Starting a subnanoscale tank tread: dynamic fluxionality of boron-based B_{10}Ca alloy cluster†

Ying-Jin Wang,‡ab Lin-Yan Feng‡a and Hua-Jin Zhai a

Alloying an elongated B_{10} cluster with Ca is shown to give rise to a dynamically fluxional B_{10}Ca cluster, the latter behaving like a tank tread at the subnanoscale. Computer global search identifies the B_{10}Ca C_{2} (‡) global-minimum structure, which is chiral in nature and retains the quasi-planar moity of bare B_{10} cluster with Ca capped at one side, forming a half-sandwich. The rotation barrier of B_{10}Ca cluster is reduced with respect to B_{10} by one order of magnitude, down to 1 kcal mol^{-1} at the PBE0/6-311+G* level, which demonstrates structural fluxionality at 600 K and beyond via molecular dynamics simulations. Structurewise, the Ca alloying in B_{10}Ca cluster generates rhombic defect holes, preactivating the species and making it flexible against deformation. Chemical bonding analyses indicate that the B_{10}Ca cluster is a charge-transfer [B_{10}]^{2−}[Ca]^{2+} complex, being doubly π/σ aromatic with the 6π and 10σ electron-counting. Such a pattern offers ideal π/σ delocalization and facilitates fluxionality. In contrast, bare B_{10} cluster has conflicting aromaticity with 6π and 8σ electrons, which is nonfluxional with a barrier of 12 kcal mol^{-1}. Double π/σ aromaticity versus conflicting aromaticity is a key mechanism that distinguishes between fluxional B_{10}Ca and nonfluxional B_{10} clusters, offering a compelling example that the concept of aromaticity (and double aromaticity) can be exploited to design dynamically fluxional nanosystems.

1. Introduction

Molecular rotors are interesting nanoscale or subnanoscale systems,1–19 in which the constituent fragments move collectively and concertedly against each other, akin to nanomachines. Such nanosystems are rare in chemistry and were primarily discovered during the past decade. Boron appears to be a magic element for nanorotors. Several types of boron-based nanorotor have been developed. Firstly, a Wankel motor B_{19}− cluster was described by Merino and coworkers5 with a negligible in-plane rotation barrier (less than 0.1 kcal mol^{-1}), immediately following an experimental study of the gas-phase B_{10} cluster with concentric dual π aromaticity via photoelectron spectroscopy (PES) in 2010.19 Similar molecular dynamics was discussed earlier in mixed B−C clusters (C_{2}B_{6}, C_{1}B_{9}^{3+}, and C_{3}B_{11}),2 although those species are highly energetic on their potential energy surfaces and thus not viable. Molecular Wankel motors have since been expanded to other systems,5–9 such as the B_{13}+ cluster. Second, Zhai and coworkers10 discovered in 2015 the so-called subnanoscale tank treads, which feature elongated rather than circular geometries with a peripheral ring gliding freely around an elongated inner core. Typical examples are the B_{11}− and B_{11} clusters, which have a rotation barrier of 0.35 and 0.60 kcal mol^{-1}, respectively. An isostructural B_{12}C model cluster12 was designed subsequently for mechanistic insights into subnanoscale tank treads. Third, a binary Mg_{2}B_{8} cluster was shown to function like a nanocompass14 and a coaxial triple-layered Be_{6}B_{11}− cluster was characterized in 2017 with dual dynamic modes: revolution (orbiting) of the peripheral B_{11} ring versus rotation (twisting) of the prismatic Be_{6} core.15 Intriguingly, the Be_{6}B_{11}− cluster mimics the dynamic Earth-Moon system at the subnanoscale and yet miniaturizes the latter in size by 18 orders of magnitude.

The ultimate mechanisms of boron-based nanorotors should lie in the electron-deficiency of boron, which leads to highly unusual geometries of boron clusters,16–26 featuring planar or quasi-planar (2D) structures up to 40 atoms for anions.26 Among other notable boron clusters are a bowl-like B_{36} borophene29 and cage-like B_{40}/B_{40} borospherenes.30,37 In terms of chemical bonding, boron clusters are governed by π/σ aromaticity, antiaromaticity, double aromaticity, and...
conflicting aromaticity, which underlie their 2D geometries (as well as their unique structural dynamics). Note that the delocalized π/σ electrons in boron clusters are clouded over a 2D plane, rather than on a single ring such as in benzene.

It is of fundamental importance to “tune” the rotation barrier of molecular rotors, which not only provides key insights into mechanisms for structural dynamics, but also facilitates control of the movements: halting or starting. Merino and coworkers showed that one C substitution stops the B19− Wankel motor, resulting in a nonfluxional B18C cluster with a barrier of 27.6 kcal mol−1. On the other hand, Popov et al.17 reported that complexion of B12 with a metal atom can reduce the rotation barrier of half-sandwich RhB12− and CoB12− clusters from 19.5 kcal mol−1 down to 8.7–8.9 kcal mol−1. Likewise, Liu et al.18 designed a metal doped IrB12− Wankel motor with a barrier of 5.0 kcal mol−1. Nevertheless, these barriers are still quite substantial. It is highly desirable to further reduce the barrier, for example, down to 1 kcal mol−1 or less.

We shall meet the above challenge using the B10Ca and B10 clusters. The clusters are chosen for a couple of reasons. First, the bare B10 cluster is known to be nonfluxional, with a triangul}
shrinkage at the other, and atom B10 being substantially popped out.33 The changes hinder the interconversion between GM and TS. The anticipated barrier lies in the distinct coordination environments: two hexacoordinate B centers in GM versus one heptacoordinate B and one pentacoordinate B in TS.

Structural parameters of B10Ca and B10 GM/TS clusters at PBE0 are presented in Fig. S2 (ESI†). All five structures have similar perimeters for the outer B8 ring: 12.5–12.8 Å. However, bond distances in the inner B2 core are slightly expanded from 1.63–1.66 Å in B10 to 1.70–1.72 Å in B10Ca. Radial B–B links are 1.79 Å in average for B10Ca GM, which are elongated relative to B10 GM (1.70 Å in average). Thus, upon Ca alloying, interior B–B links undergo discernible expansion, leading to bowl-like B10Ca clusters (GM, TS1, and TS2). Note that such changes do not necessarily imply that interior B–B links in B10Ca have lower bond orders (vide infra).

3.2. Wiberg bond indices and natural atomic charges

The NBO analyses provide quantitative data for WBIs (Fig. 2) and natural atomic charges (Fig. S3, ESI†). WBIs are highly non-uniform among peripheral and interior B–B links, between GM and TS structures, and between B10Ca and B10 clusters. We mention a couple of general observations. First, WBIs on the periphery are greater than 1.0, whereas those for interior links are below 1.0. The observation implies that the periphery is bound by two-center two-electron (2c–2e) skeleton σ bonds along with delocalized π/σ clouds. In contrast, interior B–B links are entirely held together by delocalized π/σ bonds. Second, there appears to be enhanced peripheral bonding in B10Ca GM (WBIs in total: 10.1) as compared to B10 GM (WBIs in total: 9.6). Third, WBIs of B–Ca links in B10Ca GM are minor: 0.07/0.10/0.15 for the B9–Ca/B3–Ca/B4–Ca links, suggesting negligible covalent bonding between Ca and B atoms.

Natural atomic charges in the B10Ca GM cluster (Fig. S3b, ESI†) are consistent with a charge transfer complex: [B10]2+[Ca]2+. Specifically, Ca atom has a positive charge of +1.52 |e|, with the peripheral B8 ring collectively carrying a negative charge of −1.44 |e| and the inner B2 dimer being roughly neutral. Note that atoms B4 and B8 each has −0.31 |e|, which are the most negative due to their closeness to the Ca center. The B10 GM (Fig. S3a, ESI†) shows intramolecular charge redistribution, so that the central B9B3B10B7 diamond carries negative charges (−0.22/−0.17 |e| per atom) and the four B2/B4/B6/B8 corners are positive (+0.20 |e| per atom). Going from B10 GM to B10Ca GM, the major change in charges also lies in the above four corners, where the B4/B8 centers each gains −0.51 |e| from Ca and B2/B6 each gets −0.35 |e|. Overall, the distribution of negative charges is more uniform in B10Ca GM relative to B10 GM.35

4. Discussion

4.1. Chemical bonding in the B10Ca cluster

The nature of bonding ultimately governs the dynamic fluxionality of a molecular system. Therefore, we start our discussion with chemical bonding in B10Ca cluster. Considering the electron configurations of B 2s22p2 and Ca 4s2, the B10Ca cluster has 32 valence electrons. The occupied CMOs are all based on the B10 fragment (Fig. 3), consistent with a [B10]2+[Ca]2+ charge-transfer complex; see above. The CMOs can be straightforwardly sorted into three categories. Subset (a) is primarily clouded on the periphery with major contributions from B 2s atomic orbitals (AOs). These CMOs have 0 to 4 nodal planes from bottom up (with three quasi-degenerate pairs in the middle), which follow the orbital building principles and readily recombine as eight 2c–2e Lewis B–B σ single bonds on the periphery.34

For the π framework in subset (c), three CMOs have 0 and 1 (quasi-degenerate) nodal planes, where the intrinsic nodal plane associated with pz AOs is not counted, as routine. The
three CMOs are analogous to the prototypical π sextet in benzene, thus endowing the B_{10}Ca cluster with π aromaticity according to the (4n + 2) Hückel rule. The Ca atom only contributes less than 7% in these π CMOs. For the delocalized σ framework (subset (b)), the five CMOs again follow the orbital building principles with 0 up to 2 nodal planes (including two quasi-degenerate pairs). This CMO pattern makes it imperative to claim σ aromaticity for the B_{10}Ca cluster. Indeed, the 10σ electron-counting conforms to the (4n + 2) Hückel rule. In summary, the CMO analysis indicates that the B_{10}Ca cluster possesses π/σ double aromaticity with 6π and 10σ electrons, respectively. In this model, there are no 2c–2e Lewis B–B π/σ bonds inside the peripheral B₈ ring, which explains why all those B–B links have low WBI values (less than 1.0; Fig. 2b).

The bonding picture is borne out from electron localization functions (ELFs) and AdNDP analysis (Fig. 4). The ELF and

![Fig. 2](image_url) Calculated Wiberg bond indices (WBI) from natural bond orbital (NBO) analysis at the PBE0/6-311G* level. (a) B_{10} GM and TS. (b) B_{10}Ca GM, TS₁, and TS₂.

![Fig. 3](image_url) Pictures of occupied canonical molecular orbitals (CMOs) of B_{10}Ca GM cluster, calculated at the PBE0/6-311+G* level. The CMOs are sorted to three subsets: (a) peripheral two-center two-electron (2c–2e) Lewis B–B σ single bonds; (b) global, delocalized 10σ framework; and (c) global, delocalized 6π framework.
ELF$_\sigma$ data are consistent with localized Lewis bonds on the periphery, as well as delocalized $\pi/\sigma$ frameworks. The AdNDP scheme of B$_{10}$Ca cluster (Fig. 4b) reproduces peripheral 2c–2e $\sigma$ bonds$^{33}$ and global $\pi$ bonds. For the delocalized $\sigma$ framework, we chose to partition it as one 2c–2e $\sigma$ bond on the inner B$_2$ dimer and four 3c–2e/4c–2e $\sigma$ bonds around the four corners, as hinted by the ELF$_\sigma$ data. It should be stressed that this “island” $\sigma$ scheme is only an approximation. In particular, the 2c–2e $\sigma$ bond has an occupation number (ON) of 1.61 $|e|$ only. Nonetheless, the scheme is useful in elucidating the structural dynamics (vide infra).

The CMOs, ELFs, and AdNDP schemes of TS$_1$/TS$_2$ structures of the B$_{10}$Ca cluster are shown in Fig. 5 and S4–S6 (ESI†). The basic bonding information is similar to that described above, except for a slight spatial shift of the $\pi/\sigma$ clouds, which is associated with geometric changes between GM and TS$_1$/TS$_2$ structures. For example, the AdNDP scheme for the 10$\sigma$ framework in TS$_1$ (Fig. 5b) is similar to that in the GM (Fig. 4b). However, upon clockwise rotation of TS$_1$ relative to GM, two diamond 4c–2e $\sigma$ bonds in the GM shrink spatially and become 3c–2e ones. This effectively generates a flow of $\sigma$ clouds within the islands, whose direction is anticlockwise. For a following step (not shown), certain 3c–2e $\sigma$ bonds may expand to 4c–2e ones. Therefore, during dynamic rotation of the B$_{10}$Ca cluster, delocalized $\sigma$ clouds flow (and shrink/expand) continuously like a liquid and counter the direction of geometric rotation, which are maintained approximately on the four corner islands and underlie the low rotation barrier. The delocalized $\pi$ clouds behave similarly.

For comparison, the CMOs, ELFs, and AdNDP patterns of GM/TS structures of bare B$_{10}$ cluster are shown in Fig. 6 and S7–S9 (ESI†). The B$_{10}$ GM cluster (Fig. S7b, ESI†) differs from B$_{10}$Ca GM (Fig. 3b) by one CMO less in the delocalized $\sigma$ framework, which becomes the lowest unoccupied molecular orbital (LUMO) in the former species. Based on the above understanding of B$_{10}$Ca, bare B$_{10}$ cluster should be classified as $\sigma$ antiaromatic, which has a delocalized 8$\sigma$ system and follows the 4n Hückel rule.$^{55,56}$ Indeed, the B$_{10}$ cluster is more elongated than B$_{10}$Ca, a typical consequence of $\sigma$ antiaromaticity; see Section 3.1.$^{58}$ Overall, the B$_{10}$ cluster has conflicting aromaticity with 6$\pi$ and 8$\sigma$ electrons.

It is interesting to quantitatively follow the orbital contributions of the inner B$_2$ core in B$_{10}$Ca GM and B$_{10}$ GM clusters. The B$_2$ core in B$_{10}$Ca contributes 3.32, 1.84, and 0.87 $|e|$ to subsets (a)–(c), respectively (Fig. 3), compared to 3.89, 1.02, and 1.52 $|e|$ in B$_{10}$ (Fig. S7, ESI†). In other words, the B$_2$ core has similar total contributions to $\sigma$ CMOs: 5.16 $|e|$ in B$_{10}$Ca versus 4.91 $|e|$ in B$_{10}$. However, its contributions to $\pi$ CMOs differ for the species: 0.87 $|e|$ in B$_{10}$Ca versus 1.52 $|e|$ in B$_{10}$. The latter observation suggests that the $\pi$ clouds are probably more delocalized on the plane in B$_{10}$ GM, which is a con for the dynamic fluxionality of bare B$_{10}$ cluster, in particular considering that it has a rugged shape. In contrast, the $\pi$ clouds in B$_{10}$Ca GM appear to lean to the periphery, which benefits a low dynamic barrier, as supported by the comparative ELF$_\pi$ patterns (Fig. 4a versus Fig. 6a). The above data also indicate that the B$_2$ core has collectively 5.84 $|e|$ in B$_{10}$Ca GM versus 6.43 $|e|$ in B$_{10}$ GM. Indeed, NBO data on natural atomic charges show that the B$_2$ core is practically neutral in B$_{10}$Ca and negatively charged in B$_{10}$ (by $-0.22$ $|e|$ per atom; Fig. S3, ESI†).

**Fig. 4** (a) Electron localization functions, ELF$_\sigma$, and ELF$_\pi$, of B$_{10}$Ca GM cluster. (b) Chemical bonding pattern of B$_{10}$Ca GM cluster on the basis of adaptive natural density partitioning (AdNDP) analysis, wherein the global 10$\pi$ framework is approximated to “island” $\sigma$ bonds. Occupation numbers (ONs) are indicated.
4.2. Starting a subnanoscale tank tread: distinct structural dynamics of B\textsubscript{10}Ca and B\textsubscript{10} clusters

Two key structural features of the present B\textsubscript{10}Ca and B\textsubscript{10} systems make it rather challenging to tune/control their rotation barriers. First, the clusters assume elongated (rather than circular) geometries akin to tank treads, which are generally less fluxional than molecular Wankel motors. Second, bare B\textsubscript{10} cluster is composed entirely of close-packed B\textsubscript{3} triangles without “defect” holes, which is intuitively anticipated to be rigid against deformation. Nevertheless, the energetics data in Section 3.1 clearly show that Ca alloying in the B\textsubscript{10}Ca GM cluster effectively diminishes its rotation barrier down to 0.95 and 1.25 kcal mol\textsuperscript{-1} at the PBE0 level, with respect to TS\textsubscript{1} and TS\textsubscript{2}, respectively. This is a remarkable achievement considering that bare B\textsubscript{10} cluster is nonfluxional with a barrier of 12.32 kcal mol\textsuperscript{-1}. To our knowledge, this is the first example of barrier reduction of a molecular rotor by one order of magnitude and down to \( \frac{1}{12.32} \) kcal mol\textsuperscript{-1}, a highly desirable level for dynamic fluxionality.

With the GM and TS\textsubscript{1}/TS\textsubscript{2} structures of the B\textsubscript{10}Ca cluster being identified (Fig. 1b), the dynamic structural evolution is relatively simple as illustrated in Fig. 7, assuming that the peripheral ring rotates clockwise. Starting with GM\textsubscript{1} and let the B\textsubscript{9}–B\textsubscript{2} link rotate by about 18.9°, one reaches TS\textsubscript{1} that corresponds to TS\textsubscript{1} in Fig. 1b. From GM\textsubscript{1} to TS\textsubscript{1}, rhombic B2B3B10B9 and B7B9B10B6 holes turn to squares. Going down the barrier, the B9–B2 link rotates by another 17.5° and the system returns to GM\textsubscript{2}. Due to the chiral nature of B\textsubscript{10}Ca cluster with \( C_2 \) symmetry, GM\textsubscript{1} and GM\textsubscript{2} are a pair of enantiomers and thus isoenergetic. However, the system does not recover its initial orientation. Further rotating the B9–B2 link slightly by 7.1° and making the B10–B2 and B9–B6 links, one reaches TS\textsubscript{2}. The latter differs from TS\textsubscript{1} with a higher barrier and it corresponds to TS\textsubscript{2} in Fig. 1b. From TS\textsubscript{2} to GM\textsubscript{3}, the B9–B2 link rotates another 7.1° and breaks, generating two new rhombic holes: B1B2B10B9 and B6B9B10B5. This completes a fully cycle, in which the cluster returns to its initial structure as well as orientation. During this process, the peripheral ring moves by one B–B link (or about 51°).

Despite being nonfluxional with a substantial barrier, the structural evolution of bare B\textsubscript{10} cluster can be followed on the basis of GM and TS structures identified. The process is simpler (Fig. 8). Upon rotation of the B9–B2 link clockwise by 12.0°, the B9–B3 link breaks and two rhombic B9B2B3B10 and B7B9B10B6 holes emerge, which leads to TS\textsubscript{1} that corresponds to TS in Fig. 1a. Further rotate the B9–B2 link by 38.1° and the system recovers GM\textsubscript{2}. Overall, the peripheral atoms move by one B–B link (or about 51°) from GM\textsubscript{1} to GM\textsubscript{2}.
Fig. 6  (a) ELF$_\sigma$ and ELF$_\pi$ of B$_{10}$ GM cluster. (b) AdNDP bonding pattern of B$_{10}$ GM cluster, wherein the global 8$\sigma$ framework is approximated to island $\sigma$ bonds. ONs are indicated.

Fig. 7  Structural evolution of the B$_{10}$Ca GM cluster during dynamic rotation, assuming that the peripheral B$_9$ ring moves clockwise around the B$_2$ core. Two energy barriers, TS$_{1-2}$ and TS$_{2-3}$, are to be overcome for the cluster in order to fully recover its initial structure and orientation. These barriers correspond to the TS$_2$ and TS$_3$ structures (Fig. 1b), respectively.
To demonstrate the dynamic fluxionality of B\textsubscript{10}Ca cluster, we performed BOMD simulations at selected temperatures of 300, 600, and 900 K. A typical simulation at 900 K is presented in the form of a short movie (see the ESI†), which was carried out for 50 ps using the CP2K package at the PBE0/DZVP level. Initial conditions were chosen to correspond to the microcanonical ensemble (NVE). The extracted movie roughly covers a time span of 2.5 ps. It is obvious that at 900 K the B\textsubscript{10}Ca cluster behaves vividly like a subnanoscale tank tread, with the peripheral B\textsubscript{8} ring gliding freely as a flexible chain around the B\textsubscript{2} core. In particular, the peripheral B\textsubscript{8} ring appears to be rather robust during rotation, which rotates but does not break. Similar motion is observed at 600 K except that it is slower, whereas preliminary data at 300 K do not reveal dynamic fluxionality.

As for mechanisms that distinguish between the dynamic properties of B\textsubscript{10}Ca and B\textsubscript{10} clusters, the most important one is their bonding, as detailed in Section 4.1. Note that in prior studies people have attempted to unravel the connection between aromaticity (or electron delocalization) and fluxionality, in which the magnetic response is discussed.\textsuperscript{57} The present data offer key insights using the CMOs and AdNDP analyses. While both B\textsubscript{10}Ca and B\textsubscript{10} are 6\pi aromatic, they differ in the \sigma framework: B\textsubscript{10} is \sigma antiaromatic\textsuperscript{35,36} with 8\sigma electrons and B\textsubscript{10}Ca has \sigma aromaticity (10\sigma electrons). Double \pi/\sigma antiaromaticity in B\textsubscript{10}Ca cluster makes it sort of circular with a smooth bowl-shaped surface, facilitating \pi/\sigma delocalization and structural fluxionality. In contrast, the B\textsubscript{10} cluster has a rugged surface\textsuperscript{31} and deteriorated \pi/\sigma delocalization.

In terms of geometry, the C\textsubscript{2} symmetry of B\textsubscript{10}Ca GM cluster (Fig. 1b) has two rhombic defect holes, while B\textsubscript{10} GM cluster features triangular close-packing. Such defects are known to be critical for the dynamic fluxionality of subnanoscale tank treads (and nanorotors in general).\textsuperscript{19} Thus, alloying with Ca induces structural flexibility for B\textsubscript{10}Ca cluster, which serves as “pre-activation” of the system and effectively reduces the dynamic barrier. As for the B\textsubscript{10} cluster, 8\sigma antiaromaticity leads to an elongated GM as well as substantial geometric changes between GM and TS (Fig. 1a): expansion at the left side versus shrinkage at the right. In fact, while delocalized \sigma clouds in the GM are balanced for left versus right (Fig. 6b), they become highly uneven in the TS so that the left side only has one 5c–2e \sigma bond and the right has three 3c–2e/4c–2e bonds (Fig. S9b, ESI†). Such a change is induced entirely by a rotation of the peripheral ring, which is an intriguing consequence of \sigma antiaromaticity and clearly underlies its large barrier. Lastly, we comment on a consequence of the two inequivalent TS structures of B\textsubscript{10}Ca cluster (Fig. 1b). All molecular rotors to date move randomly (clockwise or anticlockwise). However, the occurrence of two inequivalent TS structures for B\textsubscript{10}Ca hints that the system can at least be initiated with one direction preferred to another, which simplifies further control or manipulation of the dynamics.

4.3 Change in WBIs as a semi-quantitative indicator of dynamic fluxionality

To shed further light on the distinct dynamics of B\textsubscript{10}Ca and B\textsubscript{10} clusters, we make use of the WBI data from NBO analyses.
Intuitively, the barrier of a molecular rotor is connected to the WBI change between the GM and TS structures, which is denoted here as $\Delta$WBI. In the present systems, the radial links govern in-plane rotation and determine the barrier. Thus, we follow all radial B–B links. As an example, $\Delta$WBI values of the $B_{10}Ca$ cluster with respect to TS$_2$ are plotted in Fig. 9, which are compared to those of bare $B_{10}$ cluster. Here the B–B links are labeled numerically, starting from the far-left link (B9–B8) as “1” and counting clockwise up to “10” (links B9–B2 and B10–B6 included). Positive $\Delta$WBI indicates elongation and negative compression. Red dots are $\Delta$WBIs of the $B_{10}Ca$ cluster and a horizontal blue line is their average. Comparative data of the $B_{10}$ cluster are shown with filled squares.

The $B_{10}Ca$ cluster shows a smooth $\Delta$WBI curve with relatively minor and uniform values for all radial B–B links, consistent with a low barrier. In contrast, large $\Delta$WBIs and in particular spikes are present for the $B_{10}$ cluster, whose B–B links are labeled again with the far-left one (B9–B1) being “1” and counted clockwise up to “10”. Specifically, link “8” (B10–B7; Fig. 1a) has the largest $\Delta$WBI, which has to elongate substantially and eventually break in order for the cluster to reach the TS structure, thus defining the barrier height of $B_{10}$ cluster. For comparison, links B9–B3 and B10–B7 in $B_{10}Ca$ GM$_3$ (Fig. 7) are “preactivated” upon alloying with Ca atom, and they are no longer the bottleneck for fluxionality, which effectively diminish the barrier (for TS$_1$). Moreover, there is no need to break any B–B links from GM$_3$ to TS$_2$ (anticlockwise; Fig. 7), but to shrink links B9–B2 and B10–B6, which have the largest negative $\Delta$WBIs, in line with a low barrier for TS$_2$. The comparative $\Delta$WBI data of $B_{10}Ca$ and $B_{10}$ clusters distinguish between the two species, demonstrating a semi-quantitative measure for dynamic fluxionality.

5. Conclusions

We have demonstrated, via alloying with Ca atom, a means to tune the structural dynamics of a subnanoscale tank tread $B_{10}Ca$ cluster. The strategy efficiently reduces the rotation barrier by one order of magnitude, from 12 kcal mol$^{-1}$ in $B_{10}$ down to around 1 kcal mol$^{-1}$ in $B_{10}Ca$. The $B_{10}Ca$ global-minimum cluster is half-sandwich in shape, which is dynamically fluxional at 600 K and beyond, whereas bare $B_{10}$ cluster is nonfluxional. In terms of structures, Ca alloying generates two rhombic defect holes in $B_{10}Ca$ cluster, which preactivates the system for fluxionality. Chemical bonding analyses show that the $B_{10}Ca$ cluster is a charge-transfer complex and can be formulated as $[B_{10}]^{2-}[Ca]^{2+}$, featuring double (6$\pi$ and 10$\sigma$) aromaticity. The bonding pattern leads to a more circular geometry and ideal $\pi/\sigma$ delocalization, facilitating dynamic fluxionality. In contrast, the $B_{10}$ cluster has conflicting aromaticity (6$\pi$ versus 8$\sigma$), as well as a more elongated and rugged shape. Thus, our work demonstrates that $\pi/\sigma$ aromaticity can be utilized as a key concept to design dynamically fluxional nanosystems. To semi-quantitatively describe dynamic fluxionality, the change in Wiberg bond indices between global-minimum and transition-state structures serves as a useful indicator, which clearly distinguishes between a fluxional $B_{10}Ca$ cluster and a nonfluxional $B_{10}$ cluster.

Conflicts of interest

There are no conflicts to declare.
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50 The B_{10} GM cluster has a length of 4.76 Å and width of 3.13 Å (with length-to-width ratio of 1.52). Those of the B_{10} TS, B_{10}Ca TS_{1}, and B_{10}Ca TS_{2} species are 4.13/3.54, 4.01/3.38, and 4.50/3.28 Å, with ratios of 1.17, 1.19, and 1.37, respectively. Thus, there is a marked ratio change from B_{10} GM to B_{10} TS. Also, B_{10}Ca TS_{1} is particularly circular.

51 Of all five key structures in Fig. 1, the peripheral B_{8} ring is perfectly planar only in B_{10} GM. For other species, the out-of-plane distortions of central B atoms are estimated roughly. In B_{10} GM cluster, atom B9 is slightly out-of-plane toward inside (by 0.3 Å) and B10 toward the opposite side. For the B_{10} TS, B_{10}Ca TS_{1}, and B_{10}Ca TS_{2} species, atoms B9/B10 are popped out by 0.3/1.0, 0.9/0.9, and 0.8/0.8 Å, respectively.

52 There appears to be a general trend that a B center with a higher coordination and stronger bonds gets some net negative charges from surrounding B atoms. In this way, the hexacoordinate B9/B10 centers and tetracoordinate B3/B7 atoms in B_{10} GM carry negative charges (Fig. S3a; ESI†). For B_{10} TS, the B10 center remains strongly coordinated by five B atoms, whereas the B9 center is only loosely bound despite its heptacoordination nature. The latter leads to less negative charge.

53 Strictly speaking, the assignment of eight peripheral Lewis σ bonds is an approximation. The bottom five of these CMOs have nonnegligible contributions from central B9/B10 atoms (Fig. 3a). Quantitatively, the B9/B10 core accounts for 20.7% of these eight σ CMOs, which explains why their occupation numbers (ONs; Fig. 4b) in the AdNDP scheme deviate from an ideal value of 2.00 |e|.

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55 A previous work assigned the delocalized σ framework of B_{10} GM cluster as one 2c–2e σ bond in B_{2} core (although specifically called for caution) and three global σ bonds. The latter were proposed to render σ aromaticity for B_{10} cluster, due to the 6σ electron-counting. Our present analysis suggests that all these 8σ electrons should be treated collectively (Fig. S7b, ESI†) and B_{10} cluster is σ antiaromatic rather than σ aromatic. Indeed, the ELF data and our AdNDP scheme (island version; Fig. 6) do not support the idea of a 2c–2e σ bond within the B_{2} core.

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