted with 2 molar equivalents of \(n\)-butyllithium (45). The first equivalent of alkyl-lithium induces a metal-bromine exchange, and the second metalates an interannular peri-C–H bond, formally giving rise to a dianion of triphenylbenzene (46). Structurally, tpb seemed an intriguing reagent to metalate. If the reaction was in keeping with the aforementioned results, it could be envisaged that di-, tetra-, or perhaps hexadeprotonation would be possible. In reality, when the arene is reacted with 1 to 3 molar equivalents of 1, the sole isolated product is the tetra-deproto-nated variant \([\{\text{Na}_4\text{Mg}_2(\text{TMP})_6\}_2(3,3',5,5'\text{-1,3',5' triphenylbenzene}-tetra-ide})\]\text{12}\ (Fig. 2C). A twofold deprotonation of tpb was not detected, perhaps because of the more acidic nature of tpb (in comparison with \textit{meta-terphenyl}) due to the presence of the third phenyl group withdrawing electron density from the other two phenyl rings. When \text{6}\ is compared with \text{12}, there is a marked difference in the positions and orientations of the inverse crown rings (Fig. 2B versus Fig. 2C). In \text{6}, they adopt a mutual convex orientation residing in a position toward the periphery of the metalated phenyl rings. Likely because of the extra steric demand of the additional phenyl group in tpb, the inverse crowns adopt a mutual concave orientation in \text{12}, residing in a position more toward the center of the metalated rings. These pronounced structural differences highlight the conformational flexibility of the template, allowing spatially distinct arenes to be accommodated. By reacting \text{12} with an excess of iodine/THF, it can be smoothly transformed to its tetraiodo relative \(3,3',5,5'\text{-tetraiodo-1,3',5' triphenylbenzene 13}\) in 61% yield.

Finally, we turned our attention to the prototypical antiaromatic compound biphenylene. Again, reports detailing the direct metalation of biphenylene are very scarce. When reacted with \textit{n}- or \textit{t}-butyllithium and quenched with electrophiles, the major species observed are the 1-substituted biphenylenes (47). When biphenylene is reacted with a methylcyclohexane solution of 1, it was doubly metalated; however, metalation occurred simultaneously at the 1 and 4 positions to yield \([\{\text{Na}_4\text{Mg}_2(\text{TMP})_6\}_2(1,4\text{-biphenylene-di-ide})]\) \text{14}\ (Fig. 2D). This metalation pattern is identical to that obtained with naphthalene, presumably because of the similar fused-ring nature of the substrates (thus, each arene ring is initially disubstituted) (35). Complex \text{14} can be converted to 1,4-diodobiphenylene \text{15} in high yield (72%) by reaction with iodine/THF.

Organometallic species \text{2}, \text{4}, \text{6}, \text{8}, \text{10}, \text{12}, and \text{14} are soluble in [\text{D}_{12}]\text{cyclohexane}, allowing multinuclear NMR spectroscopy to be performed. A key finding is that the solid-state structures appear to be maintained in [\text{D}_{12}]\text{cyclohexane solution} (Supplementary Materials). Iodinated organic molecules \text{3}, \text{5}, \text{7}, \text{9}, \text{11}, \text{13}, and \text{15} are structurally simple; however, surprisingly, this study appears to represent their first synthetic routes, isolation, and full characterization. Access to these molecules has perhaps been hindered because most of polyaryl
species are prepared via coupling reactions [for example, Suzuki-Miyaura cross-coupling (34)], so the C–I bonds in the potential synthons are themselves susceptible to coupling chemistry, thus it is difficult to prevent undesired overcoupling from occurring. The methodology presented avoids transition metal coupling chemistry, thus the iodine atoms remain intact on the arene rings, allowing further exploitation. Given the importance of iodoarenes as coupling partners in C–C and C–heteroatom bond formation chemistry (48, 49) it is likely that these organic molecules will act as vital building blocks for the synthesis of a whole host of new functional materials. As proof of concept and illustrating the synthetic applicability of our approach, we have prepared a novel isomer of the aforementioned material CPB by converting biphenyl to 3,5-bis(N-carbazolyl)-1,1′-biphenyl 16 in high yield (81%; Fig. 4) via copper-catalyzed Ullmann-type coupling of carbazole with 3.
Fig. 4. Utilization of 1,3-diodobiphenyl for Ullmann coupling reaction with carbazole for the preparation of 16.

MATERIALS AND METHODS
General methods and chemicals
All reactions and manipulations were performed under a protective atmosphere of dry pure argon gas using standard Schlenk techniques unless otherwise stated. n-Hexane was dried by heating it to reflux over sodium benzenophene ketyl and used freshly distilled under nitrogen atmosphere. Methyleneoxohexane was distilled over sodium metal under nitrogen and then stored with activated 4 Å molecular sieves under argon before use. Anhydrous N,N′-dimethylformamide (DMF) was purchased from Sigma-Aldrich and used without further purification. [D12]cyclohexane and [D6]benzene were degassed and stored over activated 4 Å molecular sieves under argon before use. 2,2,6,6-Tetramethylpiperidine [TMP(H)] was purchased from Merck KGaA and stored over activated molecular sieves (4 Å) in an argon-filled Schlenk tube for 30 min before use. Copper iodide, carbazole, potassium carbonate, 18-crown-6, sodium thiosulfate, iodine, and hydrous magnesium sulfate, ammonium chloride, and sodium chloride were purchased from Sigma-Aldrich and titrated before use. Copper iodide, carbazole, potassium carbonate, 18-crown-6, sodium thiosulfate, iodine, anhydrous magnesium sulfate, ammonium chloride, and sodium chloride were purchased from Sigma-Aldrich and used as received.

Preparation of metalating reagent 1
Solutions of 1 (0.05 and 0.066 M in methycyclohexane) were prepared in situ according to literature methods (29, 35). Detailed procedures are given for the preparation of 2 to 17 in the Supplementary Materials.

General synthesis of organometallic reagents 2, 4, 6, 8, 10, 12, and 14
In an argon-filled Schlenk tube, freshly prepared "BuNa (640 mg, 8 mmol) was suspended in methycyclohexane (24 ml), and TMP(H) (2.04 ml, 12 mmol) was then added via syringe to give NaTMP as a pale yellow suspension, which was stirred for 30 min at ambient temperature. Commercial "Bu2Mg (4 ml, 1 M solution in n-heptane, 4 mmol) was then added via syringe to give a 0.066 M pale yellow solution of 1 in methycyclohexane. The metalating reagent 1 was stirred for 30 min at ambient temperature before use. Then, the appropriate polyarene was added, and the resulting reaction mixture was heated at the appropriate temperature. Where possible, crystals suitable for x-ray diffraction were grown (see full details in the Supplementary Materials).

General synthesis of organic compounds 3, 5, 7, 9, 11, 13, and 15
The corresponding organic substrate (biphenyl, para-terphenyl, meta-terphenyl, tpb, and diphenylmethane; 1 mmol each) was treated with an in situ solution of 1 in methycyclohexane prepared according to the synthetic organometallic protocols described in the Supplementary Materials for 2, 4, 6, 8, 10, 12, and 14. The 0.05 M solutions of 1 in methycyclohexane (20 ml) were used for the in situ preparation of 2, 4, 8, and 14, whereas the 0.066 M solutions of 1 in methycyclohexane (30 ml) were used for the in situ preparation of 6, 10, and 12. All reactions were stirred at different temperatures and for different times and in situ monitored by NMR spectroscopy in either [D12]cyclohexane or [D6]benzene until Magnusification of the corresponding aromatic substrate was achieved to give the corresponding organometallic derivatives (table S1). After the in situ metalation step was achieved, the reaction mixture was cooled to −78°C in a dry ice/acetone bath for 30 min and then added (either via cannula or disposable air-tight syringe) to an iodine solution in THF at
the same temperature. For the preparation of 3, 5, 9, and 15, 10 ml of a 1 M solution of iodine in THF was used, whereas for 7, 11, and 13, 20 ml of a 1 M solution of iodine in THF was used. The resulting reaction mixtures were stirred at −78°C for at least 3 hours and then allowed to slowly warm up to ambient temperature for a period of 24 hours. Saturated aqueous NH₄Cl solution (−10 ml) was then added, followed by saturated aqueous Na₂SO₄ solution (−10 ml) until bleaching occurred. The crude mixture was extracted with EtOAc (3 × 20 ml), and the combined organic layers were washed with brine (20 ml) and dried over anhydrous MgSO₄. The solvent was removed in a rotary evaporator, and the crude product was purified by flash column chromatography (silica gel and n-hexane/ethyl acetate) to yield the pure title compound. Alternatively, 7 can be prepared in 51% yield by reacting 6 (1 mmol) with an iodine/THF solution (20 mmol iodine/20 ml THF). The organic derivatives 3, 5, 7, 13, and 15 can be crystallized from either MeOH, EtOH, or EtOAc. Compounds 7, 11, and 13 exhibit low solubility in commonly used organic solvents, which hinders their isolation and purification in the scale used in their preparations.

Synthesis of 16
In an argon-filled Schlenk tube, 3,5-diodo-1,1′-biphenyl (150 mg, 0.37 mmol) was mixed with carbazole (140 mg, 0.83 mmol), potassium carbonate (114 mg, 0.83 mmol), copper(I) iodide (32 mg, 0.17 mmol), and 18-crown-6 (30 mg, 0.11 mmol). Then, anhydrous DMF was added (5 ml), and the reaction mixture was refluxed for 24 hours under argon atmosphere. After cooling down to ambient temperature, the reaction mixture was filtered through a short pad of Celite, and the solvent was removed under vacuum. The crude product was purified by flash column chromatography (silica gel and n-hexane/ethyl acetate) to form 16 as a pale yellow solid (0.145 g, 81%). Compound 16 can be crystallized from ethyl acetate by slow evaporation to give colorless block-like crystals suitable for an X-ray diffraction study (see the Supplementary Materials for NMR spectroscopic and mass spectral data).

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/3/6/e1700832/DC1
Detailed experimental procedures
Materials and methods
Synthetic procedures
X-ray crystallography
Supplementary text
NMR spectra
fig. 51. Sections of the 1H NMR (400.1 MHz; [D₆]benzene, 300 K) spectra showing the nonaromatic resonances for biphenylylene (top; black), reaction mixture containing two conformers of 14 (middle; blue), and isolated [Na₄Mg₂(TMPP)₂(3,3′,5′-triphenyltetra-ide)] 14 (bottom; major conformer).
fig. 52. 1H,13C-HMQC NMR (400.1 MHz; [D₆]benzene, 300 K) spectrum of [Na₄Mg₂(TMPP)₂(3,3′,5′-triphenyltetra-ide)] showing the cross peaks for the aromatic resonances.
fig. 53. 1H,13C-COSY NMR (400.1 MHz; [D₆]benzene, 300 K) spectrum of [Na₄Mg₂(TMPP)₂(3,3′,5′-triphenyltetra-ide)] showing the cross peaks for the aromatic resonances.
REFERENCES AND NOTES


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Author contributions: A.J.M.-M. carried out most of the experimental work, drafted most of the Supplementary Materials, and contributed ideas to the project. A.R.K. and I.D.H.O. ensured that the x-ray crystallography work was carried out to high level and contributed to the drafting of the manuscript. B.J.F. and S.J. carried out the initial experimental work.

C.T.O. conceived and developed the project and drafted the manuscript. Competing interests: The authors declare that they have no competing interests.

Data and materials availability: X-ray crystallographic data for compounds 2, 3, 5, 6, 12, 13, 14, 15, and 16 are freely available at the Cambridge Crystallographic Data Centre (CCDC 1516422-30) via http://www.ccdc.cam.ac.uk/data_request/cif. The research data associated with this paper are openly available at http://dx.doi.org/10.15129/b1d7b580-838c-4621-ad43-fccd2579024d. All other data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

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