

CAPVT VI

KOMPLEKSI

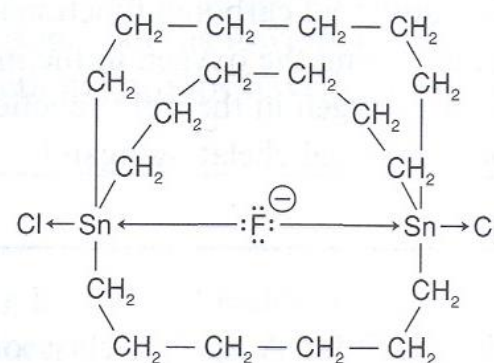
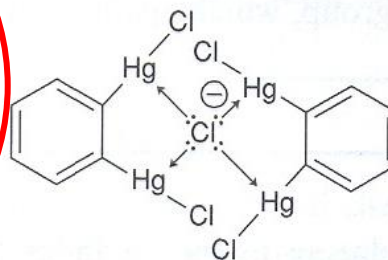
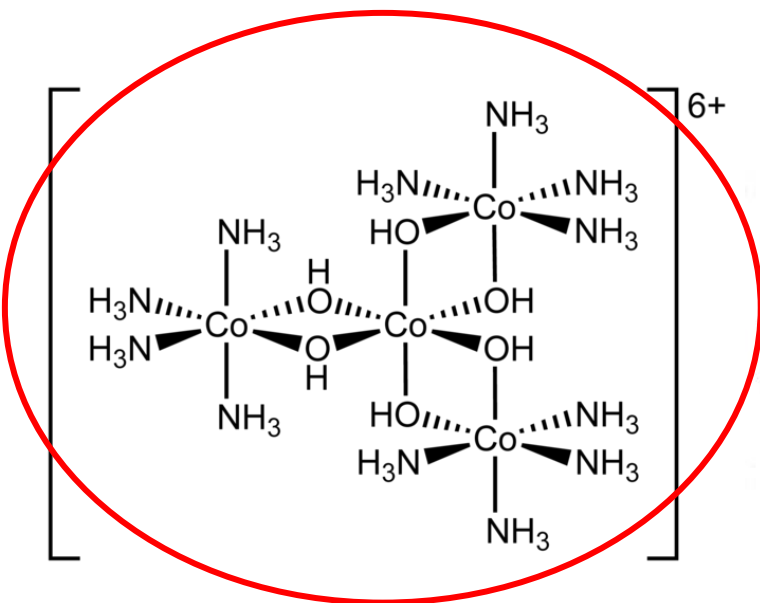
Što je kompleks?

A molecular entity formed by loose association involving two or more component molecular entities (ionic or uncharged), or the corresponding chemical species. The bonding between the components is normally weaker than in a covalent bond. The term has also been used with a variety of shades of meaning in different contexts: it is therefore best avoided when a more explicit alternative is applicable. In inorganic chemistry the term 'coordination entity' is recommended instead of 'complex'.

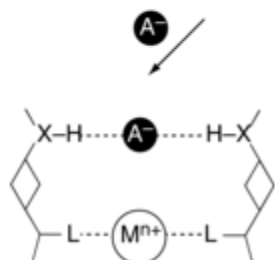
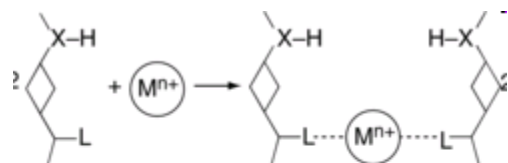
IUPAC

Spoj kiseline i baze

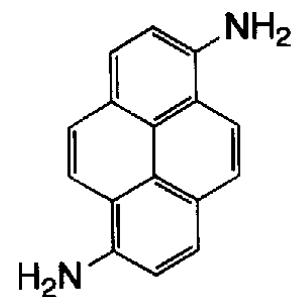
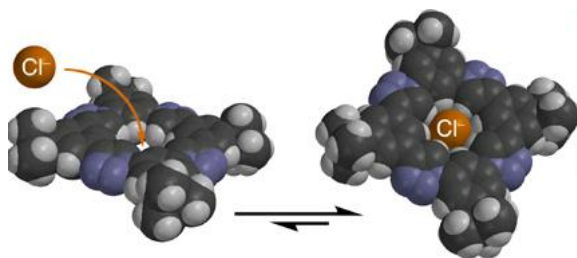
JA



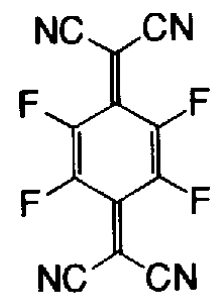
Lewisove



Brønstedove



DAP



F₄TCNQ

Usanovičeve

Koordinacijski spojevi

- Centralni atom – Lewisova kiselina
- Ligandi – Lewisove baze
 - Monodentatni, polidentatni, ambidentatni, premoščujući, kelatirajući

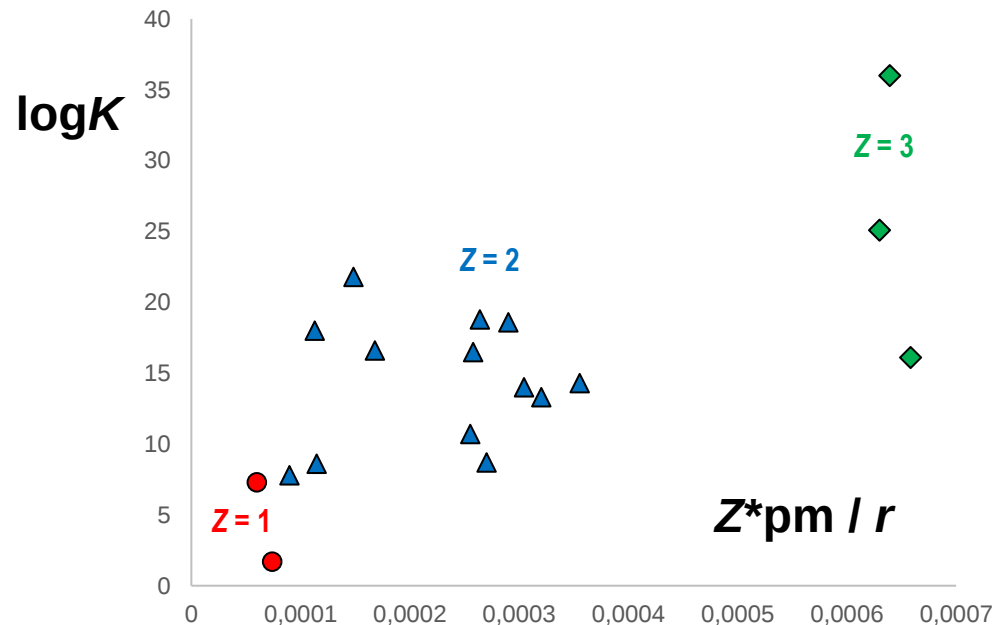
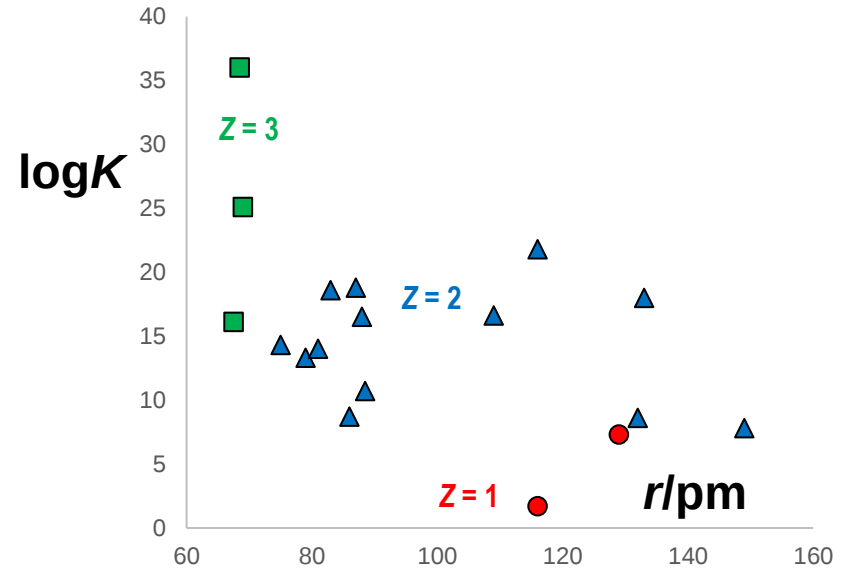
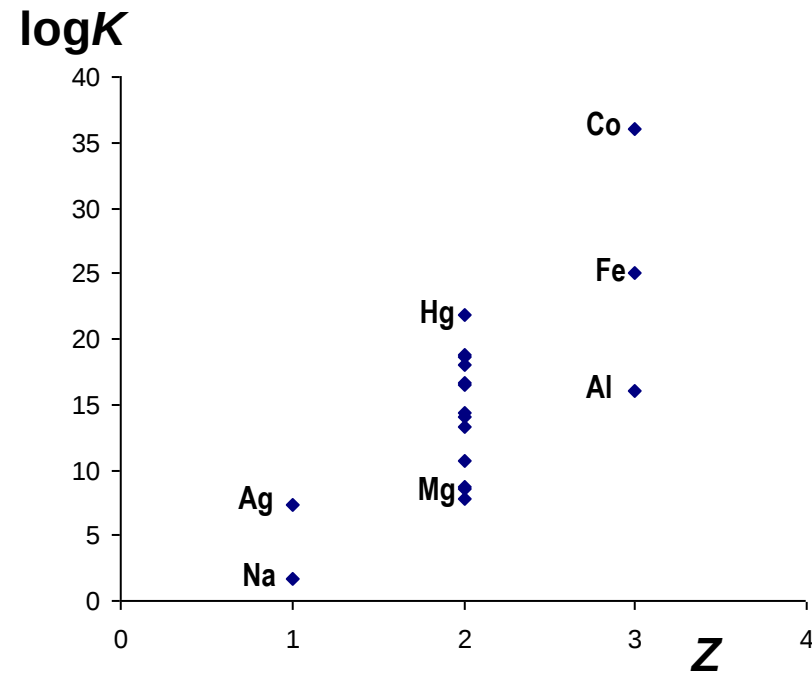
Stabilnost (*konstanta stabilnosti*)

- Ovisi o kationu i ligandu
 - Za isti ligand raste s Z^*/r (za atome pseudoplemenitoplinovske konfiguracije)
 - Irving–Williamsov niz stabilnost kompleksa za dani ligand dvovalentnog metala prve prijelazne serije raste do bakra →
$$\text{Mn(II)} < \text{Fe(II)} < \text{Co(II)} < \text{Ni(II)} < \text{Cu(II)} > \text{Zn(II)}$$
 - Za isti ligand atom s većom energijom ionizacije čini stabilniji kompleks

Konstanta stabilnosti

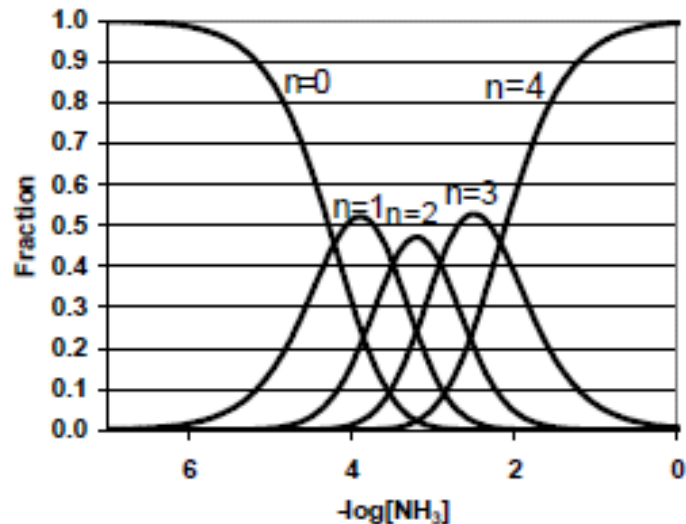
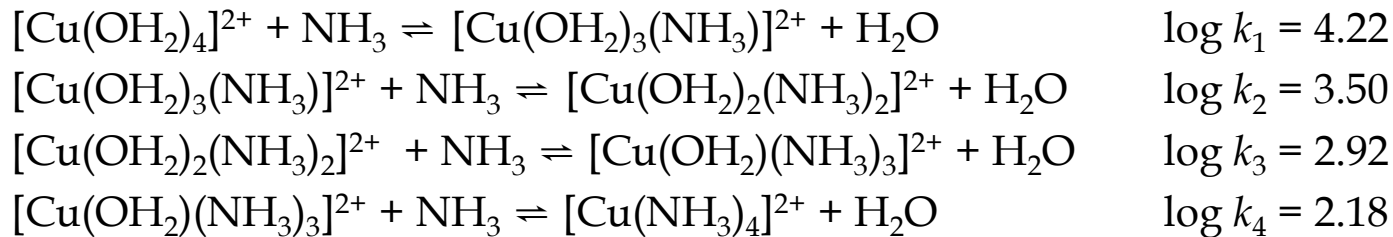
1. Ukupna (kumulativna)

Kompleksi s EDTA



2. Stupnjevita

Zamjena liganada jednog za drugim



Komponente stupnjevutih konstanti

Bjerrum (J.), 1941.

$$T_{n,n+1} = \log \frac{k_n}{k_{n+1}} = S_{n,n+1} + L_{n,n+1}$$

$$L_{n,n+1} = E_{n,n+1} + R_{n,n+1}$$

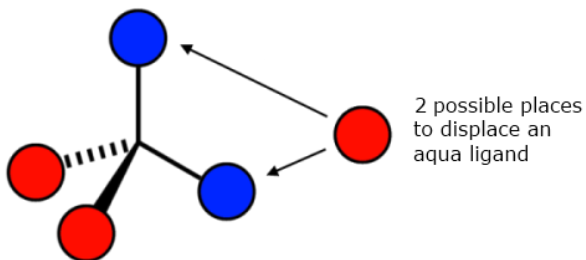
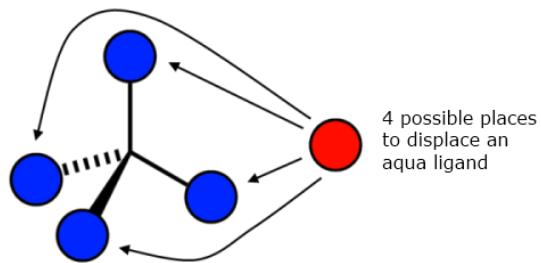
S – Statistički doprinos.

E – Elektrizitatski doprinos

R – Ostali doprinosi

$$\Rightarrow k_n = k_{Sn} k_{En} k_{Rn}$$

$$S_{n,n+1}$$



$$\frac{k_n}{k_{n+1}} = \frac{N - n - 1}{N - n} \frac{n + 1}{n}$$

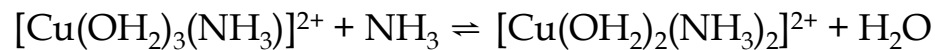
$$\frac{k_n}{k_{Sn}} = \sqrt[N]{\prod_{j=1}^N k_j}$$

N	k_{Sn}^{-1}				
1	1				
2	1/2	2			
3	1/3	1	3		
4	1/4	2/3	3/2	4	

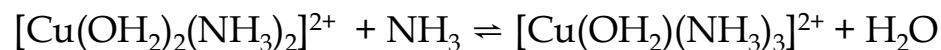
Za $[\text{Cu}(\text{NH}_3)_n]^{2+}$



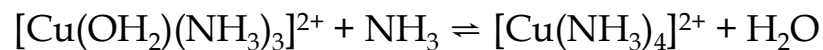
$$\log k_1 = 4.22$$



$$\log k_2 = 3.50$$

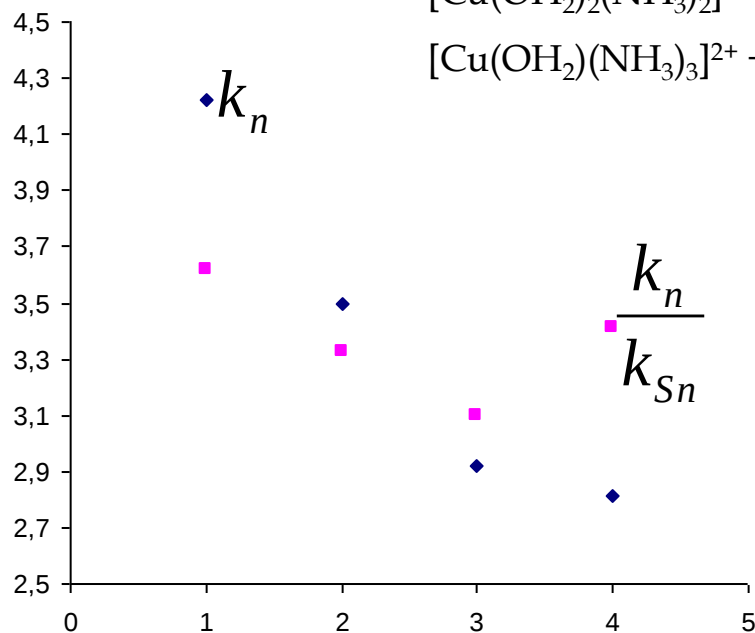


$$\log k_3 = 2.92$$

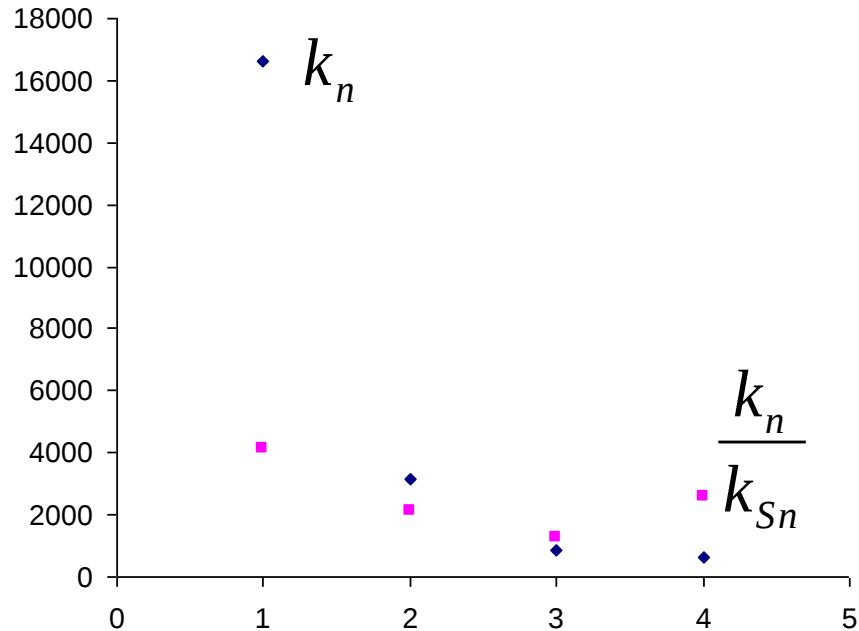


$$\log k_4 = 2.18$$

$\log k_n$



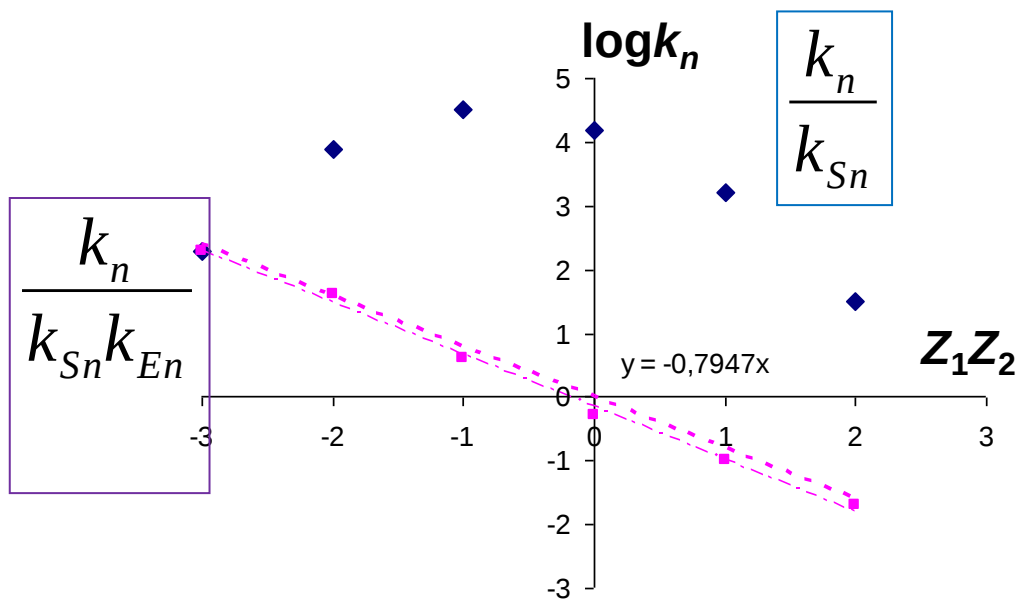
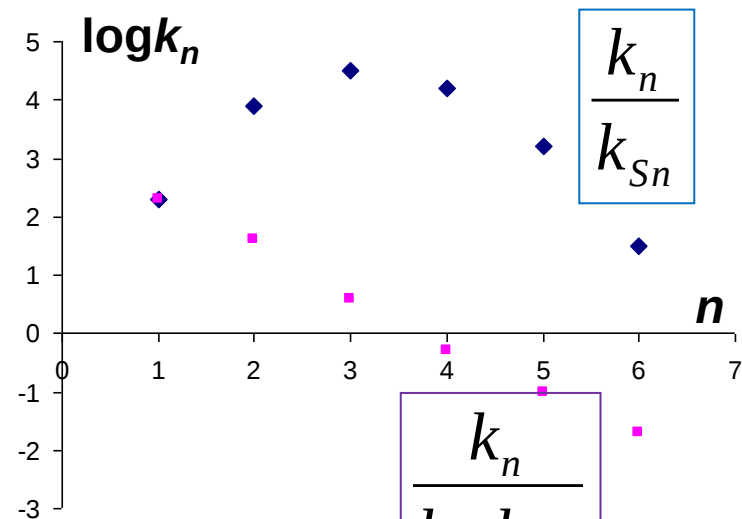
k_n



$$E_{n,n+1}$$

$$E_{n,n+1} = f \frac{\varphi_{n,n+1}}{kT}$$

$$\varphi_{n,n+1} = \frac{Z_1 Z_2}{\varepsilon r}$$



$$R_{n,n+1}$$

Ukazuje na 'kemijske' uzroke razlika stabilnosti pojedinih kompleksa:

- nagla promjena u R – mijenja se koordinacijski poliedar
- periodična promjena u R – *trans* utjecaj
- ...

Stabilnost i HSAB

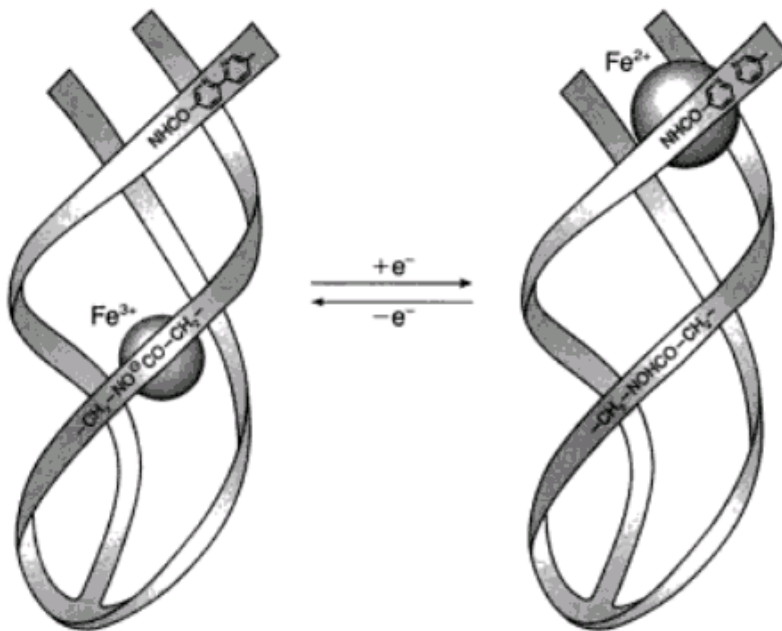
- Reakcija dvostruke izmjene tipa
$$\text{HA:SB} + \text{SA:HB} \rightarrow \text{HA:HB} + \text{SA:SB}$$

najčešće je termodinamički povoljna

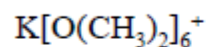
Tvrde i meke kiseline i baze u PSE

H 2.2												Borderline bases			Hard bases		He			
Li 0.98	Be 1.57	Hard acids					Hard acids					B 2.04	C 2.55	N 3.04	O 3.44	F 3.98	Ne			
Na 0.93	Mg 1.31											Borderline acids			Al 1.61	Si 1.90	P 2.19	S 2.58	Cl 3.16	Ar
K 0.82	Ca 1.00	Sc 1.36	Ti 1.54	V 1.63	Cr 1.66	Mn 1.55	Fe(+3) 1.83 (+2)	Co(+3) 1.88 (+2)	Ni 1.91	Cu(+1) 2.0	Zn 1.65	Ga 1.81	Ge 2.01	As 2.18	Se 2.55	Br 2.96	Kr 3.0			
Rb 0.82	Sr 0.95	Y 1.22	Zr 1.33	Nb 1.6	Mo 2.16?	Tc 1.9?	Ru 2.2	Rh (+3) 2.28 (+1)	Pd 2.20	Ag 1.93	Cd 1.69	In (+3) 1.78 (+1)	Sn (+4) 1.96 (+2)	Sb 2.05	Te 2.1	I 2.66	Xe 2.6			
Cs 0.79	Ba 0.89	Lu 1.27	Hf 1.3	Ta 1.5	W 2.36?	Re 1.9?	Os 2.2	Ir (+3) 2.2 (+1)	Pt 2.28	Au 2.54	Hg 2.0	Tl (+1) 1.60 (+3) 2.04	Pb (+2) 1.87 (+4) 2.33	Bi 2.02						
Fr 0.7	Ra 0.9											Soft acids								
												Borderline acids								
		La 1.10	Ce 1.12	Pr 1.13	Nd 1.14	Pm	Sm 1.17	Eu	Gd 1.20	Tb	Dy 1.22	Ho 1.25	Er 1.24	Tm 1.25	Yb					
		Ac 1.1	Th 1.3	Pa 1.5	U 1.38	Np 1.36	Pu 1.28	Am 1.3	Cm 1.3	Bk 1.3	Cf 1.3	Es 1.3	Fm 1.3	Md 1.3	No 1.3					
																	Hard acids			

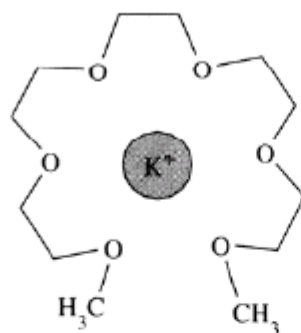
- Tvrđi kationi preferiraju koordinaciju tvrdim bazama (halogeni, O, (N)...)
- Mekani kationi preferiraju koordinaciju mekanim bazama (S, Se, P...)



Kelatni i makrociklički učinak

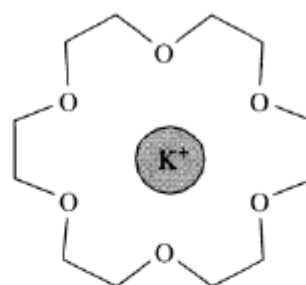


Very low stability



$[\text{K}^+ \cdot 3.11]$

$\text{Log } K (\text{MeOH}, 25^\circ\text{C})$ 2.0



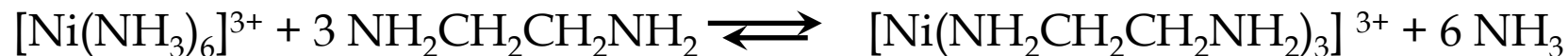
$[\text{K}^+ \cdot 3.6]$

6.1

Complex	$\Delta G^\circ (\text{J mol}^{-1})$	$\Delta H^\circ (\text{J mol}^{-1})$	$\Delta S^\circ (\text{J K}^{-1} \text{mol}^{-1})$
$[\text{K}^+ \subset 3.11]$	-11 368	-36 400	-84
$[\text{K}^+ \subset 3.6]$	-34 842	-56 000	-71

Kelatni učinak

- Spojevi s kelatirajućim ligandima stabilniji od onih s monodentatnima
 - Entropijska stabilizacija



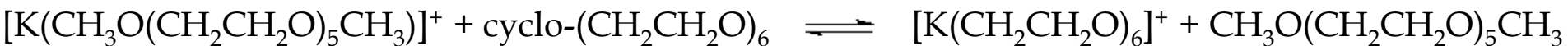
$$\Delta_r G = -67 \text{ kJ/mol}$$

$$\Delta_r H = -13 \text{ kJ/mol}$$

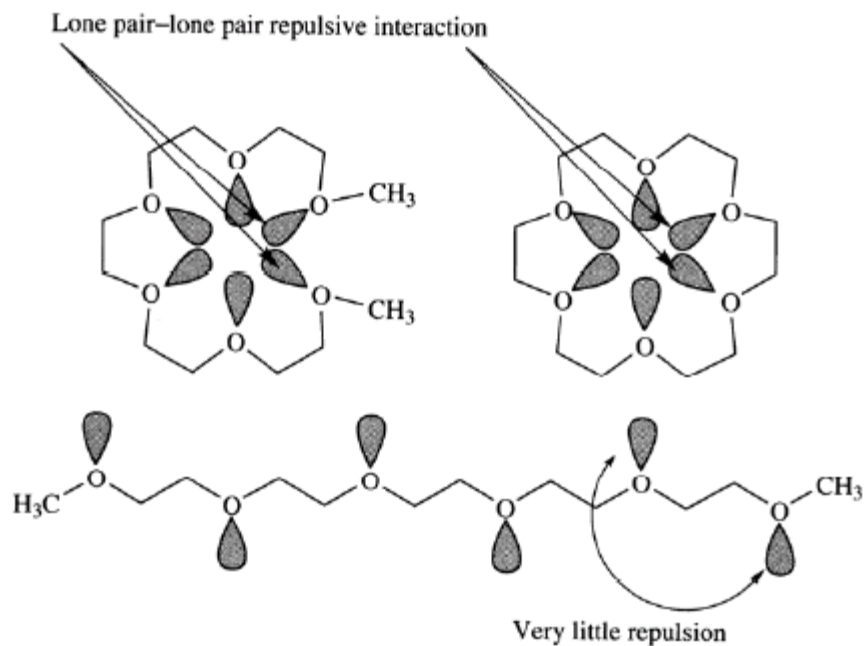
$$T \Delta_r S = 54 \text{ kJ/mol}$$

Makrociklički učinak

- Spojevi s makrocikličkim ligandima stabilniji od onih s kelatirajućim
 - Entalpijska i entropijska stabilizacija

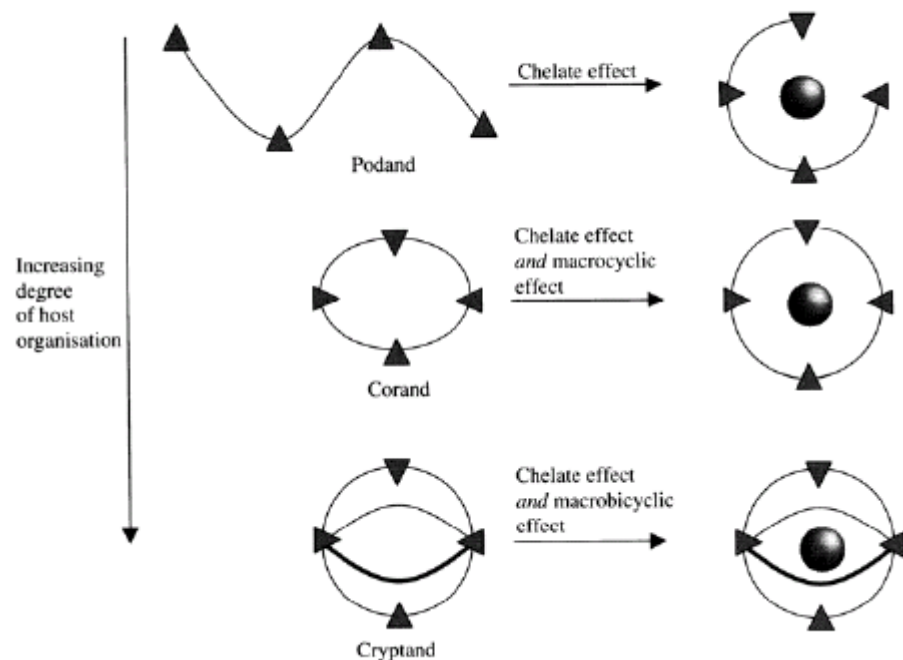


$$K \approx 10^4$$



Konformacija kelatirajućeg liganda potrebna za koordinaciju nužno uključuje približavanje veznih mjesta (donirajući elektronski parovi) – entalpijski nepovoljno)

Makrociklički ligand unaprijed ‘ukočen’ u potrebnoj konformaciji – nema entalpijske ‘kazne’ promjene konformacije



Određivanje stupnjevitih konstanti

Bodländer, Morse, Bjerrum (N.), Abegg, Jacques,
određivanje srednjeg koordinacijskog broja (do 1914.)

$$\bar{n} = \frac{c(A) - [A]}{c(M)} = \frac{[MA] + 2[MA_2] + 3[MA_3] + \dots}{[M] + [MA] + [MA_2] + \dots} = \frac{\sum_{n=1}^N n k_n [A]^n}{1 + \sum_{n=1}^N k_n [A]^n} = \sum_{n=1}^N (n - \bar{n}) k_n [A]^n$$