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<th>Unconventional Bifunctional Lewis-Brønsted Acid Activation Mode in Bicyclic Guanidine-Catalyzed Conjugate Addition Reactions</th>
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Abstract: DFT calculations have demonstrated that the unconventional bifunctional Brønsted-Lewis acid activation mode is generally applicable to a range of nucleophilic conjugate additions catalyzed by bicyclic guanidine catalysts. It competes readily with the conventional bifunctional Brønsted acid mode of activation. The optimal pro-nucleophiles for this unconventional bifunctional activation are acidic substrates with low $pK_a$, while the best electrophiles are flexible 1,4-diamide and 1,4-diester conjugated systems.

Keywords: organocatalysis; activation mode; guanidine; bifunctional activation; mechanism; DFT calculations

1. Introduction

Guanidines are versatile organocatalysts with vast applications in asymmetric synthesis [1–5]. The mechanistic aspects of guanidine-catalyzed reactions, in particular the various possible activation modes, have been reviewed recently [1,2]. Due to their strong Lewis basicity (many are in the superbase scale region), bicyclic guanidines are widely used in Brønsted base catalysis. In the Brønsted base mechanism,
guanidine abstracts a proton from the substrate and catalyzes the reaction as its conjugate acid, the guanidinium cation. Through hydrogen bonding interactions, the guanidinium ion can either activate the electrophile (mode I, Scheme 1) in a monofunctional mode or both the electrophile and nucleophile simultaneously in a bifunctional manner (mode II, Scheme 1). This bifunctional Bronsted acid activation mode was initially proposed by Corey and Grogan in asymmetric Strecker reactions using a chiral guanidine catalyst [6], and was supported by a theoretical study [7]. X-ray structure of TBD·HCl·H₂O provides further evidence to the bifunctionality of the guanidinium ion [8]. In a very recent kinetic study of three different types of bicyclic guanidine-catalyzed reactions, the bifunctionality of the guanidine catalyst is further established [9]. Our computational studies on bicyclic guanidine-catalyzed Michael [10,11] and phospha-Michael [12] reactions and isomerization reaction of alk-3-ynoates [9] have confirmed this type of bifunctional activation mode. It is worth mentioning that the guanidinium ion also plays an important role in the catalysis of phosphoric ester cleavage [13–16]. A similar bifunctional general base/general acid mechanism is believed to operate.

**Scheme 1.** Plausible mono- and bi-functional activation modes in Bronsted base catalysis of guanidine organocatalyst (\(E = \) electrophile and \(\text{Nu}^- = \) nucleophile).

The guanidinium ion is characterized by a low lying LUMO with a relatively large contribution of a vacant \(p\) orbital on the central carbon. In addition, the central carbon is strongly positively charged (e.g., NBO charge of +0.74 for [5,5] bicyclic guanidinium) [11,17] similar to that of a typical carbocation. A nucleophile (Lewis base) is expected to interact favorably with the highly electron deficient carbon of the guanidinium ion (Scheme 2). For an anionic nucleophile (e.g., PhS\(^-\)), the strong electrostatic attraction between the central guanidinium carbon and the anion provides an additional source of stabilization. As a result, guanidinium cation could readily serve as a Lewis acid catalyst via the central electrophilic carbon. This lesser known Lewis acid catalytic capability (mode III, Scheme 1) of hexyl alkyl guanidinium salts has been suggested by the authors in catalyzed reactions such as epoxide ring-opening esterification [18], lactide ring-opening polymerization [19] and the decomposition of alkyl formate [20]. Experimental support of such an activation mode is difficult to establish, but there is a wealth of evidence in X-ray crystallography that this type of “Lewis acid” interaction between the guanidinium central carbon and nucleophile exists (see references in Supporting Information).

One would envisage that this Lewis acid nucleophile activation mode could be combined with the Bronsted acid activation mode of an electrophile to provide an alternate bifunctional activation mode of guanidines as Bronsted base catalysts (mode IV, Scheme 1). Our DFT study [17] of the bicyclic guanidine-catalyzed thio-Michael reaction between thiophenol and phthalimide (Scheme 3) [21] has readily
demonstrated this alternate bifunctional Brønsted and Lewis acid activation mode. This unconventional bifunctional mode is also found to play an important catalytic role in our recent computational studies of bicyclic guanidine catalyzed-Michael addition of dimethyl malonate to 2-cyclopenten-1-one [11] and isomerization reaction of alk-3-ynoate [9].

Scheme 2. “Lewis acid” interaction between the vacant p orbital of the central electrophilic carbon of [5,5] bicyclic guanidinium ion and the lone pair of a nucleophile (Nu).

Scheme 3. Bicyclic guanidine-catalyzed thio-Michael reaction between thiophenol and phthalimide [21].

For the conjugate addition reaction studied [17], this unconventional activation mode, via pathway B (R₁ = Ph and Z = S, Scheme 4), competes readily with the conventional bifunctional Lewis acid activation mode, via pathway A (Scheme 4). Intriguingly, pathway B determines the enantioselectivity and rate of the reaction. In the catalytic cycle of the thio-Michael addition [17], the guanidine catalyst attracts a proton from thiophenol to form the thiophenolate ion (PhS⁻), which serves as a nucleophile in the subsequent conjugate addition. In the C–S bond forming step, both the electrophile and nucleophile interact with the guanidinium N–H protons simultaneously via hydrogen bond in transition state TSA (Scheme 4) in the conventional bifunctional Brønsted acid activation mode (i.e., pathway A). On the other hand, the nucleophile interacts with the cationic guanidinium carbon via a “Lewis acid” interaction while the phthalimide electrophile forms dual hydrogen bonds with both the guanidinium N–H protons in transition state TSB (Scheme 4) for the unconventional bifunctional Brønsted and Lewis acid activation mode (pathway B). Our computational study [17] showed that pathway B is significantly more favourable when the turnover frequencies of both pathways were compared and is enantioselective towards the S product, which agrees well with the experimental observation [21]. In this paper, we explore further the general application of this unconventional bifunctional activation mode to other nucleophilic conjugate additions and elucidate the factors that control the energetic preference towards this intriguing bifunctional activation mode.
2. Results and Discussion

2.1. Model Systems for Conjugate Nucleophilic Addition

Based on our mechanistic study of the bicyclic guanidine-catalyzed thio-Michael reaction [17], three criteria of the substrates appear to be important for the differential preference of the guanidinium bifunctional Brønsted-Lewis acid activation mode:

1. The electrophile has two Brønsted basic sites (i.e., hydrogen bond acceptors X and Y, see Scheme 4) for dual hydrogen bonding interaction with the two guanidinium N–H protons.
2. The nucleophile, preferably in an anionic form, should have significant Lewis basicity to interact favorably with the guanidinium central electrophilic carbon, via the “Lewis acid” interaction (Scheme 2).
3. The electrophile must have sufficient flexibility to enable the nucleophile to access the Lewis acidic site and the unsaturated carbon for conjugation addition simultaneously.

To validate above three criteria, the kinetic preference in the C–Z (where Z = carbon or heteroatom) bond forming transition states of the conventional Brønsted acid activation (pathway A) and the bifunctional Brønsted acid and Lewis activation (pathway B) was explored for nucleophilic addition of several model systems. The kinetic preference \( \Delta \Delta G^\neq \), \( G(TSA)-G(TSB) \), is taken as the difference in Gibbs free energy between the C–Z bond forming transition states of the two possible pathways, namely TSA and TSB for pathway A and B, respectively. A positive \( \Delta \Delta G^\neq \) implies that TSB is more stable than TSA, and a preference for pathway B. An unsubstituted [5,5] bicyclic guanidine catalyst (i.e., R = H,
Scheme 4) was used in this study as our main focus of this paper is on the activation mode, not stereoselectivity. It is important to note that our previous computational study has established that the rate determining step of catalytic Michael reaction is the nucleophilic addition to the carbonyl carbon instead of the protonation step [17]. Based on the super basicity of the bicyclic guanidine together with the known acidities of the pro-nucleophiles considered in the present study, it is unlikely that the activation of reactants is the rate limiting step. Therefore, it is justifiable to apply the Curtin-Hammet principle to the bicyclic guanidine-catalyzed conjugate addition studied here.

2.2. Optimal Brønsted Basic Functional Groups

Initially, various electrophiles were screened with thiophenol pro-nucleophile (N1) to identify which Brønsted basic functional groups are optimal for the bifunctional Lewis and Brønsted acid activation (Figure 1). The thio-Michael reaction of N1 with electrophile E1 was elucidated in our previous DFT study [17]. The removal of the t-butyl groups on the guanidine catalyst from Scheme 3 reduces the inclination of thio-Michael reaction between E1 and N1 to pathway B. Electrophile E2 was chosen to investigate the effect of absence on N-vinyl amide and ester moieties. Using similar nitro and ethyl ketone functional groups, a negative $\Delta \Delta G^\neq$ value of $-2.9 \text{ kJ} \cdot \text{mol}^{-1}$ was obtained, indicating a preference towards the conventional bifunctional pathway A (i.e., Brønsted acid activation). This suggests the importance of the N-vinyl amide and ester moiety in favoring the alternate activation mode. When both N-vinyl amide and ester groups are used in E3, there is a reversal of kinetically preferential towards pathway B, by 4.7 kJ·mol$^{-1}$. Substituting the ester moiety by an amide group (in E4) further amplifies the preference of pathway B ($\Delta \Delta G^\neq = +16 \text{ kJ} \cdot \text{mol}^{-1}$). However, replacing the vinyl acetamide in E3 by a vinyl ester group (in E5) does not lead to a significant change in $\Delta \Delta G^\neq (+2.6 \text{ kJ} \cdot \text{mol}^{-1})$. This may attribute to the fact that the acetamide group possesses a more rigid double bond which restricts the conformational freedom of the electrophile (Figure 2). In other words, TSB is somewhat destabilized with respect to TSA. The modification of the vinyl acetate to methyl acrylate in E6 results in a significantly lower energy pathway B ($\Delta \Delta G^\neq = +9.2 \text{ kJ} \cdot \text{mol}^{-1}$). The stronger preference for pathway B is due to the fact that TSB for the N1-E6 system possesses stronger hydrogen bonding interaction with the guanidinium ion. This argument is supported by the calculated donor and acceptor interactions in NBO analysis and the atomic charge on the carbonyl oxygen (Supporting Information).

As the unconventional bifunctional Lewis-Brønsted acid activation mode involves dual hydrogen bonding interaction between the two Brønsted basic functional groups (X and Y, Scheme 4) on the electrophile and the two Brønsted acidic N–H protons on the guanidinium ion, the stabilization interaction between the electrophile and guanidinium catalyst is expected to be a key factor for the pathway B activation. This argument is supported by comparing the binding energies of various guanidinium-electrophile hydrogen bonded complexes. The computed binding free energy provides a quantitative measure of the strength of the hydrogen bonding interaction in the guanidinium-electrophile complex. As evidenced in Figure 1, electrophiles E4 and E6, which possess the lowest binding energies, have a strong preference towards pathway B. On the other hand, systems with a low binding energy are associated with less stable TSB compared to TSA.
Figure 1. Various electrophiles (E1–E6) were screened with thiophenol (N1) for bicyclic guanidine-catalyzed conjugate addition reactions. \( \Delta \Delta G^\neq \) values (in kJ·mol\(^{-1}\)) are given in parenthesis and the binding free energies (in kJ·mol\(^{-1}\)) of the dual hydrogen bonded guanidinium-electrophile complexes are in square bracket.

Figure 2. Optimized geometries (M06-2X/6–31G\(^*\)) of TSA and TSB for conjugate addition of E3 and E5 with N1. C⋯S, S⋯H and O⋯H interaction distances are in Å.

It is important to note that noncovalent interactions between the nucleophile and electrophile are less significant compared to the substrate interactions with the guanidinium catalyst. These intermolecular interactions are the weaker C–H⋯X interactions and \( \pi–\pi \) interactions (see NBO analysis, Supporting Information). As a result, the noncovalent interactions between the electrophile and nucleophile do not have a major impact on the selectivity for the two activation modes.

2.3. Optimal Pro-Nucleophiles

Next, we examine optimal pro-nucleophiles for the preference of the bifunctional Brønsted-Lewis acid activation. We employed the best electrophile E4 to screen with different pro-nucleophiles N1 to N4.
As evidence in the large positive $\Delta \Delta G^\ddagger$ values (Figure 3), all the phenol analogues are suitable pro-nucleophiles for the preference of pathway B. The trend of $\Delta \Delta G^\ddagger$ ($N4 > N1 > N3$) indicates that the stability of the conjugate anion play an important role in the energetic preference of pathway B. This is not unexpected as the “Lewis acid” interaction (Scheme 2) involves complexation of the guanidinium cation and conjugate anion (i.e., activated nucleophile) of the pro-nucleophile.

**Figure 3.** Various pro-nucleophiles ($N1–N4$) screened with the optimal electrophile $E4$ for bicyclic guanidine-catalyzed conjugate addition reactions. $\Delta \Delta G^\ddagger$ values (in kJ·mol$^{-1}$) are given in parenthesis. The $pK_a$ values (in DMSO) were taken from Ref. [22].

Comparison of thiophenol ($N1$) with $t$-butyl thiol ($N2$) further confirms this hypothesis. As a result, the $\Delta \Delta G^\ddagger$ value correlates reasonably well with the $pK_a$ value (in DMSO) [23] of the pro-nucleophile (Figure 3). In general, pro-nucleophile with high acidity ($pK_a$) has a more stable conjugate base anion, and in turn leads to a more stable transition state (TSB) for the bifunctional Brønsted-Lewis activation. For the thiophenol analogues, the aromatic stabilization of the deprotonated aromatic conjugate anion is also a key factor for the energetic preference of pathway B. Since all the nucleophiles considered here are strong nucleophiles in anionic form, the stability of the base anion appears to be a more important factor than the nucleophilicity of the base in influencing the energetic preference towards pathway B, which involves Lewis acid interaction between the base anion and guanidinium ion. The “Lewis acid” interaction is the main source of stabilization in TSB type of transition state. The magnitude of this stabilization interaction depends on (1) donor-acceptor interaction between the lone pair of the nucleophile and the vacant $p$ orbital of the central guanidinium carbon and (2) electrostatic attraction between the anionic nucleophile and guanidinium “carbocation” ($C^+$) is expected to be large. However, it is less straightforward to quantify this electrostatic stabilization. In the cases of $N2$-$E4$-$TSB$ and $N3$-$E4$-$TSB$, the NBO atomic charges and the anion-$\cdot\cdot\cdotC^+$ distances (see Figure S1, Supporting Information) clearly demonstrate the significant stronger electrostatic attraction in $N3$-$E4$-$TSB$. C–H-$\cdots\pi$ stabilization is also evidenced in $N3$-$E4$-$TSB$ (Figure S1). The different strength of “Lewis acid” interaction together with the additional C–H-$\cdots\pi$ stabilization in $N3$-$E4$-$TSB$ explain the different preference of TSB for $N2$ and $N3$ despite both have nearly the same $pK_a$ values (Figure 3).
Despite being weak conjugate bases, the selected pro-nucleophiles studied here and/or their analogues have been shown to undergo Michael or nucleophilic addition reactions [24–26]. Experimental and computational studies of Lewis acid-base adducts by Denmark et al., indicated that a Lewis base could enhance the catalytic ability of a Lewis acid [27,28]. When a Lewis acid-base adduct is formed, the attached Lewis base will draw electron density away from the Lewis acid, increasing the partial negative charge on the Lewis base. This dispels the conventional notion that the Lewis base will lose its electron pair and become less nucleophilic. This finding further supports the unconventional bifunctional activation mode via TSB.

2.4. Bifunctional Lewis-Brønsted Acid Activation in Michael and Hetero-Michael Reactions

To explore further the general applicability of bifunctional Lewis-Brønsted acid mode of activation in Michael and hetero-Michael reactions, pro-nucleophiles N5–N8 were screened with the optimal electrophile E4 (Figure 4). All the nucleophiles chosen are capable of forming conjugate base anion with resonance stabilization, namely enolates for N5 and N6, acetate for N7 and cyanide for N8. As with several hetero-Michael reactions examined in previous section, the stability of the anion nucleophile is a key factor in governing the selectivity towards pathway B. For instance, substituting acetone (N5) with an electron withdrawing trifluoromethyl group (in N6), increases the acidity of the α-carbon hydrogen [22] and stabilizes the conjugate base enolate ion. As a consequence, there is a reversal of ΔΔG‡ on going from N5 to N6. In the same manner, the trend of ΔΔG‡ of acetone (N5), acetic acid (N7) and hydrogen cyanide (N8) can readily be explained in terms of the stability of the conjugate base anion. It is worth noting that the transition state TSB of acetate nucleophile is strongly favored compared to the cyanide system despite N7 and N8 have similar acidity (Figure 5). Interestingly, additional hydrogen bond is found in TSB for both cases, C–H···N interaction (2.46 Å) between the α-carbon hydrogen with the guanidinium ion in the N7–E4 system and normal O–H···O (2.37 Å) hydrogen bond in the N8–E4 case. NBO analysis indicates that the C–H···N stabilization energy (1.92 kJ·mol⁻¹) in the cyanide system is significantly weaker than the O–H···O stabilization (5.36 kJ·mol⁻¹) in the acetate system. It is worth noting that Michael addition of HCN with imine corresponds to Strecker reaction and guanidine is one of the most commonly used catalysts for asymmetric synthesis of α-amino acids [29]. Although the convention bifunctional activation mode has been proposed for the guanidine-catalyzed Strecker reactions [6,7], our computational results suggest that the unconventional bifunctional activation mode may play a crucial role in understanding the reactivity of stereoselectivity of guanidine-catalyzed Strecker reactions.

![Figure 4](image-url)  
**Figure 4.** Various acetyl and cyanide nucleophiles screened with the optimal electrophile E4 for bicyclic guanidine-catalyzed conjugate addition reactions. ΔΔG‡ values (in kJ·mol⁻¹) are given in parenthesis.
Figure 5. Optimized geometries (M06-2X/6–31G*) of TSB transition state for acetate (N7-E4-TSB) and cyanide (N8-E4-TSB) nucleophiles. Intermolecular interaction distances are given in Å.

2.5. Other Bicyclic Guanidine Catalyst

Another popular bicyclic guanidine catalyst, 1,5,7-triazabicyclo[4.4.0]dec-5-ene (TBD), is frequently used in the base catalysis of polymers [30,31], Michael addition [32], Henry [33] and aldol [34] reactions. Thus, we have examined also the TBD-catalyzed conjugate addition of N1 pro-nucleophile and E4 electrophile. This 6-membered ring catalyst is predicted to accommodate the bifunctional Lewis-Brønsted acid activation mode favourably ($\Delta \Delta G^\neq = +14.0$ kJ·mol$^{-1}$, Figure 6). The calculated binding energy of E4 with TBD is lower than that of the 5-membered ring bicyclic guanidine catalyst. The favourable interaction distances of TSA and TSB transition states are shown in Figure 7. The strong correlation between higher binding energy and higher selectivity towards the Lewis-Brønsted acid activation mode is again observed in this case. This result suggests that a favourable binding energy provide a good qualitative indication of a favourable Lewis-Brønsted acid activation mode in nucleophilic conjugate addition.

Figure 6. TBD-catalyzed nucleophilic addition between pro-nucleophile N1 and electrophile E4. $\Delta G^\neq$ value (in kJ·mol$^{-1}$) is given in parenthesis and binding free energy (in kJ·mol$^{-1}$) is in square bracket.
Figure 7. Optimized TSA and TSB transition states for TBD-catalyzed nucleophilic addition between N1 and E4. Interaction distances are given in Å.

3. Computational Methods

Computational Details

Equilibrium structures and transition states were fully optimized using the M06-2X [35] density functional method together with the standard 6–31G* basis set. The M06-2X functional was chosen as this empirical functional is better suited than normal hybrid DFT methods (e.g., B3LYP) in handling kinetics, thermodynamics, and noncovalent interactions [11,17,35–37]. Most importantly, our recent benchmark calculations have shown that dispersion corrections are essential in the DFT functionals to describe the long range “Lewis acid” interaction between a nucleophile and the guanidinium catalyst [17]. Frequency analyses were performed on all M06-2X/6–31G* optimized geometries to confirm the nature of the stationary points as equilibrium structures (with all real frequencies) or transition states (with only one imaginary frequency). The effect of solvation was examined by the SMD [38] implicit solvation model through M06-2X/6-311+G** single-point calculation, based on the gas-phase M06-2X/6–31G* optimized geometry. Both electrostatic and non-electrostatic terms are considered in this solvation calculations. Unless otherwise noted, the relative energies reported in the text correspond to relative free energies at 233 K (ΔG233), computed at M06-2X/6-311+G**/M06-2X/6–31G* level in diethyl ether solvent. The relative free energy (ΔGf) was computed from the equation ΔGf = ΔHf - TΔS, where ΔS is the entropy change and ΔHf = ΔH0 + (Hf - H0). Natural bond orbital (NBO) analysis was carried out at M06-2X/6–31G* level to study the charge distribution and donor-acceptor interactions, via the second-order perturbation energy analysis [39]. All calculations were performed using the Gaussian 09 suite of programs [40].

4. Conclusions

In summary, our DFT calculations have demonstrated that the unconventional bifunctional Brønsted-Lewis activation mode of guanidinium catalyst is generally applicable to various nucleophilic
conjugate additions. The optimal pro-nucleophiles are acidic substrates with low \( pK_a \) as the resonance stabilization of its conjugate base anion is important. The optimal electrophiles are flexible systems with two Brønsted basic functional groups capable of forming dual hydrogen bond with the guanidinium catalyst. The best candidates for electrophile are 1,4-diamide and 1,4-diester conjugated systems. Since the two possible bifunctional activation modes (modes II and IV, Scheme 1) favor different stereoisomers [17], one can employ the differential stabilization of the two pathways to control the stereoselectivity of nucleophilic addition product.

**Supplementary Materials**

Supplementary materials [NBO donor and acceptor interaction analysis, optimized geometries of “N2-E4-TSB and N3-E4-TSB, references for X-ray structures with guanidinium “Lewis acid” interaction, and Cartesian coordinates of all optimized geometries (in .xyz format)] can be accessed at: http://www.mdpi.com/1420-3049/20/08/15108/s1.

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**Author Contributions**

BKC conceived the ideas, performed the computational work and drafted the paper. MWW analyzed the mechanism and wrote the paper. All authors read and approved the final manuscript. MWW is the corresponding author.

**Conflicts of Interest**

The authors declare no conflict of interest.

**References**


Sample Availability: Not available.

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