Site-Selective $\text{C}_{\text{sp}}^3 - \text{C}_{\text{sp}}/\text{C}_{\text{sp}}^3 - \text{C}_{\text{sp}}^2$ Cross-Coupling Reactions Using Frustrated Lewis Pairs

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ABSTRACT: The donor−acceptor ability of frustrated Lewis pairs (FLPs) has led to widespread applications in organic synthesis. Single electron transfer from a donor Lewis base to an acceptor Lewis acid can generate a frustrated radical pair (FRP) depending on the substrate and energy required (thermal or photochemical) to promote an FLP into an FRP system. Herein, we report the $\text{C}_{\text{sp}}^3 - \text{C}_{\text{sp}}$ cross-coupling reaction of aryl esters with terminal alkynes using the $\text{B(C}_6\text{F}_5\text{)}_3/\text{Mes}_3\text{P}$ FLP. Significantly, when the 1-ethynyl-4-vinylbenzene substrate was employed, the exclusive formation of $\text{C}_{\text{sp}}^3 - \text{C}_{\text{sp}}$ cross-coupled products was observed. However, when 1-ethynyl-2-vinylbenzene was employed, solvent-dependent site-selective $\text{C}_{\text{sp}}^3 - \text{C}_{\text{sp}}$ or $\text{C}_{\text{sp}}^3 - \text{C}_{\text{sp}}^2$ cross-coupling resulted. The nature of these reaction pathways and their selectivity has been investigated by extensive electron paramagnetic resonance (EPR) studies, kinetic studies, and density functional theory (DFT) calculations both to elucidate the mechanism of these coupling reactions and to explain the solvent-dependent site selectivity.

INTRODUCTION

Frustrated Lewis pairs (FLPs) have garnered much attention over the last two decades, with numerous FLP systems being reported in the literature. The cooperative reactivity of the Lewis acidic and basic components has led to a plethora of different small-molecule activation reactions and catalysis. Current studies have focused on using FLP systems as alternatives or complementary systems to the use of transition-metal catalysts in organic synthesis. Recently, new reactivities of FLPs have been disclosed, indicating that Lewis acids and bases undergo single electron transfer (SET) events depending on the energy required (thermal or photochemical) to promote the FLP into a frustrated radical pair (FRP) system. In these instances, an electron is transferred from the donor Lewis base (LB) to the acceptor Lewis acid (LA) to generate a reactive FRP. Indeed, we and others have postulated that such radical reactivity may be taking place in the reactions of the $\text{B(C}_6\text{F}_5\text{)}_3/\text{Mes}_3\text{P}$ FLP with certain substrates. The radical reactivity of FLPs has the potential to open up new opportunities for metal-free synthesis. In a previous study of the $\text{B(C}_6\text{F}_5\text{)}_3/\text{Mes}_3\text{P}$ FLP with diaryl esters and alkenes, we observed $\text{C}_{\text{sp}}^3 - \text{C}_{\text{sp}}^2$ coupling reactions. We proposed a radical mechanism for the reaction based on the observation of $[\text{Ar}_2\text{CH}]^+$ and $[\text{Mes}_3\text{P}]^+$ in electron paramagnetic resonance (EPR) studies (Scheme 1A).

In this current study, we were interested in the reactions of FLPs with alkynes in the presence of aryl esters (Scheme 1B). The 1,2-trans-addition of the Lewis acidic and basic...
components of FLPs to alkenes is well established and has also been employed in catalytic transformations. Interestingly, when using terminal acetylenes such as phenylacetylene (PhC≡CH) with stronger bases such as tBu3P or TMP, deprotonation occurs instead of 1,2-addition to the alkyne generating [LH][PhC≡CBr(C6F5)3] salts. In transition-metal chemistry, the activation of terminal alkenes for cross-coupling reactions is commonplace in the synthetic chemist’s toolbox to construct carbon–carbon bonds. For example, palladium- or copper-catalyzed Sonogashira cross-coupling reactions is commonplace in the synthetic chemist’s toolbox to construct carbon–carbon bonds. Typically, palladium or earth-abundant metal catalysts such as iron and copper are employed for these reactions, although examples are known with other elements such as indium and Lewis acids.

Herein, we report the high reactivity of frustrated Lewis pairs in selective Csp-Csp cross-coupling reactions between aryl esters and terminal alkenes or 1-ethyl-4-vinylbenzene. We also report solvent-dependent site selectivity when using 1-ethynyl-2-vinylbenzene as a substrate leading to selective Csp-coupling depending upon the solvent employed.

**RESULTS AND DISCUSSION**

**Reaction Optimization and Development.** Initially, the FLP-mediated Csp–Csp cross-coupling reaction between bis(4-fluorophenyl)methyl-4-fluorobenzoate (1a) and phenylacetylene was investigated using a range of reaction conditions (Table 1). As expected, no reaction occurred in the absence of an FLP (Table 1, entry 1). Reaction with only the Lewis base component of the FLP (Mes3P) showed no cross-coupled product, and a stoichiometric or catalytic (10 mol %) amount of the Lewis acid B(C6F5)3 led to only 22 and 5% isolated yields of the desired Csp–Csp cross-coupled product, 2a (Table 1, entries 3 and 4). Stoichiometric amounts of both a Lewis acid and Lewis base were required for the Csp–Csp coupling reaction to attain satisfactory yields of the cross-coupled products. The B(C6F5)3/Mes3P FLP in toluene at 70 °C led to the Csp–Csp cross-coupled product, 2a, being formed in 54% yield (Table 1, entry 5). The optimum temperature was 70 °C with both lower (21 °C) and higher (110 °C) temperatures giving reduced yields (18 and 45% respectively) (Table 1, entries 6 and 7). More polar THF gave the highest yield of 83%, with trifluorotoluene (TFT) giving a 71% yield. CH2Cl2 and hexane, on the other hand, showed poorer or low yields (Table 1, entries 8–11). Interestingly, other Lewis acid boranes (BF3·OEt2 and BPh3) as well as Bronsted acids (CF3SO3H) showed no product formation when combined with Mes3P (Table 1, entries 12–14). Other basic phosphines including tBu3P, Ph3P and o-tol3P had complicated reaction mixtures with no or moderate yields of 2a being formed (Table 1, entries 15–17). Nitrogen Lewis bases including TMP (2,2,6,6-tetramethylpiperidine), DABCO (1,4-diazabicyclo(2,2,2)octane), and DMA (4-bromo dimethyl aniline) led to no product formation (Table 1, entries 18–20). The optimum conditions for the reaction were therefore chosen to be the use of the B(C6F5)3/Mes3P FLP in THF at 70 °C for 22–24 h.

**Reaction Scope.** With the optimized reaction conditions in hand, several aryl esters (1a–1l) were tested for the FLP-mediated Csp–Csp coupling reaction with various acetylenic compounds (Scheme 2). Diaryl esters bearing electron-withdrawing/π-releasing (p-F, 1a; p-Cl, 1b), neutral (p-H, 1c), and electron-donating (p-OME, 1d) groups all worked well for the reactions when coupled with aryl-substituted terminal acetylenes with electron-releasing (p-OME, p-tBu), neutral (p-H), electron-withdrawing/π-releasing (p-F, p-Cl), and electron-withdrawing (p-CF3) groups generating products 2a–q in 60–85% yields. Asymmetrical diaryl esters 1e and 1f were also found to undergo the Csp–Csp cross-coupling reaction, with several alkenes generating C–H-functionalized products 2r–w in excellent isolated yields (69–80%). We could also use alkyl/aryl esters containing just one aryl-stabilizing group. 1g could be cross-coupled with electron-neutral (phenylacetylene) and electron-releasing (4-ethynylanisole) alkenes to give 2x and 2y albeit in slightly lower yields of 61 and 65%, respectively. However, when cyclohexyl(phenyl)methyl-4-fluorobenzoate (1h) was employed, poor conversion was observed. Diaryl esters bearing strongly electron-withdrawing (p-CF3, 1i) groups also failed to react at all with terminal alkenes. We attribute this to the electron-deficient nature of the ester which is not Lewis basic enough to form an adduct with the Lewis acidic borane in the initial step of the reaction as observed by in situ 11B and 1H NMR spectroscopy. The unwillingness of ester 1j to react with arylicetlenes can also be interpreted from DFT calculations (see later and SI Figure S180).

After achieving good success for the Csp–Csp cross-coupling reaction at the benzylic position, we investigated a wider substrate scope (Scheme 3). To this end, aliphatic ester (E)-1,3-diphenylallyl-2,2-trifluorooacetate (1k) was used in the C–H functionalization. To our delight, excellent yields of products 2z–ah (72–89%) were obtained. While benzylic and allylic esters worked well, the same was not true for cross-coupling at

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**Table 1. Optimization of the Reaction Conditions for Csp–Csp Cross-Coupling Reactions**

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All of the reactions were carried out on a 0.1 mmol scale, and reported yields are isolated. All reactions were carried out for 20–22 h. 10 mol % B(C6F5)3.
Scheme 2. C$_{sp^3}$−C$_{sp}$ Cross-Coupling Reactions between Esters 1 and Acetylenes$^a$

Starting materials used in this study:

- $R^1 = p$-FC$_6$H$_4$
- $R = p$-ClC$_6$H$_4$
- $Ar = R = Ph$
- $Ar = R = p$-OMeC$_6$H$_4$
- $Ar = p$-ClC$_6$H$_4$, $R = Ph$
- $Ar = p$-MeC$_6$H$_4$, $R = Ph$
- $Ar = R = Cy$
- $Ar = R = p$-MeC$_6$H$_4$
- $Ar = R = p$-CF$_2$C$_6$H$_4$

$R^1 = p$-FC$_6$H$_4$; $R^2 = CF_3$

All reactions were performed on a 0.1 mmol scale.
the propargylic position. When the aryl/alkynyl ester 1-phenyl-3-(trimethylsilyl)prop-2-yn-1-yl 4-fluorobenzoate (1l) was employed with terminal acetylenes, an inseparable complicated reaction mixture resulted.

In our previous studies,6 we have shown that reactions of esters 1 with styrenes in the presence of the same FLP leads to $C_{sp^3}-C_{sp^2}$ coupled products. We therefore undertook an experiment to investigate the regioselectivity of the reaction by reacting the ester starting material with a 1:1 mixture of an acetylene and a styrene. For this reaction, three outcomes are theoretically possible: (i) formation of the $C_{sp^3}-C_{sp}$ coupled product, (ii) formation of the $C_{sp^3}-C_{sp^2}$ coupled product, or (iii) formation of a mixture of $C_{sp^3}-C_{sp}$ and $C_{sp^3}-C_{sp^2}$ coupled products. Using the optimized reaction conditions, an

![Scheme 3. $C_{sp^3}-C_{sp}$ Cross-Coupling Reactions between Ester 1k and Acetylenes](image)

![Table 2. Selectivity Reactions between Esters 1a and Acetylenes/Styrene](image)

![Scheme 4. Cross-Coupling Reactions between Esters 1a, c, e and 1-Ethynyl-4-vinylbenzene 4a](image)

![Table 3. Solvent-Dependent Site-Selective Studies](image)

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equimolar mixture of aryl ester 1a, 4-fluorophenylacetylene, and 4-fluorostyrene were reacted together with B(C₆F₅)₃/Mes₃P (Table 2).

Scheme 6. Products of Solvent-Dependent Site-Selective Cross-Coupling Reactions

Scheme 7. Possible Reaction Mechanisms

The crude ¹H NMR spectrum clearly showed a sharp singlet at δ = 5.09 ppm which confirmed the formation of the Csp³−Csp cross-coupled product, 2a, isolated as the major product in 78% yield (Table 2, entry 1). We were not able to detect any characteristic peaks (i.e., a doublet at δ = 4.98 ppm in the ¹H NMR spectrum) for the Csp³−Csp coupled compound. To investigate the effect of solvent on the reaction, we also conducted the reaction in TFT, where again 2a was isolated as the major product albeit with a slightly reduced yield of 72% (Table 2, entry 2). The observation of exclusive Csp³−Csp coupling is presumably a consequence of the higher reactivity of the alkyne functionality over the alkene. A similar competition experiment, using a 1:1 mixture of styrene and the internal alkyne diphenylacetylene, gave only Csp³−Csp coupling producing 3a in 63% yield (Table 2, entry 3).

Interestingly, TMS-protected alkynes behaved in the same manner as terminal alkynes, predominantly giving the Csp³−Csp coupling.
coupled product with the loss of the TMS group. Using a 1:1 mixture of styrene and trimethyl(phenylethynyl)silane afforded the Csp^3−Csp coupled product (2c, 58% isolated) as the major product with small amounts of the Csp^3−Csp_2 cross-coupled product (3a, 18% isolated) formed (Table 2, entry 4). This was also observed by in situ 'H NMR spectroscopy of the crude reaction mixture, which displayed a 1:0.4 ratio of Csp^3−Csp to Csp^3−Csp_2 cross-coupled products.

To demonstrate the scope for this selectivity, we synthesized a substrate containing both alkene and alkyne functionalities, namely, 1-ethynyl-4-vinylbenzene (4a). 4a was subsequently reacted with different aryl esters (1a,c,e) in the presence of the B(C_6F_5)_3/Mes_3P FLP. In agreement with the observation above, we observed only the formation of the Csp^3−Csp compounds as the major product (2ai−ak; 70−76%) using the optimized reaction conditions (Scheme 4). In all cases, the Csp^3−Csp_2 coupled product was either not detected or was observed in <5% yield in both THF and TFT solvents.

As for the intermolecular competition reactions, we also synthesized internal alkynes in which the acetylenic proton in 4a was replaced by a phenyl or TMS group to explore how this affected the regioselectivity of the reaction (Scheme 5). Using the optimized reaction conditions, the reaction of aryl ester 1a with 1-(phenylethynyl)-4-vinylbenzene (4b) exclusively gave the Csp^3−Csp_2 cross-coupled product in 71% isolated yield from the reaction with the alkene, whereas when 1a was reacted with trimethyl{(4-vinylphenyl)ethynyl}silane (4c), reaction at the alkyne and removal of the TMS group in situ afforded the Csp^3−Csp cross-coupled product (2ai) in 69% yield, with no significant reaction at the alkene site observed.

With these results in hand, we further explored the substrate scope using the 1-ethynyl-2-vinylbenzene (4d). 4d was reacted with ester 1a and the B(C_6F_5)_3/Mes_3P FLP under the optimized reaction conditions (THF, 70 °C, 24 h). Contrary to the reactions above, the reactions were found to be highly site-selective for the Csp^3−Csp_2 coupled product, 3c, from reaction at the alkene functional group. Examining the crude 'H NMR spectrum revealed a 0.2:1:0 ratio of the products (Csp^3−Csp coupled, 2)/Csp^3−Csp_2 coupled, 3)/(Csp^3−Csp coupled, 5) (Table 3, entry 1) with isolated yields of 79% for the Csp^3−Csp_2 coupled product, 3c, and 16% for the Csp^3−Csp coupled product, 2al (Scheme 6). Interestingly, when changing the solvent to TFT, the selectivity was completely reversed, exclusively giving Csp^3−Csp product 2al from reaction at the alkyne site.

This was observed by crude 'H NMR, indicating a 1:0:0 ratio of 2/3/5 (Table 3, entry 9) in which 2al could be isolated in 61% yield (Scheme 6). Remarkably, by simply changing the solvent we can switch the site selectivity of the reaction.

We next investigated this solvent-dependent site selectivity for a range of other esters and found the same general trend. In the following discussion, all reaction product ratios were determined via crude 'H NMR spectroscopy and are listed in Table 3, with the corresponding isolated yields for the products shown in Scheme 6. Initially, we explored the reactions in THF solvent. When electron-withdrawing 1b (p-Cl) or electron-neutral symmetrical diaryl esters 1c (p-H) and 1i (p-Me) were...
used, there was a clear preference for reaction at the alkene site leading to $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^2}$ coupled products $3d$, $3e$, and $3f$ in ratios of $0.2:1:0, 0.2:1:0$, and $0.4:1:0$ for 2/3/5, respectively (Table 3, entries 2–4). In all cases, the major and minor regioisomers could be separated. The $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^2}$ cross-coupled products were isolated in 74% ($3d$), 71% ($3e$), and 56% ($3f$) yields, and the $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^2}$ cross-coupled products were isolated in 15% ($2am$), 14% ($2an$), and 18% ($2ao$) yields. In the case of $1i$ as the starting material, double cross-coupled product $5d$ was observed in small amounts and could be separated and isolated in 10% yield (Scheme 6). Electron-rich $p$-OMe ester $1d$, on the other hand, showed no reactivity at all in THF (Table 3, entry 5). Asymmetrical diaryl ester $1e$ also gave the $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^2}$ cross-coupled product as the major isomer, albeit less selectively, showing a 0.5:1:0.1 ratio of the three products $2ar$, $3i$, and $5g$ in isolated yields of 24% ($2ar$), 40% ($3i$), and 13% ($5g$) (Table 3, entries 7 and Scheme 6). 1,3-Diphenylallyl-2,2,2-trifluoroacetate ($1k$), on the other hand, showed a preference for the double cross-coupled product, giving a 0:1:1.5 ratio of 2/3/5 with isolated yields of 22% ($3j$) and 41% ($5h$) (Table 3, entry 8 and Scheme 6).

Subsequently, we repeated the above series of reactions in TFT, and remarkably, the regioselectivity was altered and the selectivity was improved. No $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^2}$ coupled product ($3o$) or double $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^3}$ cross-coupled product ($5o$) was observed with any combination of substrates. Rather, a ratio of 1:0:0 of products 2/3/5 was observed in all cases for esters $1a$–$e$, $1g$, and $1i$ (Table 3, entries 9–15). This included $p$-OMe ester $1d$ which did not show any product formation in THF. $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^2}$ cross-coupled products $2al$–$ar$ could be isolated in 50–63% yields (Scheme 6). The only exception was 1,3-diphenylallyl-2,2,2-trifluoroacetate ($1k$), which gave a complex mixture of inseparable products when reacted with 1-ethynyl-2-vinylbenzene ($4d$) in TFT, none of which could be identified as 2, 3, or 5 (Table 2, entry 16).

Proposed Reaction Mechanism. The $\text{C}_{\text{sp}^3}-\text{C}_{\text{sp}^2}$ cross-coupling reaction could be explained by either a single- or a two-electron pathway.
First, a diamagnetic pathway could operate (Scheme 7A) in which the Lewis acid component of the FLP activates the ester carbonyl atom, leading to the formation of diaryl methylene cation $\text{I}_3$ and the borate anion $[\text{R}_1\text{CO}_2\text{B}(\text{C}_6\text{F}_5)_3]^{-}$. This was observed in our previous studies when $\text{B}(\text{C}_6\text{F}_5)_3$ was added to the diaryl ester in the presence of a nucleophile to trap the resultant carbocation. The diaryl methylene cation and the Lewis basic component of the FLP can then undergo a 1,2-FLP addition to the alkyne similar to other FLP 1,2-additions. Finally, the elimination of $[\text{Mes}_3\text{PH}]^{+}$ leads to the C–C-bonded product and salt $[\text{R}_1\text{CO}_2\text{B}(\text{C}_6\text{F}_5)_3][\text{HPMes}_3]$ ($\text{R}_1 = \text{p-FC}_6\text{H}_4$ or $\text{CF}_3$). This can be observed in the reaction by multinuclear NMR spectroscopy showing a clear $J_{\text{PH}}$ coupling of 479.5 Hz.

Alternatively, a radical pathway could operate (Scheme 7B), which may explain the necessity for using Mes$_3$P as a Lewis base rather than other phoshine or nitrogen bases. Previous studies have postulated that the B(\text{C}_6\text{F}_5)_3/Mes$_3$P FLP can undergo a single electron transfer (SET) process generating radical ion pair $[\text{B}(\text{C}_6\text{F}_5)_3]^{-}/[\text{Mes}_3\text{P}]^{+}$. Slootweg et al. later postulated that this process is promoted upon irradiation with visible light ($390$–$500$ nm). In this pathway (Scheme 7B), we propose that the first step of the reaction is identical to the diamagnetic pathway whereby B(\text{C}_6\text{F}_5)_3 activates the diaryl ester to generate diaryl methylene cation $\text{I}_3$ and borate anion $[\text{R}_1\text{CO}_2\text{B}(\text{C}_6\text{F}_5)_3]^{-}$. The Lewis base then reacts with cation $\text{I}_3$, forming a Lewis acid–base adduct. We propose that this adduct is in equilibrium with diaryl methylene radical $[\text{Ar}_2\text{CH}]^{-}$ and phosphonium radical cation $[\text{Mes}_3\text{P}]^{+}$, which are formed from the homolytic cleavage of the C–P bond. From here, diaryl methylene radical $[\text{Ar}_2\text{CH}]^{-}$ adds to arylacetylene, leading to a vinylic radical species which, upon abstraction of a hydrogen atom by $[\text{Mes}_3\text{P}]^{+}$, generates the desired C–C bonded product. To understand which pathway is operating, we undertook extensive electron paramagnetic resonance (EPR), kinetic, and density functional theory (DFT) studies to understand the reaction mechanism for the C$_{sp^3}$–C$_{sp}$ coupling reaction. 

**EPR Studies.** The knowledge that the B(\text{C}_6\text{F}_5)_3/Mes$_3$P FLP can generate radical species prompted us to undertake an EPR study to determine if radical species could be observed in these reactions. As reported previously, no EPR signal could be detected from the B(\text{C}_6\text{F}_5)_3/Mes$_3$P FLP in the absence of any substrates. Upon addition of an equimolar ratio of ester to the FLP in a TFT solution, several EPR signals arising from multiple paramagnetic species were detected at room temperature (Figure 1A). Upon comparison with previous reports, the two intense resonance lines in a 1:1 ratio (marked with asterisks) centered on $g_{\text{iso}} = 2.012$ ($B \approx 335$ mT) and separated by a phosphorus hyperfine splitting of $a_{\text{iso}}(31\text{P}) = 670$ MHz (23.8 mT) are attributed to the formation of the $[\text{Mes}_3\text{P}]^{+}$ cation. A second paramagnetic species was also detected in this sample, which is characterized by a 1:2:1 triplet centered on $g_{\text{iso}} = 2.010$ and separated by a hyperfine splitting of $670$ MHz (23.8 mT). This EPR profile must originate from two identical $I = 1/2$ nuclei, in this case associated with two equivalent $^{31}$P nuclei. The signal is therefore tentatively assigned to the formation of a $[\text{P}(\text{Mes}_3)]^{+}$ dimer formed from the association of excess Mes$_3$P with radical cation $[\text{Mes}_3\text{P}]^{+}$ as observed previously.
supported by our observation that the relative ratios of the EPR signals of [Mes₃P]⁺/[P(Mes)ₙ=2,3]₂⁺ are inter-related (i.e., when a large signal intensity of the monomer is observed, only a trace of the corresponding dimer is detected). The varying ratio of these EPR signals under different reaction conditions demonstrates the conversion between monomer and dimer via the reaction of the monomer with a second molecule of phosphine to yield the dimer radical cation, as previously observed for a series of phosphines, and possibly also the varying stabilities of the two radical species. The EPR spectrum of [(PMes₂)₂]⁺ has previously been reported, characterized by $g_{iso} = 2.006$ and $a_{iso}^{(1)P} = 474$ MHz (17 mT), and it is noted that literature examples of phosphine dimer cation radicals of divalent ($\text{R}_2\text{P}$)₂⁺ and trivalent ($\text{R}_3\text{P}$)₃⁺ systems yield very similar EPR spectra, dominated by the 1:2:1 phosphorus hyperfine splitting.

The corresponding anisotropic EPR spectra of the [Mes₃P]⁺⁺ monomer and [P(Mes)ₙ=2,3]₂⁺ dimer species in frozen solutions are shown in the SI (Figure S176), from which the principal values of the $g$ and $A$ tensors were determined. The spin Hamiltonian parameters for all of the paramagnetic species detected in this work are listed in the SI (Table S1). Importantly, contrary to our previous reports, no evidence for the formation of the carbon-based bismethoxy-diphenylmethylene radical formed upon C–O bond cleavage was observed in this case, perhaps due to the instability of the radical species.

We then probed the room-temperature EPR spectrum of the FLP in the presence of phenylacetylene (Figure 1B,C). Under these experimental conditions, we postulated that another possible mechanism could be the abstraction of the terminal hydrogen atom of the acetylene by the [Mes₃P]⁺⁺ radical cation to form the diamagnetic [Mes₃PH]⁺⁺ cation and the corresponding phenylacetylene radical. As can be seen from the wide field scanning range in Figure 1B, no evidence of monomer [Mes₃P]⁺⁺ or dimer [P(Mes)ₙ=2,3]₂⁺ was observed in this solution, suggesting the formation of the diamagnetic [Mes₃PH]⁺⁺ cation. However, no EPR evidence for the generation of the phenylacetylene radical was obtained. It is noted in previous literature studies that the terminal phenylacetylene radical is inherently unstable and is typically observed only via EPR spectroscopy under controlled conditions, such as neat liquids sealed under vacuum, or via matrix isolation methods. Notably, the remaining boron component of the FLP is not involved in the above hydrogen atom abstraction from the alkyne and may be expected to

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**Figure 5.** DFT-computed reaction pathways for the reaction of I₃ with 1-ethynyl-4-vinylbenzene (4a) and Mes₃P calculated using the SMD/B3LYP-D3/def2-TZVP//SMD/B3LYP-D3/6-31G(d) level of theory in THF.
remain in solution. The intense multiplet signal observed, reproduced in Figure 1C across a narrow field range, is therefore assigned to the boron radical anion, [B(C₆F₅)₃]⁻. A satisfactory simulation of the experimental data was achieved using ξiso = 2.0114 and incorporating a single boron nucleus, aiso¹⁹F(B) = 27 MHz, two sets of six equivalent fluorine nuclei from the ortho and meta positions, with aiso страх(¹⁹F)ortho = 18.28 MHz and aiso страх(¹⁹F)meta = 3 MHz, and three para fluorine nuclei with aiso страх(¹⁹F)para = 20.2 MHz, which agrees well with previous literature reports of this species. The corresponding anisotropic spectrum for this sample was unfortunately not resolved due to a poor-quality glass of the frozen solvent, thereby preventing the determination of the complete anisotropic spin Hamiltonian parameters for this radical anion.

Having determined the reactivity of the FLP to the individual substrates, the EPR spectrum of a full reaction mixture containing equimolar amounts of Mes₃P, B(C₆F₅)₃, and 2-vinylbenzene 4d, calculated at the SMD/B3LYP-D3/def2-TZVP//SMD/B3LYP-D3/6-31G(d) level in THF and toluene.

Figure 6. DFT calculations of the site selectivity of 13 with 1-ethyl-2-vinylbenzene 4d, calculated at the SMD/B3LYP-D3/def2-TZVP//SMD/B3LYP-D3/6-31G(d) level in THF and toluene.

experimental and simulated data, thereby this signal is tentatively attributed to a carbon-based radical, perhaps indicating that the reaction could be occurring through a radical mechanism.

**DFT and Kinetic Studies.** To examine the contrasting reaction pathways and to explain the experimental and EPR observations, we undertook a thorough DFT investigation of all potential reaction pathways. Calculations were performed at the SMD/B3LYP-D3/def2-TZVP//SMD/B3LYP-D3/6-31G(d) level of theory in THF and toluene solvent to examine the origin of the products. As previously reported by Slootweg et al., we found that the formation of the frustrated radical ion pair from the FLP is energetically unfavorable by 35.7 kcal/mol. This corroborates the observation that we and others’ fail to see any EPR signal in solutions of B(C₆F₅)₃ and Mes₃P. Very recently, Slootweg et al. showed that the coordination of B(C₆F₅)₃ to the diaryl ester increases the electron affinity of the substrate, and the energy required for SET from Mes₃P to the methylene carbon atom is 40.0 kcal/mol. This is still quite large, and our calculations have revealed that, regardless of whether a diamagnetic or paramagnetic reaction pathway is ultimately operative, the first step in the reaction is the same: B(C₆F₅)₃ activation of the ester to generate diaryl methylene cation 13 (Scheme 8) and the boron anion [R'CO₂B(C₆F₅)₃][−] with an activation energy of 10.7 kcal/mol. (See SI Figure S178 for the free-energy profile.) The energies of 13 can be noticeably varied by changing the substitution at the para position of the aryl esters. Electron-withdrawing (−p-F, 1a), electron-withdrawing/π-releasing (−p-F, 1a), and electron-releasing (−p-OMe, 1d) showed different energies for 13 (SI Figure S179–180). As expected, 13d (−9.4 kcal/mol) is energetically more favorable than 13a (13.5 kcal/mol) due to the electron-releasing substituents (p-OMe). Strongly electron-withdrawing substituents (−p-Cl) conversely make 13 formation thermodynamically less favorable. 13 is then the branching point for the single- and two-electron pathways (Figure 2). The combination of diaryl methylene cation 13 with Mes₃P can lead to three possible species in solution (Scheme 8): (i) the frustrated Lewis pair (uncoordinated 13 + Mes₃P), (ii) the Lewis acid–base adduct (14), and (iii) the frustrated radical pair (FRP, 15). The energy difference and reaction barriers between these species are very low; therefore, it is likely that all three scenarios exist in equilibrium under the reaction conditions. This supports the EPR data which shows the formation of [Mes₃P][⁺] and the [(P(Mes₃)₂)₂][⁺] dimer in the reaction of the ester with the Mes₃P/B(C₆F₅)₃ FLP.

Although the carbon-based radical could not be observed in this case, we have observed a weak carbon-based radical EPR signal when reacting other esters with the Mes₃P/B(C₆F₅)₃ FLP in the absence of irradiation or heat. We then investigated the addition of the cation (13) or radical (16) to the alkyn. Although both pathways are feasible under the reaction conditions, the cationic pathway was lower in activation energy than the radical pathway by about 26.3–18.6 = 7.7 kcal/mol. In the case of the diamagnetic pathway, the addition of the 13 cation to the alkyn generates 18 via TS5. The resulting cation in 18 is highly reactive and is rapidly trapped by the Lewis base Mes₃P generating 19. Finally, anti-elimination of [Mes₃PH][⁺] generates the cross-coupled compound and phosphonium borate salt as the final products.

The Lewis base employed is very important for the reaction to occur as seen in the screening studies. First, the ability to
form an FLP (or weak adduct) with both the Lewis acid borane and the (d)aryl methylene cation (I3) is critical, as other strong, less hindered Lewis acids such as BF₃ do not work in the reaction. Likewise, smaller phosphines or amines tend to coordinate more strongly to the carbocation (I3). In addition to being able to form an FLP, the base also functions to trap reactive I8. Indeed, one could possibly conceive that the reaction could proceed with the Lewis acid only, whereby RS⁺ accepts the proton in the last step (I8 → I10 → product, Figure 2). However, DFT calculations showed that this pathway was less favorable thermodynamically, and experimentally this reaction showed only a 22% yield with many side products formed.

We were curious to know whether both the paramagnetic and diamagnetic pathways are operative in parallel or if a diamagnetic mechanism is purely responsible for the product formation for all alkynes and the radicals observed from EPR studies were simply off-pathway intermediates. To establish this, we undertook further DFT calculations to investigate the effect of electron-withdrawing, electron-donating, and electron-neutral substituents on the ester (1) and the alkyne on the activation barrier for the reaction. The energy barriers TS₄ radical pathway and TS₅ cationic pathway were calculated (Table 4, see SI Figure S181). Initially we varied the electronic properties of the alkyne using p-XC₆H₄CH=CH (X = NO₂, CF₃, H, OMe, NMe₂) with ester 1a. As evidence from the DFT calculations, changing the substitution at the para position of the arylacetylene did not make significant difference for TS₄ radical pathway in their respective energy barrier (23.9–27.0 kcal/mol) (Table 4, entries 1–5). However, the energy barrier for TS₅ cationic pathway changed dramatically (8.2–23.2 kcal/mol). When electron-withdrawing groups (p-NO₂ and p-CF₃) on the arylacetylene were employed, the differences between TS₄ radical pathway and TS₅ cationic pathway are 0.7 and 6.1 kcal/mol (Table 4, entries 1 and 2). Electronically neutral phenylacetylene exhibits a TS₄ radical pathway → TS₅ cationic pathway difference of 7.7 kcal/mol (Table 3, entry 3). Electron-donating groups such as methoxy (TS₄ radical pathway − TS₅ cationic pathway = 12.1 kcal/mol) and N,N-dimethylaniline (TS₄ radical pathway − TS₅ cationic pathway = 18.8 kcal/mol), on the other hand, showed a significant energy difference (Table 4, entries 4 and 5). These observations can be seen in Figure 3, which shows that for TS₄ there is a negligible change in the energy barrier when changing the electronic properties of the acetylenic substrate. This is in agreement with little charge formation on the reaction center in the radical mechanism. Conversely, for TS₅, there is a strong positive correlation between the TS₅ energy barrier and the substituent σp constant. This is expected for the cationic pathway because a developing positive charge adjacent to the substituted phenyl ring will be stabilized by electron-donating groups (e.g., p-NMe₂ or p-OMe) on the acetylene. As can be seen in Figure 3, TS₅ is lower than TS₄ for all substituents explored, although the difference becomes small for strongly electron-withdrawing groups.

We subsequently computed the energy barrier for the two transition states by varying the electronic properties of the aryl ester using electron-donating (p-OMe, 1d) and electron-withdrawing (p-CF₃, 1j) esters with electron-deficient (1-ethyl-1-(trifluoromethyl)benzene) and electron-rich (1-ethyl-1-methoxybenzene) acetylenes (Table 4, entries 6–9). For both esters, a smaller change in the TS₄ radical pathway energy barrier was observed (range 22.2–27.8 kcal/mol) compared to the TS₅ cationic pathway energy barrier (range 8.1–25.5 kcal/mol) when changing the substituent on the phenylacetylene (Table 4, entries 6–9). As with ester 1a, both esters disclosed a larger energy difference between the two pathways when reacted with acetylenic compounds bearing an electron-donating group (p-OMe). Likewise, a much smaller energy difference was noted for both esters when reacted with acetylenic compounds bearing an electron-withdrawing group (p-CF₃). Interestingly, DFT studies showed that for the case of the reaction of ester 1j with 1-ethyl-1-(trifluoromethyl), the radical pathway is slightly energetically more favorable compared with the cationic pathway (Table 4, entry 8). These results suggest that for electron-withdrawing arylacetylenes both paramagnetic and diamagnetic mechanisms are potentially possible, whereas for electron-rich arylacetylenes a purely diamagnetic pathway is operative.

The key difference in the two mechanisms involves the reaction of either a cationic intermediate or a radical intermediate with the arylacetylene, generating a new cationic or radical species. Whether this new intermediate is a cationic or a neutral radical species can be probed using a Hammett plot (cf. computational studies) based on substituted arylacetylenes. To gain this insight into the reaction pathway and substituent effects also from experimental evidence, we examined competition reactions among the FLP, aryl ester 1a, and arylacetylenes p-XC₆H₄CH=CH bearing electron-withdrawing, electron-neutral, and electron-releasing groups. The Hammett plot requires relative rate constants for the reaction of different substituted alkynes that were obtained using a series of competition experiments. Initial competition experiments in the presence of 1.5 equiv of five arylacetylenes p-XC₆H₄CH=CH (X = CF₃, F, Cl, H, and OMe) were unsuccessful. The excess arylacetylene present in the reaction mixture destroyed the efficacy of the FLP system, producing a complicated reaction mixture which was not suitable for in situ NMR analysis. We therefore carried out three binary competition experiments with two alkynes being present at 1.5 equiv each (Table 5).

Using the optimized reaction conditions, three parallel reactions were carried out in which equimolar mixtures of (a) 1-ethyl-1-(4-methoxybenzene) and phenylacetylene, (b) 1-ethyl-1-(4-(trifluoromethyl)benzene and phenylacetylene, and (c) 1-ethyl-1-(4-methoxybenzene) and 1-ethyl-1-(4-(trifluoromethyl)benzene were reacted with 1 equiv of ester 1a. The ratios of Csp₂=Csp₂ cross-coupled products (2d/2c, 2b/2c, and 2d/2b) were determined from the crude reaction mixture using 19F NMR spectroscopy (Table 5, see SI, Figure S175). For entries 1 and 2 in Table 5, the relative integrals for the products were used to calculate the remaining equivalents for the alkynes after reaction. Using the approach developed by Ingold and Shaw and proposed for one-pot Hammett plots by Harper and co-workers, we obtained relative rate constants kᵣ/kᵢ for the reaction of differently substituted arylacetylenes with reaction intermediate I3 or its equilibrium species. Entry 3 confirms that in the competition between 1a and 1-ethyl-1-(4-(trifluoromethyl)benzene)/1-ethyl-1-(4-methoxybenzene, product 2b is undetectable, in agreement with the >200-fold difference in rate constants deduced from entries 1 and 2. The relative rate constants in Table 5 are normalized with respect to the unsubstituted aryl and can therefore be used directly to construct a Hammett plot (using Hammett substituent constants from ref 32). The Hammett plot (Figure 4) shows a clearly negative slope of −6.6 ± 1.7.
suggesting the formation of a positive charge on the new intermediate, which is indicative of the cationic reaction mechanism being operative. The Hammett plot is also in agreement with the Hammett-style plot constructed using the computational data (Figure 3) for the cationic mechanism. The slope of the Hammett-style plot for TSS (Figure 3) is 8.9 ± 1.2 kcal mol⁻¹. At 70 °C, this corresponds to a Hammett ρ value of 5.7 ± 0.8. The computational data is thus in excellent agreement with the experimental findings.

Finally, we turned our attention to explaining the regioselectivity with compound 4a. The DFT-computed (SMD/B3LYP-D3/de2-TZVP/SMD/B3LYP-D3/6-31G(d)) reaction pathways for the reaction of I3 with 1-ethynyl-4-vinylbenzene (4a) and Mes₃P in THF reveal that the cationic pathway is energetically more favorable (Figure 5). After the generation of I3, reaction at either the alkyne or alkene site affords the corresponding C(sp)₂=C(sp)₃ or C(sp)₃=C(sp)₂ cross-coupled product. Although the transition-state energies for the I3-alkyne adduct (14.4 kcal/mol) and I3-alkene adduct (14.3 kcal/mol) are very similar, the C(sp)₂=C(sp)₃ cross-coupled products are thermodynamically more stable (−1.4 kcal/mol) than the C(sp)₃=C(sp)₂ cross-coupled product (5.7 kcal/mol), explaining why only the C(sp)₂=C(sp)₃ cross-coupled product is observed for 1-ethynyl-4-vinylbenzene (4a) (Figure 5). To confirm this, we also executed single-point benchmark calculations for this transition state with a different method and solvent system (SI, Table S2), which showed results similar to those indicated in Figure 5.

The alternating site selectivity when using 1-ethynyl-2-vinylbenzene (4d) in differing solvents can also be highlighted in DFT studies. For the calculations, the mechanism was studied by utilizing two different solvents (toluene and THF). Experimentally, both toluene and TFT solvents lead to preferential C(sp)₂=C(sp)₃ coupling. DFT calculations for the reaction of I3 with 4d showed that the transition-state energy for the addition of the diaryl methylene cation to the alkyne or alkyne varies depending upon the solvent (Figure 6). In toluene solvent, the energy barrier for the addition of I3 to the more nucleophilic alkyne was 3.7 kcal/mol lower in energy than the transition state for addition to the alkyne (11.7 versus 15.4 kcal/mol). The converse was true for reactions in THF, whereby the addition of I3 to the alkyne was 0.5 kcal/mol lower in energy than the energy barrier for addition to the alkyne (14.7 versus 15.2 kcal/mol). We attribute this to the higher dipole moment of the calculated transition-state structure of TS3a, which can be stabilized better by the THF molecule. These results also explain why, in THF, the reactions were less selective, leading to a mixture of C(sp)₂=C(sp)₃ and C(sp)₃=C(sp)₂ products due to the small energy difference between the two pathways. The reactions in a solvent such as toluene (or TFT) are more selective toward C(sp)₂=C(sp)₃ coupling due to the larger energy difference between the two pathways.

**CONCLUSIONS**

We have demonstrated new reactivities of FLPs in the functionalization of terminal alkyynes through C(sp)₂=C(sp)₃ coupling reactions with aryl esters. DFT studies found that a diamagnetic pathway was most likely, although a low-energy single-electron pathway could operate to some extent. In particular, DFT studies indicate that the combination of the Mes₃P/diaryl methylene cation led to three species of similar energy in solution: the FLP (I3), the Lewis acid–base adduct (I₄), and the frustrated radical pair (I₅). According to the Curtin–Hammett principle, the reaction proceeds predominantly via TSS from rapidly equilibrating I₃, I₄, I₅, and I₆. These rapidly equilibrating species in solution are supported by the observation of radical species of varying stabilities and lifetimes in the reaction mixture. Thus, radical species are formed in the reaction but are not making a substantial contribution on the reaction pathway to the product, with the possible exception of arylacetylenes with strongly electron-withdrawing (e.g., p-NO₂, p-CN) substituents. These observations will be of importance when designing future reactions that can switch between one- and two-electron pathways depending upon the substrate. Moreover, we observed high site selectivity when ethynyl vinylbenzene substrates were employed in these reactions. 1-Ethynyl-4-vinylbenzene substrates reacted only at the alkyne site, but 1-ethynyl-2-vinylbenzene substrates showed high selectivities depending upon the polarity of the solvent. For 1-ethynyl-2-vinylbenzene in THF, C(sp)₂=C(sp)₃ coupling was observed, resulting in alkyne functionalization, whereas in toluene or TFT exclusive C(sp)₃=C(sp)₂ coupling and alkyne functionalization resulted. The contrasting selectivity was explained by DFT and computed transition states in the differing solvents. FLP-mediated C=C bond-forming reactions are still relatively new, but there is no doubt that advances will continue to be made in this area. This reported methodology can be utilized to generate compounds that can be subsequently employed for the synthesis of useful novel natural products where metal-free synthesis is highly desirable for avoiding metal toxicities.⁵,³³
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Notes
The authors declare no competing financial interest.

REFERENCES


