Nomenclature for boranes and related species (IUPAC Provisional Recommendations)

Michael A Beckett^{1*}, Bernd Brellochs², Igor T Chizhevsky^{3†}, Ture Damhus⁴, Karl-Heinz Hellwich⁵, John D Kennedy⁶, Risto Laitinen⁷, Warren H Powell⁸, Daniel Rabinovich⁹, Clara Viñas¹⁰, Andrey Yerin¹¹.

¹School of Chemistry, Bangor University, UK; ²Boron Hydride Highlights, Neufahrn, Germany; ³INEOS, Russian Academy of Sciences, Moscow, Russia, [†]Deceased 22nd May 2016; ⁴Novozymes A/S, Bagsvaerd, Denmark; Beilstein-Institut zur Förderung der Chemischen Wissenschaften, Frankfurt, Germany, ⁶School of Chemistry, The University of Leeds, Leeds, UK and the Institute of Inorganic Chemistry of the Czech Academy of Sciences, Řež u Prahy, The Czech Republic; ⁷Oulu University, Finland; ⁸Columbus, Ohio, USA; ⁹Charlotte, North Carlolina, USA, ¹⁰Institut de Ciencia de Materials de Barcelona (CSIC), Bellaterra, Spain; ¹¹Advanced Chemistry Development, Moscow, Russia.

Abstract: An appraisal of the current IUPAC recommendations for nomenclature of boranes and related systems has been undertaken. New developments in the field have been investigated and existing nomenclature systems have been adapted to accommodate these new developments. The principal areas considered are stoichiometric and structural nomenclature (including heteroatom and metal-atom subrogation, and substitution of hydrogen), conjoined-cage species, supra-icosahedral systems and sub-icosahedral non-standard structures. Elements of substitutive, additive and replacement nomenclature systems have been integrated into individual names to address contentious problems in boron nomenclature that have been around for a long time.

Keywords: boranes, boron hydrides, carbaboranes, heteroboranes, JUPAC Chemical Nomenclature and Structural Representation Division, 1UPAC Inorganic Chemistry Division, metallaboranes.

* email for correspondence: m.a.beckett@bangor.ac.uk

CONTENTS

BN-1 Introduction

BN-2 Stoichiometric nomenclature for boron hydrides

BN-3 Structural aspects of neutral boron hydrides (polyboranes)

- BN-4 Substitution and skeletal replacement (subrogation) of neutral boron hydrides (polyboranes) BN-4.1 Hydrogen-atom replacement BN-4.2 Subrogation (skeletal replacement)
- BN-5 Substitution and skeletal replacement (subrogation) of anionic boron hydrides (polyhydridoborates)

BN-5.1 Hydrogen-atom replacement

BN-5.2 Subrogation (skeletal replacement)

- BN-6 Substitution and skeletal replacement (subrogation) of cationic boron hydrides (polyhydridoborons)
 - BN-6.1 Hydrogen-atom replacement

BN-6.2 Subrogation (skeletal replacement)

BN-7 New developments

BN-7.1 Conjoined systems

BN-7.2 Supra-icosahedral systems

BN-7.2.1 Introduction

BN-7.2.2 Closed 13-vertex cages

BN-7.2.3 Closed 14-vertex cages

BN-7.2.4 Closed 15-vertex cages

BN-7.2.5 Closed 16-vertex cages

BN 7.2.6 Open supra-icosahedral cages

BN-7.3 Non-conventional deltahedral structures

BN-7.4 Metal-rich cages

BN-1 INTRODUCTION

The most recent IUPAC recommendations concerning boron hydrides and related compounds appeared in 2005 [1a]. These recommendations reinforced earlier IUPAC recommendations in the subject area published in 1990 [2a], but was essentially a 'holding' chapter, leaving new recommendations for a subsequent edition. The 'Introduction' [2b] to the 1990 recommendations held that the area of boron nomenclature is 'contentious' and 'specialized' and that the relevant Chapter (I-11) 'is presented as a survey of nomenclature of simpler boron systems because boranes are much discussed even at relatively elementary levels of chemistry, and because we wish to codify those basic principles which seem firmly established'. Many of these recommendations were based on a Technical Report prepared in 1972 by R. Adams [3a]. This 1972 report expands upon a brief chapter on nomenclature of boron compounds in the 1970 recommendations [4a].

The purpose of this present document is to examine existing nomenclature recommendations to ensure that they can be clearly interpreted and to introduce new recommendations, incorporating new developments which have occurred in the chemistry since these earlier reports. Each section is organized by starting with numbered Recommendations and these are followed by a more detailed commentary with relevant examples. Substituted boron hydride cages are named here by a nomenclature that encompasses features borrowed from both 'substitutive' and 'additive' IUPAC nomenclature systems.

BN-2 NOMENCLATURE FOR BORON HYDRIDES AND ANIONS

Recommendation 1: Stoichiometric names for boron hydrides and additive names for boron hydride anions, as described in the 1990 [2c] and 2005 [1b] recommendations, should continue to be used, with the one change that the ligand name 'hydro' is replaced by 'hydrido' in the additive

names.

Recommendation 2: Ambiguous class names (*e.g.* 'borates', or 'carboranes'), even if in common use, are unacceptable and should no longer be used. Certain more explicit and less ambiguous class names (*e.g.* 'hydridoborates', 'oxidoborates', '1,2-dicarbadodecaboranes') are acceptable.

Stoichiometric or compositional names of boron hydrides are, as defined in earlier IUPAC recommendations, generally useful and in most cases are easy to derive. Thus, neutral boron hydrides are named boranes, with a prefix to indicate the number of boron atoms present, and this is then followed, in parentheses, by the number of hydrogen atoms present. Illustrative examples are diborane(6), pentaborane(9) and decaborane(14) for B_2H_6 , B_5H_9 and $B_{10}H_{14}$, respectively.

Anions derived from boranes are named additively as hydridoborates borates. The numbers of hydrogen atoms (considered formally as hydride ligands) and boron atoms are specified by multiplicative prefixes, and the name has the ending 'ate' which is followed by the overall charge of the anion indicated in parentheses. The ligand name 'hydro' previously used has been changed to 'hydrido'. Thus $[B_{10}H_{10}]^{2-}$ and $[B_{12}H_{12}]^{2-}$ are named decahydridodecaborate(2–) and dodecahydridododecaborate(2–), respectively.

Note: In substitutive nomenclature anions formed by the addition of a hydride to a parent hydride are denoted by the suffix 'uide', with locant if relevant. For example, using this method tetrahydridoborate(1-) becomes boranuide, an acceptable name, and $[CH_3NHNH_3]^-$ is 2-methyl hydrazine-1-uide. This method could, in principle, be extended to other anionic borane systems, but this is not explored further in the present document.

Often, shortened names are observed in the chemical literature *e.g.* decaborate(2–), carborane. If such names are taken in context, may be satisfactory, but these abbreviated forms, when taken in isolation, are often misleading. This is particularly true for 'borates', which in addition to anionic boron hydrides has also been traditionally used for boron-oxygen anions. It should be noted that 'borate' (in the singular) is a IUPAC-accepted (abbreviated or traditional) name for trioxidoborate(3–), following the general pattern for naming the fully dehydronated anions corresponding to the classical oxoacids (sulfate, phosphate, *etc.*). More informative generic terms such as 'oxidoborates' and 'hydridoborates' should rather be used. See Section BN-4.2 for a short discussion on the use of 'carborane'.

BN-3 STRUCTURAL ASPECTS OF NEUTRAL BORON HYDRIDES (POLYBORANES)

Recommendation 3: The closed (*closo*) structures of deltahedra with 4–12 vertices, as defined as shown in Figure 1 and 1990 recommendations [2d], are used as the basis of boron-hydride nomenclature.

Recommendation 4: It is recommended that structural descriptors *closo, nido, arachno,* as defined in the 1990 recommendations [2d], are retained (Figure 2). However, the polyhedral shapes for *hypho* and *klado* are structurally difficult to visualize and use of these descriptors is no longer acceptable.

Recommendation 5: The structural descriptors for boron hydrides should be inserted into the name immediately before the prefix indicating the number of cage atoms.

Recommendation 6: Hydrogen atoms (and other groups) may be exo, endo or in bridging

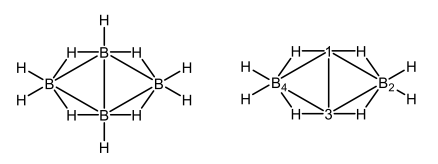
positions as defined in the 1990 recommendations [2e], and these descriptors are recommended.

Boranes generally have structures in which the boron atoms occupy vertices of closed or open deltahedra (triangulated polyhedra). These are often called 'clusters', although, in view of that these deltahedra do not contain interior atoms, 'cages' may be a more appropriate term. Both terms are in common use in the field, but for consistency 'cages' is used throughout this manuscript. These polyhedral shapes may have all of their vertices occupied by boron atoms or may be more open with some vertices absent.

IUPAC recommendations [2d] have been introduced which describe structures classed as deltahedra as *closo* and the more open deltahedral structures ('partially occupied' deltahedra) as *nido*, *arachno*, *hypho*, *or klado* depending upon the number of vertices missing from the parent *closo* cage structure. Thus, a deltahedron with one vertex missing is described as *nido*, one with two vertices missing is described as *arachno*, one with three vertices missing is called *hypho*, and one with four vertices missing is called *klado*. It is usually a vertex of highest connectivity (bonded to the greatest number of neighbours) which is removed to convert a *closo* structure to a *nido* structure, and an additional vertex adjacent to the previously highest connected vertex (thus located at the open face of the *nido*) is also removed to give rise to an *arachno* structure (Figure 2). Removal of vertices other than those of highest connectivity can result in structural isomers; in cases such other isomers have been called iso, neo, *etc*...

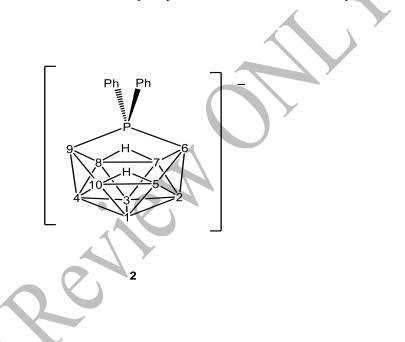
The prefixes *hypho* and *klado* are difficult to visualize from their structures; often the relationship to a *closo* deltahedral parent is not clear, and such descriptors are often given based on electron-counting schemes. Electron-counting rules [5] have been developed in parallel to these structural identifiers as an aid to understanding the structures of the boron hydrides (and their related compounds) based on their molecular (or ionic) formulae. However, a danger is that these structural descriptors are based purely on conclusions drawn from electron-counting schemes, rather than observed structure, and therefore should not be used. This matter is dealt with in more detail in Section BN-7.3.

In the simpler boranes, hydrogen atoms are attached on each boron atom by bonds that are directed radially out from the centre of the polyhedron, in positions defined as *exo*-terminal. **Unless otherwise stated, the following convention is followed for all structural diagrams within this manuscript: vertices specified by a 'number only' represent such {BH} units, whereas boron atoms with two terminal hydrogens or without terminal hydrogen atoms, and non-boron atoms, are always shown by an element symbol (with subscript numbering) and with all attached substituent groups/ligands shown**. Thus, for example, structure **1A** shows the connectivities of *arachno*-B₄H₁₀ and structure **1B** illustrates how such a structure is represented within this manuscript. In the more open *nido/arachno* structures, additional hydrogen atoms are also present and these either bridge two boron vertex atoms and are called bridging hydrogen atoms, or they may be associated solely with one boron atom and oriented tangentially towards a 'missing' vertex, when they are described as *endo*-terminal hydrogen atoms (for example, both are illustrated in structure **1A** and **1B**).



The descriptors *exo*, bridging, and *endo* are approved IUPAC descriptors in terminology and nomenclature for boron hydride cages [2e], and are in common usage. Some open-face hydrogen atoms have 'partial bridging' character intermediate between bridging and *endo*; it is recommend that such hydrogen atoms can be described for nomenclature purposes either as bridging or as *endo*, and any perceived 'partial' character discussed in the relevant research paper.

For *arachno* systems the removal of non-adjacent vertices can result in structures that have been called 'remote', *e.g.* the remote-*arachno* eleven-vertex structure of the $[(Ph_2P)B_{10}H_{12}]^-$ anion (2). In general, the use of 'remote' is not acceptable any longer and in this particular case a nomenclature based on *arachno* ten-vertex rather than *arachno* eleven-vertex can be used, with a $[\mu-6,9-(PPh_2)-arachno-B_{10}H_{12}]^-$ formulation as a basis for the full description, and a similar bridging approach to nomenclature serves adequately for several other 'remote' examples.



BN-4 SUBSTITUTION AND SKELETAL REPLACEMENT (SUBROGATION) OF NEUTRAL BORON HYDRIDES (POLYBORANES)

BN-4.1 Hydrogen atom replacement

Recommendation 7: It is recommended that the numbering schemes, as shown in Figures 1 and 2 and generally involving the clockwise-in-planes system as defined in the 1990 recommendations [2f, 2g], should continue to be used.

Recommendation 8: Open 8- and 10-vertex systems (Figure 2) have anomalous (*i.e.* not using the clockwise-in-planes) numbering systems which are in current common usage, and it is recommended that this should continue [2g].

Recommendation 9: In neutral polyborane species the replacement of a hydrogen atom by another atom or a monovalent group (a one-electron donor substitution) is denoted by using a substitutive prefix, and the replacement of a hydrogen atom by a neutral molecule (two-electron

donor substituent) is denoted by prefixing an IUPAC name for the molecule to the polyborane name, indicating the coordinating atom by the kappa (κ) convention [1c, 1d].

Note: The latter procedure resembles the treatment of ligands in additive nomenclature (except that ligands there are not replacing an atom in an initial parent structure) and we shall also call such donor substituents 'ligands' here. However, in the present document, when there are several identical and identically coordinating ligands attached to the same or different vertices in a polyborane cage, the ligand name is prefixed by a multiplicative prefix and by the locants of the vertices in question rather than incorporating these locants together with the kappa symbols as is done in [1c] when there are several atoms.

Note: Ongoing work on developing the kappa convention indicates that multiplicative ligand names cannot always be used in connection with the kappa convention and other IUPAC name types may have to be used. Also, for polydentate ligands, it may be inevitable to return to incorporating the locants together with the kappa symbols. A full explanation of such more complex situations will appear in a general treatment of the kappa convention in a future publication.

Recommendation 10: In cases where a boron cage atom no longer has a terminal hydrogen atom (or another one-electron substituent group) attached to it the term 'dehydro' preceded by that boron atom's locant must be used as a prefix immediately before the structural descriptor. The number in parentheses counting hydrogen atoms in the parent structure still includes this 'missing' hydrogen atom.

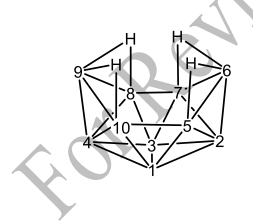
Neutral polyborane cages are typified as having triangulated polyhedra of boron atoms, with each boron atom carrying an *exo* hydrogen atom. To avoid ambiguity and for accurate definition, numbering schemes for boron vertices (and associated *exo* substituents) have been developed (Figures 1 and 2, Recommendation 7). The ten-vertex *nido* and *arachno* open structures based on the octadecahedron and the icosahedron with appropriate vertices removed are anomalous both in that the members of each *arachno/nido* pair are topologically identical (Figure 2), and in that their open faces are numbered sequentially as a 'unit' rather than in 'planes' as noted in the 1990 recommendations [2g]. The numbering system as used for *nido-* and *arachno-*decaboranes has a long history and has much literature associated with it and this anomalous numbering is retained (Figure 2 and Recommendation 8). The *nido* and *arachno* eight-vertex cage structures are also similarly anomalous in that they also have the same topology as each other, and in that the open face is numbered sequentially as a unit rather than in planes; again, because of ingrained historical usage, this 'anomalous' numbering is retained (Figure 2 and Recommendation 8). Similar considerations must also apply to the historically established open-face numbering of the '*iso-arachno*' nine-vertex structure, which is commonly referred to simply as the *arachno* structure.

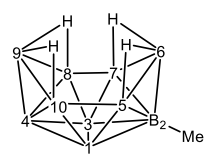
Structures that are more open also have extra hydrogen atoms which may be *bridging* and/or *endo* hydrogen atoms and these may be located by an adaption of the indicated hydrogen technique of organic substitutive nomenclature [6a]. These extra hydrogen atoms may be located by attaching a locant to an italic 'H' (*e.g.* -2H-) for a terminal hydrogen atom, and locants to the symbol ' μ H' (*e.g.* 2,3- μ H *etc.*) for bridging hydrogen atoms. In this method, each boron atom is assumed to have one hydrogen atom and only hydrogen atoms additional to these are cited. It must be noted that these cited hydrogen atoms are **not** in addition to those included in the parenthetical hydrogen atom count. *E.g. arachno*-B₄H₁₀ (**1A** and **1B**) may be named *arachno*-tetraborane(10) (stoichiometric/compositional name) or 2H,4H-1,2- μ H:1,4- μ H:2,3- μ H:3,4- μ H-*arachno*-tetraborane(10) (indicated hydrogen atom method). The 1990 edition of the IUPAC Inorganic rules [2h] introduced a shortened version for the citation of bridging hydrogen atoms illustrated by

the following format for *arachno*-B₄H₁₀, namely, 2*H*,4*H*-1,2:1,4:2,3:3,4-tetra- μ *H-arachno*-tetraborane(10). This shorter format is used in these recommendations. It should be noted for consistency within this document that bridging hydrogen atoms always follow after non-bridging hydrogen atoms within a name. This is not a formal recommendation and a full consideration of this and the treatment of other modified suffixes will be the subject of further studies.

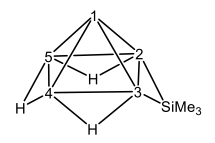
Many compounds that have cage structures can be described in terms of simple replacement of these *exo* or *endo* hydrogen atoms by various functional groups, *i.e.* they are formally substituted compounds, as exemplified by organylboranes or halogenoboranes. Bridging hydrogen atoms may also be substituted and these are indicated using the bridging symbol μ and the locants for the two boron vertices that are bridged. The method for numbering boron vertices is robust and can also be extended to polyhedra with more than twelve vertices (see Section BN-7.2). The location of hydrogen atoms that are additional to one on each boron vertex can be specified by the 'indicated hydrogen' adaption [6a].

The formal replacing group for a hydrogen atom may be either another substituent (oneelectron group), *e.g.* Cl, CH₃, Si(CH₃)₃, *etc.*, or a neutral molecule, *e.g.* NMe₃, PPh₃, SMe₂, NC₅H₅, *etc.*, which functions as a two-electron ligand. A 'one-electron group' is named as a substituent prefix in neutral polyboranes. A 'two-electron group' is a neutral group and is cited as a neutral ligand. Substitution (replacement) by a substituent (one-electron group) is straightforward, and follows standard IUPAC substitutive nomenclature practice: derivatives are named based on the parent with the number of hydrogen atoms in parentheses being that of the parent. Thus substitution of an *exo* hydrogen at the 2-position of *nido*-decaborane(14) (3) by a methyl group results in 2-methyl-5,6:6,7:8,9:9,10-tetra- μ *H-nido*-decaborane(14), (4). In a similar way, substitution of a bridging hydrogen in *nido*-pentaborane(9) by a trimethylsilyl group leads to 2,3- μ -(trimethylsilyl)-(2,3:2,5:3,4:4,5-tetra- μ *H)-nido*-pentaborane(9) (5).





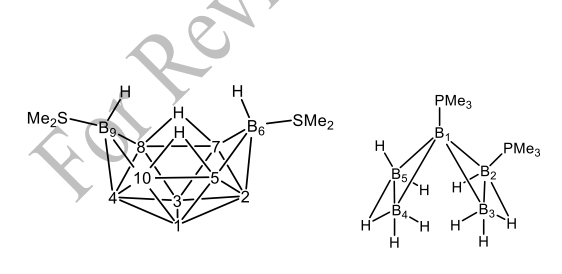
4

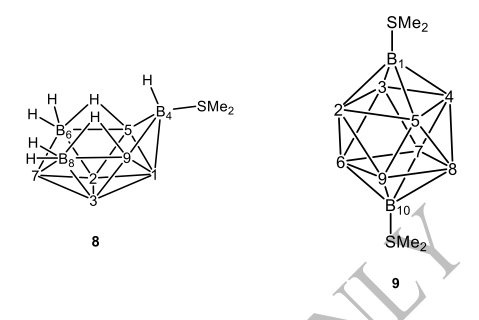


Substitutions that are not on symmetry axes or planes can lead to stereoisomers. For example the monochloro derivative of decaborane that would simply, and generally reasonably be described for most purposes, as 5-Cl-*nido*-B₁₀H₁₃, will, as generally prepared, in fact be a racemate. Separation of the enantiomers is rare, and although in principle these compounds could be named as under a system developed in organic chemistry on chiral fullerene nomenclature (see section Fu-17, [6b]), no formal recommendations on the topic are made here; these will need future further studies.

Substitution of hydrogen atoms by neutral two-electron donor groups can increase the number of electrons available for skeletal bonding. For example, replacement of two one-electron hydrogen atoms by two two-electron ligands can in some cases increase the number of internal cage bonding electrons by a count of two and in some cases induce a formal *closo* to *nido* or *nido* to *arachno* structural change, *i.e.* it can induce a rearrangement of the cage to a more open structure. Thus, for example, the conversion of $B_{10}H_{14}$ (3) to $6,9-(SMe_2)_2B_{10}H_{12}$ (6) results in a *nido*-to-*arachno* conversion. Experimentally, the reaction of excess SMe₂ with *nido*decaborane(14) (3) leads to $6,9-(SMe_2)_2$ -*arachno*- $B_{10}H_{12}$ (6) with the elimination of two hydrogen atoms as H₂. Because of this easy synthesis, such $B_{10}H_{12}L_2$ compounds, where L is a two-electron ligand, are often referred to as 'adducts' of *nido*- $B_{10}H_{14}$ despite them not being true adducts [7, 8]. The 'indicated hydrogen atoms' approach described above [6a], gives the name 6,9bis(dimethylsulfane- κS)-5,10:7,8-di- μ H-*arachno*-decaborane(12) for **6**.

Compound 7, 1,2-(PMe₃)₂B₅H₉, in the 1990 recommendations [2i] was given the addition compound name (trimethylphosphine)—*hypho*-pentaborane(9) (2/1); it is now named 1,2-bis(trimethylphosphane- κP)-3H,4H,5H-2,3:4,5-di- μ H-1-dehydro-pentaborane(10). Another familiar example, 4-(Me₂S)-*arachno*-B₉H₁₃, is named as 4-(dimethylsulfane- κ S)-6H,8H-5,6:8,9-di- μ H-*arachno*-nonaborane(13) (8). In accord with Recommendation 10, the correct name for 1,10-(Me₂S)₂-*closo*-B₁₀H₈ (9) is 1,10-bis(dimethylsulfane- κ S)-1,10-didehydro-*closo*-decaborane(10).





In terms of reactions, an addition of a two-electron ligand L with no other changes in empirical formula can indeed result in an adduct, *e.g.* B_8H_{10} to $B_8H_{10}L$. However, in many instances this can increase by two the number of electrons available for skeletal bonding, *e.g.* a change from *nido* to *arachno*, and so the term 'adduct' can also be avoided here in terms of nomenclature. A further problem with the use of 'adduct' in the general case is that it supposes a reaction – an addition – whereas nomenclature is concerned with the observed structure rather than the means of synthesizing it. A caveat here is that in some transition-element metallaborane and metallaheteroborane compounds a two-electron ligand may add to the metal atom centre and not fundamentally affect the intracage bonding so that the same cage structure (and the same skeletal bonding-electron count) are both retained. Since connectivities are being maintained this latter example would correspond to an adduct as defined by IUPAC [7, 8].

BN-4.2 Subrogation (skeletal replacement)

Recommendation 11: The structural descriptors should be inserted into the name immediately before the locant(s) and prefix(es) indicating the subrogating atom(s) present in the cage.

Recommendation 12: Subrogating atoms are indicated by using conventional skeletal replacement terms, e.g. carba for carbon, and in the conventional 'snake order' ($F \rightarrow Rn$) [1e, 9], with each 'a' term preceded by an appropriate locant.

Recommendation 13: The number of hydrogen atoms indicated in parentheses are those for the parent heteroborane, and this does not change upon substitution.

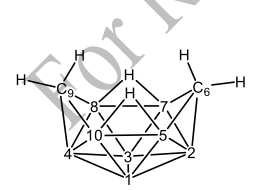
Recommendation 14: Hydrogen atoms at metal centers in metallaboranes are named as hydrido ligands and are not included in parentheses as part of the parent subrogated borane hydrogen count.

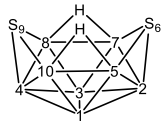
Boron atoms within a polyhedral cage framework may be replaced by atoms of other maingroup elements or by atoms of transition-metal elements, with the structural integrity of the cage being retained. Such skeletal replacement is called *subrogation*. The term subrogation is specific to skeletal replacements in boron hydride chemistry, and is not used elsewhere. The resulting structures are called *heteroboranes* in accord with the usage in the 1990 recommendations [2j]. The common variant 'heteraboranes' is not acceptable anymore. On the other hand, boranes that have boron atoms subrogated by metallic main-group or transition-element atoms are termed *metallaboranes* [2k].

Metallaheteroboranes with boron atoms subrogated by main-group and transition-element atoms are very common. The atom that subrogates the boron atom often (but not necessarily) has an *exo* group associated with it; *exo* groups may include hydrogen atoms, halogen atoms, organic groups, neutral molecules, *etc.* Subrogating atoms are indicated using the conventional replacement terms in the conventional 'snake' order ($F \rightarrow Rn$) *e.g.* oxa represents O; thia S; aza N; phospha P; arsa As; carba C; sila Si; aura Au; ferra Fe; *etc* [1e, 9]. When there is a choice of numbering the senior heteroatom is assigned the lowest locant. The name of the subrogated compound is derived from the parent unsubrogated borane with the number in parentheses of the parent hetero/metallaborane retained as a suffix.

Thus $closo-1,2-C_2B_{10}H_{12}$ is named as a dodecaborane with two vertices replaced by carbon and twelve hydrogen atoms in total, *i.e.* closo-1,2-dicarbadodecaborane(12). This last species, closo-1,2-dicarbadodecaborane(12) has been variously referred to as closo-1,2-dicarborane, ortho-dicarborane, dicarborane – and even simply carborane – as well as other variants. This casual approach to nomenclature in closo-dicarbadodecaborane(12) chemistry severely inhibits data-base searches and retrievals (see Section BN-2, and Recommendation 2).

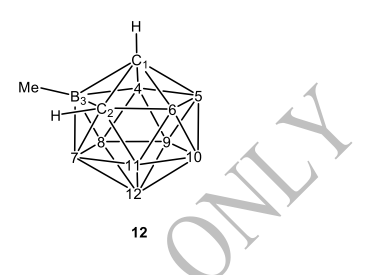
IUPAC convention [21, 3b, 4b] traditionally places the structural descriptor right in front of the borane segment of the name. However, Recommendation 11 now focuses on the 'cage' (and what it contains) and places the descriptor before the cage name *e.g. closo*-1,2-dicarbadodecaborane(12) rather than the previously accepted 1,2-dicarba*-closo*-dodecaborane(12). For the purpose of nomenclature there is an implied hydrogen atom on every boron or carbon vertex within subrogated polyboranes, and all other heteroatoms or metallic atoms do not have implied hydrogen atoms. This correctly highlights and describes the hydrogen atom count (including heteroatom substituents) for the numerical suffix in parentheses for the heteroborane parent. Therefore the compounds *arachno*-6,9-C₂B₈H₁₄ (**10**) and *arachno*-6,9-S₂B₈H₁₀ (**11**) are named as $6H,9H-5,10:7,8-di-\muH-arachno-6,9-dicarbadecaborane(14)$ and $5,10:7,8-di-\muH-arachno-6,9-dicarbadecaborane(14)$



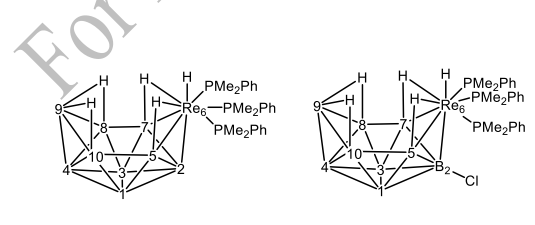




The correct name for 3-Me-*closo*-1,2-C₂B₁₀H₁₁ (12) is 3-methyl-*closo*-1,2-dicarbadodecaborane(12) (Recommendation 13) since the parent heteroborane is *closo*-1,2-C₂B₁₀H₁₂.

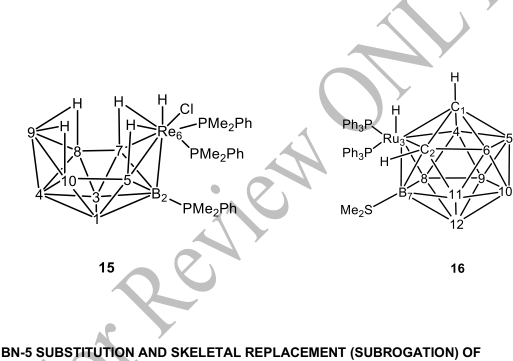


IUPAC recommendations [2m] from 1990 briefly described a method for the naming of metallaboranes possessing *exo*-hydrido ligands. Here it is 'normally not assumed', when determining the number to be placed in parentheses, that metal atoms would carry hydrogen atoms in the unmodified parent, and hence any such hydrogen atoms do need to be specified. With this system where *exo*-hydrido ligands to transition-metal atoms are not included in the hydrogen atom count for the parent metallaborane, but are indicated as hydrido ligands at the metal centre. Substitution on boron (or heteroatom) would retain the parent hydrogen atom count as is accepted for boranes and heteroboranes. This is illustrated by the following examples from *nido*-6-rhenadecaborane chemistry [10]: $6,6,6-(PMe_2Ph)_3-6-H-nido-6-ReB_9H_{13}$ (13) and $2-Cl-6,6,6-(PMe_2Ph)_3-6-H-nido-6-ReB_9H_{12}$ (14) are named $6,6,6-tris[dimethyl(phenyl)phosphane-\kappa P]-6-hydrido-5,6:6,7:8,9:9,10-tetra-µH-nido-6-rhenadecaborane(13), and <math>2-chloro-6,6,6-tris[dimethyl(phenyl)phosphane-\kappa P]-6-hydrido-5,6:6,7:8,9:9,10-tetra-µH-nido-6-rhenadecaborane(13), respectively.$



There are many metallaborane and metallaheteroborane cage compounds that include neutral molecules (two-electron donor substituents) that formally replace *exo* B-H hydrogen atoms. Thus, 6-Cl-2,6,6-(PMe₂Ph)₃-6-H-*nido*-6-ReB₉H₁₂ (**15**, an isomer of **14**) can be named 6-chlorido-2,6,6-tris[dimethyl(phenyl)phosphane- κP]-6-hydrido-5,6:6,7:8,9:9,10-tetra- μ H-2-dehydro-*nido*-6-rhenadecaborane(13). It should be noted that 'dehydro' should be part of the parent structure and thus cited immediately before the structural descriptor whereas 'hydrido' is cited alphabetically together with other ligand and/or substitutive prefixes.

Many complexes in heteroborane and metallaheteroborane chemistry that are also in this category are the so-called 'charge-compensated' complexes, *e.g.* $7-(SMe_2)-3-H-3,3-(PPh_3)_2$ -*closo*-1,2,3-C₂RuB₉H₁₀ (**16**) is named 7-(dimethylsulfane- κ S)-3-hydrido-3,3-bis(triphenylphosphane- κ P)-7-dehydro-*closo*-1,2-dicarba-3-ruthenadodecaborane(11).



ANIONIC BORON HYDRIDES (POLYHYDRIDOBORATES)

BN-5.1 Hydrogen atom replacement

As noted in Section BN-2, anionic polyboron hydride derivatives (polyhydridoborates) are named additively resulting in the 'ate' ending instead of the 'ane' ending immediately followed by the charge number in parentheses. The structural descriptors and numbering remain the same as for the corresponding neutral polyboranes. All hydrogen atoms are described as 'hydrido' ligands. The use of locants is not required if all boron sites carry a terminal hydrogen atom. Therefore closo-[B₆H₆]²⁻ is named hexahydrido-closo-hexaborate(2–). Bridging hydrogen atoms are denoted by μ -hydrido together with appropriate locants, *e.g.* 2,3- μ -hydrido. All other groups are also named as ligands. *E.g.* 2-methyl-*nido*-decaborane(14) (**3**), 2-MeB₁₀H₁₃, can be dehydronated by losing a bridging hydrogen atom as a hydron to form an anion [2-MeB₁₀H₁₂]⁻ and this is named

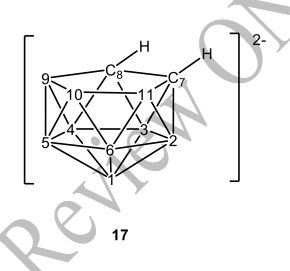
1,3,4,5,6,7,8,9,10-nonahydrido-5,6:6,7:8,9-tri-µ-hydrido-2-methanido-nido-decaborate(1-).

BN-5.2 Subrogation (skeletal replacement)

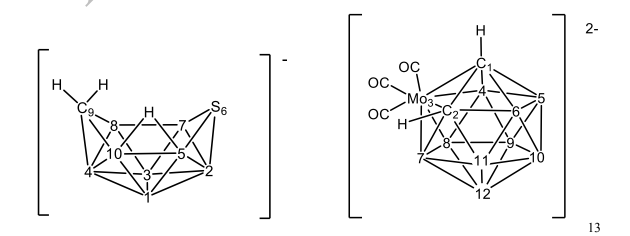
Recommendation 15: In the names of subrogated polyhydridoborates, the order of citation of the elements at cage vertices should follow the conventional 'snake' order ($F \rightarrow Rn$) and subrogating atoms are indicated using the conventional replacement terms [1e, 9], *e.g.* carba for carbon.

Note: This ordering of the element in the names of anionic subrogated polyboranes is the opposite order of citation of central atoms in additive nomenclature.

As an example the anion, nido- $[7,8-C_2B_9H_{11}]^{2-}$ [also known colloquially as the 'dicarbollide' anion] (17) is named undecahydrido-nido-7,8-dicarbaundecaborate(2-).



The method can be used to name polyhydridoborate anions with further subrogation using the element order, as specified in Recommendation 15. Thus, $[6,9-SCB_8H_{11}]^-$ (18) can be correctly named as 1,2,3,4,5,7,8,9,9,10-decahydrido-5,10- μ -hydrido-*arachno*-6-thia-9-carbadecaborate(1–) (although 6,9-CSB_8H_{11}]^- has been used in the literature) and $[3,3,3-(CO)_3-1,2,3-C_2MOB_9H_{11}]^{2-}$ is named 3,3,3-tricarbonylundecahydrido-*closo*-1,2-dicarba-3-molybdadodecaborate(2–) (19).



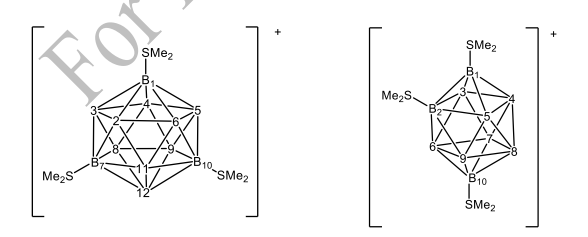
BN-6 SUBSTITUTION AND SKELETAL REPLACEMENT (SUBROGATION) OF CATIONIC BORON HYDRIDES (POLYHYDRIDOBORONS)

BN-6.1 Hydrogen atom replacement

Recommendation 16: Cationic boron hydrides are named as polyboron(n+) (n = charge on the cation) cages as described in an IUPAC publication [4c]. It is further recommended that the structural structures and numbering systems, as applied to neutral (uncharged) polyboranes and anionic polyhydridoborates, are also used for polyboron(n+) cations, and that all substituents, including hydrogen atoms, are named as ligands.

Cationic boron hydride are uncommon, but despite this, suitable nomenclature needs to be available for such compounds. A system for naming cationic polyboron cages is described in **IUPAC** recommendations 4c] where $[B_{10}H_7(NH_3)_3]^+$ is named [2n, 3c, as triammineheptahydrodecaboron(1+). It should be noted that the name is formulated as a polyboron derivative and it is immediately followed by the charge in parentheses. IUPAC now uses 'hydrido' rather than 'hydro' when naming hydride ligands in additive nomenclature and triammineheptahydridodecaboron(1+) is now the correct name. To be consistent with anionic cages, structural descriptors and numbering schemes are again needed in order to specify skeletal structures, and substituted positions and numbering systems remain the same as those used for neutral (uncharged) polyboranes and polyhydridoborates (Recommendation 16). Again, all hydrogen atoms are described as hydrido ligands, bridging hydrogen atoms are denoted by µhydrido ligands together with appropriate locants, e.g. 2,3-µ-hydrido, and all substituent groups on boron sites are named as ligands e.g. chlorido, triphenylphosphane- κP , etc.

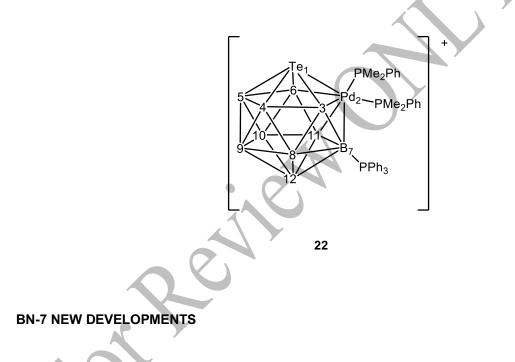
The cations present in the salts $[1,7,10-(Me_2S)_3-closo-B_{12}H_9][BF_4]$ and $[1,2,10-(Me_2S)_3-closo-B_{10}H_7][BF_4]$ are shown in structures **20** and **21**, respectively. These cations can be named as 1,7,10-tris(dimethylsulfane- κS)-2,3,4,5,6,8,9,11,12-nonahydrido-closo-dodecaboron(1+) and 1,2,10-tris(dimethylsulfane- κS)-3,4,5,6,7,8,9-heptahydrido-closo-decaboron(1+), respectively.



BN-6.2 Subrogation (skeletal replacement)

Recommendation 17: Subrogated polyboron cations are named as polyboron(n+) derivatives and subrogating atoms are indicated by using conventional replacement terms (*e.g.* carba for carbon) cited in the conventional 'snake' order (F \rightarrow Rn) [1e, 9]. The structural descriptors and numbering remain the same as for the neutral polyboranes and polyhydridoborates and all substituents, including hydrogen atoms, are named as ligands.

The cation observed in $[2,2-(PMe_2Ph)_2-7-(PPh_3)-closo-1,2-TePdB_{10}H_9][BF_4]$ can be named as 2,2-bis[dimethyl(phenyl)phosphane- κP]-3,4,5,6,8,9,10,11,12-nonahydrido-7-(triphenylphosphane- κP)-closo-1-tellura-2-palladadodecaboron(1+) (**22**) [11].



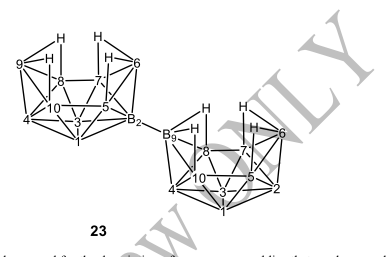
BN-7.1 Conjoined cages

Recommendation 18: The term *commo* is recommended [20] and complemented by the use of the structural terms *dicommo* and *tricommo* to describe conjoined polyhedral cages with 1, 2, or 3 atoms in common. The structural term should be placed as an infix in the name and surrounded by locants and the name of each cage in square brackets as follows: [borane]-a'*commo*-x-[borane], [borano]-a',b'*dicommo*-x,y-[borane], or [borano]-a',b',c'*-tricommo*-x,y-[borane].

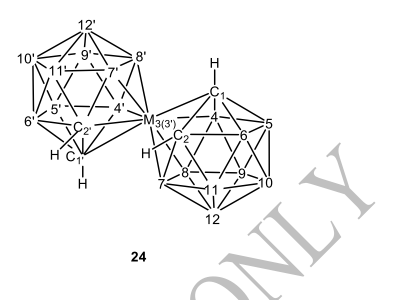
Compounds that contain two or more borane, heteroborane, metallaborane, *etc.*, cages that are joined together exhibit inter-cage conjunctions of varying intimacy. The cages can be joined by a linking (bridging) moiety, or by a σ -bond, or by using a more intimate inter-cage 'fusion' with one, two, or more atoms held in common between two cages. The term *conjuncto* has been used as a general term for all such species [20], but as the chemistry has progressed it now lacks

specificity for nomenclature purposes, and we recommend that its unqualified usage be discontinued in terms of nomenclature. The only previously recommended IUPAC term in this category is *commo* for cages with one atom in common [20].

Polyboron cages linked by a σ -bond should be named by the ring assembly methodology of organic nomenclature or by substituting one polyborane cage into another polyboron cage. Thus, the *sigma*-linked (B₁₀H₁₃)(B₁₀H₁₃) (**23**) is named 2-[5,6:6,7:8,9:9,10-tetra-µ*H-nido*-decaboran(14)-9-yl]-5,6:6,7:8,9:9,10-tetra-µ*H-nido*-decaborane(14).



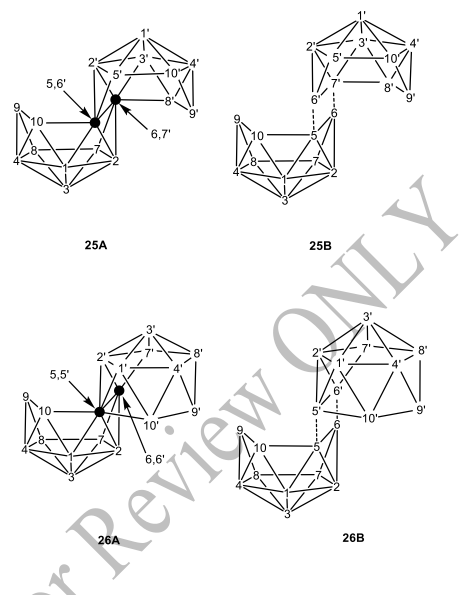
The term *commo* has been used for the description of two-cage assemblies that can be regarded structurally as consisting of two definable cages that are conjoined or fused with one atom held in common [20, 3d]. The *commo* nomenclature for the one-atom-in-common case is well entrenched in the literature and should be retained and extended to cages involving two atoms in common and three atoms in common. Hydrogen atoms attached to common atoms need to be cited, since it is otherwise assumed that they do not have hydrogen atoms attached in the count used for the number in parenthesis for the parent. Few examples are known in which the common atom is boron, but the term has been used extensively to describe the trivially-named 'bisdicarbollide' complexes of transition elements M, of nominal formulation $\{M(C_2B_9H_{11})_2\}$ (24). A terminal polyboron cage is given unprimed numbers and other polyboron cages are given serially primed numbers. Symmetrical *closo*-1,2,3-C₂MB₉H₁₁ systems are named as a 3,3'-*commo*-bis[*closo*-1,2-dicarba-3-metalladodecaborane(11)] but an alternative name (Recommendation 18) is [*closo*-1',2'-dicarba-3'-metalladodecaborane(11)]-3'-*commo*-3-[*closo*-1,2-dicarba-3-metalladodecaborane(11)] and this infix format should be used for non-symmetrical *commo* systems.



An important feature of the 'biscarbollide' and related one-metal-atom-in-common systems is the mutual contrarotation of the two subcages; a number of rotamers can therefore exist, and this can engender the usage of terms such as *gauche, transold, cisoid, eclipsed, gauchoid*, etc. Rotamer considerations, when relevant, are usually dealt with adequately as they arise in the scientific publications that report them; they are beyond the scope of this review. One possibility would be to define the structure based on the eclipsed conformation of lowest-numbered vertices (here C1 and C1' in **24** being those that are eclipsed), and introduce a clockwise rotation angle into the nomenclature relative to the nominally static basis cage (*i.e.* the unprimed cage).

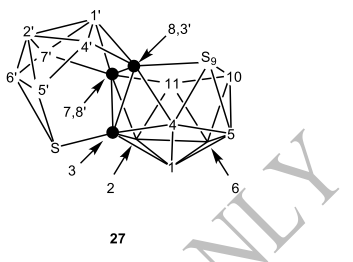
Compounds that contain two or more cages, with individual cages that are joined or fused to each other with two or more atoms held in common, are often referred to as 'macropolyhedral' species [12], a trivial, but often useful generic term, but with no role in formal nomenclature.

For cages containing a two-atoms-in-common fusion (often known as edge-fusion), which is a feature of the bulk of the so-far reported 'macropolyhedral' boranes, metallaboranes, and heteroboranes, (particularly, in the last category, thiaboranes), the term '*dicommo*' is introduced (Recommendation 18). The two known isomers of B₁₈H₂₂ can thence be described as $[5',6':8',9':9',10'-tri-\mu H-nido-decaborano(14)]-6',7'-dicommo-5,6-[6,7:8,9:9,10-tri-\mu H-nido-decaborane(14)] for the isomer commonly referred to as$ *n*-B₁₈H₂₂ (or*anti*-B₁₈H₂₂) (**25A**and**25B** $), and <math>[6',7':8',9':9',10'-tri-\mu H-nido-decaborano(14)]-5',6'-dicommo-5,6-[6,7:8,9:9,10-tri-\mu H-nido-decaborane(14)] for the isomer commonly referred to as$ *iso*-B₁₈H₂₂ (or*syn*-B₁₈H₂₂) (**26A**and**26B**). Structures**25A**and**26A**have common vertices which do not possess terminal H atoms, these vertices are signified by a black dot.

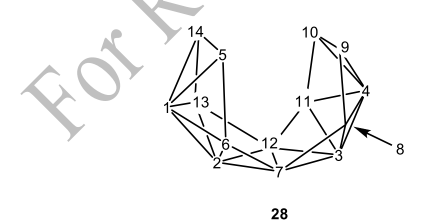


Here the trivial names *syn* and *anti* refer to the mutual disposition of the two *nido*-decaborane open faces, either on the same flank (*syn*) or on opposite flanks (*anti*) of the molecule. These mutual dispositions have been further categorised as '*transoid*', implying that the open faces are trans to each other, engendering trivial *transoid-anti* and *transoid-syn* descriptors for these two known isomers [13]. The alternative '*cisoid*' conjunctions, with the open faces *cis* to each other, have not yet been realised experimentally. However, such a '*cisoid*' conjunction has been proposed to be inherently isolatable.

There are cases where, in *addition* to the two-atoms-in-common fusion, the two subcages are conjoined with a σ -link, or by a one-atom bridge E. Organic nomenclature provides a system to describe the additional σ -link: the prefix 'cyclo' (not italicised) in front of the fused borane cages as cyclo-x,y-[borano]-a,b-*dicommo*-c,d-[borane]. Systems with bridging ligands may be similarly cited as μ -bridge-x,y-[borano]-a,b-*dicommo*-c,d-[borane], *e.g.* the S₂B₁₆H₁₆ species (**27**) can be named (5',6',3- μ 3-sulfido)[4',5':6',7'-di- μ H-nido-octaborano(8)]-8',3'-*dicommo*-8,7-[10,11- μ H-nido-9-thiaundecaborane(10)].



This *dicommo* system is generally applicable when the two subcages are recognisable in terms of known types of single cages. However, there are one or two problem compounds, and it is likely that more of these will be revealed as the area progresses. By a two-atoms-conjoined formalisation, and by hydrogen atom count, the compound $B_{14}H_{20}$ would in principle be a *dicommo nido*-eight-vertex / *arachno*-eight-vertex combination. However, this would be asymmetric, whereas the overall structure and hydrogen atom disposition are symmetrical. The overall formulation is B_nH_{n+6} , and would correspond to single-cage *arachno*, but to regard it as a fourteen-vertex fragment of a sixteen-vertex *closo* polyhedron is geometrically unrealistic. At present the species is unique and more examples of this general type would have to be isolated before a general structural pattern emerges and thus the basis for a systematic nomenclature perceived. The numbering scheme historically applied to $B_{14}H_{20}$ is as in **28** [14]. A related problem exists with $B_{12}H_{16}$ although this can at least be described *structurally* as a *dicommo* combination of a *nido*-shaped eight-vertex cage and a *nido*-shaped six-vertex cage, but precise bridging hydrogen atom locations would have to be specified.



As a logical extension of the *dicommo* approach, it is recommend that *tricommo* nomenclature is used, *e.g.* [borano]-a',b',c'-*tricommo*-x,y,z-[borane], is appropriate for cage compounds that can be interpreted in terms of recognizable single cages fused with three atoms in common (face

fusion) (Recommendation 18). This can be applied to several known compounds, but in some cases constituent cage types may become difficult to define in terms of known and definable single cage shapes, particularly so as the intercage fusion becomes more intimate.

Really intimate intercage fusion can in principle result in larger 'globular' boranes [15, 16] and related assemblies that have been called 'stuffed cages' [17]. The latter are non-hollow cages (cages with atoms inside) which are not recognisable in terms of individual single cage components and for which a systematic nomenclature is not practicable to devise until several experimentally determined examples are known and systematic structural patterns emerge. At the time of this report, only very hypothetical species such as $B_{27}H_{21}$ and $B_{84}H_{54}$ have been proposed based on speculative calculations [17], but perhaps future nomenclature in this area can utilise variants on organic diamondoid or endo-fullerene "@" nomenclature and numbering conventions. A unique species that should be mentioned in this context is B₂₀H₁₆, a long-known compound commonly named *closo*-B₂₀H₁₆ on account of its closed deltahedral configuration. However, the structure is not that of a deltahedral cage as typified by the conventional $closo-[B_nH_n]^{2-}$ structural sequence, but has a more columnar aspect, and so the use of 'closo' in terms of systematic nomenclature is inappropriate. Calculations suggest that B₂₀H₁₆ is in fact part of a structural and electronic progression based on dianionic closo-[BnHn]²⁻, neutral B20H16, dicationic [B28H20]²⁺, tetracationic [B₃₄H₂₄]⁴⁺, and so on, of which B₂₀H₁₆ and [B₂₈H₂₀]²⁺ would constitute initial members of a series of columnar structures, for which the prefix *columno* is appropriate. $B_{20}H_{16}$ would thence be *columno*-2,2,4,2,2,4,2,2- $B_{20}H_{16}$, the numbers representing the numbers of atoms in successive planes as the column is descended, this sequence being consistent with the previously described historical numbering system for this compound as in Section 11.11 of the 1970 recommendations [4d]. Alternatively, such structures could be named by the descriptors described in Section BN-7.2 using Recommendation 19.

BN-7.2 Supra-icosahedral structures

BN-7.2.1 Introduction

Recommendation 19: A new system for describing closed (*closo*) supra-icosahedral polyhedral shapes based on a published method [18] is recommended. Closed (*closo*) 13-16 vertex polyhedra are shown in Figure 3. In this system the overall idealized symmetry, the number of atoms in the cage within 'planes', their connectivities, and the number of triangulated faces may all be defined (Figure 3). Numbering, except for those described in Recommendations 20 and 21, follows the clockwise-in-plane rules as is current practice in sub-icosahedral cages, as defined in IUPAC recommendations 1990 [2f].

Currently there are no IUPAC recommendations for the naming of 'supra-icosahedral' boron hydrides (*i.e.* those with more than twelve cage vertices) or related species. Although theoreticians have long predicted supra-icosahedral boranes, metallaboranes and heteroboranes, the area is still very much under-developed from a synthetic viewpoint. However, progress in synthetic methods in recent years indicates that it is now timely to review this area. Closed (*closo*) structures are examined first and this will be followed by a discussion on open (*nido* or *arachno*) structures.

Theoretical calculations have been undertaken on deltahedral closed $[B_nH_n]^{2-}$ systems with between 13 to 24 boron atoms (*n*) [20a]. In keeping with sub-icosahedral closed cages, these

structures all contain *exo* hydrido substituents and they possess 12 vertices with a cage connectivity of five, and *n*-12 vertices with a cage connectivity of six. Continuation of the systematic nomenclature for sub-icosahedral structures would indicate that these should be correctly named as *closo* structures, and that *nido* and *arachno* structures could be derived from them using conventional vertex-subtraction rules. However, in light of synthetic work it is clear that structural aspects of supra-icosahedral cages are more complex than for the smaller sub-icosahedral cages and it is difficult to make (at present) any generalizations about what particular shapes may form the basis of a rigorous nomenclature. Thus, it has been shown that closed polyhedral shapes other than those [20a, 20b] historically calculated for $[B_n H_n]^{2-}$ are also available, and a more all-encompassing approach to nomenclature needs to be developed.

Supra-icosahedral $[B_nH_n]^{2-}$ cages have not (yet) been synthesized, but there are now known examples of 13-vertex and 14-vertex carbaboranes, 13-, 14-, and 15-vertex metallacarbaboranes, and 15-vertex and 16-vertex metallaboranes [21-24].

BN-7.2.2 Closed 13-vertex cages

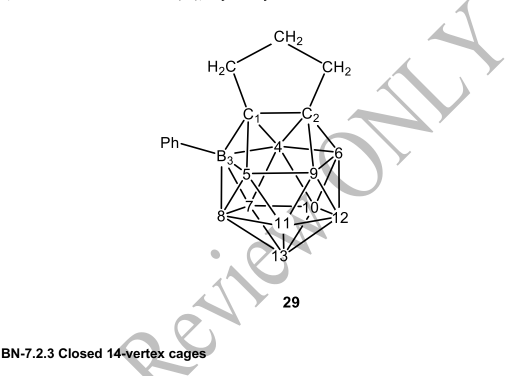
Recommendation 20: The non-standard numbering for henicosahedral 13-vertex species (Figure 3) species is well entrenched in the literature, and its continued use is recommended.

There are many known examples of closed 13-vertex {MC₂B₁₀} cage compounds, in which the cages adopt the deltahedral structure with 22 faces (docosahedron), isostructural with that calculated for [B13H13]²⁻. Recent calculations have suggested that structures of supra-icosahedral closo carbaboranes and metallacarbaboranes need not be fully deltahedral and that tubular or other polyhedral shapes might be possible [25], 13-Vertex { $C_{2B_{11}}$ } carbaboranes, e.g. [1,2- μ -{1',2'-C₆H₄(CH₂)₂]-3-Ph-closo-1,2-C₂B₁₁H₁₀, have a non-deltahedral henicosahedral (21 faces) structure, with twenty triangulated faces and one trapezoidal (C_2B_2) face [26]. This alternative closed structure needs to be differentiated from the docosahedral closed structure. This could most easily be accomplished by adopting a system first described by Powell and co-workers [18]. This system includes both the number of cage atoms found within the 'planes', and the number of triangulated faces, within the structural descriptor. Other additional qualifying descriptors, such as an idealized symmetry-point symbol, and the cage-atom connectivity (v^n) , may also be included, if necessary. For example, for the 10-vertex bicapped square antiprismatic polyhedron (hexadecahedron) the descriptor would be $[D_{2d}-(1v^4441v^4)-\Delta^{16}-closo]$ (the 'v⁵' descriptor for the 5vertex middle numbers has been omitted). For this manuscript the point group, the number of faces given in the superscript number to the Δ symbol, and the descriptor *e.g. closo* are generally included in the structural descriptor contained in square brackets (Recommendation 19).

Numbering systems in both the henicosahedral and docosahedral structures are in common usage in the literature and it is recommend that these should be formally adopted (Recommendations 19 and 20) [27]. Both structures, together with their accepted numbering patterns, are shown in Figure 3.

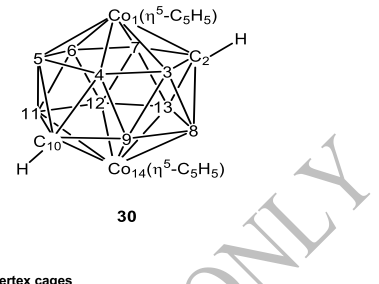
Thus $[1,2-\mu-(CH_2)_3]$ -3-Ph-*closo*-1,2-C₂B₁₁H₁₀ (**29**) with a henicosahedral structure may be named 3-phenyl-1,2-(propane-1,3-diyl)[C_s -($2v^4551$)- Δ^{20} -*closo*]-1,2-dicarbatridecaborane(13). It should be noted that this is an *ad hoc* solution that would not work for an unsymmetrical bridging *group e.g.* propane-1,2-diyl. This points to a need for a system for names for polyvalent substituent groups similar to the kappa system for ligand names *i.e.* where the bridging sites *(in casu* the atoms of the cage) can be specified individually for the various free valences. This will be the subject of further studies. Another possibility would be to extend organic fusion nomenclature to provide here the alternative name: 3-phenylcyclopenta[1,2-*b*][C_s -(2 v^4 551)- Δ^{20} -*closo*]-1,2dicarbatridecaborane(13). It should be further noted that this is also an *ad hoc* solution. In the general case, rules would have to be made for assigning letter identifiers (like '*a*' above) to all edges of the cage. This will also be the subject of further studies.

The anions with a docosahedral structure, $[B_{13}H_{13}]^{2-}$ and the 'carbons apart' metallacarbaborane $[4,4,4-(CO)_{3}-1,6,4-C_2ReB_{10}H_{12}]^-$, are named tridecahydrido $[C_{2\nu}-(1\nu^422\nu^6422)-\Delta^{22}-closo]$ tridecaborate(2-) and 4,4,4-tri(carbonyl)dodecahydrido $[C_{2\nu}-(1\nu^422\nu^6422)-\Delta^{22}-closo]-1,6$ -dicarba-4-rhenatridecaborate(1-), respectively.



The deltahedral structure calculated for the *closo* 14-vertex cage is the bicapped hexagonalantiprismatic tetracosahedron with 24 faces and D_{6d} symmetry. This structure has two BH units occupying the two cage six-connected vertices. It is recommended that the generally accepted numbering system for these tetracosahedral cages should be adopted (Recommendation 19, Figure 3) [28]. The tetracosahedron has the Powell descriptor [18] $[D_{6d}-(1v^6661v^6)-\Delta^{24}-closo]$. The largest known heteroborane to date, 2,3- μ -(CH₂)₃-2,3-*closo*-C₂B₁₂H₁₂ [22], has this structure, and can be named by bridging nomenclature as 2,3-(propane-1,3-diyl)[$D_{6d}-(1v^6661v^6)-\Delta^{24}-closo$]-2,3dicarbatetradecaborane(14), or by organic fusion nomenclature as cyclopenta[2,3-*b*][$D_{6d}-(1v^6661v^6)-\Delta^{24}-closo$]-2,3dicarbatetradecaborane(14).

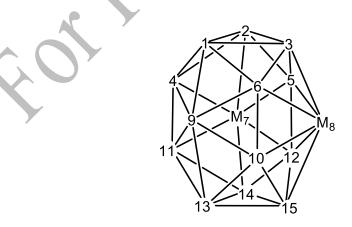
Closed isostructural 14-vertex {C₂M₂B₁₀} and {C₄M₂B₈} systems are known *e.g.* 1,14-(η^{5} -C₅H₅)₂-*closo*-2,10,1,14-C₂Co₂B₁₀H₁₂ (**30**), which is named 1,14-bis(η^{5} -cyclopentadienido)[*D_{6d}*-($1v^{6}$ 661 v^{6})- Δ^{24} -*closo*]-2,10-dicarba-1,14-dicobaltatetradecaborane(12).



BN-7.2.4 Closed 15-vertex cages

Recommendation 21: The non-standard numbering for pentacosahedral 15-vertex (Figure 3) species is well entrenched in the literature, and its continued use is recommended.

The calculated deltahedral structure for the 15-vertex [B₁₅H₁₅]²⁻ is the hexacosahedron (26 faces) with D_{3h} symmetry. This structure has been observed in species such as (η^5 -C5Me5Rh)2B13H13 [23], and 1,4-µ-(CH2)3-7-(n⁶-1-isopropyl-4-methylbenzene)-1,4,7-C2RuB12H12 [29], and the cage structure corresponds to a Powell descriptor [18] of $[D_{3h}-(333v^633)-\Delta^{26}-closo]$. The numbering scheme for these *closo* structures is shown in Figure 3 and $(\eta^5-C_5Me_5Rh)_2B_{13}H_{13}$ $[Rh(\eta^5 - C_5 Me_5)]),$ is where correctly named 7,8-bis(η^{5} -(31 М = as pentamethylcyclopentadienido)[D_{3h} -(333 v^{6} 33)- Δ^{26} -closo]-7,8-dirhodapentadecaborane(13) (Recommendation 19).

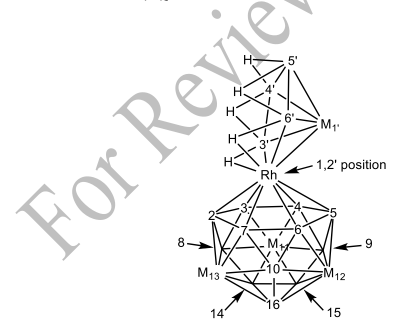


A second *closo* 15-vertex cage, $1-(\eta^6-1\text{-isopropyl-4-methylbenzene)-8,14-\mu-(CH_2)_3-8,14,1-C_2RuB_{12}H_{12}$, isomeric to that just mentioned, has a pentacosahedral (25-face) solid-state structure. This structure has a quadrilateral face and 24 triangulated faces. In solution it has effective mirrorplane symmetry (*C_s*) with a similar ($1\nu^6662$) plane arrangement, and so is numbered accordingly (Figure 3). It is recommended that this numbering scheme be retained for these *closo* pentacosahedral 15-vertex systems (Recommendation 21) [30].

BN-7.2.5 Closed 16-vertex cages



The calculated deltahedral structure for the 16-vertex closo- $[B_{16}H_{16}]^{2-}$ structure is an octacosahedron (28 faces) with T_d symmetry. This structure has been observed in a recently reported closed metallaborane cage containing a {Rh₄B₁₂} core [24] with a Powell descriptor [18] of [T_d -(1 v^6 633 v^6 3)- Δ^{28} -*closo*]. This is shown in Figure 3 together with the vertex numbering scheme (Recommendation 19). The compound containing this cage has a rhodium atom that is held in common with a *closo* 16-vertex and a *nido* six-vertex cage and the compound is a good example to illustrate the recommendation 19) nomenclature. Thus, the compound (η^5 -C₅Me₅Rh)₃B₁₂H₁₂Rh{ η^5 -C₅Me₅RhB₄H₉} (**32**, where M = [Rh(η^5 -C₅Me₅)]) is correctly named 1',11,12,13-tetrakis(pentamethylcyclopentadienido)[2',3',2',6':3',4':4',5':5',6'-penta- μ H-*nido*-1,2-dirhodahexaborane(9)]-2'-commo-1-[[T_d -(1 v^6 633 v^6 3)- Δ^{28} -closo]-1,11,12,13-tetrakis(cage)].



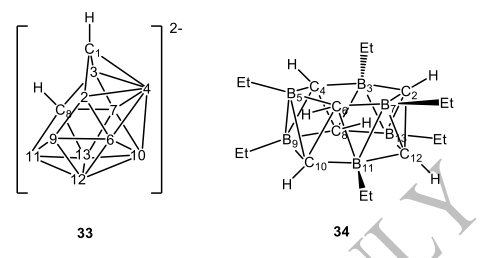
BN-7.2.6 Open supraicosahedral cages

Recommendation 22: The 'debor' method [2p, 19] is recommended for naming open supraicosahedral species.

Open (*i.e. nido* or *arachno*) structures have been reported for cages with 12-14 vertices which are based on closed supra-icosahedral polyhedra of 13-15 vertices with one or two vertices removed [21]. Open structures based on *closo* polyhedra with >15 vertices have not yet been synthesized, and are not considered further in this report.

It is expected that nido 12-vertex structures should be based on the closed 13-vertex shapes, with the highest connected vertex removed. The nido-[7,9-C2B10H12]²⁻ anion and the neutral molecule *nido*-Et₄C₄B₈H₈ have a geometry based on the docosahedral *closo* structure. This *nido* 12-vertex structure, together with alternative *nido* structures that have been observed, are shown in Figure 4. Synthesized nido 13-vertex and arachno 12-vertex cages are structurally derived from the *closo* 14-vertex tetracosahedral cage (bicapped hexagonal antiprism), with respectively one or two vertices missing. There are three different cage types reported for arachno 12-vertex systems. These structures are illustrated in Figure 4, and their shapes have been described as 'hexagonal antiprismatic', 'basket' shaped, and 'carbons adjacent with two open faces' [21]. The 'hexagonal antiprismatic' structure is related to the expected 14-vertex $[D_{6d}-(1v^6661v^6)-\Delta^{24}-closo]$ structure but the two vertices removed are both of cage-connectivity six and mutually antipodal rather than adjacent as is conventionally proposed for the generation of arachno structures. Similar examples in sub-icosahedral systems have been referred to as 'remote' arachno (see Section BN-7.3, also schematic cage structure 2), although this is not an IUPAC-accepted term. Structurally it could be interpreted as a bisnido species rather than the more open *arachno* cage. This may be a case where the electron count is the origin of the literature name rather than the observed structure, although, as recommended elsewhere in this document, structural nomenclature should be based on observed structure rather than perceived electronic considerations which may often be subjective. The 'carbons adjacent with two open faces' isomer is again derived from the deltahedral 14-vertex $[D_{6d}-(1v^{6}661v^{6})-\Delta^{24}-closo]$ structure but this time with the six-connected and a five-connected cage atom (from the lower belt, rather than the conventionally specified adjacent belt) removed. Again this might be viewed as a bisnido species rather than the more open arachno cage. The nido 13-vertex cage compound $[Ni(dppe)(Me_4C_4B_8H_8)]$ (dppe = Ph₂PCH₂CH₂PPh₂), which is also based on the expected bicapped hexagonal antiprism, has a 5-connected cage vertex removed, rather than the highest connectivity 6-connected cage vertex. This structure is illustrated in Figure 4.

The significant structural variations described in the last paragraph emphasize the need for a robust nomenclature system for such open cages. The 'debor' method [2p, 19] for describing open cages are part of IUPAC recommendations for the naming of open cages. This 'debor' method, combined with Powell descriptors [18] available for the related *closo* species from which the open structures are derived, is the preferred method of naming such open supra-icosahedral structures (Recommendation 22). In essence, the compound is named with the prefix *x*-debor, *x*,*y*-didebor or *x*,*y*,*z*,-tridebor (where *x*, *y* and *z* are locants) followed by the name of the closed structure ending with 'ane' (for a neutral borane) and the actual number of hydrogen atoms (in the unsubstituted parent) in parenthesis. Negatively charged species would need to specify hydrido positions, end with 'ate', and have the charge in parentheses.



Examples of nomenclature using the 'debor' method [2p, 19] are dodecahydrido-5-debor[$C_{2\nu}$ - $(1\nu^4 22\nu^6 422)-\Delta^{22}$ -closo]-1,8-dicarbatridecaborate(2–) for nido-[7,9-C₂B₁₀H₁₂]^{2–} (**33**) (note different numbering is used) and 3,5,7,9,11,13-hexaethyl-1,14-didebor[D_{6d} - $(1\nu^6 661\nu^6)-\Delta^{24}$ -closo]-2,4,6,8,10,12-hexacarbatetradecaborane(12) for H₆C₆B₆Et₆ (**34**). Here, the 'x-debor-' term is considered as part of the descriptor and immediately precedes it.

Overall, as more extensive supra-icosahedral chemistry is evolved, these recommendations may have to be revisited.

BN-7.3 Non-conventional deltahedral structures

Recommendation 23: The structural terms *isocloso* and *isonido* are recommended for nine-, ten- and twelve-vertex species, including their historical numbering schemes (Figure 5). Within eleven-vertex species the *isonido* descriptor with its associated numbering based on the *closo* eleven-vertex scheme (Figure 1) is also recommended (Figure 5).

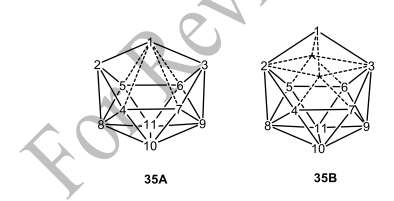
Recommendation 24: Closed geometries derived from polyhedra which are not within the conventional deltahedral set shown in Figure 1 should be described by descriptors based on the principles [2f, 18] as recommended for supra-icosahedral cages (Recommendation 19).

Recommendation 25: The 'debor' method [2p, 19] is recommended for 'non-standard' open structures, other than those highlighted in Recommendation 23.

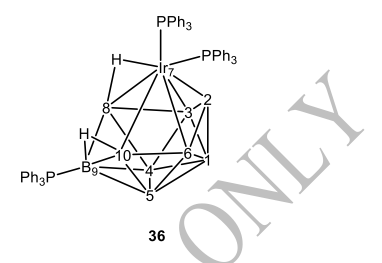
Terms such as *pseudocloso, quasicloso, hypercloso, precloso, oblato, isocloso, isonido* and *isoarachno* have also been used for the naming of cages, although they are not IUPAC recommended terms. These are discussed below. The situation is rapidly becoming confusing and unsystematic, as such terms are often introduced by authors to explain subtle structural and electronic nuances. However, it is recommended that structural descriptors in nomenclature must be based solely on the observed (idealized) structure.

The majority of known sub-icosahedral boron-hydride and boron-hydride-related species have cage geometries that have idealized structures adequately described by the structural descriptors *closo, nido,* and *arachno*, based on conventional deltahedral shapes (Figures 1 and 2; Section BN-3). The inclusion of the qualification 'idealized' indicates that not all interatomic distances and angles in subrogated boron hydrides are identical to those observed in the parent systems and due allowance must be made for this. Thus, for example, twelve-vertex metallacarbaborane derivatives such as $[3-(\eta^5-C_5Me_5)-closo-1,2,3-C_2IrB_9H_{11}]$ are clearly based on the closed deltahedral icosahedron, despite Ir-B and Ir-C interatomic distances being longer than the B-B distances observed in $[B_{12}H_{12}]^{2-}$ and C-C distances shorter. There will also be variations among the B-B distances. Such deviations from the ideal result primarily from changing atom (orbital) size and from antipodal and other directional effects within the cage. Nevertheless, the connectivities of the cage atoms and the idealized cage symmetries clearly warrant the use of the term *closo*.

Closed eleven-vertex systems such as $2,3-C_2B_9H_{11}$, $[2-CB_{10}H_{11}]^-$, and $[B_{11}H_{11}]^2$ have historically been named *closo* and assigned the octadecahedral structure shown in Figure 1. This structure has long interatomic distances between the boron atom at cage position 1 of formal cageconnectivity six and four of its neighbours in the 4, 5, 6 and 7 sites. Recent evidence, e.g. structural data (measured or calculated interatomic distances), chemical behaviour and, in some cases, fluxional behavior in solution, has been taken to indicate that they do not necessarily possess a formal closed deltahedral structure but the four long connectivities are essentially nonbonding so that the species have effectively two five-membered open faces reminiscent of a 'remote arachno' configuration (e.g. 35A and 35B) [31]. The term quasicloso has been coined for such structures. However, in essence they generally show (often time-averaged) connectivities of the closed eleven-vertex hexadecahedron ($C_{2\nu}$). It is therefore recommended that the term *closo*, with its associated numbering system, is used (Recommendation 23) for such structures and that the term *quasicloso* is not acceptable. Nevertheless, if an observed structure clearly has an open face (e.g. as determined by X-ray diffraction studies) then it should be named as such (nido or isonido). The term isocloso has also been used in certain compounds for the eleven-vertex closo structure, but such a description is based on perceived electronic structure rather than observed geometrical structure and for nomenclature this practice is not acceptable (see below).



Isocloso, isonido, and *isoarachno* have been used to describe sub-icosahedral cages with nonstandard connectivities or geometries. These may be based on unexpected closed polyhedra or have open structures based on conventional closed structures but with unusual (*i.e.* not highestconnected or non-adjacent) vertices removed. From a historical and structural point of view these terms have some merit in nomenclature provided they are not associated with electron-counting schemes. In practice, these alternative geometries have been limited to nine-, ten-, eleven- (see above for *isocloso*) and twelve-vertex species and it is recommended that the geometries shown in Figure 5 (and their associated numbering schemes) are retained (Recommendation 23). The *isonido* compound 7,7,9-(PPh)₃-*isonido*-7-IrB₉H₁₀ has an *exo* B-H hydrogen atom replaced by the neutral substituent PPh₃. This cage therefore needs to be named by the located hydrogen method (see Section BN-4.1) as 7,7,9-tris(triphenylphosphane- κP)-7:8,9:10-di- μH -9-dehydro-*isonido*-7-iridadecaborane(11) (**36**).



Using the 'debor' method [2p, 19] the *isonido* structure, 7,7,9-(PPh)₃-*isonido*-7-IrB₉H₁₀ (**36**) could equally be named as 1,1,9-tris(triphenylphosphane- κP)-9-dehydro-3-debor-1:7,6:9-di- $\mu H[C_{2\nu}-(12422)-\Delta^{18}-closo]$ -1-iridadecaborane(11), based on the *closo* structure as numbered in Figure 1, or **35A**. The 'debor' prefix is part of the cage name as are the 'indicated hydrogens'. The 'dehydro' prefix should precede the cage as an un-alphabetised detachable prefix. Should the need arise for new '*iso*' structures then the use of the 'planes' method [2f] for *closo* structures, and the 'debor' nomenclature [2p, 19] for open structures (Recommendations 24 and 25), as described in detail in the supra-icosahedral Section 7.2 should be used. This would also be an acceptable alternative nomenclature system for these nine- and ten-vertex *isonido* and *isocloso* systems. The *isocloso* ten-vertex system (Figure 5) could also be further defined as [$C_{3\nu}$ -($1\nu^{6}333$)- Δ^{16} -*closo*] if required.

Pseudocloso has been used as a descriptor for some twelve-vertex formally closed cages that contain some interatomic distances that are longer than expected. This is often held to be the result of severe steric congestion arising from substituents on neighbouring cage vertices or, in other cases, where there are no such steric constraints, from intracage electronic effects [32]. If it is accepted that such a structure is derived from an idealized *closo* compound, with interatomic distances still within an expected (perhaps on the long side) bonding range, then the compound should be named and numbered as *closo* and there is no need to introduce a new term *pseudocloso*. Authors may wish to bring attention to the so-called 'long' interatomic distances but it is recommended the term *closo* be used. If, however, the internuclear distances are such that they are outside normal bonding ranges, then structurally the compound is open and therefore needs to be named as such, *i.e. nido* (or *isonido*) and therefore the continued use of the term *pseudocloso* is no longer acceptable. As with eleven-vertex cages, the locant numbering for any *isonido* species derived by stretching the corresponding twelve-vertex *closo* should retain the conventional *closo* system.

As noted above, nomenclature is concerned with cage-atom connectivities and cage symmetry in the context of observed structures, and not concerned with perceptions about electronic

structure or bonding, save, in the last case, where an interatomic distance exceeds an accepted bonding range and thus engenders a more open cage type and thence a different structural descriptor. Wade's Rules [5] are extensively used to correlate skeletal bonding-electron counts with observed structure. In many cases, observed structures are in agreement with structures derived using Wade's Rules. However, conflict can arise if the perceived electron-count is at odds with the observed structure. In the event of such perceived conflict, it is recommended that authors should specifically state that the compound in question 'is described according to its observed structure, but that this is in conflict with the skeletal bonding-electron count as determined by and at this point the electron-counting procedure should be stated', or similar appropriate phraseology. The terms precloso and hypercloso have been used by authors to differentiate closed structures held to show the 'wrong' (unusual) rather than what may be proposed to be the 'correct' electron count according to Wade's rules. Similarly, some eleven-vertex structures with a regular standard *closo* structure have been described as 'isocloso', because of their unusual electron counts. As such, these terms are no longer acceptable and it is recommended that if the cage has the same symmetry and connectivity as the standard deltahedral structures shown in Figure 1 then it should be correctly named as *closo*, irrespective of perceived electron count.

BN-7.4 Metal-rich cages

There are many known cages which contain more metal atoms than boron atoms [33]. These cages are outside the scope of this document but will be studied in future projects dealing with metal clusters.

Membership of sponsoring bodies

Membership of the IUPAC Chemical Nomenclature and Structural Representation Division for the period 2016-2017 as follows: President: K.-H. Hellwich (Germany); Vice-President: A. Hutton (South Africa); Secretary: R. Laitinen (Finland); Titular Members: O. Achmatowicz (Poland), T. Damhus (Denmark), P. Hodge (UK), R. Macaluso (USA), J. Nagy (Hungary), M. Rogers (USA); J. Vohlidal (Czech Republic); Associate Members: M. Beckett (UK), I. Dukov (Bulgaria), G. Eller (Austria), E. Mansfield (USA), K. Taylor (USA), M. Strausbaugh (USA); National Representatives: F. Arico (Italy), A. Ferreira (Brazil), A. Fradet (France), H. Lee (Korea), T. Lowary (Canada), E. Nordlander (Sweden), M. Putala (Slovakia), A. Rauter (Portugal), J. van Lune (Netherlands), A. Yerin (Russia); Ex Officio: R. Hartshorn (New Zealand); G. Moss (UK).

Membership of the IUPAC Inorganic Chemistry Division for the period 2016-2017 as follows: President; J. Reedijk (Netherlands); Vice-President: L. Ohrstrom (Sweden); Secretary: M. Leskela (Finland); Titular Members: L. Armelao (Italy), T. Ding (China), P. Karen (Norway), R. Loss (Australia), D. Rabinovich (USA), T. Walczyk (Switzerland), M. Wieser (Canada); Associate Members: Y. Aziz (Malaysia), J. Colon (Puerto Rica), L. Meesuk (Thailand), K. Sakai (Japan), N. Trendafilova (Bulgaria); National Representatives: J. Darkwa (South Africa), M. Diop (France), J. Correia (Portugal), M. Hasegawa (Japan), S. Kalmykov (Russia), A. Kilic (Turkey), P. Knauth (France), J. Leigh (UK), S. Mathur (Germany), K. Yoon (Korea).

This manuscript was prepared in the framework of IUPAC project 2012-045-1-800.

REFERENCES

- [1] IUPAC. Nomenclature of Inorganic Chemistry, Recommendations 2005, N. G. Connelly, T. Damhus, R. M. Hartshorn, A. T. Hutton, RSC Publishing, Cambridge, UK (2005); (a) IR-6, p. 83-110; (b) IR-6.2.3.1, p. 89-90; (c) IR-9.2.4.2, p. 155-174; (d) IR-10.2.3.3, p. 210-211; (e) Table VI, p.260.
- [2] IUPAC. Nomenclature of Inorganic Chemistry, Recommendations 1990. Ed. G. J. Leigh, Blackwell Scientific Publications, Oxford, UK (1990); (a) I-11, p. 207-237; (b) p. xxxxi; (c) I-11.3.2.1 p. 217; (d) I-11.3.1.2, p. 211-214;.(e) I-11.3.2.5, p. 224;(f) I-11.3.2.3, p. 222-223;(g) I-11.3.1.2, p. 212-213;(h) I-11.3.2.4, p. 223-224;(i) I-11.4.2, p. 227-228; (j) I-11.4.3.1, p. 228; (k) I-11.4.3.3, p. 231; (l) I-11.4.3.2, p. 228; (m) I-11.4.3.3, note 110, p. 231; (n) I-11.5.2, p. 233; (o) I-11.3.12, p. 215-217; (p) I-11.3.2.7, p. 224-225.
- [3] R. M. Adams, Pure Appl. Chem. 30, (1972); (a) 683-710; (b) 697; (c) 693-694; (d) 698.
- [4] IUPAC. Nomenclature of Inorganic Chemistry Definitive Rules 1970 (2nd Ed), Butterworth and Co, London, (1971) Chapter 11, (a) 92-97; (b) p. 96; (c) p. 97; (d) p. 93-94.
- [5] K. Wade, Adv. Inorg. Chem. Radiochem. 18, 1 (1976).
- [6] (a) IUPAC. Nomenclature of Organic Chemistry. Recommendations and preferred names 2013, H. A. Favre, W. H. Powell, RSC Publishing, Cambridge, UK (2013), P-14.7.1;
 (b) W. H. Powell, F. Cozzi, G. P. Moss, C. Thilgen, R. J.-R. Hwu, A. Yerin, Pure Appl. Chem. 74, 629 (2002).
- [7] IUPAC. Compendium of Chemical Terminology. Recommendations, V. Gold, K. L. Loening, A. D. McNaught, P. Selimi, Blackwell Scientific Publication (1987), p 11.
- [8] IUPAC. Compendium of Chemical Terminology. Recommendations, (2nd Ed.), compiled by A. D. McNaught, A. Wilkinson, Blackwell Scientific Publication (1997) and XML on-line corrected version http://goldbook.iupac.org (2006) created by M. Nic, J. Jirat, B. Kosata, updates compiled by A. Jenkins.
- [9] IUPAC. Principles of Chemical Nomenclature, A Guide to IUPAC Recommendations 2011 Edition, Ed. G. J. Leigh, RSC Publishing, Cambridge UK (2011), Table 6.2, p. 63.
- [10] M. A. Beckett, N. N. Greenwood, J. D. Kennedy, M. Thornton-Pett, J. Chem. Soc., Dalton Trans. 1119 (1985).
- [11] J. P. Sheehan, T. R. Spalding, G. Ferguson, J. F. Gallagher, B. Kaitner, J. D. Kennedy, J. Chem. Soc., Dalton Trans. 35 (1993).
- [12] M. A. Beckett, J. E. Crook, N. N. Greenwood, J. D. Kennedy, J. Chem. Soc., Dalton Trans. 1879 (1986).
- [13] J. M. Oliva, J. Rué, D. Hnyk, J. D. Kennedy, V. R. Rosenfeld, Croatica Chemica Acta 86, 485 (2013).
- [14] Chemistry of the Elements, N. N. Greenwood, A. Earnshaw, Chapter 6, p. 177, Pergamon Press, 1984.
- [15] H-J. Yao, C-H. Hu, J. Sun, R-S. Jin, P-J. Zheng, J. Bould, R. Greatrex, J. D. Kennedy, D. Ormsby, M. Thornton-Pett. Collect. Czech. Chem. Commun. 64, 927 (1999).
- [16] J. Bould, W. Clegg, S. J. Teat, L. Barton, N. P. Rath, M. Thornton-Pett, J. D. Kennedy, Boron Chemistry at the Millennium, special edition of Inorg. Chim. Acta 289, 95 (1999).
- [17] D. L. V. K. Prasad, E. D. Jemmis, Phys. Rev. Lett. 100, 166504 (2008).
- [18] J. B. Casey, W. J. Evans, W. H. Powell, Inorg. Chem. 20, 1333 (1981).
- [19] J. B. Casey, W. J. Evans, W. H. Powell, Inorg. Chem. 22, 2236 (1983).
- [20] (a) L. D. Brown, W. N. Lipscomb, *Inorg. Chem.* 16, 2989 (1977); (b) J. Poater, M. Sola, C. Vinas, F. Teixidor, *Chemistry A European Journal* 22, 7347 (2016).
- [21] Carboranes, R. N. Grimes, 2011 (2nd Ed) Academic Press, Elsevier, Oxford, UK.
- [22] L. Deng, H.-S. Chan., Z. Xie, Angew. Chem. Int. Ed. 44, 2128 (2005).
- [23] D. K. Roy, B. Mondal, P. Shankhari, R. S. Angu, K. Geetharani, S. M. Mobin, S. Ghosh,

Inorg. Chem. 52, 6705 (2013).

- [24] D. K. Roy, S. K. Bose, R. S. Angu, B. Mondal, V. Ramkumar, S. Ghosh, Angew. Chem.Int. Ed. 52, 3222 (2013).
- [25] P. von Rague Schleyer, K. Najafian, A. M. Mebeb, Inorg. Chem. 37, 6765 (1998).
- [26] A. Burke, D. Ellis, B. T. Giles, B. E. Hodson, S. A. Macgregor, G. M. Rosair, A. J. Welch, Angew Chem. Int. Ed. 42, 225 (2003).
- [27] A. McAnaw, G. Scott, L. Elrick, G. M. Rosair, A. J. Welch, Dalton Trans. 42, 645 (2013).
- [28] A. McAnaw, M. E. Lopez, D. Ellis, G. M. Rosair, A. J. Welch, Dalton Trans. 42,671 (2013).
- [29] L. Deng, J. Zhang, H.-S. Chan, Z. Xie, Angew. Chem., Int. Ed. 45, 4309 (2006).
- [30] R. D. McIntosh, D. Ellis, G. M. Rosair, A. J. Welch, Angew. Chem. Int. Ed. 45, 4313 (2006).
- [31] E. G. Kononova, L. A. Leites, S. S. Bukalov, I. V. Pisareva, I. T. Chizhevsky, J. D. Kennedy, J. Bould, Eur. J. Inorg. Chem. 4911 (2007).
- [32] J. Bould, J. D. Kennedy, J. Organometal. Chem. 749, 163 (2014).
- [33] L. Barton, D. K. Srivastava, p. 275-372, Ch 8, Vol 1, in *Comprehensive Organometallic Chemistry II*, Vol. Ed., C. E. Housecroft, Eds in Chief E. W. Abel, F. G. A. Stone and G. Wilkinson, Pergamon, 1995.

Figure 1.

Conventional triangulated polyhedra with 4–12 vertices, illustrating *closo* framework cage geometries, and showing numbering conventions.

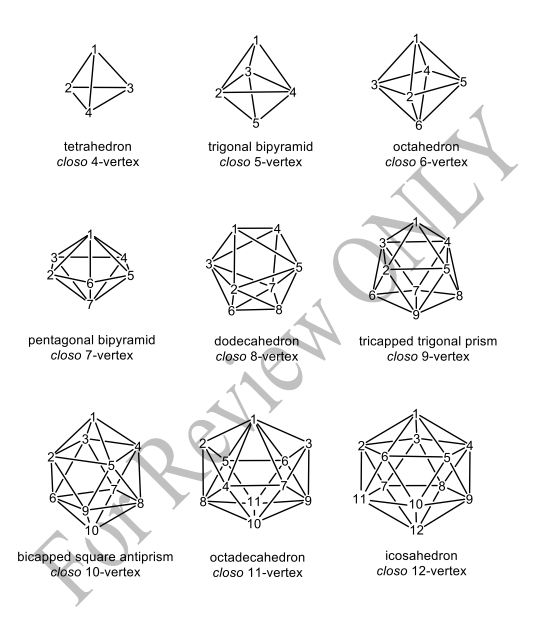


Figure 2.

Structures of commonly encountered classical Wadian *nido* and *arachno* polyhedral structures, illustrating their relationship with the *closo* structures of Figure 1 and showing numbering conventions. The *iso-arachno* 9-vertex structure is often referred to simply as *arachno*, and this is acceptable. Non-standard numbering denoted by *.

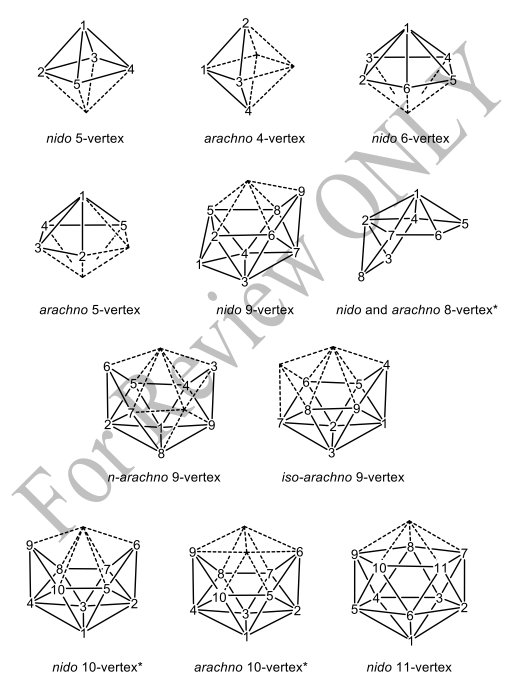


Figure 3.

Supra-icosahedral *closo*13–16 vertex structures with associated numbering schemes. The numbering schemes for 13- and 14-vertex species are well entrenched in the literature. 15- and 16-vertex structures follow conventional clockwise in planes numbering. The 13-vertex [C_s -(2551)- Δ^{20} -*closo*] and 15-vertex [C_s -(1662)- Δ^{24} -*closo*] contain quadrilateral faces and are *nido*-like but are named *closo*. Non-standard numbering denoted by *.

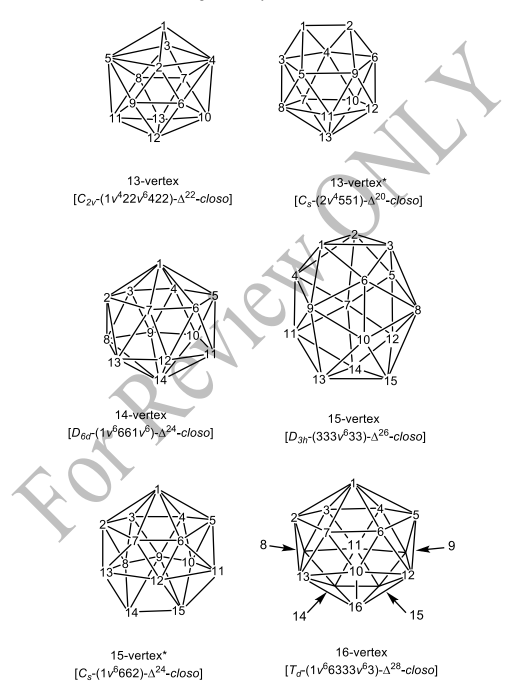


Figure 4.

12-vertex, *nido* and *arachno* and 13-vertex *nido* supra-icosahedral structures and relationships to higher *closo* geometries, shown in Figure 3.

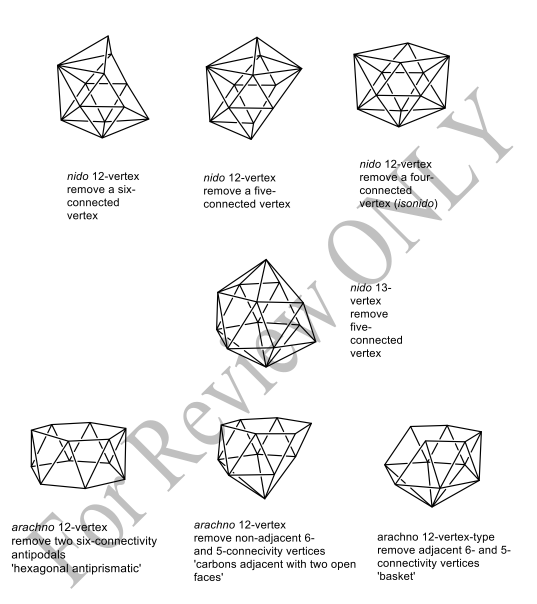


Figure 5.

9-, 10- and 12-vertex *isocloso* and ten-, eleven- and twelve-vertex *isonido* cage geometries with their associated numbering schemes. Although the twelve-vertex *isonido* and *isocloso* structures have their own idealized two-fold symmetries with a 2:2:4:2:2 stack structure, they are numbered by custom and practice as for conventional *closo*. The *isonido* eleven-vertex geometry also has an idealized two-fold symmetry, with a 1:2:2:2:2:2 stack structure, but by custom and practice is numbered as for conventional *closo*.

