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$B(C_6F_5)_3$ -catalyzed dehydrogenative cyclization of *N*-tosylhydrazones and anilines *via* a Lewis adduct: a combined experimental and computational investigation[†]

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Tris(pentafluorophenyl)borane-catalyzed dehydrogenative-cyclization of *N*-tosylhydrazones with aromatic amines has been disclosed. This metal-free catalytic protocol is compatible with a range of functional groups to provide both symmetrical and unsymmetrical 3,4,5-triaryl-1,2,4-triazoles. Mechanistic experiments and density functional theory (DFT) studies suggest an initial Lewis adduct formation of *N*-tosylhydrazone with B(C₆F₅)₃ followed by sequential intermolecular amination of the borane adduct with aniline, intramolecular cyclization and frustrated Lewis pair (FLP)-catalyzed dehydrogenation for the generation of substituted 1,2,4-triazoles.

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Introduction

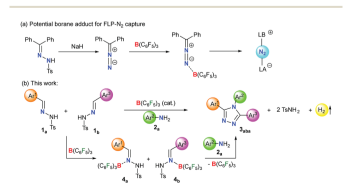
Tris(pentafluorophenyl)borane has recently emerged as a powerful Lewis acid catalyst.¹⁻³ The strong Lewis acidity^{2,3} at the boron center allows us to establish a wide range of organic transformations *via* C–B,⁴⁻⁶ C–C,⁷⁻¹⁰ C–N,^{11,12} C–O^{13,14} and C– Si¹⁵⁻¹⁸ bond formations. Pioneered by Stephan and Erker, $B(C_6F_5)_3$ has gained popularity in frustrated Lewis pair (FLP) chemistry which encompasses widespread applications in organic reactions.¹⁹⁻²² Indeed, numerous efforts have been devoted to the activation of alkenes and alkynes by FLPs for the formation of cyclic scaffolds.²³⁻²⁵ However, in many cases, the highly desirable catalytic reaction is obstructed by the initially formed stable borate adduct.²⁶⁻²⁸ Consequently, $B(C_6F_5)_3$ catalyzed cyclization leading to important heterocyclic scaffolds is rare.²⁹⁻³²

In this context, 1,2,4-triazoles are omnipresent heterocyclic motifs in numerous biologically active compounds^{33,34} and they also have widespread applications in organic light emitting diodes,^{35,36} organic photovoltaic cells, electroluminescent devices,^{37,38} bi-stable resistive memory devices,³⁹ pesticides, and medicines.⁴⁰ Given their applications, a number of strategies have been implemented for the synthesis of 1,2,4-triazoles.^{41–45} However, most of these methods are limited due to the use of a super-stoichiometric amount of reagents or oxidants, low

chemo-selectivity, narrow functional group tolerance, multiple reaction steps, and the production of copious waste. And catalytic metal- and oxidant-free, step-economical processes for an efficient synthesis of substituted 1,2,4-triazoles are in demand.

Lewis acid–base adducts are potential intermediates in a multitude of transformations.^{46–48} Recently, Stephan and coworkers have reported that diphenyldiazomethane, obtained from the corresponding *N*-tosylhydrazone in the presence of a base, readily forms adducts with $B(C_6F_5)_3$ (Scheme 1a).^{49,50} This has shifted the gear for metal-free N₂ activation closer to reality. Conversely, the direct interaction of *N*-tosylhydrazones with $B(C_6F_5)_3$ and their application in catalytic transformations have not been reported yet.

On the other hand, environmentally benign acceptorless catalytic dehydrogenation which is highly challenging even for transition-metal complexes^{51,52} is rare under metal-free conditions.^{53–56} Very recently, we have presented a manganese-



Scheme 1 Applications of borane adducts in main group chemistry.

Department of Chemical Sciences, Indian Institute of Science Education and Research Kolkata, Mohanpur-741246, India. E-mail: koley@iiserkol.ac.in; bm@iiserkol.ac.in † Electronic supplementary information (ESI) available. CCDC 1866759. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9sc02492a

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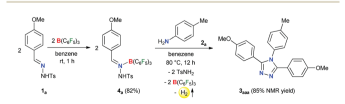
catalyzed acceptorless-dehydrogenative olefination of heteroarenes with primary alcohols.⁵⁷ Herein, we report $B(C_6F_5)_3$ catalyzed acceptorless-dehydrogenative-cyclization of *N*-tosylhydrazones **1** and anilines **2** to triaryl-1,2,4-triazoles **3** *via* a borane adduct **4** followed by sequential C–N/N–N bond formation (Scheme 1b). Furthermore, extensive DFT calculations are performed not only to understand the mechanistic features of borane catalyzed triazole formation but also to assist future development of similar classes of reactions.

Results and discussion

Initially, to check the reactivity of hydrazone 1_a with $B(C_6F_5)_3$, a stoichiometric reaction in benzene at room temperature was carried out. It leads to the formation of a Lewis adduct 4_a ($\delta^{11}B - 2.5$ ppm) after 1 h in 82% yield (Scheme 2). When 4_a was reacted with an aromatic amine 2_a at 80 °C in benzene, 3,4,5-triaryl-1,2,4-triazole 3_{aaa} was formed in 85% NMR yield after 12 h along with an equimolar amount of TsNH₂ and $B(C_6F_5)_3$.

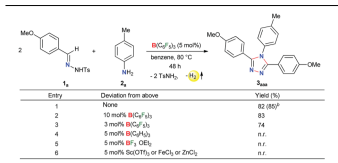
As $B(C_6F_5)_3$ was finally released from the adduct 4_a after its reaction with the amine 2_a , it was posited that catalytic turnover should be possible under thermal conditions. Interestingly, when the reaction was implemented with 5 mol% $B(C_6F_5)_3$ at 80 °C 1,2,4-triazole 3_{aaa} was obtained in 82% isolated yield along with 68% of TsNH₂ as a byproduct (Table 1, entry 1). While the increased catalyst loading did not significantly improve the yield (entry 2), a slight decrease in the product yield was observed with 3 mol% $B(C_6F_5)_3$ (entry 3). The use of less Lewis acidic boranes such as BPh₃ was unproductive (entry 4). In addition, less hindered boranes like BF₃·OEt₂ resulted in no detectable product formation (entry 5). Other Lewis acid catalysts like Sc(OTf)₃, FeCl₃ and ZnCl₂ were also not effective for this reaction (entry 6, Schemes S9 and S10 in the ESI[†]).

Next, the scope of this metal-free protocol was explored. A series of hydrazones 1 could be employed with various anilines 2 to afford symmetrical 3,4,5-triaryl-1,2,4-triazoles 3 in good to excellent yields (Table 2). The reaction of 1_a and 1_b with anilines 2_{a-f} bearing OMe-, Me-, H-, Cl- and Br-substituents at the *para*-or *meta*-position of the aryl ring afforded the triazoles **3aab**-**3bbf** in 66–87% yields. Similarly, the hydrazones 1_{c-g} having substituents with different steric and electronic properties at the *ortho*-, *meta*-, and *para*-position on the aromatic rings could readily be cyclized with 2_a to produce **3cca**-**3gga** in 71–86% yields. Electronically biased aryl rings could be installed smoothly to obtain 3_{ccb} , 3_{hbb} , and 3_{iic} in 56–78% yields. Likewise, thiophene- and fluorine-containing 1,2,4-triazoles 3_{ceh} and 3_{iig} could be synthesized in 79% and 84% yields,



Scheme 2 Borane adduct of N-tosylhydrazone 1_a and its transformation.

Table 1 Optimization of $B(C_6F_5)_3$ catalyzed cyclization of the hydrazone $\mathbf{1}_a$ with anilines $\mathbf{2}_a^a$



^{*a*} Reaction conditions: **1**_a (0.5 mmol), **2**_a (0.25 mmol), and $B(C_6F_5)_3$ (5 mol%) in 2.0 mL benzene; isolated yield. ^{*b*} NMR yield using 1,3,5-trimethoxybenzene as the internal standard. n.r. = no reaction.

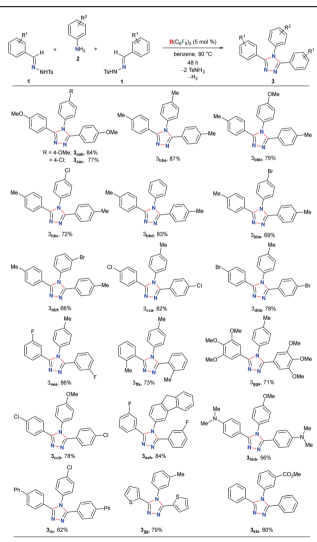


Table 2 $B(C_6F_5)_3$ catalyzed synthesis of symmetrical 1,2,4-triazoles^a

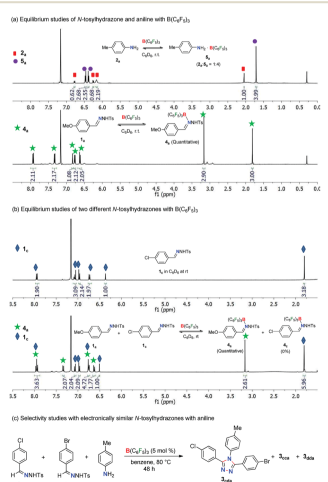
^{*a*} Reaction conditions: Table 1, entry 1. Yields of the analytically pure product.

respectively. Notably, an ester-group could also be tolerated under the reaction conditions to furnish 3_{kki} in 80% yield. On the other hand, substrates containing amides, nitro groups, olefins, primary alcohols, primary amino groups, *N*-tosylhydrazone of aliphatic aldehydes, and *N*-methanesulfonyl hydrazone were found to be unsuitable as either the starting materials remained intact or a complex mixture of products was formed (Schemes S1–S5 in the ESI[†]).

To further demonstrate the applicability of our protocol, we examined the possibility of obtaining unsymmetrical 1,2,4-triazoles by employing two different hydrazones (Table 3). In fact, the synthesis of unsymmetrically substituted 1,2,4-triazoles is considered to be highly challenging and to the best of our knowledge, their single step synthesis from readily accessible starting materials is less explored.⁴¹⁻⁴⁵ Gratifyingly, when 2_a was reacted with an equimolar mixture of 1_a and 1_c the unsymmetrical 1,2,4-triazole 3_{aca} was obtained as a major product in 77% yield and symmetrical triazoles were only obtained in minor amounts. The products could be purified via column chromatography on silica-gel using an ethyl acetate/hexane mixture as the eluent. Similar reactivity and selectivity were also observed for the synthesis of 3_{ada} and 3_{aea} . Likewise, biphenyl-, haloaromatic- and heteroaromatic-rings could be installed without any difficulty to give the triazoles 3_{aic} , 3_{aia} , 3akc, 3bcb and 3bdb in 61-80% yields. Colorless crystals of 3aic,

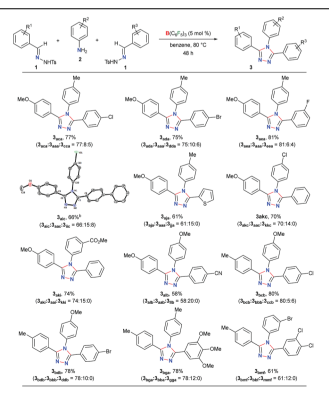
grown from a saturated benzene solution stored at room temperature, were suitable for single-crystal X-ray analysis and clearly confirmed the structure of the product (Fig. S3 in the ESI†).⁵⁸ It is noteworthy that the reaction is highly chemoselective as the reactive carbomethoxy- and cyano-groups remain unaffected under the present reaction conditions to provide 3_{aki} and 3_{alb} in 74% and 58% yields, respectively. The steric hindrance was found to exert a negligible influence on the reaction to give the desired product 3_{bmf} in good yields.

In order to obtain insight into the reaction mechanism, several equilibrium studies were performed initially with *N*-tosylhydrazones and anilines in the presence of Lewis acids (Scheme 3). At the onset, the relative Lewis basicity of aniline and *N*-tosylhydrazone towards $B(C_6F_5)_3$ was analyzed (Scheme 3a). Thus, in a stoichiometric reaction of 2_a and $B(C_6F_5)_3$ at room temperature in C_6D_6 , the ratio of 2_a and 5_a ($B(C_6F_5)_3$) adduct of 2_a) was found to be 1:4 ($2_a:5_a=1:4$). Similarly, the stoichiometric mixture of 1_a and $B(C_6F_5)_3$ in C_6D_6 at room temperature afforded quantitative formation of 4_a ($B(C_6F_5)_3$) adduct 1_a) where the ration was $1_a:4_a=0:100$. This confirms the tendency of *N*-tosylhydrazone 1 to form a stronger Lewis acid–base adduct with $B(C_6F_5)_3$ in comparison to aniline 2.



(3_{cda}:3_{cca}:3_{dda} = ~1:1:1)

Table 3 $B(C_6F_5)_3$ catalyzed synthesis of unsymmetrical 1,2,4-triazoles^{*a*}

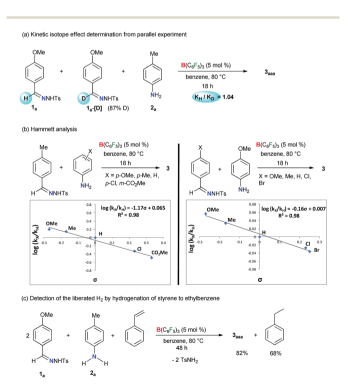


^a Reaction conditions: Table 1, entry 1 in the 0.5 mmol scale. Yields of the analytically pure product. Selectivities are given within parenthesis.
 ^b Thermal ellipsoids are shown at a 60% probability level. Ref. 58.

Moreover, a competitive equilibrium study was performed with two electronically biased N-tosylhydrazones $(1_a vs. 1_c)$ with $B(C_6F_5)_3$ (Scheme 3b). Accordingly, a stoichiometric (1:1)mixture of $\mathbf{1}_{a}$ and $\mathbf{1}_{c}$ was treated with 1 equivalent of $B(C_{6}F_{5})_{3}$ in C₆D₆ at room temperature. This selectively afforded the Lewis acid-base adduct $\mathbf{4}_a$ whereas $\mathbf{1}_c$ remained unreacted $(\mathbf{4}_a:\mathbf{1}_c=$ 1:1). Thus, *N*-tosylhydrazones having electron rich arenes will form strong Lewis acid-base adducts than hydrazones having electron deficient arenes. This is possibly a crucial factor for the selectivity observed during the synthesis of unsymmetrical 1,2,4-triazoles as shown in Table 3. Along this direction, in fact, the reaction of aniline 2_a with a mixture of 1_c and 1_d having electronically similar substituents (Cl and Br) on arenes in the presence of the B(C₆F₅)₃ catalyst provided a mixture of two symmetrical and unsymmetrical 1,2,4-triazoles (Scheme 3c) as both the hydrazones have similar probabilities for the formation of Lewis acid-base adducts with $B(C_6F_5)_3$. Thus, electronically biased hydrazones are good candidates for better selectivity of unsymmetrical triazoles.

In the case of competitive equilibrium studies of two different anilines with $B(C_6F_5)_3$, a stoichiometric (1:1:1) mixture of 2_a , 2_c and $B(C_6F_5)_3$ in C_6D_6 at room temperature gave their corresponding Lewis acid-base adduct 5_a and 5_c , respectively in an 8:1 ratio (Scheme S8 in the ESI†). This indicates that highly basic anilines will form stronger adducts with $B(C_6F_5)_3$. It is also noteworthy that other Lewis acids like Sc(OTf)₃ and BPh₃ did not form any Lewis acid-base adducts with *N*-tosylhydrazones or anilines under similar reaction conditions (Schemes S9 and S10 in the ESI†).

In addition, the kinetic isotope effect (KIE) of 1.0 measured from a parallel reaction of 2_a with 1_a and its deuterated

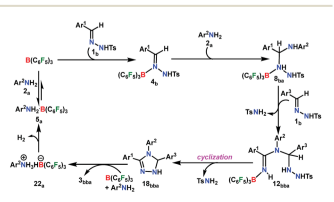


Scheme 4 Kinetic and mechanistic studies for the synthesis of 1,2,4-triazole.

analogue 1a-[D] suggests that the breakage of the imine C-H bond of 1_a is not involved in the rate-determining step (Scheme 4a). Kinetic monitoring of the reaction suggests an exponential decay of both the reactants 1_a and 2_a , whereas sigmoidal increase of 3_{aaa} was observed (Fig. S1 in the ESI[†]). The initial rate curve for triazole product formation possibly indicates that at the beginning, product formation is slower due to accumulation of reactive intermediates. Further kinetic studies revealed that the reaction is first order in the catalyst (Scheme S11 in the ESI[†]) and aniline 2_a (Scheme S12 in the ESI[†]), and zero order in the *N*-tosylhydrazone $\mathbf{1}_{a}$ (Scheme S13 in the ESI[†]). In addition, the electronic influence of the aryl-substituents on both the reactants was investigated by a Hammett correlation study (Scheme 4b). By varying different electronic groups on the arylring of 2 a small $\rho = -1.17$ was obtained. This plausibly indicates a weak resonance interaction involving a positive-charge at the N center of aniline in the rate-determining-step. On the other hand, a negligible substituent effect ($\rho = -0.16$) was determined for N-tosylhydrazones. Meanwhile, the evolution of H₂ gas as a byproduct was confirmed by the transfer hydrogenation^{59,60} of styrene under the reaction conditions (Scheme 4c). A 2:1:1 mixture of 1_a , 2_a and styrene in the presence of the $B(C_6F_5)_3$ catalyst afforded 82% of 3_{aaa} along with 68% of ethylbenzene (Scheme 4c). Moreover, we performed a number of control experiments using amidines, azines, and imines as possible reaction intermediates albeit none of them proceeded to give 1,2,4-triazoles (Schemes S14-S16 in the ESI[†]).

A plausible mechanism is proposed on the basis of the above experimental observations and previous literature reports (Scheme 5).^{54,55} To the best of our knowledge DFT calculations of N–N cyclization leading to 1,2,4 triazole fragments are obscure. Based on the proposed mechanism of the $B(C_6F_5)_3$ -catalyzed acceptorless-dehydrogenative-cyclization of *N*-tosyl-hydrazones with anilines (Scheme 5), we have performed DFT calculations to investigate the detailed reaction mechanism and to gain insight into the driving force for the formation of the 1,2,4-triazole moiety 3_{bba} . Additionally, the calculations seek to address some pertinent questions regarding the studied system: (a) the specific role of $B(C_6F_5)_3$ in the reaction, (b) the rate-limiting-step in the reaction, and (c) product distribution for unsymmetrical coupling.

The reaction is initiated with the coordination of $B(C_6F_5)_3$ to the sp² nitrogen (N¹) in *N*-tosylhydrazone (**1**_b) which results in



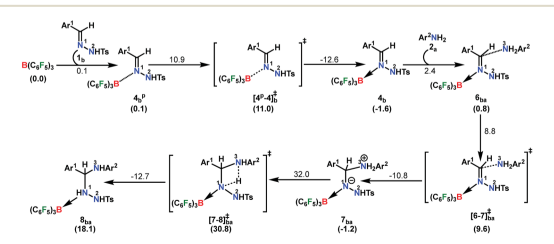
Scheme 5 Plausible reaction mechanism.

the formation of an isoenergetic encounter complex $\mathbf{4_b}^p$ (Scheme 6). The approach of the nucleophilic N^1 center in $\mathbf{1_b}$ towards the electron-deficient B center in $B(C_6F_5)_3$ furnishes the slightly more stable Lewis adduct $\mathbf{4_b}$ via a transition state $[\mathbf{4^{p-4}}]_{\mathbf{b}^{\ddagger}}$ with an activation barrier of 10.9 kcal mol⁻¹. Despite the fact that the N^2 center in $\mathbf{1_b}$ is significantly electron-rich (-0.646 e) compared to the N^1 center (-0.242 e), as obtained by the natural population analysis (NPA), $B(C_6F_5)_3$ gets coordinated to the N^1 center. This is attributed to the fact that the lone pair orbital located on the N^1 atom (HOMO-4) is significantly destabilized compared to the one on the N^2 atom (HOMO-5) by 0.4 eV (Fig. S100 in the ESI[†]). Furthermore, coordination at the N^2 center resulted in adduct $\mathbf{4b'}$, which is less stable than $\mathbf{4_b}^p$ by 2.4 kcal mol⁻¹ (Scheme S17 in the ESI[†]).

To cast light on the origin of the activation barrier and the bonding scenario in $[4^{p}-4]_{b}^{\ddagger}$, EDA-NOCV (energy decomposition analysis-natural orbital for chemical valence) analysis was performed, considering $\mathbf{1}_{\mathbf{b}}$ and $B(C_6F_5)_3$ as interacting fragments (Table S4 in the ESI[†]). Examination of the individual energy terms of the EDA reveals that the B-N¹ bond has a higher electrostatic character (ΔE_{elstat} : 39.8%) than the covalent character (ΔE_{orb} : 33.5%). Importantly, the major contribution to the total covalent interaction (ΔE_{orb}) originates from the donation of the lone pair on the N^1 center in $\mathbf{1}_{\mathbf{b}}$ to the vacant $2p_z$ orbital of boron in the $B(C_6F_5)_3$ fragment (Fig. S101 in the ESI[†]). We have calculated the associated eigenvalue of 0.49 e quantifying the amount of charge flow from donor \rightarrow acceptor fragments. Additionally, the $B(C_6F_5)_3$ fragment has the predominant contribution to the destabilizing distortion energy (ΔE_{dis}). The calculated electron density $[\rho(\mathbf{r})]$ of 0.112 at the (3, -1) bond critical point (BCP) of the $B-N^1$ bond in 4_b along with the respective Laplacian of +0.192 $[\nabla^2 \rho(\mathbf{r})]$ suggests a donoracceptor type interaction.^{61,62} Thereafter, the coordination of the substituted aniline (2_a) to 4_b affords the intermediate 6_{ba} which finally leads to a slightly more stable Zwitterionic complex 7_{ba} accompanied by a moderately low energy barrier of 8.8 kcal mol⁻¹. The imaginary mode in $[6-7]_{ba}^{\ddagger}$ portrays the formation of the C-N³ bond (1.849 Å) along with concomitant

elongation of the C-N¹ bond (1.383 Å). It is worthwhile to mention that the HOMO in 7_{ba} represents the lone pair orbital located on the N¹ atom (Fig. S100 in the ESI†). The subsequent proton transfer from N^3 to the N^1 center in 7_{ba} furnishes the significantly less stable intermediate 8ba via a four-membered transition state $[7-8]_{ba}^{\ddagger}$ (Scheme 6, Fig. 1a). The step $7_{ba} \rightarrow$ 8ba involving proton migration requires an activation barrier of 32.0 kcal mol^{-1} and thus becomes the rate-limiting step for the overall transformation.63 Indeed, this is supported by the experimental rate curve with a slower rate at the beginning of the reaction (vide supra, Fig. S1 in the ESI[†]). The single imaginary mode in $[7-8]_{ba}^{\dagger}$ depicts the synchronous breakage of N³-H (1.296 Å) and formation of N¹-H (1.353 Å) bonds. In $[7-8]_{ba}^{\dagger}$, the B-N¹ bond gets significantly elongated (1.602/1.676 Å in 7_{ba}/[7- 8_{ba}^{\dagger} and this weakening of the donor-acceptor bond is reflected in the reduced NPA charge on the B center (+0.467/ +0.488 e in $7_{ba}/[7-8]_{ba}^{\ddagger}$). It should be noted that both aniline and TsNH₂ assisted alternative intermolecular proton transfer between the two nitrogen centers $(N^3 \rightarrow N^1)$ are less favorable than the intramolecular path reported in Scheme 6 (Scheme S18a and b in the ESI[†]).

From here on, the coupling of a second N-tosylhydrazone unit is required for the progress of the reaction. This is accomplished through an initial proton transfer from the C center in $\mathbf{8}_{ba}$ to \mathbf{N}^4 in $\mathbf{1}_b$. Such a proton abstraction from the tertiary C atom is manifested with N¹-N² bond elongation. This intermolecular proton transfer is clearly favorable $(-17.5 \text{ kcal mol}^{-1})$, creating charged species $\mathbf{1}_{\mathbf{b}}^+$ and $\mathbf{9}_{\mathbf{ba}}$ respectively (Scheme 7), whereas the coordination of 1_b instead of proton transfer is highly unfavorable (refer Scheme S18c in the ESI[†]). In accordance with the experimental findings, KIE measurements suggest the non-involvement of imine C-H bond cleavage in the rate-determining step (Scheme 4a). Though obvious, it is important to note that hydrogen abstraction from electronegative N centers in 8ba is undoubtedly difficult, leading to highly unstable intermediates (Scheme S18d in the ESI[†]). Close inspection of the structural parameters in 9_{ba} indicates considerable elongation, rather than



Scheme 6 Part I: the reaction pathway for the formation of the intermediate $\mathbf{8}_{ba}$. The energy values above the arrows denote the Gibbs free energy changes (ΔG_L^S) of the individual steps. The values within parentheses are the relative ΔG_L^S energies w.r.t the starting structures. All energy terms are in kcal mol⁻¹.

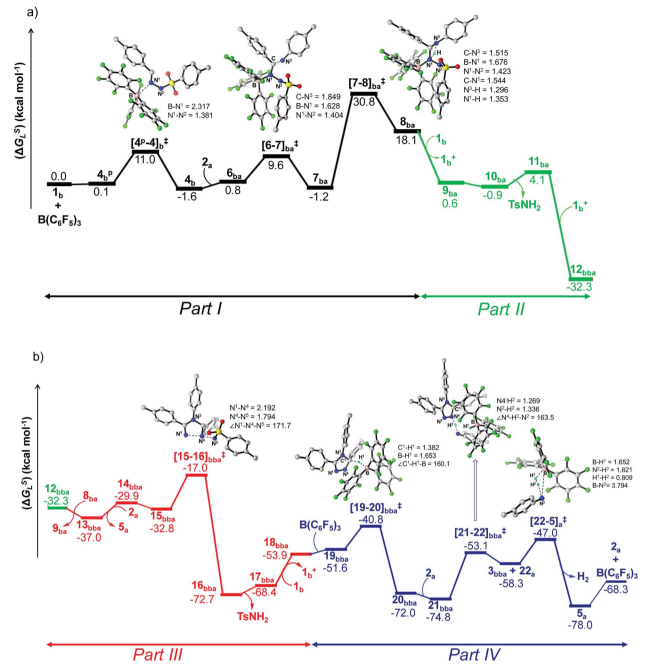
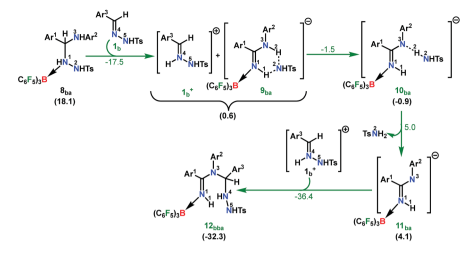


Fig. 1 Energy profile and 3D figures of optimized transition states with selected geometrical parameters for 1,2,4-triazole product $(\mathbf{3}_{bba})$ formation at the B3LYP-D3/TZVP//B3LYP/SVP level. The black, green, red and blue colored pathways represent sections (a) Part I and Part II, and (b) Part III and Part IV, respectively (refer Schemes 6–9). Bond distances are in angstroms (Å) and bond angles are in degrees (°).

dissociation of the N¹–N² bond (1.450 Å/2.987 Å = $8_{ba}/9_{ba}$) and generation of a partial double bond character in the C–N¹ bond (1.568 Å/1.323 Å = $8_{ba}/9_{ba}$). The N² center of the -NHTS unit in 9_{ba} shows significant hydrogen bonding interactions with the H² atom connected to the N³ center, as evidenced by the N³–H² (1.088 Å) and N²–H² (1.622 Å) bond lengths. The dissociation of the TsNH₂ fragment is quite evident from 10_{ba} with a shorter N²–H² distance (1.057 Å) and further elongated N¹–N² distance (3.634 Å). Complete removal of TsNH₂ will generate a highly nucleophilic N³ center in 11_{ba} which will immediately coordinate with the preformed cationic intermediate $\mathbf{1_b}^+$ to generate substantially stable $\mathbf{12_{bba}}$ (Scheme 7). The coupling of two oppositely charged species is further facilitated by the exothermicity of C–N³ bond formation.

From 12_{bba} the cyclization step is necessary to generate the triazole product 3_{bba} (Scheme 5). Under these circumstances, it might be conceivable that the liberation of a second TsNH₂ unit will facilitate N¹–N⁴ bond formation. Thus, protonation at the N⁵ center is necessary, similar to the preceding step 8_{ba} $\rightarrow 1_b^+ + 9_{ba}$ in Part II (Scheme 7). Unlike in 8_{ba} , the possibility

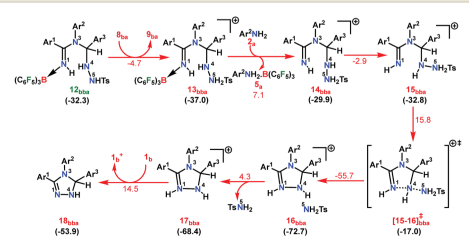


Scheme 7 Part II: the reaction pathway for the formation of the intermediate 12_{bba}. For other information refer the caption of Scheme 6.

of C-H abstraction in 12_{bba} either as a proton or hydride transfer is unfeasible (refer Scheme S18e in the ESI⁺). However, the formation of cationic species 13_{bba} after protonation along with anionic 9_{ba} generation is possible, but it is less exothermic than the previous transfer $(8_{ba} \rightarrow 1_b^+ +$ 9_{ba}; Schemes 7 and 8). Unfortunately, the addition of protons to any other electronegative center resulted in either high energy intermediates or reaction dead ends (refer Scheme S18f in the ESI[†]). In order to enhance the nucleophilicity at the N^1 center, $B(C_6F_5)_3$ was uncoordinated in the presence of an aryl amine (2_a) to generate notably unstable 14_{bba} (Scheme 8). Subsequent rearrangement to isomeric 15_{bba} provides adequate structural disposition for facile cyclization to proceed. We have explored numerous possibilities for N¹-N⁴ bond formation. However, none of them gave promising alternatives; instead the activation barriers are too high or transition states could not be optimized after numerous attempts (Scheme S18g in the ESI†). The cyclization step involving the transition state [15-16]_{bba}[‡] requires an

activation barrier of 15.8 kcal mol⁻¹ and it witnesses a progressive removal of the TsNH₂ unit.

As expected, the transition vector in [15-16]_{bba}[‡] depicts the breaking of the $N^4 - N^5$ bond (1.794 Å) with the concomitant formation of the N¹-N⁴ bond (2.192 Å). Complete removal of TsNH₂ affords the saturated triazole intermediate 17_{bba}, which is 35.6 kcal mol⁻¹ more stable than 15_{bba}. Subsequent deprotonation at the N^1 center by $\mathbf{1}_{\mathbf{b}}$ will generate the significantly less stable intermediate 18bba.⁶⁴ This step is facile with an activation barrier of only 1.6 kcal mol^{-1} (Scheme S19 in the ESI^{\dagger}). The protonated form $\mathbf{1_b}^+$ generated can have two fates: it may either participate in the preceding steps reported in Part II (Scheme 7) or can undergo an endergonic exchange of protons to a free amine $(\mathbf{2_a} + \mathbf{1_b}^+ \rightarrow \mathbf{2_a}^+ + \mathbf{1_b}; 4.8 \text{ kcal mol}^{-1})$. Generally, the intermediates formed after coupling of the second N-tosylhydrazone moiety are highly stable compared to the starting structures and the driving force for the subsequent reactions is the increasing exergonicity towards product formation (refer Fig. 1b). This statement is supported by high-resolution-massspectrometry studies which clearly detect a similar skeleton to



Scheme 8 Part III: the reaction pathway for the formation of the intermediate 18_{bba}. For other information refer the caption of Scheme 6.

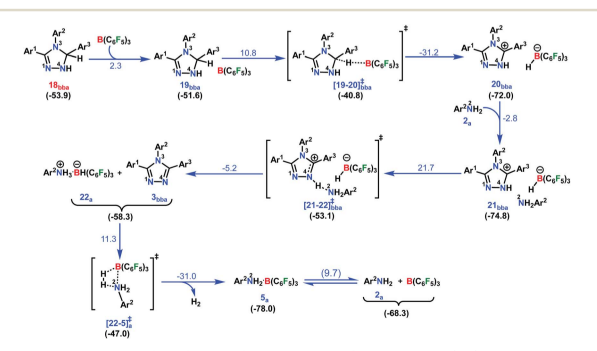
18_{bba} (Fig. S2 in the ESI[†]). In order to address the positional effect of $B(C_6F_5)_3$ in the cyclization step, we calculated two isomers in which it coordinates to other N centers (N⁴ and N⁵) in 12_{bba}. The resulting intermediates 35_{bba} and 36_{bba} are unstable and did not provide a low energy route to the cyclization step (Scheme S18h in the ESI[†]). Furthermore, in the absence of $B(C_6F_5)_3$ the cyclization step leading to 18_{bba} encounters a high transition barrier (38.2 kcal mol⁻¹; Scheme S18i in the ESI[†]) and thus underscores the significance of $B(C_6F_5)_3$ in this current transformation.

In the next step, 18_{bba} undergoes dehydrogenative aromatization^{54,55} to furnish triazole 3_{bba} and an ion pair 22_a through $B(C_6F_5)_3$ mediated hydride abstraction ($19_{bba} \rightarrow 20_{bba}$) followed by proton abstraction involving substituted anilines $(21_{bba} \rightarrow$ $3_{bba} + 22_{a}$; Scheme 9). We have calculated intrinsic activation barriers of 13.1 and 21.7 kcal mol⁻¹ for the hydride and proton abstraction steps, respectively. Thereafter, two hydrogen atoms in the ion pair 22_a produce a H₂ molecule via the fourmembered transition state $[22-5]_a^{\ddagger}$ (Scheme 9, Fig. 1b). Liberation of H₂ along with the formation of the frustrated Lewis acidbase pair (FLP) adduct 5_a is facile with a barrier of only 11.3 kcal mol⁻¹ (Scheme 9). The evolution of H₂ was also confirmed by the transfer hydrogenation of styrene (vide supra, Scheme 4c). Finally, maintaining an endoergic equilibrium, the FLP adduct regenerates $B(C_6F_5)_3$ and substrate 2_a .⁶⁵ In sum, the computational results do have concurrence with the experimental findings, particularly in understanding the dual role of $B(C_6F_5)_3$ in activating the N-tosylhydrazone towards nucleophilic attack and acceptor-less liberation of H2 with the formation of a FLP (Scheme 9). Additionally, the rate determining step involving intramolecular proton transfer (7_{ba} \rightarrow **8**_{ba}; $\Delta^{\ddagger}G_{\rm L}^{\ \rm S} = 32.0 \text{ kcal mol}^{-1}$) can be surmounted at a reaction temperature of 80 °C.66 Optimized geometries of the transition

states with selected geometrical parameters along with the energy profiles are shown in Fig. 1.

For unsymmetrical systems

An equimolar mixture of 1_a and 1_c in the presence of 2_a afforded the unsymmetrical triazole 3_{aca} as the major product (77%) compared to the symmetrical counterpart (vide supra; Table 3). To provide reasonable justification for this observation we decided to compare the relative propensity of Lewis acid-base adduct formation of N-tosylhydrazones 1_a and 1_c with $B(C_6F_5)_3$ $(B(C_6F_5)_3 \rightarrow 4_{a/c}^p \rightarrow 4_{a/c})$. As expected, the formation of 4_a is more facile than 4_c by *ca.* 3.0 kcal mol⁻¹ (Fig. S102 in the ESI[†]). This is in accordance with the experimentally observed equilibrium ratio in Scheme 3b. Eventually the activation barrier for $B(C_6F_5)_3$ coordination is favorable for the -OMe substituent by 3.6 kcal mol⁻¹ ($\Delta \Delta^{\ddagger} G_{L}^{s}$; Fig. S102 in the ESI[†]). What is more interesting is that the overall energy span for Part I is substantially higher for the -Cl substituent than -OMe (42.5 kcal mol⁻¹ vs. 33.3 kcal mol⁻¹), clearly indicating the preference for $\mathbf{1}_{a}$ to undergo $B(C_{6}F_{5})_{3}$ assisted intra-molecular proton transfer in a facile manner. Therefore, when 8aa couples with another hydrazone unit, the preferred choice will be the chloro-substituted analogue $\mathbf{1}_{c}$ as most of the $\mathbf{1}_{a}$ will be available in the adduct form 4_a . The combination of 8_{aa} with 1_c will lead directly to 12aca in a favorable fashion with an exothermicity of -50.6 kcal mol⁻¹ (Fig. S103 in the ESI[†]). From 12_{aca} , the generation of 16_{aca} requires a barrier of 20.4 kcal mol⁻¹ which is almost similar to the value obtained in the previous case (20.0 kcal mol⁻¹: $12_{bba} \rightarrow 16_{bba}$: Fig. 1b). This barrier is 1.1 kcal mol^{-1} lower than the symmetrical case (Fig. S103 in the ESI[†]) further supporting the preference for unsymmetrical triazole (3aca) formation (Fig. S103 in the ESI[†]).



Scheme 9 Part IV: the reaction pathway for the formation of the desired product 1, 2, 4-triazole complex (3_{bba}) and the hydrogen evolution step. For other information refer the caption of Scheme 6.

After the formation of the triazole ring in 16_{aca} , which is 71.5 kcal mol⁻¹ more stable than the starting materials, the subsequent $B(C_6F_5)_3$ assisted dehydrogenation follows an analogous mechanism as outlined before (Scheme 9; Fig. 1b).⁶⁷

Conclusions

In summary, we have demonstrated $B(C_6F_5)_3$ catalyzed metalfree, one-pot, dehydrogenative-cyclization of hydrazones with anilines to furnish both symmetrical and unsymmetrical 3,4,5triaryl-1,2,4-triazoles. The isolation of the *N*-tosylhydrazoneborane adduct is also reported for the first time. Mechanistic experiments and DFT calculations suggest that the $B(C_6F_5)_3$ catalyst serves a dual role: the activation of the hydrazone for the nucleophilic attack and the formation of an FLP for dehydrogenation. Calculations also reveal that the rate-determining step involves intra-molecular hydrogen transfer between the Ncenters after aniline gets bonded to the *N*-Tosylhydrazone unit. The chemo-selective, step-economical, oxidant-free and mild reaction protocol could give a potential platform for the increasing focus on main-group catalyzed chemical transformation without using transition metal.

Computational methods

All computations are performed using Gaussian 09 68 and ADF 2018.103 69 quantum codes. Geometry optimizations of the saddle points without any symmetry constraints are carried out using the B3LYP hybrid functional70 in conjunction with the SVP basis set⁷¹ in the Gaussian 09 program package. Harmonic force constants are computed at the optimized geometries to characterize the nature of the stationary points as minima (N_{img}) = 0) or transition states ($N_{img} = 1$). Transition states are located by using the linear synchronous transit (LST)⁷² scan method in which the reaction coordinate was kept fixed at different distances while all other degrees of freedom are relaxed. After the linear transit search, the transition states are optimized by using the default Berny algorithm implemented in the Gaussian 09 code. All transition states are validated by intrinsic reaction coordinate (IRC) calculations. In addition, single point calculations were performed on the B3LYP/SVP optimized structures using the dispersion corrected hybrid functional B3LYP-D3 73 in conjunction with a large basis set (triple-ζ quality split valence plus polarization, TZVP).74 The effect of solvation (benzene, dielectric constant $\varepsilon = 2.27$) was assessed by a self-consistent reaction field (SCRF) approach, using the conductor-like polarizable continuum model (CPCM).75 Tight wave function convergence criteria and an "ultrafine" (99 950) grid were used for the single point calculations. Natural bond orbital (NBO)⁷⁶ analysis was performed at the B3LYP-D3/TZVP//B3LYP/SVP level using the NBO Version 3.1 program. QTAIM (quantum theory of atoms in molecules) calculations are also performed to characterize the electron distribution around some selected bonds in the chemical species applying Bader's AIM (atoms-inmolecule) theory.77 Furthermore, to gain insight into the bonding scenario in the transition state $[4^{p}-4]_{b}^{\ddagger}$, EDA (energy decomposition analysis) calculations in conjunction with the

NOCV (natural orbital for chemical valence)⁷⁸ method are undertaken using the ADF 2018.103 package. Implementation and application of the EDA method, which was originally developed by Morokuma⁷⁹ and later modified by Ziegler and Rauk,⁸⁰ can be found elsewhere.^{81–85} The figures provided in the manuscript are generated using ChemDraw Ultra 12.0 and CYLview⁸⁶ visualization software.

Conflicts of interest

There are no conflicts to declare.

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