Metallomacrocycles: Supramolecular Chemistry with Hard and Soft Metal Cations in Action

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I. Introduction

Since the early work of Pedersen, Cram, and Lehn, the field of “supramolecular chemistry” or “host–guest chemistry” has grown enormously. Supramolecular chemistry can be defined as the chemistry beyond the covalent bond or the chemistry of associates with a well-defined structure. The structures of the associates are governed by relatively weak forces such as hydrogen bonds, ion–dipole and dipole–dipole interactions, van der Waals interactions, etc. Most of the work has concentrated on the design of selective receptors for either alkaline and alkaline earth cations. These cations are referred to as “hard” which means that they have a low polarizability. Transition metal cations are defined as “soft”, because they are relatively easy to polarize. Among the many applications of selective complexation of cations are chemical sensors, the selective removal of poisonous or radioactive metal cations from waste streams, membrane transport, immobilization of radioisotopes, and as phase-transfer catalysts.

Recently also host molecules have been designed in which one or more metal cations are incorporated to allow the selective binding of anions and nucleophilic groups of neutral molecules that can coordinate to the metal cations. Other areas of interest are model systems for (metallo)enzymes and the activation of small molecules like O₂, CO (vide infra), CO₂, and N₂. Generally organic multidentate ligands are designed in which one or more metal cations are complexed at well-defined positions. The field of homo- and hetero-dinuclear complexes of two transition metal cations has received considerable attention in order to achieve the above mentioned goals, which are often strongly related. Most of the work in this area has focused on model systems for hemocyanin and tyrosinase, both possessing two copper cations in the active site.

The present review exclusively deals with metallo-macrocycles which contain combinations of hard and soft metal cations. Metallomacrocycles are defined as metal complexes which are able to co-complex another metal cation. Many studies aim for the modulation of the redox properties of a complexed transition metal cation upon the co-complexation of a hard cation. The influence of co-complexation is not limited to complexed transition metal cations, also organic moieties that are redox active have been incorporated in ligands that can complex hard cations. Other areas of interest are allosteric effects, supramolecular aggregates, and bimetallic activation and catalysis.

In the first part hetero-di- and hetero-trinuclear complexes with bridging ligands between hard and soft cations are discussed. The second section deals with phthalocyanines, porphyrins, and related compounds, followed by the activation of carbon monoxide in hetero-dinuclear complexes. Bipyridine- and phenanthroline-containing crown ethers and ferrocenophanes and ruthenocenophanes will be discussed separately. The
Dr. Willem Verboom (1954) studied chemistry at the State University of Utrecht (The Netherlands) were he graduated in 1980 with Prof. dr. H. J. T. Bos on a thesis entitled: Thermal and photochemical rearrangements of $\gamma$-oxo-$\alpha,\beta$-unsaturated carboxamides and esters. In 1980 he joined the Laboratory of Organic Chemistry at the University of Twente where he is a senior staff member of Prof. dr. ir. D. N. Reinhoudt. At first he was involved in heterocyclic chemistry (tert-amino effect, mitomycins, etc.). At present his major research interest deals with supramolecular chemistry (synthesis of calixarenes, resorcinalarenes, (metallic)macrocycles, metallocloths, etc., and complexation studies with anions, cations, and neutral guest molecules). He is author of more than 135 scientific papers and review articles and three patents.

The last section contains miscellaneous ligands in which hard and soft metal cations are combined. The literature cited covers the period to the end of 1992.

II. Hetero-dinuclear and Hetero-trinuclear Complexes with Bridging Ligands between the Different Metal Cations

Only recently have ligands capable of positioning two or more very different metal cations, like transition metal and alkali or alkaline earth metal cations, in close proximity been described. A general approach to bringing two metal cations close together (intermetallic distance < 4 Å) is the use of a bridging ligand, which is also followed here to achieve the above mentioned goal. Ligands for the complexation of two different metal cations should possess cavities of different nature, such as tetradentate ligands for transition metal cations and crown ether type cavities for hard cations like alkali or alkaline earth metal cations.

Reinhoudt et al. have started an intensive program and synthesized a number of ligands and the corresponding complexes in which both the cavity for transition metal cations as well as the cavity for the alkali or alkaline earth metal cations are systematically varied. It was anticipated that co-complexation of a hard cation close to the complexed transition metal cation would change its (redox) properties. The complexes were studied by X-ray crystallography, polarography, cyclic voltammetry, coulometry, and EPR spectroscopy. The ultimate goal is to realize bimetallic (electro)catalysis with well-defined complexes.

The synthesis of complexes 1–5 (Chart 1) is schematically depicted in Scheme 1. The cyclization of a diamine and a dialdehyde was carried out by high dilution techniques in the presence of a barium salt, acting as a template. In case of an aromatic diamine the resulting barium complexes 1 and 2 could be isolated as orange to red crystalline materials in 53–82% yield. X-ray analysis of a number of barium complexes showed the barium cation complexed to the polyether cavity. In the case of an aliphatic diamine the resulting barium complex was not isolated but reacted in situ with nickel or copper acetate to give the dinuclear complexes 3 and
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4 as crystalline, strongly colored materials in 71–85% yield. A number of X-ray analyses have been carried out and revealed the phenolate anion as bridging ligand between the two different metal cations (an example is given in Figure 1).

Figure 1. X-ray structure of the nickel/barium complex 3-H$_2$O ($n = 1$). Only the oxygen atoms of the triflate anions are shown for clarity. (Reprinted from ref 23. Copyright 1989 American Chemical Society.)

Intermetallic distances are in the range of 3.63–3.69 Å. The nickel and copper complexes have square planar coordination as can be expected for salen- and salo-
phen-type tetradentate ligands. In the case of zinc a square pyramidal coordination was found with the salophen unit forming the ground plane and a DMF molecule at the apical position. The hard cation interacts with all oxygens of the polyether cavity and the coordination sphere is completed by counteranions and/or solvent molecules. In an analogous way the dinuclear complexes 5 with two (alkylated) oxime bonds were synthesized in 55–82% yield. X-ray analyses of a number of complexes showed the intermetallic distances were in the range of 3.69–3.73 Å. It was expected that the electron-donating properties of the nitrogen atoms would be changed by the oxygen substituent and hence the (redox) properties of the complexes.

The consequence of co-complexation of a hard cation to the transition metal complexes is an anodic shift of the reduction potential. Anodic shifts up to 300 mV were observed upon co-complexation of Ba2+. Smaller shifts (about 50 mV) were observed for the monovalent cations. Hard cations (Ba2+, K+, Na+, and Li+) complexed in the polyether cavity reduce the electron density at the phenolic oxygen atoms, and this lowers the electron-donating ability toward the transition metal ion. Consequently the transition metal ion is more positively charged, and therefore the reduction will occur at a less negative potential. The complexes of (alkylated) oximes 5 are reduced at less cathodic potentials than the complexes 3. The oxygen atom reduces the electron-donating properties of the nitrogen atom which renders the complexed transition metal cation more positively charged, and this leads to a reduction at a less negative potential. Cyclic voltammetry showed that the reduction–oxidation process is in general both electrochemically and chemically reversible. The latter observation is important for electrocatalysis, because the electrocatalyst should be stable after electrochemical reduction (activation) before reacting with a substrate. The association constants of the Li+ complexes are smaller than those of the complexes with the other hard cations \( K_{\text{assoc}} (\text{Na}^+, \text{K}^+, \text{or} \text{Ba}^{2+}) > 10^6 \text{ L mol}^{-1} \text{ in DMSO}) \).

The first example of electrocatalysis by a dinuclear complex is the conversion of benzyl bromide into phenylacetone in the presence of the nickel/barium complex 3 and acetic anhydride. This transformation does not occur in the presence of the corresponding nickel complex 3 in which only toluene and dibenzyl are formed. The reaction of the nickel/barium complex 3, benzyl bromide, and two electrons leads to an intermediate with a (relatively) stable nickel–carbon bond.

When the reaction depicted in Scheme 1 was performed with a dialdehyde that has a shorter chain between the two aryl rings, a \([2 + 2]\) instead of a \([1 + 1]\) macrocyclization occurred. This results in a macrocycle with three compartments for complexation of metal cations. The barium and cesium complexes 6 were obtained as orange crystalline compounds in 45–69% yield. According to the X-ray analysis of the barium complex 6 the barium is coordinated to all 10 of the oxygen atoms of the macrocycle, and this causes a folding of the macrocycle around barium. Consequently the two nearly planar salophen units are approximately parallel to each other. These barium and cesium complexes could be readily converted into their trinuclear complexes by reaction with 2 equiv of nickel or copper acetate. The dinickel/barium complex 6 (Figure 2) and the dicopper/barium complex 6 were analyzed by X-ray crystallography. The most interesting feature is the short intermetallic distance between the two transition metal cations \([\text{Ni}^{2+} – \text{Ni}^{2+} = 3.42 \text{ Å and Cu}^{2+} – \text{Cu}^{2+} = 3.50 \text{ Å}] \) without a bridging ligand. This very short distance results from the coordination of \( \text{Ba}^{2+} \) to the polyether cavity.

Polarography and cyclic voltammetry of the copper-containing trinuclear complexes clearly showed that the reduction process proceeds in two distinct one-electron steps. After the first one-electron reduction the redox properties of the other complexed transition metal cation is changed. This evidence suggests that the conformation of the complex in solution is similar to the conformation in the solid state.

EFR spectroscopy of the copper-containing complexes 6 (MX = Ba(CF3SO3)2; X = O and S) exhibits typical signals for an \( S = 1 \) system, which means that the copper cations are ferromagnetically coupled. From the spectra a Cu–Cu distance of 3.6 Å could be deduced, which is in reasonable agreement with the distance in the solid state. These measurements also support a folded structure with a cofacial alignment of the two salophen units.

Sessler et al. have reported the copper complex 7 (Chart 2) which potentially has a cavity for hard cations. Attempts to bind Na+ have met with little success which can be explained by the fact that one of the oxygen atoms \( \beta \) to a nitrogen atom is coordinated to the copper cation, as was observed in the X-ray structure of 7. This coordination might render the remaining cavity too small for co-complexation of a second cation (Figure 3).

Hetero-dinuclear complexes 8 (Chart 2) with two thiolate anions bridging the two different metal cations have been reported by Becher et al. They have used the same method as depicted in Scheme 1. Both \( \text{Ba}^{2+} \) and \( \text{Ti}^{4+} \) have been used as the hard cation.

Schepartz and McDevitt have reported the “self-assembly” of the ionophores 9 (Chart 2) by complexation of nickel(II) to two bidentate salicylaldimines. The

![Figure 2. X-ray structure of the dinickel/barium complex 6-H2O (X = O). (Reprinted from ref 27. Copyright 1991 American Chemical Society.)](image-url)
result of this complexation is the formation of a podand-type receptor for alkali cations. In their report no information about the structure of the nickel complexes was given which can be square planar (syn or anti) or tetrahedral. Gokel et al. showed that in toluene solution these complexes have a square planar geometry. It remained unclear whether a syn or anti coordination was obtained, but without complexation of a hard cation in the podand the anti stereochemistry is most likely. An X-ray analysis of the nickel/sodium picrate complex \( m = n = 0 \) was reported showing discrete dimers with coordinated picrate anion and one water molecule as integral parts (Figure 4). The nickel cations have an octahedral instead of square planar coordination as was also found in solutions of MeOH and CHCl₃.

Extraction experiments of alkali metal picrate salts from an aqueous phase into a chloroform phase containing the nickel complexes showed moderate to good efficiencies. Vapor pressure osmometry showed that the nickel complex \( m = 1, n = 0 \) in 0.1 M NaClO₄ solution in acetone are present as discrete Na₂Ni₂ complexes. In contrast to the more anodic reduction of the complexes 1–4 the Na₂Ni₂ complexes are reduced at more cathodic potentials than the corresponding nickel complexes. Since “self-assembly” is defined as the spontaneous formation of higher order structures, i.e. self-assembled monolayers, the above mentioned results are more appropriately described in terms of an allosteric effect. It is, however, most likely that upon co-complexation of a hard cation a conformational reorganization around the complexed cation occurs.

Kobule and Satoh have reported the synthesis of copper and zinc complexes of polyether bis(β-diketone) ligands (10, Chart 2) in 90% yield resulting in the formation of a polyether cavity for hard cations. They showed that the ligands by themselves are not able to extract alkali cations from an aqueous solution into chloroform. Reaction of the bis(β-diketone) ligands with copper(II) and zinc(II) leads to cyclic receptors for hard cations. The copper complex 10 \( n = 1 \) has a selectivity for Na⁺ over the other alkali cations, whereas the copper complex 10 \( n = 2 \) has a selectivity for K⁺ and Rb⁺ over Li⁺ and Na⁺. The complexation of copper or zinc is a good example of an allosteric effect.

III. Crown Ether-Substituted Phthalocyanines, Porphyrins, and Related Structures

Complexation of alkali and alkaline earth metal cations to phthalocyanines, porphyrins, and related square planar tetradeinate ligands with covalently attached crown ethers may lead to ordered supramolecular structures that exhibit interesting optical, electrical, and electrochemical properties.

A. Phthalocyanines

Nolte and co-workers have studied the electrical properties of the crowned phthalocyanines 11b–d (Chart 3) by performing impedance measurements as a function of co-complexation of alkaline and alkaline earth metal cations. The free ligand 11a \( n = 0–2 \) was synthesized in three steps starting from the appropriate benzocrown ethers in an overall yield of 12%. They observed that, by matching the size of the alkali metal
Chart 3

cations and the cavity of the crown ethers, complexes with a 1:1 stoichiometry (M\(\text{II}^+\) to crown ether) were obtained. Larger alkali metal cations yielded complexes with a 1:2 (M\(\text{II}^+\) to crown ether) stoichiometry. The 21-crown-7 rings always showed an M\(\text{II}^+\) to crown ether ratio of 1:1. These results are in agreement with those observed for "simple" crown ethers. With K\(^+\), Rb\(^+\), and Cs\(^+\) cofacially ordered supramolecular stacks were obtained with very different electrical properties compared to those of the crowned copper-phthalocyanines. The electrical conductivities \(\sigma\) (180 °C; \(\times 10^{-6}\) S m\(^{-1}\)) are 11c = 5 (p-type); 11c-K\(^+\) = 100 (n-type); 11c-Rb\(^+\) = 3000 (n-type); 11c-Cs\(^+\) = 700 (n-type). The co-complexation of Ba\(^{2+}\) to 11c leads to phthalocyanine networks with a much lower electrical conductivity \(\sigma\) of \(10^{-6}\) S m\(^{-1}\) (200 °C). A similar value was obtained for 11bBa\(^{2+}\).

UV-vis absorption and magnetic circular dichroism studies in chloroform/methanol mixtures of the "crowned-5" copper(II), nickel(II), zinc(II), cobalt(II), and free base phthalocyanines have been performed by Stillman and co-workers. Addition of excess potassium acetate leads to cofacially ordered dimerization. Upon dimerization the spectrum is blue-shifted which is associated with broader and decreased intensities. For the free base the Q-bands (\(\pi-\pi^*\) transition) at 660 and 700 nm shift to 640 nm (with shoulder) upon dimerization. The blue-shift is far less pronounced in the UV region. In the case of the copper(II), nickel(II), and cobalt(II) phthalocyanines the Q-band located at 670–680 nm shifts upon dimerization to 630–635 nm. Similar observations have been reported by Kobayashi and Nishiyama, by Kobayashi and Lever, and by Bekaroğlu and co-workers. On the basis of EPR spectroscopy using zero field splitting parameters a Cu–Cu distance of 4.1 Å could be deduced for 11b upon dimerization. In their studies Kobayashi and Lever showed that the cofacial dimer formation is a three-stage process. Dimerization occurs already with a K\(^+\) to phthalocyanine ratio of 1:2 (\(K\text{assoc} = 6 \times 10^9\) L\(^2\) mol\(^{-1}\) in CHCl\(_3\)). The first step occurs with 0–0.5 K\(^+\)/Pc; the second step with 0.5–1.5 K\(^+\)/Pc; and the third step with >1.5 K\(^+\)/Pc. Bekaroğlu and co-workers have studied the crown-5 free base 11a (\(n = 0\)) and copper complex 11b with small-angle X-ray diffraction, and

Chart 4
they found that the phthalocyanines are ordered in eclipsed stacks, which may provide an ion channel. Besides this stable crystalline form a meta-stable form with staggered stacks was also found. The crowded phthalocyanines are packed in a zigzag conformation with the crown ether rings parallel to each other and an angle between the crown ether ring and the phthalocyanine moiety of 30°. This packing results in a crown ether to crown ether distance of 4.2 Å and a separation of the phthalocyanine moieties by 3.4–3.5 Å. This distance is in good agreement with the van der Waals distance of 3.4 Å.

Insoluble polymer networks based on crown-6 phthalocyanines have been reported and their alkaline cation binding properties have been studied by solid/solution extraction experiments. The cobalt(II) derivatives show a large selectivity for Rb⁺ but unfortunately no explanation was given. However, it does not seem likely that binding properties of the crown-6 moieties are very different from one transition metal to the other. It is therefore possible that the observed selectivity is due to the specific structure of the polymer.

Phthalocyanines with benzo-15-crown-5 ethers attached via flexible spacers (12a and b, Chart 4) have been studied by Bekaroglu and co-workers. Co-complexation of K⁺ to the crowned phthalocyanine 12a does not lead to dimerization but to intramolecular sandwiching of the cation by the benzo-15-crown-5 rings. Upon co-complexation of K⁺, Na⁺, and Li⁺ no observable changes of the UV-vis spectrum occurred. This is in contrast to the blue-shifts observed for the crowded phthalocyanines discussed above, but consistent with the fact that sandwiching of K⁺ does not lead to phthalocyanine dimers. If only four instead of eight benzo-15-crown-5 moieties are present (12b) dimerization occurs upon co-complexation of K⁺, Rb⁺, and Cs⁺. The absorption at 670 nm of the monomer decreases in intensity, whereas the intensity of the absorption at 640 nm of the dimer increases. Liquid–liquid extraction experiments of the free base, the nickel complex, and copper complex show selectivity of K⁺ and Rb⁺ over Li⁺, Na⁺, and Cs⁺. The copper complex has a larger affinity for the alkali metal cations than the free base or the nickel complex.

The electronic absorption and magnetic circular dichroism spectra of a bisphthalocyanine connected “in-plane” by an 18-crown-6 ring (13, Chart 4) are not affected by complexation of alkaline or alkaline earth metal cations.

B. Porphyrins

The first iron(III) porphyrin with four 15-crown-5 rings at the meso positions (14e, M = Fe(III), Chart 5) was synthesized by Kobayashi and Osa starting from 4-formylbenzo-15-crown-5 and pyrrole in 6% yield. Small but significant changes in the absorption spectra were observed upon complexation of Na⁺ or K⁺, whereas Li⁺ and Cs⁺ have no effect. With F⁻ as the counterion high spin complexes were obtained, but CN⁻ as axial ligand gives a low-spin system. The redox potentials are hardly affected upon co-complexation of hard cations. Bogat-skii and Zhilina have reported the synthesis of meso-tetrakis(benzo-18-crown-6)porphyrin (14a) starting from 4-formylbenzo-18-crown-6 and 4-formylcatechol. Unfortunately no spectroscopic stud-

Chart 5

- R₁ = R₂ = R₃ = R₄ = 4-benzo-18-crown-6
- R₁ = R₂ = R₃ = R₄ = 4-benzo-15-crown-5
- R₁ = R₂ = phenyl, R₃ = R₄ = 4-benzo-15-crown-5
- R₁ = phenyl, R₂ = R₃ = R₄ = 4-benzo-15-crown-5
- R₁ = R₂ = R₃ = R₄ = 4-benzo-15-crown-5
- M = 2H, Fe(III), Zn(II), Cu(II), VO(V), Mg(II), Mn(III)

Chart 6

Related to the above-described porphyrins and phthalocyanines with crown ether moieties are the tetraaza-[14]annulene derivatives 15 and 16 (Chart 6). Kruse and Breitmaier reported the nickel(II) and

C. Related Structures
cobalt(II) complexes 16 together with the co-complexation of KI and NaI. Sakata and Annoura\textsuperscript{64} reported the ethyl derivative 15 and showed by EPR spectroscopy that cofacial dimerization occurs upon co-complexation of K\textsuperscript{+} or NH\textsubscript{4}\textsuperscript{+}, leading to a triplet state with axial symmetry. The co-complexation of Na\textsuperscript{+} in the crown ether rings has no effect on the UV-vis spectra, but the co-complexation of K\textsuperscript{+} or NH\textsubscript{4}\textsuperscript{+} has a profound effect. The hard cation induced dimers exhibit a reduced intensity of the absorption band in the 450–510-nm region, broadening of the absorptions, and a blue-shift of the intense band in the 375–430-nm region.
The azo-coupled di(tri)topic ligands 22-24 (Chart 9) were obtained by reacting the nickel or copper complex of bis(acetylametoneimine) with the diazonium compound derived from aminobenzo-crown ethers. Electrical conductivity measurements in CH₃Cl₂ showed cation selectivity of K⁺ > Na⁺ > NH₄⁺ and Ca²⁺ > Mg²⁺. The extraction efficiencies of the perchlorate salt from an aqueous layer into a CH₃Cl₂ layer are comparable to those of the “free” crown ethers, which means that the complexed transition metals hardly change the binding affinities of the crown ethers. The UV-vis and ¹H NMR spectra were not affected by co-complexation of hard cations in the crown ether rings.

Platinum complexes with benzocrown ether rings (25, Chart 10) have been synthesized by reacting PtCl₄L₂ (L = 4-cyanobenzo-15-crown-5) with hydrazine. The co-complexation of Na⁺ was deduced from IR spectroscopy and elemental analysis. Cyclopalladation of 4-(CH₂=NOH) or C(CH₃)=NOH benzocrown ether leads to the chloro-bridged dipalladium complexes 26. Mononuclear palladium complexes can be derived by reaction of 26 with PPh₃ or PMe₂Ph. Potentially ditopic copper(II) and cobalt(III) complexes (27 and 28) have been reported, but no studies of co-complexation of hard cations are available. Sessler et al. reported the synthesis of the dysprosium complex 29 starting from the appropriate dialdehyde and 4,5-crowned 1,2-benzenediamine. NMR spectroscopy revealed a change of the chemical shift of 1.89 ppm upon addition of 1 equiv of the dysprosium complex. The dimethyl ether analogue of the dysprosium complex induces a chemical shift of 1.08 ppm, an indication that, despite the 2⁺ charge, the crowned dysprosium complex is capable of complexing Na⁺.

**IV. Activation of Carbon Monoxide**

Bifunctional activation of carbon monoxide may be achieved by dinuclear complexes that can coordinate to both the carbon and the oxygen atom. Transition metal affinities for the carbon atom in carbon monoxide, whereas the oxygen atom is more likely to bind to alkali or alkaline earth metal cations.

Powell et al. have demonstrated that the co-complexation of Li⁺ in metalcrown ethers, in which the Li⁺ is also coordinated to one of the CO ligands (30-37, Chart 11), can activate this CO ligand toward nucleophilic attack. There are no reports on the reaction of RLi with cis-M(CO)₂(PR₃)₂ (M = Cr, Mo, W), but the metalcrown ethers 30-37 react with RLi leading to heterodinuclear complexes with an acyl or a benzoyl ligand which is both coordinated to the transition metal and to Li⁺. A systematic study showed that the reactivity is strongly dependent on the substituents of the phosphorus atoms. Stronger Li⁺ binding to the metalcrown ether leads to a larger reactivity. The metalcrown ether should coordinate to Li⁺ in such a way that the Li⁺ can also coordinate to the oxygen atom of CO. It was further observed that the donor sites in the metalcrown ether should be as basic (hard) as possible. The molybdenum complexes 38 also react with RLi (R = n-Bu, t-Bu, Et₂N, t-Pr₂N) to give complexes with a R⁻-C₅H₄N ligand of which the carbonyl carbon is coordinated to the molybdenum and the cyanobenzyl oxygen to the Li⁺. The Li⁺ is bound to the crown ether. The coordination of Li⁺ and the molybdenum has been proven by a number of X-ray
Figure 5. X-ray structure of 33 (X = O) reacted with PhLi. (Reprinted from ref 81c. Copyright 1990 Munksgaard.)

Two representative examples are shown in Figures 5 and 6.

Figure 6. X-ray structure of 32 [X-X = MeNCH2CH2NMe, A = CH2N(Me)(CH2)3N(Me)CH2] reacted with PhLi. (Reprinted from ref 81c. Copyright 1990 Munksgaard.)

Cobalt and rhodium complexes of the monoazacrown ether 39a with co-complexed Na⁺ or Li⁺ are able to
accelerate the hydroformylation of alkenes compared with the cobalt and rhodium complexes of 39a,\textsuperscript{82-84} For the analogous Fe(II) complexes 39b,\textsubscript{c} it has been shown that the association constant of 39b-Na\textsuperscript{+} is larger than that of 39c-Na\textsuperscript{+} due to coordination of the carbonyl oxygen atom of the acyl ligand in addition to the donor atoms of the crown ether. Besides a larger thermodynamic stability of 39b-Na\textsuperscript{+} also the kinetic stability is larger.

Rhodium(I) complexes with an intra-annular CO ligand containing also a diazacrown ether ring (40) have been reported by Carroy and Lehn,\textsuperscript{85} The co-complexation of hard cations to the diazacrown ring did not lead to a significant change of the \textsuperscript{13}CO signal in the NMR spectrum and also the vibrational frequency \nu-(CO) was unchanged. The CO ligand is not activated by co-complexation of hard cations which is possibly due to co-complexation of the hard cations “on top” instead of inside the ligand.

V. Bipyridine- and Phenanthroline-Containing Ligands

Rebek et al.\textsuperscript{86-88} have synthesized molecules 41 (Chart 12) capable of allosteric behavior due to both a 2,2'-bipyridyl and a crown ether cavity. The two sites, although separated and electronically “insulated”, do not behave independently. Chelation of a transition metal at the bipyridyl site forces restrictions on the conformational freedom of the macrocycle which is reflected in its selectivity for the complexation of alkaline metal cations. The importance of this type of compound for enzymatic catalysis and details on the complexation are given in a recent review article.\textsuperscript{89} Allosteric bircrown ether ligands containing a 2,2'-bipyridyl fragment have been developed by Beer et al.\textsuperscript{90,91} Chelation of a transition metal at the bipyridyl site forces the bipyridyl function toward coplanarity, restricting the conformational freedom of the two benzocrown ether moieties but favoring the formation of intramolecular sandwich complexes with spherical alkali metal cations (42, Chart 13). The results of this study have been summarized in a review on redox responsive macrocyclic receptor molecules.\textsuperscript{92} Recently Beer et al.\textsuperscript{93,94} also prepared alkynyl- (43) and vinyl-linked (44, 45) benzo- and azacrown ether bipyridyl complexes. Heteropolymetallic ruthenium(II)–sodium complexes, in which each crown ether binds one sodium cation, were isolated from complexation reactions of the Ru(II) complexes of 42–44 with excess amounts of sodium hexafluorophosphate. Fluorescence spectroscopy revealed the recognition of Li\textsuperscript{+}, Na\textsuperscript{+}, or Mg\textsuperscript{2+} by the vinyl-linked crown ether bipyridyl Ru(II) complexes. The metal-to-ligand charge transfer emission maxima for the complexes are shifted to significantly lower wavelengths and increase in intensity upon the addition of the cation. The Ru(II) complexes of 42–44 could be easily electropolymerized onto platinum and optically transparent conducting glass electrodes to form films.
Although electrochemical recognition by such modified electrodes has not been observed, such systems show promise as novel spectrochemical sensors. By using step-by-step lithiation of 4,4'-dimethyl-2,2'-bipyridine Beer et al. prepared in a one-pot synthesis the polytopic 2,2'-bipiridyl ligand 45 containing both a benzocrown ether and a ferrocene moiety. Preliminary coordination studies reveal that 45 forms a heteropoly-metallic ruthenium(II)–ferrocene–sodium cation complex, of which the photo- and electrochemical properties have not yet been published.

Another type of allosteric ionophores are compounds 46 (Chart 14) consisting of two substituted bipyridine units connected at the α-position with a (poly)-ethyleneoxy chain. These compounds can be prepared in 50–60% yield by treatment of mono(bromomethyl)bipyridine derivatives with ethylene glycol oligomers in the presence of potassium hydroxide in dry dioxane as a solvent. A K⁺/Na⁺ selectivity of about 10 was observed in transport experiments across CH₂Cl₂ as a liquid membrane using ionophore 46 in the presence of Cu⁺. 500-MHz ¹H NMR spectroscopy proved the formation of a cyclic pseudocrown ether structure upon the addition of Cu⁺ to ionophore 46, which may explain the transport selectivity found. Similar results were obtained with 47 in which the 2,2'-bipyridine moieties are connected with a diazacrown ether. Ionophores 47 selectively complex Cu⁺ and Zn²⁺ under simultaneous formation of a pseudocryptand which gives rise to selective complexes of alkaline metal cations.

The water-soluble ruthenium(2+) complexes 48 (Chart 14) have recently been described by Dürre et al. Addition of alkali and alkaline earth cations to a solution of the ruthenium complex in acetonitrile leads to an increase of both the fluorescence quantum yield and the lifetime of the excited state. Analysis of titration data showed that 1:1 complexes are formed, which was not expected. The ruthenium complex 48 (n = 1) has a selectivity of Ca²⁺ and Ba²⁺ over Li⁺, Na⁺, and Mg²⁺. The ruthenium complex 48 (n = 2) has a large selectivity of Ca²⁺ over Ba²⁺, Li⁺, Na⁺, and Mg²⁺. Wytko and Weiss reported the synthesis of the macrocyclic hard and soft ligand 49 containing an o-phenanthroline subunit and an azacrown ether. The biphenyl spacer ensures the rigidity of the molecular framework and, consequently, the high degree of preorganization of the ligand. The formation of copper(I)– and rhodium(I)–alkal complex is still under investigation. Recently the same group published the synthesis of a related molecule viz. a highly rigid phenanthroline-capped porphyrin which contains two different soft donor sites.

VI. Ferrocenophanes and Ruthenocenophanes

Macroyclic compounds containing a ferrocene moiety, so-called ferrocenophanes, have received considerable attention. Their ability to complex metal cations offers the possibility for the interaction between two
metal centers. In particular, such compounds yield *redox-active host molecules* that can be used for selective recognition of cations. As outlined in the Introduction, only systems having a cavity for the complexation of hard cations will be discussed. Several corresponding thiaferrocenophanes and related compounds have been described in the literature, which are very suitable for the complexation of heavy metal cations.108–108

Biernat and Wilczewska106 reported the reaction of 1,1′-dihydroxyferrocene (50) with dichloro ethers 51 in the presence of sodium hydride to give the polyoxaferrocenophanes 52 in 17–36% yield (Scheme 2). The same reaction was performed by Akabori et al.107 using in situ prepared 50 from 1,1′-diacetoxyferrocene (53) and potassium hydride as a base to afford compounds 52 in increased yields of 25–60%.

The yields are dependent on the size of the ring and decrease in the order n = 4 > 3 > 2 > 1 > 0. It has been postulated that the potassium cation acts as a template in the cyclization. These polyoxaferrocenophanes extract to some extent alkali and alkaline earth metal cations from aqueous solutions. The extraction is less efficient from the corresponding benzocrown ethers, indicating that the incorporation of the soft iron atom of the ferrocene unit into the macrocyclic polyether ring has a considerable influence on the complexing ability with hard alkali and alkaline earth metal cations. On the other hand, a significant selectivity of 52 (n = 3, 4) was found toward the thallium(I) ion. H NMR and Mössbauer spectroscopy of the complexes of 52 (n = 4) with LiSCN and KSCN suggest a certain interaction between the iron atom of the ferrocene unit and the complexed cation. However, an X-ray structure determination of the complex of 52 (n = 3) with NaSCN (Figure 7) revealed that the iron atom of the ferrocene does not take part directly in complex formation and that the sodium cation forms a contact ion pair with the nitrogen atom of the counteranion.108 In a similar reaction Akabori et al.106 also prepared the acyclic ferrocene-containing polyethers 54 in 53–85% yield. Compared with 52 they even exhibited less extraction of alkali metal cations although to some extent they extract alkaline earth cations. The extraction of monovalent cations by 54 (n = 3) increases in the order Na+ < Li+ < K+ < Cs+ < Tl+ < Ag+.

When the reaction of 1,1′-diacetoxyferrocene (53) with dichloro ethers 51 was carried out in the presence of 18-crown-6 as phase-transfer catalyst the 1,1′-bis-[α-chloropoly(ethylenoxy)]ferrocenes (55) were obtained in 26–54% yield together with small amounts of the polyoxaferrocenophanes 52.110 Reaction of 55 with sodium sulfide in DMF at reflux temperature gave polyoxathiaferrocenophanes 56 in 21–66% yield. Due to the presence of the soft sulfur atom, the extraction of alkali metal cations was lower than in the case of ferrocenophanes 52. However, compound 56 (n = 3) extracts efficiently thallium(I) cations. Lariat ferrocencrown ethers corresponding to 52 have been synthesized and these hardly extract alkali metal cations.111

Another class of ferrocenophanes are those compounds in which the polyether chain via a XCH2 bond is connected with the ferrocene moiety. Czech et al.112 investigated the reaction of 1,1′-bis-[α-hydroxyethyl]-ferrocenes (57) with dithiols 59 in CH2Cl2 in the presence of a catalytic amount of trifluoroacetic acid under conditions of high dilution (Scheme 3). Reaction of 57 with 2,2′-dimercaptopodiethyl ether 59 (n = 0) gave the mononuclear oxadithia[9]ferrocenophane 60 (n = 0) and dinuclear dioxa-thiatetra[9,9]ferrocenophanes 61 (n = 0), the latter being the major reaction product. When R = H in both cases mixtures of separable diastereomers were obtained. However, when the reaction was performed with dithiols 59 (n = 1, 2) having a longer ethylenoxy chain mononuclear 60 was the
predominant reaction product (e.g. \( R = H, n = 1, 2, 89\% \)), while only very low yields were isolated of 61,113,114 it was proven that the first step of the reaction of 57 with dithiols 59 is a very fast formation of the cyclic ethers 58. Starting from a mixture of diastereomeric 57 first a mixture of trans- and cis-58 was formed giving rise to rac- and meso-60, respectively. The structure of meso-60 \((n = 2)\) has been proven with X-ray crystallography.115

Recently, Petter et al.116 have discovered that this result is not limited to thioethers. Starting from 1,1'-bis(methoxymethyl)ferrocene (62) the corresponding polyoxaferrocenophanes 63 \((n = 2-5)\) were obtained in 14–31\% yield (Scheme 4).

Oxferrocenophanes 63 \((n = 2-4)\) were also prepared by Akabori et al.117 by reaction of 57 \((R = H)\) with the appropriate oligoethylene glycol ditosylates or the corresponding dihalides using potassium tert-butoxide as a base; besides a low yield of 63 \((3–13\%)\), very small amounts of the dinuclear product were also obtained. From extraction experiments it was found that oxiferrocenophanes 63 show little or no complexing ability toward alkali metal cations. This in contrast to thallium(I) and silver cations for which a high complexing ability was found. These results correspond with those obtained for the ferrocenophanes 52. It is noteworthy that compounds 63 are able to form a stable complex with silver without decomposition, whereas compounds 52 decomposed rapidly in the presence of silver ions. This indicates that the distance between the iron atoms of the ferrocene nucleus and the complexed silver cation controls the rate of oxidation of the ferrocene moiety.

In compounds like 63, the macrocycle is formed at the 1,1'-positions of the ferrocene ring. Izumi et al.118 have prepared ferrocenophanes 64 and 65 (Chart 15) which contain crown ether type units connected at the 1,2- or 1,3-positions of one of the cyclopentadienyl rings. Compounds 64 and 65 were obtained in 6–17\% yield by reaction of 1,2- or 1,3-bis(hydroxymethyl)ferrocene, respectively, with oligoethylene glycol ditosylates using potassium tert-butoxide as a base. In extraction experiments 64 and 65 also exhibit a high selectivity toward thallium cation (vide supra). Although the extraction of alkali metal cations is modest, 1,2-ferrocenophanes 64 show a higher complexing ability than 1,3-ferrocenophanes 65, even though the latter have a somewhat larger ring size. This may be explained by assuming that the proton at the 2-position of the cyclopentadienyl ring of 65 is sterically hindering in the complexation process.

Bartsch et al.119 investigated ferrocene crown compounds 69 and 71 in which the ferrocene nucleus bears one and two 18-crown-6 units, respectively. (Hydroxymethyl)ferrocene 66 was reacted with 3-mercaptopropionald (67) in the presence of trifluoroacetic acid to give 68 in 82\% yield which was cyclized with the ditosylate of pentaethylene glycol in the presence of potassium tert-butoxide to ferrocene crown 69 which was obtained in 20\% yield (Scheme 5). The corresponding ferrocene bis(oxo) 71 was prepared in 88\% yield by reaction of bis(hydroxymethyl)ferrocene (57, \( R = H \)) with 2 equiv of (mercaptomethyl)-18-crown-6 (70). Extraction experiments with KSCN showed a potassium loading of 44\% and 128\% for 69 and 71,
respectively, indicating that 71 adopts conformations in which the two crown ether units act independently rather than to form sandwich complexes. For the ferrocene bis-crown 71 a selectivity was found in the order $K^+ > Rb^+ > Na^+$ and $Li^+$ being undetectable.

A third type of ferrocenophanes comprises compounds in which the ferrocene unit is connected via a carbonyl moiety to the polyether chain. Oepen and Vögtle\textsuperscript{120} reported that in the presence of pyridine reaction of 1,1'-bis(chlorocarboxyl)ferrocene (72) with diols 73 ($X=(\text{CH}_2\text{OCH}_2)_n$; $n=0-3$) in benzene led to the formation of oxaferrrocenophane 75a in 29–36\% yield (Scheme 6). Izumi et al.\textsuperscript{121} reinvestigated this reaction in the absence of pyridine and found in all cases in addition to 75a the 1,1'-ferrocenediyl diolcohol derivatives 74a as the major reaction product. Similarly, however, in the presence of triethylamine, compounds 74b–e (2–9\% yield) and 75b–e (19–67\% yield) were prepared. The dinuclear ferrocenophanes 76a–e were obtained by reaction of 74a–e with an additional equivalent of 72 in yields of 23–29\% (a,d) and 5\% (b,c,e). Ferrocenophanes 75a and 76a showed low complexing ability with alkali, alkaline earth, and transition metal cations without any difference in the extractability, analogously as found for 52 (vide supra).

Dinuclear ferrocenophanes 76c and 76e containing an additional soft donor atom exhibited high complexing ability toward silver and mercury (under acidic conditions) cations without decomposition of the ferrocene unit. In contrast to many other ferrocenophanes, both 75 and 76 showed a poor complexing ability toward thallium(I) cation probably due to the incompatability of the guest and the hole size of the host. A series of pyridine-containing ferrocenophanes 79 and the corresponding ruthenophanes 81, both with different sizes of the cavity, have been prepared in the same way as described above (Scheme 7) and their complexation behavior studied.\textsuperscript{122} The pyridinometallocenophanes 79 and 81 showed a similar low extraction of alkali and alkaline earth metal cations and a high extractability toward heavy-metal cations, in particular for silver, indium ($\text{In}^{3+}$) and zirconium ($\text{Zr}^{4+}$) cations. In general, ferrocenophanes 79 extract alkali and alkaline earth cations better than ruthenocenophanes 81.

Ferrocenocryptands 82 (Chart 16) have been studied by Hall et al.\textsuperscript{92,123} Extraction studies with alkali and alkaline earth metal cations indicated that 82 shows a specificity which follows the order $Ca^{2+} > K^+ > Mg^{2+} > Na^+ > K^+$. The interaction of 82 ($n=1$) with a range

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{chart16}
\caption{Chart 16}
\end{figure}
of divalent cations has been examined by $^{13}$C NMR spectroscopy, and this revealed the formation of 1:1 and 2:1 (host/guest) complexes in a ratio dependent on the conditions. Several 1:1 complexes (with Mg$^{2+}$, Ca$^{2+}$, Sr$^{2+}$, Ba$^{2+}$, Zn$^{2+}$) have been isolated and characterized confirming the conclusions reached by $^{13}$C NMR spectroscopy. Hall et al. also reported the first solid-state crystal structure of an asymmetric ferrocene containing cryptand [82 (n = 0)]. Recently Gokel et al. investigated the behavior of ferrocenylidimethyl-2,2'-cryptand 83 by reduction of 82 (n = 1) with LiAlH$_4$. The crystal structure of 83 (n = 1)-H$_2$O (Figure 8) contrasts sharply with that of 82 (n = 1)-H$_2$O$_2$ in that in the former two hydrogen bonds hold a single water molecule to the external face of the diaza-18-crown-6 subunit, whereas two water molecules are present in the structure of 82. In the latter one, water is coordinated to a carbonyl group and the other appears to fill a molecular void outside the ring. Compound 83 exhibits an electrochemical behavior that is strongly dependent on alkali and alkaline earth metal cations present. This is due to a combination of nitrogen atom basicity and greater conformational mobility of 83 compared with 82 (n = 1). Cyclic voltammetry of 83-H$_2$O in the presence of e.g. sodium perchlorate revealed that the oxidation of the Na$^+$ complex takes place at more positive potentials than the oxidation of the free ligand. From this it can be concluded that the ferricinium (oxidized) form of the complex is destabilized by repulsive electrostatic interactions with the coordinated cation. The X-ray analysis of 83 (n = 1)-NaClO$_4$ showed that the Na$^+$ is coordinated to the donor atoms of the crown ether at a distance of 4.39 Å from the Fe atom (Figure 9).

Beer et al. have extensively studied the electrochemical recognition of charged species by redox-active ferrocene or ruthenocene containing macrocycles either by “through space” electrostatic perturbation and/or via various conjugated bond linkages linking the heteroatoms of the ionophore to the redox center. A combination of different factors contribute to the experimentally observed magnitude and type (one or two waves) of the shift in the respective redox couples of these host systems produced by charged-guest binding. These include (i) the redox-active host–guest complementarity, (ii) the polarizing power of the guest, (iii) the proximity of the host binding site to the redox center, and (iv) the nature of the bond linkage from the host binding site to the redox center. The results of their investigations and those of a few other groups have recently been reviewed and consequently are not incorporated in this review. Recently Beer et al. also reported redox-responsive receptors 86 in
which a ferrocene moiety is linked to a benzo-15-crown-5
cation binding unit via a conjugated Schiff base bond.
Compound 86 was prepared in 55% yield by reaction of
4-ferrocenylaniline 84 and 4-formylbenzo-15-crown-5
(85) (Scheme 8). With the use of cyclic voltammetry
the reversible redox couple of 86 shifts anodically upon
the addition of Na⁺, K⁺, or Mg²⁺ cations. The magnitudes
of these shifts reflect the different polarizabilities
of the coordinated metal cations. Mg²⁺ with the
largest charge/radius ratio gives the greatest anodic shift
and K⁺ the smallest.

In the framework of a study to develop a new class
of redox catalytic systems capable of promoting redox
reactions on a guest substrate via guest inclusion and
Lewis acid catalytic activation, compound 88 was
designed containing four benzo-15-crown-5 moieties
and eight ferrocenyl redox-active centers. Reaction
of resorcinol and 4-formylbenzo-15-crown-5 (85) under
acidic conditions gave resorcin[4]arene 87 (Chart 17)
which upon treatment with excess of (chlorocarbonyl)-
ferrocene afforded 88 in 22% overall yield. Preliminary
coordination studies with the sodium cation reveal that
one Na⁺ is bound in each of the respective benzo-15-
crown-5 moieties. Cyclic voltammetric results suggest
the ferrocene moieties present in 88 are oxidized in one
step.

VII. Miscellaneous

The [η⁶-phenylmethylsila-14-crown-5]chromium tricarbo
nyl complex 89 (Chart 18) was obtained by reacting
the corresponding silacrown ether with chromium
diacetylcarbonyl. Co-complexation of alkali metal cations
induced shifts in the ¹³C NMR absorptions of the
OCH₂ groups (Li⁺, Na⁺ > K⁺). The corresponding
[η⁶-phenyldimethoxymethylsila]chromium tricarbonyl
did not show shifts of the O¹⁴CH₂ absorptions
upon adding alkali metal cations. Cyclic voltammetry
in CH₂CN showed an irreversible oxidation–reduction
process (EC mechanism) leading to [(CH₂CN)₃Cr-
(CO)₅]⁺ and the silacrown ether. Shifts of the anodic
peak up to 110 mV were recorded after adding excess
of MPF₆ (Li⁺ > Na⁺ > K⁺). For the dimethoxy
derivative shifts of 20–25 mV were observed. The mono-
(chromium tricarbonyl) and bis(chromium tricarbonyl)
derivatives of dibenzo-18-crown-6 90 have been reported
by Pannell et al.¹³³ The introduction of the electron-
withdrawing chromium tricarbonyl groups decreases
the association constants of the complexes with Na⁺
and K⁺, as was determined by the picrate extraction
method. Unexpectedly, the selectivity of K⁺ over Na⁺
was reversed for the bis(chromium tricarbonyl) ana-
logue. Yellow chromium tricarbonyl complexes of
dibenzocrown ethers 91 have been obtained by reaction
of the benzocrown ethers with chromium hexacar-
bonyl.¹³⁴ The green NaSCN complex has been reported
[chromium tricarbonyl].

Balch et al.¹³⁵,¹³⁶ synthesized N,N'-bis(Ph₃PCH₂)-
diaza-18-crown-6 starting from diaza-18-crown-6. From
this ditopic receptor they prepared a number of hetero-
dinuclear complexes 92, which have a metal–metal bond,
except for the potassium/iridium complex. The intermetallic
distance determined by X-ray analyses (Figures 10 and 11)
are Tl-Ir 2.875 Å, Tl-Pt 2.911 Å, Pb-Ir 3.12 Å, Sn-Ir 2.92 Å,
and K-Ir 3.33 Å. The

Chart 18

Figure 10. X-ray structure of 92 (M₁ = Pt, X = Y = CN⁻; M₂ = Tl⁺). (Reprinted from ref 135. Copyright 1990
American Chemical Society.)
Figure 11. X-ray structure of 92 (M₁ = Ir; X = CO; Y = I⁻; M₂ = K⁺). (Reprinted from ref 136b. Copyright 1991 American Chemical Society.)

Chart 19

C₁₇

Chart 20

C₁₇

C₁₇

Chart 21

C₁₇

C₁₇

C₁₇

thallium/iridium complex 92 undergoes an oxidative addition with Cl₂ or I₂ leading to the hetero-dinuclear complex 93 with an X ligand at both apical positions.

The tetraacid precursor for the complexes 94 (Chart 19) was synthesized by reacting the tetraaza-crown ether with BrCH₂COO⁻Na⁺ in a mixture of CH₃CN and water. After complexation of a paramagnetic cation in the tetraacid compartment a shift reagent for hard cations was obtained. The dysprosium complex 94 induced relatively small NMR shifts (0.18-2.57 ppm) of the co-complexed alkali cations. The authors concluded that it is unlikely that this shift reagent is useful for in vivo NMR studies.¹³⁷

The hetero-tritopic receptors 95a-e (Chart 20) were synthesized by reacting 4-formylbenzo-15-crown-5 with the appropriate diamine.¹³⁸-¹⁴⁰ The corresponding hetero-metallic complexes with Cu⁺ or Ag⁺ and Na⁺ or K⁺ could be isolated. FAB-MS spectrometry and ¹³C NMR spectroscopy of the copper-containing complexes showed a Cu to Na ratio of 1:2 and a Cu to K ratio of 1:1. The X-ray analysis of [95b-Cu⁺-K⁺][PF₆]₂ showed that the copper is complexed in a distorted tetrahedral geometry with the potassium sandwiched by two crown ether rings. The Ag⁺-containing complexes showed for both Na⁺ and K⁺ a 1:2 ratio. Sandwitching of K⁺ is not observed possibly due to the larger steric demand of Ag⁺ compared to Cu⁺. The tetratopic receptor 96 was obtained by reacting tren with 4-formylbenzo-15-crown-5.¹⁴¹ The 96-CuPF₆(NaPF₆)₃ complex has been prepared.

The ditopic receptors 97 and 98 (Chart 21) were synthesized by reacting the appropriate monocrown ethers with 3,7-dithia-1,9-nonanediol and 3,6-dithia-1,8-octanediol, respectively.¹⁴²,¹⁴³ The hetero-dinuclear complex 97-NaSCN-Cu(SCN)₂ has been reported. The tritopic receptor 99 was obtained by reacting the corresponding monocrown-diol with 2,6-bis(chloromethyl)pyridine under high-dilution conditions in THF.¹⁴³,¹⁴⁴ The hetero-trinuclear complex 99-(KSCN)₂-Co(SCN)₂ has been isolated.

Crown ether receptors with dit(p²-cyclopentadienyl)-tungsten(IV) and -molybdenum(IV) as redox center
Metallomacrocycles

Chart 22

Chart 23

Chart 24

4-thiobenzo-15-crown-5, or the potassium salt of 4-carboxybenzo-15-crown-5 (leading to 101). The reversible one-electron oxidation in CH₃CN is shifted anodically (50–80 mV) upon the addition of Li⁺, Na⁺, or K⁺. The differences between the various alkali metal cations are not large.145,146

Jones et al.147,148 synthesized a number of Mo(III)-crown ether complexes (102–105, Chart 23) starting from Mo(NO)LHal₂ (L = tris(3,5-dimethylpyrazolyl)-hydroborate) and the appropriate ethylene glycols and amino-substituted benzocrown ethers. They observed electrochemically irreversible reduction/oxidation processes which were shifted anodically up to 320 mV upon co-complexation of Na⁺ or K⁺.

Oxidative addition of Pt(stilbene)(PPh₃)₃ to 4-iodobenzocrown ethers leads to trans-PtIr(PPh₃)₂ (106, Chart 24).134 Addition of 1 equiv of NaI did not show significant changes of the chemical shifts but the J(Pt-P) coupling constant decreased by 29–35 Hz. Addition of more equivalents of NaI did not lead to a further decrease indicating a 1:1 stoichiometry of the platinum/sodium complex.

VIII. Summary

It has been shown in this review that complexes with both hard and soft metal cations can be used for bimetallic catalysis and activation, the formation of supramolecular aggregates, allosteric effects, and changing the redox properties of the transition metal cation. The molecular structures used to achieve these goals reflect the variety of approaches used. Although in some cases potential applications have been suggested, until now they have not evolved into prototypes. The authors feel that supramolecular structures with several different metal cations at well-defined places will be the major challenge for the next decade.

IX. References

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