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Rationalizing the diverse reactivity of [1.1.1]propellane through  $\sigma$ - $\pi$ -delocalization

This work provides a unified framework that explains the stability and broad reactivity of [1.1.1]propellane. It challenges classical strain relief arguments often used to rationalize propellane reactivity and demonstrates that reaction pathways and barriers are governed by delocalization of electron density within the cage upon reaction. These results bridge the gap between theory and real-life applications, changing our understanding of the way strain can manifest itself in organic chemistry.

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# Rationalizing the diverse reactivity of [1.1.1]propellane through $\sigma$ - $\pi$ -delocalization†

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[1.1.1]Propellane is the ubiquitous precursor to bicyclo[1.1.1]pentanes (BCPs), motifs of high value in pharmaceutical and materials research. The classical Lewis representation of this molecule places an inter-bridgehead C–C bond along its central axis; ‘strain relief’-driven cleavage of this bond is commonly thought to enable reactions with nucleophiles, radicals and electrophiles. We propose that this broad reactivity profile instead derives from  $\sigma$ - $\pi$ -delocalization of electron density in [1.1.1]propellane. Using *ab initio* and DFT calculations, we show that its reactions with anions and radicals are facilitated by increased delocalization of electron density over the propellane cage during addition, while reactions with cations involve charge transfer that relieves repulsion inside the cage. These results provide a unified framework to rationalize experimental observations of propellane reactivity, opening up opportunities for the exploration of new chemistry of [1.1.1]propellane and related strained systems that are useful building blocks in organic synthesis.

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## Introduction

[1.1.1]Propellane (tricyclo[1.1.1.0<sup>1,3</sup>]pentane **1**, Fig. 1) is a molecule that has long fascinated organic and theoretical chemists due to its unique structure and intriguing reactivity. In recent years, **1** has also become established as a key precursor to bicyclo[1.1.1]pentanes (BCPs),<sup>1,2</sup> motifs that are attractive bioisosteres for aryl,<sup>3–8</sup> alkynyl<sup>9</sup> and *tert*-butyl<sup>10</sup> groups in drug design due to the enhanced pharmacokinetic profile of the BCP spacer unit compared to the parent functional group (Fig. 1a). BCPs are also useful motifs for organic materials, including rod-like one-dimensional polymers,<sup>11</sup> supramolecular spacer units,<sup>12</sup> liquid crystals<sup>13</sup> and FRET sensors.<sup>14</sup> Such applications have stimulated the development of a number of methods to access BCPs in a single step from **1** via radical<sup>15–24</sup> and anionic<sup>25–29</sup> intermediates (Fig. 1b). Limitations nonetheless remain, in particular due to the harsh reaction conditions often associated with many of these processes, which can restrict functional group tolerance. A third reactivity mode of **1** involves facile cationic addition, a process that fragments the cage to an *exo*-methylene cyclobutane.<sup>30–32</sup>

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### a. Applications of bicyclo[1.1.1]pentanes (BCPs)



### b. [1.1.1]Propellane as a ubiquitous source of BCPs



Fig. 1 (a) Applications of bicyclo[1.1.1]pentanes (BCPs) in pharmaceutical and materials settings. (b) The omniphilic reactivity of [1.1.1]propellane with anions/organometallics (blue arrows), radicals (red arrows) and cations (grey arrows).

This diverse 'omniphilic' reactivity has traditionally been ascribed to the high strain energy of [1.1.1]propellane (total strain energy  $\sim 100$  kcal mol $^{-1}$ ),<sup>33</sup> albeit only  $\sim 30$  kcal mol $^{-1}$  of this strain is thought to be released on cleavage of the inter-bridgehead C1–C3 bond.<sup>2,34</sup> Nonetheless, this omniphilicity is not generally observed with other ring-opening reactions where relief of ring strain is a driving force (such as in cyclopropane,  $E_{\text{strain}} = \sim 28$  kcal mol $^{-1}$ ),<sup>35</sup> which could suggest other factors govern the reactivity of **1**.<sup>36</sup>

To date, a handful of experimental studies have employed density functional theory (DFT) to investigate the pathways of addition of radicals and anions to [1.1.1]propellane, offering useful insight to support reaction development.<sup>17,18,20,28,29</sup> However, no in-depth theoretical analysis has tackled the origin of the omniphilic reactivity of **1**. Here we combine *ab initio* approaches with electron density difference analysis and Distortion/Interaction Analysis (DIA)<sup>37–39</sup> to develop a unified model for the reactivity of [1.1.1]propellane. Contrary to classical explanations, we demonstrate that its chemistry is not determined simply by geometric 'strain relief'; instead, it is the change in electron delocalization over the cage that defines the observed activation profiles. These results bridge the gap between the theoretical understanding of [1.1.1]propellane as a 'scientific curiosity' to its applications in real-life settings.<sup>2</sup>

## Results and discussion

### The structure of [1.1.1]propellane

Before considering the reactivity of [1.1.1]propellane, we first sought to characterize the nature of its inter-bridgehead C1–C3 bond in the ground state (Fig. 2a). The geometry of **1** has been determined experimentally by low-temperature X-ray diffraction<sup>40</sup> and gas-phase electron diffraction;<sup>41</sup> from the latter, the C1–C3 distance was found to be 1.594 Å which, being similar to the C–C bond length in ethane (1.524 Å),<sup>42</sup> is suggestive of a C1–C3 bonding interaction. The nature of this bond has been much debated over the past 30 years, with theoretical<sup>43–55</sup> and experimental<sup>30,41,43,56–59</sup> studies offering contrasting evidence. For example, despite a closed-shell singlet ground state and a calculated bond energy of  $\sim 65$  kcal mol $^{-1}$ ,<sup>43</sup> photoelectron spectroscopy reveals the HOMO (C1–C3  $\sigma$  MO) to be non-bonding due to poor overlap of atomic orbitals between the 'inverted' carbon atoms.<sup>59</sup> This result is corroborated by synchrotron experiments that reveal depletion of electron density at the bond critical point.<sup>57</sup>

Such studies led to the proposal that the inverted C1–C3 bond in **1** should be best described as a charge-shift bond, with its stability attributed to resonance stabilization between the covalent ([C1 $\cdots$ C3], repulsive) and ionic ([C1 $^{(-)}$ ... $^{(+)}$ C3] and [C1 $^{(+)}$ ... $^{(-)}$ C3], attractive) structures.<sup>47,60</sup> This type of bonding arises through significant Pauli repulsion pressure between each of the C1/3–C2 'wing' bonds and the C1–C3 bond,<sup>61,62</sup> and accounts for both the observed closed-shell singlet ground state and the unusual positive sign of the Laplacian of the electron density at the critical point of the C1–C3 bond. However, the consequences of the unique electronic configuration of **1** on its reactivity remains an unanswered question.



Fig. 2 (a) Top: populations and symmetries of MOs in the (2,2) active space with their corresponding atomic orbital composition. Bottom left: effect of  $\sigma$ - $\pi$  delocalization on key geometric features of **1**. Inset: [CASSCF(2,2)]–[RHF] electron density difference plot for **1**, isovalue = 0.002 a.u. (b) DFT electron density difference plot for **1**: [BLYP]–[CASSCF(2,2)] (left), [B3LYP]–[CASSCF(2,2)] (center), [B2PLYP]–[CASSCF(2,2)] (right); isovalue of 0.01 a.u. Geometry optimized at CASSCF(2,2). All calculations use the def2-QZVPP basis set.

To further quantify the effects of the electronic repulsion within the propellane cage, we employed the complete active space (CASSCF) method (with a (2,2) active space) for the formal inclusion of static correlation. This allows electrons to populate both the C1–C3  $\sigma$  and  $\sigma^*$  orbitals (Fig. 2a). In 'normal' closed-shell organic molecules,  $>1.98$  electrons populate the occupied molecular orbitals (MOs), however in **1** we found this value to be 1.91 electrons, indicating that static correlation effects are important. Such behavior is typically only seen in systems with near-degenerate frontier orbitals, which is certainly not the case here ( $E_{\text{gap}} = 13.0$  eV, HF/def2-QZVPP). We propose that delocalization of electron density from the C1–C3  $\sigma$  to  $\sigma^*$  reduces the unusually high electronic repulsion inside the propellane cage; the effect of this delocalization can be seen in the stabilization of the structure of **1** by 19.2 kcal mol $^{-1}$ .

An electron density difference plot (Fig. 2a, inset) comparing the Hartree–Fock and CASSCF(2,2) densities reveals that, using CASSCF(2,2), charge is depleted from the center of the cage (red lobes), and is instead delocalized onto the bridge carbon atoms





(blue lobes). This is consistent with the depleted electron density that is observed at the critical point of the C1–C3 bond.

This electronic reorganization can be considered as ' $\sigma$ - $\pi$ -delocalization' ( $\sigma \rightarrow [\sigma^* + \pi]$ ), where electrons from the C1–C3  $\sigma$  MO partially populate the C1–C3  $\sigma^*$  orbital, which is of the correct symmetry to overlap with the  $\pi$  system formed from p orbitals on the bridge carbon atoms (Fig. 2a, top).<sup>63</sup> This effect can be observed through lateral compression of the cage to maximize this 'side-on' orbital overlap, decreasing the C2–C4 distance from 2.234 to 2.219 Å and increasing the C1–C3 distance from 1.540 to 1.589 Å, which is in excellent agreement with experiment (Fig. 2a, bottom left).

Overall, this description of the electronic structure of **1** is equivalent to the charge-shift bonding model introduced by Shaik and co-workers with a localized representation of the orbitals (see ESI Fig. S4†).<sup>47</sup> Moreover, the use of electron density difference plots provides an intuitive pictorial representation of the role of delocalization in **1**. It reveals that electron density is not only expelled from the cage along the C1–C3 axis, as was previously thought, but is also delocalized onto the bridge carbon atoms. This delocalization mechanism both reduces Pauli repulsion between the C1–C3 and C1–C2 bonding electrons, and stabilizes the cage towards fragmentation of the C1–C2 bond.

To reduce the cost and complexity of the calculations, we also sought a suitable DFT method to accurately model these delocalization effects and the reactivity of **1**. The performance of DFT depends directly on the electron density, and the challenge in describing **1** arises from low density in the C1–C3 region.<sup>57</sup> This problem can be qualitatively depicted using density difference plots (Fig. 2b), which were calculated for a range of DFT functionals and compared to those obtained with CASSCF(2,2). While the inexpensive GGA functional (BLYP)<sup>64,65</sup> accurately predicts the C1–C3 bond length to  $<0.01$  Å, it poorly reproduces the CASSCF(2,2) density, and substantially overdelocalizes electron density across the cage. The incorporation of some exact exchange into the functional (B3LYP)<sup>66</sup> overlocalizes electron density along the C1–C3 axis, resulting in a bond that is too short, and likely too strong. However, further augmentation to a double-hybrid GGA functional through the inclusion of MP2-like correlation (B2PLYP)<sup>67</sup> results in a lengthened C1–C3 bond, and a better match with the CASSCF(2,2) electron density. Further analysis of the singlet-triplet gap and vertical ionization potential for each of these functionals reveals double-hybrid GGA functionals to give the best performance of those tested (see ESI Fig. S13†). These results highlight how accuracy in geometries and energetics are not necessarily linked to accuracy in electronic structure.

A benchmark study on the reactivity of [1.1.1]propellane with anions, radicals and cations reactions also revealed that double-hybrid DFT was required to accurately describe the geometric and electronic features of this system (see ESI Fig. S17–S22†). Among them, B2GP-PLYP-D3BJ with the triple- $\zeta$  quality basis set def2-TZVP was found to afford good geometries and energies within 1 kcal mol<sup>−1</sup> of the reference coupled-cluster method DLPNO-CCSD(T).<sup>68–70</sup> Solvent effects, accounted with the SMD implicit model,<sup>71</sup> were found to be particularly important for

anionic and cationic reactions, but have a much smaller effect on radical reactivity.<sup>72</sup>

### The reactivity of [1.1.1]propellane with anions

Having identified the beneficial role of  $\sigma$ - $\pi$ -delocalization and a suitable computational methodology, we set out to explore the reactivity of **1** under a variety of conditions, and compare theoretical predictions with experimental reality. This began with a study of anionic addition, for which a number of examples of have been described that access aryl-, alkyl- and amino-substituted BCPs.<sup>25–29</sup> Given the apparent strain relief associated with this chemistry, it is perhaps surprising that extended reaction times ( $\sim 16$  h) and elevated temperatures ( $>50$  °C) are often required for successful addition. Walsh *et al.* have studied the related addition of dithiane anions,<sup>29</sup> complementing their synthetic work with DFT calculations at the M06-2X level of theory,<sup>73</sup> which revealed carbanion addition to be highly *endergonic*. These examples led us to question the role of strain relief in the reaction of anions with **1**, and in particular how a thermodynamically disfavored process could occur in a system thought to be primed to react.

Addition of an anion to **1** is analogous to a standard S<sub>N</sub>2 reaction, where the leaving group in this case is the distal C3 atom in the cage. However, the inverted geometry of the bridgehead carbon atom in **1** uniquely circumvents the need for planarization in the transition state,<sup>39</sup> which might suggest that the activation barrier has a greater contribution from a change in electronic structure than from geometric distortion.<sup>23,26,27,33</sup> To investigate the origin of this barrier to addition, a Distortion/Interaction Analysis (DIA) was performed on a model hydride addition to **1**.<sup>37–39</sup> This approach separates the overall energy of the addition [ $\Delta E(\text{total})$ ] into geometric distortion [ $\Delta E(\text{dist})$ ] and electronic interactions [ $\Delta E(\text{int})$ ] of the approaching species. On going from the reactant state to the transition state (TS),  $\Delta E(\text{int})$

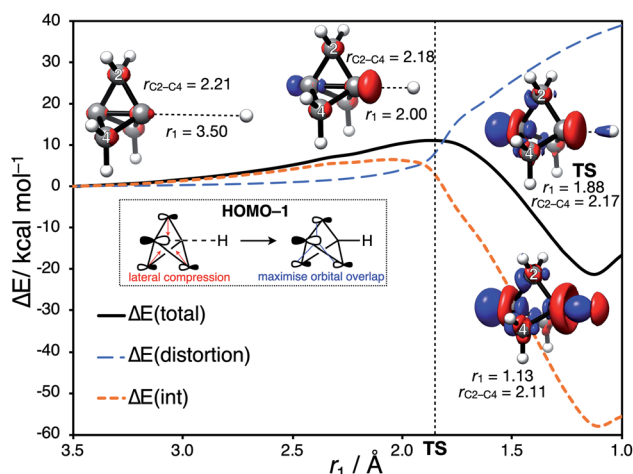


Fig. 3 Potential energy surface (PES), distortion (blue)/interaction (orange) energies and electron density difference plots for the addition of H<sup>−</sup> to **1**, calculated at the SMD(THF)-DLPNO-CCSD(T)/ma-def2-QZVPP//CASSCF(4,3)/ma-def2-QZVPP level of theory. Inset: cage compression maximizes orbital overlap in the HOMO−1.



was found to be positive, and to dominate over  $\Delta E(\text{dist})$  (70% contribution of  $\Delta E(\text{int})$  up to the TS) (Fig. 3). This is highly unusual: to our knowledge,  $\Delta E(\text{int})$  has not been observed to be positive and greater than  $\Delta E(\text{dist})$  for any other system.<sup>39</sup> Prior to the TS, the population of the  $\sigma$  and  $[\sigma^* + \pi]$  MOs remains constant while electron density is polarized by the approaching anion (Fig. 3, blue lobes, and ESI Fig. S7 and S8†), suggesting that electrostatic repulsion between the anion and **1** is the dominant factor causing the barrier to addition.

Significant structural distortion only occurs *after* the transition state, which suggests that geometric strain relief is not responsible for the reactivity of **1**. In fact, to maximize  $\sigma$ - $\pi$ -delocalization, the cage is laterally compressed to increase the overlap of each bridge carbon p orbital with the C3 p orbital in the HOMO-1 (Fig. 3, inset), which is a strain-increasing process.

To explore the implications of these findings in a chemically established setting, we modelled the addition of turbo-Hauser amides ( $\text{R}_2\text{NMgCl} \cdot \text{LiCl}$ ,  $\text{R} = \text{alkyl}$ ) to form 1-amino-BCPs (Fig. 4).<sup>26</sup> The reaction of **1** with amide anions was computed using  $\text{NH}_2^-$ ; the formation of the BCP anion adduct **2** was found to be reversible and endergonic ( $\Delta G^\ddagger = 21.3 \text{ kcal mol}^{-1}$ ,  $\Delta G^\circ = +1.3 \text{ kcal mol}^{-1}$ ), despite the formation of C-N in place of the weaker C1-C3 bond (balanced with the formation of a C-centered anion). DIA at the transition state (**TS**<sub>12</sub>) reveals  $\Delta E(\text{dist})$  and  $\Delta E(\text{int})$  are both positive, with  $\Delta E(\text{int})$  around twice the magnitude of  $\Delta E(\text{dist})$  (+8.3 and +4.1  $\text{kcal mol}^{-1}$  respectively). Only on complexation of the bicyclo[1.1.1]pentyl anion to a  $\text{MgCl}^+$  ion does the overall reaction become highly exergonic ( $\Delta G^\circ = -64.8 \text{ kcal mol}^{-1}$ ). This description appears consistent with the relatively harsh reaction conditions often required for anionic additions, as significant electronic

repulsion must be overcome prior to the development of the stabilizing  $\sigma$ - $\pi$ -delocalization.

### The reactivity of [1.1.1]propellane with radicals

Our attention next turned to the reactivity of **1** with radicals, chemistry that is of much utility in the synthesis of highly-functionalized BCPs.<sup>15–21,23,24</sup> Such reactions proceed through addition of a radical to the C1-C3 bond to give a bridgehead BCP radical that subsequently reacts either *via* atom transfer, or addition to a radical trap (such as an azodicarboxylate, a further molecule of **1**, or an organometallic species). Alkoxycarbonyl, alkyl and aryl radical additions have been studied using DFT (B3LYP, M06-2X,  $\omega$ B97X-D, B2PLYP)<sup>66,67,73,74</sup> by the Uchiyama group<sup>17</sup> and ourselves,<sup>18,20</sup> where the focus has lain on the fate of the bicyclo[1.1.1]pentyl radical.

We instead sought to study the initial radical addition to **1**, where a similar reactivity mode to the anionic addition was anticipated (Fig. 5). As in the anionic regime, the barrier for the addition of  $\cdot\text{CH}_3$  is dominated by the positive  $\Delta E(\text{int})$  term (66% contribution up to the TS, Fig. 5a).  $\Delta E(\text{dist})$  again increases most significantly *after* the TS, showing that this reactivity profile cannot be explained by simple geometric distortion. However, compared with the anionic addition, radical addition features a decreased electrostatic penalty in bringing two neutral species together; as a result, the additional electron density from approach of the radical species is incorporated inside the cage, rather than expelled from the back of the cage (Fig. 5a,  $r_1 = 2.25 \text{ \AA}$ ).

To understand how the repulsion from the injection of an additional electron is overcome during the reaction, we examined the change in CASSCF(3,3) MO coefficients along the reaction coordinate (Fig. 5b). At large separation, the orbitals resemble those of the isolated species;  $\phi_1$  and  $\phi_3$  correspond to the C1-C3  $\sigma$  and  $[\sigma^* + \pi]$  orbitals of **1** respectively (*vide supra*), and  $\phi_2$  corresponding to the p orbital of the radical species. As the two species approach one another, the MOs mix such that  $\phi_1$  is the bonding combination,  $\phi_2$  is non-bonding, and  $\phi_3$  is the antibonding combination of the  $\sigma$  orbitals in the forming and breaking bonds (Fig. 5b, bottom). The populations of these MOs are described by the coefficients  $C_i$  ( $i = 1, 2, 3$ ), which change over the course of the reaction.  $C_1$  [ $(\phi_1)^2(\phi_2)^1(\phi_3)^0$ ] (blue) and  $C_2$  [ $(\phi_1)^0(\phi_2)^1(\phi_3)^2$ ] (grey), which correspond to the  $\sigma \rightarrow [\sigma^* + \pi]$  delocalization in **1** itself, dominate at large separations, but decrease approaching the TS, where a new configuration  $C_3$  [ $(\phi_1)^1(\phi_2)^1(\phi_3)^1$ ] (red) emerges (Fig. 5c). This corresponds to the singlet diradical state of **1** interacting with the approaching radical species.

These results indicate that cleavage of the C1-C3 bond by a radical species requires access to an open-shell doublet electronic configuration. **1** itself is not diradicaloid in nature,<sup>54</sup> and as such perturbation of the electronic structure of **1** by an approaching radical necessarily results in a 'more diradical' configuration, which minimizes electronic repulsion inside the cage at the TS (Fig. 5c). This description is consistent with the equivalent charge-shift representation of the C1-C3 bond, where the covalent-ionic resonance interaction decreases upon



Fig. 4 Reaction free energy profile for turbo-Hauser amide addition to **1**, calculated at SMD(THF)-DLPNO-CCSD(T)/ma-def2-QZVPP//SMD(THF)-B2GP-PLYP-D3BJ/ma-def2-TZVP. Electron density difference plots calculated at SMD(THF)-DLPNO-CCSD(T)/ma-def2-QZVPP; isovalue = 0.01 a.u.; blue lobes indicate gain and red lobes indicate loss of electron density compared with the isolated species.





Fig. 5 (a) PES (DLPNO-CCSD(T)/def2-QZVPP//CASSCF(3,3)/def2-QZVPP) and distortion/interactions energies for the addition of  $\cdot\text{CH}_3$  to **1**. (b) (Top) PES (CASSCF(3,3)/def2-QZVPP) for the addition of a methyl radical to **1**, showing the proportion of each leading electronic configuration as a function of the forming C–C bond ( $r_1$ ). (Bottom) LCAO-MO depiction of the frontier molecular orbitals involved in the radical addition process to **1**, and the leading electronic configurations for the reaction.

interaction with a radical species, causing an increase in  $\Delta E(\text{int})$  prior to the TS.

To test the consequences of this proposal in a chemically established setting, the addition of  $\text{I}_2$  to **1** was selected as a model reaction, not least given its utility as a method for the titration of solutions of **1** (Fig. 6).<sup>75</sup> This reaction is presumed to occur *via* addition of  $\text{I}^\cdot$  to **1** to form an iodobicyclo[1.1.1]pentyl radical **3**, which is trapped by  $\text{I}_2$  to afford 1,3-diiodoBCP **4**, regenerating  $\text{I}^\cdot$  as a chain carrier. However, computational study of this reaction revealed that the initial addition of  $\text{I}^\cdot$  does not proceed to the expected carbon-centered radical; remarkably, barrierless exergonic association ( $\Delta G^\circ =$



Fig. 6 Proposed mechanism for the diiodination of **1** (top), and free energy profile for the diiodination of **1**, calculated at SMD( $\text{Et}_2\text{O}$ )-DLPNO-CCSD(T)/def2-QZVPP//SMD( $\text{Et}_2\text{O}$ )-B2GP-PLYP-D3BJ/def2-TZVP. Spin density of **3'** is plotted at an isovalue of 0.01 a.u.

$-2.1 \text{ kcal mol}^{-1}$ ) of the iodine radical to **1** is favored over full C1–C3 bond cleavage, with only a slight increase in the C1–C3 bond length observed in **3'** compared with **1** ( $\Delta r_2 = 0.02 \text{ \AA}$ ). Spin density is delocalized in the radical adduct **3'** (Fig. 6), despite the lack of distortion of the geometry of **1**; this causes C1–C3 bond weakening in order to accommodate the unpaired electron, resulting in facile iodine atom abstraction from  $\text{I}_2$  through the distal carbon atom, to form **4** and an iodine radical. We propose that full C1–C3 cleavage during radical addition can only occur if the formation of the new C–X bond is sufficiently strong to outweigh the loss of  $\sigma$ - $\pi$ -delocalization energy; in this example, the formation of a weak C–I bond results in the C1–C3 bond remaining intact. Should a strong C–X bond be formed upon addition, full cleavage of the inter-bridgehead C–C bond is seen (see ESI Fig. S29† for addition of halide and chalcogen hydride radicals to [1.1.1]propellane). These results suggest that **1** can undergo reversible radical reactions, which again challenges the role of strain relief as a reaction driving force.

### The reactivity of [1.1.1]propellane with cations

In both radical and anionic reactions, the propellane cage must accommodate an increase in electron density (Fig. 7). This raised the question of the consequence of *removing* electrons from the cage, as would be expected in cationic activation. The reactions of **1** with cations is an underexplored field; despite the observation of rapid protonation by acetic acid,<sup>30</sup> few cation-promoted reactions have been reported.<sup>31,32</sup> Cationic bicyclo[1.1.1]pentyl adducts are known to rapidly fragment to bicyclo[1.1.0]butyl-1-carbinyl cations (Fig. 7a),<sup>76</sup> which suggests that a significant change in the electronics of the cage occurs as addition takes place.



Fig. 7 (a) Addition of a cation to **1** results in cage fragmentation. (b) Electron density difference calculated at [CASSCF(2,3)/def2-QZVPP] – [CASSCF(2,2)/def2-QZVPP + HF/def2-QZVPP] level of theory for the adduct, [1.1.1]propellane and methyl cation respectively; isovalue = 0.01 a.u. (c) PES and distortion/interaction energies for the addition of a methyl cation to **1**, calculated at SMD(Et<sub>2</sub>O)-DLPNO-CCSD(T)/def2-QZVPP//CASSCF(2,3)/def2-QZVPP.

Analysis of the electron density distribution on a model system (methyl cation addition to **1**) revealed contrasting behavior to that seen with anions and radicals. While in the latter cases transition state barriers arise from the increase in electron repulsion during the approach of the two species, for cation addition, electron density is lost from the cage (Fig. 7b). In this case, charge transfer from **1** to the cation was found to be barrierless (Fig. 7c), and accompanied by a large negative  $\Delta E(\text{int})$ .

Intriguingly, the C1–C3 bond was found to *decrease* in length over the course of the addition, which contrasts with the instinctive expectation that bond cleavage should be accompanied by bond lengthening. Upon reaction at C1, the frontside C1–C2 bonds lengthen, while the backside C3–C2 bonds shorten. A More O'Ferrall–Jencks plot for the addition process as a function of the forming C1–CH<sub>3</sub> length ( $r_1$ ) and the breaking C1–C2 length ( $r_2$ ) reveals a relatively flat bifurcated potential energy surface, where initial charge transfer that cleaves the C1–C3 bond is accompanied by either delayed donation of electron density from the C1–C2 bond to stabilize the forming cation through cage fragmentation (Fig. 8, path A), or the formation of a bicyclo[1.1.1]pentyl cation that then spontaneously fragments (path B). Both paths lead to a non-classical bicyclo[1.1.0]butyl-1-carbinyl cation **5** with a partially-formed C1–C3 bond.<sup>77</sup> No local minimum or shoulder was observed for the bicyclo[1.1.1]pentyl cation, suggesting that this species is not a viable intermediate in the cationic reactivity of **1**.

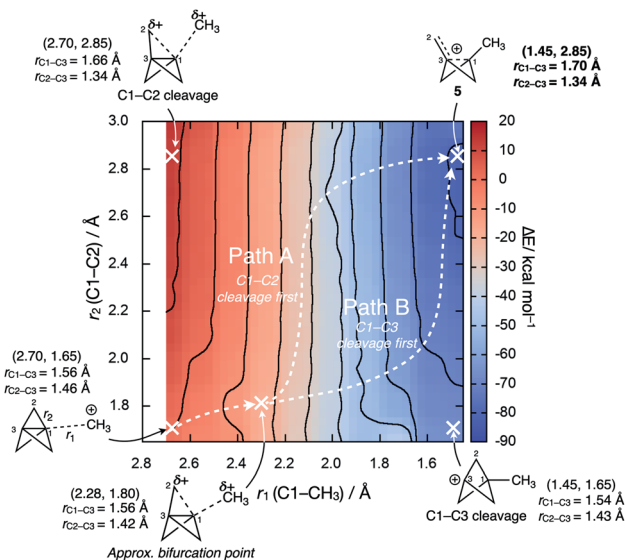


Fig. 8 More O'Ferrall–Jencks plot for the addition of a methyl cation to **1** as a function of the forming C1–CH<sub>3</sub> bond distance ( $r_1/\text{\AA}$ ) and the breaking C1–C2 bond distance ( $r_2/\text{\AA}$ ) calculated at the SMD(Et<sub>2</sub>O)-B2GP-PLYP-D3BJ/def2-TZVP level of theory.  $\times$  denotes points of interest including the approximate bifurcation point, and their corresponding ( $r_1$ ,  $r_2$ ) coordinates. Key distances for stationary point **5** are shown in bold.

We suggest that, ironically, C1–C2 fragmentation occurs due to the loss of Pauli repulsion inside the cage, as the resultant decreased population of the  $[\sigma^* + \pi]$  MO ( $\sigma$ – $\pi$  delocalization) weakens the C1–C2 bonds; the C1–C3 bond is also weakened and decreases in length as electron density is donated to the cation. This observation contrasts with analogous results from Jemmis and co-workers in calculations of halogen-bonded complexes of **1**, where the removal of electron density from the cage was suggested to *strengthen* the C1–C3 bond (B3LYP-D3BJ/QZ4P).<sup>45</sup>

### A unified model for [1.1.1]propellane reactivity

The above results provide a unified framework that explains the omniphilic reactivity of [1.1.1]propellane with anions, radicals and cations. Two distinct modes of reactivity are defined, with the main features summarized as follows:

(1) The *ground state* structure of **1** is stabilized by partial population of the C1–C3  $\sigma$ -antibonding/bridge  $\pi$ -bonding orbitals. The resultant  $\sigma$ – $\pi$ -delocalization relieves Pauli repulsion inside the cage (Fig. 9a). This leads to a moldable electron density that allows **1** to engage with electron-rich, electron-deficient and open-shell species.

(2) *Anionic and radical additions* to **1** involve an increase in electronic repulsion inside an already electron-rich cage, causing a barrier to addition. To offset this effect, stabilization after the transition state occurs through  $\sigma$ – $\pi$ -delocalization of electron density over the cage, which is maximized through compression of the cage (Fig. 9b, blue [anion] and red [radical] lines).







Fig. 9 (a) Summary of the omniphilic reactivity profile of **1**, explained through the change in the electron density distribution over the cage. (b) Compression/expansion of the propellane cage as a function of internuclear distance ( $r_1$ ) and the change in C2–C4 distance with respect to isolated [1.1.1]propellane ( $\Delta r_{C2-C4}$ ) (x denotes the position of the transition state for each reaction).

(3) *Cationic additions* are dominated by loss of electron density from the electron-rich cage through charge transfer to the cation. This process is barrierless due to the reduction of Pauli repulsion. However, as the  $\sigma$ – $\pi$ -delocalization that results from this repulsion is itself responsible for the structural integrity of the propellane cage (Fig. 9b, purple line), this loss of stabilization can result in spontaneous fragmentation to form a bicyclo[1.1.0]butyl-1-carbinyl cation.

These results provide a basis to develop new reactions of **1**, such as processes that capitalize on the unique structural distortion of the cage upon *loss* of electronic repulsion. We also predict that ‘reversible addition’ reactivity could feature in the formation of radical-pair complexes with species incapable of overcoming the delocalization energy of **1**. Moreover, we suggest that substitution of the bridge carbon atoms, either as heteroatoms or with carbon-based substituents, will modify the degree of  $\sigma$ – $\pi$ -delocalization both in the ground state and during reactions. The consequences of substitution on strain and electronic structure in [1.1.1]propellanes have been suggested to derive from changes in electronegativity; however, the ability of substituents to engage in  $\sigma$ – $\pi$ -delocalization is an alternative effect to consider,<sup>37,78,79</sup> for example stabilizing addition pathways by increased delocalization.

## Conclusions

The classical Lewis representation of [1.1.1]propellane places an inter-bridgehead bond along its central axis, which is thought to enable reactions through ‘rapid strain relief’. Here, we have developed a unified framework that challenges this view, and explains the omniphilic reactivity profile of **1**. We show that the ground state electronic structure of **1** is best described as a delocalized electron density spanning the entire cage. The result is a moldable electron density that imparts a broad reactivity profile on **1**: anionic and radical additions are favored by stabilization of adducts through increased  $\sigma$ – $\pi$ -delocalization, while reactions with cationic species are driven by charge transfer that relieves Pauli repulsion. We anticipate that this understanding will open up possibilities for the exploration of new reactivity of [1.1.1]propellane and related systems, and will provide a general framework to understand the behavior of

strained cage-like molecules, which continue to be of both fascination and utility in organic synthesis.

## Conflicts of interest

There are no conflicts to declare.

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