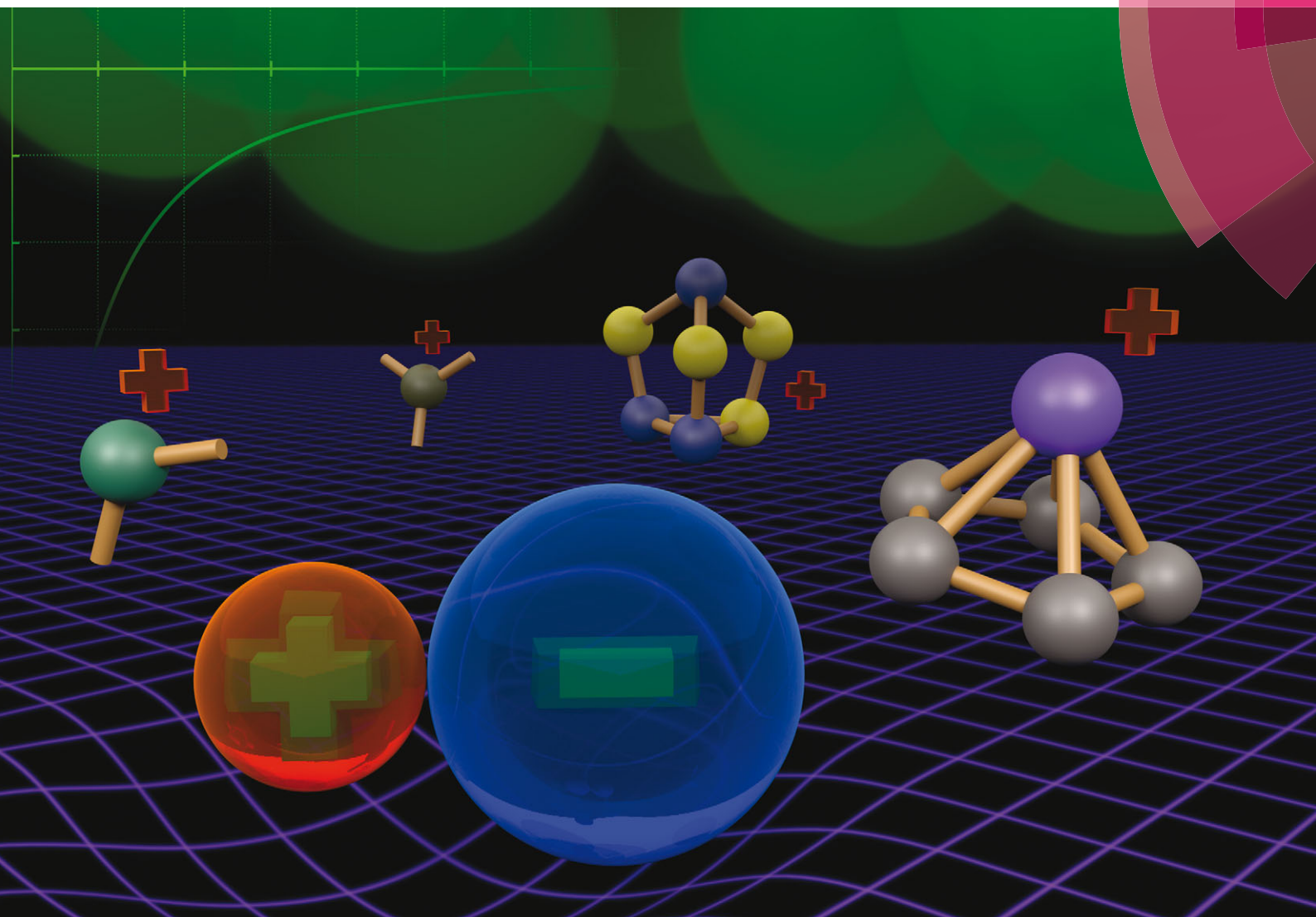


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Ingo Krossing *et al.*
Reactive p-block cations stabilized by weakly coordinating anions

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Reactive p-block cations stabilized by weakly coordinating anions

Tobias A. Engesser, Martin R. Lichtenhaler, Mario Schleep† and Ingo Krossing*

The chemistry of the p-block elements is a huge playground for fundamental and applied work. With their bonding from electron deficient to hypercoordinate and formally hypervalent, the p-block elements represent an area to find terra incognita. Often, the formation of cations that contain p-block elements as central ingredient is desired, for example to make a compound more Lewis acidic for an application or simply to prove an idea. This review has collected the reactive p-block cations (rPBC) with a comprehensive focus on those that have been published since the year 2000, but including the milestones and key citations of earlier work. We include an overview on the weakly coordinating anions (WCAs) used to stabilize the rPBC and give an overview to WCA selection, ionization strategies for rPBC-formation and finally list the rPBC ordered in their respective group from 13 to 18. However, typical, often more organic ion classes that constitute for example ionic liquids (imidazolium, ammonium, etc.) were omitted, as were those that do not fulfill the – naturally subjective – “reactive”-criterion of the rPBC. As a rule, we only included rPBC with crystal structure and only rarely refer to important cations published without crystal structure. This collection is intended for those who are simply interested what has been done or what is possible, as well as those who seek advice on preparative issues, up to people having a certain application in mind, where the knowledge on the existence of a rPBC that might play a role as an intermediate or active center may be useful.

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Introduction

Main group chemistry continues to reside at the heart of fundamental as well as applied chemistry. As such, recent years have seen an enormous growth of concepts that shed new light

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Tobias A. Engesser

On the basis of his interest for phosphorus compounds he stayed for three months in the group of Prof. Christopher “Kit” Cummins at the Massachusetts Institute of Technology in 2015.

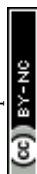
Dipl.-Chem. Tobias Engesser obtained his intermediate diploma at the Technische Universität Karlsruhe (now KIT) and continued his studies at the Albert-Ludwigs-Universität Freiburg, where he also wrote his diploma thesis about reactive tellurium cations and started his PhD in 2011 at the Institute of Inorganic Chemistry under the supervision of Prof. Ingo Krossing. His research interests are reactive phosphorus cations and gold(I) starting materials, both in



Martin R. Lichtenhaler

University of California, Berkeley as a postdoc in 2016. His research interests are organometallic and polymer chemistry, catalysis, molecular modelling and materials science.

Martin R. Lichtenhaler (1986) received his Diploma (2010) and PhD (2015) from the Albert-Ludwigs-Universität Freiburg, Germany. As a member of the Krossing group, he has developed a novel class of highly efficient, main group metal-based olefin polymerization catalysts and discovered unprecedented cationic clusters of univalent indium. After a research stay with the New Technology Office of Merck Ltd. Japan, he is likely to join the



on hitherto undiscovered, or more correctly, underdeveloped areas of main group chemistry. Thus, the availability of stable singlet carbenes¹ as strong donors offered tremendous new perspectives as did the establishment of the frustrated Lewis pairs (FLP) concept² or the systematic investigation of (often low valent) cationic mixed main group-transition metal salts.^{3–5} In the framework of those approaches, next to other fundamental⁶ and applied questions,^{7–10} also the stabilization or use of reactive p-block cations (rPBC) with weakly coordinating anions (WCAs) was one focus that led to fascinating new rPBC. This review gives a comprehensive overview on recent rPBC developments since about 2000, but also cites all-time classics in the field. It also includes the fascinating class of transition metal substituted rPBC for which the assignment of the positive charge to one specific moiety is often not clear.

Scope of this review

Many of the p-block elements have relatively high ionization potentials and electronegativities. Thus, most of the stable examples base on delocalization and other electronic or steric effects. In addition, rPBC are often very electrophilic and/or oxidizing. Therefore, chemically stable and inert weakly coordinating anions (WCAs) and solvents are needed to access their salts. These ingredients allowed the syntheses of a large number of fundamentally interesting rPBC of the groups 13–18 in the condensed phase. We discuss typical synthesis routes, give a brief overview of the WCAs, and describe the rPBC ordered according to their main group as well as cation class. However, typical, often more organic ion classes that constitute for example ionic liquids (imidazolium, ammonium, *etc.*) were omitted, as were those that do not fulfill the – naturally subjective – “reactive”-criterion of the rPBC. As a rule, we only included rPBC with crystal structure and only rarely refer to important cations published without crystal structure.

Handling of substance classes with recent reviews

Some of the substance classes, which fit into this review were just recently and sometimes very comprehensively reviewed (*cf.* our contribution describing the advances in the synthesis of homopolyatomic cations of the non-metals since 2000¹¹). To reduce the overlap, we decided to give an overview on general aspects such as WCAs in Table 1 and include a short table with relevant reviews for each main group at the beginning of each main group chapter and only list the compounds in these cases. Therefore, we mainly list, but do not describe the cations of this category in the chapters of their corresponding element. Nevertheless, the scope of this review is rather large, which in any case precludes extensive discussions and mainly serves as an overview on what is known.

Reactive p-block cations

The rPBC in this article need a WCA as counterion and, therefore, we first briefly describe typical WCAs and give some advice on their selection before turning to typical ionization and synthetic procedures for rPBC preparations. Thereafter, the ordering of the cation classes for the individual sections is described, and finally the rPBC are grouped according to the main group of the relevant cationic entry. In addition, first applications emerged for rPBC salts and will be highlighted in the respective cation sections.

WCA overview

Because of their potential in fundamental and applied chemistry,^{12–15} a great variety of different WCA types are currently known (Fig. 2) and was frequently reviewed (Table 1).

But which out of the multitude of published WCAs shown in Fig. 2 should be used for a given problem...? Is there one best WCA that fulfills all needs...?



Mario Schleep

Dipl.-Chem. Mario Schleep is a PhD student in the group of Prof. Ingo Krossing at the University of Freiburg, where he received his Diploma in 2012. During his studies, he has undertaken a research stay dedicated to poly-metallic chromium and vanadium clusters in the group of Prof. Eric McInnes at the University of Manchester (England) for two semesters. While dealing with electrolytes for lithium ion batteries during his thesis, his

current research focuses on the synthesis of reactive tin(II) cations stabilized by weakly coordinating anions.



Ingo Krossing

Ingo Krossing studied chemistry in Munich (LMU) and finished his PhD thesis 1997 (with Prof. H. Nöth). From 1997 to 1999, he worked as Feodor Lynen postdoc with Prof. J. Passmore at UNB, Canada. In 1999, he started his independent career as a Liebig- and DFG-Heisenberg-Fellow at the Universität Karlsruhe (TH) (mentor: Prof. H. Schnöckel). 2004 he changed as assistant professor to the Ecole Polytechnique Federale de Lausanne (EPFL),

before being appointed Chair of Inorganic Chemistry at the Albert-Ludwigs-Universität Freiburg in 2006. His research interests cover ionic systems from reactive cations to ionic liquids, as well as electrochemical energy storage. With an ongoing ERC Advanced Grant he develops absolute acidity and redox scales.



Table 1 General reviews with focus on WCAs

Year	Topic	Title	Ref.
1993	WCAs	The search for larger and more weakly coordinating anions	15
1998	WCAs	Carboranes: a new class of weakly coordinating anions for strong electrophiles, oxidants, and superacids	16
2004	WCAs	Noncoordinating anions—fact or fiction? A survey of likely candidates	13 and 12
2006	WCAs	Chemistry with weakly-coordinating fluorinated alkoxyaluminate anions: gas phase cations in condensed phases?	14, 15 and 17
2006	WCAs	Chemistry of the carba- <i>clos</i> -dodecaborate(−) anion, $[\text{CB}_{11}\text{H}_{12}]^-$	18
2008	WCAs	π -Complexation of post-transition metals by neutral aromatic hydrocarbons: the road from observations in the 19th century to new aspects of supramolecular chemistry	19
2013	WCAs	Weakly coordinating anions: halogenated borates and dodecaborates	20
2013	WCAs	Weakly coordinating anions: fluorinated alkoxyaluminates	21
2013	WCAs	Weakly coordinating anions: highly fluorinated borates	22
2015	WCAs	Taming the cationic beast: novel developments in the synthesis and application of weakly coordinating anions (Publication in progress by IK)	23

Clearly holds: the more reactive the rPBC are, the more demanding is the task for the anions, to meet the requirements for a successful stabilization in the condensed phase. Some of this reactivity may be dampened kinetically by the use of suitable bulky ligands, *e.g.* for the silylium ions. However, there is not one ultimate WCA that fulfills all requirements to allow for use with all in here described rPBC. Typically, rPBC follow at least one of the following classifications:

- Being a strong electrophile, thus having a strong tendency to coordinate an anion or solvent. Silylium ions SiR_3^+ are good examples for this. This coordination is often the entrance towards an anion degradation by heterolytic cleavage of a bond in the WCA.
- Being a strong oxidant, thus needing anions and solvents compatible with this need. Halogen and noble gas cations are typical examples.
- Being a weakly bound complex, in which the interesting main group particle can easily be displaced by anion or solvent, just as in many metal–non-metal clusters. This includes protonated, weakly basic molecules that tend to pass the proton to more basic and more coordinating anions or solvents.

Thus, the demand for very weak coordination behaviour is only medium for several very oxidizing cations, but the necessity of the WCA being stable against oxidation is a prerequisite of highest importance. For example, the typical counterions of group 16 to 18 rPBC are fluorometallates like $[\text{MF}_6]^-$ or $[\text{M}_2\text{F}_{11}]^-$ ($\text{M} = \text{As}, \text{Sb}$) compatible with (i) the oxidizing power of the cation and (ii) the typically used super acid solvents. However, despite the fact that fluoroantimonates allow for the synthesis of tremendously oxidizing cations like $[\text{Xe}_2]^+$, they fail to stabilize the extreme electrophiles $[\text{SiR}_3]^+$ and form F-SiR_3 and antimony pentafluoride. On the other hand, with some steric protection at the silylium ion, already the $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ WCA suffices to stabilize for example the $[\text{Si}(\text{Mes})_3]^+$ cation. By contrast, and due to the aromatic system, $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ is not compatible with the only mildly oxidizing $[\text{NO}]^+$ or $[\text{NO}_2]^+$ cation. Some thoughts that allow for the selection of a suitable WCA for a given problem may be summarized by the triangle shown in Fig. 1.

With Fig. 1 in mind, a personal selection of the “best WCAs” includes $[\text{1-H-CB}_{11}\text{Me}_5\text{Br}_6]^-$,²⁴ $[\text{1-Et-CB}_{11}\text{F}_{11}]^-$,²⁵ $[\text{CB}_{11}(\text{CF}_3)_{12}]^-$,²⁶ $[\text{Sb}_4\text{F}_{21}]^-$,²⁷ $[\text{Sb}(\text{OTeF}_5)_6]^-$,²⁸ $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$,^{29–31} $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ ^{32–34} and



Fig. 1 Triangle delineating the independent demands of a rPBC that lead to different mixtures of the WCA properties necessary for its successful stabilization.

$[\text{B}(\text{CF}_3)_4]^-$.³⁵ A recent noteworthy addition overcoming the frequent disorder of the also towards fluoride abstraction less stable $[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$ anion is the $[\text{B}(\text{Ar}^{\text{Cl}})_4]^-$ WCA.³⁶

Other aspects that will influence the choice, are the synthetic availability of the entire WCA class, or the specific starting material necessary to ionize the system of interest. In this respect, most of the WCAs known so far also do have disadvantages: the carborates are hard to synthesize and have often low yields. $[\text{CB}_{11}(\text{CF}_3)_{12}]^-$ is even explosive, as is the LiC_6F_5 intermediate needed for the $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ synthesis. In addition, starting materials such as solvent free Ag^+ salts or $[\text{NO}]^+$, $[\text{NO}_2]^+$ are not accessible as salts of $[\text{B}(\text{C}_6\text{F}_5)_4]^-$. Anions with multiple $-\text{CF}_3$ groups often tend to disorder in the solid state, which sometimes makes it hard to solve or refine the crystal structure. The problems associated with the refinement of structures containing the $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ WCA even led to the development of the software tool DSR.³⁷ It allows for the simplified refinement of such disordered structures and is now implemented with standard programs like OLEX2.³⁸

Therefore, the search for new useful anions is still in progress. With the ammoniated chloroborate cluster anion $[\text{1-Me}_3\text{N-B}_{12}\text{Cl}_{11}]^-$ another promising candidate that refined earlier ideas by S. Strauss *et al.*,³⁹ was just recently presented by Jenne *et al.* in 2014.⁴⁰ The positive charge of the ammonium function leads to an overall -1 charge and makes it possible to use the in 30 g scale accessible $-\text{B}_{12}\text{Cl}_{11}$ cluster residue. Important starting materials $\text{M}^+[\text{1-Me}_3\text{N-B}_{12}\text{Cl}_{11}]^-$ ($\text{M}^+ = \text{Na}^+$, $[\text{HNMe}_3]^+$, $[\text{HNOct}_3]^+$, $[\text{NO}]^+$, $[\text{CPh}_3]^+$, $[\text{N}^t\text{Bu}_4]^+$, $[\text{Et}_3\text{Si}]^+$) have been described facilitating the application.^{40,41} More details on typical WCA starting materials to introduce a counterion into the given system can be found in





Fig. 2 Some of the weakly coordinating anions discussed in this review.

the synthesis section below as well in the numerous WCA reviews cited in Table 1.

Synthesis routes to reactive main group cation salts

At the beginning, each proposal to prepare a target-rPBC needs to consider the choice of the WCA as delineated in the preceding section, as well as the available starting materials, ionization method and reaction medium.

WCA starting materials. A suitable starting material, should be accessible in good yields and contain a useful cation that typically acts as either a strong oxidant (*e.g.* $[\text{O}_2]^+$,⁴² $[\text{NO}]^+$,²⁹ $[\text{NO}_2]^+$,⁴³ $\text{N}(\text{arene})_3^+$,⁴⁴) a halide (*e.g.* Li^+ ,⁴⁵ Na^+ , Ag^+ ,⁴⁶), hydride- or alkyl-abstractor ($[\text{CPh}_3]^+$,⁴⁷), a Brønsted acid ($[\text{H}(\text{OEt}_2)_2]^+$,⁴⁸ $[\text{H}(\text{NMe}_2\text{Ph})]^+$) or a metal cation, if a simple metal complex is desired as product (*e.g.* $\text{Cu}^{+49,50}$) (Table 2). Neutral Lewis acids for bond heterolysis are available in great variety and include the classical simple halides $\text{M}^{\text{III}}\text{X}_3$ and $\text{M}^{\text{V}}\text{X}_5$ ($\text{M}^{\text{III}} = \text{B}, \text{Al}, \text{Ga}$; $\text{M}^{\text{V}} = \text{P}, \text{As}, \text{Sb}, \text{Bi}$; $\text{X} = \text{F}, \text{Cl}, \text{Br}, \text{I}$; not all combinations useful), the rather fine tunable $\text{B}(\text{aryl})_3$ acids (aryl = fluorinated,⁵¹ chlorinated⁵² or fluoroalkylated⁵³ aromatic residue), or aluminum based systems like $\text{Al}(\text{C}_6\text{F}_5)_3$,⁵⁴ and $\text{Al}(\text{OR}^{\text{F}})_3$.⁵⁵ Also the ion-like $\text{R}_3\text{Si}(\text{WCA})$ compounds have frequently been used.^{56,57} Recent systematic work analyzed the potency of a given Lewis acid *versus* fluoride, chloride, hydride and methanide as a base. It includes benchmark Lewis acidity values for a smaller set of simple MX_n acids.⁵⁸ Neutral Brønsted acids like HF, HNTf_2 and derivatives thereof,⁵⁹ or combinations of Brønsted and Lewis acids like $\text{HBr}/n\text{AlBr}_3$,^{60,61}

are suitable for protonations. Novel, and in large quantity available very strong acids like $\text{R}^{\text{HF}}\text{OSO}_3\text{H}$ ⁶² should also be mentioned.

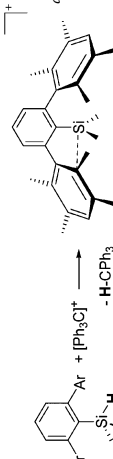
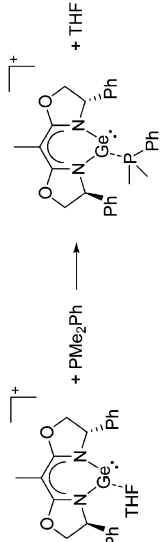
Suitable media/solvents. Since the synthesis of reactive ions is aspired, a suitable reaction medium should favorably be polar but not itself be a base or a nucleophile. This often rules out classical polar solvents that are itself good donors such as ethers or nitriles. Often chlorinated solvents CH_2Cl_2 ($\epsilon_r = 8.9$), 1,2- $\text{Cl}_2\text{C}_2\text{H}_4$ ($\epsilon_r = 10.4$) or Cl-Ph ($\epsilon_r = 5.7$) tend to be good choices that nevertheless are incompatible with strong electrophiles like the silylium ions $[\text{SiR}_3]^+$. Fluorinated arenes like F-Ph ($\epsilon_r = 5.5$) and 1,2- $\text{F}_2\text{C}_6\text{H}_4$ ($\epsilon_r = 13.4$) are good additions that became cheaper (but not cheap) over the last decade. However, they are incompatible with oxidants like $[\text{NO}]^+$ or $[\text{NO}_2]^+$ due to nitration/nitrosation reactions. Especially for non-metal cations, often superacids or SO_2 ($\epsilon_r = 16.3$), SO_2ClF ($\epsilon_r = \text{n.a.}$) *etc.* are the solvents of choice. ILs⁶³ like acidic $\text{BMIM}[\text{AlCl}_4]$ ^{64,65} and others were shown in recent years to be very promising media for rPBC cation synthesis.^{65–67} Especially for group 15 cations, solvent free reactions using $\text{Me}_3\text{Si-OSO}_2\text{CF}_3$ or MX_3 ($\text{M} = \text{Al}, \text{Ga}$; $\text{X} = \text{Cl}, \text{Br}$) were shown to provide quantitative yields of the desired salts. By contrast, several of such reactions do work only incomplete or not at all in solution.⁶⁸ Similarly, protonations with $\text{HBr}/n\text{AlBr}_3$ turned out to be best done solvent free.^{60,61}

The recently established concepts of absolute acidity,⁶⁹ absolute reductivity⁷⁰ and their two-dimensional combination as the protoelectric potential map⁷⁰ can be used to understand protonation and/or redox chemistry over medium/solvent and





Table 2 Acronym (Acr.) and type of the classified synthesis routes leading to rPBC

Acr.	Type	Example	Ref.
Com	Complexation reaction	$\text{Ag}^+ + \xrightarrow{2 \text{ P}_4} \left[\begin{array}{c} \text{P} \quad \text{P} \\ \diagdown \quad \diagup \\ \text{Ag} \\ \diagup \quad \diagdown \\ \text{P} \quad \text{P} \end{array} \right]^+$	72
Ox	Oxidation reaction; including $1e^-$ and $2e^-$ oxidations.	$\text{Ag}^+_{(\text{solv})} + \text{Ga}^0_{(\text{s})} \xrightarrow{1 e^- \text{ Ox.}} \text{Ga(arene)}^+_{(\text{solv})} + \text{Ag}^0_{(\text{s})}$ $\text{S}_8 + 3 \text{ AsF}_5 \xrightarrow{2 e^- \text{ Ox.}} \text{S}_8^{2+}([\text{AsF}_6^-])_2 + \text{AsF}_3$	30 and 31
Lewis	Lewis acid induced halogen bond heterolysis with neutral Lewis acids, including ion-like compounds.	$\text{Mes}_2\text{B-F} + \xrightarrow{\text{Et}_3\text{Si}(\text{HCB}_{11}\text{Cl}_{11})} [\text{Mes}_2\text{B}]^+[\text{HCB}_{11}\text{Cl}_{11}]^- + \text{Et}_3\text{Si-F}$	32, 33 and 73
Salt	Salt elimination reaction	$[\text{BCl}_3(\text{NHC})] + \text{Na}^+ \longrightarrow [\text{BCl}_2(\text{NHC})]^+ + \text{NaCl}$	74
Hyd	Hydride metathesis reaction with neutral or ionic H^- -acceptor		76
Alk	Alkyl metathesis reaction with neutral or ionic R^- -acceptor	$\text{AlMe}_3 + \text{B}(\text{C}_{12}\text{F}_9)_3 \xrightarrow{\text{in Et}_2\text{O}} [\text{Me}_2\text{Al}(\text{OEt}_2)]^+ [\text{MeB}(\text{C}_{12}\text{F}_9)_3]^-$	77
Ins	Insertion reaction	$\text{P}_4 + \text{NO}^+ \longrightarrow [\text{P}_4\text{NO}]^+$	78
Prot	Protonation reaction	$\text{AlCp}_3 + [\text{H}(\text{OEt}_2)_2]^+ \xrightarrow{-2 \text{ Et}_2\text{O}} [\text{AlCp}_2]^+ + \text{Cp-H}$ $[\text{Al}(\text{OR}^{\text{PF}})_4]^- \xrightarrow{[\text{Al}(\text{OR}^{\text{PF}})_4]^-}$	79
Lig	Ligand exchange reaction		80
Ion	Ionization	$\text{Pr}_3\text{Si}(\text{HCB}_{11}\text{Cl}_{11}) + \text{C}_6\text{H}_4\text{Cl}_2 \longrightarrow [\text{Pr}_3\text{Si}(\text{HCB}_{11}\text{Cl}_{11})]^+ [\text{HCB}_{11}\text{Cl}_{11}]^-$	81
Other	Other reaction not classified as one of the above	—	—

^a This type of reaction is sometimes referred to as Bartlett-Condon-Schneider (BCS) type hydride transfer reaction.⁷⁵

even phase boundaries. This also includes ILs and therefore a thermodynamically sound pH definition has been introduced for IL media.^{61,71}

Ionization protocols. Overall, we have categorized the rPBC included with the tables in the following sections by an acronym describing the synthetic approach used for their preparation. The synthesis routes are collected, explained and abbreviated in Table 2. Almost all the early approaches to reactive main group cations used halide abstractors as the Lewis acids AsF_5 or SbF_5 , which form the conjugated $[\text{AsF}_6]^-$ or $[\text{SbF}_6]^-$ WCAs through the reaction. The trityl cation is a hydride abstractor, which is especially in case of silanes as starting materials very useful to produce silylium cations. Most of the metal–non-metal complexes were synthesized by complexation of a non-metal molecule (*e.g.* P_4 , S_8 , Cl_2 , Xe) with a metal salt of a WCA. The coinage metals Cu^I , Ag^I and Au^I with their d^{10} electron configuration induce positive charge on the main group elements, stabilize the almost undistorted non-metal clusters, and provide insights in their bonding situation. If the cation is a strong oxidant, it is also possible to oxidize neutral substrates directly to give reactive cations, which are in turn stabilized by the corresponding WCA. An interesting recent addition are the transfer oxidation of *e.g.* the simple diorganodichalcogenides R_2E_2 ($\text{E} = \text{S}, \text{Se}$) with the combination of XeF_2 (primary oxidant and source of fluoride) and a Lewis acid.⁸²

On the representations of the cation chemical structures

Note that the structural diagrams used throughout this review obey a distance criterion for interactions, but not necessarily a 2e2c-understanding of every interaction line. However, at least in the organic residue we attempted to follow an electron precise 2e2c picture. Necessarily, this gets difficult for structures with N-heterocyclic carbenes that formally allow for a neutral dative (imidazolyl) as well as ionic (imidazolium) description (Fig. 3).

For simplicity, we chose the representation shown in the box in Fig. 3 and adopted similar drawings for related cases throughout. Thus, we only use arrows for relatively weak interactions with the bonding situation in ammine-borane $\text{H}_3\text{B} \leftarrow \text{:NH}_3$ being the prototype as suggested by A. Haaland,⁸³ and later contributions.⁸⁴ For thoughts on these ongoing discussions, see these recent publications.⁸⁵ Only if the positive charge can clearly (and not just

formally) be attributed to one atom, we assigned the charge to this atom. More common is the case in the box in Fig. 3, in which the charge may be delocalized to quite a series of atoms and therefore we placed the charge at the upper right corner.

Ordering of the cation classes

The rPBC were as far as possible ordered according to accepted cation classes that may either refer to the number of valence electrons (*i.e.* the onium/enium/inium-series) or to the structure. In each subchapter, we intend to go from homoatomic, to binary and then to more complex cation compositions. The not always consistently used classification according to onium- (8 VE), enium- (6 VE) and inium-cations (4 VE) presents some problems. Note, that the coordination number of a group 14 onium ion may not always be four, as the σ -donation of π -density of a donating double bond may increase the coordination number to 5 as, for example, in the 2-norbornyl cation, a carbonium ion.⁸⁶ Similar considerations hold for other donor coordinated onium- and enium-ions. Thus, we typically include the coordination number in the cation classification, for example as ligand substituted, (CN = 2). By contrast to these cation assignments, the group 15 to 18 cations were in addition classified by the oxidation state of the central atoms. This is often used for such rPBC. In addition, we included ion-like compounds that were initially defined for silylium compounds with coordinated counterions that structurally have to be addressed as a tight ion-pair but from the reactivity still bear a considerable amount of reactivity related to the free cation, *e.g.* see the ion like silylium compounds $\text{R}_3\text{Si}(\text{WCA})$ in Fig. 4. Related cases were published for coordinated aluminum cations, *e.g.* $\text{R}_2\text{Al}(\text{WCA})$, and were used in a similar manner.⁸⁷ For ion-like compounds we keep the notation with the anion in parentheses and no charges written, as in $\text{R}_3\text{Si}(\text{WCA})$ and $\text{R}_2\text{Al}(\text{WCA})$. Heteropolyatomic clusters were discussed in the group of their most electropositive element (*e.g.* $[\text{P}_3\text{Se}_4]^+$ in group 15, but $[\text{S}_4\text{N}_4]^{2+}$ in group 16).

In the following chapters we describe the rPBCs of the Group 13 to 18 elements and give selected representative examples for each cation type. However, for reasons of legibility, the full tables that comprehensively cover the rPBC entries of the groups, are collected in landscape format at the end of this document.

Group 13 cations

Traditionally, group 13 chemistry is dominated by compounds in the +III oxidation state.⁸⁸ Of those, the simple halides are commonly applied as Lewis acid catalysts and initiators (*e.g.* BF_3) and usually associated with anion formation (*e.g.* $[\text{BF}_4]^-$). However, discrete trivalent group 13 cations have been found to be more reactive, owing to their greater electrophilicity if paired with coordinative unsaturation.^{10,89} Except for boron, it has become increasingly



Fig. 3 Possible descriptions of NHC-containing structures exemplified for the simple $[\text{Cl}_2\text{B}-\text{i}^t\text{Bu}]^+$ cation.

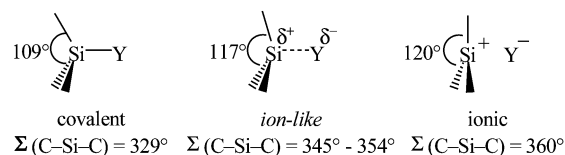


Fig. 4 Definition of ion-like compounds as exemplified for silylium ions.



Table 3 Review articles including cationic group 13 compounds

Year	Group	Title	Ref.
1985	13	Arene complexes of univalent gallium, indium and thallium	100 and 101
1998	13	Cationic group 13 complexes	102
2004	13	From group 13-group 13 donor-acceptor bonds to triple-decker cations	94
2005	13	Borinium, borenium, and boronium ions: synthesis, reactivity, and applications	89
2007	13	Development of the chemistry of indium in formal oxidation states lower than +III	103
2008	13	Borylene transfer from transition metal borylene complexes	13 and 12
2008	13	Synthesis, characterization, and applications of group 13 cationic compounds	95
2009	13	Highly electrophilic main group compounds: ether and arene thallium and zinc complexes	90
2009	13	Transition metal borylene complexes: boron analogues of classical organometallic systems	104
2010	13	Electron-precise coordination modes of boron-centered ligands	105
2011	13	Coordination chemistry of group 13 monohalides	96
2011	13	New light on the chemistry of the group 13 metals	88
2011	13	The chemistry of the group 13 metals in the +I oxidation state	106
2011	13	Mixed or intermediate valence group 13 metal compounds	107
2011	13	Coordination and solution chemistry of the metals: biological, medical and environmental relevance	108
2012	13	Cationic tricoordinate boron intermediates: borenium chemistry from the organic perspective	109
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possible to stabilize group 13 cations in their +I oxidation state, *e.g.* by employing bulky substituents and/or WCAs^{31,90–92} (for thallium, this is the favored oxidation state due to the inert pair effect⁹³). Featuring a lone pair of electrons and empty p-orbitals, the +I cations are ambiphilic and can function both as Lewis base or acid, thus offering unique reactivities and selectivities in organometallic chemistry,^{94–97} as well as organic⁹⁸ and polymer^{7,99} syntheses. Overall, different aspects of the chemistry of cationic group 13 compounds were reviewed and these contributions are compiled in Table 3. In this context, this section intends to give a comprehensive overview of reactive group 13 cations of the larger WCAs since about 2000. Due to the large scope of this chapter, we mainly omit rPBC with the simple halometallate based counterions and only include those in special cases of high relevance.

Boron cations

For a long time, boron cations have remained a chemical curiosity due to their redox lability. However and partly owing to the developments in the field of WCAs, more and more boron-based cations are being reported. Overall, the cations can be classified according to the coordination number at boron: *i.e.*, di-, tri-, and tetra-coordinated boron cations are referred to as borinium, borenium and boronium cations. To this day, the boron cations have been most notably reviewed by Nöth (1985; a milestone in cationic boron chemistry),¹¹² Piers (2005; structural and bonding aspects)⁸⁹ and Vedejs (2012; reactivities and applications).¹⁰⁹

Alkyl-/aryl substituted (CN = 2). To our knowledge, there is only one contribution to this class of compounds: *i.e.*, the recently reported [Mes₂B]⁺ borinium cation with the very good [HCB₁₁Cl₁₁][−]/[B(C₆F₅)₄][−] WCAs.^{32,113} Herein, the boron atom adopts a linear di-coordinated structure and the Mes substituents are aligned orthogonal to each other, allowing for a perfect shielding as well as π -donation into the empty p-orbitals of the highly

electrophilic borinium cation (*cf.* the modelled delocalized molecular orbitals). The [Mes₂B]⁺ cation is likely to become a textbook compound as it is the first borinium cation that does not rely on strongly π -donating heteroatom substituents (*cf.* the earlier reported [(^tBu₃PN)₂B]⁺ cation¹¹⁴ in the section ligand substituted (CN = 2) in Table 8).

Ligand substituted (CN = 3). Tricoordinate borenium cations are not as electron deficient as the borinium cations and therefore more stable. Nonetheless, the cations can only be isolated in the solid state if chelating (*e.g.* phthalocyanine¹¹⁵ and catecholborane^{116,117}) or strongly σ -donating ligands (*e.g.* N-heterocyclic carbene I^tBu⁷⁴ or hexaphenylcarbodiphosphorane^{118,119}) are applied. For the synthesis and reactivity of the [BCl₂(I^tBu)]⁺ cation, the nature of the WCA is crucial. Hence and though the cation can be prepared in the presence of [AlCl₄][−], [OTf][−] or [B(Ar^{Cl})₄][−], only the latter allows for a structure with no notable cation–anion contact. This leads to an increased reactivity of the [BCl₂(I^tBu)]⁺[B(Ar^{Cl})₄][−] salt.⁷⁴

Ligand substituted (CN = 4). Some of the tetra-coordinated boronium cations directly derive from the corresponding borenium cations: *i.e.*, the tricoordinate [PMAF–9BBN]⁺ cation (only stable in solution, as monitored by ¹¹B NMR spectroscopy) reacts with 1-MIM in a ligand exchange and addition reaction to form the tetra-coordinated [(1-MIM)₂(9BBN)]⁺ cation (Fig. 5).²¹²

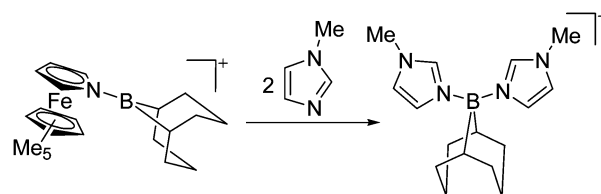


Fig. 5 Synthesis of the [(1-MIM)₂(9BBN)]⁺ boronium cation *via* a borenium cation precursor.





Fig. 6 (a) Orbital interaction between borylenes and transition-metal fragments; (b) and (c) exemplarily selected transition-metal substituted borinium cations.

For the isolation of the discrete $[\text{BH}_2(\text{PR}_2\text{H})_2]^+$ cation, the nature of the WCA is again essential: compared to $[\text{OTf}]^-$, $[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$ features no hydrogen bond with the cation, thus allowing for increased reactivities.¹²⁰

Transition-metal substituted. The number of transition-metal substituted boron cations is much higher than the one of related alkyl-/aryl- or heteroatom substituted compounds. Numerous contributions have been made by Braunschweig and Aldridge and both authors recently reviewed the chemistry of transition-metal borylene complexes.^{3,5,104} The d-orbitals of the transition-metals allow for stabilizing σ - and π -interactions with the orbitals of boron (Fig. 6) and of all the ligands the $\text{FeCp}(\text{CO})_2/\text{FeCp}'(\text{CO})_2/\text{FeCp}^*(\text{CO})_2$ substituents protrude: *e.g.*, various linear borinium cations, such as $[\text{CpFe}(\text{CO})_2\text{B}(\text{NCy}_2)]^+$, and borenium cations derived thereof, such as $[(\text{CpFe}(\text{CO})_2)\text{B}(\text{NCy}_2)-(4\text{-Pic})]^+$, have been isolated.¹²

Another notable substance class are the cationic T-shaped platinum boryl complexes that are usually accessible *via* salt metathesis reactions: *e.g.*, $[(\text{Cy}_3\text{P})_2(\text{MeCN})\text{Pt}(\text{B}\equiv\text{O})]^+$ can be synthesized by reacting $(\text{Cy}_3\text{P})_2\text{Pt}(\text{B}\equiv\text{O})(\text{Br})$ with the halide abstracting reagent $\text{Ag}^+[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$.¹²¹ Employing a ferrocenyl ligand on the other hand, Braunschweig *et al.* were able to isolate a rare example of a structurally characterized boron dication: $[\text{FcB}(\text{Pic})_3]^{2+}$ (Fig. 7).¹²²

Multinuclear. Due to the pronounced electron deficiency of boron there are not many contributions to the field of cationic multinuclear boron-based rPBC. The neutral diborane $[\text{HB}(\mu\text{-hpp})_2]$ complex however, is an excellent precursor for hydride abstractions and *via* unexpected boron-boron coupling reactions the unprecedented tetraborane dication $[\text{B}_4\text{H}_2(\mu\text{-hpp})_4]^{2+}$ was isolated (Fig. 8).¹²³

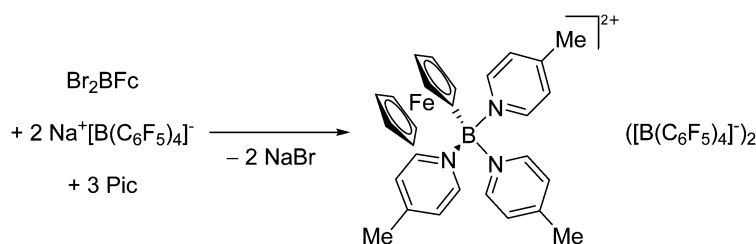


Fig. 7 Synthesis of the boron dication $[\text{FcB}(\text{Pic})_3]^{2+}$ *via* bromide abstraction and subsequent complexation.



Fig. 8 The tetraborane dication $[\text{B}_4\text{H}_2(\mu\text{-hpp})_4]^{2+}$. The bonding properties in the rhomboid B_4 core of the product can be described as two B–B units connected by 3c-2e bonds, sharing a short diagonal.

Multinuclear transition-metal substituted. Compared to their mononuclear congeners, both the ligands and coordination modes in the multinuclear borinium, borenium and boronium cations are very similar: (i) linear in the $[(\text{CpFe}(\text{CO})\{\text{B}(\text{NCy}_2)\})_2(\mu\text{-dmpe})]^{2+}$ complex,¹²⁴ (ii) trigonal-planar in the $[(\text{Cy}_3\text{P})_2\{\text{Pt}(\text{B}(\text{Br}))\}_2(\mu\text{-Ph})]^{2+}$ dication¹²⁵ and (iii) tetrahedral in $[(\text{bipy})(\text{Me})\text{B}]_2(\mu\text{-Fc})]^{2+}$.^{126,127} The aggregation usually occurs *via* bi-functional ligands like dmpe or *via* the transition-metal ligand itself (Fig. 9).

Aluminum cations

Among the group 13 cations, the lower- and higher-coordinated derivatives of aluminum have been of significant interest as they feature increased Lewis acidities and ligand labilities, thus allowing for higher catalytic activities compared to their neutral analogs.⁹⁵ While Atwood (1998)¹⁰² and Dagorne (2008)⁹⁵ have given a good overview on cationic aluminum species from a fundamental perspective, Sarazin and Carpentier (2015)¹⁰ recently

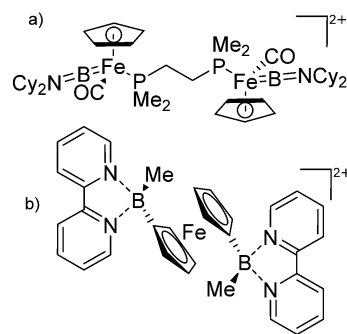


Fig. 9 Dicationic (a) $[(\text{CpFe}(\text{CO})\{\text{B}(\text{NCy}_2)\})_2(\mu\text{-dmpe})]^{2+}$ ¹²⁴ and (b) $[(\text{bipy})(\text{Me})\text{B}]_2(\mu\text{-Fc})]^{2+}$ ^{126,127} complexes.



Fig. 10 The ion-like $(\text{Me}_2\text{Al})_2\text{B}_{12}\text{Cl}_{12}$ salt.⁸⁷ For clarity, all BCl moieties of the perchlorinated *closio*-dodecaborate that feature no contact to the “[Me_2Al]⁺” cation have been omitted.

reviewed various discrete cationic aluminum complexes that are able to catalyze ring-opening polymerizations.

Alkyl or aryl substituted. The synthesis of di-coordinated alkyl complexes of aluminum $[\text{R}_2\text{Al}]^+$ ($\text{R} = \text{Me}, \text{Et}, 2,6\text{-Mes}_2\text{C}_6\text{H}_3$) is only viable, if extremely weakly coordinating anions (e.g. borate⁸⁷ and carboranes¹²⁸) and/or bulky substituents¹²⁹ are applied. In the case of “[Me_2Al]⁺” and “[Et_2Al]⁺”, the Lewis acidity of the aluminum cations is so significant that the latter feature distinct contacts to the corresponding WCAs and should therefore be described as ion-like compounds (Fig. 10). However, preliminary investigations showed that ion-like $(\text{Et}_2\text{Al})_2\text{B}_{12}\text{Cl}_{12}$ is a very active initiator for the cationic polymerization of isobutylene.¹³⁰

The $[(2,6\text{-Mes}_2\text{C}_6\text{H}_3)_2\text{Al}]^+$ cation on the other hand, is a discrete and therefore almost linear di-coordinate aluminum cation that features no contact to the WCA $[\text{B}(\text{C}_6\text{F}_5)_4]^-$.¹²⁹ The occurrence of the highly Lewis acidic aluminum cation is attributable to the intrinsic stabilization effect of the 2,6- $\text{Mes}_2\text{C}_6\text{H}_3$ ligand: *i.e.*, bending of the flanking Mes-moieties towards the aluminum center.

Cyclopentadienyl complexed. This class of compounds is to some extent related to the just mentioned alkyl substituted $[(2,6\text{-Mes}_2\text{C}_6\text{H}_3)_2\text{Al}]^+$ complex. Hence, the Cp ligands are η^5 - but not σ -bonding, and allow for the synthesis of discrete aluminum cations with different WCAs as counterions: $[(\eta^5\text{-Cp})_2\text{Al}]^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$,⁷⁹ $[(\eta^5\text{-Cp}')_2\text{Al}]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$ ¹³¹ and $[(\eta^5\text{-Cp}^*)_2\text{Al}]^+[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$.^{132,133} Moreover, the salts offer insights into the relationship between the nucleophilicity of Cp, Cp' and Cp*, the corresponding WCAs and the resultant Lewis acidities and reactivities of the aluminum cations: *i.e.*, with increasing nucleophilicity of the Cp ligands ($\text{Cp} < \text{Cp}' < \text{Cp}^*$) the WCAs can be less coordinating $[\text{MeB}(\text{C}_6\text{F}_5)_3]^- > [\text{B}(\text{C}_6\text{F}_5)_4]^- > [\text{Al}(\text{OR}^{\text{PF}})_4]^-$. The more interacting anions induce decreased Lewis acidities and lower reactivities of the aluminum cations for the initiation of olefin polymerizations: $[(\eta^5\text{-Cp})_2\text{Al}]^+ > [(\eta^5\text{-Cp}')_2\text{Al}]^+ > [(\eta^5\text{-Cp}^*)_2\text{Al}]^+$.⁷⁹

Ligand substituted (CN = 2). The above mentioned σ -coordinated $[\text{R}_2\text{Al}]^+$ complexes are either stabilized by intermolecular interactions with the corresponding WCAs or intramolecularly by two bulky terphenyl ligands. Within this context, Sekiguchi *et al.* were able to contribute another cationic di-coordinated, yet differently intramolecularly stabilized aluminum species: the $[\text{tBu}_2\text{MeSi-Al-Si}^t\text{Bu}_2\text{-SiMe}^t\text{Bu}_2]^+$ cation.¹³⁴ As supported by the solid-state structure and theoretical calculations, the stabilizing element is a σ - π hyperconjugation of the aluminum cation and the neighboring Si-Si σ bond (Fig. 11).

Ligand substituted (CN = 3). Tricoordinate aluminum cations are a bit less electrophilic than their di-coordinated congeners

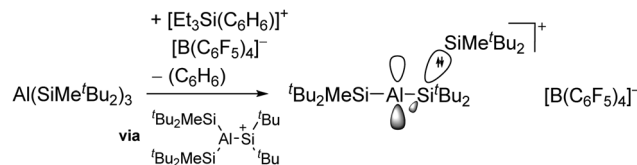


Fig. 11 Synthesis of the $[\text{tBu}_2\text{MeSi-Al-Si}^t\text{Bu}_2\text{-MeSi}^t\text{Bu}_2]^+$ cation via demethylation and subsequent migration of a $^t\text{Bu}_2\text{MeSi}$ group.

but nevertheless still very reactive. The few examples that have been reported, require chelating and sterically demanding β -diketiminato ligands, thus allowing for the successful synthesis of cationic $[(\beta\text{-diketiminato})\text{Al-H}]^+$ ¹³⁵ and $[(\beta\text{-diketiminato})\text{Al-Me}]^+$ ¹³⁶ complexes, respectively (Fig. 12).

Ligand substituted (CN = 4). In their recent review on group 13 cations, Dagorne and Atwood state that “*four-coordinate cations are most common ... as they incorporate an electronically saturated metal center*”.⁹⁵ In all compounds the aluminum cations are coordinated in a tetrahedral fashion with at least one coordination site being occupied by a heteroatom (N, O, P). Moreover, the vast majority of aluminum cations are incorporated into heterocycles, which derive from chelating ligands, such as Pytsi,¹³⁷ hpp,¹³⁸ BOX,¹³⁹ $^i\text{Pr}_2\text{-ATI}$ ^{140,141} and SchNMe_2 .¹⁴² The usual synthesis routes are alkyl or hydride abstractions. On the other hand, there are a few examples where aluminum is coordinated by four discrete ligands: $[\text{Me}_2\text{Al}(\text{OEt}_2)_2]^+$,⁷⁷ $[\text{Me}_2\text{Al}(\text{THF})_2]^+$,¹⁴³ $[\text{Me}_2\text{Al}(\text{NPhMe}_2)_2]^+$ ¹⁴⁴ and $[\text{H}_2\text{Al}(\text{NMe}_3)_2]^+$ ¹⁴⁵ (cf. Fig. 13 for the complex synthesis of the $[\text{H}_2\text{Al}(\text{NMe}_3)_2]^+$ cation and the *in situ* generation of the corresponding WCA).

Ligand substituted (CN ≥ 5). As mentioned in the previous sub-chapter, chelating ligands are of significant importance in



Fig. 12 (a) The $[(\beta\text{-diketiminato})\text{Al-H}]^+$ cation derives from the reaction of a *N*-imidoylamidine ligand with $\text{AlH}_3\text{-NMe}_2\text{Et}$ and $[\text{Ph}_3\text{C}]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$.¹³⁵ (b) The $[(\beta\text{-diketiminato})\text{Al-Me}]^+$ cation is formed by reacting the neutral precursor $(\beta\text{-diketiminato})\text{Al}(\text{Me})_2$ with the demethylating reactants $[\text{CPh}_3]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$ and $\text{B}(\text{C}_6\text{F}_5)_3$, respectively.¹³⁶

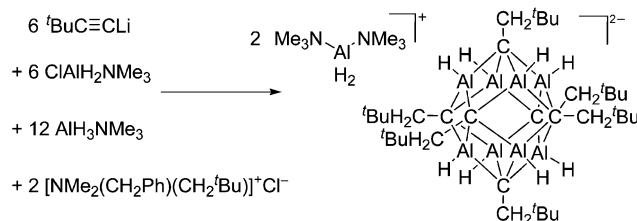


Fig. 13 Salt metathesis and hydroalumination reactions lead to the formation of the weakly coordinating carbaalane cluster that allows for the synthesis of two equivalents of the $[\text{H}_2\text{Al}(\text{NMe}_3)_2]^+$ cation.¹⁴⁵

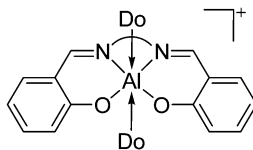


Fig. 14 Octahedral or quadratic-pyramidal coordinated $[\text{SalenAl}(\text{Do})_n]^+$ cations ($\text{Do} = \text{Et}_2\text{O}$, THF) with $n = 1, 2$.

terms of stabilizing cationic highly coordinated ($\text{CN} \geq 5$) aluminum cations. Of all the different chelates, the Salen derivatives¹⁴⁶ protrude, allowing for the synthesis of distorted square pyramidal/octahedral aluminum cations that interact with one¹⁴⁷ or two^{148–151} equivalents of Lewis base, such as Et_2O and THF (Fig. 14).

Multinuclear. A common structural motif of dicationic and dinuclear aluminum cations are the often centrosymmetric $[\text{Al}_2\text{O}_2]$ -rhomboids^{141,152–154} as seen in the recently reported $[\{(\text{OSSO})\text{Al}\}_2]^{2+}$ cation (Fig. 15).¹⁵⁵

On the other hand, there are various dinuclear, yet singly charged aluminum cations in which the latter usually feature different coordination modes. Notable contributions to this field of research have been made by Jordan *et al.*, such as the cationic aluminum aminotroponimate¹⁴¹ and amidinate¹⁵⁶ complexes in Table 8.

AlCp* substituted. The coordination chemistry of low-valent group 13 organyls such as AlCp^* to transition-metals is a growing field in inorganic chemistry, though more contributions were reported using the heavier homologue GaCp^* (see below). Nonetheless, Fischer *et al.* were able to isolate the cationic $[\text{Rh}(\text{COD})(\text{AlCp}^*)_3]^+$ complex by reacting $[\text{Rh}(\text{COD})_2]^+$ with three equivalents of AlCp^* .¹⁵⁷

Gallium cations

As mentioned above, gallium in its +I oxidation state is thermodynamically unstable and usually disproportionates into the metal and the +III ions. Notable contributions to the field of reactive gallium cations therefore allow for the stabilization of the +I oxidation state of gallium.³¹

Alkyl or aryl substituted. The isolation of the linear di-coordinated $[(2,6\text{-Mes}_2\text{C}_6\text{H}_3)_2\text{Ga}]^+$ cation¹⁵⁸ was performed by Wehmschulte *et al.* as a test run for the above mentioned

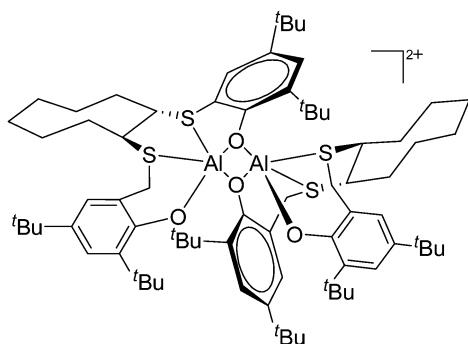


Fig. 15 In the $[\{(\text{OSSO})\text{Al}\}_2]^{2+}$ cation one aluminum atom is coordinated in a trigonal-bipyramidal and the other in a distorted-square-pyramidal fashion. The cationic species is a potential catalyst for the ring opening polymerization of propylene oxide.¹⁵⁵

structurally related $[(2,6\text{-Mes}_2\text{C}_6\text{H}_3)_2\text{Al}]^+$ cation.¹²⁹ Hence, the bowl-shaped terphenyl substituents are potential ligands in terms of shielding highly electrophilic cations. Moreover, both syntheses were only possible due to the presence of very good WCAs, such as the $[\text{Li}\{\text{Al}(\text{OR}^{\text{HF}})_4\}_2]^-$ and $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ anion.

Cyclopentadienyl complexed. Partial protolysis of GaCp^* with $[\text{H}(\text{OEt}_2)_2]^+[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$ yields the bipyramidal double-cone structured $[\text{Ga}_2(\eta^5\text{-Cp}^*)]^+$ cation.¹⁵⁹ The latter can be seen as a GaCp^* -substituted “naked” Ga^+ cation, thus reacting as a gallium(i) source with ligands such as DDP (Fig. 16).

The coordination mode of the Cp^* ligands in the $[(\eta^1\text{-Cp}^*)(\eta^3\text{-Cp}^*)\text{Ga}]^+$ cation on the other hand differs.¹⁶⁰ Hence, the originally expected η^5, η^5 -ferrocene-like structure that was also observed for the aluminum analogue is likely perturbed by the more interacting $[\text{BF}_4]^-$ counterion.

Arene complexed. Cationic arene complexes of univalent gallium are known for more than 30 years and Schmidbaur *et al.* have made notable contributions to this field of research.^{100,101} Yet, the reported compounds feature strong cation–anion interactions and are labile towards com- and disproportionations. More recently, Krossing *et al.* developed a simple oxidative route to $[\text{Ga}(\eta^6\text{-arene})_n]^+$ complexes of the weakly coordinating $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ anion with $n = 2, 3$ (Fig. 17).^{31,91,92} The arene complexes have proven to be a powerful starting material for further gallium(i) chemistry (*e.g.* various ligand exchange reactions) but also highly efficient catalyst for the polymerization of isobutylene.^{7,8,99}

Ligand substituted ($\text{CN} = 2$). The arene ligands in the $[\text{Ga}(\eta^6\text{-arene})_n]^+$ cations with $n = 2, 3$ can be substituted by electron-rich analogues. In addition, σ -donating ligands such as carbenes IR ($\text{R} = \text{Pr}$, Mes)¹⁶¹ or phosphines P^tBu_3 ¹⁶² can also be applied, yielding bent $[\text{Ga}(\text{IR})_2]^+$ and $[\text{Ga}(\text{P}^t\text{Bu}_3)_2]^+$ complexes (*cf.* the stereoactive electron lone pair at the gallium(i) cation). Another notable di-coordinated gallium(iii) cation is the linear $[\text{Bu}_3\text{Si-Ga-Si}^t\text{Bu}_3]^+$ complex (Fig. 18), which could be isolated in the presence of the $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ WCA, but not the simple $[\text{GaCl}_4]^-$ anion.¹⁶³

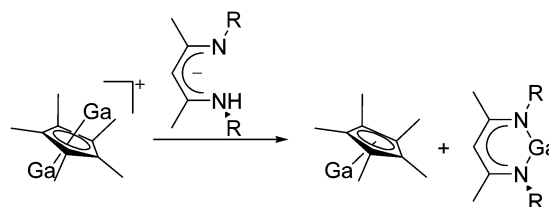


Fig. 16 The $[\text{Ga}_2(\eta^5\text{-Cp}^*)]^+[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$ salt cleanly reacts as a gallium(i) source with ligands such as DDP .

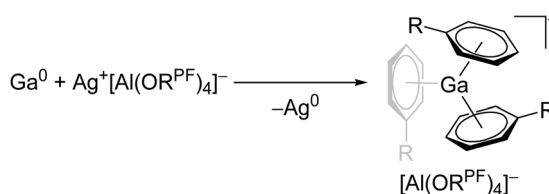


Fig. 17 Oxidative access to $[\text{Ga}(\eta^6\text{-arene})_n]^+$ complexes with $n = 2, 3$ ($\text{R} = \text{F}$, Me).



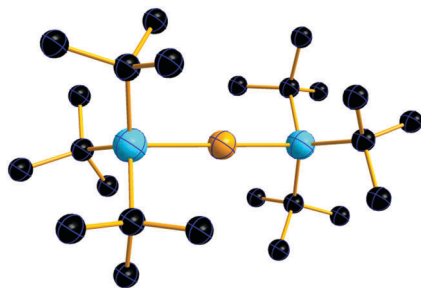


Fig. 18 Molecular structure of the $[\text{Bu}_3\text{Si-Ga-Si}^t\text{Bu}_3]^+$ cation. A. Budanow, T. Sinke, J. Tilmann, M. Bolte and M. Wagner, Two-coordinate gallium ion $[\text{Bu}_3\text{Si-Ga-Si}^t\text{Bu}_3]^+$ and the halonium ions $[\text{Bu}_3\text{Si-X-Si}^t\text{Bu}_3]^+$ ($\text{X} = \text{Br}, \text{I}$): sources of the supersilyl cation $[\text{Bu}_3\text{Si}]^+$, *Organometallics*, 2012, **31**, 7298–7301. Data from this reference were used to draw this figure and the hydrogen atoms were omitted for clarity.¹⁶³

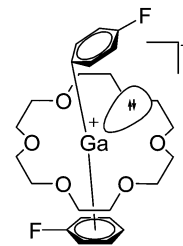


Fig. 21 The $[\text{Ga}(\text{[18]crown-6})(\eta^6\text{-}/\eta^1\text{-C}_6\text{H}_5\text{F}_2)]^+$ cation. The $\eta^6\text{-}$ and $\eta^1\text{-}$ coordination modes could be an indication for a stereoactive lone pair on the side of the weaker and only $\eta^1\text{-}$ coordinated $\text{C}_6\text{H}_5\text{F}$ molecule.¹⁶⁸

Ligand substituted (CN = 3). This class of tricoordinate gallium(i) cations again derives from the above mentioned $[\text{Ga}(\eta^6\text{-arene})_n]^+$ cations with $n = 2, 3$. The coordination mode for the gallium(i) cations is trigonal-pyramidal due to the stereoactive lone pair electrons. Besides sterically less demanding phosphines, N-heterocyclic arenes like pyrazine and DTBMP (a $\sigma\text{-}$, and not a $\pi\text{-}$ donating ligand, proving its perception of being non-nucleophilic wrong) were also applied as potential ligands.¹⁶⁴ Due to the bifunctionality of pyrazine, both the monomeric $[\text{Ga}(\text{pyrazine})_3]^+$ complex and the one-dimensional coordination polymer $[\{\text{Ga}(\mu\text{-pyrazine})_2(\eta^1\text{-pyrazine})\}]^+_\infty$ were isolated (Fig. 19).

Ligand substituted (CN = 4). Using the BOX ligand, Dagorne *et al.* isolated tetra-coordinate neutral gallium complexes.¹⁶⁵ The latter were easily ionized by applying $[\text{CPh}_3]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$ or $\text{B}(\text{C}_6\text{F}_5)_3$ in NMe_2Ph . Interestingly, the trityl cation functions as hydride and $\text{B}(\text{C}_6\text{F}_5)_3$ as methyl abstracting reactant (Fig. 20).

Ligand substituted (CN ≥ 5). Cationic penta- and hexa-coordinated gallium complexes are synthesized *via* protonation^{166,167} or complexation.¹⁶⁸ Within this context, the $[\text{Ga}(\text{[18]crown-6})(\text{C}_6\text{H}_5\text{F})_2]^+$ complex is of special interest as the gallium(i) cation

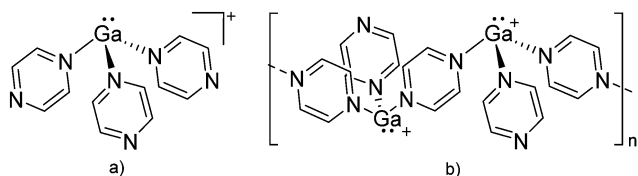


Fig. 19 (a) Monomeric $[\text{Ga}(\text{pyrazine})_3]^+$ complex and (b) one-dimensional coordination polymer $[\{\text{Ga}(\mu\text{-pyrazine})_2(\eta^1\text{-pyrazine})\}]^+_\infty$. The propagation of the polymer into the second dimension was not possible as each cationic strand is surrounded by strands of the corresponding $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ anions.¹⁶⁴

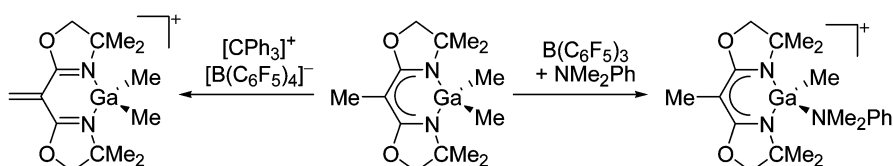


Fig. 20 Hydride vs. methyl abstraction of neutral BOX ligated gallium complexes.¹⁶⁵

features no contact to the corresponding $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ anion and the $\text{C}_6\text{H}_5\text{F}$ ligands coordinate in different fashions (Fig. 21).¹⁶⁸

As the N-heterocyclic arenes are potential ligands for univalent gallium (see above), Krossing *et al.* additionally reacted the chelating bipy with the $[\text{Ga}(\eta^6\text{-C}_6\text{H}_5\text{F})_2]^+$ complex. Instead of witnessing a simple ligand exchange reaction, they isolated the paramagnetic and distorted octahedral $[\text{Ga}^{\text{III}}\{\text{bipy}\}_3]^+_{2+}$ complex due to the non-innocence of the bipy ligand.¹⁷ This is reminiscent to transition metal chemistry where for example the $[\text{Ru}^{\text{III}}\{\text{bipy}\}_3]^+_{2+}$ complex features similar bonding.

Transition-metal substituted. Similar to the transition-metal substituted boron cations, $\text{CpFe}(\text{CO})_2$ (FP) and $\text{Cp}^*\text{Fe}(\text{CO})_2$ (FP*) are the most important ligands in terms of stabilizing di-, tri- and tetra-coordinated gallium cations: *cf.* the $[(\text{FP}^*)_2\text{Ga}]^+$,¹⁶⁹ $[(\text{FP}^*)_2\text{Ga}(4\text{-Pic})]^+$,¹⁷⁰ $[(\text{FP}^*)\text{Ga}(\text{phen})(\text{Y})]^+$ ($\text{Y} = \text{Cl}, \text{S}^p\text{Tol}$)¹⁷¹ cations (Fig. 22).

Multinuclear. There are not many contributions to this field of research and some cationic multinuclear gallium complexes are a product of hydrolysis.^{172,173} Two remarkable examples however are the dinuclear $[(\eta^6\text{-C}_6\text{H}_5\text{F})\text{Ga}(\mu\text{-}\eta^6\text{-}m\text{-TP})_2\text{Ga}(\eta^6\text{-C}_6\text{H}_5\text{F})]^+_{2+}$ complex^{8,99} in which the gallium(i) cations are solely π -coordinated by arene ligands as well as the σ -coordinated amidinate-bridging $[\text{BuC}(\text{N}^i\text{Pr})_2]\text{GaMe}[\text{BuC}(\text{N}^i\text{Pr})_2]\text{GaMe}_2]^+$ cation.¹⁵⁶

Multinuclear transition-metal substituted. A notable class of contributions are the β -diketiminate/THF coordinated gallium cations that can be bridged by a gold atom¹⁷⁴ or a $\{\text{ZnCl}(\text{THF})_2\}_2$ -rhomboid.¹⁷⁵ Reaction of $\text{Cp}^*\text{Fe}(\eta^5\text{-P}_5)$ with the $[\text{Ga}(\text{o-C}_6\text{H}_4\text{F}_2)]^+$ complex on the other hand resulted in aggregation and the formation of a cationic one-dimensional coordination polymer (Fig. 23).¹⁷⁶

GaCp* substituted. As of today, GaCp^* is a widely used ligand concerning cationic transition-metal complexes, thus leading to an enormous variety of cationic gallium species. This area of research has been intensively reviewed by Fischer *et al.*¹¹⁰ and we would like to refer to the multiple entries in Table 8 of this review. Yet, some of the compounds also include “naked” and bridging gallium atoms:



Fig. 22 (a) The linear di-coordinated cation $[(FP^*)_2(\mu\text{-Ga})]^+$ derives from a salt metathesis of $(FP^*)_2\text{GaCl}$ and $\text{Na}^+[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$. The Fe–Ga–Fe moiety features a significant π bonding component (population analysis).¹⁶⁹ (b) The $[(FP^*)_2(\mu\text{-Ga})\text{-}(4\text{-Pic})]^+$ cation is an addition product of $[(FP^*)_2\text{Ga}]^+$ and 4-Pic and the second structurally characterized complex containing a cationic tricoordinate gallium centre.¹⁷⁰ (c) Applying the chelating phen ligand, Ueno *et al.* isolated the tetra-coordinated $[(FP^*)\text{Ga}(\text{phen})(Y)]^+$ ($Y = \text{Cl}, S^*\text{Tol}$) cations, *i.e.* the first transition-metal complex with a thiolate group on the gallium atom.¹⁷¹

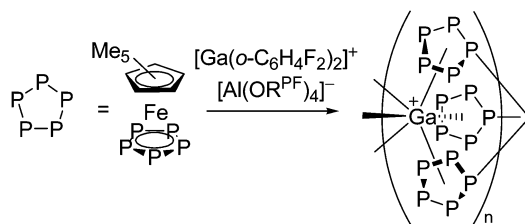


Fig. 23 Reaction of $[\text{Ga}(\text{o-C}_6\text{H}_4\text{F}_2)_2]^+$ and $\text{Cp}^*\text{Fe}(\eta^5\text{-P}_5)$ results in aggregation and formation of a cationic one-dimensional coordination polymer.

e.g. $[(\text{GaRu}(\text{PCy}_3)_2(\text{GaCp}^*)_2)]^+$,¹⁷⁷ $[(\text{GaM}(\text{GaCp}^*)_4)]^+$ ($\text{M} = \text{Ni},^{178} \text{Pt}^{179,180}$), $[(\text{Cp}^*\text{Ga})_4\text{Rh}\{\text{Ga}(\text{Me})\}]^+$,¹⁸¹ $[(\text{Cp}^*\text{Ga})_4\text{Rh}\{\text{Ga}(\text{Me})(\text{py})\}]^+$,¹⁸¹ $[\{\text{Ru}(\text{GaCp}^*)_3\}\text{-}[(\text{CH}_2)_2\text{C}\{\text{CH}_2(\mu\text{-Ga})\}]_2]^+$,¹⁷⁷ $[(\text{GaCp}^*)_4\text{Pt}\{\text{Pt}(\text{H})(\text{GaCp}^*)_3\}\text{-}(\mu\text{-Ga})]^{2+}$.¹⁸⁰ Contrary to GaCp^* (a strong σ -donor and weak π -acceptor, *cf.* similarity to the boron related compounds in Fig. 6), the “naked” gallium cations function as pure acceptor ligands, with significant components of σ - and π -symmetry contributing to the M–Ga linkages.^{179,180}

Indium cations

Compared to the lighter homologue gallium, well-defined indium(i) halides exist, though they are practically insoluble in organic solvents. The synthesis of $\text{In}^+[\text{OTf}]^-$ by Macdonald *et al.* as a soluble alternative is therefore an important development concerning the indium(i) chemistry.¹⁸²

Cyclopentadienyl complexed. Using the just mentioned $\text{In}^+[\text{OTf}]^-$ salt as starting material and reacting it with manganocene, the inverse sandwich complex $[\text{In}_2(\eta^5\text{-Cp})]^+$ was successfully synthesized.¹⁸³ Interestingly, the counterion is the complex $[\text{Cp}_3\text{In}^{\text{III}}\text{-Cp-In}^{\text{III}}\text{Cp}_3]^-$ ion, deriving from a partial oxidation of the starting material. The formation of the mixed valence species seems to be preferred over an alternative indium(ii) species. Reacting InCp^* (a hexamer in the solid state) with a mixture of $\text{B}(\text{C}_6\text{F}_5)_3$ and $\text{H}_2\text{O}\cdot\text{B}(\text{C}_6\text{F}_5)_3$, the first indium-based triple-decker cation



Fig. 24 Reducing the size of the counterion from $[(\text{C}_6\text{F}_5)_3\text{BO}(\text{H})\text{B}(\text{C}_6\text{F}_5)_3]^-$ to $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ “squeezes” the toluene molecules from the triple-decker cation, yielding the inverse sandwich complex $[\text{In}_2(\eta^5\text{-Cp})]^+$.

$[(\eta^6\text{-Tol})\text{In}(\mu\text{-}\eta^5\text{-C}_5\text{Me}_5)\text{In}(\eta^6\text{-Tol})]^+$ was formed.¹⁸⁴ Reducing the size of the counterion from $[(\text{C}_6\text{F}_5)_3\text{BO}(\text{H})\text{B}(\text{C}_6\text{F}_5)_3]^-$ to $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ on the other hand, results in the formation of the inverse sandwich complex $[\text{In}_2(\eta^5\text{-Cp})]^+$ in which the indium(i) cations are not capped by toluene molecules but rather interact with the $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ anions (Fig. 24).¹⁸⁵

Arene complexed. By reacting elemental indium with $\text{Ag}^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$, Krossing *et al.* expanded the above mentioned oxidative route to gallium(i) salts towards the synthesis of $[\text{In}(\text{arene})_n]^+$ complexes with $n = 2, 3$.¹⁶² Identical compounds can also be synthesized by using the salt metathesis reactions of Scheer *et al.*, with insoluble InCl as starting material.¹⁸⁶

Ligand substituted (CN = 2). These $[\text{In}(\text{arene})_n]^+$ complexes with $n = 2, 3$ are an ideal starting material for further indium(i) chemistry: *e.g.* the arene ligands can be substituted for N-heterocyclic carbenes such as iPr .¹⁶¹ Salt metathesis reactions on the other hand are still very important: *i.e.*, using the isosteric and isoelectronic terphenyl Mes_2py ligand, Aldridge *et al.* were able to isolate mixed-leptic $[\text{In}(\text{Mes}_2\text{py})(\eta^6\text{-C}_6\text{H}_5\text{F})]^+$ (both σ - and π -coordinated) and homo-leptic $[\text{In}(\text{Mes}_2\text{py})_2]^+$ complexes (only σ -coordinated, though the flanking mesityl rings of the Mes_2py ligands also partly π -coordinate).¹⁸⁷ The latter features an indium(i) cation wholly encapsulated by two Mes_2py ligands and remarkably long In–N distances, which the authors explain with an energy mismatch between the (low lying) pyridine ligand donor and (high energy) metal acceptor orbitals.

Ligand substituted (CN = 3). Besides complexation and ligand exchange reactions of $\text{In}^+[\text{OTf}]^-$ and $[\text{In}(\text{arene})_n]^+$ ($n = 2, 3$) with ligands such as bis(imino)pyridines^{188,189} and PPh_3 ,¹⁶² tricoordinate indium cations can also be isolated by thermolysis of $[(^i\text{Pr}_2\text{-ATI}(\text{CPh}_3))\text{InMe}_2]^+$, a cationic tetra-coordinated indium precursor (Fig. 25, conversion of (a) to (b)).¹⁹⁰

Ligand substituted (CN = 4). The cationic diimine $[(^i\text{Pr}_2\text{-ATI}(\text{CPh}_3))\text{InMe}_2]^+$ complex was synthesized by reacting the neutral $(^i\text{Pr}_2\text{-ATI})\text{InMe}_2$ precursor with the ionizing $[\text{Ph}_3\text{C}]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$ salt (Fig. 25).¹⁹⁰ Surprisingly, the latter does not function as methyl abstracting reactant but rather adds to the C5 carbon of $(^i\text{Pr}_2\text{-ATI})\text{InMe}_2$. Reacting $(^i\text{Pr}_2\text{-ATI})\text{InMe}_2$ with the protonating $[\text{HNMe}_2\text{Ph}]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$ on the other hand, results in CH_4 formation and the labile adduct $[(^i\text{Pr}_2\text{-ATI})\text{In}(\text{Me})(\text{NMe}_2\text{Ph})]^+$ (Fig. 25).¹⁹⁰

Ligand substituted (CN ≥ 5). Compared with the lighter homologues aluminum and gallium, indium shows a tendency to expand its coordination sphere.^{95,102} Protonolysis of the neutral $\text{In}(\text{CH}_2\text{SiMe}_3)_3$ complex in THF therefore yields a penta-coordinated indium cation: $[\text{In}(\text{CH}_2\text{SiMe}_3)_2(\text{THF})_3]^+$.¹⁹¹ Moreover, $\text{In}^+[\text{OTf}]^-$ ^{192,193}





Fig. 25 Formation of tetra- and tricoordinate cationic diimine substituted indium complexes: (a) $[(t\text{Pr}_2\text{-ATI}(\text{CPh}_3))\text{InMe}_2]^+$, (b) $[(t\text{Pr}_2\text{-ATI})\text{InMe}]^+$ and (c) $[(t\text{Pr}_2\text{-ATI})\text{In}(\text{Me})(\text{NMe}_2\text{Ph})]^+$. For each complex, the counterion is $[\text{B}(\text{C}_6\text{F}_5)_4]^-$.

and $[\text{In}(\text{arene})_n]^+{}^{168}$ ($n = 2, 3$) can be reacted with the crown ether [18]crown-6, yielding cationic indium complexes with similar structures to the gallium congener (*cf.* Fig. 21) and strong anion-cation interactions in the case of the $[\text{OTf}]^-$ anion. Reacting $\text{In}^+[\text{OTf}]^-$ with [15]crown-5 on the other hand, the sandwich complex $[\text{In}([\text{15}]\text{crown-5})_2]^+$ was isolated.¹⁹⁴

Transition-metal substituted. The class of cationic transition-metal substituted indium compounds very much relates to the related gallium structures: *i.e.*, the $[\text{InPt}(\text{PPh}_3)_3]^+$ complex with a “naked” Pt-substituted indium cation^{179,180} as well as the di- and tricoordinate $[(\text{FP}^*)_2\text{In}]^+$ and $[(\text{FP}^*)_2\text{In}(\text{THF})]^+$ complexes.¹⁹⁵ Reacting the chelating phen ligand with the $[\text{In}(\text{C}_6\text{H}_5\text{F})_2]^+$ complex in the presence of silver salt, Krossing *et al.* isolated the silver bound indium dication¹⁷ $[(\text{phen})_2\text{In-Ag}(\eta^3\text{-C}_6\text{H}_5\text{F})]^{2+}$ that is related to the $[\text{InPt}(\text{PPh}_3)_3]^+$ complex.^{179,180} In this complex the tetragonal-pyramidal $[\text{In}(\text{phen})_2]^+$ cation reacts as a Lewis basic donor (*cf.* the stereoactive 5s lone pair at indium), while the $[\text{Ag}(\eta^3\text{-C}_6\text{H}_5\text{F})]^+$ complex is the corresponding Lewis acidic acceptor.

Multinuclear. For the synthesis of multinuclear indium cations, the $[\text{In}(\text{arene})_n]^+{}^{168}$ complexes with $n = 2, 3$ are a powerful starting material. Hence a dicationic $[(\text{PPh}_3)_3\text{In}]_2(\mu\text{-PPh}_3)]^{2+}$ complex in which one PPh_3 ligand bridges both indium(i) cations was isolated.¹⁶² Applying the non-innocent and chelating bipy and phen ligands on the other hand, Krossing *et al.* surprisingly isolated the first cationic tri- and tetra-nuclear indium clusters: $[\text{In}_3(\text{bipy})_{5-6}]^{3+}$ and $[\text{In}_4(\text{Do})_6]^{4+}$ (Do = phen, bipy) (Fig. 26).¹⁷ This result very much differs from the above mentioned synthesis of the paramagnetic $[\text{Ga}^{\text{III}}(\text{bipy})_3]^+{}^{2+}$ complex and can be attributed to the higher redox-stability of indium compared to gallium. In addition and to our knowledge, these are the first higher charged clusters that have been reported to this day: *i.e.*, for cluster formations usually reductive syntheses are applied, yielding neutral and anionic clusters.

Thallium cations

In contrast to the lighter homologues, thallium's thermodynamic most stable oxidation state is +I. Hence, various syntheses of unsubstituted thallium(i) cations of different WCAs have been reported: *i.e.*, the protonation of TlOEt using $[\text{H}(\text{OEt}_2)_2]^+[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-/[\text{B}(\text{C}_6\text{F}_5)_4]^-$,^{196,197} the Lewis acid base reaction of $\text{Tl}^+[\text{OTeF}_5]^-$ and $\text{B}(\text{OTeF}_5)_3$ ¹⁹⁸ and the salt metathesis of TlF and $\text{Li}^+[\text{Al}(\text{OR}^{\text{HF}})_4]^-$.^{199,200} The thallium(i) salts are relatively stable (*cf.* the silver congeners decompose upon exposure to light) and mainly used as reactants to introduce WCAs (*e.g.* salt metathesis reactions). $\text{Tl}^+[\text{Al}(\text{OR}^{\text{HF}})_4]^-$ could only be isolated, if the precursors were applied in an exact 1:1 stoichiometry. An

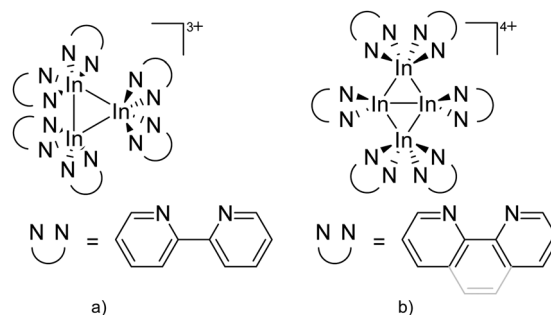


Fig. 26 Cationic (a) $[\text{In}_3(\text{bipy})_{5-6}]^{3+}$ and (b) $[\text{In}_4(\text{Do})_6]^{4+}$ (Do = bipy, phen) complexes synthesized *via* ligand exchange reactions and aggregations.

excess of TlF however, led to the formation of the cationic multinuclear $[\text{Tl}_3\text{F}_2\text{Al}(\text{OR}^{\text{HF}})_3]^+$ complex.²⁰⁰

Arene complexed. Various cationic thallium(i) arene complexes have been reported. While the di- and tricoordinate $[\text{Tl}(\eta^6\text{-arene})_n]^+$ complexes (arene = $\text{C}_6\text{H}_5\text{Me}$, $n = 2, 3$;²⁰¹ Mes, $n = 2$;²⁰² C_6Me_6 , $n = 2$;²⁰³) are structurally related to the lighter homologues, C_6Me_6 additionally allows for the first mono-coordinated $[\text{Tl}(\eta^6\text{-C}_6\text{Me}_6)]^+$ complex (DFT calculations gave a remarkable $\text{Tl-C}_6\text{Me}_6$ π -bonding energy of 163 kJ mol^{-1}).²⁰⁴

Ligand substituted (CN = 2). Reacting $\text{Tl}^+[\text{OTeF}_5]^-$ with $\text{B}(\text{OTeF}_5)_3$ in 1,2-dichloroethane, the solvent functions as chelating ligand, thus forming the five-membered TlCl_2C_2 -ring in $[\text{Tl}(1,2\text{-Cl}_2\text{C}_2\text{H}_4)]^+$ (after silver and ruthenium, thallium was at that time the third reported metal atom coordinated by a simple chlorocarbon).¹⁹⁸ By contrast, from CH_2Cl_2 the “naked” $\text{Tl}^+[\text{B}(\text{OTeF}_5)_4]^-$ salt was isolated.

Ligand substituted (CN = 3). Similar to the lighter homologues gallium and indium, the $[\text{Tl}(\eta^6\text{-C}_6\text{H}_5\text{R})_2]^+$ ($\text{R} = \text{F}, \text{Me}$) bent-sandwich complexes can interact with N-heterocyclic ligands such as Mes_2Py , thus forming tricoordinate $[\text{Tl}(\text{Mes}_2\text{py})(\eta^6\text{-C}_6\text{H}_5\text{R})_2]^+$ complexes.¹⁸⁷ On the other hand tri-dentate chelating ligands like $\text{timtmb}^{\text{rBu}}$ ²⁰⁵ and bis(imino)pyridines²⁰⁶ can be applied to isolate tricoordinate thallium(i) cations (Fig. 27).

Ligand substituted (CN = 4). The protonation of TlOEt with $[\text{H}(\text{OEt}_2)_2][\text{H}_2\text{N}[\text{B}(\text{C}_6\text{F}_5)_3]_2]^-$ in Et_2O yielded the tetrahedral coordinated cationic $[\text{Tl}(\text{OEt}_2)_4]^+$ complex, which shows no contact to the corresponding WCA.²⁰³

Ligand substituted (CN ≥ 5). 2,5-Bis(2-pyridyl)-1-phenylphosphole (NPPH) exhibits a rich coordination chemistry towards thallium(i) cations and dependent on the nature of the solvents and WCAs, different structures were obtained: *i.e.*,



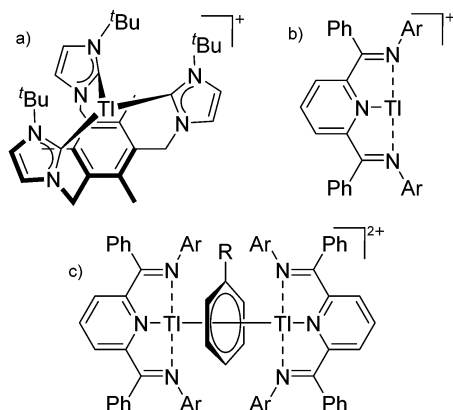


Fig. 27 Both the (a) $[Tl(timtm-bu)]^+$ and (b) $[ArN=CPh]_2(NC_5H_3)Tl]^+$ complexes derive from $Tl^+[OTf]^-$ and are synthesized via complexation reactions of the corresponding ligands. In addition, the inverted sandwich structure (c) $[ArN=CPh]_2(NC_5H_3)Tl]_2(\mu-\eta^6-C_6H_5R)]^{2+}$ ($Ar = 2,6-Et_2C_6H_3$, $2,5-tBu_2C_6H_3$; $R = H, Me$) was isolated.

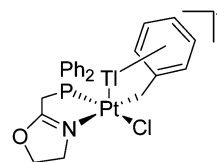


Fig. 29 The first fully characterized metal-metal bonded Tl–Pt–Cl complex. If $Ag^+[OTf]^-$ and $Ag^+[BF_4]^-$ is applied, the expected chloride abstraction takes place.²⁰⁸

reacting $Tl^+[Al(OR^{PF})_4]^-$ with NPPh in CH_2Cl_2/C_6H_5Me , the tetra-coordinated and C_6H_5Me -capped $[Tl(NPPh)_2(\eta^6-C_6H_5Me)]^+$ complex formed, whereas in CH_2Cl_2/n -pentane the dinuclear and dicationic $[Tl_2(NPPh)_4]^{2+}$ was isolated (Fig. 28). If $Tl^+[PF_6]^-$ was applied as starting material a coordination polymer with strong cation–anion interactions was formed.

An even higher coordinated thallium cation is the $[Tl([18]crown-6)]^+$ complex, which features a similar structure as the $[18]crown-6$ complexes of the lighter homologues gallium and indium.²⁰³

Transition-metal substituted. Reacting the above mentioned $[Tl(\eta^6-C_6H_5Me)_n]^+$ complexes ($n = 2, 3$) with $FeCp_2$, Sarazin *et al.* were able to isolate the $[Tl_2(FeCp_2)_3][H_2N\{B(C_6F_5)_3\}_2]^-$ salt that contains the mono- and di-coordinated $[Tl(FeCp_2)_n]^+$ complexes with $n = 1, 2$ in a 1 : 1 ratio.²⁰³ Increasing the amount of $FeCp_2$ from 1 to 2.2 equivalents, only the $[Tl(FeCp_2)]^+$ complex was isolated.²⁰⁴ In contrast to the lighter homologues, the reaction of $Cp^*Fe(\eta^5-As_5)$ with $Tl^+[PF_6]^-$ and $Li^+[FA\{OC_6F_{10}(C_6F_5)_3\}_3]^-$ did not result in aggregation and formation of a cationic one-dimensional coordination polymer, but rather yielded the pseudo-trigonal-planar $[Tl\{(\eta^5-As_5)FeCp^*\}_3]^+$ complex.¹⁷⁶ Performing a similar chemistry in the presence of the very good WCA $[Al(OR^{PF})_4]^-$ however, one-dimensional polymers were isolated (*cf.* Fig. 23), proving

the importance of the WCA.²⁰⁷ Reacting the neutral $Pt(CH_2Ph)-Cl(PCH_2-ox)$ complex with $Tl^+[PF_6]^-$, Braunstein *et al.* did not isolate any chloride abstraction product but a “trapped” thallium(i) cation: the cationic $[P(Ph_2)CH_2Ox\{Cl\}(Tl)Pt-CH_2Ph]^+$ complex.²⁰⁸ Herein, the ligand functions as a chelate and interacts with thallium *via* a Pt–Tl bond and a η^6 -benzyl coordination (Fig. 29).

Multinuclear. Some of the cationic multinuclear thallium complexes have already been mentioned in the text above. A further example is the $[Tl(\beta\text{-triketimine})_2]^{2+}$ complex that features Tl– η^6 -aryl and weak thalophilic interactions, allowing to overcome the Coulomb repulsion of both cations (*cf.* Fig. 28).¹⁴ The reaction of the P_n -ligand $\{CpMo(CO)_2\}_2(P_2)$ with $Tl^+[Al(OR^{PF})_4]^-$, yields the dicationic $[Tl_2\{CpMo(CO)_2\}_2P_2]^{2+}$ complex that features a distorted Tl_2P_4 ring (Fig. 30).²⁰⁷

Reacting $RuCl_2(DMeOPrPE)_2$ with $Tl^+[PF_6]^-$ an “arrested” chloride abstraction occurs.²⁰⁹ In the resultant one-dimensional coordination polymer, the thallium(i) cations are coordinated in an unusual octahedral fashion with a stereoactive $6s^2$ lone pair at thallium.

Group 14 cations

Already in 1887, Henderson described the synthesis of trityl malonate starting from triphenylmethyl bromide and ethylic sodiomalonate²⁴⁶ and 15 years later, Bayer and Villiger realized that the yellow color of a solution of triphenylmethane in concentrated sulfuric acid is the result of the formation of a carbocation.²⁴⁷ Despite these early discoveries, it took another 63 years until the structure of this cation could be determined.²⁴⁸ While the first structure determination succeeded with $[ClO_4]^-$, the structure of the trityl cation is nowadays known with several different anions (*e.g.* ref. 249) and it has become a common

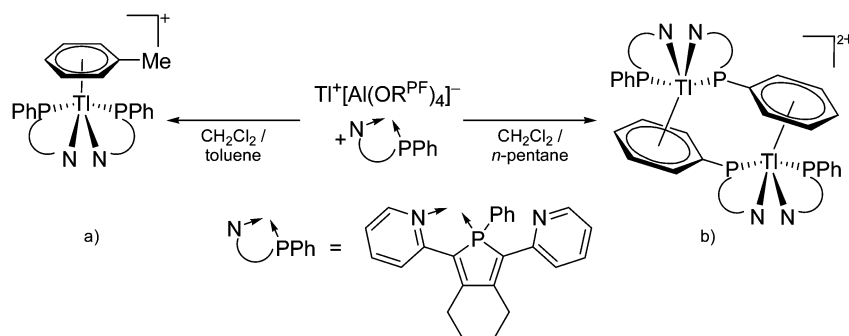


Fig. 28 Solvent effects on the formation of cationic thallium(i) complexes of NPPh. (a) If toluene is applied, the solvent-stabilized penta-coordinated $[Tl(NPPh)_2(\eta^6-C_6H_5Me)]^+$ complex forms. (b) If non-coordinating n -pentane is applied two $[Tl(NPPh)_2]^+$ cations aggregate *via* their phenyl substituents, forming the dinuclear and dicationic $[Tl_2(NPPh)_4]^{2+}$ complex.

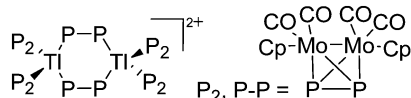


Fig. 30 Formation of the $[\text{Ti}_2((\text{CpMo}(\text{CO})_2)_2)_6]^{2+}$ complex.

reagent for the generation of various other cations. Only one earlier structure determination of a carbocation was published: the structure of triphenylcyclopropenium perchlorate in 1963.²⁵⁰ Since then, many rPBCs of group 14 were synthesized and characterized. Carbocations have drawn a lot of interest, due to their role as intermediates in organic chemistry. Silylium ions are more electrophilic and more reactive than their carbon analogues, so that the structural characterization of a truly free silylium ion was only achieved in 2002²⁴ and is still in the focus of interest. But also the heavier elements of group 14 were subject to extensive research and today a multitude of interesting rPBCs are known, part of which have been reviewed in the articles included with Table 4.

Carbon

Homopolyatomic and cluster cations. In group 14, carbon is the only element for which homopolyatomic cations are known in condensed phase. While $[\text{C}_7]^{+}$,⁴⁴ and $[\text{C}_{60}]^{+}$,²⁵⁹ are already known for more than ten years, there is only one more recently published compound of that class. In $[\text{C}_{60}]^{2+}[(\text{AsF}_6)^-]_2$,²⁶⁰ the cations build a 1D polymeric structure, in which the $[\text{C}_{60}]^{2+}$ cations are connected alternately by single C–C bonds and four-membered carbon rings. Along with the before mentioned $[\text{C}_{60}]^{+}$, the protonated buckminsterfullerene $[\text{HC}_{60}]^{+}$ was published²⁵⁹ and by oxidation of the $[\text{C}_{59}\text{N}]_2$ dimer, $[\text{C}_{59}\text{N}]^{+}$ was synthesized and structurally characterized.²⁶¹

Carbonium ions. As mentioned before, the classification according to onium-, enium and inium-cations is not always consistent and in literature, the term carbonium ion is often used to describe what is mostly a carbenium ion. A prototype for a carbonium ion is the 2-norbornyl cation, whose structure has been controversially discussed. In 2013, 49 years after its first preparation under stable ion conditions,²⁶² its structure could be determined by scXRD.⁸⁶ This finally provided a crystallographic proof that the 2-norbornyl cation adopts the non-classical structure (Fig. 31). It remains the only structurally characterized non-classical



Fig. 31 Non-classical vs. classical structure of the 2-norbornyl cation.

carbonium ion. Substituted relatives exhibit distinctly distorted structures that are better classified as carbenium ions.²⁶³

Carbenium ions. The first simple structurally characterized alkyl cation, was the *tert*-butyl cation with $[\text{Sb}_2\text{F}_{11}]^-$ as the counterion,²⁶⁴ and later also with $[\text{HCB}_{11}\text{Me}_5\text{Cl}_6]^-$.²⁶⁵ In the same publication, two more carbocations with slight variations in the alkyl chains were presented (Fig. 32).²⁶⁵ Recent additions include the super-acidic room temperature ionic liquid $[(\text{CH}_3)_3\text{C}]^+[\text{Al}_2\text{Br}_7]^-$ ²⁶⁶ and an additional structure of the *tert*-butyl cation with the $[\text{HCB}_{11}\text{Cl}_{11}]^-$ anion.²⁶⁷ In 2000, ion-like $(\text{CH}_3)_2\text{CF}(\text{AsF}_6)$ was the first structurally characterized example of a fluorinated carbocation and was published together with a higher substituted variant.²⁶⁸ In both compounds, each cation is stabilized by two stronger contacts to the anion. The higher substituted $[(m\text{-CF}_3\text{C}_6\text{H}_4)(\text{C}_6\text{H}_5)\text{CF}]^+$ derivative, contains the less coordinating $[\text{As}_2\text{F}_{11}]^-$ anion in the structure with only weak interaction between the ions.²⁶⁸ With $[\text{HCB}_{11}\text{I}_{11}]^-$, two more fluoro-substituted carbocations and one with fluorine substituted aryl residues could be isolated (see Fig. 32).⁵⁶ Apart from $[\text{CF}_3]^+$, all $[\text{CX}_3]^+$ cations are now synthesized and structurally characterized (see Fig. 32). First, $[\text{Cl}_3]^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ was published in 2003²⁶⁹ and shortly after $[\text{CCl}_3]^+$ and $[\text{CBr}_3]^+$ with $[\text{Sb}(\text{OTeF}_5)_6]^-$ as the counterion.²⁷⁰ In addition, the latter was used to stabilize related $[\text{C}(\text{OTeF}_5)_3]^+$.²⁷⁰ Later, also $[\text{CCl}_3]^+$ and $[\text{CBr}_3]^+$ were synthesized with the $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ and the $[(\text{R}^{\text{PF}}\text{O})_3\text{Al}-\text{F}-\text{Al}(\text{OR}^{\text{PF}})_3]^-$ counterions.²⁷¹ In all of those compounds containing $[\text{CX}_3]^+$ cations, still some weak interactions between cation (mainly halogen atoms) and anion exist. These interactions are weaker between $[\text{Br}-\text{C}(\text{SBr})_2]^+$ and the mentioned alkoxyaluminate, due to delocalization of the charge.¹¹⁷ Although comparable, far stronger interaction between cation and anion was found in $[(\text{MeO})(\text{MeS})\text{CSH}]^+[\text{SbF}_6]^-$.²⁷² However, there is no close contact between the carbon atom and the fluorine atoms of the anion. Instead, the anion forms hydrogen bonds to the thiol group of the cation.

In 2004, the structure of the benzonorbornenyl cation was published, with an intramolecular stabilization of the cationic center by the aromatic ring.²⁷³ Intermolecular stabilized carbenium cations are known of the $[\text{Cl}_3]^+$ with the weak bases PX_3

Table 4 Review articles including cationic group 14 compounds

Year	Title	Ref.
1995	Modern approaches to silylium cations in condensed phase	251
2005	Cations of group 14 organometallics	252
2005	Carbon, silicon, germanium, tin and lead	253
2010	Silylium ions in catalysis	254
2010	H^+ , CH_3^+ , and R_3Si^+ carborane reagents: when triflates fail	255
2011	N-heterocyclic carbene analogues with low-valent group 13 and group 14 elements: syntheses, structures, and reactivities of a new generation of multitailented ligands	256
2013	Catenated compounds – group 14 (Ge, Sn, Pb)	257
2015	Cations and dications of heavier group 14 elements in low oxidation states	258





Fig. 32 Structurally characterized carbenium ion salts.



Fig. 33 Structurally characterized ligand-stabilized carbenium ions and vinyl cation salts.

(X = Cl, Br, I) and AsI_3 (Fig. 33).²⁷⁴ Only two related vinyl cations are known (see Fig. 33).^{275,276} Both are β -substituted by two silyl groups, which help to stabilize the positive charge.

Delocalized (cyclic) carbocations. Only shortly after the first structural characterization of an alkyl cation, the first structure determination of an arenium ion – $[\text{C}_6\text{Me}_7]^+[\text{AlCl}_4]^-$ – was published.²⁷⁷ To date, more structurally characterized arenium ions with several WCAs are known (Fig. 34).

An exception is the radical cation $[\text{C}_6\text{F}_6]^{+\bullet}$ in the solid state structures with $[\text{Sb}_2\text{F}_{11}]^-$ and $[\text{Os}_2\text{F}_{11}]^-$: it yields two different forms.²⁷⁸ One cation can be described as a quinoidal cation and the other as a bisallyl cation (see Fig. 35) and both are separated by a barrier of around 13 kJ mol^{-1} according to calculations.

Shortly after the publication of the radical cation of the hexafluorobenzene, some more related structures were presented. Among them, the other perhalogenated benzene radical cations^{279,280} and some partially and mixed substituted analogs, including the $[\text{C}_6\text{F}_5-\text{C}_6\text{F}_5]^{+\bullet}$ ^{280,281} (Fig. 36). The only other example displaying both a quinoidal and a bisallyl cationic form is $[2,4,6\text{-}^t\text{Bu}_3\text{C}_6\text{H}_2\text{NH}_2]^{+\bullet}$.²⁸² At 123 K, this cation adopts the bisallylic structure but upon heating, a transition to the quinoidal form occurs.

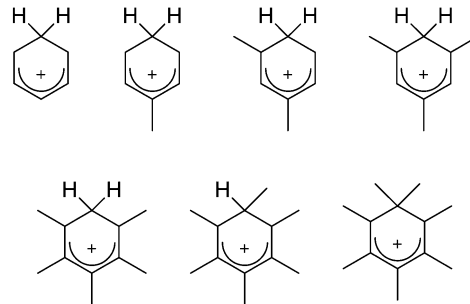
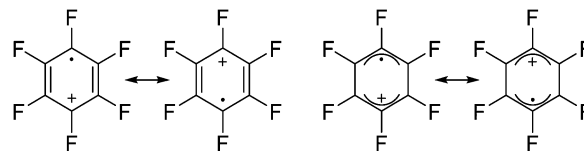


Fig. 34 Protonated and methylated structurally characterized arenium ion salts.

Fig. 35 Lewis structures of the canonical forms of the quinoid and the bisallyl cationic form of $[\text{C}_6\text{F}_6]^{+\bullet}$.

A different type of delocalized cations are the allyl cations amongst which the cyclopropenyl cations take a special position. Already since 1986, two examples, $[(\text{Cy})_3\text{C}_3]^+$ and $[(\text{Cy})_2(\text{Ph})\text{C}_3]^+$, are known²⁸³ and in the same year, an allyl cation stabilized by an hydroxyl group has been published (Fig. 37).²⁸⁴ In 2002, the structure of $[\text{C}_5\text{Me}_5\text{H}_2]^+$ was determined although it was by mistake addressed as an $[\text{C}_5\text{Me}_5]^+$ cation, probably due to its unexpected formation during the reaction of $\text{C}_5\text{Me}_5\text{H}$ with $[\text{Ph}_3\text{C}]^+$.²⁸⁵ Finally a silyl stabilized allyl cation was characterized, which formed *via* an interesting mechanism that starts with the formation of a silylium cation (Fig. 37).²⁸⁶

Ion-like carbon compounds. As mentioned before, in the analog structure of $[(m\text{-CF}_3\text{-C}_6\text{H}_4)(\text{Ph})\text{CF}]^+[\text{AsF}_6]^-$ instead of $[\text{AsF}_{11}]^-$, stronger interactions to the anions are present.²⁶⁸ The same applies to the related $\text{Me}_2\text{CF}(\text{AsF}_6)$.²⁶⁸ Also known is the ion-like $(\text{Me}_2\text{CH})(\text{HCB}_{11}\text{Me}_5\text{Br}_6)$, which displays a covalent C–Br distance of about 210 pm.²⁸⁷ Along with the latter, the preparation of $\text{H}_3\text{C}(\text{HCB}_{11}\text{Me}_5\text{Br}_6)$ and $\text{H}_3\text{C}(\text{HCB}_{11}\text{Me}_5\text{Cl}_6)$ was

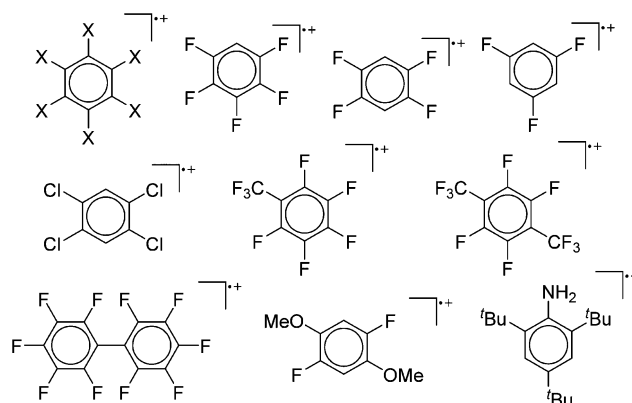


Fig. 36 Structurally characterized substituted benzene radical cation salts.



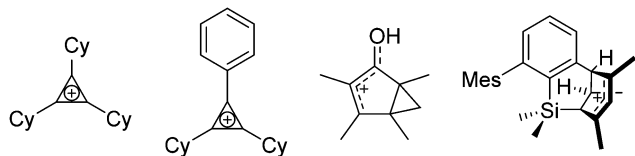


Fig. 37 Structurally characterized delocalized cation salts.

reported, but no structural data from XRD was presented. In 2010, the strongly methylating ion-like $\text{Me}_2\text{B}_{12}\text{Cl}_{12}$ was structurally characterized with a C–Cl bond length of 182 pm.²⁸⁸

Silicon

Silylium ions (CN = 3). Silylium ions are certainly amongst the most electrophilic cations known and thus exhibit an enormous Lewis acidity. Most of them are either stabilized by bulky ligands, or display a strong interaction with the corresponding WCA and have therefore to be categorized as ion-like compounds. In addition, the first claimed “stable silyl cation” $[\text{Et}_3\text{Si}]^+$ in 1993 contained a coordinating toluene ligand – a feature typical for many silylium ions.²⁸⁹ In order to obtain a truly tricoordinate silylium ion without stabilization through the anion or an additional ligand, bulkier substituents were needed. Hence, the first structurally characterized compound featuring a free silylium ion was $[\text{Mes}_3\text{Si}]^+[\text{HCB}_{11}\text{Me}_5\text{Br}_6]^-$ ²⁴ and in 2013, $([\text{Pemp}_3\text{Si}]^+)_2[\text{B}_{12}\text{Cl}_{12}]^{2-}$ was published (Fig. 38).²⁹⁰ The latter was afterwards also synthesized and characterized with $[\text{Al}(\text{OR}^{\text{PF}_6})_4]^-$.²⁹¹ In all three structures, the cation has no closer contacts to the anion.

Delocalized cyclic cations. Despite the early characterization of cyclopropenyl cations, the first example for a comparable silicon ion was published only in 2000 (Fig. 39, left).²⁹² In this compound, however, it is not the three-membered silicon-ring with a delocalized π -system but rather a silicon butterfly with one Si–Si– σ -bond stabilizing the positive charge. A direct equivalent of a cyclopropenyl cation was finally published in 2005 (Fig. 39).²⁹³ One more example is known with the positive charge being partially delocalized over four silicon atoms.²⁹⁴ An example for a Si(n) cation with 6π -aromaticity provides the silyliumylidene-like species introduced by Driess *et al.* (Fig. 39, right).²⁹⁵ This compound is stabilized by delocalization so that, although produced through protonation with $[\text{H}(\text{OEt}_2)_2]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$, no ether molecule remains coordinated to the cation.

Ligand-stabilized silicon cations. Already in 1983, pyridine stabilized $[\text{Me}_3\text{Si}]^+$ was reported.²⁹⁶ Yet, this compound is stable to such an extent that Br^- and I^- are sufficient as anions and that it



Fig. 38 Structurally characterized tricoordinate silylium ions.

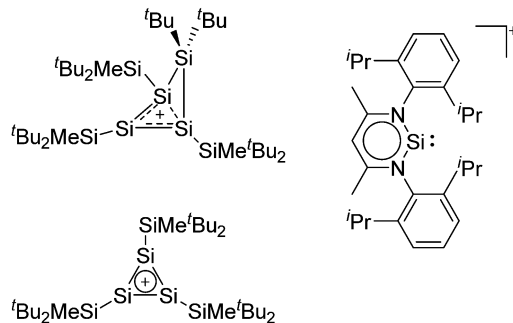


Fig. 39 Delocalized cyclic silicon centered cations.

can be prepared just by reacting Me_3SiX with pyridine. Many of these $[\text{R}_3\text{Si-L}]^+$ ions stabilized by different σ -donors are known: with nitriles,^{297–299} pyridine,^{296,300} water,³⁰¹ *o*-dichlorobenzene,⁸¹ sulfur dioxide⁸¹ and bipyridine.³⁰² Even though 2,6-bis(2,6-difluorophenyl)-phenyldimethylsilylium ion has no additional ligand acting as a σ -donor, the cationic center is stabilized by one fluorine of each 2,6-difluorophenyl-substituent (Fig. 40).³⁰³ With the stronger stabilizing DMAP, the dication $[\text{Me}_2\text{Si}(\text{dmap})_2]^{2+}$ has been synthesized.³⁰²

$[\text{R}_3\text{Si-L}]^+$ ions with π -donor ligands L = arenes like the before mentioned $[\text{Et}_3\text{Si}(\text{C}_7\text{H}_8)]^+$ are less stabilized than those with σ -donors (Fig. 42).²⁸⁹ Several different arene adducts of $[\text{Me}_3\text{Si}]^+$ were reported by Schulz and Villinger *et al.* (Fig. 42).³⁰⁴ As can be seen in Fig. 41, some of these compounds are coordinated by a second arene molecule binding in an η^6 -fashion to the proton *ipso* to the silylium center. This shows that these arene adducts are also very strong cationic Brønsted acids.

Comparable to the before mentioned 2,6-bis(2,6-difluorophenyl)phenyldimethylsilylium ion without any additional ligand, a 2,6-diaryphenyldimethylsilyl cation is existing, which is stabilized by intramolecular π -donation (Fig. 42).⁷⁶

Compounds of the type $[\text{R}_3\text{Si-X-SiR}_3]^+$ have to be treated as a special case of ligand stabilization. The first example of this type is the initially as $[\text{Et}_3\text{Si}]^+$ misinterpreted $[\text{Et}_3\text{Si-H-SiEt}_3]^+$,³⁰⁵ whose structure determination has been published about two years ago.³⁰⁶ $[\text{Me}_3\text{Si-H-SiMe}_3]^+$ ⁸¹ is also known as well as the analogous $[\text{Me}_3\text{Si-X-SiMe}_3]^+$ compounds with $\text{X} = \text{F}, \text{Cl}, \text{Br}, \text{I}$ ³⁰⁷ and trifluoromethanesulfonate.³⁰⁸ The X-bridged species are typically addressed as halonium ions, but it appears more reasonable to address them as ligand-stabilized silylium ions (see Fig. 43).

Calculations state that the positive charge is still located at the silicon atoms and F, Cl and Br are negatively charged.³⁰⁷ Only in the case of iodine, a small positive charge is located at the bridging atom.³⁰⁷ Additionally, bisilylated pseudohalonium cations $[\text{Me}_3\text{Si-X-SiMe}_3]^+$ with $\text{X} = \text{CN}, \text{OCN}, \text{SCN}$, and NNN are known.³⁰⁹ Of these, only in $[(\text{Me}_3\text{Si})_2\text{NNN}]^+$ both silyl groups are attached to the same atom,³⁰⁹ so that the structure of the cation is analog to the protonated hydrogen azide³¹⁰ (see Fig. 44 and Table 5 for $[\text{H}_2\text{N}_3]^+[\text{SbF}_6]^-$). Some more examples with bridged SiR_3 -groups, in which both groups are connected with each other, are known (Fig. 44).^{311–315}

A special case of intramolecular ligand stabilization can be observed in $[\text{FcSiMe}^t\text{Bu}]^+$.³¹⁶ Here the silicon is dipped towards the iron atom due to two $3c2e$ bonds between C_{ipso} , Si and Fe and C'_{ipso} , Si and Fe, respectively (Fig. 45).³¹⁶

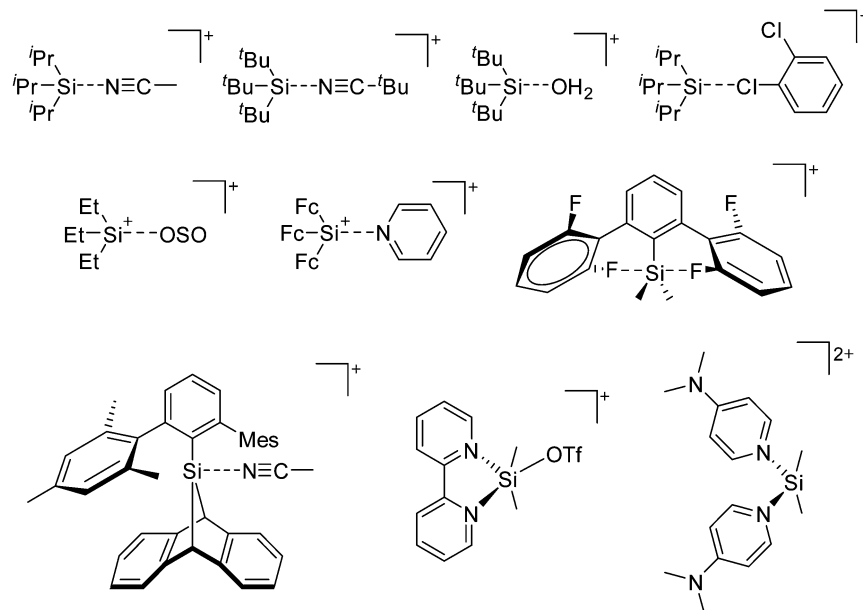


Fig. 40 Structurally characterized silylium ions stabilized by σ -donors.

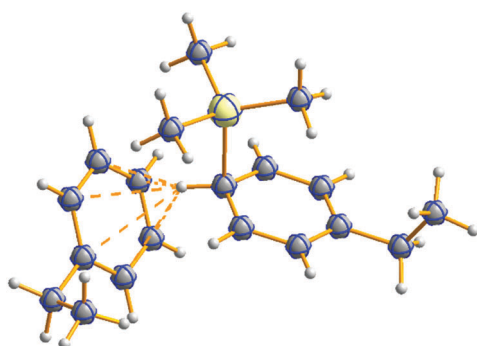


Fig. 41 Molecular structure of $[\text{Me}_3\text{Si}(\text{C}_8\text{H}_{10})\cdot(\text{C}_8\text{H}_{10})]^+$. The weak interaction between the stabilized cation and the adjacent ethylbenzene is indicated by the dashed bonds. M. F. Ibad, P. Langer, A. Schulz and A. Villinger, *J. Am. Chem. Soc.*, 2011, **133**, 21016–21027. Data from this reference were used to draw this figure.

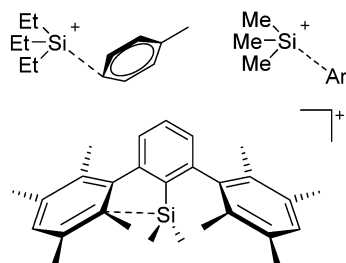


Fig. 42 Structurally characterized silylium ions stabilized by internal or external π -donors. Ar = benzene, toluene, ethylbenzene, *n*-propylbenzene, and iso-propylbenzene, *o*-xylene, *m*-xylene, *p*-xylene, 1,2,3-trimethylbenzene, 1,2,4-trimethylbenzene, mesitylene.

More intramolecular σ -donor stabilized silylium ions are known: $[\text{RSi}(\text{R}')_2]^+$ or $[\text{RSi}(\text{R}')(\text{R}'')]^+$ with R being a pincer ligand can be seen as an extra class of ligand-stabilized silicon cations.

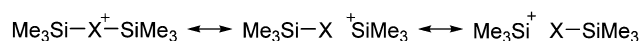


Fig. 43 Canonical structures of the halogen-bridged bis-silylium ions.

In 2009, several silylium ions with OCO and SCS pincer ligands were published by Jutzi *et al.* (Fig. 46).³¹⁷

All before mentioned ligand-stabilized silicon cations contain an inter- or intramolecularly by additional donor atoms stabilized $[\text{R}_3\text{Si}]^+$ cation. Two more different types of ligand-stabilized silicon cations were published with silicon in oxidation state +IV. Both were synthesized by oxidation of silicon(II) cations through elemental sulfur (Fig. 47).^{318,319} These cations containing subvalent silicon are very rare and most of the known examples are bearing a cyclopentadienyl substituent (see Cyclopentadienyl substituted cations). However, with well stabilizing ligands, two $[\text{LSiCl}]^+$ cations were synthesized (Fig. 47).^{318,320} Both are prepared just by adding the chelating ligand to NHC-SiCl_2 . The NHC ligand is being replaced by L and yields the $[\text{LSiCl}]^+$ cation with chloride as the anion. This shows that the silicon cationic center is largely stabilized by coordination. By using well stabilizing NHCs, it was possible to generate an $[(\text{L})(\text{L}')\text{Si}]^+[\text{I}]^-$ and even the dication $[\text{L}_3\text{Si}]^{2+}[\text{I}]^{2-}$.³²¹ In addition, two related silicon(II) monocations $[\text{RSi}(\text{L})_n]^+$ were structurally characterized in which the residue R is not a halogen atom (Fig. 48).³¹⁹

Cyclopentadienyl substituted cations. So far, two cyclopentadienyl substituted silicon cations without any further stabilization through additional ligands were structurally characterized. First $[(\text{C}_5\text{Me}_5)\text{Si}]^+$ with $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ was published in 2004³²² and two years later, the synthesis and characterization of $[(\text{C}_5^t\text{Pr}_5)\text{Si}]^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ was presented.³²³ However, the latter structure determination was of poor quality and did not allow to obtain any exact structural parameters. Additionally, two ether stabilized $[(\text{C}_5\text{Me}_5)\text{Si}]^+$ cations are known (Fig. 49).³²⁴



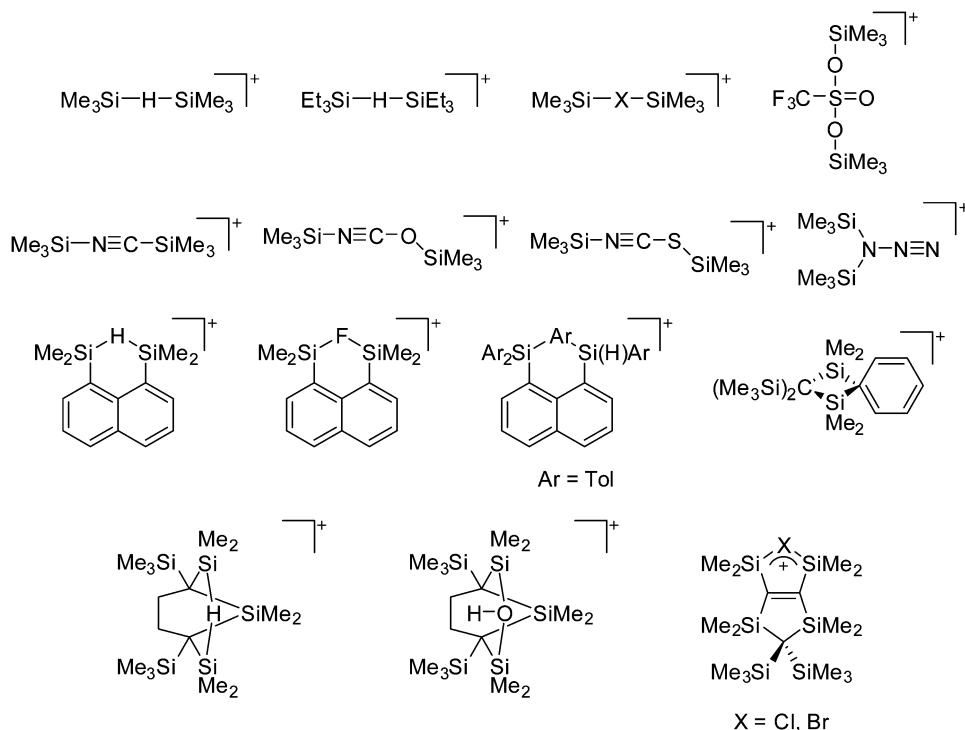


Fig. 44 Structurally characterized bridged bisilylium ion salts.



Fig. 45 Molecular structure of $[\text{FcSiMe}_4\text{Bu}]^+$. K. Mütter, R. Fröhlich, C. Mück-Lichtenfeld, S. Grimme and M. Oestreich, *J. Am. Chem. Soc.*, 2011, **133**, 12442–12444. Data from this reference were used to draw this figure.



Fig. 47 Structurally characterized ligand-stabilized silathionium cations.

first structurally characterized $\text{R}_3\text{Si}(\text{WCA})$ was the iso-propyl substituted compound in 1993.³²⁵ Today, at least one example with the most common alkyl substituents ^tBu , ^iPr , Et and Me is known.^{81,297,326–330} Additionally, with $^t\text{Bu}_2\text{MeSi}(\text{CB}_{11}\text{H}_6\text{Br}_6)$ one mixed substituted ion-like silylium compound was published.³²⁸ Another example might be $\text{Fc}_3\text{Si}(\text{OTf})$. However, its Si–O interaction is with 175 pm in the range of a normal Si–O bond.³³¹

Germanium, tin and lead

Cluster cations. As for silicon, no homopolyatomic cations comparable to the fullerene ions are known for germanium, tin and lead. Nevertheless, one example of a germanium cluster exists (Fig. 50).³³² The cluster is composed of ten germanium atoms, of which seven bear substituents. The remaining three unsubstituted germanium atoms carry the positive charge, which is evenly distributed.

Enium ions. Just as for silicon, enium ions of Ge, Sn and Pb need substituents with a high steric demand to shield the cationic center. The first example of the heavier elements of group 14 – $[\text{Bu}_3\text{Sn}]^+[\text{CB}_{11}\text{Me}_{12}]^-$ – does have, as expected,

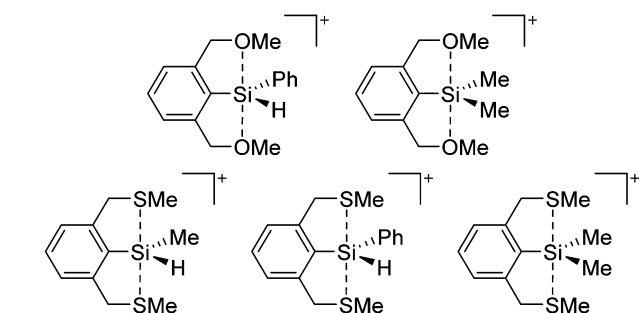


Fig. 46 Structurally characterized silylium ion salts with intramolecular stabilization by pincer ligands.

Ion-like silylium compounds. In alkylsilylium ion salts without an additional stabilizing ligand, interactions between the cation and the corresponding WCA can be observed. The

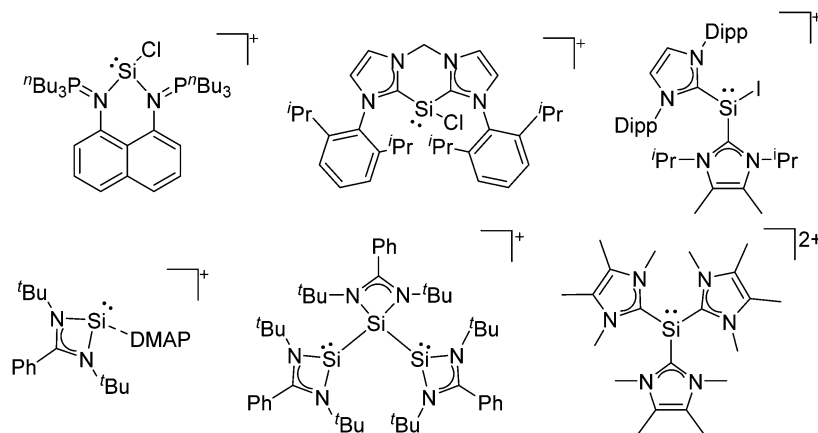


Fig. 48 Structurally characterized ligand-stabilized cations of subvalent silicon.

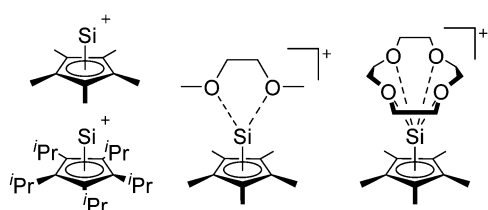


Fig. 49 Structurally characterized cyclopentadienyl substituted silicon cation salts with and without additional ligands.

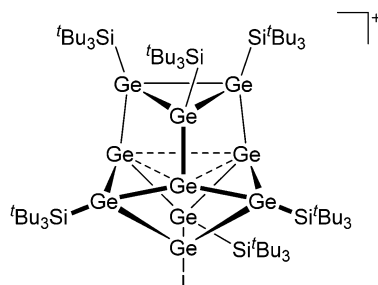


Fig. 50 Lewis structure of the 5-iodo-2,4,6,8,9,10-hexakis(tert-butylsilyl)heptacyclo[4.4.0.0.1.3.0.2.5.0.3.9.0.4.7.0.8.10]decagerman-7-ylum ion.

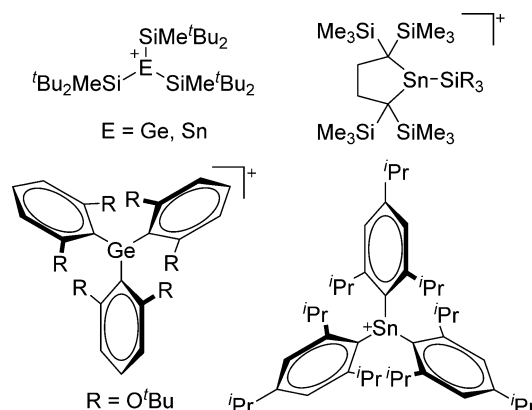


Fig. 51 Structurally characterized enium ions of germanium and tin.

interactions between cation and the $[\text{CB}_{11}\text{Me}_{12}]^-$ WCA.³³³ At about the same time, Lambert *et al.* and Sekiguchi *et al.* published the first examples of free enium ions of germanium and tin. While Lambert relied on bulky aryl substituents to synthesize $[(\text{Tipp})_3\text{Sn}]^+$,³⁴ Sekiguchi deployed silyl groups and managed to produce $[(\text{Bu}_2\text{MeSi})_3\text{Ge}]^+$ ³³⁴ and $[(\text{Bu}_2\text{MeSi})_3\text{Sn}]^+$,³³⁵ all with $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ as their counterpart (Fig. 51). Although enium ions with aryl substituents have always been under the first examples for carbon, silicon and tin, it kept lacking an example for germanium until in 2009 $[\text{Ge}\{2,6\text{-O}^t\text{Bu}_2\text{C}_6\text{H}_3\}_3]^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ was synthesized and characterized.³³⁶ However, the cationic center is stabilized by contacts to the oxygen atoms of the *tert*-butoxy residues at 286 and 288 pm.³³⁶ More recently, a mixed substituted enium ion of tin has been published (Fig. 51).³³⁷ Examples for lead are still missing and the only formally $[\text{R}_3\text{Pb}]^+$ containing ion-like

substance $\text{Et}_3\text{Pb}(\text{HCB}_{11}\text{H}_5\text{Br}_6)$ has like its Si, Ge and Sn analogs stronger interactions between the ions.³³⁸

Delocalized cyclic cations. The germanium compound $[\text{Ge}_3(\text{Si}^t\text{Bu}_3)_3]^+$, has been published long before the first silicon analog of a cyclopropenyl cation.³³⁹ Although it is known with a few different anions,^{340,341} it is still the only example of delocalized cyclic cations of the heavier elements of group 14 (similar to Fig. 39).

Ligand-stabilized. Far less ligand-stabilized cations of Ge, Sn and Pb in oxidation state +IV are known than of Si. $[\text{Me}_3\text{Sn}(\text{OPPh}_3)_2]^+[(\text{MeSO}_2)_2\text{N}]^-$ and $[\text{Ph}_3\text{Sn}(\text{OPPh}_3)_2]^+[(\text{MeSO}_2)_2\text{N}]^-$ were synthesized already in 1994³⁴² and six years later, the $[\text{Bu}_3\text{E}(\text{NC}^t\text{Bu})]^+$ cations were synthesized with $\text{E} = \text{Ge}$ and Sn , but only for the germanium compound the crystal structure is known.²⁹⁸ In addition, together with the analogous silicon complex, $[\text{Me}_2\text{Ge}(\text{bipy})(\text{OTf})]^+[\text{OTf}]^-$ has been published.³⁰² Interesting is however, that the corresponding substances with DMAP coordinating to germanium and the ones with DMAP or bipyridine coordinating to tin have to be described as ion-like, since in all of them both $[\text{OTf}]^-$ anions do have close contacts to the cationic center.³⁰² As already stated for silicon, symmetrical compounds of the type $[\text{R}_3\text{E-X-ER}_3]^+$ are somewhat special since the positive charge is evenly distributed and it is not possible to speak of an cation and a ligand anymore. Contrary to silicon, only one cation belonging to this type is known for the heavier homologues (Fig. 52).³⁴³ Additionally, for germanium



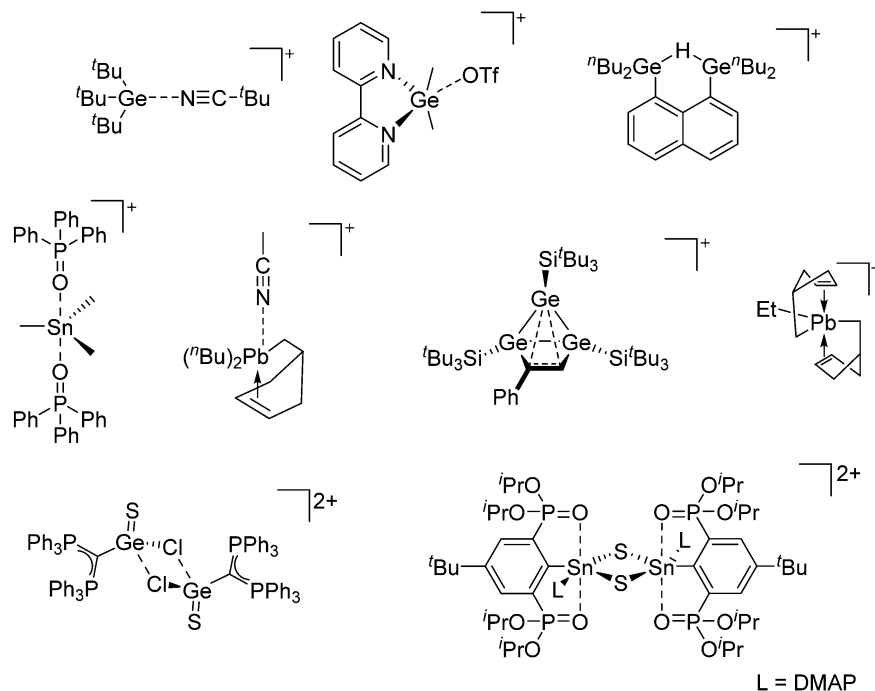


Fig. 52 Ligand-stabilized cations of germanium, tin and lead in oxidation state IV.

and tin ligand-stabilized dimeric cations are known, both synthesized by oxidation of their E(II) precursors through elemental sulfur (Fig. 52).^{344,345}

Norbornyl cations with the heavier group 14 elements were classified in here as ligand-stabilized cations, although one may address them as onium ions. Although the heavier norbornyl cation analogues were all published – also with silicon – no crystal structure could be determined.³⁴⁶ However, by addition of acetonitrile to the norbornyl cations, the stronger σ -donor replaces the weaker π -donating C=C double bond. An exception is the plumbanorbornyl cation, which gets coordinated by acetonitrile additionally and remains coordinated by the alkene (scXRD).³⁴⁶ A comparable π -stabilization as in the norbornyl cations can be found in the 1,4,5-trigermabicyclo[2.1.0]pent-2-en-5-ylum ion, in which the cationic center is coordinated intramolecularly by a C=C double bond.³⁴⁷ Another unique π -stabilization can be observed in bis(cyclopentenemethyl)plumbylium.³⁴⁸ This cation is intramolecularly stabilized by the C=C double bonds of the two cyclopentene substituents (Fig. 52).

Hard to classify are two germanium cations stabilized by a monoanionic bidentate bis(NHC)borate ligand (Fig. 53).³⁴⁹ Both originate from the attempt to synthesize a germanium dication stabilized by the before mentioned ligand through the reaction of LGeH with $[\text{Ph}_3\text{C}]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$. Instead of delivering the desired germanium dication, two different products were obtained. In one, instead of abstracting the hydride, the trityl cation attacks the lone pair of the Ge(II) cation, forming the adduct. In the other, the hydride is indeed abstracted by the trityl cation, but the resulting germanium dication is coordinated by unreacted starting material.

Apart from those examples, the ligand-stabilized cations of the heavier group 14 elements are in oxidation state +II. Already

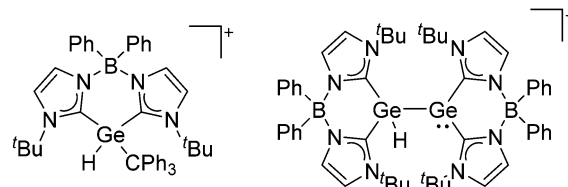


Fig. 53 Germanium cations stabilized by a monoanionic bidentate bis(NHC)borate ligand.

as early as 1989, $[\text{Sn}([15]\text{crown-5})_2]^{2+}$ has been published along with its crystal structure.³⁵⁰ This cation is accessible directly through the reaction of SnCl_2 with two equivalents of the crown ether, which is why $[\text{SnCl}_3]^-$ serves as the counterion. In this or a similar fashion it has been possible to synthesize a portfolio of different crown ether complexes of tin(II) and lead(II).^{351–353}

To isolate the first related Ge(II) compound, better stabilizing ligands were needed. By employing NHC ligands, a germanium dication was isolated (Fig. 54).³⁵⁴ The germanium center is highly stabilized by its ligands, and – although iodides are the counterions – only weak interactions between the ions are present. Another germanium containing dication was synthesized with the encapsulating cryptand[2.2.2],³⁵⁵ and a few years later the analogous tin complex.³⁵⁶ Today, quite a few different crown ether complexes of germanium are known as well (Table 9).³⁵⁷ By using other well stabilizing chelating N-donor ligands, it was also possible to isolate $[(\text{L})\text{Ge}]^{2+}$ cations.³⁵⁸

The autoionization reaction used for the preparation of many of the crown ether complexes has also been applied to synthesize most of the structurally characterized $[(\text{L})\text{EX}]^+[\text{WCA}]^-$ compounds of germanium and tin (Fig. 55).^{344,359–362,389} With even stronger donating ligands, comparable salts $[(\text{L})\text{GeCl}]^+\text{Cl}^-$ were

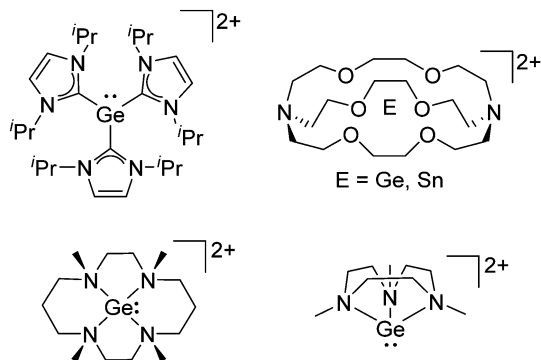


Fig. 54 Structurally characterized dicationic compounds of germanium and tin.

prepared.^{363,364} These compounds are strongly stabilized so that even halides are sufficient as anions (Fig. 56). Related $[RE(L)]^+$ cations with the residue R not being an halogen atom are also known. In these cations, the residue is capable to stabilize the cationic center by an additional σ - or π -donation (Fig. 57).^{80,345,365–368} In case of bulky residues it was possible to work without an additional ligand and to obtain the free $[RE]^+$ cations (Fig. 57).^{365,366,369} For lead, one additional $[RE(L)]^+$ cation is known with R being a bulky aryl ligand and with a toluene molecule coordinating to the lead atom.³⁷⁰

A rather special case is $[Sn(C_7H_8)_3]^{2+}$, in which a tin(II) cation is coordinated by three toluene molecules.³⁷¹ Although lots of arene complexes of tin(II) are known, almost all of them do still have strong interactions to the anions, mostly halides and/or $[AlCl_4]^-$ (see for some examples ref. 19). An exception is the Sn(II) complex with [2.2.2]paracyclophane.³⁷² Only one of the two $[AlCl_4]^-$ ions is coordinated to the tin atom, the other one does not have interactions with



Fig. 56 Ligand-stabilized cations $[L)GeCl]^+$ with chloride as their counterion.

the cation. However, $[Sn(C_7H_8)_3]^{2+}([B(C_6F_5)_4]^-)_2$ is the first example of a tin(II) complex with independent arenes and without additional stabilization by the anion (Fig. 58).

Cyclopentadienyl substituted cations. The tin analog of the $[(C_5Me_5)Si]^+$ cation was already published in 1979,³⁷³ about 25 years before the silicon compound was characterized by XRD. This is due to the fact, that $[(C_5Me_5)Sn]^+$ could be synthesized as its $[BF_4]^-$ salt, which is not possible in case of $[(C_5Me_5)Si]^+$ because of its instantaneous decomposition.³⁷⁴ In $[(C_5Me_5)Sn][BF_4]$ are still some stronger interactions present between the fluorine atoms and tin. In 2005, the structure of the $[(C_5Me_5)Sn][B(C_6F_5)_4]$ was determined in which these interactions are a lot weaker³⁷⁵ and with the same anion, $[(C_5Me_5)Pb]^+$ was synthesized and structurally characterized.³⁷⁵ The sole exception is germanium, whose $[(C_5Me_5)Ge]^+$ was only characterized by XRD with $[BF_4]^-$ ³⁷⁶ and $[SnCl_3]^-$ ³⁷⁷ as its counterion and not with any larger WCA. In addition, interesting triple-decker cations are known for tin and lead: $[(C_5Me_5)_2Sn_2(\mu-Me_5C_5)]^+$ was first synthesized and structurally characterized with the $[Ga(C_6F_5)_4]^-$ anion,³⁷⁸ its structure was subsequently published with $[B(C_6F_5)_4]^-$ together with the analogous lead compound (Fig. 59).³⁷⁵

Ion-like compounds of germanium, tin and lead. As for silicon, alkyl substituted enium ions of the heavier group 14 elements without any additional ligand need stabilizing

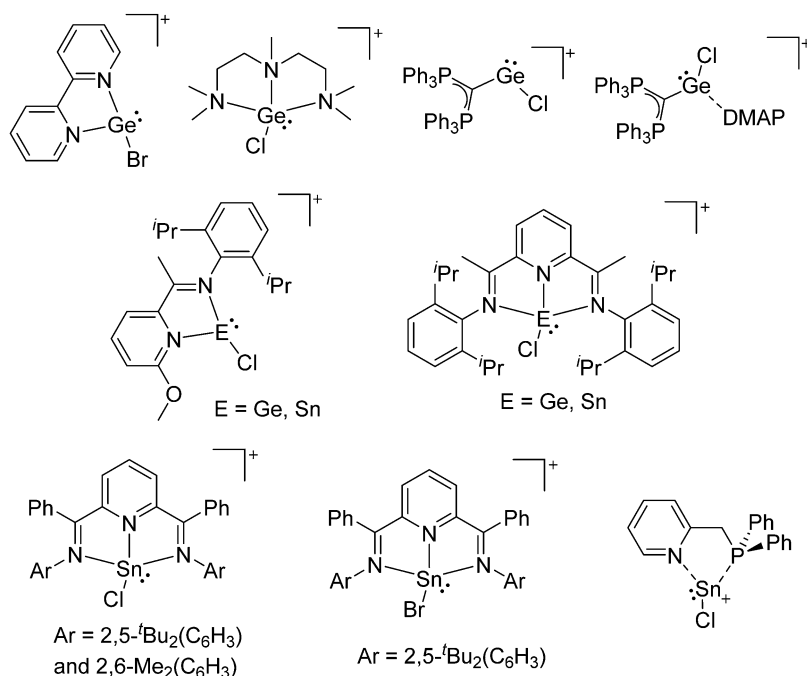


Fig. 55 Ligand-stabilized cations $[L)EX]^+$ of germanium and tin.



Fig. 57 Structurally characterized $[RE(L)]^+$ and $[RE]^+$ cations of germanium, tin and lead.

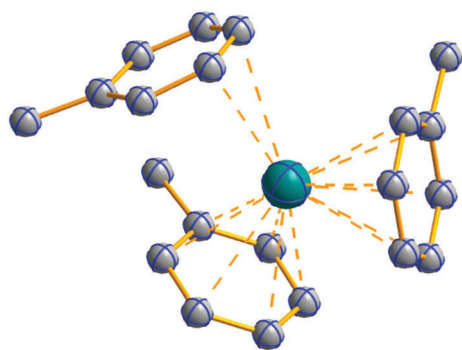


Fig. 58 Molecular structure of $[Sn(C_7H_8)_3]^{2+}$. A. Schäfer, F. Winter, W. Saak, D. Haase, R. Pöttgen and T. Müller, *Chem. – Eur. J.*, 2011, **17**, 10979–10984. Data from this reference were used to draw this figure.

interactions with the WCA. However, far less examples are known for $E = Ge, Sn$ and Pb , although already in 2000, the first example was published with $^nBu_3Sn(CB_{11}Me_{12})$.³³³ The other known examples are the $Et_3E(HCB_{11}H_5Br_6)$ compounds already mentioned before.³³⁸

Transition-metal substituted cations of germanium, tin and lead. Other than for silicon, more transition metal coordinated cations are known for the heavier elements of group 14, especially for tin. *Via* a salt elimination reaction, the complex cation $[(dppe)_2W \equiv Sn-C_6H_3-2,6-Me_2]^+$ was synthesized with $[PF_6]^-$ as its counterion in which the $W-Sn-C$ angle is close to 180° .³⁷⁹ A similar germanium compound was published one

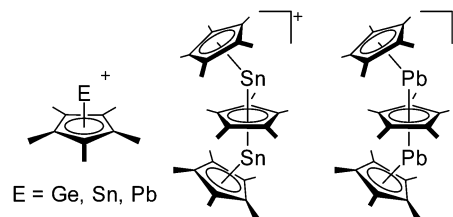


Fig. 59 Structurally characterized cyclopentadienyl substituted cations of germanium, tin and lead.

year after, in 2004. In $[(MeCN)(dppe)_2W \equiv Ge-(\eta^1-Cp^*)]^+$, the germanium is substituted by a Cp^* and the tungsten atom is coordinated additionally by an acetonitrile molecule.³⁸⁰ As WCA serves $[B(C_6F_5)_4]^-$ in this case. A new complex cation featuring a $Sn-Pt$ bond was published in 2010. In $trans-[Pt(Me)(SnCl_2)(2-PyPPh_2)_2][BF_4]$, the tin atom is pentacoordinated and adopts a trigonal-bipyramidal geometry.³⁸¹ Along with that, comparable compounds were synthesized with the remaining group 10 metals, but no crystal structure determination was performed. By using an OCO-pincer ligand, a chromiumpentacarbonyl coordinated tin(II) cation was synthesized. Two variants were published, $\{2,6-(MeOCH_2)_2C_6H_3\}(H_2O)SnCr(CO)_5(OTf)$, in which the tin is coordinated additionally by a water molecule and $\{2,6-(MeOCH_2)_2C_6H_3\}(THF)_2SnCr(CO)_5^+[CB_{11}H_{12}]^-$, in which the tin is hexacoordinated with two THF molecules complementing the coordination sphere.³⁸² The former has indeed no contact between the triflate and the tin atom, but a strong hydrogen bond



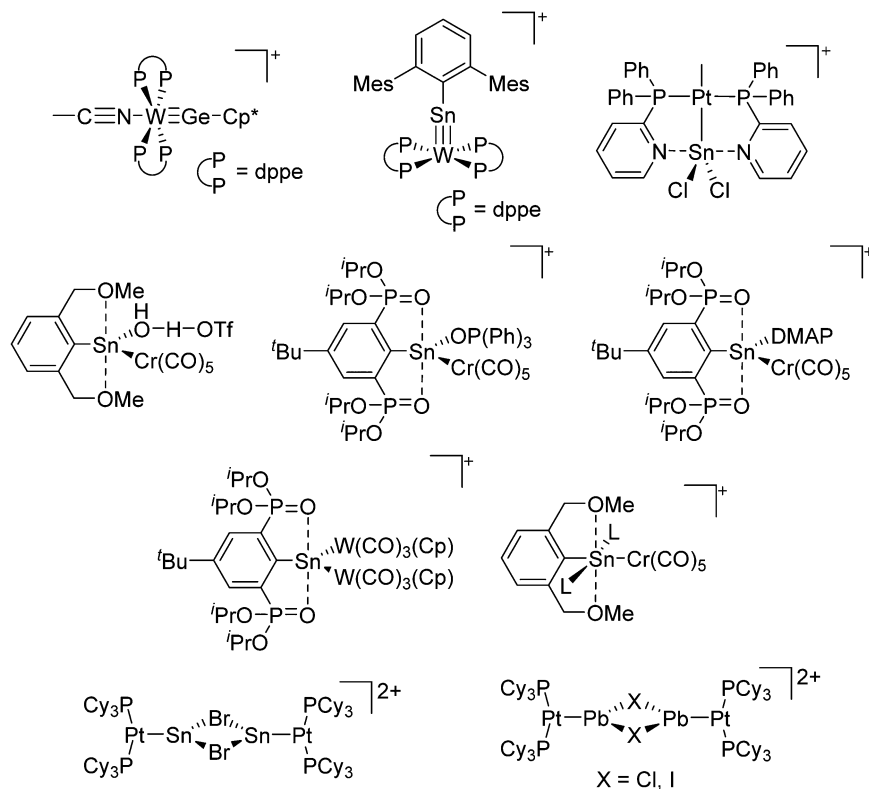


Fig. 60 Structurally characterized transition metal substituted cations of germanium, tin and lead.

between the coordinated water molecule and the anion is existing, with an O–O distance of about 261 pm. Two more chromiumpentacarbonyl coordinated tin(II) cations were published in 2013, both also with a pincer-type ligand (Fig. 60).³⁸³ The same ligand was used to prepare the $[\text{R}(\text{Sn}\{\text{W}(\text{CO})_3\text{Cp}\}_2)]^+$, with $\text{R} = \text{R} = 4\text{-}^t\text{Bu-2,6-}\{\text{P}(\text{O})(\text{O}^i\text{Pr})_2\}_2\text{C}_6\text{H}_2$ and $[\text{W}(\text{CO})_3\text{Cp}]^-$ as its counterion.³⁸⁴ Recently, new platinum-coordinated cations of tin and lead were published. Starting from $(\text{Cy}_3\text{P})_2\text{Pt}(\text{SnBr}_2)$, $[(\text{Cy}_3\text{P})_2\text{Pt-SnBr}]^+$ was synthesized with two different anions.³⁸⁵ The analogous dimeric lead cation $[(\text{Cy}_3\text{P})_2\text{Pt-PbCl}]^+$ was accessible by using $(\text{Cy}_3\text{P})_2\text{Pt}(\text{PbCl}_2)$ as a starting material.³⁸⁵ Through further reaction with AlX_3 , $\{(\text{Cy}_3\text{P})_2\text{Pt}\}_2\text{Sn}(\text{AlBr}_4)_2$ respectively $\{(\text{Cy}_3\text{P})_2\text{Pt}\}_2\text{Pb}(\text{AlCl}_4)_2$ were synthesized.³⁸⁵ However, in both dications some interactions between the ions are present. Additionally, the dimeric lead cation was also synthesized with iodine instead of chlorine.³⁸⁵

Group 15 cations

Of all the pnictogen elements, especially phosphorus has a rich cation chemistry. The analogy between CR_4 and $[\text{PR}_4]^+$ displays the possibility of creating a large variety of cationic phosphorus frameworks (Fig. 61).

Over the last decades a multitude of catenated phosphorus cations were synthesized. The classical phosphino-phosphonium cation (Fig. 62, left), which can be synthesized through halide abstraction from PR_2Cl and formal insertion/coordination of the resulting $[\text{PR}_2]^+$ (see section “Oxidation state +III” below) into a $\text{R}_2\text{P-R}$ bond/to PR_3 stands for an entire substance class of compounds typically containing organic residues R .³⁹⁰ However,

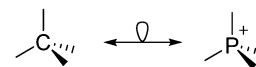


Fig. 61 Analogy between cationic phosphorus atom and carbon.

we refer the interested reader to the multitude of recent reviews especially on these cations,^{391,392} the analogous interpnictogen cations (Fig. 63)³⁹³ and other types of cationic pnictogen compounds (Table 5).

Homopolyatomic cations. Except for the long-known bismuth cations, all homopolyatomic pnictogen cations were synthesized in the last 16 years. In late 1999 the third all-nitrogen molecule $[\text{N}_5]^+$ – besides N_2 and N_3^- – was prepared through a reaction of $[\text{N}_2\text{F}]^+[\text{AsF}_6]^-$ and HN_3 .⁴⁰⁰ The obtained compound $[\text{N}_5]^+[\text{AsF}_6]^-$ is explosive but an anion exchange led to the more stable $[\text{N}_5]^+[\text{SbF}_6]^-$ and to the crystal structure of $[\text{N}_5]^+[\text{Sb}_2\text{F}_{11}]^-$.⁴⁰¹ In 2004 the reduction of SbCl_3 with $[\text{Ga}]^+[\text{GaCl}_4]^-$ in a GaCl_3 -benzene solution led to the square-antiprismatic *arachno*- $[\text{Sb}_8]^{2+}$ Wade cluster cation.⁴⁰² Recently also the first – formally electron precise and Zintl type – phosphorus cation $[\text{P}_9]^+$ was synthesized through the reaction of P_4 and the nitrosyl salt of the $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ WCA.²⁹ By contrast, bismuth has a rich cation chemistry. The first structure of a bismuth cation was measured already in 1962. Most of them were synthesized through high temperature solid state reactions. The newer room temperature approaches are based on ionic liquids.⁴⁰³ Normally the clusters formed are badly soluble, but there is evidence that the use of very weakly coordinating anions like $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ can lead to $[\text{Bi}_n]^+$ clusters, which are soluble in





Fig. 62 Examples for catenated phosphorus cations in the formal phosphino-phosphonium or diphosphonium form. Also cyclic versions are available.



Pn = As, Sb

Fig. 63 Example for a catenated interpnictogen cation.

solvents like CH_2Cl_2 or SO_2 .⁴⁰⁴ Besides the lighter noble gases and fluorine, only arsenic has still no homopolyatomic cation. Yellow arsenic (As_4), which has now a relatively stable storage form (see section “Metal–pnictogen complexes”) might be a good starting point for a future synthesis (Fig. 64).

Metal–pnictogen complexes. There are still only a few complexes with pnictogen modifications as ligands in the literature. Early examples of tetrahedro- P_4 complexes like $(\text{PPh}_3)_2\text{Rh}^{\text{I}}\text{Cl}(\eta^2\text{-P}_4)$ are better viewed as phosphide complex $(\text{PPh}_3)_2\text{Rh}^{\text{III}}(\text{P}_4^{2-})$. By contrast, the d^{10} -metal cation Ag^+ is ideal for the stabilization of the non-metallic clusters and the electronic structure of the ligand stays relatively unaffected (see also chapter chalcogen cations). In 2001, the WCA $[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ made it possible to crystallize the $[\text{Ag}(\text{P}_4)_2]^+$ complex and later through salt metathesis with CuI also the copper complex $[\text{Cu}(\text{P}_4)_2]^+$ was accessible. In 2012 the gold complex was obtained as $[\text{GaCl}_4]^-$ salt and completed the whole series $[\text{M}(\text{P}_4)_2]^+$ ($\text{M} = \text{Cu}, \text{Ag}, \text{Au}$). Recently light-stable (!) $[\text{Ag}(\text{As}_4)_2]^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$ was synthesized, which finally serves as a good storage form of yellow arsenic (As_4). As such, As_4 is both thermally and photochemically unstable. The salt made it possible to transfer the As_4 tetrahedron to gold in $[\text{Ph}_3\text{PAu}(\text{As}_4)]^+$ and opens new possibilities in the synthesis of arsenic complexes (Fig. 65).⁴⁰⁵

Diazonium cations and heavier homologues. There is a multitude of crystal structures of different cluster or cluster-like cations that contain pnictogen atoms.^{391–393,397} We decided to give

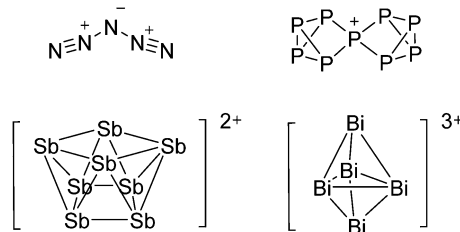


Fig. 64 Examples for homopolyatomic pnictogen cations (only for bismuth, several entries are known).

an overview and to list parent (model)-compounds like $[\text{N}_2\text{Ph}]^+$ and $[\text{N}_2\text{Mes}]^+$ in case of diazonium cations⁴⁰⁶ as examples for the entire diazonium substance classes. The heavier homologues of the diazonium cations $[\text{RNPn}]^+$ ($\text{Pn} = \text{P}, \text{As}$) need sterically demanding groups like Mes^* ($2,4,6\text{-}^t\text{Bu}_3\text{C}_6\text{H}_2$) to protect the highly reactive triple bonds (Fig. 66).^{407,408}

Cluster and cage cations. The reaction of phosphorus halides PX_3 with halide abstractors led to very reactive carbene-analogous $[\text{PX}_2]^+$ cations (see chapter “Phosphenium ions”), which are able to formally insert in the $\text{X}_2\text{P-X}$ bonds of a second equivalent to form $[\text{P}_2\text{X}_5]^+$ clusters or in the P–P bond of white phosphorus to produce $[\text{P}_5\text{X}_2]^+$ for instance. Insertion in P_4S_3 or halide abstraction from $\text{P}_4\text{S}_3\text{I}_2$ followed by rearrangements led to the phosphorus–sulphur cluster cations $[\text{P}_5\text{S}_2\text{X}_2]^+$ ¹³⁴ and $[\text{P}_7\text{S}_6\text{I}_2]^+$.²³⁴ Ref. 134 contains investigations on the nature of this formal insertion reaction, which is not as simple as thought and rather follows a concerted, orbital controlled mechanism (Fig. 67).

The binary group 15 and 16 cations have also a strong tendency to form clusters. The newer examples like the antimony–chalcogen cations $[\text{Sb}_{10}\text{Se}_{10}]^{2+}$ and $[\text{Sb}_7\text{Te}_8]^{5+}$ were synthesized in ionic liquids or GaCl_3 melts.^{409,410} Very recently $[\text{P}_3\text{Se}_4]^+$, the first binary P–Se-cation was characterized by six different groups with three different approaches.⁴¹¹ It is accessible from solution, but also through solid state syntheses.⁴¹² In 2004 the synthesis of the sulphur- and selenium–bismuth cations from a chloroaluminate melt completed the series of the heterocubane cluster cations $[\text{Bi}_4\text{Ch}_4]^{4+}$ ($\text{Ch} = \text{S}, \text{Se}, \text{Te}$) (Fig. 68).^{413,414}

$(4n + 2)\pi$ -cations. The pnictogen cations with planar delocalized π -systems can be described as (pseudo-)aromatic systems. The four-membered rings were all synthesized through halide abstraction from the neutral rings with two halogen atoms. The five-membered As–N ring was prepared through cycloaddition of the highly reactive

Table 5 Review articles including cationic group 15 compounds

Year	Title	Ref.
2004	Homoatomic cages and clusters of the heavier group 15 elements. Neutral species and cations	394
2008	Catena–phosphorus cations	392
2011	Homo- and heteroatomic polycations of groups 15 and 16. recent advances in synthesis and isolation using room temperature ionic liquids	66
2012	Multiple-charged P_1 -centered cations: perspectives in synthesis	395
2013	Catenated phosphorus compounds	391
2013	Recent advances in the syntheses of homopolyatomic cations of the non-metallic elements C, N, P, S, Cl, Br, I and Xe	11
2013	Catenated compounds – group 15 (As, Sb, Bi)	396
2014	Interpnictogen cations: exploring new vistas in coordination chemistry	393
2014	The chemistry of cationic polyphosphorus cages – syntheses, structure and reactivity	397
2015	Coordination chemistry of homoatomic ligands of bismuth, selenium and tellurium	398 and 399



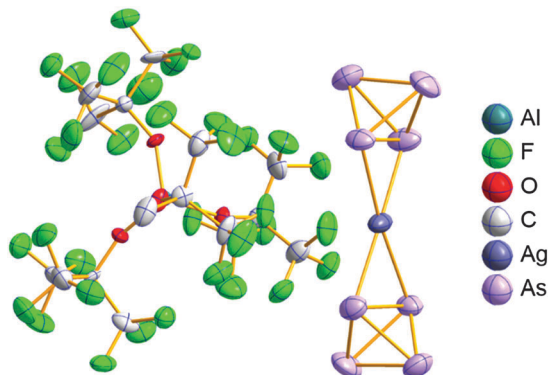


Fig. 65 Molecular structure of $[\text{Ag}(\text{As}_4)_2]^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$. C. Schwarzmaier, M. Sierka, M. Scheer, *Angew. Chem., Int. Ed.*, 2013, **52**, 858–861. C. Schwarzmaier, M. Sierka and M. Scheer, *Angew. Chem.*, 2013, **125**, 891–894. Data from this reference were used to draw this figure. The disorder of the anion was omitted for clarity.

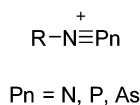


Fig. 66 The diazonium cation and its heavier homologues. R e.g. Mes*.

$[\text{AsNMe}_3]^+$, which reacts as dienophile with the 1,3 dipole trityl azide N_3CPh_3 (Fig. 69).

$\pi^*-\pi^*$ -complexes. Like the chalcogen compounds, the pnictogen cations containing $\pi^*-\pi^*$ -interactions can be described as dimers of chalcogen radicals, whose half-occupied interacting orbitals have π^* -character. The interannular $\pi^*-\pi^*$ -bonds between the “monomers” are relatively weak. They were synthesized through halide abstraction from the chlorides of the monomers (Fig. 70).⁴¹⁵

Radical cations. Radical cations of pnictogens can be obtained through direct oxidation of Pn_2 fragments with stabilizing ligands like N-heterocyclic carbenes (NHC) or cyclic alkylaminocarbenes (CAAC). As one-electron-oxidants the trityl salt of $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ was used (Fig. 71).^{208,416,417}

Bulky arylphosphines and -diphosphines (Fig. 72) can also be oxidized to their radical cations, if the cation is stabilized by a WCA.^{418,419}

The delocalization of the single electron over a ring system also leads to stabilization. The four-membered radical cation ring systems with different pnictogen atoms (Fig. 73) were obtained through direct oxidation with silver and nitrosyl salts of WCAs.⁴²⁰

Formal oxidation state +I. In some compounds like $[\text{P}_3\text{Ph}_6]^+$ ⁴²¹ the oxidation state of the central pnictogen “P⁺” can be described as +I, which is for example supported by the unusual high field shift in ³¹P-NMR of the central phosphorus atom in these cations (−210 to −270 ppm).⁴²² In case of the ligand-stabilized arsenic cation $[\text{AsDppDIMPY}]^+$ this is also supported by the synthesis: $(\text{DppDIMPY} = [\alpha, \alpha' - \{2, 6 - \text{Pr}_2\text{PhNC}(\text{Me})\}_2(\text{C}_5\text{H}_5\text{N})])$. The reduction of AsCl_3 with SnCl_2 led to a cation with a planar carbenoid-like structure.⁴²³ Under the same conditions with a different ligand an arsa-carbenoid of type $[\text{As}(\text{NR})_2\text{C}_2\text{H}_2]^+$ was obtained (Fig. 78). This displays the difficulty of a clear assignment of oxidation states in such systems (Fig. 74).

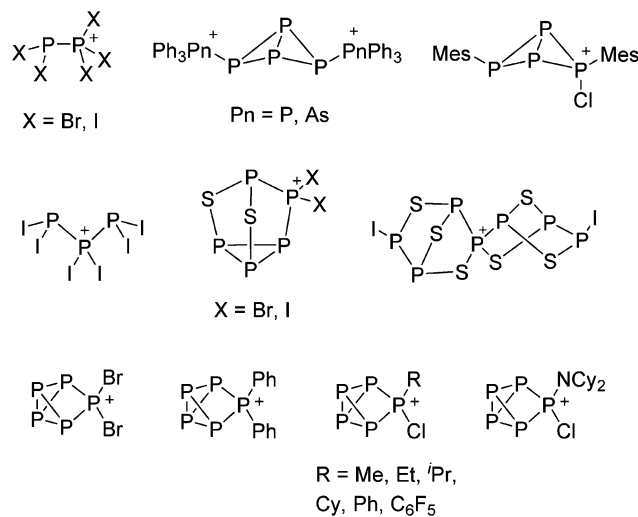


Fig. 67 Typical examples for phosphorus cations.

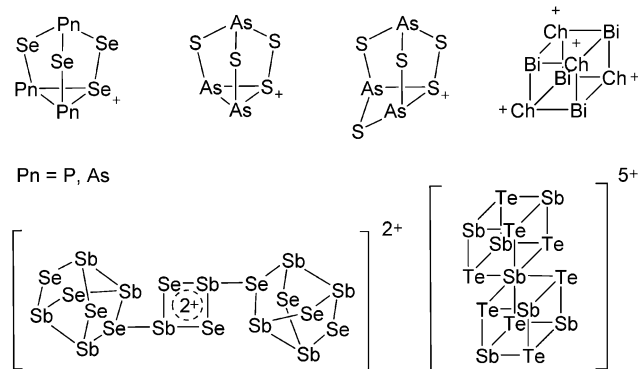


Fig. 68 Examples for binary pnictogen containing cations.

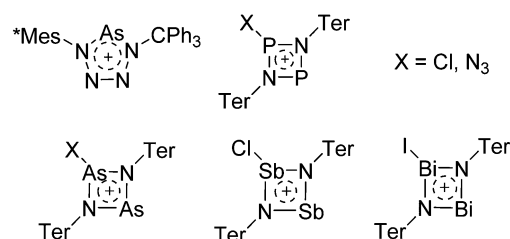


Fig. 69 Heteroatomic, cationic aromatic 6 π -systems containing pnictogen atoms.

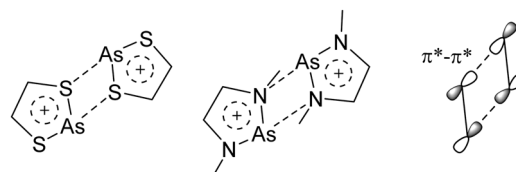


Fig. 70 Arsenic cations containing $\pi^*-\pi^*$ -interactions.

Phosphenium ions (oxidation state +III). The chemistry of the highly reactive phosphenium ions $[\text{P}(\text{R/Y})_2]^+$ (Fig. 75) was part of many studies in the past. The stability increases with the π -donor-ability



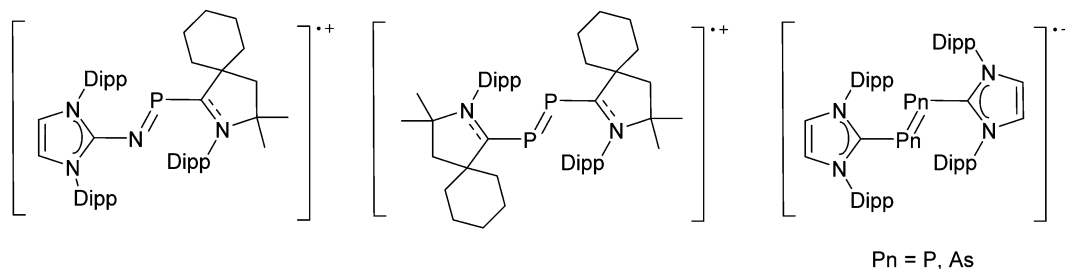


Fig. 71 Cationic phosphorus radicals stabilized by NHCs or CAACs.

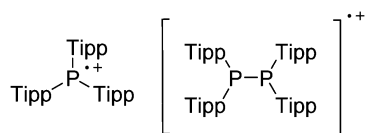


Fig. 72 Phosphorus radical cations.



Fig. 75 Structurally characterized simple phosphonium ions.

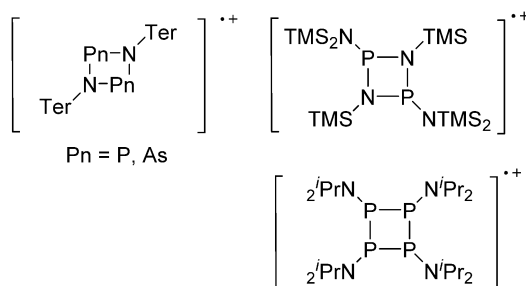


Fig. 73 Cyclic pnictogen radical cations.

of the substituent and the Lewis acidity with a stronger negative inductive effect (FIA: $[\text{P}(\text{NH}_2)_2]^+ < [\text{P}(\text{Cl})(\text{NH}_2)]^+ < [\text{P}(\text{Cl})_2]^+$).⁴²⁴

For most of the reactive phosphorus cations, the decomposition is normally accompanied by the formation of strong P–X bonds (X = F, Cl...). This makes it necessary to use weakly coordinating anions stable against electrophilic cations. For the homoleptic halogen substituted cations, extremely weak anions are needed. The first examples of the less, but still highly reactive mixed amino-halogen substituted phosphonium cations were published already in 1976. Through the use of a halide-abstractor (MCl_3 , M = Al, Ga, Fe) it was possible to prepare $[\text{P}(\text{NR}_2)\text{Cl}]^+$ (R = Me, Et, ^iPr) (X = $[\text{AlCl}_4]^-$) but no crystallographic data was obtained. It was not until 2012 that the first crystal structure of a halogen and a pseudohalogen mono-substituted phosphonium cation was determined. The structures of $[\text{P}(\text{NR}_2)\text{X}]^+$ (R = TMS; X = Cl, N_3 , NCO, NCS) and (R = ^iPr ; X = Cl, N_3) were determined by scXRD. All cations were stabilized with the $[\text{GaCl}_4]^-$ anion. Especially the

azidophosphonium compound turned out to be a versatile starting material for further chemistry, and made it possible to derive more complex phosphorus-centered cations like iminophosphorane-substituted-phosphonium salts $[\text{Pr}_2\text{NPNP}(\text{Cl})_2\text{NR}_2][\text{GaCl}_4]^-$ [R = ^iPr , SiMe_3] (Fig. 76) – for instance through the reaction with the corresponding chlorophosphane R_2NPCl_2 .

Miscellaneous cations in oxidation state +III. There are also some examples of the heavier homologues in oxidation state +III (Fig. 77). They were typically synthesized through halide abstraction with Lewis acids.

Another example of ligand-stabilized pnictogen cations are the N-heterocyclic carbenoid rings $[\text{Pn}(\text{NR})_2\text{C}_2\text{H}_2]^{2+}$ (Pn = P, As, Sb), which are formally 1,4-diaza-1,3-butadiene complexes of a pnictogen cation in oxidation state +III, but the delocalization of the positive charge supports also a description as a neutral pnictogen atom. In case of the 1,3,2-diazaphospholidinium rings $[\text{Pn}(\text{NR})_2\text{C}_2\text{H}_4]^+$ (Pn = P, As) the double bond between C4 and C5 is missing.

Reactive pnictonium cations (oxidation state +V). The halo-pnictonium cations $[\text{PnX}_4]^+$ with pnictogen atoms in oxidation state +V (Fig. 79), have a very different presence in the literature. For phosphorus, all four cations (X = F, Cl, Br, I) have been synthesized but for $[\text{PF}_4]^+$ no crystal structure was determined. A multitude of structures of $[\text{PX}_4]^+$ with different anions was characterized, but only a few of $[\text{AsX}_4]^+$ and $[\text{SbX}_4]^+$ are known. The cations are normally prepared from PnX_3 , X_2 and a Lewis acid.

But there are also some newer, highly oxidized cations in the literature: the formal $[\text{PnPh}_3]^{2+}$ cations (Fig. 80), which have a



Fig. 74 Phosphorus and arsenic cations in formal oxidation state +I.

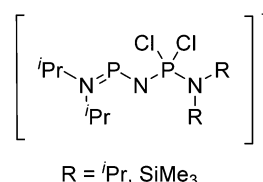


Fig. 76 Iminophosphorane-substituted phosphorus cation.

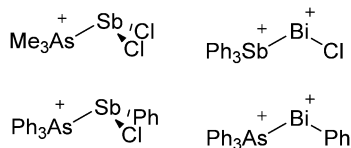


Fig. 77 Pnictogen cations with pnictogen atoms in formal oxidation state +III.

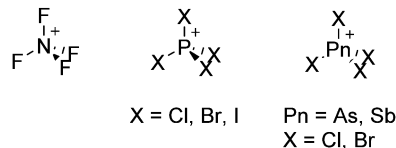


Fig. 79 Classical halo-pnictonium cations in oxidation state +V.

strong contact to the anion, serve as useful starting materials for further coordination chemistry of Pn^{V} compounds.⁴²⁵ The ligand stabilized formal “ PO^+ ” cations were prepared through the oxidation of a phosphorus carbenoid (Fig. 80, see also the phosphorus carbenoids above) with the amine-*N*-oxides Me_3NO and pyO .⁴²⁶ The charge of the carbene-stabilized formal “ $[\text{PFPh}_2]^{2+}$ ”, which was prepared from the carbene-stabilized “ $[\text{PF}_2\text{Ph}_2]^+$ ” through fluoride abstraction is likely partially localised on the strongly bound ligand (Fig. 80).⁴²⁷

Protonated cations. With the super acidic system HF/MF_5 ($\text{M} = \text{As}, \text{Sb}$) it is possible to protonate hydrazoic or phosphoric acid for instance and obtain the aminodiazonium $[\text{H}_2\text{N}_3]^+$ and phosphatidium (tetrahydroxyphosphonium) cation $[\text{P}(\text{OH})_4]^+$ (Table 10). The structure determinations of the $[\text{SbF}_6]^-$ salts revealed the structures of the cations (Fig. 81).⁴²⁸

Group 16 cations

Hundreds of chalcogen cations are known to the literature (see Table 6 for reviews). The relatively strong $\text{Ch}-\text{Ch}-$ and $\text{Ch}-\text{X}$ -single bonds ($\text{Ch} = \text{S}, \text{Se}, \text{Te}$; $\text{X} = \text{F}, \text{Cl}, \text{Br}, \text{I}$) led to a great diversity of reactive compounds, which include homo- and heteropolyatomic clusters, radical cations and a large number of different $[\text{ChX}_3]^+$ structures for instance. To avoid the formation of the more stable neutral compounds, weakly coordinating anions are needed to stabilize the reactive chalcogen cations.

Homopolyatomic cations. The first observation of homopolyatomic cations were the colored solutions of elemental sulfur, selenium and tellurium in sulfuric acid in the 18th and 19th century. Over the next centuries, the nature of these solutions stayed unclear and it was not before the middle of the 20th century that the use of superacidic media and better analytical methods made it possible to characterize the responsible species. Since then, starting with $[\text{O}_2]^+[\text{PtF}_6]^-$ in 1962,⁵¹⁰ the crystal structures of a multitude of different homopolyatomic chalcogen cations were measured in the last 50 years (Fig. 82). All of them have more or less weakly coordinating anions as counter ions. In some cases, cationic clusters with unusual bonding situations including *trans*-annular interactions and negative hyperconjugation were found that presented quite a challenge for theory (*e.g.* the

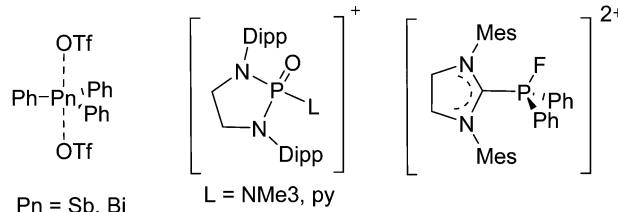
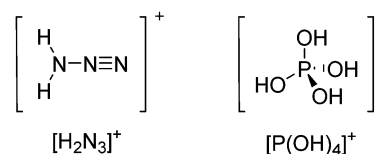


Fig. 80 Examples for recent pnictogen cations in oxidation state +V.

Fig. 81 Structures of $[\text{H}_2\text{N}_3]^+$ and $[\text{P}(\text{OH})_4]^+$ in their $[\text{SbF}_6]^-$ salts.

$[\text{S}_8]^{2+}$ dication).^{11,503,511} The cations were mostly synthesized under superacidic conditions or through solid state or solvothermal reactions at higher temperature. Either the elemental chalcogen is directly oxidized with strong oxidants like MF_5 ($\text{M} = \text{As}, \text{Sb}$) or WCl_6 or a combination of the elemental chalcogen, chalcogen halides ChX_4 (*e.g.* SeCl_4 , TeBr_4) and a strong Lewis acid undergo a synproportionation.

Metal-chalcogen complexes. The use of very weakly coordinated metal salts $\text{M}[\text{WCA}]$ (*e.g.* $\text{M}^+ = \text{Cu}^+, \text{Ag}^+$) made it possible to obtain complexes with very weak ligands like the elemental modifications of the chalcogen elements. Stable, metastable and hitherto unknown modifications were prepared, for example $[\text{Cu}(\text{S}_{12})(\text{S}_8)]^+$,⁵¹² $[\text{Cu}_2\text{Se}_{19}]^{2+}$ (Fig. 83)⁵⁰ and $[\text{Ag}_2\text{Se}_6]^{2+}$ (Fig. 84).⁵¹³ In all such complexes, extensive charge delocalization from the metal cation to the chalcogen ring took place as evidenced by cation-anion contacts as well as accompanying quantum chemical calculations.

Clusters/cluster-like cations. Chalcogens have a strong tendency to form clusters. There are examples for chains, rings and cages with almost every combination of the groups 15 and 16 (Fig. 68 and 85). The clusters often have delocalized charges, positive and

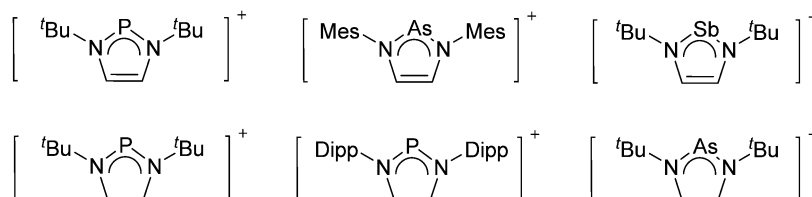


Fig. 78 Examples for pnictogen carbenoids.



Table 6 Review articles including cationic group 16 compounds

Year	Title	Ref.
2000	Recent advances in the understanding of the syntheses, structures, bonding and energetics of the homopolyatomic cations of groups 16 and 17	503
2003	Homoatomic sulfur cations	504
2004	Cages and clusters of the chalcogens	505
2006	Synthesis, reactions and structures of telluronium salts	506
2011	Homo- and heteroatomic polycations of groups 15 and 16. Recent advances in synthesis and isolation using room temperature ionic liquids	66
2013	Catenated sulfur compounds	507
2013	Catenated compounds – group 16 (Se, Te)	508
2013	Recent advances in the syntheses of homopolyatomic cations of the non-metallic elements C, N, P, S, Cl, Br, I and Xe	11
2013	RCN ^{SSS} ⁺ : a novel class of stable sulfur rich radical cations	509
2015	Coordination chemistry of homoatomic ligands of bismuth, selenium and tellurium	398 and 399

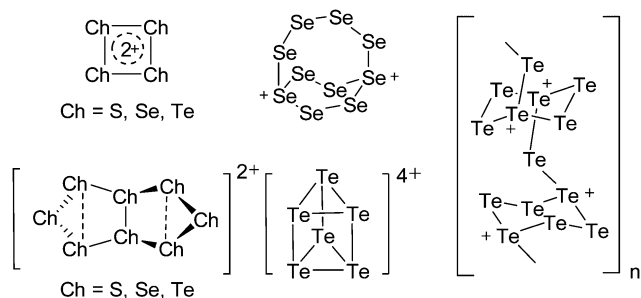


Fig. 82 Selected examples for homopolyatomic chalcogen cations.

negative hyperconjugation, or show pseudo-aromaticity (Fig. 86) or π^* - π^* -interactions (Fig. 87). The clusters were often synthesized through direct oxidation of neutral clusters like S_4N_4 or mixtures of the elements (e.g. Se and Te) with strong oxidants like MF_5 in SO_2 . $[NS]^+$, a useful starting material for the syntheses of further rPBC, can be obtained by halide abstraction from trichlorocyclotrithiazene ($NSCl$)₃ with $[Ag]^+[WCA]^-$.⁵¹⁴

(4n + 2) π -cations. Some planar cationic conjugated π -systems containing chalcogen atoms can be described as (pseudo-)aromatic systems. The four-membered rings are related to the homopolyatomic cations Ch_4^{2+} and were synthesized through direct oxidation of mixtures of the elemental chalcogens instead of

the pure elements. The five-membered rings were obtained by cycloadditions of $[NS]^+$ and $[NS_2]^+$ ⁵¹⁵ or in case of the selenium containing rings with Se, $[Se_8]^{2+}$ or EtSeCl and $[NS]^+$ as starting materials.^{516–518} $[S_4N_3]^+$, a 10 π -system, was synthesized through the reaction of S_4N_4 with Se_2Cl_2 and is stabilized by the polymeric $[SeCl_5]^-$ anion.⁵¹⁹

π^* - π^* -complexes. The chalcogen cations containing π^* - π^* -interactions can be described as dimers of chalcogen radicals, whose half-occupied interacting orbitals have π^* -character. The two $[Ch_2I_4]^{2+}$ cations (Ch = S, Se) have a very similar structure, but include different orbital interactions. In case of $[S_2I_4]^{2+}$ two 2e4c-bonds were formed through the π^* of the diatomic molecules $([I_2 \cdots S_2 \cdots I_2]^{2+})$.⁵²⁰ In $[Se_2I_4]^{2+}$ the two delocalized π^* orbitals of the “monomer” $[SeI_2]^+$ are overlapping.⁵²¹ In the last 10 years, chalcogen systems, which are analogous to the $[I_4]^{2+}$ cation were characterized by scXRD. The isolobality of $[Ch_2R_2]^+\bullet$ and $[X_2]^+\bullet$ leads to the same rectangular structural motif with two long π^* - π^* - and two short σ -interactions. Overall those cations are typically diamagnetic in the solid state (Fig. 88).

Radical cations. The chalcogen elements have a rich radical cation chemistry. Most of the cations contain (pseudo-)aromatic systems or Ch–Ch fragments, over which the unpaired electron is delocalized. Strong one-electron oxidants like $[NO]^+$ or XeF_2

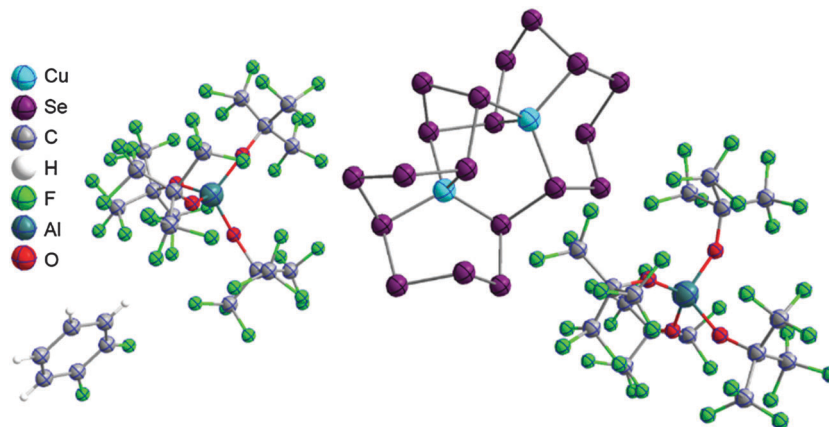
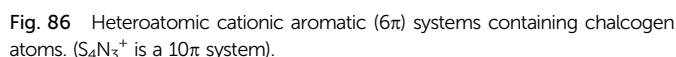
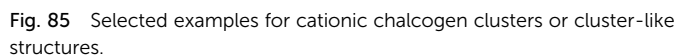
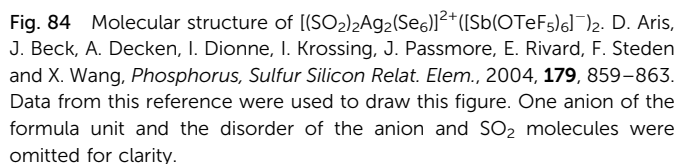


Fig. 83 Molecular structure of $[Cu_2Se_{19}]^{2+}([Al(OC(CF_3)_3)_4]^-)_2 \cdot C_6F_4H_2$. (a) J. Schaefer, A. Steffani, D. A. Plattner and I. Krossing, *Angew. Chem., Int. Ed.*, 2012, **51**, 6009–6012; *Angew. Chem.*, 2012, **124**, 6112–6115. Data from this reference were used to draw this figure.



combined with a Lewis acid were frequently used to synthesize the cations. There are also examples of diradicals like $(\text{CNS}_3\bullet^+)_2$, obtained by the reaction of homopolyatomic sulfur cations (a formal $[\text{S}_3]^+$ equivalent) and dicyanogen.^{522,523} We also refer the reader to a recently published comprehensive review about $[\text{RCNSSS}]^{\bullet+}$ radical cations and their properties.⁵⁰⁹



Oxidation state +II. Chalcogen cations $[\text{RCh}]^+$ in the formal oxidation state +II are only known in combination with stabilizing donor ligands. It was for example possible to stabilize the formal selenium cation $[\text{RSe}]^+$ with the two amine-arms of a pincer ligand⁵²⁴ (Fig. 89). Two different Te(II) cations with the strong donors DMAP and the carbene ⁱPrIm as ligands (Fig. 89) were synthesized with the useful starting material $[(\text{Dipp}_2\text{BIAN})\text{Te}]^{2+}[(\text{OTf})^-]_2$, a base stabilized “Te(OTf)₂”, which was presented in 2009 and can also be understood as a tellurium analogue of a carbene.⁵²⁵ The thi-, selen- and tellur-iridium cations (in the figure drawn as coordination complexes of $[\text{RCh}]^+$ ions) were all synthesized with starting materials that contain Ch–Ch bonds like Me_2S_2 ⁸² or $[\text{Se}_3\text{Me}_3]^+$ ⁵²⁶ or already oxidized chalcogens like $[\text{PhTe}]^+[\text{SbF}_6]^-$ ⁵²⁶ and alkynes.

Another example of ligand-stabilized chalcogen cations are the N-heterocyclic carbenoidic rings $[\text{R}_2\text{C}_2\text{N}_2\text{Ch}]^{2+}$ (Ch = S, Se, Te), which are formally 1,4-diaza-1,3-butadiene complexes of a chalcogen cation in oxidation state +II, but the delocalization of the positive charge supports a description as a chalcogen in oxidation state +IV (Fig. 90).⁵²⁷ The carbene-analogues were prepared through complexation of an *in situ* generated Ch^{2+} dication, which can be obtained through halide abstraction from SCL_2 ,⁵²⁸ SeCl_4 ⁵²⁹ or $(\text{Dipp}_2\text{BIAN})\text{TeI}_2$ ⁵²⁵ (Dipp_2BIAN = 1,4-(2,6-diisopropyl)phenylbis(arylimino)-acenaphthene).

Oxidation state +IV. The $[\text{ChX}_3]^+$ cations were one of first structurally characterized reactive chalcogen cations (Fig. 91). The first experiments mainly on $[\text{TeCl}_3]^+$ were published already in the 1950s.⁵³⁰ Now, a multitude of crystal structures can be found in the literature. Due to the relatively simple vibrational spectra of the four-atomic molecules, they can serve as a probe for the coordination power of the anion. The stronger the secondary interaction is, the weaker are the intramolecular Ch–X bonds and thus they get red-shifted.⁵³¹ There are also over 60 crystal structures of triorganyltelluronium $[\text{TeR}_3]^+$ cations with different anions. We decided only to list the “classics” $[\text{TePh}_3]^+$ ⁵³² and $[\text{Te}(\text{C}_6\text{F}_5)_3]^+$ ⁵³³ and to refer to a recently published review about the chemistry and structures of these cations.⁵⁰⁶ Some newer examples of compounds with chalcogen atoms in oxidation state +IV are the triazidetelluronium cation $[\text{Te}(\text{N}_3)_3]^+$ ⁵³⁴ and a fluoride bridged version of $[\text{TeCl}_3]^+$.²⁸

Oxidation state +VI. To our knowledge, the only chalcogen cation with a chalcogen atom in oxidation state +VI is $[\text{TePh}_5]^+$.

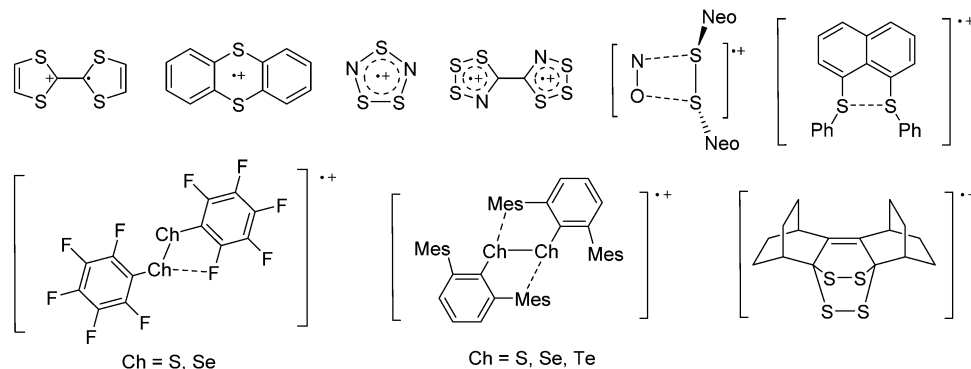


Fig. 88 Selected chalcogen radical cations, which were stabilized by WCAs.

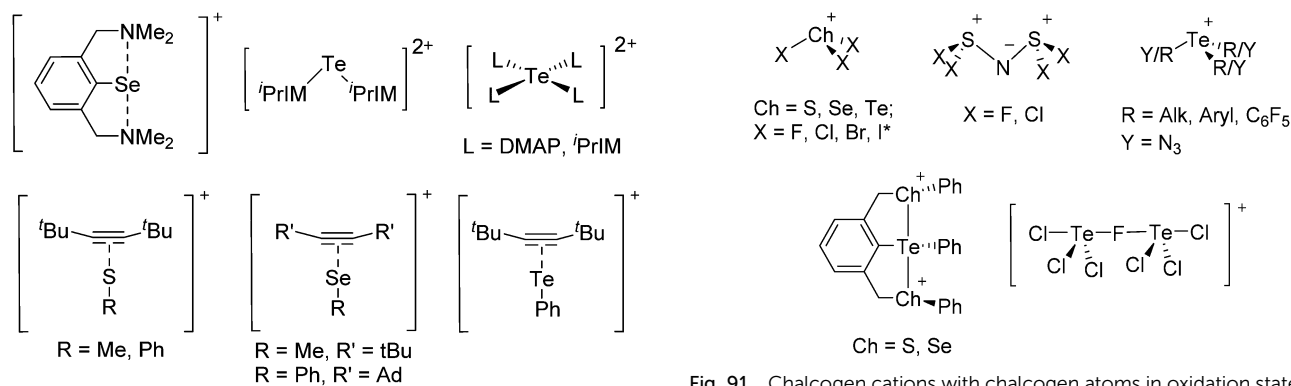


Fig. 89 Selected chalcogen cations with chalcogen atoms in the formal oxidation state of +II.

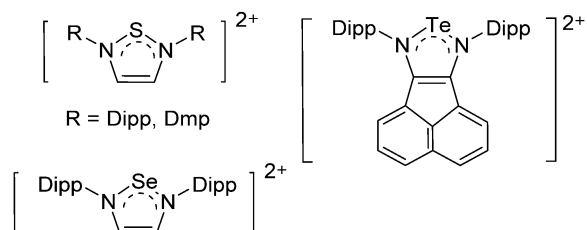


Fig. 90 Carbene-analogous chalcogen dications.

It was obtained through halide abstraction from TePh_5Cl with silver triflate and crystallized with the classical WCAs $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ ⁵³² and $[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$ (Fig. 92).⁵³⁵

Protonated chalcogen cations. The use of super-acidic conditions, makes it possible to obtain the protonated forms of very weakly basic molecules like trifluorosulfonic acid (Fig. 93).⁵³⁶ In some cases, it is not possible to isolate the neutral form of a molecule, but the conjugated positively charged acid (*e.g.* carbonic acid⁵³⁷). We decided to list the protonated carbonic acid together with the sulphur species like protonated sulphuric acid in this chapter (Table 11).⁵³⁸

Group 17 cations

The electronegative halogen atoms are strongly oxidizing and have high ionization energies as well as electron affinities. This makes halogen cations good electrophiles, which need very

Fig. 91 Chalcogen cations with chalcogen atoms in oxidation state +IV. * Every combination but $[\text{Si}_3]^{+}$ is known.



Fig. 92 $[\text{TePh}_5]^+$, a cation with a chalcogen atom in oxidation state +VI.

oxidation-resistant WCAs to be stabilized in condensed phases. Concerning the synthesis of these highly reactive compounds, there are many similarities to the noble gas cations. Almost every halogen cation, for which a crystal structure is known, was synthesized through a halide abstraction by a strong Lewis acid like MF_5 ($\text{M} = \text{As}, \text{Sb}$). The starting materials are normally neutral inter-halogen compounds like ClF_3 , BrF_5 , IF_7 , I_2Cl_6 , or IBr , and the majority of the obtained cations contains halogen atoms in oxidation state +III, +V and +VII (Table 7).

Fluorine cations

Because of the high electronegativity and ionization potential of fluorine, it would be very difficult to oxidize it and obtain



Fig. 93 Examples for protonated molecules obtained under superacidic conditions.

Table 7 Review articles including cationic group 17 compounds

Year	Title	Ref.
2000	Recent advances in the understanding of the syntheses, structures, bonding and energetics of the homopolyatomic cations of groups 16 and 17	503
2008	Polyvalent perfluoroorgano- and selected polyfluoroorgano-halogen(III and V) compounds	640
2013	Recent advances in the syntheses of homopolyatomic cations of the non-metallic elements C, N, P, S, Cl, Br, I and Xe	11

actual fluorine cations in the condensed phase. By contrast, in the gas phase this is possible.⁶⁴¹ Some bulk cationic compounds contain fluorine, but it is very unlikely that the positive charge is actually localized on the fluorine atom. Such compounds like for example the in Fig. 94 shown compound that was published as a formal disilylfluoronium ion are discussed in the chapters of the element, which has a larger positive charge density (here silicon).⁶⁴²

It should be mentioned that also electrophilic “F⁺” or “N–F⁺” reagents like “Selectfluor” belong to this class of compounds that are very useful for organic transformations and compatible with solvents like CH₂Cl₂.⁶⁴³

Chlorine, bromine and iodine cations

Homopolyatomic cations. Two homopolyatomic chlorine cations are known in the solid state and its crystal structures of [Cl₃]⁺ and [Cl₄]⁺ were published in 1999⁴² and 2000.⁶⁴⁴ [Cl₂]⁺, which would be the lighter homologue to the known [Br₂]⁺ and [I₂]⁺, was only detected in the gas phase. [Cl₃]⁺ can be synthesized through a reaction of ClF with AsF₅, with the adduct $\delta^+ \text{Cl}-\text{F}^{\delta-} \rightarrow \text{AsF}_5$ as intermediate. This leads to a more activated ClF with a more positively charged chlorine atom and results in the formation of [Cl₂F]⁺[AsF₆][−], which can be described as a formal “Cl⁺” stabilized by a second equivalent of ClF. When elemental chlorine is used instead of a second equivalent of ClF, [Cl₃]⁺ is formed. [Cl₄]⁺, which is a homopolyatomic cation but also a $\pi^*-\pi^*$ -complex (see below: $\pi^*-\pi^*$ -complexes of group 17) can be obtained by direct oxidation of chlorine with the strong one-electron oxidant IrF₆. For bromine, three cations are known. Of those, [Br₂]⁺ was one of the first homopolyatomic cations of the non-metals for which the crystal structure was determined. It was already in 1968 that Edwards *et al.* stabilized it with the very good [Sb₃F₁₆][−] WCA.⁶⁶³ The structures of the other two known cations [Br₃]⁺ and [Br₅]⁺ were measured relatively late in 1991. [Br₅]⁺ can be synthesized by oxidation of elemental bromine with the strong oxidant [XeF]⁺²¹¹ and the only measurable crystals of a [Br₃]⁺ salt were obtained from a 20 year old [BrF₂]⁺[AsF₆][−] solution.⁶⁴⁵ Iodine has five known cations. [I₂]⁺, [I₃]⁺ and [I₅]⁺ are isostructural to the

lighter homologues and can all be obtained by oxidation from I₂ with the strong Lewis acids MF₅ (M = As, Sb). [I₄]²⁺ was synthesized through the entropically unfavorable dimerization of the paramagnetic [I₂]^{•+} radical cation at low temperature and can be described as an rectangular planar diamagnetic $\pi^*-\pi^*$ -complex (see below: $\pi^*-\pi^*$ -complexes of group 17) (Fig. 95).

Metal-halogen complexes. The very weak coordination power of the neutral dihalogen molecules X₂ (F₂, Cl₂, Br₂, I₂) made it very difficult to obtain metal-halogen-complexes. Only for diiodine, which is the strongest donor, it was possible to get polymeric [Ag(I₂)]_n⁺ cations through the reaction of [Ag]⁺[MF₆][−] (M = As, Sb) and I₂ in liquid SO₂.⁶⁴⁶ Also one example of a neutral complex with diiodine as ligand is known: [Rh₂(O₂CCF₃)₄(I₂)]·I₂.⁶⁴⁷ Dibromine and dichlorine complexes remained unknown. However, very recently,⁶⁴⁸ the use of the very weakly coordinating solvent perfluorohexane and one of the weakest anions [Al(OR^{PF})₄][−] led to the isolation of the first dichlorine and dibromine complexes [Ag(X₂)]⁺ (X = Cl, Br, also I) (Fig. 96). Moreover, diiodine turned out to have a rich coordination chemistry and formed three further structures with Ag₂I₂-moiety as well as isolated [Ag₂(I₂)₄]²⁺ as well as [Ag₂(I₂)₆]²⁺ dications well separated from the counterion (Fig. 96). It should be noted that the [Ag(X₂)]⁺ cations are structurally related to the [X₃]⁺ cations and the polymeric [Ag(I₂)]_n⁺ cations bear some similarity to [I₅]⁺.

$\pi^*-\pi^*$ -complexes. The three halogen containing cations that can be described as $\pi^*-\pi^*$ complexes, can be understood as adducts of two homonuclear diatomic (radical) cations. According to its synthesis from chlorine and [O₂]⁺[SbF₆][−], [Cl₂O₂]^{•+} can be described as complex of the paramagnetic [O₂]^{•+} cation with Cl₂. For [Cl₄]^{•+}, which is made from Cl₂ and the strong oxidant IrF₆, an initially formed paramagnetic [Cl₂]^{•+} could react with another equivalent of Cl₂. In the case of [I₄]²⁺, which was obtained from an [I₂]^{•+} solution at low temperature, a dimerization of the two paramagnetic [I₂]^{•+} units to give the [I₄]²⁺ dimer is obvious. All three syntheses have in common that at least one molecule with a half-filled π^* orbital as HOMO is involved, for [I₄]²⁺ even both “starting materials”. The formed $\pi^*-\pi^*$ interaction includes two quarter bonds for [Cl₂O₂]^{•+} (243/241 pm, Σ vdW radii: 327 pm) and [Cl₄]^{•+} (293 pm, Σ vdW radii: 350 pm) and two half bonds for [I₄]²⁺, where the interaction leads to 2e-4c $\pi^*-\pi^*$ bond (328 pm, Σ vdW radii: 396 pm, Fig. 97).^{11,503}

Oxidation state +I. The isolated halogens X in this oxidation state would either correspond to a triplet state X⁺ or the triplet dimer [X = X]²⁺ – isoelectronic to the dichalcogens O₂ to Te₂. Neither the monomer, nor the dimer cation are hitherto known in condensed phase. Only one cation type with a formal oxidation state of +I is known: [I(Donor)₂]⁺, in which the electrophilicity of I⁺

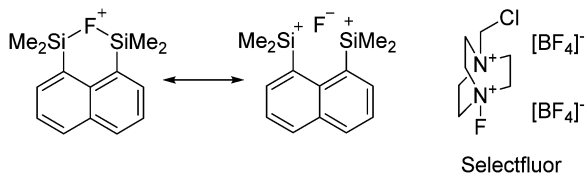


Fig. 94 Left: Formal disilylfluoronium ion that bears a larger positive charge density at the silicon atoms. Therefore, it is discussed with the silylium cations in the section on group 14 rPBC. Right: The Selectfluor reagent.



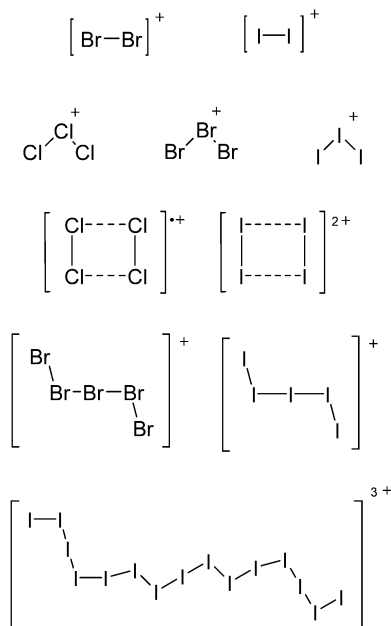


Fig. 95 Structurally characterized homopolyatomic cation of chlorine, bromine and iodine.

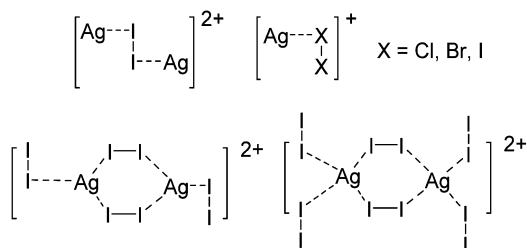


Fig. 96 Structurally characterized metal complexes with dihalogen molecules as ligands.

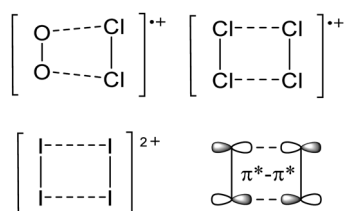


Fig. 97 Structurally characterized halogen cations, which include $\pi^*-\pi^*$ -interactions. Bond lengths: $[\text{Cl}_2\text{O}_2]^+$: O–O 121, Cl...O 241/243, Cl–Cl 191; $[\text{Cl}_4]^+$: Cl–Cl 194, Cl...Cl 293; $[\text{I}_4]^{2+}$: I–I 258, I...I 328 [pm].

is dampened by coordination of neutral donor molecules. Thus, $[\text{I}_3]^+[\text{AsF}_6]^-$ reacted with acetonitrile to form the structurally characterized salt $[\text{I}(\text{NCMe})_2]^+[\text{AsF}_6]^-$ (Fig. 98).⁶⁴⁹ The closely related $[\text{I}(\text{py})_2]^+[\text{BF}_4]^-$ is a useful reagent in organic chemistry.⁶⁵⁰ For the latter it is disputable, if this is better assigned as being a pyridinium cation.

Oxidation state +III. The known interhalonium cations with the central halogen atoms in oxidation state +III are $[\text{ClF}_2]^+$, $[\text{BrF}_2]^+$, $[\text{ICl}_2]^+$, $[\text{IBrCl}]^+$ and $[\text{IBr}_2]^+$ (Fig. 99). All were synthesized

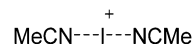


Fig. 98 $[\text{I}(\text{NCMe})_2]^+$ cation in $[\text{I}(\text{NCMe})_2][\text{AsF}_6]$ with iodine in the formal oxidation state +I.⁶⁴⁹

by halide abstraction from a neutral interhalogen. Others, like $[\text{Cl}_2\text{F}]^+$ and $[\text{IF}_2]^+$ are also accessible, but no crystal structures are known.⁶⁵¹ The cations $[\text{I}_3\text{Cl}_2]^+$ and $[\text{I}_3\text{Br}_2]^+$ can also be understood as interhalonium compounds with two iodine atoms in oxidation state +III and one in –I and are in some way the lighter homologues of the homopolyatomic cation I_5^+ .^{652–654} Two recently published examples are the dialkyl chloronium cations $[\text{ClR}_2]^+$ (R = Me, Et), which were synthesized through alkylation of chloromethane and –ethane, both stabilized by the very good carborate WCA $[\text{CHB}_{11}\text{Cl}_{11}]^-$ (Fig. 100). The superacid $\text{H}(\text{CHB}_{11}\text{Cl}_{11})$ was used to generate ion-like methyl and ethyl cations “R⁺” by protonation of the chloroalkanes RCl , which than react with a second equivalent of RCl to the chloronium cations.⁶⁵⁵

In case of bromine and iodine, also some new examples were synthesized during the last 20 years. The stable cyclic bromonium and iodonium ions of sterically hindered olefins (Fig. 101) were stabilized by the $[\text{OTf}]^-$ anion and can be seen as stable intermediates of the halogenation of olefins.⁶⁵⁶

Oxidation state +V. There are also some examples of cations in oxidation state +V. Besides the classical $[\text{XF}_4]^+$ cations (X = Cl, Br, I), the two oxocations $[\text{ClO}_2]^+$ and $[\text{BrO}_2]^+$ as well as since 2008 also one example of a cation with a $\text{X}^{\text{V}}\text{-carbon}$ bond were structurally characterized. This electrophilic $[\text{IF}_2(\text{C}_6\text{F}_5)_2]^+$ cation was synthesized as $[\text{BF}_4]^-$ salt through halide abstraction and ligand exchange from $\text{C}_6\text{F}_5\text{IF}_4$ and $\text{C}_6\text{F}_5\text{BF}_2$.⁶⁵⁷ Examples without crystal structures include $[\text{OCIF}_2]^+$ ⁶⁵⁸ and $[\text{OBrF}_2]^+$ (Fig. 102).⁶⁵⁹

Oxidation state +VII. In 2004, with the octahedral complexes of the series $[\text{XF}_6]^+$ (X = Cl, Br, I) the first structures of cations with oxidation state +VII were published (Fig. 103). All three cations were crystallized with the anion $[\text{Sb}_2\text{F}_{11}]^-$ and have very weak contact to the anion (Fig. 104). It is also possible to obtain the $[\text{AsF}_6]^-$ and $[\text{SbF}_6]^-$ salts, but the differentiation between the octahedral $[\text{XF}_6]^+$ and $[\text{MF}_6]^+$ would not be easily done by X-ray crystallography (e.g., for $[\text{BrF}_6]^+[\text{AsF}_6]^-$ or $[\text{IF}_6]^+[\text{SbF}_6]^-$). The use of $[\text{Sb}_2\text{F}_{11}]^-$ allowed for a clear differentiation between cations and anions (Table 12).⁶⁶⁰

Group 18 cations

Due to the high ionization potential of the noble gases, their cations all need weakly coordinating and very oxidation resistant

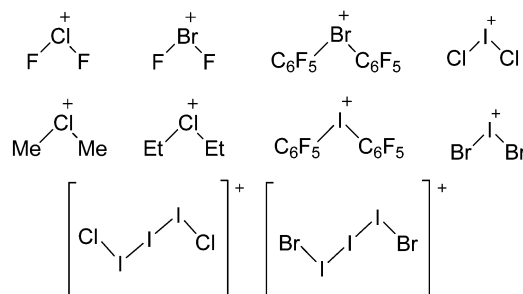


Fig. 99 Structurally characterized halonium cations.

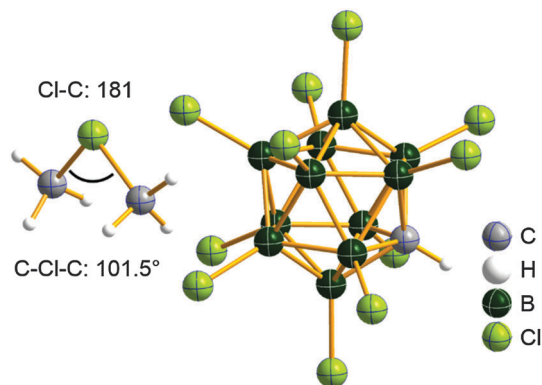


Fig. 100 Molecular structure of $[\text{ClMe}_2]^+[\text{CHB}_{11}\text{Cl}_{11}]^-$. E. S. Stoyanov, I. V. Stoyanova, F. S. Tham and C. A. Reed, *J. Am. Chem. Soc.*, 2010, **132**, 4062–4063. Data from this reference were used to draw this figure. Bond lengths in [pm].



Fig. 101 Stable $[\text{OTf}]^-$ salts of cyclic bromonium and iodonium ions.⁶⁵⁶

anions to be stabilized. The history of noble gas compounds started in the early 1960s, when Bartlett obtained $[\text{Xe}]^+[\text{PtF}_6]^-$ from a reaction of $[\text{O}_2]^+[\text{PtF}_6]^-$ and xenon – the noble gas with the lowest ionization potential. Over the next decades the composition of this product stayed unclear. In 2000 a very comprehensive review about the nature of the product was published with the result that it is very likely a $[\text{XeF}]^+$ salt of a polymeric (weakly coordinating) $[\text{PtF}_5]_n^-$ anion.⁶⁸¹ All of the hitherto known and structurally characterized noble gas cations are included with Table 13.

Krypton cations

The only other noble gas besides xenon, for which cations are known, is krypton. The oxidation state +II is the only one known yet. The diversity of the amount of compounds and the number of hitherto realized Kr–X bonds is much smaller. Only two different fluorine-containing cations are known ($[\text{KrF}]^+$ and $[\text{Kr}_2\text{F}_3]^+$) and some nitrile complexes, which can be described as Kr–N compounds. $[\text{KrF}]^+$ is a very strong oxidative fluorinating reagent⁶⁸² and made it possible to synthesize $[\text{XeF}_5]^+$ from Xe,⁶⁸³ $[\text{ClF}_6]^+$ from ClF_5 ,⁶⁸⁴ $[\text{BrF}_6]^+$ from BrF_5 ⁶⁸⁵ and $[\text{O}_2]^+$ from O_2 .⁶⁸⁶ There are only seven crystal



Fig. 102 Structurally characterized halogen cations with halogen atoms in oxidation state +V.^{657,660–662}

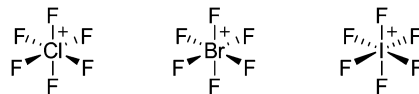


Fig. 103 Structurally characterized halogen cations with halogen atoms in oxidation state +VII.⁶⁶⁰



Fig. 104 Molecular structure of $[\text{BrF}_6]^+[\text{Sb}_2\text{F}_{11}]^-$. J. F. Lehmann, G. J. Schrobilgen, K. O. Christe, A. Kornath and R. J. Suontamo, *Inorg. Chem.*, 2004, **43**, 6905–6921. Data from this reference were used to draw this figure. Bond lengths in [pm].

structures of krypton cations in the literature and five of them with the classical $[\text{AsF}_6]^-$ and $[\text{SbF}_6]^-$ anions.

Xenon cations

The lower ionization potential calls for a richer chemistry of xenon. Thus, since the birth of noble gas chemistry, a multitude of different xenon cations containing the element in the oxidation state II, IV and VI were synthesized.

Homopolyatomic cations. The two known homopolyatomic cations are $[\text{Xe}_2]^+$ and $[\text{Xe}_4]^+$. It was only possible to obtain a crystal structure from $[\text{Xe}_2]^+$, but because of its importance $[\text{Xe}_4]^+$, which was assigned based on spectroscopic and computational evidence as being stable at higher Xe pressure, is also mentioned.

Fluoroxenon cations and related. The majority of such cations include Xe–F bonds ($[\text{XeF}_m]^+$, $m = 1, 3, 5$; $[\text{Xe}_2\text{F}_n]^+$, $n = 3, 11$, Fig. 105), but over the decades a lot of different cations with Xe–X (X = F, Cl, O, N, C) bonds were obtained and are included with Table 13 and in part also with Fig. 106. Some of them were already published in the late sixties,^{687–689} but also recently with $[\text{XeF}]^+[\text{SbF}_6]^-$ and $[\text{XeF}_3]^+[\text{SbF}_6]^-$ some new structures were presented^{690,691} and in 2015 the crystal structures of $[\text{XeF}_5]^+$ with the classical WCAs $[\text{SbF}_6]^-$ and $[\text{Sb}_2\text{F}_{11}]^-$ were measured for the first time,⁶⁹² which shows that the investigation of xenon cations is still in progress.

Most of the syntheses use Lewis acids to abstract fluoride from the neutral xenon fluoride (XeF_2 , XeF_4 , XeF_6). Most structures contain the conjugated weakly coordinating anions of these Lewis acids ($[\text{MF}_6]^-$, M = As, Sb, Au, Ru). Another mentionable approach is the reaction of XeF_2 with the Lewis acid $\text{B}(\text{C}_6\text{F}_5)_3$, which led to the $[\text{F}_5\text{C}_6\text{Xe}]^+$ cation containing the first Xe–C bond and the unsymmetric anion $[(\text{F}_5\text{C}_6)_2\text{BF}_2]^-$.⁶⁹³ Through the use of the dioxidifluoride XeO_2F_2 as starting



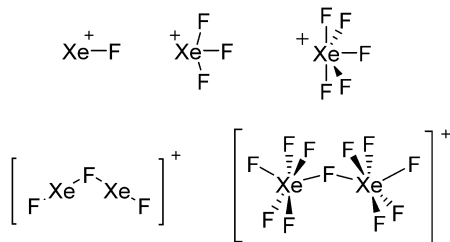


Fig. 105 Fluoroxenon cations.

material, it was also possible to obtain the mixed cation $[\text{XeO}_2\text{F}]^+$ or the fluoride bridged $[\text{F}(\text{XeO}_2\text{F})_2]^+$ (Fig. 107).⁶⁹⁴

Metal-xenon complexes. There are also some examples of metal-xenon complexes ($\text{M} = \text{Au}^{\text{I}}, \text{Au}^{\text{II}}, \text{Au}^{\text{III}}$ and Hg^{II}), which can be described as part of the stabilization of modifications of main group elements as metal complexes $[\text{M}_m(\text{E}_n)]^+$. Formally, Xe is isoelectronic to iodide I^- , and thus complex formation appeared to be difficult, but feasible. The syntheses proceed under the superacidic conditions of the systems HF/MF_5 ($\text{M} = \text{As}, \text{Sb}$). In case of these compounds, the role of the anions can be described as unreactive, but not really weakly coordinating. Most of them have relatively short contacts to the anions (shorter than the sum of the van der Waals-radii) and show typical coordination spheres with the anions included (e.g. square planar $\text{trans-Au}^{\text{II}}\text{Xe}_2(\text{SbF}_6)_2$) (Fig. 108).⁶⁹⁵

Only the $[\text{AuXe}_4]^{2+}$ dication exists in a truly ionic structure with two $[\text{Sb}_2\text{F}_{11}]^-$ counterions in the lattice (Fig. 109).

Conclusion

Sparked by the availability of new WCAs and new WCA starting materials in combination with novel concepts like FLP and others, the number of rPBC exploded over the last one to two decades. Noteworthy additions were found for each p-block element and, despite their quite high moisture and air sensitivity, true applications of rPBC salts emerged.

Where will this lead to over the next one or two decades...? To our understanding the blue sky synthesis of rPBC salts barely accessible with good/novel WCAs in combination with suitable media will continue to function as an "eye-opener" of what is possible. Many surprising discoveries will force us to sharpen our use of bonding concepts or lead to novel applications. It is often the

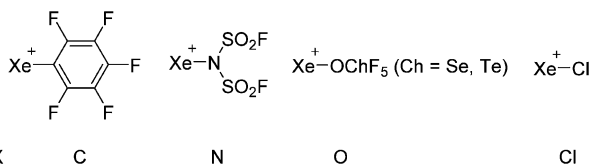
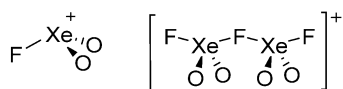
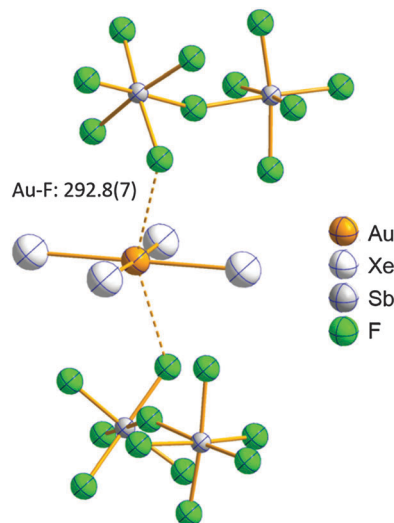
Fig. 106 Examples of Xe-X bonds ($\text{X} = \text{C}, \text{Cl}, \text{N}, \text{O}$) in cations.

Fig. 107 Examples of Xe-O-F cations.

Fig. 108 Molecular structure of $\text{trans-Au}^{\text{II}}\text{Xe}_2(\text{SbF}_6)_2$. T. Drews, S. Seidel and K. Seppelt, *Angew. Chem.*, 2002, **114**, 470–473; *Angew. Chem., Int. Ed.*, 2002, **41**, 454–456. Data from this reference were used to draw this figure.Fig. 109 Molecular structure of tetragonal $[\text{Au}^{\text{II}}\text{Xe}_4]^{2+}([\text{Sb}_2\text{F}_{11}]^-)_2$. T. Drews, S. Seidel and K. Seppelt, *Angew. Chem.*, 2002, **114**, 470–473; *Angew. Chem., Int. Ed.*, 2002, **41**, 454–456. Data from this reference were used to draw this figure.

combination of structural knowledge ("Wow, this crazy cation is stable...? I would have never thought so.") that leads to the right moment of wonder and then inspiration ("Hm, if this cation is really straight forward accessible, one could use its electrophilic/acidic/oxidizing/activating properties in application XY"). In the 21st century, it is our duty as creative scientists to use this potential from fundamentals to the first application. Do not hesitate to really seek for application of your rPBC salt, as rarely others will pick up on these ideas, since the activation barrier for synthesizing a to this application group unknown rPBC is simply too high. So do not give up until you have demonstrated a possible application – yourself or through collaborations – to a level that others will continue. And on the other hand this compilation of rPBC should encourage application based groups to identify interesting cations that may have an application. Contact the people, the chances are very good that through an informal collaboration showing a proof-of-principle new and relevant application areas may be developed.

In this respect, we are looking forward to all the scientific creativity that is breaking loose, and to realize what potentially could be done with the rPBC. This is an integral element of innovation and the justification for preparing blue sky or simply beautiful and esoteric compounds.





Table 8 Group 13 cations, their counterions (WCA) as well as the synthesis routes. The entries are ordered as follows: (i) from boron- to thallium-based cations, (ii) from unsubstituted, via alkyl/aryl, Cp, arene and ligand to transition-metal substituted cations, (iii) from low to high CNs, (iv) from mono- to multinuclear group 13 complexes. Note that the structural diagrams obey a distance criterion for interactions, but not necessarily a 2e2c-understanding of every interaction line. See comment in the Introduction

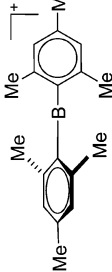
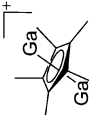
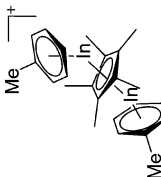
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
Unsubstituted					
In ⁺	[OTf] [−]	Prot	InCp ⁺ + H ⁺ [WCA] [−]	Soluble in organic solvents in contrast to the In(i) halides	182
Tl ⁺	[B(Ar ^{CF3}) ₄] [−] / [B(C ₆ F ₅) ₄] [−]	Prot	TlOEt + [H(OEt) ₂] ⁺ [WCA] [−]	—	196 and 197
Tl ⁺	[B(OTeF ₅) ₄] [−]	Lewis	Tl ⁺ [OTeF ₅] [−] + B(OTeF ₅) ₃ in CH ₂ Cl ₂ / 1,2-C ₂ Cl ₃ F ₃	—	198
Tl ⁺	[Al(OR ^{PF}) ₃] [−] / [Al(OR ^{HFP}) ₄] [−] / [Al(OR ^{MeF}) ₄] [−]	Salt	TlF + Li ⁺ [WCA] [−]	—	199 and 200
Alkyl/aryl substituted					
[Mes ₂ B] ⁺	[HCB ₁₁ Cl ₁₁] [−] / [B(C ₆ F ₅) ₄] [−]	Salt	Mes ₂ BF + Et ₃ Si(HCB ₁₁ Cl ₁₁) / [Et ₃ Si(Mes)] ⁺ [WCA] [−]		32 and 113
(R ₂ Al) (R = Me, Et) (Et ₂ Al)	[B ₁₂ Cl ₁₂] ^{2−} / [CB ₁₁ H ₆ X ₆] [−] (X = Cl, Br)	Alk Alk	R ₂ Al + {[CPh ₃] ⁺ }[B ₁₂ Cl ₁₂] ^{2−} / Et ₃ Al + [CPh ₃] ⁺ [CB ₁₁ H ₆ X ₆] [−]	Ion-like compound Ion-like compound	87 128
[(2,6-Mes ₂ C ₆ H ₃) ₂ Al] ⁺	[B(C ₆ F ₅) ₄] [−]	Hyd	(2,6-Mes ₂ C ₆ H ₃) ₂ AlH + [CPh ₃] ⁺ [WCA] [−]	Related structure to the [Mes ₂ B] ⁺ cation, though the Mes moieties of the 2,6-Mes ₂ C ₆ H ₃ substituent additionally shield the aluminum cation	129
[(2,6-Mes ₂ C ₆ H ₃) ₂ Ga] ⁺	[Li{Al(OR ^{HFP}) ₄ }] [−]	Salt	(2,6-Mes ₂ C ₆ H ₃) ₂ GaCl + 2Li ⁺ [WCA] [−]	Similar structure as the [(2,6-Mes ₂ C ₆ H ₃) ₂ Al] ⁺ cation	158
Cyclopentadienyl complexed					
[(η ⁵ -Cp) ₂ Al] ⁺	[Al(OR ^{PF}) ₄] [−]	Prot	AlCp ₃ + [H(OEt) ₂] ⁺ [WCA] [−]	—	79
[(η ⁵ -Cp) ₂ Al] ⁺	[B(C ₆ F ₅) ₄] [−]	Alk	Cp ₃ Al + [CPh ₃] ⁺ [WCA] [−]	—	131
[(η ⁵ -Cp) ₂ Al] ⁺	[Ph(Me)B(η ⁵ -C ₅ H ₄) ₂]ZrCl ₂] [−]	Alk	Cp ⁺ ₂ AlMe + {Ph(SMe ₂)B(η ⁵ -C ₅ H ₄) ₂ }ZrCl ₂ + [Ph ₃ P=N=PPh ₃] ⁺ Cl [−]	—	132
[(η ⁵ -Cp) ₂ Al] ⁺	[MeB(C ₆ F ₅) ₃] [−]	Alk	Cp ⁺ ₂ AlMe + B(C ₆ F ₅) ₃	—	133
[(η ⁵ -Cp) ₂ (Et ₂ O) ₂ Al] ⁺	[Al(OR ^{PF}) ₄] [−]	Prot	AlCp ₃ + [H(OEt) ₂] ⁺ [WCA] [−]	Et ₂ O can coordinate the [(η ⁵ -Cp) ₂ Al] ⁺ cation	79
[Ga ₂ (η ⁵ -Cp ⁺) ⁺	[B(Ar ^{CF3}) ₄] [−]	Prot	[H(OEt) ₂] ⁺ [WCA] [−] + GaCp ⁺		159
[(η ¹ -Cp*)(η ³ -Cp*)Ga] ⁺	[BF ₄] [−]	Prot	Cp ⁺ ₃ Ga + HBF ₄	cf. [B(η ⁵ /η ¹ -Cp ⁺) ₂] ⁺ and [Al(η ⁵ /η ⁵ -Cp ⁺) ₂] ⁺	160
[In ₂ (η ⁵ -Cp)] ⁺	[Cp ₃ In-Cp-InCp ₃] [−]	Com	In ⁺ [OTf] [−] + Cp ₂ Mn in C ₆ H ₅ Me	Inverted sandwich structure (cf. the related [Ga ₂ (η ⁵ -Cp ⁺) ⁺ cation)	183
[In ₂ (η ⁵ -Cp)] ⁺	[B(C ₆ F ₅) ₄] [−]	Prot	[(C ₆ H ₅ Me)H] ⁺ [WCA] [−] + InCp ⁺	Similar structure to the [Ga ₂ (η ⁵ -Cp ⁺) ⁺ cation	185
[(μ-η ⁵ -C ₅ Me ₅)In ₂ (η ⁶ -Tol) ₂] ⁺	[(C ₆ F ₅) ₃ BO(H)B(C ₆ F ₅) ₃] [−]	Prot, Com	(Cp ⁺ In) ₆ + B(C ₆ F ₅) ₃ + H ₂ O·B(C ₆ F ₅) ₃		184 and 185

Table 8 (continued)

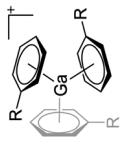
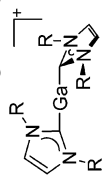
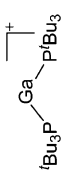
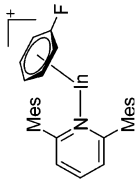
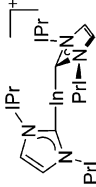
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
Arene complexed					
$[\text{Ga}(\eta^6\text{-C}_6\text{H}_5\text{R})_n]^+$ (R = F, Me; n = 2, 3)	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Ox	$\text{Ga}^0 + \text{Ag}^+[\text{WCA}]^-$ in arene		31, 91 and 92
$[\text{Ga}(\eta^6\text{-arene})_n]^+$ (n = 2, 3)	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Com	$[\text{Ga}(\text{C}_6\text{H}_5\text{F})_{2-3}]^+[\text{WCA}]^- + \text{arene}$ (arene = Mes, p-Xyl, C ₆ Me ₆)	Bent-sandwich (2 ligands) or tubby coordinated complex (3 ligands)	7
$[\text{Ga}(\eta^6\text{-DPE})]^+$	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Com	$[\text{Ga}(\text{C}_6\text{H}_5\text{F})_{2-3}]^+[\text{WCA}]^- + \text{DPE}$	First structurally characterized bent-sandwich ansa-arene complex	8 and 99
$[\text{In}(\eta^6\text{-C}_6\text{H}_5\text{F})_n]^+$ (n = 2, 3)	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Ox	$\text{In}^0 + \text{Ag}^+[\text{WCA}]^-$ in C ₆ H ₅ F	Bent-sandwich complex (cf. gallium analogue)	162
$[\text{In}(\eta^6\text{-o-C}_6\text{H}_4\text{F}_2)_2]^+$	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Salt	$\text{InCl} + \text{Li}^+[\text{WCA}]^-$ in o-C ₆ H ₄ F ₂	Bent-sandwich complex (cf. gallium analogue)	186
$[\text{Ti}(\eta^6\text{-C}_6\text{Me}_6)]^+$	$[\text{H}_2\text{N}\{\text{B}(\text{C}_6\text{F}_5)_3\}_2]^-$	Other	$[\text{Ti}(\text{C}_6\text{Me}_6)_2]^+$ in Et ₂ O + C ₆ H ₅ Me, vacuum	First example of a mono-η ⁶ -coordinated thallium complex	204
$[\text{Ti}(\eta^6\text{-C}_6\text{H}_5\text{Me})_2]^+$	$[\text{HCB}_3\text{H}_5\text{Br}_6]^-$	Salt	$\text{Cs}^+[\text{HCB}_3\text{H}_5\text{Br}_6]^- + \text{TlF}$	Bent-sandwich complex (cf. gallium analogue)	201
$[\text{Ti}(\eta^6\text{-C}_6\text{H}_5\text{Me})_3]^+$	$[\text{H}_2\text{N}\{\text{B}(\text{C}_6\text{F}_5)_3\}_2]^-$	Com	$[\text{Ti}(\text{OEt}_2)_2]^+[\text{WCA}]^- + \text{C}_6\text{H}_5\text{Me}$	Tubby coordinated complex (cf. gallium analogue)	203
$[\text{Ti}(\eta^6\text{-Mes})_2]^+$	$[\text{B}(\text{OTeF}_5)_4]^-$	Lewis, Com	$\text{Ti}^+[\text{OTeF}_5]^- + \text{B}(\text{OTeF}_5)_3$ in Mes	Tubby coordinated complex (cf. gallium analogue)	202
$[\text{Ti}(\eta^6\text{-C}_6\text{Me}_6)_2]^+$	$[\text{H}_2\text{N}\{\text{B}(\text{C}_6\text{F}_5)_3\}_2]^-$	Com	$[\text{Ti}(\text{OEt}_2)_3]^+[\text{WCA}]^- + \text{C}_6\text{Me}_6$	Tubby coordinated complex (cf. gallium analogue)	203
Ligand substituted (CN = 2)					
$[\text{Cp}^*\text{B}(\text{IMes})]^2+$	$[\text{AlCl}_4]^-$	Lewis	$\text{Cp}^*\text{BCl}_2(\text{IMes}) + 2\text{AlCl}_3$		210
$[(^t\text{Bu}_3\text{PN})_2\text{B}]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Hyd	$(^t\text{Bu}_3\text{PN})_2\text{BH} + [\text{Ph}_3\text{C}]^+[\text{WCA}]^-$	$^t\text{Bu}_3\text{P}=\text{N}-\text{B}-\text{N}=\text{P}^t\text{Bu}_3$	114
$[\text{Bu}_2\text{MeSi}-\text{Al}-\text{Si}^t\text{Bu}_2-\text{Si}^t\text{Bu}_2\text{Me}]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Alk	$\text{Al}(\text{SiMe}^t\text{Bu}_2)_3 + [\text{Et}_3\text{Si}]^+[\text{WCA}]^-$	Hyperconjugation with a neighboring Si-Si bond	134
$[\text{Ga}(\text{IR})_2]^+$ (R = Pr, Mes)	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Com	$[\text{Ga}(\text{C}_6\text{H}_5\text{F})_2]^+[\text{WCA}]^- + \text{IR}$		161
$[\text{Bu}_2\text{MeSi}-\text{Ga}-\text{Si}^t\text{Bu}_2-\text{Si}^t\text{Me}^t\text{Bu}_2]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Alk	$\text{Ga}(\text{SiMe}^t\text{Bu}_2)_3 + [\text{Et}_3\text{Si}(\text{C}_6\text{H}_5)]^+[\text{WCA}]^-$	Stabilized by hyperconjugation with a neighboring Si-Si bond	134
$[\text{Bu}_3\text{Si}-\text{Ga}-\text{Si}^t\text{Bu}_3]^+$	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Salt	$(^t\text{Bu}_3\text{Si})_2\text{GaCl} + \text{Ag}^+[\text{WCA}]^-$	Linear arrangement	163
$[\text{Ga}(\text{P}^t\text{Bu}_3)_2]^+$	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Com	$[\text{Ga}(\text{C}_6\text{H}_5\text{F})_2]^+[\text{WCA}]^- + \text{P}^t\text{Bu}_3$		162
$[\text{In}(\text{Mes}_2\text{py})(\eta^6\text{-C}_6\text{H}_5\text{F})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt, Com	$\text{InBr} + \text{Na}^+[\text{WCA}]^- + \text{Mes}_2\text{py}$		187
$[\text{In}(\text{IPr})_2]^+$	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Com	$[\text{In}(\text{C}_6\text{H}_5\text{F})_2]^+ + \text{IPr}^-$		161



Table 8 (continued)

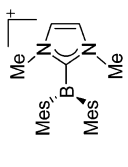
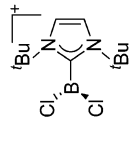
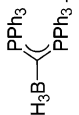
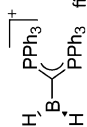
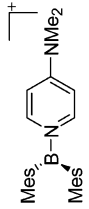
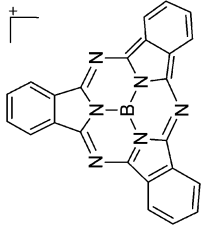
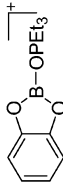
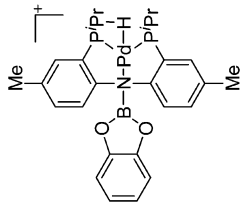
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
[In(Mes ₂ py) ₂] ⁺	[B(Ar ^{CtF3}) ₄] ⁻	Salt	In ⁺ Br ⁻ + Na ⁺ [WCA] ⁻ + 2Mes ₂ py		187
[Ti(1,2-Cl ₂ C ₂ H ₄) ⁺	[B(OTeF ₅) ₄] ⁻	Lewis, Com	Ti ⁺ [OTeF ₅] ⁻ + B(OTeF ₅) ₃ in 1,2-C ₂ H ₄ Cl ₂		198
Ligand substituted (CN = 3)					
[BMes ₂ (IME) ⁺	[OTf] ⁻	Salt	Mes ₂ BF + [Me ₃ Si] ⁺ [OTf] ⁻ + [Ag(IME) ₂] ⁺ [Ag ₂ I ₃] ⁻		211
[BCl ₂ (t ⁱ Bu)] ⁺	[B(Ar ^{Cl}) ₄] ⁻	Salt	BCl ₃ (t ⁱ Bu) + Na ⁺ [WCA] ⁻		74
[{(PPh ₃) ₂ C}BH ₂] ⁺	[HB(C ₆ F ₅) ₃] ⁻	Hyd	 + B(C ₆ F ₅) ₃	 first dihydroboreonium cation	118 and 119
[BMes ₂ (DMAP)] ⁺ /[B(Ar ^N) ₂ (DMAP)] ⁺ Ar ^N = 4-(Me ₂ N)-2,6-Me ₂ -C ₆ H ₃	[OTf] ⁻	Salt	Mes ₂ BF + Me ₃ Si-OTf + Ar ^N ₂ BF + DMAP		57
[B(SubPc)] ⁺ [(Sub = C ₂ H ₄ N ₂)	[HCB ₁₁ Me ₅ Br ₆] ⁻	Salt	B(SubPc)Cl + Et ₃ Si(HCB ₁₁ Me ₅ Br ₆)		115
[CatB(O=Pt ₃)] ⁺	[HCB ₁₁ H ₅ Br ₆] ⁻	Salt, Com	Ag ⁺ [WCA] ⁻ + CatBBr + OPEt ₃		116
[(CatB)(PNP)PdH] ⁺	[B(Ar ^{CtF3}) ₄] ⁻ /[CB ₁₁ H ₁₂] ⁻	Other	[(PNP)Pd(THF)] ⁺ [WCA] ⁻ + CatBH		117



Table 8 (continued)

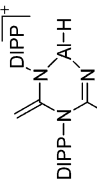
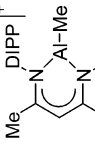
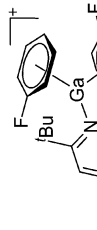
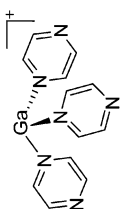
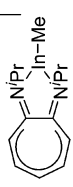
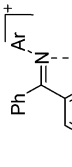
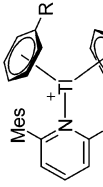
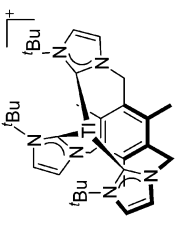
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\text{ArN}(\text{C}(\equiv\text{CH}_2)\text{NAr})(\text{C}(\text{Me})\text{NAr})\text{AlH}]^+$ (Ar = DIPP)	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Hyd	$\text{ArN}(\text{CMeNAr})_2 + \text{AlH}_3 \cdot \text{NMe}_2 \cdot \text{Et} + [\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		135
$\{[\text{HC}(\text{CMeNAr})_2]\text{AlMe}\}^+$ (Ar = DIPP)	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$ $[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$	Alk	$\{[\text{HC}(\text{CMeNAr})_2]\text{AlMe}_2 + [\text{CPh}_3]^+[\text{WCA}]^- / \text{B}(\text{C}_6\text{F}_5)_3$		136
$[\text{Ga}(\eta^6\text{-C}_6\text{H}_5\text{F}_2)(\text{DTBMP})]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{Ga}(\eta^6\text{-}\mu\text{-C}_6\text{H}_3\text{F}_3)_n]^+[\text{WCA}]^- + \text{DTBMP}$ ($n = 2, 3$)		164
$[\text{Ga}(\text{pyrazine})_3]^+ / \{[\text{Ga}(\mu\text{-pyrazine})_2]^\pm (\eta^1\text{-pyrazine})\}^\pm\}_\infty$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{Ga}(\text{C}_6\text{H}_5\text{F})]^+[\text{WCA}]^- + \text{pyrazine}$ ($n = 2, 3$)		164
$[\text{Ga}(\text{PPh}_3)_3]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{Ga}(\text{C}_6\text{H}_5\text{Me})_2]^+[\text{WCA}]^- + \text{PPh}_3$		31, 91 and 92
$[(^1\text{Pr}_2\text{-ATI})\text{InMe}]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	Thermolysis of $[(^1\text{Pr}_2\text{-ATI})(\text{CPh}_3)\text{InMe}_2]^+[\text{WCA}]^-$		190
$[\{\text{ArN}(\text{CPh})_2(\text{NC}_5\text{H}_3)\text{In}\}^+ (\text{Ar} = 2,4\text{-}^t\text{Bu}_2\text{C}_6\text{H}_3, 2,5\text{-}^t\text{Bu}_2\text{C}_6\text{H}_3, 2,6\text{-Et}_2\text{C}_6\text{H}_3, 2,6\text{-}^1\text{Pr}_2\text{C}_6\text{H}_3)]$	$[\text{OTf}]^-$	Com	$\text{In}^+[\text{WCA}]^- + \text{bis(imino)pyridine ligand}$		188 and 189
$[\text{In}(\text{PPh}_3)_3]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{In}(\text{C}_6\text{H}_5\text{F})_4]^+[\text{WCA}]^- + 3 \text{ PPh}_3$ ($n = 2, 3$)	Trigonal pyramidal (cf. gallium analogue)	162
$[\text{Ti}(\text{Mes}_2\text{py})(\eta^6\text{-C}_6\text{H}_3\text{R})_2]^+$ (R = F, Me)	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt, Com	$\text{TiCl} + \text{Na}^+[\text{WCA}]^- + \text{Mes}_2\text{py}$ in $\text{C}_6\text{H}_5\text{R}$		187
$[\text{Ti}(\text{timtmb}^{\text{tBu}})]^+$	$[\text{OTf}]^-$	Com	$\text{Ti}^+[\text{WCA}]^- + \text{timtmb}^{\text{tBu}}$		205



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\{\text{ArN}=\text{CPh}\}_2(\text{NC}_6\text{H}_3)\text{Ti}]^+$ (Ar = 2,6-Et ₂ C ₆ H ₃ , 2,5-'Bu ₂ C ₆ H ₃)	[OTf] [−]	Com	Ti ⁺ [WCA] [−] + bis(imino)pyridine ligand		206
Ligand substituted (CN = 4)					
$[\{(\text{PPh}_3)_2\text{C}\}\text{BH}_2(\text{DMAP})]^+$	$[\text{HB}(\text{C}_6\text{F}_5)_3]^-$	Com	$[\{(\text{PPh}_3)_2\text{C}\}\text{BH}_2]^+[\text{WCA}]^- + \text{DMAP}$		118 and 119
$[\text{BH}_2(\text{PR}_2\text{H})_2]^+$ (R = 'Bu, Cy, Ph)	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt	$[\text{BH}_2(\text{PR}_2\text{H})]^+\text{Br}^- + \text{Na}^+[\text{WCA}]^-$		120
$[(1\text{-MIM})_2(9\text{BBN})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$[\text{PMAF-9BBN}]^+[\text{WCA}]^- + 1\text{-MIM}$		212
$[\text{Me}_2\text{Al}(\text{OEt}_2)_2]^+$ $[\text{Me}_2\text{Al}(\text{THF})_2]^+$ $[\text{Me}_2\text{Al}(\text{NPhMe}_2)_2]^+$ $[\text{H}_2\text{Al}(\text{NMe}_3)_2]^+$	$[\text{MeB}(\text{C}_{12}\text{F}_9)_3]^-$ $[\{\text{Me}_2\text{Si}(\text{NDIPP})_2\}_2\text{Zr}_2\text{Cl}_3]^-$ $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ $[(\text{AlH})_6(\text{CCH}_2'\text{Bu})_6]^{2-}$	Alk Alk Prot, Com Other	AlMe ₃ + B ₄ (C ₁₂ F ₉) ₃ in Et ₂ O Al ₂ Me ₆ + {Me ₂ Si(NDIPP) ₂ }ZrCl ₃ (THF) ₂ Al ₂ Me ₆ + [HNMe ₂ Ph] [−] [WCA] [−] 'Bu≡CLi + AlH ₃ ·NMe ₃ + ClAlH ₂ ·NMe ₃ + [^t BuCH ₂ (Bz)NMe ₂] [−] Cl [−]	— — — —	77 143 144 145
$[(\text{Pytsi})\text{AlMe}]^+$	$[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$	Alk	(Pytsi)AlMe ₂ + B(C ₆ F ₅) ₃		137
$[\text{H}_2\text{C}\{\text{hpp}\}_2\text{AlMe}_2]^+$	$[\text{BPh}_4]^-$	Prot	$[\{\text{hpp}\}\text{H}_2\text{C}\{\text{hpp}\}\text{H}]^+[\text{WCA}]^- + \text{AlMe}_3$		138
$[\{\text{H}_2\text{C}=\text{C}(\text{BOX-Me}_2)_2\}_2\text{Al}(\text{Me})_2]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Hyd	$\{\text{BOX-Me}_2\}\text{Al}(\text{Me})_2 + [\text{CPh}_3]^+[\text{WCA}]^-$		139
$[\{\text{BOX-Me}_2\}\text{Al}(\text{Me})(\text{NMe}_2\text{Ph})]^+$	$[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$	Alk	$\{\text{BOX-Me}_2\}\text{Al}(\text{Me})_2 + \text{B}(\text{C}_6\text{F}_5)_3$ in NMe ₂ Ph		139



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[(6-(CH_2NMe_2)_2-2-CPh_3-4-Me-C_6H_2O)Al(C_6H_5O)Al(tBu)(NMe_2Ph)]^+$	$[HB(C_6F_5)_3]^-$	Hyd	$\{6-(CH_2NMe_2)_2-2-CPh_3-4-Me-C_6H_2O\}Al-(tBu)_2 + B(C_6F_5)_3 + NMe_2Ph$		213 and 214
$[(HC(CPhNSiMe_3)_2)Al(Do)Me]^+$ (Do = Et_2O, THF)	$[B(C_6F_5)_4]^-$ / $[MeB(C_6F_5)_3]^-$	Prot/Alk, Com	$\{HC(CPhNSiMe_3)_2\}AlMe_2 + [HNMe_2Ph]^+[WCA]^- + Et_2O/B(C_6F_5)_3 + THF$		215
$[(ArN)C(Me)CHPPh_2(NAr)AlMe(OEt_2)]^+$ (Ar = DIPP)	$[B(C_6F_5)_4]^-$	Alk, Com	$(ArN)C(Me)CHPPh_2(NAr)Me_2 + [Ph_3C]^+[WCA]^-$ in Et_2O		216
$[(^iPr_2-ATT)Al(Et)(Do)]^+$ (Do = ClPh, NCMe)	$[B(C_6F_5)_4]^-$	Alk, Com	$(^iPr_2-ATT)AlEt_2 + [CPh_3]^+[WCA]^-$ in PhCl / + MeCN		140 and 141
$[(SchNMe_2)AlMe]^+$	$[BPh_4]^-$	Salt	$(SchNMe_2)AlMeCl + Na^+[WCA]^-$		142
$[(^i\eta^2-O,P-(2-PPh_2-4-Me-6-tBu-C_6H_2O))_2Al]^+$	$[MeB(C_6F_5)_3]^-$	Alk	$\{^i\eta^2-O,P-(2-PPh_2-4-Me-6-tBu-C_6H_2O)\}_2AlMe + B(C_6F_5)_3$		217
$[H_2C=C(BOX-Me_2)_2]Ga(Me)_2]^+$	$[B(C_6F_5)_4]^-$	Hyd	$\{BOX-Me_2\}Ga(Me)_2 + [CPh_3]^+[WCA]^-$		165
$[BOX-Me_2]Ga(Me)]^+$	$[MeB(C_6F_5)_3]^-$	Alk	$\{BOX-Me_2\}Ga(Me)_2 + B(C_6F_5)_3$ in NMe_2Ph		165
$[(^iPr_2-ATT)Ga(Me)(ClPh)]^+$	$[B(C_6F_5)_4]^-$	Alk, Com	$(^iPr_2-ATT)GaMe_2 + [CPh_3]^+[WCA]^-$ in PhCl		140



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\{1,2-(\text{N}^i\text{Pr})_2-5\text{-CPh}_3\text{-cyclohepta-3,6-diene}\}\text{InMe}_2]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$(^i\text{Pr}_2\text{-ATI})\text{InMe}_2 + [\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		190
$[(^i\text{Pr}_2\text{-ATI})\text{In}(\text{Me})(\text{NMe}_2\text{Ph})]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Prot	$(^i\text{Pr}_2\text{-ATI})\text{InMe}_2 + [\text{HNNMe}_2\text{Ph}]^+[\text{WCA}]^-$		190
$[\text{Ti}(\text{OEt}_2)_4]^+$	$[\text{H}_2\text{N}\{\text{B}(\text{C}_6\text{F}_5)_3\}_2]^-$	Prot	$\text{TiOEt} + [\text{H}(\text{OEt}_2)_2]^+[\text{WCA}]^-$ in Et_2O		203
Ligand substituted (CN = 5)					
$[\{\text{Salen}^{\text{CF}_3}\}\text{Al}(\text{OEt}_2)]^+$	$[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$	Alk	$\{\text{Salen}^{\text{CF}_3}\}\text{AlMe} + \text{B}(\text{C}_6\text{F}_5)_3$ in Et_2O		147
$[\text{Ga}(\eta^1\text{-C}_3\text{H}_5)_2(\text{THF})_n]^+ (n = 2, 3)$	$[\text{B}(\text{C}_6\text{F}_5)_4]^- / [\text{B}(\text{Ar}^{\text{Cl}})_4]^-$	Prot	$\text{Ga}(\eta^1\text{-C}_3\text{H}_5)_3(\text{THF}) + [\text{HNNMe}_2\text{Ph}]^+[\text{WCA}]^-$		166
$[\text{In}(\text{CH}_2\text{SiMe}_3)_2(\text{THF})_3]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Prot	$\text{In}(\text{CH}_2\text{SiMe}_3)_3 + [\text{HNNMe}_2\text{Ph}]^+[\text{WCA}]^-$ in THF		191
$[\text{Ti}(\text{NPPh})_2(\eta^6\text{-C}_6\text{H}_5\text{Me})]^+$ NPPh = 2,5-bis(2-pyridyl)-1-phenylphosphole	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{N}^+ \text{PPh} =$		218
Ligand substituted (CN ≥ 6)					
$[\text{DoAl}(\text{MeOH})_2]^+$ (Do = Salen, Acen)	$[\text{BPh}_4]^-$	Salt, Com	$\text{DoAlCl} + \text{Na}^+[\text{WCA}]^- + \text{MeOH}$		148 and 149



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\text{salpen}^t\text{Bu})\text{Al}(\text{THF})_2]^+$	$[\text{BPh}_4]^-$	Salt, Com	$\text{Salpen}^t\text{Bu})\text{AlCl} + \text{Na}^+[\text{WCA}]^- + \text{THF}$		150 and 151
$[\text{SchNMMe}_2)\text{Al}(\text{OPh})(\text{THF})_2]^+$	$[\text{BPh}_4]^-$	Com	$[\text{SchNMMe}_2)\text{AlPh}]^+[\text{WCA}]^- + \text{O}_2 \text{ in THF}$		142
$[\text{GaH}(\text{THF})_4(\text{OTf})]^+$	$[\text{Ga}(\text{THF})_4(\text{OTf})_2]^-$	Prot	$\text{GaCp}^* + \text{HOSO}_2\text{CF}_3 \text{ in THF}$		167
$[\text{Ga}^{\text{III}}\{(\text{bipy})_3\}^*]^{2+}$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{N} \text{---} \text{N} = \text{C}_6\text{H}_4\text{---} \text{N} \text{---} \text{N}$ ($n = 2, 3$)		17
$[\text{Ga}([18]\text{crown-6})(\eta^6\text{-}\eta^1\text{-C}_6\text{H}_5\text{F})_2]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{Ga}(\eta^6\text{-C}_6\text{H}_5\text{F})]^+[\text{WCA}]^- + [18]\text{crown-6}$ ($n = 2, 3$)		168
$[\text{In}([18]\text{crown-6})]^+$	$[\text{OTf}]^-$	Com	$\text{In}^+[\text{WCA}]^- + [18]\text{crown-6}$	No coordinated solvent, but a strong anion-cation interaction: <i>cf.</i> In-O = 227.2 pm and 278.5 pm (sum of the van der Waals radii 345 pm) Similar structure to the gallium analogue (see above)	192 and 193
$[\text{In}([18]\text{crown-6})(\eta^6\text{-}\eta^1\text{-C}_6\text{H}_5\text{F})_2]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{In}(\eta^6\text{-C}_6\text{H}_5\text{F})_n]^+[\text{WCA}]^- + [18]\text{crown-6}$ ($n = 2, 3$)		168
$[\text{In}([15]\text{crown-5})_2]^+$	$[\text{OTf}]^-$	Com	$\text{In}^+[\text{WCA}]^- + [15]\text{crown-5}$	Sandwich complex	194
$[\{\text{HC}(3,5\text{-Me}_2\text{pz})_3\}_n\text{Ti}]^+$ ($n = 1, 2$)	$[\text{PF}_6]^-$	Com	$\text{Ti}^+[\text{WCA}]^- + \text{HC}(3,5\text{-Me}_2\text{pz})_3$		219
$[\text{Ti}([18]\text{crown-6})]^+$	$[\text{H}_2\text{N}\{\text{B}(\text{C}_6\text{F}_5)_3\}_2]^-$	Com	$[\text{Ti}(\text{C}_6\text{H}_5\text{Me})_2]^+[\text{WCA}]^- + [18]\text{crown-6}$	Similar to gallium analogue, yet featuring significant Ti-F interactions to two counteranions	203



Table 8 (continued)

Cation	WCA	Synthesis	Class ^a	Comment/structure	Ref.
Transition-metal substituted					
$[(FP^*)(BMes)]^+$	$[B(Ar^{CF_3})_4]^-$	$(FP^*)(BMes)Br + Na^+[WCA]^-$	Salt		220
$[CpFe(CO)(PCy_3)(BNCMe_2)]^+$	$[B(Ar^{CF_3})_4]^-$	$CpFe(CO)(PCy_3)(B(Cl)NMe_2) + Na^+[WCA]^-$	Salt		221
$[CpM(CO)(R)\{B(NCy_2)\}]^+$ (M = Fe, Ru; Do = CO, PMe ₃ , PPh ₃)	$[B(Ar^{CF_3})_4]^-$	$CpM(CO)(R)\{B(NCy_2)Cl\} + Na^+[WCA]^-$	Salt		12 and 124
$[(Cy_3P)_2(MeCN)Pt(B \equiv O)]^+$	$[B(Ar^{CF_3})_4]^-$	$(Cy_3P)_2Pt(B \equiv O)(Br) + Ag^+[WCA]^- + MeCN$	Salt, Com		121
$[(OC)_5Mn]_2(\mu-B)]^+$	$[B(Ar^{CF_3})_4]^-$	$\{(OC)_5Mn\}_2(\mu-BBr) + Na^+[WCA]^-$	Salt		222
$[(FP')_2(\mu-B)]^+$	$[B(Ar^{CF_3})_4]^-$	$(FP')_2B(Cl) + Na^+[WCA]^-$	Salt		222
$[Fe(NC_5H_2Me_2)BPh]^+$	$[Al(OR^{PF})_4]^-$	 $Ag^+/Li^+[WCA]^-$	Salt		223
$[(FP)\{B(N^iPr_2)(OPPh_3)\}]^+$	$[B(Ar^{CF_3})_4]^-$	$[(FP)B(N^iPr_2)]^+[WCA]^- + Ph_3PO$	Com		224



Table 8 (continued)

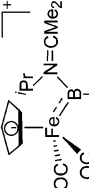
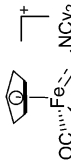
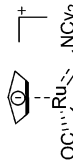
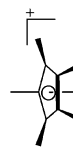
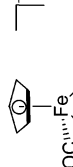
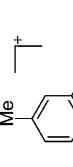
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[(FP)B\{N(^iPr)(CMe_2)(Do)\}^+]$ (Do = Ph ₂ C=O, Me ₂ C=N ⁱ Pr)	$[B(Ar^{CF_3})_4]^-$	Com, other	$[(FP)B=O] + Do + [WCA]^-$	 including a Meerwein-Ponndorf β-hydride transfer	225
$[(FP)B(NCy_2)(Do)]^+$ (Do = C ₅ H ₄ PPh ₃ , 4-Pic)	$[B(Ar^{CF_3})_4]^-$	Com	$[(FP)B=O] + Do + [WCA]^-$		12
$[CpRu(CO)_2(B(NCy_2)(4-Pic))]^+$	$[B(Ar^{CF_3})_4]^-$	Salt, Com	$CpRu(CO)_2(B(NCy_2)Cl) + Na^+[WCA]^- + 4-Pic$		124
$[(FP^*)B(Cl)(LB)]^+$ (Do = 3,5-lutidine, PMe ₃ , IMe)	$[B(Ar^{CF_3})_4]^-$	Salt	$(FP^*)B(Cl)_2(Do) + Na^+[WCA]^-$		226
$[(FP^*)B(nacnac)]^+$	$[B(Ar^{CF_3})_4]^-$	Salt, other	$Cp^*Fe(CO)_2(B(NCy_2)Cl) + Na^+[WCA]^-$		227
$[(FP)C(NCy)_2BNR_2]^+$ (R = ⁱ Pr, Cy)	$[B(Ar^{CF_3})_4]^-$	Ins	$[(FP)(BNR_2)_2][WCA]^- + RN=C=NR$ (substoichiometric)		228
$[(H)(PNP)Pd(BCat)]^+$	$[B(Ar^{CF_3})_4]^-$	Other	$[(BCat)(PNP)Pd(BCat)][WCA]^- + H_2O$		117



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[(R_3P)_2Pt(B(Fe)Br)]^+$ ($R = iPr, Cy$)	$[B(Ar^{CF_3})_4]^-$	Salt	$(R_3P)_2Pt(Br)\{B(Fe)Br\} + Na^+[WCA]^-$		229 and 230
$[(Cy_3P)_2Pt(B(X)X')]^+$ ($X = Br; X' = ortho\text{-}tolyl, iBu, NMe_2, [B(Ar^{CF_3})_4]^- [B(C_6F_5)_4]^-$ Pip, Br; $XX' = (NMe_2)_2, CatB$)	$[B(Ar^{CF_3})_4]^- [B(C_6F_5)_4]^-$	Salt	$(Cy_3P)_2Pt(Br)\{B(X)X'\} + Na^+[WCA]^-$		231
$[(Cy_3P)_2Pt(Br)\{B(NC_5H_4-4-R)X\}]^+$ ($R = Me, X = NMe_2, Pip, Br;$ $R = iBu, X = Pip$)	$[B(Ar^{CF_3})_4]^-$	Salt	$(Cy_3P)_2Pt(Br)\{B(Br)\{B(NC_5H_4-4-R)X\} + Na^+[WCA]^-$		125
$[(Cy_3P)_2Pt(B(Br)(NMe_2))-(NCMe)]^+$	$[B(Ar^{CF_3})_4]^- [B_{12}Cl_{12}]^{2-}$	Com/salt, Com	$(Cy_3P)_2Pt(B(Br)(NMe_2)) + NCMe/(Cy_3P)_2Pt(B(Br)(NMe_2))Br +\{Na^+\}_2[WCA]^{2-} + MeCN$		230
$[(Cy_3P)_2Pt(BCl_2)]^+$	$[B(Ar^{CF_3})_4]^-$	Salt	$(Cy_3P)_2Pt(BCl_2)Cl + Na^+[WCA]^-$		230
$[Cp^*Ru(P^iPr_3)(BH_2Mes)]^+$	$[B(C_6F_5)_4]^-$	Salt	$Cp^*Ru(P^iPr_3)(BH_2MesCl) +Li^+[WCA]^- \cdot 2.5OEt_2$		232
$[(PMAF)_2BH_2]^+$	$[B(C_6F_5)_4]^-$	Hyd, Com	$PMAF-BH_3 + [CPh_3]^+[WCA]^- + PMAF$		212
$[Rh(PPh_3)_2(\kappa^1-\eta\text{-}PPh_2BH_2\cdot PPh_3)]^+$	$[B(Ar^{CF_3})_4]^-$	Salt, Com	$ClRh(PPh_3)_3 + Na^+[WCA]^- + H_3B\cdot PPh_2H$		120
$[FeBMe(bipy)]^+$	$[PF_6]^-$	Salt, Com	$FeBBMe + bipy + [NH_4]^+[WCA]^-$		126



Table 8 (continued)

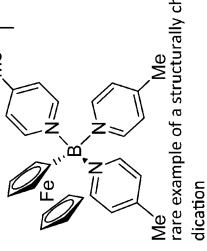
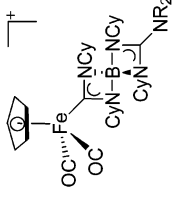
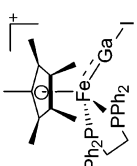
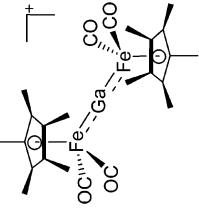
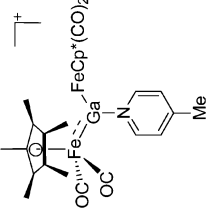
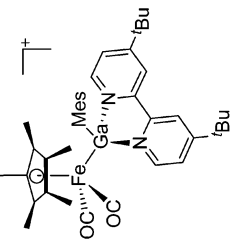
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\text{FeB}(\text{Pic})_3]^{2+}$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt, Com	$\text{Br}_2\text{BFc} + 2\text{Na}^+[\text{WCA}]^- + 3\text{Pic}$	 rare example of a structurally characterized boron dication	122
$[(\text{FP})\{\text{C}(\text{NCy})_2\text{B}(\text{NCy})_2\text{CNR}_2\}]^+$ (R = Pr, Cy)	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Ins	$[(\text{FP})\{\text{B}(\text{NR}_2)\}]^+ + \text{CyN}=\text{C}=\text{NCy}$		124 and 228
$[(\text{dppe})\text{Cp}^*\text{FeGa}]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt	$(\text{dppe})\text{Cp}^*\text{FeGaI}_2 + \text{Na}^+[\text{WCA}]^-$		233
$[(\text{FP}^*)_2\text{Ga}]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt	$(\text{FP}^*)_2\text{GaCl} + \text{Na}^+[\text{WCA}]^-$		169
$[(\text{FP}^*)_2\text{Ga}(4\text{-Pic})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt, Com	$(\text{FP}^*)_2\text{GaCl} + \text{Na}^+[\text{WCA}]^- + 4\text{-Pic}$		170
$[(\text{FP}^*)_2\text{Ga}(\text{Mes})(\text{dtbpy})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt, Com	$(\text{FP}^*)_2\text{Ga}(\text{Mes})\text{I} + \text{Na}^+[\text{WCA}]^- + \text{dtbpy}$		234



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[(FP^*)Ga(phen)(Y)]^+$ (Y = Cl, <i>S</i> ^p Tol) $[BPh_4]^-$		Salt, Com/Lewis	$2(FP^*)GaCl_2 + Na^+[WCA]^- + phen/[FP^*]Ga(phen)(Cl)]^+ + Me_3Si^pTol$		171
$[(FP)Ga(OEt_2)\{(NCy)_2C^tBu\}]^+$	$[B(Ar^{CE_3})_4]^-$	Salt, Com	$(FP)Ga(Cl)\{(NCy)_2C^tBu\} + Na^+[WCA]^-$ in Et_2O		235
$[(FP)_2Ga(bipy)]^+$	$[Cl_2Ga(FP)_2]^-$	Lewis, Com	$2ClGa(FP)_2 + bipy$		236
$[InPt(PPh_3)_3]^+$	$[B(Ar^{CE_3})_4]^-$	Com	$In^+[WCA]^- + Pt(PPh_3)_4$		179 and 180
$[(phen)_2In-Ag(\eta^3-C_6H_5F)]^{2+}$	$[Al(OR^{PF})_4]^-$	Com	$N \equiv N =$		17
				first cationic indium-silver compound	
$[(FP^*)_2In]^+$	$[B(Ar^{CE_3})_4]^-$	Salt	$(FP^*)_2InCl + Na^+[WCA]^-$		195
$[(FP^*)_2In(THF)]^+$	$[B(Ar^{CE_3})_4]^-$	Com	$[(FP^*)_2In]^+[WCA]^- + THF$		195
$[Ti(\eta^5-FeCp_2)]^+$	$[H_2N\{B(C_6F_5)_3\}_2]^-$	Others	$[Ti(\eta^6-C_6H_5Me)_3]^+[WCA]^- + 2.2FeCp_2$		204

Table 8 (continued)

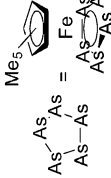
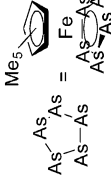
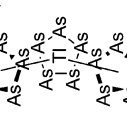
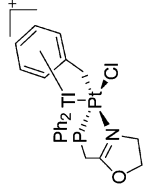
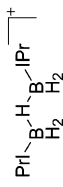
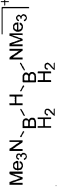
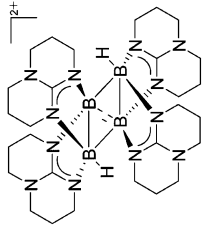
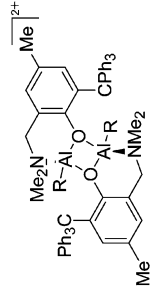
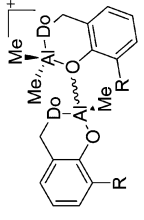
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\text{Ti}_2(\eta^5\text{-FeCp}_2)_3]^{2+}$	$[\text{H}_2\text{N}\{\text{B}(\text{C}_6\text{F}_5)_3\}_2]^-$	Com	$[\text{Ti}(\eta^6\text{-C}_6\text{H}_5\text{Me})_2][\text{WCA}]^- + \text{FeCp}_2$ $\text{Cp}^*\text{Fe}(\eta^5\text{-As}_5) + \text{Ti}^+[\text{PF}_6]^- + \text{Li}^+[\text{WCA}]^-$ 	 1:1 mixture of $[\text{Ti}(\eta^5\text{-FeCp}_2)]^+$ and $[\text{Ti}(\eta^5\text{-FeCp}_2)_2]^+$	203
$[\text{Ti}\{\{(\eta^5\text{-As}_5)\text{FeCp}^*\}_3\}^+]$	$[\text{FAl}\{\text{OC}_6\text{F}_{10}(\text{C}_6\text{F}_5)_3\}_3]^-$	Salt, Com		 trigonal planar complex with no short distances to the next units	176
$[\{\text{P}(\text{Ph}_2)\text{CH}_2\text{ox}\}(\text{Cl})(\text{Ti})\text{Pt-CH}_2\text{Ph}\}]^+ [\text{PF}_6]^-$		Other	$\text{Ti}^+[\text{WCA}]^- + \{\text{P}(\text{Ph}_2)\text{CH}_2\text{ox}\}\text{Pt}(\text{Cl})\text{-CH}_2\text{Ph}\}$		208
Multinuclear					
$[\{\text{IPr}(\text{H}_2\text{B})_2(\mu\text{-H})\}^+]$	$[\text{HB}(\text{C}_6\text{F}_5)_3]^-$	Hyd	$\text{IPr} + \text{B}(\text{C}_6\text{F}_5)_3$		118 and 119
$[\{\text{Me}_3\text{N}(\text{H}_2\text{B})_2(\mu\text{-H})\}^+]$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Hyd	$\text{Me}_3\text{N-BH}_3 + [\text{CPh}_3][\text{WCA}]^-$		237
$[\text{B}_4\text{H}_2(\mu\text{-hpp})_4]^{2+}$	$[\text{HB}(\text{C}_6\text{F}_5)_3]^-$	Hyd, Com	$[\text{HB}(\mu\text{-hpp})]_2 + \text{B}(\text{C}_6\text{F}_5)_3$		123
$[\{6\text{-(CH}_2\text{NMe}_2\text{)}_2\text{-2-CPh}_3\text{-4-Me-C}_6\text{H}_2\text{O}\}\text{Al}(\text{R})\}_2]^{2+}$ (R = C ₆ H ₁₃)	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com	$[\{6\text{-(CH}_2\text{NMe}_2\text{)}_2\text{-2-CPh}_3\text{-4-Me-C}_6\text{H}_2\text{O}\}\text{Al}(\text{R})\}_2]^{2+} + 1\text{-hexene}$		152
$[\{2\text{-(CH}_2\text{Do)}_6\text{-R-C}_6\text{H}_5\text{O}\}\text{AlMe-}\{2\text{-(CH}_2\text{Do)}_6\text{-R-C}_6\text{H}_5\text{O}\}\text{AlMe}_2\}^+]$ (R = Ph, 'Bu; Do = NMe ₂ , NC ₄ H ₈ , NC ₃ H ₁₀)	$[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$	Alk	$\{2\text{-(CH}_2\text{Do)}_6\text{-R-C}_6\text{H}_5\text{O}\}\text{AlMe}_2 + \text{B}(\text{C}_6\text{F}_5)_3$		238 and 214





Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\{\text{MeC}(\text{NR})_2\}_2\text{Al}_2\text{Me}_3]^+$ ($\text{R} = \text{iPr}, \text{Cy}$) $[\text{B}(\text{C}_6\text{F}_5)_4]^-$ / $[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$		Alk, Com	$\{\text{MeC}(\text{NR})_2\}\text{AlMe}_2 + [\text{CPh}_3]^+[\text{WCA}]^-$ / $\text{B}(\text{C}_6\text{F}_5)_3$		156
$[\text{AlEt}(\mu-\eta^2, \eta^1-\text{iPr}_2\text{-ATT})-(\mu\text{-Et})\text{AlEt}_2]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com	$[(\text{iPr}_2\text{-ATT})\text{Al}(\text{Et})][\text{WCA}]^- + \text{AlEt}_3$		141
$[\{(\text{iPr}_2\text{-ATT})\text{AlMe}_2(\mu\text{-Me})\}^+]$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Alk, Com	$(\text{iPr}_2\text{-ATT})\text{AlMe}_2 + [\text{CPh}_3]^+[\text{WCA}]^-$		141 and 239
$[\{(\text{iPr}_2\text{-ATT})\text{Al}(\mu\text{-O}^i\text{Pr})\}_2]^{2+}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com	$[(\text{iPr}_2\text{-ATT})\text{Al}(\text{Et})][\text{WCA}]^- + \text{acetone}$		141 and 153
$[\text{Me}_2\text{Al}(\mu\text{-OSi}(\text{R}^{123})_3)_2\text{AlMe}-(\text{NMe}_2\text{Ph})]^+$ ($\text{R}^1, \text{R}^2 = \text{Me}; \text{R}^3 = \text{Me}, \text{iBu}$)	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Prot	$\text{Me}_2\text{Al}(\mu\text{-OSiR}_3)_2\text{AlMe}_2 + [\text{HNMe}_2\text{Ph}]^+[\text{WCA}]^-$		154
$[\{(\text{iPr}_2\text{-ATT})\text{Al}-(\mu\text{-C}\equiv\text{C}^i\text{Bu})\}_2]^{2+}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com	$[(\text{iPr}_2\text{-ATT})\text{Al}(\text{Et})][\text{WCA}]^- + \text{tert-butyl acetylene}$		141 and 153
$[\{(\text{tacn})\text{AlMe}_2\}_2]^{2+}$	$[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$	Alk	$[(\text{tacn})\text{AlMe}_2]_2 + \text{B}(\text{C}_6\text{F}_5)_3$		240
$[\{(\text{OSSO})\text{Al}\}_2]^{2+}$	$[\text{MeB}(\text{C}_6\text{F}_5)_4]^-$	Alk	$(\text{OSSO})\text{AlMe} + \text{B}(\text{C}_6\text{F}_5)_3$		155



Table 8 (continued)

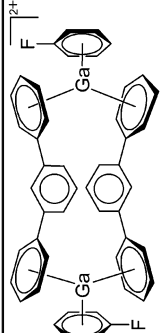
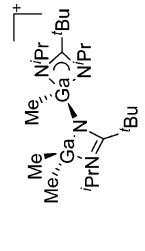
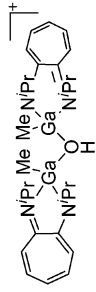
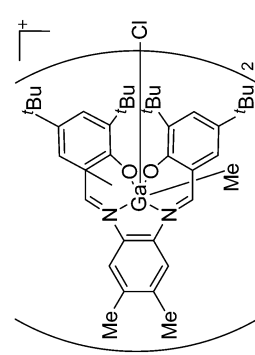
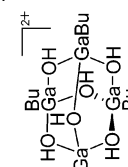
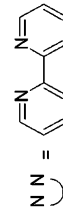
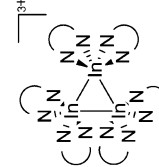
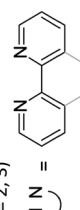
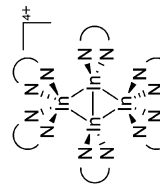
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[(\eta^6\text{-C}_6\text{H}_5\text{F})\text{Ga}(\mu\text{-}\eta^6\text{-}m\text{-TP})_2\text{-Ga}(\eta^6\text{-C}_6\text{H}_5\text{F})]^{2+}$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{Ga}(\eta^6\text{-C}_6\text{H}_5\text{F})_n][\text{WCA}]^- + m\text{-TP} (n = 2, 3)$	 first structurally characterized dicationic ansa-arene complex	8 and 99
$[(\text{BuC}(\text{N}^i\text{Pr})_2)_2\text{GaMe}(\text{BuC}(\text{N}^i\text{Pr})_2)_2\text{GaMe}_2]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Alk, Com	$\{\text{BuC}(\text{N}^i\text{Pr})_2\}_2\text{GaMe}_2 + [\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		156
$[(\text{Pr}_2\text{-ATI})\text{GaMe}(\text{NMe}_2\text{Ph})]^{+}[\text{WCA}]^- + \text{H}_2\text{O}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other			172
$[(\text{Salomphen})\text{Ga}(\mu\text{-Cl})]^+$	$[\text{BPh}_4]^-$	Salt	$(\text{Salomphen})\text{GaCl} + \text{Na}^+[\text{BPh}_4]^-$		241
$[(\text{BuGa})_4(\mu\text{-OH})_6]^{2+}$	$[\text{HCB}_{11}\text{Br}_6\text{Me}_5]^-$	Other	$[(2,6\text{-Mes}_2\text{C}_6\text{H}_3)\text{GaBu}]^+[\text{WCA}]^- + \text{H}_2\text{O}$		173
$[(\text{PPh}_3)_3\text{In}]_2(\mu\text{-PPh}_3)]^{2+}$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{In}(\text{C}_6\text{H}_5\text{F})_n][\text{WCA}]^- + \text{PPh}_3 (n = 2, 3)$	One PPh_3 moiety functions as a bridge between both In^+ cations	162
$[\text{In}_3(\text{bipy})_{5-6}]^{3+}$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{In}(\text{C}_6\text{H}_5\text{F})_n][\text{WCA}]^- + \text{bipy} (n = 2, 3)$ $\text{N} \text{---} \text{N} =$ 	 first cationic indium clusters	17
$[\text{In}_4(\text{Do})_6]^{4+}$ (Do = bipy, phen)	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{In}(\text{C}_6\text{H}_5\text{F})_n][\text{WCA}]^- + \text{phen} (n = 2, 3)$ $\text{N} \text{---} \text{N} =$ 	 first cationic indium cluster	17



Table 8 (continued)

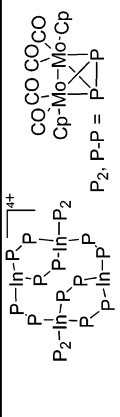
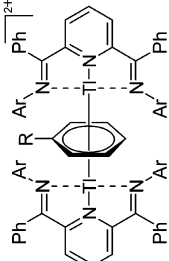
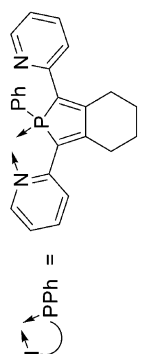
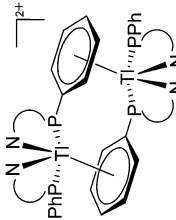
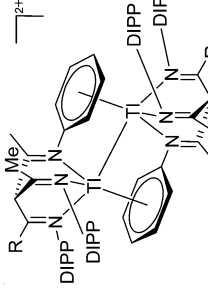
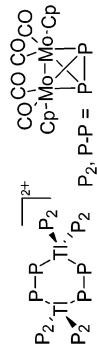
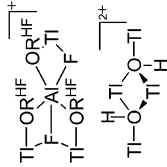
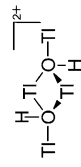
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\text{In}_4\{(\text{CpMo}(\text{CO})_2)_2\text{P}_2\}_8]^{4+}$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$[\text{In}(\text{o-C}_6\text{H}_4\text{F}_2)_2][\text{WCA}]^- + \{\text{CpMo}(\text{CO})_2\}_2(\text{P}_2)$		186
$[\{(\text{ArN}=\text{CPh})_2(\text{NC}_5\text{H}_3)\text{Ti}\}_2(\mu-\eta^6\text{-C}_6\text{H}_5\text{R})]^{2+}$ (Ar = 2,6-Et ₂ -C ₆ H ₃ , 2,5'-Bu ₂ -C ₆ H ₃ ; R = H, Me)	$[\text{OTf}]^-$	Com	$2[(\text{ArN}=\text{CPh})_2(\text{C}_5\text{H}_3\text{N})\text{Ti}][\text{WCA}]^- + \text{C}_6\text{H}_5\text{R}$		206
$[\text{Ti}_2(\text{NPPH})_4]^{2+}$ NPPH = 2,5-bis(2-pyridyl)-1-phenylphosphole	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{Ti}^+[\text{WCA}]^- + \text{NPPH in CH}_2\text{Cl}_2 / n\text{-pentane}$ 	 If $[\text{PF}_6]^-$ is employed as WCA, a one-dimensional coordination polymer forms	218
$[\text{Ti}(\beta\text{-triketimine})_2]^{2+}$ (R = Me, ^t Bu) $[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$		Com	$\text{Ti}^+[\text{WCA}]^- + \beta\text{-triketimine}$		14
$[\text{Ti}_2\{(\text{CpMo}(\text{CO})_2)_2\text{P}_2\}_6]^{2+}$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\{\text{CpMo}(\text{CO})_2\}_2(\text{P}_2) + \text{Ti}^+[\text{WCA}]^-$		207
$[\text{Ti}_3\text{F}_2\text{Al}(\text{OR}^{\text{HF}})_3]^+$	$[\text{Al}(\text{OR}^{\text{HF}})_4]^-$	Salt, other TIF + 2Li ⁺ [WCA] ⁻			200
$[\text{Ti}_4(\mu\text{-OH})_2]^{2+}$	$[\text{H}_2\text{N}\{\text{B}(\text{C}_6\text{F}_5)_3\}_2]^-$	Other	$[\text{Ti}(\text{OEt})_2][\text{WCA}]^- + \text{H}_2\text{O}$		203
$[\{\text{Ti}(\text{OR})_4(\mu\text{-Cl})_2\}_n]^+$	$[\text{PF}_6]^-$	Other	$\text{RuCl}_2(\text{DMcOPrPE})_2 + \text{Ti}^+[\text{WCA}]^-$	“Arrested” chloride abstraction yielding a one-dimensional coordination polymer	209



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
Multinuclear transition-metal substituted					
$[(\mu\text{-dmpe})\{\text{CpFe}(\text{CO})\{\text{B}(\text{NCy}_2)\}_2\}^{2-}]^{2+}$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com, salt	$(\text{FP})\{\text{B}(\text{NCy}_2)\}_2\text{Cl} + \text{dmpe} + \text{Na}^+[\text{WCA}]^-$		124
$[(\text{BCat})(\text{PNP})\text{Pd}(\text{BCat})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^- / [\text{CB}_{11}\text{H}_{12}]^-$	Other	$[(\text{PNP})\text{Pd}(\text{THF})]^+[\text{WCA}]^- / (\text{PNP})\text{Pd}(\text{CB}_{11}\text{H}_{12}) + \text{CatB-BCat}$		117
$[(\text{Cy}_3\text{P})_2\{\text{Pt}(\text{BBr})\}_2(\mu\text{-C}_6\text{H}_4)]^{2+}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Salt	$\{(\text{Cy}_3\text{P})_2\text{Pt}(\text{BBr})\}_2(\mu\text{-C}_6\text{H}_4) + \text{K}^+[\text{WCA}]^-$		125
$[(\text{Cy}_3\text{P})_2\text{PtB}(\mu\text{-O})_2]^{2+}$	$[\text{Al}(\text{OR}^{\text{PF}_3})_4]^-$	Salt	$(\text{Cy}_3\text{P})_2\text{BrPt}(\text{B}\equiv\text{O}) + \text{Ag}^+[\text{WCA}]^-$		242
$[(\text{bipy})(\text{Me})\text{B}(\mu\text{-Fc})]^{2+}$	$[\text{PF}_6]^-$	Salt, Com	$\text{Fc}(\text{BBRMe})_2 + 2\text{bipy} + [\text{NH}_4]^+[\text{WCA}]^-$		126 and 127
$[(\text{FP})\text{Ga}(\text{Mes})_2(\mu\text{-Cl})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Salt, Com	$(\text{FP})\text{Ga}(\text{Mes})(\text{Cl}) + \text{Na}^+[\text{WCA}]^-$		195
$[(\text{FeCp}(\text{CO})_2)\text{Ga}\{(\text{NCy})_2\text{-C}^t\text{Bu}\}]^{2-}(\mu\text{-OH})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Other	$[\text{FP}\text{Ga}(\text{OEt}_2)\{(\text{NCy})_2\text{-C}^t\text{Bu}\}]^+[\text{WCA}]^- + \text{H}_2\text{O}$		235



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\{\text{Ga}(\text{P}_5\text{FeCp}^*)_3\}_n]^{2+}$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{P}_5\text{FeCp}^* + [\text{Ga}(\sigma\text{-C}_6\text{H}_4\text{F}_2)]^+[\text{WCA}]^-$ 	 one-dimensional coordination polymer	176
$[\{(\text{DDP})(\text{THF})\text{Ga}\}_2\text{Au}]^{+}$	$[\text{B}(\text{Ar}^{\text{CE}_3})_4]^-$	Salt, Com	$\{(\text{DDP})\text{Ga}\}_2\text{AuCl} + \text{Na}^+[\text{WCA}]^-$ in THF		174
$[\{(\text{THF})(\text{DDP})\text{GaZn}(\text{THF})\}_2]^{2-}$ $[\mu\text{-Cl}]_2^{2+}$	$[\text{B}(\text{Ar}^{\text{CE}_3})_4]^-$	Salt, Com	$(\text{DDP})(\text{Cl})\text{GaZn}(\text{Cl})(\text{THF})_2 + \text{Na}^+[\text{WCA}]^-$ in THF		175
$[\{\text{In}\{\eta^5\text{-E}_5\}\text{FeCp}^*\}_3]^{1+}$ (E = P, As)	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{E}_5\text{FeCp}^* + \text{In}^+[\text{WCA}]^-$ 	One-dimensional coordination polymer/similar structure to $[\{\text{Ga}(\text{P}_5\text{FeCp}^*)_3\}_n]^{2+}$	176
$[\{\text{Tl}\{\eta^5\text{-E}_5\}\text{FeCp}^*\}_3]^{1+}$ (E = P, As)	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{E}_5\text{FeCp}^* + \text{Tl}^+[\text{WCA}]^-$ 	One-dimensional coordination polymer/similar structure to $[\{\text{Ga}(\text{P}_5\text{FeCp}^*)_3\}_n]^{2+}$	207 and 176
ECp* substituted (E = Al, Ga)					
$[\text{Rh}(\text{COD})(\text{AlCp}^*)_3]^{+}$	$[\text{B}(\text{Ar}^{\text{CE}_3})_4]^-$	Com	$[\text{Rh}(\text{COD})_2]^+[\text{WCA}]^- + 3\text{AlCp}^*$		157
$[\text{Cp}^*\text{Fe}(\text{GaCp}^*)_3]^{+}$	$[\text{B}(\text{Ar}^{\text{CE}_3})_4]^-$	Com	$[\text{Fe}(\text{MeCN})_6]^{2+}[\text{WCA}]^- + 4\text{GaCp}^*$		243



Table 8 (continued)

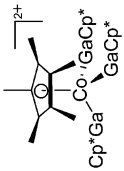
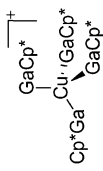
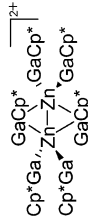
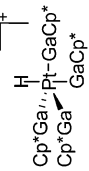
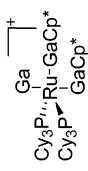
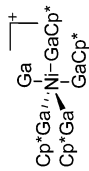
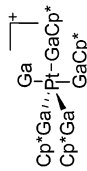
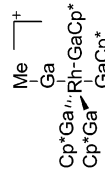
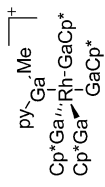
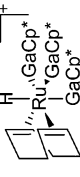
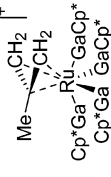
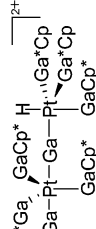
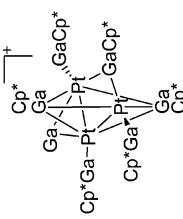
Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[\text{Cp}^*\text{Co}(\text{GaCp}^*)_3]^{2+}$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Ox, Com	$[\text{Co}(\text{MeCN})_6]^{2+}[\text{WCA}]^-_2 + 4\text{GaCp}^*$		243
$[\text{Cu}(\text{GaCp}^*)_4]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$[\text{Cu}(\text{MeCN})_4]^+[\text{WCA}]^- + 4\text{GaCp}^*$		243
$[\text{Zn}(\text{GaCp}^*)_4]^{2+}$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Prot, Com	$\text{ZnMe}_2 + [\text{H}(\text{OEt}_2)_2]^+[\text{WCA}]^- + 4\text{GaCp}^*$		175
$[\text{Zn}_2(\text{GaCp}^*)_6]^{2+}$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Other	$\text{Zn}_2\text{Cp}^*_2 + [\text{Ga}_2\text{Cp}^*]^+[\text{WCA}]^-$ mechanism unclear	 stabilized by dispersion forces	244
$[\text{Rh}(\text{COD})(\text{GaCp}^*)_3]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$[\text{Rh}(\text{COD})_2]^+[\text{WCA}]^- + 3\text{GaCp}^*$		157
$[\{\text{Rh}(\text{NBD})(\text{PCy}_3)_2(\text{GaCp}^*)_2\}]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$[\text{Rh}(\text{NBD})(\text{PCy}_3)_2]^+[\text{WCA}]^- + 2\text{GaCp}^*$		157
$[\text{Pt}(\text{H})(\text{GaCp}^*)_4]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Prot	$\text{Pt}(\text{GaCp}^*)_4 + [\text{H}(\text{OEt}_2)_2]^+[\text{WCA}]^-$		180
$[(\text{Ga})\text{Ru}(\text{PCy}_3)_2(\text{GaCp}^*)_2]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Other	$\text{Ru}(\text{PCy}_3)_2(\text{GaCp}^*)_2(\text{H})_2 + [\text{Ga}_2\text{Cp}^*]^+[\text{WCA}]^-$		177
$[(\text{Ga})\text{Ni}(\text{GaCp}^*)_4]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Other	$\text{Ni}(\text{GaCp}^*)_4 + [\text{FeCp}_2]^+[\text{WCA}]^-$		178
$[(\text{Ga})\text{Pt}(\text{GaCp}^*)_4]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$\text{Pt}(\text{GaCp}^*)_4 + [\text{Ga}_2\text{Cp}^*]^+[\text{WCA}]^-$		179 and 180
$[(\text{Cp}^*\text{Ga})_4\text{Rh}(\text{Ga}(\text{Me}))]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Prot	$(\text{Cp}^*\text{Ga})_4\text{Rh}(\eta^1\text{-Cp}^*\text{GaMe}) + [\text{H}(\text{OEt}_2)_2]^+[\text{WCA}]^-$		181



Table 8 (continued)

Cation	WCA	Class ^a	Synthesis	Comment/structure	Ref.
$[(\text{Cp}^*\text{Ga})_4\text{Rh}\{\text{Ga}(\text{Me})(\text{py})\}]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$[(\text{Cp}^*\text{Ga})_4\text{Rh}(\text{GaMe})]^+[\text{WCA}]^- + \text{py}$		181
$[\text{Ru}(\text{COD})(\text{H})(\text{GaCp}^*)_3]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$[\text{Ru}(\text{COD})(\text{H})(\text{DMH})_3]^+[\text{WCA}]^- + 3\text{GaCp}^*$		245
$[\text{Ru}(\text{GaCp}^*)_4\{\eta^3\text{-(CH}_2)_2\text{C}(\text{Me})\}]^{2+}$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Prot	$\text{Ru}(\text{GaCp}^*)_3(\text{TMM}) + [\text{H}(\text{OEt}_2)_2]^+[\text{WCA}]^-$ $\text{TMM} = \eta^1\text{-C}(\text{CH}_2)_3$		177
$[\{\text{Ru}(\text{GaCp}^*)_3\{[(\text{CH}_2)_2\text{C}\{\text{CH}_2(\mu\text{-Ga})\}]_2\}^+\}]^{2+}$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Com	$\text{Ru}(\text{GaCp}^*)_3(\text{TMM}) + [\text{Ga}_2\text{Cp}^*]^+[\text{WCA}]^-$		177
$[\{(\text{GaCp}^*)_4\text{Pt}\{\text{Pt}\{(\text{H})(\text{GaCp}^*)_3\}(\mu\text{-Ga})\}^+\}]^{2+}$	$\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Prot, Com	$\text{Pt}(\text{GaCp}^*)_4 + [\text{H}(\text{OEt}_2)_2]^+[\text{WCA}]^-$		180
$[\text{Pt}_3(\text{GaCp}^*)_6(\mu\text{-Ga})]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Other	$\text{Pt}(\text{GaCp}^*)_4 + [\text{FeCp}_2]^+[\text{WCA}]^-$ (substoichiometric)	 metal-rich molecules as discrete models for intermetallic phases	178

^a Classification according to the introduction (Table 2).



Table 9 Overview on structurally characterized cations of group 14

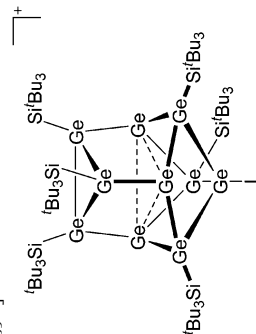
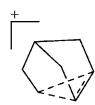
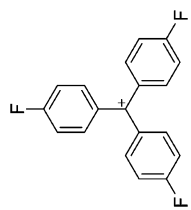
Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
Homopolyatomic and cage cations					
$[\text{C}_{76}]^+$	$[\text{HCB}_{11}\text{H}_3\text{Br}_6]^-$	Ox	$\text{C}_{76} + [\text{Ar}_3\text{N}]^+[\text{WCA}]^-$		44
$[\text{C}_{60}]^+$	$[\text{HCB}_{11}\text{H}_5\text{Cl}_6]^-$	Ox	$\text{C}_{60} + [\text{Ar}_3\text{N}]^+[\text{WCA}]^-$		259
$[\text{C}_{60}]^{2+}$	$[\text{AsF}_6]^-$	Ox	$\text{C}_{60} + 3 \text{AsF}_5$	Polymeric	260
$[\text{HC}_{60}]^+$	$[\text{HCB}_{11}\text{H}_5\text{Cl}_6]^-$	Prot	$\text{C}_{60} + \text{H(WCA)}$		259
$[\text{C}_{59}\text{N}]^+$	$[\text{Ag}(\text{HCB}_{11}\text{H}_5\text{Cl}_6)_2]^-$	Ox	$(\text{C}_{59}\text{N})_2 + 2[\text{HBPC}]^+[\text{WCA}]^-$		261
					
	$[\text{B}(\text{C}_6\text{F}_4\text{H})_4]^-$	Other	$\text{tBu}_3\text{Si}-\text{Ge}(\text{tBu}_3)-\text{Ge}(\text{tBu}_3)-\text{Si}(\text{tBu}_3)-\text{I}$ $\text{tBu}_3\text{Si}-\text{Ge}(\text{tBu}_3)-\text{Ge}(\text{tBu}_3)-\text{Si}(\text{tBu}_3)-\text{I} + \text{KI} + \text{K}^+[\text{WCA}]^-$		332
Onium ions					
	$[\text{Al}_2\text{Br}_7]^-$	Lewis	 $\text{Br} + 2 \text{AlBr}_3$		86
Enium ions					
$[(\text{CH}_3)_3\text{C}]^+$	$[\text{HCB}_{11}\text{Me}_3\text{Cl}_6]^-$	Hyd	$n\text{-BuH} + \text{Me}^+[\text{WCA}]^-$ or $n\text{-BuH} + \text{Me(WCA)}$		265
$[(\text{CH}_3)_3\text{C}]^+$	$[\text{Al}_2\text{Br}_7]^-$	Lewis	$n\text{-BuBr} + 2 \text{AlBr}_3$		266
$[(\text{CH}_3)_3\text{C}]^+$	$[\text{HCB}_{11}\text{Cl}_{11}]^-$	Other	Thermal decomposition of $[\text{Et}_2\text{Cl}][\text{CHB}_{11}\text{Cl}_{11}]$ $n\text{-pentane} + \text{Me(WCA)}$ or Me(WCA)		267
	$[\text{HCB}_{11}\text{Me}_3\text{Br}_6]^-$	Hyd	 Me(WCA)		265
	$[\text{HCB}_{11}\text{Me}_3\text{Br}_6]^-$	Hyd	 Me(WCA) or Me(WCA) + 		265
	$[\text{HCB}_{11}\text{I}_{11}]^-$	Lewis	$p\text{-CH}_3\text{-C}_6\text{F}_4\text{-CF}_3 + \text{Et}_3\text{Si(WCA)} + \text{PhF}$		56



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[HCB ₁₁ I ₁₁] [−]	Lewis	<i>p</i> -CH ₃ -C ₆ F ₄ -CF ₃ + Et ₃ Si(WCA) + PhF		56
	[As ₂ F ₁₁] [−]	Lewis	C ₆ H ₅ CF ₃ + AsF ₅	Excess AsF ₅	268
	[HCB ₁₁ I ₁₁] [−]	Lewis	<i>p</i> -F-C ₆ H ₄ CF ₃ + Et ₃ Si(WCA) + PhF		56
	[HCB ₁₁ I ₁₁] [−]	Lewis	CH ₃ CF ₃ + Et ₃ Si(WCA) + PhF		56
[Cl ₃] ⁺ [CCl ₃] ⁺ [CCl ₃] ⁺ [CBr ₃] ⁺ [CBr ₃] ⁺ [C(OTeF ₅) ₃] ⁺	[Al(OR ^{PF}) ₄] [−] [Sb(OTeF ₅) ₆] [−] [Al(OR ^{PF}) ₄] [−] ; [(R ^{PF} O) ₃ Al-F-Al(OR ^{PF}) ₃] [−] [Sb(OTeF ₅) ₆] [−] [Al(OR ^{PF}) ₄] [−] ; [(R ^{PF} O) ₃ Al-F-Al(OR ^{PF}) ₃] [−] [Sb(OTeF ₅) ₆] [−]	Salt Ox Salt Ox Ox	Cl ₄ + Ag ⁺ [WCA] [−] CCl ₄ + [XeOTeF ₅] ⁺ [WCA] [−] CCl ₄ + Ag ⁺ [WCA] [−] CBr ₄ + [XeOTeF ₅] ⁺ [WCA] [−] CBr ₄ + Ag ⁺ [WCA] [−] CBr ₄ + [XeOTeF ₅] ⁺ [WCA] [−]		269 270 271 270 271 270
	[Al(OR ^{PF}) ₄] [−]	Ox	CS ₂ + [AsBr ₄] ⁺ [WCA] [−] and CS ₂ + Br ₂ + Ag ⁺ [WCA] [−]		117
	[B(C ₆ F ₅) ₄] [−]	Hyd			275
	[B(C ₆ F ₅) ₄] [−]	Hyd			276
[Mes ₃ Si] ⁺ [Pemp ₃ Si] ⁺ [Pemp ₃ Si] ⁺	[HCB ₁₁ Me ₃ Br ₆] [−] [B ₁₂ Cl ₁₂] ^{2−} [Al(OR ^{PF}) ₄] [−]	Other Hyd Hyd	Mes ₃ Si(CH ₂ CH=CH ₂) + Et ₃ Si(WCA) 2Pemp ₃ MeSiH + [Ph ₃ C] ₂ ⁺ [WCA] [−] 1.5Pemp ₃ MeSiH + [Ph ₃ C] ⁺ [WCA] [−]		24 290 291



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
$\text{Bu}_2\text{MeSi}^+\text{Ge}^-\text{SiMe}^-\text{Bu}_2$ 	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Ox	$\text{Bu}_2\text{MeSi}^+\text{Ge}^-\text{SiMe}^-\text{Bu}_2$ $+ [\text{Ph}_3\text{C}]^+\text{[WCA]}^-$		334
	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Salt	$(\text{Ar})_3\text{GeBr} + \text{Ag}^+\text{[WCA]}^-$	R = O'Bu	336
$\text{Bu}_2\text{MeSi}^+\text{Sn}^-\text{SiMe}^-\text{Bu}_2$ 	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Ox	$\text{Bu}_2\text{MeSi}^+\text{Sn}^-\text{SiMe}^-\text{Bu}_2$ $+ [\text{Ph}_3\text{C}]^+\text{[WCA]}^-$		335
$[(\text{Tripp})_3\text{Sn}]^+$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$(\text{allyl})(\text{Tripp})_3\text{Sn} + [\text{E}]^+\text{[WCA]}^-$	$[\text{E}]^+$ not exactly defined, likely $[\text{Et}_3\text{Si}(\text{C}_6\text{H}_5)]^+$ or comparable	34
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com, Lig	Me_3Si Me_3Si	R = Et, iPr	337
Delocalized (cyclic) cations $[\text{C}_6\text{H}_7]^+$	$[\text{HCB}_{11}\text{Me}_3\text{Br}_6]^-$	Prot	$\text{C}_6\text{H}_6 + \text{H(WCA)}$		386
	$[\text{HCB}_{11}\text{H}_3\text{Br}_6]^-$	Prot	$\text{C}_6\text{H}_6 + \text{H(WCA)}$		387
	$[\text{HCB}_{11}\text{H}_3\text{Br}_6]^-$	Prot	$\text{C}_6\text{Me}_2\text{H}_4 + \text{H(WCA)}$		387
	$[\text{HCB}_{11}\text{H}_3\text{Br}_6]^-$	Prot	$\text{C}_6\text{Me}_3\text{H}_3 + \text{H(WCA)}$		387
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Prot	$\text{Et}_3\text{Si(WCA)} + \text{HCl} + \text{C}_6(\text{Me})_5\text{H}$	C-H...F-C interactions	388

Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[HCB ₁₁ H ₅ Br ₆] [−]	Prot	C ₆ Me ₆ + H(WCA)		387
[C ₆ Me ₇] ⁺	[AlCl ₄] [−]	Other	C ₆ Me ₆ + CH ₃ Cl + AlCl ₃	Crystallized out of HF In HF	277
[C ₆ F ₆] ⁺	[Sb ₂ F ₁₁] [−]	Ox	C ₆ F ₆ + [O ₂] ⁺ [WCA] [−]		278
[C ₆ F ₆] ⁺	[Os ₂ F ₁₁] [−]	Ox	C ₆ F ₆ + OsF ₆ + SbF ₅		278
[C ₆ Cl ₆] ⁺	[Sb ₂ F ₁₁] [−]	Ox	C ₆ F ₆ + SbF ₅		279
[C ₆ Br ₆] ⁺	[As ₂ F ₁₁] [−]	Ox	C ₆ Br ₆ + [O ₂] ⁺ [AsF ₆] [−] + HSO ₃ F		280
[C ₆ I ₆] ⁺	[AsF ₆] [−]	Ox	C ₆ I ₆ + AsF ₅	In HF	279
[C ₆ I ₆] ⁺	[SbF ₆] [−]	Ox	C ₆ I ₆ + SbF ₅	In HF	279
[C ₆ I ₆] ⁺	[OTf] [−]	Other	[C ₆ I ₆][AsF ₆] + HOTf		279
[C ₆ HF ₅] ⁺	[AsF ₆] [−]	Ox	C ₆ HF ₅ + [O ₂] ⁺ [WCA] [−]		279
	[SbF ₆] [−]	Ox	C ₆ H ₂ F ₄ + [O ₂] ⁺ [WCA] [−]		279
	[AsF ₆] [−]	Ox	C ₆ H ₃ F ₃ + [O ₂] ⁺ [WCA] [−] + AsF ₅		279
	[SbF ₆] [−]	Ox	C ₆ H ₃ F ₃ + [O ₂] ⁺ [Sb ₂ F ₁₁] [−]	In HF	279
	[Sb ₂ F ₁₁] [−]	Ox	C ₆ F ₅ (CF ₃) + [O ₂] ⁺ [WCA] [−]	In HF	280
	[Sb ₂ F ₁₁] [−]	Ox	C ₆ F ₄ (CF ₃) ₂ + SbF ₅	In HF	280
	[Sb ₂ F ₁₁] [−]	Ox	C ₆ H ₂ Cl ₄ + [O ₂] ⁺ [WCA] [−]		280



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[Sb ₃ F ₁₆] [−]	Ox	[C ₆ F ₅ –C ₆ F ₅] + [O ₂] ⁺ [WCA] [−] + SbF ₅		280
	[Nb ₂ F ₁₁] [−]	Ox			281
	[SbF ₆] [−]	Ox			282
[Cy ₃ C ₃] ⁺ [Cy ₂ PhC ₃] ⁺	[SbF ₆] [−] [BF ₄] [−]	Salt Lewis	[Cy ₃ C ₃]Cl + Ag ⁺ [WCA] [−] [Cy ₂ (Ph)C ₃]F + BF ₃	+ Ag ⁺ [WCA] [−]	283 283
	[SbCl ₆] [−]	Prot		+ HCl + SbCl ₅	284
	[HCB ₁₁ Me ₃ Br ₆] [−]	Hyd	ArMe ₂ SiH + [Ph ₃ C] ⁺ [WCA] [−]		286
	[B(C ₆ F ₅) ₄] [−]	Other	^t Bu ₂ MeSi ₄ SiMe ^t Bu ₂ ^t Bu ₂ MeSi–Si ⁺ –Si [−] SiMe ^t Bu ₂ + [Et ₃ Si(C ₆ H ₅)] ⁺ [WCA] [−]		292
	[B(C ₆ F ₄ R) ₄] [−]	Other	^t Bu ₂ MeSi ₄ SiMe ^t Bu ₂ ^t Bu ₂ MeSi–Si ⁺ –Si [−] SiMe ^t Bu ₂ + [Ph ₃ C] ⁺ [WCA] [−]	R = 4-SiMe ^t Bu	293
	[Zr ₂ Cl ₇ Cp* ₂] [−]	Lewis			294





Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Prot	 + $[\text{H}(\text{OEt})_2]^+ [\text{WCA}]^-$		295
	$[\text{B}(\text{C}_6\text{H}_5)_4]^-$, $[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$, $[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$\text{tBu}_3\text{Si}-\text{Ge}(\text{tBu})_2-\text{Si}(\text{tBu})_3$ + $[\text{Ph}_3\text{C}]^+ [\text{WCA}]^-$	R = 4-Si(Me) ₂ (tBu)	339–341
Ligand-stabilized cations 	$[\text{Sb}_2\text{F}_{11}]^-$	Lewis	 + 2 SbF ₅		273
$[\text{I}_3\text{C-PX}_3]^+$ $[\text{I}_3\text{C-PI}_3]^+$ $[\text{I}_3\text{C-AsI}_3]^+$	$[\text{Al}(\text{OR}^{\text{PF}_6})_4]^-$ $[(\text{R}^{\text{PF}_6})_3\text{Al-F-Al}(\text{OR}^{\text{PF}_6})_3]$ $[\text{Al}(\text{OR}^{\text{PF}_6})_4]^-$	Comp Comp Comp	$[\text{Cl}_3]^+ [\text{WCA}]^- + \text{PX}_3$ $[\text{Cl}_3]^+ [\text{WCA}]^- + \text{PI}_3$ $[\text{Cl}_3]^+ [\text{WCA}]^- + \text{AsI}_3$	X = Cl, Br, I	274 274 274
	$[\text{HCB}_9\text{H}_4\text{Br}_5]^-$	Hyd	$\text{tPr}_3\text{SiH} + [\text{Ph}_3\text{C}]^+ [\text{WCA}]^- + \text{MeCN}$		297
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$\text{tBu}_3\text{Si-Si}^+\text{tBu}_3 + 2[\text{Ph}_3\text{C}]^+ [\text{WCA}]^- + \text{tBuCN}$		298
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com	 + MeCN		299
$[\text{Fe}_3\text{SiPy}]^+$	$[\text{B}(\text{Ar}^{\text{CF}_3})_4]^-$	Lig	$[\text{Fe}_3\text{Si}(\text{THF})]^+ + \text{py}$		300
	$[\text{HCB}_{11}\text{H}_5\text{Br}_6]^-$	Ion	$\text{tBu}_3\text{Si}(\text{WCA}) + \text{H}_2\text{O}$		301
	$[\text{HCB}_{11}\text{Cl}_{11}]^-$	Ion	$\text{tPr}_3\text{Si}(\text{WCA}) + \text{C}_6\text{H}_4\text{Cl}_2$		81

Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[HCB ₁₁ Me ₃ Br ₆] [−]	Ion	Et ₃ Si(WCA) + SO ₂		81
	[OTf] [−]	Ion	Me ₂ Si(OTf) ₂ + bipy		302
	[B(C ₆ F ₅) ₄] [−]	Hyd	Me ₂ ArSiH + [Ph ₃ C] ⁺ [WCA] [−]		303
	[OTf] [−]	Ion	Me ₂ Si(OTf) ₂ + 2DMAP		302
	[B(C ₆ F ₅) ₄] [−]	Hyd	Et ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]	Ar = benzene, toluene, ethylbenzene, <i>n</i> -propylbenzene, and iso-propylbenzene, <i>o</i> -xylene, <i>m</i> -xylene, <i>p</i> -xylene, 1,2,3-trimethylbenzene, 1,2,4-trimethylbenzene, mesitylene	289
[Me ₃ Si(Ar)] ⁺	[B(C ₆ F ₅) ₄] [−]	Lig	[Me ₃ SiHSiMe ₃] ⁺ [WCA] [−] + arene		304
	[B(C ₆ F ₅) ₄] [−]	Hyd			76
[Me ₃ SiHSiMe ₃] ⁺	[HCB ₁₁ HCl ₁₁] [−]	Hyd	2Me ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		81
[Et ₃ SiHSiEt ₃] ⁺	[B(C ₆ F ₅) ₄] [−]	Hyd	2Et ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		306
[Me ₃ SiXSiMe ₃] ⁺	[B(C ₆ F ₅) ₄] [−]	Ion	Me ₃ SiX + Me ₃ Si(WCA)	X = F, Cl, Br, I	307
[Me ₃ Si-CN-SiMe ₃] ⁺	[B(C ₆ F ₅) ₄] [−]	Ion	Me ₃ SiCN + Me ₃ Si(WCA)		309
[Me ₃ Si-OCN-SiMe ₃] ⁺	[B(C ₆ F ₅) ₄] [−]	Ion	Me ₃ SiOCN + Me ₃ Si(WCA)		309
[Me ₃ Si-SCN-SiMe ₃] ⁺	[B(C ₆ F ₅) ₄] [−]	Ion	Me ₃ SiSCN + Me ₃ Si(WCA)		309
[(Me ₃ Si) ₂ NNN] ⁺	[B(C ₆ F ₅) ₄] [−]	Ion	Me ₃ SiNNN + Me ₃ Si(WCA)		309
	[B(C ₆ F ₅) ₄] [−]	Ion	Me ₃ Si(OTf) + Me ₃ Si(WCA)		308





Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Hyd	$\text{Me}_2\text{HSi}-\text{SiHMe}_2$ + $[\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		311
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$\text{Me}_2\text{Si}-\text{H}-\text{SiMe}_2$ + $\text{C}_6\text{H}_5\text{CF}_3$ or $\text{C}_{10}\text{H}_{21}\text{F}$ or $[\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		311
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Hyd	$\text{Ar}_2\text{HSi}-\text{SiHAr}_2$ + $[\text{Ph}_3\text{C}]^+[\text{WCA}]^-$	Ar = Tol	312
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$(\text{Me}_3\text{Si})_3\text{CSiMePhH}$ + $[\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		313
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$\text{Me}_3\text{Si}-\text{SiMe}_3$ + $[(\text{Me}_5\text{C}_6)_3\text{Si}]^+[\text{WCA}]^-$ or $[\text{Et}_3\text{Si}(\text{C}_6\text{H}_5)]^+[\text{WCA}]^-$		314
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$\text{Me}_3\text{Si}-\text{SiMe}_3$ + $[\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		314
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$[(\text{Me})_3\text{Si}]_3\text{C}_2\text{X}$ + H_2O		315
	$[\text{B}_{12}\text{Cl}_{12}]^{2-}$	Hyd	$2\text{FcMe}^+\text{BuSiH}$ + $[\text{Ph}_3\text{C}]^+[\text{WCA}]^-$		316

Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
	[OTf] ⁻	Lewis			317
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Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[OTf] [−]	Ox		[WCA] [−] + 1/8 S ₈	319
	Cl [−]	Ion		+	320
	I [−]	Ion			321
	[OTf] [−]	Ox		+ [Me ₃ Si-DMAP] [−] [WCA] [−] + 2 DMAP	319
		Other		⁺ [WCA] [−] +	319
	I [−]	Ion			321
	[B(C ₆ F ₅) ₄] [−]	Ox		⁺ Bu ₃ Ge-Ge ⁺ Bu ₃ + 2[Ph ₃ C] ⁺ [WCA] [−] + ⁺ BuCN	298



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[OTf] [−]	Ion	Me ₂ Ge(OTf) ₂ + bipy		302
	[B(C ₆ F ₅) ₄] [−]	Hyd	ⁿ Bu ₂ GeH + naphthalene		343
	[AlCl ₄] [−]	Ox	Ph ₃ P=CH-Ge(OTf)(Cl) + [Ph ₃ C] ⁺ [WCA] [−]		344
	[B(C ₆ F ₅) ₄] [−]	Ox or Com	Ph ₃ P=CH-Ge(OTf)(H) + [Ph ₃ C] ⁺ [WCA] [−]		349
	[B(C ₆ F ₅) ₄] [−]	Hyd	Ph ₃ P=CH-Ge(OTf)(H) + [Ph ₃ C] ⁺ [WCA] [−]		349
	I [−]	Ion	Ph ₃ P=CH-Ge(OTf)(H) + 2		354
	[OTf] [−]	Ion, Lig	Ph ₃ P=CH-Ge(OTf)(H) + cryptand[2.2.2]		355
	[GeCl ₃] [−]	Ion	GeCl ₂ dioxane + [12]crown-4		357
	[OTf] [−]	Ion	GeCl ₂ dioxane + [12]crown-4 + 2Me ₃ Si(OTf)		357
	[GeCl ₃] [−]	Ion	2GeCl ₂ dioxane + [15]crown-5		357
	[OTf] [−]	Ion	GeCl ₂ dioxane + [15]crown-5 + 2Me ₃ Si(OTf)		357
	[GeCl ₃] [−]	Ion	2GeCl ₂ dioxane + 1.5[18]crown-6		357



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[GeCl ₃] [−]	Ion	2 + GeCl ₂ -dioxane		360
	[GeCl ₃] [−]	Ion	+ 2 GeCl ₂ + + AlCl ₃		361
	[AlCl ₄] [−]	Lewis	+ 2 GeCl ₂ + + AlCl ₃		344
	[B ₁₂ Cl ₁₂] ^{2−}	Com	+ DMAP + 0.2 K ₂ [WCA] [−]		344
	[HO{B(C ₆ F ₅) ₃] ₂] [−]	Lewis	+ H ₂ O + B(C ₆ F ₅) ₃		369
	[B(C ₆ F ₅) ₄] [−]	Prot	+ [Et ₃ Si(C ₆ H ₅)] ⁺ [WCA] [−]	Protonated by acidified benzene	365
	[B(C ₆ F ₅) ₄] [−]	Prot	+ [H(OEt) ₂] ⁺ [WCA] [−]		365
	[SbF ₆] [−]	Salt	+ Ag ⁺ [WCA] [−] + THF		80



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[SbF ₆] [−]	Lig	[WCA] [−] + py		80
	[SbF ₆] [−]	Lig	THF + PMe ₂ Ph		80
	[Al(OR ^{PF}) ₄] [−]	Salt	Ar-N-Ge ⁺ -Me ₃ Si + Ag ⁺ [WCA] [−] or Li ⁺ [WCA] [−]		366
	[Al(OR ^{PF}) ₄] [−]	Lig			366
	[B(C ₆ F ₅) ₄] [−]	Lewis	^t Bu ₃ Si ⁺ Br + ^t Bu ₃ Si [−] Ge ⁺ Si [−] Ph + [Et ₃ Si(C ₆ H ₅)] ⁺ [WCA] [−]		347
	[MeSO ₂] ₂ N] [−]	Ion	Me ₃ SnN(SO ₂ Me) ₂ + 2OPPh ₃		342
	[B(Ph) ₄] [−]	Ox	^t Bu-P(=O)(ⁱ PrO)-C ₆ H ₄ -Sn ⁺ (DMAP)-C ₆ H ₄ -P(=O)(ⁱ PrO)- ^t Bu + 1/8 S ₈	L = DMAP	345
	[SnCl ₃] [−]	Ion	25SnCl ₂ + 2[15]crown-5		350
	[OTf] [−]	Ion	Sn(OTf) ₂ + 2[12]crown-4		352
	[OTf] [−]	Ion	Sn(OTf) ₂ + 2[15]crown-5		352
	[OTf] [−]	Ion	Sn(OTf) ₂ + [18]crown-6		352
	[PF ₆] [−]	Other	[Sn([18]crown-6)] ²⁺ [WCA] ₂ [−] + KF		353



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[OTf] [−]	Ion	Sn(OTf) ₂ + cryptand[2.2.2] or 2SnCl ₂ + cryptand[2.2.2] + 4Me ₃ Si(OTf)		356
	[SnCl ₃] [−]	Ion	2SnCl ₂ + cryptand[2.2.2]		356
	[SnBr ₃] [−]	Ion	2SnCl ₂ + cryptand[2.2.2] + 4Me ₃ SiBr		356
	[B(C ₆ F ₅) ₄] [−]	Com			346
	[B(C ₆ F ₅) ₄] [−]	Other	Et ₂ Pb[] ₂ + [Ph ₃ C] ⁺ [WCA] [−]		348
	[SnCl ₃] [−]	Salt	+ SnCl ₂ + THF		389
	[GeBr ₃] [−]	Ion	GeBr ₂ + bipy	Bulk product is GeBr ₂ (bipy)	359
	[GeCl ₃] [−]	Ion	GeCl ₂ -dioxane + pmdta		359
	Cl [−]	Ion	GeCl ₂ -dioxane + +		363



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	Cl ⁻	Ox	+ 1/8 S ₈		363
	Cl ⁻	Ion	GeCl ₂ -dioxane 		364
	[GeCl ₃] ⁻	Ion	GeCl ₂ -dioxane + Me ₄ -cyclam		358
	[GeBr ₃] ⁻ + Br ⁻	Ion	GeBr ₂ + Me ₃ -tacn		358
	[SnCl ₃] ⁻	Ion	2 SnCl ₂ +		360
	[SnCl ₃] ⁻	Ion			361
	[SnBr ₃] ⁻	Ion	SnBr ₂ + Ar-N ₂ -Ar 	Ar = 2,5'-Bu ₂ (C ₆ H ₃)	362
	[SnCl ₃] ⁻	Ion	SnCl ₂ + Ar-N ₂ -Ar 	Ar = 2,5'-Bu ₂ (C ₆ H ₃) and 2,6-Me ₂ (C ₆ H ₃)	362



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Salt	$\text{Ar}-\text{N}^+\text{SiMe}_3 + \text{Ag}^+[\text{WCA}]^- \text{ or } \text{Li}^+[\text{WCA}]^-$		366
	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Lig	$\text{SiMe}_3 + \text{DMAP}$		366
	$[\text{MeB}(\text{C}_6\text{F}_5)_3]^-$	Lewis	$\text{Ar}-\text{N}^+\text{Me} + \text{B}(\text{C}_6\text{F}_5)_3$		367
	$[\text{SbF}_6]^-$	Salt	$\text{Ph} + \text{Ag}^+[\text{WCA}]^-$		368
	$[\text{B}(\text{Ph})_4]^-$	Salt	$i\text{-PrO}-\text{P}(=\text{O})-\text{O}-\text{Sn}^+-\text{Cl} + \text{Na}^+[\text{WCA}]^-$		345
	$[\text{B}(\text{Ar}^{\text{C}_6\text{F}_5})_4]^-$	Salt	$i\text{-PrO}-\text{P}(=\text{O})-\text{O}-\text{Sn}^+-\text{Cl} + \text{Na}^+[\text{WCA}]^-$		345
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Other	$(\text{Tipp})_2\text{Sn} + [\text{Et}_3\text{Si}(\text{C}_7\text{H}_8)]^+[\text{WCA}]^-$		371



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	$[\text{Pb}(\text{NO}_3)_3][12\text{crown-4}]^+$	Ion	$\text{Pb}(\text{NO}_3)_2 + [12\text{crown-4}]$		351
	$[\text{Pb}(\text{NO}_3)_3][15\text{crown-5}]^+$	Ion	$\text{Pb}(\text{NO}_3)_2 + 2[15\text{crown-5}]$		351
	$[\text{Pb}(\text{NO}_3)_3(\text{benzo-}[15\text{crown-5}])]^+$	Ion	$\text{Pb}(\text{NO}_3)_2 + \text{benzo-}[15\text{crown-5}]$		351
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Salt	$\text{Ar-N}(\text{PbCl})\text{Ar} + \text{Li}^+[\text{WCA}]^-$		367
	$[\text{B}(\text{C}_6\text{F}_5)_3(\text{CH}_3)]^-$	Lewis	$\text{Tipp-Pb-Tipp} + \text{B}(\text{C}_6\text{F}_5)_3$		370
Cyclopentadienyl substituted cations					
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Prot	$(\text{Me}_5\text{C}_5)_2\text{Si} + [\text{Me}_5\text{C}_5\text{H}_2]^+[\text{WCA}]^-$		322
	$[\text{Al}(\text{OR}^{\text{Pr}})_4]^-$	Prot	$(\text{Me}_5\text{C}_5)(\text{C}^{\text{Pr}}\text{Pt}_5\text{C}_3)\text{Si} + [\text{H}(\text{OEt}_2)_2]^+[\text{WCA}]^-$		323
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com	$[(\text{Me}_5\text{C}_5)\text{Si}]^+[\text{WCA}]^- + \text{dme}$		324
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Com	$[(\text{Me}_5\text{C}_5)\text{Si}]^+[\text{WCA}]^- + [12\text{crown-4}]$		324
	$[\text{BF}_4]^-$	Prot	$(\text{C}_5\text{Me}_5)_2\text{Ge} + \text{H}(\text{WCA})$		376
	$[\text{SnCl}_3]^-$	Lewis	$(\text{C}_5\text{Me}_5)\text{GeCl} + \text{SnCl}_2$		377
	$[\text{BF}_4]^-$	Prot	$(\text{C}_5\text{Me}_5)_2\text{Ge} + \text{H}(\text{WCA})$		373



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	[B(C ₆ F ₅) ₄] [−]	Salt	(C ₅ Me ₅)SnCl + Li ⁺ [WCA] [−]		375
	[B(C ₆ F ₅) ₄] [−]	Salt	(C ₅ Me ₅)PbCl + Li ⁺ [WCA] [−]		375
	[Ga(C ₆ F ₅) ₄] [−]	Other	(C ₅ Me ₅) ₂ Sn + Ga(C ₆ F ₅) ₃		378
	[B(C ₆ F ₅) ₄] [−]	Com	[(C ₅ Me ₅)Sn] ⁺ [WCA] [−] + (C ₅ Me ₅) ₂ Sn		375
	[B(C ₆ F ₅) ₄] [−]	Com	[(C ₅ Me ₅)Pb] ⁺ [WCA] [−] + (C ₅ Me ₅) ₂ Pb		375
Ion-like compounds					
Me ₂ (B ₁₂ Cl ₁₂)		Salt	[Li] ₂ ⁺ [WCA] [−] + 2.2MeF + 2.6AsF ₅	1 : 1 mixture with [Li] ₂ ⁺ [WCA] [−]	288
(Me ₂ CH)(HCB ₁₁ Me ₅ Br ₆)		Lewis	(H ₃ C) ₂ CHCl + (H ₃ C)(HCB ₁₁ H ₃ Br ₆)		287
Me ₂ CF(AsF ₆)		Lewis	(H ₃ C) ₂ CF ₂ + AsF ₅		268
(<i>m</i> -CF ₃ -C ₆ H ₄)(Ph)CF(AsF ₆)		Lewis	C ₆ H ₄ CF ₃ + AsF ₅		268
Me ₃ Si(HCB ₁₁ F ₁₁)		Hyd	Me ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		326
Me ₃ Si(C ₂ H ₅ CB ₁₁ F ₁₁)		Hyd	Me ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		326
Me ₃ Si(FAI(OR ^{PF}) ₃)		Other	Ag ⁺ [Al(OR ^{PF}) ₄] [−] + Me ₃ SiCl or AlEt ₃ + 3HOR ^{PF} + Me ₃ SiF		327
Et ₃ Si(HCB ₁₁ H ₃ Br ₆)		Hyd	Et ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		328
Et ₃ Si(HCB ₁₁ Cl ₁₁)		Hyd	Et ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		81
ⁱ Pr ₃ Si(HCB ₁₁ H ₃ Br ₆)		Hyd	ⁱ Pr ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		325
ⁱ Pr ₃ Si(HCB ₁₁ H ₅ Cl ₆)		Hyd	ⁱ Pr ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		329
ⁱ Pr ₃ Si(HCB ₁₁ H ₅ I ₆)		Hyd	ⁱ Pr ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		329
ⁱ Pr ₃ Si(HCB ₉ H ₄ Br ₅)		Hyd	ⁱ Pr ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		297
^t Bu ₃ Si(HCB ₁₁ H ₃ Br ₆)		Hyd	^t Bu ₃ SiH + [Ph ₃ C] ⁺ [WCA] [−]		328
^t Bu ₃ Si(FAI(OR ^{PF}) ₃)		Other	^t Bu ₃ SiH + 15 (^t Bu) ₃ SiF + Ag ⁺ [Al(OR ^{PF}) ₄] [−]		330
^t Bu ₂ MeSi(HCB ₁₁ H ₅ Br ₆)		Hyd	^t Bu ₂ MeSiH + [Ph ₃ C] ⁺ [WCA] [−]		328
Fc ₃ Si(OTf)		Prot	+ HOTf		331
Et ₃ Ge(HCB ₁₁ H ₃ Br ₆)		Hyd	Et ₃ GeH + [Ph ₃ C] ⁺ [WCA] [−]		338
Et ₃ Sn(HCB ₁₁ H ₅ Br ₆)		Lewis	Et ₃ SnCl + (Et) ₃ Si(WCA)		338
ⁿ Bu ₃ Sn(CB ₁₁ Me ₁₂)		Ox	ⁿ Bu ₃ Sn ₂ + 2 CB ₁₁ Me ₁₂		333
Et ₃ Pb(HCB ₁₁ H ₅ Br ₆)		Lewis	Et ₃ PbCl + (Et) ₃ Si(WCA)		338



Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
Transition-metal substituted cations					
	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Salt	$\text{Cl}-\text{M}=\text{Ge}-\text{Cp}^* + \text{Li}^+[\text{WCA}]^- + \text{MeCN}$ $\text{P} = \text{dppe}$		380
	$[\text{PF}_6]^-$	Salt	$\text{Sn}(\text{Mes})_2 + \text{P}(\text{dppe})_2 + \text{Cl}^-$ $\text{P} = \text{dppe}$		379
	$[\text{BF}_4]^-$	Com	$\text{Ph}_3\text{P}-\text{Pt}(\text{Ph})_2-\text{P}(\text{Ph})_3 + \text{SnCl}_2 \text{ or } [\text{PtMeCl}(\text{cod})] + 2 \text{ 2-PyPPh}_2 + \text{SnCl}_2 + [\text{NBu}_4]^+[\text{BF}_4]^-$ $[\text{WCA}]^-$		381
		Salt	$\text{Sn}(\text{OMe})_2 + \text{Ag}^+[\text{WCA}]^- + \text{H}_2\text{O}$		382
	$[\text{CB}_{11}\text{H}_{12}]^-$	Salt	$\text{Sn}(\text{OMe})_2 + \text{Ag}^+[\text{WCA}]^- + \text{H}_2\text{O}$	L = THF	382
	$[\text{ClO}_4]^-$	Ion	$\text{Sn}(\text{OMe})_2 + \text{Ag}^+[\text{WCA}]^- + \text{H}_2\text{O}$		383

Table 9 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
	$[\text{ClO}_4]^-$	Ion		+ DMAP	383
	$[\text{W}(\text{CO})_5\text{Cp}]^-$	Prot		+ $\text{HW}(\text{CO})_5(\text{Cp})$	384
	$[\text{AlBr}_4]^-$	Lewis		+ AlBr_3	385
	$[\text{B}(\text{Ar}^{\text{Cl}})_4]^-$	Salt		+ $\text{Na}^+[\text{WCA}]^-$	385
$\{(\text{Cy}_3\text{P})_2\text{Pt}\}_2\text{Sn}(\text{AlBr}_4)_2$		Lewis		+ AlBr_3	385
	$[\text{AlCl}_4]^-$	Lewis		+ AlCl_3	385
	$[\text{B}(\text{Ar}^{\text{Cl}})_4]^-$	Salt		+ $\text{Na}^+[\text{WCA}]^-$	385
	$[\text{AlCl}_4]^-$	Salt		+ 2	385
$\{(\text{Cy}_3\text{P})_2\text{Pt}\}_2\text{Pb}(\text{AlCl}_4)_2$		Lewis		+ AlCl_3	385



Table 10 Overview on selected and structurally characterized pnictogen cations

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
Homopolyatomic cations					
[N ₅] ⁺	[Sb ₂ F ₁₁] [−]	Other	[N ₂ F] ⁺ [SbF ₆] [−] + HN ₃ in aHF	2.5 equiv. of P ₄ , no X-ray	401
[P ₉] ⁺	[Al(OR ^{PF₆}) ₄] [−]	Ox	P ₄ + [NO] ⁺ [Al(OC(CF ₃) ₃) ₄] [−]		29
[Sb ₈] ²⁺	[GaCl ₄] [−]	Lewis	SbCl ₃ + Ga ⁺ [GaCl ₄] [−] in GaCl ₃ /C ₆ H ₆		402
[Bi ₂] ²⁺	[Ag ₃ Bi ₃ Br ₁₅] ^{3−} · 6[Br] [−]	Lewis, Ox	Bi + BiBr ₃ + Ag	HTS (350 °C)	429
2[Bi ₅] ²⁺ · [Bi ₆] ²⁺	2[<i>trans</i> -Bi ₂ Br ₁₂] ^{2−} · [<i>trans</i> -Bi ₆ Br ₁₃] ^{2−}	Lewis, Ox	Bi + Ir + BiBr ₃	HTS (1000 °C)	430
[Bi ₅] ³⁺	[AlCl ₄] [−]	Lewis, Ox	Bi + BiCl ₃ + AlCl ₃	HTS	431
[Bi ₅] ³⁺	[AlCl ₄] [−]	Lewis, Ox	Bi + BiCl ₃ + [BMIM][Cl]/AlCl ₃	Ionic liquid based synthesis	403
[Bi ₅] ³⁺ · 2SO ₂	[AlX ₄] [−] (X = Br, I)	Lewis, Ox	Bi + BiX ₃ + AlX ₃	HTS (490 °C (I), 520 °C (Br))	432
[Bi ₈] ²⁺	[AsF ₆] [−]	Lewis, Ox	Bi + AsF ₅ in SO ₂	No X-ray	196
[Bi ₈] ²⁺	[AlCl ₄] [−]	Lewis, Ox	Bi + BiCl ₃ + AlCl ₃	HTS	433
[Bi ₉] ⁵⁺	[Ta ₂ O ₉ Br ₇] [−]	Lewis, Ox	Bi + BiBr ₃ + TaBr ₅	HTS (570 °C), traces of H ₂ O	434
	4[BiCl ₅] ^{2−} · [Bi ₂ Cl ₈] ^{2−}	Lewis, Ox	Bi + BiCl ₃	HTS (325 °C), Bi ₆ Cl ₇	435 and 236
[Bi ₉] ⁵⁺	[Bi] ³⁺ · 3[HFCl ₆] ^{2−}	Lewis, Ox	Bi + BiCl ₃ + HFCl ₄	HTS	436
[Bi ₉] ⁵⁺	[Bi] ³⁺ · 3[NbCl ₆] ^{2−}	Lewis, Ox	Bi + BiCl ₃ + NbCl ₅	HTS (550 °C), Nb(v) to Nb(iv) reduction	434
[Bi ₉] ⁵⁺	[Sn ₇ Br ₂₄] ^{10−}	Lewis, Ox	Bi + BiBr ₃ + Sn	HTS (250 °C), C _{4v} symmetric	437
Metal–nonmetal–cluster complexes					
[Cu(P ₄) ₂] ⁺	[Al(OR ^{PF₆}) ₄] [−]	Com	CuI + [Ag] ⁺ [Al(OR ^{PF₆}) ₄] [−] + P ₄		438
[Cu(P ₄) ₂] ⁺	[GaCl ₄] [−]	Com	CuCl + GaCl ₃ + P ₄		439
[Ag(P ₄) ₂] ⁺	[Al(OR ^{PF₆}) ₄] [−]	Com	Ag ⁺ [Al(OR ^{PF₆}) ₄] [−] + P ₄		72
[Ag(P ₄) ₂] ⁺	[GaCl ₄] [−]	Com	AgCl + GaCl ₃ + P ₄		439
[Au(P ₄) ₂] ⁺	[GaCl ₄] [−]	Com	AuCl + GaCl ₃ + P ₄		439
[Cp ⁺ M(dppe)(P ₄)] ⁺ (M = Fe, Ru)	[BPh ₄] [−]	Salt	[Cp ⁺ M(dppe)Cl] + P ₄ + Na ⁺ [BPh ₄] [−]		440
[CpOs(PPh ₃) ₂ (P ₄)] ⁺	[OTf] [−]	Salt	[CpOs(PPh ₃) ₂ Cl] + Ag ⁺ [OTf] [−]		441
[{CpRu(PPh ₃) ₂ (P ₄)] ²⁺	[OTf] [−]	Com, salt	[CpRu(PPh ₃) ₂ Cl] + P ₄ + Ag ⁺ [OTf] [−]		442
[{CpRu(PPh ₃) ₂ }] ²⁺	[OTf] [−]	Com	[CpOs(PPh ₃) ₂ (P ₄)] ⁺ + [{CpRu(PPh ₃) ₂ }] ⁺	Bridging end-on/end-on	441
[CpOs(PPh ₃) ₂ (P ₄)] ²⁺	[Al(OR ^{PF₆}) ₄] [−]	Com	Ag ⁺ [Al(OR ^{PF₆}) ₄] [−] + As ₄		405
[AuPPh ₃ (As ₄)] ⁺	[Al(OR ^{PF₆}) ₄] [−]	Com, salt	[Ag(As ₄) ₂] ⁺ [Al(OR ^{PF₆}) ₄] [−] + AuPPh ₃ Cl		405
[Cp ⁺ Ru(dppe)(As ₄)] ⁺	[Al(OR ^{PF₆}) ₄] [−]	Com, salt	[Ag(As ₄) ₂] ⁺ [Al(OR ^{PF₆}) ₄] [−] + Cp ⁺ Ru(dppe)Cl		443
[Ag(P ₄ S ₃) ₆] ⁺ (n = 1, 2)	[Al(OR ^{PF₆}) ₄] [−]	Com	Ag ⁺ [Al(OR ^{PF₆}) ₄] [−] + P ₄ S ₃		444
[Ag ₂ (P ₄ S ₃) ₆] ⁺	[Al(OR ^{PF₆}) ₄] [−]	Com	Ag ⁺ [Al(OR ^{PF₆}) ₄] [−] + P ₄ S ₃		170
[{CpRu(PPh ₃) ₂ }(P ₄ S ₃)] ²⁺	[OTf] [−]	Com, salt	[CpRu(PPh ₃) ₂ Cl] + P ₄ S ₃ + Ag ⁺ [OTf] [−]		442
Clusters, cluster-like and catenated cations					
[N ₂ Ph] ⁺	[BF ₄] [−]	Other	PhNH ₂ + NaNO ₂ in HCl _(aq) and Na ⁺ [BF ₄] [−]	N–N triple bond	445
	[BF ₄] [−]	Other	H ₂ NC ₆ (CF ₃) ₅ + [NO] ⁺ [BF ₄] [−]		446
Mes – N ⁺ ≡ N	[OsO ₂ (NO ₃) ₂ (Mes)] [−]	Other	[N ₂ Mes] ⁺ [NO ₃] [−] + [OsO ₂ (NO ₃)(Mes)] ⁺		447
	[GaCl ₄] [−]	Other	Hg(N ₂ (TMS) ₃) ₂ + Ag ⁺ [GaCl ₄] [−] or Bi(N ₂ (TMS) ₃) ₂ + GaCl ₃ + Cl ₂	LTS (−80 °C)	448

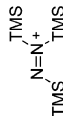
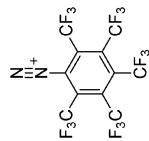




Table 10 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
Mes* – N ⁺ ≡P	[AlCl ₄] [–]	Lewis	Mes*·NPCl + AlCl ₃	N–P triple bond	407
	[GaCl ₄] [–]	Other	[P(N ⁱ Pr ₂)N ₃] ⁺ [GaCl ₄] [–] + PCy ₃		449
	[GaCl ₄] [–]	Other	[P(N ⁱ Pr ₂)N ₃] ⁺ [GaCl ₄] [–] + P(NR ₂)Cl ₂	N ₂ as leaving group	449
	[SbF ₆] [–]	Salt	[ⁱ BuIM]NPCl(C(PPh ₃) ₂) ⁺ [Cl] [–] + Ag ⁺ [SbF ₆] [–]	Dicationic iminophosphorane	450
[P ₂ Ph ₅] ⁺	[OTf] [–]	Lewis	Ph ₂ PCl + TMSOTf + PPh ₃	First homoleptic phosphine-phosphonium	451
[P ₂ Me ₆] ²⁺	[OTf] [–]	Lewis	P ₂ Me ₄ + MeOTf		452
[P ₂ Br ₅] ⁺	[Al(OR ^{PF}) ₄] [–]	Salt	PBr ₃ + Ag ⁺ [Al(OR ^{PF}) ₄] [–]		453
[P ₂ I ₅] ⁺	AlI ₄ [–]	Lewis	PI ₃ + AlI ₃		454
[P ₂ I ₅] ⁺	[Al(OR ^{PF}) ₄] [–]	Salt	PI ₃ + Ag ⁺ [Al(OR ^{PF}) ₄] [–]		453
	[AlCl ₄] [–]	Lewis	PPh ₃ + PCl ₃ + AlCl ₃	[P(PPh ₃) ₂] ⁺	421
	[OTf] [–]	Lewis	P ₂ Me ₄ + PMe ₂ Cl + TMSOTf	[(PMe ₂) ₃] ⁺	455
	[AlCl ₄] [–]	Prot	[P(PPh ₃) ₂] ⁺ [AlCl ₄] [–] + AlCl ₃ + HCl		456
[P ₃ I ₆] ⁺	[F(Al(OR ^{PF}) ₃) ₂] [–]	Salt	P ₄ + I ₂ + Ag ⁺ [Al(OR ^{PF}) ₄] [–]	[(P ₃) ₃] ⁺	457
[PMe(AsMe ₃) ₂] ⁺	[OTf] [–]	Other	MePCl ₂ + AsMe ₃ + TMSOTf	[PRL ₂] ⁺	458
[P ₄ Ph ₈] ²⁺	[OTf] [–]	Lewis	PPhCl ₂ + PPh ₃ + TMSOTf		459

Table 10 (continued)

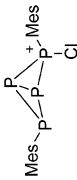
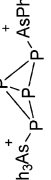
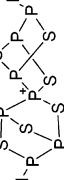
Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
$[P_4NO]^+$	$[Al(OR^{PF_6})_4]^-$	Other	$P_4 + [NO]^+[Al(OR^{PF_6})_4]^-$	Insertion in P_4 , no X-ray	78
	$[GaCl_4]^-$	Lewis	$ClP(PMes^*)_2PCl + GaCl_3$	Bicyclic phosphine-phosphonium	460
$[(pPh)_2(PnPPh_3)_2]^{2+}$ (Pn = As, Sb) $[P_5Br_2]^+$	$[AlCl_4]^-$ $[Al(OR^{PF_6})_4]^-$	Lewis Lewis, other	$PPhCl_2 + PnPPh_3 + AlCl_3$ $P_4 + PBr_3 + Ag^+[Al(OR^{PF_6})_4]^-$	Insertion in P_4	461 453 and 462
$[P_5Ph_2]^+$, $[P_7Ph_6]^{3+}$	$[GaCl_4]^-$	Lewis, other	$P_4 + Ph_2PCl + GaCl_3$	Different stoichiometries, insertion in P_4 , HTS (60–70 °C)	68
$[P_5RCl]^+$ (R = Me, Et, ⁱ Pr, Cy, Ph, C ₆ F ₅)	$[GaCl_4]^-$	Lewis, other	$P_4 + R_2PCl + GaCl_3$	Insertion in P_4	463
$[P_5R_2]^+$ (R = Me, Et, ⁱ Pr, Cy, Mes, Dipp)	$[GaCl_4]^-$	Lewis, other	$P_4 + R_2PCl + GaCl_3$	Insertion in P_4	464
$[P_5(NCy_2)Cl]^+$	$[GaCl_4]^-$	Lewis, other	$P_4 + P(NCy_2)Cl_2 + GaCl_3$	Insertion in P_4	465
 X = Br, I	$[Al(OR^{PF_6})_4]^-$	Salt, other	$P_4S_3 + PX_3 + Ag^+[Al(OR^{PF_6})_4]^-$	Initial insertion in P_4S_3	134
	$[AlCl_4]^-$	Other	$[P_4(AsPh_3)_2]^{2+}([AlCl_4]^-)_2 + PPh_3$	Ligand exchange	466
	$[AlCl_4]^-$	Other	$AsPh_3 + PCl_3 + AlCl_3$		466
	$[Al(OR^{PF_6})_4]^-$	Salt	$P_4S_3I_2 + Ag^+[Al(OR^{PF_6})_4]^-$		234
	$[OTf]^-$	Other	$(AsPh_3)(OTf)_2 + PCl_3 + Ph_3As$		467
	$[AlCl_4]^-$	Lewis, other	$P_{(red)} + Se + SeCl_4 + BMimCl/AlCl_3$	Ionic liquid based synthesis, rhombohedral and orthorhombic modification	412



Table 10 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
[P ₃ Se ₄] ⁺	[AlCl ₄] ⁻	Lewis, other	P ₃ Se ₄ + Me ₅ C ₆ Br + AlCl ₃	In solution, orthorhombic modification	412
[P ₃ Se ₄] ⁺	[Ga ₂ Cl ₇] ⁻	Lewis, other	PCl ₃ + SeTMS ₂ + GaCl ₃	In solution	412
[P ₃ Se ₂ Ph ₂] ⁺	[GaCl ₄] ⁻	Ox	[P ₃ Ph ₂] ⁺ [GaCl ₄] ⁻ + Se	HTS (160 °C)	464
[P ₅ Se ₂ Cy ₂] ⁺	[GaCl ₄] ⁻	Ox	[P ₅ Cy ₂] ⁺ + GaCl ₃ + Se	HTS (140 °C)	464
Mes ⁺ -N≡As	[GaCl ₄] ⁻	Lewis	Mes ⁺ NaCl + GaCl ₃	N-As triple bond	408
[As ₃ N ₃ Ph ₃ Cl ₂] ⁺	[GaCl ₄] ⁻	Lewis	N ₂ As ₂ Ph ₂ Cl ₂ + GaCl ₃	As ₃ N ₃ ring	468
[As ₂ Me ₆] ²⁺	[OTf] ⁻	Other	Me ₃ As + PCl ₃ + TMSOTf		458
[As ₂ Ph ₆] ²⁺	[AlCl ₄] ⁻	Ox	AsPh ₃ + PCl ₃ + AlCl ₃		466
		Ox	As ₄ S ₄ + AsF ₅ in SO ₂		469
[As ₃ S ₄] ⁺	[SbF ₆] ⁻	Ox	As ₄ S ₄ + SbF ₅ in SO ₂		469
		Ox, Lewis	As + AsCl ₃ + S + AlCl ₃	HTS (80 °C)	470
	[SbF ₆] ⁻	Ox	As + Se + SbF ₅ in SO ₂		469
[As ₃ Se ₄] ⁺	[AlCl ₄] ⁻	Ox, Lewis	As + AsCl ₃ + Se + AlCl ₃	HTS (80 °C)	470
[Sb ₂ Se ₂] ⁺	[AlCl ₄] ⁻	Ox	Sb + Se + BMImCl/AlCl ₃	Ionothermal, HTS (160 °C)	67
		Ox, others	Sb + Se + SeCl ₄ + BMImCl/AlCl ₃	Ionic liquid based synthesis	409
	[AlCl ₄] ⁻	Ox, Lewis	Sb + Te + SbCl ₃ + NaCl + AlCl ₃	HTS (130 °C), polymeric	15
[Sb ₂ Te ₆] ⁺	3[Ga ₂ Cl ₇] ⁻ · 2[GaCl ₄] ⁻ / 2[Ga ₂ Cl ₇] ⁻ · 3[GaCl ₄] ⁻	Ox, Lewis	Sb + Te + SbCl ₃ + GaCl ₃	AlCl ₃ /GaCl ₃ melt	410
		Ox, Lewis	Sb + Te + SbCl ₃ + MCl ₃ + NaCl	AlCl ₃ /GaCl ₃ melt	410



Table 10 (continued)

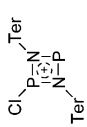
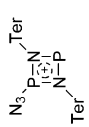
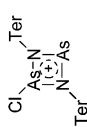
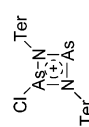
Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
$[\text{Bi}_2\text{X}_4]^{2+}$ (X = Cl, Br) $[\text{Bi}_4\text{OF}_2\text{Cl}_6(\text{C}_6\text{Me}_6)_4]^{2+}$	$[\text{AlX}_4]^-$ $[\text{Al}(\text{OR}^{\text{PR}})_4]^-$	Lewis Other	$\text{BiX}_3 + \text{AlX}_3$ $[\text{Bi}_5]^{+}[\text{AsF}_6]^- + \text{Li}^{+}[\text{Al}(\text{OR}^{\text{PR}})_4]^- + \text{C}_6\text{Me}_6$	Evidence for soluble $[\text{Bi}_n]^{x+}$ salts, partial decomposition of anion and solvent	471 404
 + Ch—Bi Ch = S, Se	$[\text{AlCl}_4]^-$	Ox, Lewis	Bi + BiCl ₃ + Ch in AlCl ₃ /NaCl	HTS (130 °C), heterocubane, series Bi ₄ Ch ₄ ⁴⁺ (Ch = S, Se, Te) complete	413
 + Te—Bi	$[\text{AlCl}_4]^-$	Ox, Lewis	Bi + BiCl ₃ + Te in AlCl ₃ /NaCl	HTS (130 °C), heterocubane	414
$[\text{PdBi}_{10}]^{4+}$ (4n + 2)π-cations	$([\text{BiBr}_4]^-)_{\infty}$	Ox	Bi ₂ Pd + Bi + Br ₂	HTS (1000 °C) Pd@Bi ₁₀ ⁴⁺	472
	$[\text{GaCl}_4]^-$	Lewis	ClP(μ-NTer) ₂ Pd + GaCl ₃	P ₂ N ₂ ring	473
	$[\text{N}_3(\text{GaCl}_3)_2]^-$	Other	$[\text{ClP}(\mu\text{-NTer})_2\text{P}]^{+}[\text{GaCl}_4]^- + \text{TMSN}_3 + \text{GaCl}_3$	P ₂ N ₂ ring	473
$^{*}\text{Mes-N}^{+}(\text{CPh}_3)\text{N-N}$	$[\text{GaCl}_4]^-$	Other	$[\text{Mes}^{*}\text{-NAs}]^{+}[\text{GaCl}_4]^- + \text{Ph}_3\text{CN}_3$	Cycloaddition	408
	$[\text{GaCl}_4]^-$	Lewis	ClAs(μ-NTer) ₂ AsCl + GaCl ₃	As ₂ N ₂ ring	474
	$[\text{OTf}]^-$	Salt	ClAs(μ-NTer) ₂ AsCl + Ag ⁺ [OTf] ⁻	As ₂ N ₂ ring	474



Table 10 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
	$[N_3(GaCl_3)_2]^-$	Lewis	$[ClAs(\mu-NTer)_2As]^+[GaCl_4]^- + TMSN_3 + GaCl_3$	As ₂ N ₂ ring	474
$[As(\mu-NTer)_2As]^{2+}$	$[OTf]^-$	Salt	$ClAs(\mu-NTer)_2AsCl + Ag^+[OTf]^-$	As ₂ N ₂ ring, 4π system, 2 equiv. of Ag ⁺ [OTf] [−]	474
$[ClSb(\mu-NTer)_2Sb]^+$	$[GaCl_4]^-$	Lewis	$ClSb(\mu-NTer)_2SbCl + GaCl_3$	Sb ₂ N ₂ ring	475
$[Sb(\mu-NTer)_2Sb]^{2+}$	$[OTf]^-$	Salt	$ClSb(\mu-NTer)_2SbCl + Ag^+[OTf]^-$	Sb ₂ N ₂ ring, 4π system, 2 equiv. of Ag ⁺ [OTf] [−]	475
$[Bi(\mu-NTer)_2Bi]^+$	$[B(C_6F_5)_4]^-$	Salt	$IBi(\mu-NTer)_2BiH + [Ag(Tol)_3]^+[B(C_6F_5)_4]^-$	Bi ₂ N ₂ ring	475
$[Bi(\mu-NTer)_2Bi]^{2+}$	$[OTf]^-$	Salt	$ClBi(\mu-NTer)_2BiCl + Ag^+[OTf]^-$	Bi ₂ N ₂ ring, 4π system, 2 equiv. of Ag ⁺ [OTf] [−]	475
$\pi^*-\pi^*$ -complexes					
	$[GaCl_4]^-$	Lewis	$AsS_2(CH)_2Cl + GaCl_3$		415
	$[MCl_4]^-$ (M = Al, Ga)	Lewis	$As(NMe)_2(CH)_2Cl + MCl_3$		415
Radicals					
	$[B(C_6F_5)_4]^-$	Ox	$[P(CN(Dipp)C_{10}H_{18})N(C(N(Dipp))(C_2H_5))] + [CPb_3]^+[B(C_6F_5)_4]^-$	CN(Dipp)C ₁₀ H ₁₈ =CAAC, cyclic alkylaminocarbene	416
	$[SbF_6]^-$	Ox	$[(NTMS)_2(PNTMS)_2]_2 + [NO]^+[SbF_6]^-$		420

Table 10 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
$\begin{array}{c} \text{Tiipp} \\ \\ \text{P}^{\text{++}} \\ \\ \text{Tiipp} \end{array}$	$[\text{X}]^- = [\text{SbF}_6]^- , [\text{Al}(\text{OR}^{\text{PR}})_4]^-$	Ox	$\text{P}^{\text{Tiipp}}_3 + \text{Ag}^+[\text{X}]^-$		418
$\left[\begin{array}{c} \text{Tiipp} \quad \text{Tiipp} \\ \quad \\ \text{P}^{\text{++}} - \text{P}^{\text{++}} \\ \quad \\ \text{Tiipp} \quad \text{Tiipp} \end{array} \right]^{\text{++}}$	$[\text{Al}(\text{OR}^{\text{HIT}})_4]^-$	Ox	$\text{P}_2^{\text{Tiipp}}_4 + \text{Ag}^+[\text{Al}(\text{OR}^{\text{HIT}})_4]^-$		419
$\left[\begin{array}{c} \text{P} - \text{N}^{\text{Ter}} \\ \quad \\ \text{P} - \text{N} \\ \quad \\ \text{Ter} \end{array} \right]^{\text{++}}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Ox	$[\text{P}_2(\text{NTer})_2] + [\text{Ag}(\text{Tot})_3]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$		476
$\left[\begin{array}{c} \text{Dipp} \quad \text{Dipp} \\ \quad \\ \text{P} = \text{P} \\ \quad \\ \text{Dipp} \quad \text{Dipp} \end{array} \right]^{\text{++}}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Ox	$[\text{P}_2(\text{C}(\text{N}(\text{Dipp}))_2\text{C}_2\text{H}_2)_2] + [\text{CPh}_3]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$	$\text{C}(\text{N}(\text{Dipp}))_2\text{C}_2\text{H}_2=\text{NHC}$, N-heterocyclic carbene	208
$\left[\begin{array}{c} \text{Dipp} \quad \text{Dipp} \\ \quad \\ \text{P} = \text{P} \\ \quad \\ \text{Dipp} \quad \text{Dipp} \end{array} \right]^{\text{++}}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Ox	$[\text{P}_2(\text{CN}(\text{Dipp})\text{C}_{10}\text{H}_{18})_2] + [\text{CPh}_3]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$	$\text{NC}_{10}\text{H}_{18}$ = tetramethylamide	477
$\left[\begin{array}{c} \text{Dipp} \quad \text{Dipp} \\ \quad \\ \text{P} = \text{P} \\ \quad \\ \text{Dipp} \quad \text{Dipp} \end{array} \right]^{\text{++}}$	$[\text{Al}(\text{OR}^{\text{PR}})_4]^-$	Ox	$[\text{P}_4(\text{N}^{\text{PR}})_4] + [\text{NO}]^+[\text{BF}_4]^- + \text{Li}^+[\text{Al}(\text{OR}^{\text{PR}})_4]^-$		420
$\left[\begin{array}{c} \text{P} - \text{N}^{\text{Ter}} \\ \quad \\ \text{P} - \text{N} \\ \quad \\ \text{Ter} \end{array} \right]^{\text{++}}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Ox	$[\text{AsP}(\text{NTer})_2] + [\text{Ag}(\text{Tot})_3]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$		476
$\left[\begin{array}{c} \text{Dipp} \quad \text{Dipp} \\ \quad \\ \text{As} = \text{As} \\ \quad \\ \text{Dipp} \quad \text{Dipp} \end{array} \right]^{\text{++}}$	$[\text{GaCl}_4]^-$	Ox	$[\text{As}_2(\text{C}(\text{N}(\text{Dipp}))_2\text{C}_2\text{H}_2)_2] + \text{GaCl}_3$		417





Table 10 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
$\left[\begin{array}{c} \text{As-N}^{\text{Ter}} \\ \text{N-As} \\ \text{Ter} \end{array} \right]^{++}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Ox	$[\text{As}_2(\text{N}^{\text{Ter}})_2] + [\text{Ag}(\text{ToI})_3] [\text{B}(\text{C}_6\text{F}_5)_4]^-$		476
Oxidation state +I 	$[\text{SbCl}_5 \cdot \text{THF}]^-$	Lewis	$\text{AsCl}_3 + \text{SnCl}_2 + \text{DppDIMPY}$	$[\text{AsL}_3]^+, \text{DppDIMPY} = [\alpha, \alpha' - \{2, 6 - \text{Pr}_2\text{PhN} - \text{C}(\text{Me})_2\}_2(\text{C}_3\text{H}_3\text{N})]$	423
Oxidation state +III $[\text{P}(\text{N}^{\text{Pr}})_2]^+$	$[\text{GaCl}_4]^-$	Lewis	$\text{P}(\text{N}^{\text{Pr}})_2\text{Cl} + \text{GaCl}_3$	$[\text{PX}_2]^+$	478 and 479
$[\text{P}(\text{NCy}_2)\text{Cl}]^+$	$[\text{GaCl}_4]^-$	Lewis	$\text{P}(\text{NCy}_2)\text{Cl}_2 + \text{GaCl}_3$	$[\text{PX}_2]^+$	465
$[\text{P}(\text{N}^{\text{Pr}})_2\text{Cl}]^+$	$[\text{GaCl}_4]^-$	Lewis	$\text{P}(\text{N}^{\text{Pr}})_2\text{Cl}_2 + \text{GaCl}_3$	$[\text{PX}_2]^+$	449
$[\text{P}(\text{NTMS}_2)\text{Cl}]^+$	$[\text{GaCl}_4]^-$	Lewis	$\text{P}(\text{NTMS}_2)\text{Cl}_2 + \text{GaCl}_3$	$[\text{PX}_2]^+$	480
$[\text{P}(\text{N}^{\text{Pr}})_2\text{N}_3]^+$	$[\text{GaCl}_4]^-$	Other	$[\text{P}(\text{N}^{\text{Pr}})_2\text{Cl}]^+ [\text{GaCl}_4]^- + \text{TMSN}_3$	$[\text{PX}_2]^+$	449
$[\text{P}(\text{NTMS}_2)\text{X}]^+ (\text{X} = \text{N}_3, \text{NCO}, \text{NCS})$	$[\text{GaCl}_4]^-$	Other	$[\text{P}(\text{NTMS}_2)\text{Cl}]^+ [\text{GaCl}_4]^- + \text{TMSX}$	$[\text{PX}_2]^+$	480 and 481
$[\text{P}(\text{NTMS}_2)\text{OTMS}]^+$	$[\text{GaCl}_4]^-$	Other	$[\text{P}(\text{NTMS}_2)\text{Cl}]^+ [\text{GaCl}_4]^- + \text{TMSNO}$	$[\text{PX}_2]^+$	481
$[\text{PCp}^*\text{Cl}]^+$	$[\text{Cl}(\text{Al}(\text{OR}^{\text{Pr}})_3)_2]^-$	Lewis	$\text{PCp}^*\text{Cl}_2 + \text{PhF} \cdot \text{Al}(\text{OR}^{\text{Pr}})_3$	$[\text{PR}_2]^+$	482
$[\text{PCp}^*_2]^+$	$[\text{Cl}(\text{Al}(\text{OR}^{\text{Pr}})_3)_2]^-$	Lewis	$\text{PCp}^*_2\text{Cl} + \text{PhF} \cdot \text{Al}(\text{OR}^{\text{Pr}})_3$	$[\text{PR}_2]^+$, phosphocenium ion	482
$[\text{AsCp}^*\text{Cl}]^+$	$[\text{Cl}(\text{Al}(\text{OR}^{\text{Pr}})_3)_2]^-$	Lewis	$\text{AsCp}^*\text{Cl}_2 + \text{PhF} \cdot \text{Al}(\text{OR}^{\text{Pr}})_3$	$[\text{AsRX}]^+$	482
$[\text{SbCl}_2(\text{AsMe}_3)]^+$	$[\text{OTf}]^-$	Lewis	$\text{SbCl}_3 + \text{AsMe}_3 + \text{TMSOTf}$	$[\text{SbX}_2\text{L}]^+$	483
$[\text{SbPhCl}(\text{AsPh}_3)]^+$	$[\text{AlCl}_4]^-$	Lewis	$\text{SbPhCl}_2 + \text{AsPh}_3 + \text{AlCl}_3$	$[\text{SbRXL}]^+$	483
$[\text{BiPh}(\text{AsPh}_3)]^+$	$[\text{OTf}]^-$	Lewis	$\text{BiCl}_2\text{Ph} + \text{AsPh}_3 + \text{TMSOTf}$	$[\text{BiXL}]^{2+}$	484
$[\text{BiCl}_2(\text{AsPh}_3)]^+$	$[\text{AlCl}_4]^-$	Lewis	$\text{BiCl}_3 + \text{AsPh}_3 + \text{AlCl}_3$	$[\text{BiXL}]^{2+}$	484
$[\text{BiCl}_2(\text{SbPh}_3)]^+$	$[\text{OTf}]^-$	Lewis	$\text{BiCl}_3 + \text{SbPh}_3 + \text{TMSOTf}$	$[\text{BiX}_2\text{L}_2]^+$	483
$[\text{BiCl}_2(\text{AsPh}_3)_2]^+$	$[\text{AlCl}_4]^-$	Lewis	$\text{BiCl}_3 + \text{SbPh}_3 + \text{AlCl}_3$	$[\text{BiX}_2\text{L}]^+$	484
$[\text{BiCl}_2(\text{SbPh}_3)_2]^+$	$[\text{AlCl}_4]^-$	Lewis	$\text{BiCl}_3 + \text{SbPh}_3 + \text{AlCl}_3$	$[\text{BiX}_2\text{L}]^+$	483
$[\text{Bi}(\text{N}_2\text{TMS}_3)_2]^+$	$[\text{GaCl}_4]^-$	Lewis	$\text{Bi}(\text{N}_2\text{TMS}_3)_2\text{Cl} + \text{GaCl}_3$	$[\text{BiX}_2]^{2+}$	485
$\left[\begin{array}{c} \text{tBu-N} \\ \text{P} \\ \text{tBu-N} \end{array} \right]^+$	$[\text{BF}_4]^-$	Salt	$\text{PCl}(\text{N}^{\text{tBu}})_2\text{C}_2\text{H}_2 + \text{Ag}^+ [\text{BF}_4]^-$	P carbenoid	486
$\left[\begin{array}{c} \text{tBu-N} \\ \text{P} \\ \text{tBu-N} \end{array} \right]^+$	$[\text{PF}_6]^-$	Salt	$\text{PCl}(\text{N}^{\text{tBu}})_2\text{C}_2\text{H}_2 + \text{Ag}^+ [\text{PF}_6]^-$	P carbenoid	487
$\left[\begin{array}{c} \text{Me}_2\text{N-N} \\ \text{P} \\ \text{N-Me}_2 \end{array} \right]^+$	$[\text{GaCl}_4]^-$	Lewis	$\text{PCl}(\text{NMe})_2\text{C}_2\text{H}_4 + \text{GaCl}_3$	P carbenoid	479
$\left[\begin{array}{c} \text{tBu-N} \\ \text{P} \\ \text{tBu-N} \end{array} \right]^+$	$[\text{GeCl}_5]^- \cdot [\text{Cl}]^-$	Other	$\text{Ge}(\text{N}^{\text{tBu}})_2\text{C}_2\text{H}_4 + \text{PCl}_3$	P carbenoid	488



Table 10 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
	[PF ₆] [−]	Salt	PCl[(^t Bu) ₂ C ₂ H ₄ + Ag ⁺ [PF ₆] [−]]	P carbenoid	487
	[GaCl ₄] [−]	Lewis	PCl(NDipp) ₂ C ₂ H ₄ + GaCl ₃	P carbenoid	426
	[SbCl ₅ ·THF] [−]	Lewis	AsCl ₃ + SnCl ₂ + (MesN) ₂ C ₂ H ₂	As carbenoid	423
	[GeCl ₅] [−] [Cl] [−]	Other	Ge(^t Bu) ₂ C ₂ H ₄ + AsCl ₃	As carbenoid	488
	[GaCl ₄] [−]	Lewis	As(ⁱ PrN) ₂ C ₁₀ H ₆ Cl + GaCl ₃	As carbenoid	489
	[AlCl ₄] [−]	Lewis	AsCl(HN)SC ₆ H ₄ + AlCl ₃	As carbenoid	490
	[AlCl ₄] [−]	Lewis	AsClS ₂ C ₆ H ₃ CH ₃ + AlCl ₃	As carbenoid	490
	[GaCl ₄] [−]	Lewis	AsCl[(NMe) ₂ C ₃ H ₆ + GaCl ₃	As carbenoid	491
	[OTf] [−]	Other	Sb(ⁱ PrN) ₂ C ₁₀ H ₆ (NMe ₂) + HOTf	Not planar through the ligand NHMe ₂	489



Table 10 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment ^b	Ref.
$\left[\begin{array}{c} \text{Bu} \\ \diagup \\ \text{N} \cdots \text{Sb} \cdots \text{Bu} \\ \diagdown \\ \text{N} \end{array} \right]^+$	$[\text{Sb}_2\text{Cl}_8]^{2-}$	Lewis	$[\text{Sb}(\text{tBuN})_2\text{C}_2\text{H}_2]^+ + \text{SbCl}_3$	Sb carbenoid	492
Oxidation state +V					
$[\text{NF}_4]^+$	$[\text{BF}_4]^-$, $[\text{SbF}_6]^-$, $[\text{Sb}_2\text{F}_{11}]^-$	Lewis	$\text{NF}_3 + \text{F}_2 + \text{BF}_3$ or SbF_5	No X-ray For more structures see ref. 495	493
$[\text{PF}_4]^+$	$[\text{Sb}_3\text{F}_{16}]^-$	Lewis	$\text{PF}_5 + \text{SbF}_5$		494
$[\text{PCl}_4]^+$	$[\text{SnCl}_6]^{2-}$	Lewis	$\text{PCl}_5 + \text{SnCl}_4$		496
$[\text{PBr}_4]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Salt	$\text{PBr}_3 + \text{Br}_2 + \text{Ag}^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$		453
$[\text{PI}_4]^+$	$[\text{AlCl}_4]^-$	Lewis	$\text{PI}_3 + \text{ICl} + \text{AlCl}_3$ in CS_2		495
$[\text{PI}_4]^+$	$[\text{AlBr}_4]^-$	Lewis	$\text{PI}_3 + \text{IBr} + \text{AlBr}_3$ in CS_2		495
$[\text{PI}_4]^+$	$[\text{AlI}_4]^-$	Lewis	$\text{PI}_3 + \text{I}_2 + \text{AlI}_3$ in CS_2		497
$[\text{PI}_4]^+$	$[\text{GaI}_4]^-$	Lewis	$\text{PI}_3 + \text{I}_2 + \text{GaI}_3$ in CS_2		495
$[\text{PI}_4]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Salt	$\text{PI}_3 + \text{I}_2 + \text{Ag}^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$		453
$[\text{AsCl}_4]^+$	$[\text{AsF}_6]^-$	Lewis, Ox	$\text{AsCl}_3 + \text{Cl}_2 + \text{AsF}_5$		498
$[\text{AsCl}_4]^+$	$[\text{As}(\text{OTeF}_5)_6]^-$	Lewis, Ox	$\text{AsCl}_3 + \text{ClOTeF}_5 + \text{As}(\text{OTeF}_5)_5$	499	
$[\text{AsBr}_4]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Salt	$\text{AsBr}_3 + \text{Br}_2 + \text{Ag}^+[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	500	
$[\text{AsBr}_4]^+$	$[\text{FAs}(\text{OTeF}_5)_5]^-$	Lewis, Ox	$\text{AsBr}_3 + \text{BrOTeF}_5 + \text{AsF}(\text{OTeF}_5)_4$	499	
$[\text{SbCl}_4]^+$	$[\text{Sb}_2\text{F}_{11}]^-$	Lewis	$\text{SbCl}_5 + \text{SbF}_5$	501	
$[\text{SbCl}_4]^+$	$[\text{Sb}(\text{OTeF}_5)_6]^-$	Ox	$\text{Sb}(\text{OTeF}_5)_3 + \text{Cl}_2$	502	
$[\text{SbBr}_4]^+$	$[\text{Sb}(\text{OTeF}_5)_6]^-$	Ox	$\text{Sb}(\text{OTeF}_5)_3 + \text{Br}_2$	502	
$\left[\begin{array}{c} \text{Dipp} \\ \diagup \\ \text{N} \cdots \text{P} \cdots \text{L} \\ \diagdown \\ \text{N} \end{array} \right]^+$ L = NMe ₃ , py	$[\text{GaCl}_4]^-$	Ox	$[(\text{CH}_2)_2(\text{NDipp})_2\text{P}]^+ + \text{OL}$	‘‘PO’’ cation	426
$\left[\begin{array}{c} \text{Mes} \\ \diagup \\ \text{N} \cdots \text{R} \cdots \text{Ph} \\ \diagdown \\ \text{N} \end{array} \right]^{2+}$	$[\text{B}(\text{C}_6\text{F}_5)_4]^-$	Lewis	$[(\text{SiMe}_3)_2\text{PF}_2\text{Ph}_2]^+ + [\text{Et}_3\text{Si}(\text{Tol})]^+[\text{B}(\text{C}_6\text{F}_5)_4]^-$	‘‘PFPPh ₂ ’’ ²⁺ , cation	427
$[\text{Ph}_3\text{Pn}]^{2+}$ (Pn = Sb, Bi)	$[\text{OTf}]^-$	Salt	$\text{Ph}_3\text{PnCl}_2 + \text{Ag}^+[\text{OTf}]^-$	Strong contact to the anion	425
Protonated cations					
H_2N_3^+	$[\text{SbF}_6]^-$	Prot	$\text{HN}_3 + \text{HF/SbF}_5$	310	428
$\text{P}(\text{OH})_4^+$	$[\text{SbF}_6]^-$	Prot	$\text{H}_3\text{PO}_4 + \text{HF/SbF}_5$		

^a Classification according to the introduction: Lewis = Lewis acid halogen bond heterolysis, Ox = oxidation, Com = complexation reaction, Prot = protonation, other = all other reactions not classified. ^b HTS = high temperature synthesis.



Table 11 Overview on structurally characterized chalcogen cations

Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
Homopolyatomic cations					
[O ₂] ⁺	[PtF ₆] ⁻	Ox	O ₂ + PtF ₆	Neutron diffraction	510
[O ₂] ⁺	[PtF ₆] ⁻	Ox	O ₂ + PtF ₆		539
[S ₄] ²⁺	[AsF ₆] ⁻	Ox	S + AsF ₅ in SO ₂		540
[S ₄] ²⁺ ·4[S ₇ I] ⁺	[AsF ₆] ⁻	Ox, Lewis	S + I ₂ + AsF ₅ in SO ₂		540 and 541
[S ₄] ²⁺	[Sb ₉ F ₃₉] ²⁻ (= [(SbF ₆) ₅ (Sb ₂ F ₄)(Sb ₂ F ₅)] ²⁻)	Ox	S + SbF ₅ in SO ₂	Traces of Br ₂ were added	542
[S ₄] ²⁺ ·AsF ₃	[AsF ₆] ⁻	Ox	S + AsF ₅ + AsF ₃ in HF	Traces of Br ₂ were added	543
[S ₈] ²⁺	[AsF ₆] ⁻	Ox	S + HF/AsF ₅		544
[S ₈] ²⁺	[SbF ₆] ⁻	Ox	S + SbF ₅ in SO ₂		542
[S ₈] ²⁺	[AsF ₆] ⁻	Ox	[S ₈] ²⁺ ·([AsF ₆]) ₂ in SO ₂ /SO ₂ ClF		73
[S ₈] ²⁺	[AsF ₆] ⁻	Ox	S + AsF ₅ in SO ₂ /SO ₂ ClF		545
[S ₁₉] ²⁺	[SbF ₆] ⁻	Ox	S + SbF ₅ in SO ₂		542
[Se ₄] ²⁺	[AlCl ₄] ⁻	Lewis, Ox	Se + SeCl ₄ + AlCl ₃	HTS (130 °C)	546
[Se ₄] ²⁺	[SeF ₆] ⁻	Lewis, Ox	Se + SeCl ₄ + MCl ₄		547
[Se ₄] ²⁺	[SbF ₆] ⁻	Ox	Se + SbF ₅ in SO ₂		546
[Se ₄] ²⁺	[MoOCl ₄] ⁻	Ox	S + Se + SbF ₅ in SO ₂	HTS (190 °C)	548
[Se ₄] ²⁺	[Mo ₂ O ₂ Cl ₈] ⁻ ·[MCl ₆] ⁻ (M = Zr, Hf)	Lewis, Ox	[Se ₄] ²⁺ ·[Mo ₂ O ₂ Cl ₈] ²⁻ + [Se ₄] ²⁺ ·[MCl ₆] ²⁻ or Se, SeCl ₄ + MoOCl ₄ + MCl ₄	HTS (120 °C)	549
[Se ₈] ²⁺	[AlCl ₄] ⁻	Lewis, Ox	Se + SeCl ₄ + AlCl ₃	Crystals from vapour-phase transport	550
[Se ₈] ²⁺ ·[Te ₆] ¹⁺ ·SO ₂	[AsF ₆] ⁻	Ox	Se + Te + AsF ₅ in SO ₂		551
[Se ₁₀] ²⁺	[SO ₃ F] ⁻	Ox	Se + SbF ₅ in SO ₂		552
[Se ₁₀] ²⁺	[Bi ₄ Cl ₁₄] ⁻	Lewis, Ox	Se + AsF ₅ in SO ₂		553
[Se ₁₇] ²⁺	[WCl ₆] ⁻	Ox	Se + SeCl ₄ + BiCl ₃	HTS (90 °C)	554
[Se ₁₇] ²⁺	[NbCl ₆] ⁻	Lewis, Ox	Se + SeCl ₄ + NbCl ₅ in SnCl ₄	Solventothermal (150 °C)	555
[Se ₁₇] ²⁺	[TaBr ₆] ⁻	Lewis, Ox	Se + SeBr ₄ + TaBr ₅ in SiBr ₄	Solventothermal (150 °C)	556
[Te ₄] ²⁺	[AlCl ₄] ⁻ [Al ₂ Cl ₇] ⁻	Lewis, Ox	Te + TeCl ₄ + AlCl ₃	HTS (250 °C)	557
[Te ₄] ²⁺	[SbF ₆] ⁻	Ox	Te + SbF ₅ in SO ₂	Germanium was added to obtain mixed cations	546
[Te ₄] ²⁺	[WCl ₆] ⁻	Ox	Te + WCl ₆	HTS (190 °C)	558
[Te ₄] ²⁺	[WCl ₆] ⁻	Ox	Te + WCl ₆	Traces of Br ₂ were added, β-mod.	559
[Te ₄] ²⁺	[Zr ₂ Br ₁₀] ²⁻	Lewis, Ox	Te ₂ Br + ZrBr ₄	HTS (210 °C)	560
[Te ₄] ²⁺	[HfCl ₆] ⁻	Lewis, Ox	Te + TeCl ₄ + HfCl ₄	HTS (200 °C)	235
[Te ₄] ²⁺	[MCl ₆] ⁻ (M = Nb, Ta), [TaBr ₆] ⁻ , [Ta ₂ Cl ₁₀ O] ²⁻	Lewis, Ox	Te + TeCl ₄ + MCl ₅	HTS (170 °C)	561
[Te ₄] ²⁺	[Bi ₆ Cl ₁₀] ²⁻ [Bi ₂ Br ₃] ³⁻	Lewis, Ox	Te + TeX ₄ + BiX ₃ (X = Cl, Br)	HTS (170 °C)	562
[Te ₄] ²⁺	[Nb ₂ Cl ₁₀ O] ²⁻	Lewis, Ox	Te + TeCl ₄ + NbCl ₅ + NbOCl ₃	HTS (200 °C)	563
[Te ₄] ²⁺	[MoCl ₄ O] ⁻	Ox	Te + MoOCl ₄	HTS (250 °C)	564
(Te ₁₀) ²⁺ ·Te ₄ ²⁺	[Bi ₄ Cl ₁₆] ²⁻	Lewis, Ox	Te + TeCl ₄ + BiCl ₃	HTS (150 °C)	565
[Te ₆] ²⁺	[MCl ₆] ²⁻ (M = Zr, Hf)	Lewis, Ox	Te + TeCl ₄ + MCl ₄	HTS (220 °C)	103
[Te ₆] ²⁺	[WCl ₄ O] ⁻	Ox	Te + WOCl ₄	HTS (150 °C)	566
[Te ₆] ²⁺	[NbCl ₄ O] ⁻	Lewis, Ox	Te + TeCl ₄ + NbOCl ₃	HTS (200 °C)	567
[Te ₆] ²⁺ ·2AsF ₃	[AsF ₆] ⁻	Ox	Te + AsF ₅ in SO ₂	Reduction of Te ₄ ²⁺	568
(Te ₂) ²⁺ ·n	[AsF ₆] ⁻	Other	[Te ₄] ²⁺ ·([AsF ₆]) ₂ + Fe(CO) ₅ in SO ₂		569
(Te ₂) ²⁺ ·n	[Be ₂ Cl ₆] ²⁻	Lewis, Ox	Te + TeCl ₄ + BeCl ₂		565
(Te ₂) ²⁺ ·n	[WBr ₄ O] ⁻ [Br] ⁻	Ox	Te + WOBr ₄ /WBr ₅		570
(Te ₂) ²⁺ ·n	[WCl ₄ O] ⁻ [Cl] ⁻	Ox	Te + WOCl ₄ /WCl ₅	HTS (230 °C)	571
(Te ₂) ²⁺ ·n	[NbCl ₄ O] ⁻ [Cl] ⁻	Ox	Te + TeCl ₄ + NbOCl ₃	HTS (150 °C)	571
(Te ₂) ²⁺ ·n	[NbBr ₄ O] ⁻ [Br] ⁻	Ox	Te + TeCl ₄ + NbOBr ₃	HTS (225 °C)	101
[Te ₈] ²⁺	[WCl ₆] ⁻	Ox	Te + WCl ₆	HTS (220 °C)	101
[Te ₈] ²⁺	[ReCl ₆] ²⁻	Lewis, Ox	Te + TeCl ₄ + ReCl ₄	HTS (200 °C)	572
				HTS (230 °C)	573

Table 11 (continued)

Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
[Te ₈] ²⁺	[Bi ₄ Cl ₁₄] ²⁻	Lewis, Ox	Te + TeCl ₄ + BiCl ₃	HTS (160 °C)	574
[Te ₈] ²⁺	[U ₂ Br ₁₀] ²⁻	Lewis, Ox	Te + TeBr ₄ + UBr ₅ in SiBr ₄	Solventothermal 200 °C	575
[Te ₈] ²⁺	[Ta ₄ O ₄ Cl ₁₆] ¹⁴⁻	Lewis, Ox	Te + TeCl ₄ + TaCl ₅ + TaOCl ₃ + [BMIM] ⁺ Cl ⁻	Ionic liquid based synthesis	576
[Te ₈] ¹⁴⁺	[(VOCl ₄ O) ²⁻] _n	Ox	Te + VOCl ₃	HTS (270 °C), cubic	577
Metal-nonmetal-cluster complexes					
[Cu(S ₁₂)(S ₈)] ⁺	[Al(OR ^{pp}) ₄] ⁻	Com	Cu ⁺ + S		512 and 578
[Cu(S ₁₂)(CH ₂ Cl ₂)] ⁺	[Al(OR ^{pp}) ₄] ⁻	Com	Cu ⁺ + S		512 and 578
[Ag(S ₈)] ⁺	[Al(OR ^{pp}) ₄] ⁻	Com	Ag ⁺ + S		579
[Cu ₂ Se ₁₀] ²⁺	[Al(OR ^{pp}) ₄] ⁻	Com	Cu ⁺ + Se _(red)		50
[(Ag ₂ (Se ₆)) ²⁺] _∞	[AsF ₆] ⁻	Com	Ag ⁺ + Se		513
[Ag(Se ₆)]	[Ag ₂ (SbF ₆) ₃] ⁻	Com	Ag ⁺ + Se		513
[Ag ₂ (Se ₆)(SO ₂) ₂] ²⁺	[Sb(OTeF ₅) ₃] ⁻	Com	Ag ⁺ + Se _(grey)		513
[Ag ₂ (Se ₆)(SO ₂) ₄] ²⁺	[Al(OR ^{pp}) ₄] ⁻	Com	Ag ⁺ + Se _(grey)		580
[Ag ₂ Se ₁₂] ²⁺	[FAI(OC(C ₂ F ₅)(C ₆ F ₅)) ₃] ⁻ , [Al(OR ^{pp}) ₄] ⁻	Com	Ag ⁺ + Se _(red)		176 and 581
Clusters/cluster-like					
NS ⁺	[AlCl ₄] ⁺	Lewis	(NSCl) ₃ + AlCl ₃		517
[NS ₂] ⁺	[AlCl ₄] ⁺	Other	S ₄ N ₄ + AlCl ₃		582
[S ₄ N ₄] ²⁺	[SbCl ₆] ⁻	Ox	S ₄ N ₄ /S ₃ N ₃ Cl ₃ + SbCl ₅ in SO ₂		583
[S ₄ N ₄] ²⁺	[Sb ₃ F ₁₄] ⁻ , [SbF ₆] ⁻	Ox	S ₄ N ₄ + SbF ₅ in SO ₂		584
[S ₃ N ₃] ⁺	[SbCl ₆] ⁻	Lewis	S ₃ N ₃ Cl ₃ + SbCl ₅ in SOCl ₂		585
[S ₃ Cl ₃] ⁺	[AsF ₆] ⁻	Other	[SbCl ₃] ⁺ [AsF ₆] ⁻ + S ₈ in SO ₂		586
[S ₂ N ₂ O ₂] ²⁺	[SbF ₆] ⁻ , [Sb ₂ F ₁₁] ⁻	Other	[NS ₂] ⁺ [AsF ₆] ⁻ + (CN) ₂ in SO ₂		587
[S ₄ Te ₄] ²⁺ ·SO ₂	[AsF ₆] ⁻	Ox	S + Te + AsF ₅ in SO ₂		588
[S ₃ Te ₃] ²⁺	[AsF ₆] ⁻	Ox	[SeCl ₃] ⁺ [AsF ₆] ⁻ + Se in SO ₂		585
[Se ₃ Cl ₃] ⁺	[AsF ₆] ⁻	Other	[Se ₄] ²⁺ [(AsF ₆)] ₂ + Ph ₂ Se ₂		589
[Se ₆ Ph ₂] ²⁺ ·2SO ₂	[AsF ₆] ⁻	Ox	Te + I ₂ + AsF ₅ in SO ₂ or [SeI ₃] ⁺ [AsF ₆] ⁻ ₂ + [Se ₈] ²⁺ [(AsF ₆)] ₂		590
[Se ₆ I ₂] ²⁺ ·2SO ₂	[WCl ₆] ⁻	Ox	Te + I ₂ + WCl ₆	HTS (150 °C)	566
[Te ₆ I ₂] ²⁺	[(MOX ₄)] ⁻ (M = Mo, X = Cl, Br; M = W, X = Br)	Other	Te ₂ Br + MoOBr ₃ , TeCl ₄ + MoNCl ₂ /MoOCl ₃ , Te + WBr ₅ /WOBr ₃		591
[Se ₄ Te ₂] ²⁺	[MF ₆] ⁻ (M = As, Sb)	Ox	Se + Te + MF ₅ in SO ₂		588
[Se ₄ Te ₃] ²⁺	[MOCl ₄] ⁻	Ox	Se + [Te ₆] ²⁺ [(MOCl ₄)] ₂	HTS (190 °C)	592
[Se ₆ Te ₃] ²⁺ ·[Se ₈ Te ₂] ²⁺ ·2SO ₂	[AsF ₆] ⁻	Ox	S + Se + Te + AsF ₅ in SO ₂	Heterocubane	593
[Se ₆ Te ₂] ²⁺ ·SO ₂	[MF ₆] ⁻ (M = As, Sb)	Ox	Se + Te + MF ₅ in SO ₂	Isostructural to Se ₁₀ ²⁺	593
[Se ₄ Te ₃] ²⁺ ·SO ₂	[AsF ₆] ⁻	Ox	Te + [Se ₈] ²⁺ [(AsF ₆)] ₂ in SO ₂		594
(4n + 2)π-cations					
[S ₂ N ₃] ⁺	[Hg ₂ Cl ₆] ²⁻	Lewis	NSCl + HgCl ₂		595
[S ₂ N ₂] ²⁺	[MF ₆] ⁻ (M = As, Sb)	Other	[SN] ⁺ [MF ₆] ⁻ + [S ₂ N] ⁺ [MF ₆] ⁻ in SO ₂	Cycloaddition	515
[S ₄ N ₃] ⁺	[(SeCl ₅)] [∞]	Ox	S ₄ N ₄ + Se ₂ Cl ₂ in SOCl ₂	10π-aromatic	519
[S ₃ Se] ²⁺	[Sb ₃ F ₁₆] ⁻ ·3[SbF ₆] ⁻	Ox	S + Se + SbF ₅ in SO ₂	Traces of Br ₂ were added, contains disordered mixture of [S _x Se _{x-4}] ²⁺	596
[SSe ₂ N ₂] ²⁺	[AsF ₆] ⁻	Ox	[(SSe ₂ N ₂) ₂] ²⁺ [(AsF ₆)] ₂ + AsF ₅ in SO ₂		597
[S ₂ SeN ₂] ²⁺	[AsF ₆] ⁻	Other	[NS] ⁺ [AsF ₆] ⁻ + [Se ₈] ²⁺ [AsF ₆] ⁻		516
[S ₂ SeN ₂ Cl] ⁺	[AlCl ₄] ⁻	Other	[NS] ⁺ [AlCl ₄] ⁻ + Se/EtSeCl		517 and 518
[Se ₂ Te ₂] ²⁺	[Sb ₃ F ₁₄] ³⁻ ·[SbF ₆] ⁻	Ox	Se + Te + SbF ₅ in SO ₂		598
[SeTe ₃] ⁺	[Sb ₃ F ₁₄] ³⁻ ·[SbF ₆] ⁻	Ox	Se + Te + SbF ₅ in SO ₂	Contains disordered mixture of [SeTe ₃] ²⁺ , [Te ₄] ²⁺ and [Se ₂ Te ₂] ²⁺	598





Table 11 (continued)

Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
$\pi^*-\pi^*$ -complexes					
	$[\text{AsF}_6]^-$	Ox	$\text{S}_4\text{N}_4 + \text{AsF}_5$ in SO_2		599
$[\text{S}_6\text{N}_4]^{2+}$	$[\text{S}_2\text{O}_2\text{F}]^-, [\text{SO}_3\text{F}]^-$	Ox	$\text{S}_4\text{N}_4 + \text{HSO}_3\text{F}$ in SO_2		600
	$[\text{AsF}_6]^-$	Ox	$\text{S} + \text{I}_2 + \text{AsF}_5$ in SO_2		520
	$[\text{Sb}_2\text{F}_{11}]^+$	Ox	$\text{Se} + [\text{I}_2][\text{Sb}_2\text{F}_{11}]^-$ in SO_2		521
	$[\text{MF}_6]^-$ (M = As, Sb)	Other	$\text{S}_4\text{N}_4 + [\text{Se}_4]^{2+}([\text{MF}_6]^-)_2$ in SO_2		601
	$[\text{BF}_4]^-$	Ox	$\text{Me}_2\text{Se}_2 + \text{XeF}_2 + \text{BF}_3 \cdot \text{OEt}_2$		82
$[(\text{MeSe})_4]^{2+}$	$[\text{OTf}]^-$	Ox	$\text{Et}_2\text{Te}_2 + [\text{NO}]^+[\text{OTf}]^-$		602
	$[\text{OTf}]^-$	Ox	$\text{Et}_2\text{Te}_2 + [\text{NO}]^+[\text{OTf}]^-$		82 and 602
Radicals					
	$[\text{AsF}_6]^-$	Ox	$\text{S}_4\text{N}_4 + [\text{Te}_4]^{2+}([\text{AsF}_6]^-)_2$ in SO_2		600
	$[\text{NTf}_2]^-$	Ox	$\text{TTF} + 0.5\text{XeF}_2 + \text{TMSNTf}_2$	TTF = Tetrathiafulvalene	82



Table 11 (continued)

Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
	$[\text{SbCl}_6]^-$	Ox	$(\text{C}_8\text{H}_{10})_2\text{S}_2 + \text{SbCl}_5$	Strong transannular interactions	603
	$[\text{OTf}]^-$	Ox	$(\text{NeoS})_2 + [\text{NO}]^+$	Diorgano disulfide-nitrosonium adduct	604
	$[\text{FA}(\text{OR}^{\text{PF}})_3]^-$	Ox	$\text{C}_{12}\text{H}_8\text{S}_2 + \text{XeF}_2 + \text{Al}(\text{OC}(\text{CF}_3)_3)_3$	$\text{C}_{12}\text{H}_8\text{S}_2 = \text{thianthrene}$	82
	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Ox	$1,8\text{-(SPh)}_2\text{Nap} + [\text{NO}]^+$	S-S-3e-σ-bond Nap = naphthalene	605
	$[\text{AsF}_6]^-$	Ox	$\text{S} + \text{AsF}_5 + (\text{CN})_2 \text{ in } \text{SO}_2$	Traces of Br_2 were added	522
$(\text{CNS}_3^{**})_2$	$[\text{SbF}_6]^-$, $[\text{Sb}_2\text{F}_{11}]^-$	Ox	$\text{S} + \text{SbF}_5 + (\text{CN})_2 \text{ in } \text{SO}_2$	Traces of Br_2 were added	523
	$[\text{Sb}_2\text{F}_{11}]^-$	Ox	$(\text{C}_6\text{F}_5\text{S})_2 + \text{SbF}_5$		606
	$[\text{As}_2\text{F}_{11}]^-$	Ox	$(\text{C}_6\text{F}_5\text{Se})_2 + \text{AsF}_5$		606
	$[\text{SbF}_6]^-$	Ox	$(2,6\text{-Mes}_2\text{C}_6\text{H}_3\text{Ch})_2 + [\text{NO}]^+$		606
Oxidation state +II $[\text{Me}_2\text{S-SMe}]^+$ $[\text{MeS-S}(\text{Me})\text{-SMe}]^+$	$[\text{SbCl}_6]^-$ $[\text{SbCl}_6]^-$	— Ox, Lewis	— $\text{S}_2\text{Me}_2 + \text{SbCl}_5$	$=[\text{S}_3\text{Me}_3]^+$	607 607

Ch = S, Se, Te



Table 11 (continued)

Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
$\left[\begin{array}{c} \text{R} \backslash \\ \text{N} \cdots \text{S} \cdots \text{N} / \text{R} \end{array} \right]^{2+}$ R = Dipp, Dmp	[OTf] [−]	Com	S(OTf) ₂ + (NR) ₂ C ₂ H ₂	Sulfur carbenoid, oxidation state unclear	528
$\left[\begin{array}{c} \text{Dipp} \backslash \\ \text{N} \cdots \text{S} \cdots \text{N} / \text{Dipp} \end{array} \right]^{2+}$	[B(C ₆ F ₅) ₄] [−]	Other	[S(NDipp) ₂ C ₂ H ₂] ²⁺ ([OTf] [−]) ₂ + K ⁺ [B(C ₆ F ₅) ₄] [−]	Sulfur carbenoid, salt metathesis, oxidation state unclear	528
$\left[\begin{array}{c} \text{cis/trans-} \\ \text{Bu} \cdots \text{Se} \cdots \text{Bu} \\ \\ \text{Me} \end{array} \right]^+$	[X] [−] = [SbCl ₆] [−] , [BF ₄] [−]	Other	[S ₃ Me ₃] ⁺ [X] [−] + C ₂ H ₂ ^t Bu ₂	Thiiranium ion	608
$\left[\begin{array}{c} \text{Bu} \cdots \text{Se} \cdots \text{Bu} \\ \\ \text{Me} \end{array} \right]^+$	[X] [−] = [BF ₄] [−] , [PF ₆] [−]	Other	[S ₃ Me ₃] ⁺ [X] [−] + C ₂ ^t Bu ₂	Thiirenium ion	608
[MeSC ₂ ^t Bu ₂] ⁺ ·CH ₂ Cl ₂	[CHB ₁₁ Cl ₁₁] [−]	Ox	Me ₂ S ₂ + C ₂ ^t Bu ₂ + XeF ₂ + Me ₃ Si(CHB ₁₁ Cl ₁₁)		82
$\left[\begin{array}{c} \text{Bu} \cdots \text{Se} \cdots \text{Bu} \\ \\ \text{Ph} \end{array} \right]^+$	[B(C ₆ F ₅) ₄] [−]	Ox	Ph ₂ S ₂ + C ₂ ^t Bu ₂ + XeF ₂ + [Me ₃ Si(Tol)] ⁺ [B(C ₆ F ₅) ₄] [−]	Thiirenium ion	82
[MeSe-Se(Me)-SeMe] ⁺	[SbCl ₆] [−]	Ox, Lewis	Se ₂ Me ₂ + SbCl ₅	=[Se ₃ Me ₃] ⁺	607
$\left[\begin{array}{c} \text{Bu} \cdots \text{Se} \cdots \text{Bu} \\ \\ \text{Me} \end{array} \right]^+ \cdot 0.5 \text{CH}_2\text{Cl}_2$	[SbCl ₆] [−]	Other	[Se ₃ Me ₃] ⁺ [SbCl ₆] [−] + C ₂ ^t Bu ₂	Selenirenium ion	526
[PhSeC ₂ Ad ₂] ⁺ ·CH ₂ Cl ₂	[SbCl ₆] [−]	Other	[Phse] ⁺ [SbCl ₆] [−] + C ₂ Ad ₂		526
$\left[\begin{array}{c} \text{Dipp} \backslash \\ \text{N} \cdots \text{Se} \cdots \text{N} / \text{Dipp} \end{array} \right]^{2+}$	[SnCl ₆] ^{2−}	Ox, Lewis	(NDipp) ₂ C ₂ H ₂ + SnCl ₂ + SeCl ₄	Selenium carbenoid	529
$\left[\begin{array}{c} \text{Bu} \cdots \text{Se} \cdots \text{Bu} \\ \\ \text{Ph} \end{array} \right]^+ \cdot 1.5 \text{CCl}_4$	[SbF ₆] [−]	Other	[PhTe] ⁺ [SbF ₆] [−] + C ₂ ^t Bu ₂	Tellurirenium ion	526
$\left[\begin{array}{c} \text{NMMe}_2 \\ \\ \text{Se} \\ \\ \text{Ph} \end{array} \right]^+$	[PF ₆] [−]	Other	C ₆ H ₃ (CH ₂ NMe ₂) ₂ SeMe + ^t BuOCl + K ⁺ [PF ₆] [−]		524

Table 11 (continued)

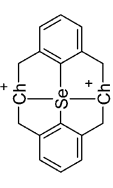
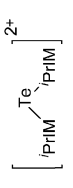
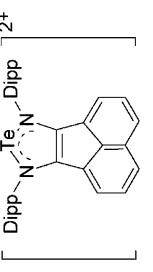
Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
	[PF ₆] [−]	Ox	(C ₆ H ₅ (CH ₂) ₂ Ch) ₂ Se + [NO] ⁺ [PF ₆] [−]		609
	[OTf] [−]	Other	[(Dipp ₂ BIAN)Te] ²⁺ ([OTf] [−]) ₂ + DMAP	Ligand exchange	525
	[OTf] [−]	Other	[(Dipp ₂ BIAN)Te] ²⁺ ([OTf] [−]) ₂ + 2 ⁱ PrIM	Ligand exchange, anions coordinating	525
	[OTf] [−]	Other	[(Dipp ₂ BIAN)Te] ²⁺ ([OTf] [−]) ₂ + ⁱ PrIM	Ligand exchange, 4 equivalents of ⁱ PrIM	525
	[OTf] [−]	Hal	(Dipp ₂ BIAN)TeI ₂ + Ag ⁺ [OTf] [−]	Tellurium carbenoid, Dipp ₂ BIAN = 1,4-(2,6-diisopropyl)phenyl-bis(arylimino)acenaphthene	525
Oxidation state +IV					
[SF ₃] ⁺	[BF ₄] [−]	Lewis	SF ₄ + BF ₃	Crystals through sublimation	610
[SF ₃] ⁺	[GeF ₆] ^{2−}	Lewis	SF ₄ + GeF ₄		611
[SbCl ₃] ⁺	[ICl ₄] [−]	Ox, Lewis	S + Cl ₂ + I ₂		612
[SbCl ₃] ⁺	[ICl ₄] [−]	Ox, Lewis	S + Cl ₂ + I ₂	Second modification	613
[SbCl ₃] ⁺	[UCl ₆] [−]	Lewis	SOCl ₂ + UCl ₅		614
[SbCl ₃] ⁺	[AlCl ₄] [−]	Lewis	SOCl ₂ + AlCl ₃		615
[SbCl ₃] ⁺	[SbCl ₆] [−]	Ox, Lewis	As ₄ S ₄ + SbCl ₅ in SO ₂	HTS (100 °C), large excess of ICl ₃ was added	531
[SbCl ₃] ⁺	[MoOCl ₄] [−]	Ox	S + MOCl ₄	Attempt to prepare [SBr ₃] ⁺ [SbCl ₆] [−]	616
[SBr _{1.2} Cl _{1.8}] ⁺	[SbCl ₆] [−]	Ox, Lewis	S + Br ₂ + SbCl ₅ in SO ₂		531
[SBr ₃] ⁺	[AsF ₆] [−]	Lewis	S + Br ₂ + AsF ₅ in SO ₂		617 and 618
[(SX ₂) ₂ N] ⁺ (X = F, Cl)	[AsF ₆] [−]	Ox	[S ₂ N] ⁺ [AsF ₆] [−] + X ₂ in SO ₂		619
[SeF ₃] ⁺	[NbF ₆] [−] , [Nb ₂ F ₁₁] [−] , [TaF ₆] [−]	Lewis	SeF ₄ + NbF ₅	First structure with a [ChX ₃] ⁺ cation (Ch = S, Se, Te; X = F, Cl, Br, I)	620
[SeCl ₃] ⁺	[AlCl ₄] [−]	Lewis	SeCl ₄ + AlCl ₃ in SO ₂ Cl ₂		621
[SeCl ₃] ⁺	[SbCl ₆] [−]	Ox, Lewis	As + Se + SbCl ₅ or SeCl ₄ + SbCl ₅ in SO ₂	Melt	531
[SeCl ₃] ⁺	[MoOCl ₄] [−]	Others	[Se ₄] ²⁺ [MoOCl ₄] ^{2−} in SOCl ₂	Decomposition at 150 °C, β-modification	622
[SeCl ₃] ⁺	[AuCl ₄] [−]	Lewis	SeCl ₄ + AuCl ₃		623
[SeBr ₃] ⁺	[AsF ₆] [−]	Ox, Lewis	Se + Br ₂ + AsF ₅ in SO ₂		624
[SeBr ₃] ⁺	[SbF ₆] [−]	Ox, Lewis	[Se ₄] ²⁺ [(SbF ₆] [−]) ₂ + Br ₂ + AsF ₅ in SO ₂	Small amount of AsF ₃ was added	624
[SeBr ₃] ⁺	[AlBr ₄] [−]	Lewis	SeBr ₄ + AlBr ₃	HTS (150 °C)	625
[SeI ₃] ⁺	[AsF ₆] [−]	Ox, Lewis	[Se ₄] ²⁺ [(SbF ₆] [−]) ₂ + I ₂ in AsF ₃		617 and 618
[SeI ₃] ⁺	[SbF ₆] [−]	Ox, Lewis	Se + I ₂ + SbF ₅ in SO ₂		626
[TeF ₃] ⁺	[Sb ₂ F ₁₁] [−]	Lewis	TeF ₄ + SbF ₅		531
[TeF ₃] ⁺	[SO ₄] ^{2−}	Ox, Lewis	Te + Br ₂ + AsF ₅ in SO ₂		



Table 11 (continued)

Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
[TeCl ₃] ⁺	[AlCl ₄] ⁻	Lewis	TeCl ₄ + AlCl ₃	Monoclinic	627
[TeCl ₃] ⁺	[AlCl ₄] ⁻	Lewis	S ₇ TeCl ₂ + AlCl ₃	TeCl ₃ [AlCl ₄] (triclinic)	531
[TeCl ₃] ⁺	[AsF ₆] ⁻	Lewis	TeCl ₄ + AsF ₅		531
[TeCl ₃] ⁺	[SbF ₆] ⁻	Lewis	TeF ₄ + SbF ₅ in CH ₂ Cl ₂	Solvent decomposition	531
[TeCl ₃] ⁺	[AuCl ₄] ⁻	Lewis	TeCl ₄ + AuCl ₃		628
[TeCl ₃] ⁺	[MoCl ₆ O] ⁻	Lewis	TeCl ₄ + MoOCl ₃	HTS (180 °C)	629
[TeCl ₃] ⁺	[MoCl ₆] ²⁻	Lewis	TeCl ₄ + MoCl ₄	HTS (195 °C)	630
[TeCl ₃] ⁺	[Re ₂ Cl ₆] ²⁻	Lewis, Ox	Te + TeCl ₄ + ReCl ₅	HTS (150 °C), β-modification	630
[TeCl ₃] ⁺	[MoCl ₆] ²⁻	Lewis	TeCl ₄ + MoCl ₄	HTS (300 °C)	631
[TeCl ₃] ⁺	[MCl ₆] ⁻ (M = Nb, Ta)	Lewis	TeCl ₄ + MCl ₅	α- and β-modification	632
[TeCl ₃] ⁺	[WCl ₆] ⁻	Lewis	TeCl ₄ + WCl ₆		632
[TeBr ₃] ⁺	[AsF ₆] ⁻	Ox, Lewis	Te + Br ₂ + AsF ₅ in SO ₂	HTS (160 °C)	624
[TeBr ₃] ⁺ · 1/2Br ₂	[AuBr ₄] ⁻	Ox, Lewis	Te + Au + Br ₂	Decomposition above 250 °C	633
[TeBr ₃] ⁺	[Zr ₂ Br ₉] ⁻	Other	[Te ₄] ¹²⁺ [Zr ₂ Br ₁₀] ¹²⁻		560
[TeBr ₃] ⁺	[MBr ₆] ⁻ (M = Ta, W)	Lewis	TeBr ₄ + MBr ₅		632
[TeI ₃] ⁺	[AsF ₆] ⁻	Ox, Lewis	Te + I ₂ + AsF ₅ in SO ₂		634
[TeI ₃] ⁺	[SbF ₆] ⁻	Ox, Lewis	Te + I ₂ + SbF ₅ in SO ₂		617 and 618
[TeI ₃] ⁺	[AlI ₄] ⁻	Lewis, Ox	Te + I ₂ + AlI ₃		625
[TeI ₃] ⁺	[ML ₄] ⁻ (M = Ga, In)	Lewis, Ox	Te + M + I ₂		635
[TeI ₃] ⁺ · 1/2SO ₂	[AsF ₆] ⁻	Ox, Lewis	Te + I ₂ + AsF ₅ in SO ₂	HTS (150 °C)	625
[TeI ₃] ⁺	[Al(OR) ^{PR}] ₄] ⁻	Salt	TeX ₄ + Ag ⁺ [Al(OR) ^{PR}] ₄] ⁻	HTS (350 °C)	636
[TePh ₃] ⁺	[B(C ₆ F ₅) ₄] ⁻	Other	[TePh ₅] ⁺ [B(C ₆ F ₅) ₄] ⁻	Hemisolvate of [TeI ₃] ⁺ [AsF ₆] ⁻	637
[Te(C ₆ F ₅) ₃] ⁺	[OTf] ⁻	Other	Te(C ₆ F ₅) ₄ + TMSOTf	Good solubility	532
[Te(N ₃) ₃] ⁺	[SbF ₆] ⁻	Ox	[Te ₄] ¹²⁺ [(SbF ₆) ⁻] ₂ + KN ₃ in SO ₂	Thermal decomposition at 150 °C	533
[F(TeCl ₃) ₂] ⁺	[Sb(OTeF ₅) ₆] ⁻	Ox, Lewis	TeBr ₄ + Ag ⁺ [Sb(OTeF ₅) ₆] ⁻ in SO ₂ ClF	Side products may be explosive!	534
					28
	[BF ₄] ⁻	Ox	(C ₆ H ₅ (CH ₂ CHPh) ₂)TePh + [NO] ⁺ [BF ₄] ⁻		638
	[OTf] ⁻	Ox	(C ₆ H ₅ (CH ₂ SPh) ₂)TePh + ^t BuOCl + O(Tf) ₂		638
	[B(C ₆ F ₅) ₄] ⁻	Hal	TePh ₃ Cl + Ag ⁺ [OTf] ⁻ + Li ⁺ [B(C ₆ F ₅) ₄] ⁻		532
	[ClO ₄] ⁻	Hal	TePh ₅ Cl + Ag ⁺ [ClO ₄] ⁻		535
	[B(Ar ^{CF₃}) ₄] ⁻	Hal	TePh ₅ Cl + Ag ⁺ [OTf] ⁻ + [B(Ar ^{CF₃}) ₄] ⁻		535

Oxidation state +VI

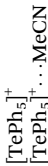
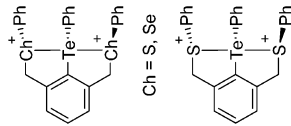




Table 11 (continued)

Cation	Anion	Class ^a	Synthesis	Comment ^b	Ref.
Protonated cations					
$\left[\begin{array}{c} \text{OH} \\ \\ \text{HO}-\text{C}^+-\text{OH} \end{array} \right]^+$	$[\text{AsF}_6]^-$	Prot	$\text{OC}(\text{OTMS})_2 + \text{HF}/\text{AsF}_5$		537
$\text{H}-\text{O}-\text{S}^+(\text{OH})_2$	$[\text{Ge}_3\text{F}_{16}]^{4-}$	Prot	$\text{Me}_2\text{SO} + \text{HF}/\text{GeF}_4$		639
$\left[\begin{array}{c} \text{O} \quad \text{F} \\ \quad \\ \text{HO}-\text{S}^+-\text{C}^+-\text{F} \\ \quad \\ \text{HO} \quad \text{F} \end{array} \right]^+$	$[\text{SbF}_6]^-$	Prot	$\text{OTf}_2 + \text{HF}/\text{SbF}_5$		536
$\left[\begin{array}{c} \text{O} \\ \\ \text{HO}-\text{S}^+(\text{OH})_2 \end{array} \right]^+$	$[\text{SbF}_6]^-$	Prot	$\text{SO}_2(\text{OTMS})_2 + \text{HF}/\text{SbF}_5$		538

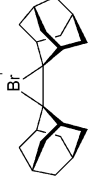
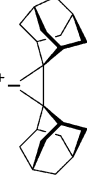
^a Classification according to the introduction (Table 2): Com = complexation reaction, Lewis = Lewis acid halogen bond heterolysis, Prot = protonation, Ox = oxidation, other = all other reactions not classified. ^b HTS = high temperature synthesis.

Table 12 Overview on all structurally characterized halogen cations

Cation	Anion	Class ^a	Synthesis	Comment	Ref.
Homopolyatomic cations					
$[\text{Cl}_3]^+$	$[\text{AsF}_6]^-$	Lewis	$[\text{Cl}_3\text{F}]^+[\text{AsF}_6]^- + \text{Cl}_2 + \text{AsF}_5$		42
$[\text{Cl}_3]^+$	$[\text{X}]^- = [\text{SbF}_6]^- , [\text{Sb}_2\text{F}_{11}]^- , [\text{Sb}_3\text{F}_{16}]^-$	Other	$[\text{Cl}_2\text{O}_2]^+[\text{X}]^- + \text{Cl}_2$ in HF at RT		42
$[\text{Br}_2]^+$	$[\text{Sb}_3\text{F}_{16}]^-$	Ox	$\text{Br}_2 + \text{HSO}_3\text{F}/\text{SbF}_5/3\text{SO}_3$		663
$[\text{Br}_3]^+$	$[\text{AsF}_6]^-$	Other	$[\text{BrF}_2]^+[\text{AsF}_6]^-$ decomposition	Store $[\text{BrF}_2]^+[\text{AsF}_6]^-$ for 20 years (!)	645
$[\text{Br}_5]^+$	$[\text{MF}_6]^-$ (M = As, Sb)	Ox	$[\text{XeF}]^+[\text{MF}_6]^- + \text{Br}_2$		211
$[\text{I}_2]^+$	$[\text{Sb}_2\text{F}_{11}]^-$	Ox	$\text{I}_2 + \text{SbF}_5$		15
$[\text{I}_3]^+$	$[\text{AsF}_6]^-$	Ox	$\text{I}_2 + \text{AsF}_5$		15
$[\text{I}_5]^+$	$[\text{AsF}_6]^-$	Ox	$\text{I}_2 + \text{AsF}_5$		664
$[\text{I}_{15}]^{3+}$	$[\text{SbF}_6]^-$	Ox	$\text{I}_2 + \text{SbF}_5$		15
Metal–nonmetal-cluster complexes					
$[\text{Ag}(\text{Cl}_2)]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{Ag}^+[\text{Al}(\text{OR}^{\text{PF}})_4]^- + \text{Cl}_2$	Only stable at low temperature	
$[\text{Ag}(\text{Br}_2)]^+$	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{Ag}^+[\text{Al}(\text{OR}^{\text{PF}})_4]^- + \text{Br}_2$		
$[\text{Ag}_2(\text{I}_2)_x]^{2+}$ (x = 1, 4, 6)	$[\text{Al}(\text{OR}^{\text{PF}})_4]^-$	Com	$\text{Ag}^+[\text{Al}(\text{OR}^{\text{PF}})_4]^- + \text{I}_2$		
$[\text{Ag}(\text{I}_2)]_n^{n+}$	$[\text{MF}_6]^-$ (M = Sb, As)	Com	$\text{Ag}^+[\text{MF}_6]^- + \text{I}_2$	First complex with halogen as donor	646
$\pi^*-\pi^*$ -complexes					
$[\text{Cl}_4]^+$	$[\text{IrF}_6]^-$	Ox	$\text{Cl}_2 + \text{IrF}_6$	Radical, homo-polyatomic cation	644
$[\text{Cl}_2\text{O}_2]^+$	$[\text{X}]^- = [\text{SbF}_6]^- , [\text{Sb}_2\text{F}_{11}]^-$	Com	$[\text{O}_2]^+ + \text{Cl}_2$		42
$[\text{I}_4]^{2+}$	$[\text{MF}_6]^-$ (M = As, Sb), and $[\text{Sb}_3\text{F}_{14}]^-$ $[\text{SbF}_6]^-$	Com	$2[\text{I}_2]^+$	Also homopoly-atomic cation	665
Oxidation state +I					
$[\text{I}(\text{NCMe})_2]^+$	$[\text{AsF}_6]^-$	Other	$[\text{I}_3]^+ + \text{MeCN}$	$[\text{I}_3]^+$ as I^+ donor	649



Table 12 (continued)

Cation	Anion	Class ^a	Synthesis	Comment	Ref.
Oxidation state +III					
[ClF ₂] ⁺	[AsF ₆] ⁻	Lewis	ClF ₃ + AsF ₅		666
[ClF ₂] ⁺	[SbF ₆] ⁻	Lewis	ClF ₃ + SbF ₅		667
[ClF ₂] ⁺	[RuF ₆] ⁻	Lewis	ClF ₃ + RuF ₅		668
[ClMe ₂] ⁺ [ClEt ₂] ⁺	[CHB ₁₁ Cl ₁₁] ⁻	Prot	H(CHB ₁₁ Cl ₁₁) + RCl (R = Me, Et)	Protonation of RCl	655
[BrF ₂] ⁺	[SbF ₆] ⁻	Lewis	BrF ₃ + SbF ₅		669
[Br(C ₆ F ₅) ₂] ⁺	[BF ₄] ⁻	Lewis	BrF ₃ + (C ₆ F ₅) ₂ BF ₃		670
	[OTf] ⁻	Lewis	C ₂₀ H ₂₈ + Br ₂ + MeOTf	Stable bromonium ion C ₂₀ H ₂₈ = adamantylideneadamantane	656
[ICl ₂] ⁺	[SbF ₆] ⁻	Lewis	I ₂ Cl ₆ + SbF ₅		671
[IBr ₂] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	IBr + SbF ₅		672
[IBr _{0.75} Cl _{1.25}] ⁺	[SbCl ₆] ⁻	Lewis	IBr + Cl ₂ + SbF ₅		672
[I ₃ Cl ₂] ⁺	[X] ⁻ = [SbCl ₆] ⁻ , [AlCl ₄] ⁻	Lewis, other	I ₂ + SbCl ₅		652–654
[I ₃ Br ₂] ⁺	[SbCl ₆] ⁻	Lewis, other	I ₂ + SbCl ₅		653
[I(C ₆ F ₅)R] ⁺	[BF ₄] ⁻	Lewis	I(C ₆ F ₅)F ₂ + RBF ₂ R = C ₆ H ₅ , <i>o</i> -C ₆ H ₄ F, <i>m</i> -C ₆ H ₄ F, <i>p</i> -C ₆ H ₄ F, 2,4,6-C ₆ H ₂ F ₃ , C ₆ F ₅	Strong interactions with the anions	673
	[OTf] ⁻	Lewis	C ₂₀ H ₂₈ + I ₂ + Ag ⁺ [OTf] ⁻	Stable iodonium ion C ₂₀ H ₂₈ = adamantylideneadamantane	656
Oxidation state +V					
[ClF ₄] ⁺	[SbF ₆] ⁻	Lewis	ClF ₅ + SbF ₅		661
[ClO ₂] ⁺	[SbF ₆] ⁻	Lewis	ClO ₂ F + HF/SbF ₅		674
[ClO ₂] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	ClO ₂ F + 2SbF ₅	Also an oxidation product of [ClF ₂] ⁺	675
[ClO ₂] ⁺	[X] ⁻ = [BF ₄] ⁻ , [GeF ₅] ⁻	Lewis	ClO ₂ F + BF ₃ , GeF ₄		611 and 676
[ClO ₂] ⁺	[ClO ₄] ⁻	Ox	ClO ₂ + O ₃	Cl ₂ O ₆	677
[ClO ₂] ⁺	[RuF ₆] ⁻	Lewis Ox	ClO ₂ F + 2RuF ₅ ClF ₃ + HF/RuO ₄		668
[BrF ₄] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	BrF ₅ + 2SbF ₅		662 and 678
[BrO ₂] ⁺	[SbF ₆] ⁻	Lewis, other	BrO ₃ F + SbF ₅	Reduction of Br ^{VII}	674
[IF ₄] ⁺	[SbF ₆] ⁻	Lewis	IF ₅ in HF/SbF ₅		662 and 679
[IF ₄] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	IF ₅ + 2HF/SbF ₅		662 and 680
[I(C ₆ F ₅) ₂ F ₂] ⁺	[BF ₄] ⁻	Lewis	C ₆ F ₅ IF ₄ + C ₆ F ₅ BF ₂		657
Oxidation state +VII					
[ClF ₆] ⁺	[Sb ₂ F ₁₁] ⁻	Ox, Lewis	ClF ₅ + F ₂ + SbF ₅		660
[BrF ₆] ⁺	[Sb ₂ F ₁₁] ⁻	Ox, Lewis	BrF ₅ + F ₂ + SbF ₅		660
[IF ₆] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	IF ₇ + SbF ₅		660

^a Classification according to the introduction (Table 2): Lewis = Lewis acid halogen bond heterolysis, Ox = oxidation, Com = complexation reaction, Other = all other reactions not classified.



Table 13 Cationic noble gas compounds with weakly coordinating anions

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
Homopolyatomic cations					
[Xe ₃] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	[XeF] ⁺ /Xe + SbF ₅	First homopoly-atomic cation	27
[Xe ₄] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	[XeF] ⁺ [Sb ₂ F ₁₁] ⁻ , SbF ₅ + Xe	No X-ray	696
Metal-nonmetal-cluster complexes					
[AuXe ₄] ²⁺	[Sb ₂ F ₁₁] ⁻	Lewis, Com	AuF ₃ , Xe in HF/SbF ₅	First xenon metal complex, gold(II) 2 modifications: triclinic and tetragonal	695 and 697
[trans-AuXe ₂] ²⁺	[SbF ₆] ⁻	Com	HAuCl ₄ , XeF ₂ , Xe in HF/SbF ₅	Gold(II)	695
[cis-AuXe ₂] ²⁺	[Sb ₂ F ₁₁] ⁻	Com	[AuXe ₄] ²⁺ [Sb ₂ F ₁₁] ⁻ 2-Xe at RT	Gold(II)	695
[trans-AuXe ₂ F] ²⁺	[SbF ₆] ⁻ + [Sb ₂ F ₁₁] ⁻	Com	Au, XeF ₂ , Xe in HF/SbF ₅	Gold(III)	695
[Au ₂ Xe ₂ F] ³⁺	[SbF ₆] ⁻	Com	Au, XeF ₂ , Xe in HF/SbF ₅	Gold(III)	695
[F ₃ AsAuXe] ⁺	[Sb ₂ F ₁₁] ⁻	Com	AuF ₃ , Xe, AsF ₃ in HF/SbF ₅		698
[HgXe] ²⁺	[SbF ₆] ⁻ + [Sb ₂ F ₁₁] ⁻	Com	HgF ₂ , Xe in HF/SbF ₅		698
Oxidation state +II					
[KrF] ⁺	[MF ₆] ⁻ (M = As, Sb, Bi)	Lewis	KrF ₂ + MF ₅	First krypton cation	699
[KrF] ⁺	[AuF ₆] ⁻	Lewis	KrF ₂ + Au		682
[Kr ₂ F ₃] ⁺	[SbF ₆] ⁻ [AsF ₆] ⁻	Lewis	KrF ₂ + MF ₅		699
[Kr ₂ F ₃] ⁺ [KrF] ⁺					
[XeF] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	XeF ₂ + 2SbF ₅		687
[XeF] ⁺	[AuF ₆] ⁻	Lewis	XeF ₂ + RuF ₅		700
[XeF] ⁺	[AsF ₆] ⁻	Lewis	XeF ₂ + AsF ₅		701
[XeF] ⁺	[N(SO ₂ F) ₂] ⁻	Lewis	XeF ₂ + HN(SO ₂ F) ₂	First Xe-N bond, strong interaction with the anion	702
[XeF] ⁺ ·HF					
[XeF] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	XeF ₂ + 2SbF ₅		27
[XeF] ⁺	[X] ⁻ = [AsF ₆] ⁻ , [SbF ₆] ⁻ , [Sb ₂ F ₁₁] ⁻ , [BiF ₆] ⁻ , [Bi ₂ F ₁₁] ⁻	Lewis	XeF ₂ + AsF ₅ , SbF ₅ , BiF ₅	Better structures for [XeF] ⁺ [X] ⁻ : [X] ⁻ = [MF ₆] ⁻ , [SbF ₆] ⁻ , [Sb ₂ F ₁₁] ⁻	691
[Xe ₂ F ₃] ⁺	[AsF ₆] ⁻	Lewis	2XeF ₂ + AsF ₅	Monoclinic structure	703 and 704
[Xe ₂ F ₃] ⁺	[AuF ₆] ⁻	Lewis	XeF ₂ + AuF ₅		686
[Xe ₂ F ₃] ⁺	[MF ₆] ⁻ (M = As, Sb)	Com	[XeF] ⁺ as starting material		705
[Xe(N(SO ₂ F) ₂)] ⁺	[Sb ₃ F ₁₆] ⁻	Lewis	3 step synthesis with AsF ₅ and SbF ₅	[Xe ₂ F ₃] ⁺ [AsF ₆] ⁻ trigonal structure	706
[XeC ₆ F ₃] ⁺	[(F ₃ C) ₂ BF ₂] ⁻	Lewis	XeF ₂ + B(C ₆ F ₅) ₃		693
[(MeCN)XeC ₆ F ₅] ⁺ [XeC ₆ F ₃] ⁺	[BX] ⁻ (X = CF ₃ , C ₆ F ₅) [BX] ⁻ (X = CF ₃ , CN)	Lewis, Com	XeF ₂ + C ₆ F ₅ BF ₂	First Xe-C bond	707
[(C ₆ F ₅ Xe) ₂ Cl] ⁺	[AsF ₆] ⁻	Other	[XeC ₆ F ₃] ⁺ + TMSCl	Salt metatheses with [XeC ₆ F ₃] ⁺ [BF ₄] ⁻	
[XeOCH ₃] ⁺ (Ch = Se, Te)	[AsF ₆] ⁻	Lewis	FXeOCH ₃ + AsF ₅	First Xe-Cl bond	708
[XeOTeF ₅] ⁺ ·SO ₂ ClF	[Sb(OTeF ₅) ₆] ⁻	Other	Xe(OTeF ₅) ₂ + Sb(OTeF ₅) ₃	First Xe-O bond	709
[XeCl] ⁺	[Sb ₂ F ₁₁] ⁻	Other	[XeF] ⁺ + [Cl] ⁻	OTeF ₅ ⁻ abstraction	710
[XeN(H)TeF ₃] ⁺	[AsF ₆] ⁻	Lewis	[F ₃ TeNH ₃] ⁺ [AsF ₆] ⁻ + XeF ₂	Ligand exchange	711
[(F ₃ SN)XeF] ⁺	[AsF ₆] ⁻	Com	F ₃ SN + [XeF] ⁺	First Xe-N(sp ³) bond	712
[XeNSF ₄] ⁺	[AsF ₆] ⁻	Other	In solid state and HF/BrF ₃ solution	First Xe-N(sp) bond	713
[(F ₃ SN)XeNSF ₄] ⁺	[AsF ₆] ⁻	Com	[F ₃ SNXeF] ⁺ + F ₃ SN	Rearrangement of [F ₃ SNXeF] ⁺ [AsF ₆] ⁻	714
[Xe ₃ OF ₃] ⁺	[MF ₆] ⁻ (M = As, Sb)	Other	H ₂ O + XeF ₂ /[XeF] ⁺	Hydrolysis to XeFOH followed by a reaction with [Xe ₂ F ₃] ⁺	715
Oxidation state +IV					
[XeF ₃] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	XeF ₄ + 2SbF ₅		717
[XeF ₃] ⁺	[BiF ₆] ⁻	Lewis	XeF ₄ + BiF ₅		718
[XeF ₃] ⁺	[Sb ₂ F ₁₁] ⁻	Lewis	XeF ₄ + SbF ₅	Better structure	691
[XeF ₃] ⁺ ·HF [H ₃ F ₄] ⁺ ·2[(XeF ₃)] ⁺ ·HF	[Sb ₂ F ₁₁] ⁻ [SbF ₆] ⁻ ·2[Sb ₂ F ₁₁] ⁻ [SbF ₆] ⁻	Lewis	XeOF ₂ ·xHF + HF/SbF ₅		719

Table 13 (continued)

Cation	Anion	Class. ^a	Synthesis	Comment	Ref.
Oxidation state +VI					
$[\text{XeF}_5]^+$	$[\text{PtF}_6]^-$	Lewis	$\text{XeF}_6 + \text{PtF}_5$	First xenon cation structure	688 and 689
$[\text{XeF}_5]^+$	$[\text{AsF}_6]^-$	Lewis	$\text{XeF}_6 + \text{AsF}_5$		704
$[\text{XeF}_5]^+$	$[\text{SbF}_6]^-$	Ox	$[\text{XeF}_5]^+[\text{SbF}_6]^- + \text{F}_2$ in aHF	Crystals from a $\text{O}_2\text{SbF}_6/\text{XeF}_5/\text{SbF}_6$ mixture	692
$[\text{XeF}_5]^+$	$[\text{Sb}_2\text{F}_{11}]^-$	Other	$[\text{XeF}_5]^+[\text{SbF}_6]^-$		692
$[\text{XeF}_5]^+$	$[\text{RuF}_6]^{2-}$	Lewis	$\text{XeF}_6 + \text{RuF}_5$		700
$[\text{XeF}_5]^+$	$[\text{PdF}_6]^{2-}$	Lewis	$2\text{XeF}_6 + \text{PdF}_4$		720
$[\text{XeF}_5]^+$	$[\text{Ti}_4\text{F}_{19}]^{3-}$	Lewis	$\text{XeF}_6 + \text{TiF}_4$	XeF_6 from XeF_2 , F_2 and UV radiation	721
$[\text{XeF}_5]^+$	$[m\text{-F}(\text{OsO}_3\text{F}_2)_2]^- [\text{OsO}_3\text{F}_2]^-$	Lewis	$\text{XeF}_6 + (\text{OsO}_3\text{F}_2)_\infty$		690
$[\text{XeF}_5]^+$	$[\text{Cu}(\text{SbF}_6)_3]^-$	Other	$[\text{XeF}_5]^+[\text{SbF}_6]^- + \text{Cu}^+[\text{SbF}_6]^-$ in aHF	Anion exchange	43
$[\text{XeF}_5]^+$	$[\text{AgF}_4]^-$	Lewis	$\text{AgF}_2 + \text{KrF}_2 + \text{XeF}_6$ in aHF		722
$[\text{XeF}_5]^+$	$[\text{AuF}_4]^-$	Lewis	$\text{XeF}_6 + \text{BrF}_3/\text{AuF}_3$		722
$[\text{XeF}_5]^+$	$[\text{AsF}_6]^-$	Com	$\text{XeF}_6 + [\text{XeF}_5]^+[\text{AsF}_6]^-$		723
2XeF_2	$[\text{AsF}_6]^-$		$[\text{AsF}_6]^-$		
$[\text{XeF}_5]^+$	$[\text{SbF}_6]^-$	Lewis	$\text{XeF}_6 + [\text{H}_3\text{O}]^+[\text{SbF}_6]^-$	α - and β -modification of $[\text{XeO}_2\text{F}]^+[\text{SbF}_6]^-$	694
$[\text{XeF}_5]^+$	$[\text{SbF}_6]^-$	Other	$[\text{XeF}_5]^+[\text{SbF}_6]^- + [\text{NO}_2]^+[\text{SbF}_6]^-$		43
$[\text{XeF}_5]^+$	$[\text{AuF}_6]^-$	Lewis	$2\text{XeF}_6 + \text{AuF}_5$		724
$[\text{Xe}_2\text{F}_{11}]^+$	$[\text{OsO}_3\text{F}_2]^-$	Lewis	$\text{XeF}_6 + (\text{OsO}_3\text{F}_2)_\infty$		690
$[\text{XeO}_2\text{F}]^+[\text{F}(\text{XeO}_2\text{F})_2]^+$	$[\text{MF}_6]^-$ (M = As, Sb)	Lewis	$\text{XeO}_2\text{F}_2 + \text{MF}_5$ (M = As, Sb)		694

^a Classification according to the introduction (Table 2): Lewis = Lewis acid halogen bond heterolysis, Com = complexation reaction, other = all other reactions not classified.

Abbreviations

CN	Coordination number
DFT	Density functional theory
ϵ_r	Relative permittivity of a solvent (static dielectric constant)
FLP	Frustrated Lewis pairs
IL	Ionic liquid
n.a.	Not available
rPBC	Reactive p-block cations
WCA	Weakly coordinating anion
Ar^{CF_3}	3,5-(CF_3) ₂ C_6H_3
Ar^{Cl}	3,5- Cl_2 - C_6H_3
9BBN	9-Bora[3.3.1]bicyclononane
bipy	1,2-Bipyridine
BOX	Bis(oxazoline)
CatBH	Catecholborane/1,3,2-benzodioxaborole
COD	1,5-Cyclooctadiene
Cp	C_5H_5
Cp'	$\text{C}_5\text{Me}_4\text{H}$
Cp*	C_5Me_5
Cy	Cyclohexyl
DDP	2-(DIPP)amino-4-(Dipp)imino-2-pentene
Dipp	2,6- ⁱ Pr_2 - C_6H_3
DMAP	4-Dimethylaminopyridine
DMeOPrPE	1,2-(Bis(dimethoxypropyl)-phosphino)ethane
DMH	1,1- $\text{Me}_2\text{N}_2\text{H}_4$
Dmp	2,6-Dimethyl-phenyl
dmpe	1,2-Bis(dimethylphosphino)ethane
Do	Donor
DPE	1,2-Diphenylethane
dppe	1,2-Bis(diphenylphosphino)ethane
DTBMP	2,6-Di- <i>tert</i> -butyl-4-methylpyridine
dtbpy	4,4'-Di- <i>tert</i> -butyl-2,2'-bipyridyl
Et	Ethyl
Fc	Ferrocenyl
FP	$\text{CpFe}(\text{CO})_2$
FP'	$\text{Cp}'\text{Fe}(\text{CO})_2$
FP*	$\text{Cp}^*\text{Fe}(\text{CO})_2$
hppH	1,3,4,6,7,8-Hexahydro-2 <i>H</i> -pyrimido-[1,2- <i>a</i>]-pyrimidine
IMe	1,3-Bis(methyl)imidazol-2-ylidene
IMes	1,3-Bis(2,4,6-trimethylphenyl)imidazol-2-ylidene
ⁱ Pr	iso-Propyl
ⁱ Pr ₂ -ATI	<i>N,N'</i> -Diisopropylaminotroponimate
IPr	1,3-Bis(2,6-diisopropylphenyl)imidazol-2-ylidene
^t Bu	1,3-Bis(<i>tert</i> -butyl)imidazol-2-ylidene
Me	Methyl
Me ₄ -cyclam	<i>N,N',N'',N'''</i> -Tetramethyl-1,4,8,11-tetraazacyclo-tetradecane
Mes	2,4,6- $\text{Me}_3\text{C}_6\text{H}_2$
$\text{Me}_3\text{SiS}^p\text{Tol}$	1-SSiMe ₃ -4-Me- C_6H_4
Me ₃ -tacn	<i>N,N',N'''</i> -Trimethyl-1,4,7-triaza-cyclononane
nacnac	(NMesCMe) ₂ CH
NBD	2,5-Norbornadiene

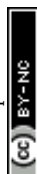
NPPh	2,5-Bis(2-pyridyl)-1-phenylphosphole
1-MIM	N-Methylimidazole
m-TP	meta-Terphenyl
OSSO	trans-1,2-Cyclooctanediyl-bridged[OSSO]-type bis(phenolate)
OR ^{PF}	–OC(CF ₃) ₃
OR ^{HT}	–OC(CH ₃)(CF ₃) ₂
OR ^{HF}	–OC(H)(CF ₃) ₂
OR ^{MeF}	–OC(CH ₃)(CF ₃) ₂
Ph	–C ₆ H ₅
Phen	1,10-Phenanthroline
4-Pic	4-Methylpyridine
Pip	Piperidyl
PMAF	Pentamethylazaferrocene
pmdta	N,N,N',N',N''-Pentamethyldiethylenetriamine
PNP	Bis(2- ⁱ Pr ₂ P ⁴ -Me-phenyl)amido
Py	Pyridine
Pytsi	C(SiMe ₃) ₂ SiMe ₂ (2-C ₅ H ₄ N)
p-Xyl	para-Xylene
R	Typical univalent organic residue
Salen	N,N'-Ethylenebis(2-hydroxyphenyl)imine
Salen ^{CF₃}	N,N'-Ethylenebis(2-hydroxy-2-(CF ₃) ₂ -ethyl)imine
Salomphen	N,N'-(4,5-Dimethyl)phenylene-bis(2-hydroxyphenyl)imine
Salpen	N,N'-Propylenebis(2-hydroxyphenyl)imine
Sch	Tridentate Schiff base
SubPc	Subphthalocyanine
taen	1,4- ⁱ Pr ₂ -1,4,7-Triaza-cyclononane
^t Bu	tert-Butyl
Tf	–SO ₂ CF ₃
THF	Tetrahydrofuran
timtmb ^{tBu}	1,3,5-{Tris(3-tert-butylimidazol-2-ylideno)methyl}-2,4,6-trimethylbenzene
TMM	η ⁴ -C(CH ₂) ₃
Tol	Toluene
Tipp	2,4,6- ⁱ Pr ₃ -C ₆ H ₂
X	Halogen

Acknowledgements

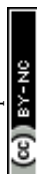
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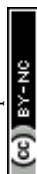
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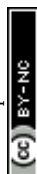
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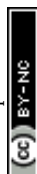
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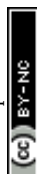
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