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Lewis acid mediated, mild C–H aminoalkylation of azoles *via* three component coupling†Marion H. Emmert,^{†*a} Cyndi Qixin He,^{†*b} Akshay A. Shah^{†c} and Stephanie Felten^{†a}

This manuscript reports the development of a mild, highly functional group tolerant and metal-free C–H aminoalkylation of azoles *via* a three-component coupling approach. This method enables the C–H functionalization of diverse azole substrates, such as oxazoles, benzoxazoles, thiazoles, benzothiazoles, imidazoles, and benzimidazoles. DFT calculations identify a key deprotonation equilibrium in the mechanism of the reaction. Using DFT as a predictive tool, the C–H aminoalkylation of initially unreactive substrates (imidazoles/benzimidazoles) can be enabled through an *in situ* protecting/activating group strategy. The DFT-supported mechanistic pathway proposes key interactions between the azole substrate and the Lewis acid/base pair TBSOTf/EtNⁱPr₂ that lead to azole activation by deprotonation, followed by C–C bond formation between a carbene intermediate and an iminium electrophile. Two diverse approaches are demonstrated to explore the amine substrate scope: (i) a DFT-guided predictive analysis of amine components that relates reactivity to distortion of the iminium intermediates in the computed transition state structures; and (ii) a parallel medicinal chemistry workflow enabling synthesis and isolation of several diversified products at the same time. Overall, the presented work enables a metal-free approach to azole C–H functionalization *via* Lewis acid mediated azole C–H deprotonation, demonstrating the potential of a readily available, Si-based Lewis acid to mediate new C–C bond formations.

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Introduction

Functionalized azole scaffolds are common target molecules in drug discovery. However, efficient diversifying routes towards these building blocks are underdeveloped. Typical approaches to access such chemical matter include multi-step, *de novo* syntheses of the azole core through condensation approaches.¹ Under the framework of a recent drug discovery program, azoles with pendant amino groups in alpha-position became an important structural class (Scheme 1). However, the rapid diversification of the chemical matter was limited by the lengthy *de novo* syntheses as well as by substrate-dependent synthetic success of assembling different azole cores from alpha-bromo ketones, NH₃ (or a surrogate), and complex α -amino acids

(Scheme 1, left side).¹ Therefore, we envisioned a different disconnection: direct C–H functionalization of the azole core with iminium electrophiles, thereby opening the door to one-step access to the desired structures and 3-component couplings, if the iminium electrophile can be accessed *in situ* from aldehyde and amine precursors (Scheme 1, right side).²

This strategy would also contribute to establishing more general and synthetically useful methods for azole C–H alkylation, which are generally challenging transformations. In fact, most azole C–H alkylation methods are promoted by transition metals and require forcing conditions, such as high temperatures, stoichiometric amounts of strong bases, and high catalyst loadings.^{3–5} Although various metal-free azole C–H alkylations are reported in the literature,^{2b,3,6} no examples of direct C–H aminoalkylations of azoles under mild reaction conditions have been described.

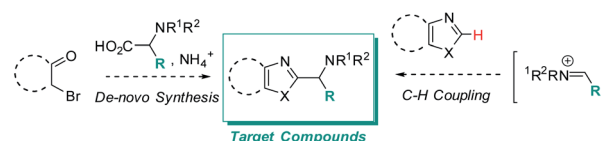
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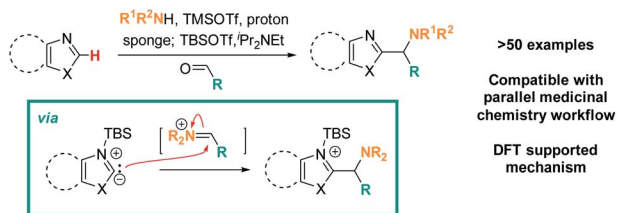
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† Electronic supplementary information (ESI) available: Detailed experimental procedures, characterization data for new compounds, details of DFT calculations. See DOI: 10.1039/d0sc06868c

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



Scheme 1 *De novo* synthesis of alpha-amino azoles requiring pre-functionalized building blocks (left) and target transformation (right) to allow rapid diversification *via* C–H functionalization and *in situ* iminium formation.



Scheme 2 Lewis-acid promoted, redox-neutral, one-pot, three-component azole C-H aminoalkylation & proposed mechanism.

Addressing these limitations, this manuscript describes a Lewis acid promoted direct C-H functionalization of azole substrates, which proceeds under mild conditions (Scheme 2). Trapping the *in situ* generated iminium electrophile with an azole nucleophile produces a diversity of heterocyclic structures in a three-component coupling approach.

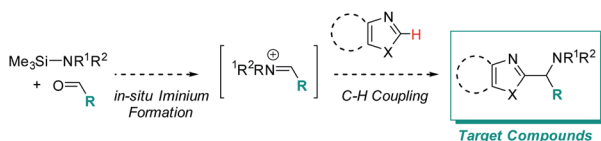
The mildness of the protocol is showcased by its ability to proceed in the presence of reactive functional groups such as amides, heterocycles, alkynes, and even enolizable ketones. Furthermore, its compatibility with a parallel medicinal chemistry workflow has been demonstrated, allowing access to compound arrays in parallel. Data from reaction optimization and DFT calculations support the need for Lewis acid and amine base to act in concert for successful C-H aminoalkylation, supporting the intermediacy of a silylated carbene as the key reactive species in these reactions.

Results and discussion

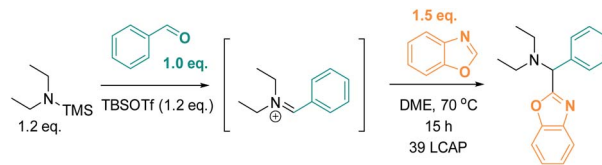
Reaction design and optimizations

We initially reasoned that synthesis of the highly functionalized target compounds could be achieved by direct C-H functionalization of azoles^{4–6} with an *in situ* formed iminium electrophile (Scheme 3); the procedure for forming the iminium intermediates was inspired by conditions used in Doyle's reductive three-component coupling.⁷ A reaction optimization screen (solvents, temperature, metal catalyst precursors, ligands, bases, acids; for details see ESI†) *via* high throughput experimentation led to the discovery that C-H aminoalkylation of benzoxazole at the 2-position was successful (39 LCAP; LCAP = LC area percent) when the azole was reacted with the *in situ* formed iminium electrophile in DME – even in the absence of any metal catalyst (Scheme 4).

A multidimensional optimization of reactant loadings (Scheme 5; 1.0 to 3.0 eq. TBSOTf, 1.0 to 2.5 eq. benzoxazole, 1.0 to 2.0 eq. TMS-NEt₂) revealed that all varied reaction components showed a significant influence on the reaction outcome.



Scheme 3 Desired target compounds and reaction design for C-H aminoalkylation.



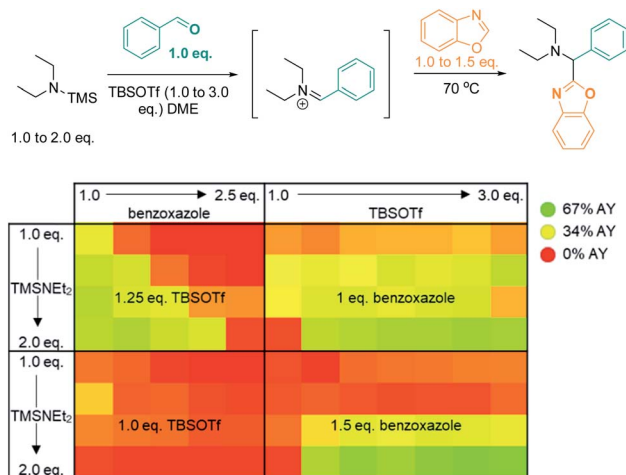
Scheme 4 Metal-free formation of desired C-H aminoalkylation product. LCAP = LC area percent.

The highest assay yields (65–66% AY; determined by calibrated UPLC analysis) were obtained under conditions with an excess of both basic and acidic reactants (2.5 or 3.0 eq. TBSOTf, 2.0 eq. TMS-NEt₂, 1.5 eq. benzoxazole). These findings imply that high concentrations of acidic reagent (TBSOTf) and basic reagents promote high yields. As both basic components of the reaction mixture (TMS-NEt₂, benzoxazole) contribute to the skeletal atoms of the desired product, reducing the required stoichiometry (ideally to 1 : 1) is highly desirable.

We reasoned that substituting the excess of basic reactants with an exogenous base may result in even better conversion and higher efficiency. To test this hypothesis, we surveyed the effects of five base additives (pyridine, EtNⁱPr₂, DBU, NMO, 2,6-lutidine; Scheme 6) at two different loadings (0.5 eq. and 1.0 eq.) in the presence of two concentrations of TMS-NEt₂ (1.25 eq. and 2.0 eq.). In this series, most of the tested conditions afforded yields close to 50% AY. The highest yield was obtained with the sterically bulky base EtNⁱPr₂ (1.0 eq.) in combination with 1.25 eq. TMS-NEt₂ (96% AY), while other common bases (pyridine, 2,6-lutidine) were detrimental to the reactivity under analogous conditions (22% and 17% AY, respectively; 1.0 eq. base, 1.25 eq. TMS-NEt₂).

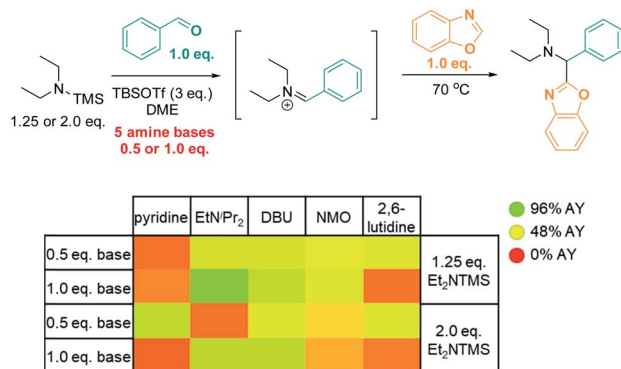
Mechanistic hypothesis

Based on the data presented above, both TBSOTf and bases promote the reaction. Among the tested bases, EtNⁱPr₂ is the most bulky base, but not the most basic (pK_a's of the corresponding protonated ammonium acids: 5.2 – pyridine;⁸ 6.6 –



Scheme 5 Optimization *via* high-throughput screening: influence of 3 variables (benzoxazole, TBSOTf, and Et₂NTMS loading) on product yield. AY = assay yield, as determined by calibrated UPLC analysis (210 nm), using biphenyl as internal standard.





Scheme 6 High throughput screen: base optimization. AY = assay yield, as determined by calibrated UPLC analysis (210 nm), using biphenyl as internal standard.

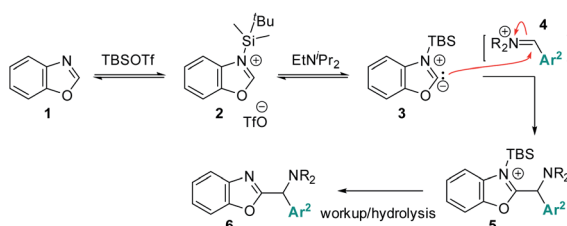
2,6-lutidine;⁸ 9.2 – NMO;⁹ 10.5 – EtNⁱPr₂;¹⁰ 12 – DBU¹¹). This suggests that steric hindrance around the base lone pair is beneficial for the overall reaction outcome. The influence of steric bulk on chemical reactivity of acids and bases has been documented particularly in investigations of Frustrated Lewis Pairs (FLPs);¹² the results from the amine screen described in the last paragraph thus led us to hypothesize that steric factors between EtNⁱPr₂ and TBSOTf may weaken the interaction between this Lewis base and acid pair and in turn enable efficient reactivity.

Based on this reasoning, our mechanistic hypothesis (Scheme 7) assigns important roles to EtNⁱPr₂ and TBSOTf: we propose that the TBSOTf first reacts with the azole substrate **1** to form silylated intermediate **2**, which then undergoes deprotonation. In a subsequent reaction step inspired by the mechanism of the Stetter reaction,¹³ the formed carbene **3** undergoes C–C bond formation with the pre-formed iminium electrophile **4**. Hydrolysis of **5** during workup and/or isolation leads to the final product **6**.

Preliminary experimental studies that react **1** with TBSOTf, ⁱPr₂NEt, and TfoH–D₁ indeed detect mass spectroscopic evidence for **2** as well as deuterium incorporation in **1**. These data (see ESI† for details) support the accessibility of a deprotonated species such as **3** under the tested conditions.

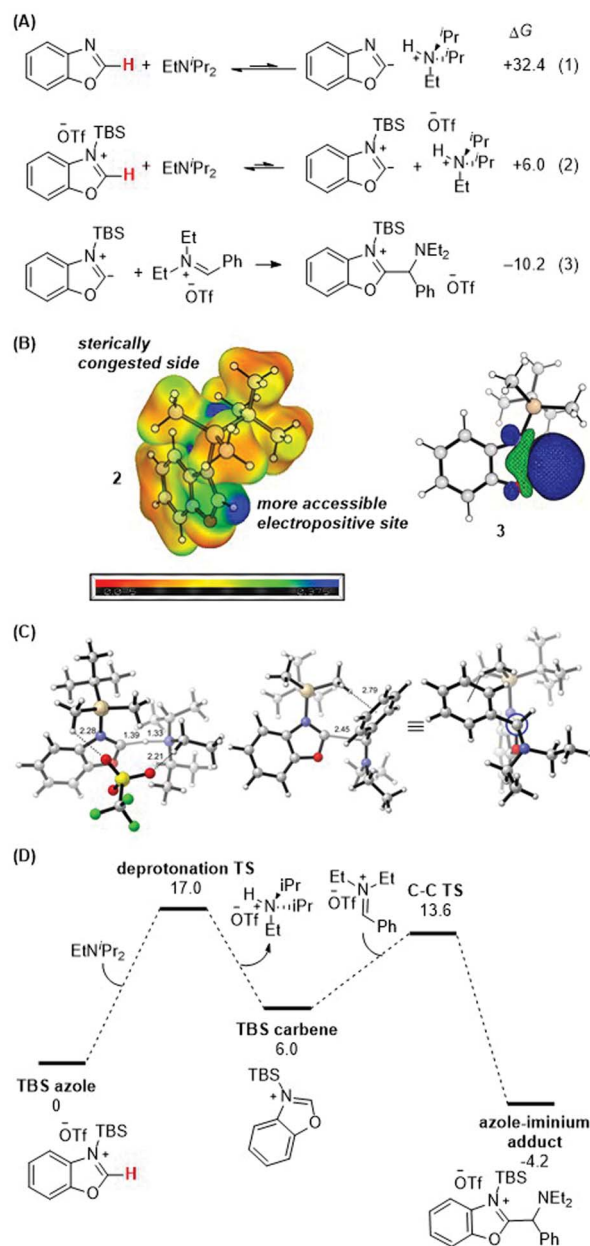
DFT support for deprotonation pathway

DFT calculations were performed to gain further insight into the thermodynamic and kinetic features of the proposed



Scheme 7 Proposed reaction mechanism via Lewis acid mediated azole deprotonation.

mechanistic pathway. Scheme 8A details the obtained thermodynamics of deprotonation and C–C bond formation. Without silylation at the N-atom of benzoxazole, deprotonation of the C–H bond at the 2-position is highly endergonic (eqn (1); $\Delta G = +32.4 \text{ kcal mol}^{-1}$) and thus unlikely to occur. Silylation with TBS (eqn (2)) activates this position and drastically shifts the deprotonation equilibrium by over 26 kcal mol^{−1} ($\Delta G = +6.0 \text{ kcal mol}^{-1}$). Finally, the energetics of the C–C bond formation step (eqn (3); $\Delta G = -10.2 \text{ kcal mol}^{-1}$) are exergonic,

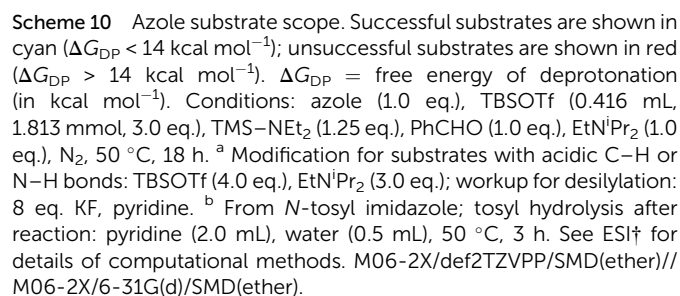
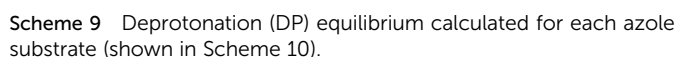


Scheme 8 (A) Calculated thermodynamics for deprotonation equilibria and C–C bond formation with benzoxazole and silylated benzoxazole. (B) HOMO of silylated benzoxazole. (C) Lowest-energy transition state of C–C bond formation. (D) Energy profile. See ESI† for details of computational methods. M06-2X/def2TZVP/SMD(ether)//M06-2X/6-31G(d)/SMD(ether).



Finally, the lowest energy transition states for the deprotonation and the C-C bond formation steps were computed (Scheme 8C and D). The proton transfer transition state was located with an activation barrier of 17.0 kcal mol⁻¹ and is the rate-determining step of the reaction. The deprotonation barrier excluding the triflate counterion in the model is 2.1 kcal mol⁻¹ lower. Upon formation of the carbene, the C-C bond forming TS between the silylated carbene intermediate (with TBS as silyl group and benzoxazole as azole) and the iminium electrophile [PhCH=NEt₂]⁺ is 13.6 kcal mol⁻¹. The TS structure features a forming C-C bond distance of 2.5 Å and a favorable electrostatic interaction between the α-C-H of the TBS group and the phenyl group of the iminium reactant at 2.8 Å. Higher energy conformers are reported in the ESI.†

When considering the calculated free energies of deprotonation ΔG_{DP} for the respective silylated substrates in combination with the experimental findings, a clear pattern emerges:



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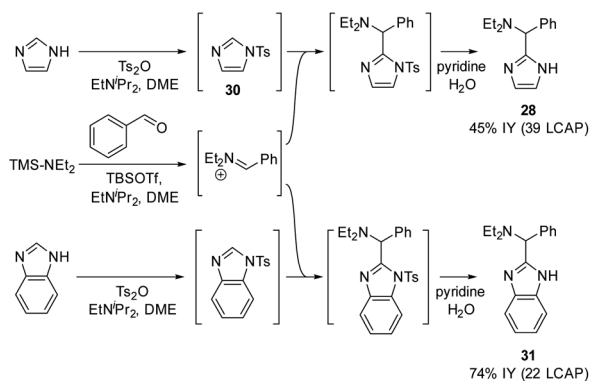
unfavorable deprotonation kinetics and thermodynamic equilibrium.

To circumvent this significant limitation in the azole scope, we considered activating protecting groups for imidazoles and benzimidazoles that may be able to shift the deprotonation equilibrium below the threshold energy of 14 kcal mol⁻¹. Indeed, we discovered that urea derivative **29** (Scheme 10; ΔG_{DP} 14.0 kcal mol⁻¹) afforded reasonable crude yields of the desired coupling product (**41** LCAP); however, isolation by column chromatography on silica led to decomposition of the product, resulting in a low isolated yield (3%). This observation suggests that electron-withdrawing protecting groups (such as acyl groups) can activate imidazole or benzimidazole substrates for C–H deprotonation and subsequent aminoalkylation.

In an extension of the activating protecting group strategy, subjecting *N*-tosyl imidazole (ΔG_{DP} 10.5 kcal mol⁻¹) to the standard conditions afforded an 88% isolated yield of the desired imidazole product **28**. To enable analogous reactivity with more diverse imidazoles and benzimidazoles that might not be as easily available as tosylated precursors, we further developed a procedure (Scheme 11) combining *in situ* tosylation, C–H aminoalkylation, and detosylation in one pot. With this sequence, imidazole and benzimidazole coupling products **28** and **31** were synthesized, starting from unprotected imidazole and benzimidazole substrates in moderate to good yields (45% and 74%, respectively) without the need to isolate the *N*-tosyl intermediates. These results demonstrate that the use of electron-withdrawing protecting groups can be implemented successfully to enable the C–H aminoalkylation of less activated azoles.

Predicting amine reactivity *via* distortion analysis of the transition state

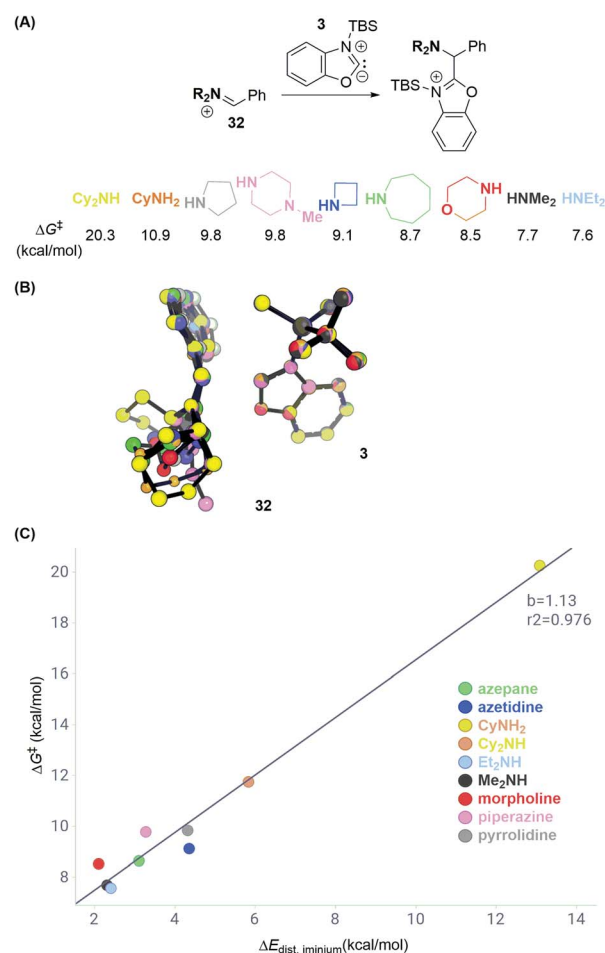
Having explored different azole substrates in C–H aminoalkylation, we set out to elucidate how structural features of the amine component may affect the reactivity. Given that the deprotonation barrier of a given silylated azole can be overcome, and the resulting silylated carbene is sufficiently stabilized, such as in the case of benzoxazole, comparing the activation energies of the C–C bond forming step across various electrophiles should enable predictions of reactivity patterns.



Scheme 11 *In situ* tosylation strategy.

We first calculated transition state energies for reactions between a series of differently substituted iminium electrophiles (**32**; Scheme 12A) and carbene intermediate **3** (Table S2†). The substitution patterns for **32** were chosen to reflect common amine cores in pharmaceutical chemistry and to vary between low and high steric demand. DFT calculations predict, accordingly, activation energies that span from 7.6 to 20.3 kcal mol⁻¹ (with respect to the silylated carbene and iminium). Interestingly, the C–C bond forming step for the dicyclohexyl iminium electrophile is 12.6 kcal mol⁻¹ higher than that for diethyl iminium, which is higher than the deprotonation barrier in the reaction pathway (Scheme 8D).

A distortion–interaction analysis¹⁴ (Scheme 12B and C) reveals that the total distortion energies ΔE_{dist} for the components in each reaction are unique and correlate with the activation free energies $\Delta G^\ddagger$ of C–C bond formation. In contrast, the values of the interaction energy term between **3** and each of the iminium intermediates **32** lie within 0.5 kcal mol⁻¹. Scheme 12B depicts an overlay of all computed transition state



Scheme 12 Distortion–interaction analysis of reactions with iminium electrophiles. (A) Transition state energies $\Delta G^\ddagger$ for series of iminium electrophiles. (B) Overlay of transition state structures. (C) Distortion energies ΔE_{dist} versus $\Delta G^\ddagger$ of C–C bond formation. See ESI† for details of computational methods. M06-2X/def2TZVPP/SMD(ether)//M06-2X/6-31G(d)/SMD(ether).



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