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Tris((4-BMes₂)phenyl)methanide: a carbanion with a delocalised triple quinoidal structure

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Herein, we report that incorporation of strongly electron-accepting BMes₂ (Mes = 2,4,6-trimethylphenyl) groups at the three *para*-positions of a trigonal [CPh₃][−] framework leads to extensive delocalisation of the negative charge over the three aryl branches and onto the boron centres. Single-crystal structure analysis reveals that all three branches of **CB-Mes-1** exhibit short C(center)–C(phenylene) and C(phenylene)–B bonds, and significant C–C bond length alternation within the central phenylene ring, indicative of a unique triple quinoidal structure in a single system. The HOMO of **CB-Mes-1** is delocalised over the whole molecule, further indicating its quinoidal structure. Upon deprotonation of the neutral HC((C₆H₄)-4-B(Mes)₂)₃ precursor (**CB-Mes**), the resulting **CB-Mes-1** shows a 439 nm (18 400 cm^{−1}) red shift in its absorption from 316 nm in the UV to 755 nm in the NIR. Compared with trigonal N-centred analogues and [CPh₃][−], **CB-Mes-1** displays a significantly raised HOMO energy level, accompanied by a dramatic red shift of the maximum absorption band with a large oscillator strength.

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Introduction

Unveiling electron delocalisation is fundamental to understanding the structure, reactivity, and properties of organic compounds.^{1–13} Trimethylenemethane (TMM), first recognised by Moffitt and Coulson in 1948, is the most typical representative of trigonal systems.¹⁴ It features high symmetry and reactivity, along with a triplet open-shell diradical ground state.¹⁵ Thus, TMM is a representative non-Kekulé system characterised by the presence of at least two atoms not involved in π -bonding, resulting in four unpaired π electrons that are localised rather than delocalised over the entire molecule (Fig. 1a).¹⁶ Shiotani *et al.* reported that the TMM radical cation shows an apparent *D*_{3h} structure. Nevertheless, this is a result of intramolecular dynamics among the three energetically equivalent *C*_{2v} structures, and its unpaired electron primarily resides in the π orbitals of the two equivalent terminal carbon atoms.¹⁷ This finding highlights the fact that high molecular symmetry does not necessarily ensure uniform electron delocalisation.

Compared to TMM, the trigonal triphenylmethyl [CPh₃] cation, radical, and anion systems (**A–C**, respectively, Fig. 1b) are more stable owing to the resonance stabilisation arising from charge delocalisation over the phenyl rings, and offer more peripheral sites for functionalisation.^{18,19} In the unsubstituted CPh₃ systems, the distances from the central carbon atom to its three neighbouring carbon atoms in these molecules exhibit appreciable double-bond character.^{20–22} For [CPh₃][−] (**C**), as it interacts with metal cations in most cases, the central carbon atom adopts sp² hybridisation and exhibits delocalisation of the negative charge over the phenyl rings.^{23–25} However, the “free” [CPh₃][−] ([CPh₃][−][Me₄N⁺]) exhibits a distinct geometry in the solid state; one phenyl ring is nearly coplanar with the central carbon framework (11.5°), one falls within the expected range (26.1°), and the third is much more twisted with respect to the central plane (61.1°). As a result, the co-planar ring displays a large extent of charge delocalisation, whereas charge delocalisation is nearly shut off for the highly twisted phenyl ring.²⁶ In terms of the functionalised trigonal [CPh₃][−] trityl system, three electron-donating dimethylamino groups at the *para*-positions of the phenyl rings in **A** give rise to a triply quinoidal structure, in which the C–N bond lengths are 1.364 Å (**D**).²⁷ The central carbon atom of CPh₃ systems can be replaced by a boron atom and the aryl rings modified with electron donors. For example, in the trigonal carbazolyl-substituted tri-arylborane **E**, the ground-state dipole moment of **E** is 1.7 D, and it exhibits a symmetry lower than *D*₃ in the ground state, as two branches of the molecule contribute only marginally to the overall dipole moment.²⁸ This demonstrates that the electron-

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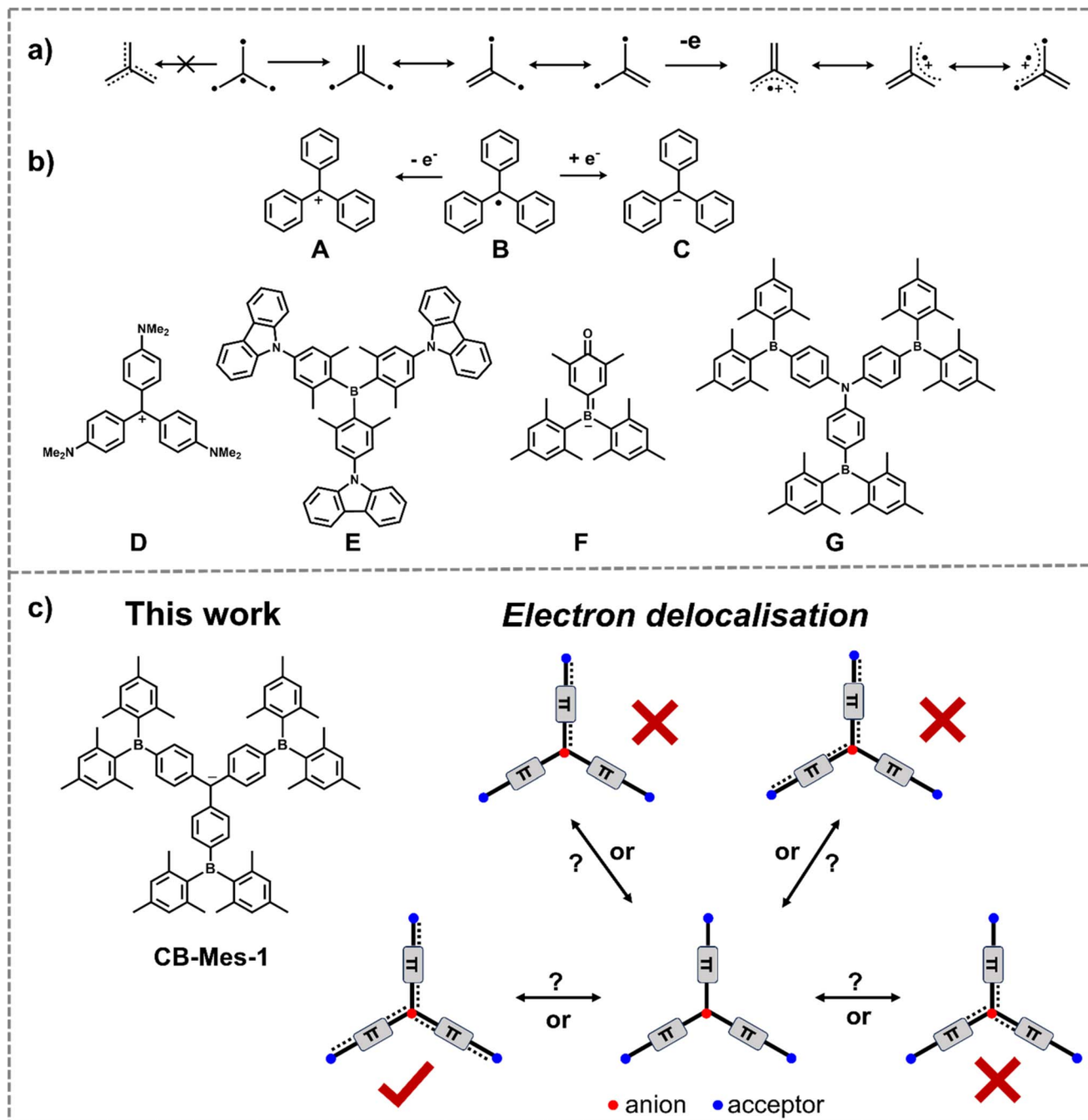
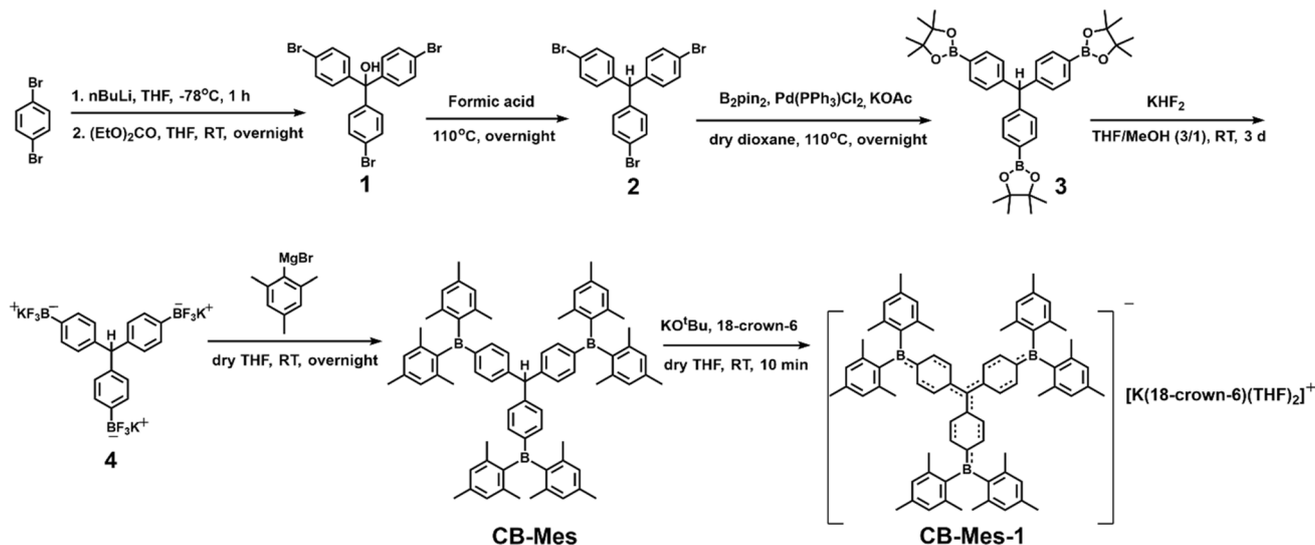


Fig. 1 (a) The Kekulé structures of TMM, and delocalisation pattern of the TMM radical cation (right); (b) [CPh₃] systems (A–D), boron-centred systems (E, F), and nitrogen-centred system (G); (c) the electron delocalisation profile of CB-Mes-1.

donating influence of the carbazoyl substituents does not distribute evenly over the three aryl moieties in this case. The strong electron-donating O⁻ group at the *para*-position of the phenyl ring attached to the boron centre in F, leads to a quinoidal structure, as observed in a few other cases.²⁹ Likewise, the carbanion [Mes₂BCH₂]⁻ shows B=C double bond character (Mes = 2,4,6-trimethylphenyl).²² The NPh₃ core is isoelectronic with [CPh₃]⁻ and, with three BAR₂ (Ar = -Mes (G), ⁻Fmes, and -C₆F₅) groups as strong acceptors, those compounds exhibit delocalised HOMOs.^{30,31}

However, no study has clarified how the central negative charge in [CPh₃]⁻ responds to a simultaneous reduction in the electron density of all three aryl substituents (Fig. 1c). Herein, we installed three strongly π-accepting BMes₂ groups^{32–37} at the *para*-positions of the phenyl rings in the trigonal [CPh₃]⁻ framework *via* synthesis of the neutral compound HC[1,4-C₆H₄B(Mes₂)₃]₃, **CB-Mes**, and, following deprotonation, its corresponding carbanion **CB-Mes-1** (Fig. 1c). Structural and electronic properties were elucidated by single-crystal X-ray diffraction, photophysical measurements, and DFT



Scheme 1 Synthetic routes to **CB-Mes** and **CB-Mes-1**.

calculations. The results demonstrate that the p_z -electrons on the central carbon atom are fully delocalised over the entire molecule, giving rise to a triple quinoidal resonance structure.

Results and discussion

The synthetic routes to **CB-Mes** and **CB-Mes-1** are shown in Scheme 1. Compounds **1–3** were synthesised according to literature procedures.^{38,39} Compound **3** was transformed into its potassium trifluoroborate salt **4** in 94% yield using KHF_2 under an ambient atmosphere in a THF/methanol mixture. It was then reacted with the Grignard reagent MesMgBr to give the neutral **CB-Mes** compound in 14% yield. Owing to the strong electron accepting ability of the BMe_2 moieties, the central carbon atom

was easily deprotonated by treatment with KO^tBu in THF, affording the corresponding carbanion (**CB-Mes-1**). This process was monitored by *in situ* ^1H NMR spectroscopy (Fig. S1). The ^1H NMR spectra of **CB-Mes** and **CB-Mes-1** reveal two doublets and one singlet in the aromatic region with an intensity ratio of 1 : 1 : 2, consistent with C_3 symmetry in solution. Interestingly, all proton signals of **CB-Mes-1** shift downfield compared to those of **CB-Mes**. The upfield shift of the ^{11}B NMR signal of **CB-Mes-1** ($\delta = 66.1$ ppm) relative to that of **CB-Mes** ($\delta = 73.3$ ppm) reveals an increase in the electron density at the three boron centres, indicating the possibility of a triple quinoidal resonance structure.

Single crystals suitable for X-ray diffraction analysis were obtained by slow evaporation of a THF solution of **CB-Mes** and

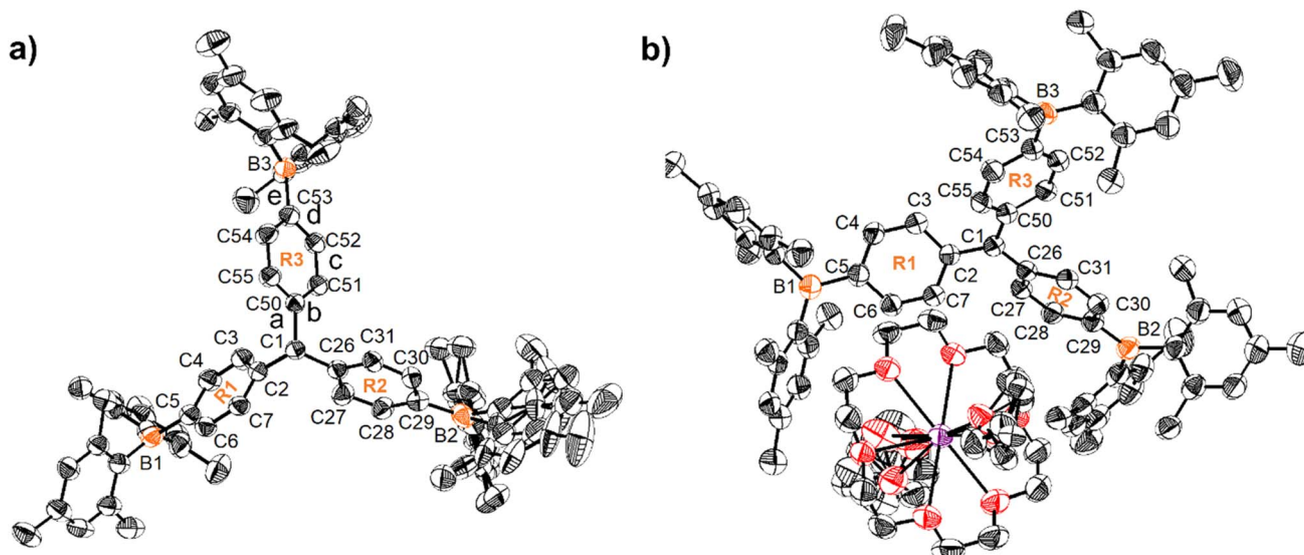


Fig. 2 Molecular structures of (a) **CB-Mes** and (b) **CB-Mes-1**, determined by single-crystal X-ray diffraction with ellipsoids drawn at 50% probability. H atoms and uncoordinated THF molecules are omitted for clarity.



Table 1 Bond lengths [Å], angles [°], dihedral angles [°], and torsion angles between phenylene rings R1, R2, and R3 [°] from single-crystal diffraction data and from DFT calculations on model compounds (calc.)^a

Compound	CB-Mes	CB-Mes-1
C1–C2	1.518(5)	1.446(4)
C1–C26	1.527(5)	1.449(4)
C1–C50	1.521(4)	1.463(4)
Mean bond length a	1.522(3)	1.453(2)
C–C (calc.)	1.531	1.449
C2–C3	1.395(5)	1.431(4)
C26–C27	1.380(5)	1.409(4)
C50–C51	1.397(5)	1.417(4)
C7–C2	1.395(5)	1.411(4)
C31–C26	1.400(5)	1.423(4)
C55–C50	1.386(5)	1.409(4)
Mean bond length b	1.392(2)	1.417(2)
C–C (calc.)	1.402	1.420
C3–C4	1.377(6)	1.370(4)
C27–C28	1.397(6)	1.376(4)
C51–C52	1.377(5)	1.370(4)
C6–C7	1.388(5)	1.366(4)
C30–C31	1.375(5)	1.374(4)
C54–C55	1.384(5)	1.374(4)
Mean bond length c	1.383(2)	1.372(2)
C–C (calc.)	1.393	1.386
C4–C5	1.411(5)	1.412(4)
C28–C29	1.379(6)	1.419(4)
C52–C53	1.409(5)	1.402(4)
C5–C6	1.410(5)	1.410(4)
C29–C30	1.413(6)	1.404(5)
C53–C54	1.403(5)	1.414(4)
Mean bond length d	1.404(2)	1.410(2)
C–C (calc.)	1.411	1.413
B1–C5	1.555(6)	1.547(5)
B2–C29	1.544(17) ^b	1.545(5)
B3–C53	1.561(5)	1.542(5)
Mean bond length e	1.553(6)	1.545(3)
B–C (calc.)	1.567	1.535
C7–C2–C1–C26	88.0(4)	27.1(5)
C7–C2–C1–C26 (calc.)	80.4	30.6
C31–C26–C1–C50	72.4(4)	28.4(4)
C31–C26–C1–C50 (calc.)	80.7	30.7
C55–C50–C1–C2	81.8(4)	27.3(5)
C55–C50–C1–C2 (calc.)	81.2	30.5
C2–C1–C26	112.2(3)	120.4(3)
C2–C1–C26 (calc.)	112.9	120.0
C2–C1–C50	111.1(3)	120.3(3)
C2–C1–C50 (calc.)	112.9	120.0
C26–C1–C50	111.5(3)	119.2(3)
C26–C1–C50 (calc.)	112.8	120.0
sum of the C–C1–C angles	334.8(2)	359.9(2)
sum of the C–C1–C angles (calc.)	338.6	360.0
∠(R1,R2)	80.6(1)	52.5(1)
∠(R2,R3)	88.3(1)	50.8(1)
∠(R1,R3)	83.5(1)	45.6(1)

^a Esd's of the mean values were calculated from the esd's of the individual values using the formula $\sqrt{\sum_{i=1}^n \sigma_i^2/n}$. Values obtained from

DFT calculations (calc.) employed model compounds **CB-Mes'** and **CB-Mes-1'** in which the Mes groups were replaced with phenyl groups. Bond types **a**, **b**, **c**, **d**, and **e** are shown in Fig. 2a. ^b Values are given for the major part (68% occupancy) of the disorder.

by diffusion of pentane into a THF solution of **CB-Mes-1**, respectively. Their structures are shown in Fig. 2, and selected bond lengths and angles are given in Table 1. **CB-Mes** and **CB-Mes-1** adopt familiar propeller-like geometries, and crystallise in the monoclinic *Cc* and triclinic *P* $\bar{1}$ space groups, respectively. The torsion angles between the planes of the phenylene rings describing their relative orientations in **CB-Mes** are 80.6(1)°, 88.3(1)°, and 83.5(1)°, which significantly decrease in **CB-Mes-1** to 52.5(1)°, 50.8(1)°, and 45.6(1)° (Table 1). In addition, a much smaller deviation between the central carbon atom (C1) and the plane defined by C2, C26, and C50 was observed in **CB-Mes-1** (0.030(3) Å), compared to **CB-Mes** (0.451(4) Å). The average C(central)–C(phenylene) bond length (**a**) shortens from 1.522(3) Å (**CB-Mes**) to 1.453(2) Å (**CB-Mes-1**), and the sum of the C–C1–C angle increases from 334.8(2)° (**CB-Mes**) to 359.9(2)° (**CB-Mes-1**). These results indicate that the central carbon atom is sp³-hybridised in **CB-Mes**, but sp²-hybridized in **CB-Mes-1**, as reported for unsubstituted [CPh₃][−].³⁸ The average C(phenylene)–B bond appears to be slightly shortened from 1.553(6) Å in **CB-Mes** to 1.545(3) Å in **CB-Mes-1**, although the two values are within 3 esd's of one another. This value is slightly larger than that of linear boron-containing anions because, in **CB-Mes-1**, the electrons are distributed over the three BMes₂ groups.^{29,40} This C(phenylene)–B bond is also longer than those in the one-electron-reduced 2-(BMes₂)pyrene (1.514(6) Å) and the two-electron-reduced 2,7-bis(BMes₂)pyrene (1.510(3) Å), in which the negative charge is predominantly localised on the boron centres, but is comparable to those observed in the one-electron-reduced species of the latter (1.547(6) Å) and in 1,4-bis(BMes₂)phenylene (1.532(4) Å).² Compared to **CB-Mes**, the mean bond lengths **b** and **d** (Table 1; for labelling, see Fig. 2a) in **CB-Mes-1** become longer, whereas the average length of the **c** bonds decreases. Considering the short C(central)–C(phenylene) and C(phenylene)–B bonds and the significant C–C bond length alternation within the three phenyl rings, **CB-Mes-1** could be described as having a triple quinoidal structure. The optimised structures of **CB-Mes** and **CB-Mes-1** obtained from density functional theory (DFT) calculation show similar characteristics to those observed in their single crystal structures (Table 1).

As the NPh₃ core is isoelectronic with [CPh₃][−], we selected model compounds N-(1,4-C₆H₄-BPh₂)₃ (**H**) and N-[1,4-C₆H₄-B(C₆F₅)₂]₃ (**I**)³¹ as representative N-containing analogues (Fig. S4) for comparison. DFT, TD-DFT, and NBO calculations on **H** and **I** were conducted at the same level as for **CB-Mes'** and **CB-Mes-1'**, and the data are summarised in Table S3. The calculated **e** bond length in **CB-Mes-1'** (1.535 Å) is shorter than that in **H** (1.564 Å). In addition, the strong electron-withdrawing effect of the –C₆F₅ groups leads to a shortening of the **e** bond from 1.564 Å in **H** to 1.539 Å in **I**, whereas the C/N_{central}–C_{phenylene} (**a**) bond lengths show only slight changes between **H** and **I**. These results indicate that the electron-donating ability of the central carbanion is significantly stronger than that of the N atom. Natural bond orbital (NBO) calculations reveal the presence of a C(phenylene)-to-B donor-acceptor interaction in **CB-Mes-1'** associated with a second-order energy of 150.7 kcal mol^{−1}. Consistent with these



structural trends, the Wiberg bond index (WBI) values of the **e** bond in **CB-Mes'** (0.908), **H** (0.918), and **I** (0.974) are all smaller than that in **CB-Mes-1'** (1.021). Moreover, the WBI value of the **a** bond (1.867) in **CB-Mes-1'** is almost twice those observed in **CB-Mes'** (0.986), **H** (1.013), and **I** (1.016). Natural population analysis (NPA) calculations on **CB-Mes** reveal that the charge on the boron atom is $0.881 e^-$, which decreases to $0.802 e^-$ upon deprotonation. However, **H** and **I** exhibit boron charges that are $0.072 e^-$ and $0.079 e^-$, respectively, more positive than that of **CB-Mes-1'**. These indicate increased electron densities in the C(phenylene)-B bonds and strong C(phenylene)-B interactions in **CB-Mes-1'**.

The torsion angles between the three phenyl rings in $[\text{CPh}_3]^-$ show significant differences ($62.8(1)^\circ$, $51.9(1)^\circ$, and $43.8(1)^\circ$, Table S2) and, while two of them are fairly similar to that in **CB-Mes-1**, the third one is somewhat larger. Moreover, the bond lengths within the phenyl ring with the smallest torsion angle in $[\text{CPh}_3]^-$ exhibit a pronounced difference relative to the other two rings. These observations indicate that the electrons are not equally distributed among the three branches. In contrast, the three *p*-BMes₂ groups in **CB-Mes-1** exert a remarkable influence on the molecular geometry, enhancing the uniformity of electron delocalisation. Compared to $[\text{CPh}_3]^-$, **CB-Mes-1** exhibits decreased torsion angles and increased C-C bond length alternation in the three phenyl groups, attributed to its triple quinoidal resonance structure. In addition, the C(phenylene)-B bonds in **CB-Mes-1** are significantly shorter than those in linear donor- π -BMes₂ structures, such as (*p*-Me₂N-phenyl) dimesitylborane, illustrating that, although each boron center carries only one-third of the negative charge, the electron-donating ability of the carbanion is stronger than that of NMe₂.⁴¹

The harmonic oscillator model of aromaticity (HOMA) was calculated for **CB-Mes'** and **CB-Mes-1'** based on their DFT-optimised geometries to elucidate variations of aromaticity. The HOMA value of the central phenyl rings in **CB-Mes-1'** (0.82) is smaller than that in **CB-Mes'** (0.96), further verifying its triple quinoidal resonance structure.

Cyclic voltammetry (CV) and differential pulse voltammetry (DPV) measurements in THF (Fig. S5) reveal three reversible reductions at -2.72 , -2.83 , and -2.89 V (vs. Fc/Fc⁺ in THF),

which can be ascribed to the sequential reduction of the three BMes₂ groups in **CB-Mes**. The small redox potential gaps suggest a lack of substantial electron delocalisation in the neutral species. We also attempted to record the CV of **CB-Mes-1**; however, unfortunately, it is highly unstable in common electrolyte solutions such as $[\text{Bu}_4\text{N}]^+[\text{PF}_6]^-$ and $[\text{Bu}_4\text{N}]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$.

The UV-vis absorption spectra of **CB-Mes** and **CB-Mes-1** were measured in dry THF solutions under an argon atmosphere. **CB-Mes** shows two absorption bands in the ultraviolet region, with the maximum at 316 nm ($\epsilon = 5.4 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$), which results from the three BMes₂ groups (Fig. 3a and b).⁴² Its fluorescence spectrum displays a single, broad emission band at 386 nm, with an absolute fluorescence quantum yield (Φ) of 0.23 in THF (Fig. 3c and Table S4). As the solvent polarity increases, the absorption of **CB-Mes** shows only slight changes, whereas the emission spectra exhibit a pronounced solvent-dependent behaviour, with the maximum shifting from 361 nm (hexane) to 399 nm (acetonitrile), indicating a charge transfer process in **CB-Mes**. For the anion **CB-Mes-1**, a broad absorption band is observed at 755 nm ($\epsilon = \text{ca. } 10^4 \text{ M}^{-1} \text{ cm}^{-1}$), which is significantly red-shifted compared to **CB-Mes**. This is attributed to its triple quinoidal resonance structure. Compared with calculated values for N-centred model compounds **H** and **I**, **CB-Mes-1** displays a significantly raised HOMO energy level (-2.89 eV vs. -7.19 eV for **H** and -8.21 eV for **I**; see Table S3), accompanied by a dramatic red shift of the maximum absorption band with a large oscillator strength.

Density functional theory (DFT) calculations on **CB-Mes'** and **CB-Mes-1'** were performed at the ω B97X-D/6-31G+(d,p) level (Fig. 4). For computational simplicity, the Mes groups in **CB-Mes** and **CB-Mes-1** were replaced with phenyl groups. Due to the *C*₃ symmetry of the optimised structures, the LUMO and LUMO + 1 of both molecules are degenerate and delocalised over the entire molecule. The HOMO of **CB-Mes'** is mainly localised on the three central phenyl rings with a small contribution from the central CH group, whereas in **CB-Mes-1'** there is a large contribution from the central carbon atom and delocalisation through the phenylene rings and onto the three boron atoms as well, further supporting the quinoidal structure. The HOMO of **CB-Mes-1'** is similar to that of $\text{N}[\text{C}_6\text{H}_4\text{B}(\text{Mes})_2]_3$.^{30,31,43} The

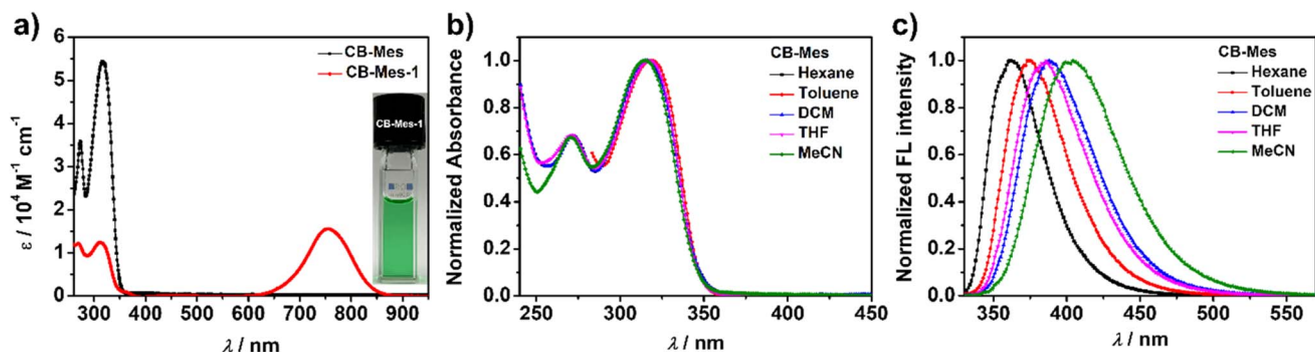


Fig. 3 (a) Absorption spectra of **CB-Mes** and **CB-Mes-1** in THF; (b) normalised absorption and (c) emission spectra of **CB-Mes** in different solvents.



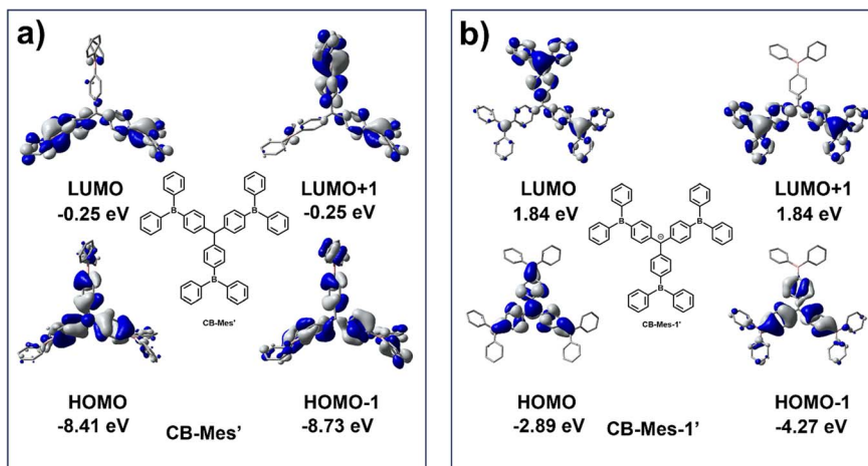


Fig. 4 Selected Kohn–Sham molecular orbitals and energy levels of (a) **CB-Mes'** and (b) **CB-Mes-1'**.

positions of the absorption bands are accurately reproduced by time-dependent density functional theory (TD-DFT) calculations at the B3LYP/6-31+G(d,p) level. **CB-Mes'** (312 nm) and **CB-Mes-1'** (745 nm) exhibit strongly allowed transitions from S_0 to the lowest degenerate levels ($S_{1,2}$), which are attributed to the degenerate $\text{HOMO} \rightarrow \text{LUMO}$ and $\text{HOMO} \rightarrow \text{LUMO} + 1$ transitions (Tables S5 and S6).

Conclusion

In summary, we have demonstrated that installing three strongly electron-accepting BMe_2 substituents at the *para*-positions of a trigonal CPh_3^- framework induces a remarkable structural and electronic evolution upon deprotonation of neutral **CB-Mes**. The resulting anion, **CB-Mes-1**, exhibits nearly similar geometries in its three branches, all of them showing shortened key C–C and C–B bonds and bond length alternation in the central phenyl rings, collectively indicative of a triple quinoidal delocalised structure. DFT calculations and spectroscopic evidence support an extensive redistribution of the negative charge over the entire π -framework, in stark contrast to the localised charge observed in classical CPh_3^- species in the solid state. In addition, this special geometry leads to a significant red shift ($18\,400\text{ cm}^{-1}$) in the absorption of **CB-Mes-1**, formed by deprotonation of **CB-Mes**. These findings not only fill a missing conceptual gap regarding electron distribution in functionalised trigonal carbanions, but also provide a general molecular design principle for creating electronically adaptive anionic π -architectures.

Author contributions

T. B. M., L. J., and Y. Z. contributed to the conception and design of the experiments and wrote the manuscript. Y. Z. synthesized and characterized the compounds and conducted the photophysical measurements and theoretical studies. J. K. performed the single-crystal X-ray diffraction data collection and A. F. refined the structures and analysed the resulting data.

I. K. performed the electrochemistry supervised by H. B. S. Y. provided valuable advice during data analysis and manuscript preparation and T. B. M., L. J., and S. Y. supervised the project. All authors analysed and discussed the results and participated in preparing the final draft of the manuscript.

Conflicts of interest

The authors declare no conflicts.

Data availability

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information: including experimental details, synthesis, NMR spectra, UV-vis spectra, computational details, crystallographic information, and additional characterization data. See DOI: <https://doi.org/10.1039/d5sc10121b>.

CCDC 2504657 (for **CB-Mes**) and 2504659 (for **CB-Mes-1**) contain the supplementary crystallographic data for this paper.^{44a,b}

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