



Cite this: *Chem. Commun.*, 2025, 61, 19493

Received 24th August 2025,  
Accepted 10th November 2025

DOI: 10.1039/d5cc04816h

rsc.li/chemcomm

# Controlling Lewis acid-catalysed bicyclobutane cycloadditions: carbonyl substituents dictate electrophilic vs. nucleophilic addition pathways

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**Combined DFT and DLPNO-CCSD(T) mechanistic studies reveal that carbonyl substituents of bicyclobutane (BCB) dictate the reaction pathways in Lewis acid-catalysed BCB-imine cycloadditions, toggling between electrophilic and nucleophilic additions, as well as concerted and stepwise ring-opening/addition mechanisms.**

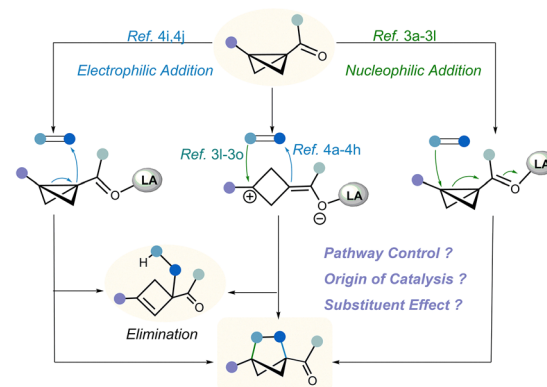
Driven by the “escape from flatland” concept in medicinal chemistry, three-dimensional bridged cyclic hydrocarbons have received growing interest in recent years.<sup>1</sup> These strained architectures, including bicyclo[2.1.1]hexanes (BCHs) and bicyclo[3.1.1]heptanes (BCHepts), offer superior properties such as improved solubility and better metabolic stability compared to traditional aromatic frameworks. In particular, BCHs have emerged as valuable bioisosteres capable of mimicking *ortho*- and *meta*-substituted benzenes while providing distinct three-dimensionality.

The synthesis of BCHs has been significantly advanced through cycloaddition chemistry of bicyclo[1.1.0]butanes (BCBs) *via* thermo-driven,<sup>2</sup> Lewis-acid catalysed,<sup>3,4</sup> photocatalysed,<sup>5</sup> or radical-based<sup>6</sup> catalysed reactions. Among them, Lewis acid catalysis has proven to be a versatile and efficient strategy to access BCHs from BCBs.<sup>7</sup> Continuous contributions from groups including Leitch, Studer, Glorius, Deng, Feng, Biju, Bach, Shi, Zhou, Li, and Han have significantly advanced this field,<sup>3,4</sup> enabling cycloaddition of BCBs with a diverse range of unsaturated substrates. Notably, recent advances now enable asymmetric synthesis of enantioenriched BCH.<sup>8</sup> Despite significant advances in the synthetic applications of Lewis acid-catalysed BCB cycloadditions, the underlying reaction mechanisms remain poorly understood.<sup>3,4,9</sup>

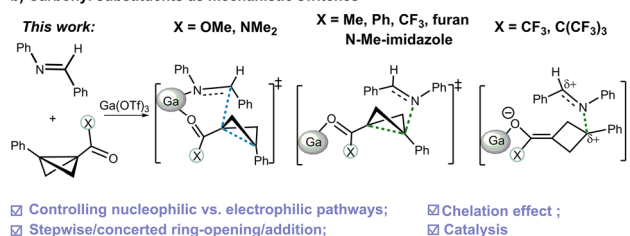
Competing hypotheses (Scheme 1) suggest that Lewis acid coordination may either: (1) enhance the electrophilic character of the BCB moiety, facilitating addition with various

nucleophiles including indoles and ynamides (nucleophilic addition pathway),<sup>3</sup> or (2) activate the BCB as a nucleophile to react with electron-deficient partners (electrophilic addition pathway).<sup>4</sup> The latter pathway requires particular attention as the cycloaddition competes with potential E1 elimination.<sup>4a,4b</sup> Notably, many<sup>3l-o,4a-h</sup> of these studies have proposed that Lewis acid activation might induce direct cleavage of the bridge C–C bond in BCB, generating a zwitterionic enolate intermediate that subsequently attacks unsaturated substrates through a stepwise mechanism rather than a concerted ring-opening/addition pathway. Given these mechanistic complexities, our ability to selectively steer these competing pathways remains limited.

a) Proposed mechanisms for Lewis acid-catalyzed BCB cycloaddition



b) Carbonyl substituents as mechanistic switches



**Scheme 1** Mechanistic considerations of Lewis-acid catalysed cycloaddition of BCB.

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Current experimental approaches to controlling BCB cycloaddition pathways primarily focus on modulating the electronic properties of the unsaturated coupling partners.<sup>3a-d</sup> However, our recent studies indicated that the electronic characteristics of BCB active species play a key role in pathway selection.<sup>10</sup> Building upon this insight, we now computationally demonstrate that modifying the BCB carbonyl substituents (X = OMe, NMe<sub>2</sub>, Me, Ph, CF<sub>3</sub>, C(CF<sub>3</sub>)<sub>3</sub>, N-Me-imidazole and furan) is a feasible strategy to control reaction pathways using the same coupling partner.

Our mechanistic investigation begins with the Ga(OTf)<sub>3</sub>-catalysed cycloaddition of BCB ester **2a** with *N*-aryl imine **1a**, given the rich experimental data. Calculations using the DLPNO-CCSD(T)/def2-TZVP(SMD)||SMD(THF) B3LYP-D3/6-31G(d) method confirm the electrophilic addition pathway,<sup>4a</sup> but initiates with favorable bi-coordination of both substrates to Ga, forming stable intermediate **IN1<sub>2a</sub>** (Fig. 1a). However, contrary to prior proposals, the ring-opening and addition proceeds concertedly *via* **TS1E<sub>2a</sub>** ( $\Delta G^\ddagger = 16.4$  kcal mol<sup>-1</sup>) to give **IN2<sub>2a</sub>**, not through a stepwise process. The alternative nucleophilic addition pathway proves energetically less favorable (**TS1N<sub>2a</sub>**,  $\Delta G^\ddagger = 24.3$  kcal mol<sup>-1</sup>) due to the requirement of imine dissociation from Ga(III) to attack the distal BCB carbon atom.<sup>11</sup> This dissociation is thermodynamically unfavourable, as evidenced by the complex formed by **Ga/2a** and the uncoordinated imine lying 9.7 kcal mol<sup>-1</sup> higher in energy than **IN1<sub>2a</sub>**.

While the major product for **2a** is the cycloadduct, intermediate **IN3<sub>2a</sub>** can undergo competitive E1 elimination (Fig. S2). For **2a**, the elimination product is thermodynamically disfavored by 3.0 kcal mol<sup>-1</sup> relative to the cycloaddition

product. Nevertheless, *tert*-butyl substitution on the imine stabilizes the E1 product by 7.8 kcal mol<sup>-1</sup> lower than the cycloaddition product, and drives the reaction exclusively toward E1 elimination. These calculations are in perfect agreement with experimental observations, which provides robust validation of our calculation model.

Remarkably, introducing a methyl substituent at the BCB carbonyl group induces a complete mechanistic inversion, shifting the dominant pathway from electrophilic to nucleophilic addition (Fig. 1b). This nucleophilic pathway proceeds through a concerted ring-opening/nucleophilic addition TS (**TS1N<sub>2b</sub>**,  $\Delta G^\ddagger = 17.7$  kcal mol<sup>-1</sup>), which serves as the rate-determining step. Nevertheless, the competing electrophilic pathway (**TS1E<sub>2b</sub>**) becomes disfavored by 3.9 kcal mol<sup>-1</sup>. Compared to BCB ester, this ketone derivative clearly exhibits significantly enhanced reactivity toward nucleophilic attack—despite similarly unfavorable imine dissociation—highlighting the critical role of substituent effects in driving the observed mechanistic switch. Further evaluations of NMe<sub>2</sub>- and Ph-substituted BCBs suggest the electronic control of pathway selectivity. The Ph-system displays comparable pathway selectivity and activation barriers to the Me-system, whereas the amide analogue with strong electron-donating character follows the behavior observed for BCB ester (Fig. S4 and S5).

Our distortion/interaction-activation strain (D/IAS) analysis reveals that while the Ga-coordinated **2b**/imine system exhibits faster-growing distortion energy along the nucleophilic addition reaction coordinate compared to the BCB ester analogue (Fig. 2), this penalty is more than compensated by significantly stronger inter-fragment interactions, making nucleophilic addition the dominant pathway for BCB ketone. Further energy decomposition analysis (EDA)<sup>12</sup> pinpoints the enhanced interactions arising from superior electrostatic interactions, and pronounced induction (including polarization and charge transfer) between the Ga-activated **2b** and imine components. Frontier molecular orbital (FMO) analysis suggests that the interaction energy differences mainly originate from fundamental electronic restructuring. The LUMO of the Lewis acid-activated BCB ketone is substantially

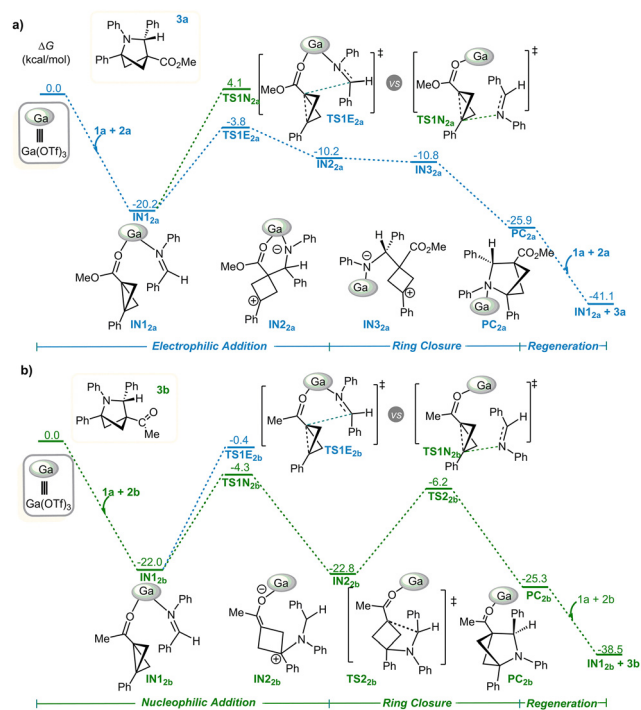


Fig. 1 Free energy profile (in kcal mol<sup>-1</sup>) of Lewis acid-catalysed cycloaddition of BCB substrate. (a) X = OMe; (b) X = Me.

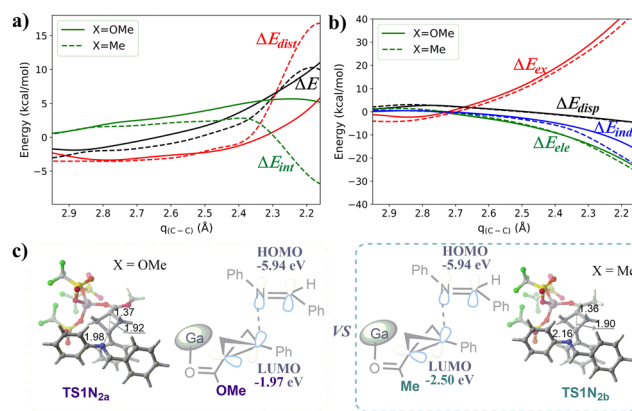


Fig. 2 Energy decomposition analysis for BCB ketone and ester reactivity. (a) Comparison of distortion and interaction energies. (b) Breakdown of interaction energy components. (c) Frontier orbital interaction and transition state geometries with key bond lengths (Å).



stabilized ( $-2.50$  eV) relative to the ester analogue ( $-1.97$  eV), reflecting the weaker electron-donating ability of Me than OMe. This stabilization narrows the HOMO(imine)–LUMO(Ga-BCB) gap from  $3.97$  (OMe) to  $3.44$  eV (Me). This orbital energy modulation directly correlates with the lower nucleophilic addition barrier for BCB ketones ( $17.7$  kcal mol $^{-1}$ ) compared to esters ( $24.3$  kcal mol $^{-1}$ ). Collectively, these findings establish a clear structure-mechanism relationship: subtle modifications at the BCB carbonyl group (ester  $\rightarrow$  ketone) can profoundly redirect the reaction pathways by tuning frontier orbital energies and intermolecular interaction patterns.

Comparative analysis of OMe- and Me-substituted BCBs reveals another intriguing finding concerning the viability of the proposed stepwise ring-opening/addition pathway.<sup>31-o,4a-h</sup> For **2a** and **2b**, the putative Ga-bound ring-opened enolate intermediates and associated transition states are energetically disfavoured relative to the concerted pathway, ruling out the stepwise mechanism (Fig. 3 and Fig. S6). Nevertheless, we observe a progressive increase in thermodynamic stability of the intermediate across the series (OMe  $\rightarrow$  Me), suggesting a potential carbonyl substituent-driven “stepwise-concerted” pathway shift by modifying the electron donating or withdrawing ability. Calculations using the CF $_3$ -substituted BCB show that the energy barrier difference between the two pathways is rather small ( $\sim 1.1$  kcal mol $^{-1}$ ), rendering them degenerate, while C(CF $_3$ ) $_3$  completely suppresses the concerted path, enabling only the stepwise mechanism. These findings establish that carbonyl substituents can also serve as powerful mechanistic switches between concerted and stepwise pathways.

Next, we studied BCB functionalized with an *N*-methyl-imidazole group (**2c**)-transforming it into a privileged bidentate substrate that is widely utilized in cycloaddition reactions.<sup>37</sup> The reaction profile of **2c** exhibits unique mechanistic features (Fig. S7), favouring a nucleophilic addition pathway analogous to BCB ketones but distinguished by its metal coordination behaviour (Fig. 4). The bidentate coordination mode (*via* BCB carbonyl oxygen and imidazole nitrogen) of substrate **2c** generates an exceptionally stable intermediate (IN1 $_{2c}$ ,  $-42.1$  kcal mol $^{-1}$ ), in which imine is also coordinated to Ga. Nucleophilic addition

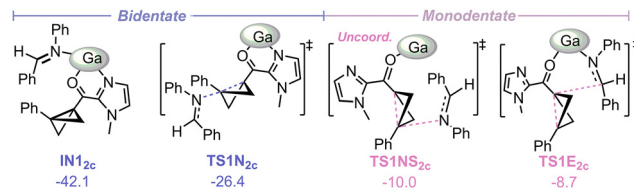


Fig. 4 Free energy (kcal mol $^{-1}$ ) of key species in Lewis acid-catalysed cycloaddition of BCB substrate **2c** with chelating substituent *N*-Me-imidazole.

proceeds *via* a dissociative transition state (TS1N $_{2c}$ ,  $\Delta G^\ddagger = 15.7$  kcal mol $^{-1}$ ) where the imine dissociates to attack the BCB. The bidentate chelation is critical, as it drastically stabilizes the pathway, with the monodentate coordination exhibiting a  $16.4$  kcal mol $^{-1}$  higher barrier. This stabilization as well creates a  $17.7$  kcal mol $^{-1}$  energy gap between nucleophilic and electrophilic pathways, effectively suppressing the latter. Comparably, a weaker-coordinating furan substituent forms a less stable complex ( $-26.3$  kcal mol $^{-1}$ ) and narrows the pathway gap to just  $2.6$  kcal mol $^{-1}$  (Fig. S8), confirming that strong chelators preferentially stabilize nucleophilic addition transition states.

Finally, we elucidate the catalytic origin of Lewis acid catalysis in the nucleophilic addition pathway. In the absence of Ga(OTf) $_3$ , the nucleophilic addition TSs of **2b/2c** are energetically disfavoured by  $23.8$ – $27.3$  kcal mol $^{-1}$ . Our D/IAS calculations demonstrate that the driving force of catalysis originates from the significantly enhanced intermolecular interactions between activated BCB and imine (Fig. 5a). EDA identifies the stabilization mechanism as a combination of reduced exchange repulsion and enhanced induction interactions for **2b** and **2c** (Fig. 5b and Fig. S9). Orbital analysis further rationalizes these effects: Ga(III) coordination lowers both the HOMO and LUMO energies of the BCB substrate. This dual orbital tuning creates an optimal electronic environment by: (i) minimizing HOMO–HOMO repulsion between reactants (consistent with the Pauli repulsion-lowering catalysis strategy proposed by Hamlin *et al.*<sup>13</sup>), which accounts for  $\sim 60\%$  of the total catalytic stabilization, and (ii) enhancing HOMO(imine)–LUMO(BCB) overlap, facilitating efficient electron transfer (Fig. 5c). Thus, the catalytic role of Ga in the nucleophilic addition pathway arises from the synergistic effects of Pauli repulsion-lowering and LUMO-lowering action. Notably, while the LUMO orbital shows significant distribution on the carbonyl carbon site, calculations confirm that nucleophilic attack at this position is kinetically disfavoured, with an energy barrier consistently  $> 6.0$  kcal mol $^{-1}$  higher than that at the bridgehead carbon across all BCB derivatives (Fig. S10).

In summary, this communication demonstrates how carbonyl substituents of BCB dictate reaction pathways in Lewis acid-catalysed BCB-imine cycloadditions, establishing a framework for controlling strained ring reactivity through rational substituent design.

We thank the National Natural Science Foundation of China (12504333), and Guangdong Basic and Applied Basic Research Foundation (2022A1515110835 and 2025A1515010088) for funding this work. The computational resources are supported by SongShan Lake HPC Center (SSL-HPC) in Great Bay University.

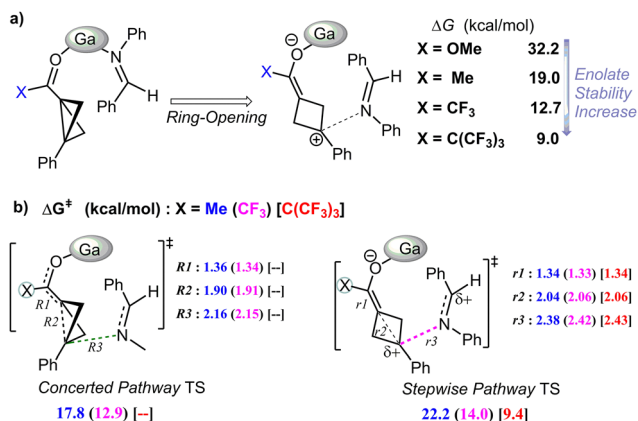


Fig. 3 (a) Thermodynamics of direct ring opening of BCB; (b) overall barrier of the nucleophilic addition transition state.





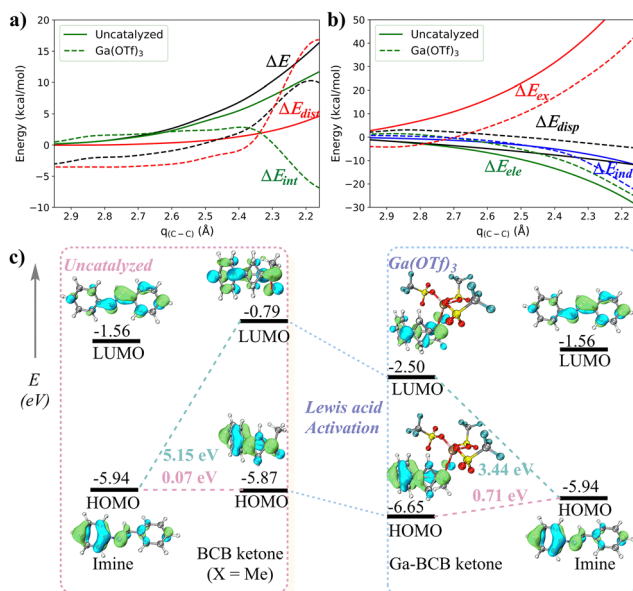


Fig. 5 (a) D/IAS and (b) EDA along the reaction coordinate of nucleophilic addition for uncatalysed and catalysed cycloadditions of BCB ketone, and (c) frontier orbital interaction diagram.

## Conflicts of interest

There are no conflicts to declare.

## Data availability

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information: computational details, free energy profiles of varying BCB derivatives, absolute energies and coordinates. See DOI: <https://doi.org/10.1039/d5cc04816h>.

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