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Planarity Is Not Plain: Closed- vs Open-Shell Reactivity of a Structurally Constrained, Doubly Reduced Arylborane toward Fluorobenzenes

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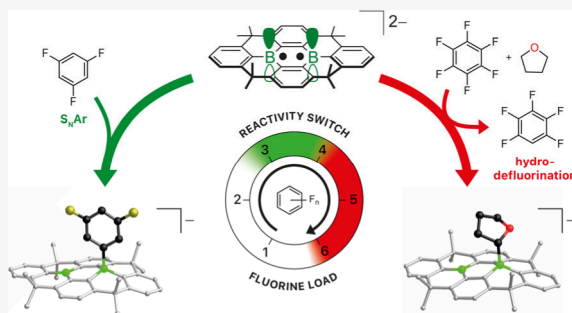
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ABSTRACT: The ability to activate small molecules is imparted to 9,10-dihydro-9,10-diboraanthracenes (DBAs) through the injection of two electrons. We report on the activation of fluorobenzenes $C_6F_nH_{6-n}$ by the doubly reduced, structurally constrained DBA $[1]^{2-}$ in THF (n : 1,3,4,5,6). Compound **1** is a 9,10-diphenyl DBA, forced into planarity by methylene bridges between the phenyl substituents and the DBA core. This rigidity results in enhanced stability under ambient conditions and an elevated planar-to-pyramidal reorganization energy upon boron tetracoordination, unlocking new reactivity. The dianion salts $M_2[1]$ were synthesized in excellent yields by stirring neutral **1** with alkali metals M in THF (M : Li, Na, K); comproportionation of $Li_2[1]$ with **1** generates the blue radical salt $Li[1]$, characterized by EPR spectroscopy and X-ray diffraction. While $Li_2[1]$ is inert toward C_6FH_5 up to 120 °C, it reacts with 1,3,5- $C_6F_3H_3$ at 100 °C to yield a $B(sp^2)/B(sp^3)$ adduct with a difluorophenyl ligand ($Li[2]$). Treatment of $Li_2[1]$ with 1 eq. of C_6F_5H or C_6F_6 induces selective monohydrodefluorination, occurring in parallel with the formation of a unique $B(sp^2)/B(sp^3)$ tetrahydrofuran-2-yl adduct ($Li[3]$). The three isomers of $C_6F_4H_2$ represent intermediate cases, where the competition between trifluorophenyl- and tetrahydrofuran-2-yl-adduct formation is governed by the relative positions of the F substituents and the nature of the counteranion (M^+ : Li^+ , K^+). Through experimental and quantum-chemical studies, we unveil the underlying reaction mechanisms and show that $Li_2[1]$ acts either as a B-centered nucleophile in an S_NAr -type conversion (low benzene fluorination) or as a reducing agent in a single-electron transfer/H atom abstraction sequence (high benzene fluorination).



INTRODUCTION

Despite scattered earlier examples,^{1–5} the advent of frustrated Lewis pairs (FLPs)^{6–8} marked the pivotal paradigm shift, after which element–element-bond activation was no longer considered the exclusive domain of transition-metal complexes. More recently, it has also been demonstrated that the FLP concept can be complemented by moving on to subvalent p-block compounds.^{9–17} In this regard, extended π -electron systems containing embedded trigonal-planar-coordinated boron atoms are particularly promising platforms due to their high affinity for injected electrons and ability to delocalize charge.^{18–22} The 9,10-dihydro-9,10-diboraanthracenes (DBAs) that our group has been primarily employing incorporates two closely spaced boron atoms, enabling their cooperative interaction. In the past few years, we disclosed that the doubly reduced forms $[R_2\text{-DBA}]^{2-}$ readily activate a number of small molecules,²³ among them $H-H$,^{24,25} $H-Bpin$,²⁶ $H-CCtBu$,²³ $tBuO-OtBu$,²⁷ and $F-Ar^F$ (best performing B-bonded substituents R : H, Me; Ar^F : fluoroaryl).²⁸

Taking the nonpolar substrate H_2 as the model system, the activation mechanism was elucidated through quantum-

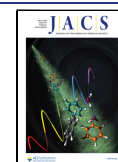
chemical calculations.^{24,25} It was found that the HOMO and LUMO of $[R_2\text{-DBA}]^{2-}$ possess the same local symmetries at the B atoms as the LUMO and HOMO of H_2 , respectively, pointing to a possible concerted oxidative addition reaction. As the polarity of the bond to be activated increases (cf. $F-Ar^F$), it becomes increasingly plausible that the concerted mechanism shifts toward an S_NAr -type scenario involving a nucleophilic B site. Following this rationale, we achieved the synthesis of the fluorophenyl borates $Li[A]$ and $Li[B]$ by treating $Li_2[Me_2\text{-DBA}]$ with 1,3,5- $C_6F_3H_3$ and C_6F_6 , respectively (Figure 1a).²⁸ A strong indication of an S_NAr -type mechanism as opposed to oxidative $F-C$ addition across both B sites is provided by the reactivity of the borafluorene dianion

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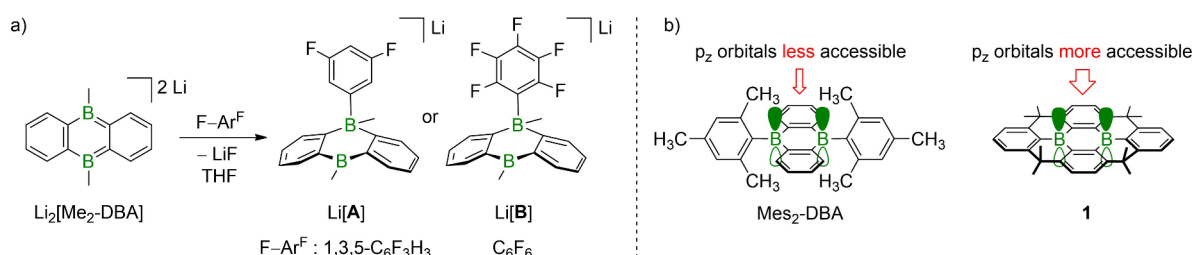


Figure 1. a) Activation of C–F bonds in 1,3,5- $\text{C}_6\text{F}_3\text{H}_3$ or C_6F_6 by $\text{Li}_2[\text{Me}_2\text{-DBA}]$ via an $\text{S}_{\text{N}}\text{Ar}$ -type mechanism. b) Kinetic stabilization of $\text{Mes}_2\text{-DBA}$ blocks substrate access to the $\text{B}(\text{p}_z)$ orbitals; stabilization of **1** by structural constraint retains substrate access to the $\text{B}(\text{p}_z)$ orbitals.

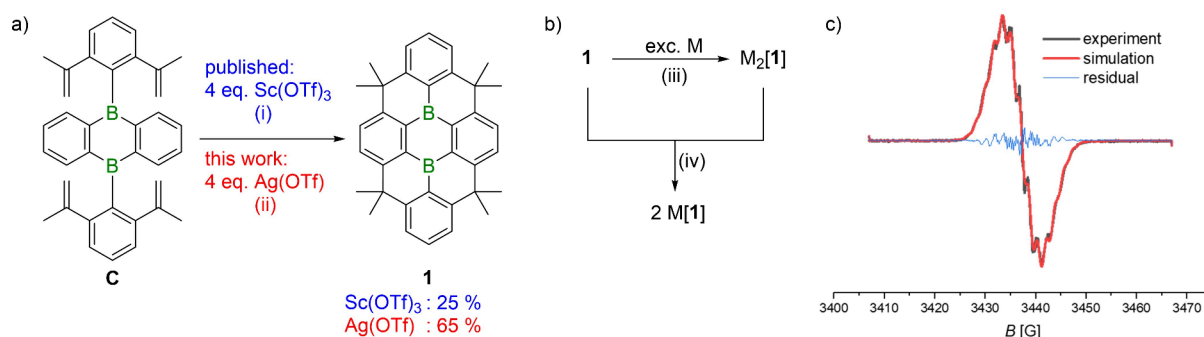


Figure 2. a) Replacing the Lewis acid catalyst $\text{Sc}(\text{OTf})_3$ with $\text{Ag}(\text{OTf})$ in the Friedel–Crafts-type cyclization of **C** improves the yield of **1** to 65%. b) Synthesis of the DBA-dianion salts $\text{M}_2[\text{1}]$ via two-electron reduction of **1** with excess alkali metal (M : Li, Na, K). Generation of the radical-anion salt $\text{M}[\text{1}]$ by comproportionation of **1** and $\text{M}_2[\text{1}]$ (M^+ : Li^+ , K^+). c) Experimental (black) and simulated (red) EPR spectrum of $\text{Li}[\text{1}]$ in THF ($c = 0.49 \text{ mmol L}^{-1}$) at room temperature. The difference between the experimental and simulated spectra likely arises from couplings of Me protons that could not be included into the simulation. Reaction conditions: (i) 1,2-dichloroethane, 83°C , 24 h; (ii) 1,2-dichloroethane, 100°C , 2 d; (iii) THF, room temperature, 3 h, quantitative; (iv) THF, room temperature, instantaneous conversion, quantitative.

salt $\text{Li}_2[\text{H-BFlu}]$: Despite possessing only a single B center, $\text{Li}_2[\text{H-BFlu}]$ exhibits the same reactivity toward F-Ar^{F} as $\text{Li}_2[\text{Me}_2\text{-DBA}]$.^{28,29} Although the considerable potential of $[\text{R}_2\text{-DBA}]^{2-}$ for small molecule activation and even transition metal-free catalysis is rapidly emerging, an important aspect remains unresolved to-date: An efficient reaction requires unimpeded access of the substrate to the B_2 core, which necessitates that the B atoms carry only small substituents such as $\text{R} = \text{H}$ or Me . However, handling even neutral H_2 - and Me_2 -DBAs requires considerable experience, as they are chemically sensitive due to the lack of steric protection. The most obvious solution—replacing H/Me with Mes (mesityl)—proves ineffective, as it not only prevents the undesired attack of, e.g., $\text{H}_2\text{O}/\text{O}_2$, but also blocks any intended interactions at the B sites, even with the smallest substrate, H_2 (Figure 1b).^{25,30} An alternative strategy is to apply the ‘concept of structural constraint’. In the present case, this leads us to compound **1** (Figure 1b),³¹ which has already demonstrated superior chemical stability in the context of emissive materials research. The stability of **1** arises from the rigidly fixed cyclic skeleton around the B atoms and renders any shielding of the $\text{B}(\text{p}_z)$ orbitals unnecessary. Thus, in its dianionic form $[\text{1}]^{2-}$, the compound may also be an attractive candidate for mediating chemical conversions.

Herein, we investigate the suitability of $[\text{1}]^{2-}$ for reactions with fluorobenzenes, because (i) this allows a direct comparison with the behavior of structurally unconstrained dianions $[\text{R}_2\text{-DBA}]^{2-}$, (ii) the available variety of isomeric partially fluorinated benzenes enables the investigation of reactivity trends in response to subtle substrate modifications, (iii) targeted derivatization of F-Ar^{F} can provide value-added products. First, we establish that both the radical $[\text{1}]^{\bullet-}$ and the

closed shell species $[\text{1}]^{2-}$ are accessible in essentially quantitative yields and can be fully characterized. Second, we disclose an unprecedented reactivity shift from the formation of fluorophenyl borates analogous to $\text{Li}[\text{A}]/\text{Li}[\text{B}]$ to hydrodefluorination products, with the crossover point governed by the fluorine load of the substrate and the polarizability of the counteranion M^+ in $\text{M}_2[\text{1}]$ (M^+ : hard Li^+ or soft K^+ ; cf. Figure 3). Third, we prove that a nucleophilic aromatic substitution ($\text{S}_{\text{N}}\text{Ar}$) pathway competes with a single-electron-transfer-induced (SET) radical scenario.

RESULTS AND DISCUSSION

Synthesis of **1, $[\text{1}]^{\bullet-}$, $[\text{1}]^{2-}$.** The structurally constrained, doubly B-doped polycyclic aromatic hydrocarbon **1** (Figure 2a) can be synthesized via an eight-step sequence, as reported by Yamaguchi et al.³¹ Each of the first seven steps achieves yields $>60\%$, thus requiring no further optimization. However, in the final step—a 4-fold Friedel–Crafts-type cyclization on the precursor **C**—the reaction proceeds with only a 25% yield, despite extensive screening of various (Lewis) acids, which identified the employed $\text{Sc}(\text{OTf})_3$ as the most effective catalyst among them.³² Given that compound **1** is central to this study, improving its yield through additional catalyst screening was imperative.³² It was thereby discovered that the soft Lewis acid $\text{Ag}(\text{OTf})$ mediates the reaction with yields of 60–65%, regardless of the specific batch used. Notably, the best-performing catalyst, $\text{Ag}(\text{OTf})$, induces the most pronounced downfield shifts of the propen-2-yl proton signals of **C** in the room-temperature ^1H NMR spectrum, indicating a significant $\text{Ag}^+\cdots\text{olefin } \pi$ -interaction prior to the high-temperature C–C bond formations (Figure S2).

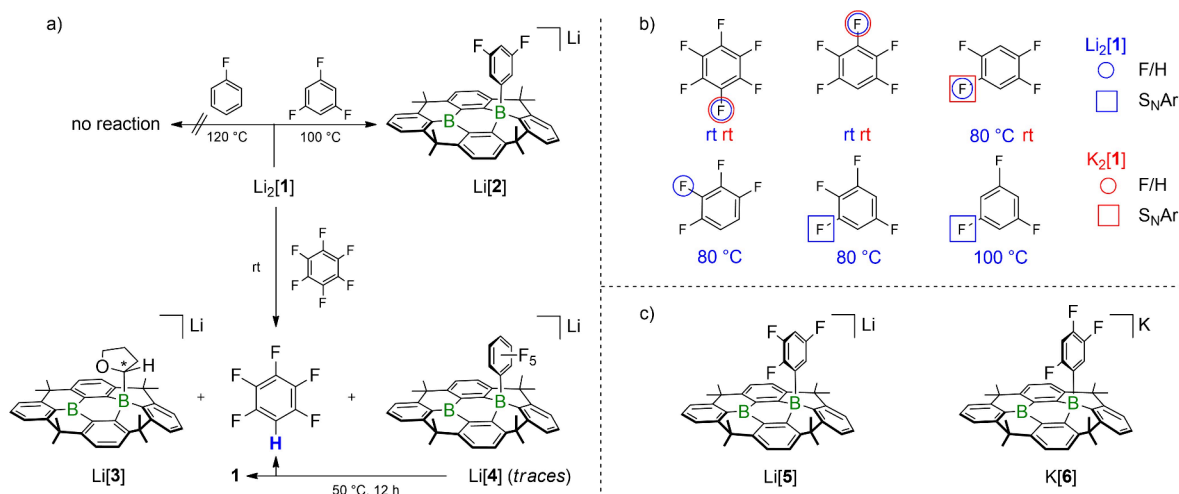


Figure 3. a) Reactivity of $\text{Li}_2[1]$ in THF toward C_6F_6 , $1,3,5\text{-C}_6\text{F}_3\text{H}_3$, and C_6F_6 . b) Fluoroarenes employed and types of their conversions using $\text{Li}_2[1]$ or $\text{K}_2[1]$. Circles: hydrodefluorinations (F/H) mediated by the Li^+ (blue) or K^+ salt (red). Squares: $\text{S}_\text{N}\text{Ar}$ -type reactions of the Li^+ (blue) or K^+ salt (red). c) Products of $\text{S}_\text{N}\text{Ar}$ -type reactions: $\text{Li}_2[1]$ and $1,2,3,5\text{-C}_6\text{F}_4\text{H}_2$ give $\text{Li}[5]$ (75%); $\text{K}_2[1]$ and $1,2,4,5\text{-C}_6\text{F}_4\text{H}_2$ give $\text{K}[6]$ (quantitative).

The cyclic voltammogram of **1** shows two reversible reduction events at $E_{1/2} = -2.04$ and -2.56 V (vs FcH/FcH^+ , $[\text{nBu}_4\text{N}][\text{PF}_6]$, THF),³¹ suggesting that chemical one- and two-electron reduction of **1** should be feasible on a preparative scale. Considering that our previous reactivity studies on $[\text{DBA}]^{2-}$ dianions revealed decisive differences depending on the choice of the counteranion,^{24–26} we performed the corresponding reduction experiments using excess Li, Na, and K metal (Figure 2b). In all cases, quantitative conversions were confirmed by ^1H NMR spectroscopy, affording the corresponding dianion salts $\text{M}_2[1]$. These salts, which have distinct colors in THF solution—dark green (Li^+), brown (Na^+), and purple (K^+)—all appear dark green in the solid state. The radical-anion salts $\text{M}[1]$ were prepared via comproportionation reactions between equimolar amounts of **1** and $\text{M}_2[1]$ (M^+ : Li^+ , K^+ ; Figure 2b). Both salts are dark blue in THF and nearly black in the solid state.

Upon exposure to ambient air, $\text{M}[1]$ and $\text{M}_2[1]$ undergo oxidation to **1** without the formation of any NMR-detectable decomposition products. This is a decisive difference compared to $\text{M}[\text{R}_2\text{-DBA}]$ and $\text{M}_2[\text{R}_2\text{-DBA}]$ (R : H, Me), where also the neutral DBAs are only stable under inert conditions.³³ We note in this context the related radical species $[\text{1C}]^\bullet$, which is formally derived by replacing one of the B atoms in $[\text{1}]^{2-}$ with C(sp^2).³⁴ $[\text{1C}]^\bullet$ is so remarkably inert that it could even be purified by column chromatography on silica gel in air. Even at elevated temperatures, neither $\text{M}[1]$ nor $\text{M}_2[1]$ react with the THF solvent over the long-term; moreover, neither H^\bullet nor H^+ abstraction occurs in the presence of 1,4-cyclohexadiene. Yet, upon the addition of $\text{HSn}(\text{nBu})_3$, $\text{M}[1]$ affords the $\text{B}(\text{sp}^2)\cdots\text{B}(\text{sp}^3)\text{H}$ single hydrioborate $[\text{1}\cdot\text{H}]^-$, while $\text{M}_2[1]$ is converted to the corresponding symmetrical $\text{HB}(\text{sp}^3)\cdots\text{B}(\text{sp}^3)\text{H}$ double hydrioborate $[\text{1}\cdot 2\text{H}]^{2-}$ (see the Supporting Information and Figures S16 – S20 for more details).

The diamagnetic species $\text{M}_2[1]$ were characterized by (heteronuclear) NMR spectroscopy (M^+ : Li^+ , Na^+ , K^+ ; see the Supporting Information for the fully assigned spectra) and single-crystal X-ray diffraction (M^+ : Li^+ , K^+ ; Figures S111, S112). In line with DFT calculations, the obtained results consistently point toward the conclusion that, despite

incorporation of the boron-bonded C_6H_3 rings into the conjugation pathway, even in $[\text{1}]^{2-}$ most of the added charge still resides on the DBA unit.

An EPR spectrum confirming the open-shell electronic structure of $[\text{1}]^{\bullet-}$ was recorded on $\text{Li}[1]$ in THF at room temperature ($c = 0.49$ mmol L^{-1} ; Figure 2c). The observed spectrum was simulated by assuming hyperfine coupling of the unpaired electron ($g_{\text{iso}} = 2.0033 \pm 0.0003$) to two magnetically equivalent B nuclei with isotropic coupling constants of $a(^{11}\text{B}) = 1.6 \pm 0.4$ G and $a(^{10}\text{B}) = 0.5 \pm 0.4$ G. We further assumed coupling to sets of four, four, and two magnetically equivalent protons with $a(^1\text{H}) = 2.0 \pm 0.4$ G, 1.5 ± 0.4 G, and 0.7 ± 0.4 G, respectively (weak coupling of the odd electron to the CH_3 groups was also observed, but not included in the simulation; see the Supporting Information, section 7.1.1, for the computed spin density of $[\text{1}]^{\bullet-}$).

Single-crystal X-ray diffraction of two radical-anion salts, $[\text{Li}(12\text{-c-4})_2][1]$ and $[\text{K}(2.2.2\text{crypt})][1]\cdot 2\text{THF}$, confirmed the molecular structure of $[\text{1}]^{\bullet-}$ (Figure S113 and S122, respectively). The salts form solvent-separated ion pairs in the crystal lattice.

Reactivity of $\text{M}_2[1]$ toward Fluorobenzenes $\text{C}_6\text{F}_n\text{H}_{6-n}$. With $[\text{1}]^{2-}$ in hand, we next examined whether this dianion continues to function as a B-centered nucleophile or if the reorganization energy required for the planar-to-pyramidal conformational transition of the B site becomes prohibitively high due to the structurally constrained framework. We note in this context that **1** is inert toward hydrolysis yet reacts smoothly with appropriate F^- sources to form the corresponding mono- and difluoroborates $[\text{1}\cdot\text{F}]^-$ and $[\text{1}\cdot 2\text{F}]^{2-}$, respectively.^{31,35–37} In the following, the reactions of $\text{M}_2[1]$ (M^+ : Li^+ , K^+) with equimolar amounts of $\text{C}_6\text{F}_n\text{H}_{6-n}$ in THF are described.³⁸ Given the rather complex overall scenario (Figure 3), we will, for the sake of clarity, first provide an overview of the product distributions and discuss analytical details of the isolated products in a subsequent section. A third section is dedicated to mechanistic considerations.

Similar to the structurally unconstrained $\text{Li}_2[\text{Me}_2\text{-DBA}]$,²⁸ $\text{Li}_2[1]$ does not react with C_6F_6 up to $T = 120$ °C. With $1,3,5\text{-C}_6\text{F}_3\text{H}_3$, both species undergo a quantitative $\text{S}_\text{N}\text{Ar}$ -type reaction to furnish the difluorophenylborates $\text{Li}[\text{A}]$ ($T = 50$

°C; Figure 1a) and Li[2] ($T = 100$ °C; Figure 3a).³⁹ However, a striking contrast between the two species emerges in their room-temperature reactions with C_6F_6 : While $Li_2[Me_2-DBA]$ continues to undergo nucleophilic F substitution, furnishing the respective pentafluorophenylborate Li[B] (Figure 1a),²⁸ the reaction with $Li_2[1]$ instead leads to the formation of the tetrahydrofuran-2-yl adduct Li[3] and the monohydrodefluorination product C_6F_5H (Figure 3a). Here, the pentafluorophenylborate Li[4] is only formed in trace amounts; upon heating to 50 °C, Li[4] is slowly converted to C_6F_5D and neutral **1** (in situ 1H NMR spectroscopy in THF- d_8). Upon switching the substrate of $Li_2[1]$ from C_6F_6 to C_6F_5H , the qualitative outcome remains the same, since Li[3] and 1,2,4,5- $C_6F_4H_2$ are obtained as the by far main products at room temperature (Figure 3b). Reducing the fluorine load of the substrate further made it necessary to consider all three isomers of $C_6F_4H_2$ and to apply different temperatures: the less thermodynamically stable⁴⁰ 1,2,4,5- and 1,2,3,4-isomers underwent clean hydrodefluorination (Figure 3b); as indicated by a color change from green ($Li_2[1]$) to blue, the reactions set in at room temperature and 50 °C, respectively, and required 80 °C to reach completion (judged by full decolorization and ^{19}F NMR spectroscopy). In the case of the thermodynamically most favored 1,2,3,5-isomer,⁴⁰ mainly S_NAr reactivity with F^-/B substitution took place in an overall less selective conversion (75%, $T = 80$ °C; Li[5], Figure 3c).

To investigate the extent to which the softer K^+ cation influences the reaction outcome, selected experiments were repeated with $K_2[1]$. Notably, a pronounced counteraction effect was evident for the substrate 1,2,4,5- $C_6F_4H_2$, which showed a reactivity switch from hydrodefluorination (as seen with $Li_2[1]$) to an S_NAr pathway ($K[6]$; Figure 3c). To determine whether this distinct behavior arises from differences in ion-pairing, specifically the formation of contact vs solvent-separated ion pairs by $Li_2[1]$ and $K_2[1]$ in THF solution,⁴¹ we conducted the experiment with $Li_2[1]$ in the presence of excess crown ether 12-c-4 at room temperature. Under these conditions, selective nucleophilic F substitution rather than hydrodefluorination occurred also with the Li^+ salt.

At this stage, the following conclusions can be drawn: (i) Contact ion-pair formation sterically hinders substrate access to the reactive B sites of $[1]^{2-}$. (ii) The suppressive effect on S_NAr reactivity is more pronounced than on hydrodefluorination reactivity. (iii) When both reaction pathways are sterically feasible, the S_NAr pathway seems to be preferred, consistent with results for the unconstrained $[R_2-DBA]^{2-}$ analogues. (iv) Provided that the reactant $Li_2[1]$ is not present in excess, the hydrodefluorination reaction on hexa-, penta-, and tetrafluorobenzenes removes precisely one F atom with complete regioselectivity (Figure 3b). (v) The formation of the thermally stable (at least up to 100 °C) tetrahydrofuran-2-yl adduct Li[3] as the almost exclusive byproduct of the hydrodefluorination reaction is truly remarkable in light of the known susceptibility of THF to decomposition via ring-fragmentation upon α -metalation.⁴² Other examples of authentic α -metalated THF heterocycles are rare: Mulvey et al. achieved the targeted α -zincation of THF by exploiting the unique reactivity of a bimetallic sodium dialkyl(amido)zincate (50% yield after 14 d).⁴³ Soon after, the same group reported the 'structurally engineered deprotonation/alumination of THF' by means of a lithium aluminate base aggregate (35%).⁴⁴ A second example of an α -aluminated THF molecule stems from Driess et al. and was plausibly formed through

deprotonation of THF by an elusive, in situ-formed, donor–acceptor-stabilized $Al(I)-H$ complex.¹³ Finally, Okuda et al. synthesized a tetrahydrofuran-2-yl zirconium(III) complex from a corresponding $[Zr(III)-SiH_2Ph]$ precursor and THF (61% yield after 48 h).⁴⁵

Product Characterization. All NMR spectra were recorded in THF- d_8 at room temperature. The 1H NMR spectrum of the D_{2h} -symmetric dianion $[1]^{2-}$ is characterized by one singlet resonance for the eight equivalent Me groups, alongside one singlet (4H), one doublet (4H), and one triplet (2H), all assignable to the aryl protons. The presence of the 3,5- $C_6F_2H_3$ substituent in $[2]^-$ lowers the symmetry of the anion, resulting in six aryl signals from the B_2 -PAH framework and four distinct Me resonances (each integrating to 6H). A $^1H-^1H$ NOESY experiment revealed that the most magnetically shielded Me group is the one in closest spatial proximity to the difluorophenyl substituent. The two 1H resonances of the 3,5- $C_6F_2H_3$ ring are upfield shifted relative to the signal of 1,3,5- $C_6F_3H_3$, with multiplicities matching those of the analogous DBA derivative $Li[A]$.²⁸ The presence of 20 B_2 -PAH and 4 difluorophenyl signals in the $^{13}C\{^1H\}$ NMR spectrum further supports the proposed molecular structure of Li[2]. Its ^{11}B NMR spectrum shows only a resonance for the tetracoordinate B site ($\delta = -16.7$); as is often observed for B-PAHs, the signal corresponding to the tricoordinate B atom is broadened beyond detection.⁴⁶ A key characteristic of the tetrahydrofuran-2-yl adduct Li[3] is the further increase in the number of distinct Me resonances to eight in both the 1H and $^{13}C\{^1H\}$ NMR spectra; all other H and C atoms of the B_2 -PAH scaffold are also magnetically unique. This arises from the B-bonded, stereogenic, and configurationally stable α -C atom, which imparts C_1 symmetry to $[3]^-$. Concomitantly, seven 1H and four ^{13}C signals are detected for the tetrahydrofuran-2-yl substituent. The most striking chemical shift difference within this signal set is observed for the O-bonded α -C atoms: while $\delta(^{13}C) = ca. 67$ for the O-CH₂-C fragment closely matches that of free THF, $\delta(^{13}C) = 91.7$ for the O-C*H-B fragment is significantly downfield shifted (the C*H proton resonates at 3.06 ppm). The B_2 -PAH unit of Li[5] (Figure 3c), the major product of the S_NAr substitution reaction on 1,2,3,5- $C_6F_4H_2$ (75%), shows NMR features nearly identical to those of Li[2]. The $^{19}F\{^1H\}$ NMR spectrum of Li[5] displays two doublets ($^3J_{FF} = 24.4$ Hz, $^4J_{FF} = 17.5$ Hz) and a corresponding doublet of doublets. The 1H and $^{19}F\{^1H\}$ NMR spectra also provide evidence for the formation of Li[3] and of hydrodefluorinated species as side products of this reaction. Additionally, X-ray crystallography revealed the generation of (at least) one isomeric substitution product cocrystallizing with Li[5] (Figure S116). Based on these observations, we conclude that for $Li_2[1]$, 1,2,3,5- $C_6F_4H_2$ marks the crossover point between the S_NAr vs hydrodefluorination scenarios. Although the position of the B atom on the respective trifluorophenyl ring differs between compounds $K[6]$ and Li[5] (Figure 3c), the spatial relationship of the F atoms remains identical. Consequently, the qualitative NMR characteristics of both compounds are so similar that a detailed discussion of the $K[6]$ spectra is not warranted (the Supporting Information contains the fully assigned NMR spectra and the solid-state structure of $K[6]$). The fluorobenzenes obtained in this work are all known. In addition to in situ GC-MS, 1H and ^{19}F NMR spectra were recorded (see the Supporting Information for plots of the spectra); the obtained chemical shifts and signal multiplicities align with the literature data.⁴⁷

In addition to NMR characterization, the molecular structures of $\text{Li}[2]$, $\text{Li}[3]$, $\text{Li}[5]$, and $\text{K}[6]$ were confirmed by single-crystal X-ray diffraction (see the [Supporting Information](#) for full details). As representative examples, $[\text{Li}(\text{thf})_4][2] \cdot 0.2 \text{ THF}$ and $[\text{Li}(\text{thf})_3][3] \cdot \text{THF}$ are discussed in more detail ([Figure 4](#)). $[\text{Li}(\text{thf})_4][2] \cdot 0.2 \text{ THF}$ crystallizes as

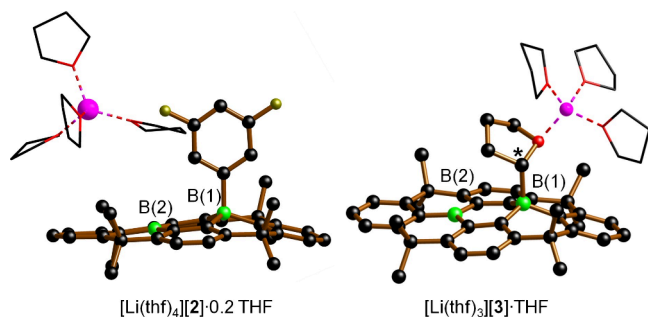


Figure 4. Molecular structures of $[\text{Li}(\text{thf})_4][2] \cdot 0.2 \text{ THF}$ and $[\text{Li}(\text{thf})_3][3] \cdot \text{THF}$ in the solid state. H atoms and noncoordinating solvent molecules are omitted for clarity. Li, purple; B, green; C, black; O, red; F, yellow-green.

solvent-separated ion pairs. Each anionic moiety $[2]^-$ comprises one tetracoordinate B(1) and one tricoordinate B(2) site, with the former bearing the 3,5- $\text{C}_6\text{F}_2\text{H}_3$ ligand. The F^- ion released during $\text{Li}[2]$ formation does obviously not remain coordinated to the second available Lewis-acidic center.

It is revealing to compare selected structural parameters of $[2]^-$ with those of the analogous, structurally unconstrained

$[\text{Li}(12\text{-c-4})_2][\text{A}]$ (cf. [Figure 1a](#)).²⁸ Reference values from $[\text{A}]^-$ are given in curly brackets after each metrical value of $[2]^-$: The average B(2)–C bond length of $[2]^-$ is 1.53 Å {1.56 Å}, and the sum of bond angles about B(2) amounts to 360° {360°}, in line with a planar, sp^2 -hybridized atom. Conversely, the average length of the three endocyclic B(1)–C bonds amounts to 1.59 Å {1.64 Å}, indicating significant elongation due to rehybridization; the corresponding three endocyclic C–B(1)–C bond angles sum to 336° {332°} (ideal tetrahedron: 329°). The B(1)–C bond to the terminal difluorophenyl substituent measures 1.683(8) Å {1.665(2) Å}. Overall, B(1) in $[2]^-$ is significantly pyramidalized, albeit to a lesser degree than the corresponding B site in $[\text{A}]^-$. Moreover, the difference between endocyclic and exocyclic B–C bonds is more pronounced in $[2]^-$ than in $[\text{A}]^-$. Although the observed effects may seem to be straightforwardly attributable to the structural constraint of **1**, this is not necessarily the whole story, as its F^- -adduct is even slightly more pyramidalized than $[\text{A}]^-$.^{28,31} We therefore propose that the steric repulsion of the bulky Lewis base also matters, as reflected particularly in the long B(1)– $\text{C}_6\text{F}_2\text{H}_3$ bond of $[2]^-$. Compound $[\text{Li}(\text{thf})_3][3] \cdot \text{THF}$ crystallizes as a contact-ion pair, with the $[\text{Li}(\text{thf})_3]^+$ cation bonded to the O atom of the B(1)-appended tetrahydrofuran-2-yl substituent. The bond connecting B(1) to the stereogenic C* atom of the tetrahydrofuran-2-yl residue measures 1.698(5) Å. Consistent with Bent's rule,⁴⁸ the C*–O/C*– CH_2 bonds are significantly longer than the analogous bonds involving the other α -C atom, i.e., 1.477(4)/1.543(5) vs 1.434(5)/1.498(6) Å.⁴⁹ A structurally comparable product $[\text{Li}(\text{thf})_4][3^{\text{Me}}]$ was obtained when the reaction between $\text{Li}_2[1]$ and C_6F_6 was carried out in 2,5-dimethyltetrahydrofur-

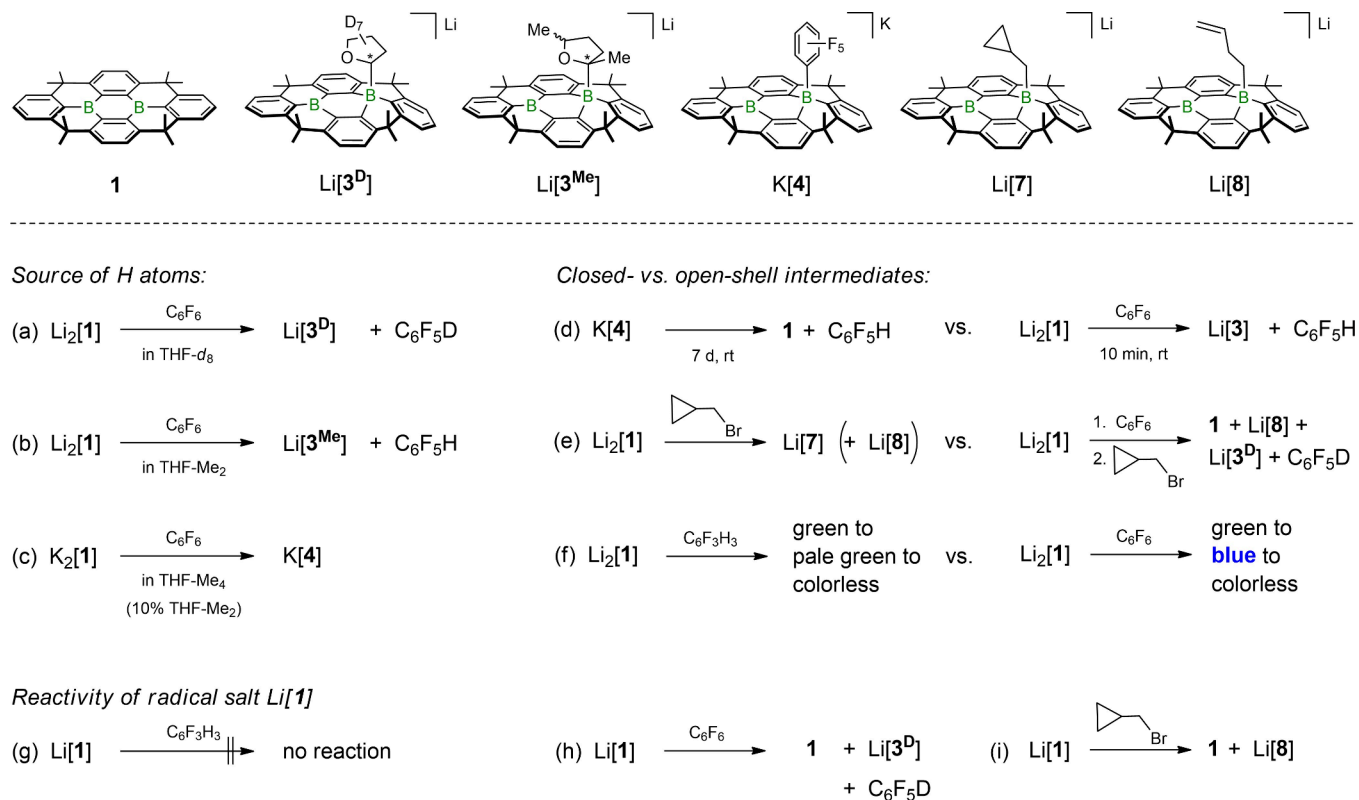


Figure 5. Experiments (a)–(i) provide insight into key elementary steps of the reaction mechanism underlying the reaction of $\text{Li}_2[1]$ with C_6F_6 , leading to the formation of $\text{Li}[3]$ and $\text{C}_6\text{F}_5\text{H}$. All experiments were performed in THF or THF- d_8 , unless stated otherwise. (h) **1**, $\text{Li}[3^{\text{D}}]$, and $\text{C}_6\text{F}_5\text{D}$ were formed in a 1:1:1 ratio; (i) **1** and $\text{Li}[8]$ were formed in a 1:1 ratio.

an (THF-Me₂; Figure 5). As the main difference, the 2,5-dimethyltetrahydrofuran-2-yl substituent no longer coordinates to the Li⁺ cation in the solid state. Instead, the crystal lattice is composed of solvent-separated ion pairs with [Li(thf)₄]⁺ cations, likely as a result of the steric shielding effect imparted by the two Me groups adjacent to the O-donor atom (Figure S115).

Mechanistic Investigations. Most of the compounds newly synthesized during the mechanistic studies closely resemble those reported above, rendering a detailed discussion of their NMR and solid-state structural characteristics redundant. Given the complexity of the reaction mechanisms and the numerous details involved, we aim to keep the focus entirely on this aspect and refer to the [Supporting Information](#) for a compilation of all analytical data relevant to confirming the nature of the additional products (note that all products discussed in the following have been structurally characterized by single-crystal X-ray diffraction). Based on the experimental evidence gathered from the reactions of Li₂[Me₂-DBA] (and Li₂[H-BFlu]) with F-Ar^F, an S_NAr-type mechanism involving B-centered nucleophiles is a plausible postulate.²⁸ While all available findings suggest that the transformation of Li₂[1] with 1,3,5-C₆F₃H₃ to afford Li[2] follows a similar pathway, the reaction of Li₂[1] with C₆F₆ takes a different route, furnishing the tetrahydrofuran-2-yl adduct Li[3] and C₆F₅H (Figure 3a).

This process therefore warrants more detailed scrutiny (Figure 5). We began by exploring the source of the H atoms in the hydrodefluorination products: (a) Replacing THF with THF-*d*₈ resulted in the preferential formation of C₆F₅D over C₆F₅H. While the NMR signature of the B₂-PAH byproduct remained essentially identical to that of [3][−], we no longer detected ¹H resonances for the tetrahydrofuran-2-yl substituent, confirming its identity as a C₄D₇O moiety in Li[3^D]. (b) Switching from THF to THF-Me₂ still yielded the 2,5-dimethyltetrahydrofuran-2-yl analogue Li[3^{Me}] of Li[3], along with C₆F₅H. (c) Removal of the remaining two α-H atoms using 2,2,5,5-tetramethyltetrahydrofuran (THF-Me₄) required the more soluble K₂[1] instead of Li₂[1] and 5–10 vol % of THF-Me₂ as solubilizing agent. Under these conditions, the C₆F₅ adduct K[4] (Figure 5; Figure S119) was obtained as the dominant initial product, which subsequently converted to neutral 1 and C₆F₅H over 7 d at room temperature or within 1 d at 50 °C. These results lead to the following conclusions: (a) The THF solvent is the source of the H atom in C₆F₅H. (b) H atom abstraction from the solvent remains the preferred—albeit somewhat slowed down—reaction, even when the site of attack is sterically shielded in THF-Me₂; likewise, the increased steric bulk does not prevent the formation of the B–C bond in Li[3^{Me}] (Figure S115). (c) K[4] becomes isolable only if the majority of the solvent molecules lack α-H atoms (THF-Me₄). Although compounds M[4], once generated, degrade slowly via α-deprotonation of THF, yielding C₆F₅H and the extremely short-lived⁵⁰ tetrahydrofuran-2-yl anion [C₄H₇O][−], the absence of [3][−] in this process suggests that hydrodefluorination within the M₂[1]/C₆F₆/THF system does not proceed via M[4] as the typical pathway. Instead, the following observations point toward a different key intermediate (Figure 5): (d) While in situ ¹⁹F NMR monitoring of a solution of K[4] in THF confirms the gradual accumulation of C₆F₅H over time, the overall reaction is 3 orders of magnitude slower than the formation of C₆F₅H directly from the M₂[1]/C₆F₆/THF mixture. (e) Treatment of a THF-*d*₈ solution of Li₂[1]

with 1.2 eq. of the fast radical clock (bromomethyl)-cyclopropane [(C₃H₅)CH₂Br] yields the (C₃H₅)CH₂ adduct Li[7] along with the ring-opened H₂C=C(H)CH₂CH₂ adduct Li[8] (85% and 15%, respectively; Figure S120). This result confirms that [1]^{2−} can indeed behave as a B-centered nucleophile in a predominantly closed-shell reaction.⁵¹ However, when 1 eq. of C₆F₆ was added to the mixture prior to introducing the radical clock (keeping all other conditions unchanged), exclusively the ring-opened Li[8] was formed besides 1, Li[3^D], and C₆F₅D, providing evidence for the presence of radicals in the sample.⁵¹ (f) For the mixtures Li₂[Me₂-DBA]/1,3,5-C₆F₃H₃, Li₂[Me₂-DBA]/C₆F₆, or Li₂[1]/1,3,5-C₆F₃H₃, the original green color of the dianion salt gradually fades as the reaction progresses. In stark contrast, the green solution of Li₂[1] in THF instantaneously turns deep blue when C₆F₆ is added, before decolorizing within minutes (the products Li[3] and C₆F₅H are colorless). The observed color change in (f) is particularly diagnostic, as the emerging blue tint of the reaction mixture appears visually identical to that of the radical-anion salt Li[1] (see above). This prompted us to explore the possibility of a radical pathway. Given that the UV–vis spectral differences between K[1] and K₂[1] are more distinct than those between Li[1] and Li₂[1] (Figure S105), we conducted subsequent experiments using the K⁺ salts, which show the same reactivity toward C₆F₆ as the Li⁺ salts. Absorption bands at 940 and 705 nm are particularly useful for distinguishing K[1] from K₂[1] in THF, as they have minimal overlap with the absorption bands of the respective counterpart (cf. the normalized spectra shown in bold black and red lines in Figure 6).⁵² The preparation of K₂[1]/C₆F₆/

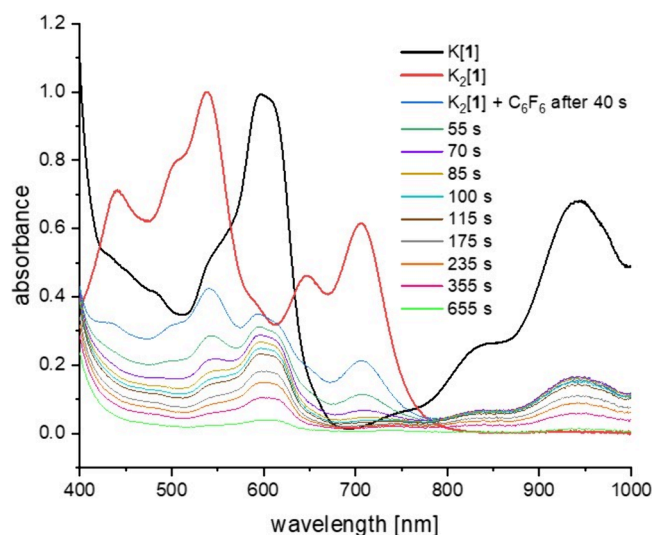


Figure 6. Normalized UV–vis absorption spectra of K[1] (bold black line) and K₂[1] (bold red line) in THF. Time-dependent UV–vis spectroscopic monitoring of an equimolar mixture of K₂[1] and C₆F₆ in THF, recorded from 40 s after mixing (blue line).

THF samples under strictly inert conditions, their transfer to the cuvette, and the initiation of the UV–vis measurement typically required 40 s. Consequently, the first recorded spectrum reflects the chemical composition of the reaction mixture at room temperature after this time, revealing the presence of both unconsumed K₂[1] and newly formed K[1] (blue line in Figure 6). After an additional 75 s, the absorption bands of K₂[1] had largely disappeared, while those of K[1]

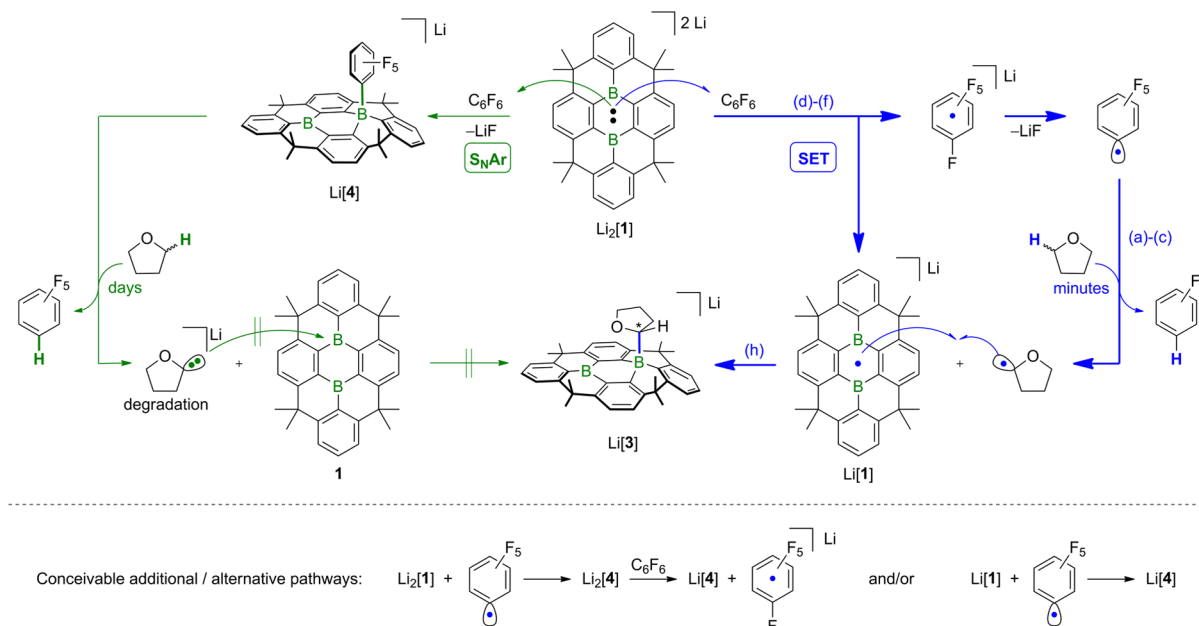


Figure 7. Proposed mechanistic scenario for the reaction of $\text{Li}_2[1]$ with C_6F_6 in THF. Blue arrows: Single-electron transfer (SET) pathway via the blue radical salt $\text{Li}[1]$ and $[\text{C}_6\text{F}_6]^\bullet$ (major process). The signature product of this sequence is $\text{Li}[3]$; labels (a)–(f), (h) refer to the experiments outlined in Figure 5 that support the proposed individual step. Green arrows: Nucleophilic aromatic substitution ($\text{S}_\text{N}\text{Ar}$) pathway leading to the formation of $\text{Li}[4]$ (minor process); subsequent slow deprotonation of THF gradually converts $\text{Li}[4]$ to **1** and $\text{C}_6\text{F}_5\text{H}$. Formation of $\text{Li}[4]$ through coupling between $[1]^{2-}$ or $[1]^\bullet$ and $[\text{C}_6\text{F}_5]^\bullet$ cannot be excluded.

remained at approximately their original intensity (brown line). After a further 9 min, however, the spectral features of $\text{K}[1]$ are also barely discernible (green line). Taken together, this supports the view that $\text{K}_2[1]$ undergoes continuous conversion to $\text{K}[1]$, which reacts in parallel at a comparable rate, resulting in a steady concentration of $\text{K}[1]$. Once the conversion of $\text{K}_2[1]$ to $\text{K}[1]$ ceases due to the depletion of $\text{K}_2[1]$, the concentration of $\text{K}[1]$ begins to decline as well. The final product, tetrahydrofuran-2-yl adduct $\text{K}[3]$, has no absorption bands of appreciable intensity in the wavelength range of 400–1000 nm. An analogous UV–vis monitoring of the $\text{K}_2[1]/1,2,4,5\text{-C}_6\text{F}_4\text{H}_2/\text{THF}$ system, which quantitatively generates the $2,4,5\text{-C}_6\text{F}_3\text{H}_2$ adduct $\text{K}[6]$ at room temperature (Figure 3c), revealed a gradual fading of the $\text{K}_2[1]$ bands without the intermediate emergence of the UV–vis signature of $\text{K}[1]$ (Figure S108). We consider the results of the two UV–vis monitoring studies as strong evidence that the reaction of $1,2,4,5\text{-C}_6\text{F}_4\text{H}_2$ with $\text{K}_2[1]$ follows a closed-shell pathway, whereas an open-shell mechanism dominates in the case of C_6F_6 .

How does the reactivity pattern differ when the radical-anion salt $\text{Li}[1]$ is used directly, rather than being generated in situ from $\text{Li}_2[1]/\text{C}_6\text{F}_6$? The following series of experiments provided insights (Figure 5): (g) $\text{Li}[1]$ remains inert toward $1,3,5\text{-C}_6\text{F}_3\text{H}_3$ up to temperatures of 120 °C (THF- d_6 ; the ^1H and ^{19}F NMR signals of $1,3,5\text{-C}_6\text{F}_3\text{H}_3$ are detectable in the otherwise NMR-silent sample and do not change over time; the dark blue color of $\text{Li}[1]$ persists). (h) With the higher fluorinated substrate C_6F_6 , however, a room-temperature reaction occurs, furnishing **1**, $\text{Li}[3]$, and $\text{C}_6\text{F}_5\text{D}$ in a 1:1:1 ratio. According to in situ ^{19}F NMR spectroscopy, the use of 0.5 eq. C_6F_6 results in its quantitative conversion to $\text{C}_6\text{F}_5\text{D}$, whereas the use of 1.0 eq. C_6F_6 generates an equimolar mixture of $\text{C}_6\text{F}_5\text{D}$ and unconsumed C_6F_6 . (i) The radical clock $(\text{C}_3\text{H}_5)\text{CH}_2\text{Br}$ reacts with $\text{Li}[1]$ to produce equimolar

amounts of **1** and the ring-opened adduct $\text{Li}[8]$. Experiments (g) and (h) indicate that $\text{Li}[1]$ is not sufficiently reducing to transfer an electron to $1,3,5\text{-C}_6\text{F}_3\text{H}_3$, yet it is a strong enough electron donor to reduce C_6F_6 (computed adiabatic electron affinities: -0.565 vs 0.400 eV).⁵³ In turn, some $\text{Li}[1]$ molecules are converted to **1**, while others form $\text{Li}[3]$. The invariant outcome of experiment (h), regardless of whether 0.5 or 1.0 eq. C_6F_6 is added, indicates that the SET from $\text{Li}[1]$ to C_6F_6 is the rate-determining step. Otherwise, the expected extent of $\text{Li}[1] \rightarrow \text{1}$ oxidation would exceed 50% as soon as more than 0.5 eq. C_6F_6 is introduced. A similar reasoning holds for experiment (i): Initially, $\text{Li}[1]$ undergoes an SET with $(\text{C}_3\text{H}_5)\text{CH}_2\text{Br}$. Following Br^- elimination from $[(\text{C}_3\text{H}_5)\text{CH}_2\text{Br}]^\bullet$, the resulting $[(\text{C}_3\text{H}_5)\text{CH}_2]^\bullet$ radical rearranges to the $[\text{H}_2\text{C}=\text{C}(\text{H})\text{CH}_2\text{CH}_2]^\bullet$ radical, which afterward recombines with unconsumed $\text{Li}[1]$ to furnish $\text{Li}[8]$.

Experiments (a)–(i) outlined in Figure 5 provide a consistent picture of all key steps underlying the reactions of $\text{M}_2[1]$ with fluorobenzenes. These reactions can proceed via either closed-shell $\text{S}_\text{N}\text{Ar}$ or open-shell SET/HAT mechanisms (HAT: hydrogen-atom transfer⁵⁴). The most important parameter that defines the boundary between these two scenarios is the fluorine load on the benzene ring: an increasing degree n of benzene fluorination stabilizes the π^* acceptor orbital, thereby leading to more positive adiabatic electron affinities (AEAs) of $\text{C}_6\text{F}_n\text{H}_{6-n}$. According to quantum-chemical calculations and photoelectron spectroscopy, only C_6F_6 and $\text{C}_6\text{F}_5\text{H}$ possess AEAs of appreciably positive values (>0.2 eV).^{53,55–57} This aligns with our observation that the deep blue radical intermediate $[1]^\bullet$ is detected during the reactions of $\text{M}_2[1]$ with C_6F_6 and $\text{C}_6\text{F}_5\text{H}$, but not in those of $\text{Li}_2[1]$ with trifluorinated $1,3,5\text{-C}_6\text{F}_3\text{H}_3$ (computed ΔG_SET values for the corresponding SETs: -6.5 , -4.3 , and $+9.5$ kcal/mol; Figure S125). The borderline nature of tetrafluorinated $1,2,4,5\text{-C}_6\text{F}_4\text{H}_2$ becomes apparent from the fact that its hydro-

defluorination to 1,2,4- $\text{C}_6\text{F}_3\text{H}_3$ with $\text{Li}_2[1]$ involves $[1]^{\bullet-}$ ($\Delta G_{\text{SET}} = +4.0$ kcal/mol), whereas no $[1]^{\bullet-}$ is observed during the formation of $[6]^-$ from $[\text{Li}(12\text{-c-4})_n]_2[1]$ or $\text{K}_2[1]$.⁵⁸ Upon electron uptake, $\text{C}_6\text{F}_n\text{H}_{6-n}$ lose their planar conformations. Specifically, C_6F_6 undergoes out-of-plane distortion along normal mode-like coordinates, resulting in the C_{2v} -symmetric $[\text{C}_6\text{F}_6]^{\bullet-}$ anion: Two *para*-positioned F atoms bend upward, whereas the remaining four exhibit a slight downward bend. Consequently, the singly occupied molecular orbital (SOMO) exhibits significant $\pi/\sigma^*(\text{C}-\text{F})$ mixing, which explains why the injected electron can efficiently induce C–F-bond cleavage to release F^- and afford the $[\text{C}_6\text{F}_5]^{\bullet}$ radical (see the Supporting Information for an NBO analysis).⁵⁶

In summary, considering all experimental and computational data, we propose that the reaction of $[1]^{2-}$ with C_6F_6 is initiated by an SET, generating blue $[1]^{\bullet-}$ and $[\text{C}_6\text{F}_6]^{\bullet-}$ (blue pathway in Figure 7). While $[1]^{\bullet-}$ itself is stable in THF over the long-term, $[\text{C}_6\text{F}_6]^{\bullet-}$ rapidly releases one F^- ion to form the neutral $[\text{C}_6\text{F}_5]^{\bullet}$ radical,^{59–61} driven by LiF precipitation. In a subsequent HAT step, the $[\text{C}_6\text{F}_5]^{\bullet}$ radical abstracts an α -H atom from THF, yielding the hydrodefluorination product $\text{C}_6\text{F}_5\text{H}$, along with a $[\text{C}_4\text{H}_7\text{O}]^{\bullet}$ radical, which then recombines with the long-lived $[1]^{\bullet-}$ to furnish $\text{Li}[3]$.⁶² A quantum-chemical evaluation of the entire reaction sequence reveals an essentially barrierless and strongly exergonic process (Figure S140). The trace amounts of side product $\text{Li}[4]$ originate either from an $\text{S}_{\text{N}}\text{Ar}$ attack of green $[1]^{2-}$ on C_6F_6 (green pathway in Figure 7), from an attack of $[\text{C}_6\text{F}_5]^{\bullet}$ on the closed-shell arene $[1]^{2-}$ (followed by one-electron oxidation of the primary product $\text{Li}_2[4]$ by C_6F_6), or from $[\text{C}_6\text{F}_5]^{\bullet}/[1]^{\bullet-}$ coupling as a minor competing pathway to the HAT step. Switching from C_6F_6 to 1,3,5- $\text{C}_6\text{F}_3\text{H}_3$ shuts down the SET pathway due to the negative AEA of the new substrate. Consequently, the $\text{Li}[4]$ analogue $\text{Li}[2]$ is obtained selectively along the $\text{S}_{\text{N}}\text{Ar}$ pathway.

The two most closely related HDF/HAT reactions involving fluoroarenes reported to date have been disclosed by Weaver et al.⁶³ and Zhang et al.⁶⁴ In both cases, the reactions proceed via light-driven processes employing Ir- or pyrene-based photocatalysts, respectively. In the initial step, these photocatalysts mediate electron transfer from the sacrificial electron donor *i*Pr₂NEt to the respective fluoroarene acceptor. The resulting fluoroarene radical anion then undergoes a similar transformation cascade as shown for $\text{Li}[\text{C}_6\text{F}_6]$ in Figure 7. However, unlike our system, the H atom donors are not the solvent but the $[\text{iPr}_2\text{NEt}]^{\bullet+}$ radical cations generated in the course of the photoinduced SET process. Moreover, HDF reactions driven by $\text{Li}_2[1]$ proceed within minutes and even in the dark, whereas the photocatalysis protocols require continuous irradiation over 24 h (see the Supporting Information for more details). Of the two literature processes, only Zhang's—like ours—avoids the use of a transition metal. Notably, however, the two reference processes are truly catalytic, achieving turnover numbers >20,000, whereas the $\text{M}_2[1]/\text{C}_6\text{F}_n\text{H}_{6-n}$ reaction under current conditions proceeds in a stoichiometric manner. Nevertheless, the structural constraint that stabilizes **1** enables its easy regeneration from the side product $\text{Li}[3]$ in the presence of weak H^+ donors, indicating the potential of our protocol for future development into a catalytic process.

CONCLUSION

The search for transition metal-free homogeneous catalysts is both intellectually appealing and practically important. We previously demonstrated that doubly reduced 9,10-dihydro-9,10-diboraanthracenes ($[\text{R}_2\text{-DBA}]^{2-}$; R: H, Me) provide a promising compound class for activating element–element bonds. A critical limitation of $\text{R}_2\text{-DBAs}$, however, is their air and moisture sensitivity, as the two B sites must remain accessible to the substrate and therefore should not be sterically protected. To overcome this issue, we introduce the structurally constrained DBA derivative **1** for element–element bond-activation, where the B atoms are embedded in a planar framework, ensuring ambient stability. Fluorobenzenes were selected as substrates due to the robust nature of their C–F bonds, which makes them challenging to address.

Despite its rigid skeleton, and similar to the unconstrained $[\text{Me}_2\text{-DBA}]^{2-}$, $[1]^{2-}$ acts as a B-centered nucleophile toward moderately fluorinated benzenes. For example, 1,3,5- $\text{C}_6\text{F}_3\text{H}_3$ undergoes $\text{S}_{\text{N}}\text{Ar}$ -type reactions to afford the difluorophenyl borate $[2]^-$. In contrast to $[\text{Me}_2\text{-DBA}]^{2-}$, the reactivity strikingly switches when $[1]^{2-}$ reacts with highly fluorinated benzenes: In THF, $[1]^{2-}$ mediates hydrodefluorination of C_6F_6 to $\text{C}_6\text{F}_5\text{H}$, accompanied by the formation of the tetrahydrofuran-2-yl adduct $[3]^-$ via a radical pathway. This high-yielding open-shell mechanism, unprecedented for $[\text{R}_2\text{-DBA}]^{2-}$ derivatives, greatly expands their reactivity spectrum. The pronounced hydrodefluorination (regio)selectivities observed for C_6F_6 , $\text{C}_6\text{F}_5\text{H}$, as well as 1,2,4,5- and 1,2,3,4- $\text{C}_6\text{F}_4\text{H}_2$ suggest potential applications in the targeted synthesis of partially fluorinated arenes or in the degradation of organofluorine pollutants. The chemical stability of **1** allows its recycling by simple exposure of $[3]^-$ to air. Additionally, $[3]^-$ itself is notable as an extremely rare α -metalated THF derivative with potential as a tetrahydrofuran-2-yl transfer reagent in synthetic applications. Comparable adducts cannot be obtained via Lewis acid–base pairing between **1** and $[\text{C}_4\text{H}_7\text{O}]^-$, as α -deprotonated THF undergoes instantaneous fragmentation. The α -radical, in contrast, has a sufficiently long lifetime to form $[3]^-$ through an open-shell pathway—a vastly underexplored method for B–C-bond formation. The unique reactivity of $[1]^{2-}$ showcases the power of structural constraint in tuning main-group element reactivity and provides new perspectives for catalyst design beyond transition metal chemistry.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/jacs.5c05588>.

All experimental and computational details (PDF)

Calculated_structures.xyz (TXT)

Accession Codes

Deposition Numbers 2434035–2434046 and 2448823 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via the joint Cambridge Crystallographic Data Centre (CCDC) and Fachinformationszentrum Karlsruhe Access Structures service.

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Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

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ABBREVIATIONS

PAH, polycyclic aromatic hydrocarbon

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