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From inverse sandwich $Ta_2B_7^+$ and Ta_2B_8 to spherical trihedral $Ta_3B_{12}^-$: prediction of the smallest metallo-borospherene†

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Transition-metal-doped boron nanoclusters exhibit interesting structures and bonding. Inspired by the experimentally discovered inverse sandwich D_{6h} Ta_2B_6 and spherical trihedral D_{3h} $La_3B_{18}^-$ and based on extensive first-principles theory calculations, we predict herein the structural transition from perfect di-metal-doped inverse sandwich D_{7h} $Ta_2B_7^+$ (1) and D_{8h} Ta_2B_8 (2) to tri-metal-doped spherical trihedral D_{3h} $Ta_3B_{12}^-$ (3). As the smallest metallo-borospherene reported to date, $Ta_3B_{12}^-$ (3) contains three octa-coordinate Ta atoms as integral parts of the cage surface coordinated in three equivalent η^8 - B_8 rings which share two eclipsed equilateral B_3 triangles on the top and bottom interconnected by three B_2 units on the waist. Detailed orbital and bonding analyses indicate that both $Ta_2B_7^+$ (1) and Ta_2B_8 (2) possess $\sigma + \pi$ dual aromaticity, while $Ta_3B_{12}^-$ (3) is $\sigma + \pi + \delta$ triply aromatic in nature. The IR, Raman, and UV-vis or photoelectron spectra of the concerned species are computationally simulated to facilitate their future spectroscopic characterizations.

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1. Introduction

Boron exhibits strong propensity to form multicentre-two-electron bonds (mc-2e bonds) in both polyhedral molecules and bulk allotropes due to its prototypical electron-deficiency.¹⁻⁴ Joint photoelectron (PE) spectroscopy experimental and first-principles theory investigations in the past two decades have unveiled a rich landscape for B_n ^{-/0} boron nanoclusters from planar or quasi-planar structures ($n = 3-38, 41-42$) to cage-like borospherenes ($n = 39, 40$).³⁻⁷ Cage-like D_{2d} $B_{40}^{-/0}$ are the first borospherenes discovered in 2014,⁵ while C_3/C_2 B_{39}^- are the first axially chiral borospherenes confirmed in experiments as the global minima (GM) of the B_n^- monoanions.⁶ These discoveries mark the onset of borospherene chemistry parallel to fullerene chemistry. The borospherene family has been systematically expanded by our group at first-principles theory levels to the cage-like B_n^q series in different charge states ($n = 36-42, q = n-40$) which all possess twelve delocalized π bonds on the surface and follow the universal $\sigma + \pi$ double delocalization bonding pattern.⁵⁻¹⁰ Sea-shell-like C_2 $B_{28}^{-/0}$ and C_s B_{29}^- were later observed as minor isomers of the monoanions in PES experiments^{11,12} and sea-shell-like C_s B_{29}^+ , C_2 B_{31}^+ , C_2 B_{32} , C_2 B_{34} , C_2 B_{35}^+ , and cage-like C_s B_{39}^+ have also been predicted recently in theory.¹³⁻¹⁶

Transition-metal (TM)-doping induces dramatic structural changes and leads to earlier planar → tubular → spherical structural transitions in boron clusters due to effective TM-B coordination interactions, as evidenced by the experimentally observed pyramidal C_{8v} $Ta\oplus B_8^-$,¹⁷ C_s $Ta\oplus B_6^-$,¹⁸ and perfect planar D_{10h} $Ta\oplus B_{10}^-$ which has the highest coordination number of CN = 10 in planar species¹⁹ and the theoretically predicted half-sandwich TaB_{12}^- .²⁰ A family of transition-metal-centred double-ring tubular boron complexes in staggered motifs were recently observed in experiments, including D_{8d} $Co\oplus B_{16}^-$,²¹ D_{9d} $Rh\oplus B_{18}^-$,²² and C_s $Ta\oplus B_{20}^-$,²³ with C_s $Ta\oplus B_{20}^-$ ($B_2-[Ta\oplus B_{18}]^-$) behaving like a tubular molecular rotor at 900 K facilitated by prototypical multicentre fluxional bonds (FBs) which form and break constantly.²⁴ A series of perfect di-metal-doped inverse sandwich complexes have also been observed in PE experiments, including D_{6h} $Ta_2B_6^{-/0}$,²⁵ D_{8h} La_2B_8 , and D_{8h} Pr_2B_8 .²⁶ Our group recently reported the smallest core-shell-like tubular molecular rotor C_{2h} $La_2[B_2\oplus B_{18}]$ which contains a B_2 core rotating inside a B_{18} tube almost freely.²⁷ The first experimentally observed tri-metal-doped inverse triple-decker C_{2v} $La_3B_{14}^-$ exhibits a tilted $La-B_8-La-B_8-La$ triple-decker structure with two conjoined η^8 - B_8 rings sharing a B-B unit on one edge.²⁸ With increasing cluster size, the first perfect spherical trihedral metallo-borospherenes D_{3h} $Ln_3B_{18}^-$ ($Ln = La, Tb$) were very recently discovered in a joint PE experimental and first-principles theory investigation.²⁹ These metallo-borospherenes represent a new class of unusual geometry with tunable magnetic or catalytic properties, with three deca-coordinate Ln centres as integral parts of the cage surface coordinated in three equivalent η^{10} - B_{10} decagons which share two eclipsed

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triangular B_6 units interconnected by three B_2 units. Metallo-borospherenes were previously proposed in theory for MB_{40} ($M = Be, Mg$) and Ni_nB_{40} ($n = 1-4$) which contain hepta-coordinate metal centres on the cage surface of B_{40} , with the corresponding metallo-borophenes with hepta-coordinate metal centres as precursors.^{30,31} However, it remains unknown at current stage what is the possible smallest size of metallo-borospherenes and if such coordination-stabilized metallo-borospherenes can be characterized in experiments.

Based on extensive first-principles theory calculations, we predict in this work the possibility of the perfect di-TM-doped inverse sandwich $D_{7h} Ta_2B_7^+$ (**1**) and $D_{8h} Ta_2B_8$ (**2**) and, more importantly, the smallest tri-TM-doped cage-like metallo-borophene $D_{3h} Ta_3B_{12}^-$ (**3**) which contains three octa-coordinate Ta centres as integral parts of the cage surface coordinated in three equivalent η^8 - B_8 octagons that share two equilateral B_3 triangles interconnected by three B_2 units. Detailed analyses indicate that both $Ta_2B_7^+$ (**1**) and Ta_2B_8 (**2**) are $\sigma + \pi$ dually aromatic in nature, while $Ta_3B_{12}^-$ (**3**) is the first transition-metal-doped boron complex reported to date with $\sigma + \pi + \delta$ triple aromaticity.

2. Theoretical procedure

Extensive global minimum (GM) searches were performed on $Ta_2B_7^+$, Ta_2B_8 , and $Ta_3B_{12}^-$ using the Tsinghua Global Minimum (TGMin) package^{32,33} based on a constraint Basin-Hopping algorithm,³⁴ with more than 2000 singlet or triplet stationary points explored for each cluster at PBE/DZVP level.³⁵ Low-lying isomers were then fully optimized at the PBE0 (36) level with the 6-311+G* basis set³⁷ for B and Stuttgart relativistic small-core pseudopotential^{38,39} for Ta using the Gaussian 16 program suite,⁴⁰ with vibrational frequencies checked to make sure that all isomers reported are true minima. The 15 lowest-lying isomers were subsequently fully re-optimized at both PBE0 and BP86 (ref. 41 and 42) levels with the basis sets of aug-cc-pVTZ for B^{43,44} and Stuttgart+2f1g for Ta. Relative energies for the four lowest-lying isomers were further refined using the more accurate coupled cluster method with triple excitations CCSD(T)⁴⁵⁻⁴⁷ implemented in Gaussian 16 with the same basis sets. Natural bonding orbital (NBO) analyses were performed using the NBO 6.0 program⁴⁸ and detailed bonding analyses carried out utilizing the adaptive natural density partitioning (AdNDP) approach.^{49,50} Molecular dynamics (MD) simulations were performed on $Ta_2B_7^+$ (**1**), Ta_2B_8 (**2**), and $Ta_3B_{12}^-$ (**3**) for 30 ps using the CP2K software suite at different temperatures.⁵¹ The anisotropy of the current-induced density (ACID) analysis⁵² was realized using the ACID code, with the ring-current maps generated using POV-Ray 3.7.⁵³ The iso-chemical shielding surfaces (ICSSs)^{54,55} were generated with the Multiwfn 3.7 code.⁵⁶ The UV-vis and PES spectra were simulated using the time-dependent TD-DFT-PBE0 approach.⁵⁷

3. Results and discussions

3.1 Structures and stabilities

We start from $Ta_2B_7^+$, the smallest di-metal-doped Ta-B complex concerned in this work. Extensive GM searches

indicate that inverse sandwich $D_{7h} Ta_2B_7^+$ (**1**, 1A_1) is the deep-lying GM of the monocation which lies 1.58 eV lower than the second lowest-lying isomer $C_{2v} Ta_2B_7^+$ (1A_1) at CCSD(T) level (Fig. S1†), with the large HOMO-LUMO gap of $\Delta E_{gap} = 3.77$ eV at PBE0. The monocation contains a perfect B_7 ring sandwiched between two Ta atoms along the C_7 molecular axis at the two ends (similar to the experimentally observed inverse sandwich $D_{6h} Ta_2B_6$ which has a perfect B_6 ring sandwiched between two Ta atoms²⁴), with the B-B bond length of $r_{B-B} = 1.54$ Å, Ta-B distance of $r_{Ta-B} = 2.28$ Å, and Ta-Ta distance of $r_{Ta-Ta} = 2.86$ Å (which is obviously larger than the proposed Ta-Ta single bond length of 2.46 Å⁵⁸). With one more B atom added in, the perfect inverse sandwich $D_{8h} Ta_2B_8$ (**2**, $^1A_{1g}$) is achieved which is the deep-lying GM of the neutral lying 0.62 eV lower in energy than the second lowest-lying isomer $C_s Ta_2B_8$ ($^1A'$) at CCSD(T) (Fig. S2†). It encompasses a perfect B_8 ring sandwiched between two Ta atoms along the C_8 molecular axis at the two ends, with the HOMO-LUMO gap of $\Delta E_{gap} = 2.15$ eV at PBE0. With a B_8 octagon as the ligand, the Ta-Ta distance in Ta_2B_8 (**2**) is effectively shortened to $r_{Ta-Ta} = 2.52$ Å which is close to the proposed average Ta-Ta single bond length.⁵⁸ The highly stable inverse sandwich $Ta_2B_7^+$ (**1**) and Ta_2B_8 (**2**) may serve as building blocks to form tri-metal-doped Ta-B complexes, similar to the situation in the experimentally observed triple-decker $C_{2v} La_3B_{14}^-$ (**28**) and spherical trihedral $D_{3h} La_3B_{18}^-$.²⁹

Following the direction, we obtained the lowest-lying $C_{2v} Ta_3B_{10}^+$ (1A_1) for $Ta_3B_{10}^+$, $C_s Ta_3B_{11}^-$ ($^1A'$) for $Ta_3B_{11}^-$, and $C_s Ta_3B_{11}^-$ ($^2A'$) for $Ta_3B_{11}^-$ which all possess incomplete spherical-trihedral structures with one edge broken (Fig. S3†). The smallest perfect spherical trihedron was achieved at $D_{3h} Ta_3B_{12}^-$ (**3**, 1A_1) which, as the GM of the monoanion, lies 0.24 eV lower than the second lowest-lying $C_{2v} Ta_3B_{12}^-$ (1A_1) and 1.06 eV lower than the third lowest-lying $C_s Ta_3B_{12}^-$ ($^1A'$) (which both belong to metallo-borospherenes) at the most reliable CCSD(T) level (Fig. S4†). As shown in Fig. 1, $Ta_3B_{12}^-$ (**3**) contains three equivalent octa-coordinate Ta centres as integral parts of the cage surface coordinated in three equivalent η^8 - B_8 rings which share two eclipsed equilateral B_3 triangles at the top and bottom interconnected by three B_2 units on the waist. It has the calculated B-B bond lengths of $r_{B-B} = 1.58-1.74$ Å, Ta-B coordination distances of $r_{Ta-B} = 2.26-2.27$ Å, and elongated Ta-Ta distances

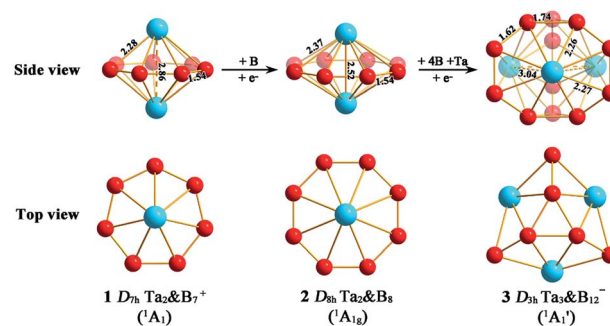


Fig. 1 Optimized structures of $D_{7h} Ta_2B_7^+$ (**1**), $D_{8h} Ta_2B_8$ (**2**), and $D_{3h} Ta_3B_{12}^-$ (**3**) at PBE0//B/aug-cc-pVTZ/Ta/Stuttgart+2f1g level, with bond lengths indicated in Å.



of $r_{\text{Ta-Ta}} = 3.04 \text{ \AA}$, respectively, with the large HOMO(e'')-LOMO(e') gap of $\Delta E_{\text{gap}} = 2.50 \text{ eV}$ at PBE0 which is comparable with the corresponding value of 2.70 eV calculated for the experimentally observed D_{3h} $\text{La}_3\text{B}_{18}^-$ (29) at the same theoretical level. $\text{Ta}_3\text{B}_{12}^-$ (3) can be constructed from Ta_2B_8 (2) by adding one B_4 edge ($\text{B}-\text{B}-\text{B}-\text{B}$) perpendicular to the B_8 ring in the front and one Ta atom coordinated to the newly formed B_8 ring, forming three interconnected Ta_2B_8 inverse sandwiches around the C_3 main molecular axis (Fig. 1). In comparison with D_{3h} $\text{Ln}_3\text{B}_{18}^-$ ($\text{Ln} = \text{La}, \text{Tb}$) which possess a perfect cage-like D_{3h} B_{18} ligand with three equivalent η^{10} - B_{10} decagons sharing two eclipsed B_6 triangles at the top and bottom interconnected by three B_2 units on the waist,²⁹ $\text{Ta}_3\text{B}_{12}^-$ (3) has a perfect cage-like D_{3h} B_{12} ligand with three equivalent η^8 - B_8 octagons sharing two eclipsed B_3 triangles at the top and bottom interconnected by three B_2 units on the waist. $\text{Ta}_3\text{B}_{12}^-$ (3) has the same symmetry (D_{3h}) as $\text{Ln}_3\text{B}_{18}^-$, but smaller in size with six less B atoms in the ligand. Such a structural change can be understood based on the fact that Ta ([Xe]5d³6s²) has a smaller atomic radius (1.46 Å) than that (1.83 Å) of La ([Xe]5d¹6s²),⁵⁹ but possesses two more 5d valence electrons than the latter. The Ta centres appear to have the right atomic radii and electronic configurations to match the D_{3h} B_{12} ligand both electronically and geometrically. $\text{Ta}_3\text{B}_{12}^-$ (3) is therefore the smallest metallo-borospherene reported to date. Perfect spherical trihedral D_{3h} $\text{Nb}_3\text{B}_{12}^-$ and D_{3h} $\text{V}_3\text{B}_{12}^-$ also appear to be true minima of the systems at both PBE0 and BP86 levels (Fig. S5†), with the HOMO-LUMO gaps of $\Delta E_{\text{gap}} = 2.59$ and 2.67 eV at PBE0, respectively.

Extensive MD simulations (Fig. S6†) indicate that Ta_2B_7^+ (1), Ta_2B_8 (2), and $\text{Ta}_3\text{B}_{12}^-$ (3) are dynamically stable at 1200 K, 700 K, and 1200 K, with the small calculated average root-mean-square-deviations of RMSD = 0.11, 0.10, and 0.11 Å and maximum bond length deviations of MAXD = 0.27, 0.24, and 0.29 Å, respectively. No high-lying isomers were observed during the MD simulation. Ta-B complexes with larger HOMO-LUMO energy gaps appear to be dynamically more stable than the ones with narrower energy gaps.

3.2 Natural orbital and bonding analyses

The high stabilities of these high-symmetry complexes originate from their unique electronic structures and bonding patterns. Detailed natural bonding orbital (NBO) analyses show that the Ta centres in Ta_2B_7^+ (1), Ta_2B_8 (2), and $\text{Ta}_3\text{B}_{12}^-$ (3) possess the electronic configurations of Ta [Xe]6s^{0.21}5d^{3.46}, Ta [Xe]6s^{0.22}5d^{3.64}, and Ta [Xe]6s^{0.24}5d^{3.72}, natural atomic charges of $q_{\text{Ta}} = +1.34|e|$, $+1.12|e|$ and $+1.01|e|$, and total Wiberg bond indexes of $\text{WBI}_{\text{Ta}} = 4.85, 4.80$, and 5.07, respectively, indicating that each Ta centre in these complexes donates its 6s² electrons almost completely to the B_n ligand ($n = 7, 8, 12$), while, in return, accepts roughly one electron in its partially filled 5d orbitals from the B_n ligands *via* p → d back-donations. The formations of effective Ta-B coordination interactions in 1–3 are strongly supported by the calculated Ta-B bond orders of $\text{WBI}_{\text{Ta-B}} = 0.59, 0.40$, and 0.50–0.53 (Ta-B interactions within TaB_7 or TaB_8 pyramids) which appear to be systematically higher than the typical Cr-C coordination bond orders of

$\text{WBI}_{\text{Cr-C}} = 0.34$ in D_{6h} $\text{Cr}(\text{C}_6\text{H}_6)_2$. As mentioned above, a Ta centre in $\text{Ta}_3\text{B}_{12}^-$ (3) has the total bond order of $\text{WBI}_{\text{Ta}} = 5.07$ which is also obviously higher than the corresponding value of $\text{WBI}_{\text{Cr}} = 4.12$ calculated for Cr in D_{6h} $\text{Cr}(\text{C}_6\text{H}_6)_2$.

Detailed AdNDP bonding analyses unveil both the localized and delocalized bonds of the concerned systems. As shown in Fig. 2a, Ta_2B_8 (2) possesses 8 2c-2e B-B σ bonds on the periphery of the η^8 - B_8 ligand with the occupation numbers of ON = 1.82|e| and 1 2c-2e Ta-Ta σ bond between the two Ta centres with ON = 2.00|e| in the first row. The remaining 16 valence electrons form 8 totally delocalized 10c-2e coordination bonds between the η^8 - B_8 ligand and two Ta coordination centres with ON = 2.00|e|, including 3 10c-2e σ bonds in the second row, 3 10c-2e π bonds in the third row, and 2 10c-2e (p-d) δ bonding interactions in the fourth row. Both the

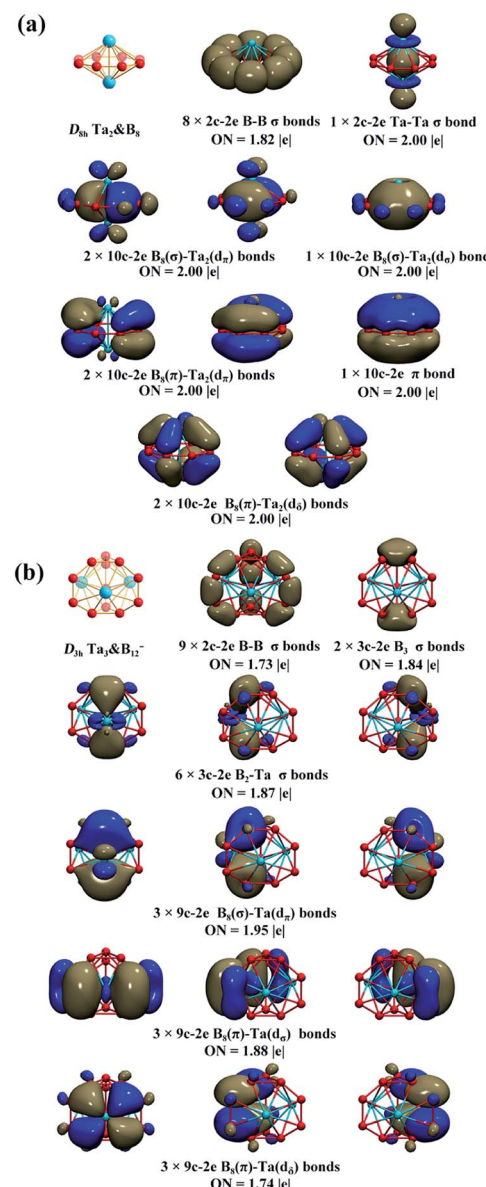


Fig. 2 AdNDP bonding patterns of (a) D_{8h} Ta_2B_8 (2) and (b) D_{3h} $\text{Ta}_3\text{B}_{12}^-$ (3), with the occupation numbers (ON) indicated.



delocalized 6e σ -system and delocalized 6e π -system match the $4n + 2$ aromatic rule ($n = 1$) and together they render $\sigma + \pi$ dual aromaticity to Ta_2B_8 (2). Similarly, both the experimentally observed D_{6h} Ta_2B_6 (ref. 25) and D_{7h} Ta_2B_7^+ (1) are $\sigma + \pi$ doubly aromatic in nature, as clearly indicated in their canonical molecular orbitals (CMOs) depicted in Fig. S7.[†]

The AdNDP bonding pattern of spherical trihedral $\text{Ta}_3\text{B}_{12}^-$ (3) (Fig. 2b) exhibits certain similarity with that of the inverse sandwich Ta_2B_8 (2), but it is more complex and intriguing. $\text{Ta}_3\text{B}_{12}^-$ (3) possesses 9 localized 2c-2e σ B-B bonds in the three B_2 units and between the B_2 units and the six apexes of the two B_3 triangles and 2 equivalent 3c-2e σ bonds on the two B_3 triangles at the top and bottom in the first row and 6 equivalent 3c-2e B_2 -Ta σ bonds between the three Ta atoms and the two B_3 triangles at the top and bottom in the second row. The remaining 18 valence electrons form 9 delocalized 9c-2e bonds evenly distributed on three equivalent C_{2v} TaB_8 octagonal pyramids (which are similar to the experimentally observed C_{8v} TaB_8^- octagonal pyramid¹⁷) around the C_3 main molecular axis, including 3 equivalent 9c-2e $\text{B}_8(\sigma)$ -Ta(d_{π}) bonds in the third row, 3 equivalent 9c-2e $\text{B}_8(\pi)$ -Ta(d_{σ}) bonds in the fourth row, and 3 equivalent $\text{B}_8(\pi)$ -Ta(d_{δ}) bonds in the fifth row. Each TaB_8 octagonal pyramid thus possesses 1 9c-2e $\text{B}_8(\sigma)$ -Ta(d_{π}) bond, 1 9c-2e $\text{B}_8(\pi)$ -Ta(d_{σ}) bond, and 1 $\text{B}_8(\pi)$ -Ta(d_{δ}) bond. Overall, the $\text{Ta}_3\text{B}_{12}^-$ (3) spherical trihedron has 3 equivalent ($d-p$) σ bonds, 3 equivalent ($d-p$) π bonds, and 3 equivalent ($d-p$) δ bonds evenly distributed on three equivalent TaB_8 octagonal pyramids around the C_3 molecular axis, forming one delocalized 6e σ -system, one delocalized 6e π -system, and one delocalized 6e δ -system on the cage surface each matching the $4n + 2$ rule ($n = 1$) independently. Such a delocalized bonding pattern renders $\sigma + \pi + \delta$ triple aromaticity to the smallest metallo-borospherene of $\text{Ta}_3\text{B}_{12}^-$ (3) which is a highly stable D_{3h} spherical trihedron with 18 delocalized electrons evenly distributed on the cage surface.⁶⁰

The aromatic nature of Ta_2B_7^+ (1), Ta_2B_8 (2), and $\text{Ta}_3\text{B}_{12}^-$ (3) is further evidenced by their calculated nucleus-independent chemical shift (NICS) values of NICS = -110.5 , -101.9 , and -86.6 ppm at the geometrical centres, respectively. Based on the calculated NICS-ZZ components, Fig. 3 plots their iso-chemical-shielding surfaces^{54,55} with Z-axis parallel to the designated C_7 , C_8 , or C_2 molecular axes to illuminate the chemical shielding around the TaB_7 or TaB_8 pyramids in these complexes. Obviously, the space inside the spherical trihedron surrounded by the delocalized Ta-B_n coordination bonds in horizontal direction or within about 1.0 Å above the Ta centres in vertical direction belong to chemical shielding regions (highlighted in yellow) with negative NICS-ZZ values, indicating that the aromatic contribution mainly comes from Ta(5d)-B(2p) coordination interactions between the Ta centres and B_n ligands around them ($n = 7$ or 8), while the chemical de-shielding areas (highlighted in green) with positive NICS values are located outside the B_n ring in horizontal direction. The ICSSs of these complexes appear to be analogous to that of the prototypical aromatic benzene.⁵⁴⁻⁵⁶

Fig. S8[†] depicts the corresponding ring current maps of Ta_2B_7^+ (1), Ta_2B_8 (2), and $\text{Ta}_3\text{B}_{12}^-$ (3). Consistent magnetic

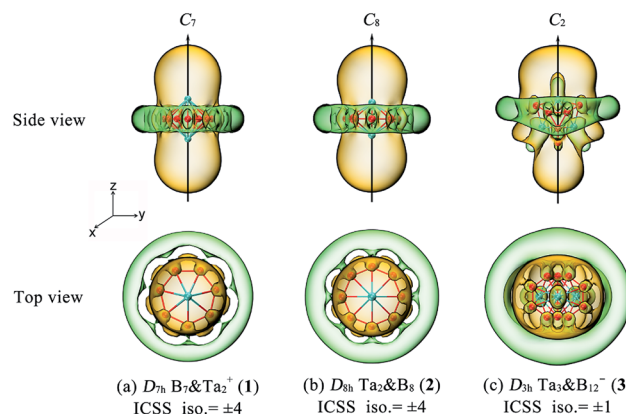


Fig. 3 Calculated iso-chemical shielding surfaces (ICSS) of (a) Ta_2B_7^+ (1), (b) Ta_2B_8 (2), and (c) $\text{Ta}_3\text{B}_{12}^-$ (3), with the corresponding NICS-ZZ components indicated. The C_7 axis of Ta_2B_7^+ (1), C_8 axis of Ta_2B_8 (2), and C_2 axis of $\text{Ta}_3\text{B}_{12}^-$ (3) are designated as z axis in vertical direction to compare the shielding effects around the TaB_n pyramids ($n = 7, 8$). Yellow regions stand for chemical shielding areas, while green areas represent chemical de-shielding areas.

responses occur in 1, 2, and 3 in an external magnetic field in the vertical direction parallel to the assigned molecular axes, irrespective of the electron type, similar to the situations in both benzene^{52,53} and double-ring tubular B_{20} (61) which are known to possess typical planar or tubular aromaticity. Such diatropic ring currents well support the aromatic nature of these Ta-doped boron complexes.

3.3 Simulated IR, Raman, and PE spectra of $\text{Ta}_3\text{B}_{12}^-$ (3)

We computationally simulate the IR, Raman, and UV-vis or PES spectra of Ta_2B_7^+ (1), Ta_2B_8 (2), and $\text{Ta}_3\text{B}_{12}^-$ (3) at PBE0 level to facilitate their future spectral characterizations. PE measurements and infrared photodissociation (IR-PD) spectra in combination with first-principles theory calculations have proven to be powerful approaches in characterizing novel clusters in gas phases.^{3-7,17-28,62} As shown in Fig. 4, the high-symmetry $\text{Ta}_3\text{B}_{12}^-$ (3) exhibits relatively simple IR and Raman spectral patterns, with the major IR active peaks at 507 (e'), 559 (e'), 668 (a₂'), and 881 (e') cm^{-1} (Fig. 4a) and main Raman spectral features at 209 (a₁'), 881 (e'), 998 (e''), and 1201 (a₁') cm^{-1} (Fig. 4b), respectively. The first Raman active symmetrical vibrational mode (a₁') at 209 cm^{-1} corresponds to typical “radial breathing mode” (RBM) of the metallo-borospherene which may be used to characterize the hollow structures of single-walled boron nanoclusters in experiments.⁶³

The calculated PE spectrum of $\text{Ta}_3\text{B}_{12}^-$ (3) in Fig. 4c exhibits major spectral features at 2.95 (2E''), 3.17 (2A₁'), 3.66 (2A₂''), 3.98 (2E'), 4.17 (2A₁''), 5.32 (2E'), 5.68 (2E'), and 5.98 (2A₂''), respectively, corresponding to vertical electronic transitions from the ground state of the anion (¹A₁') to the excited states of the neutral at the ground-state geometry of the anion. The open-shell neutral Ta_3B_{12} has a slightly distorted C_{2v} Ta_3B_{12} (2A₂) ground-state structure due to Jahn-Teller effect, with the calculated electron affinity of EA = 2.87 eV. The simulated IR, Raman, and UV-vis spectra of Ta_2B_7^+ (1) and Ta_2B_8 (2) are depicted in Fig. S9.[†]



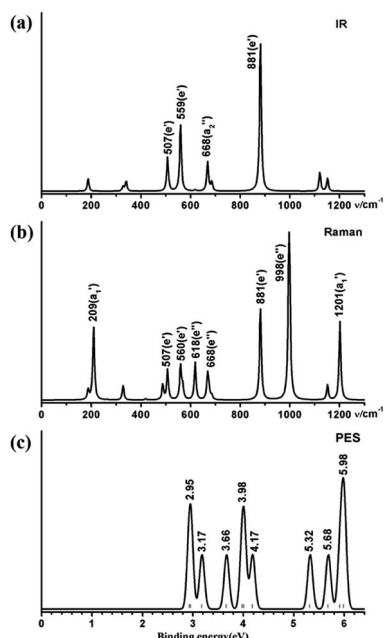


Fig. 4 Simulated (a) IR, (b) Raman, and (c) PES spectra of D_{3h} $Ta_3B_{12}^-$ (3) at PBE0//B/aug-cc-pVTZ/Ta/Stuttgart+2f1g level.

4. Conclusions

Extensive first-principles theory calculations performed in this work indicate a structural transition from perfect inverse sandwich D_{7h} $Ta_2B_7^+$ (1) and D_{8h} Ta_2B_8 (2) to the smallest metallo-borospherene D_{3h} $Ta_3B_{12}^-$ (3). As the first transition-metal-doped boron complex reported to date with $\sigma + \pi + \delta$ triple aromaticity, $Ta_3B_{12}^-$ (3) possesses three octa-coordinate Ta centres as integral parts of the cage surface coordinated in three equivalent η^8 -B₈ octagons. The results obtained in this work suggest that a large family of coordination-stabilized multi-TM-doped TM_nB_n metallo-borospherenes with tunable magnetic and electronic properties may exist in experiments in which the transition metal dopants and B_n ligands match both geometrically and electronically.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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