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Quantitative syntheses of permethylated closo-1,10- $R_2C_2B_8Me_8$ (R = H, Me) carboranes. Egg-shaped hydrocarbons on the Frontier between inorganic and organic chemistry†

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Electrophilic methylation of the closo-1,10-R₂C₂B₈H₈ (1) (R = H or Me) dicarbaboranes at higher temperatures or thermal rearrangement of the 1,6-R₂C₂B₈Me₈ (3) compounds at 400–500 °C generated the B-permethylated derivatives closo-1,10-R₂C₂B₈Me₈ (2) in quantitative (>95%) yields. The compounds exhibit extreme air stability as a consequence of a rigid, egg shaped hydrocarbon structures incorporating inner 1,10-C₂B₈ carborane core.

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Introduction

Methods for cage substitution on the cage of closo-1,10-C₂B₈H₁₀ (1a) generally parallel those employed for the larger C₂B₁₀H₁₂ icosahedral carboranes.1 The CH hydrogens in 1a are sufficiently protonic in character to undergo lithiation with butyllithium in ethereal solvents, generating the mono- and dilithio derivatives.2 The C-lithiated species afford the main entry to alkyl, aryl, carboxyl, silyl, and other C-substituted derivatives via treatment with appropriate reagents. For example, exopolyhedral metal complexes,3,4 and the silyl-linked mixed-carboranes⁵ have also been prepared by this route along with numerous C-metallated compounds containing main-group metals. The only B-substitution processes so far reported are, however, the direct reaction between 1a and Cl₂ affording the Bperchloro species 1,10-H₂C₂B₈Cl₈ (ref. 2) and those leading to a series of halo derivatives 1,10-H₂C₂B₈H₇-2-X.6 The successful B-methylation experiments in the 12-vertex carborane series, 7-9 together with those achieved by our group in the B-methylation of $\mathit{closo}\text{-}1,6\text{-}C_2B_8H_{10},^{10}$ $\mathit{arachno}\text{-}6,9\text{-}C_2B_8H_{14},^{11}$ prompted us to extend boron-methylation strategy to the most stable members of the 10-vertex closo series, closo-1,10- $R_2C_2B_8H_{10}$ (1) (where R = H or Me), which have been now relatively easily available.12 In

this article we would like to present electrophilic reactions with methylation agents leading to quantitative permethylation of Bvertexes under the formation of rigid, hydrocarbon-boron structures of egg shape that exhibit outstandingly high stability.

Results and discussion

The electrophilic CH3I/AlCl3 methylation of carborane closo-1,10-H₂C₂B₈H₈ (1a) (Scheme 1) led on heating at 115 °C for 15 h to exclusive formation of the B-permethylated dicarbaborane closo-1,10-H₂C₂B₈Me₈ (2a) in practically quantitative yield (>95%). The scheme also shows that the $CH_3OTf/HOTf$ (Tf = SO₂CH₃) methylation proceeded excellently at 165 °C for 48 h, giving again a quantitative yield of 2a (>95%). It is, however, interesting that the CH3I/AlCl3 methylation (115 °C, 15 h) of closo-1,10-Me₂C₂B₈H₈ (1b) has completely failed, while the CH₃OTf/HOTf methylation of **1b** proceeded smoothly at 175 °C for 48 h, giving again a quantitative yield of the decamethylated closo-1,10-Me₂C₂B₈Me₈ (2b). This finding is in accord with that observed in the comparable 12-vertex closo-1,12-Me₂C₂B₁₀Me₁₀ series (reflux, 20 h, 91% yield).9 As also observed for the latter species, an attempt at methylating the CH1,10 vertexes in 2a via the Li²⁺ salt as in Scheme 1 has failed, too. The difficulty in forcing this reaction to completion must be due to the relative lack of reactivity of the CH vertices present in 2a, which is exacerbated by the steric protection afforded to each CH vertex by the methyl groups of the four surrounding BMe, vertices.

Another straightforward route leading to quantitative formation of permethylated dicarbaboranes 2 consists in thermal isomerisation of *closo*-1,6-R₂C₂B₈Me₈ (3) carboranes (R = H 3a or Me 3b)¹⁰ by heating at 400–500 $^{\circ}$ C in a sealed tube for 2 h. A similar thermal isomerisation principle could be also applied to the substituted derivatives, for example to *closo*-1,6-

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Paper

Scheme 1 Quantitative syntheses of permethylated derivatives of the closo-1,10-R₂C₂B₈H₈ family

H₂C₂B₈Me₇-8-OTf (4a), 10 which underwent cage rearrangement to afford 97% of the closo-1,10-H₂C₂B₈Me₇-2-OTf (5a) upon a similar heating, as shown in Scheme 2.

The structures of derivatives 2a and 5a were determined by an X-ray diffraction study (see Fig. 1 and 2). Both carboranes adopt the expected bicapped Archimedean antiprismatic geometry with two unsubstituted apical CH1,10 vertexes along with eight or seven BMe groups, respectively. The structure 5a unambiguously confirms the B2-substitution with the O-SO₂-CF₃ group and permethylation in all other B-positions of the cluster. Unfortunately, the crystals of the permethylated 2b were not found suitable for crystallographic studies and the 2a/2b pair was therefore geometry optimized at the MP2/TZVP level (Fig. 3 and 4).

The optimization revealed that the comparable B-B, C-B, and B-Me bonding vectors are very similar to those found crystallographically for 2a and 5a. The computation has also led to a good agreement between theoretical and experimental $\delta(^{11}B)$ values for **2a** and **2b** (max. deviation less than 3 ppm), for individual values see ESI.†

The HF/cc-pVTZ calculations of the electrostatic potential (ESP) surface show that the parent 1a has hydridic hydrogen atoms, which can form dihydrogen bonds,13 the hydridic Bbound hydrogens in 2a and 2b are now replaced by methyl groups of amphiphilic character.14 The hydrogen atoms of the methyl groups have positive ESP surface and the exo-skeletal carbon atoms have negative ESP surface (see Fig. 5). From the viewpoint of electron transmission, Me groups behave as weak H₃C H₃C CH₃ CH₃ CH₃ H₃C B B CH₃ CH

Scheme 2 Quantitative synthesis of the permethylated derivative closo-1,10-R₂C₂B₈Me₇-2-OTf.

electron acceptors, when compared to toluene, xylene, hexamethylbenzene etc. This is in accord with the 1,6-isomers of the same molecular shape and the same also applies to CMe methyl groups; the lower electron density at C1,10 in 2b in relation to 2a (see C1···C10 body diagonals) may perhaps prove this concept. The electron transmission thus follows the established electronegativity concept (C in CH₃ is more electronegative than C (cage)) reflecting the fact that exo-skeletal substituents are bound via classical 2-centre 2-electron bonds. This agrees with the concept elaborated by Viñas et al. on a hexamethylated closo-H₂C₂B₁₀H₄Me₆ system. Conceivably, the whole MMR spectra of et and et are significantly paramagnetically deshielded to high frequencies (et et downfield shifted) with respect to those of parent compounds, for the computed MMR spectra see ESI.

The constitution of all compounds isolated in this study is also in agreement with the results of multinuclear ¹¹B, [¹¹B-¹¹B]-COSY,¹⁷ ¹H, and ¹³C NMR measurements that led to complete assignments of individual cage BMe, BX, CH, and CMe units (for hardcopies of the NMR measurements, see ESI, Fig. S1–S13†). The multinuclear (¹¹B, ¹H and ¹³C) NMR spectra

for all compounds isolated in this work are compared and depicted in Fig. 6, a general feature of the persubstituted compounds being the identity of the ¹¹B and ¹¹B–{¹H}-decoupled NMR spectra that exhibit only singlet resonances.

The ¹¹B NMR spectra of the D_{4h} -symmetry derivatives ${\bf 2a}$ and ${\bf 2b}$ (Fig. 3, S7 and S10†), exhibit only one singlet due to equivalency of all B-positions; the corresponding ¹H spectra (Fig. 6, S8 and S10†) consist of two sharp 1:12 or 1:4 singlets for ${\bf 2a}$ and ${\bf 2b}$, respectively, attributed to CH1,10 (or Me1,10) and BMe resonances. The ¹³C–{¹H} NMR spectra of carboranes of structure 2 (Fig. 6, S9 and S11†) show one lower-field C1,10 singlet along with a very broad (J_{C-B} coupling) high-field BMe resonance of relative areas 1:4, whereby that of ${\bf 2b}$ shows an additional C-Me signal. For NMR spectra of the key permethylated derivatives ${\bf 2a}$ and ${\bf 2b}$, see Fig. 7.

On the other hand, the 11 B NMR spectrum of the C_s -symmetry compound 5a (Fig. 3 and S12†) displays 3:2:2:1 patterns of singlets with one coincidental overlap. The 1 H spectrum of 5a (Fig. 6 and S13†) shows two different CH signals and four well resolved 2:2:2:1 patterns of BMe resonances in the high-field; the 13 C- 1 H 1 NMR spectrum contains two different low-field CH resonances together with a typical low-

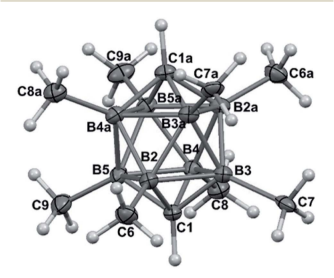


Fig. 1 ORTEP (30% probability level) representation of the molecular structure of closo-1,10-H₂C₂B₈Me₈ (2a). Selected bond lengths (Å): C1-B2 1.600(4), C1-B4 1.595(4), B3-B2 1.870(4), B2-B3a 1.821(4), mean B-CH₃ 1.639(4); angles (°): B5-B2-B3 89.99(18), C1-B2-B3a 107.2(2), B2-B3a-B3 61.70(17), B5-C1-B2 71.39(17).

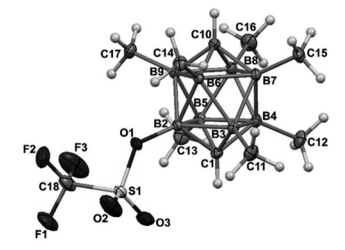


Fig. 2 ORTEP (30% probability level) representation of the molecular structure of closo-1,10-H₂C₂B₈Me₇-2-OTf (5a). Selected bond lengths (Å): C1-B2 1.584(2), C1-B4 1.598(3), B2-B3 1.851(3), B2-B6 1.806(3), mean B-CH₃ 1.578(3), B2-O 1.453(2), mean C10-B 1.601(3), angles (°): B5-B2-B3 91.96(11), C1-B2-B6 109.72(13), B2-B6-B3 61.03(10), B2-C1-B5 70.64(12).

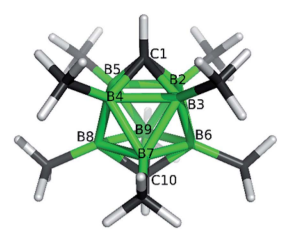


Fig. 3 Molecular model for closo-1,10-H₂C₂B₈Me₈ (2a) as obtained from the MP2/TZVP optimization. The selected internal coordinates are (distances in Å, angles in °): C1–B2 1.602, C1···C10 3.329, B2–B3 1.866, B2–B6 1.818, B6–B7 1.866.

field CF₃ quartet, apart from a broad, high-field BMe signal of intensity 7 (see Fig. S13†).

The 19 F NMR spectrum of **5a** shows one singlet resonance at -76.0 ppm, as expected.

Experimental

Materials and methods

All the reactions were carried out under argon atmosphere. Dichloromethane and hexane were dried over CaH₂ and freshly distilled before use. Other conventional chemicals were of reagent or analytical grade and were used as purchased. NMR spectroscopy was performed at 400 and 600 Mz (for ¹H), inclusive of standard [¹¹B-¹¹B]-COSY¹⁷ experiments (all theoretical cross-peaks were observed) leading to complete assignments of

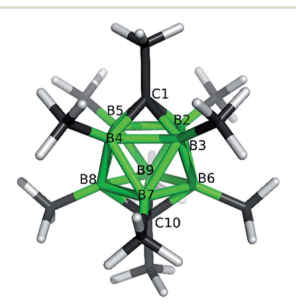


Fig. 4 Molecular model for closo-1,10-Me₂C₂B₈Me₈ (**2b**) as obtained from the MP2/TZVP optimization. The selected internal coordinates are (distances in Å, angles in °): C1–B2 1.607, C1···C10 3.367, B2–B3 1.856. B2–B6 1.812. B6–B7 1.858.

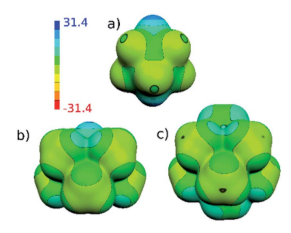


Fig. 5 Computed (HF/cc-pVTZ) electrostatic potential (ESP) surface for (a) **1a**, (b) **2a** and (c) **2b**. The color range of the ESP in kcal mol⁻¹.

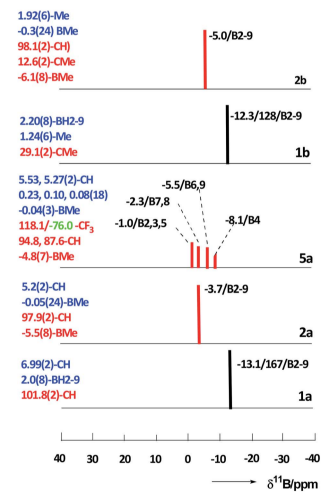


Fig. 6 Stick diagrams comparing the ^{11}B NMR chemical shifts and relative intensities for closo compounds 1,10-H₂C₂B₈H₈ (1a), 1,10-H₂C₂B₈Me₈ (2a), 1,10-H₂C₂B₈Me₇-2-OTf (5a), 1,10-Me₂C₂B₈H₈ (1b), and 1,10-Me₂C₂B₈Me₈ (2b) (red sticks = singlets, black = doublets). The data for individual cluster positions are ordered as: $\delta(^{11}B)/^{1}J_{\rm BH}$ (where applicable)/assignment. ^{1}H (blue text) and $[^{13}C,^{19}F]$ (red and green text) NMR resonances (δ in ppm/TMS) are summarized on the left side of each diagram with relative intensities other than 1 in parentheses.

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B2-9 -3.7**BMe** -0.05 CH1.10 CH1,10 BMe 52 118.0 -0.4B2-9-CH₄ B2-9 -0.23-50 C1,10-Me 12.6 C1.10-CH3 1.92 C1,10

Fig. 7 192.6 MHz 11 B, 600 MHz 1 H, and 150.9 MHz 13 C NMR spectra of closo-1,10-H $_2$ C $_2$ B $_8$ Me $_8$ (2a) (top) and closo-1,10-Me $_2$ C $_2$ B $_8$ Me $_8$ (2b) (bottom).

all resonances to individual cage B-vertexes. Chemical shifts are given in ppm to high-frequency (low field) of $\Xi=32.083971$ MHz (nominally $F_3B\cdot OEt_2$ in $CDCl_3$) for ^{11}B (quoted $\pm~0.5$ ppm), $\Xi=25.144$ MHz for ^{13}C (quoted $\pm~0.5$ ppm), and $\Xi=100$ MHz for ^{1}H (quoted $\pm~0.05$ ppm), Ξ is defined as in ref. 18 and the solvent resonances were used as internal secondary standards. The starting carboranes of structures 1 and 3 were prepared according to the reported methods. 10,12

Dimethylation of closo-1,10-C₂B₈H₁₀ (1a) on carbon vertices

A solution of 1a (120 mg, 1 mmol) in dry $\rm Et_2O$ (ca. 10–20 ml) was cooled to $-78~^{\circ}C$ and then treated dropwise with 2.5 M LiBu (solution in hexane) (1 ml, 2.5 mmol) under stirring. The off-white slurry of the $\rm Li^+$ salt was stirred for additional 1 h prior to addition of methyl triflate (MeOTf, m.w. 164.1) drop by drop, (410 mg, 2.5 mmol) under cooling down in an dry-ice bath. The mixture was then left stirring for additional 2 h at room temperature. After adding 5 % hydrochloric acid (10 ml) under repeated cooling and shaking, the $\rm Et_2O$ layer was separated and evaporated to provide a crude product $\it closo$ -1,10-Me $_2\rm C_2B_8H_8$ (1b) in practically a quantitative yield, as assessed by NMR spectroscopy (see Fig. S4–S6†).

$closo-1,10-R_2C_2B_8Me_8$ (2) (where R = H 2a or Me 2b)

- (a) Methylation with MeOTf. A solution of carboranes 1a or 1b (reaction scale ~ 1.5 mmol) in neat MeOTf (5 ml) was treated with three drops of HOTf and the mixture was heated for 48 hours at 165 °C in a thick-walled reaction vessel equipped with a Teflon screw cap. The volatiles were evaporated and the residue extracted with hexane, the extract was filtered through a plug of silica gel, evaporated, and then sublimed at 150–180 °C (bath) to isolate white crystals of carboranes 2a and 2b in practically quantitative yields (97–98%). For NMR spectra, see Fig. 6 and S7–S12.†
- (b) Methylation with MeI. A solution of carboranes 1a or 1b (reaction scale ~ 1.0 mmol) in neat MeI (10 ml) was treated with anhydrous $AlCl_3$ (ca. 14 mg, 0.14 mmol) and heated at 115 °C (oil bath) for 15 hours in a thick-walled thick-walled reaction vessel equipped with a Teflon screw cap. The volatiles were evaporated the residue subjected to extraction with pentane, filtered through a plug of silica gel, and vacuum sublimed as sub (a) to isolate 2a (yield 97%). Compound 2b has not been formed at all and 1b was recovered from the pentane extract in $\sim 80\%$ yield.

(c) Thermal rearrangement of *closo*-1,6-R₂C₂B₈Me₈ (3) compounds (where R = H or Me). Compounds 3a or 3b (reaction scale ~ 0.5 mmol) were heated in a sealed tube at 400–500 °C (heating gun) for 2 hours. Sublimation as in preceding experiments led to essentially quantitative isolation of carboranes 2a and 2b. For 2a: MS (ESI $^-$): m/z (max.) calcd 232.28, found 232.25; for C₁₀H₂₆B₈ (m.w. 232.80) calcd 51.59 %C, 11.26 %H; found 51.21 %C, 11.15 %H. For 2b MS (ESI $^-$): m/z (max.) calcd 260.31, found 260.31; for C₁₂H₃₀B₈ (m.w. 260.85) calcd 55.25 %C, 11.59 %H; found 54.80 %C, 11.19 %H. For NMR spectra, see Fig. 6 and S7–S12.†

Thermal rearrangement of closo-1,6-R₂C₂B₈Me₇-8-OTf (4a)

Compound **4a** (74 mg, 0.2 mmol) was heated in a sealed ampoule at 400–500 °C (heating gun) for 2 hours. The resulting product was identified as closo-1,10-R₂C₂B₈Me₇-2-OTf (**5a**) and isolated on sublimation (bath temperature ~ 150 °C) in essentially quantitative yield. For **5a** MS (ESI⁻): m/z (max.) calcd 366.20, found 366.19; for C₁₀H₂₃B₈O₃SF₃ (m.w. 366.84) calcd 32.74 %C, 6.32 %H; found 32.28 %C, 6.15 %H. For NMR spectra, see Fig. 6, S13 and S14.†

Computational details

Magnetic shielding was calculated using the GIAO-MP2 method incorporated into Gaussian09 (ref. 19) utilizing the IGLO-II basis with the MP2/TZVP geometry and frozen core electrons. Electrostatic potentials were computed at the HF/cc-pVDZ level using Gaussian09 and Molekel4.3 (ref. 20) programs. It has recently been shown that this basis set size is sufficient for these purposes.²¹

X-ray crystallography

The X-ray data for the derivatives 2a and 5a (colourless crystals by slow evaporation of a hexane solution) were collected at 150(2) K with a Bruker D8-Venture diffractometer equipped with Mo (Mo/ K_{α} radiation; $\lambda = 0.71073$ Å) microfocus X-ray (IµS) source, photon CMOS detector and Oxford Cryosystems cooling device was used for data collection. The frames were integrated with the Bruker SAINT software package using a narrow-frame algorithm. Data were corrected for absorption effects using the Multi-Scan method (SADABS). Obtained data were treated by XT-version 2014/5 and SHELXL-2014/7 software implemented in APEX3 v2016.5-0 (Bruker AXS) system.22 Hydrogen atoms were mostly localized on a difference Fourier map, however to ensure uniformity of treatment of crystal, all hydrogen were recalculated into idealized positions (riding model) and assigned temperature factors $H_{iso}(H) = 1.2U_{eq}$ (pivot atom) or of $1.5U_{\rm eq}$ (methyl). H atoms in methyl groups were placed with C-H distances of 0.96 while the hydrogen atoms of the C-H in the carborane cage were assigned according to the maxima on the difference Fourier map. $R_{\text{int}} = \sum |F_0|^2 - F_0|^2$ $F_{\text{o,mean}}^2 | / \sum F_{\text{o}}^2$, $S = \left[\sum (w(F_{\text{o}}^2 - F_{\text{c}}^2)^2) / (N_{\text{diffrs}} - N_{\text{params}}) \right]^{1/2}$ for all data, $R(F) = \sum_{i=1}^{\infty} ||F_o|| - ||F_c|||/\sum_{i=1}^{\infty} ||F_o||$ for observed data, $WR(F^2) = [\sum_{i=1}^{\infty} (w(F_o^2 - F_c^2)^2)/(\sum_{i=1}^{\infty} w(F_o^2)^2)]^{1/2}$ for all data. Crystallographic data for structural analysis have been deposited with the Cambridge Crystallographic Data Centre, CCDC no. 1573271 and 1573272 for 2a and 5a, respectively. Crystallographic data for 2a: $C_{10}H_{26}B_8$, M=232.79, orthorhombic, *Pbcn a* = 8.5140(13), *b* = 15.098(3), *c* = 12.670(2) Å, $\beta = 90^{\circ}, Z = 4, V = 1628.6(5) \text{ Å}^3, D_c = 0.949 \text{ g cm}^{-3}, \mu = 0.045$ mm^{-1} , $T_{\text{min}}/T_{\text{max}} = 0.6416/0.7456$; $-10 \le h \le 11$, $-19 \le k \le 11$ 19, $-15 \le l \le 16$; 14 313 reflections measured ($\theta_{\text{max}} = 27.55^{\circ}$), 1875 independent ($R_{\text{int}} = 0.0566$), 136 with $I > 2\sigma(I)$, 91 parameters, S = 1.078, R_1 (obs. data) = 0.0977, wR_2 (all data) = 0.2866; max., min. residual electron density = 0.792, -0.319 $e\mathring{A}^{-3}$. Crystallographic data for 5a: $C_{10}H_{23}B_8F_3O_3S$, M=366.82, monoclinic, $P2_1/c$, a = 13.0608(9), b = 9.8980(6), c =16.0303(9) Å, $\beta = 111.202(2)$ °, Z = 4, V = 1932.1(2) Å³, $D_c =$ 1.261 g cm⁻³, $\mu = 0.200 \text{ mm}^{-1}$, $T_{\min}/T_{\max} = 0.6787/0.7456$; $-16 \le h \le 16$, $-12 \le k \le 12$, $-20 \le l \le 19$; 47 313 reflections measured ($\theta_{\text{max}} = 27.54^{\circ}$), 4445 independent ($R_{\text{int}} =$ 0.0300), 3971 with $I > 2\sigma(I)$, 241 parameters, S = 1.049, R_1 (obs. data) = 0.0502, $wR_2(all data) = 0.1431$; max., min. residual electron density = 0.729, -0.628 eÅ^{-3} .

Conclusions

There are not too many reactions in the area of carborane chemistry that proceed quantitatively.1 To these rare cases belong syntheses leading to permethylated derivatives of closo-1,10-R₂C₂B₈H₈ (1) reported this work. It was shown that all Bpositions in structures 1 can be furnished with methyl substituents, via electrophilic reactions with MeOTf or MeI reagents. In quantitative yields proceed also the 1,6- → 1,10-carbon rearrangement reactions of the isomeric compounds closo-1,6- $R_2C_2B_8Me_8$ (3). Moreover, the permethylated compounds, such as 2a and 2b, can be, in fact, envisaged as egg (or ellipsoid) shaped hydrocarbons (see Fig. 5) of remarkable air stability due to the protective sheath of the surrounding methyl groups. For example, the persubstituted 2a can be stored in air for at least a month without any noticeable change, while the unprotected 1a is decomposed in air within a couple of hours, especially in a solution. The less stability of the unprotected intermediatesized carborane 1a derives from its non-icosahedral constitution (though it exhibits features of 3D aromaticity²³). The quantitative yields and relative easiness of the synthesis predestinate these persubstituted derivatives for using in designed syntheses in specific areas of carborane chemistry as multipurpose reagents, for example in cluster-insertion/ expansion or cage-degradation processes. Apart from this, such compounds are expected to exhibit extreme hydrophobicity, which can be made use of in various directions of chemical or biochemical research. Relevant experiments aimed at extension of permethylation chemistry are therefore in progress in our laboratories.

Conflicts of interest

There are no conflicts to declare.

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Acknowledgements

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