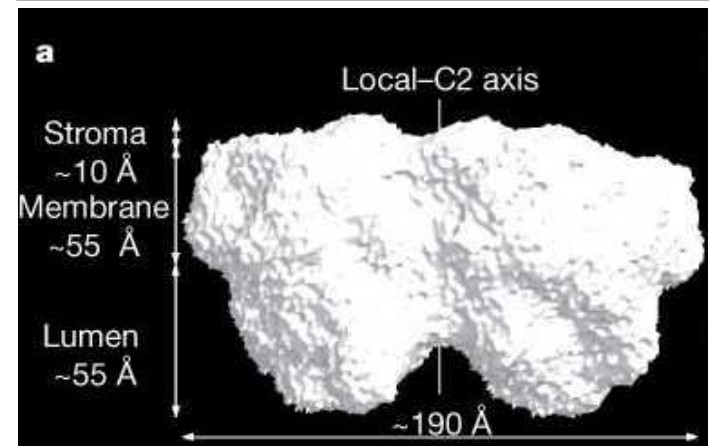
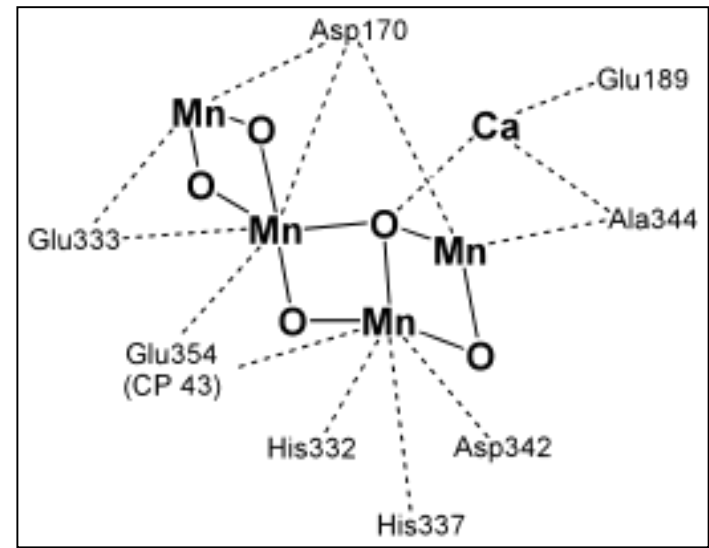


Bioinorganic Chemistry

Content

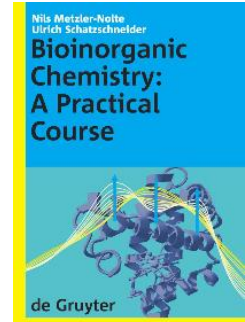
1. What is bioinorganic chemistry?
2. Evolution of elements
3. Elements and molecules of life
4. Phylogeny
5. Metals in biochemistry
6. Ligands in biochemistry
7. Principles of coordination chemistry
8. Properties of bio molecules
9. Biochemistry of main group elements
10. Biochemistry of transition metals
11. Biochemistry of lanthanides and actinides
12. Modell complexes
13. Analytical methods in bioinorganic
14. Applications areas of bioinorganic chemistry



"Simplicity is the ultimate sophistication" Leonardo Da Vinci

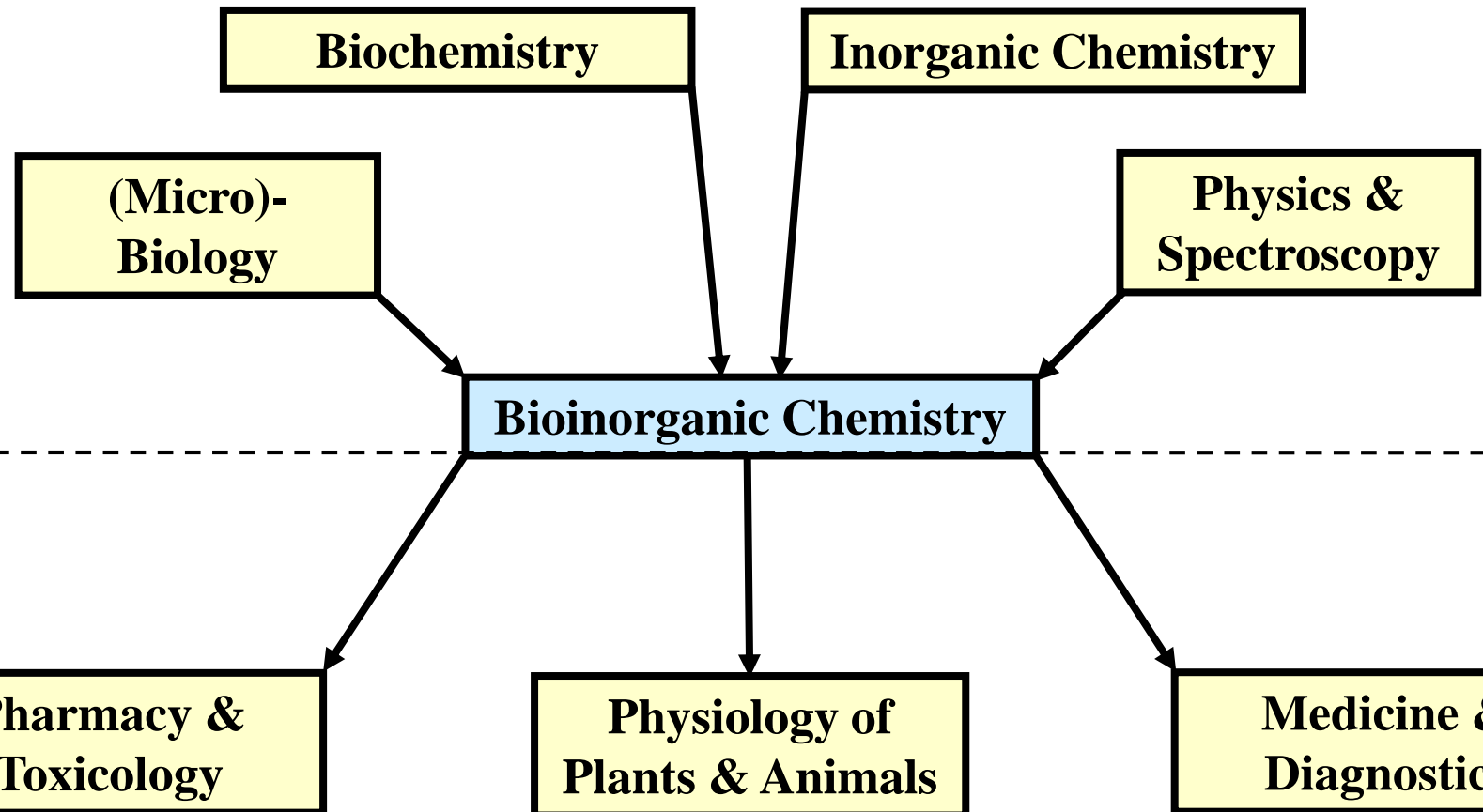
Literature

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1. What is Bioinorganic Chemistry?

A Highly Interdisciplinary Science at the Verge of Biology, Chemistry, Physics, and Medicine



2. Evolution of the Elements

Most Abundant Elements in the Universe according to Atom Number Fractions are

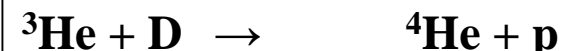
1. Hydrogen	88.6%	Reactive: H ₂ , H ₂ O, CH ₄ → biology
2. Helium	11.3%	Noble gas → atmospheres
3. Oxygen	0.063%	Reactive: O ₂ , H ₂ O → biology
4. Carbon	0.035%	Reactive: CO ₂ , CH ₄ → biology
5. Nitrogen	0.011%	Reactive: N ₂ , NH ₃ → biology
6. Neon	0.010%	Noble gas → atmospheres
7. Magnesium	0.0032 %	Oxides → planetary crusts
8. Silicon	0.0029 %	Silicates → planetary crusts

Elements from Lithium onwards, which are regarded as metals by astronomers, contribute to only about 0.1%

Whilst hydrogen, helium, and traces of lithium have been formed during the big bang, all other heavier elements up to iron had to be generated by fission within the stars. Even heavier elements were formed only in supernovae (SN) events or in super giants.



⁴He production in stars and during the big bang



2. Evolution of Elements

Table of Isotopes of the Light Elements (Stable Isotopes are Drawn in Blue)

10								^{17}Ne	^{18}Ne
9								^{16}F	^{17}F
8						^{13}O	^{14}O	^{15}O	^{16}O
7						^{12}N	^{13}N	^{14}N	^{15}N
6				^9C	^{10}C	^{11}C	^{12}C	^{13}C	^{14}C
5				^8B	^9B	^{10}B	^{11}B	^{12}B	^{13}B
4			^6Be	^7Be	^8Be	^9Be	^{10}Be	^{11}Be	^{12}Be
3			^5Li	^6Li	^7Li	^8Li	^9Li		
2		^3He	^4He	^5He	^6He		^8He		
1	^1H	^2H	^3H						
	0	1	2	3	4	5	6	7	8

Beryllium ^9Be is the first element with solely one stable isotope (pure element): Very toxic...

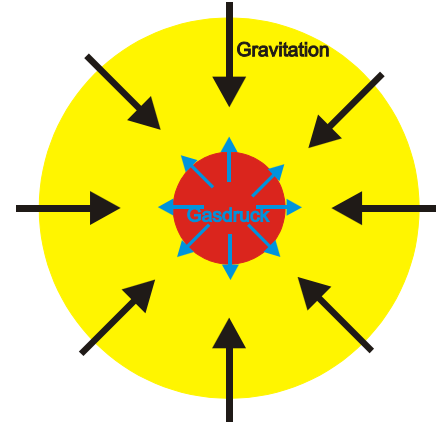
Fluorine ^{19}F is the first biochemical relevant element with solely one stable isotope...

Phosphorus ^{31}P is the only stable isotope and the most critical element for biology...

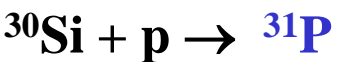
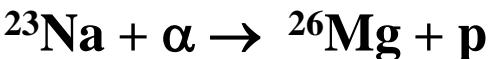
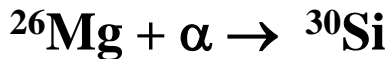
2. Evolution of Elements

Formation of Moderately Heavy Elements (Stellar Synthesis)

Several fission processes lead to a number of products:

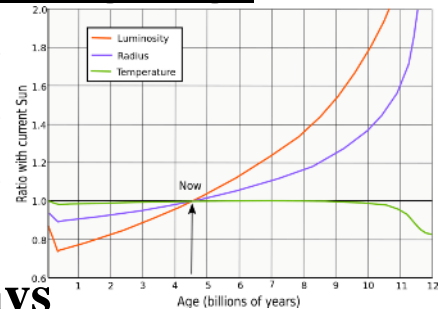


Formation of phosphorus (rare)



Pre-supernova burning stages of a star with 25 times the solar mass

Burn process	T [10 ⁹ K]	Main products	Duration of burning stage
H	0.02	^4He , ^{14}N	$7 \cdot 10^6$ a
He	0.2	^{12}C , ^{16}O , ^{20}Ne	$5 \cdot 10^5$ a
C	0.8	^{20}Ne , ^{23}Na , ^{24}Mg	$6 \cdot 10^2$ a
Ne	1.5	^{20}Ne , ^{23}Na , ^{24}Mg	1 a
O	2.0	^{28}Si , ^{32}S , ^{40}Ca	180 days
Si	3.5	^{54}Fe , ^{56}Ni , ^{52}Cr	1 day!



2. Evolution of Elements

Formation of the Heavy Elements

S(slow)-process (in red supergiants):

- Kinetics: Time of β -decay must be orders of magnitude higher than the period till the next capture of a neutron
- Starting points are seed cores such as ^{56}Fe
- They capture neutrons $\rightarrow ^{59}\text{Fe}$ and decompose via a β -decay to ^{59}Co
- This process is repeating itself \rightarrow the process moves along the stability valley of the table of isotopes

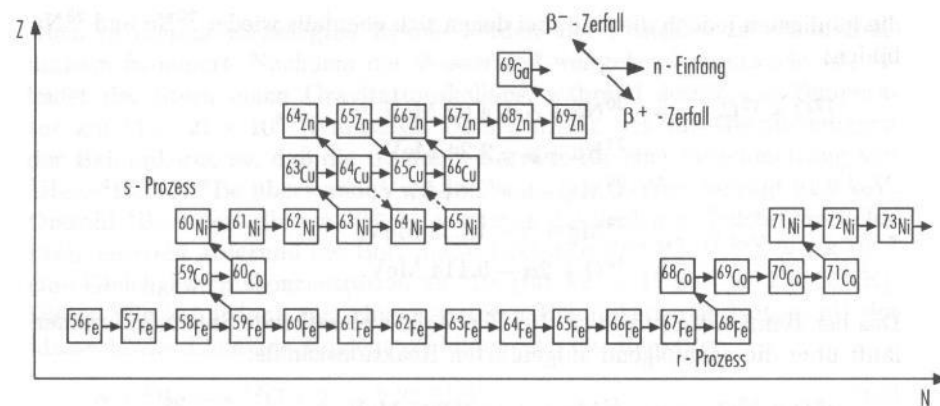
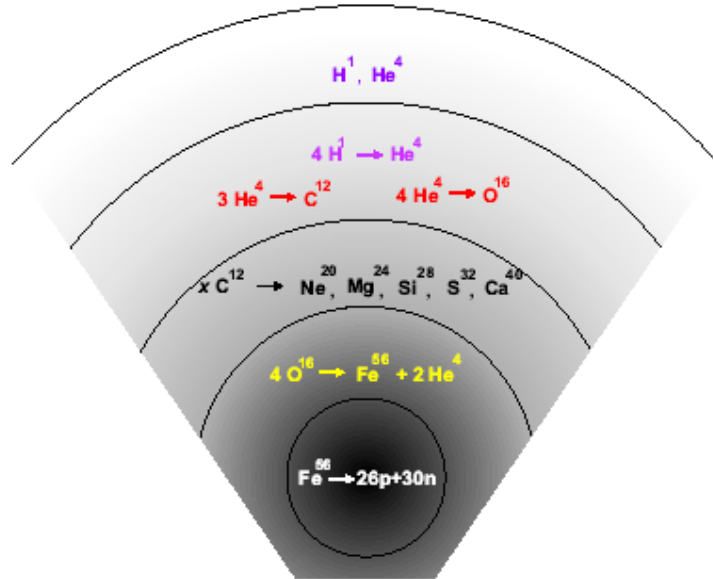


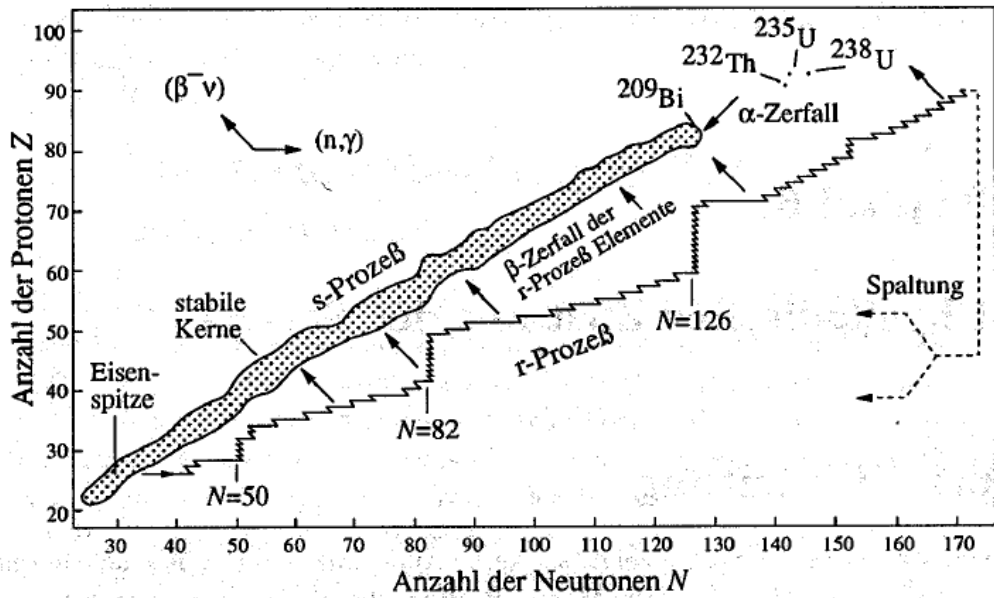
Bild 9.7. Verzweigung der s- und r-Prozesse zur Elementbildung oberhalb Eisen

2. Evolution of Elements

Formation of the Heavy Elements

R(rapid)-process (in supernovae, SN):

- Requires extremely high flux of neutrons to compensate for the β -decay
- The core is enriched with neutrons (20 - 30 neutrons) until it reaches the “neutron drip line“. By spontaneous emission of neutrons the core remains in that waiting state until it decomposes via the β -decay
- Such neutron density (10^{24} cm^{-3}) is reached by photo disintegration within the core of SN



2. Evolution of Elements

1	Groups												18	2 He			
1 H	2 Be	3 Li	4 Be	5	6	7	8	9	10	11	12	13 B	14 C	15 N	16 O	17 F	2 He
11 Na	12 Mg	3	4	5	6	7	8	9	10	11	12	13 Al	14 Si	15 P	16 S	17 Cl	18 Ar
19 K	20 Ca	21 Sc	22 Ti	23 V	24 Cr	25 Mn	26 Fe	27 Co	28 Ni	29 Cu	30 Zn	31 Ga	32 Ge	33 As	34 Se	35 Br	36 Kr
37 Rb	38 Sr	39 Y	40 Zr	41 Nb	42 Mo	43 Tc	44 Ru	45 Rh	46 Pd	47 Ag	48 Cd	49 In	50 Sn	51 Sb	52 Te	53 I	54 Xe
55 Cs	56 B	57 La	72 Hf	73 Ta	74 W	75 Re	76 Os	77 Ir	78 Pt	79 Au	80 Hg	81 Tl	82 Pb	83 Bi	84 Po	85 At	86 Rn
87 Fr	88 Ra	89 Ac	104 Rf	105 Db	106 Sg	107 Bh	108 Hs	109 Mt	110 Ds	111 Rg	112 Cn						
			58 Ce	59 Pr	60 Nd	61 Pm	62 Sm	63 Eu	64 Gd	65 Tb	66 Dy	67 Ho	68 Er	69 Tm	70 Yb	71 Lu	
			90 Th	91 Pa	92 U	93 Np	94 Pu	95 Am	96 Cm	97 Bk	98 Cf	99 Es	100 Fm	101 Md	102 No	103 Lr	

Synthesis within 15 min after the big bang stellar synthesis

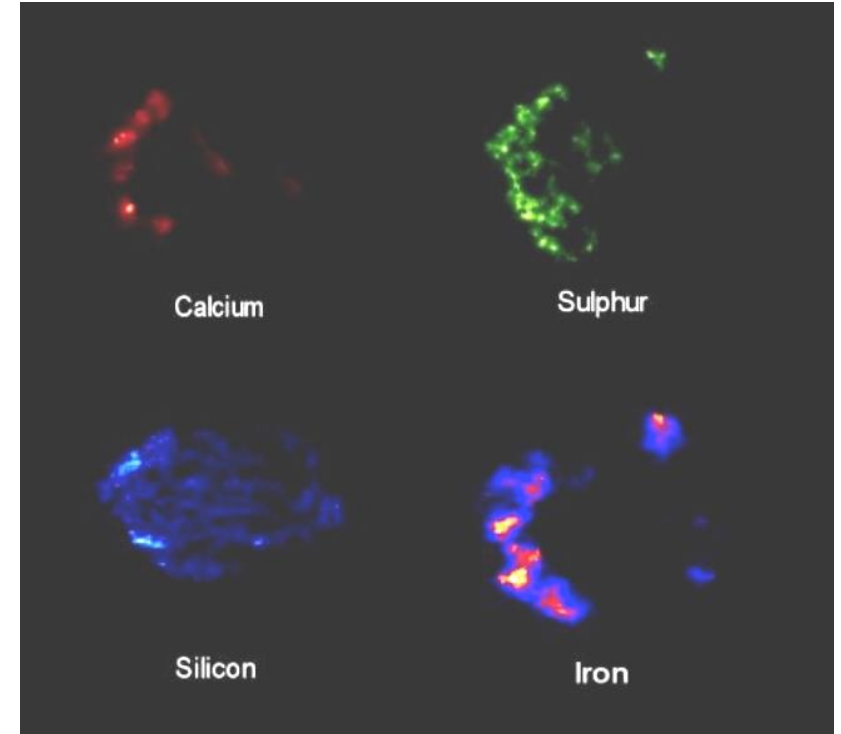
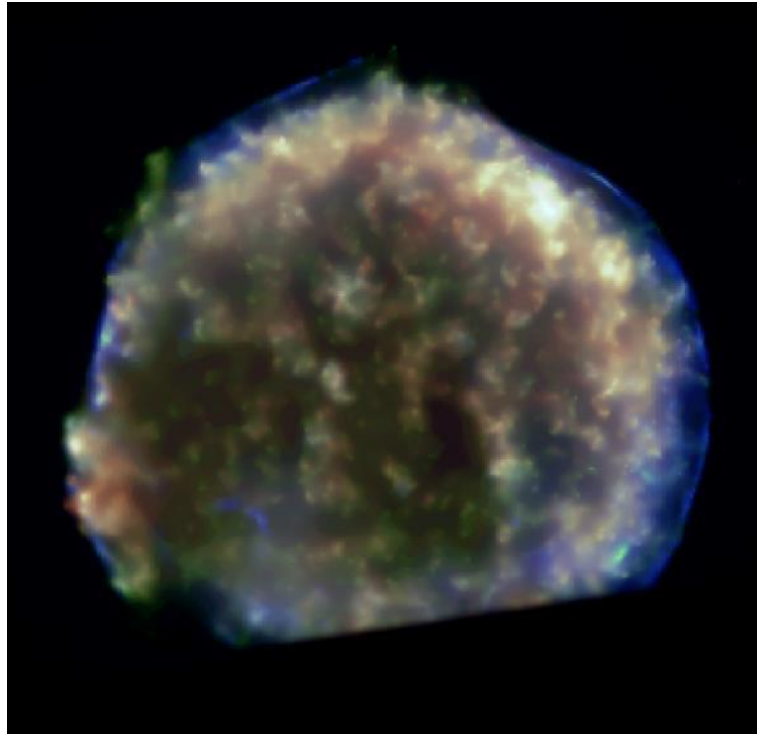
Stellar formation via the s(slow)-process (neutron capture and β -decay)

Formation via the (rapid)-process during supernovae explosions of the type II

2. Evolution of Elements

Distribution within the Interstellar Medium

1. Supernovae explosions, e.g. in Supernova Type Ia of white dwarfs (SN1572, observed by Danish astronomer Tycho Brahe)
2. T-Tauri stars → strong stellar winds, e.g. in the columns of creation (JWST)



X-ray images of SN1572 by space telescope on board of satellite “Chandra”

2. Evolution of Elements

Elemental Composition of the Sun and of Carbonaceous Chondrites (C1) Typical for Stars in the Milky Way

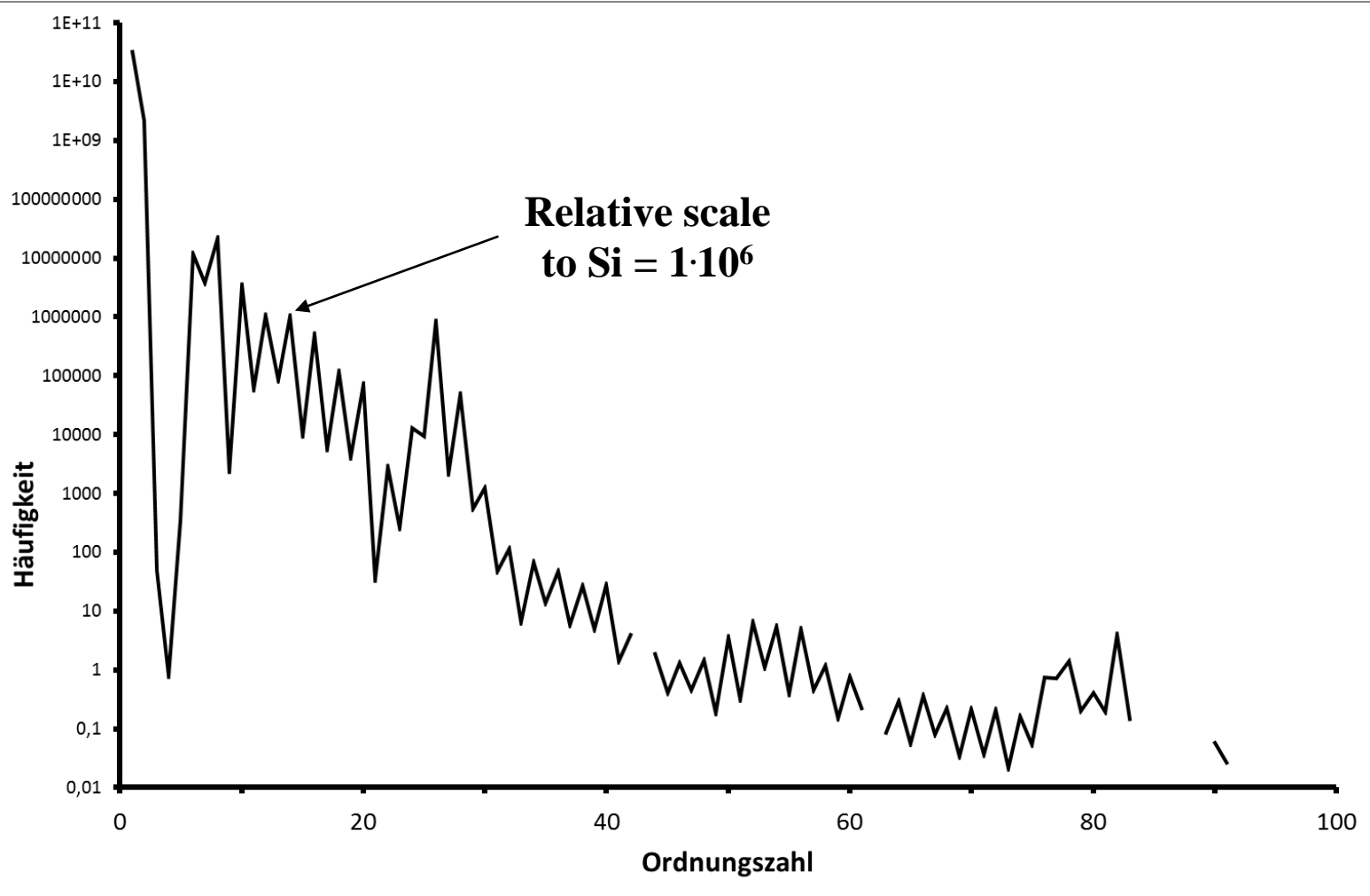
Tabelle 4.9.1. Elementhäufigkeiten $\log N$ im Sonnensystem: Sonne (\odot) nach H. Holweger (1985) und *kohlige Chondrite vom Typ C1* nach E. Anders und M. Ebihara (1982). Normierung auf Wasserstoff $\log N(H) = 12.0$. Anpassung der solaren und meteoritischen Häufigkeitsverteilungen bei Silizium $\log N(Si) = 7.6$. Bestimmung der Sonnenhäufigkeiten aus der Photosphäre mit Ausnahme von He, Ne, Ar (Korona bzw. Protuberanzen) und Ti (Sonnenflecken). Meteorite: C1-Chondrite bis auf Be, B, Br, Rh, I, für die andere Chondrite herangezogen wurden. Für Kr, Xe, Hg geschätzte Werte aus Interpolationen. Radioaktive Elemente: Th, U Angabe der *heutigen* Häufigkeiten; bei der Entstehung des Sonnensystems vor $4.5 \cdot 10^9$ a (2.8.24) waren die Häufigkeiten um $\delta \log N = 0.2$ (Th) bzw. 0.3 (U) höher

	\odot	C1		\odot	C1		\odot	C1		\odot	C1
1 H	12.0	—	22 Ti	5.1	5.0	44 Ru	1.8	1.9	66 Dy	1.1	1.2
2 He	11.0	—	23 V	4.1	4.1	45 Rh	1.1	1.1	67 Ho	0.3	0.6
3 Li	1.1	3.4	24 Cr	5.8	5.7	46 Pd	1.7	1.7	68 Er	0.9	1.0
4 Be	1.2	1.5	25 Mn	5.4	5.6	47 Ag	0.9	1.3	69 Tm	0.3	0.1
5 B	2.5	3.0	26 Fe	7.6	7.6	48 Cd	1.9	1.8	70 Yb	1.1	1.0
6 C	8.6	—	27 Co	4.9	5.0	49 In	1.7	0.9	71 Lu	0.8	0.2
7 N	8.0	—	28 Ni	6.2	6.3	50 Sn	1.9	2.2	72 Hf	0.9	0.8
8 O	8.9	—	29 Cu	4.2	4.3	51 Sb	1.0	1.1	73 Ta	—	0.0
9 F	4.6	4.5	30 Zn	4.6	4.7	52 Te	—	2.3	74 W	1.1	0.7
10 Ne	7.6	—	31 Ga	2.9	3.2	53 I	—	1.6	75 Re	—	0.3
11 Na	6.3	6.4	32 Ge	3.5	3.7	54 Xe	—	(2.2)	76 Os	1.4	1.5
12 Mg	7.5	7.6	33 As	—	2.4	55 Cs	—	1.2	77 Ir	1.4	1.4
13 Al	6.4	6.5	34 Se	—	3.4	56 Ba	2.1	2.2	78 Pt	1.8	1.7
14 Si	7.6	7.6	35 Br	—	2.7	57 La	1.1	1.3	79 Au	1.1	0.9
15 P	5.4	5.6	36 Kr	—	(3.3)	58 Ce	1.6	1.7	80 Hg	—	(1.3)
16 S	7.2	7.3	37 Rb	2.6	2.5	59 Pr	0.7	0.8	81 Tl	0.9	0.9
17 Cl	—	5.3	38 Sr	3.0	3.0	60 Nd	1.4	1.5	82 Pb	1.9	2.1
18 Ar	6.7	—	39 Y	2.2	2.3	62 Sm	0.8	1.0	83 Bi	—	0.8
19 K	5.1	5.2	40 Zr	2.6	2.6	63 Eu	0.5	0.6	90 Th	0.2	0.1
20 Ca	6.4	6.4	41 Nb	1.4	1.5	64 Gd	1.1	1.1	92 U	—	-0.4
21 Sc	3.1	3.1	42 Mo	1.9	2.0	65 Tb	0.2	0.4			

Normalised by hydrogen by $N(H) = 12.0$

2. Evolution of Elements

Elemental Composition of the Sun



Source:
Wikipedia

The “bio” elements F, P, Cl and K are rather rare!

3. Elements and Molecules of Life

Elemental Composition of the Solar Systems

About 4.7 billion years ago:

81 stable elements existing in the protoplanetary (solar) nebula. That means all elements till Bi, except Tc and Pm, since they only possess short-lived isotopes

About 26 elements in living organisms:

1. Necessary in quantitative amounts: 11 elements

C, H, O, N, S, P, Na, Mg, Cl, K, Ca

2. In smaller amounts needed: 8 elements

Mn, Fe, Co, Ni, Cu, Zn, I, Mo

3. Elements, occurring only in some species: 8 elements

B, F, Si, V, Cr, Se, Sn, W

	C	H	O	N	S	P
Carbohydrates	X	X	X			
Lipides	X	X	X	X		X
Proteins	X	X	X	X	X	
Nucleotides	X	X	X	X		X
Porphyrines	X	X	X	X		

3. Elements and Molecules of Life

**Most Abundant Elements of Earth's Crust
(Atmo-, Bio-, Hydro-, Cryo- and Lithosphere)
According to Weight Fractions:**

1. Oxygen	48.9%
2. Silicon	26.3%
3. Aluminium	7.7%
4. Iron	4.7%
5. Calcium	3.4%
6. Sodium	2.6%
7. Potassium	2.4%
8. Magnesium	<u>1.9%</u> 97.9%



All other elements of the periodic table add up to only 2.1%:

H: 1400 ppm

S 350 ppm

C 200 ppm

Cu 60 ppm

Co 25 ppm

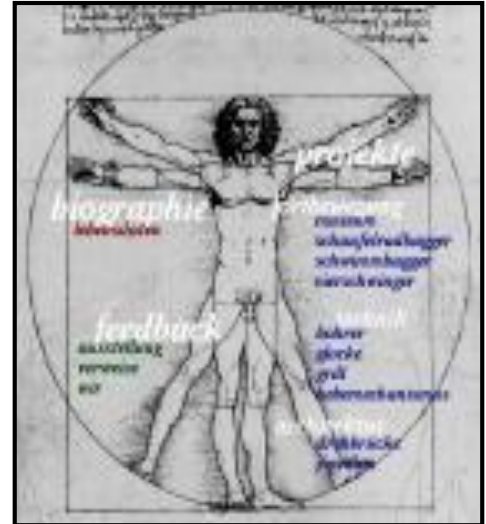
3. Elements and Molecules of Life

Most Abundant Elements in the Human Body According to Weight Fraction

1. Oxygen	65.4%
2. Carbon	18.1%
3. Hydrogen	10.1%
4. Nitrogen	3.0%
5. Calcium	1.5%
6. Phosphorus	1.0%
7. Sulphur	<u>0.25%</u>
	99.35%

All other elements of the periodic table contribute to solely 0.65% to the mass of humans

Trace Elements	Daily demand of the human body
Iron	10 - 20 mg
Zinc	7 - 10 mg
Manganese	2 - 5 mg
Copper	1 – 1.5 mg
Molybdenum	0.05 – 0.1 mg
Vanadin	0.01 - 0.03 mg
Cobalt	0.003 mg



3. Elements and Molecules of Life

Composition of the Human Body (70 kg) by the Elemental Weight Fraction

- oxygen 43 kg
- carbon 16 kg
- hydrogen 7 kg
- nitrogen 1.8 kg
- calcium 1.0 kg
- phosphorus 780 g
- potassium 140 g
- sulphur 140 g
- sodium 100 g
- chlorine 95 g
- magnesium 19 g
- iron 4.2 g
- fluorine 2.6 g
- zinc 2.3 g
- silicon 1.0 g
- rubidium 0.68 g
- strontium 0.32 g
- bromine 0.26 g
- lead 0.12 g
- copper 72 mg
- aluminium 60 mg
- cadmium 50 mg
- cerium 40 mg
- barium 22 mg
- iodine 20 mg
- tin 20 mg
- titanium 20 mg
- boron 18 mg
- nickel 15 mg
- selenium 15 mg
- chromium 14 mg
- manganese 12 mg
- arsenic 7 mg
- lithium 7 mg
- caesium 6 mg
- mercury 6 mg
- germanium 5 mg
- molybdenum 5 mg
- cobalt 3 mg
- antimony 2 mg
- silver 2 mg
- niobium 1.5 mg
- zirconium 1 mg
- lanthanum 0.8 mg
- gallium 0.7 mg
- tellurium 0.7 mg
- yttrium 0.6 mg
- bismuth 0.5 mg
- thallium 0.5 mg
- indium 0.4 mg
- gold 0.2 mg
- scandium 0.2 mg
- tantalum 0.2 mg
- vanadium 0.11 mg
- thorium 0.1 mg
- uranium 0.1 mg
- samarium 50 µg
- beryllium 36 µg
- tungsten 20 µg

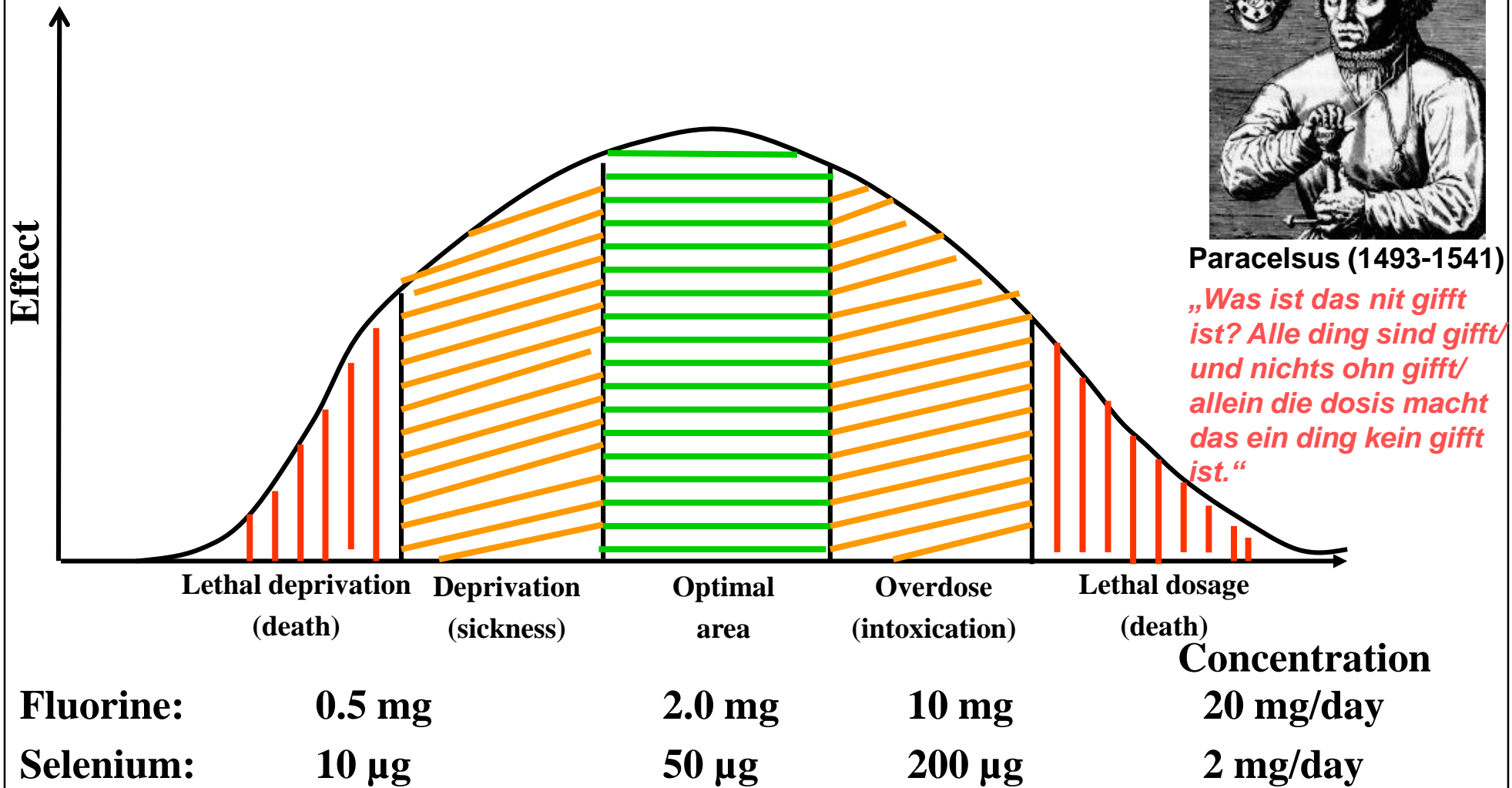
John Emsley, "The Elements", 3rd ed. Clarendon Press, Oxford, 1998

3. Elements and Molecules of Life

1	Groups												18				
1 H	2 He													2 He			
3 Li	4 Be													10 Ne			
11 Na	12 Mg	3	4	5	6	7	8	9	10	11	12	13 Al	14 Si	15 P	16 S	17 Cl	18 Ar
19 K	20 C	21 Sc	22 Ti	23 V	24 Cr	25 Mn	26 Fe	27 Co	28 Ni	29 Cu	30 Zn	31 Ga	32 Ge	33 As	34 Se	35 Br	36 Kr
37 Rb	38 Sr	39 Y	40 Zr	41 Nb	42 Mo	43 Tc	44 Ru	45 Rh	46 Pd	47 Ag	48 Cd	49 In	50 Sn	51 Sb	52 Te	53 I	54 Xe
55 Cs	56 Ba	57 La	72 Hf	73 Ta	74 W	75 Re	76 Os	77 Ir	78 Pt	79 Au	80 Hg	81 Tl	82 P	83 Bi	84 Po	85 At	86 Rn
87 Fr	88 Ra	89 Ac	104 Rf	105 Db	106 Sg	107 Bh	108 Hs	109 Mt	110 Ds	111 Rg	112 Cn						
			58 Ce	59 Pr	60 Nd	61 Pm	62 Sm	63 Eu	64 Gd	65 Tb	66 Dy	67 Ho	68 Er	69 Tm	70 Yb	71 Lu	
			90 Th	91 Pa	92 U	93 Np	94 Pu	95 Am	96 Cm	97 Bk	98 Cf	99 Es	100 Fm	101 Md	102 No	103 Lr	

3. Elements and Molecules of Life

Concentration and Physiological Effect of a Substance



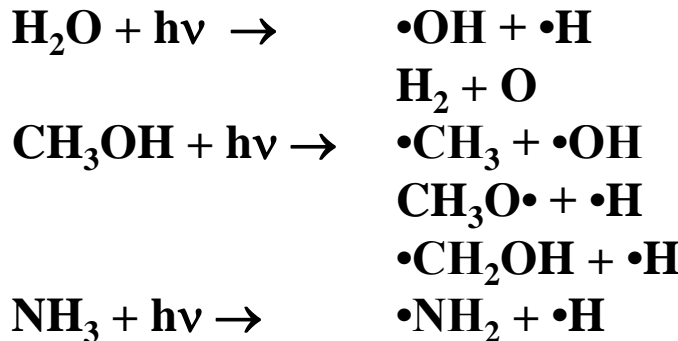
Paracelsus (1493-1541)

*„Was ist das nit gifft
ist? Alle ding sind gifft/
und nichts ohn gifft/
allein die dosis macht
das ein ding kein gifft
ist.“*

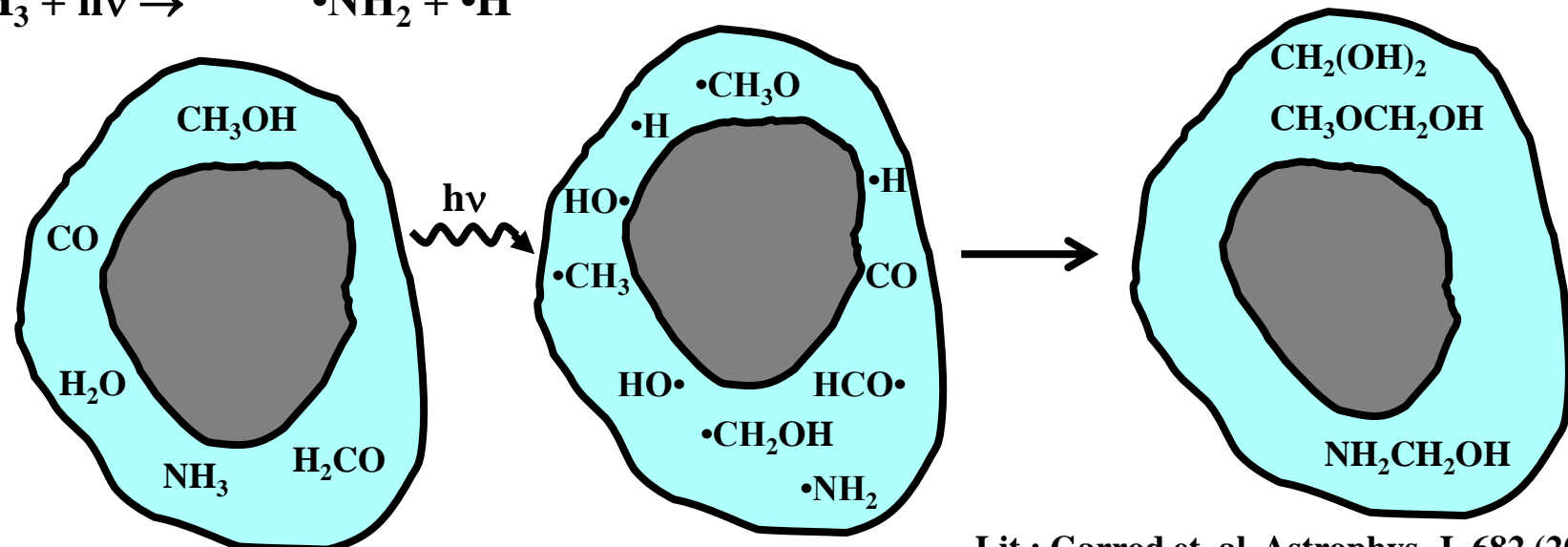
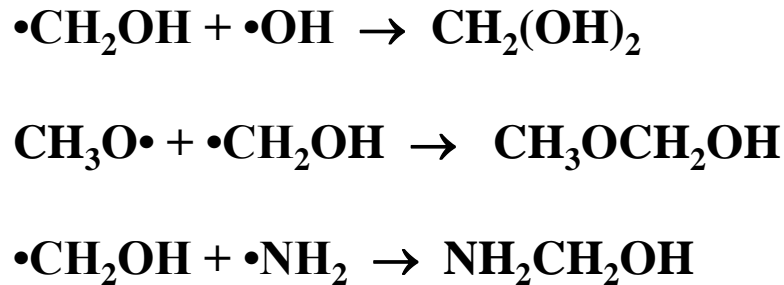
3. Elements and Molecules of Life

Formation of Amino acids is crucial to life

Activation by photolysis reactions



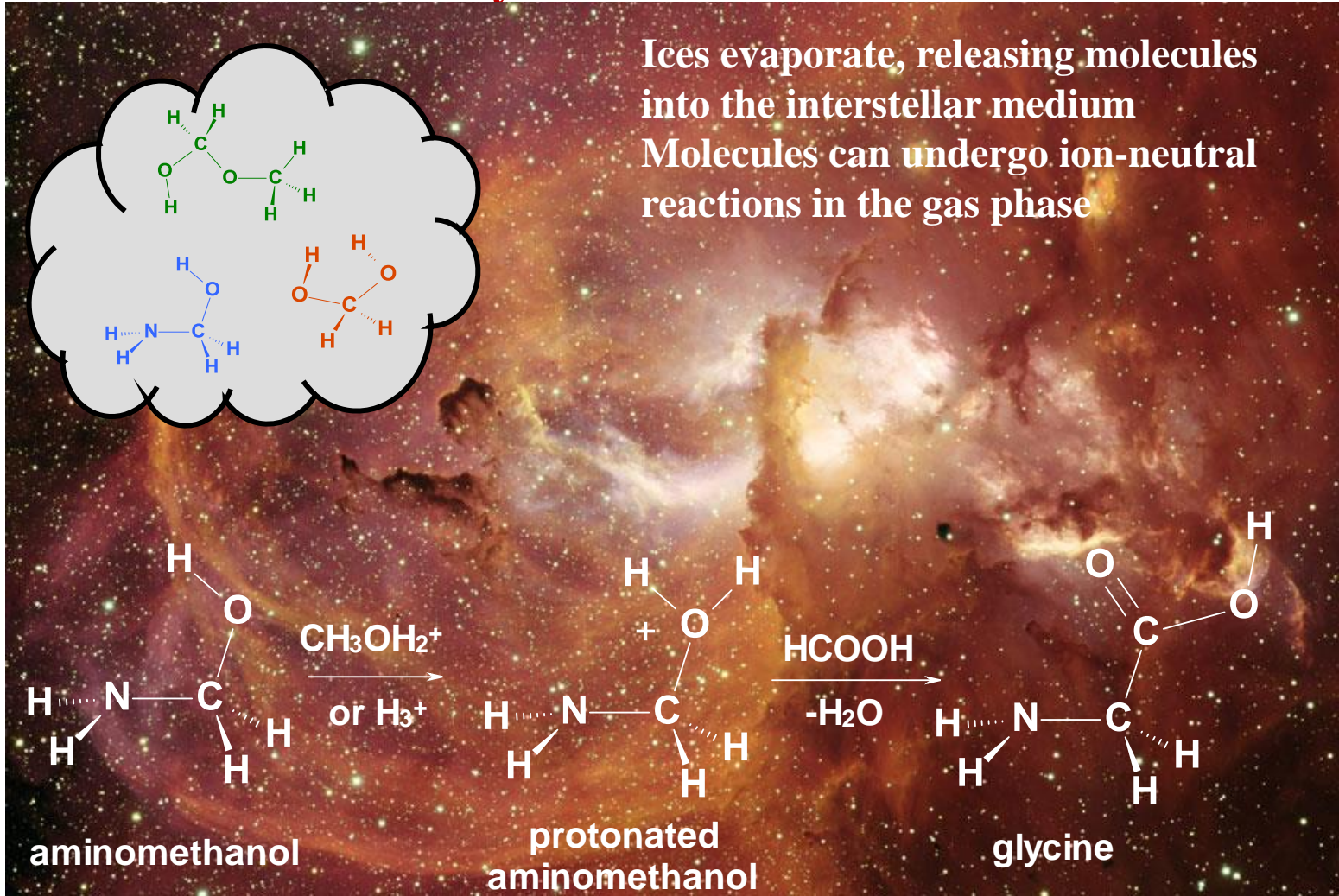
Radical-Radical recombination reactions



Lit.: Garrod et. al. *Astrophys. J.* 682 (2008) 283-302

3. Elements and Molecules of Life

Probiotic Astrochemistry

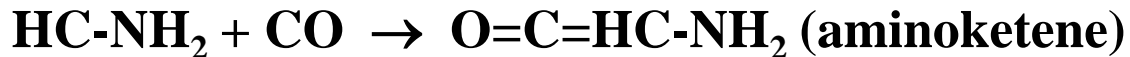
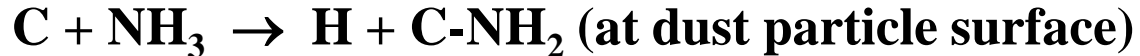


Photograph: T.A. Rector and T. Abbott, U. Alaska and NOAO, AURA, NASA, NGC 3582

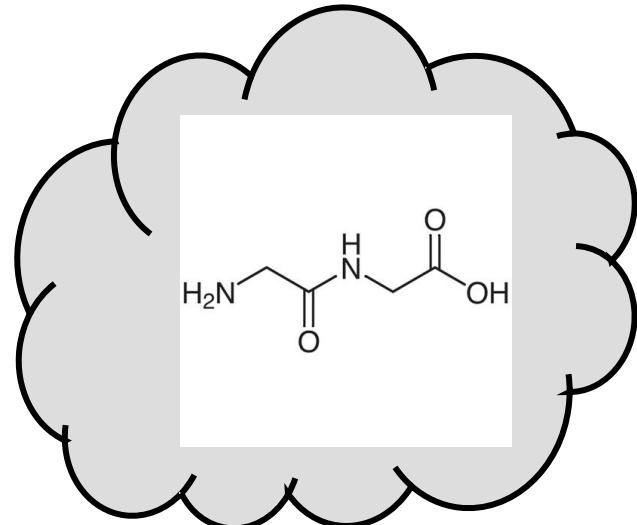
3. Elements and Molecules of Life

Probiotic Astrochemistry

Formation of peptides in space without condensation reaction at dust particles of the interstellar medium (ISM), T ~ 10-20 K:



→ polypeptides?



Lit.: Nature Astronomy 6 (2022) 381

3. Elements and Molecules of Life

Probiotic Geochemistry

Primordial atmosphere

$\text{H}_2\uparrow$, $\text{He}\uparrow$, CH_4 , N_2 , NH_3 , H_2O



1st atmosphere (4 bill. years ago)

80% H_2O

10% CO_2

5-7% H_2S

Traces of N_2 , H_2 , CO , He , NH_3

2nd atmosphere (3 bill. years ago)

N_2

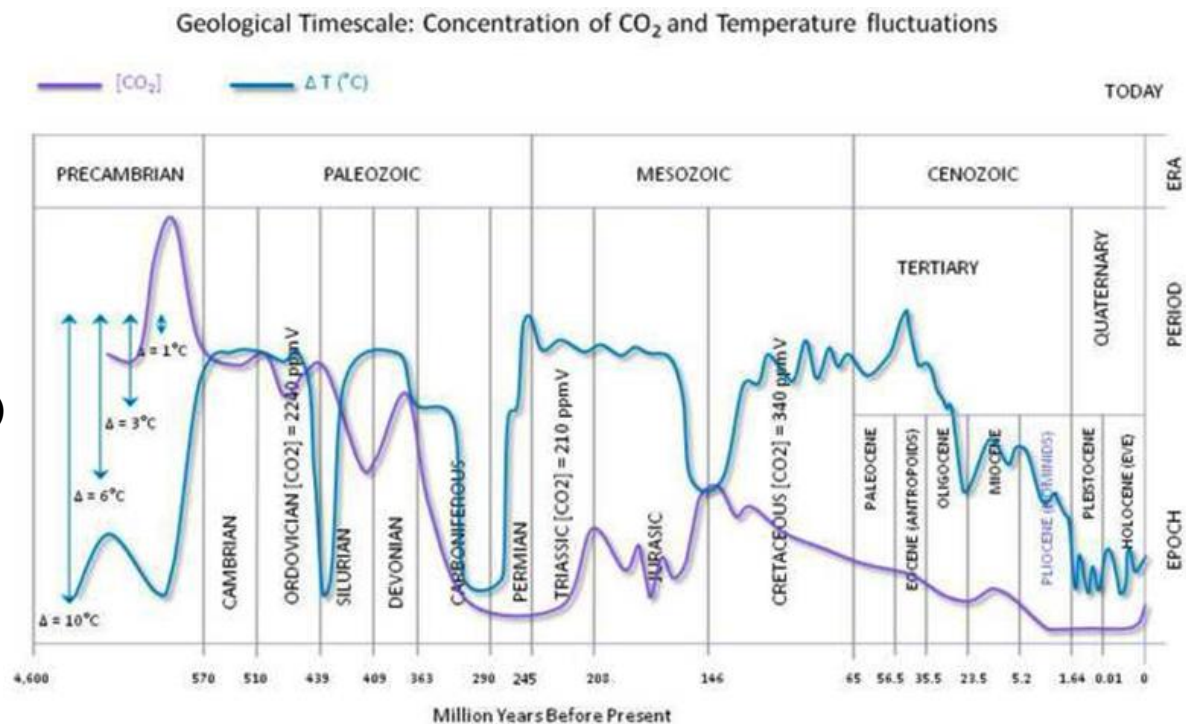
Traces of CO_2 , H_2O , and Ar

Present atmosphere

78% N_2 , 21% O_2

0.93% Ar

Traces of CO_2 , H_2O , O_3 , CH_4



1- Analysis of the Temperature Oscillations in Geological Eras by Dr. C. R. Scotese © 2002. 2- Ruddiman, W. F. 2001. Earth's Climate: past and future. W. H. Freeman & Sons. New York, NY. 3- Mark Pagani et al. Marked Decline in Atmospheric Carbon Dioxide Concentrations During the Paleocene. Science; Vol. 309, No. 5734; pp. 600-603. 22 July 2005.
Corrected on 07 July 2008 (CO2: Ordovician Period).

4. Phylogeny

The Distribution of Elements in the Terrestrial Atmos-, Bio-, Hydro, Cryo-, and Lithosphere Differs Significantly from the Stellar Distribution of Elements

Earth's core

Heavy elements \Rightarrow Fe, Ni and other metals along with C as carbides

Lithosphere

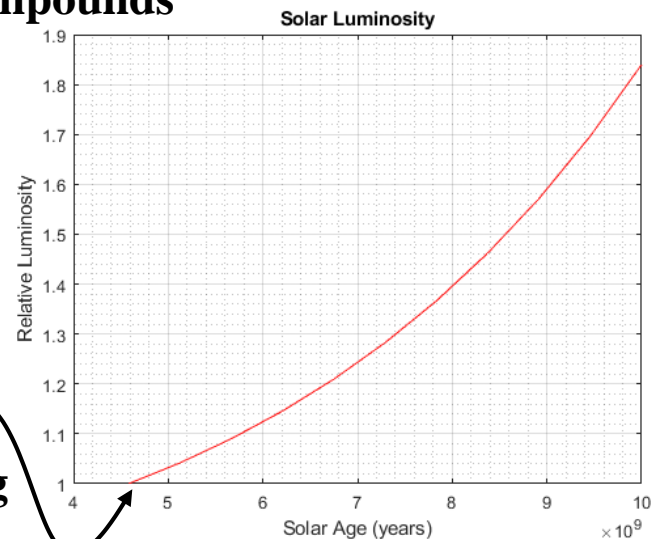
lighter elements \Rightarrow silicates, aluminosilicate, Mg- and Ca-compounds

Primordial atmosphere

- Rapid emission of H₂ due to too low mass of the earth
- Photolysis of water vapour: $2 \text{H}_2\text{O} \rightarrow 2 \text{H}_2 + \text{O}_2$
- Reductive: CH₄, N₂, NH₃, H₂O, PH₃, H₂S, CO₂

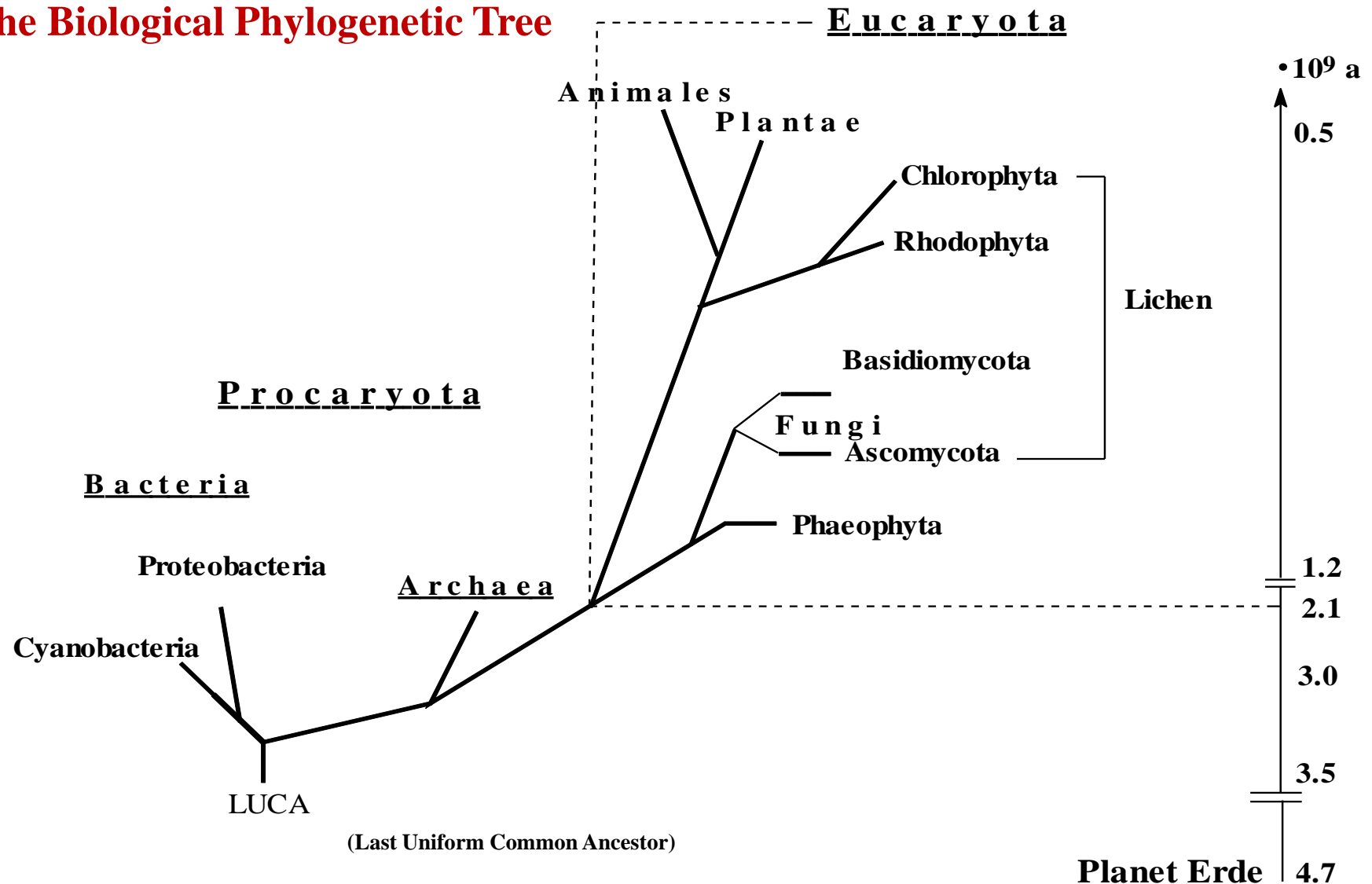
Present atmo-, hydro-, cryosphere, and lithosphere

- N₂ \rightarrow NO_x \rightarrow NO₂/NO₃⁻ (fertiliser) through lightning
- CO₂ \rightarrow CO₃²⁻ \rightarrow carbonates↓ (e.g. dolomite)
- CO₂ \rightarrow C (fossil fuels) + O₂ through biological activity \rightarrow O₃ (ozone layer)
- H₂O(g) \rightarrow H₂O(l) "oceans" \rightarrow H₂O(s) "ice caps"



4. Phylogeny

The Biological Phylogenetic Tree



4. Phylogeny

From Prokaryotes to Eukaryotes

Prokaryotes

Bacteria and archaea

→ mostly unicellular but bigger agglomerates possible

Eukaryotes

All higher organisms

Plants

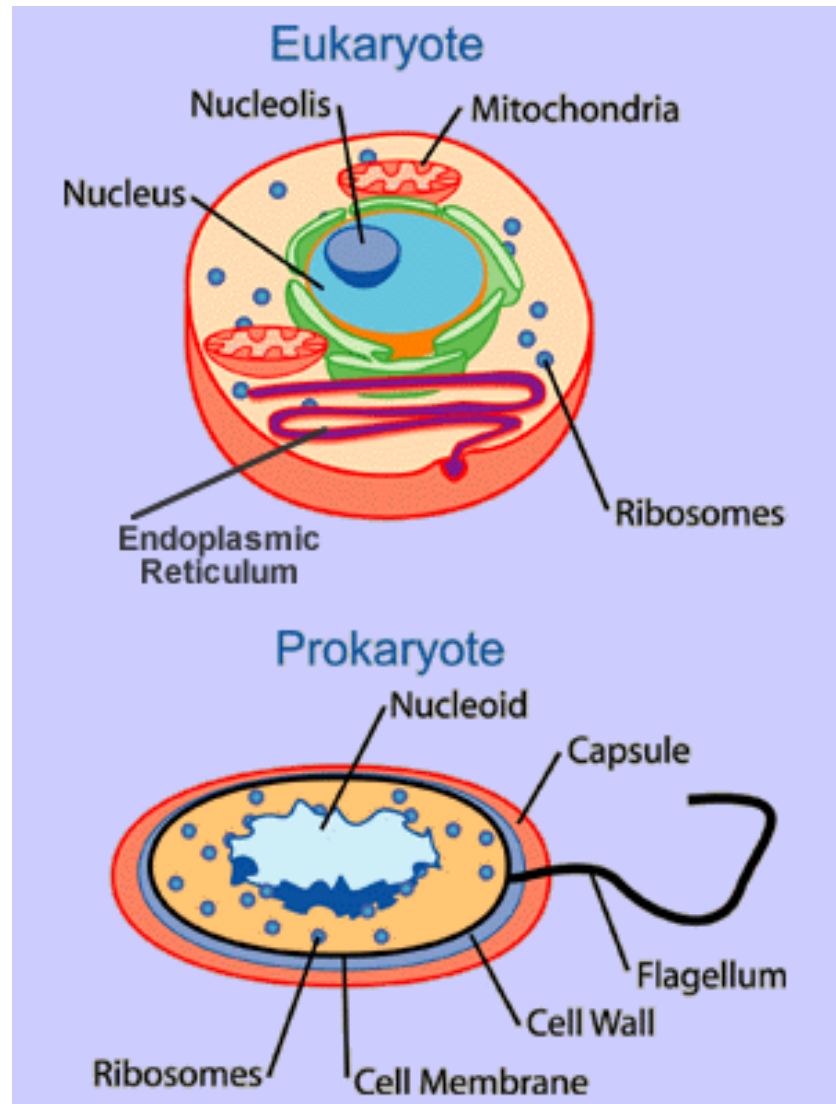
Animals

Fungi

Algae

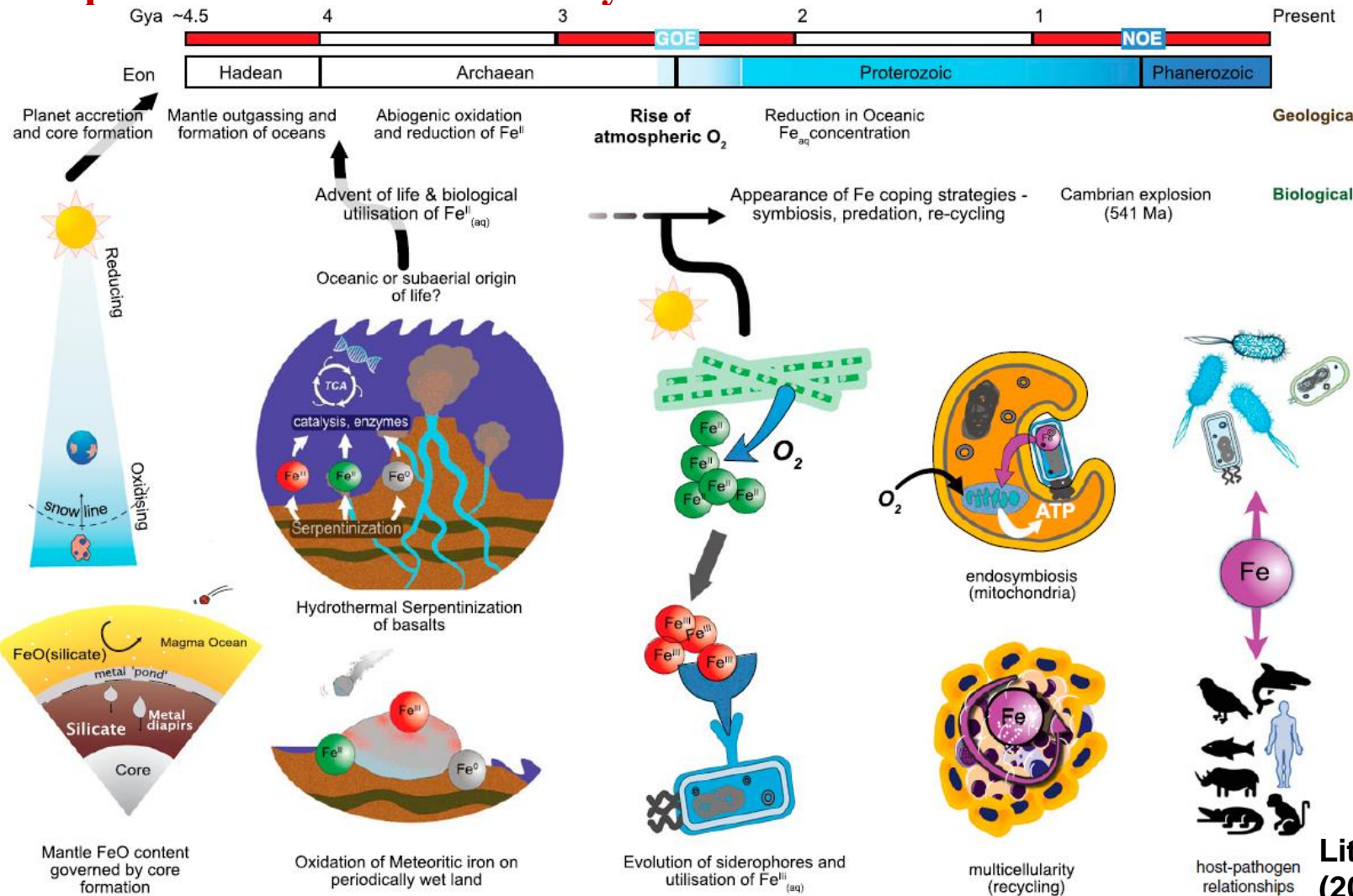
→ cell core and mitochondria/chloroplasts

→ mostly polycellular



4. Phylogeny

Temporal Variation of Planetary Fe Concentration: Main Driver of Evolution?

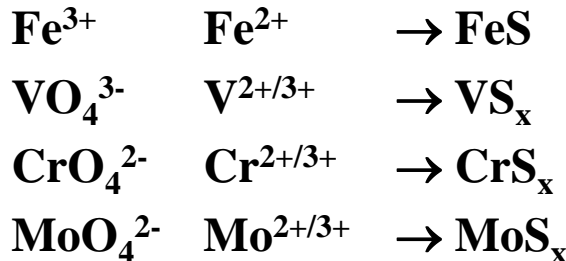


Lit.: PNAS 118
(2021) e2109865118

4. Phylogeny

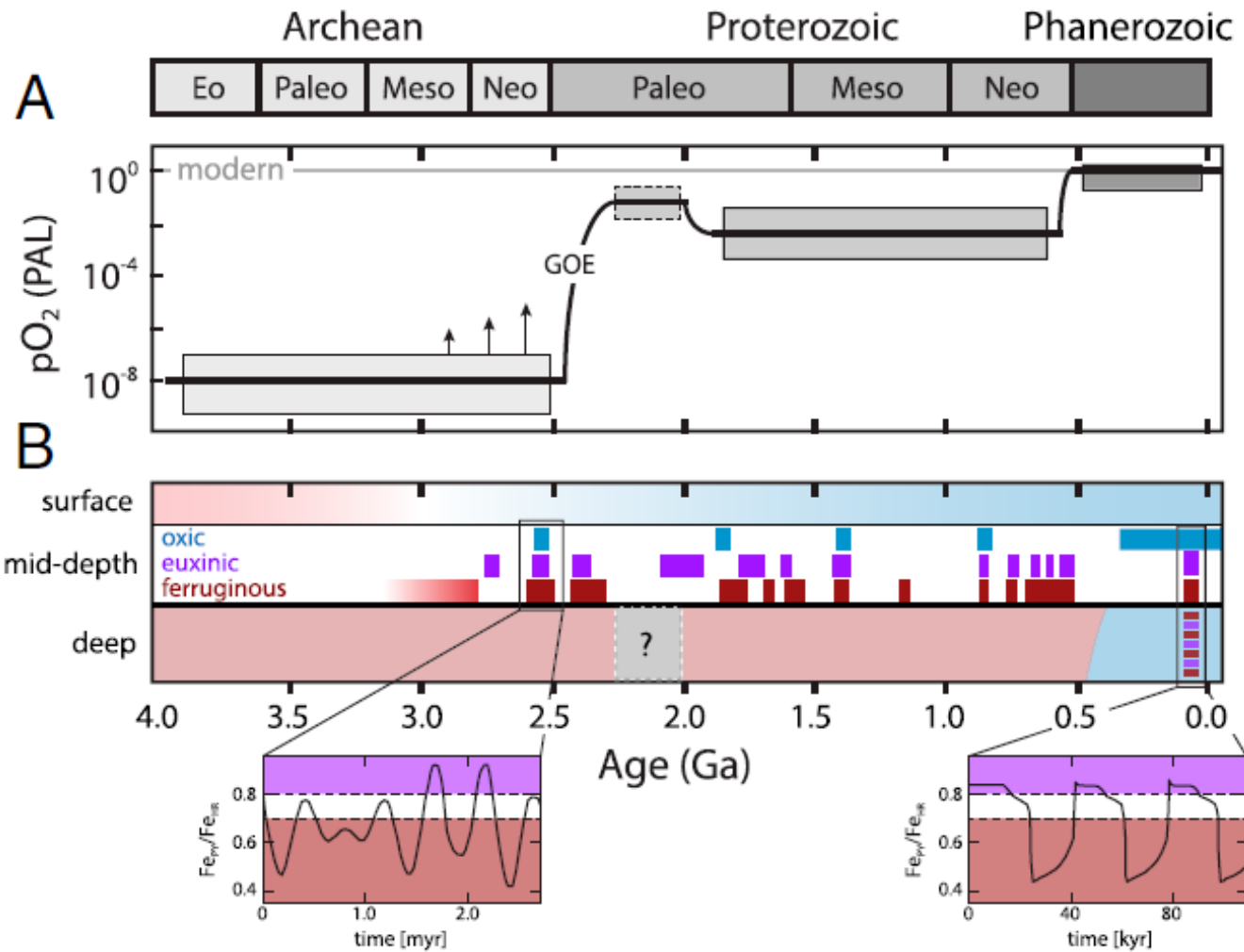
Temporal Variation of Planetary Metal Concentration in Ocean Water

oxic euxinic



Euxenic conditions causes severe depletion of transition metals in ocean water

e.g. at P/T mass extinction about 252 mill. years ago



Lit.: PNAS 117 (2020) 33043

4. Phylogeny

Key Points so far

- Many (transition) metals are essential for life
- Evolution is driven by presence or absence of transition metals
- Organisms make economic use of available resources, but also have developed mechanisms to accumulate certain elements
- Despite the low amount of metal ions present in living systems, they are enormously important for virtually all life processes
- Both deficiency and overload / excess lead to illness
- Dissipation of “toxic” metals, such as Pb into the biosphere is a threat to many ecosystems

5. Metals in Biochemistry

Essential Metals for Life (Aqueous Chemistry)

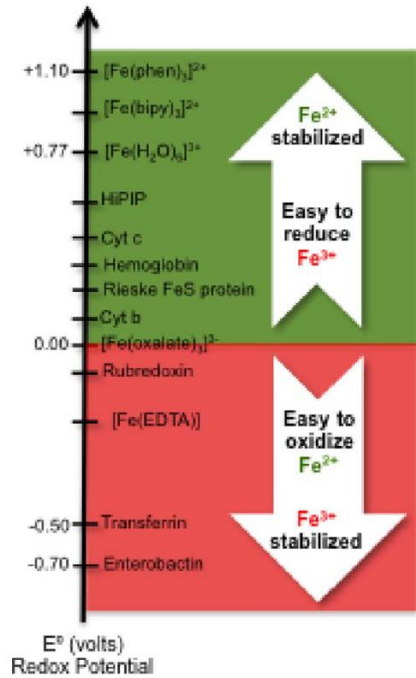
Alkaline metals	Na, K	readily soluble in water, acids, and bases
Alkaline earth metals	Mg, Ca	readily soluble in acids
Main group metals	Sn, Se	relatively poor solubility
Transition metals	V, Cr, Mn, Fe, Co, Ni, Cu, Zn, Mo, W	moderate and variable solubility
Rare earth metals	LREE (La-Nd)? Eu?	poor solubility some hyperaccumulators known, e.g. wheat, dycranopteris (ferns)



5. Metals in Biochemistry

Functions of Metals for Life

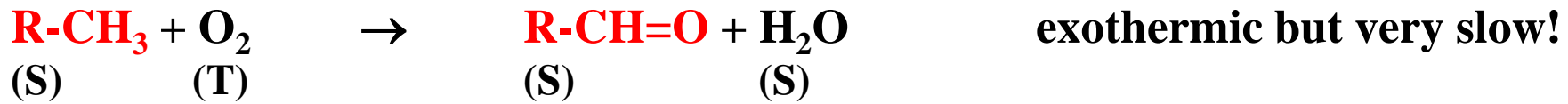
- Metal cations can adopt several coordination numbers
 - bonding and activation of substrates to enhance Lewis acidity
- Metal cations can adopt several coordination geometries
 - fine tuning of electronic properties such as redox potentials
- Ligands can be substituted without changing the structure
 - fast ligand exchange, i.e. fast but unspecific catalysis
- (Transition) metal cations can change size without changing the oxidation state
 - changing number of ligands, activation of inactive ligands
 - high spin Fe^{2+} (78 pm) in $[\text{Fe}(\text{por})(\text{H}_2\text{O})]$ vs. low spin Fe^{2+} (55 pm) in $[\text{Fe}(\text{por})(\text{O}_2)]$
- (Transition) metal cations can adopt many oxidation states and transfer „spin“
 - atom transfer reactions, redox reactions, electron storage
 - activation of ${}^3\text{O}_2$ (triplet)



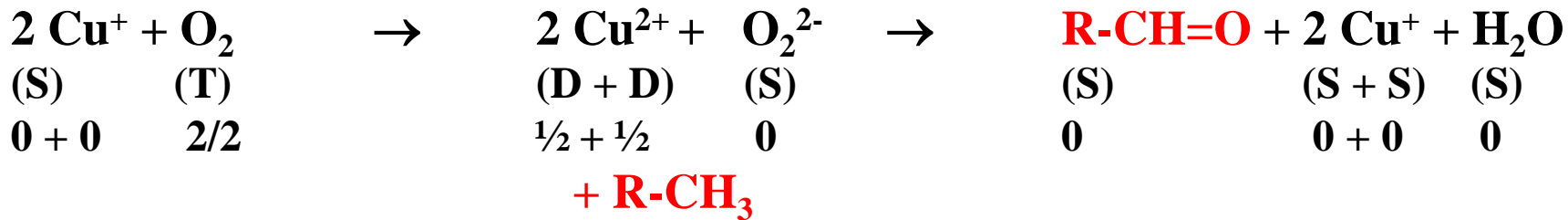
5. Metals in Biochemistry

Functions of Metals for Life: Transfer of Spin for the Activation of Oxygen

Most chemical reactions between organic molecules and oxygen are „forbidden“ and thus very slow:



Activation of ${}^3\text{O}_2$ (triplet) possible by spin transfer to a metal cation:



→ Metals eases spin-exchange → catalysis of “spin-forbidden” chemical reactions

5. Metals in Biochemistry

Biomolecules Comprising Metal Ions

Structural proteins

- Protein structure
- Scaffolding functions

Ca, Zn

Mg, Ca

Transportation and storage proteins

- e⁻-transport
- O₂-transport
- Mⁿ⁺-transport

Fe, Cu

V, Fe, Cu

Fe, Zn

Enzymes

- Hydrolases
- Oxidoreductases
- Isomerases/synthetases

Mg, Zn

V, Cr, Mn, Fe, Co, Ni, Cu, Mo, W

Co

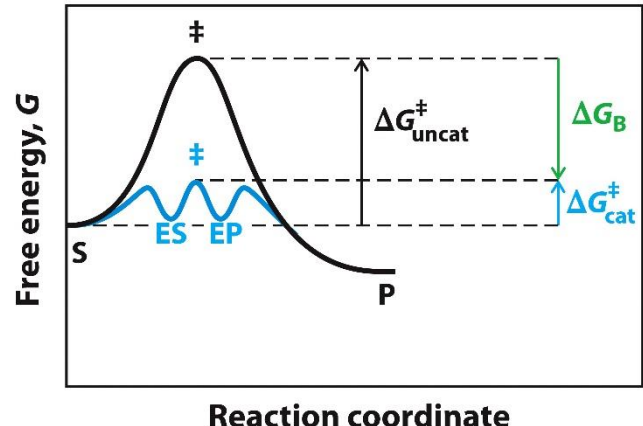
Non-enzymes

- M-transport
- Energy conversion
- Photosynthesis

Na, K, Fe

Mg

Mg

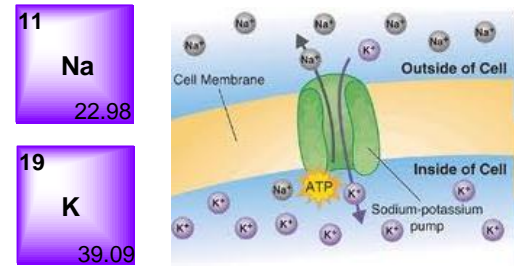


5. Metals in Biochemistry

The Alkaline Metal Cations Na^+ and K^+

Functions in biochemistry

- Maintenance of membrane potentials through concentration gradients of Na^+ and K^+ in cooperation with Cl^- and Ca^{2+} via the (muscular) cell membrane \Rightarrow signal transmission, kidney function
- Ion transport occurs via ion channels (passive or active)



Ion	Extracell. [mM]	Intracell. [mM]	Ratio	Membrane potential [mV]
Na^+	145	12	12	+68
K^+	4	155	0.026	-99
Cl^-	1.5	$< 10^{-7}$	> 15000	$> +128$
Ca^{2+}	123	4.2	30	-90

Membrane potential: $E = RT/zF \cdot \ln[c(\text{M}^{n+})_{\text{ec}}/c(\text{M}^{n+})_{\text{ic}}]$

with $F = \text{Faraday constant} = 96485 \text{ As/mol}$, $T = 310 \text{ K}$

Free enthalpy: $\Delta G = -z \cdot F \cdot E$

5. Metals in Biochemistry

The Alkaline Metal Cations Na^+ and K^+

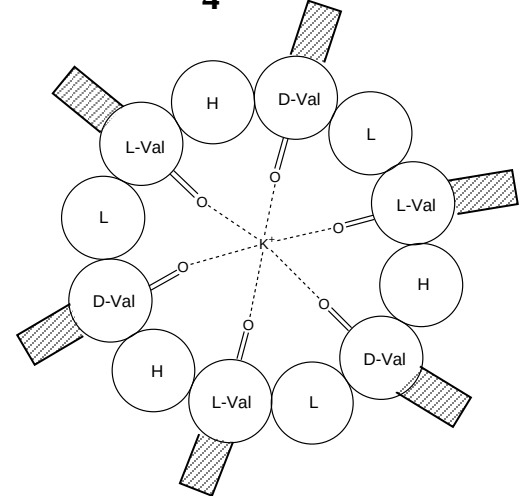
Na^+ functions (in extracellular fluid)

- Electrical impulses along nerve systems (see above)
- Osmotic balance „sodium pump”
- Acid-base balance
- Conformation of proteins and nucleic acids



K^+ functions (in intracellular fluid)

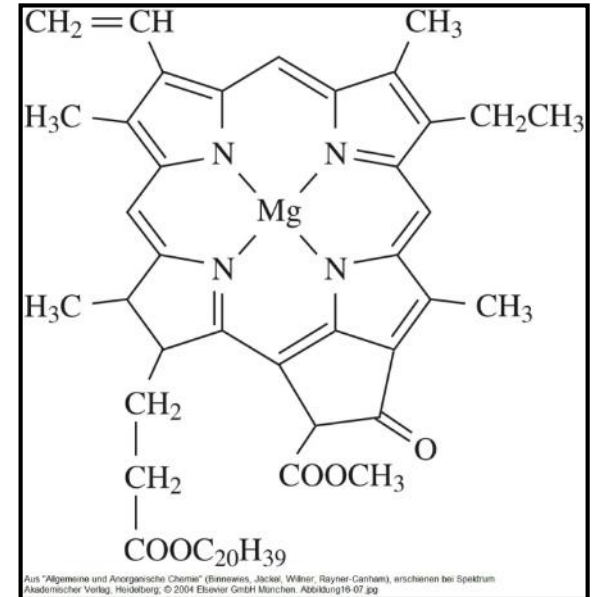
- Enzyme activator
- Conformation of proteins and RNA
- Secretion of gastric acid
- Transmembrane potentials
- Cyclic antibiotics: **Valinomycin, Monactin, Nonactin**



5. Metals in Biochemistry

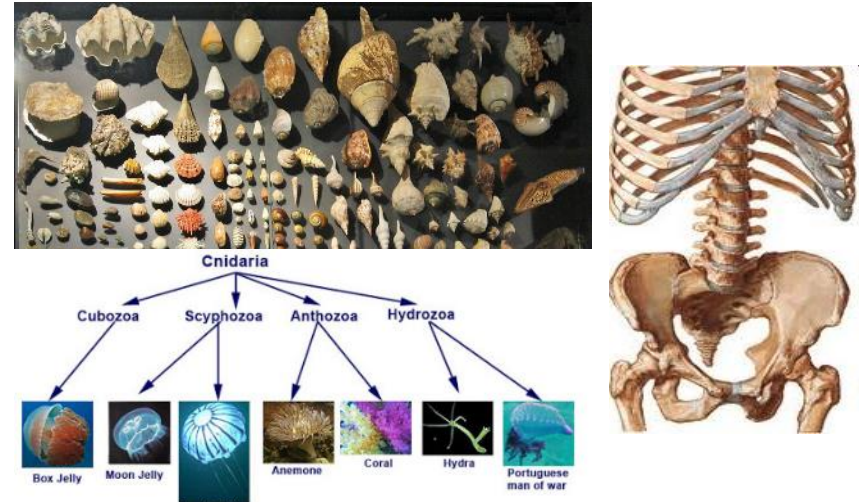
Magnesium Cations

- Metal centres in chlorophyll (photosynthesis)
- In active centres of ATPases and other enzymes
⇒ PCR (Polymerase Chain Reaction)
- Intracellular fluids



Calcium Cations

- Extracellular fluids
- Of importance for blood coagulation and muscle contraction
- Exoskeleton: CaCO₃
 - Mollusca (scallops, snails)
 - Cnidaria (corals, jelly fish)
- Endoskeletons: Ca₅(PO₄)₃X with X = OH, F, Cl
 - Chordata or vertebrates (vertebrates)
 - Cephalopoda



5. Metals in Biochemistry

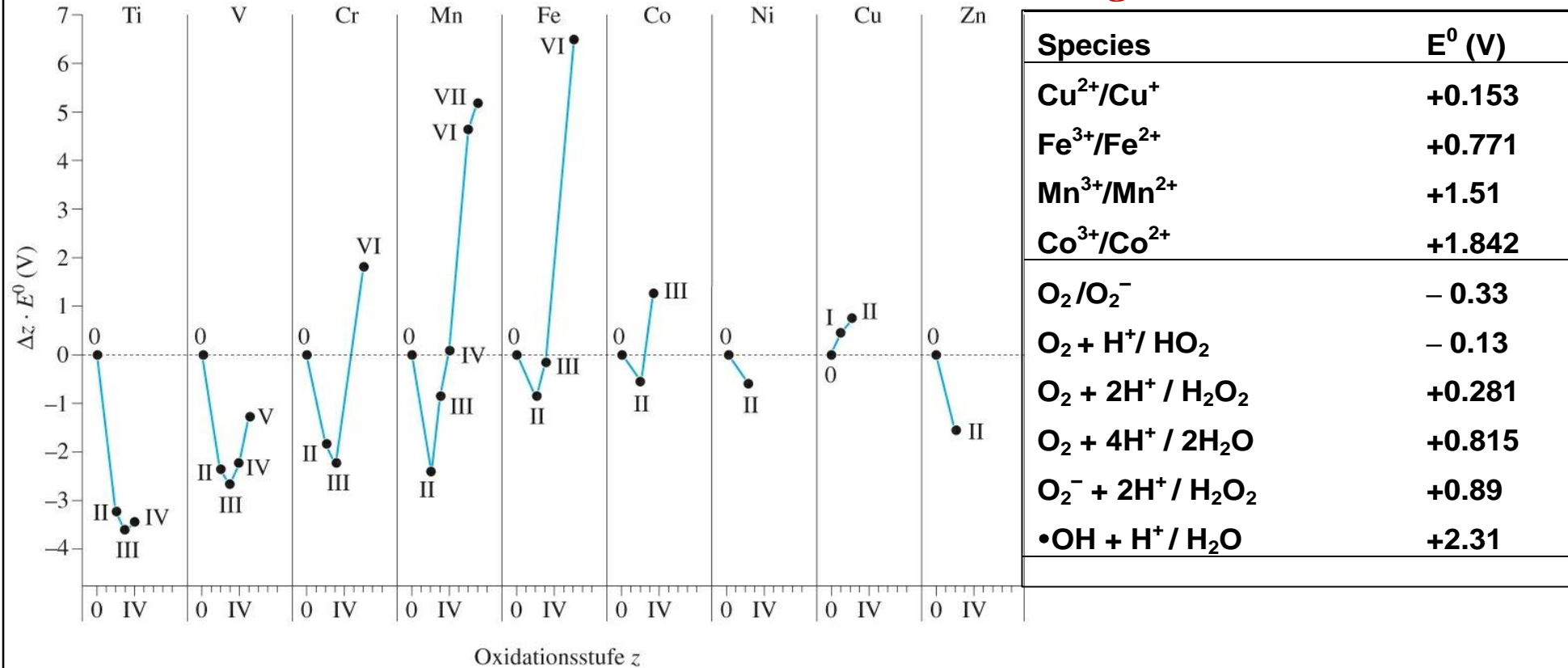
Oxidation States of Transitions Metals (TM)

3d-Elements	Sc +3	Ti +2, +3, +4	V +2,+3, +4, +5	Cr +2, +3, +6	Mn +2, +3 +4, +7	Fe +2, +3	Co +2, +3	Ni +2	Cu +1, +2	Zn +2
4d-Elements	Y +3	Zr +4	Nb +5	Mo +4, +6	Tc* +7	Ru +2, +3, +4	Rh +3	Pd +2	Ag +1, +2	Cd +2
5d-Elements	La +3	Hf +4	Ta +5	W +4, +6	Re +4, +7	Os +4, +8	Ir +3, +4	Pt +2, +4	Au +1, +3	Hg +1, +2

- All TM, which are sufficiently abundant within earth's crust and possess relatively stable oxidation states, are readily soluble and thus biologically available and are of importance as trace elements
- Many metals in high oxidation states form poorly soluble oxides $\Rightarrow \text{TiO}_2, \text{ZrO}_2, \text{HfO}_2, \text{Nb}_2\text{O}_5, \text{Ta}_2\text{O}_5, \text{MnO}_2, \text{RuO}_2, \text{OsO}_4, \text{IrO}_2$
- Metals, that are most stable at high oxidation states, occur in the earth's crust as poorly soluble oxides and are thus not biologically available (Ti, Zr, Hf)

5. Metals in Biochemistry

Oxidation States of 3d-Metals in Acidic Solution (Frost-Diagram)



- The metals are reducing agents, with titanium being the strongest reducing agent and copper in cationic form as (half) noble metal being slightly oxidising.
- Cr^{VI} is a strong and Mn^{VI} , Mn^{VII} and Fe^{VI} are extremely strong oxidising agents
- Relatively stable oxidation states: $\text{Ti}^{\text{III/IV}}$, $\text{V}^{\text{III/IV}}$, Cr^{III} , $\text{Mn}^{\text{II/III/IV}}$, $\text{Fe}^{\text{II/III}}$, $\text{Co}^{\text{II/III}}$, Ni^{II} , $\text{Cu}^{\text{I/II}}$, Zn^{II}

5. Metals in Biochemistry

Mn²⁺-Ions: Labile Complexes with Highly Variable Coordination Sphere

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.97 ^[hs-6o]
Configuration	[Ar]3d ⁵
CFSE/Dq	0
Preferred coordination	(N,O) ₆
Species at pH 7	[Mn(H ₂ O) ₆] ²⁺
pK _A of aqua-ion	10.6
+ NH ₃	Mn(OH) ₂
+ H ₂ S + metal salt/NH ₃ -solution	MnS
+ CN ⁻	ls-[Mn(CN) ₆] ⁴⁻



5. Metals in Biochemistry

Mn³⁺-Ions: Complexes with Strong Distortion (Jahn-Teller-Ion)

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.785 ^[hs-6]
Configuration	[Ar]3d ⁴
CFSE/Dq	> 0
Preferred coordination	O ₄₋₆ , JT-distorted
Species at pH 7	Mn(OH) ₃ ; MnOOH
pK _A of aqua-ion	0.7
+ NH ₃	Mn(OH) ₃ , MnOOH?
+ H ₂ S + metal salt/NH ₃ -solution	?
+ CN ⁻	ls-[Mn(CN) ₆] ³⁻



5. Metals in Biochemistry

Mn⁴⁺-Ions: Kinetically Stable Complexes with High Oxidation Strength

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.67 ^[60]
Configuration	[Ar]3d ³
CFSE/Dq	-12
Preferred coordination	O ₆
Species at pH 7	MnO(OH) ₂ , MnO ₂
pK _A of aqua-ion	-
+ NH ₃	-?
+ H ₂ S + metal salt/NH ₃ -solution	?
+ CN ⁻	-?



5. Metals in Biochemistry

Fe²⁺-Ions: Predominantly High-Spin Complexes

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.77 ^[hs-4t] , 0.78 ^[hs-4sp] , 0.92 ^[hs-6o] , 0.75 ^[ls-6o]
Configuration	[Ar]3d ⁶
CFSE/Dq	hs: -4, ls: -24
Preferred coordination	hs: (N,O) ₅₋₆ , S ₄ , ls: N ₆ , N ₅ (O,S)
Dissolved species at pH 7	[Fe(H ₂ O) ₅₋₆] ²⁺
pK _A of aqua-ion	9.5
+ NH ₃	almost [Fe(NH ₃) ₄₋₆] ²⁺ , Fe(OH) ₂
+ H ₂ S + metal salt/NH ₃ -solution	FeS
+ CN ⁻	[Fe(CN) ₆] ⁴⁻



Abb. "Bioinorganische Chemie" (Binnagow, Jüstel, Wöhrel, Bönnighausen), erschienen bei Spektrum
Akademischen Verlag Heidelberg © 2004 Elsevier GmbH München; Fotodok 41050 pg

5. Metals in Biochemistry

Fe³⁺-Ions: Formation of Poorly Soluble Fe(OH)₃ Favours the Formation of Highly Stable Complexes

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.63 ^[hs-4t] , 0.785 ^[hs-6o] , 0.69 ^[ls-6o]
Configuration	[Ar]3d ⁵
CFSE/Dq	hs: 0, [ls-6o]: -20
Preferred coordination	hs: O ₆ , ls: (N,O) ₆
Dissolved species at pH 7	Fe(OH) ₃
pK _A of aqua-ion	2.2
+ NH ₃	Fe(OH) ₃
+ H ₂ S + metal salt/NH ₃ -solution	Fe(OH) ₃
+ CN ⁻	ls-[Fe(CN) ₆] ³⁻



Zu "Allgemeine und Anorganische Chemie" (Binnwies, Jäckel, Wilher, Raynor-Carham), erschienen bei Spektrum
Akademischer Verlag, Heidelberg, © 2004 Elsevier GmbH München. FeO3.jpg

5. Metals in Biochemistry

Ni²⁺-Ions: Kinetically Stable Octahedral Complexes

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.69 ^[4] , 0.83 ^[60]
Configuration	[Ar]3d ⁸
CFSE/Dq	-12
Preferred coordination	(N,O) ₆
Solution at pH 7	[Ni(H ₂ O) ₆] ²⁺
pK _A of aqua-ion	9.9
+ NH ₃	[Ni(NH ₃) ₆] ²⁺
+ H ₂ S + metal salt/NH ₃ -solution	NiS
+ CN ⁻	[Ni(CN) ₄] ²⁻ , [Ni(CN) ₅] ³⁻



Zu "Allgemeine und Anorganische Chemie" (Binnigstet, Jackel, Witte, Rayner-Gerhart), erschienen bei Spektrum
Akademischer Verlag, Heidelberg. © 2004 Elsevier GmbH München. [Ni(209)]504.jpg

5. Metals in Biochemistry

Cu⁺-Ions: Kinetically Labile Complexes of Tetrahedral Structure

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.74 ^[4t] , 0.91 ^[6o]
Configuration	[Ar]3d ¹⁰
CFSE/Dq	0
Preferred coordination	N ₄ , S ₄
Solution at pH 7	Disproportion
pK _A of aqua-ion	—
E ^{0'}	0.1 V
+ NH ₃	[Cu(NH ₃) ₄] ⁺
+ H ₂ S + metal salt/NH ₃ -solution	Cu ₂ S
+ CN [−]	[Cu(CN) ₄] ^{3−}

5. Metals in Biochemistry

Cu²⁺-Ions: Complexes with Strong Distortion (Jahn-Teller-Ion)

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.71 ^[4] , 0.79 ^[5] , 0.87 ^[60]
Configuration	[Ar]3d ⁹
CFSE/Dq	> 0
Preferred coordination	N/O ₄₋₆ , JT-distorted
Solution at pH 7	[Cu(H ₂ O) ₅] ²⁺
pK _A of aqua-ion	8.0
E ^{0'}	0.1 V
+ NH ₃	[Cu(NH ₃) ₄ (H ₂ O) ₂] ²⁺
+ H ₂ S + metal salt/NH ₃ -solution	“CuS“ contains Cu ^{II} and S ²⁻ next to Cu ^I and S ₂ ²⁻
+ CN ⁻	Reduction to [Cu(CN) ₄] ³⁻

CuCl₂ (Dr. J.N. Keil 2022)



5. Metals in Biochemistry

Zn²⁺-Ions: Labile Complexes with Variable Coordination Geometry

Properties	Typical Values / Products
Ionic radius/ \AA [coordination]	0.74 ^[4t] , 0.82 ^[5] , 0.88 ^[6o]
Configuration	[Ar]3d ¹⁰
CFSE/Dq	0
Preferred coordination	N ₄ , (N,O) ₅₋₆ , S ₄
Dissolved species at pH 7	[Zn(H ₂ O) ₅₋₆] ²⁺
pK _A of aqua-ion	9.0
+ NH ₃	[Zn(NH ₃) ₄] ²⁺
+ H ₂ S + metal salt /NH ₃ -solution	ZnS
+ CN ⁻	[Zn(CN) ₄] ²⁻

5. Metals in Biochemistry

Coordination of Biochemically Relevant Metal Cations

Cation	CN	Geometry	Biochemical ligands
Na^+	6	octahedral	O: ether, hydroxyl, carboxylate
K^+	6-8	flexible	O: ether, hydroxyl, carboxylate
Mg^{2+}	6	octahedral	O: carboxylate, phosphate
Ca^{2+}	6-8	flexible	O: carboxylate, carbonyl, phosphates
$\text{Mn}^{2+}(\text{d}^5)$	6	octahedral	O: carboxylate, phosphates N: imidazole
$\text{Mn}^{3+}(\text{d}^4)$	6	tetragonal	O: carboxylate, phosphate, hydroxide
$\text{Fe}^{2+}(\text{d}^6)$	4	tetrahedral	S: thiolate
	6	octahedral	O: carboxylate, alkoxide, oxide, phenolates N: imidazole, porphyrin
$\text{Fe}^{3+}(\text{d}^5)$	4	tetrahedral	S: thiolate
	6	octahedral	O: carboxylate, alkoxide, oxide, phenolates N: imidazole, porphyrin
$\text{Co}^{2+}(\text{d}^7)$	6	octahedral	O, carboxylate N, imidazole

5. Metals in Biochemistry

Coordination of Biochemically Relevant Metal Cations

Cation	CN	Geometry	Biochemical ligands
$\text{Ni}^{2+}(\text{d}^8)$	4	square-planar	S: thiolate N: imidazole, polypyrrole rare!
$\text{Cu}^+(\text{d}^{10})$	6	octahedral	
	4	tetrahedral	S: thiolate, thioether N: imidazole
$\text{Cu}^{2+}(\text{d}^9)$	4	tetrahedral	S: thiolate, thioether N: imidazole
	4	square-planar	O: carboxylate N: imidazole
	6	tetragonal (distorted octahedral)	O: carboxylate N: imidazole
$\text{Zn}^{2+}(\text{d}^{10})$	4	tetrahedral	O: carboxylate, carbonyl S: thiolate N: imidazole
	5	square-pyramidal	O: carboxylate, carbonyl N: imidazole

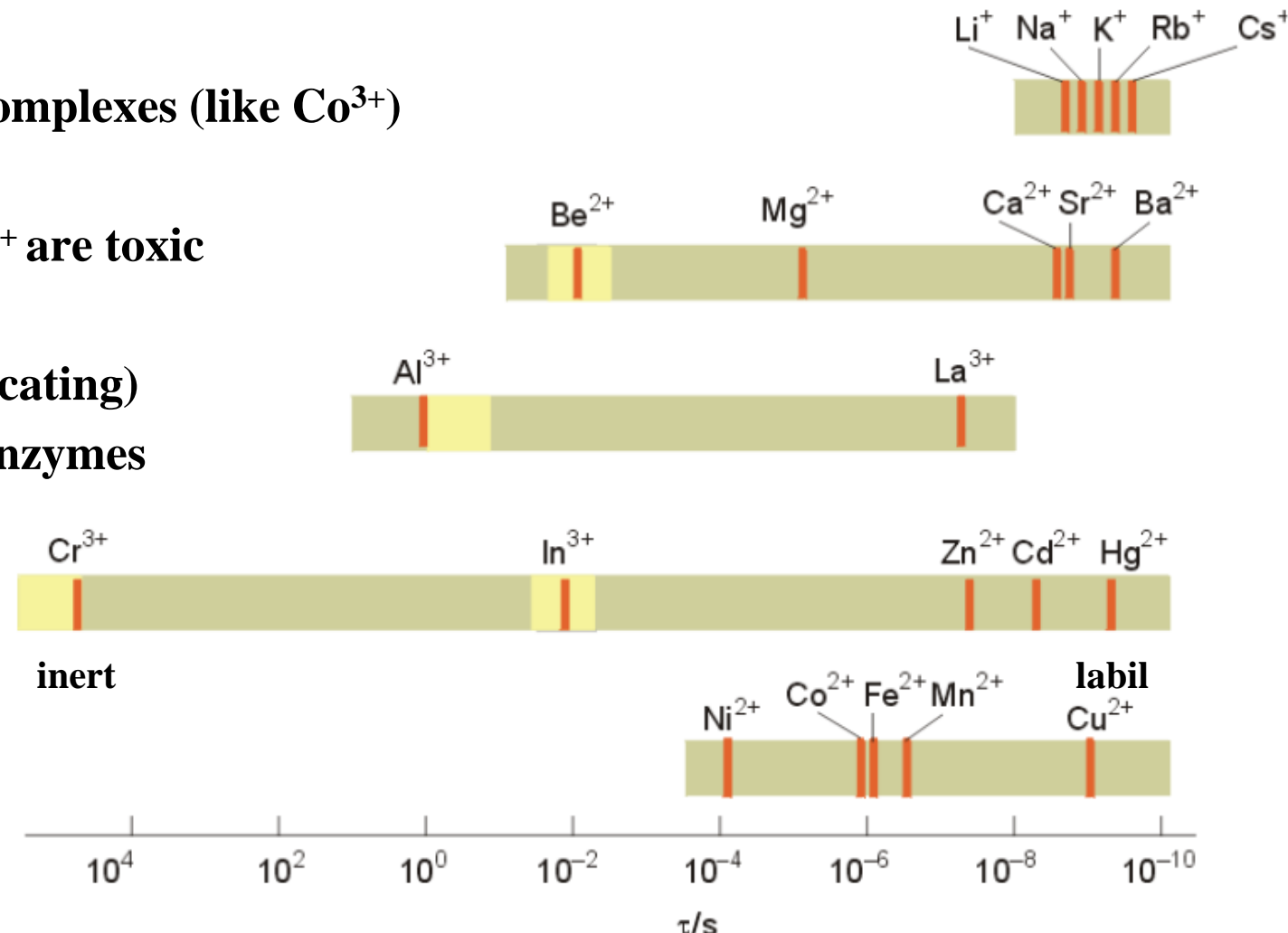
5. Metals in Biochemistry

Kinetic Aspects – Lifetime of Aqua Ligands to Metal Bonds [s]

- Cr^{3+} forms inert complexes (like Co^{3+})

- In^{3+} , Cd^{2+} and Hg^{2+} are toxic

- Be^{2+} blocks (intoxicating)
 Mg^{2+} -containing enzymes



6. Ligands in Biochemistry

Small Inorganic Molecules \Rightarrow “Hard Ligands“

H_2O	
OH^-	
O^{2-}	
CO_2/CO_3^{2-}	
O_2^{2-}	
HO_2^-	
O_2^-	
O_2	
NO	
NO_2	
CO	
S^{2-}	
CN^-	
OCN^-	
SCN^-	
N_3^-	
$N=C=N^2-$	

Aquo (aqua)

Hydroxy (hydroxido)

Oxo

Carbonato

Peroxy

Hydroperoxy

Superoxy

Dioxygenyl

Nitrosyl

Nitrito

Carbonyl

Sulfido

Cyanido

Cyanato

Thiocyanato

Azido

Cyanamide

C source

Oxidant

Oxidant

Radical

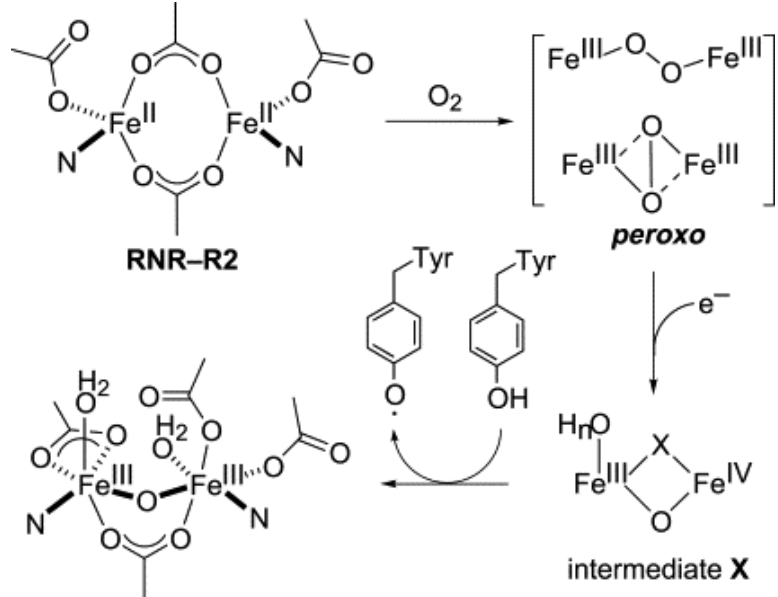
Radical

Radical

Radical

Strong bonding to Fe^{2+}

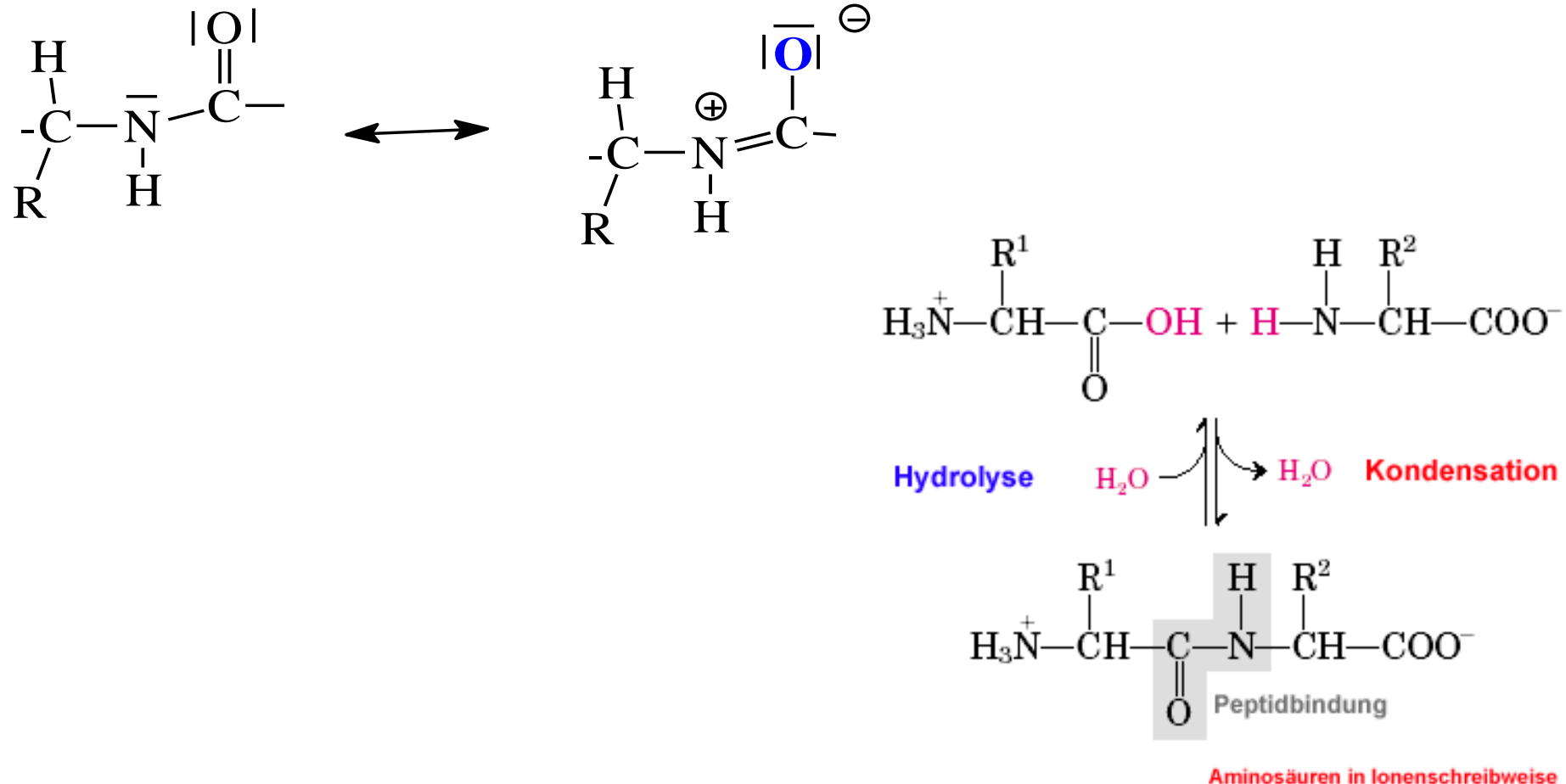
Strong binding to Ca^{2+} , Zn^{2+}



$CaCN_2$ as a fertiliser

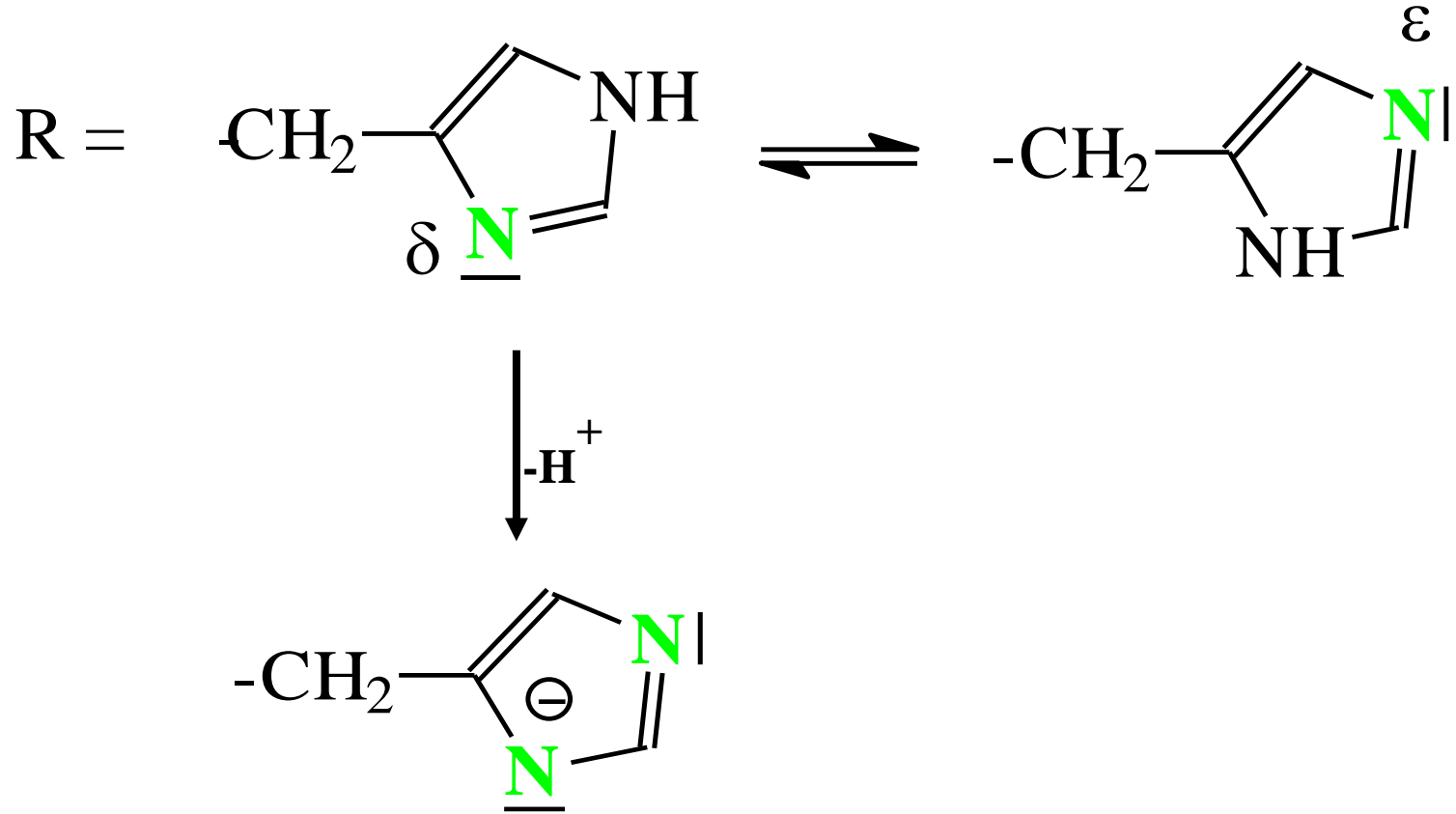
6. Ligands in Biochemistry

Polypeptides and the Peptide Bond R-NH-CO-R' \Rightarrow "Hard Ligands"



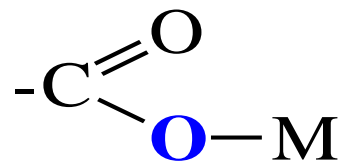
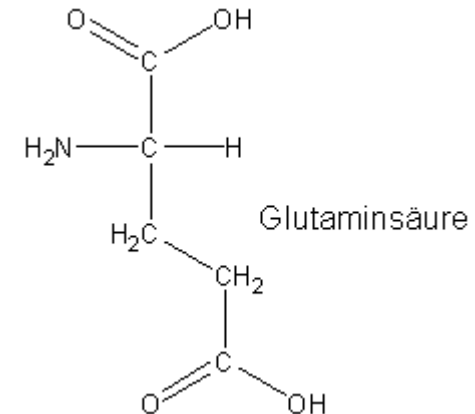
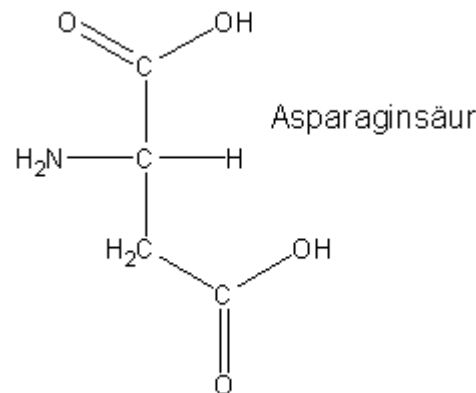
6. Ligands in Biochemistry

Histidine (His, H) \Rightarrow Neutral or Basic Ligands for Zn, Cu, Mn, Fe or Ni

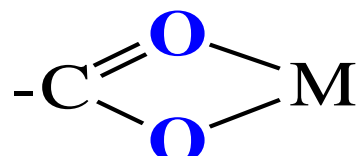


6. Ligands in Biochemistry

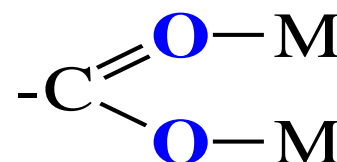
Aspartic Acid (Asp, D) and Glutamic Acid (Glu, E) \Rightarrow Acidic Ligands with High Affinity to Mg^{2+} and Ca^{2+}



end-on
einzähnig



side-on
(zeizähnig,
chelatartig)



end-on
verbrückend

6. Ligands in Biochemistry

Cysteine (Cys, C) and Glutamic Acid (Glu, E)

⇒ Soft Ligands with Affinity to Zn, Cu, Fe and Ni

Organic group R

-CH₂-SH **Cysteine (Cys, C)**

-CH₂-SeH **Selenocysteine (Sec, U)**

-CH₂-CH₂-S-CH₃ **Methionine (Met, M)**

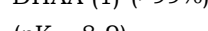
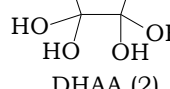
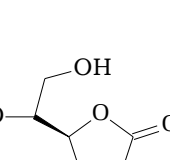
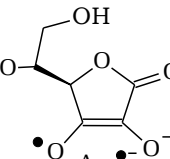
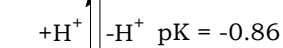
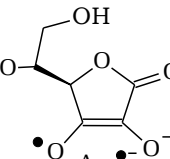
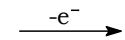
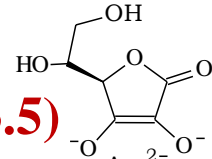
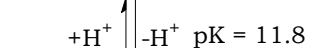
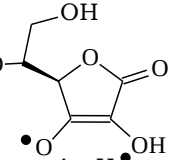
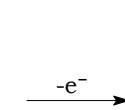
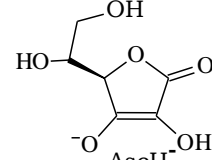
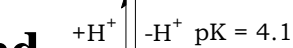
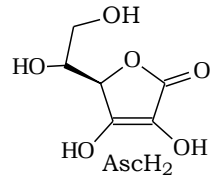
→ Relevant for protein folding

→ R-S-S-R or R-Se-Se-R bridges

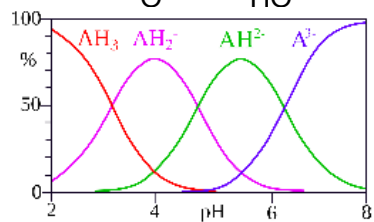
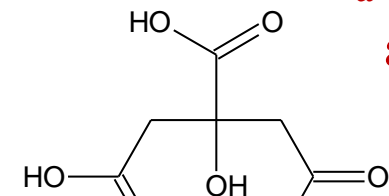
6. Ligands in Biochemistry

Ascorbic Acid (Vitamin C)

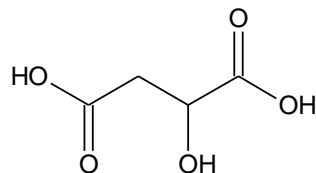
- Chelating ligand, e.g. for Fe^{3+}
- Reductive, i.e. it is a redox active ligand
- Radical scavenger
- Temperature sensitive
- Acidic character: $\text{pK}_a \sim 4.1$



Citric Acid ($\text{pK}_a \sim 3.1$)



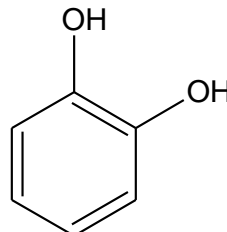
and Malic Acid ($\text{pK}_a \sim 3.5$)



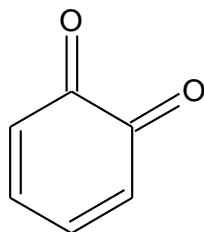
6. Ligands in Biochemistry

Siderophores \Rightarrow “Hard Ligands for Fe³⁺“

Catechol (o-hydroxyphenyl)

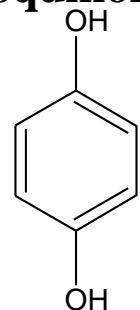


o-Quinone

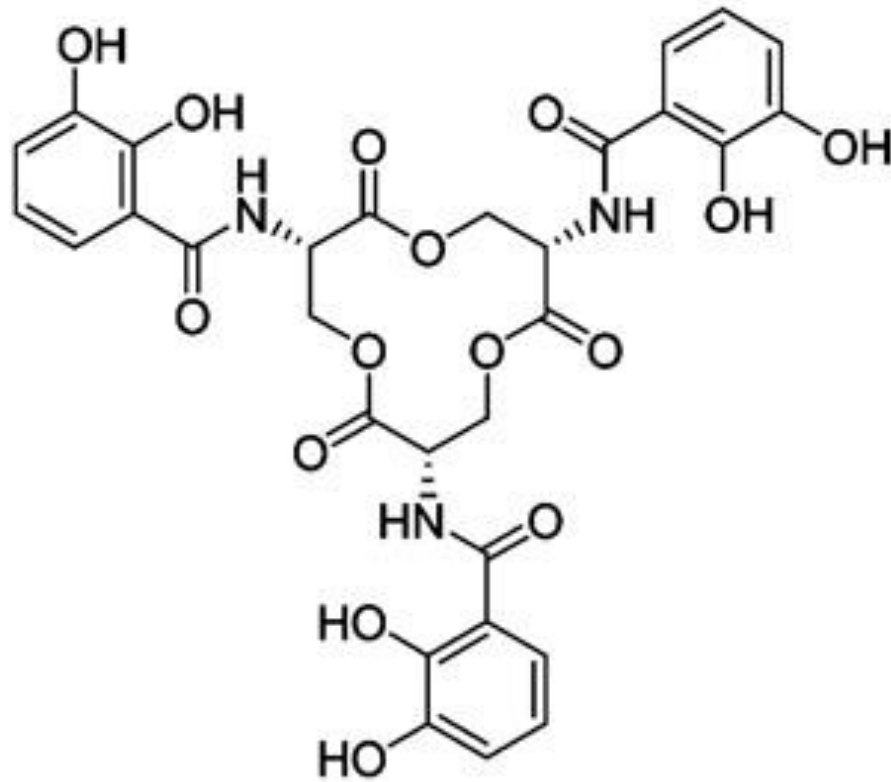
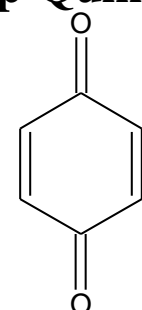


Oxidation results in semi-quinone and
then o-quinone, meaning this is a redox-active
Ligand, as well

Hydroquinone (p-hydroxyphenyl)



p-Quinone



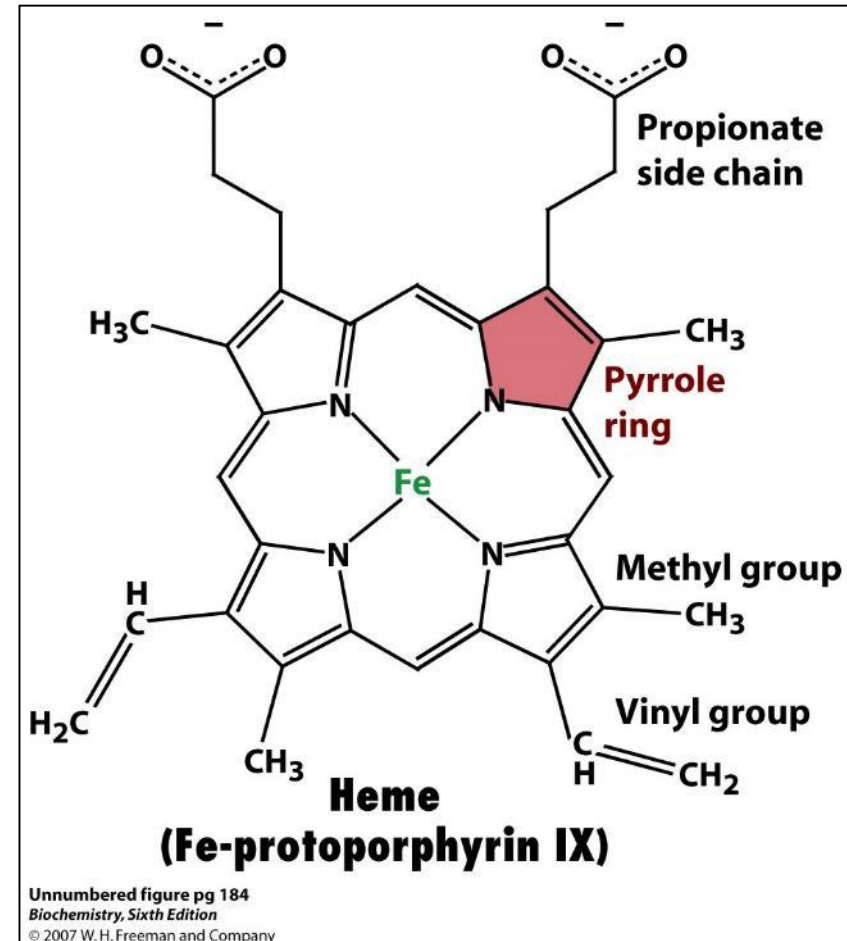
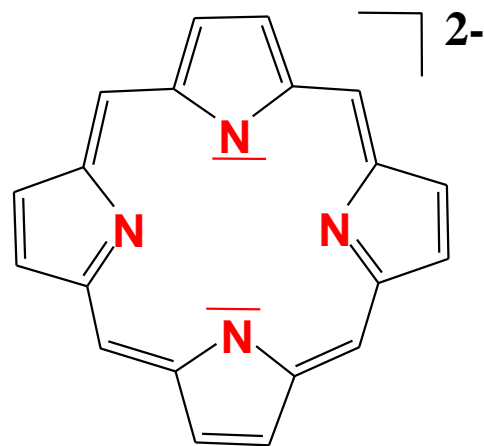
Enterobactin (E. coli)

$$\log K' (\text{pH } 7) = 25$$

6. Ligands in Biochemistry

Porphyrins → Macrocyclic Ligands in Heme Proteins

Hemoglobin (Hb), Myoglobin (Mb) $\Rightarrow \text{Fe}^{2+}$
Chlorophyll $\Rightarrow \text{Mg}^{2+}$



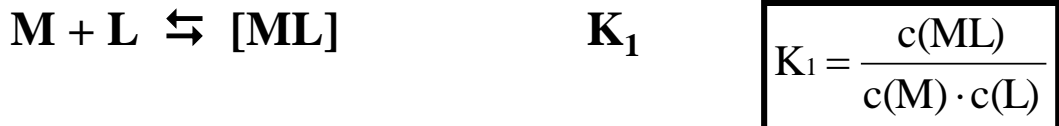
7. Principles in Coordination Chemistry

Thermodynamic Complex Stability

Complex equilibria in solution (cleavage of ligands)



Formation of a complex normally proceeds stepwise



and so on



total formation constant = $K_F = \beta_n$

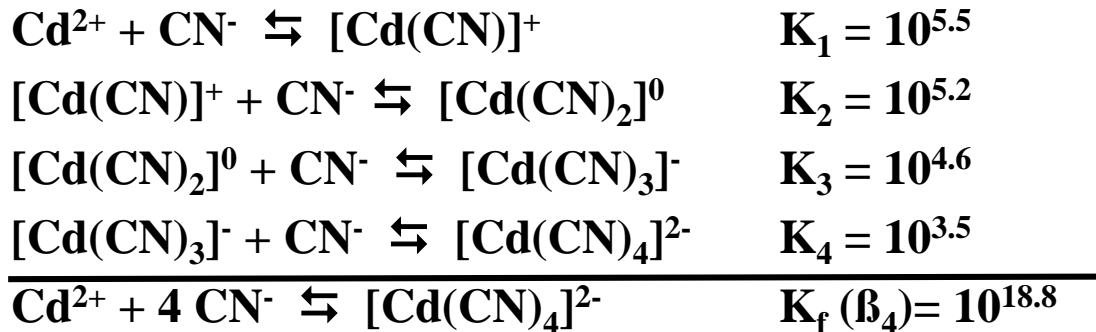
$$K_F = \beta_n = \prod_{i=1}^n K_i = \frac{c(\text{ML}_n)}{c(\text{M}) \cdot c(\text{L})^n}$$

Free reaction enthalpy $\Delta G_r^\circ = -R \cdot T \cdot \ln K$

7. Principles in Coordination Chemistry

Complex Stability utilizing $[Cd(CN)_4]^{2-}$ as an Example

Stepwise formation of complexes with Cd^{2+} and CN^- :



⇒ The complex formation constant K_n often declines with increasing degree of substitution!

Cause for that behaviour

- Sterical hindrance
- Coulomb-effect during the incorporation of charged ligands, i.e. CN^-
- Reduction of entropy through increased degree of order, i.e. $\Delta S_r^0 < 0$

7. Principles in Coordination Chemistry

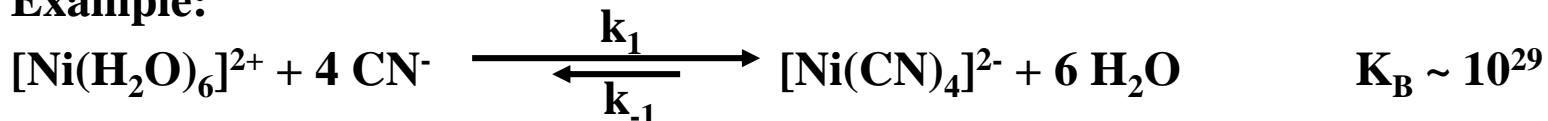
Thermodynamic and Kinetic Complex Stability

The **thermodynamic stability** is described by the complex formation constant K_f or β .

The higher this number the more stable the complex is (**unstable – stable**)

Free reaction enthalpy $\Delta G_r^0 = -RT \cdot \ln K_f$

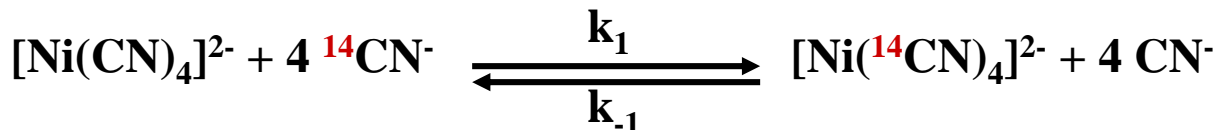
Example:



The equilibrium favours the right side, which means the complex is thermodynamically stable

Never the less ligand exchange happens fast, i.e. **kinetic complex stability is low (labile – inert)**

Free activation enthalpy ΔG_r^\neq



Eyring-equation:

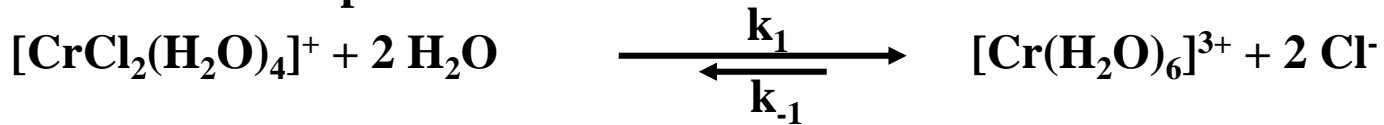
$$k = \frac{k_B \cdot T}{h} \cdot e^{\frac{-\Delta G^\#}{RT}}$$

$\tau_{1/2} = 30 \text{ s (fast)}$

7. Principles in Coordination Chemistry

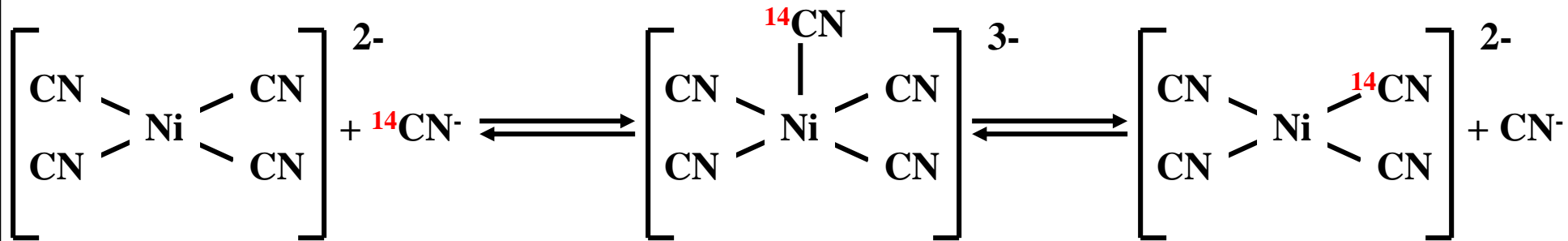
The Kinetic Complex Stability, i.e. the Reactivity of a Complex is Defined by the Structure and the Possible Reaction Pathway

Octahedral complexes



⇒ Very slow ligand exchange although the hexaquochromium(III)-complex is more stable

Square-planar complexes



⇒ Very fast ligand exchange although the thermodynamic driving force is zero

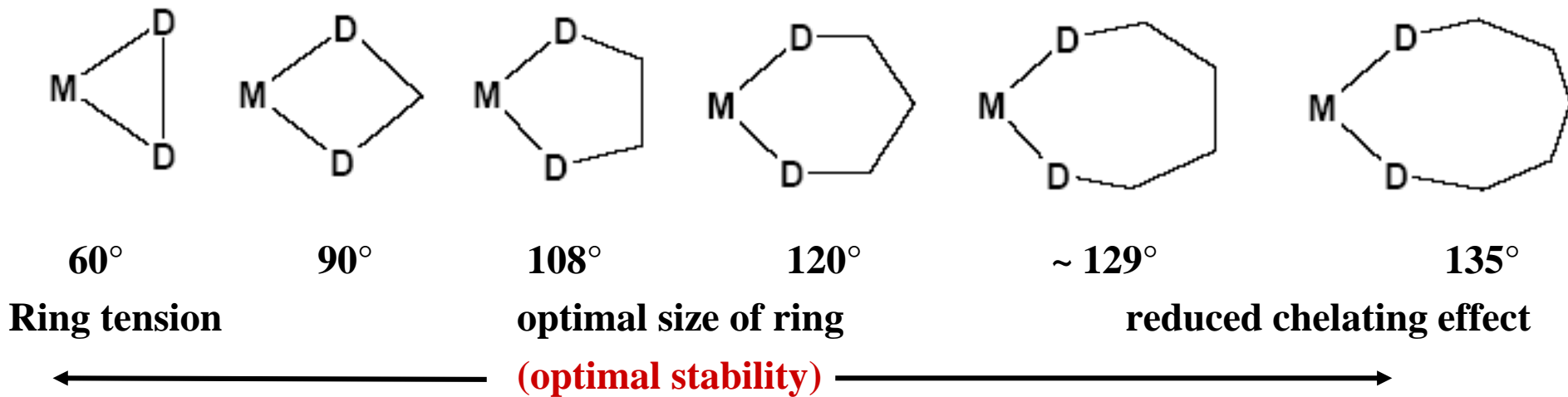
7. Principles in Coordination Chemistry

Chelating Ligands Form Extremely Stable Complexes

Bi-dental ligands are also called chelating ligands (Greek: *chele* = claw, “chelicerata”).

The ligand takes the metal in its “claws”. If thereby (chelated) rings with 5 or 6 members are formed, they are more stable than complexes formed by mono-dental ligands, because they are favoured by enthalpy. Furthermore during the chelating process, non-chelating ligands are set free, thus increasing entropy.

⇒ The chelating effect is thus also an entropic effect!



7. Principles in Coordination Chemistry

Chelating Ligands vs. Non-chelating Ligands

Complex formation by chelating ligands results in more stable complexes as in the case with mono-dental ligands



Formation of $[\text{Ni}(\text{NH}_3)_6]^{2+}$ -complex \Rightarrow particle number remains the same

Formation of $[\text{Ni}(\text{en})_3]^{2+}$ -complex \Rightarrow particle number increases $\Rightarrow \Delta S^\circ > 0$

Complex formation with a chelating ligand leads to increased entropy!

ΔH is comparable for both cases

$$\Delta G^\circ = \Delta H^\circ - T\Delta S^\circ \text{ and } \Delta G^\circ = -RT \cdot \ln K_K \\ (\Delta \Delta G^\circ = 0 - T\Delta \Delta S^\circ)$$

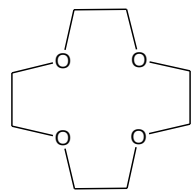
with K_K = complex formation constant

Formation of $[\text{Ni}(\text{en})_3]^{2+}$ \Rightarrow more negative $\Delta G^\circ \Rightarrow$ larger K_K

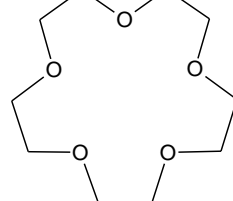
7. Principles in Coordination Chemistry

Macrocyclic Ligands (suitable for model complexes!)

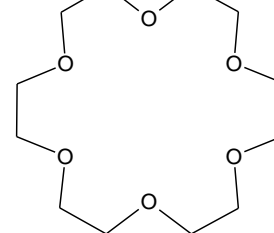
Cyclic chelating ligands, that form highly stable complexes, due to their rigidity and dentality



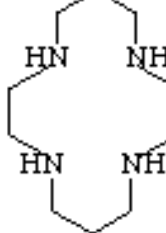
12-Crown-4



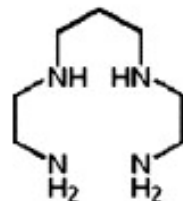
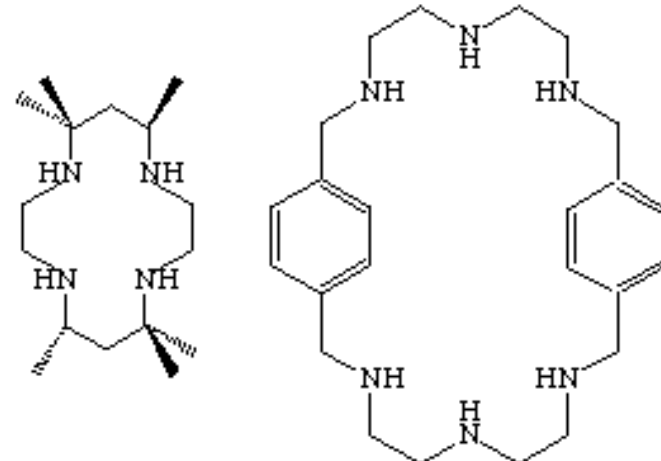
15-Crown-5



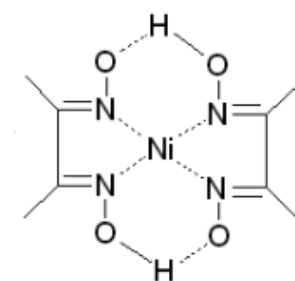
18-Crown-6



[14]aneN₄



2,3,2-tet



Complex



$\log K_{\text{NiL}}$

15.8



22.2



14.6

7. Principles in Coordination Chemistry

Dependence of Stability Constants of Metal Complexes

1. Central atoms



⇒ Correlates with decreasing cation radius / increasing ionic charge density
(Irving-Williams stability series)

2. Ligands

- Chelating effect, macrocyclic effect
- Polarizability (hard vs. soft), backbonding

Hard and Soft Acids and Bases HSAB concept (R.G. Pearson 1963)

- Metal atoms = Acids (electron acceptors)
- Ligands = Bases (electron donators)
- High stability: soft metal atoms - soft ligands
 - hard metal atoms - hard ligands
- Low stability: soft metal atoms - hard ligands
 - hard metal atoms - soft ligands

7. Principles in Coordination Chemistry

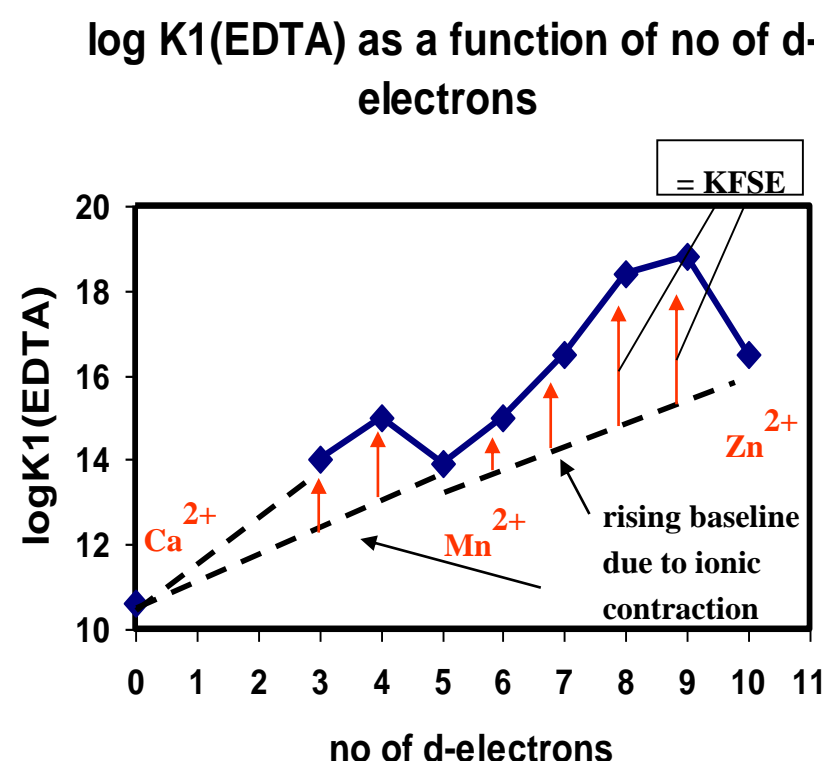
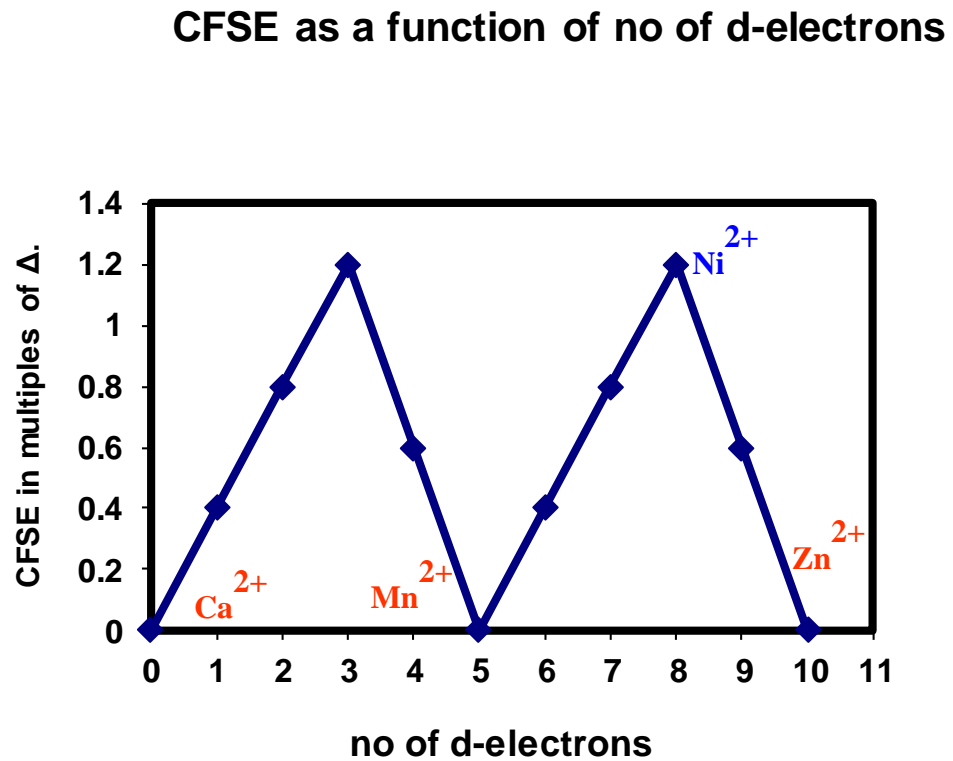
HSAB Concept: Classification of Metal Atoms (Acids) and Ligands (Bases)

	Bases	Acids
Hard	NH_3 , R-NH_2 , N_2H_4 , H_2O , OH^- , O^{2-} , R-OH , RO^- , R_2O , CO_3^{2-} , R-COO^- , NO_3^- , PO_4^{3-} , SO_4^{2-} , ClO_4^- , F^- , Cl^- <i>poorly deformable electron shells</i>	H^+ , Li^+ , Na^+ , K^+ , Ba^{2+} , Mg^{2+} , Ca^{2+} , Sr^{2+} , Ti^{3+} , Ti^{4+} , Zr^{4+} , VO^{3+} , Cr^{3+} , Cr^{6+} , Mn^{2+} , Mn^{4+} , Mn^{7+} , Fe^{3+} , Co^{3+} , Al^{3+} , Ga^{3+} , In^{3+} <i>are highly polarizing</i>
Intermediates	N_3^- , N_2 , Ph-NH_2 , NO_2^- , Br^- $\text{C}_5\text{H}_5\text{N}$, SO_3^{2-} , imidazole, aniline	Fe^{2+} , Co^{2+} , Ni^{2+} , Cu^{2+} , Zn^{2+} , Rh^{3+} , Ir^{3+} , Ru^{3+} , Sn^{2+} , Pb^{2+}
Soft	H^- , R^- , CN^- , CO , SCN^- , R_3P , RSH , R_2S , RS^- , S_2O_3^- , I^- , RNC , $(\text{RS})_2\text{PO}_2^-$ <i>easily deformable electron shells</i>	Pd^{2+} , Pt^{2+} , Cu^+ , Ag^+ , Au^+ , Hg^+ , Hg^{2+} , Tl^+ , Me^0 , Cd^{2+} <i>are weakly polarizing</i>

7. Principles in Coordination Chemistry

Dependence of Stability Constants of Metal Complexes

3. Crystal field stabilisation energy (illustrated by pseudo-octahedral EDTA-complexes)



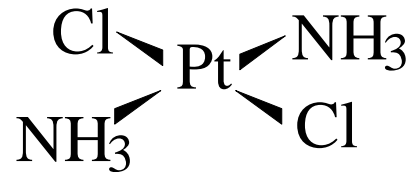
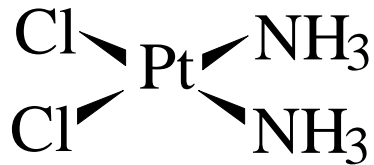
7. Principles in Coordination Chemistry

Dependence of Stability Constants of Metal Complexes

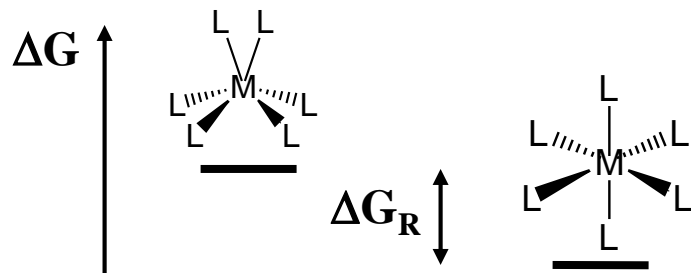
4. Coordination geometry

CN = 4: Tetrahedral coordination is favoured by d⁰, d⁵, d⁷ and d¹⁰

Square-planar coordination is favoured by d⁸ and d⁹ → cis/trans-isomerism



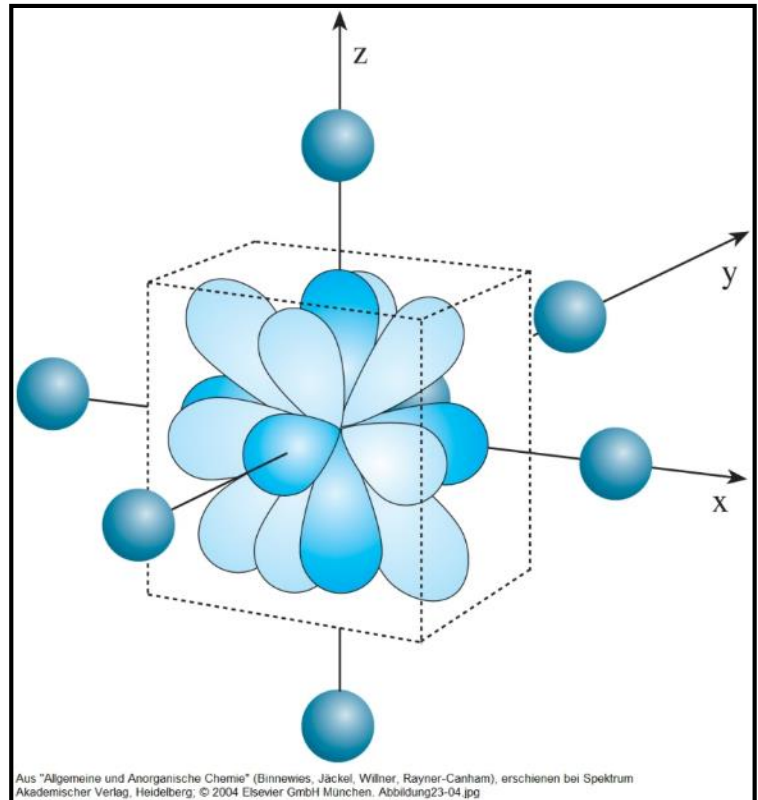
CN = 6: Octahedral more stable than trigonal-prismatic coordination



7. Principles in Coordination Chemistry

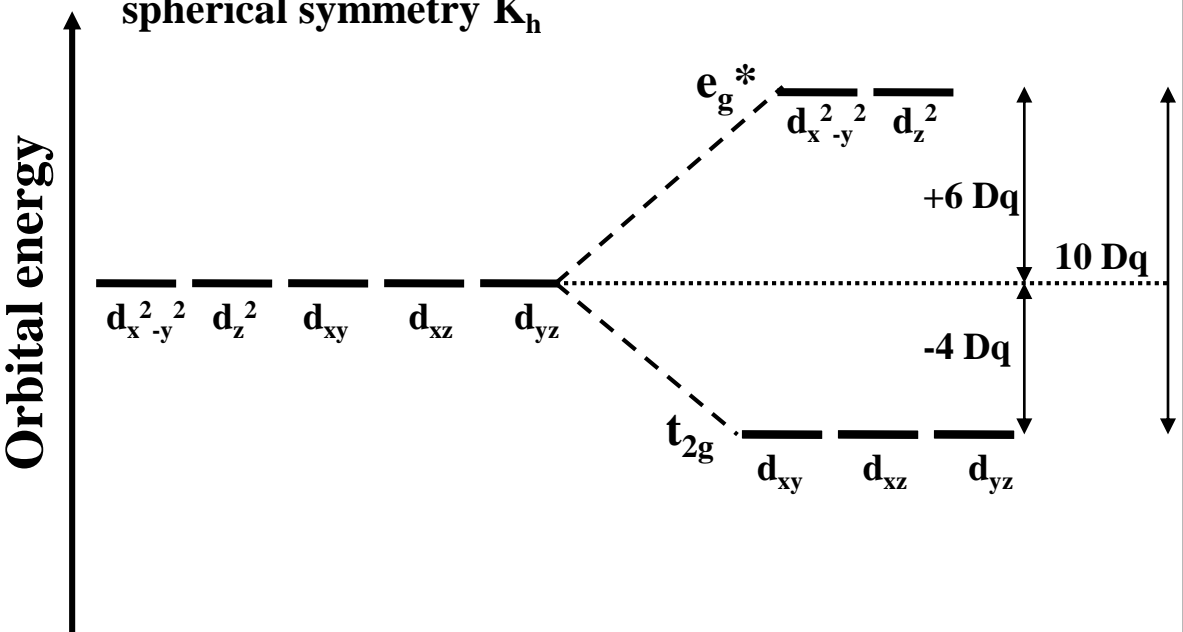
Crystal Field Splitting in Octahedral Field

$$Dq = z \cdot e^2 \cdot r^4 / (6 \cdot a \cdot R^5)$$



Ion in field of
spherical symmetry K_h

Ion in octahedral field O_h



Aus "Allgemeine und Anorganische Chemie" (Binnewies, Jäckel, Wilner, Rayner-Canham), erschienen bei Spektrum Akademischer Verlag, Heidelberg. © 2004 Elsevier GmbH München. Abbildung 23-04.jpg

The energy gap between the t_{2g} and e_g^* -orbitals is called **10 Dq** or Δ_0

7. Principles in Coordination Chemistry

Crystal Field Stabilisation Energy (CFSE)

Crystal field stabilisation energy in octahedral crystal field

$$\text{CFSE} = x(-4 Dq_o) + y(+6 Dq_o) + P \quad \text{with} \quad \begin{aligned} P &= \text{spin-pairing energy} \\ x &= \text{number of electrons in } t_{2g} \\ y &= \text{number of electrons in } e_g^* \end{aligned}$$

d^n	CFSE high-spin	CFSE low-spin	ΔCFSE	Examples
0	$0 Dq_o$	$0 Dq_o$	-	$\text{Sc}^{3+}, \text{Y}^{3+}, \text{Ln}^{3+}, \text{Ti}^{4+}$
1	$-4 Dq_o$	$-4 Dq_o$	-	Ti^{3+}
2	$-8 Dq_o$	$-8 Dq_o$	-	V^{3+}
3	$-12 Dq_o$	$-12 Dq_o$	-	$\text{Cr}^{3+}, \text{Mo}^{3+}, \text{W}^{3+}$
4	$-6 Dq_o$	$-16 Dq_o + 1 P$	$-10 Dq_o + 1 P$	Mn^{3+}
5	$0 Dq_o$	$-20 Dq_o + 2 P$	$-20 Dq_o + 2 P$	$\text{Mn}^{2+}, \text{Fe}^{3+}, \text{Ru}^{3+}$
6	$-4 Dq_o + 1 P$	$-24 Dq_o + 3 P$	$-20 Dq_o + 2 P$	$\text{Fe}^{2+}, \text{Co}^{3+}, \text{Ru}^{2+}, \text{Ir}^{3+}$
7	$-8 Dq_o + 2 P$	$-18 Dq_o + 3 P$	$-10 Dq_o + 1 P$	Co^{2+}
8	$-12 Dq_o + 3 P$	$-12 Dq_o + 3 P$	-	Ni^{2+}
9	$-6 Dq_o + 4 P$	$-6 Dq_o + 4 P$	-	Cu^{2+}
10	$0 Dq_o + 5 P$	$0 Dq_o + 5 P$	-	$\text{Cu}^+, \text{Zn}^{2+}$

Low-spin complexes are favoured if $10 Dq > P \Rightarrow \Delta_O \text{total is always negative!}$

7. Principles in Coordination Chemistry

Spin-Pairing Energy (P)

$$P_{\text{total}} = P_c + P_e \text{ with}$$

P_c = Coulomb's repulsive energy

Electronic repulsion in an orbital

$3d > 4d > 5d$, since orbitals become ever more diffuse

P_e = Loss of exchange energy (quantum mechanical part)

~ number of possibilities n to arrange electrons with parallel spin in pairs

$$P_e = [n(n-1)/2] * E_{\text{ex}} \text{ with } E_{\text{ex}} = \text{average exchange energy}$$

d^n	P_e high-spin [E_{ex}]	P_e low-spin [E_{ex}]	ΔP_e [E_{ex}]	ΔCFSE	ΔP_e per 10 Dq_0 [E_{ex}]
4	 $4(4-1)/2 = 6$	 $3(3-1)/2 = 3$	3	-10 Dq_0	3
5	 $5(5-1)/2 = 10$	 $3(3-1)/2 + 2(2-1)/2 = 4$	6	-20 Dq_0	3
6	 $5(5-1)/2 = 10$	 $3(3-1)/2 + 3(3-1)/2 = 6$	4	-20 Dq_0	2
7	 $5(5-1)/2 + 2(2-1)/2 = 11$	 $4(4-1)/2 + 3(3-1)/2 = 9$	2	-10 Dq_0	2

7. Principles in Coordination Chemistry

Spin-Pairing Energy – Relevance for Biochemically Important TM-Cations

- For ions with d^4 - or d^5 -configuration the loss of exchange energy with regard to 10 Dq is most pronounced
- Ions with d^6 - or d^7 -configuration form low-spin complexes even for weak ligand field:
 P_{ges} for $d^6 < d^7 < d^4 < d^5$
- P_{ges} for d^7 -ions is somewhat higher than for d^6 -ions, since P_c is greater

d^n	Free Ion	P_c [cm $^{-1}$]	P_{ex} [cm $^{-1}$]	P_{ges} [cm $^{-1}$]	
4	Cr $^{2+}$	5950	14475	20425	Numbers according to L.E. Orgel J. Phys. Chem. 23 (1955) 1819
	Mn $^{3+}$	7350	17865	25125	
5	Mn $^{2+}$	7610	16215	23825	J. Inorg. Nucl. Chem. 2 (1956) 229
	Fe $^{3+}$	10050	19825	29875	
6	Fe $^{2+}$	7460	11690	19150	Numbers for complexed ions are 15-30% smaller due to the nephelauxetic effect of the ligands!
	Co $^{3+}$	9450	14175	23625	
7	Co $^{2+}$	8400	12400	20800	

Fe $^{2+}$ forms low-spin complexes, even in weak crystal fields,
whereas Fe $^{3+}$ often forms high-spin complexes despite its higher ionic charge!

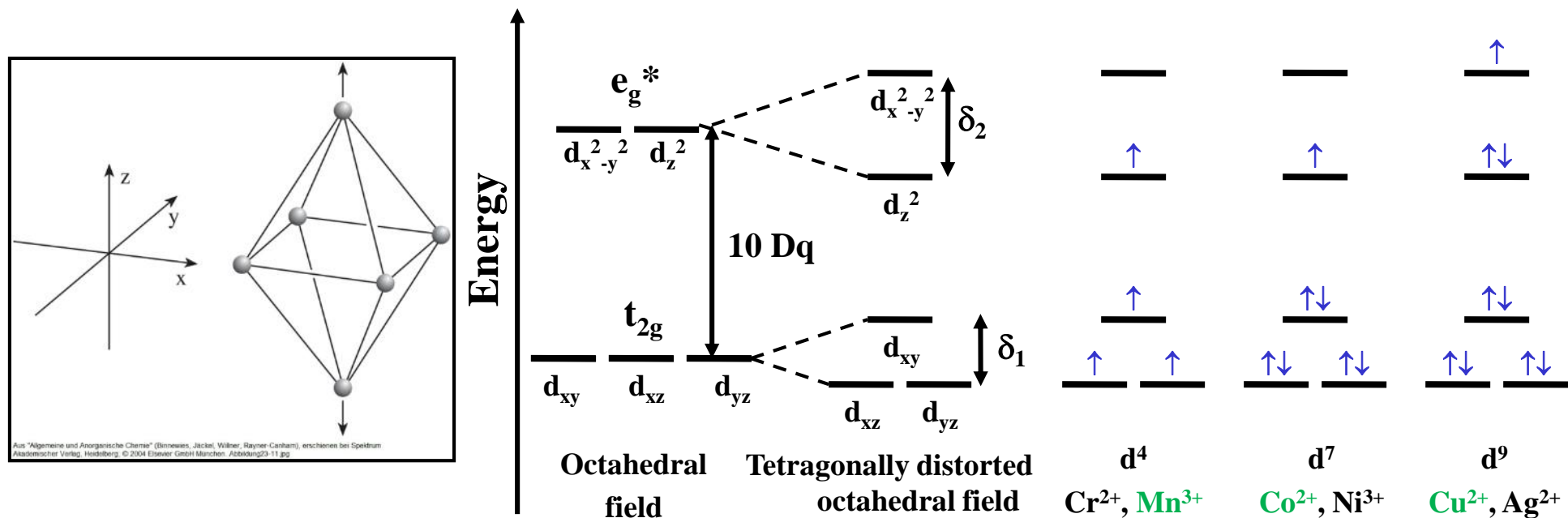
7. Principles in Coordination Chemistry

Crystal Field Splitting in Octahedral Field

Cause: Degeneracy of electronic states

Jahn-Teller-Theorem (Hermann Arthur Jahn and Edward Teller, 1937)

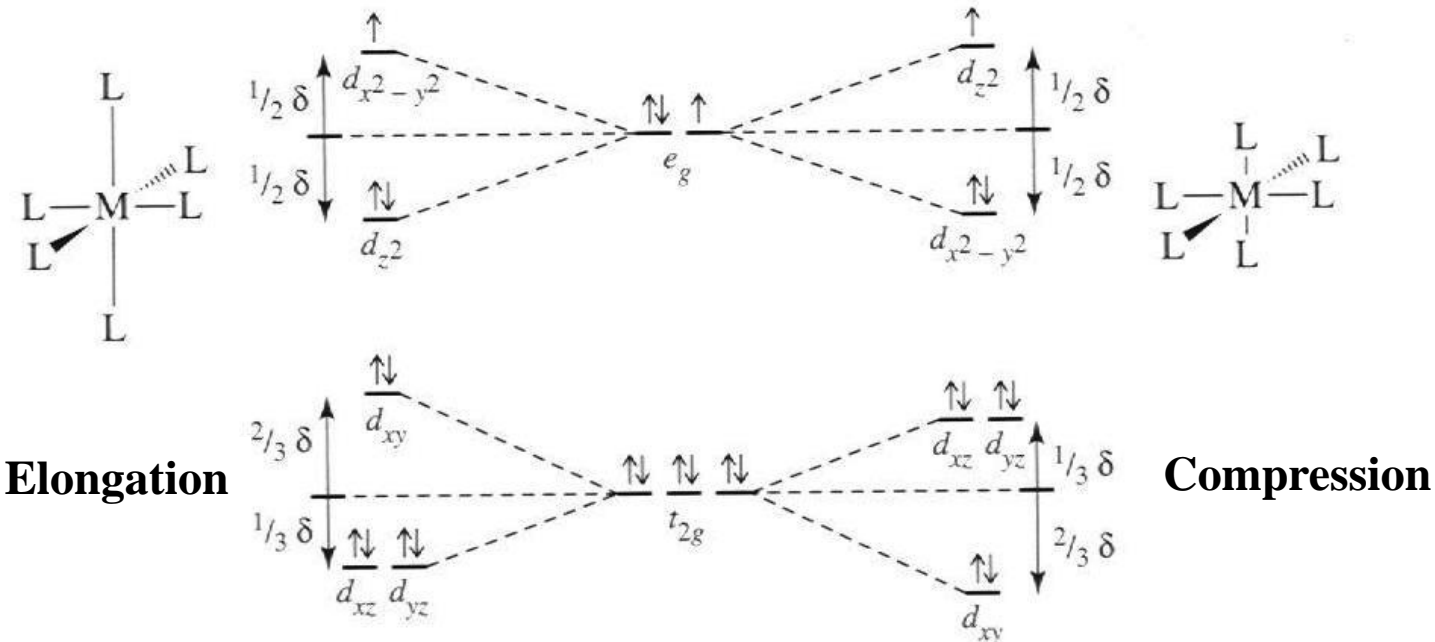
“Every non-linear molecule, which is in an electronically degenerate state, is prone to distortion lowering the symmetry and thus counteracting the electronic degeneracy“
⇒ Additional energy gain for d^4 (h.s.), d^7 (l.s.), and d^9 -configurations



7. Principles in Coordination Chemistry

Crystal Field Splitting in Tetragonally Distorted Octahedral Field

Tetragonally distorted octahedral crystal field as a consequence of the Jahn-Teller effect



Electronic-configuration	nd ¹	nd ²	nd ³	nd ⁴	nd ⁵	nd ⁶	nd ⁷	nd ⁸	nd ⁹	nd ¹⁰
High-spin J.T.	Weak	Weak	-	Strong	-	weak	Weak	-	Strong	-
Low-spin J.T.	Weak	Weak	-	Weak	Weak	-	Strong	-	Strong	-

7. Principles in Coordination Chemistry

Static vs. Dynamic Jahn-Teller-Effect in Octahedral Field

Static Jahn-Teller-Effect

Prerequisite: Electronic degeneracy in e_g^* -level

Detection: RSA, IR, UV/Vis

Example: $K_2Na[MnF_6]$ Mn^{III}: [Ar]3d⁴ h.s. elongated octahedron
 $[Cu(NH_3)_6]^{2+}$ Cu^{II}: [Ar]3d⁹ elongated octahedron

Dynamic Jahn-Teller-Effect

Prerequisite: Electronic degeneracy in t_{2g} -level

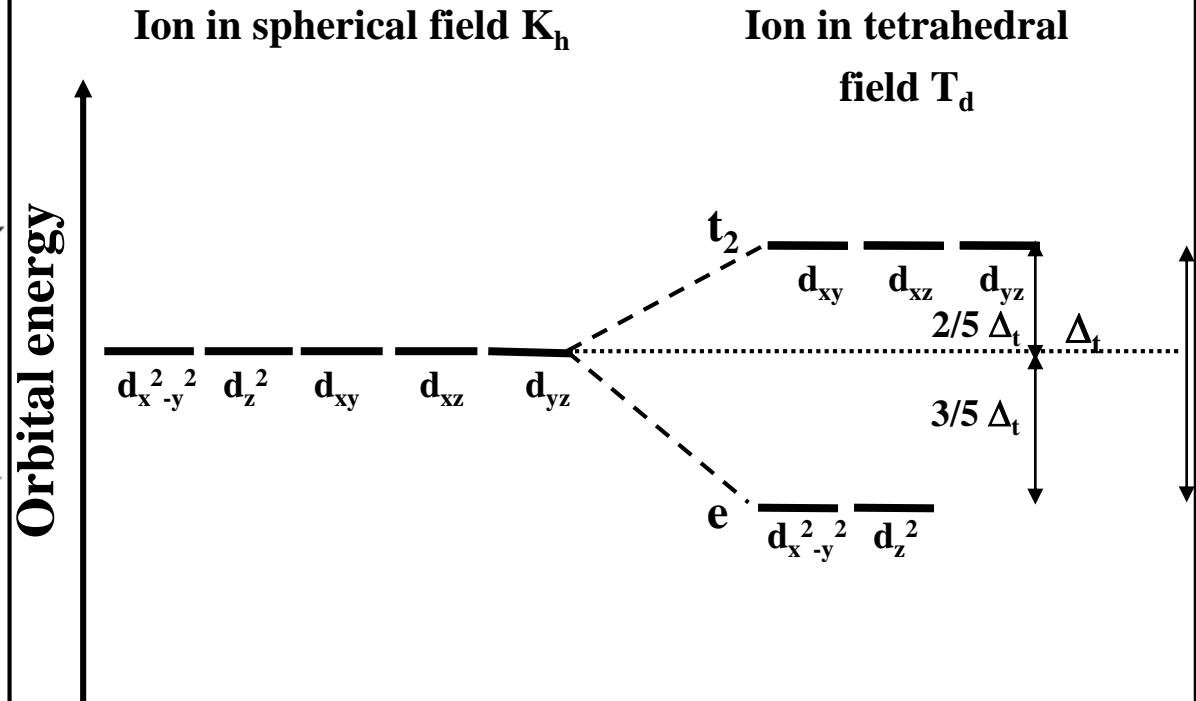
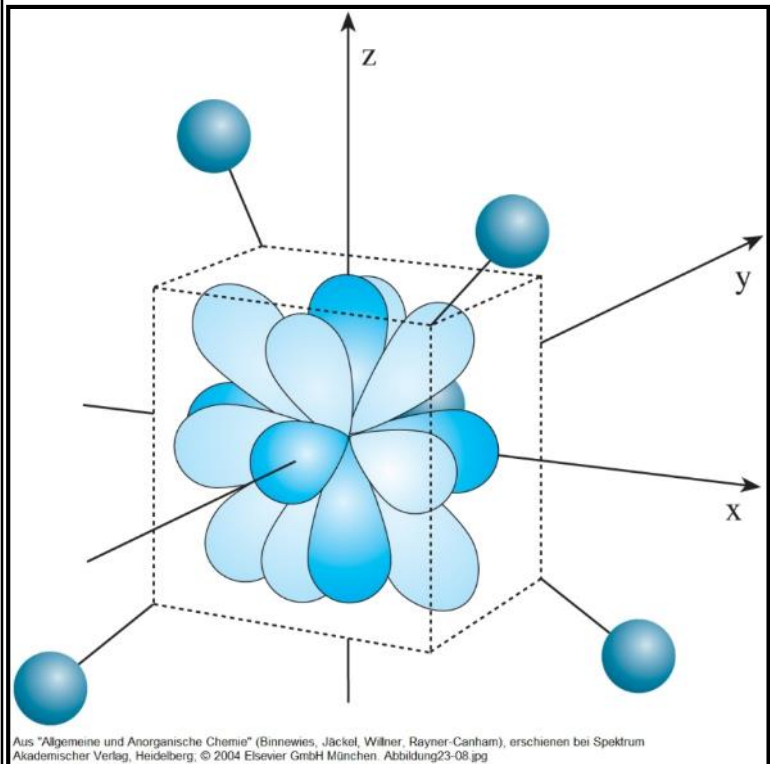
Detection: Difficult at room temperature, since δ is in the range of $k_B T$ ($\sim 200 \text{ cm}^{-1}$)

Example: $K_2Na[TiF_6]$ Ti^{III}: [Ar]3d¹ regular octahedron
crystallises at room temperature in cubic structure
⇒ Jahn-Teller polarons in $BaTiO_3$!

7. Principles in Coordination Chemistry

Crystal Field Splitting in Tetrahedral Field

Tetrahedral Crystal Field



The energy gap between the t_2 and the e -orbitals Δ_t is only $4/9$ of Δ_o , because only four instead of six ligands are present and those are not situated on the axes of the d-orbitals.

7. Principles in Coordination Chemistry

CFSE in Octahedral and Tetrahedral Field

Crystal field stabilisation energy in tetrahedral vs. octahedral crystal field

Calculation with $\Delta_t = 4/9\Delta_o$

“site preference”

d ⁿ	CFSE(tetrahedral)	CFSE(octahedral)	ΔCFSE(octahedr. – tetrahedr.)
1	-2.67 Dq _o	-4 Dq _o	-1.33 Dq _o
2	-5.33 Dq _o	-8 Dq _o	-2.67 Dq _o
3	-3.55 Dq _o	-12 Dq _o	-8.45 Dq _o
4	-1.78 Dq _o	-6 Dq _o (h.s.)	-4.22 Dq _o
		-16 Dq _o + 1 P (l.s.)	-14.22 Dq _o + 1 P
5	0 Dq _o	0 Dq _o (h.s.)	0 Dq _o
		-20 Dq _o + 2 P (l.s.)	-20 Dq _o + 2 P
6	-2.67 Dq _o	-4 Dq _o (h.s.)	-1.33 Dq _o
		-24 Dq _o + 2 P (l.s.)	-21.33 Dq _o + 2 P
7	-5.33 Dq _o	-8 Dq _o (h.s.)	-2.67 Dq _o
		-18 Dq _o + 1 P (l.s.)	-12.67 Dq _o + 1 P
8	-3.55 Dq _o	-12 Dq _o	-8.45 Dq _o
9	-1.78 Dq _o	-6 Dq _o	-4.22 Dq _o
10	0 Dq _o	0 Dq _o	0 Dq _o

7. Principles in Coordination Chemistry

CFSE in Tetrahedral Field

Some general rules

The magnitude of the CFSE in the tetrahedral field is only 4/9 of that in the octahedral field!

- Only high-spin complexes
- Ions with electronic configurations, leading to high CFSE, e.g. with [Ar]3d³-, [Ar]3d⁵(low-spin)- or [Ar]3d⁶(low-spin)-configuration, favour, if possible, octahedral coordination polyhedra
⇒ aqua complexes

Tetrahedral coordination polyhedra are observed for:

- Bulky ligands, i.e. proteins
- Ligands with double or triple bonds to the metal centre, e.g. oxy- and nitride ligands

Rule: There is no electron configuration where the electronic stabilisation is higher for the tetrahedral than for the octahedral coordination (site preference) ⇒ Octahedral geometry is preferred

Exception: d⁵ (high-spin) and d¹⁰, because CFSE in both octahedron and tetrahedron are zero

7. Principals in Coordination Chemistry

Crystal Field Splitting in Different Symmetries

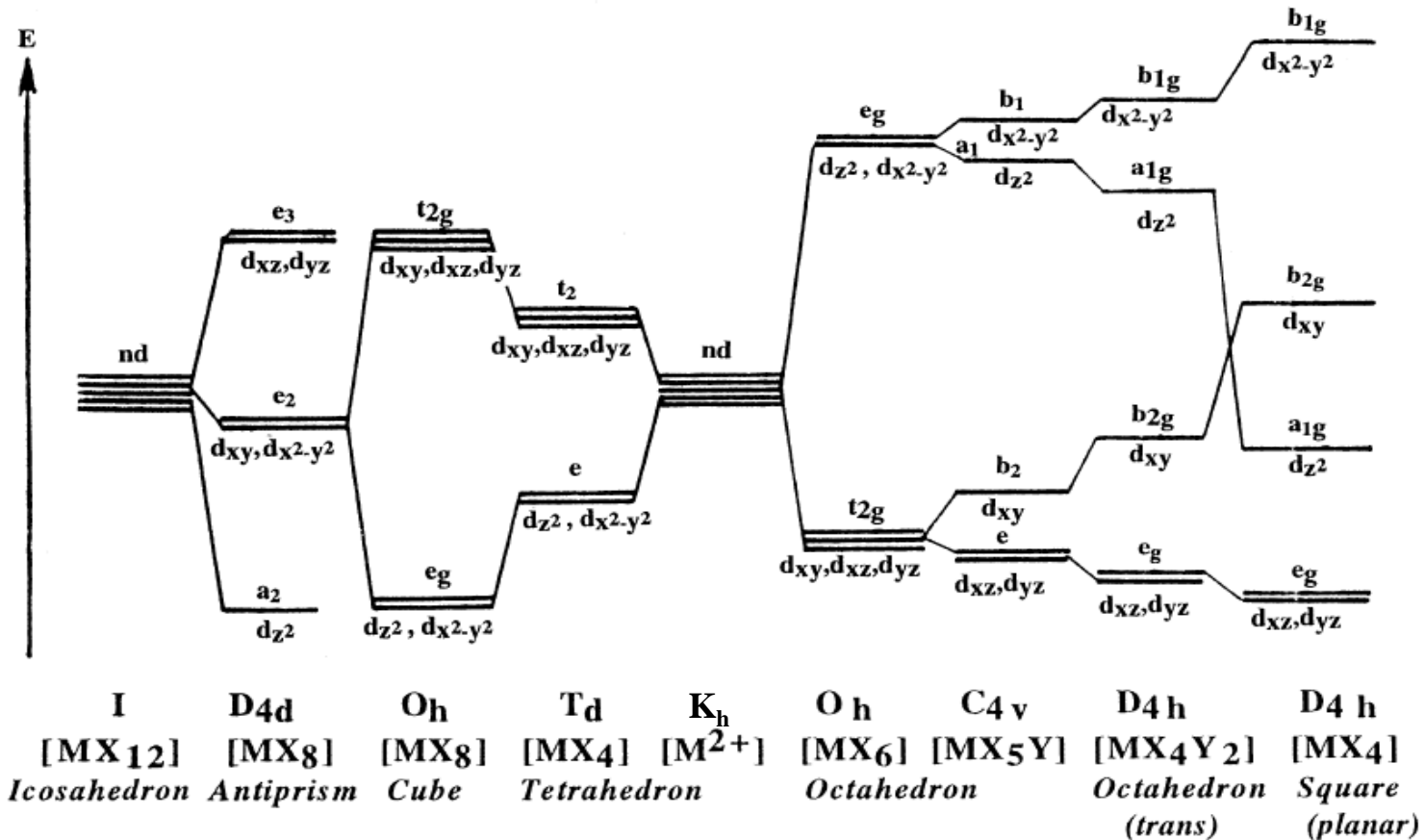
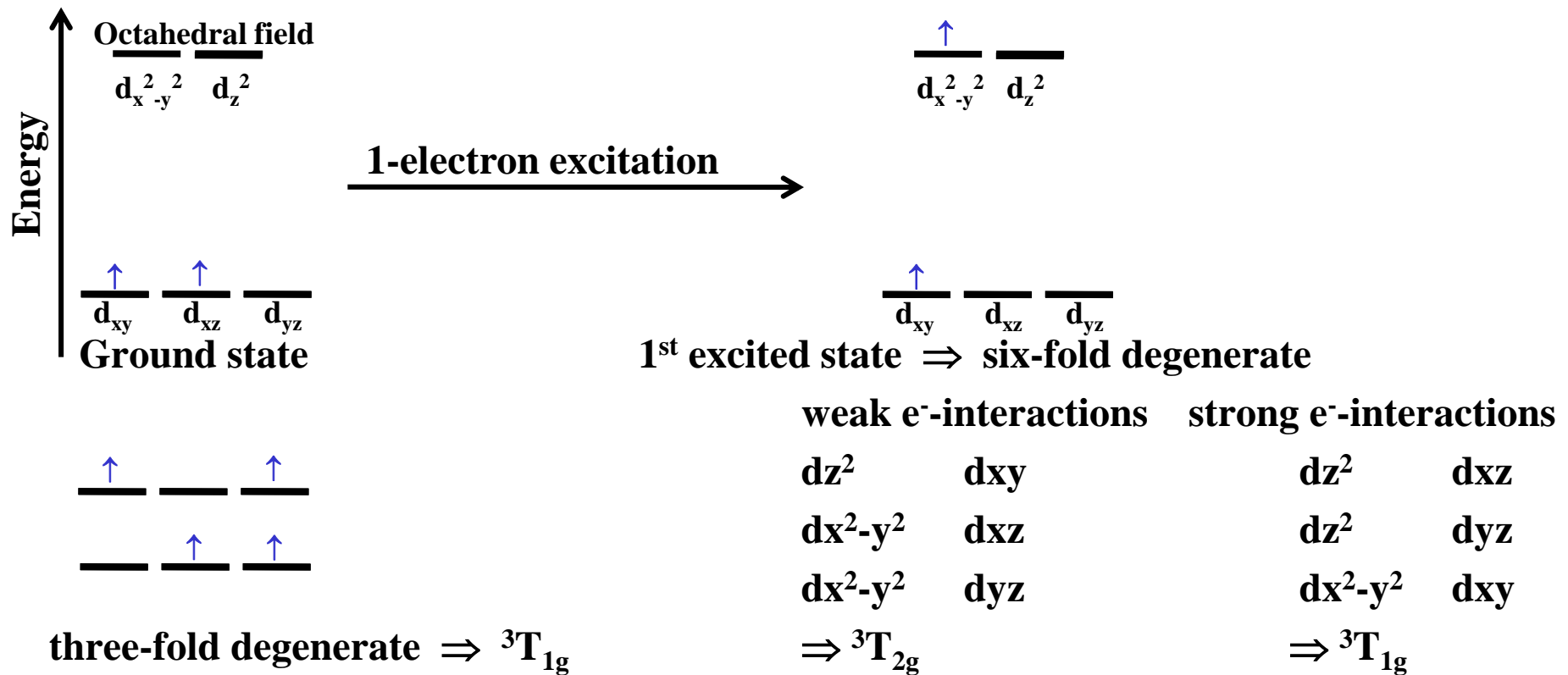


FIGURE 4.5. Splitting of d orbital energy levels in ligand fields of different symmetries. In MX_5Y and MX_4Y_2 complexes the splitting of the T_{2g} and E_g terms can be inverted depending on the ratio of field strengths X/Y . (After Schmidtke [4.12].)

7. Principles in Coordination Chemistry

Description of Electronic States in Multiple-Electron-Atoms

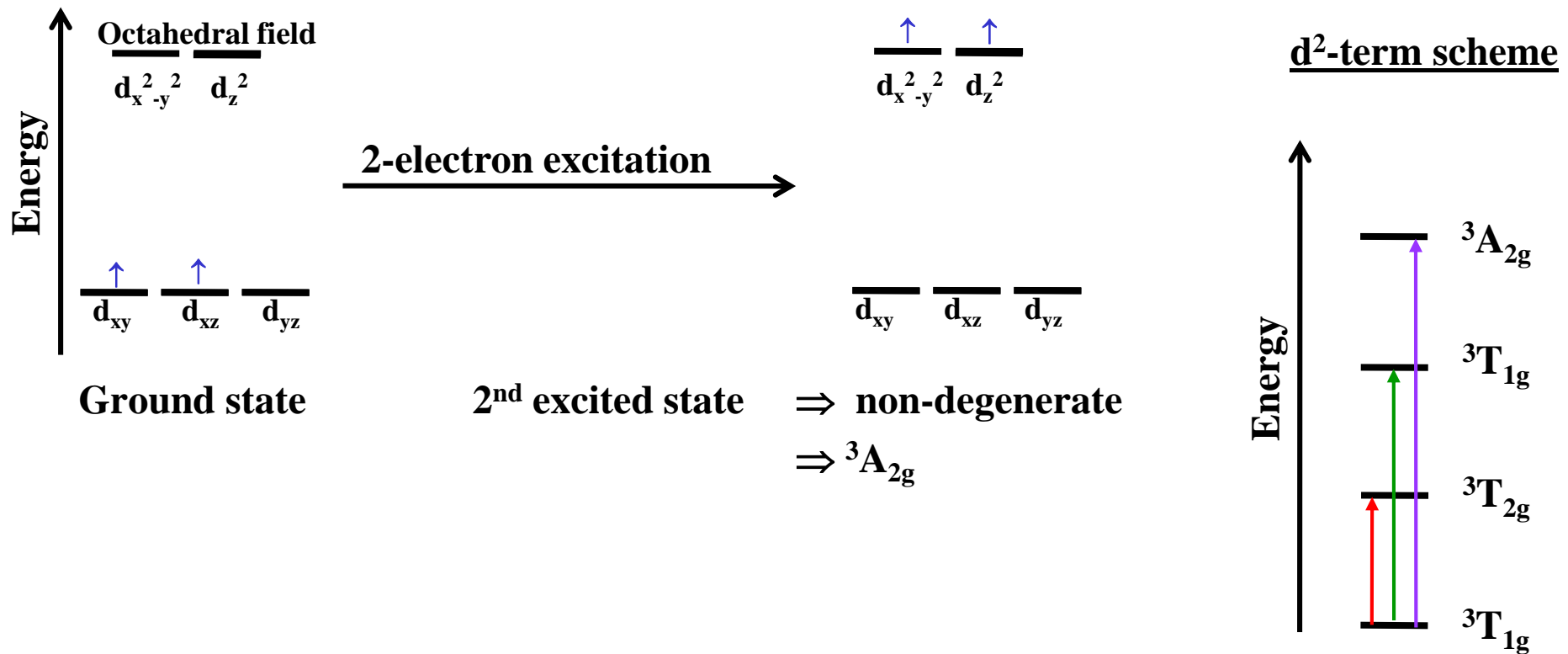
Method of the strong field: LS-coupling is considerably smaller than crystal field splitting \Rightarrow true for elements of the 3d-series up to bromine



7. Principles in Coordination Chemistry

Description of Electronic States in Multiple-Electron-Atoms

Method of the strong field: LS-coupling is considerably smaller than crystal field splitting \Rightarrow true for elements of the 3d-series up to bromine

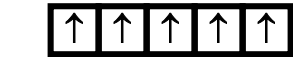


7. Principles in Coordination Chemistry

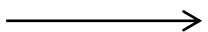
Description of Electronic States in Multiple-Electron-Atoms

Method of the weak field: LS-coupling is notably stronger than the crystal field splitting \Rightarrow true from bromine on, thus for the elements of the 4d- and 5d-series as well as the lanthanides (\rightarrow Dieke-diagram) and the actinides

Electron configuration



ml -2 -1 0 1 2



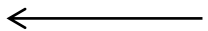
orbital and spin momentum L and S

$$L = |\sum l_i| \text{ and } S = \sum s_i$$

Coulomb-interactions

Spin-orbit coupling

Crystal field splitting



Crystal field energy terms

$\Rightarrow A, B, E, T$

Total angular momentum J

$$J = |L - S| \dots |L + S|$$

7. Principles in Coordination Chemistry

Description of Electronic States in Multiple-Electron-Atoms

Orbital angular momentum $L = |\sum l_i|$

$ M_L $	0	1	2	3	4	5	6	7	8	9	...
	S	P	D	F	G	H	I	K	L	M	...

Spin angular momentum $S = \sum s_i$

$ M_S $	0	1I_2	1	3I_2	2	5I_2	...
$2S+1$	1	2	3	4	5	6	...
	Singlet	Doublet	Triplet	Quartet	Quintet	Sextet	...

$$2S+1L$$

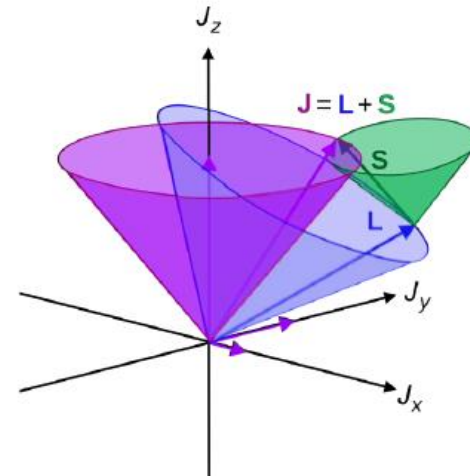
Total angular momentum $J = |L - S| \dots |L + S|$

results from different orientations

of L and S towards each other

\Rightarrow Russell-Saunders (RS) coupling

$$2S+1L_J$$



http://upload.wikimedia.org/wikipedia/commons/thumb/3/30/LS_coupling.svg/2000px-LS_coupling.svg.png

7. Principles in Coordination Chemistry

Quantum Mechanical Micro States

$$\text{Number \#} = \frac{n!}{e!h!}$$

with

n = maximal number of electrons in sub-shell
(sum of e + h)

e = number of electrons of corresponding configuration

h = number of holes of corresponding configuration

p-shell \Rightarrow n = 6 e

	1	2	3	4	5	6
#	6	15	20	15	6	1

Discussion of three electron configurations

1. Elemental carbon [He]2s²2p²
2. 3d-transition metal ions [Ar]3dⁿ
3. Lanthanide ions Ln³⁺ [Xe]4fⁿ

7. Principles in Coordination Chemistry

Examples for Micro States: p²-Configuration, i.e. 2 Electrons in p-Orbitals

L	+2	0	-2	+1	0	-1	+1	0	-1
+1	↑↓			↑	↑	↓		↓	
0		↑↓		↑		↑	↑		↓
-1			↑↓		↑	↑		↓	↓
S	0	0	0	+1	+1	+1	-1	-1	-1

L	+1	0	-1	+1	0	-1
+1	↑	↑		↓	↓	
0	↓		↑	↑		↓
-1		↓	↓		↑	↑
S	0	0	0	0	0	0

⇒ 15 micro states

7. Principles in Coordination Chemistry

RS-Terms for the $2p^2$ -Configuration

$$L = 2 \quad M_L = -2, -1, 0, 1, 2$$

$$S = 1 \quad M_S = -1, 0, 1$$

$$M_L = 2 \text{ and } M_S = 0 \Rightarrow ^1D$$

L =	+2	+1	0	-1	-2
S = 0	X	X	X	X	X

$$M_L = 1 \text{ and } M_S = 1 \Rightarrow ^3P$$

L =	+1	0	-1
S = +1	X	X	X
S = 0	X	X	X
S = -1	X	X	X

$$M_L = 0 \text{ and } M_S = 0 \Rightarrow ^1S$$

L =	0
S = 0	X

Hund's rules (3 different ones)

\Rightarrow Energetic order for $2p^2$ config.:

1st S as high as possible
2nd L as high as possible
 $^3P < ^1D < ^1S$

3rd Hund's rule for J:
 $J = |L - S|$ most stable for shells filled less than half

7. Principles in Coordination Chemistry

RS-Terms for the d^n -Configurations

$$2S+1L_J$$

d^n	-2	-1	0	1	2	L	S	Ground term h.s. (l. s.)
d^1		↑				2	1/2	2D
d^2	↑	↑				3	1	3F
d^3	↑	↑	↑			3	3/2	4F
d^4	↑	↑	↑	↑		2	2	5D (3H)
d^5	↑	↑	↑	↑	↑	0	5/2	6S (2I)
d^6	↑↓	↑	↑	↑	↑	2	2	5D (1I)
d^7	↑↓	↑↓	↑	↑	↑	3	3/2	4F (2H)
d^8	↑↓	↑↓	↑↓	↑	↑	3	1	3F
d^9	↑↓	↑↓	↑↓	↑↓	↑	2	1/2	2D

7. Principals in Coordination Chemistry

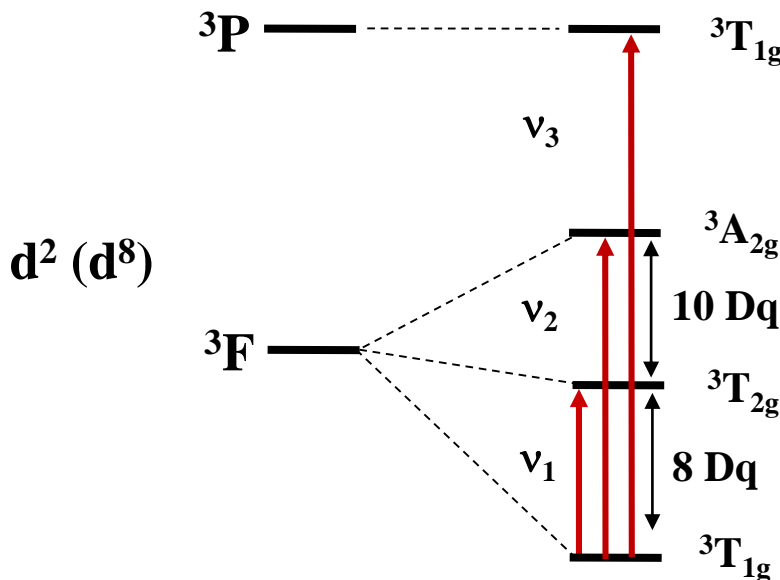
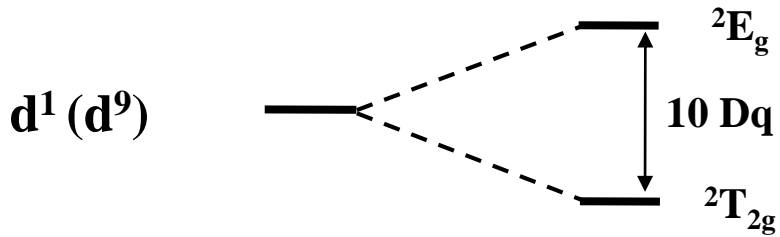
RS-Terms for all dⁿ-Configurations

All Russell Saunders terms for 3dⁿ free ion configurations

Configuration	# Micro-states	# Energy levels	Ground state terms	Excited energy terms
d ¹ , d ⁹	10	1	² D	-
d ² , d ⁸	45	5	³ F	³ P, ¹ G, ¹ D, ¹ S
d ³ , d ⁷	120	8	⁴ F	⁴ P, ² H, ² G, ² F, 2x ² D, ² P
d ⁴ , d ⁶	210	16	⁵ D	³ H, ³ G, 2x ³ F, ³ D, 2x ³ P, ¹ I, 2x ¹ G, ¹ F, 2x ¹ D, 2x ¹ S
d ⁵	252	16	⁶ S	⁴ G, ⁴ F, ⁴ D, ⁴ P, ² I, ² H, 2x ² G, 2x ² F, 3x ² D, ² P, ² S
d ¹⁰	1	1	¹ S	-

7. Principles in Coordination Chemistry

Splitting of RS-Terms in Crystal Field \Rightarrow Splitting Terms

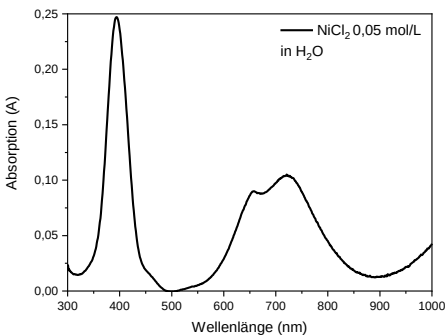
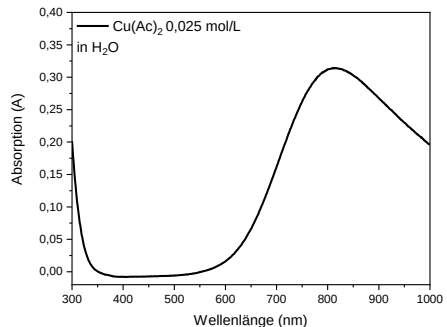


1 band

$$\Delta E = 10 \text{ Dq}$$

Inverse splitting in tetrahedral field

V⁴⁺



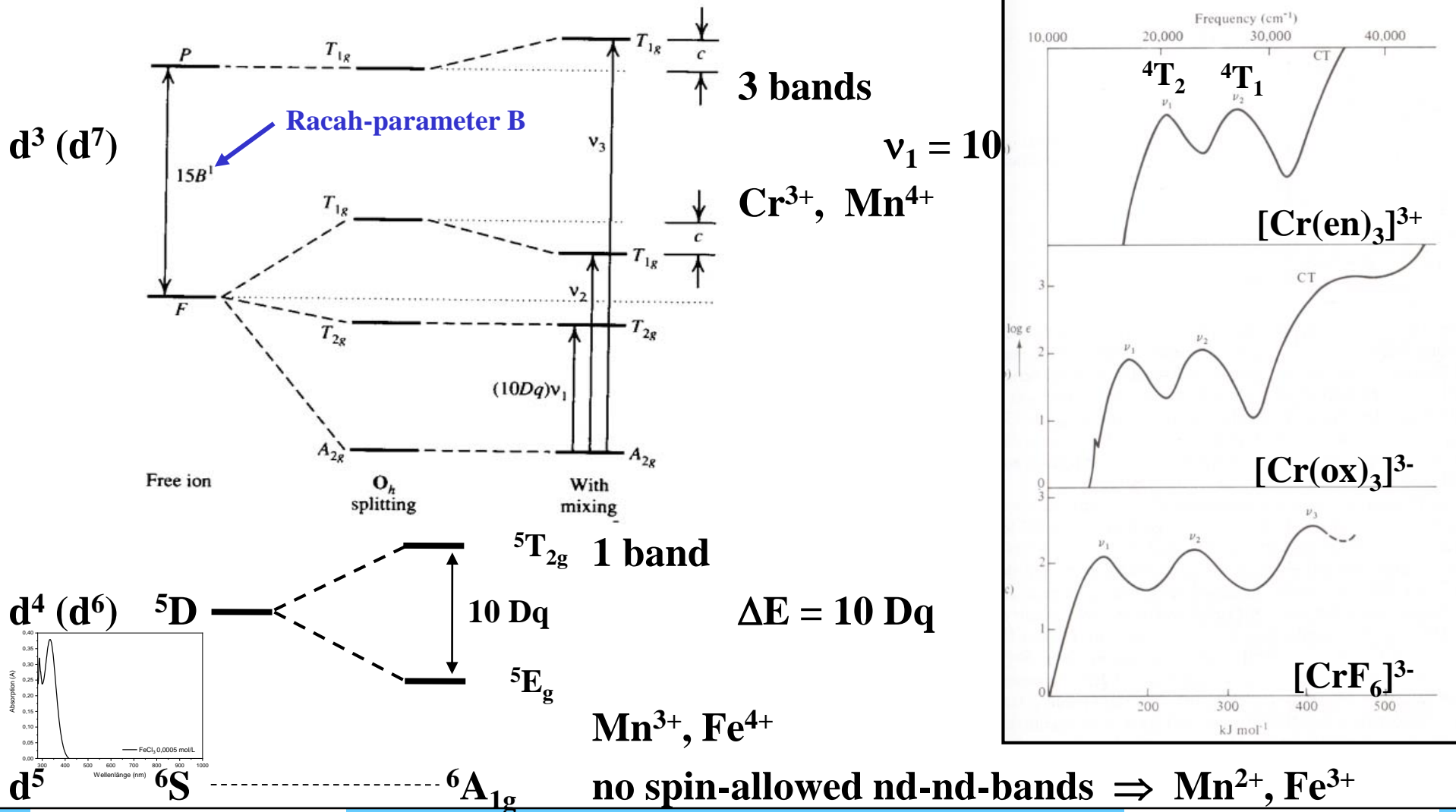
3 bands

\Rightarrow determination of 10 Dq possible only by subtraction $= v_2 - v_1$

V³⁺, Mn⁵⁺

7. Principles in Coordination Chemistry

Splitting of RS-Terms in Crystal Field \Rightarrow Splitting Terms



7. Principals in Coordination Chemistry

Splitting of RS-Terms in Crystal Field \Rightarrow Calculation of 10 Dq

Octahedron d^1, d^4, d^6 and d^9
1 band $10 Dq = \nu$

Octahedron d^2
3 bands $10 Dq = \nu_2 - \nu_1$ and B by calculation

Octahedron d^7
3 bands $10 Dq = \nu_2 - \nu_1$ and B by calculation

Octahedron d^3 and d^8
3 bands $10 Dq = \nu_1$ and B by calculation

$$\frac{B}{Dq} = \frac{\left(\frac{\Delta E}{Dq}\right)^2 - 10 \frac{\Delta E}{Dq}}{15\left(\frac{\Delta E}{Dq} - 8\right)}$$

$$Dq = E(^4T_2) / 10$$
$$\Delta E = E(^4T_1) - E(^4T_2)$$

7. Principals in Coordination Chemistry

Interelectronic Repulsion: Racah-Parameter A, B, and C (Giulio Racah 1909 - 1965)

Means to describe the Interelectronic repulsion or Coulomb-repulsion between the terms, with B being the most important Racah-parameter, because it directly describes the splitting between the RS-terms.

Free M^{n+} -ion

$$B \sim 500 - 1100 \text{ cm}^{-1}$$

$$C \sim 4 B \text{ (approximation!)}$$

Complexed M^{n+} -ion

B is ca. 30% smaller due to the nephelauxetic effect,
i.e. the delocalisation of metal-centred electrons to
the ligands $\Rightarrow B'$

Nephelauxetic ratio

$$\beta = B'/B \quad \text{with } (1-\beta) = h_L * k_M$$

h_L = nephelauxetic parameter of ligands

k_M = nephelauxetic parameter of the metals

$$A = F_0 - 49 F_4$$

$$B = F_2 - 5 F_4$$

$$C = 35 F_4$$

(with $F_{0,2,4}$ = Slater-integrals)

7. Principles in Coordination Chemistry

Nephelauxetic Effect ~ Electron Density Between Metal Ions and Ligands

- Quantification of effect by parameter $\beta = B'/B$
- Ionic charge density and polarizability of the ligands by parameter h_L

Ligand	h_L
F^-	0.8
H_2O	1.0
DMF	1.2
$(NH_2)_2CO$	1.2
NH_3	1.4
en	1.5
$C_2O_4^{2-}$	1.5
Cl^-	2.0
CN^-	2.1
Br^-	2.3
N_3^-	2.4
I^-	2.7

7. Principles in Coordination Chemistry

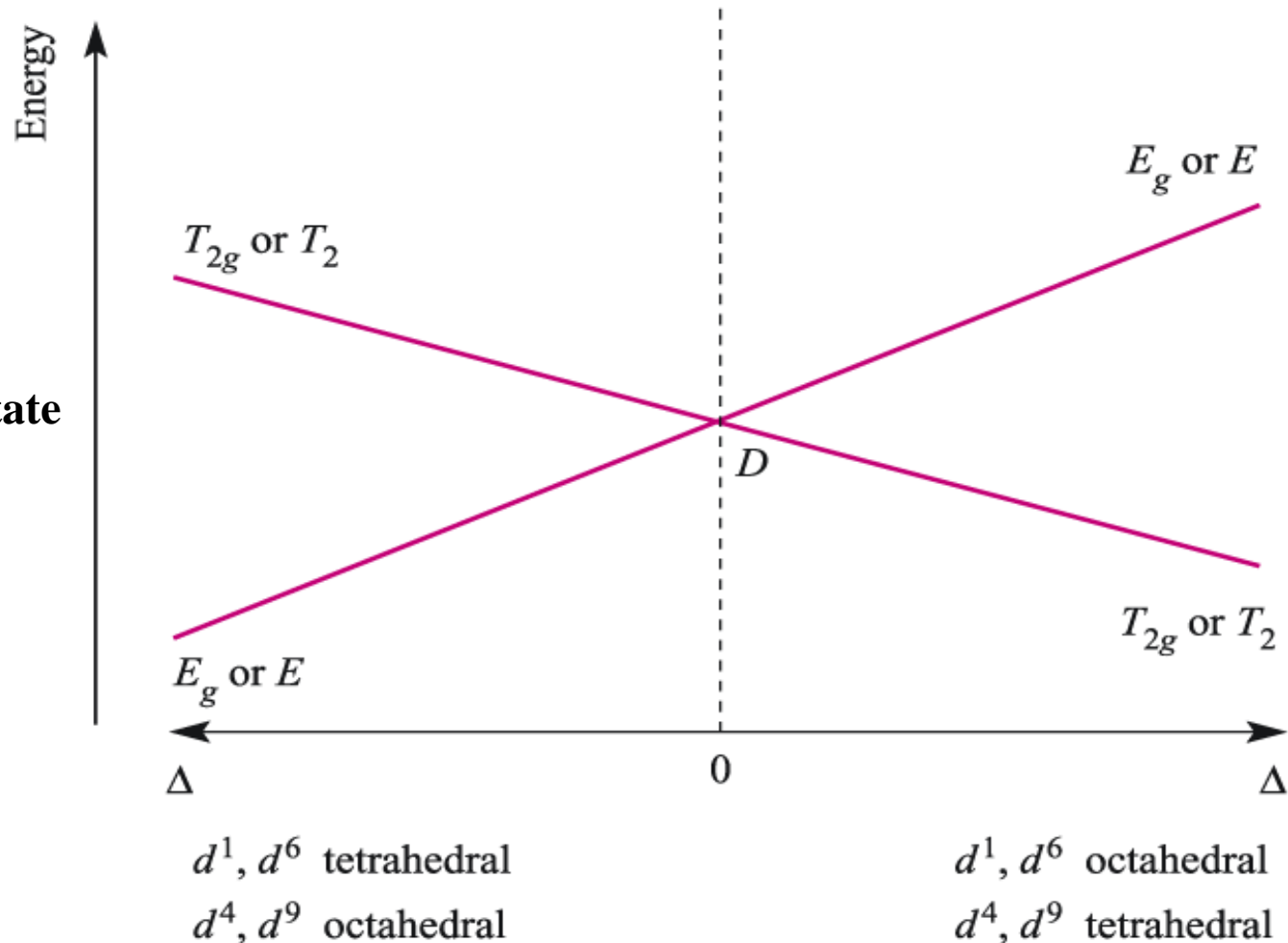
Energies of Splitting Terms as Function of the Field Strength \Rightarrow Orgel-Diagram

Low-spin configurations
are not taken into account
(weak crystal field!)

Only terms with same spin
multiplicity as the ground state
are considered

“1 electron configuration”
 d^1 and d^6

“1 hole configuration”
 d^4 and d^9



7. Principles in Coordination Chemistry

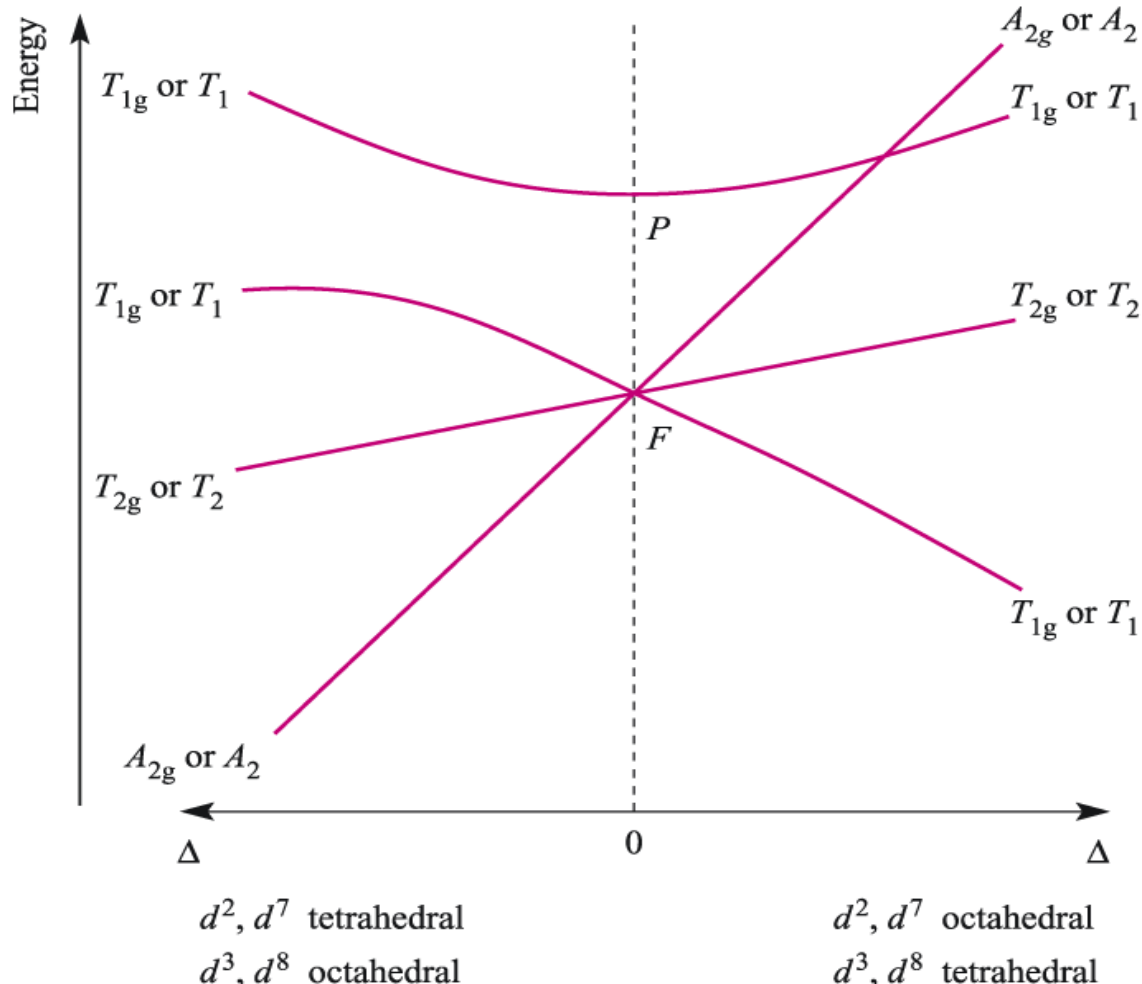
Energies of Splitting Terms as Function of the Field Strength \Rightarrow Orgel-Diagram

Low-spin configurations
are not taken into account
(weak crystal field!)

Only terms with same spin
multiplicity as the ground state
are considered

“2 electron configuration”
 d^2 and d^7

“2 hole configuration”
 d^3 and d^8



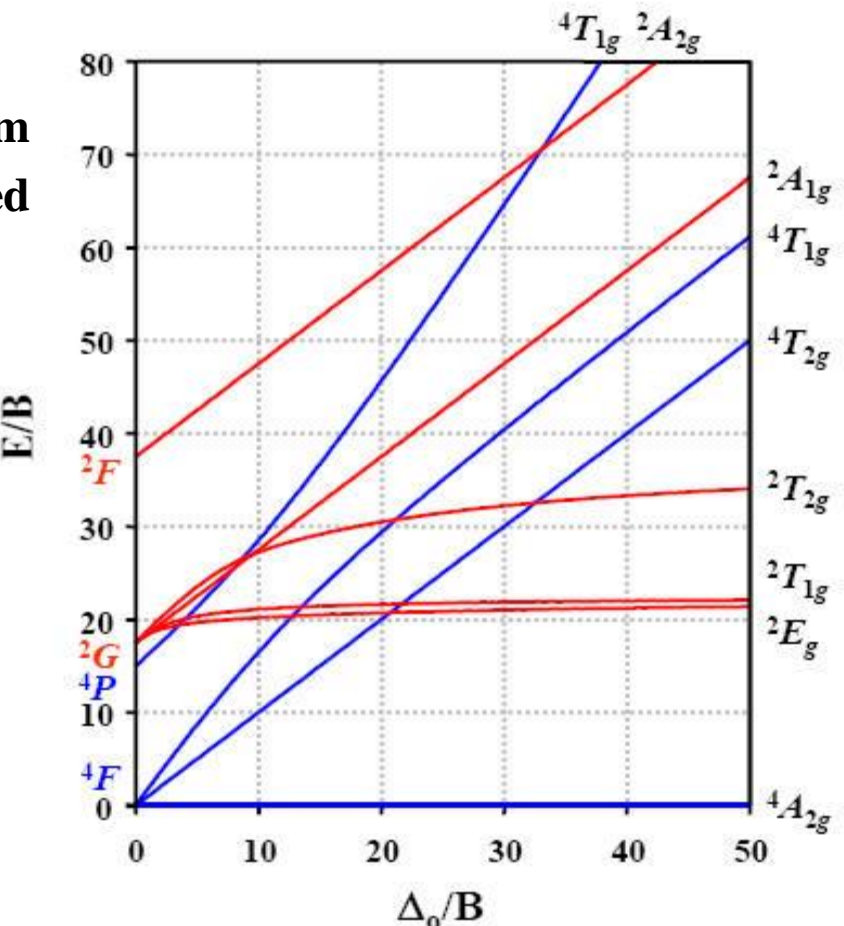
7. Principles in Coordination Chemistry

Tanabe-Sugano-Diagram (Term Correlations)

- Also low-spin terms and other spin multiplicities are taken into account
- The abscissa equals the energy of the ground term
- The energy of the crystal field terms is normalised by B
- The magnitude of B depends on the ion

Configuration	Ion	B [cm ⁻¹]	C [cm ⁻¹]
3d ³	Cr ³⁺	918	3850
	Mn ⁴⁺	1064	
3d ⁴	Cr ²⁺	830	3430
	Mn ³⁺	1140	3675
3d ⁵	Mn ²⁺	960	3325
3d ⁶	Fe ²⁺	1058	3901
	Co ³⁺	1100	

d^3 Tanabe-Sugano Diagram



7. Principles in Coordination Chemistry

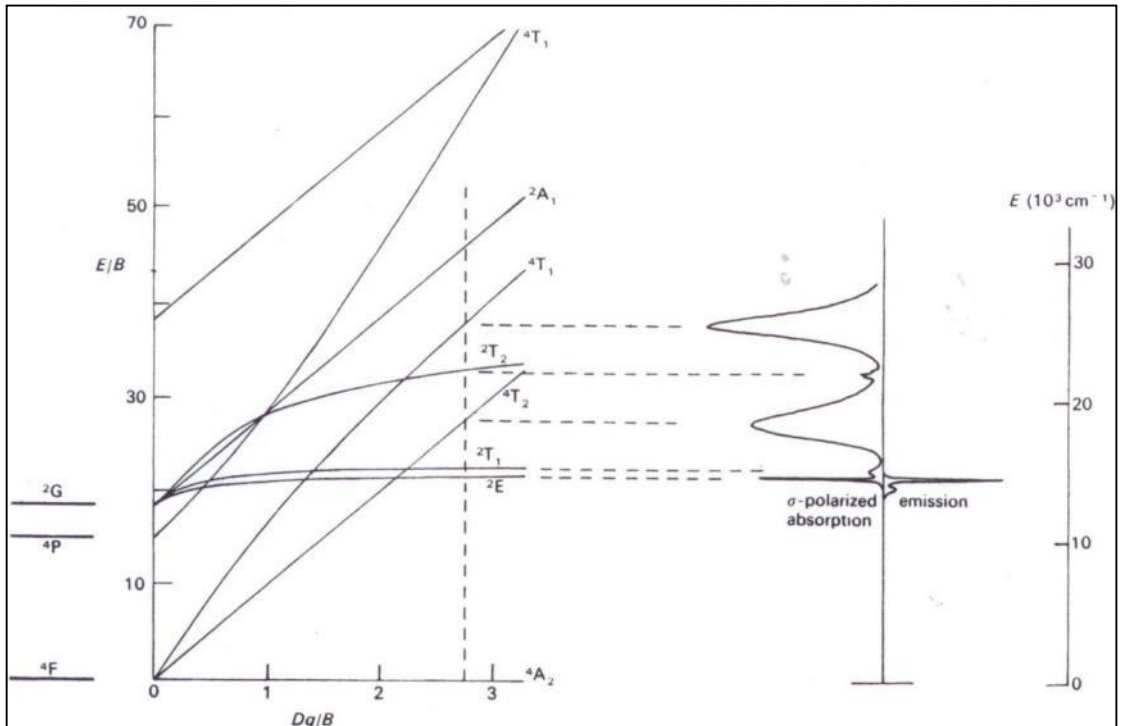
Optical Spectra of 3d-Ions

Energetic positions of terms

- Coulomb-interaction $\sim 10000 \text{ cm}^{-1}$
- Spin-orbit-coupling $\sim 100 \text{ cm}^{-1}$
- Crystal field splitting $\sim 1000 \text{ cm}^{-1}$

Shape of optical transitions

- Parallel terms:
Sharp lines
- Terms with different slopes:
Broad bands



Selection rules

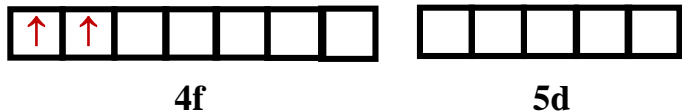
- All $d^n \rightarrow d^n$ transitions are parity-forbidden ($g \leftrightarrow g$)
- Transitions between different spins are also spin-forbidden
- Specific symmetric selection rules according to group theory (Ref.: F.A. Cotton, "Chemical Applications of Group Theory")

7. Principles in Coordination Chemistry

Ln^{3+} -Ions

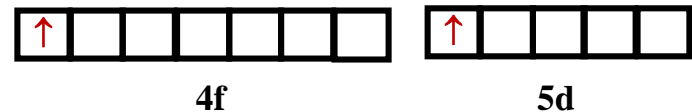
Pr^{3+} ground state

$[\text{Xe}]4\text{f}^2 \rightarrow 13 \text{ SLJ-levels (91 microstates)}$



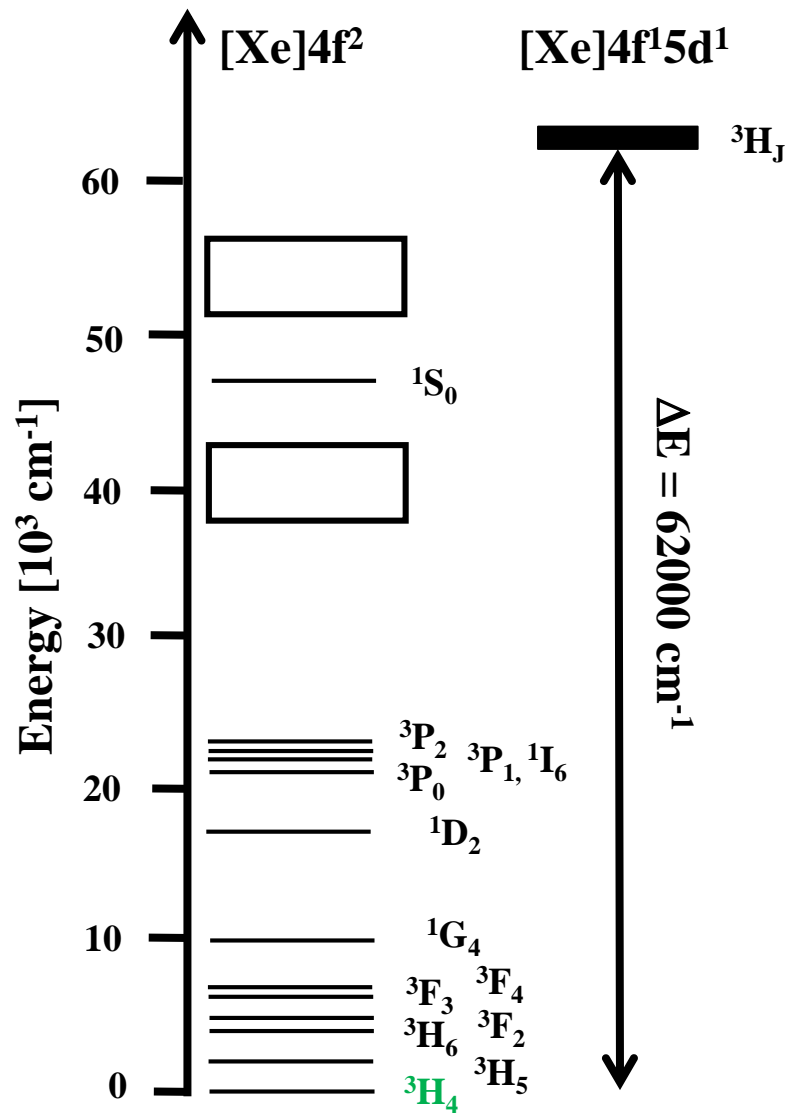
Pr^{3+} 1st excited state

$[\text{Xe}]4\text{f}^15\text{d}^1 \rightarrow 2 \text{ SLJ-levels}$



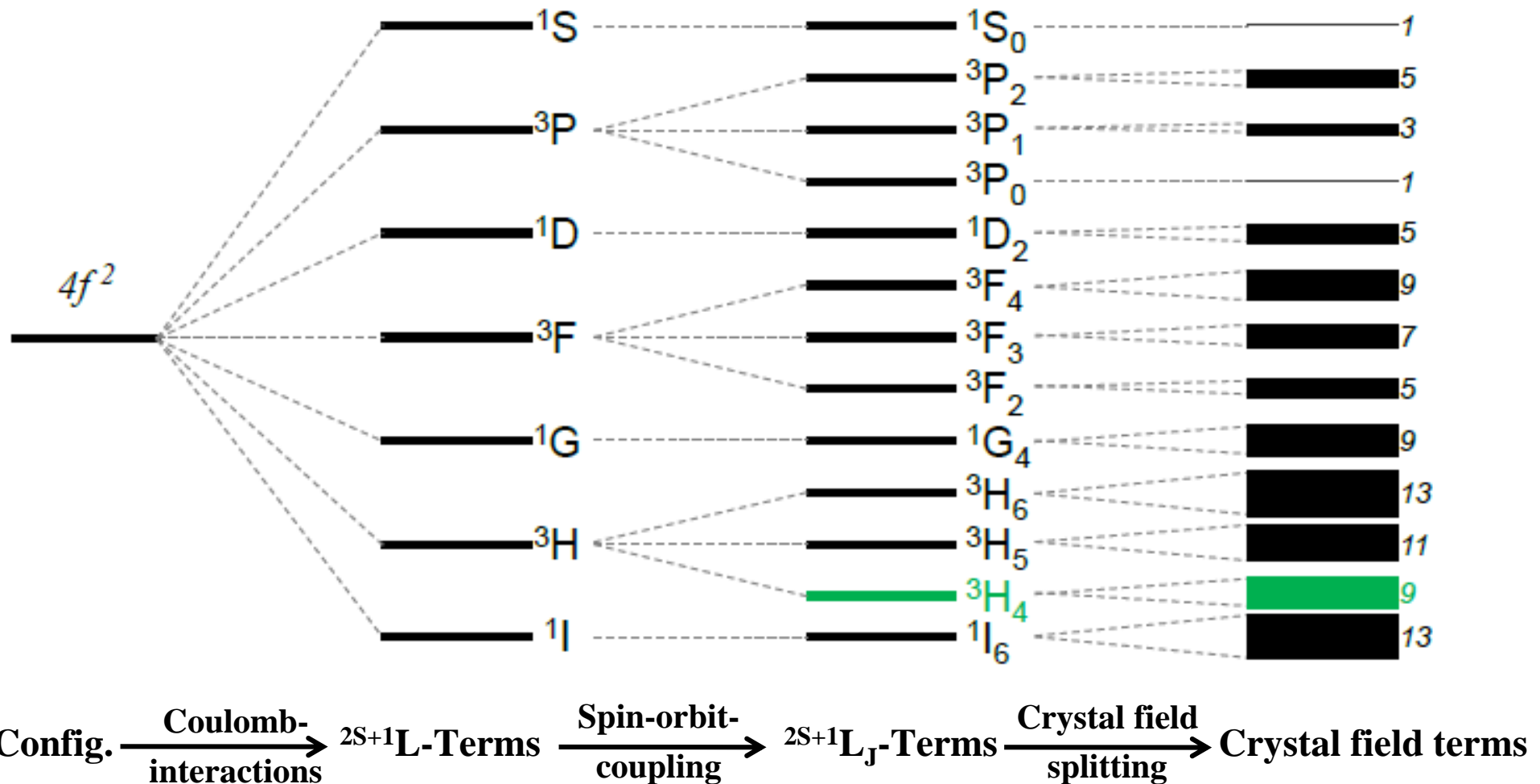
$\Rightarrow [\text{Xe}]4\text{f}^2 - [\text{Xe}]4\text{f}^2$ transitions

$\Rightarrow [\text{Xe}]4\text{f}^2 - [\text{Xe}]4\text{f}^15\text{d}^1$ transitions



7. Principles in Coordination Chemistry

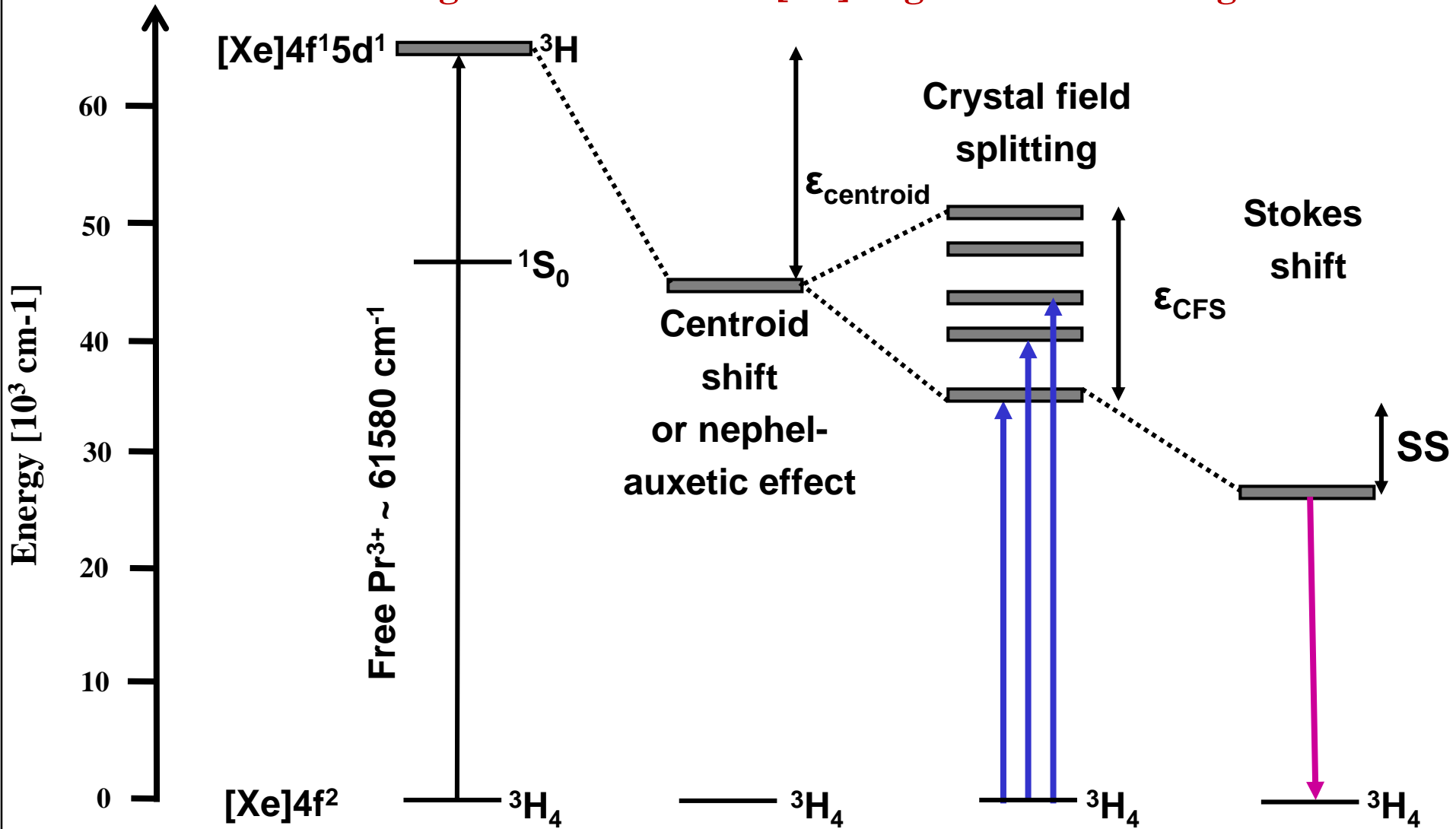
Pr³⁺-Ion with 4f²-Configuration → 91 Microstates



Energetic ordering is subject to the three Hund rules

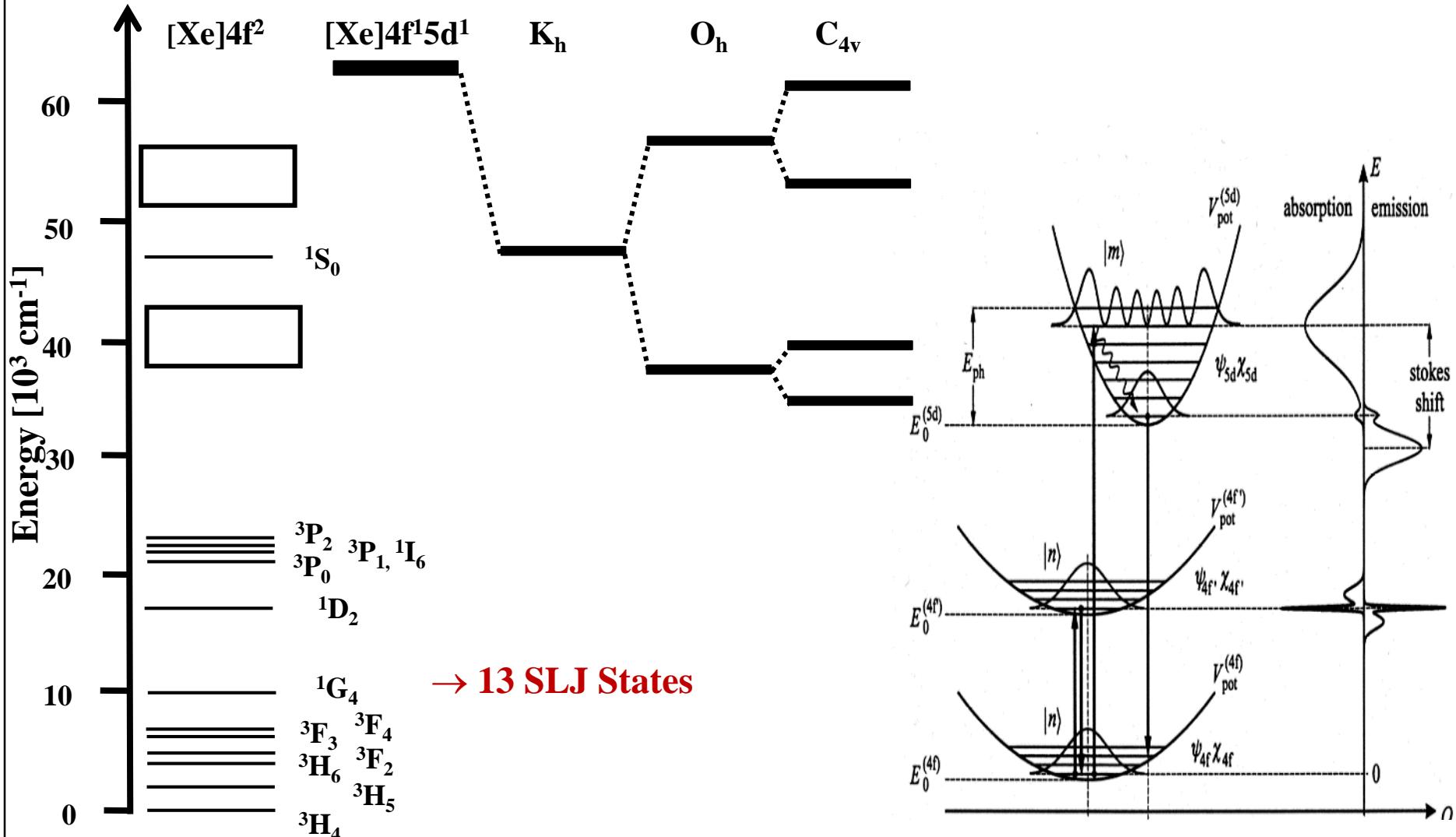
7. Principles in Coordination Chemistry

Ln³⁺-Ions with 4fⁿ-Configuration: Pr³⁺ with [Xe]4f² ground state configuration



7. Principles in Coordination Chemistry

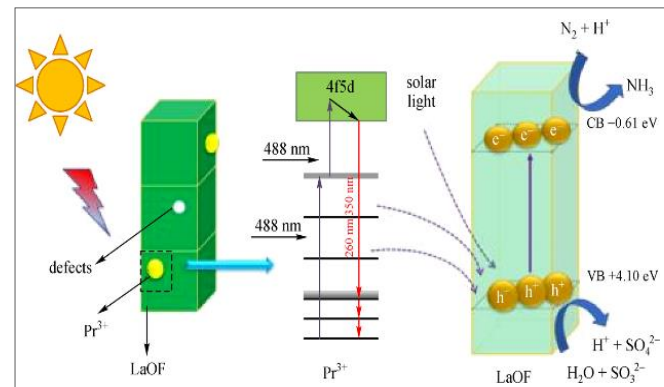
Ln³⁺-Ions with 4fⁿ-Configuration: Pr³⁺ with [Xe]4f² ground state configuration



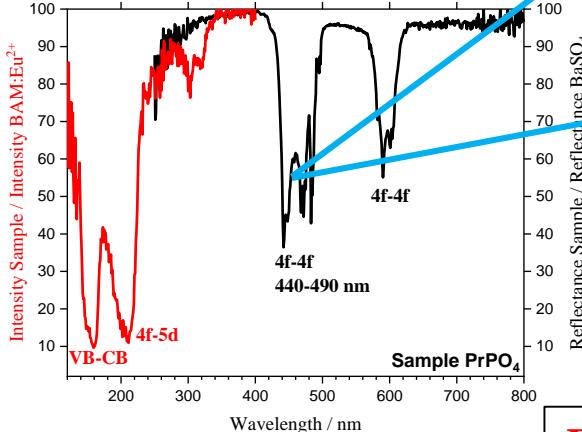
7. Principles in Coordination Chemistry

Ln^{3+} -Ions with $4f^n$ -Configuration: Pr^{3+} with $[\text{Xe}]4f^2$ ground state configuration

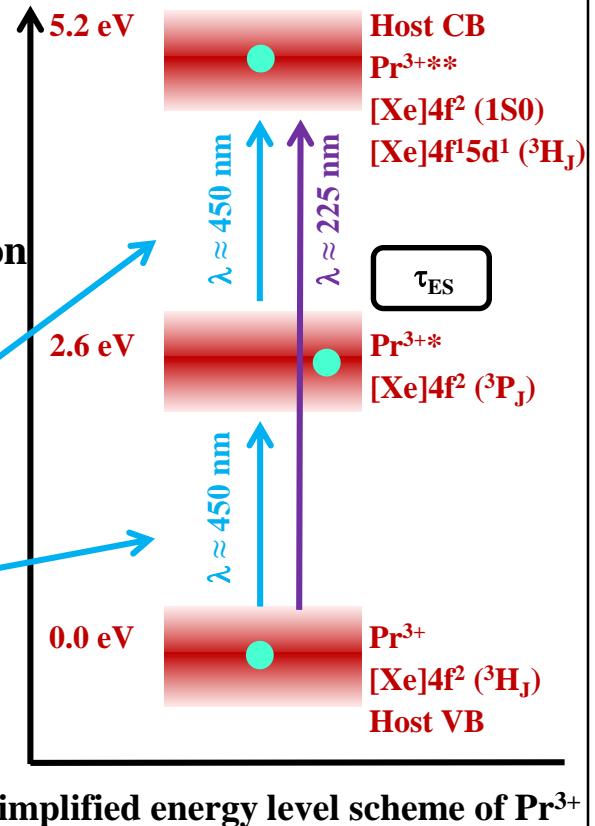
- Haber-Bosch process (α -Fe) $\text{N}_2 + 3 \text{H}_2 \rightleftharpoons 2 \text{NH}_3$ (400 - 500 °C)
- Mo/Fe Nitrogenase (Fe^{n+}) $2 \text{N}_2 + 10 \text{H}^+ + 8 \text{e}^- \rightleftharpoons 2 \text{NH}_4^+ + \text{H}_2$ (RT)
- Heterogeneous photocatalysis by up-conversion induced photoionisation
Semiconductor (SC) + blue laser \rightarrow SC* \rightarrow SC** [$\text{e}^-(\text{CB}) + \text{h}^+(\text{VB})$]
 $\text{e}^-(\text{CB}) + \text{N}_2 \rightarrow (\text{N}_2)^-$
 $6(\text{N}_2)^- + 6 \text{H}_2\text{O} \rightarrow 6 \text{OH}^- + 5 \text{N}_2 + 2 \text{NH}_3$



Photocatalytical NH_3 formation at LaOF:Pr



Reflection spectrum of PrPO_4



Simplified energy level scheme of Pr^{3+}

Bioinorganic chemistry driven idea:
Can Pr^{3+} serve for the N_2 fixation?

Lit.: LaOF-Pr MW hydrothermal synthesis for photocatalytic N fixation, Front Mater Science 14 (2020) 43

7. Principles in Coordination Chemistry

Crystal Field and Ligand Field

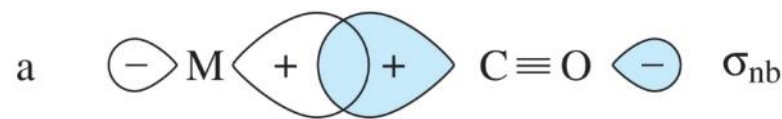
Back-binding to ligands

- Typical for CO, NO, O₂, CN⁻, some biochemically important molecules!
- Formally, metals have low oxidation state or high electron density, respectively which is spread over the ligands through back-transfer of charge

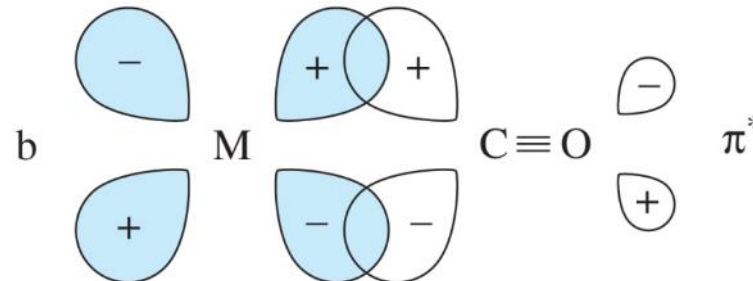
Example: M(CO)₆ “metal carbonyl”



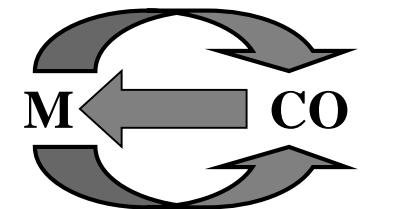
a) σ -donor-bond



b) π -acceptor-bond



Aus "Allgemeine und Anorganische Chemie" (Binnewies, Jackel, Willner, Rayner-Canham), erschienen bei Spektrum Akademischer Verlag, Heidelberg, © 2004 Elsevier GmbH München. Abbildung 23-29 jpg



“synergetic effect”

7. Principles in Coordination Chemistry

From Crystal to Ligand Field

Explanation of ligand ordering in the spectrochemical series



Strong ligands

π -back binding

Weak ligands

no π -back binding

The spectrochemical series does not correlate with the charge of the ligands but with the ability of the ligands to delocalise electron density from the metal atom and thus to enhance the positive charge density or the effective field strength at the metal atom.

**π -acceptor-ligands:
(back-binding)**

Stabilise metals in low oxidations states \Rightarrow CO, NO, CN⁻, CN₂²⁻, O₂, N₂

π -donor-ligands: St

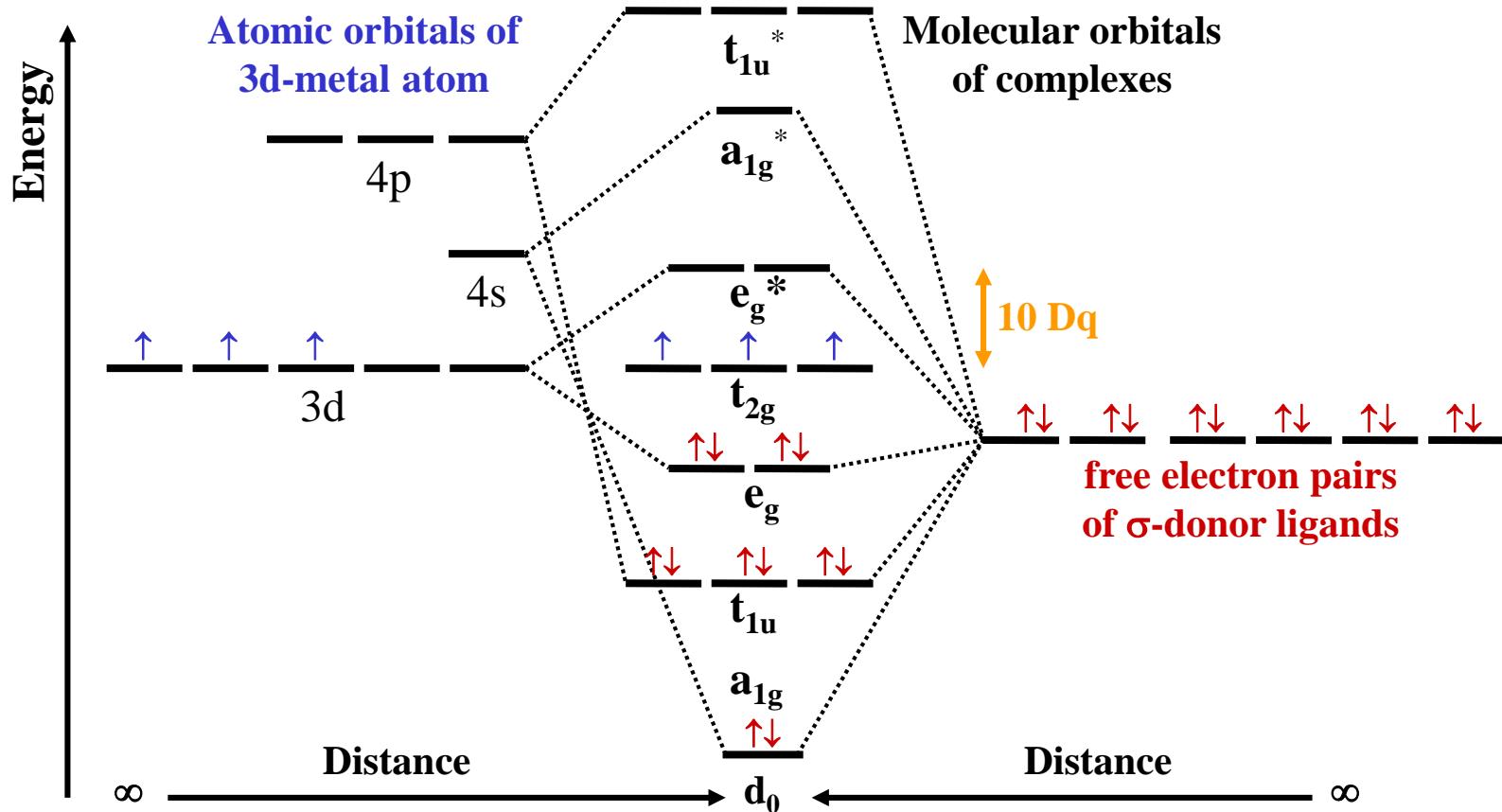
(Metal-ligand-multiple bonds)

Stabilise metals in high oxidation states \Rightarrow O²⁻, N³⁻

7. Principles in Coordination Chemistry

Molecular Orbital (MO) Theory

- ⇒ Overlap of metal and ligand orbitals leads to formation molecular orbitals
- ⇒ Example: octahedral complex built up by **6 σ-donor ligands** and **3d-metal atom**

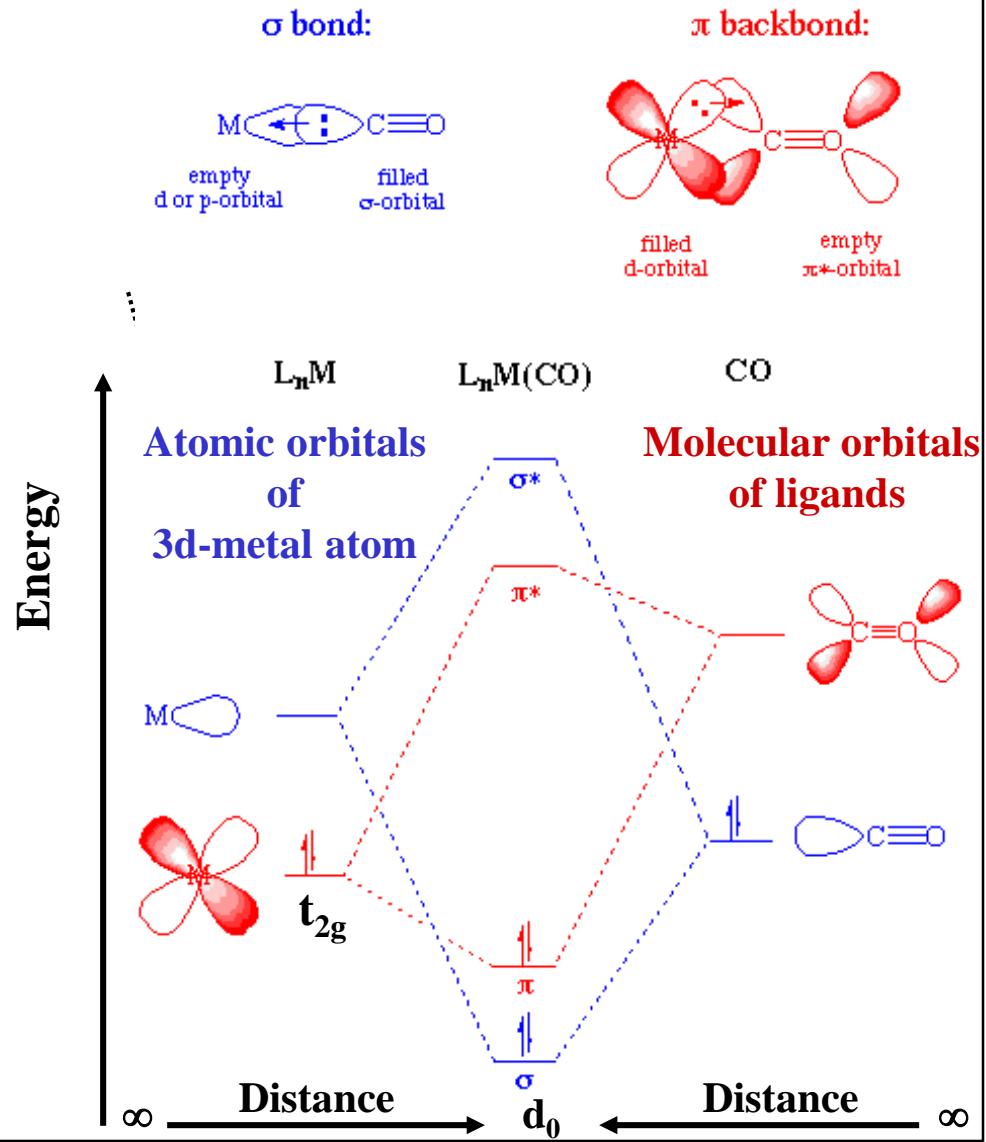


7. Principles in Coordination Chemistry

Molecular Orbital (MO) Theory

Effects of π -back binding

- Strengthening of metal-ligand-bond
- Enhanced crystal field splitting by lowering the energy of the t_{2g} -orbitals
- Weakening of intra-ligand bonding through transfer of electron density into anti-bonding molecular orbitals of the ligand
 - ⇒ decreased vibrational frequencies
 - ⇒ increased reactivity of the ligands (activation)
 - ⇒ catalytic and enzymatic reactions



7. Principles in Coordination Chemistry

Molecular Orbital (MO) Theory

Explanation of ligand ordering in the spectrochemical series



Strong ligands

unoccupied π^* -orbitals

π -acceptor ligands

no suitable p-orbitals

pure σ -donor ligands

Weak ligands

occupied p-orbitals

π -donor ligands

Type of ligand	Effect on metal-ligand-bond	Cristal field splitting
π -acceptor	highly stabilizing	high
σ -donor	stabilizing	intermediate
π -donor	destabilizing	small

8. Properties of Biomolecules

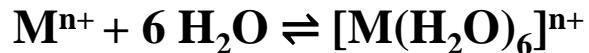
Biological Molecules – Overview and Quantities in a Typical Eukaryotic Cell

• Water		70%
• Proteins	→ Polypeptides → Amino acids	15%
– Structural proteins		
– Transportational proteins		
– Storage proteins		
– Sphere proteins (enzymes)		
• Nucleic acids	→ Polynucleotides → Phosphate + Bases + Desoxyribose	
– DNA		1%
– RNA		6%
• Starch/cellulose	→ Polysaccharides → aldoses/ketoses	3%
• (Phospho)lipids	→ Glycerine + fatty acids (+ phosphates)	2%
• “Monomers”	→ Prosthetic groups, Co-factors	2%
• Inorganic ions	→ Na^+ , K^+ , Mg^{2+} , Ca^{2+} , $\text{Fe}^{\text{n}+}$, $\text{Mn}^{\text{n}+}$, $\text{Co}^{\text{n}+}$, $\text{Cu}^{\text{n}+}$, Zn^{2+} , F^- , Cl^- , HCO_3^- , PO_4^{3-} , MoO_4^{2-} , WO_4^{2-}	1%
• Other “inorganics”	→ O_2 , CO_2 , CO, NO, CN^- , OCN^- , H_2O_2 ,	ppm-ppb

8. Properties of Biomolecules

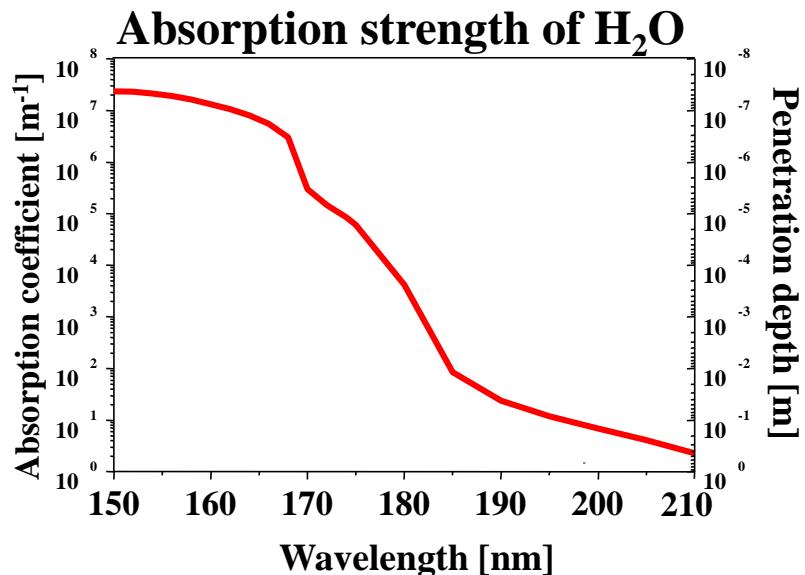
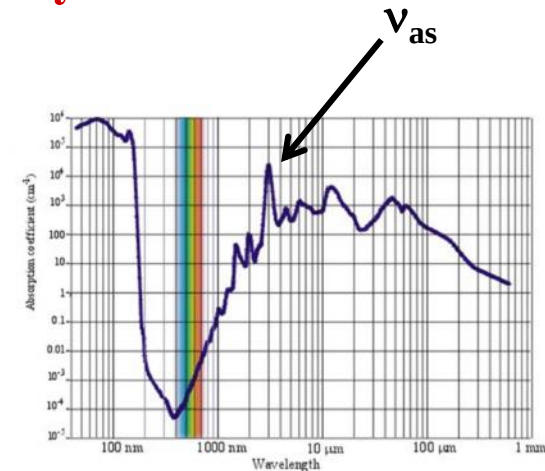
Water → Solvents ⇒ Biochemistry = “Aqueous Chemistry“

- Auto-proteolysis: $2 \text{H}_2\text{O} \rightleftharpoons \text{H}_3\text{O}^+ + \text{OH}^-$ pK_a (25 °C) = 14.0
- High dipole moment: $\mu = q \cdot d = 1.85 \text{ Debye [Cm]}$
⇒ High polarity and strong H-bridges
- Optical transparency about 200 – 800 nm
⇒ absorption in the IR- as well as VUV/EUV-range
- Metal cations increase acidity of water



Metal cation pK_a (25 °C, 0.1 M)

Ca^{2+}	13.4
Mn^{2+}	11.1
Cu^{2+}	10.7
Zn^{2+}	10.0
Fe^{3+}	2.2



8. Properties of Biomolecules

Proteins: Structure

Proteins are built up from one or multiple polypeptide sequences, which, themselves, are formed by the combination of amino acids.

The reaction leads to acid amides and is called peptide bond.

Amino acids (AS) -OOC-CH(R)-NH_3^+

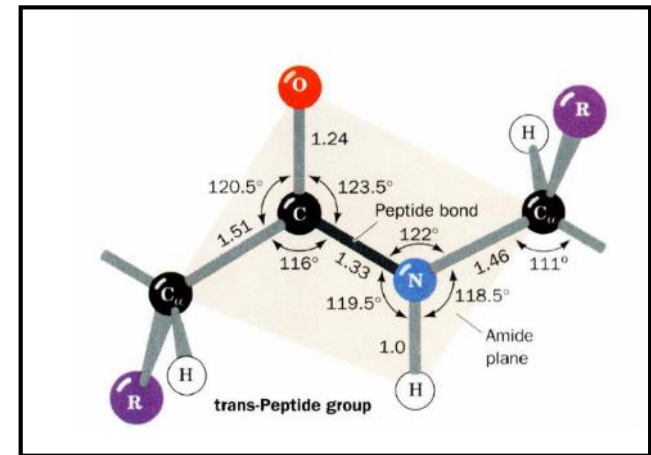
↓
Polypeptides (100 – 100000 AS, $M = 10^5 – 10^8 \text{ g/mol}$)

↓
Proteins (one or multiple polypeptide chains)

→ Metal proteins

↓
Holo protein (= protein + prosthetic group)

→ Metal holo proteins

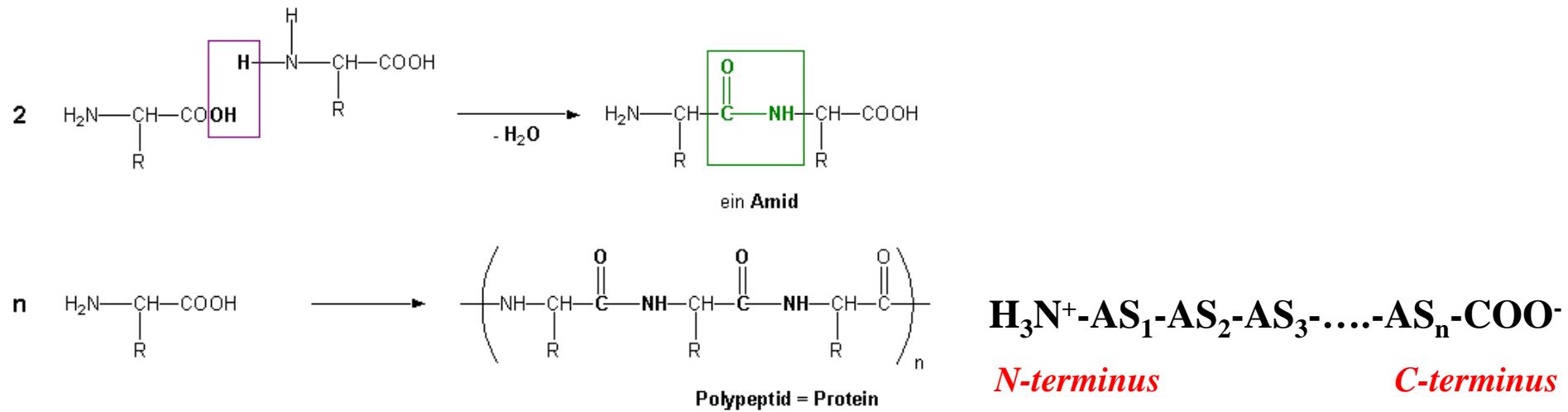


8. Properties of Biomolecules

Proteins: Formation and Structure

1. Linkage of amino acids to polypeptides (primary structure)

The synthesis takes place in the ribosomes (AS-sequence is determined by m-RNA)



2. Folding of polypeptides to 3-dimensional constructions (secondary and tertiary structure)

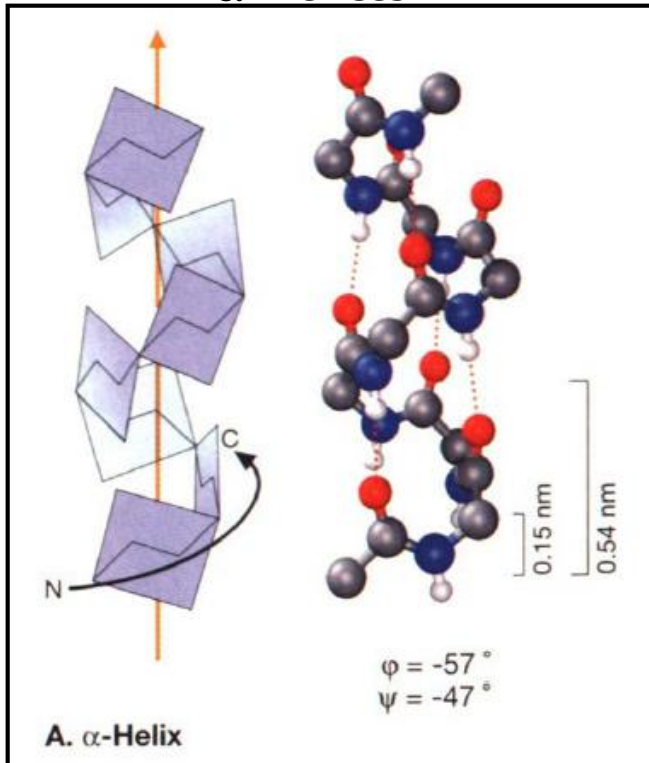
- by van-der-Waals interactions (steric)
 - by ionic interactions (electrostatic) \Rightarrow **stabilisation via metal cations**
 - by hydrogen bonds (weak bonding)
 - by disulphide bridges R-S-S-R (strong bonding)

8. Properties of Biomolecules

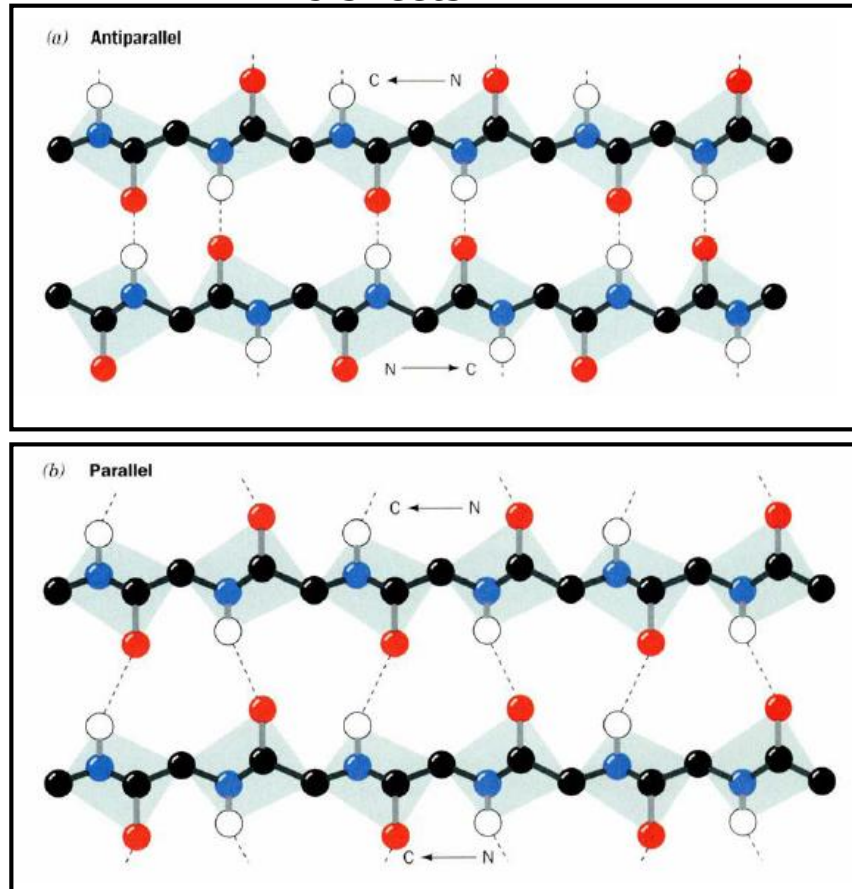
Proteins: Secondary Structure

Hydrogen bridges and other interactions lead to secondary structures

α -Helices



β -sheets



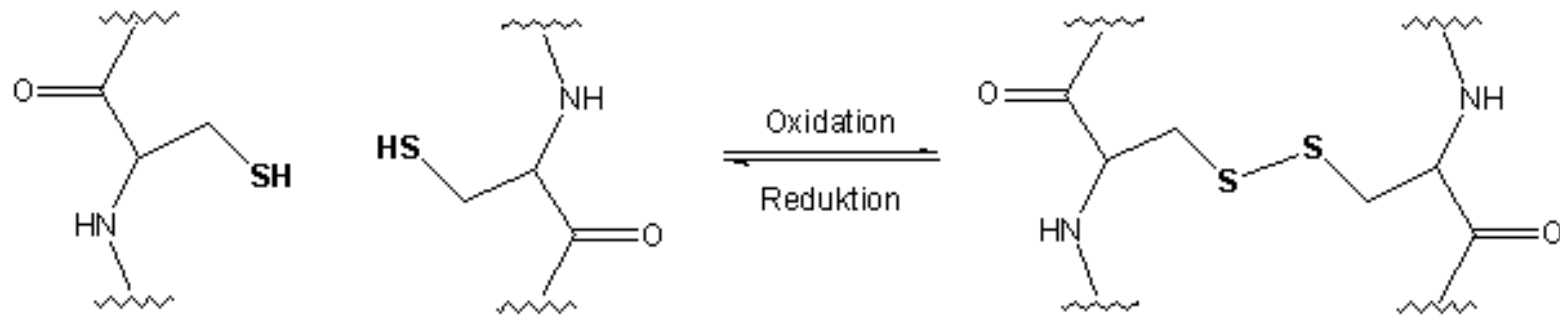
Collagens (triple helix)

α -keratins (2 x double helix)

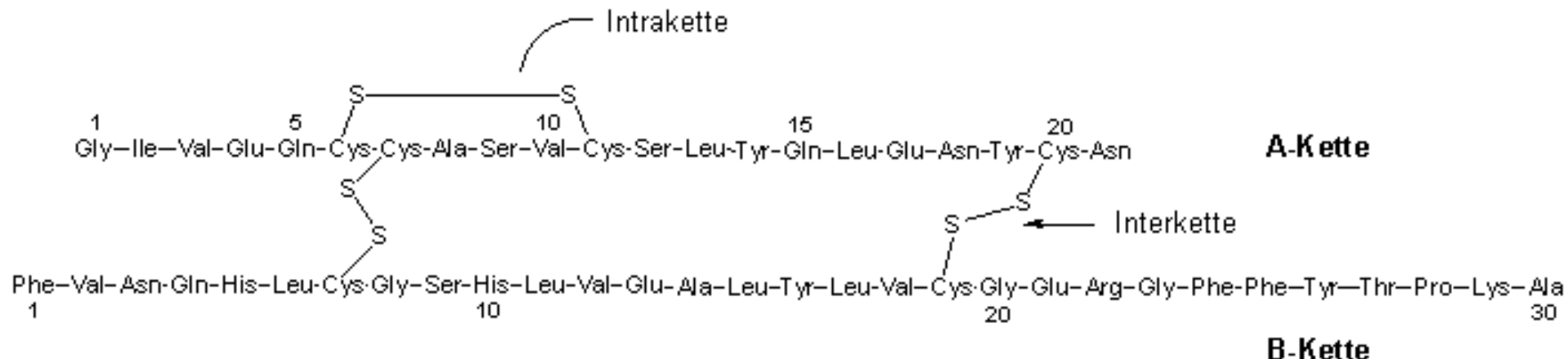
8. Properties of Biomolecules

Proteins: Disulphide Bridges R-S-S-R'

Enzymatic formation:



Disulphide bridges can link AS within a chain or between two chains



8. Properties of Biomolecules

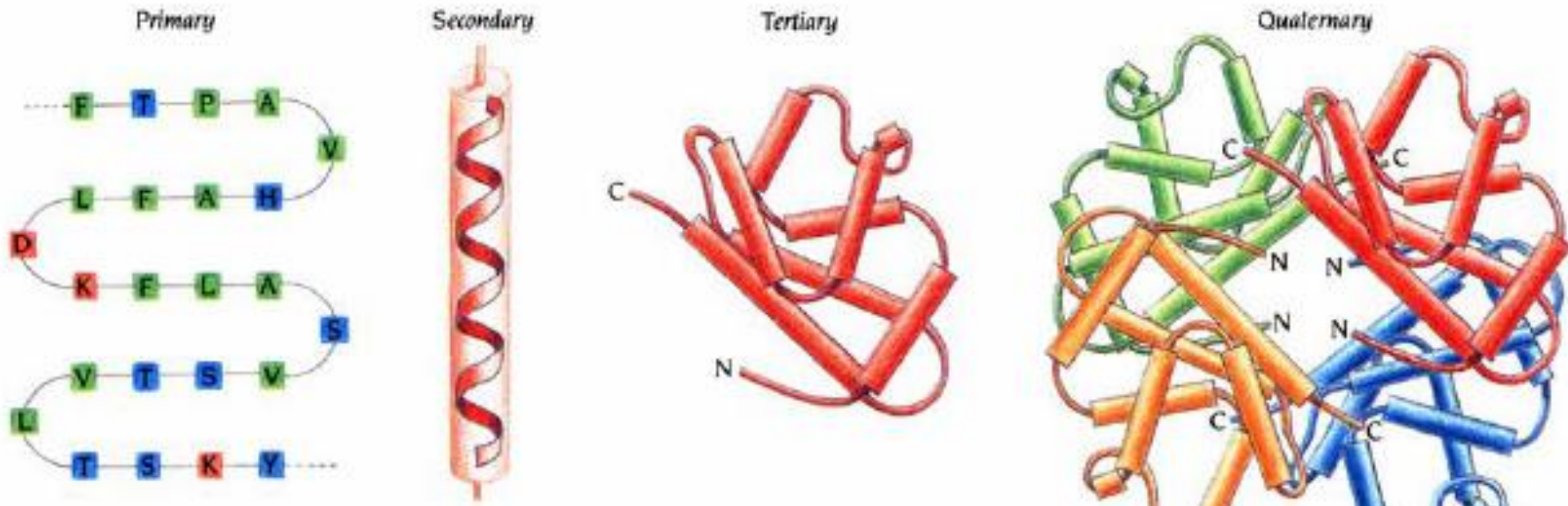
Proteins: Tertiary and Quaternary Structure

Tertiary structure:

three dimensional structure of a single polypeptide chain

Quaternary structure:

arrangement of multiple polypeptide chains in a protein



8. Properties of Biomolecules

Proteins: Properties

Physical

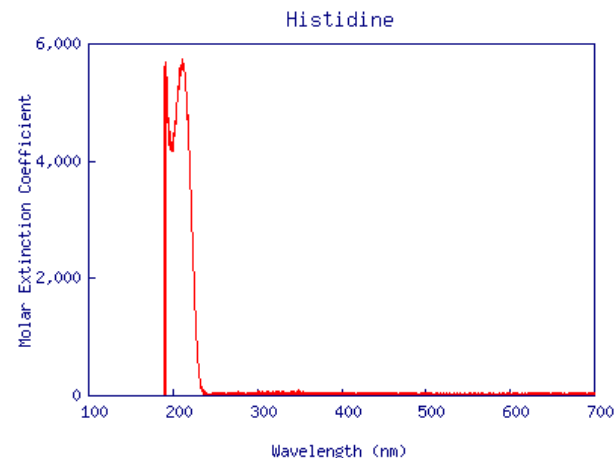
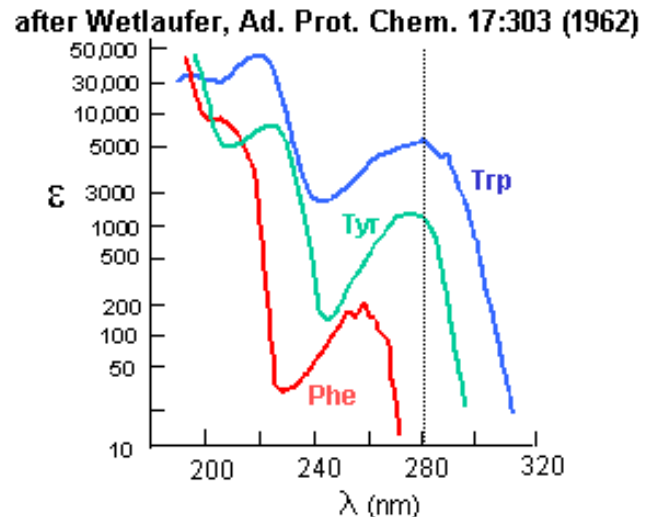
- Stability: low (enzymes) till high (horn)
- Temperature sensitivity \Rightarrow denaturation upon heating
- Some proteins are insoluble,
some give colloidal solutions

Solutions of proteins are optically active

- Absorption in UV-range
 - Aliphatic < 240 nm
 - Aromatic < 320 nm

Chemical

- Hydrolysis upon heating in combination with acids or bases:
Polypeptide \rightarrow amino acids
- Cleavage by proteases (at defined interfaces)
 \Rightarrow sequence analysis



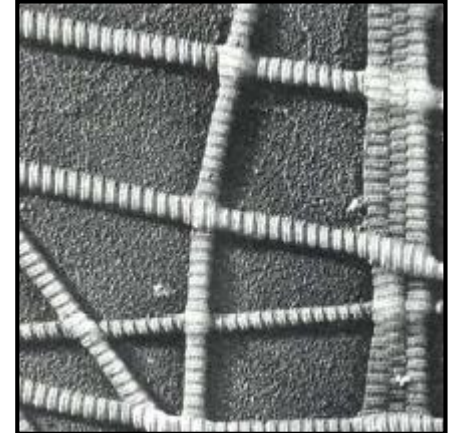
8. Properties of Biomolecules

Proteins: Functions

Structural proteins

- Collagen
- α -keratins, β -keratins

TEM-images
of collagen fibres

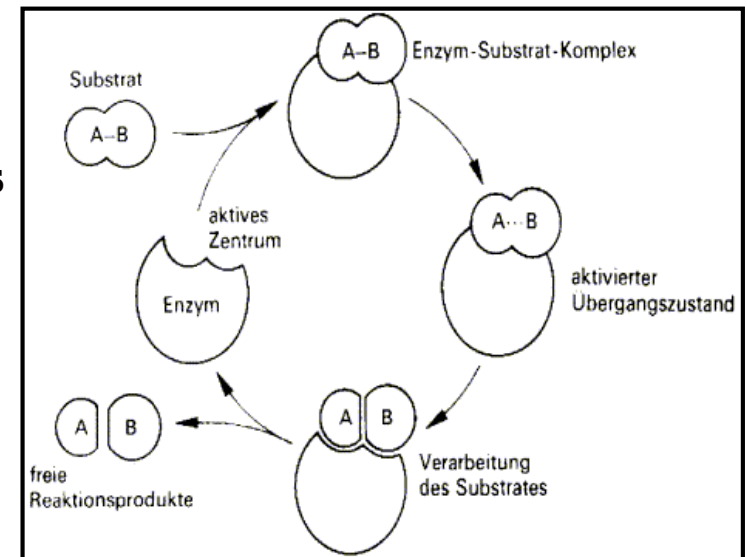


Transportation and storage proteins

- Oxygen transport: heme- and myoglobin
- Storage of iron: ferritin and ferredoxine

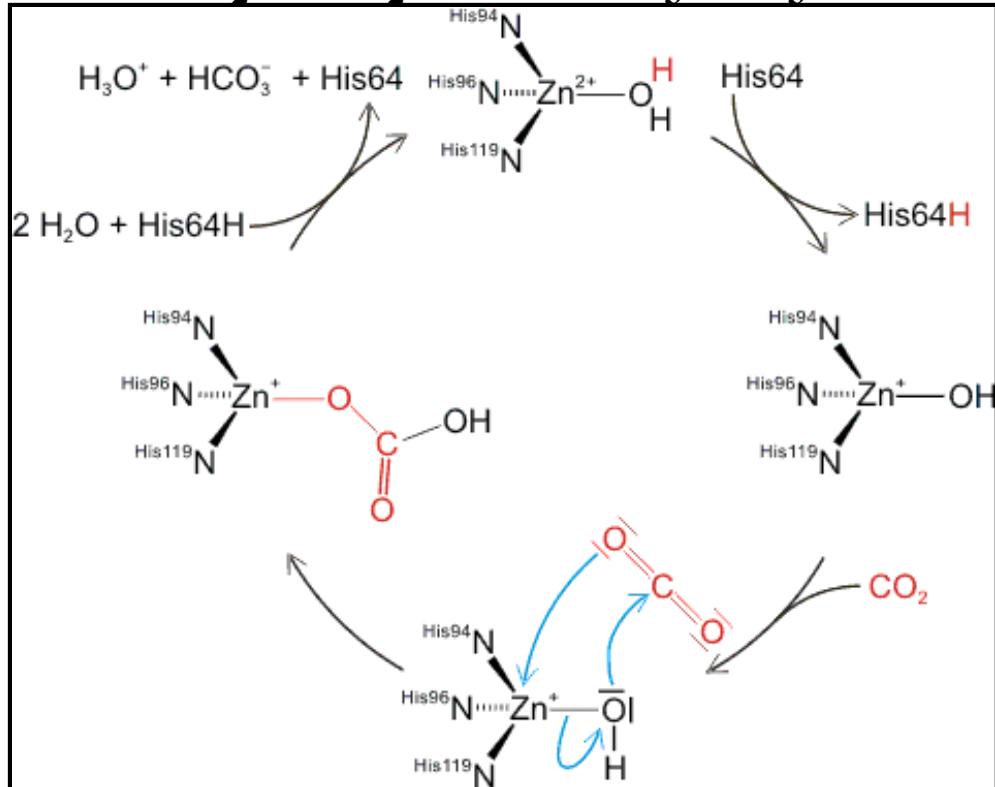
Sphere proteins (enzymes)

- Oxidoreductases → redox reactions (e.g. catalysis)
- Transferases → transfer of small molecular groups (e.g. hexokinases)
- Hydrolases → hydrolysis of proteins, sugars and lipids (e.g. amylases, ureases, trypsin)
- Lyases → addition reactions of small molecules at double bonds (e.g. citrate synthase)
- Ligases → linkage of small molecules to bigger units (e.g. DNA ligase)
- Isomerases → alteration of molecular constitution (e.g. phosphoglucomutase)

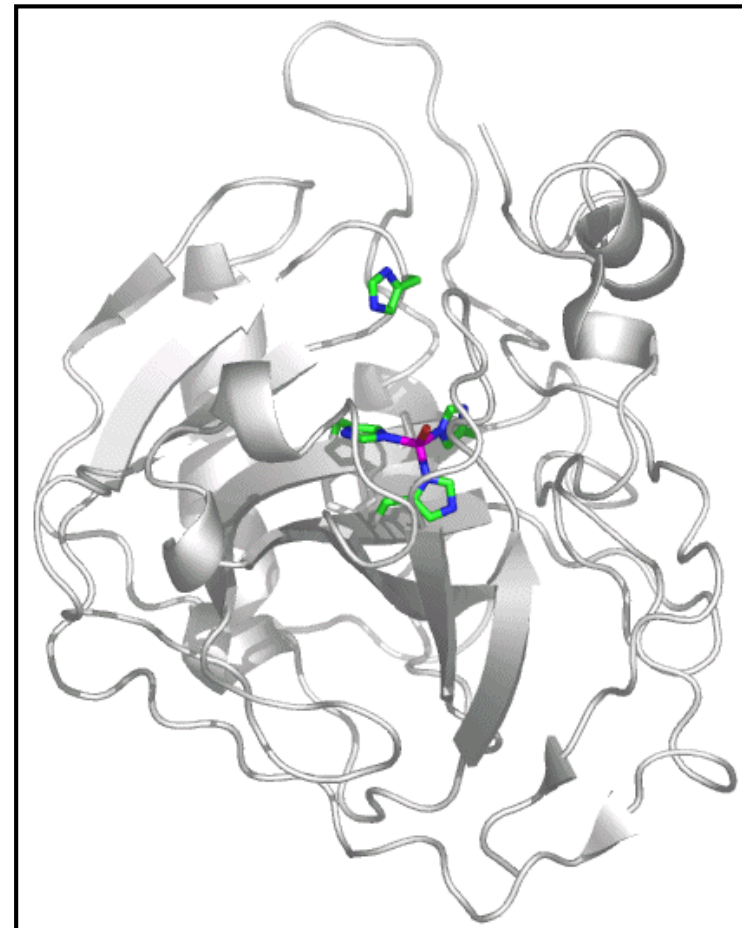


8. Properties of Biomolecules

Proteins: Function of the Zn²⁺-Enzyme, Carbonic Anhydrase



A lot of other Zn-enzymes also catalyse hydrolysis of polar bonds, such as proteases and esterases

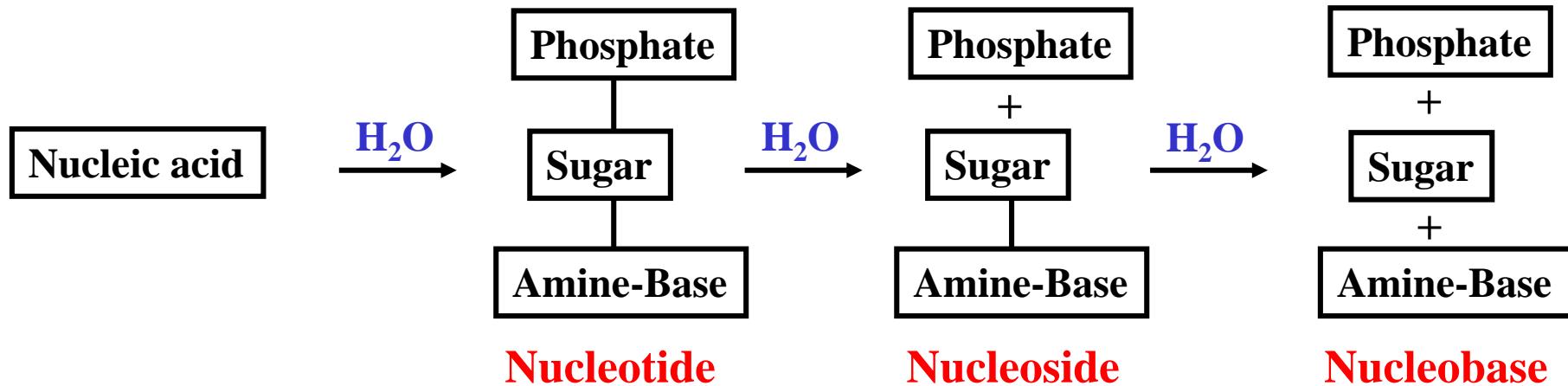


Single-stranded protein consisting of 260 amino acids

8. Properties of Biomolecules

Nucleic Acids: Building Blocks

During hydrolysis of a nucleic acid sugar, phosphate and amine-bases are formed:

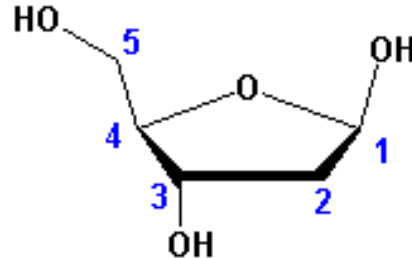


<u>Nucleic acids</u>	<u>Sugar</u>	<u>Amine bases</u>
DNA	2-desoxyribose	cytosine, thymine, adenine, guanine
RNA	D-ribose	cytosine, uracil, adenine, guanine

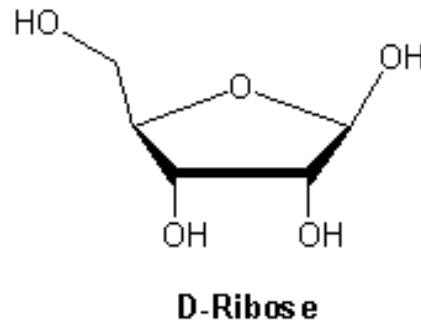
8. Properties of Biomolecules

Nucleic Acids: Building Blocks

Sugar

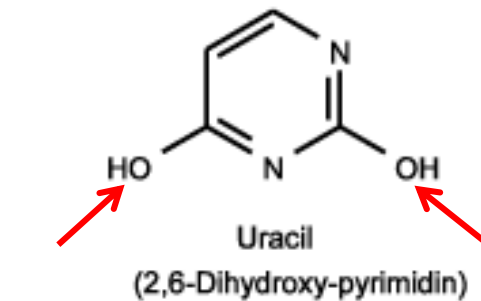
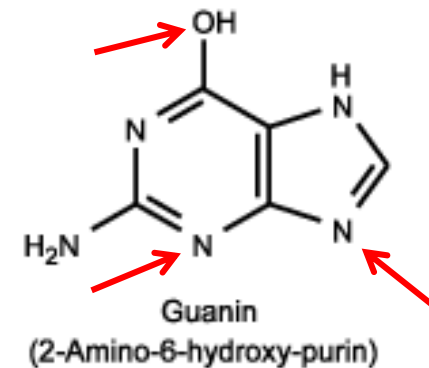
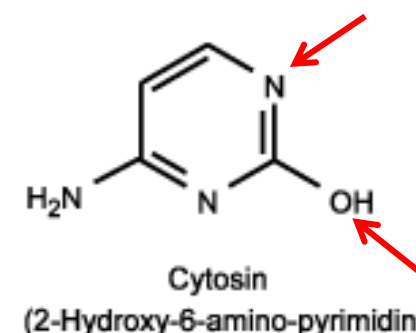
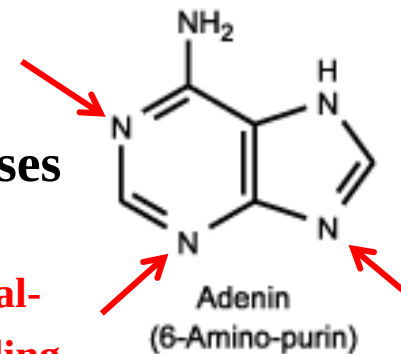


2-deoxyribose



Amine bases

Metal-binding-positions



⇒ Unstable against hydrolysis

8. Properties of Biomolecules

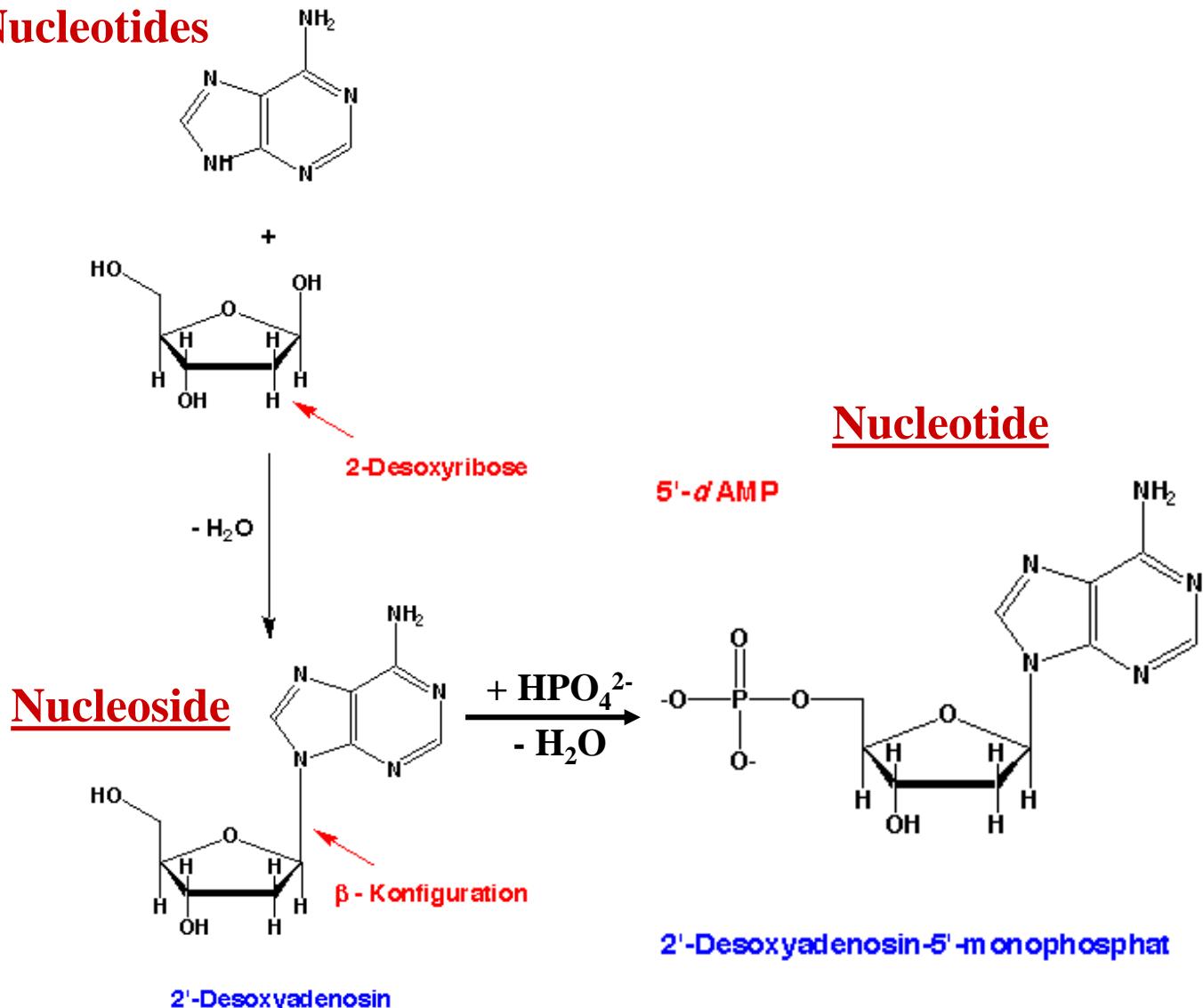
DNA: Nucleosides and Nucleotides

dAMP =
desoxyadenosine-
monophosphate

dGMP =
desoxyguanosine-
monophosphate

dCMP =
desoxycytidine-
monophosphate

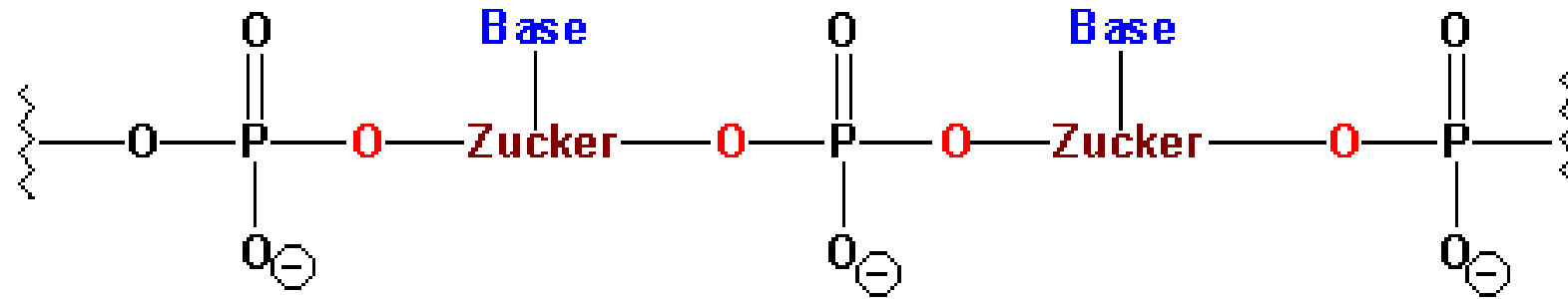
dTMP =
desoxythymidine-
monophosphate



8. Properties of Biomolecules

DNA: Primary Structure (Sequence)

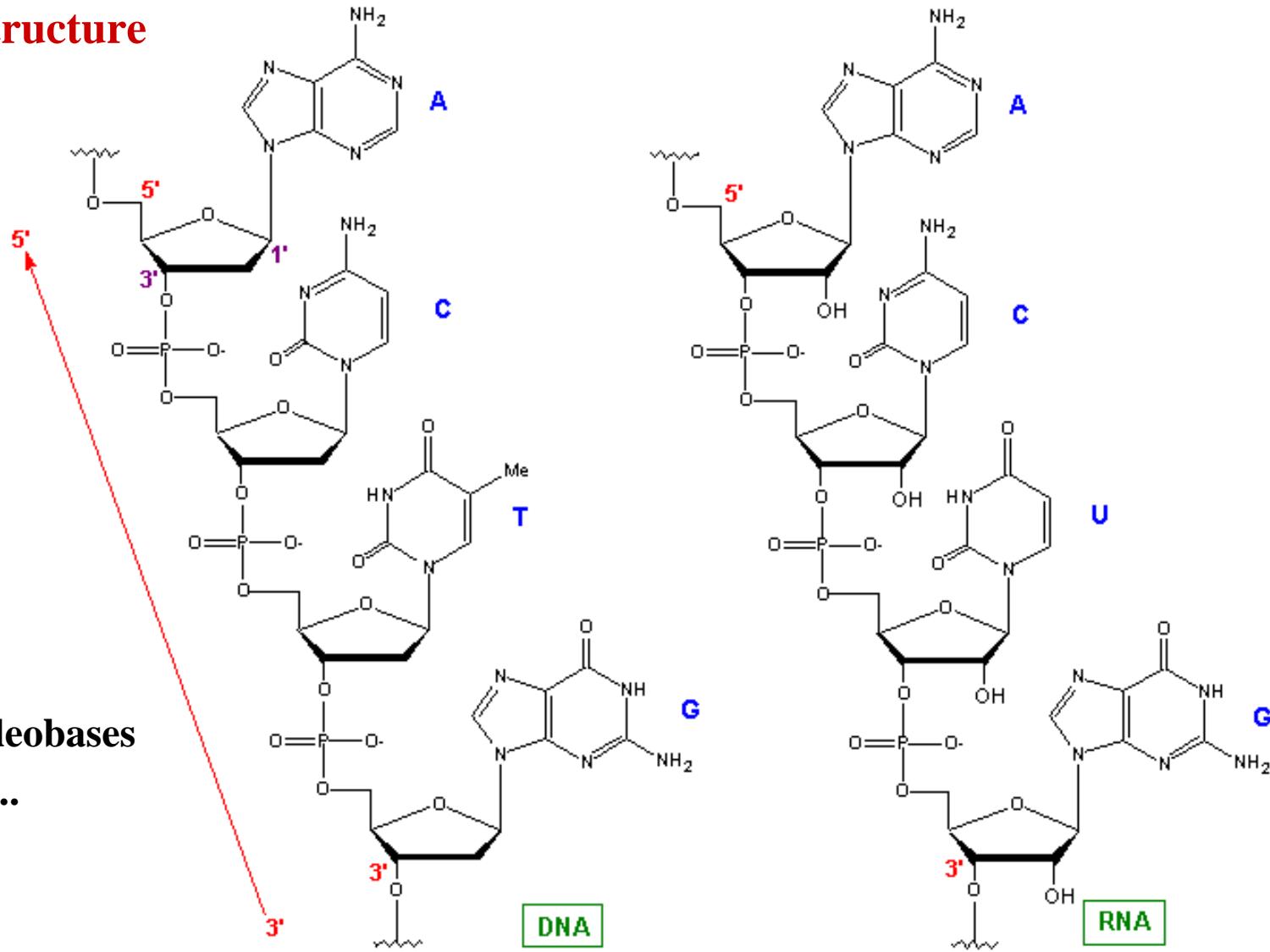
A polymer chain is formed through the continuous linkage of phosphate ester bridges between the C5 of the sugar unit from one nucleotide to the C3 of another sugar



One end of the polymer chain possesses a free hydroxyl group at the C3' (3'-end) and the other possesses another phosphate unit at C5 (5'-end)

8. Properties of Biomolecules

DNA and RNA: Structure



Hetero atoms of nucleobases

→ Cu²⁺, Cr³⁺, Pt²⁺, ...

Phosphate backbone

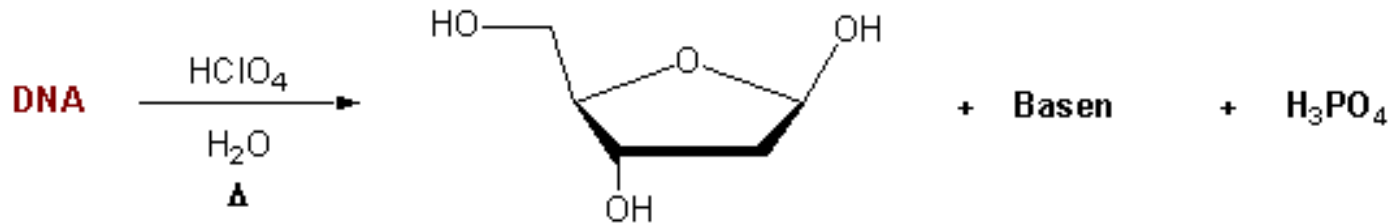
→ Mg²⁺, Na⁺, K⁺

8. Properties of Biomolecules

DNA and RNA: Chemical Properties

For acidic hydrolysis

A strong acid is needed



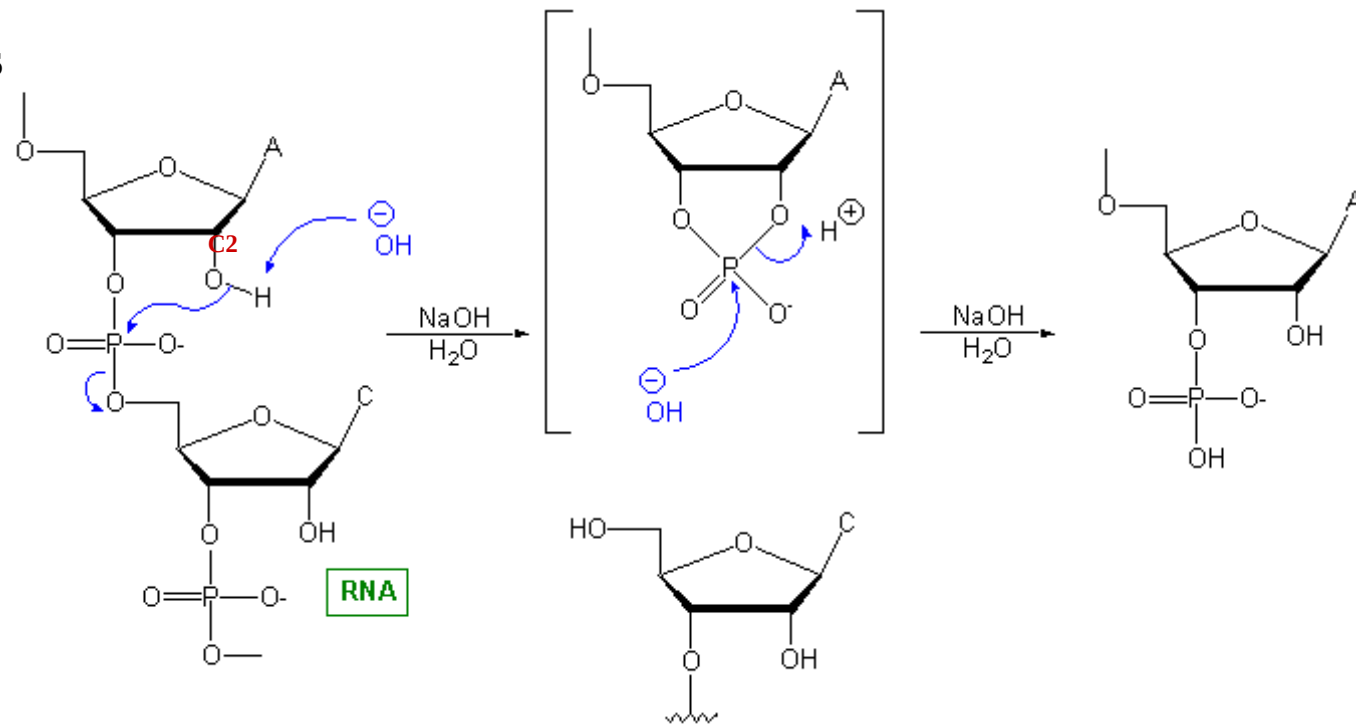
In basic conditions DNA

is relatively stable, whereas

RNA is rapidly cleaved
(cyclic phosphor-
acid ester)

DNA misses the hydroxyl

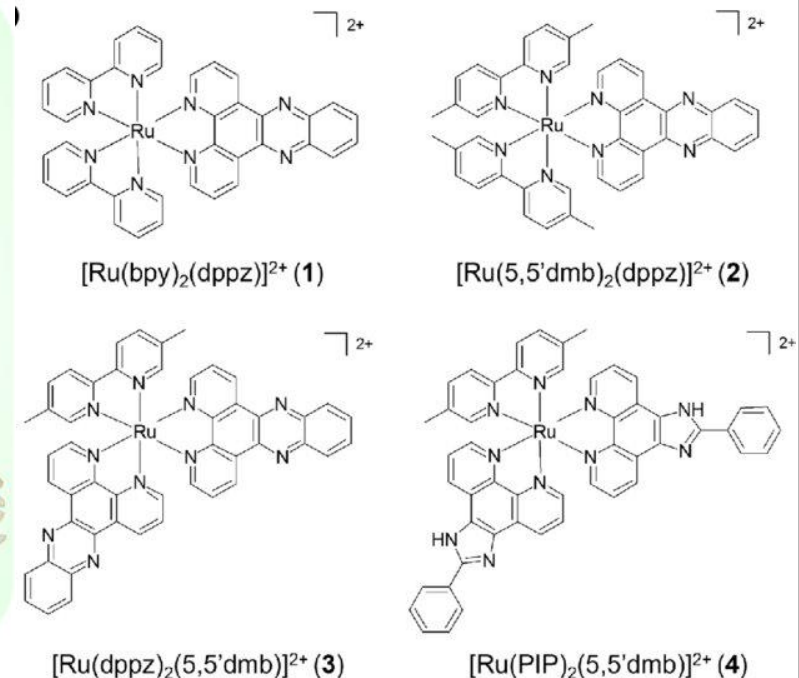
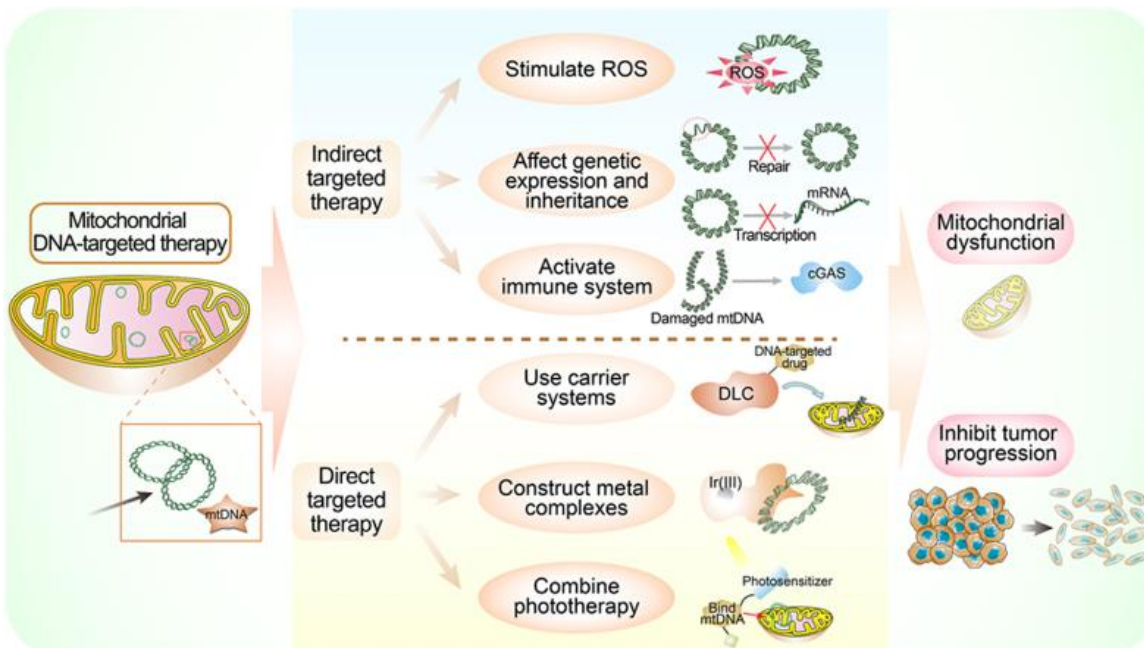
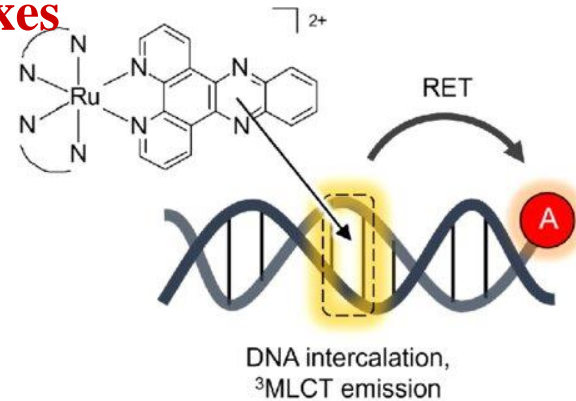
Group at C2



8. Properties of Biomolecules

DNA and RNA: Interaction with Ru²⁺ or Ir³⁺ complexes

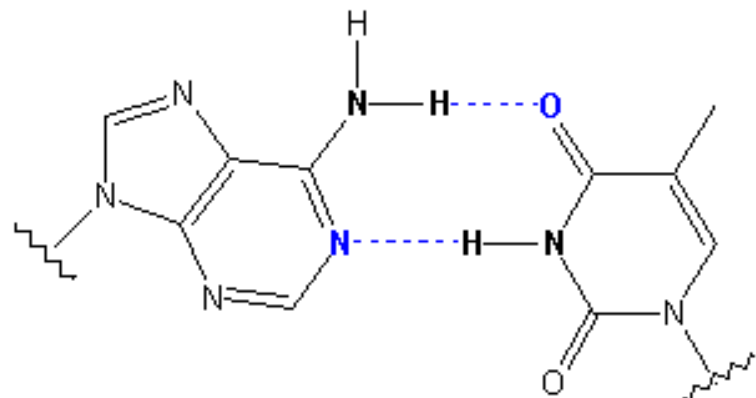
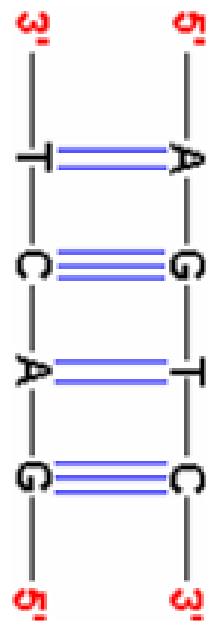
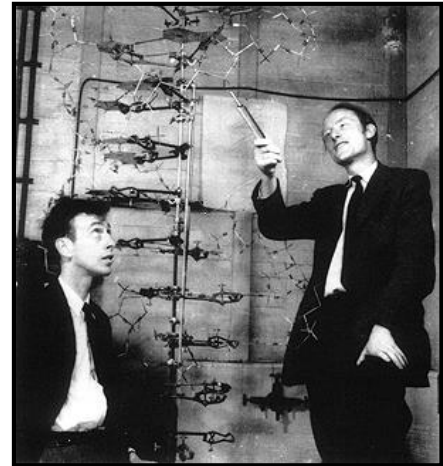
Goal: Gene therapy, e.g. of cancer by intercalation of Ru²⁺ complexes into mt-DNA of tumour cells



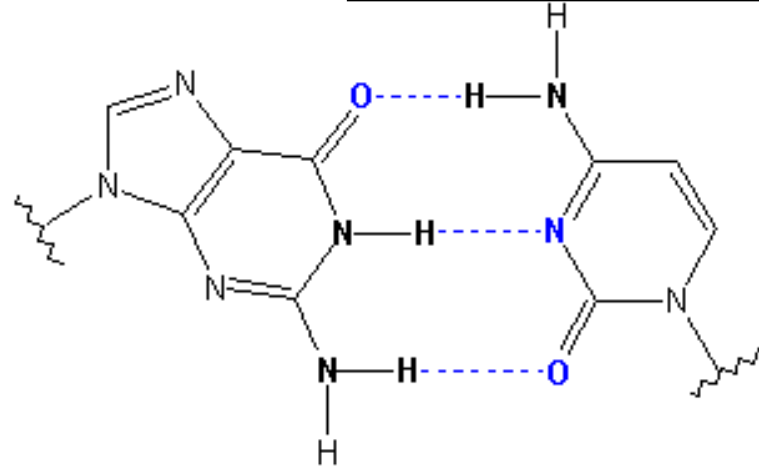
8. Properties of Biomolecules

DNA: Secondary Structure Suggested in 1953 by James D. Watson & Francis Crick

- DNA is a double helical structure, consisting of two strands with complimentary base sequences
- The ratio of A to T and G to C is always one to one
- The bases A and T as well as the bases G and C are linked via hydrogen bonds



$\Delta G^\circ \sim -1.2 \text{ kcal/mol}$

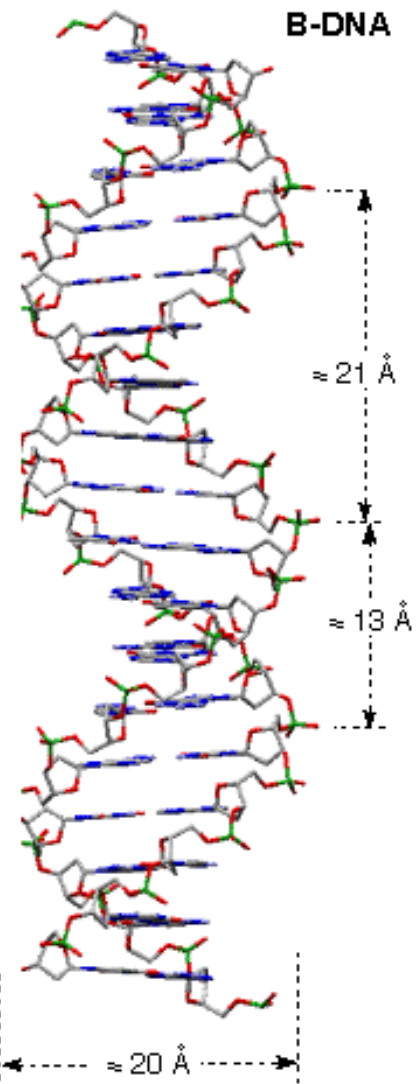
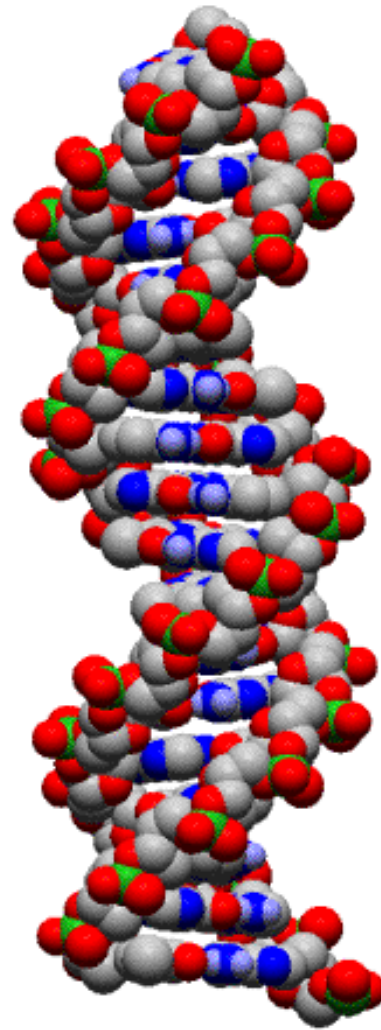
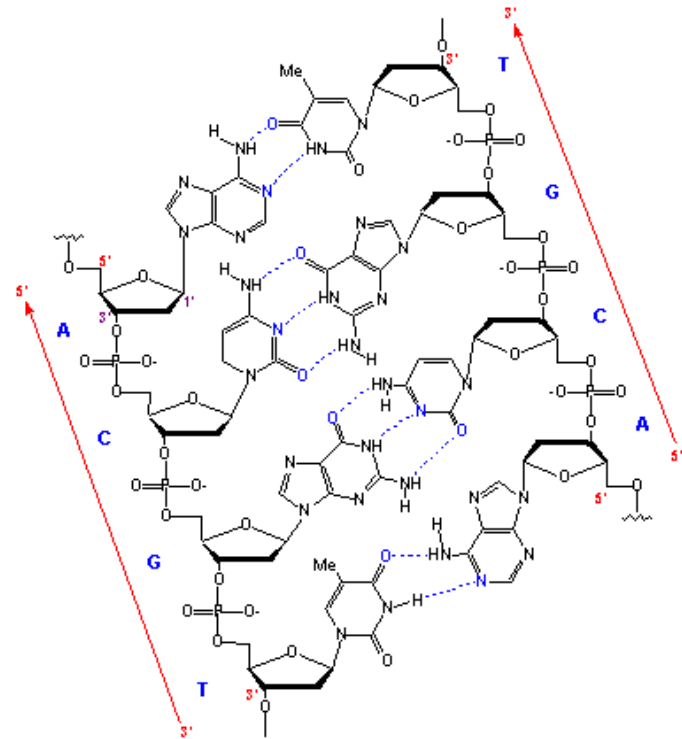


$\Delta G^\circ \sim -2.4 \text{ kcal/mol}$

8. Properties of Biomolecules

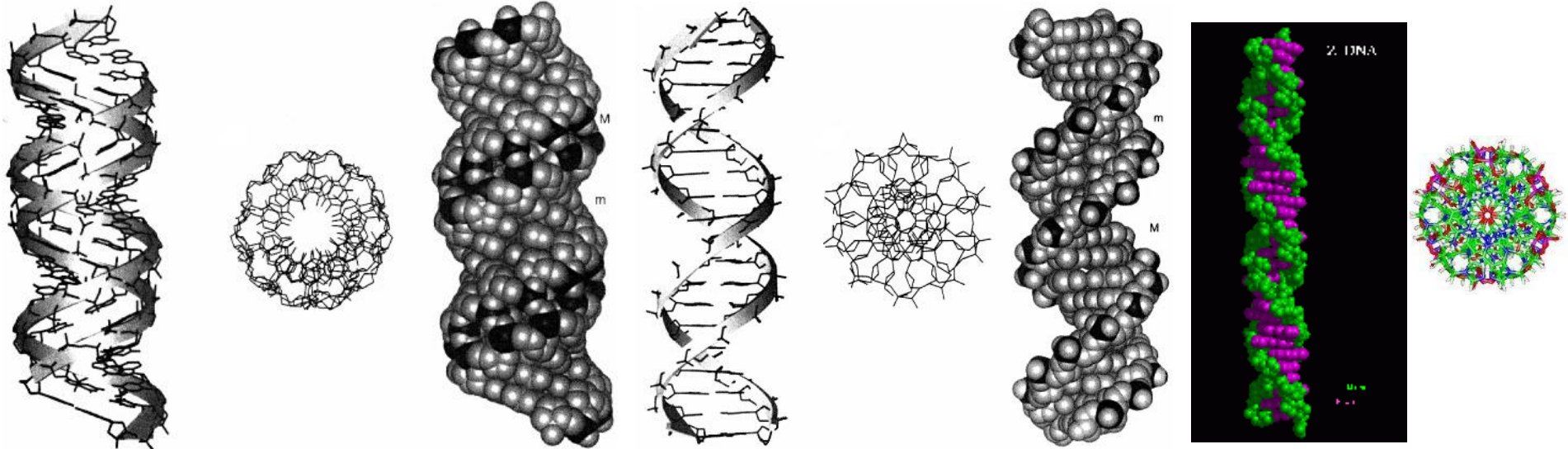
DNA: Secondary Structure Is a Double Helix

Due to structural reasons, the arrangement, where hydrogen bonds are optimally formed and sterical hindrance is minimized, is a double helix



8. Properties of Biomolecules

DNA: Secondary Structure A-, B-, and Z-DNA



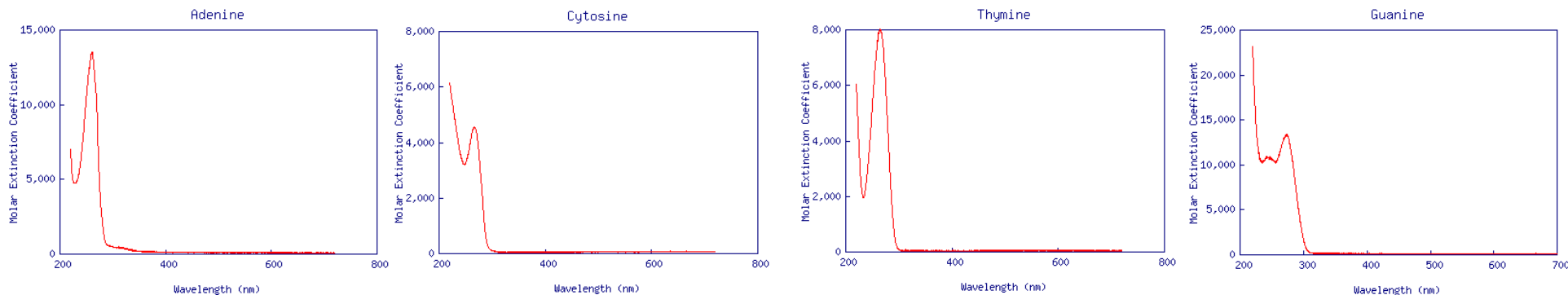
Parameter	A-DNA	B-DNA	Z-DNA
Form	broad	intermediate	narrow
Increase per Bp [nm]	0.23	0.34	0.38
Helix diameter [nm]	2.6	2.4	1.8
Sense of rotation	right	right	left
Bp per helical turn	11	10.4	12
Pitch [nm]	2.5	3.5	4.7
Angle Bp towards helical axis	19°	1°	9°

Source: Neidle, Stephen, *Nucleic Acid Structure and Recognition*, Oxford University Press, 2002, p. 36.

8. Properties of Biomolecules

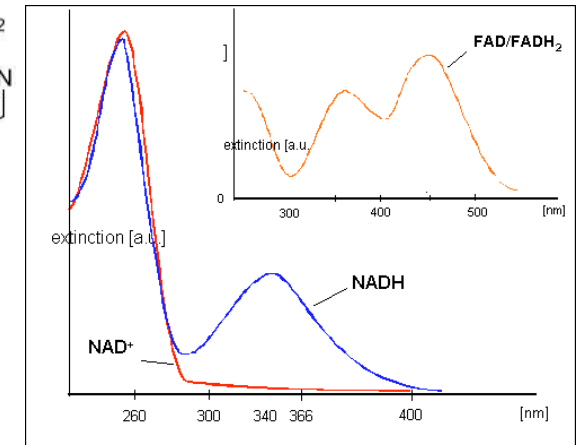
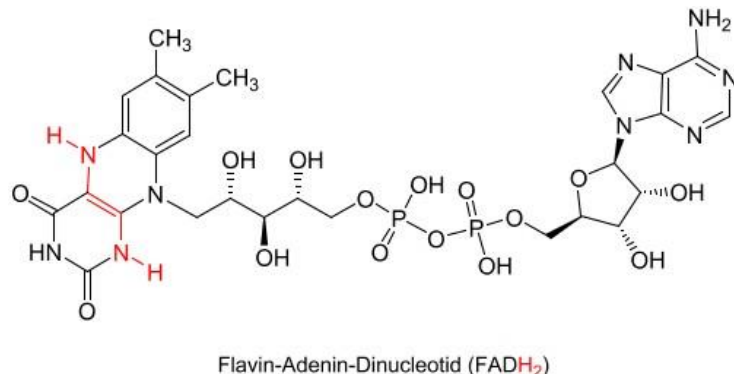
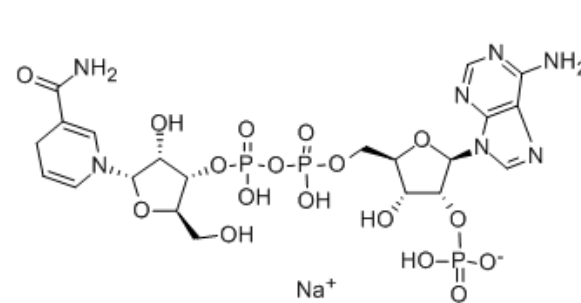
Spectroscopic Properties of Nucleo Bases

Absorption bands at 265 nm (A, T, C, G) and at 240 nm (G)



Other biomolecules, that are absorbing even in the near-UV or blue spectral range are

NAD(P)H or FADH₂



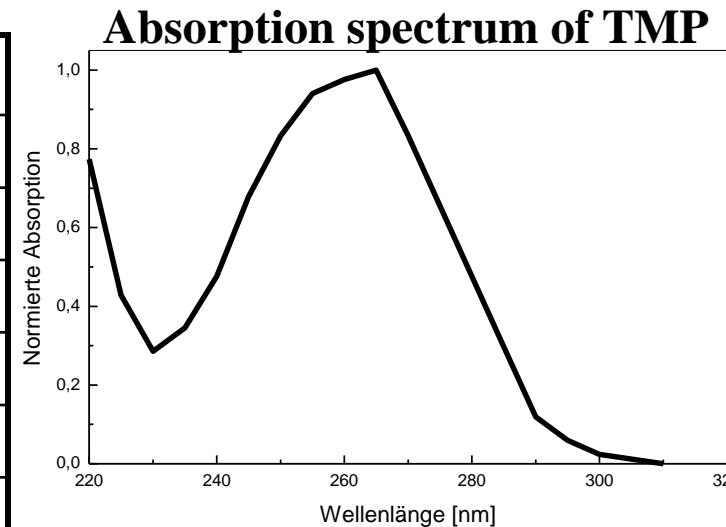
8. Properties of Biomolecules

Spectroscopic Properties of Nucleo Bases

Species	λ_{\max} [nm]	ϵ [$\text{l mol}^{-1}\text{cm}^{-1}$]	Transition
Adenine	260	13400	n- π^* , π - π^*
Guanine	275	8100	n- π^* , π - π^*
Cytosine	267	6100	n- π^* , π - π^*
Thymine	264	7900	n- π^* , π - π^*
AMP	260	15500	n- π^* , π - π^*
ss-poly-AMP	260	10600	n- π^* , π - π^*
ds-poly-AMP	258	9600	n- π^* , π - π^*

Information from the UV-absorption spectrum

- AT- and GC-content
- Single or double helix
- Thermal stability of DNA
- Melting point of DNA
(temperature when double helix is cleaved)

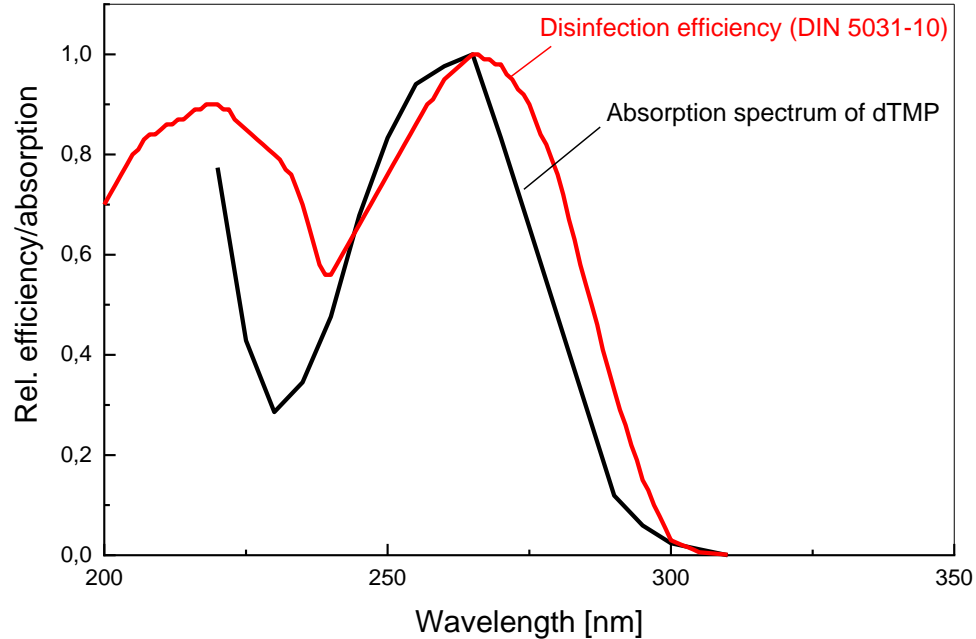


Chargaff's DNA Data Base Composition in Various Species (%)

Species	A	T	G	C
<i>Homo sapiens</i>	31.0	31.5	19.1	18.4
<i>Drosophila melanogaster</i>	27.3	27.6	22.5	22.5
<i>Zea mays</i>	25.6	25.3	24.5	24.6
<i>Neurospora crassa</i>	23.0	23.3	27.1	26.6
<i>Escherichia coli</i>	24.6	24.3	25.5	25.6
<i>Bacillus subtilis</i>	28.4	29.0	21.0	21.6

8. Properties of Biomolecules

Spectroscopic Properties of Nucleo Bases



Nucleotide

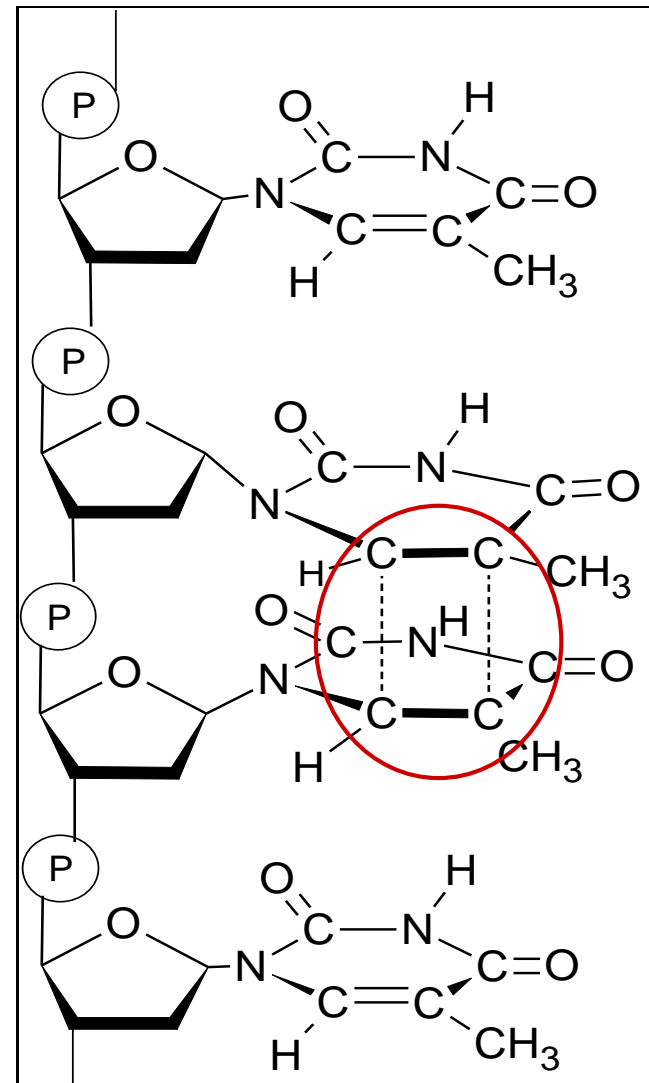
Extinction coefficient ϵ at 260 nm

dAMP $15200 \text{ lmol}^{-1}\text{cm}^{-1}$

dTMP $8400 \text{ lmol}^{-1}\text{cm}^{-1}$

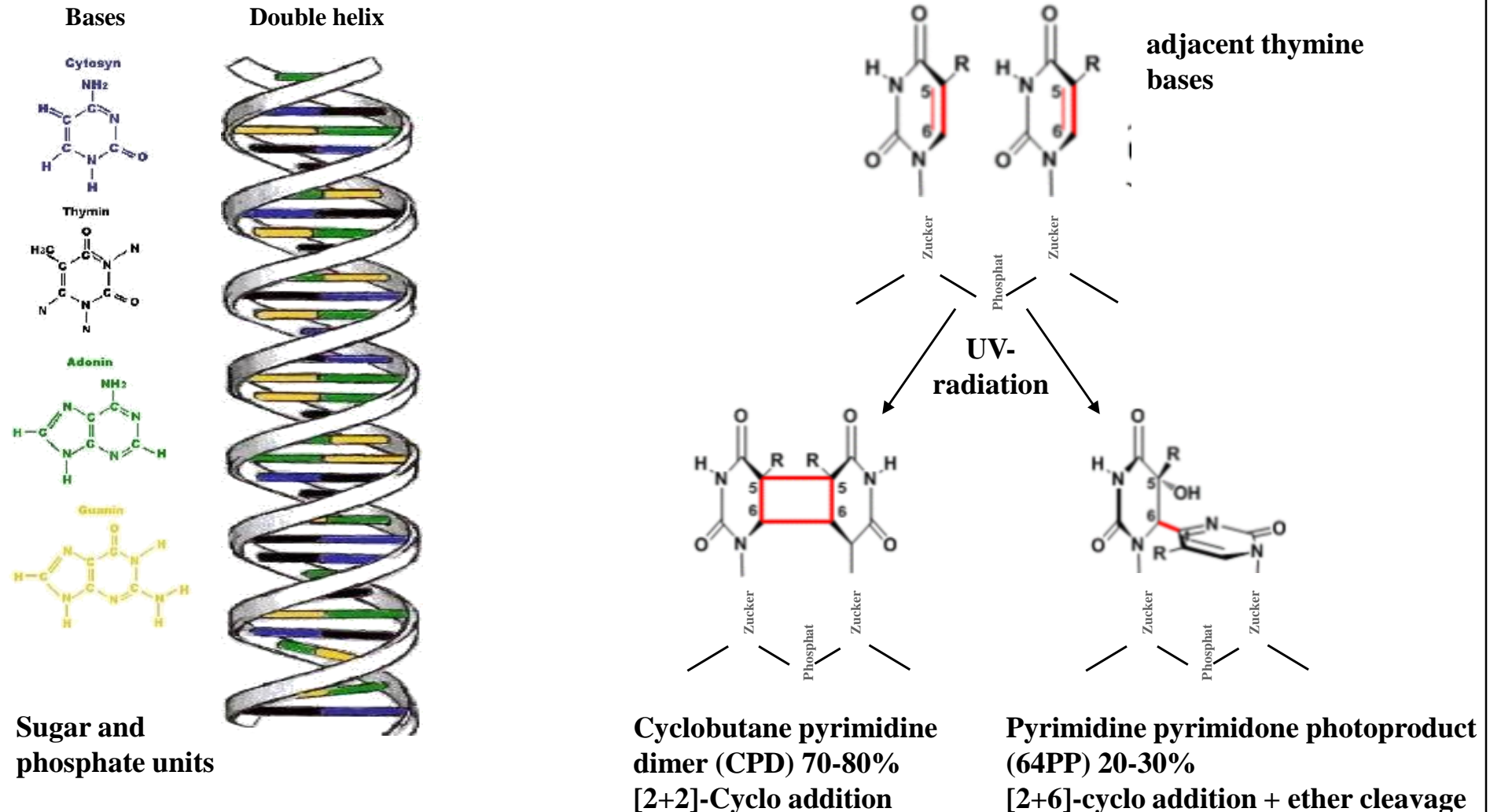
dGMP $12000 \text{ lmol}^{-1}\text{cm}^{-1}$

dCMP $7100 \text{ lmol}^{-1}\text{cm}^{-1}$



8. Properties of Biomolecules

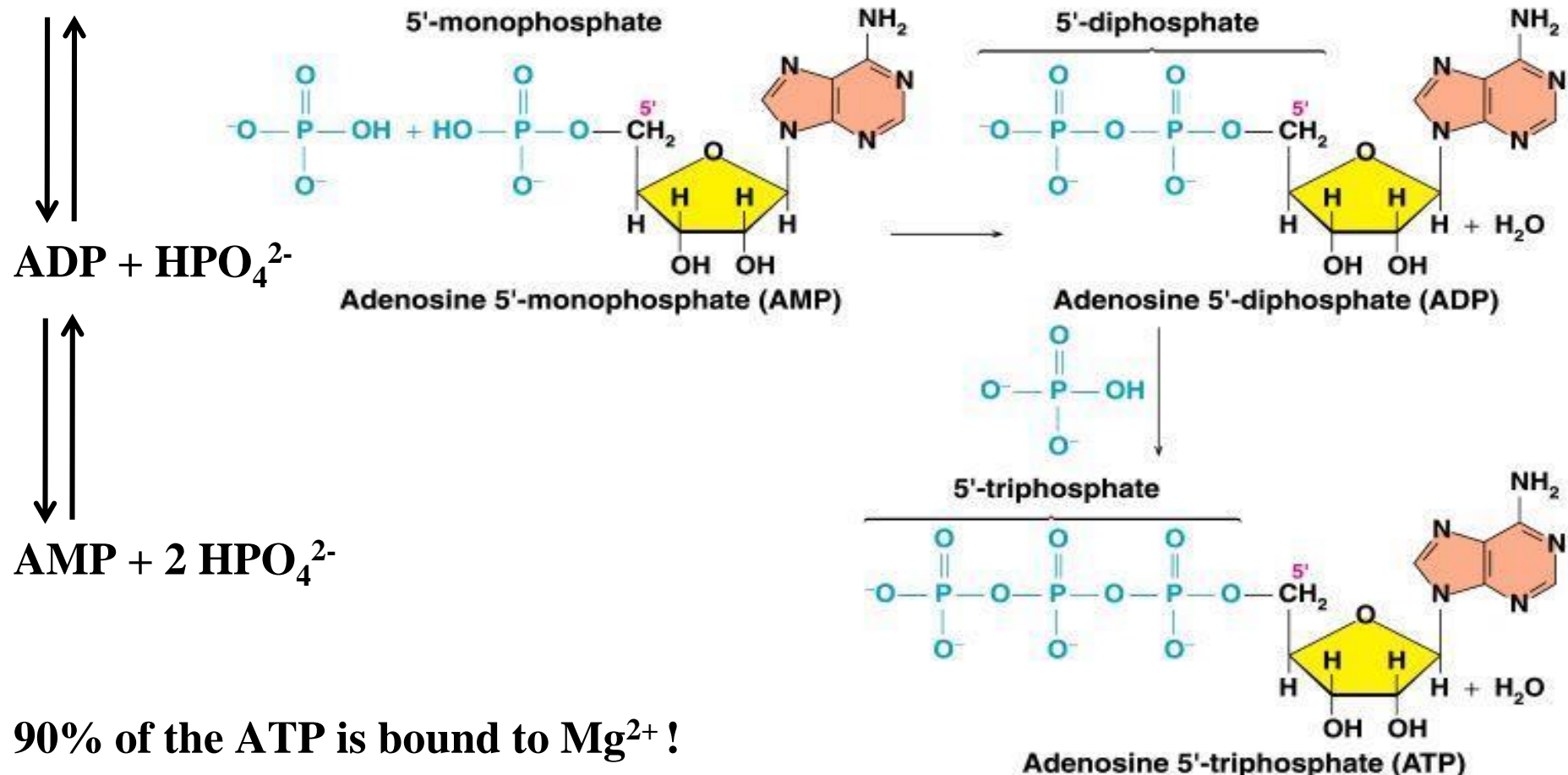
Spectroscopic Properties of Nucleo Bases



8. Properties of Biomolecules

Nucleoside-5'-monophosphates Form Diphosphates and Triphosphates, too

⇒ ATP is the most important energy source for all cellular activities



⇒ 90% of the ATP is bound to Mg^{2+} !

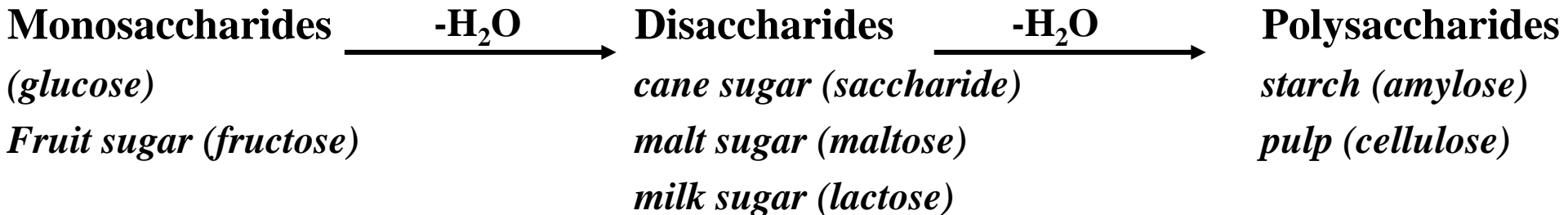
8. Properties of Biomolecules

Carbohydrates: Definition and Nomenclature

Carbohydrates include are huge group of natural substances, such as sugars, starch and celluloses. The name can be derived formally from the general formula

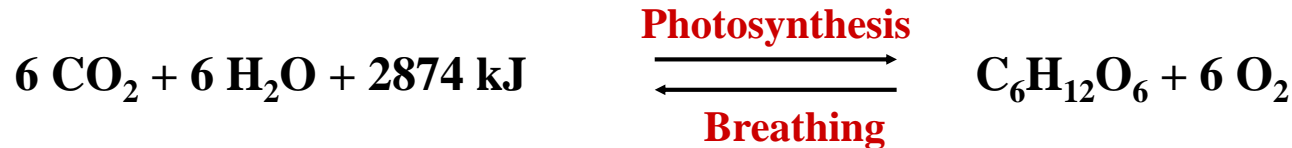


Depending on the chain length the following is discriminated: trioses ($x = 3$), tetroses ($x = 4$), pentoses ($x = 5$) and hexoses ($x = 6$), also called monosaccharides.

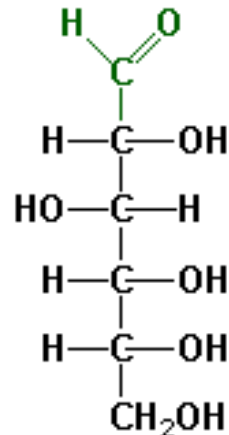


8. Properties of Biomolecules

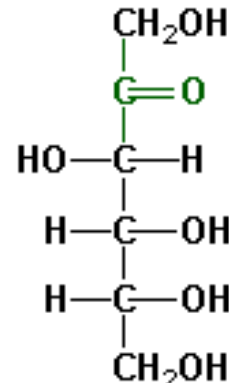
Carbohydrates: Synthesis and Properties



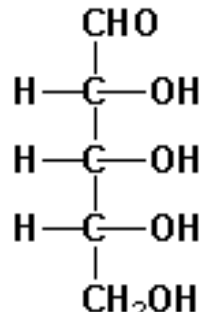
Carbohydrates are polyvalent alcohols, possessing an **aldehyde-** or **keto-group**



Glucose
(eine **Aldohexose**)



Fructose
(eine **Ketohexose**)

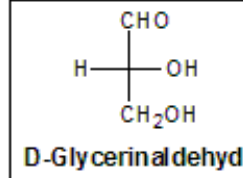


Ribose
(eine **Aldopentose**)

8. Properties of Biomolecules

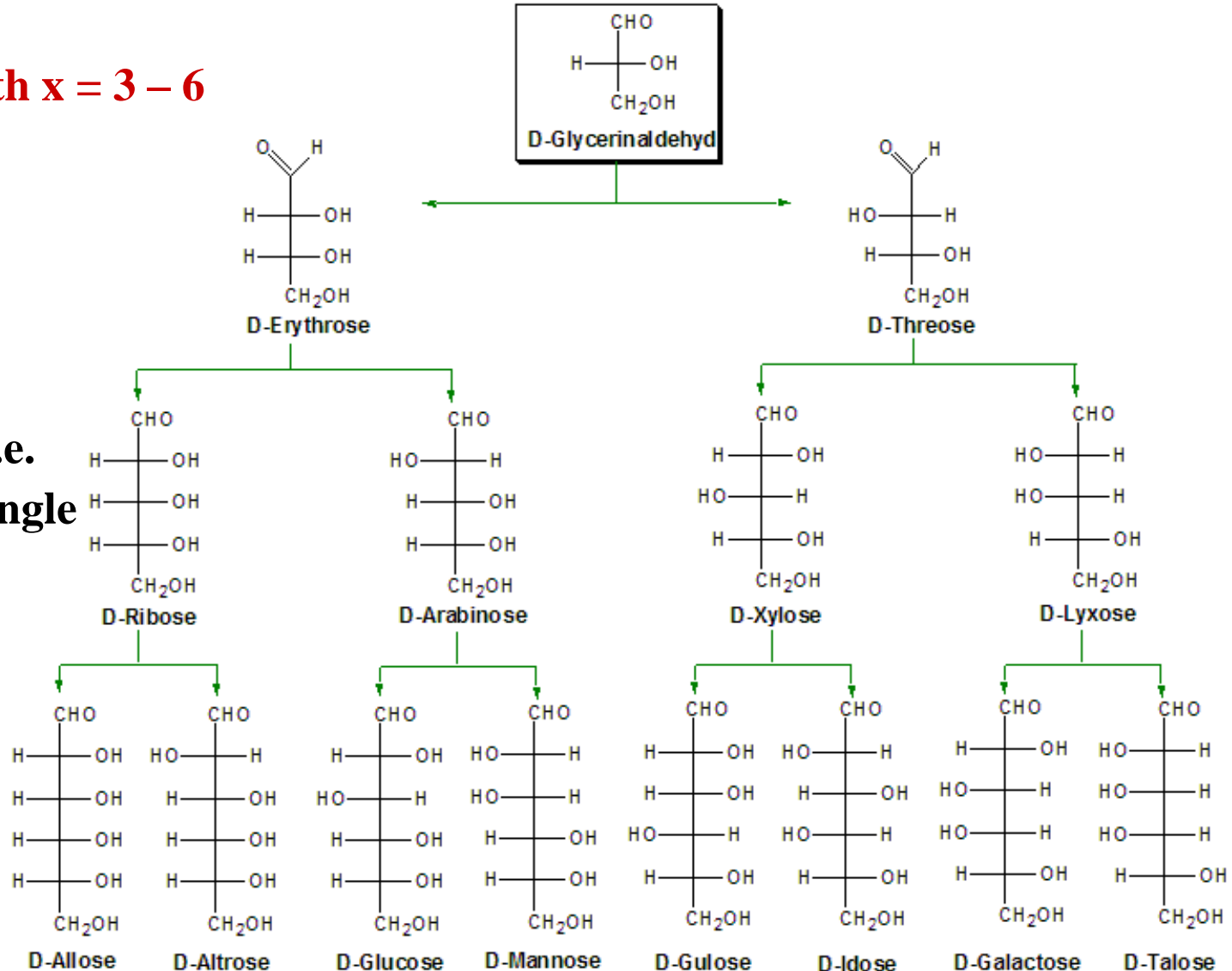
Carbohydrates:

Aldoses $C_x(H_2O)_n$ with $x = 3 - 6$



Optical properties

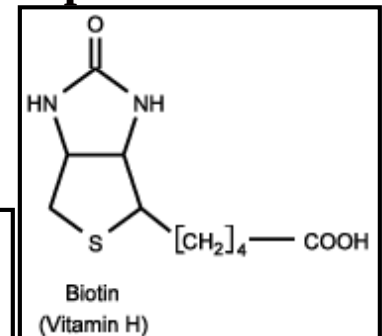
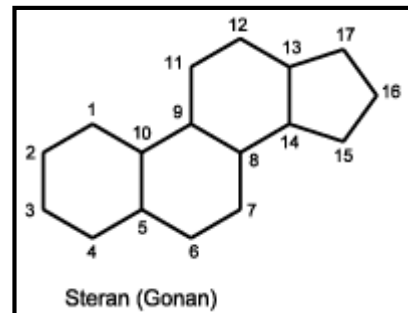
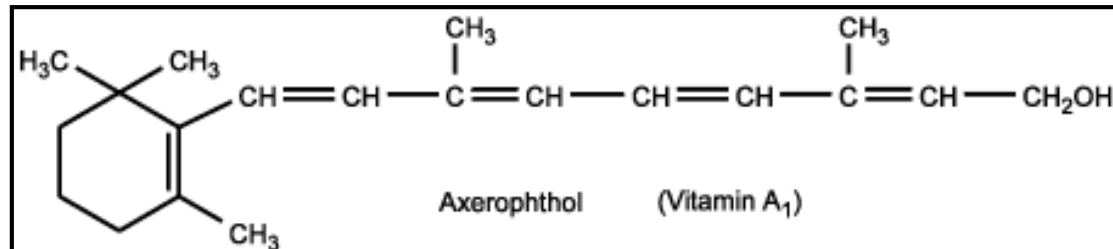
- Transparent till ca. 210 nm
- Optically active, i.e. change of phase angle of polarized light
- Exhibit circular-dichroism



8. Properties of Biomolecules

More Biomolecules: Overview

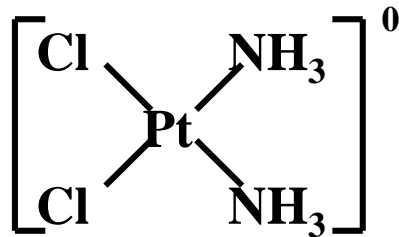
Class	Example	Function
Lipids	Fat	(Cell)membranes
Phospholipids	Lecithin	(Cell)membranes
Terpenes	Isoprene	Phytonutrients, vitamins, hormone, pigments
Steroids	Sterane	Vitamins, hormones
Heterocycles	Biotin	Vitamins, co-factors
Porphyrins	Heme	Vitamins, pigments, enzymes, transport proteins
Complexes	cis-platinum [Gd(dota)]	Therapy Diagnostics



8. Properties of Biomolecules

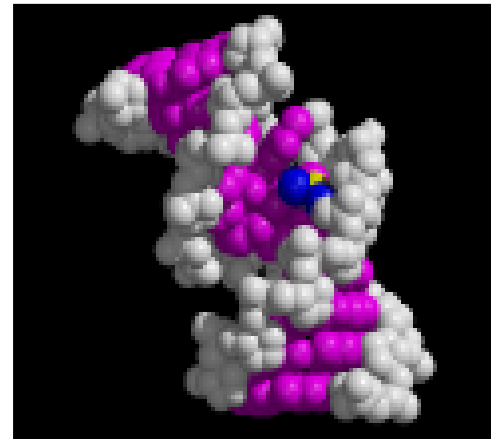
More Biomolecules: cis-Platinum

Cis-diamine-dichloro-platinum(II) (Peyrone's salt)



The square-planar cis-platinum disturbs the structure of DNA and leads to the dying of rapidly growing tumour cells

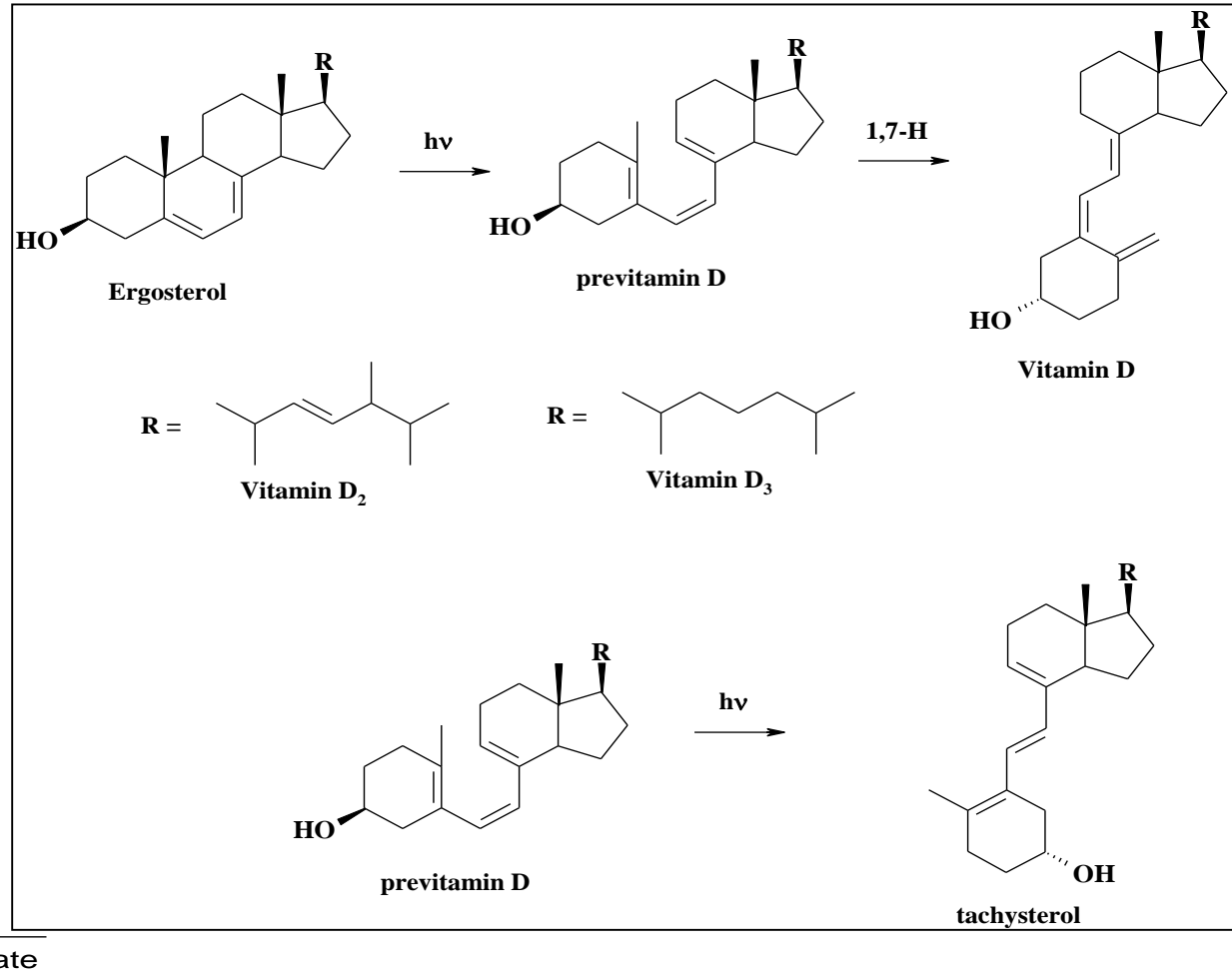
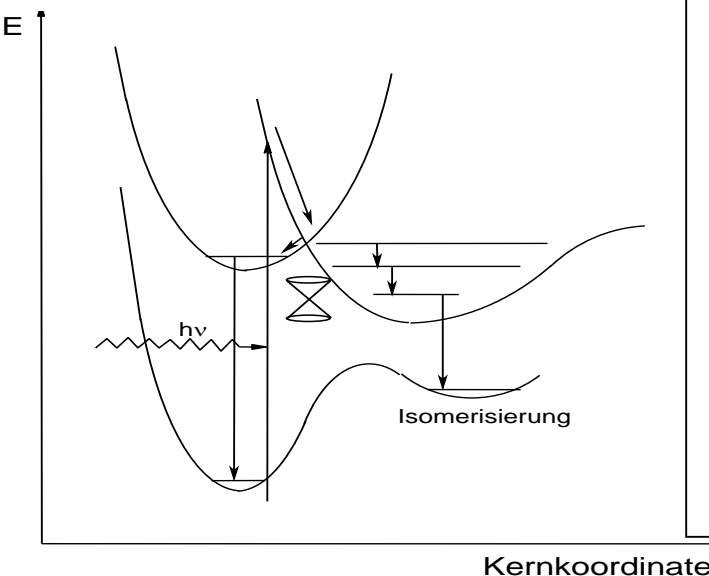
⇒ Chemotherapeutic agent for bronchial carcinoma and tumours in the genitourinary system



8. Properties of Biomolecules

More Biomolecules: Vitamin D

Formation by photo-
Isomerisation reaction
precalciferol (previtamin D)
+ $h\nu$ (282 nm)
→ calciferol (vitamin D)



8. Properties of Biomolecules

More Biomolecules: Luciferine (D-LH₂)

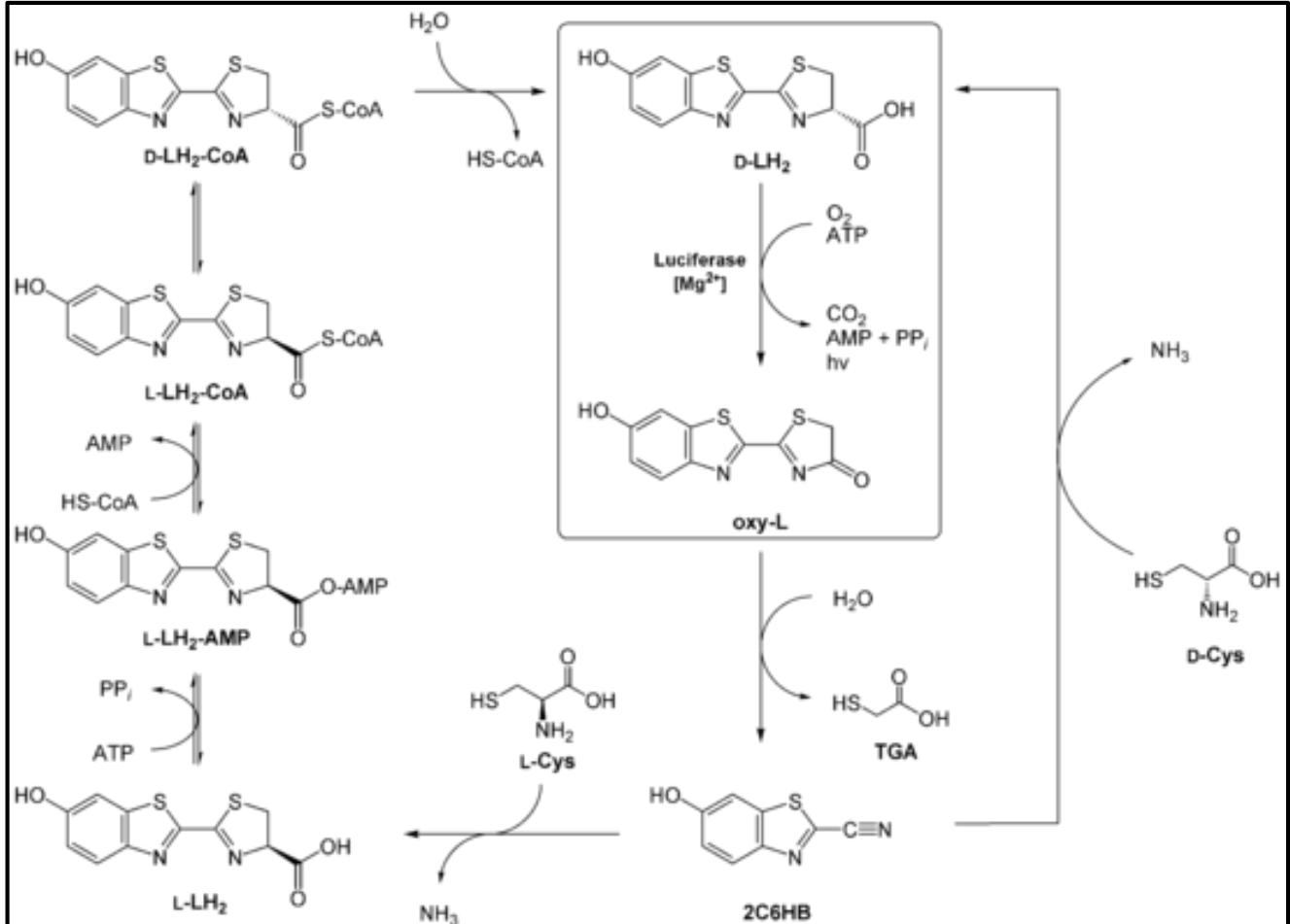
Oxidation is catalysed by Luciferase, which is Mg²⁺ dependent

Firefly luciferine is found in Lampyridae species

Benefits

- Attraction
- Communication
- Defence

Application: Luciferine assays for ATP analysis



9. Biochemistry of Main Group Elements

The Alkali Metal Cations

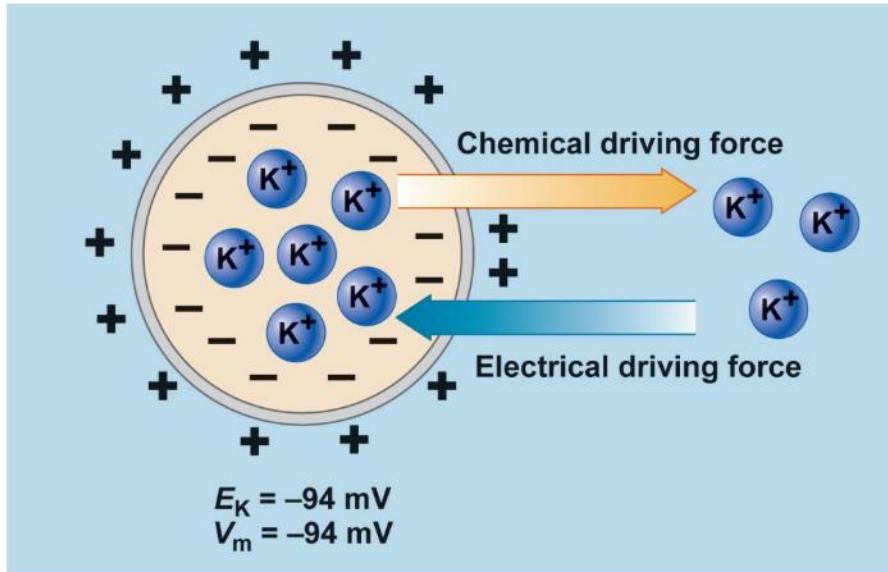
Ion	CN	Ionic radius [pm]	Geometry	Ligands	CFSE
Li⁺	4 - 8	73 – 106	variable	O	0
Na⁺	4 – 12	113 – 153	variable	O	0
K⁺	4 – 12	151 – 178	variable	O	0
Rb⁺	6 – 12	166 – 186	variable	O	0
Cs⁺	6 – 12	181 – 202	variable	O	0
Fr⁺	6 – 12	194 (CN = 6)	(radioactive)	O	0

9. Biochemistry of Main Group Elements

The Alkali Metal Cations

Functions

- Osmotic control
- Electrolytic equilibria
- Ionic current
- Control of ionic channels (“gating”)
- Structural stabilisation, e.g. of enzymes like pyruvate kinase



(a)

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Typical mammal cell: ~100 mV along 5 nm thick membrane \Rightarrow 200000 Vcm⁻¹

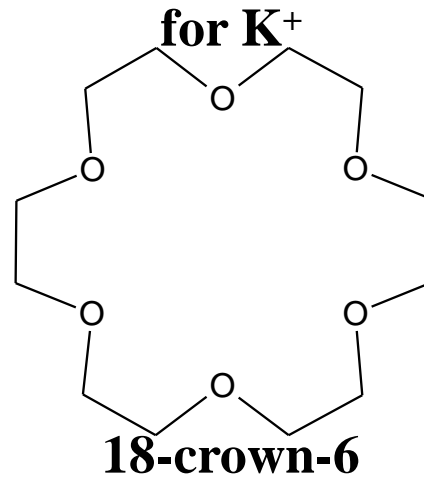
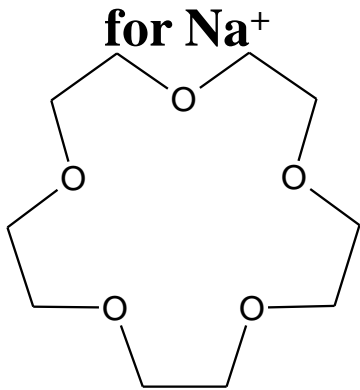
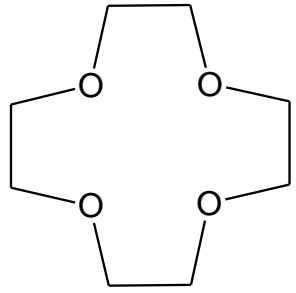
Ion	Extracellular [mM]	Intracellular [mM]	Membrane-potential [mV]
Na ⁺	150	12	+68
K ⁺	4	140	-99

9. Biochemistry of Main Group Elements

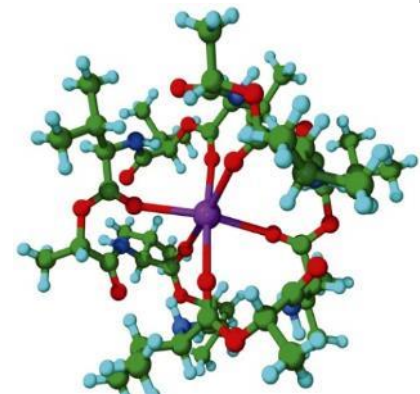
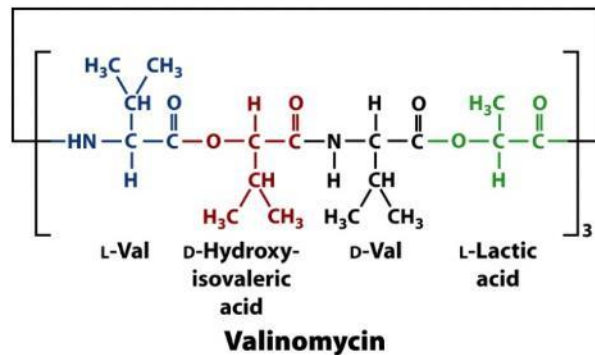
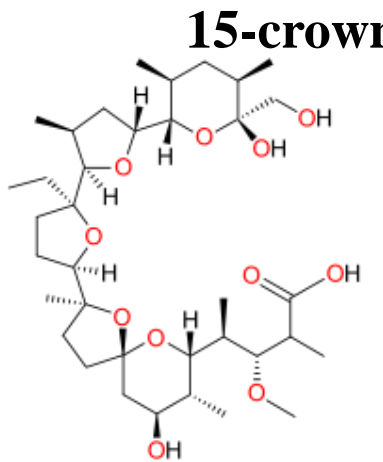
The Alkali Metal Cations

Transportation

a) Ionophors →



12-crown-4



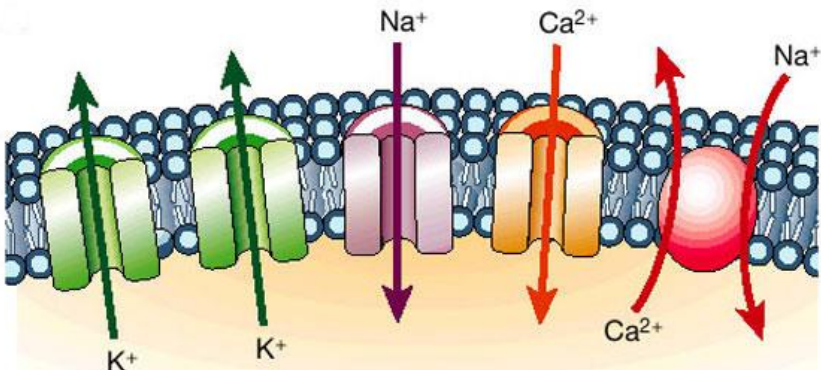
9. Biochemistry of Main Group Elements

The Alkali Metal Cations

Transportation

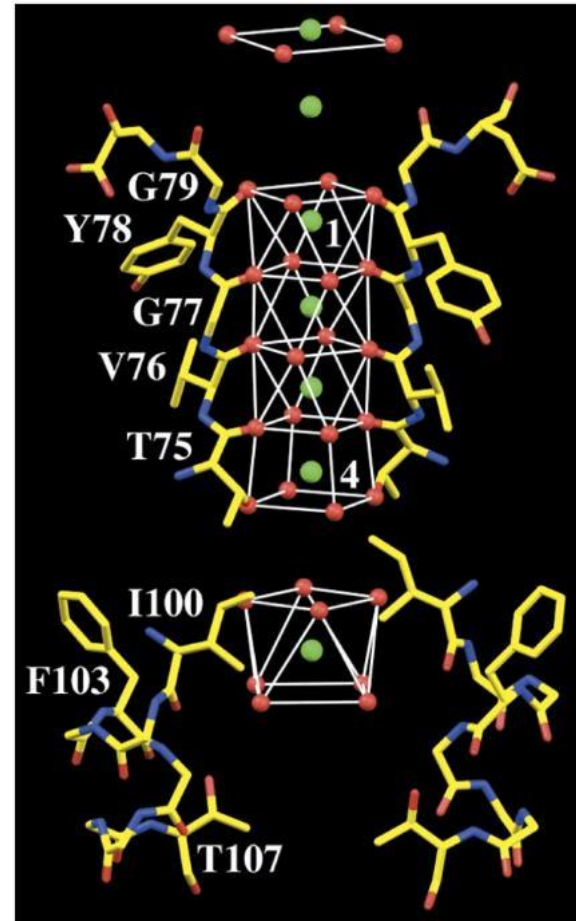
b) Ion channels

- Integral membrane proteins of high selectivity
- Form pores in membranes, which allow the transportation of ions along electrochemical potentials
- Can be opened or closed, e.g. through neurotransmitters (ligands or Ca^{2+})



Example: KcsA K^+ ion channel

10000fold selectivity for K^+ vs. Na^+ homotetramer, i.e. four identical protein units



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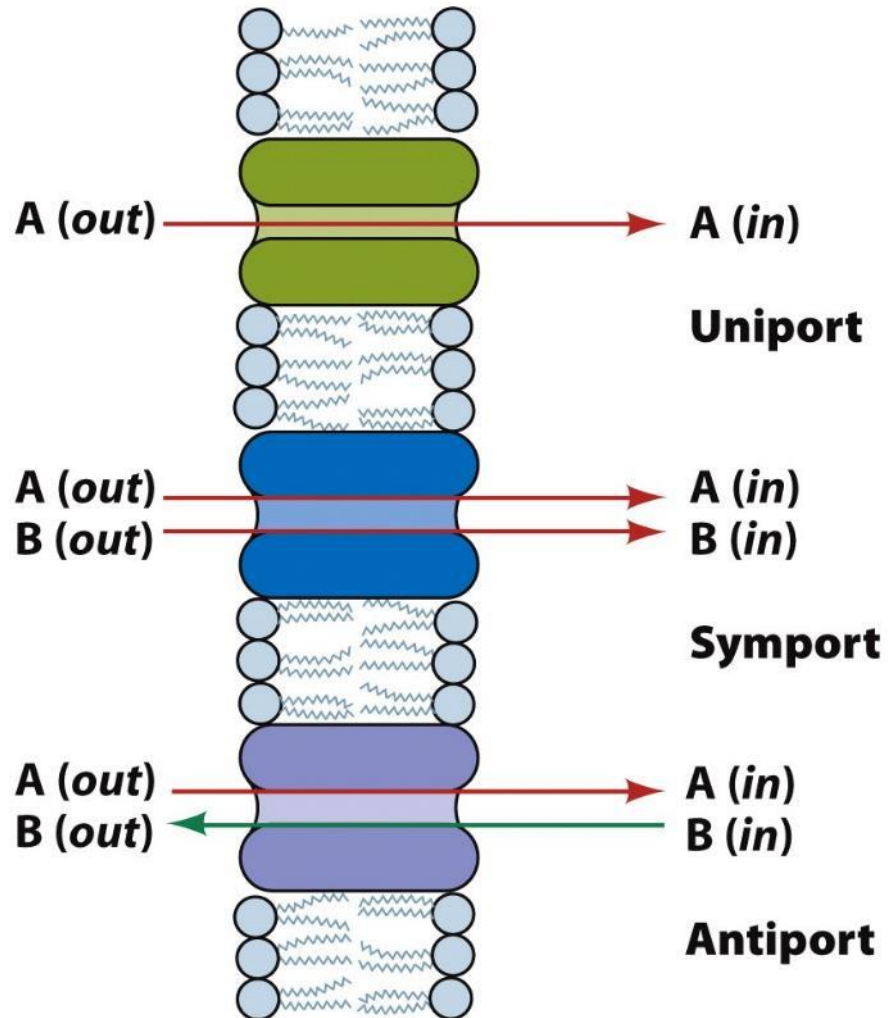
9. Biochemistry of Main Group Elements

The Alkali Metal Cations

Transportation

b) Ion channels

Uniport: $\text{Na}^+ \rightarrow$, Glucose \rightarrow



Symport: $\text{Na}^+/\text{Glucose} \rightarrow$

Antiport: $\text{Na}^+ \rightarrow / \leftarrow \text{K}^+$
 $\text{Na}^+ \rightarrow / \leftarrow \text{Ca}^{2+}$

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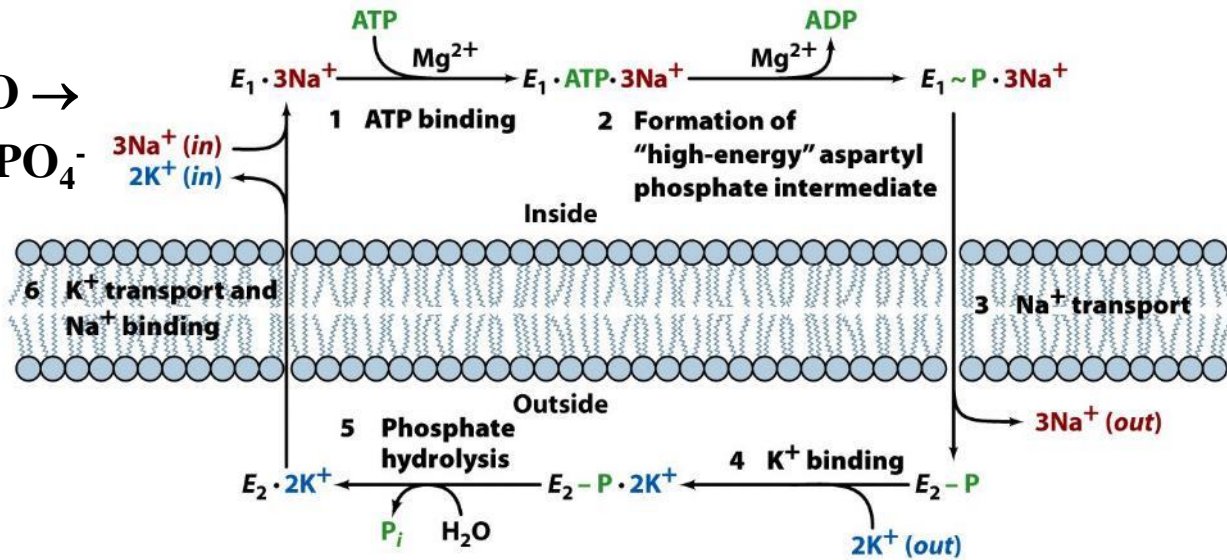
9. Biochemistry of Main Group Elements

The Alkali Metal Cations

Transportation

c) Ion pumps → Na^+/K^+ -pump (ATPase)

- Maintenance of resting potential
- Regulation of cellular volume
- Signal transduction and integration



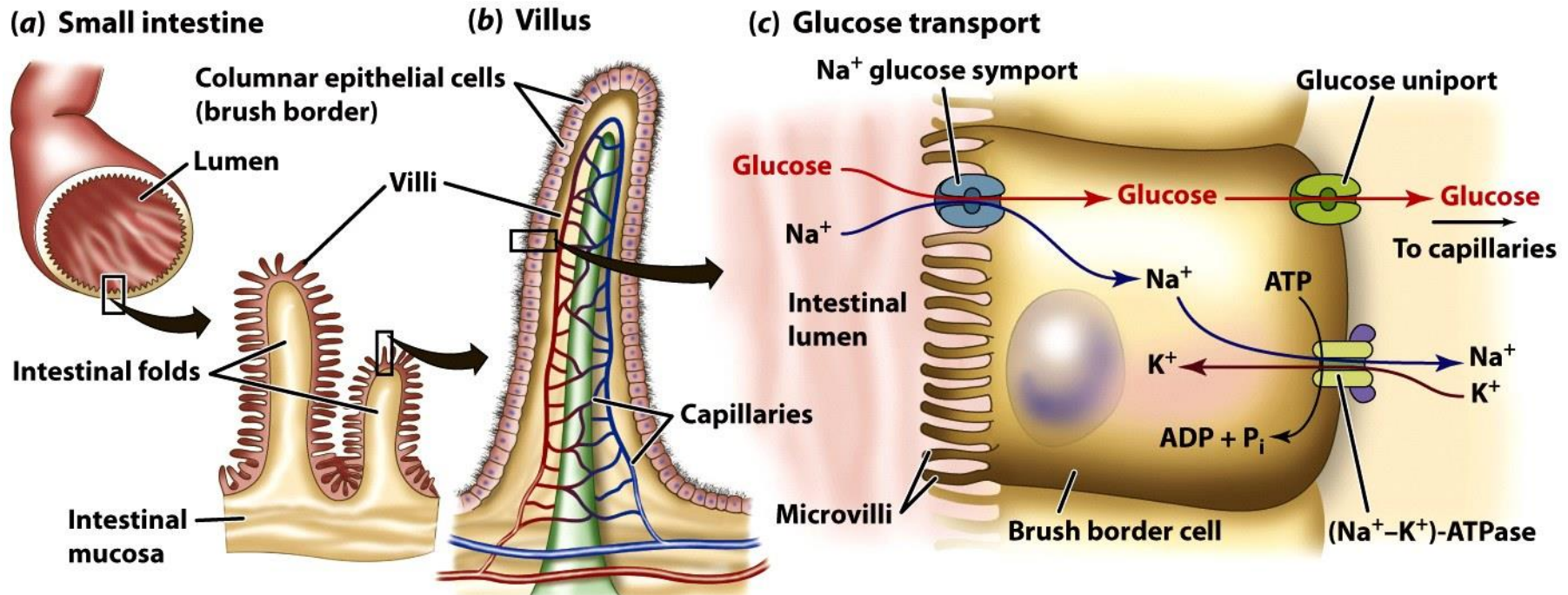
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9. Biochemistry of Main Group Elements

The Alkali Metal Cations

Transportation

c) Ion pumps → Na^+/K^+ -pump (ATPase) → uptake of glucose in small intestine



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9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Ion	CN	Ionic radius [pm]	Geometry	Ligands	CFSE
Be²⁺	3 - 6	30 - 59	variable	O	0
Mg²⁺	4 – 8	71 - 103	variable	N, O	0
Ca²⁺	6 – 12	114 - 148	variable	O	0
Sr²⁺	6 – 12	132 - 158	variable	O	0
Ba²⁺	6 – 12	149 - 175	variable	O	0
Ra²⁺	8 – 12	162 – 184	(radioactive)	O	0

9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Overview of functions

- | | |
|-----------|----------------------------------|
| Mg^{2+} | Phosphate metabolism |
| | Protein-/ nucleic acid structure |
| | Central atom in chlorophyll |
| Ca^{2+} | Muscle contraction |
| | Cellular signals |
| | Enzyme activation |
| | Blood coagulation |
| | Mineralisation (endoskeleton) |
| | Morphogenesis |
| | Genetic regulation |



Demineralised bones possess collagen,
wherein the crystals are embedded

<u>Ion</u>	<u>Extracellular [mM]</u>	<u>Intracellular [mM]</u>
Mg^{2+}	1.5	2.5
Ca^{2+}	2.5	0.1

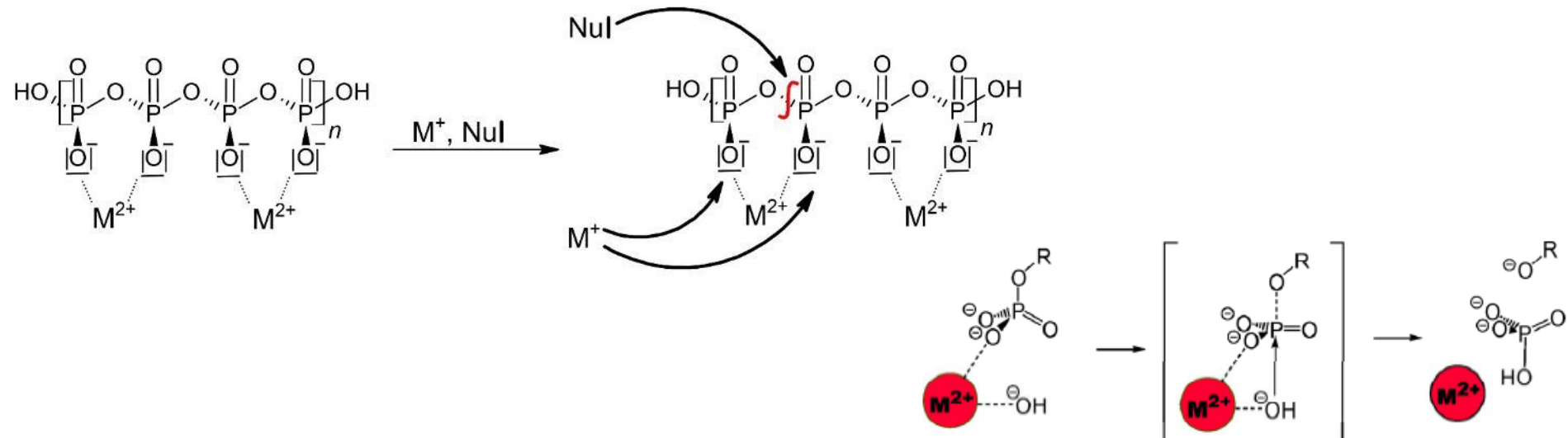
9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Mg^{2+} is very hard and shows high affinity to phosphate $\rightarrow Mg(NH_4)PO_4 / MgKPO_4$

Functions in detail

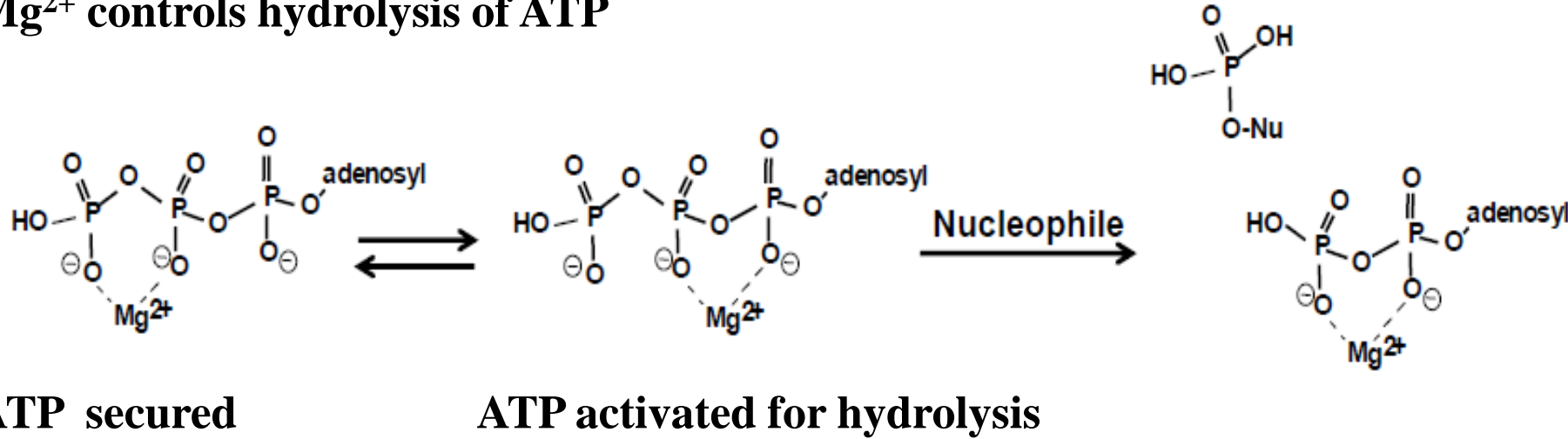
- Charge compensation, e.g. for ATP (reduction of negative charge density)
- Polarisation to enhance nucleophilic character: $Mg^{2+} + H_2O \rightarrow [Mg\cdots OH]^+ + H^+$
- Stereo chemical fixation of reactants during phosphate cleavage \rightarrow ATP/ADP
- Catalyst for polyphosphate decomposition?



9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Mg^{2+} controls hydrolysis of ATP

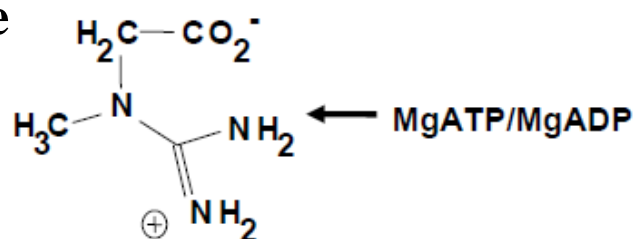


ATP secured

ATP activated for hydrolysis

Phosphate transfer to glyceride \rightarrow 2-phosphoglyceride

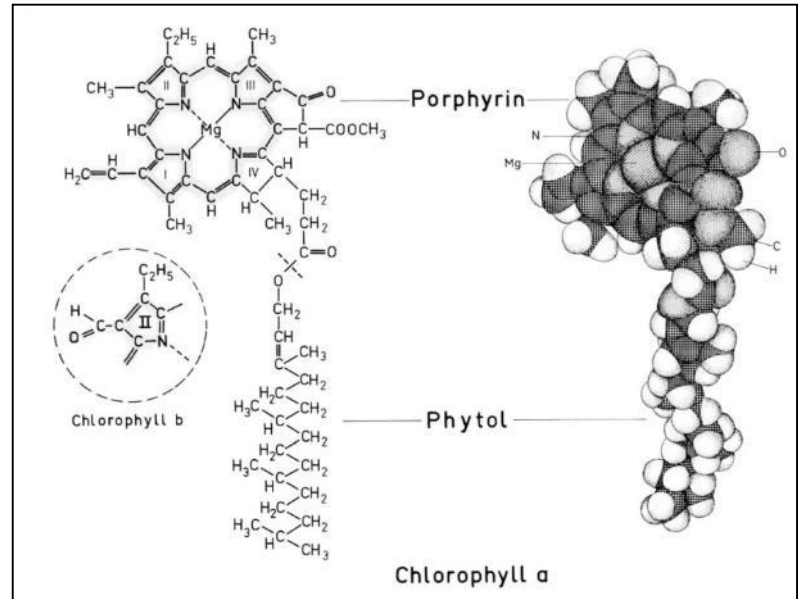
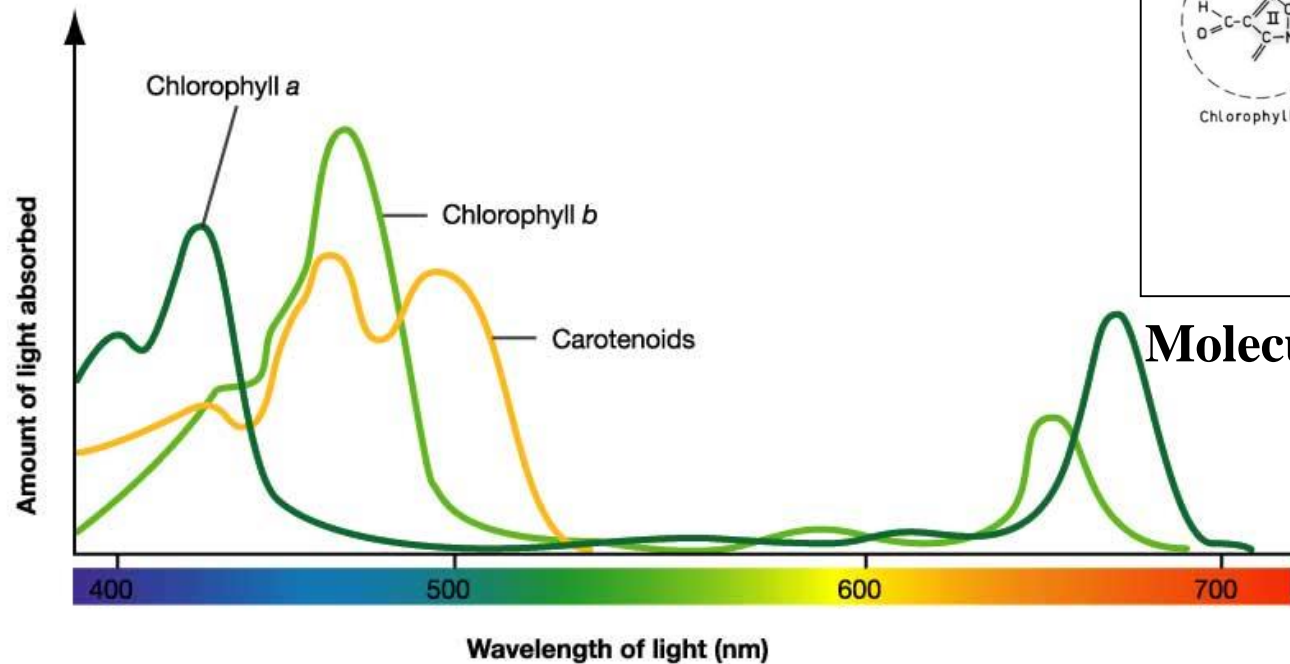
Phosphate transfer to creatine \rightarrow creatine phosphate



9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Mg^{2+} is the metal centre complexed by porphyrin ligands in chlorophyll a/b



Molecular structure of chlorophyll

Absorption spectrum of the most important pigments in photosynthesis

9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Fluorescence

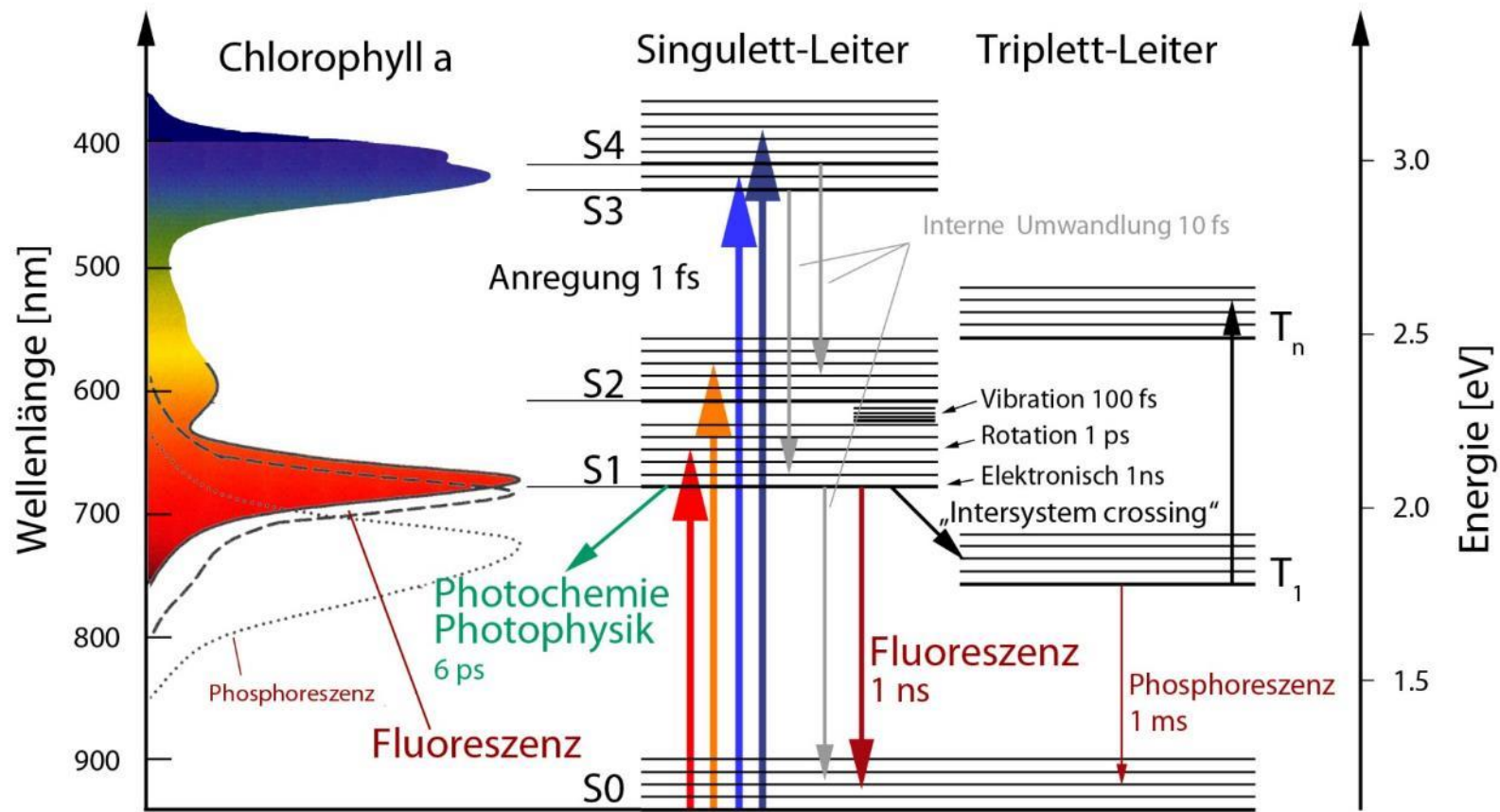
Intact leafs

~ 3 - 7%

i.e. ET to
reactive
center

Isolated
chlorophyll

~ 30%



9. Biochemistry of Main Group Elements

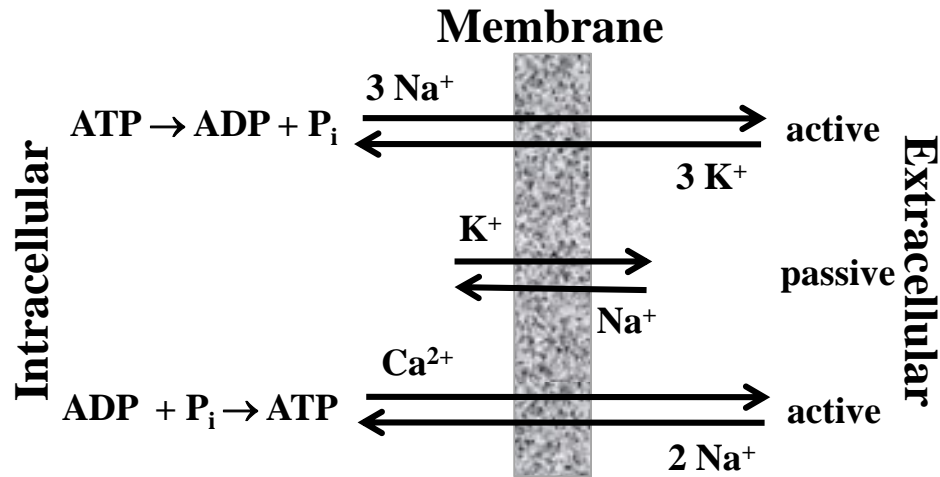
The Alkaline Earth Metal Cations

Ca^{2+} exhibits a broad variety of functions

- Structural functions: protein folding and build-up of skeleton
 - Exoskeleton
 - Mollusca (mussels, snails)
 - Cnidaria (corals)
 - Arthropods (insects, spiders, scorpions, crayfish...)
 - Endoskeletons
 - Vertebrates (bones and teeth)
 - Cephalopods (cuttlebones)
- Trigger and activation functions
 - Bonding to μ_2 -carboxylates of proteins
 - Labile complexes allow fast changes of structure (muscle contraction)
- Electrolyte transportation



Cuttlebones of *Sepia officinalis*,
(Source: Wikipedia)

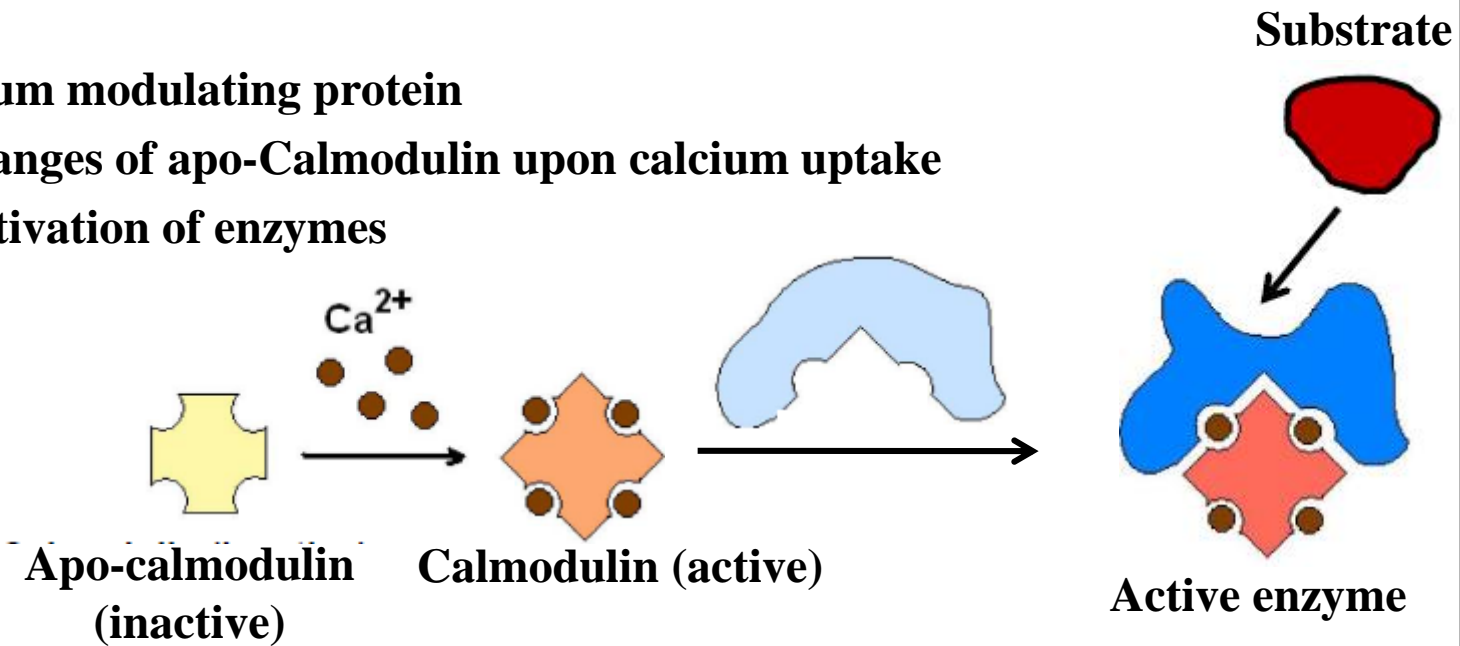


9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Activation

- Calmodulin = calcium modulating protein
- Conformational changes of apo-Calmodulin upon calcium uptake
- Recognition and activation of enzymes



Muscle contraction

- Depolarisation of cell membrane by opening of Na-ion channels
- Liberation of Ca^{2+} from acidic storage protein: calsequestrin
- Calsequestrin contains up to 50 Ca^{2+} -binding sites, i.e. carboxylate groups: Glu, Asp
- Uptake of Ca^{2+} by troponin C, which is coupled with the ATP-hydrolyses

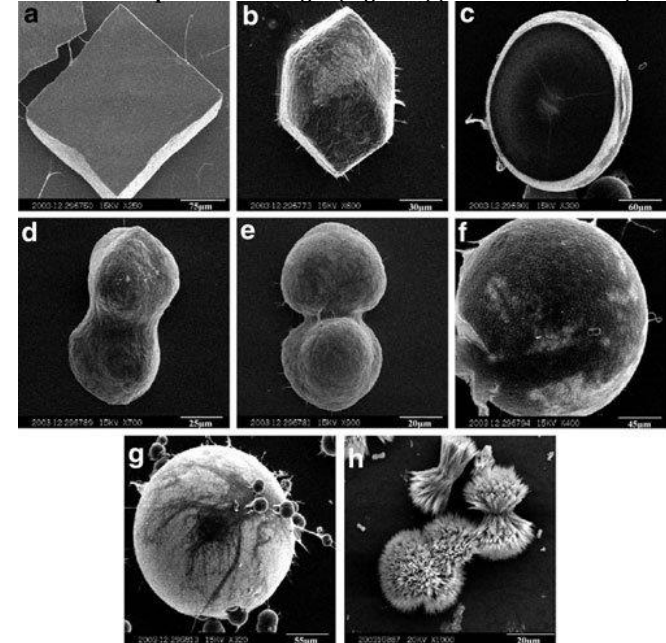
9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Biomineralisation (CaCO_3)

- In mussels, snails, otoliths, ...
- Morphological control and orientation by organic ligands
 - Carboxyl groups (glu, asp, ...)
 - Oxidised carbohydrates
 - Collagene
- Lab examples
 - Spindle-shaped calcite crystals in presence of malonic acid
 - Disc-shaped vaterite crystals through stearic acid

SEM morphologies of CaCO_3 crystals precipitated in the solution, collagen concentration: 0.1 g/l. (a) Irregular rhombohedral calcite crystal grown in the solution without magnesium. (b) Irregular lumpish crystals with lamellar growth structure ($\text{Mg}/\text{Ca}:1$). (c, d, e) Discoid and dumbbell calcium carbonate crystals. (f) Spherical aragonite crystals at higher Mg^{2+} concentration ($\text{Mg}/\text{Ca}:5$). (g) Spherical aragonite crystals with more regular shape ($\text{Mg}/\text{Ca}:5$, collagen concentration: 0.4 g/l). (h) Aragonite crystals with needlelike shape without collagen ($\text{Mg}/\text{Ca}:5$) (Lit.: Jiao et al. 2006)

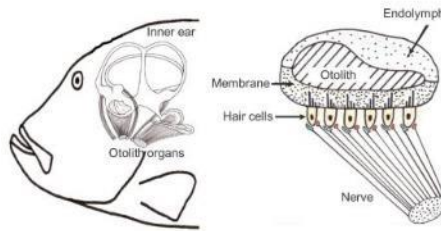


Modification	CaCO_3 (aragonite)	CaCO_3 (calcite)	CaCO_3 (vaterite)
Crystal system	Orthorhombic	Trigonal	Hexagonal
Space group	Pnma (#62)	R-3ch (#167)	P63/mmc (#194)
Coordination number	9	6	8
Formula unit/unit cell	4	6	2

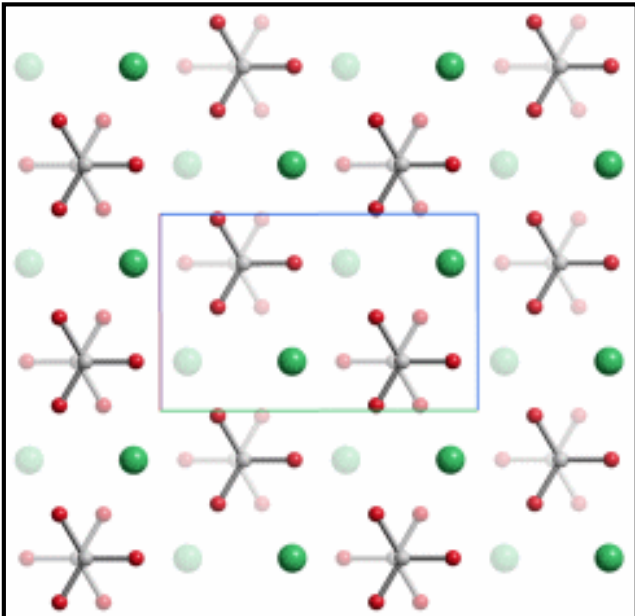
9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Biomineralisation of CaCO_3

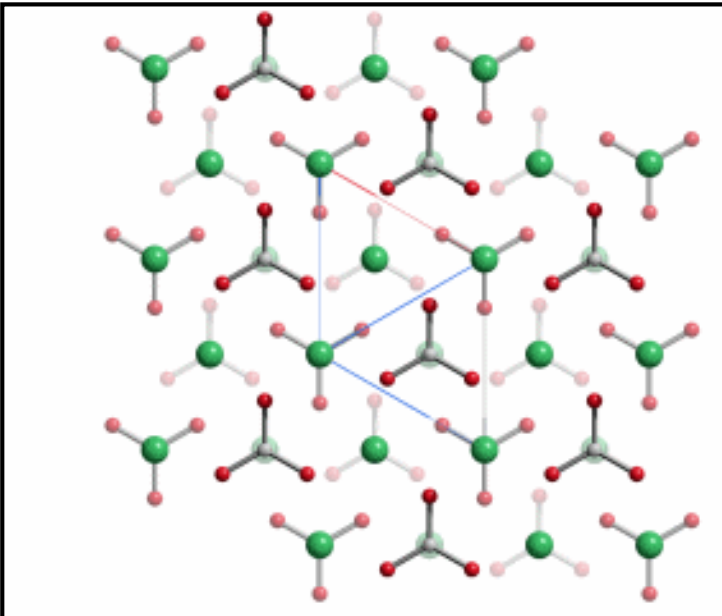


Aragonite (meta-stable)



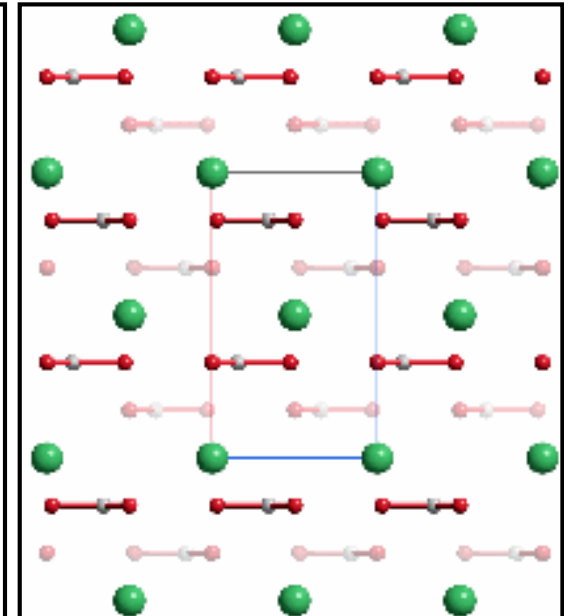
Fish-otholiths, corals, pearls

Calcite (stable)



Mussel shells

Vaterite (meta-stable)



Spheroliths

9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Biomineralisation of $\text{Ca}_5[\text{PO}_4]_3\text{X}$ with $\text{X} = \text{OH}$

- In bones of vertebra and cephalopods
- Collagen serves as template, i.e. it defines the orientation of the apatite crystals
- Binding of Ca^{2+} via carboxylates groups of osteocalcin and via phosphoproteins

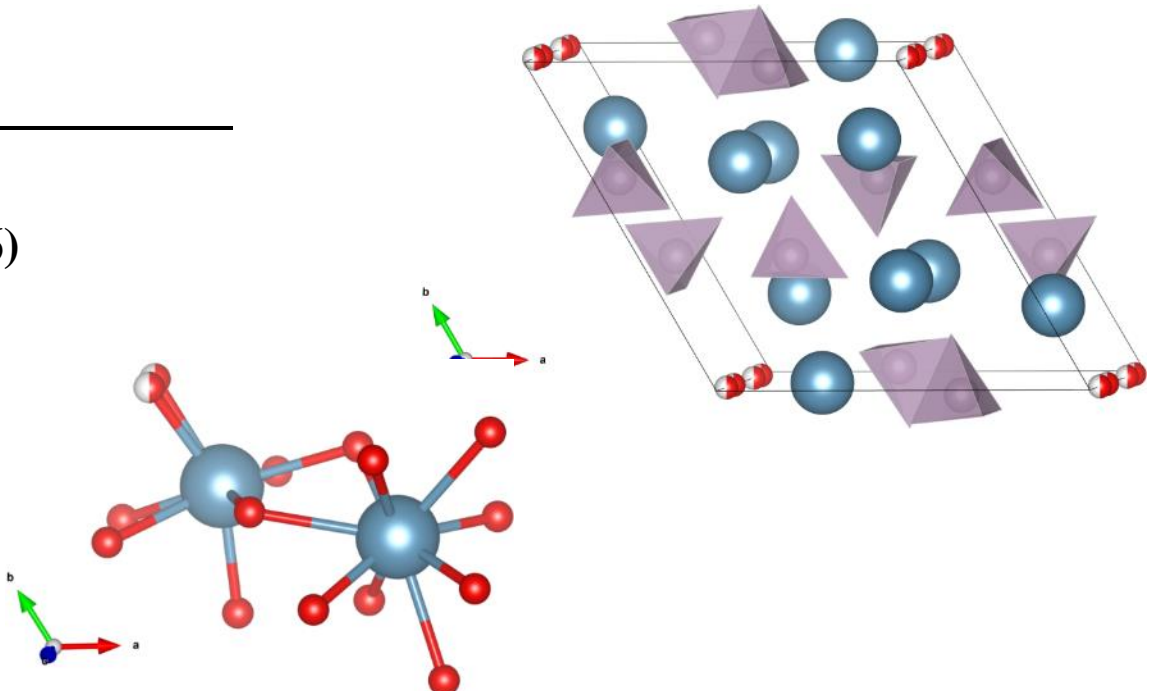
Hydroxyapatite $\text{Ca}_5[\text{PO}_4]_3\text{OH}$

Crystal system Trigonal

Space group Pnma (#176)

Coordination number 8 and 9

Formula unit per unit cell 2

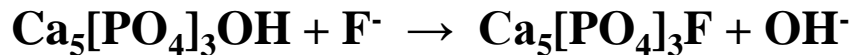


9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

Biomineralisation of $\text{Ca}_5[\text{PO}_4]_3\text{X}$ with $\text{X} = \text{F}$

- In teeth (enamel) of vertebrates and cephalopods
- Is formed by fluorination of hydroxyapatite (toothpaste contains $\text{Na}_2\text{PO}_3\text{F}$)

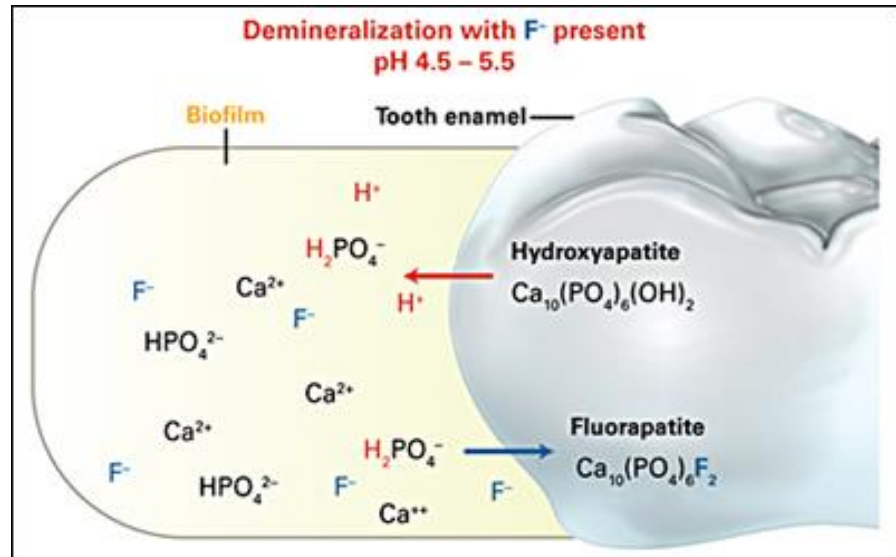


Fluoroapatite $\text{Ca}_5[\text{PO}_4]_3\text{F}$

Crystal system	Trigonal
Space group	Pnma (#176)
Coordination number	7 and 9
Formula unit per unit cell	2

Pyrophosphates $\alpha\text{-Ca}_2\text{P}_2\text{O}_7$ and $\beta\text{-Ca}_2\text{P}_2\text{O}_7$

- Crystal-induced arthropathy

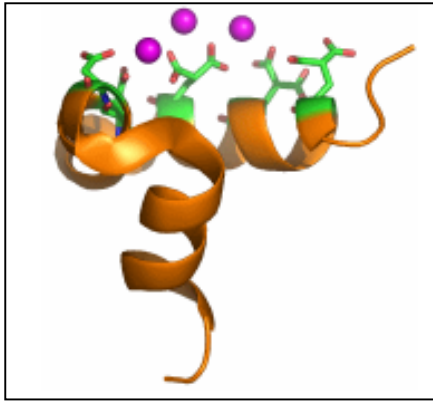


9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

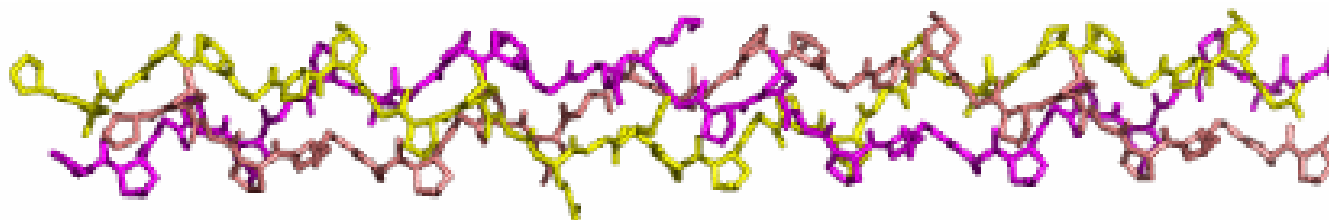
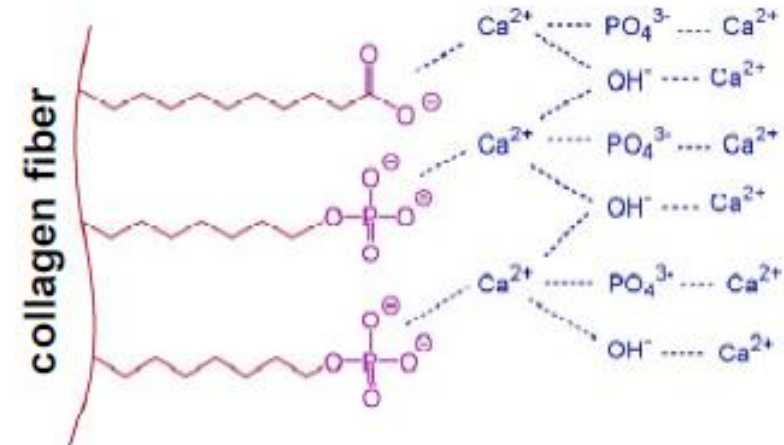
Osteocalcin

- Fixates apatite



Collagen

- Three left-handed helices, combined to a right-handed super-helix
- Composite material without binding sites for Ca^{2+}
- Apatite crystals are incorporated parallel to the collagen helix



9. Biochemistry of Main Group Elements

The Alkaline Earth Metal Cations

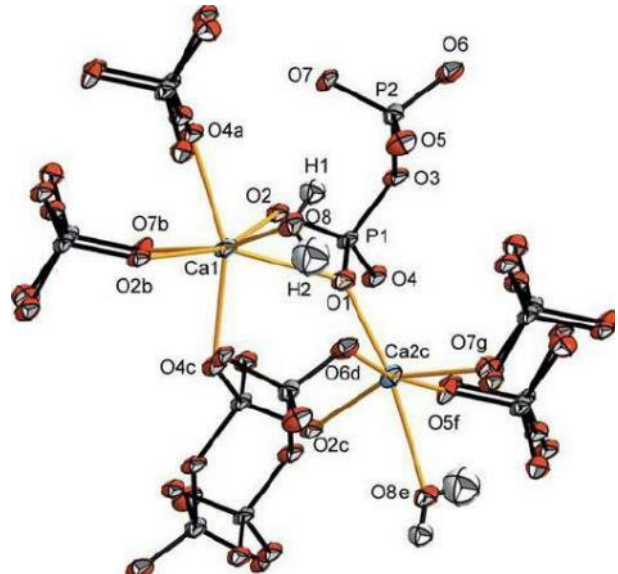
Calciumpyrophosphates

Formula	$\text{Ca}_2\text{P}_2\text{O}_7 \cdot 2\text{H}_2\text{O}$ (CPPD)	$\text{Ca}_2\text{P}_2\text{O}_7 \cdot \text{H}_2\text{O}$
Crystal system	triclinic	monoclinic
Space group	P1	P2 ₁ /n
Z	2	4
CN		7 (Ca1), 6 (Ca2)

Causes “Crowned Dens Syndrome”: Severe neck pain

Literature

- Acta Cryst. B31 (1975) 1730
- J. Bone Joint Surg. Am. 89 (2007) 2732
- Acta Cryst. C70 (2014) 862



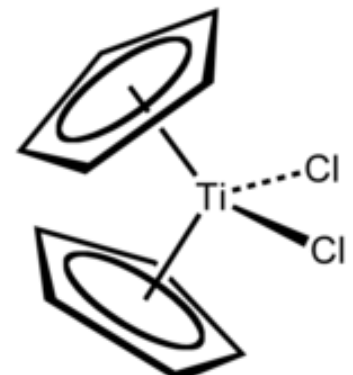
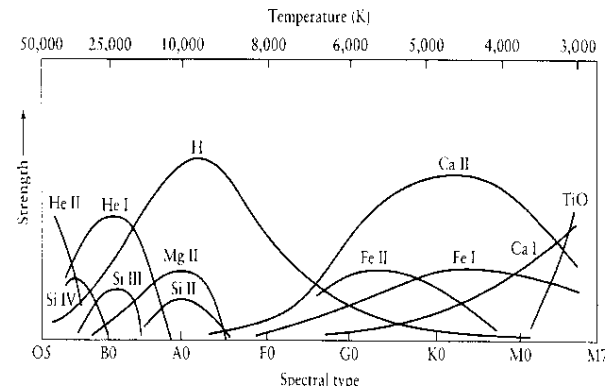
10. Biochemistry of Transition Metals

Titanium Group

Titanium



Ti^{4+} [Ar] **TiO_2 is poorly soluble and non-toxic even if orally ingested**
 $[\text{Ti}(\text{cp})_2\text{Cl}_2]$ (titanocene dichloride) and $\text{cis},\text{cis},\text{cis}-[\text{Ti}(\text{bzac})_2(\text{OEt}_2)]$ (bidotitanium) are used as cancerostatica (with bzac = 1-phenylbutane-1,3-dion)
Forms $[\text{TiO}]^{2+}$ unit



Zirconium

Zr^{4+} [Kr] **ZrO_2 , ZrSiO_4 and zirconate (MZrO_3) are poorly soluble**

Hafnium

Hf^{4+} [Xe] **Hafnium is scarce in the earth's crust and HfO_2 is also poorly soluble, which is why hafnium is of no importance for the biosphere**

10. Biochemistry of Transition Metals

The Vanadium Group

Vanadium

V^{2+}	$[Ar]3d^3$	$[V(H_2O)_6]^{2+}$	violet
V^{3+}	$[Ar]3d^2$	$[V(H_2O)_6]^{3+}$	green
V^{4+}	$[Ar]3d^1$	$[VO(H_2O)_4]^{2+}$	blue
V^{5+}	$[Ar]$	VO_4^{3-} or $[VO_2]^+$	colourless to pale yellow



Niobium

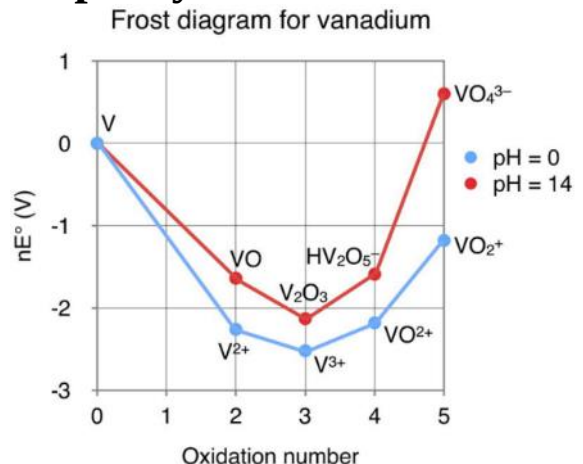
Nb^{5+}	$[Kr]$
-----------	--------

Nb_2O_5 and niobates ($MNbO_4$) are poorly soluble

Tantalum

Ta^0	$[Xe]5d^5$
--------	------------

metallic tantalum is high-melting and chemically inert permanent implants such as bone nails, springs or plates
 Ta_2O_5 and niobate ($MNbO_4$) are poorly soluble

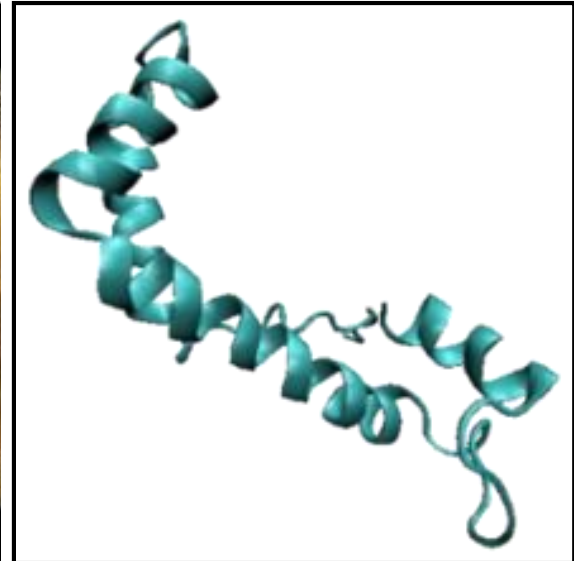
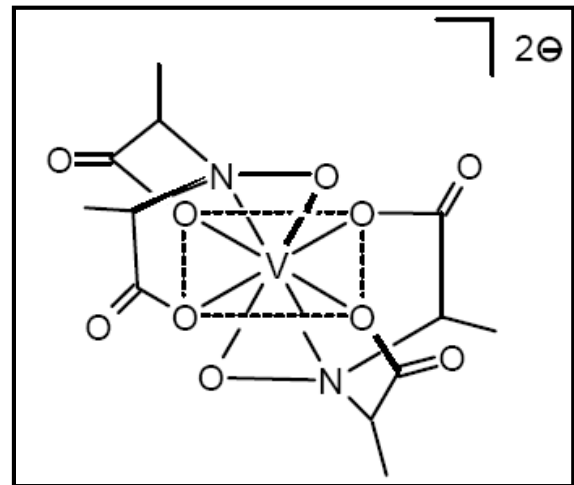
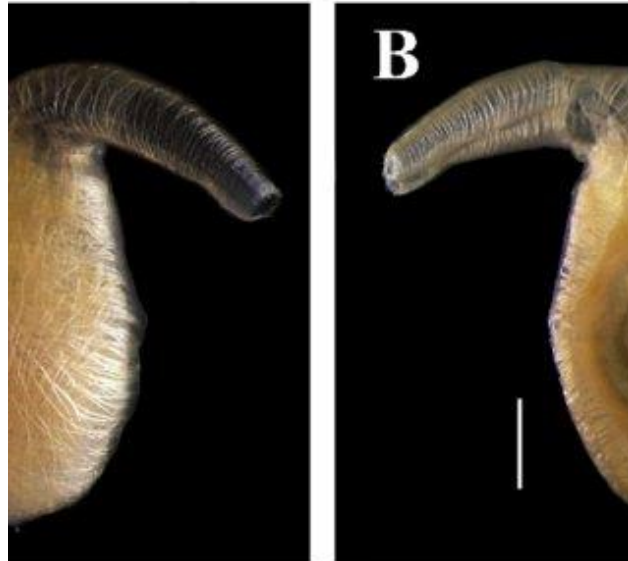


10. Biochemistry of Transition Metals

The Vanadium Group

Enrichment

- **Amavadin**
In fly agarics (*amanita muscaria*), V^{4+} is accumulated in the bottom of the toadstool
- Storage protein vanabin is responsible for the enrichment of VO^{2+} in vanadocytes of sea squirts (*ascidia gemmata*)
→ accumulation

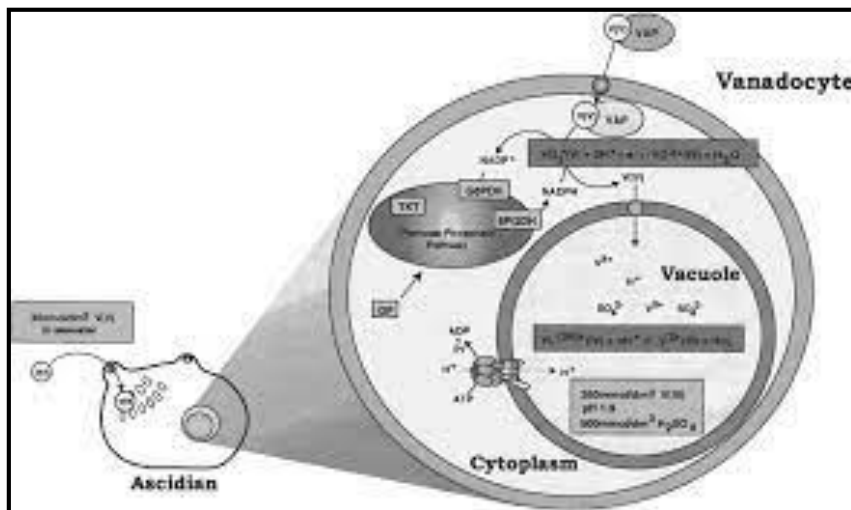


10. Biochemistry of Transition Metals

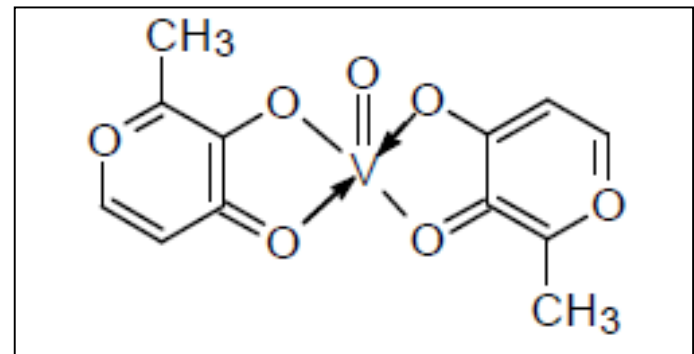
The Vanadium Group

Oxygen control?

- Tunicates comprise vanadocytes, in which V^{3+} is enriched for oxygen storage/transport?
Today: $\sim 35 \text{ nmol/l } VO_4^{3-}$ in seawater



- Insulin mimetics
 VOSO_4
 $[\text{VO(acac)}_2]$
Bis(maltolato)oxovanadium

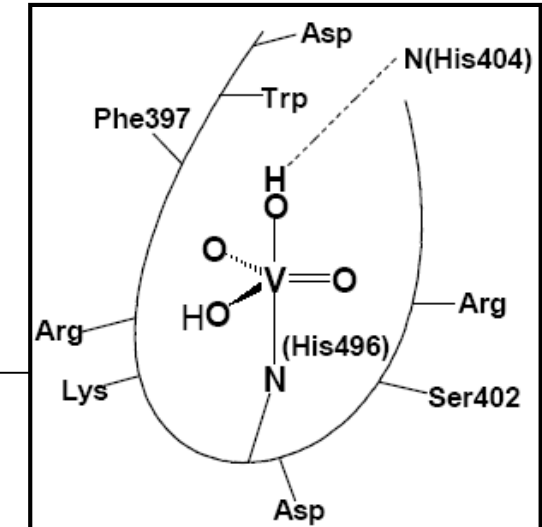
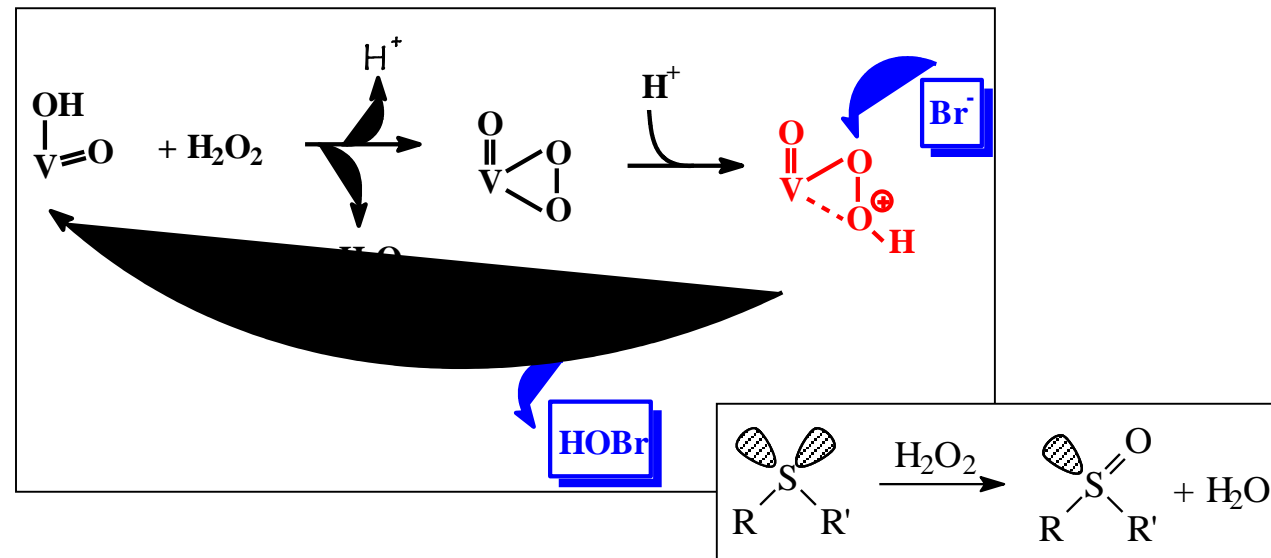
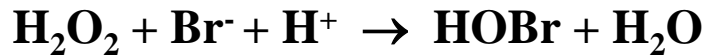


10. Biochemistry of Transition Metals

The Vanadium Group

Metal enzymes

- Haloperoxidases (e.g. in knotted wrack, *ascophyllum nodosum*) contain in their activated form vanadin(V)-ions in trigonal-bipyramidal coordination

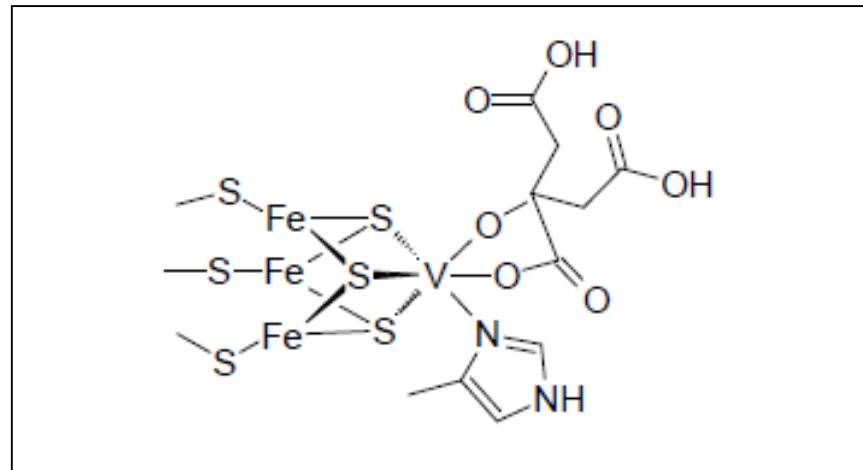


10. Biochemistry of Transition Metals

The Vanadium Group

Metal enzymes

- Nitrogenases
 - e.g. from *azotobacter chroococcum* and *azotobacter vinelandii*
 - some mutants contain Fe-V-cluster
 - Mo can be replaced by V
 - diagonal relationship V/Mo



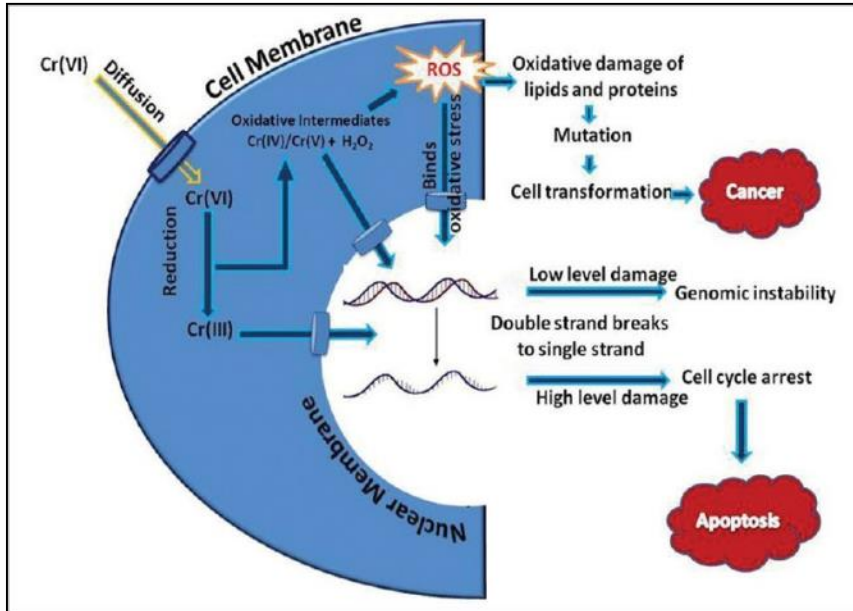
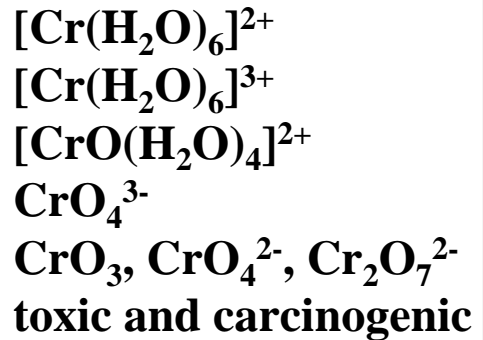
- $\text{N}_2 + 14 \text{ H}^+ + 12 \text{ e}^- + 40 \text{ MgATP} \rightarrow 2 \text{ NH}_4^+ + 3 \text{ H}_2 + 40 \text{ MgADP} + 40 \text{ HPO}_4^{2-}$
- Model complexes for fixation of nitrogen
 - $[\text{V}(\text{N}(\text{CH}_2\text{CH}_2\text{S})_3)] + \text{N}_2 + 4 \text{ H}^+ + 4 \text{ e}^- \rightarrow [\text{V}(\text{N}(\text{CH}_2\text{CH}_2\text{S})_3(\text{N}_2\text{H}_4)]$
 - $[\text{V}(\text{O}(\text{CH}_2\text{CH}_2\text{S})_2]$

10. Biochemistry of Transition Metals

The Chromium Group

Chromium

Cr^{2+}	$[\text{Ar}]3\text{d}^4$
Cr^{3+}	$[\text{Ar}]3\text{d}^3$
Cr^{4+}	$[\text{Ar}]3\text{d}^2$
Cr^{5+}	$[\text{Ar}]3\text{d}^1$
Cr^{6+}	$[\text{Ar}]$



Molybdenum

Mo^{4+}	$[\text{Kr}]4\text{d}^2$
Mo^{6+}	$[\text{Kr}]$

involved in 2-electron reductions (nitrate reductases)
 MoO_4^{2-} and $\text{Mo}_7\text{O}_{24}^{6-}$

Tungsten

W^{4+}	$[\text{Xe}]5\text{d}^2$
W^{5+}	$[\text{Xe}]5\text{d}^1$
W^{6+}	$[\text{Xe}]$

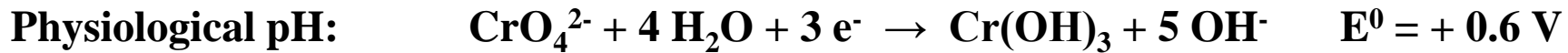
WO_4^{4-}
 WO_4^{3-}
 WO_4^{2-}
tungsten blue
tungstates are poorly soluble

10. Biochemistry of Transition Metals

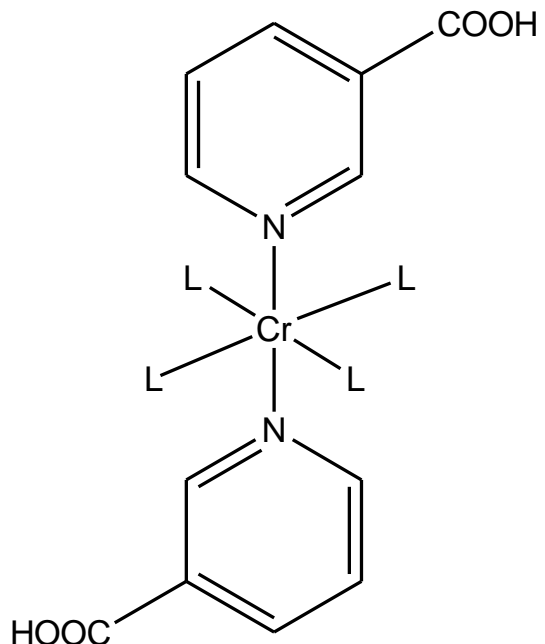
The Chromium Group

Chromium

- Chromium(VI) is carcinogenic, because it can oxidize the OH-groups of deoxyribose of DNA and also proteins



- Chromium(III) regulates the blood sugar level glucose-tolerance factor together with insulin and glucagon
- Cr^{3+} is transported by transferrin (Fe-transporter)
- Chromium deprivation may foster high blood pressure



10. Biochemistry of Transition Metals

The Chromium Group

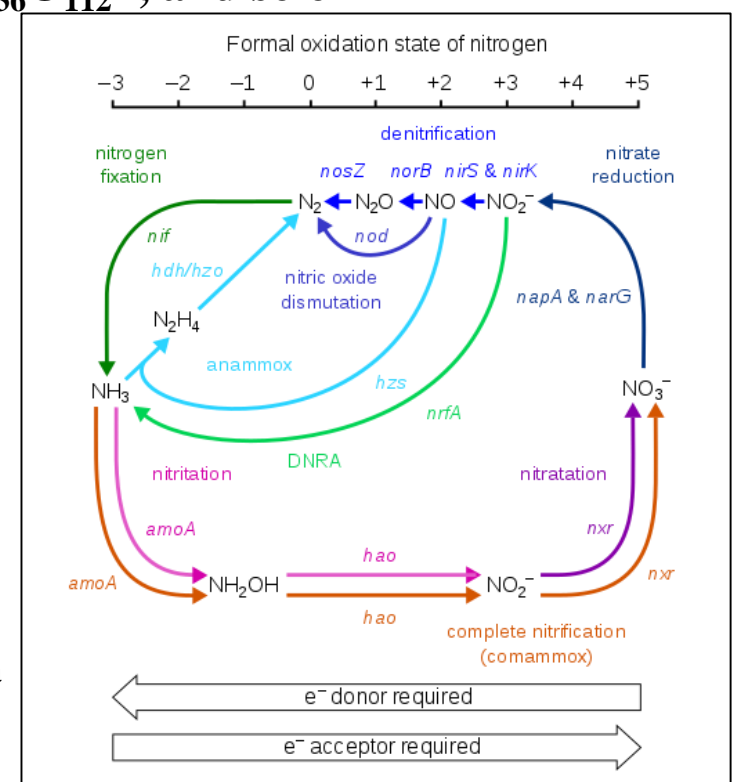
Molybdenum

- Only element of 2nd transition metal series of biological importance
- As molybdate MoO_4^{2-} readily soluble and available through sea water
- Forms polyoxymolybdates → $\text{Mo}_7\text{O}_{24}^{6-}$, $\text{Mo}_8\text{O}_{26}^{4-}$, $\text{Mo}_{36}\text{O}_{112}^{8-}$, and so on
- Biochemically relevant oxidation states: IV, V, VI
⇒ 1- or 2-electron-transfer-reactions
- Take part in nitrogen fixation
- Coordination by O-, S- and N-ligands
- Relevant enzymes
 - Nitrogenases
 - Nitrate reductases
 - Aldehyde oxidases
 - Oxytransferases

Azotobacter



Sources:
Wikipedia

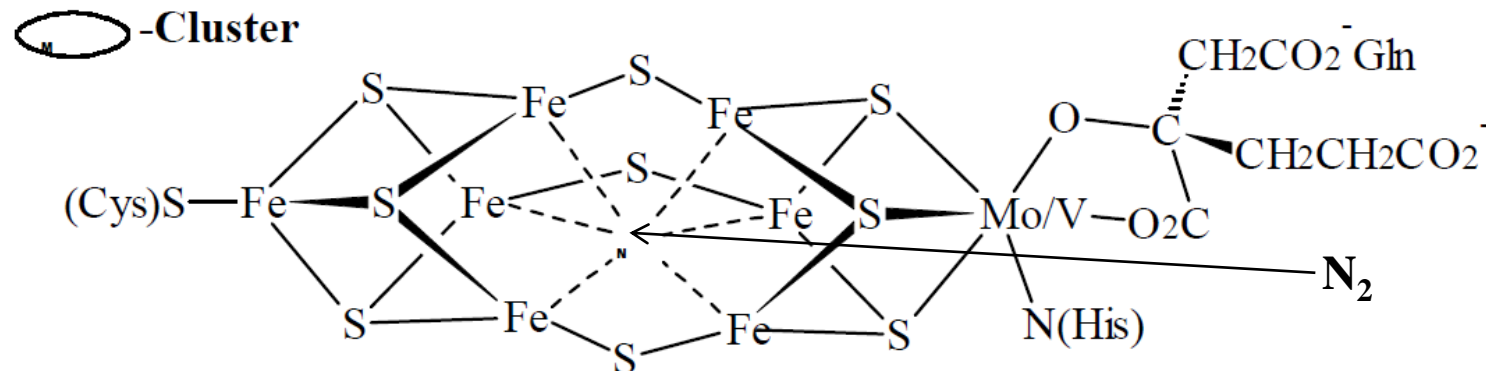


10. Biochemistry of Transition Metals

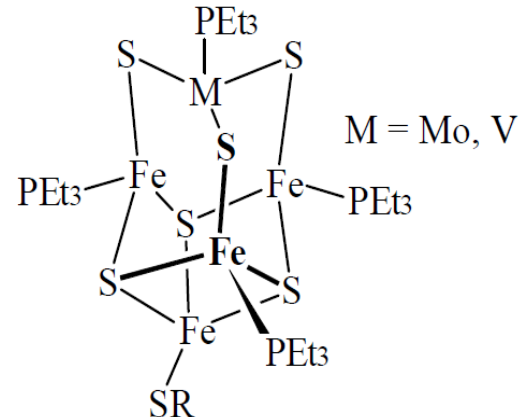
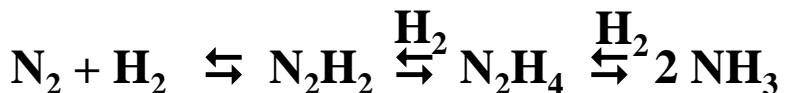
The Chromium Group

Nitrogen fixation by nitrogenases

Haber-Bosch	$N_2 + 3 H_2 \rightleftharpoons 2 NH_3$	400 – 500 °C
V-nitrogenase	$2 N_2 + 14 H^+ + 12 e^- \rightleftharpoons 2 NH_4^+ + 3 H_2$	RT
Mo-nitrogenase	$2 N_2 + 10 H^+ + 8 e^- \rightleftharpoons 2 NH_4^+ + H_2$	RT



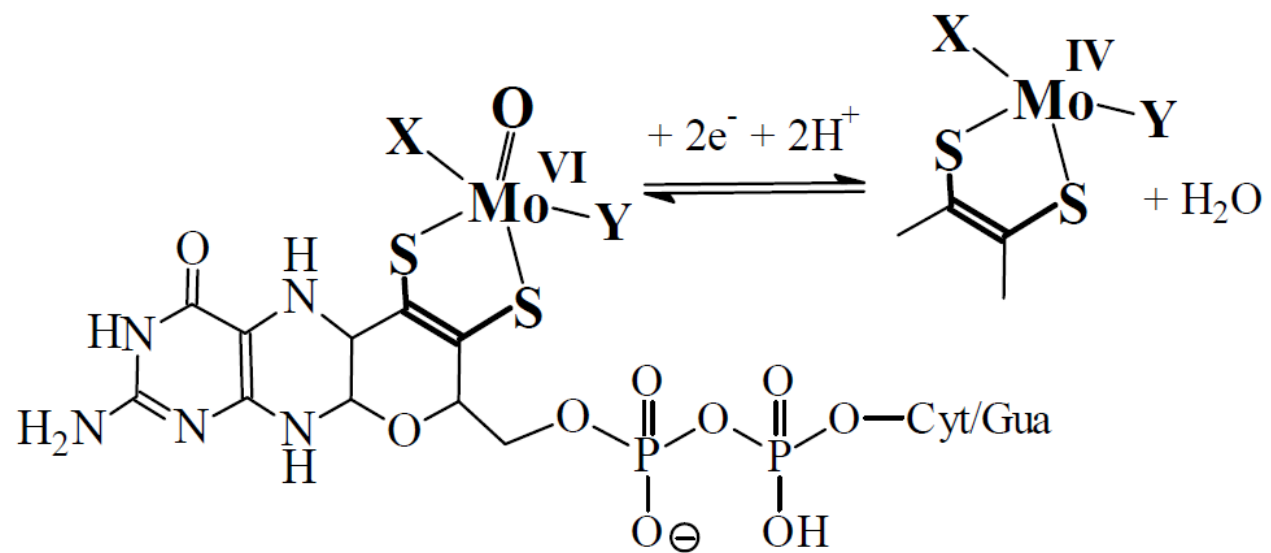
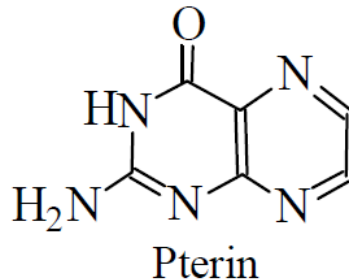
Structural models to elucidate the function



10. Biochemistry of Transition Metals

The Chromium Group

Oxytransferases with molybdenum (molybdopterines)



→ 2-electron-transfer-reactions

DMSO reductase: $(\text{CH}_3)_2\text{SO}_2 \rightarrow$ methylsulphonic acid $(\text{CH}_3)_2\text{SO} \rightarrow$ dimethyl sulphide $(\text{CH}_3)_2\text{S}$ “indicator for biological activity in sea birds“

10. Biochemistry of Transition Metals

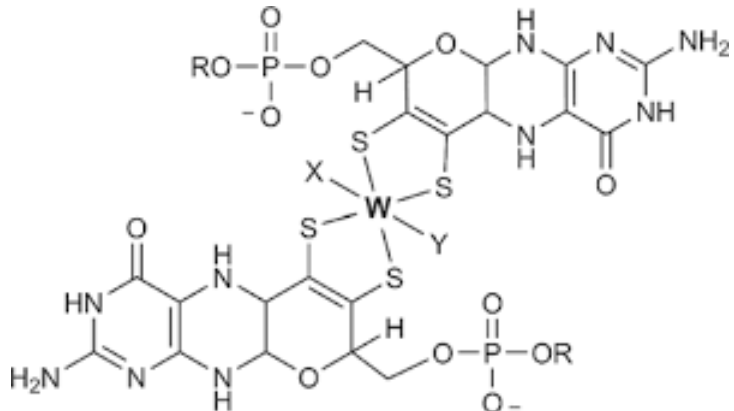
The Chromium Group

Tungsten

- Sole element of 3rd transition metal series (5d) of biological importance
- Metal enzymes in hyper thermal archaeabacteria are stable up to 110 °C, since the strong metal-ligand-interactions stabilise these enzymes
- Stability of W-O-bonds → see tungstates
- Example: Acetylene hydratase $\text{C}_2\text{H}_2 + \text{H}_2\text{O} \rightarrow \text{CH}_3\text{CHO}$ (acetaldehyde)

Literature

- Coord. Chem. Rev. 255 (2011) 1039
- J. Mol. Microbiol. Biotechnol. 26 (2016) 119



10. Biochemistry of Transition Metals

The Manganese Group

Manganese

Mn^{2+}	[Ar]3d ⁵
Mn^{3+}	[Ar]3d ⁴
MnO_4^{4-}	[Ar]3d ³
MnO_4^{3-}	[Ar]3d ²
MnO_4^{2-}	[Ar]3d ¹
MnO_4^-	[Ar]

pale-rose
red
brown
blue
green
violet

most stable (in the acidic pH range)
tends to disproportionate
does not disproportionate
tends to disproportionate
tends to disproportionate
strong oxidizing agent

Technetium

Radioactive and extremely scarce

TcO_4^-	[Kr]	colourless
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activity ~ 30 millicuries

$^{99\text{m}}\text{Tc}$ ($t_{1/2} = 6 \text{ h}$) in coordinated form is used in diagnostic nuclear medicine, as citrate or diphosphonato methane complex: Single Photon Emission Computed Tomography (SPECT)

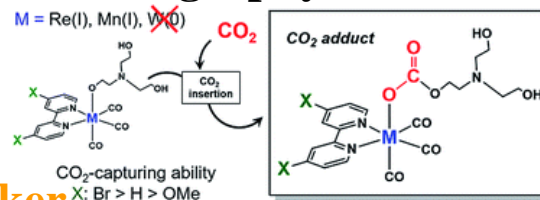
Rhenium

Rhenium is extremely scarce and rhenium oxide is poorly soluble

Re^+	[Xe]4f ¹⁴ 5d ⁶ l.s.	yellow to red
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colourless

CO_2 activation, marker
weakly oxidizing



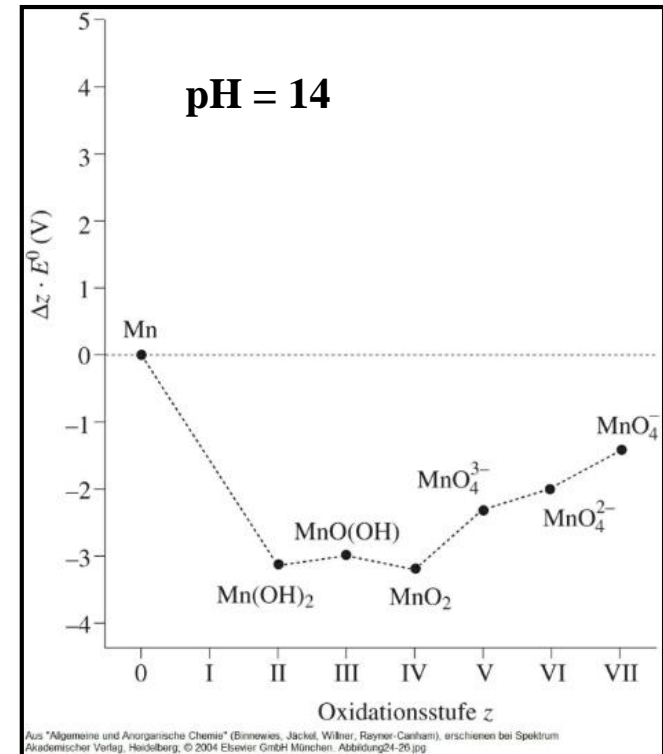
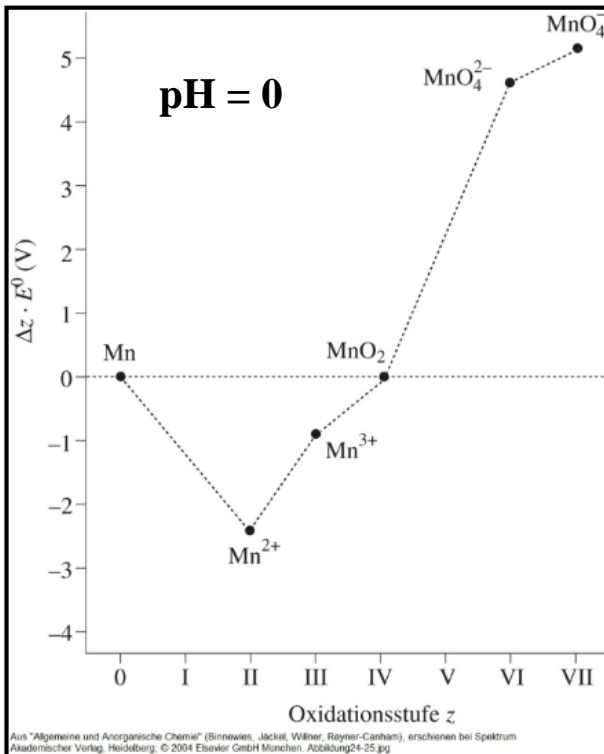
10. Biochemistry of Transition Metals

The Manganese Group

Manganese redox chemistry

- Metallic manganese tends to be oxidised $\text{Mn} \rightleftharpoons \text{Mn}^{2+} + 2 \text{e}^- \quad E^\circ = -1.19 \text{ V}$
- In acidic solution Mn^{2+} is the most stable oxidation state
- In alkaline solution Mn^{4+} is the most stable species but Mn^{2+} and Mn^{3+} possess similar stability

Frost diagram for manganese in acidic (right) and alkaline (left) solution

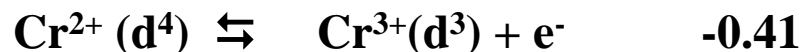
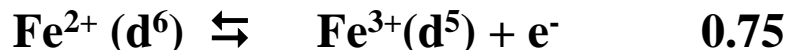
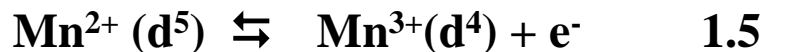


10. Biochemistry of Transition Metals

The Manganese Group

Oxidation state +II ($[Ar]3d^5$)

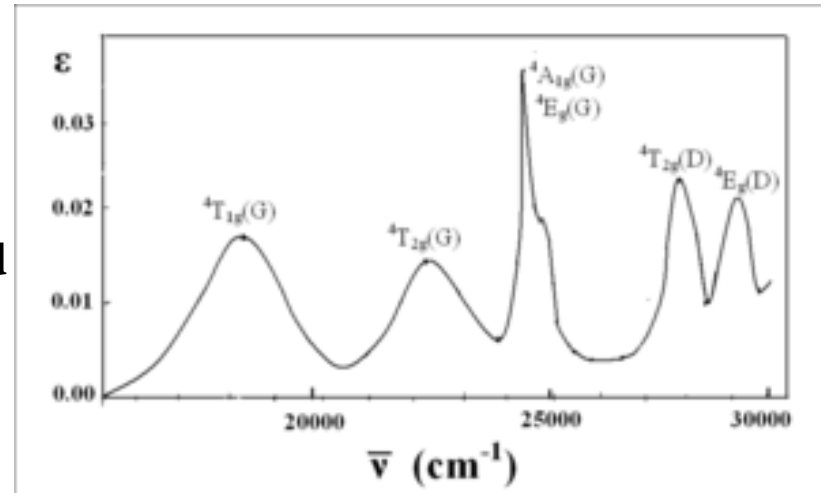
- Mn²⁺ is relatively stable in comparison to other divalent TM-ions and not a reducing agent in acidic solution: E⁰ [V] at pH 0



- Manganese(II)-salts or solutions are only weakly coloured since the absorption in the visible range is only possible via spin-forbidden 3d-3d-transitions (d⁵, high-spin)



- Strongly coloured low-spin complexes are formed only with very strong ligands, e.g.



10. Biochemistry of Transition Metals

The Manganese Group

Oxidation state +III ([Ar]3d⁴)

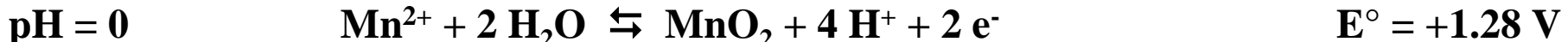
- Dissolution of braunite Mn₂O₃ in conc. H₂SO₄
$$\text{Mn}_2\text{O}_3 + 6 \text{H}^+ + 9 \text{H}_2\text{O} \rightleftharpoons 2 [\text{Mn}(\text{H}_2\text{O})_6]^{3+}$$
results in a solution of the garnet red hexaaquamanganese(III)-ions
- Manganese(III)-ions tend to disproportionation
$$2 \text{Mn}^{3+} + 2 \text{H}_2\text{O} \rightleftharpoons \text{Mn}^{2+} + \text{MnO}_2 + 4 \text{H}^+$$
if no reducing agent is present
- The stable, dark red manganese(III)-acetate is formed upon exposure of permanganate to manganese(II)-acetate in glacial acetic acid:
$$3 \text{KMnO}_4 + 12 \text{Mn(OAc)}_2 + 11 \text{HOAc} + 3 \text{H}^+ \rightarrow 5 [\text{Mn}_3\text{O(OAc)}_6]\text{OAc} \downarrow + 7 \text{H}_2\text{O} + 3 \text{K}^+$$
(HOAc = CH₃-COOH)
- Mixed-valent compounds are strongly coloured → MMCT (Intervalence compounds)
$$[\text{L}_2\text{Mn}^{\text{II}}\text{Mn}^{\text{III}}(\mu\text{-OH})_3]^{2+} \quad \text{with L = 1,4,7,-Trimethyl-1,4,7-triaza cyclononane}$$
$$[\text{L}_2\text{Mn}^{\text{III}}\text{Mn}^{\text{IV}}(\mu\text{-O})_2(\mu\text{-OH})]^{2+}$$

10. Biochemistry of Transition Metals

The Manganese Group

Oxidation state +IV ([Ar]3d³)

$\text{MnO}_2 \cdot \text{H}_2\text{O} = \text{MnO(OH)}_2$ is a strong oxidizing agent in acidic solution



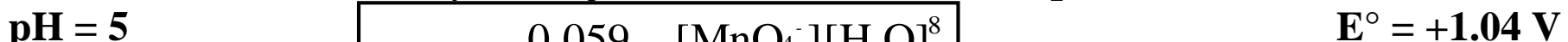
⇒ lab-synthesis of chlorine



⇒ O₂ oxidizes manganese(II)-hydroxide to MnO₂·H₂O

Oxidation state +VII ([Ar]3d⁰)

The violet permanganate ion MnO₄⁻ is a strong oxidizing agent in acidic solution



Permanganate can be formed through oxidation of Mn²⁺ with PbO₂ in acidic environment

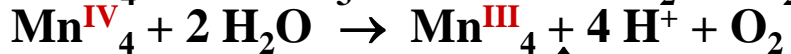


10. Biochemistry of Transition Metals

The Manganese Group

Manganese in the biosphere

- Manganese is the key element in the light reaction of photosynthesis, i.e. it is needed to cleave water in the oxygen-evolving cluster:



→ Photosystem II (light reaction)

- Arginase

Nitrogen-containing metabolite
→ urea synthesis $\text{H}_2\text{N}-\text{CO}-\text{NH}_2$

- Superoxide dismutase

Decomposition of superoxide radical O_2^-



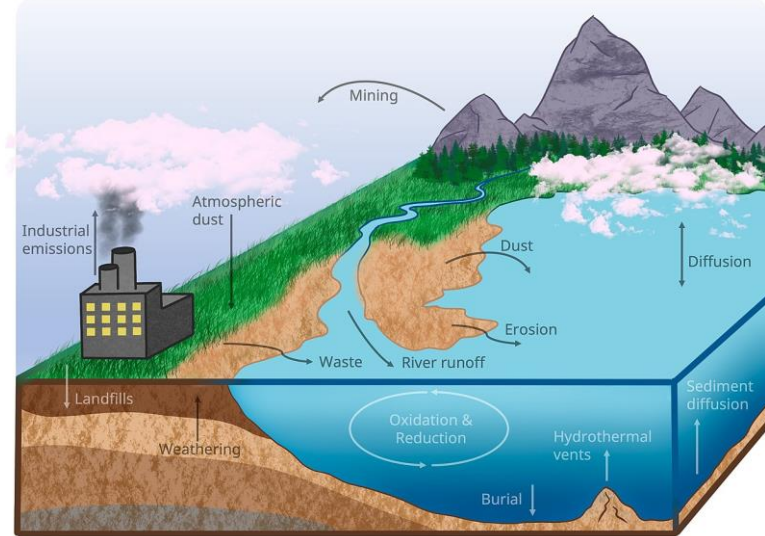
Mn besides Zn, Cu, Fe and Se is a co-factor for anti-oxidative acting enzymes

- Pyruvate carboxylase

Conversion of pyruvate in oxaloacetate by activation of HCO_3^-



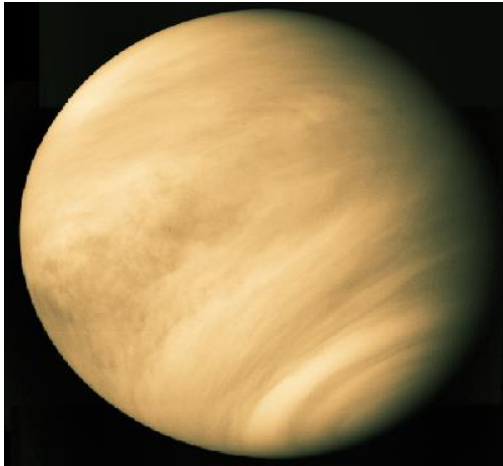
Enzyme-biotin- CO_2 + pyruvate → enzyme-biotin + oxaloacetate



10. Biochemistry of Transition Metals

Photosynthesis: The Energetic Base of the Biosphere (Mn Catalysed H₂O Cleavage)

Venus



2.61 kW/m²

Albedo = 0.76

→ T_e = 232 K

96% CO₂ + 3% N₂ +
SO₂ + H₂O + Ar (ppm)

93 bar → T_{eff} = 740 K

Earth



1.37 kW/m² = 1.56·10¹⁸ kWh/a

Albedo = 0.30

→ T_e = 255 K

78% N₂ + 21% O₂ + 0.9% Ar
+ CO₂ + H₂O + CH₄ (ppm)

1 bar → T_{eff} = 288 K

Life = aqueous chemistry

Water → H₂ and O₂ → energy!

3 O₂ → 2 O₃ by VUV photolysis

Mars



0.59 kW/m²

Albedo = 0.15

→ T_e = 213 K

95% CO₂ + 3% N₂ +
1.5% Ar + H₂O (ppm)

5.6 mbar → T_{eff} = 225 K

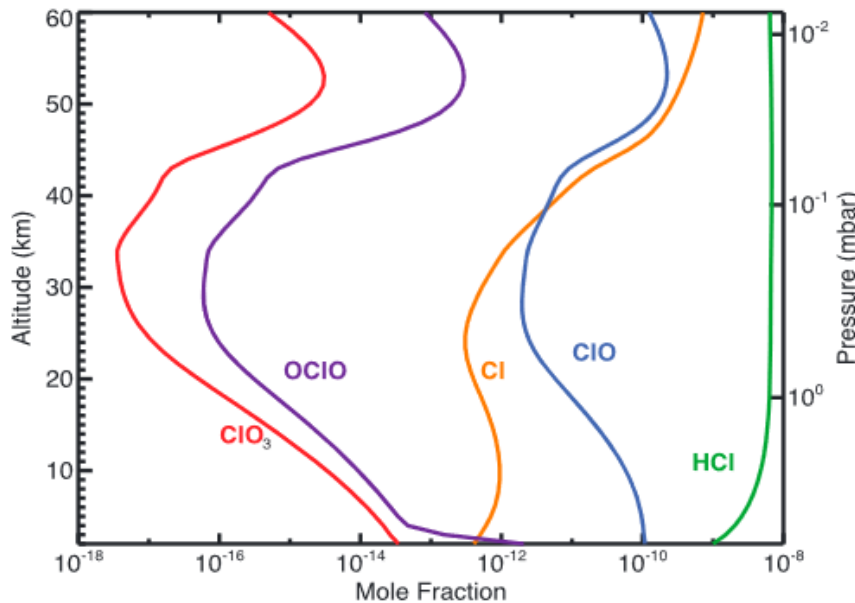
Remark: O₂ converted to
Ca(ClO₄)₂ by radiolysis

10. Biochemistry of Transition Metals

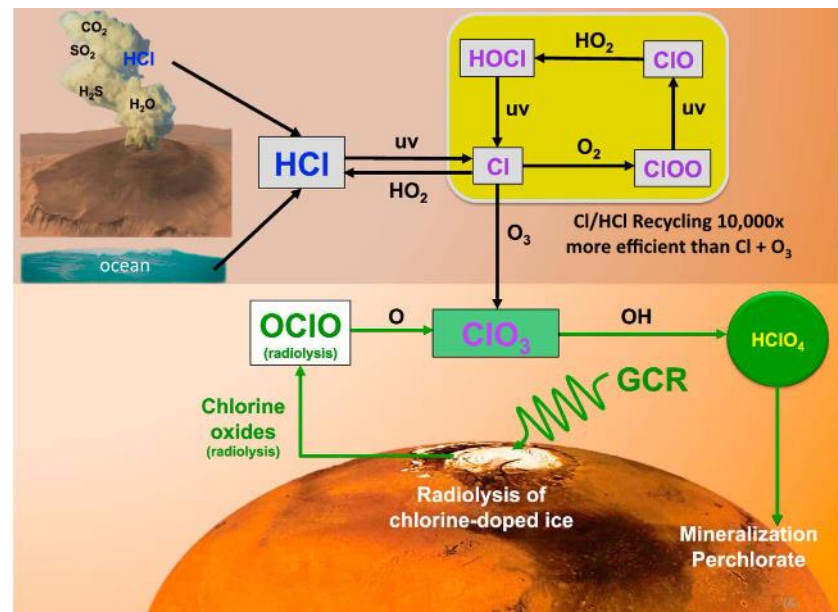
Photosynthesis: Possible on surface of Mars?

No, since the ground layer is full of perchlorates due to mineralisation of HClO_4 formed by radiolysis of chlorine doped water ice $\rightarrow \text{Ca}(\text{ClO}_4)_2$

Chlorine species in Martian atmosphere



Conversion of HCl towards HClO_4



Lack of a Martian magnetic field caused $\text{H}_2\text{O}/\text{H}_2$ loss due to solar wind

Lit.: Perchlorate formation on Mars through surface radiolysis, J. Geophys. Res. Planets 121 (2016) 1472

10. Biochemistry of Transition Metals

Photosynthesis: Almost All Energy Consumed by Living Organisms Stems from Solar Energy (Exception: Thermophiles in the Deep Sea)

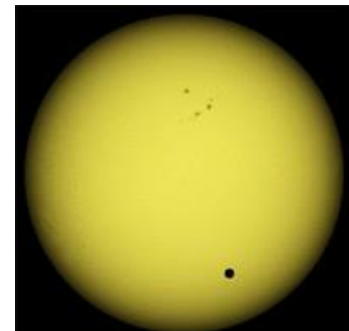
Energy source in solar system: The sun

Luminosity (radiation flux) $3.8 \cdot 10^{26} \text{ W}$

Annual radiation power $1.24 \cdot 10^{34} \text{ J}$ (presently!)

Habitable zone

Venus (early stage of solar system), earth (today), mars (late phase...)



Venus transit

Planet	Perihelion- and aphelion-distance in astronomic units	Solar radiation maximum und minimum (W/m^2)
Mercury	0.3075 – 0.4667	14,446 – 6,272
Venus	0.7184 – 0.7282	2,647 – 2,576
Earth	0.9833 – 1.017	1,413 – 1,321
Mars	1.382 – 1.666	715 – 492
Jupiter	4.950 – 5.458	55.8 – 45.9
Saturn	9.048 – 10.12	16.7 – 13.4
Uranus	18.38 – 20.08	4.04 – 3.39
Neptune	29.77 – 30.44	1.54 – 1.47

10. Biochemistry of Transition Metals

Photosynthesis: Almost All Energy Consumed by Living Organisms Stems from Solar Energy (Exception: Thermophiles in the Deep Sea)

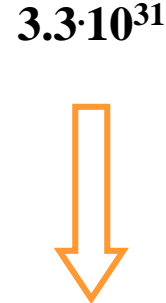
Energy flux in solar system

	Radiation flux [W]	Irradiance I [W/m ²]	Energy flux / day [J]
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Sun	$3.8 \cdot 10^{26}$	$6.37 \cdot 10^7$	$3.3 \cdot 10^{31}$
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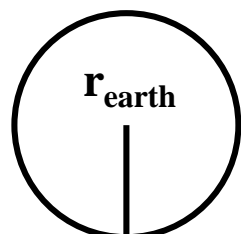
$$(r_{\text{Sun-earth}}^2/r_{\text{sun}}^2 \sim 46200)$$



Earth	$7.0 \cdot 10^{17}$	1420 (Aphelion)	$6.0 \cdot 10^{22}$
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1328 (Perihelion)

1367 (Extraterrestrial solar constant) = I_C



solar
radiation

$$\text{Global } I_C = \pi r_{\text{earth}}^2 / 4\pi r_{\text{earth}}^2 = I_C / 4$$
$$\text{Global albedo (reflectance) } a_g = 0.3$$

10. Biochemistry of Transition Metals

Photosynthesis: Solar Irradiation on Earth

- Extraterrestrial solar constant
- Global extraterrestrial irradiation
- Earth's surface
- Global absorbed solar energy per year
- Annual biomass production

$$I_C = 1367 \text{ W/m}^2$$

$$I_{CG} = (1-a_g) \cdot 1/4 \cdot I_C = 239 \text{ W/m}^2 \text{ (absorbed)}$$
$$510 \cdot 10^{12} \text{ m}^2 \text{ (510 Mill. km}^2\text{)}$$

$$3.86 \cdot 10^{24} \text{ J} \Rightarrow 1.2 \cdot 10^5 \text{ TJ/s (TW)}$$

$$1.7 \cdot 10^{14} \text{ kg}$$

$$\Rightarrow \sim 1.0 \cdot 10^{17} \text{ kJ } (\Delta G^0(\text{hexose}) = 2872 \text{ kJ/mol})$$

Primary energy consumption (1998)

- World **14 TW ($\cong 1.2 \cdot 10^{10}$ t coal)**
- USA **3 TW**
- FRG **0.5 TW**
- Photovoltaics (energy efficiency ~ 15%)
 $510 \times 10^{12} \text{ m}^2 \cdot (14 \text{ TW} / 1.2 \cdot 10^5 \text{ TW}) / 0.15 = 4 \times 10^{11} \text{ m}^2 \approx 0.4 \text{ Mill. km}^2 \approx 0.08\% \text{ of Earth s.}$

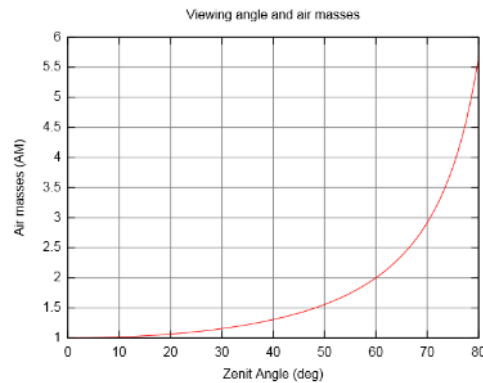
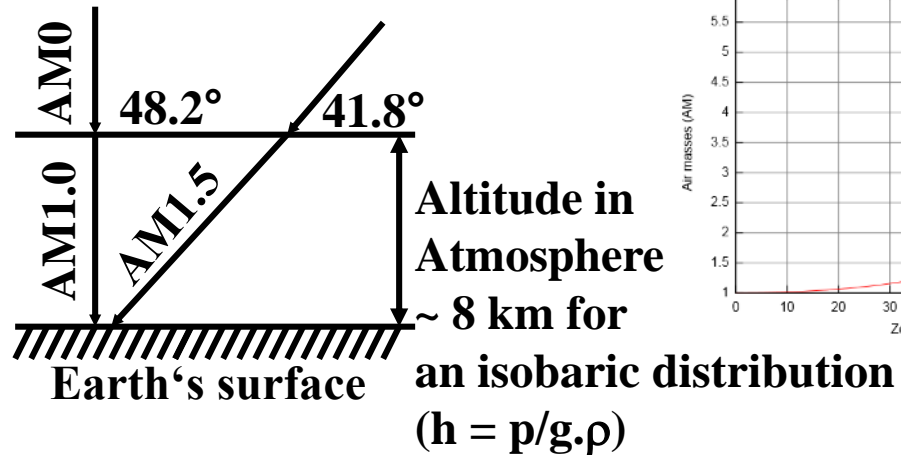
$$\Rightarrow \sim 4.4 \cdot 10^{17} \text{ kJ } (2016: 16 \text{ TW} \Rightarrow \sim 5.2 \cdot 10^{17} \text{ kJ })$$

Fossil fuels

- Global resources $m_{O_2} = 10^{15} \text{ t (O}_2 \text{ in atmosphere)} \rightarrow 400 \cdot 10^{12} \text{ t C}$
- Known resources $10.4 \cdot 10^{12} \text{ t C } (\sim 2.5\%)$

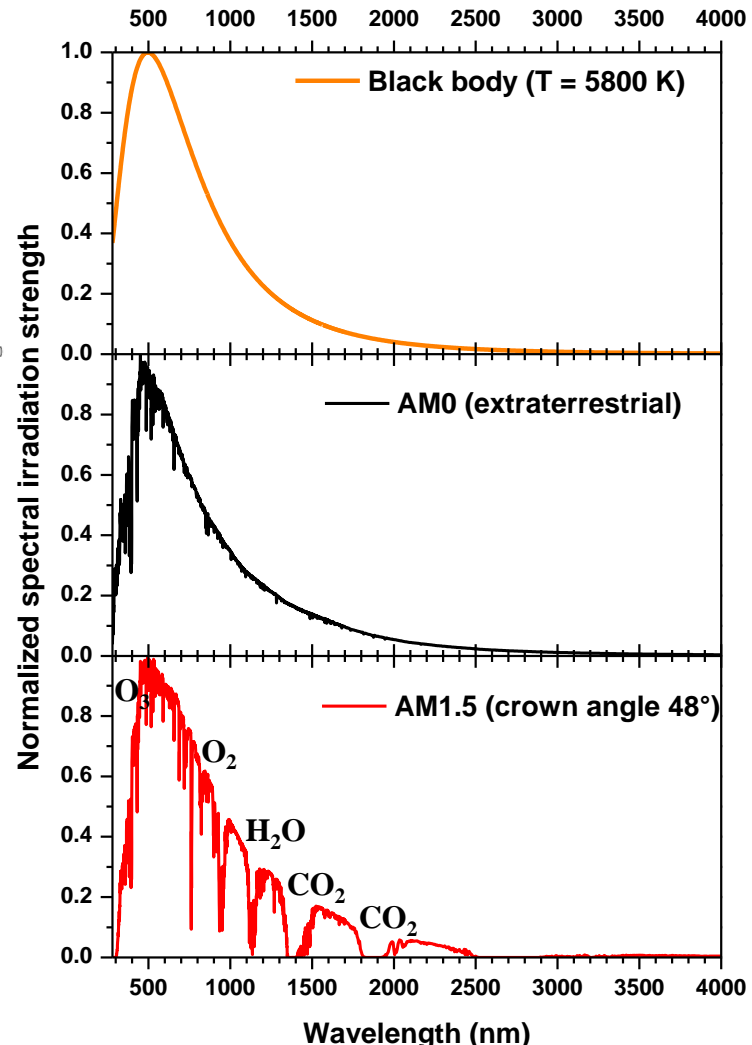
10. Biochemistry of Transition Metals

Photosynthesis: Spectral Distribution of Radiation upon Earth's Surface



AM = Air mass	AM	1.5 (global)
UV ~	5%	
VIS ~	51%	
IR ~	44%	

Solar energy production in thin layers, e.g. leafs, require strong absorbents (antenna molecules)
Absorption constant $\epsilon > 1 \cdot 10^4 \text{ cm}^{-1} \text{M}^{-1}$ 400 – 700 nm
⇒ chlorophyll, carotin, phycocyanin, phycoerthrin



10. Biochemistry of Transition Metals

Photosynthesis: Absorption of Irradiated Solar Energy

$$A + R + T = 1$$

with A = Absorption, R = Reflectance (albedo), T = Transmission

Irradiation of earth

$$T = 0 \Rightarrow A = 1 - R$$

Surface of	R (albedo)	Solar energy use	Absorber	Absorption process
Earth	10 – 25%	Si-solar cells	Silicon	VB – CB transitions
Sand	25 – 40%	Grätzel-solar cells	Ru^{2+} complexes	Metal-to-Ligand-Charge-Transfer (MLCT)
Grass	15 – 25%			
Forrest	10 – 20%	Chloroplasts	Chlorophyll	$\pi - \pi^*$, $n - \pi^*$
Snow	75 – 95%		β -carotin & other accessory pigments	
Sea	10%			
Earth (global)	30%			

High absorption strength only by allowed optical transitions

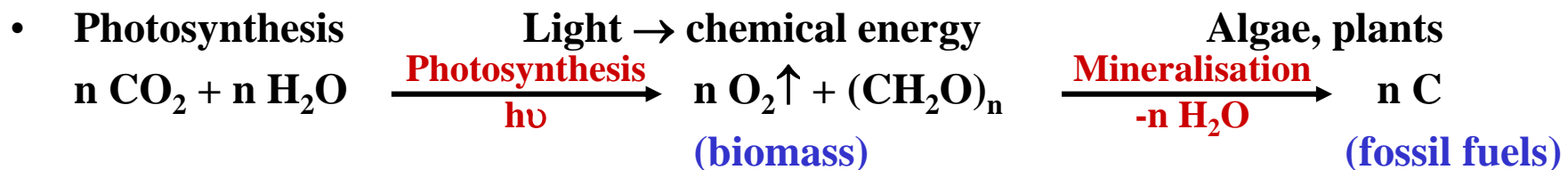
10. Biochemistry of Transition Metals

Photosynthesis: Applications of Irradiated Solar Energy

1. Absorption	Energy transfer via excitons	Excited species
2. Light reaction	Energy uptake	Formation of H ₂ and ATP
3. Dark reaction	Energy storage	Biomass, batteries, etc.

Technical and biological applications

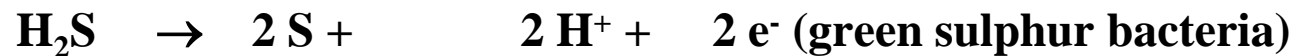
• Solarthermics	Light → thermal energy	Solar collectors
• Photovoltaics	Light → electrical energy	Solar cells 1. Semiconductor (Si, Ge) 2. Liquids (Grätzel)
• Photosynthesis	Light → chemical energy	Algae, plants Mineralisation -n H ₂ O



10. Biochemistry of Transition Metals

Photosynthesis: Energy Production in Autotrophic Organisms

- Light reaction photolysis of a hydrogen donor (energy uptake)



Photosynthetic Active Radiation (PAR) = 400 – 700 nm (170 – 300 kJ/mol)

- Dark reaction Synthesis of carbohydrates (energy storage)



Photosynthetic CO₂ fixation in glucose requires about 470 kJ/mol per C-Atom

Photochemical work

$$W = I \cdot A \cdot \Phi$$

with

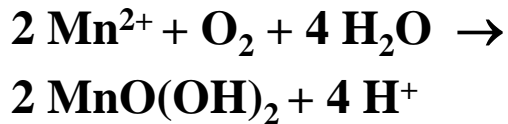
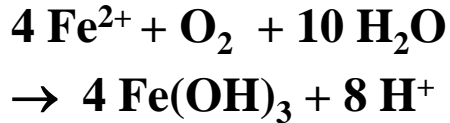
I = Irradiance [W/m²]

A = Absorption [0.0 ... 1.0]

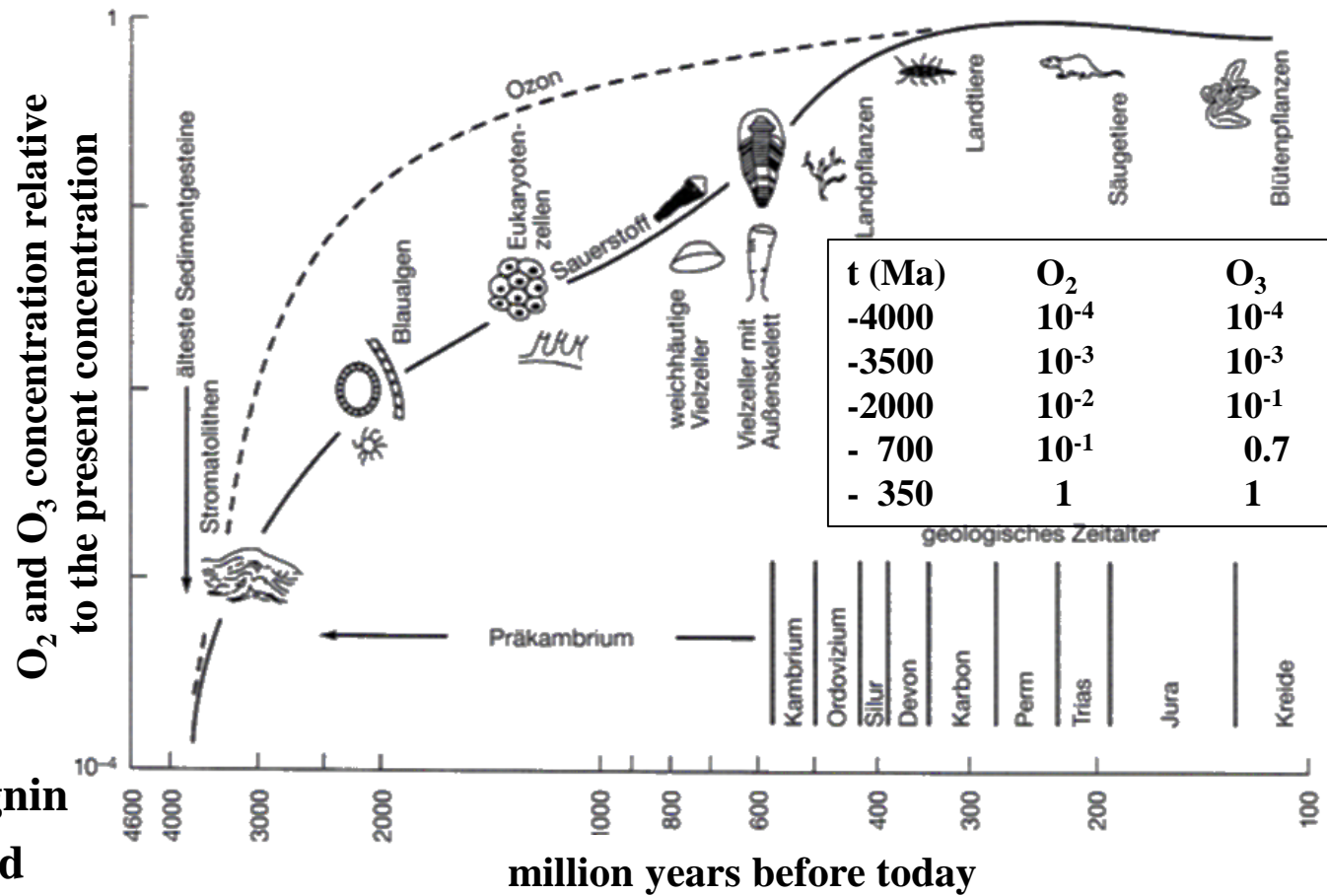
Φ = Quantum yield [0.0 ... 1.0]

10. Biochemistry of Transition Metals

Photosynthesis: Oxygen production as a side effect → O₃ layer formation



→ Fe and Mn deficiency
→ Self inhibition of photosynthesis



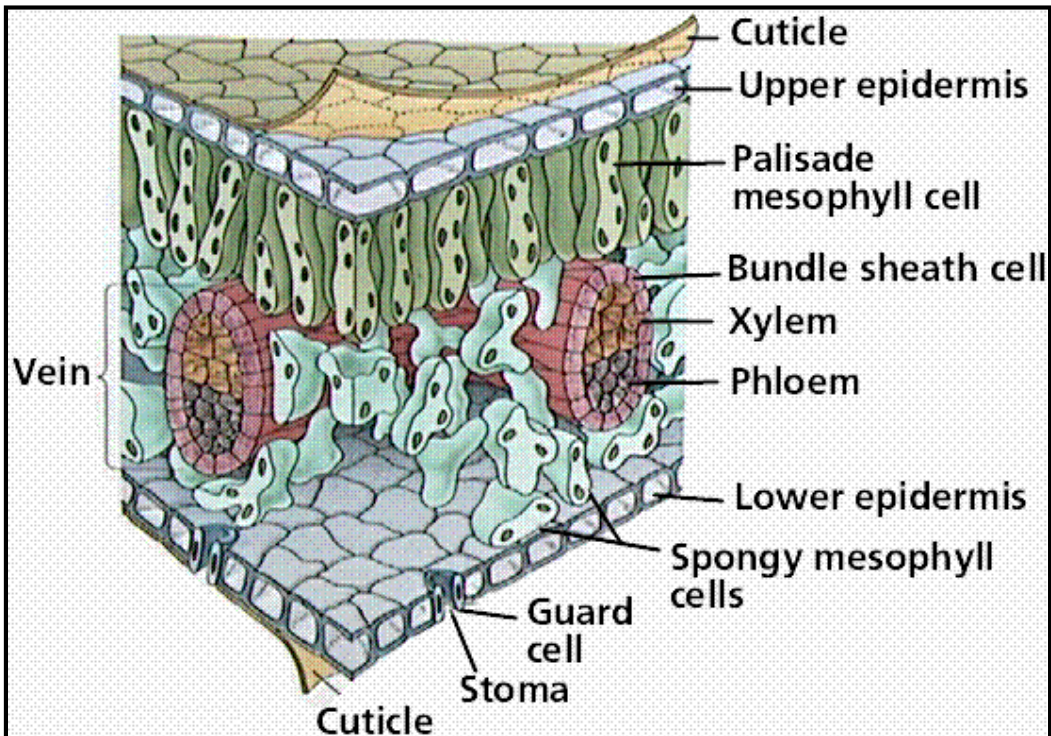
Carbon age: Invention of lignin
→ strong CO₂ reduction and
O₂ concentration rose to 30-35%

Source: Graedel Crutzen 1994

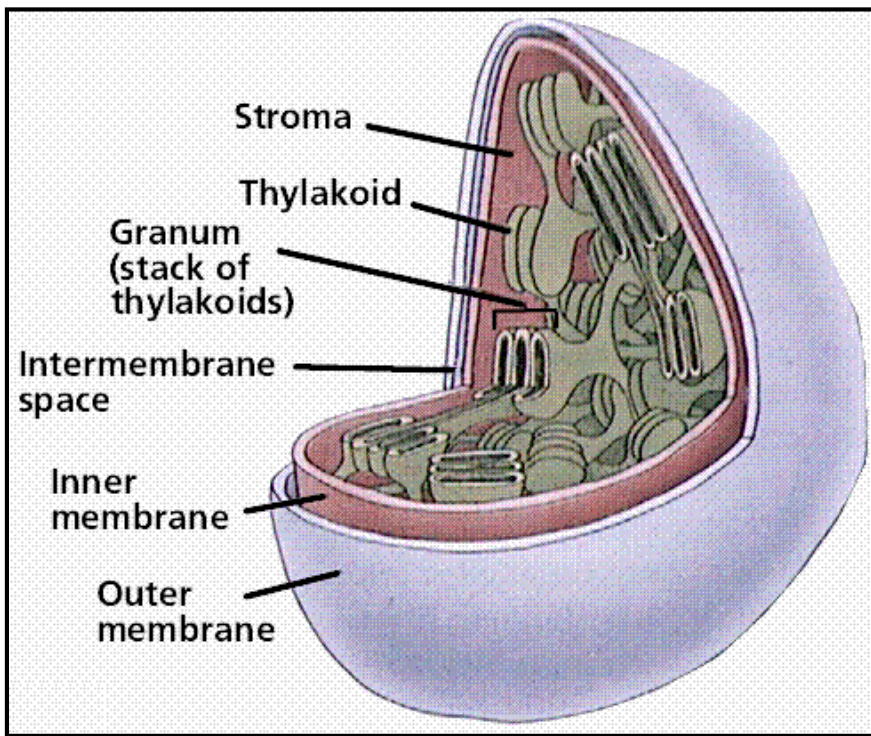
10. Biochemistry of Transition Metals

Photosynthesis: Location and Structure of Chloroplasts

Cross-section of a leaf of a higher plant



Schematic built of a chloroplast



Photosynthetic activity takes place in thylakoid membranes

- Membrane potential ~ 0.2 V
- Lipid/protein-ratio ~ 1:1

10. Biochemistry of Transition Metals

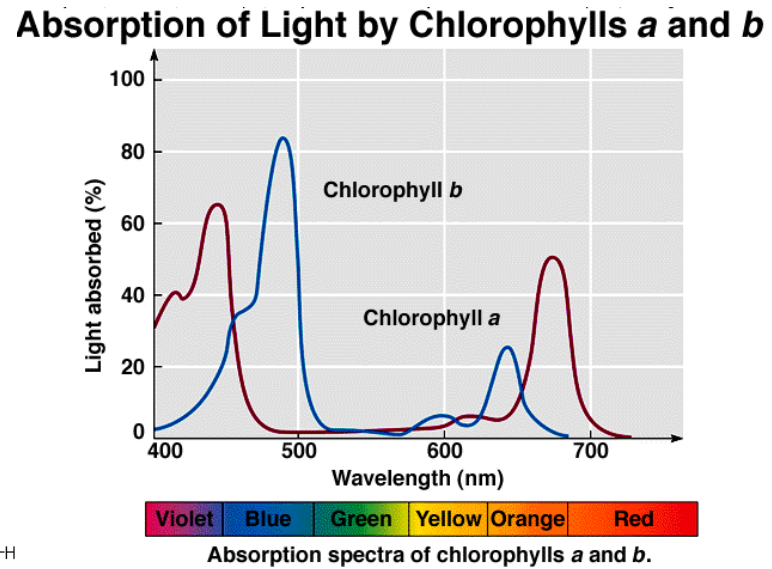
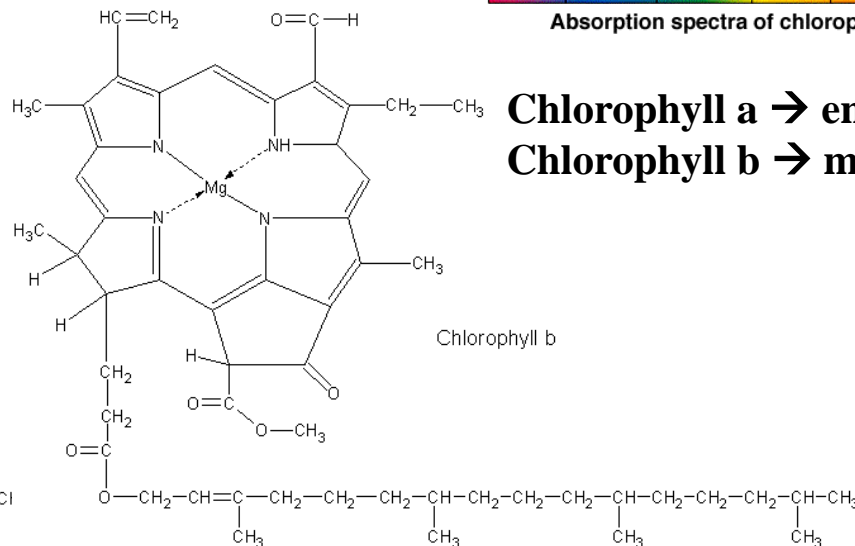
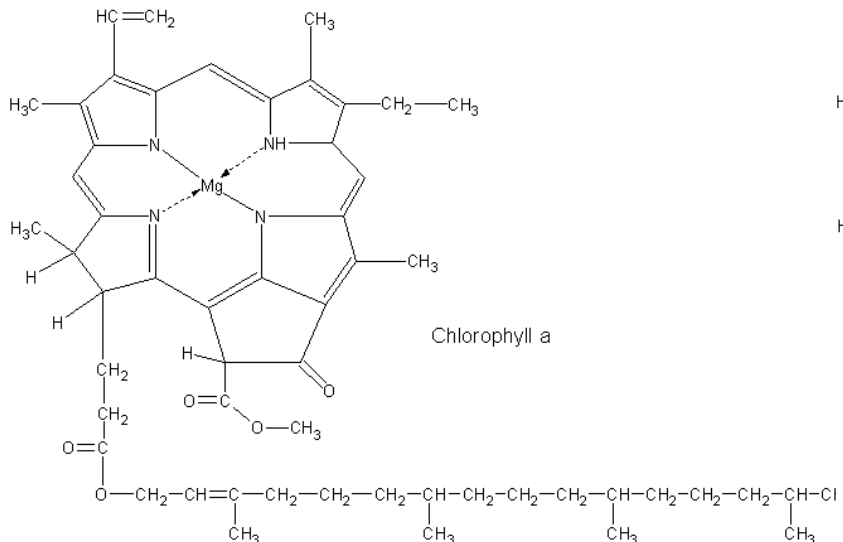
Photosynthesis: Antenna Dyes

Dye

Substances absorbing light selectively, resulting in a subtractive colour spectrum

Chlorophyll a, b

Green dye, present in all photosynthetically active cells, and absorbing in the blue and red spectral range



Chlorophyll a → energy transfer
Chlorophyll b → main absorber

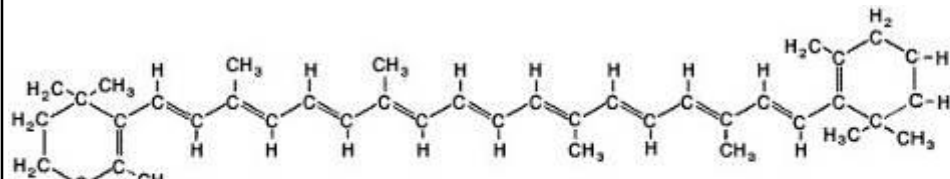
10. Biochemistry of Transition Metals

Photosynthesis: Antenna Dyes

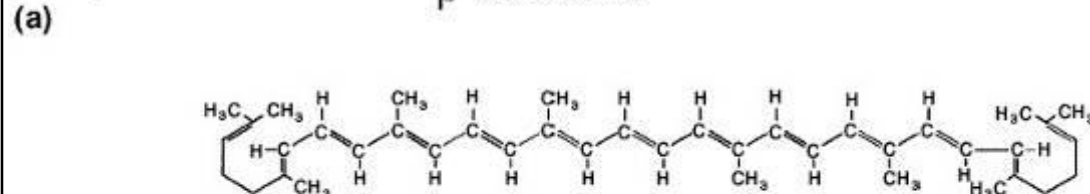
Accessory pigments

Dyes which completes the absorption spectrum of chlorophyll in the visible range:
chlorophyll a, chlorophyll b, carotenoids, ...

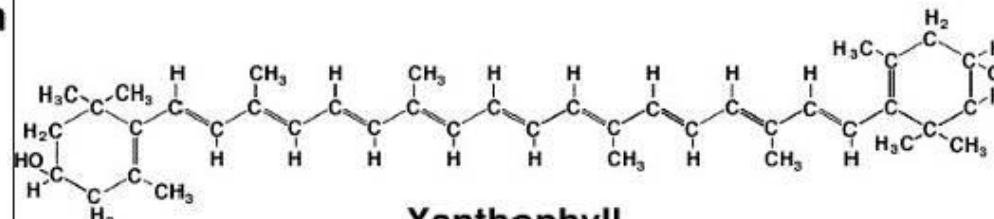
β -Carotin, lycopene, phycocyanin, phycoerythrin, ...



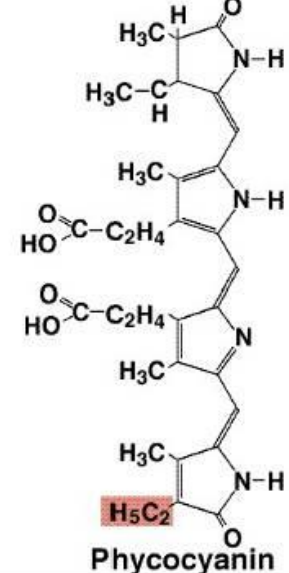
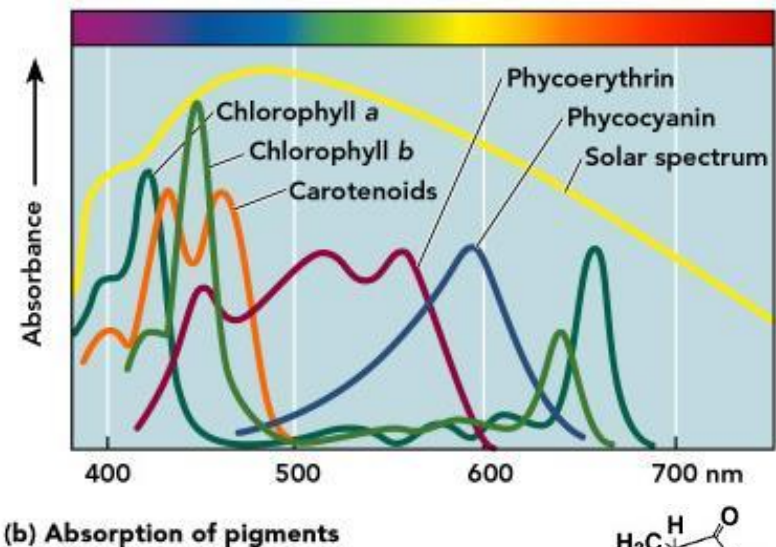
β -carotene



Lycopene



Xanthophyll



Phycocyanin

10. Biochemistry of Transition Metals

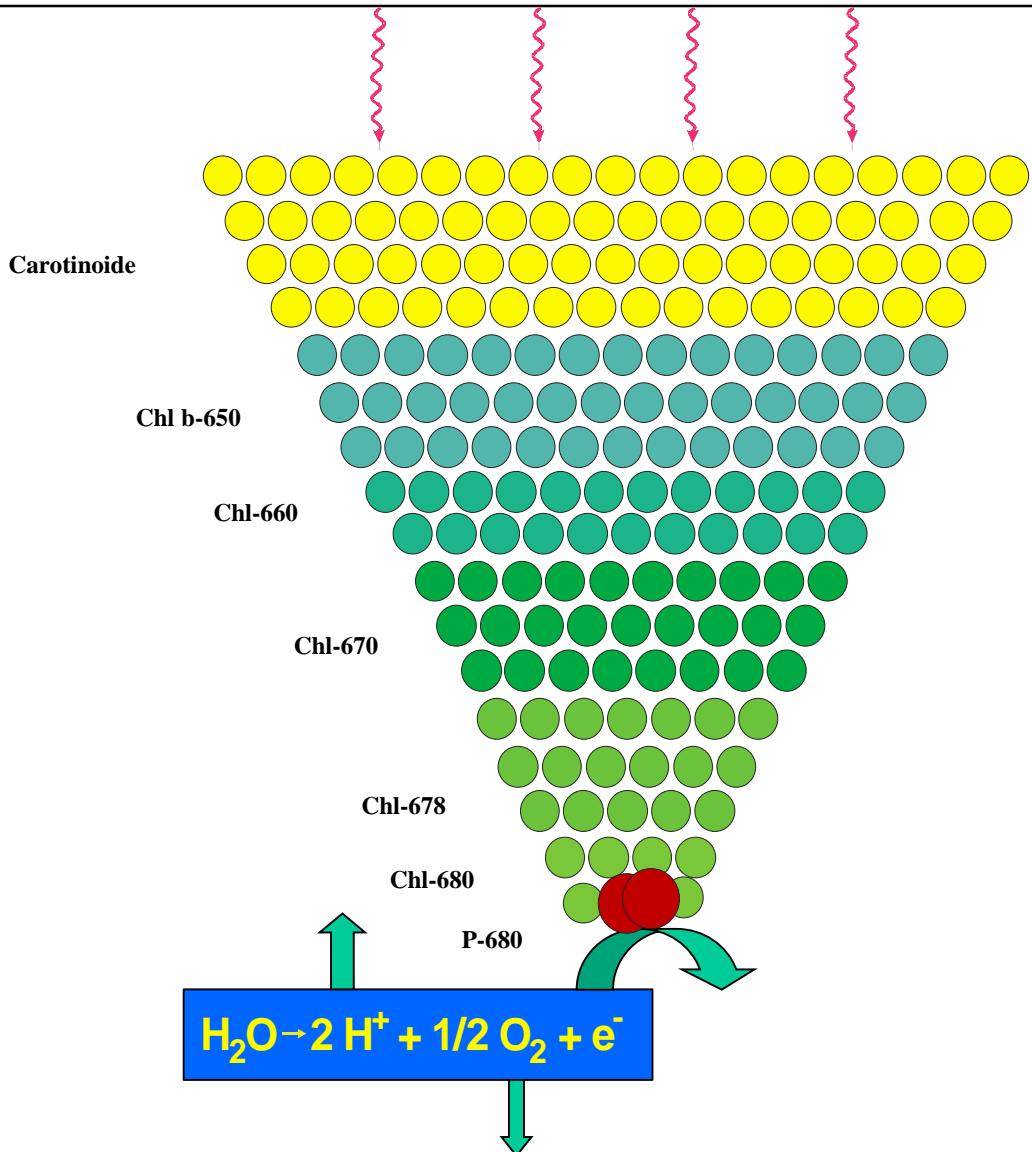
Photosynthesis: Antenna Dyes

Energy transfer (ET) to
reaction centre **Photosystem II**
(PS II)

Problem: Formation of singlet Oxygen,
which may destroy proteins

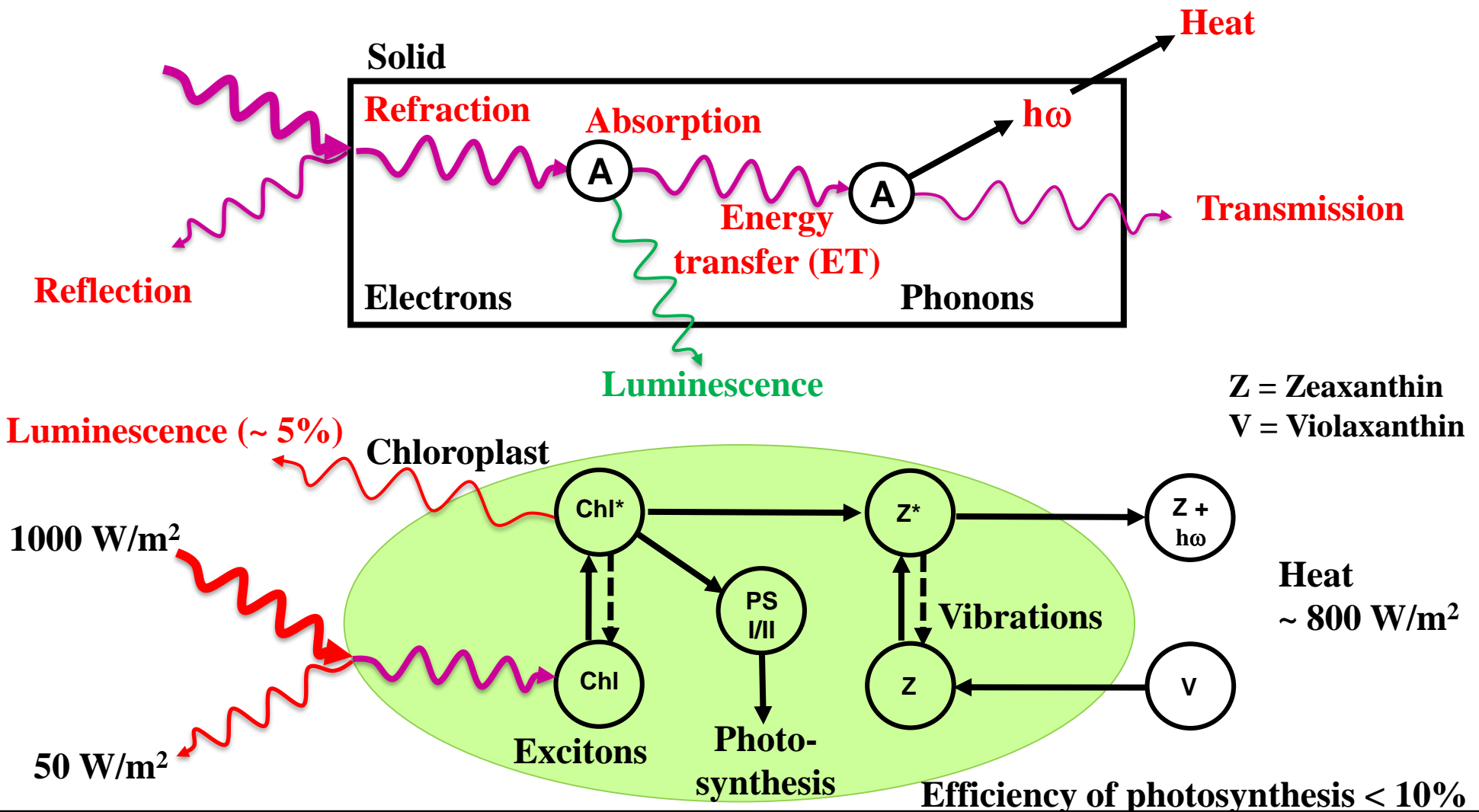
- $^1\text{Chl} + \text{hv} \rightarrow ^1\text{Chl}^*$
- $^1\text{Chl}^* \rightarrow ^3\text{Chl}^*$
- $^3\text{Chl}^* + ^3\text{O}_2 \rightarrow ^1\text{Chl} + ^1\text{O}_2$

Solution: Mg^{2+} incorporation
prevents formation of $^3\text{Chl}^*$
due its weak spin-orbit coupling



10. Biochemistry of Transition Metals

Photosynthesis: Absorption of Irradiated Solar Energy



10. Biochemistry of Transition Metals

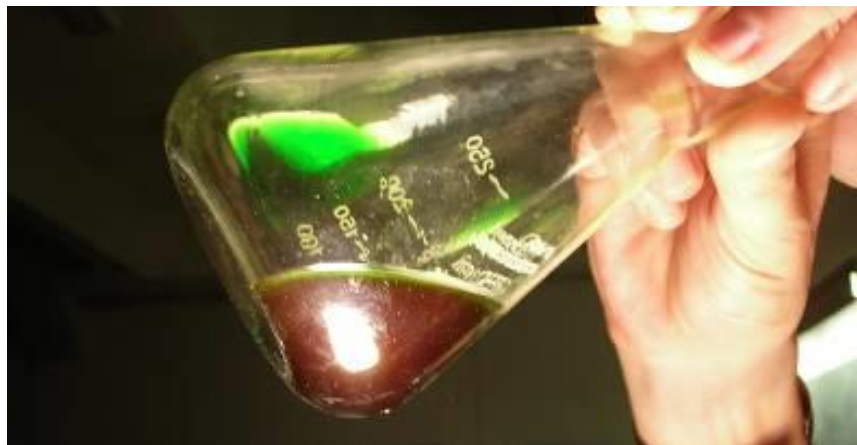
Photosynthesis: Time scales

- Absorption photon: ground-state → singlet state 10^{-15} s
- Higher singlet-state → singlet state $10^{-14} - 10^{-13}$ s
- Lowest singlet state → ground state $10^{-11} - 10^{-9}$ s
- Lifetime triplet state $10^{-4} - 10^{-2}$ s
(not relevant)
- Transfer of energy between adjacent molecules 10^{-10} s
- Transfer of energy to trigger chemical reactions
(ET to reaction centre) 10 ms

10. Biochemistry of Transition Metals

Photosynthesis: Fluorescence of Chlorophyll

- Pure chlorophyll + blue light
 - intensive red fluorescence
 - no energy migration
- Chlorophyll in chloroplast + light
 - weak fluorescence
 - Energy transfer to PSI/II



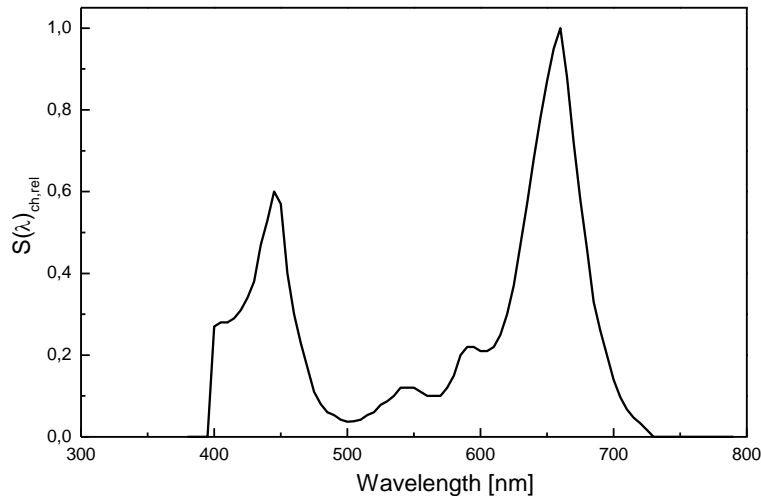
10. Biochemistry of Transition Metals

Photosynthesis: Action Spectra

Absorbed energy in PSI and II

1. Heat dissipation 80%
2. Luminescence 3-7%
3. Chemical reactions 10-15%
e.g. chlorophyll and sugar synthesis

Chlorophyll synthesis action spectra



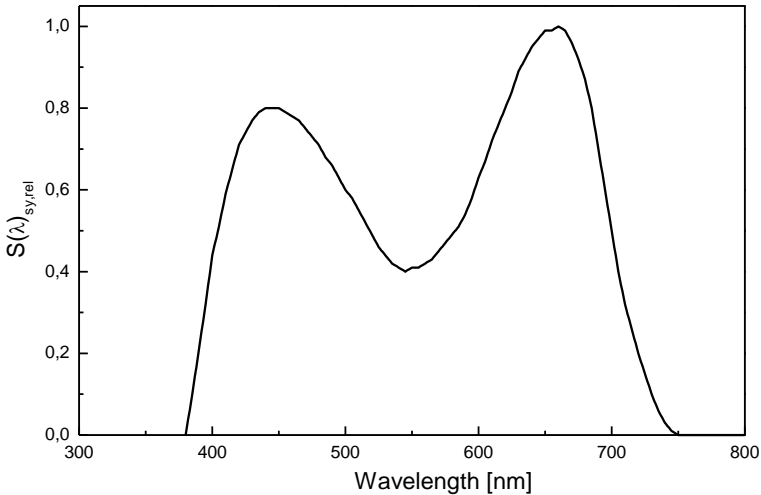
Action spectra

Relative activity of different photon energies
to achieve a measurable effect

→ Similar absorption spectrum compared to that
of chlorophyll

Taken from DIN 5031-10

Photosynthesis action spectra



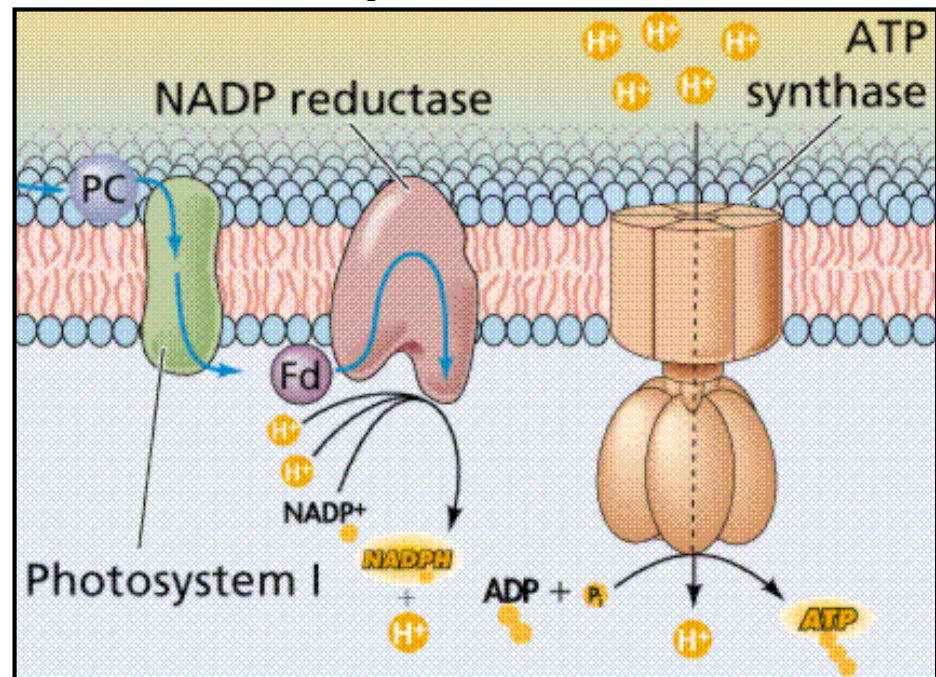
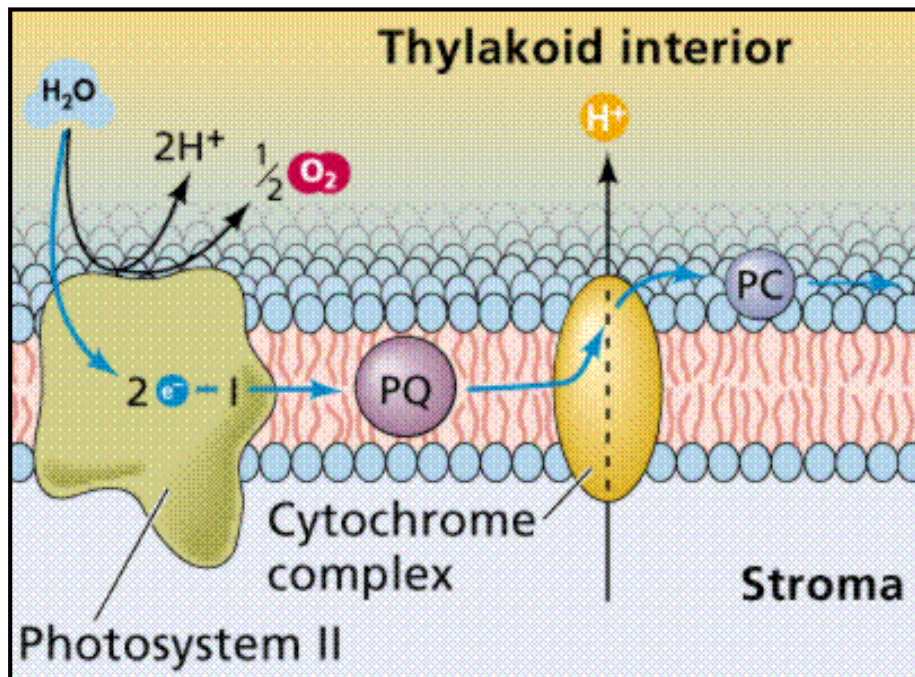
10. Biochemistry of Transition Metals

Photosynthesis: Emerson-Experiment

⇒ The light reaction consists of coupled photo systems: photosystem I and II

Photosystem II: Absorption up to 680 nm (P 680)

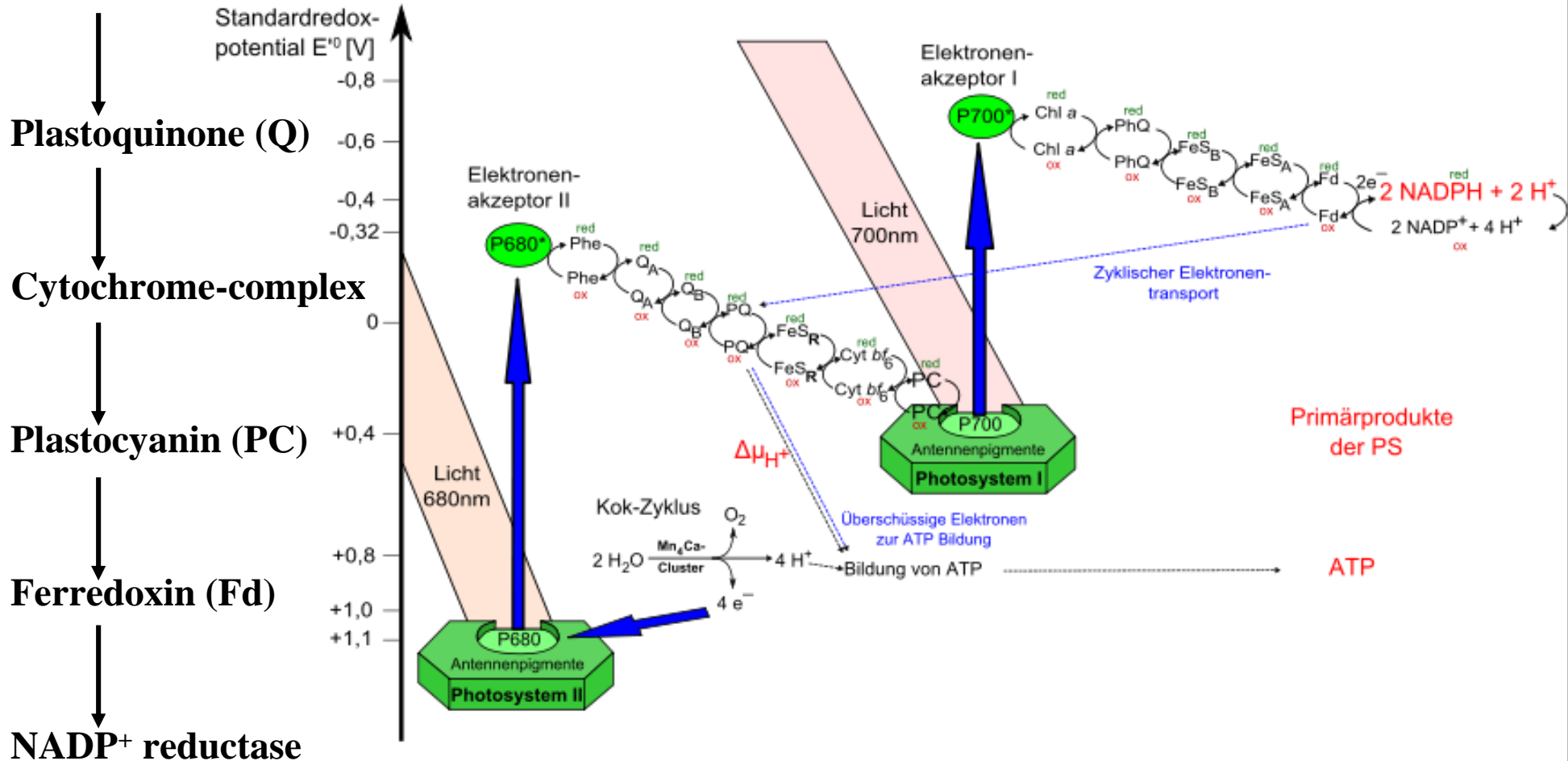
Photosystem I: Absorption up to 700 nm (P 700)



10. Biochemistry of Transition Metals

Photosystem I and II: Electron Flux

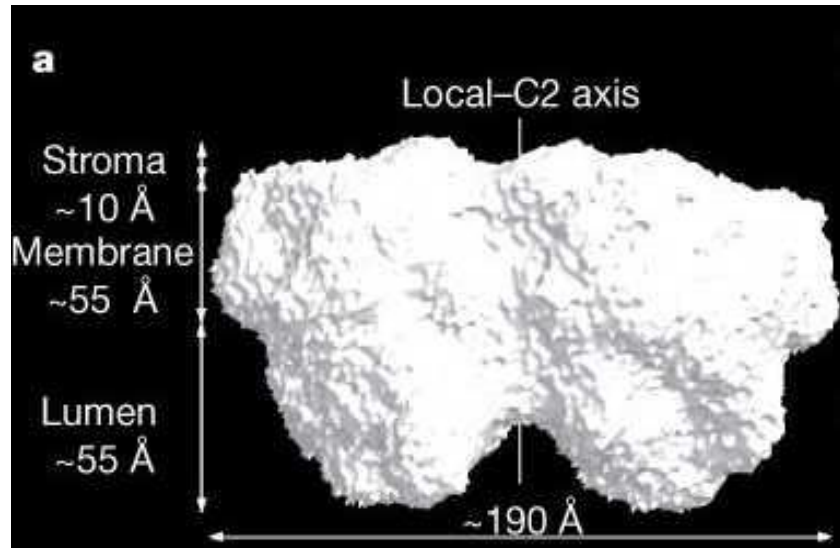
Mn-cluster



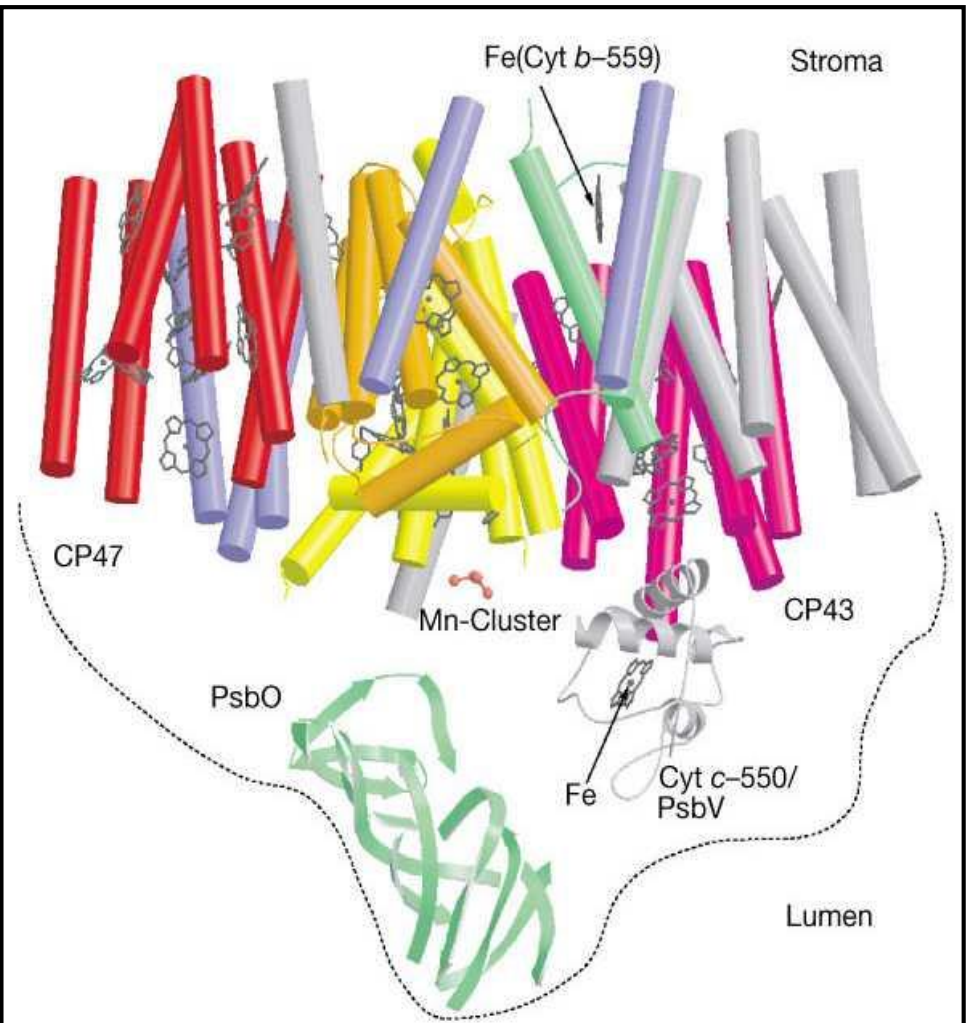
10. Biochemistry of Transition Metals

Photosynthesis: Structure of Photosystems II (Dimer), M = 350 kDa (cyanobacteria)

Electron density on the surface of PS II:
(cyanobacterium: *synechococcus elongatus*)



- Central Mn-cluster contains four Mn-ions, Ca^{2+} and Cl^-
- 2 Mn-Mn distances of 2.7 Å
- 1 Mn-Mn distances of 3.3 Å

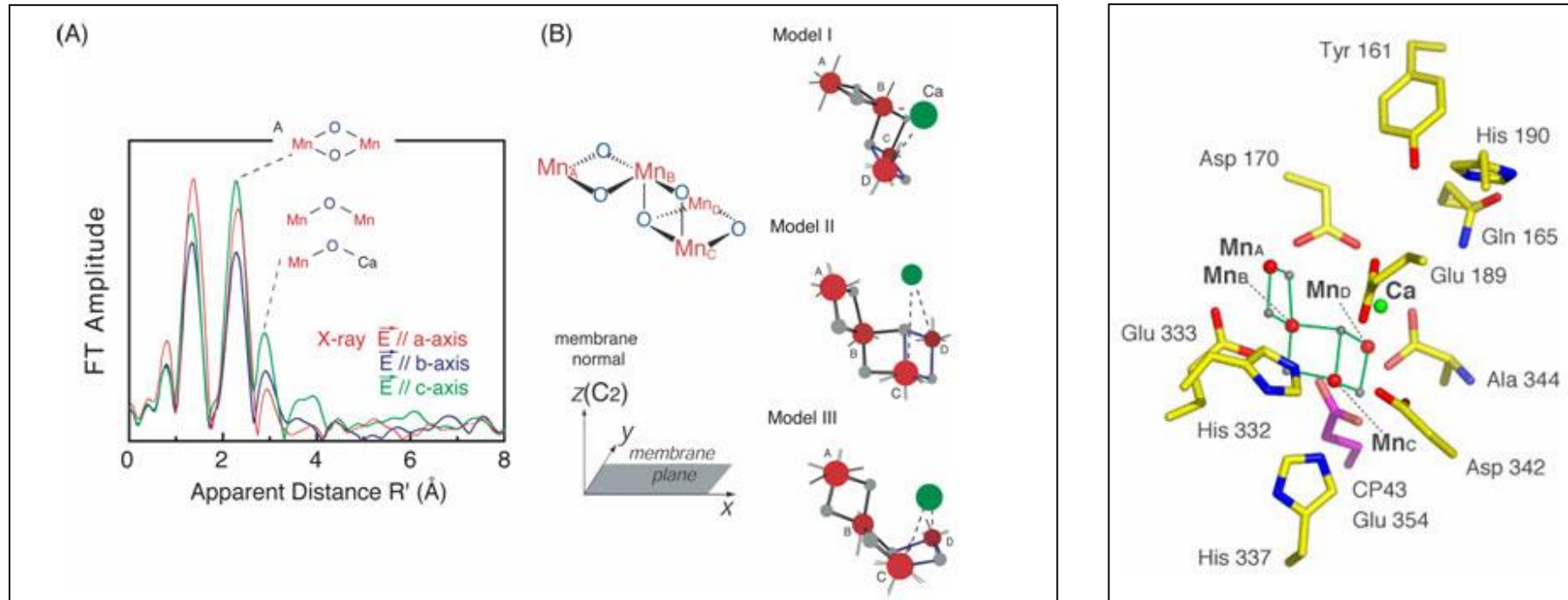


10. Biochemistry of Transition Metals

Photosynthesis: Structure of Photosystem II – Spectroscopic Analysis

a) Extended X-ray Absorption at Fine Structure (EXAFS)

→ determination of distances between heavy atoms (Mn- and Ca-ions)



b) EPR-spectroscopy → oxidation state of the manganese atoms

c) T-dependent magnetic susceptibility → magnetic coupling (super-exchange)

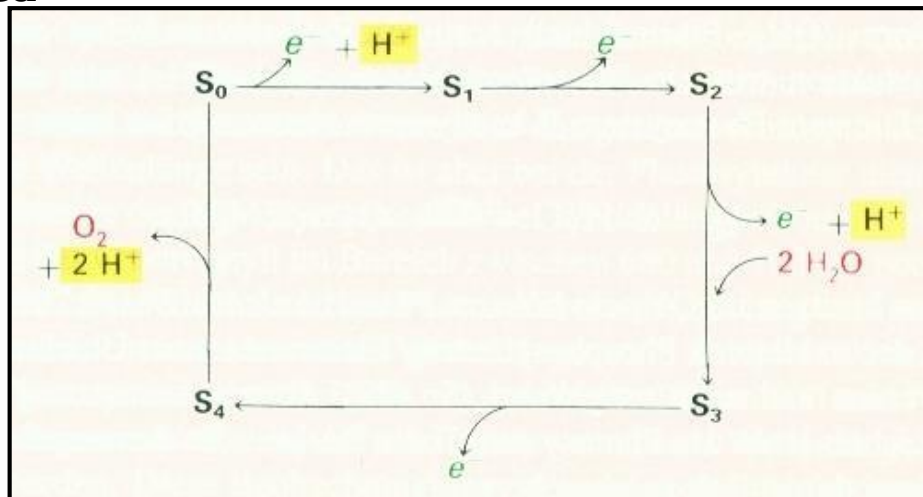
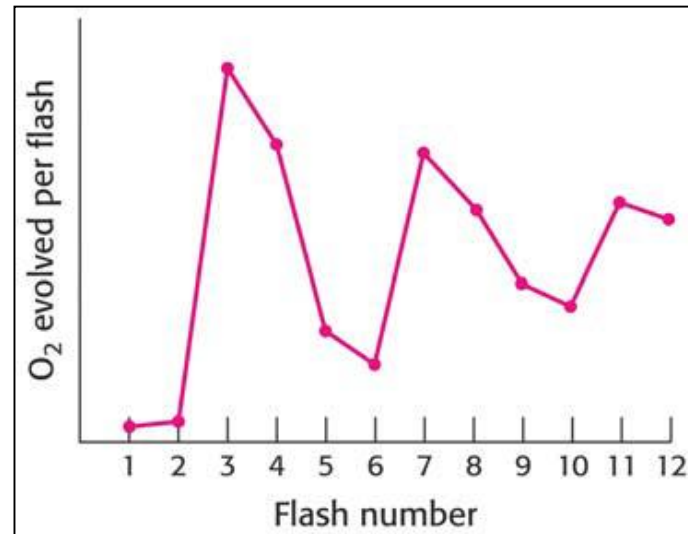
10. Biochemistry of Transition Metals

Photosynthesis: Oxidation State of Mn-Cluster (Oxygen Evolving Center OEC)

d) Photolysis experiments

Findings:

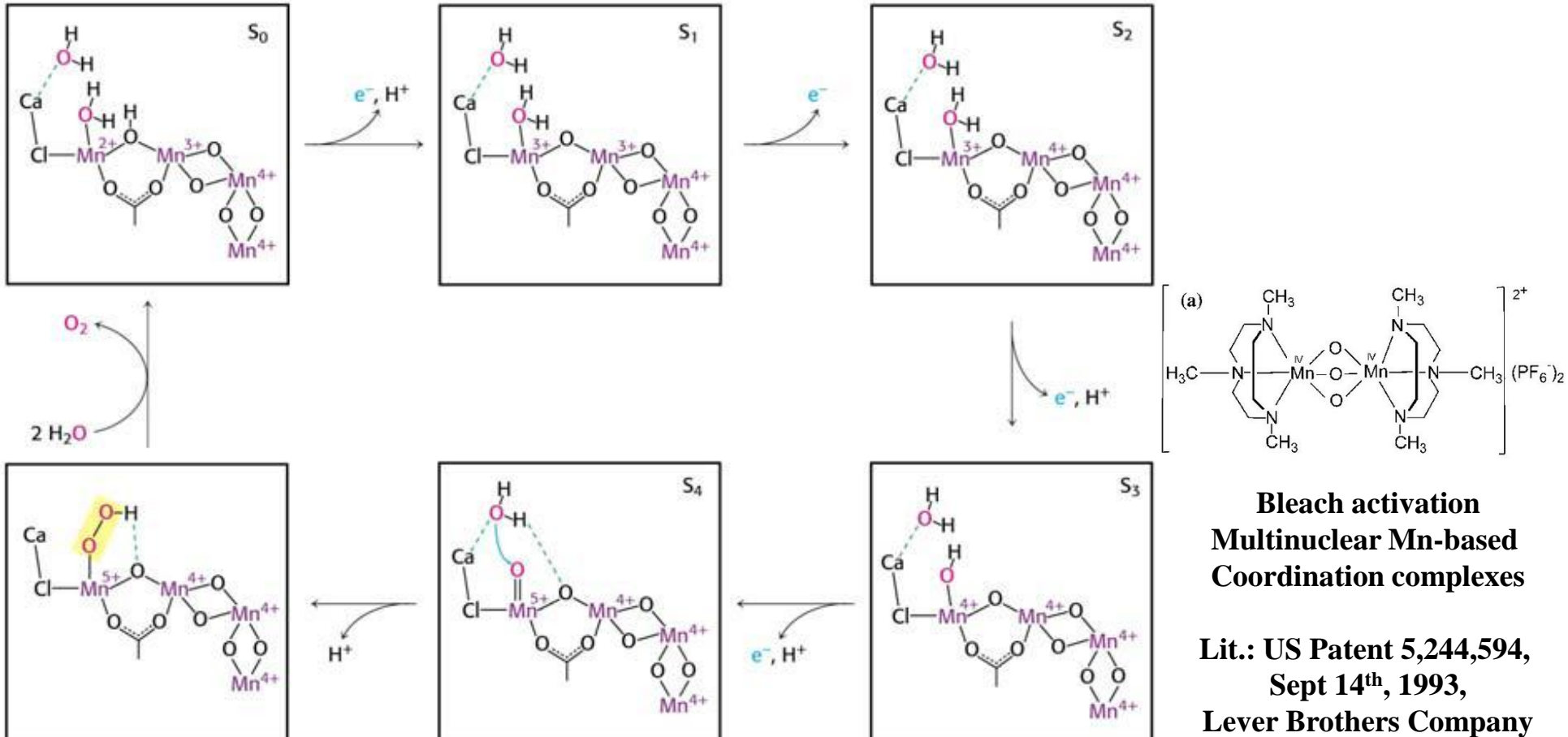
- Oxygen evolution is oscillating ($S_0 \dots S_4$)
- Oxidation occurs upon excitation of P680 to $P680^*$
- $P680^*$ oxidises the OEC (via a Tyr-side chain)
- Mn^{2+} is gradually oxidised to Mn^{5+}
- O_2 is probably released from an in-situ formed peroxy-unit
(hints from structures of model complexes)



10. Biochemistry of Transition Metals

Photosynthesis: Mechanism of Water Cleavage in the Mn-Cluster

→ The Mn-cluster acts as a “homogeneous catalyst”: Models exist for oxo-bridged Mn⁴⁺ ions



10. Biochemistry of Transition Metals

Photosynthesis: Formation of the Biochemical Energy Source ATP by a Electrochemical Gradient



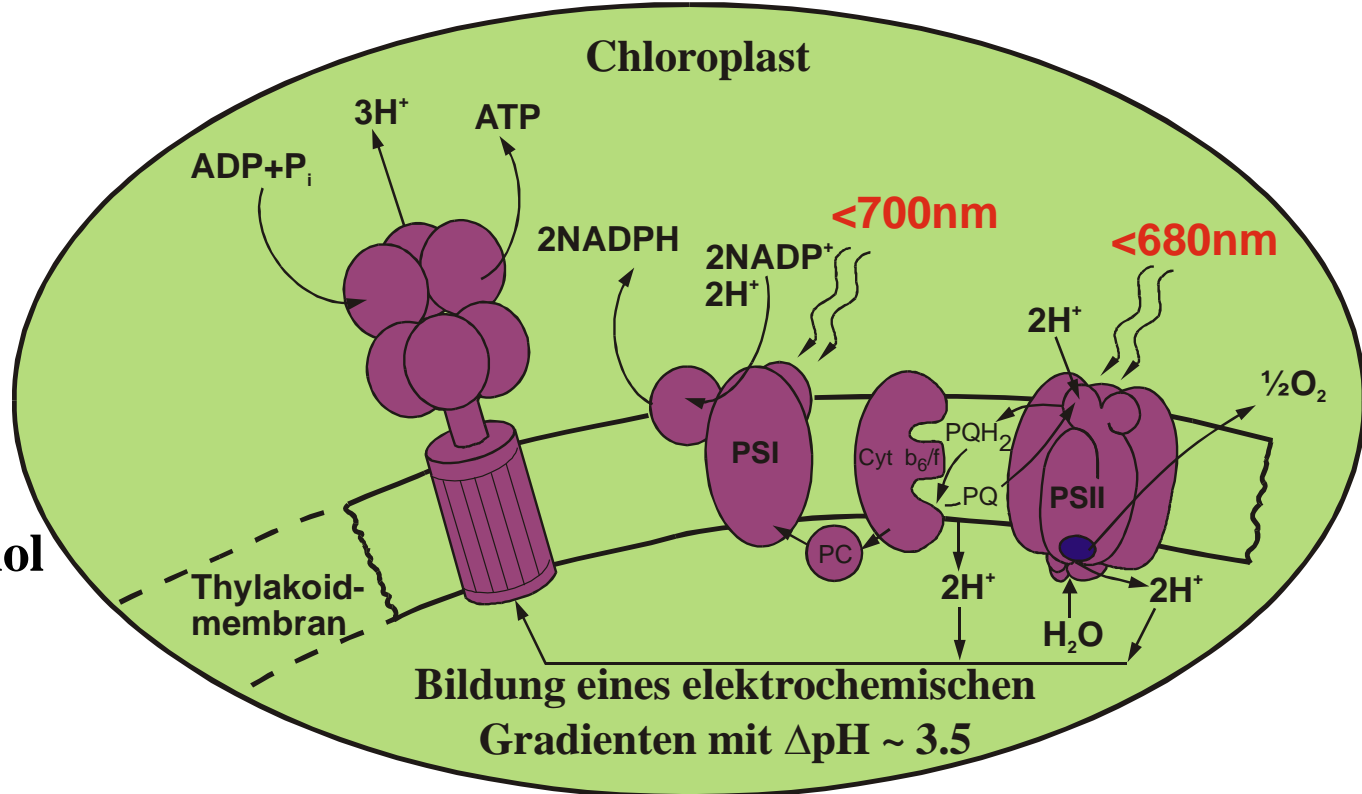
$$\rightarrow \Delta \text{pH} \sim 3.5$$

$$\begin{aligned}\Delta E &= E_1 - E_2 \\ &= 0.059/z \cdot \log(c_1/c_2) \\ &= 0.059 \cdot \Delta \text{pH} \\ &= 0.207 \text{ V}\end{aligned}$$

$$\Delta G = -n \cdot F \cdot \Delta E = -20 \text{ kJ/mol}$$

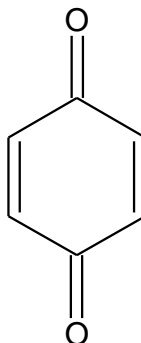
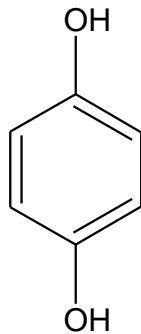


$$\Delta G = 30.5 \text{ kJ/mol}$$



10. Biochemistry of Transition Metals

Photosynthesis: The mobile Electron Transport System Quinone / Hydroquinone



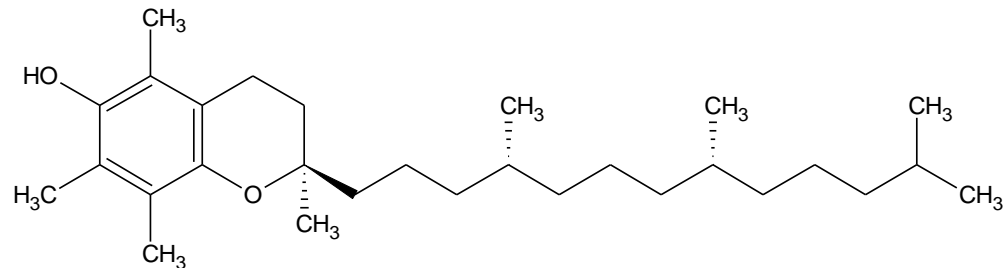
$$E = E^0 + \frac{0.059}{z} \lg \frac{[\text{quinone}][\text{H}_3\text{O}]^2}{[\text{hydroquinone}]} = E^0 + \frac{0.059}{z} \lg \frac{[\text{quinone}]}{[\text{hydroquinone}]} - 0.059 \cdot \text{pH}$$

Biochemically important quinone/hydroquinone-systems

Plastoquinone

Ubiquinone (coenzyme Q)

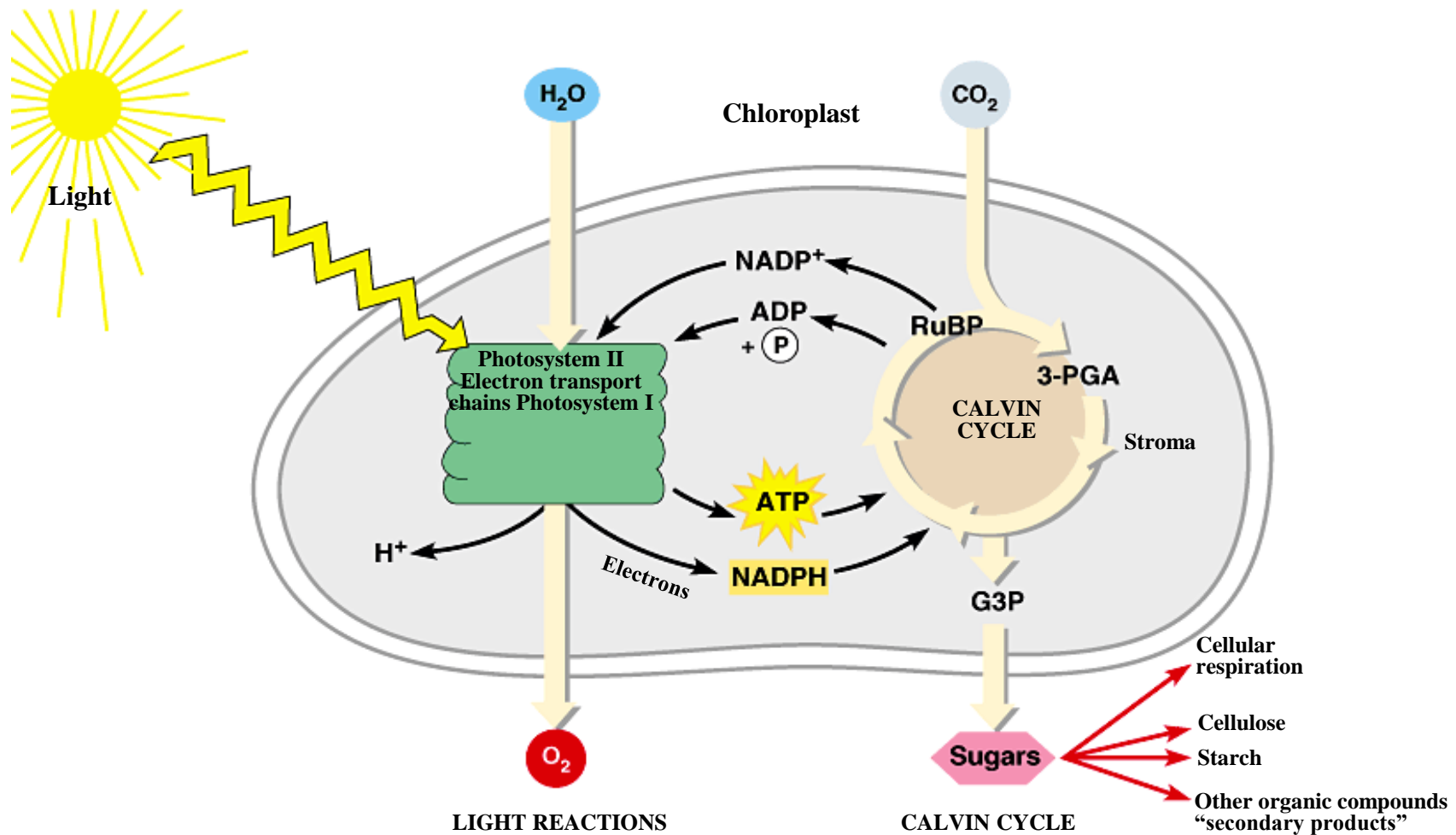
Tocopherol (vitamin E) →



⇒ Electron transport between PSI and PSII

10. Biochemistry of Transition Metals

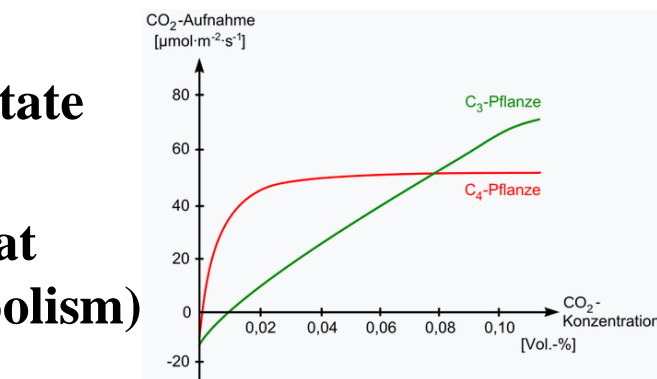
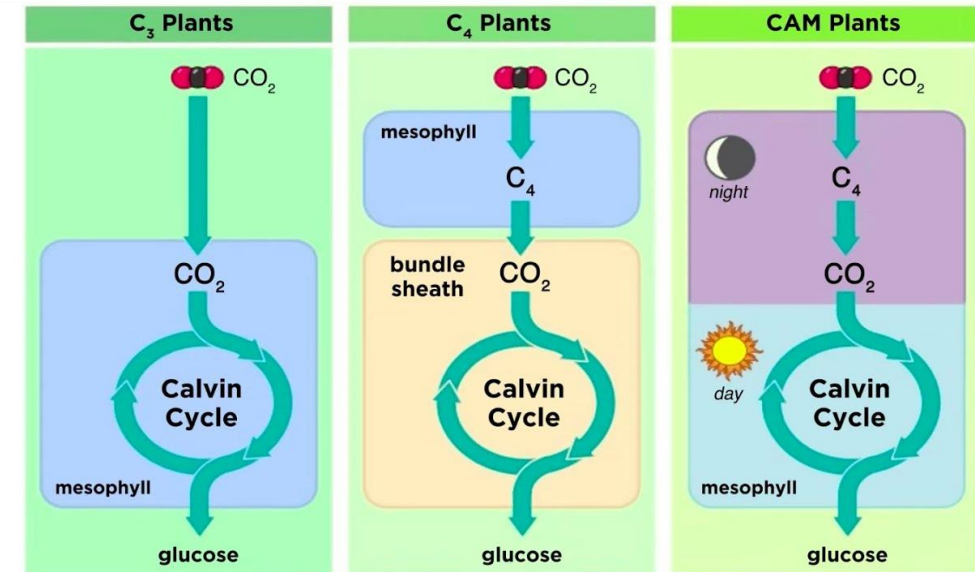
Photosynthesis: Summary of All Relevant Processes



10. Biochemistry of Transition Metals

Photosynthesis: The Dark Reaction (Calvin Cycle)

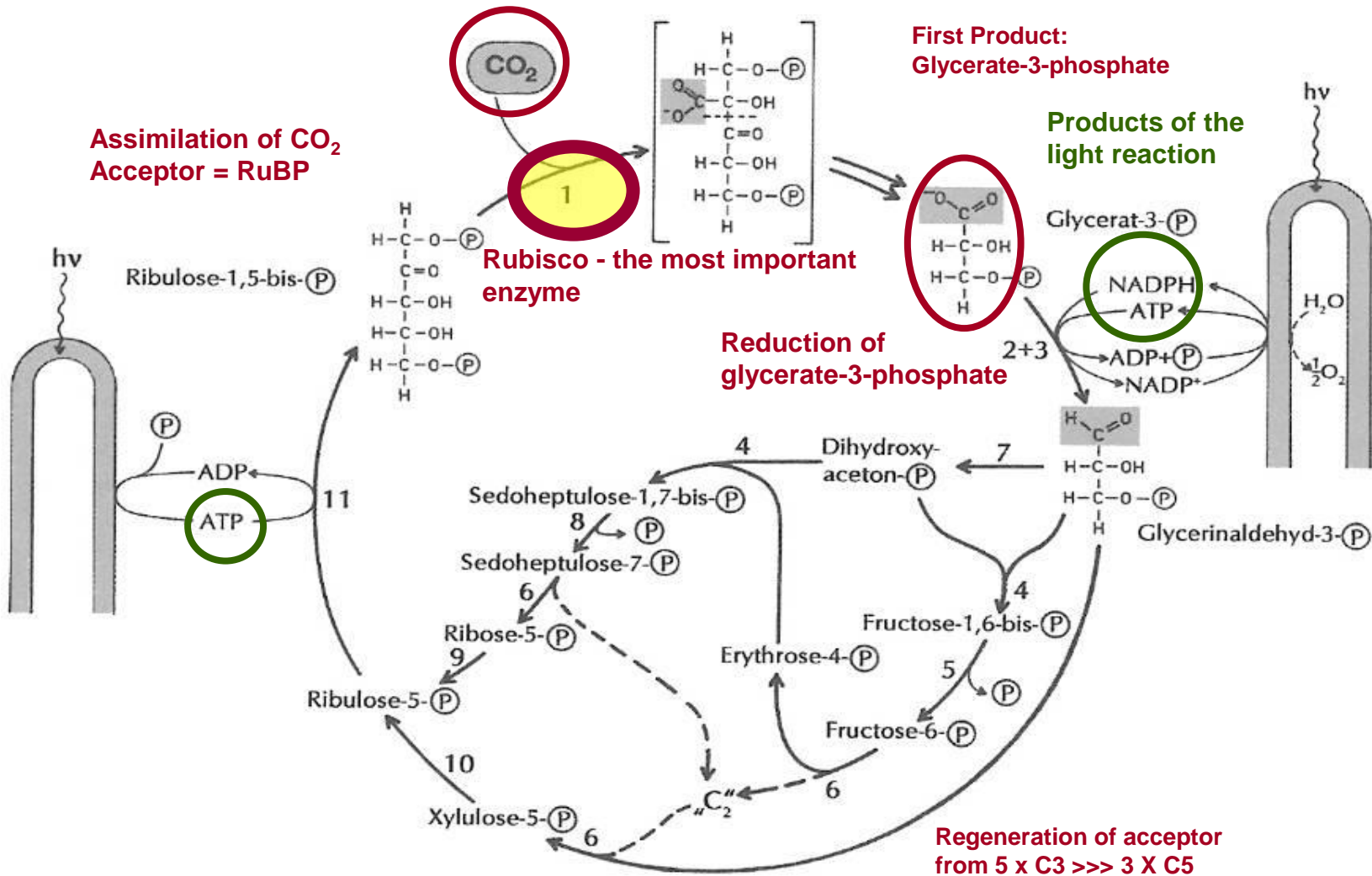
- Takes place in the stroma of the chloroplasts
- CO₂ fixation by
 - C₃-plants, e.g. elm (90%)
Ribulose-1,5-biphosphate
→ 2 Glycerat-3-phosphate
 - C₄-plants, e.g. corn (2%)
Phosphoenolpyruvate → Oxalacetate
 - CAM-plants, e.g. succulents (8%)
Phosphoenolpyruvate → Oxalacetate
(CAM = Crassulacean Acid Metabolism)



- Follows different biochemical synthesis routes: Spatial and temporal separation

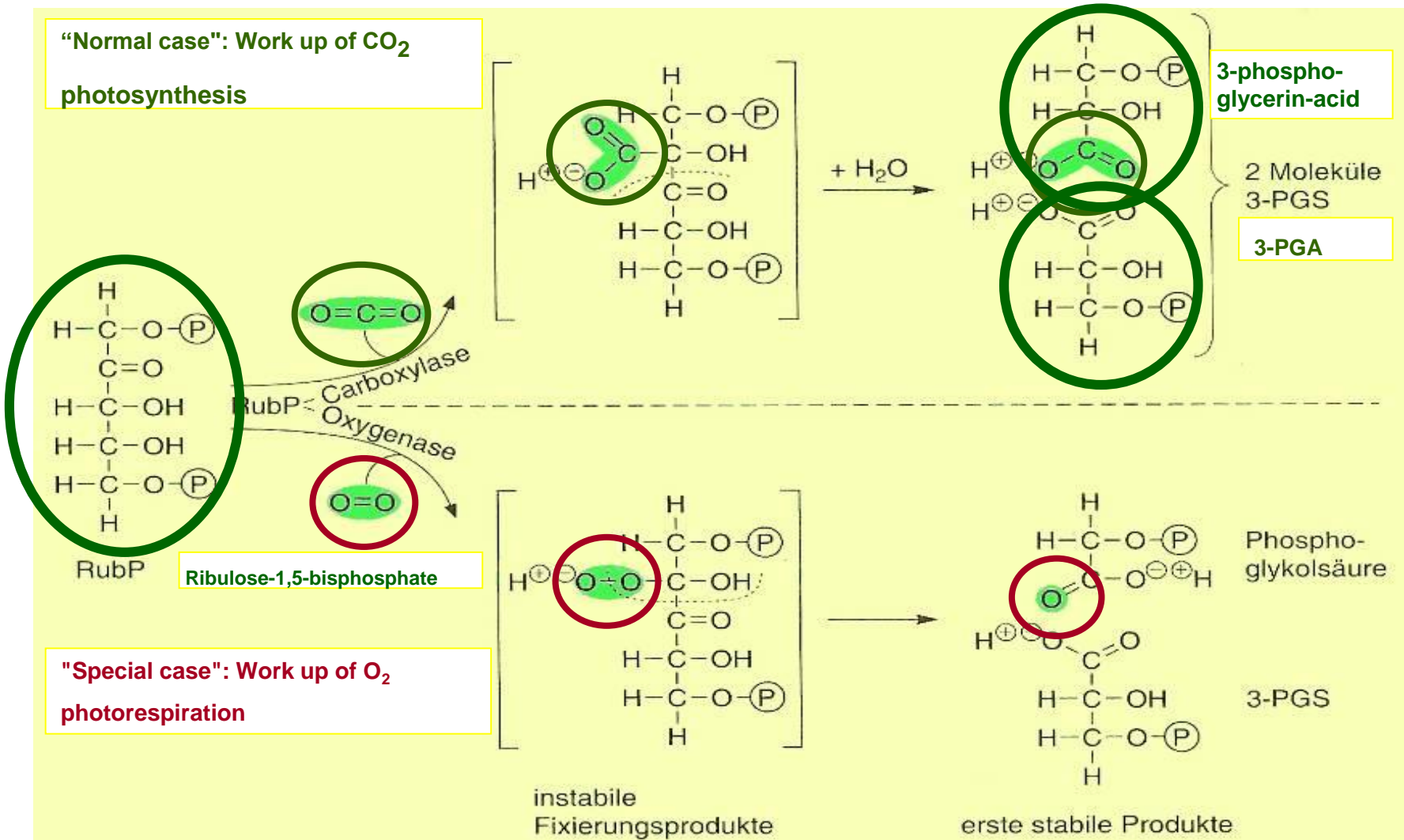
10. Biochemistry of Transition Metals

Photosynthesis: The Dark Reaction (Calvin Cycle)



10. Biochemistry of Transition Metals

RuBisCo – The Most Important Enzyme



10. Biochemistry of Transition Metals

Photorespiration

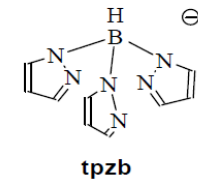
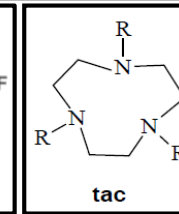
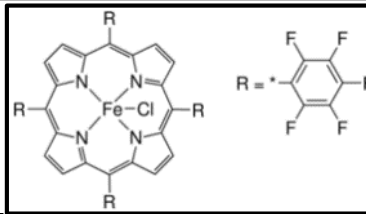
- RuBisCo catalyses two enzymatic reactions
 - Carboxylation
 - Addition of CO_2 to RuBP
 - Preferred under normal conditions
 - Photorespiration
 - Oxidation of RuBP by addition of O_2
 - Preferred when stoma is closed
 - Takes place when CO_2 -partial pressure is low and that of O_2 is high
- CO_2 and O_2 compete for the binding to RuBP!

10. Biochemistry of Transition Metals

The Iron Group

Iron

Fe ²⁺	[Ar]3d ⁶	moderately reductive agent
Fe ³⁺	[Ar]3d ⁵	relatively redox stable but kinetically labile
Fe ⁴⁺	[Ar]3d ⁴	[L ₂ Fe ₂ (cat) ₂ (μ-N)] ⁺ are strong oxidising agents
Fe ⁵⁺	[Ar]3d ³	[Fe ^{III} (porphyrin)N ₃] → [Fe ^V (porphyrin)N] + N ₂
Fe ⁶⁺	[Ar]3d ²	Na ₂ FeO ₄ , K ₂ FeO ₄ , BaFeO ₄ are strong oxidising agents Fe ³⁺ + 12 H ₂ O ⇌ FeO ₄ ²⁻ + 8 H ₃ O ⁺ + 3 e ⁻ E ⁰ = +2.20 V



Ruthenium

Very scarce

Ru ²⁺	[Kr]4d ⁶ l.s.	Ruthenium(II)-complexes as antenna in Grätzel cells
Ru ³⁺	[Kr]4d ⁵ l.s.	Ruthenium(II/III)-complexes as cancerostatics
Ru ⁴⁺	[Kr]4d ⁴ l.s.	RuO ₂
Ru ⁸⁺	[Kr]	RuO ₄ yellow, strongly oxidising

Osmium

Very scarce

Os ⁸⁺	[Xe]4f ¹⁴	OsO ₄ yellow, extremely toxic (oxidises 1,2-dioles)
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10. Biochemistry of Transition Metals

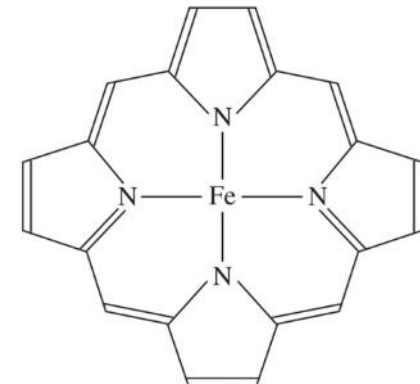
The Iron Group

Bioinorganic Chemistry of Iron

Iron is essential for the oxygen transport and for many electron transfer reactions

Heme-protein (iron-porphyrin-complexes)

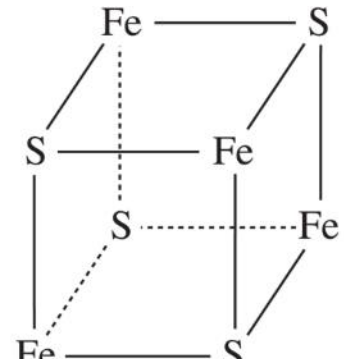
		$E^0 [V]$
• Hemoglobin	O ₂ transport	0.17
• Myoglobin	O ₂ storage	0.05
• Cytochrome (a,b,c)	Electron transfer	0.04 – 0.40
• Oxygenases	Oxygenation	
• Oxidases	O ₂ reduction to O ₂ ⁻ , O ₂ ²⁻ , O ²⁻	
• Peroxidases	Oxidation with H ₂ O ₂	
• Catalases	H ₂ O ₂ disproportionation to H ₂ O and O ₂	



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Non-heme-protein (iron-sulphur-cluster)

	$E^0 [V]$
• Rubridoxin	-0.06
• Ferredoxin	-0.42
• Nitrogenases	N ₂ reduction to NH ₃
• Transferrin	Iron transport
• Ferritin	Iron storage



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10. Biochemistry of Transition Metals

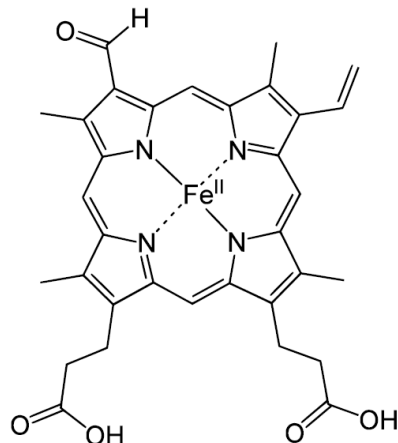
The Iron Group

Fe (and Cu): The basis of Oxygen transport proteins

Non-cellular

Chlorocruorine Annelida **18 kDa**

Erythrocytochrome Annelida, arthropoda **several MDa**

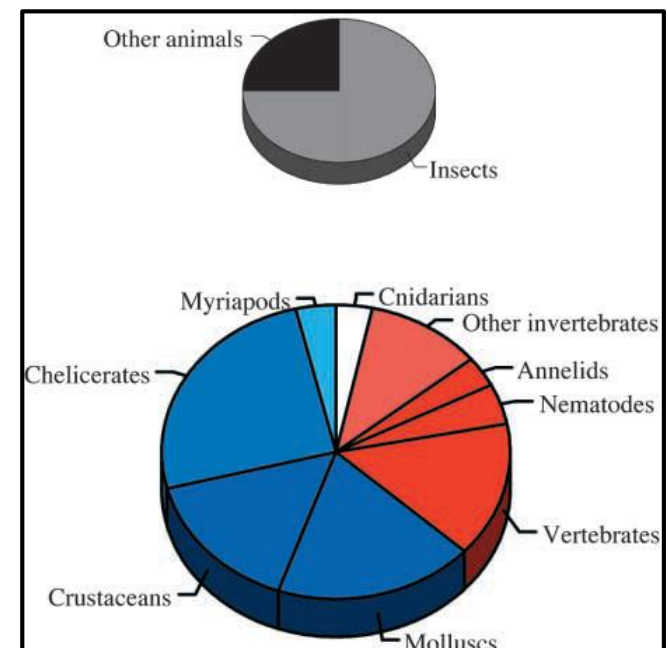


Cellular

Hemerythrins Annelida (sipunculidae) **8 x 13-14 kDa**

Myoglobin Vertebrata **17 kDa**

Hemoglobin Vertebrata **65 kDa**

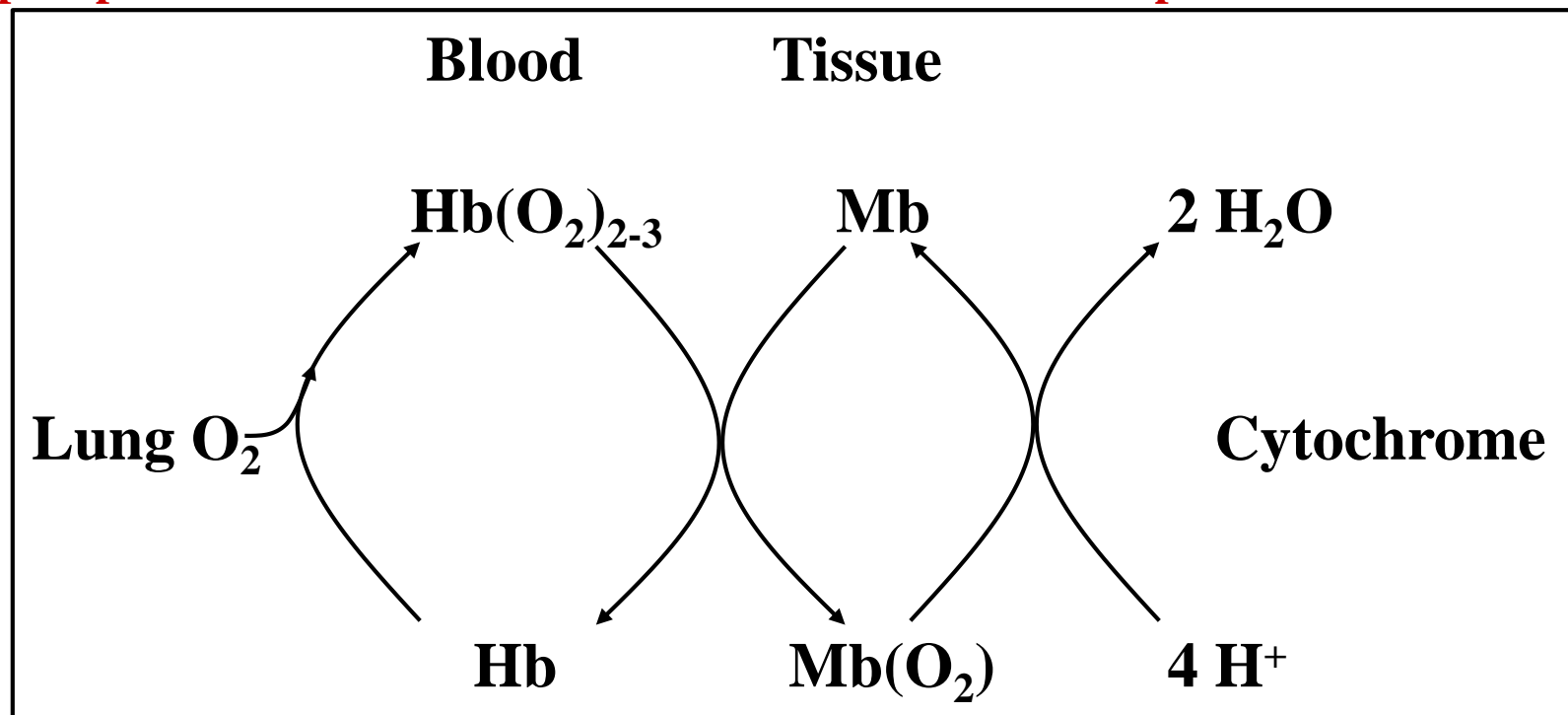


10. Biochemistry of Transition Metals

Fe-proteins: From Oxygen Transport to the Respiratory Chain

The adult human contains about 4 g Fe, with 75% thereof being bound in erythrocytes of the hemoglobin (as in other vertebrates)

Oxygen transport chain (lung - blood - tissue) → respiratory chain (mitochondria)
Fe-transport protein **Fe-redox protein**



10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins: Hemoglobin and Myoglobin

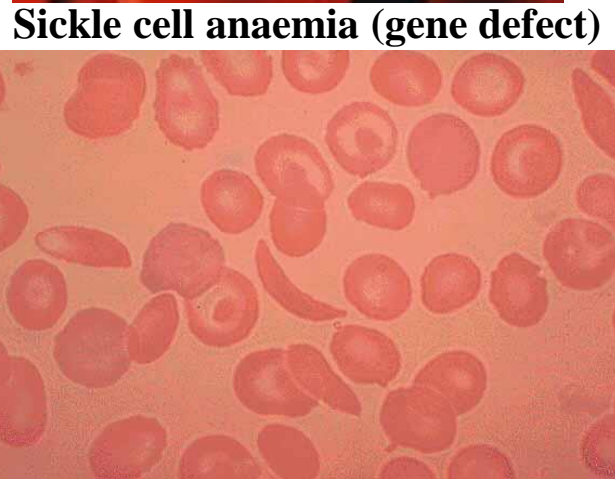
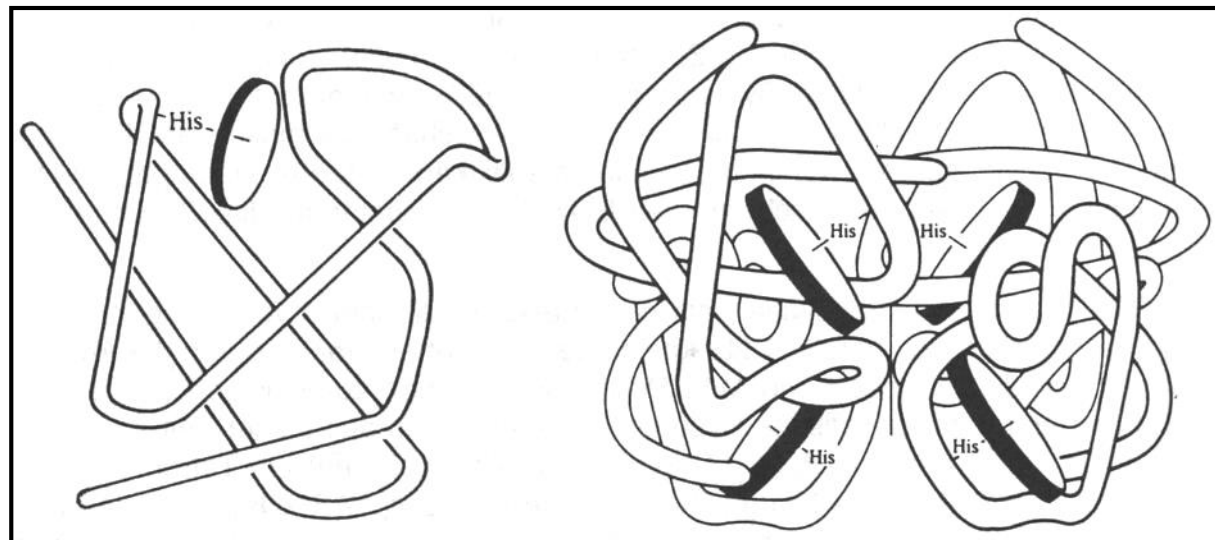
In vertebrates and some arthropods

Tetramer $\alpha_2\beta_2$ (Hb) and monomer (Mb)
with a single Fe²⁺ ion per moiety

Erythrocytes ~ $3 \cdot 10^8$
hemoglobin molecules

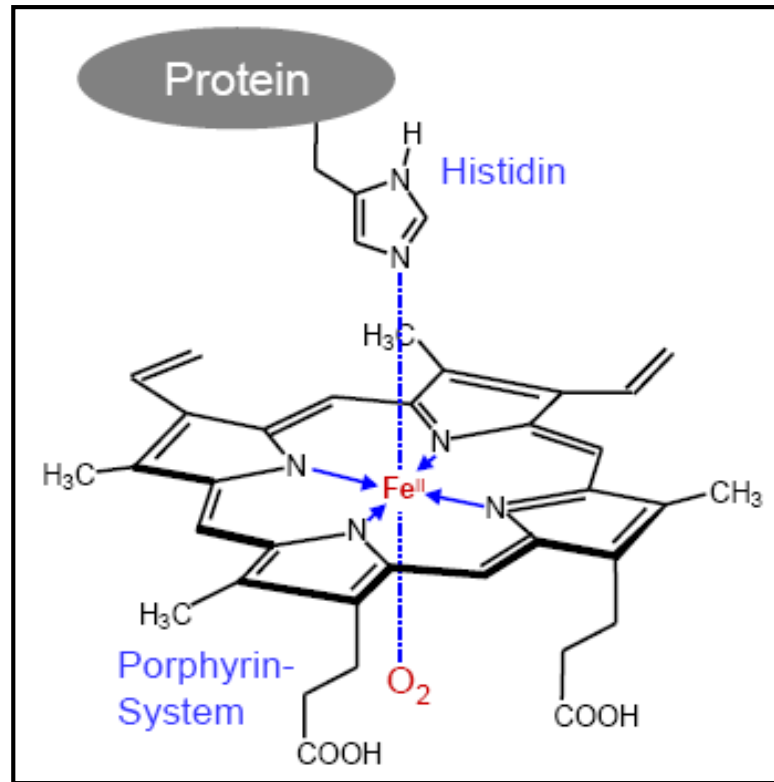
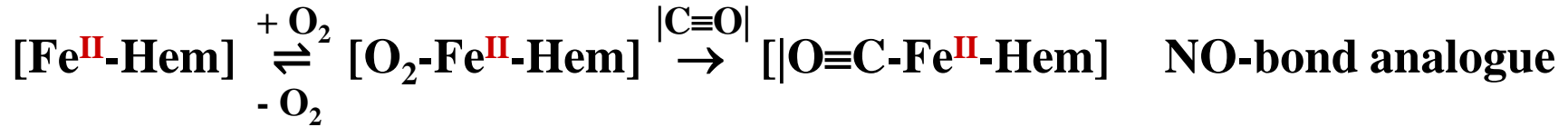


Structure of myoglobin (Mb) and hemoglobin (Hb)

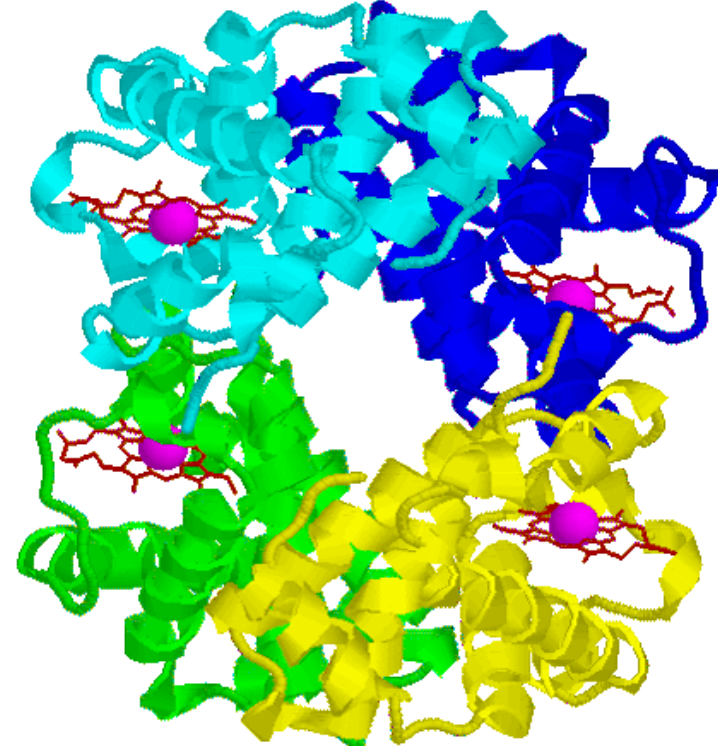


10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins: Hemoglobin



Hemoglobin is a heterotetramer $\alpha_2\beta_2$ ($M = 65 \text{ kD}$)



10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins: Hemoglobin

Function and structure of heme

Oxyhemoglobin

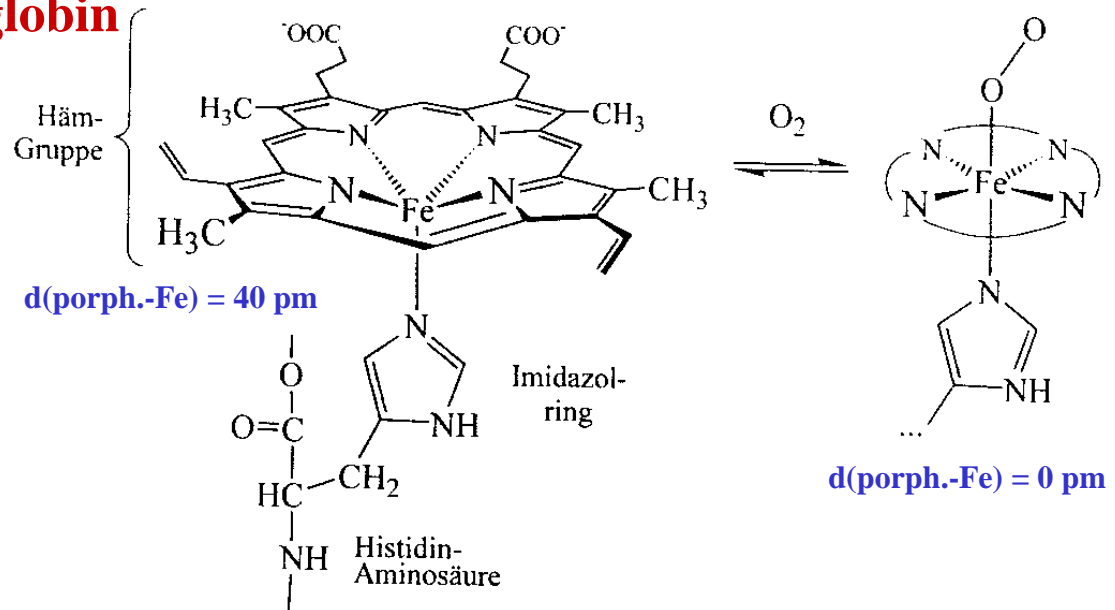
Fe²⁺, [Ar]3d⁶ l.s., diamagnetic

r = 55 pm ⇒ in-plane structure

Desoxyhemoglobin

Fe²⁺, [Ar]3d⁶ h.s., paramagnetic

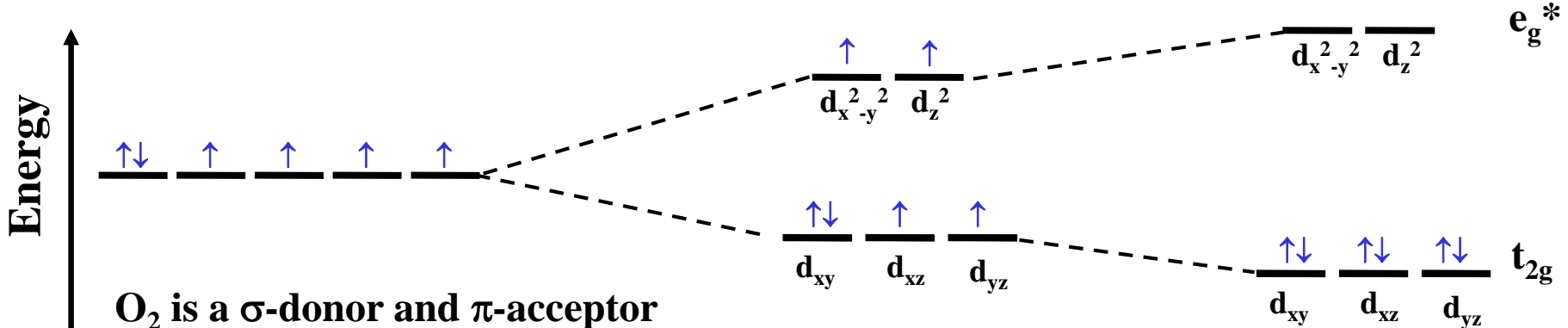
r = 78 pm ⇒ out-of-plane structure



Free Fe²⁺

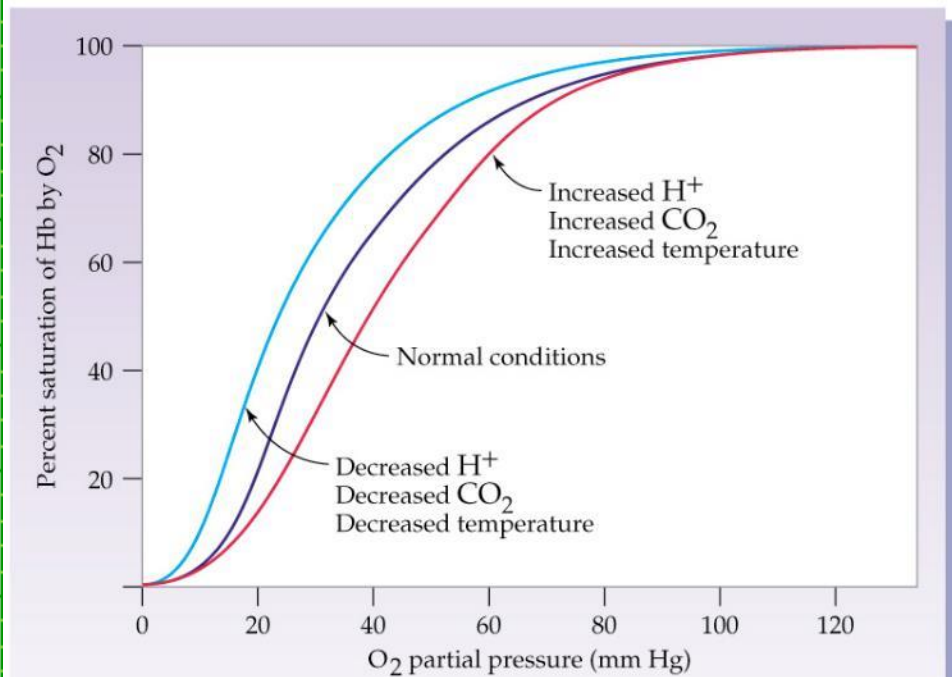
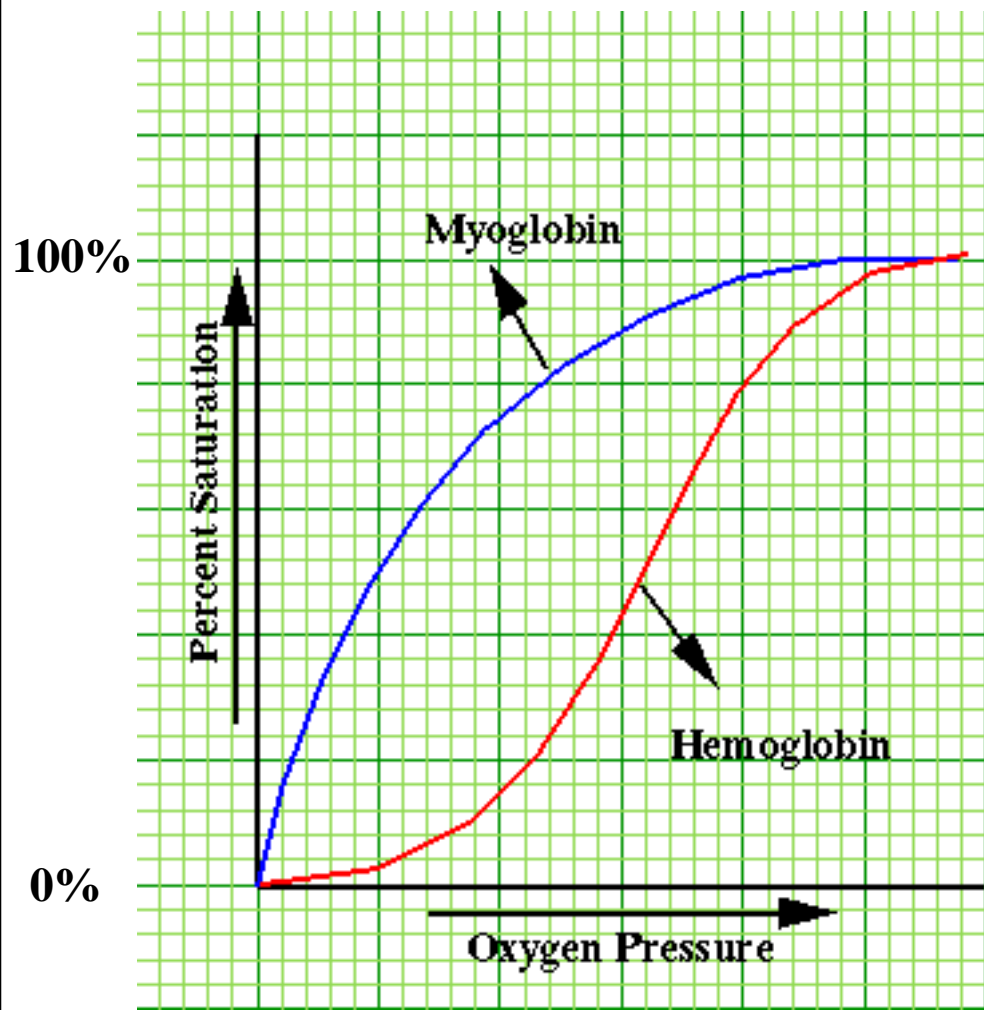
Fe²⁺ in desoxyhemoglobin

Fe²⁺ in oxyhemoglobin



10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins: Hemoglobin and Myoglobin – Oxygen Affinity

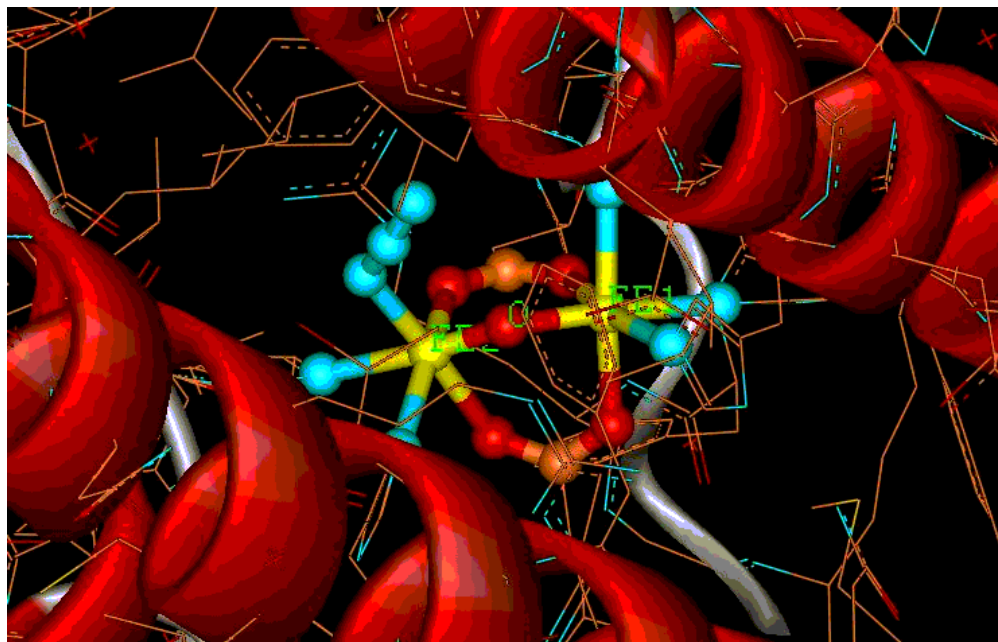


10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins: Hemerythrin

In sipunculidae (splashworms, marine)

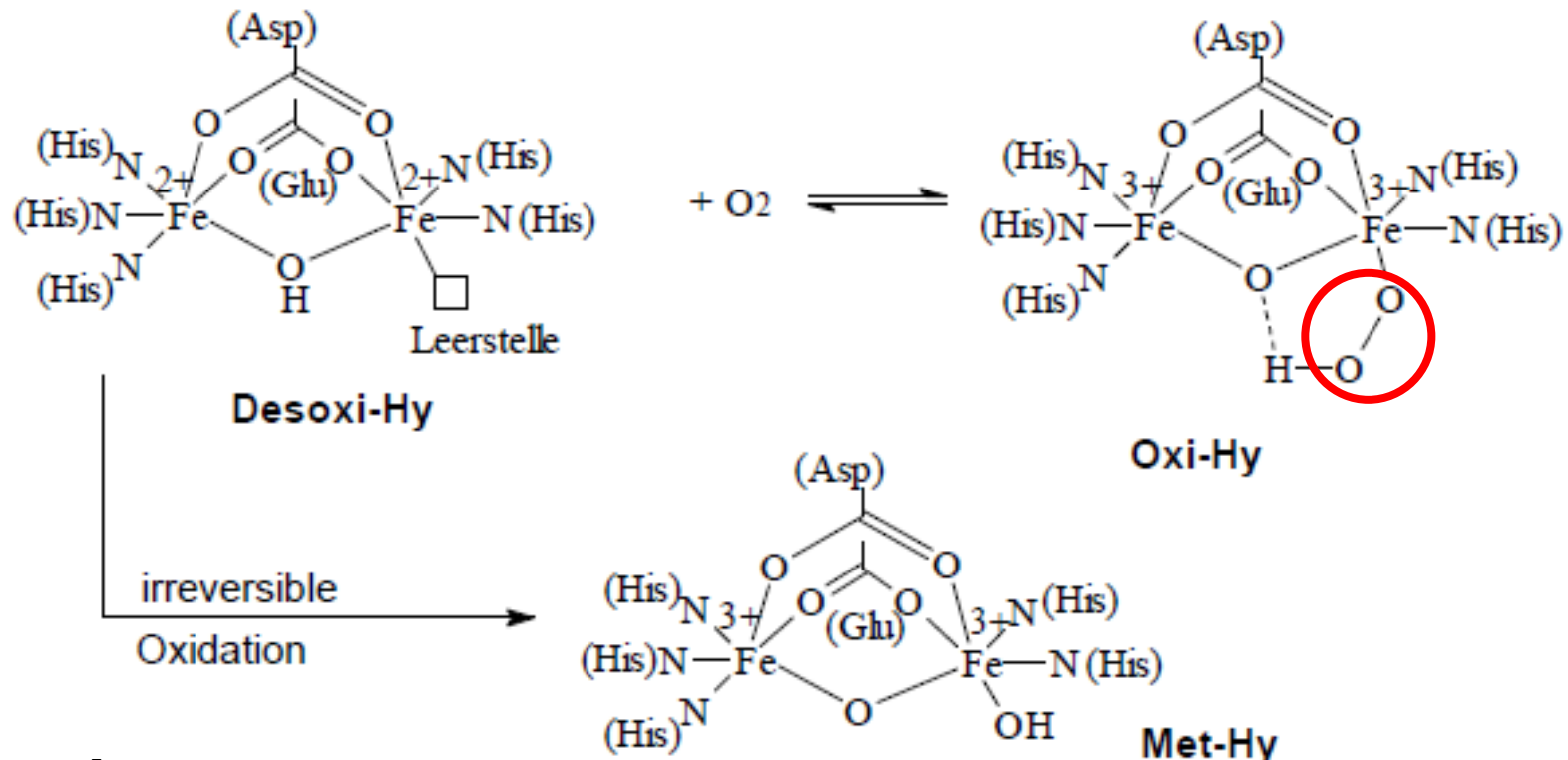
Octamers with two Fe^{2+/3+} per sub-unit (D₄-symmetry)



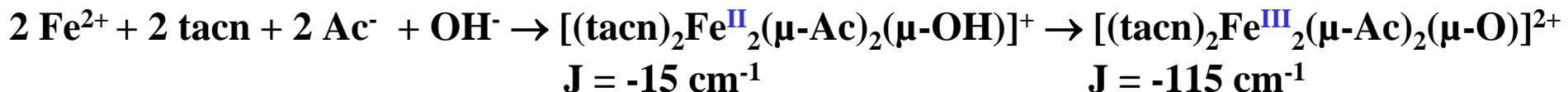
10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins: Hemerythrin (Hy)

Structure of O₂-binding unit



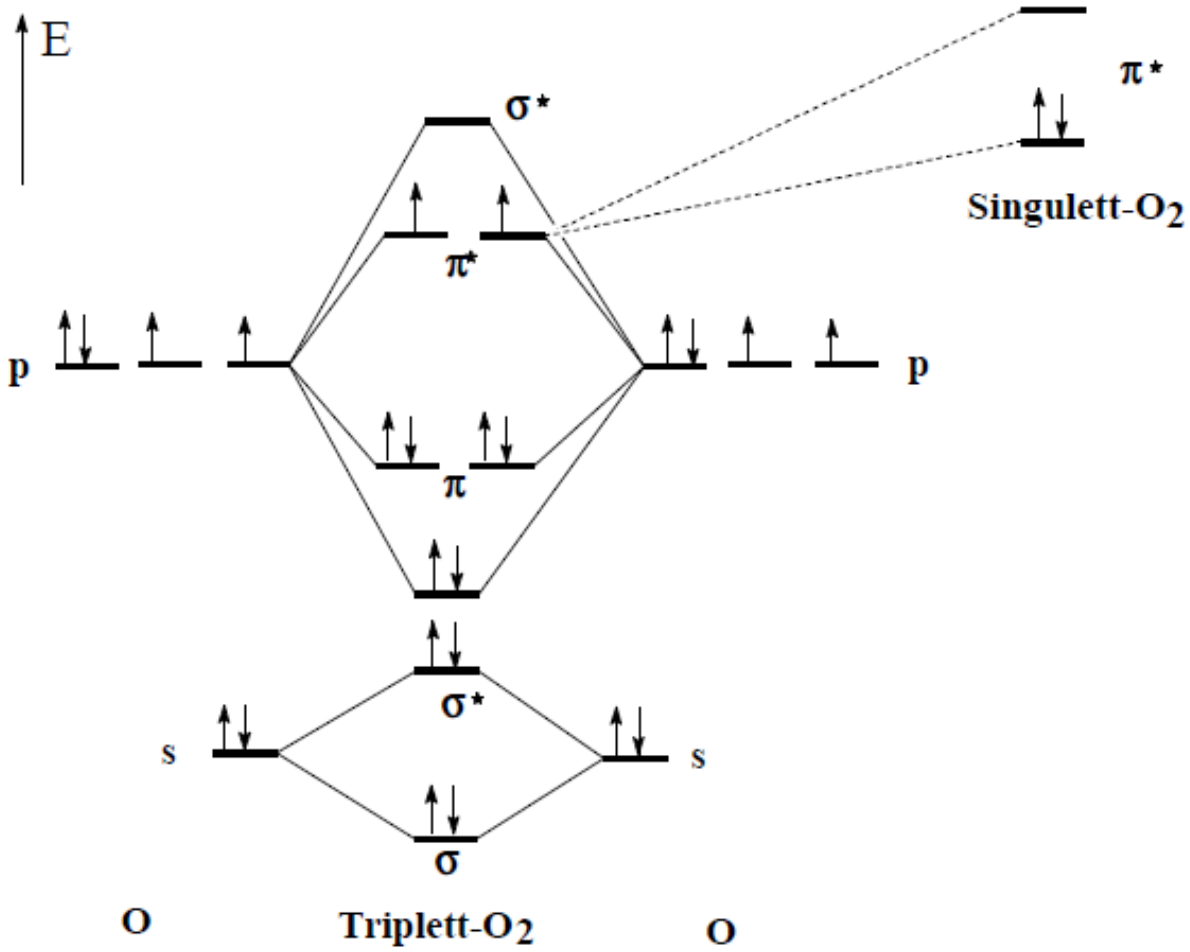
Modell complexes



10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins: Oxygen Reduction

Molecule	Bond order	d(O-O) [pm]	v(O-O) [cm ⁻¹]
O ₂ ⁺ (dioxygenyl cation)	2.5	112	1860
³ O ₂ (triplet-oxygen)	2.0	121	1555
O ₂ ⁻ (superoxide anion)	1.5	133	1145
O ₂ ²⁻ (peroxide anion)	1.0	149	770
Mb-O ₂ (oxygenated myoglobin)	~2.0	122	1107



Upon bonding of O₂ to Hb
or Mb the O₂-bond is only slightly
weakened, which means O₂ is not
being reduced during its transport

10. Biochemistry of Transition Metals

O₂-Transport-Fe-Proteins:

**Electronic configuration of Iron
in haemoglobin (Hb) and oxidised
haemoglobin (Hi)**

**Determination by Mössbauer
and EPR Spectroscopy with**

S = Total spin

δ [mms⁻¹] = Isomerie shift

Δ [mms⁻¹] = Quadrupol splitting

Table 13.2 Mössbauer spectra for haemoglobin derivatives [10]

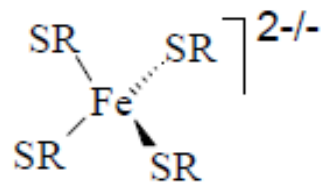
Compound*	<i>S</i>	<i>T/K</i>	Δ /(mm s ⁻¹)	δ (Fe) /(mm s ⁻¹)
HbCO	0	195	0·36	0·18
		4	0·36	0·26
Hb reduced	2	195	2·40	0·90
		4	2·40	0·91
HbNO	?	195–1·2	magnetically broadened	
HbO ₂	0	195	1·89	0·20
		77	2·19	0·26
		1·2	2·24	0·24
HiF	$\frac{5}{2}$	195–1·2	magnetically broadened	
HiH ₂ O	$\frac{5}{2}$	195	2·00	0·20
HiOH	$\frac{1}{2}$?	195	1·57	0·18
		77	1·9	0·2
HiN ₃	$\frac{1}{2}$	195	2·30	0·15
HiCN	$\frac{1}{2}$	195	1·39	0·17

* The abbreviation Hb is used for a Fe(II) haemoglobin compound and Hi for a Fe(III) haemoglobin compound.

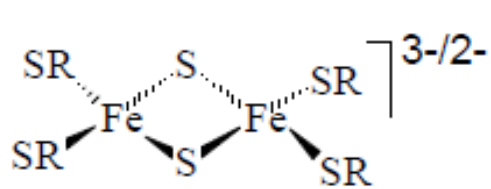
10. Biochemistry of Transition Metals

Fe-Sulphur Proteins

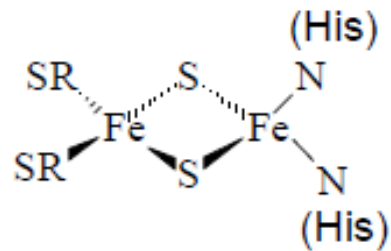
Cluster with 1, 2, 3 or 4 iron atoms as well as cys- and/or his-ligands



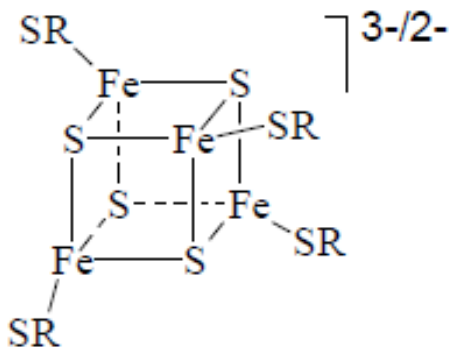
Rubredoxin



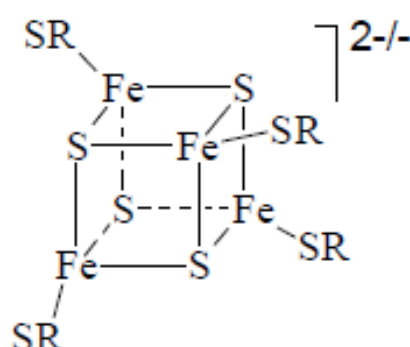
[2Fe-2S]-Ferredoxin



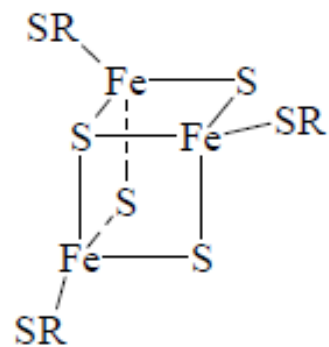
Rieske-Zentrum



[4Fe-4S]-Ferredoxin



HiPIP



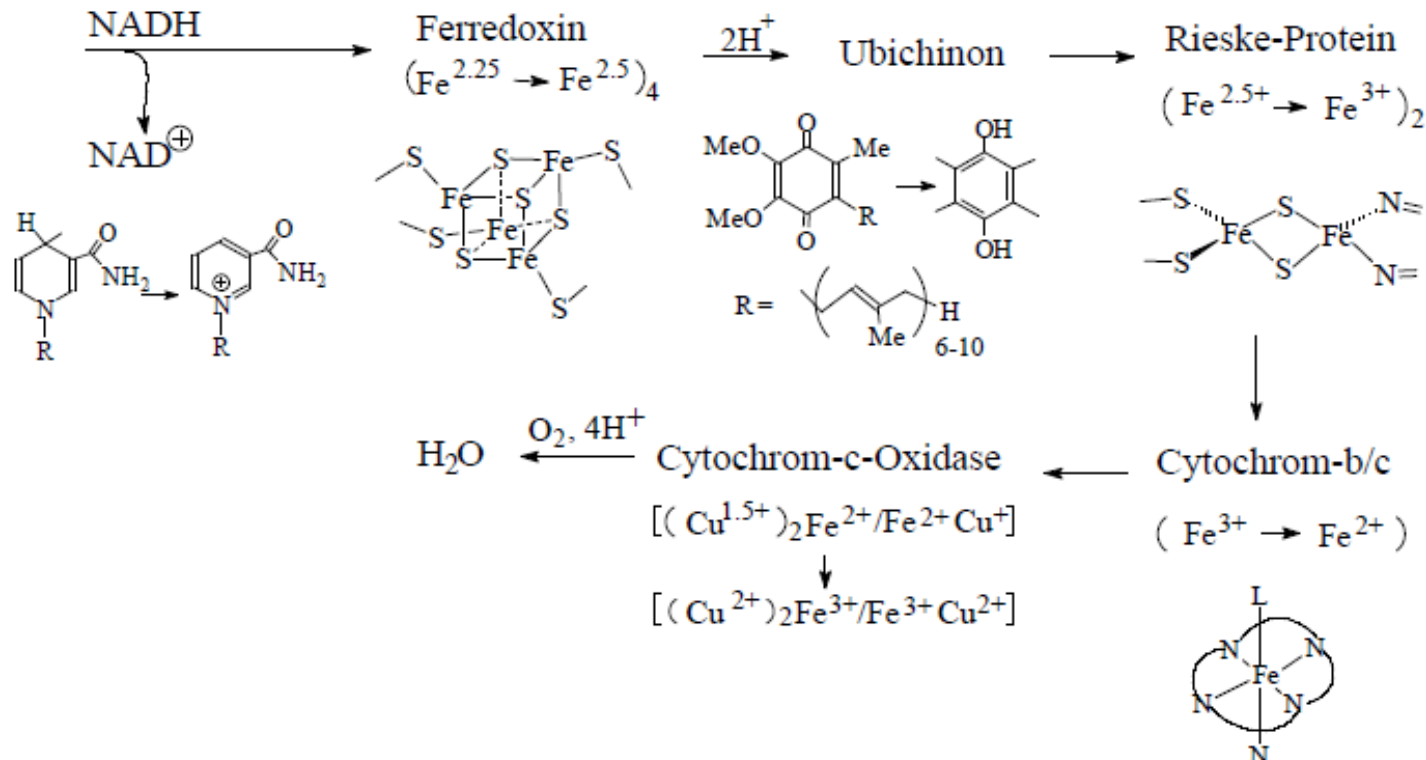
[3Fe-4S]-Ferredoxin

10. Biochemistry of Transition Metals

Fe-Redox Proteins: Mitochondrial Respiratory Chain

Balance: $O_2 + 2 \text{ NADH} + 2 \text{ H}^+ \rightleftharpoons 2 \text{ H}_2\text{O} + 2 \text{ NAD}^+$ $E^\circ = -1.13 \text{ V} \Rightarrow \Delta G = -218 \text{ kJ/mol}$

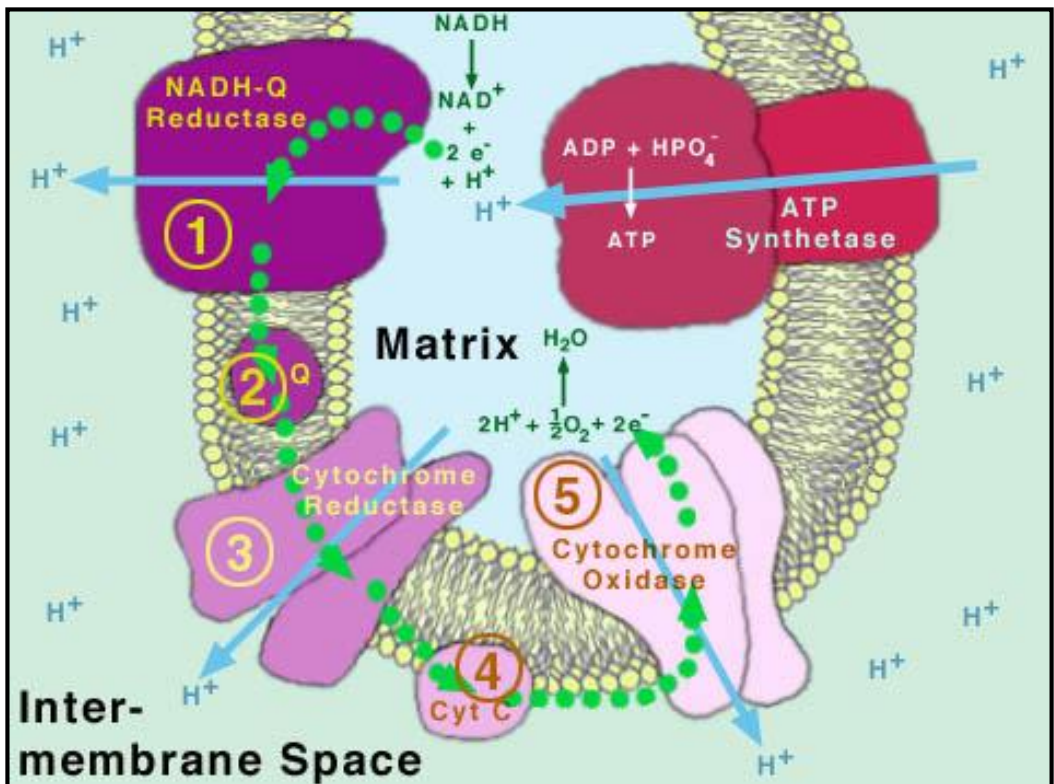
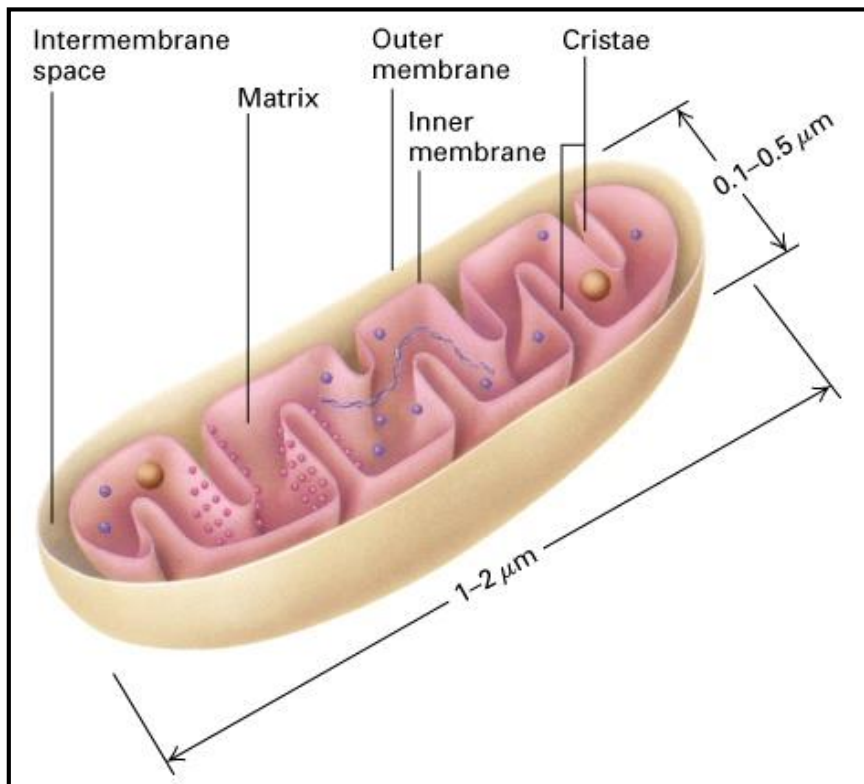
Mechanism of oxygen reduction in mitochondrial respiratory chain (5 steps)



10. Biochemistry of Transition Metals

Fe-Redox Proteins: Mitochondrial Respiratory Chain

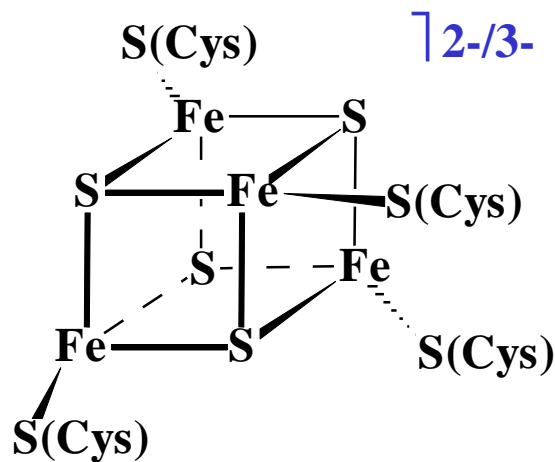
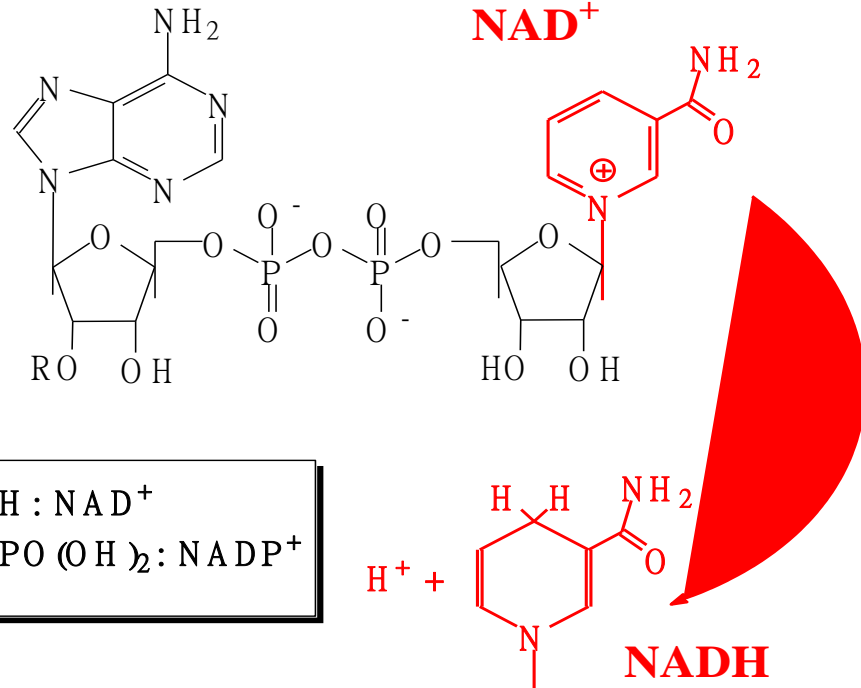
Location: Intermembrane space of eukaryotic mitochondria



10. Biochemistry of Transition Metals

Fe-Redox Proteins: Mitochondrial Respiratory Chain

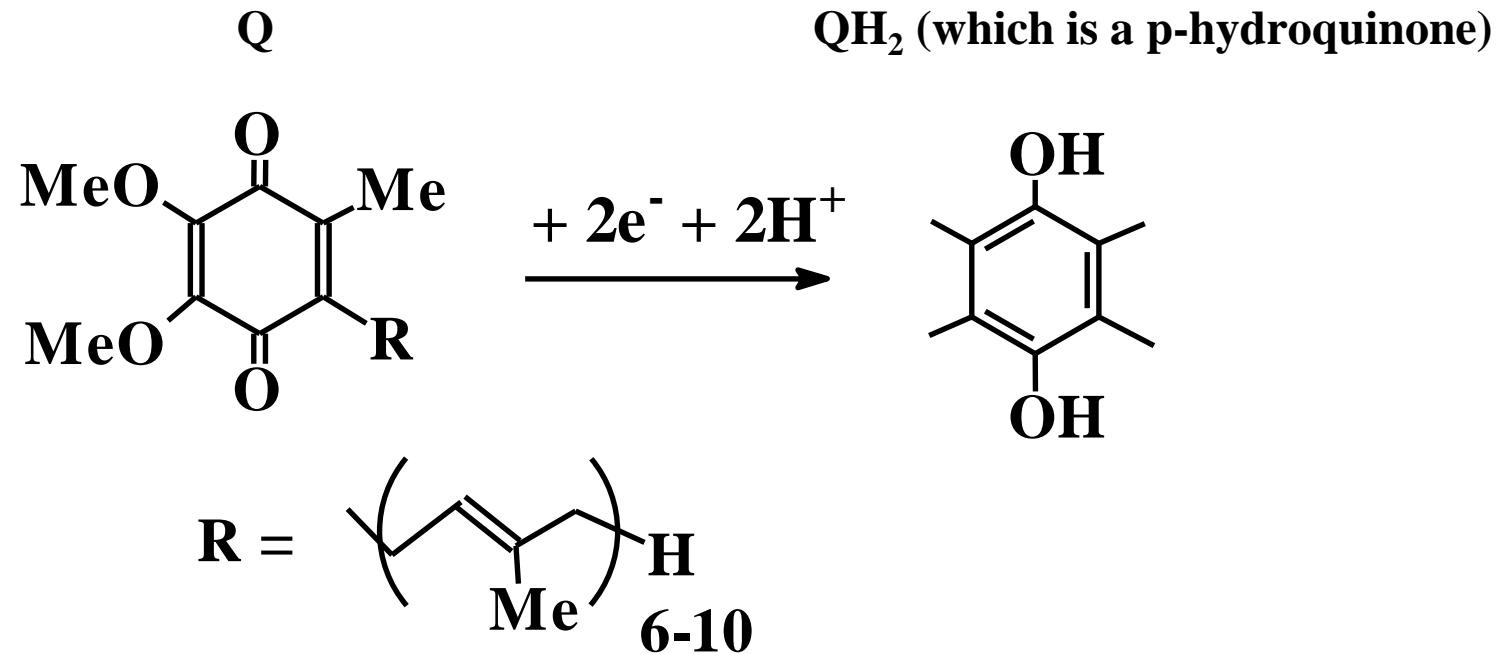
1st Step Transfer of reduction equivalents from NADH to a ferredoxin



10. Biochemistry of Transition Metals

Fe-Redox Proteins: Mitochondrial Respiratory Chain

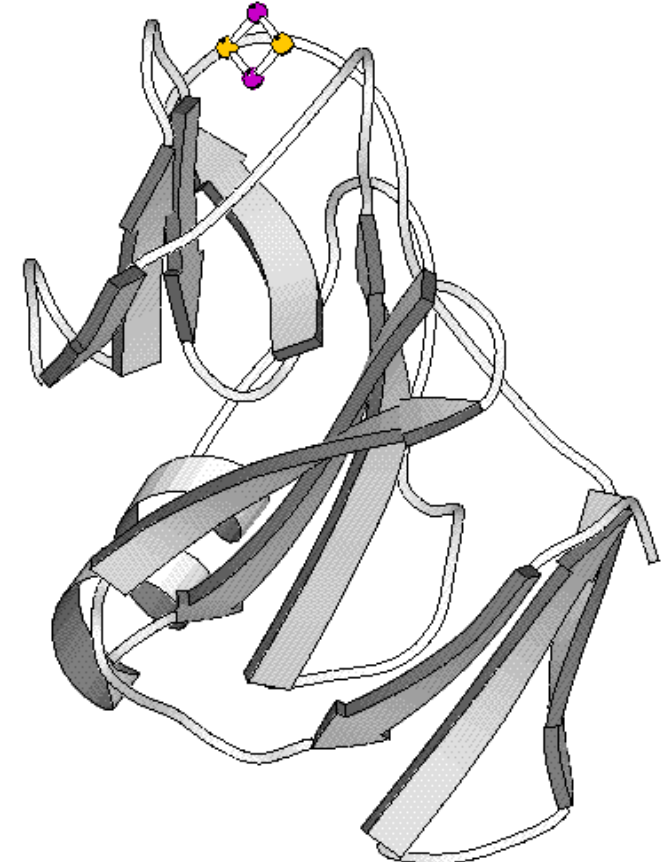
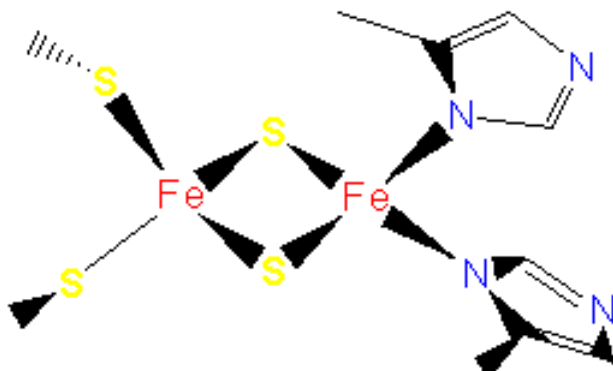
2nd Step Transfer of electrons to ubiquinone (Q, which is a quinone)



10. Biochemistry of Transition Metals

Fe-Redox Proteins: Mitochondrial Respiratory Chain

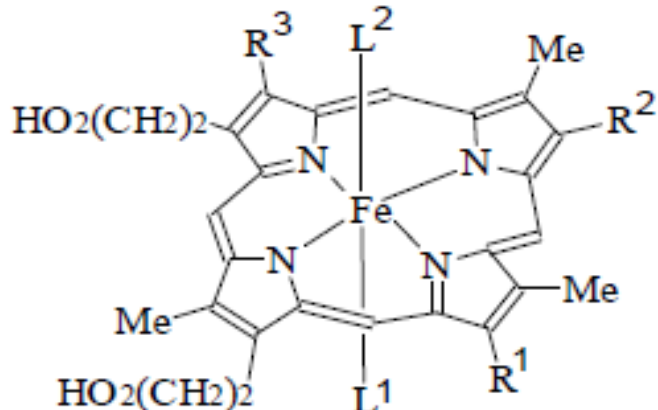
3rd Step Transfer of electrons from ubihydroquinone to a **Rieske-protein**



10. Biochemistry of Transition Metals

Fe-Redox Proteins: Mitochondrial Respiratory Chain

4th Step Transfer of electrons from Rieske-protein to cytochrome-b and -c



Cytochrom a: R¹ = Vinyl, R² = C₁₇H₃₄OH, R³ = Formyl
L¹ = L² = His

Cytochrom b: R¹ = R² = Vinyl, R³ = Me
L¹ und L² frei oder His

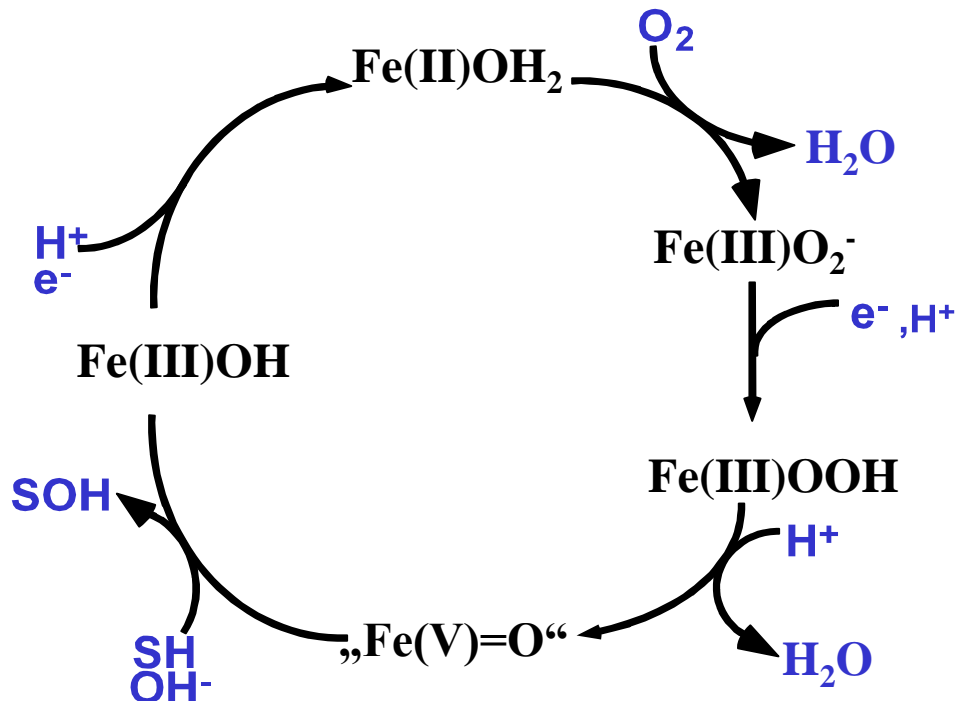
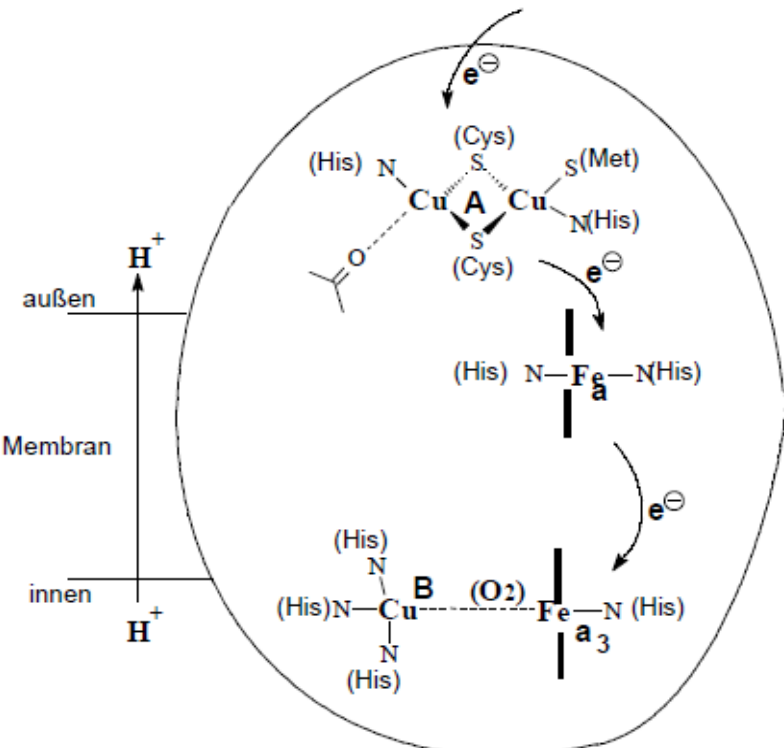
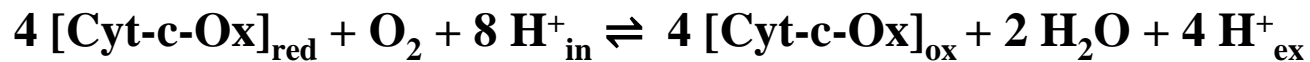
Cytochrom c: R¹ = R² = -CH(Me)-S-CH₂-C(O)NH-
R³ = ; L¹ = His, L² = Met

Cytochrom P₄₅₀: R¹ = R² = Vinyl, R³ = Me
L¹ = Cys, L² = H₂O

10. Biochemistry of Transition Metals

Fe-Redox Proteins: Mitochondrial Respiratory Chain

5th Step Transfer of electrons from cytochrome-c to cytochrome-c-oxidase and reduction of oxygen to water (using proton pumps)

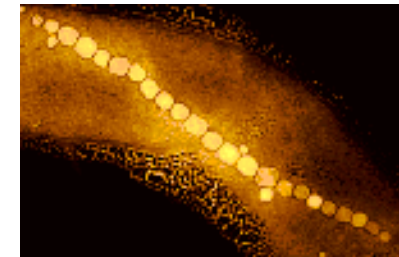
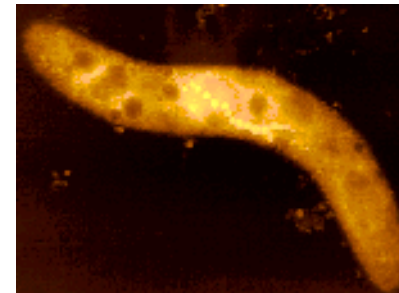


10. Biochemistry of Transition Metals

Fe-Biomineralisation and Fe-Storage

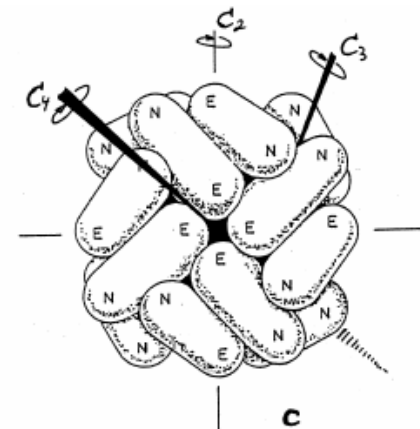
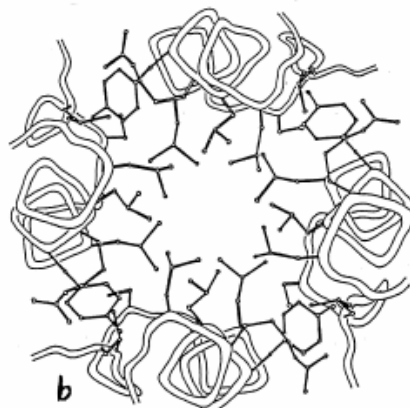
Formation of mineralised Fe-salts by living creatures

- $\alpha\text{-FeO(OH)}$ **Goethite** **limpets (radula)**
- Fe_3O_4 **Magnetite** **magnetotactic bacteria**
- Fe_3S_4 **Greigite** **magnetotactic bacteria**
- FeS_2 **Pyrite** **sulphate reducing bacteria**



Ferritins contain goethite-like material of the following composition

“ $8 \text{ FeO(OH)} \cdot \text{FeO}(\text{H}_2\text{PO}_4)$ ”



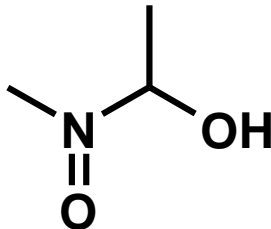
10. Biochemistry of Transition Metals

Fe-Transport: Siderophores

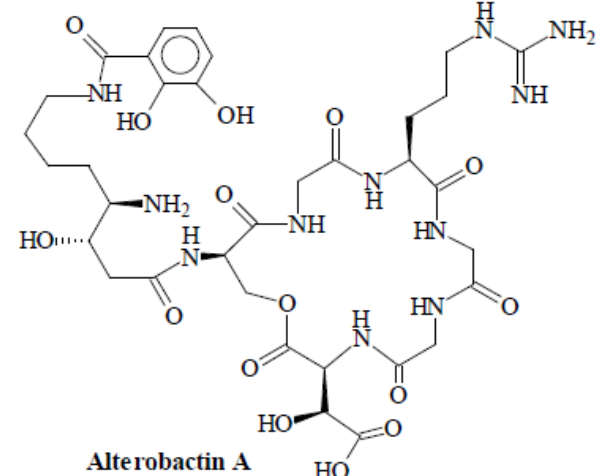
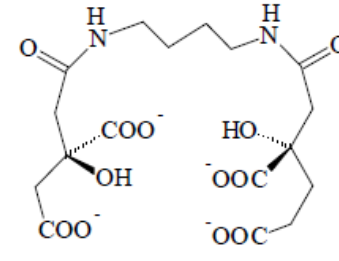
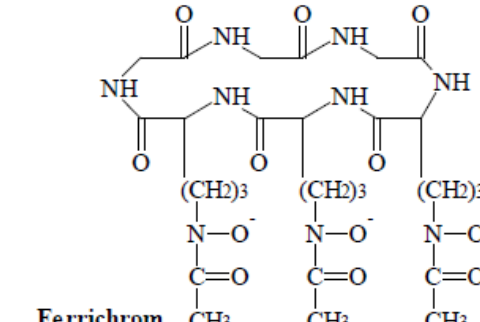
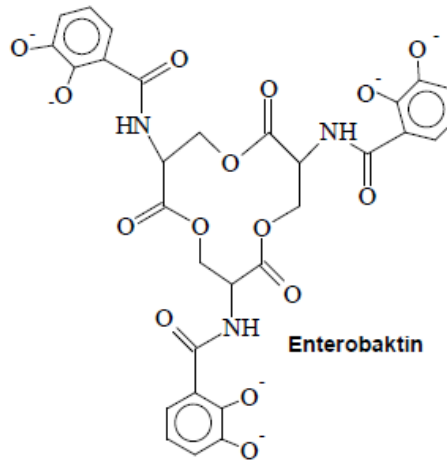
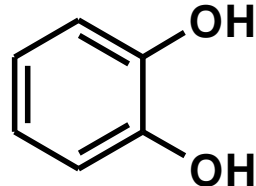
Siderophores are released by many micro organisms and plants into their surrounding aqueous medium in order to mobilise Fe^{3+} by complexation from poorly soluble iron hydroxide deposits in soil

Ligands ($\log K \sim 20 \dots 50$)

- Hydroxamates



- Catecholates



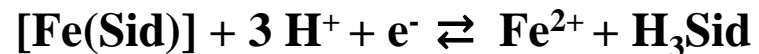
10. Biochemistry of Transition Metals

Fe-Transport: Siderophores

Equilibria:

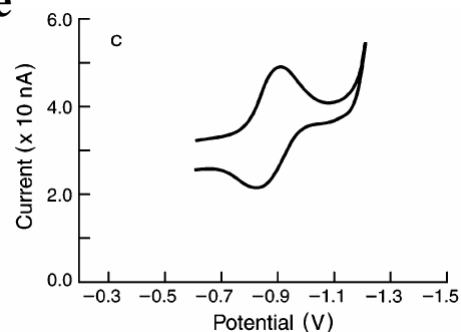


Soil



Plant cell

Siderophor	Log K	E° [V] at pH 7	Ligand type
Mugenein acid	18.1	-0.102	Carboxylate, Amino-N
Aerobactin	22.5	-0.336	Hydroxamate, Carboxylate
Coprogen	30.2	-0.447	Hydroxamate
Deferoxamin B	30.5	-0.468	Hydroxamate
Ferrichrom A	32.0	-0.448	Hydroxamate
Enterobactin	~ 49	-0.790	Catecholate
Alterobactin A	~ 51	-0.972	Catecholate Hydroxo Carboxylate



→ Analytical method: Cyclic Voltammetry (CV), which deliver voltage-current diagrams

10. Biochemistry of Transition Metals

The Cobalt Group

Cobalt

Co^+	$[\text{Ar}]3\text{d}^8$	strong reducing agent, high-spin or low-spin
Co^{2+}	$[\text{Ar}]3\text{d}^7$	weak reducing agent, high-spin or low-spin
Co^{3+}	$[\text{Ar}]3\text{d}^6 \text{ l.s.}$	relatively redox stable, very high kinetic stability, l.s.-complexes, except $[\text{CoF}_6]^{3-}$

Rhodium

Extremely scarce

Rh^{2+}	$[\text{Kr}]4\text{d}^7 \text{ l.s.}$	
Rh^{3+}	$[\text{Kr}]4\text{d}^6 \text{ l.s.}$	$[\text{Rh}(\text{bqdi})(\text{NH}_3)_4]^{3+}$ bind to DNA and are able to cleave it through photo activation (with bqdi = benzoquinone diimine or other diimines)

Iridium

Extremely scarce

Ir^{3+}	$[\text{Xe}]4\text{f}^{14}5\text{d}^6 \text{ l.s.}$	Ir^{3+} -complexes are kinetically extremely stable (OLEDs)
Ir^{4+}	$[\text{Xe}]4\text{f}^{14}5\text{d}^5 \text{ l.s.}$	IrO_2

10. Biochemistry of Transition Metals

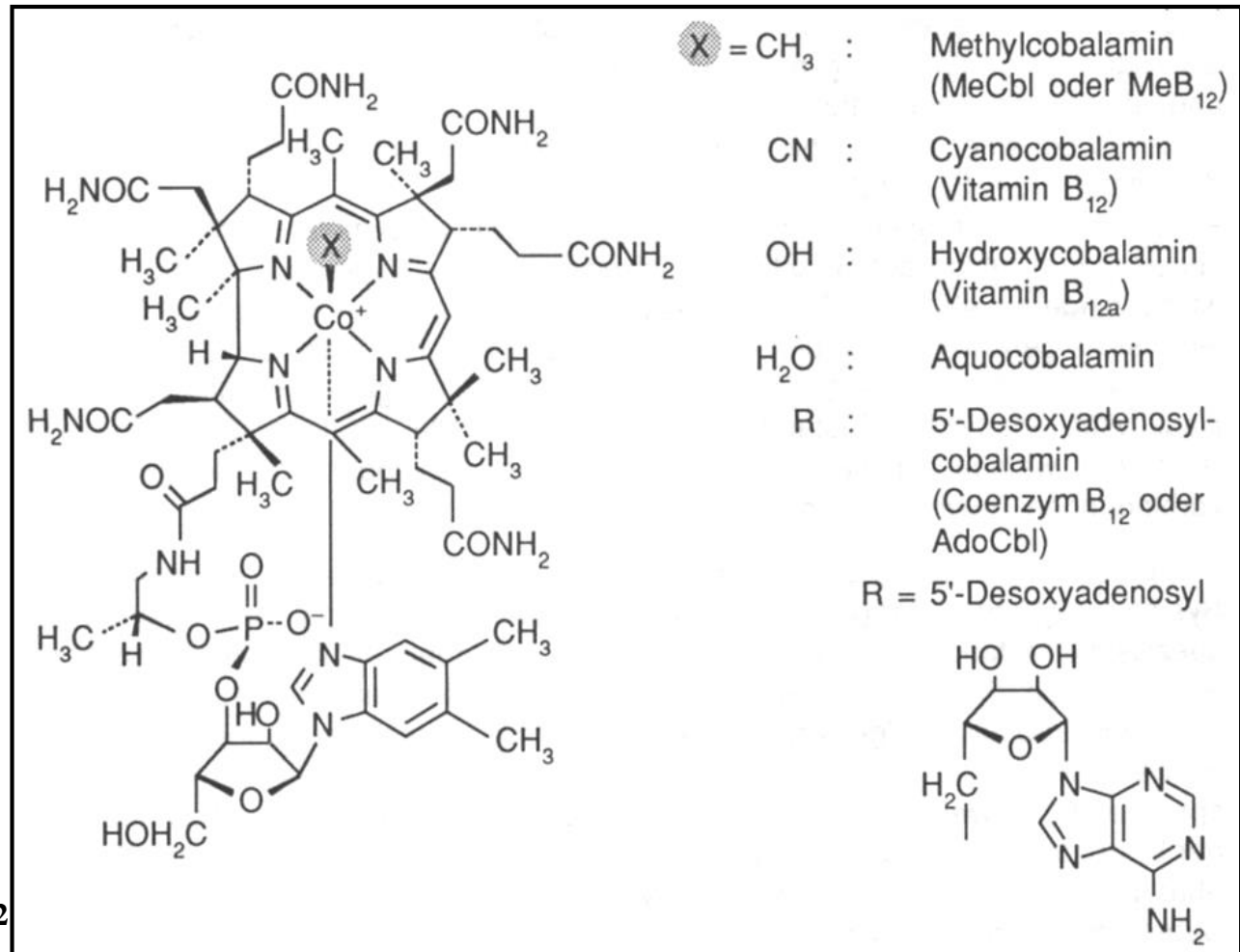
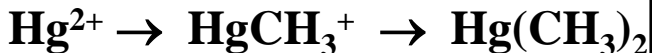
Cobalt Proteins

The adult human contains ~ 2.5 mg of cobalt, primarily bound in the macrocycle complex called cobalamin (vitamin B₁₂). The macrocycle ligand is a corrin ring

Alkyl cobalamins take part in redox reactions, alkylations and rearrangements.

Co^{III}-, Co^{II}- and Co^I-species may participate in 1-electron reductions and oxidations

Methylations (even of Hg²⁺):



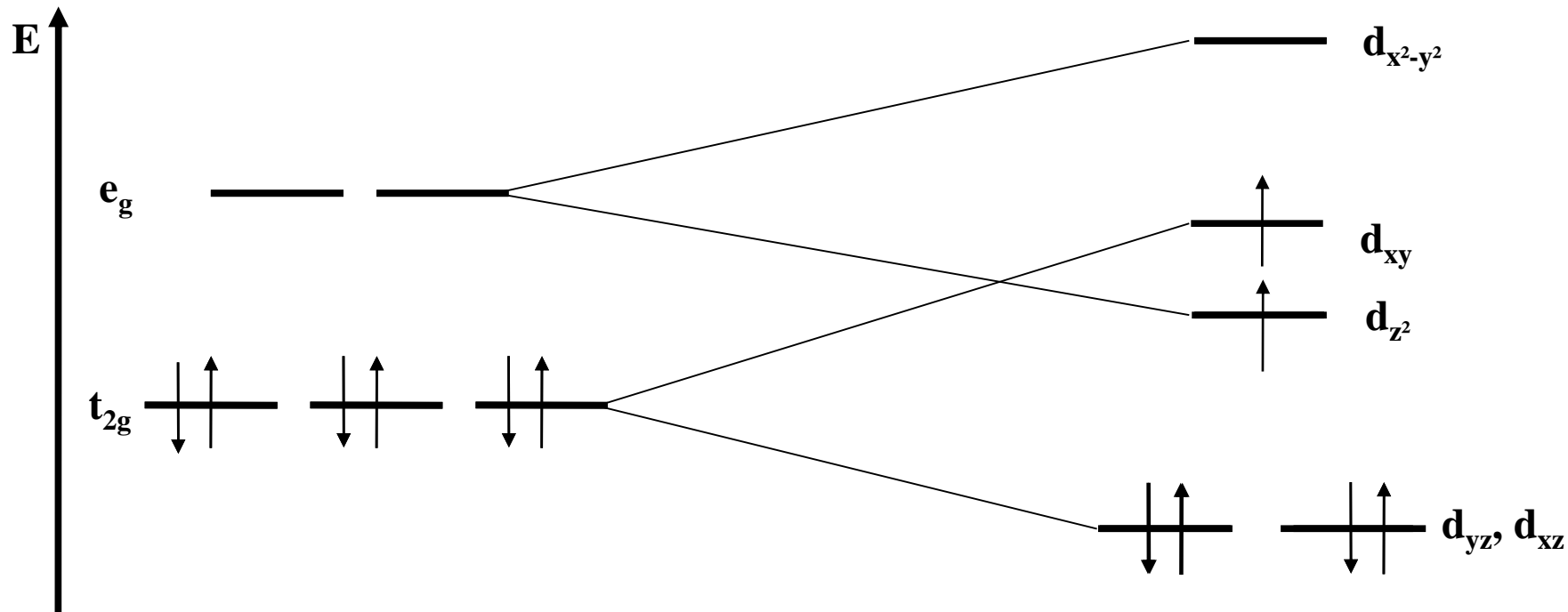
10. Biochemistry of Transition Metals

Cobalamin: Some Facts

- Cobalt is the most scarce element of the 3d-series, that means it is a trace element (Mn: 1060 ppm; Fe: 62000 ppm; Co: 29 ppm; Ni: 99 ppm; Cu: 68 ppm). Obviously, Co has been “chosen” due to a specific functionality
- The corrin ring (15 atoms) is smaller as the porphyrin ring (16 atoms, i.e. there is one CH₂-unit less in the ring). Co-porphyrin complexes are thus no model complexes for cobalamines. The bonding plane is distorted planar.
- The only known stable metal-organic compound in living creatures is the coenzyme MeB₁₂, with MeB₁₂ being stable in water
- Vitamin B₁₂ was discovered as treatment for pernicious anaemia in the 20's of the 20th century. In the beginning, essences of animal liver was used, which core part is made up of cobalt
- Not until 1948, the synthesis of cyanocobalamin = vitamin B₁₂ was successful (does not occur naturally in the body but exhibits therapeutic properties)
- Coenzyme B₁₂ is produced by animals and is stored in the liver
- The structure of vitamin B₁₂ and shortly after that of coenzyme B₁₂ was discovered by Dorothy Crowfoot-Hodgkins (Nobel price 1964)

10. Biochemistry of Transition Metals

Cobalamin: Crystal Field Splitting in Co^{3+} -Complexes $[\text{Ar}]3d^6$



Octahedral
diamagnetic

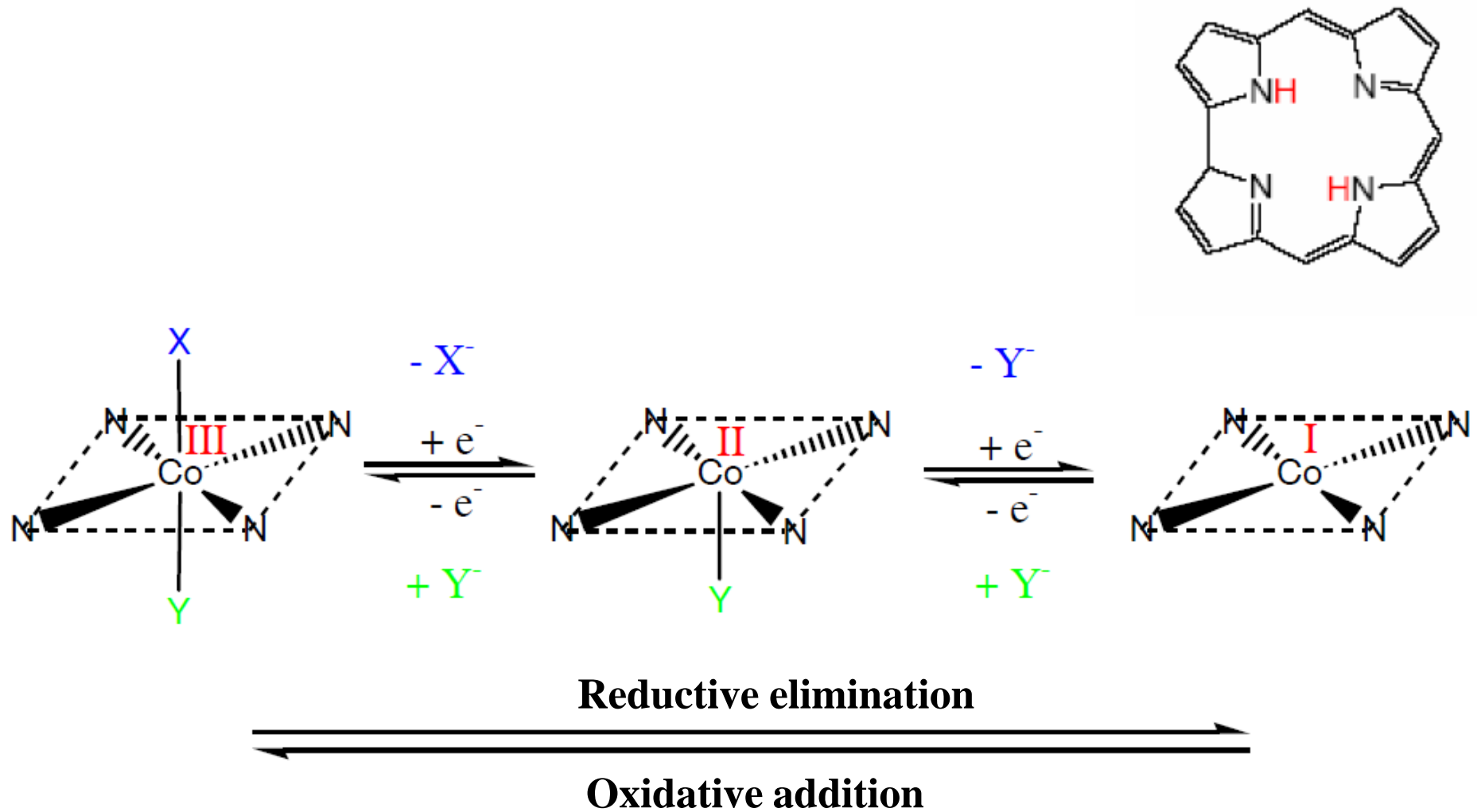
$$\text{CFSE} = -24 \text{ Dq(octahedral)}$$

Square-planar
paramagnetic

1- and 2-electron reduction feasible

10. Biochemistry of Transition Metals

Cobalamin: One-Electron-Reduction and One-Electron-Oxidation



10. Biochemistry of Transition Metals

Co-C-Bond Cleavage

Heterolytic cleavage

Only in presence of
reaction partner



By substitution
with for example
 H_2O to Co(III) and
a carbanion R^-

(EPR inactive)

Formation of a
supernucleophil
Co(I) and a
carbocation R^+

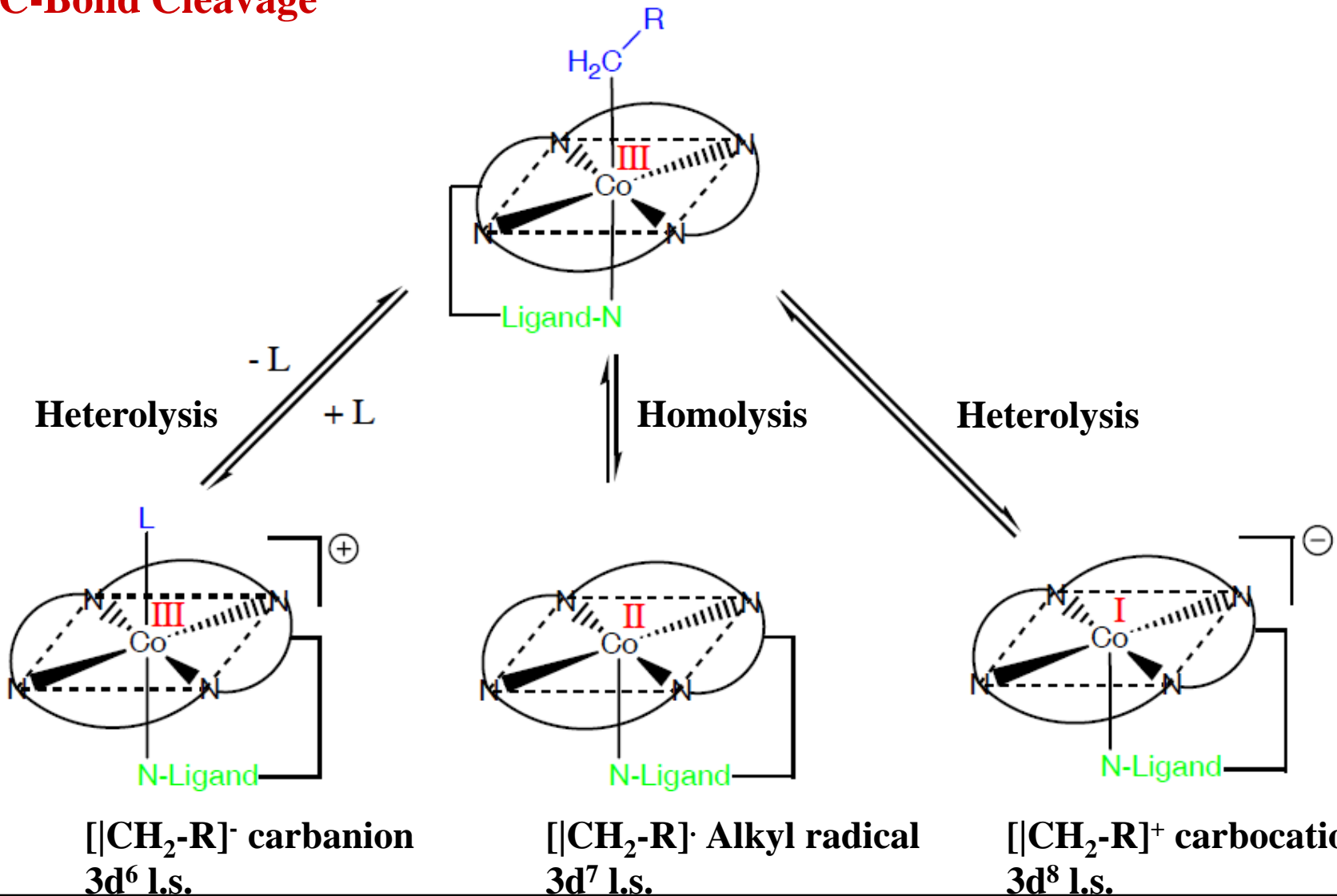
(EPR inactive)

Homolytic cleavage

Formation of a reactive primary
alkyl radical and low-spin $[\text{Ar}]3\text{d}^7$
cobalt(II)
(EPR active)

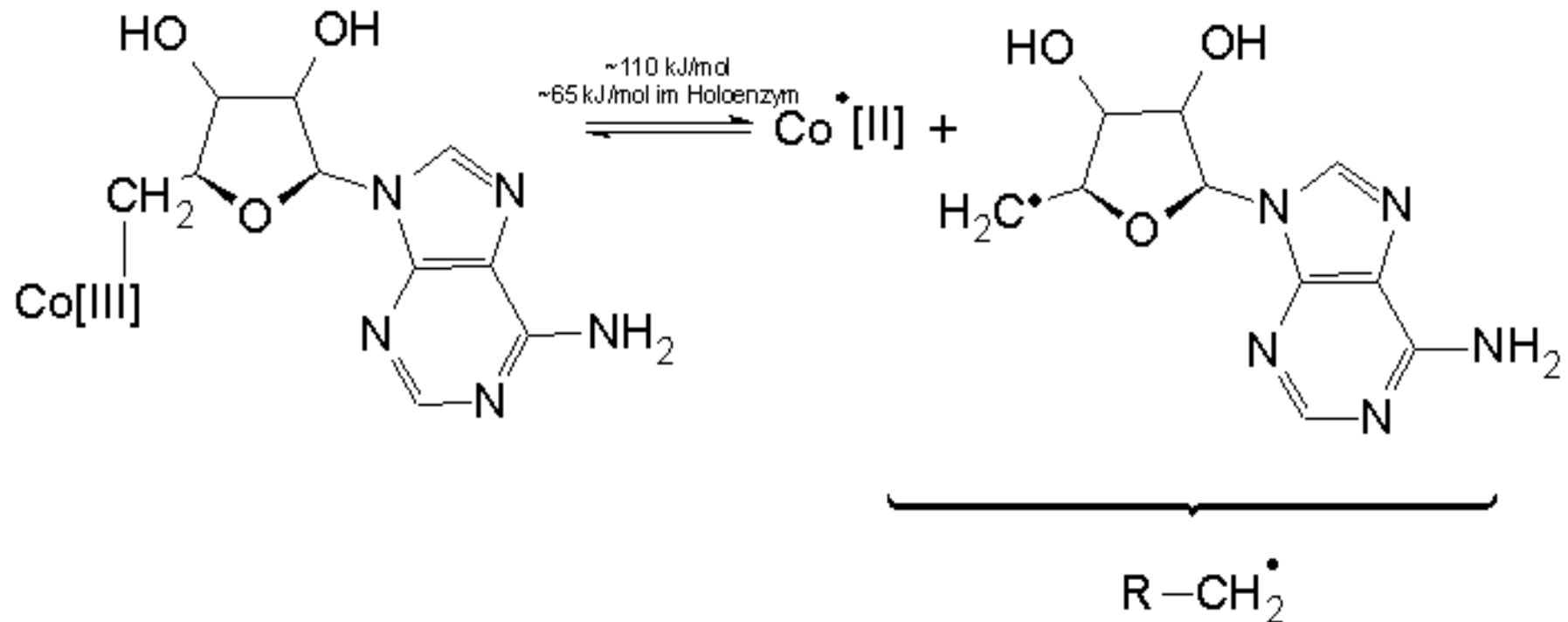
10. Biochemistry of Transition Metals

Co-C-Bond Cleavage



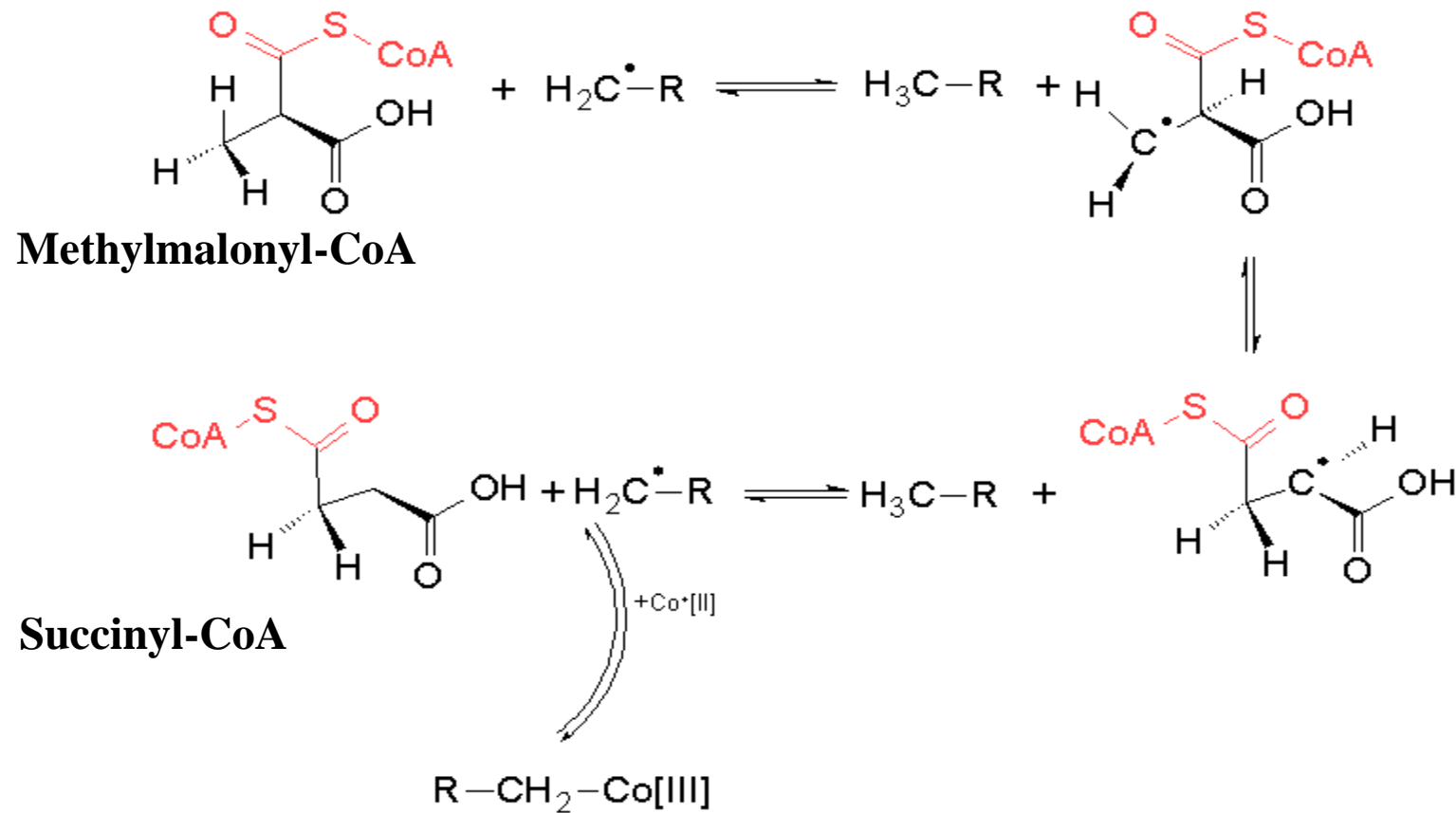
10. Biochemistry of Transition Metals

Homolytic Bond Cleavage



10. Biochemistry of Transition Metals

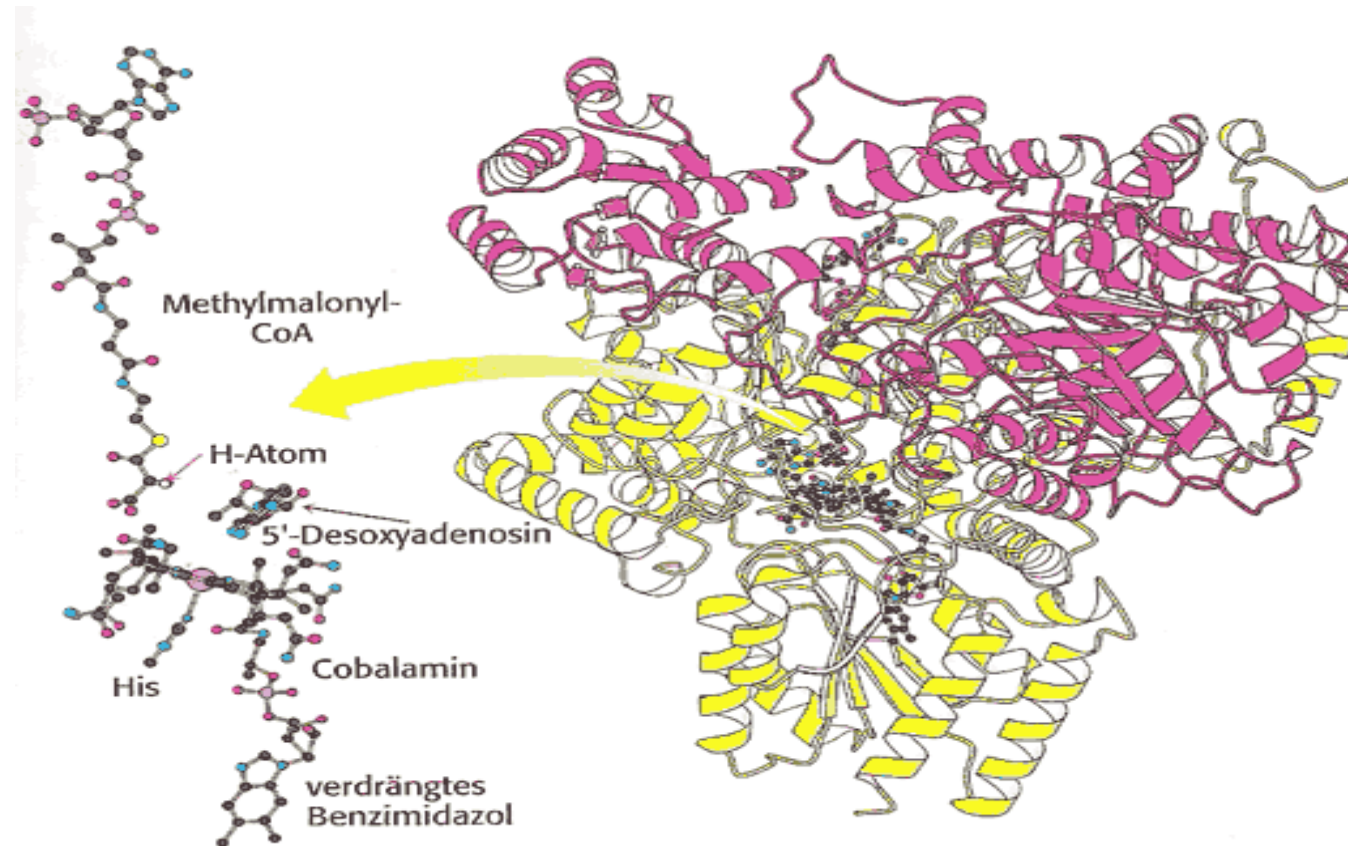
Methylmalonyl-CoA Mutase



Only indirect involvement of the Co-complex

10. Biochemistry of Transition Metals

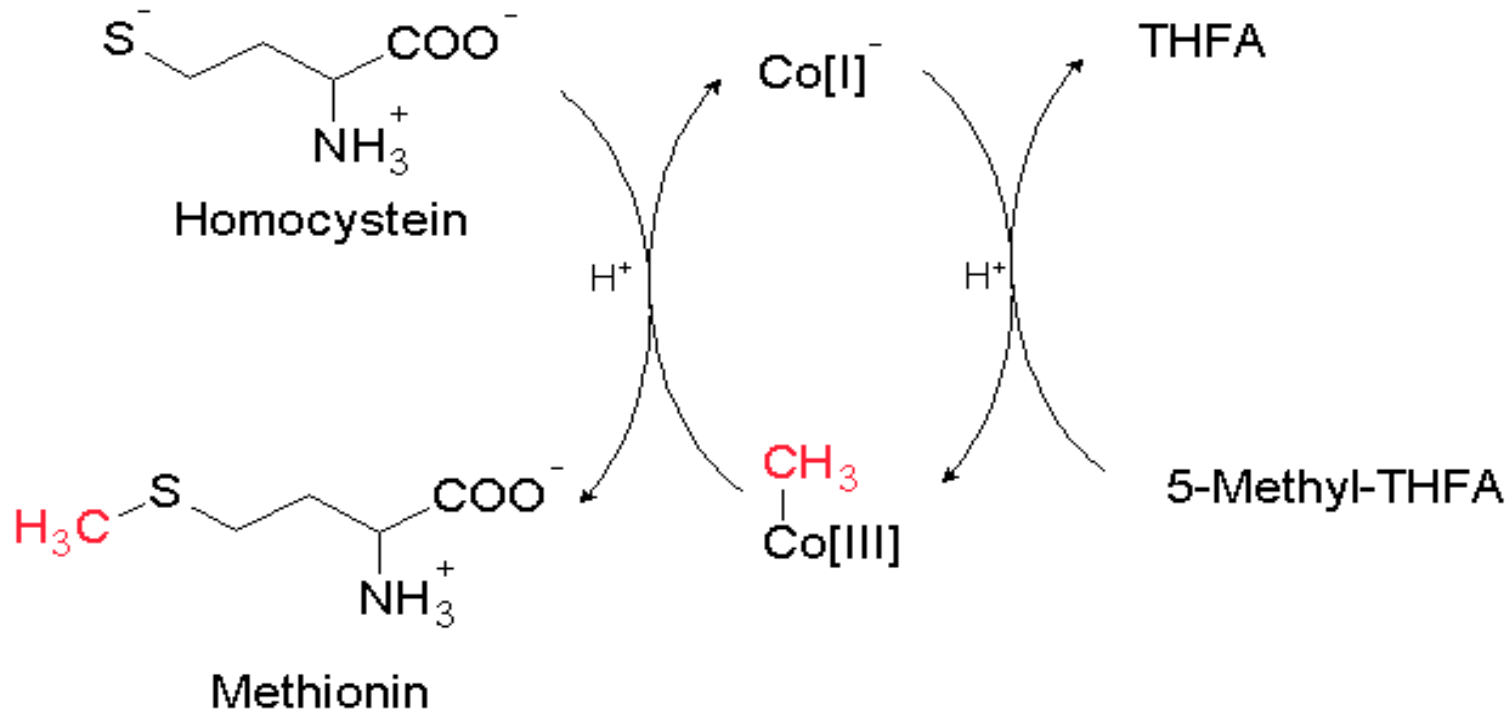
Methylmalonyl-CoA Mutase



Source: <http://www1.tu-darmstadt.de/fb/ch/akplenio/moproc/metalloproteine/cobalamin/cob3.htm>

10. Biochemistry of Transition Metals

Methylation by Tetrahydrofolic Acid (THFA) via Methylcobalamin MeB₁₂



10. Biochemistry of Transition Metals

The Nickel Group

Nickel

Ni^+

[Ar]3d⁹

strong reducing agent

Ni^{2+}

[Ar]3d⁸ l.s.

stable

Ni^{3+}

[Ar]3d⁷ l.s.

strong oxidising agent, low-spin

Palladium

Extremely scarce

Pd^{2+}

[Kr]3d⁸ l.s.

Pd^{4+}

[Kr]3d⁶ l.s.

Platinum

Extremely scarce

Pt^{2+}

[Xe]4f¹⁴3d⁸ l.s.

Cis-platinum $[\text{PtCl}_2(\text{NH}_3)_2]$ is cancerostatic

Pt^{4+}

[Xe]4f¹⁴3d⁶ l.s.

Pt⁴⁺-complexes are kinetically extremely stable (OLEDs)

10. Biochemistry of Transition Metals

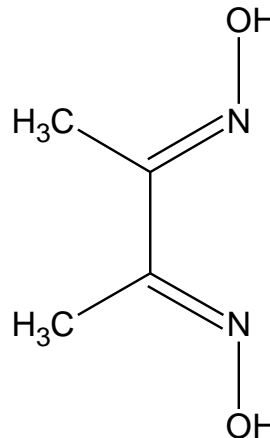
The Nickel Group

Nickel(II) chemistry

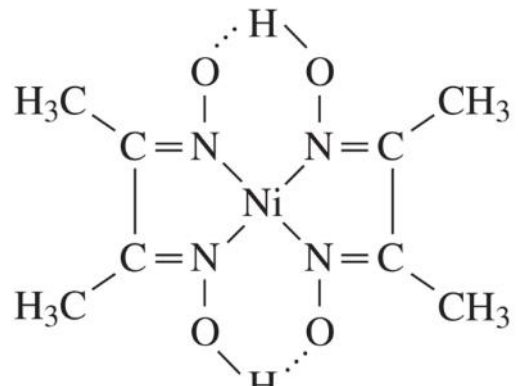
Hydrolysis



Coordination compounds



Dimethylglyoxime (H_2dmg)



Bis(dimethylglyoximato)nickel(II)

10. Biochemistry of Transition Metals

The Nickel Group

Nickel(II)-complexes

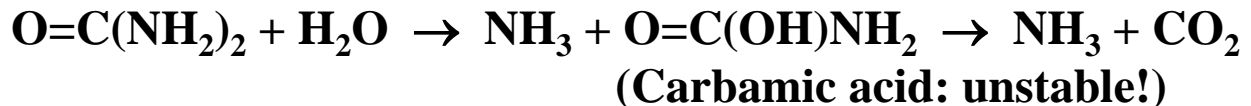
Typical for Ni^{2+} are octahedral, square-planar and tetrahedral complexes

Octahedral	Square-planar	Tetrahedral
H_2O NH_3 Ethylendiamine	Strong ligands, such as CN^- or Hdmg that force a square-planar arrangement	Cl^- Br^- I^-
Green, blue to violet Paramagnetic	Yellow, red Diamagnetic	Blue Paramagnetic

10. Biochemistry of Transition Metals

Nickel(II)-Proteins

- a) Ureases: Catalyse the decomposition of urea



- b) Ni-Fe-CO-hydrogenases:



- c) Acetyl-CoA-synthetases:

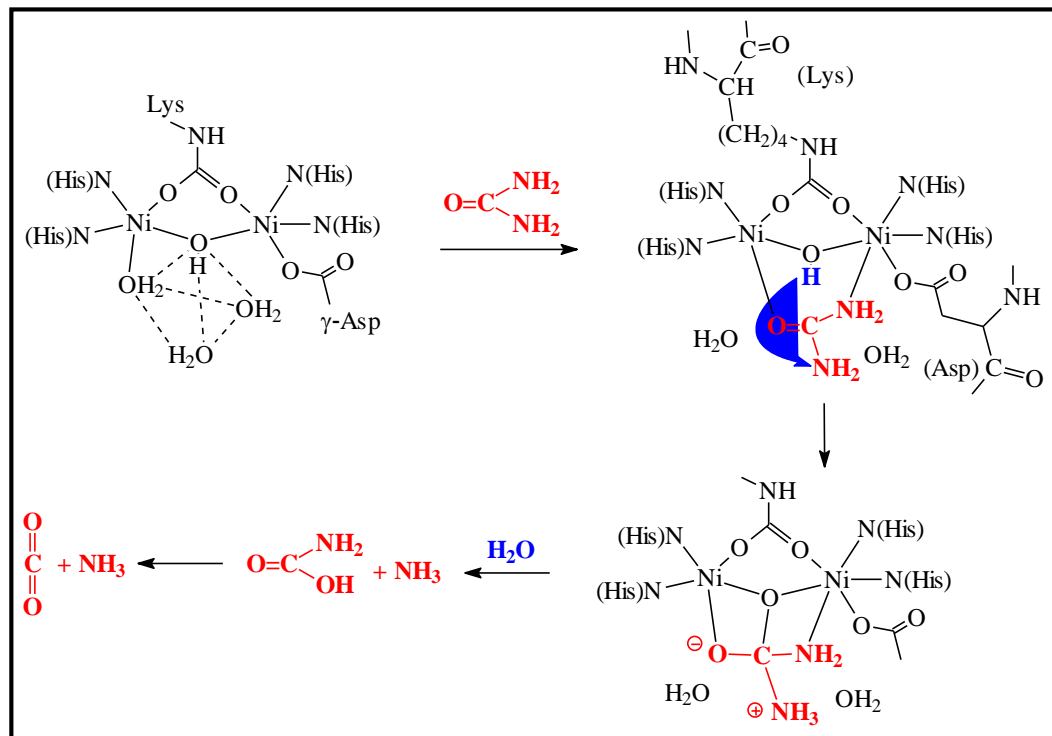


(Acetyl-CoA)

⇒ Acetyl-CoA = Precursor for acetylation reactions

- d) Ni superoxid dismutase

→ decomposition of O_2^- Radicals



10. Biochemistry of Transition Metals

The Copper Group

Copper

Cu^0	$[\text{Ar}]3\text{d}^{10}5\text{s}^1$
Cu^+	$[\text{Ar}]3\text{d}^{10}$
Cu^{2+}	$[\text{Ar}]3\text{d}^9$
Cu^{3+}	$[\text{Ar}]3\text{d}^8$

colloidal copper is antiseptic
tends to disproportionate, labile complexes
moderate oxidising agents(glucose-detection)
strong oxidising agent, stable l.s.-complexes

Silver

Scarce

Ag^0	$[\text{Kr}]4\text{d}^{10}5\text{s}^1$
Ag^+	$[\text{Kr}]4\text{d}^{10}$
Ag^{2+}	$[\text{Kr}]4\text{d}^9$

colloidal silver acts antimicrobial
strong oxidising agent
very strong oxidising agent



Gold

Extremely scarce

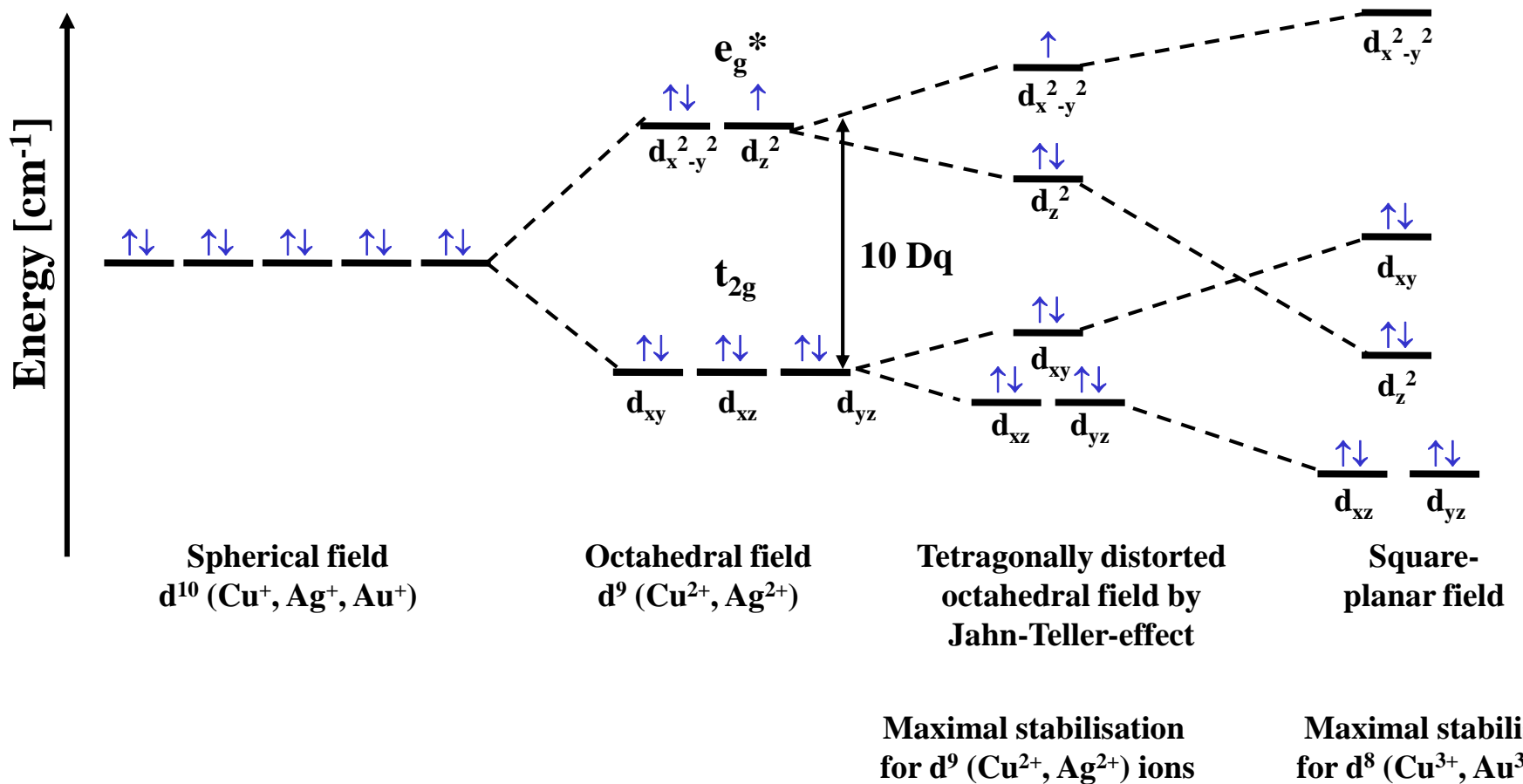
Au^0	$[\text{Xe}]4\text{f}^{14}5\text{d}^{10}6\text{s}^1$
Au^+	$[\text{Xe}]4\text{f}^{14}5\text{d}^{10}$
Au^{3+}	$[\text{Xe}]4\text{f}^{14}5\text{d}^8$

“Beaten gold“ is approved for food → E175
 $\text{Na}[\text{Au}(\text{CN})_2]$ → Treatment of rheumatoid arthritis
 $\text{H}[\text{AuCl}_4]$ → Extraction of purple of Cassius

10. Biochemistry of Transition Metals

The Copper Group

Crystal field splitting and stereo chemistry



10. Biochemistry of Transition Metals

Copper Proteins

Some functions

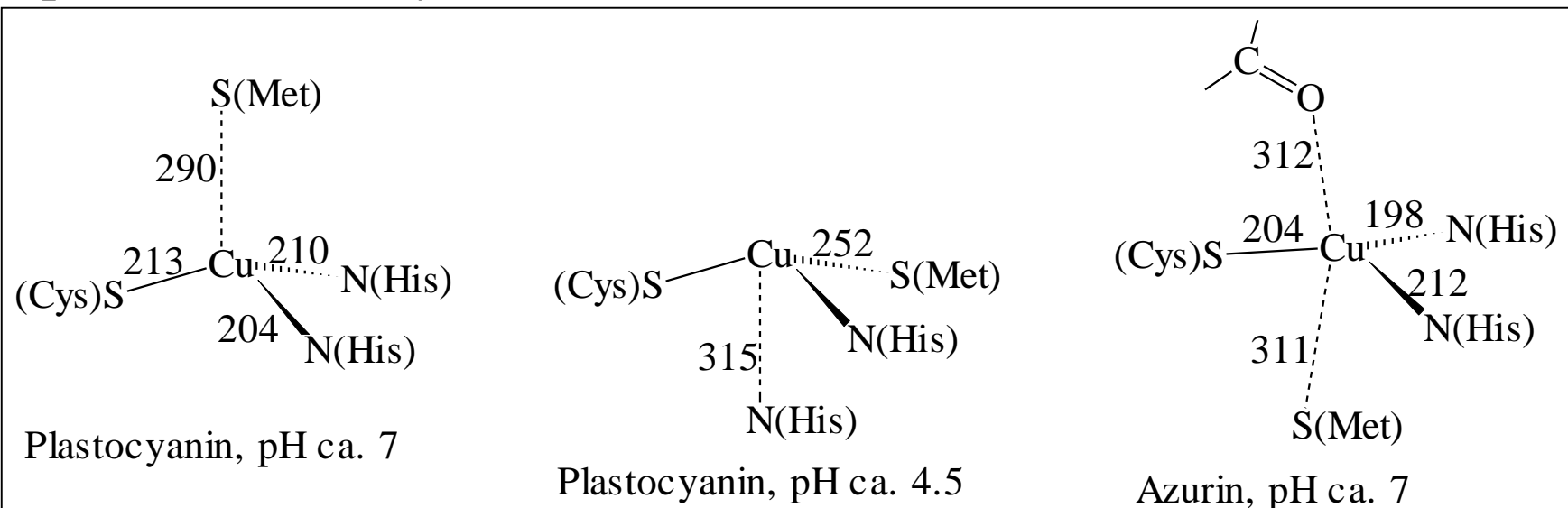
- Electron transport, e.g. in respiratory chain (cytochrome-*c*-oxidase) and in the electron transport chain of PSII to PSI: plastocyanine + azurin
- Oxygen transport: haemocyanines
- Regulation of iron and copper resorption: ceruloplasmin
- One-electron redox processes, e.g. in nitrite reductase: $\text{NO}_2^- \rightarrow \text{NO}$
- Two-electron redox processes, e.g. in galactose oxidase: $\text{RCH}_2\text{OH} \rightarrow \text{RCHO}$
- Disproportionation: detoxification of superoxide anion radical by CuZn-superoxide dismutase: $2 \text{O}_2^- + 2 \text{H}^+ \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$
- Oxygenation of organic substrates, e.g. by tyrosinase:
Tyrosine \rightarrow dopa \rightarrow indolquinone \rightarrow melanin
- Acetyl-coenzym-A-synthetase: $\text{CO} + \{\text{CH}_3\} + \text{CoA} \rightarrow \text{CH}_3\text{-C(O)-CoA}$
- As bio mineral: Atacamite = $\text{Cu}(\text{OH})_2 \cdot \text{Cu}(\text{OH})\text{Cl}$, which stem from teeth of marine blood worms (Genus glycera): Cu^{2+} with melanin



10. Biochemistry of Transition Metals

Copper Proteins: Type I “Blue Cu-Proteins”

- Structure: Trigonal coordination geometry; Cys, His and Met as ligands
- Opt. spectra: LMCT-bands ($\text{Cys}^- \rightarrow \text{Cu}^{2+}$) at 600 nm ($\epsilon \sim 3000 \text{ M}^{-1}\text{cm}^{-1}$)
- EPR-spectra: 4 hyper-fine lines by coupling with cores ($A = 5 \text{ mT}$)
 ^{63}Cu ($I = 3/2$, N = 70%)
 ^{65}Cu ($I = 3/2$, N = 30%)
- Function: Mostly electron transport
- Example: Plastocyanine, azurin



10. Biochemistry of Transition Metals

Copper Proteins: Type II

Structure:

Tetragonal coordination geometry, i.e.
3 His- as well as additional O- and N-functional ligands

Opt. spectra:

3d-3d absorption bands

EPR spectra:

4 hyperfine lines by coupling with cores ($A = 18 \text{ mT}$)

^{63}Cu ($I = 3/2$, N = 70%)

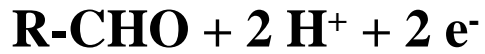
^{65}Cu ($I = 3/2$, N = 30%)

Function:

Oxydases and
oxygenases

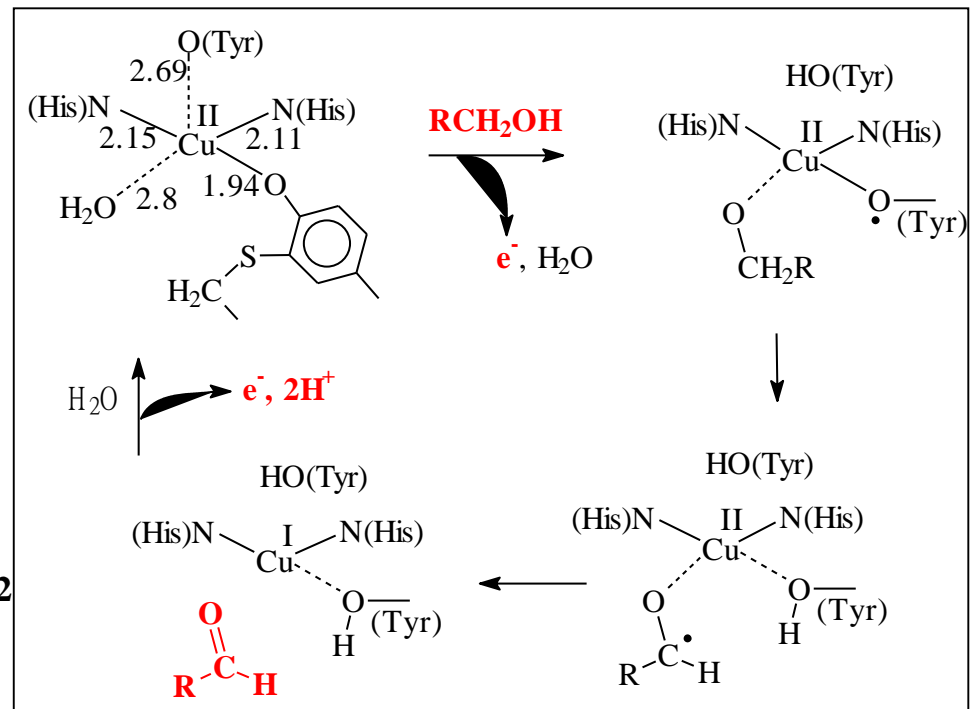
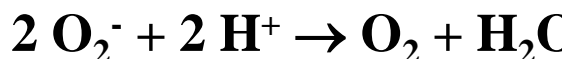
Example:

Galactose oxidase:



CuZn-

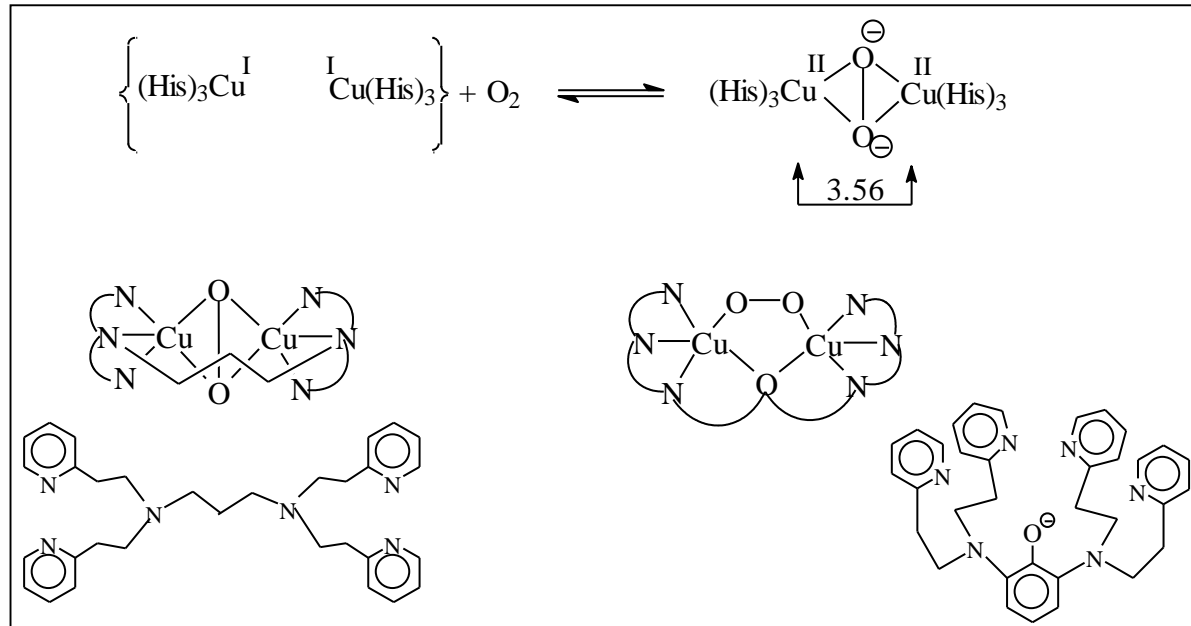
Superoxide dismutase:



10. Biochemistry of Transition Metals

Copper Proteins: Type III (according to Robin and Day)

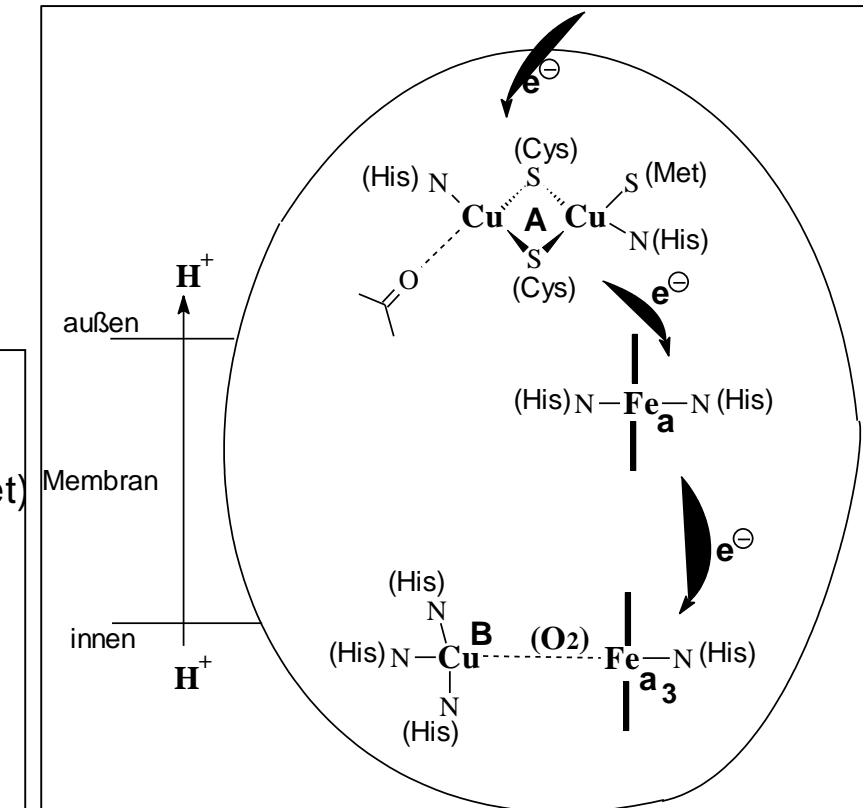
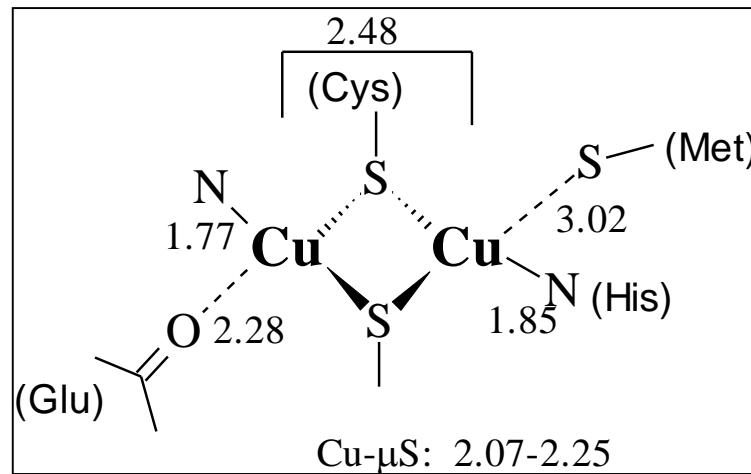
- Structure: Two copper centres of trigonal coordination
3 His-ligands, one peroxide bridge and possibly additional oxy bridges
- Opt. spectra: LMCT-bands ($O_2^{2-} \rightarrow Cu^{2+}$) at 600 nm ($\epsilon \sim 1000 M^{-1}cm^{-1}$)
- EPR spectra: EPR-inactive due to anti-ferromagnetic interactions between Cu^{2+} -ions
- Function: O_2 -transport
- Example: Haemocyanine
 $2 Cu^+ + O_2 \rightleftharpoons 2 Cu^{2+} + O_2^{2-}$
“oxidative addition”



10. Biochemistry of Transition Metals

Copper Proteins: Type A

- Structure: Dual-core, cysteine-bridged copper(I)-centres
- Opt. spectra: Rose due to MLCT
- EPR spectra: EPR-inactive → [Ar]3d¹⁰-configuration
- Function: Electron transport
- Example: N₂O-reductase
Cytochrome-c oxidase
(Cu^A-centre: see below)



10. Biochemistry of Transition Metals

The Zinc Group

Zinc

Zn^0

[Ar]3d¹⁰5s²

Zn^{2+}

[Ar]3d¹⁰

strong reducing agents

(distorted) tetrahedral labile complexes

Cadmium

Scarce

Cd^0

[Kr]4d¹⁰5s²

Cd^{2+}

[Kr]4d¹⁰

inhaled during smoking (20 cigarettes ~ 1 µg Cd)

redox-stable, octahedral labile complexes

Mercury

Extremely scarce

Hg^0

[Xe]4f¹⁴5d¹⁰6s²

Hg^+

[Xe]4f¹⁴5d¹⁰6s¹

Uptake via respiratory system (MAK-value = 0.1 mg/m³)

[Hg-Hg]²⁺ is diamagnetic, Hg_2Cl_2 is white and decomposes upon irradiation → Hg (calomel = nicely black), laxative which damages the kidneys

HgS: hexagonal black + cubic red modification

HgO: amorphous yellow + crystalline red modification

10. Biochemistry of Transition Metals

Zinc Proteins

Human: **2-2.5 g Zn per 70 kg body weight**

Transport: **Resorbed Zn binds to serum albumin and transferrin**

Proteins: **Carbonic anhydrase, carboxypeptidase, zinc finger, DNA-repair protein, etc. (several 100 known!)**

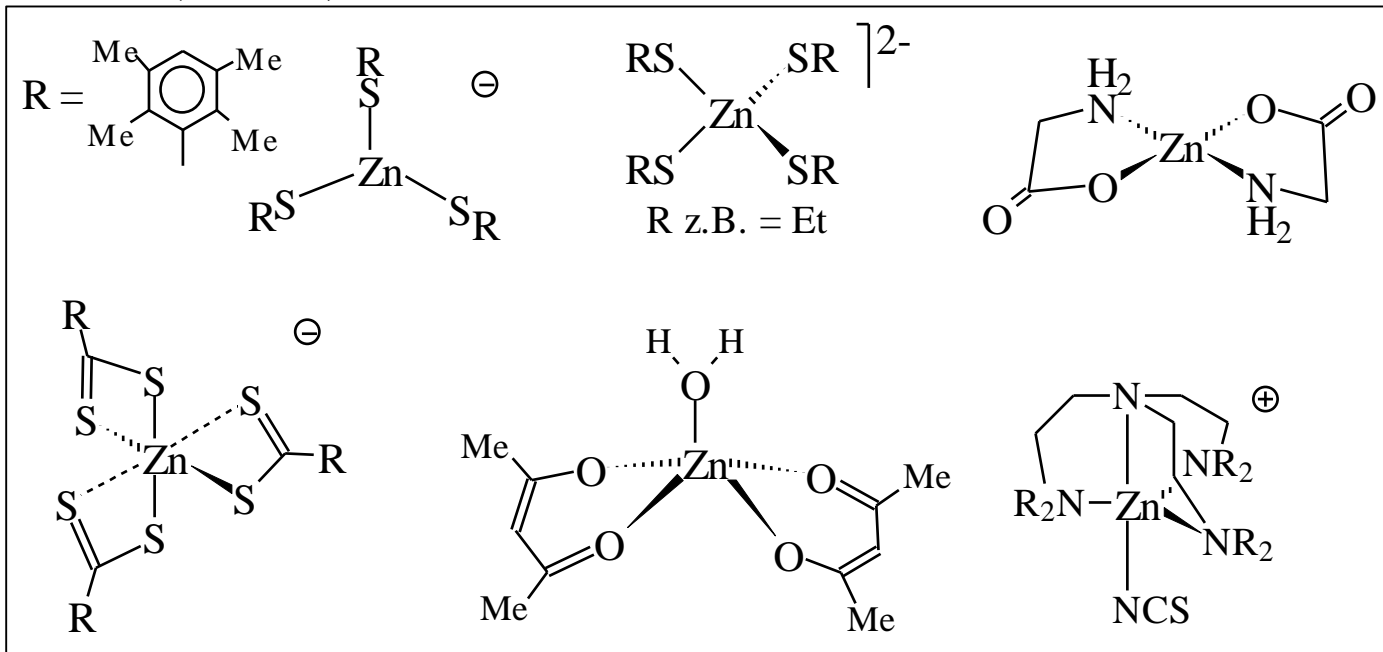
Some functions

- **Catalytic function:** hydrolases (peptidases, phosphatases, lipases), synthetases, isomerases, ligases
- **Structural functions:** stabilisation of tertiary structure of proteins
- **Hormonal regulation:** the hexameric storage modification of insulin is stabilized by three Zn^{2+} - ions coordinated to His, with three aqua ligands completing the coordination sphere of zinc and resulting in a CN of 6
- **Ada DNA repair-protein:** a zinc centre coordinating to four Cys, demethylises methyl phosphate
- **Zinc storage:** by thioneines (heavy metal-binding proteins)

10. Biochemistry of Transition Metals

Zinc Proteins

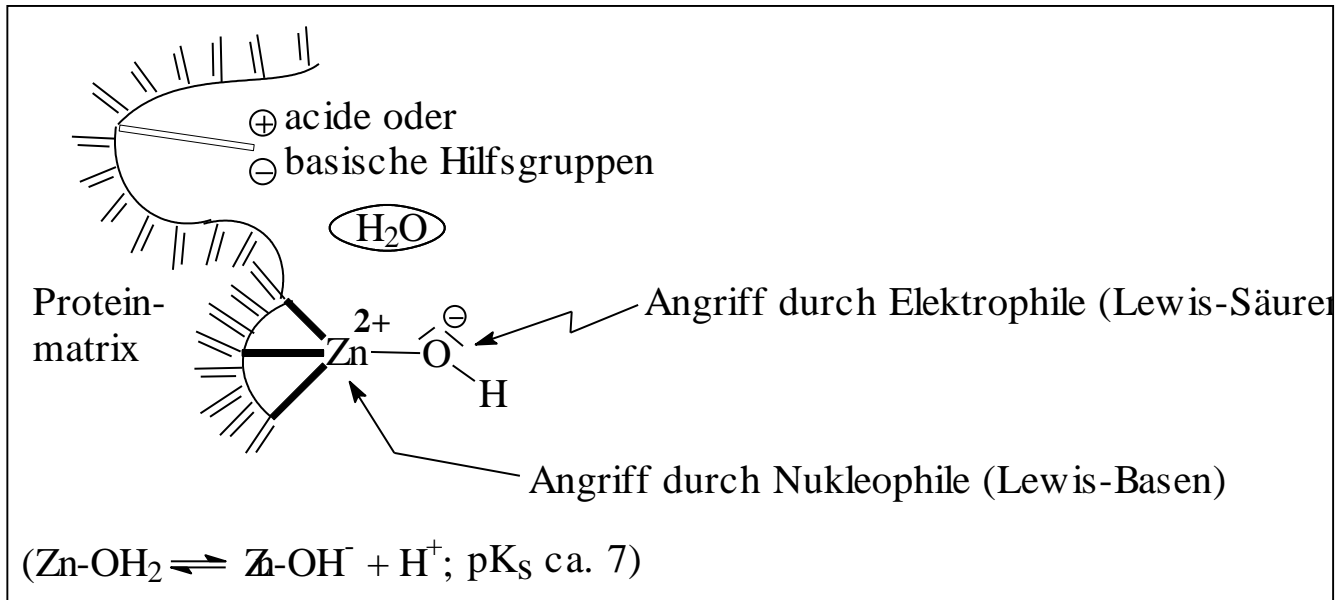
CN	Coordination geometry	Ligands
3	Trigonal –pyramidal (seldom)	S
4	Distorted tetrahedral, distorted tetragonal	S, O, N
5	Distorted tetragonal-pyramidal, distorted trigonal bipyramidal	O, N
6	octahedral (seldom)	S



10. Biochemistry of Transition Metals

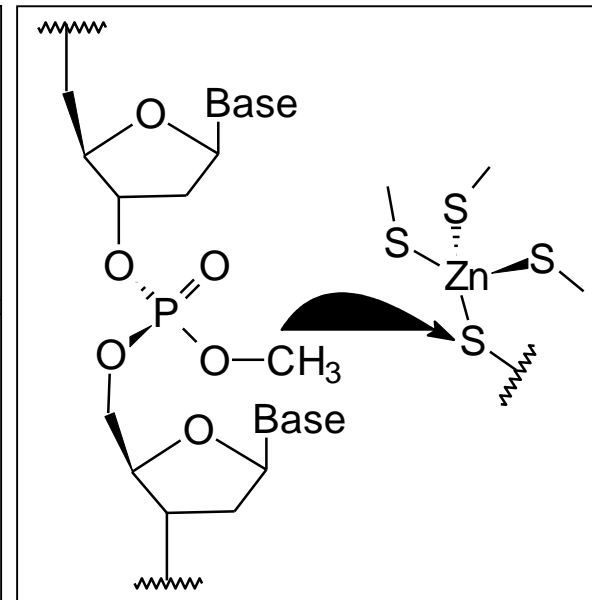
Zinc Proteins

Substrate activation



e.g. in alcohol dehydrogenase

Transfer of methyl groups



in Ada DNA repair-protein

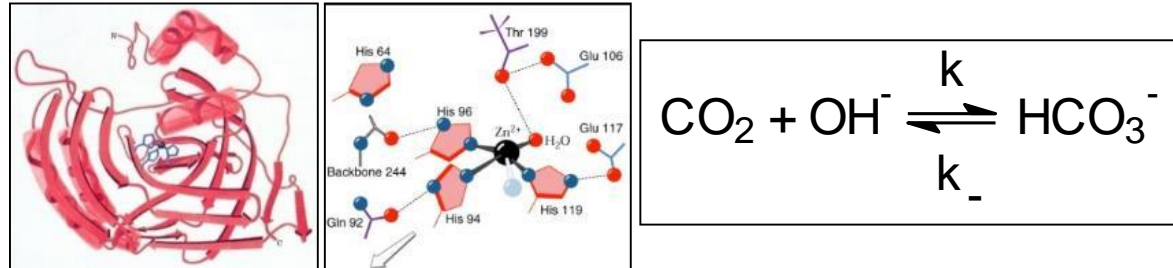
10. Biochemistry of Transition Metals

Zinc Proteins: Carbonic Anhydrase

Velocity

- Without a catalyst

$$k = 8.5 \cdot 10^3 \text{ M}^{-1}\text{s}^{-1}, k_- = 2 \cdot 10^{-4} \text{ s}^{-1}$$



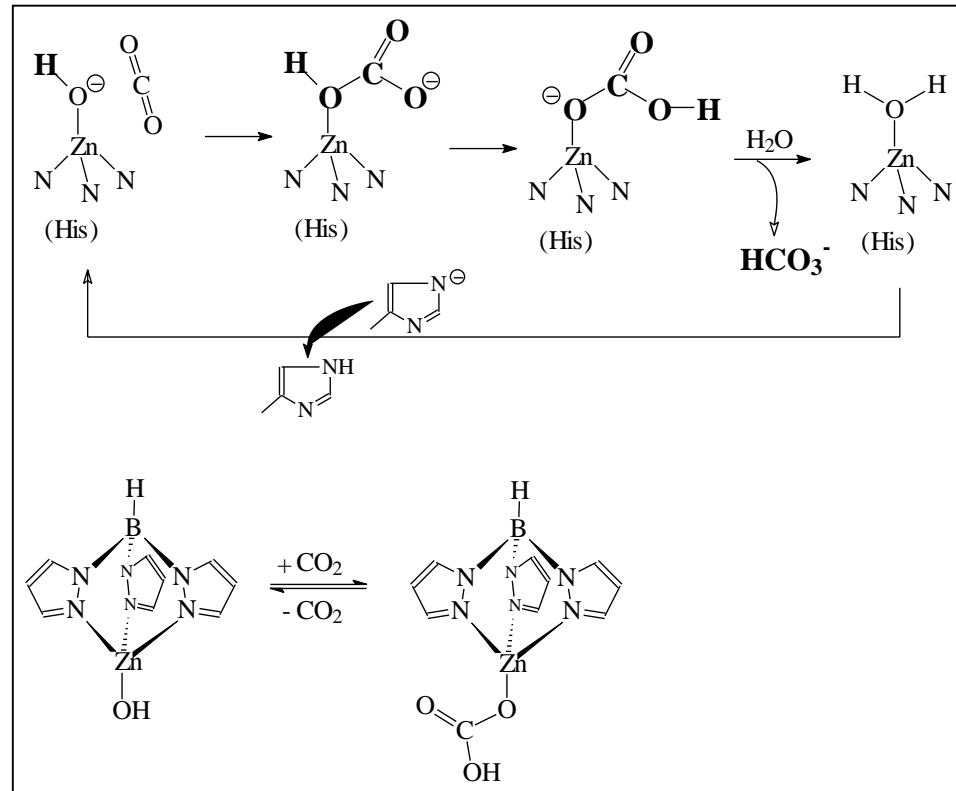
- Enzymatic

Equilibrium adjustment $\sim 10^7$ times faster

- Purpose is the transformation of CO_2 to HCO_3^- at place of origin



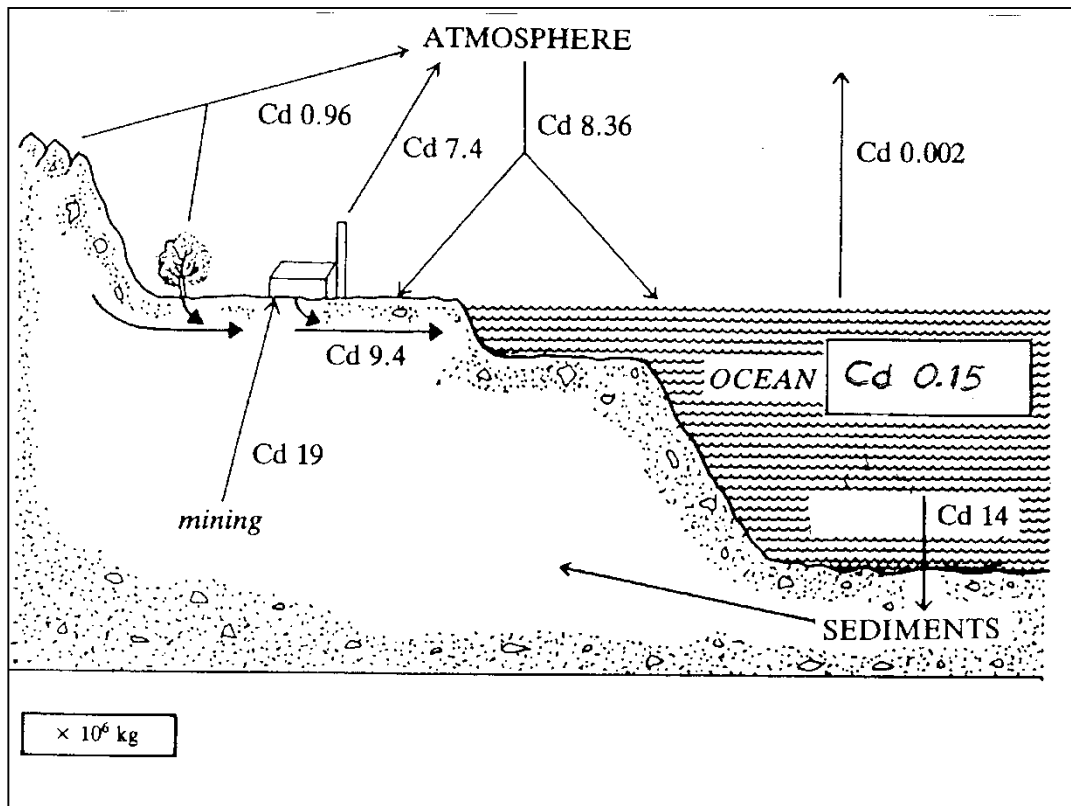
- Modell complex: $[\text{Zn}(\text{tpzb})(\text{OH})]$ with tpzb = trispyrazolyl borate



10. Biochemistry of Transition Metals

Cadmium: Biological Aspects of Cd²⁺

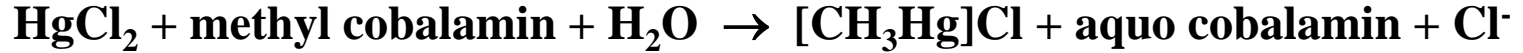
- Relation to zinc → deactivation of zinc enzymes through expulsion of Zn²⁺ from the active centre
- Similar ionic radius to Ca²⁺
→ Interference with Ca²⁺-balance,
e.g. disruption of Ca²⁺-ATPase
and construction of bones
- Acute Cd-intoxications can be
treated by glutathione



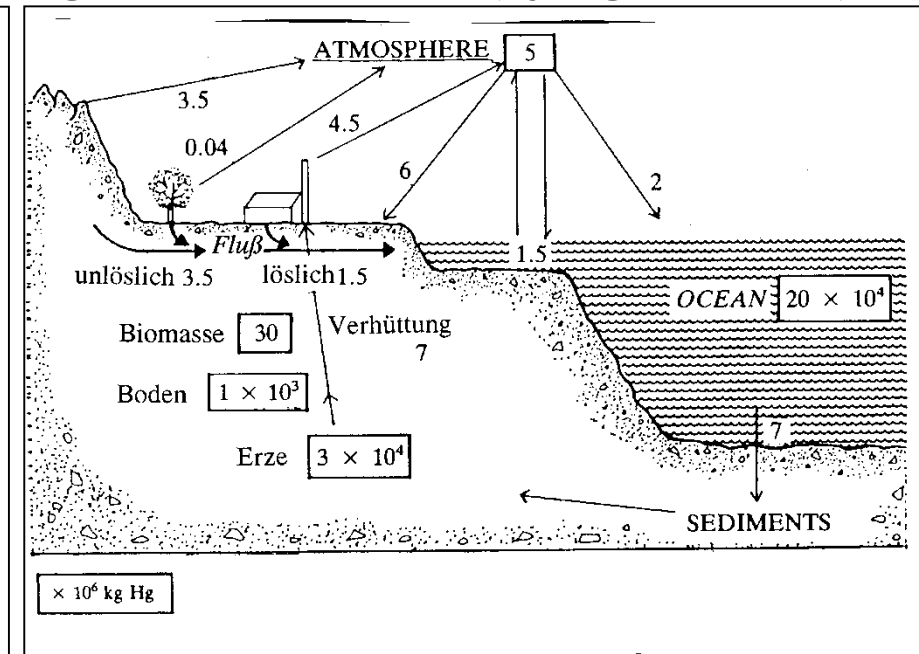
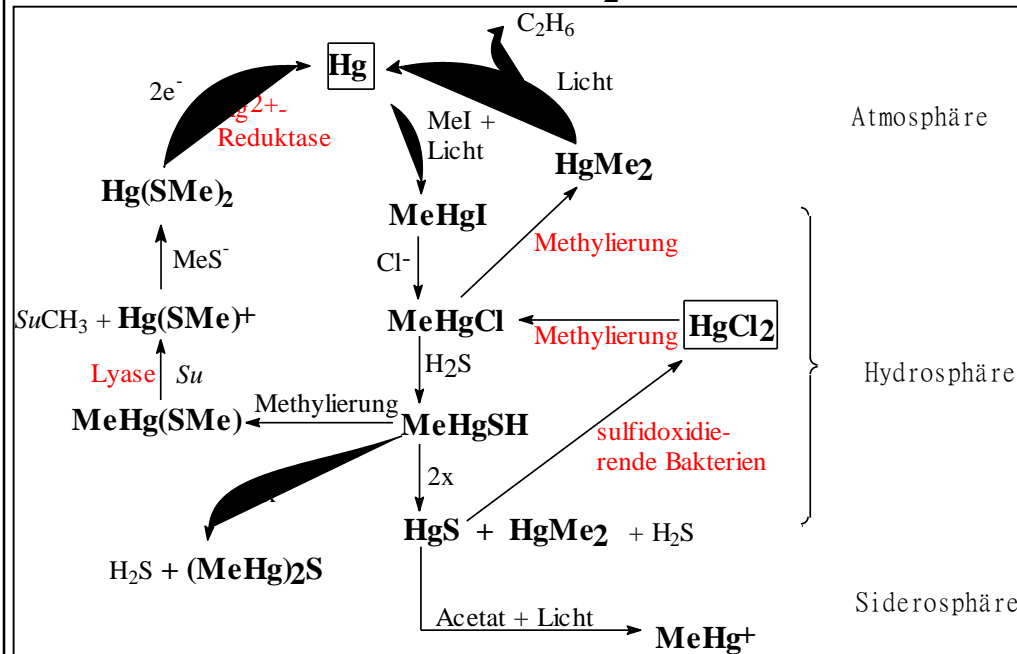
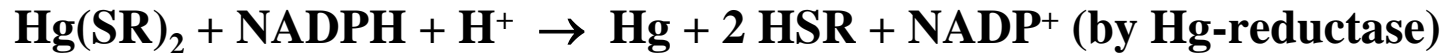
Source: P. O'Neill, Environmental Chemistry, 2nd Ed., Chapman & Hall, London 1993

10. Biochemistry of Transition Metals

Mercury: Biological Aspects of Hg and Hg²⁺

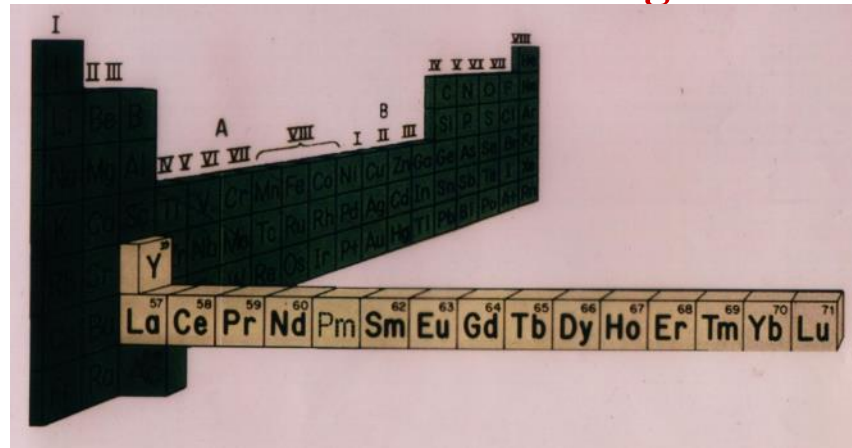


Follow-up reactions: $\text{MeHg}^+ + \text{SR}^- \rightarrow \text{MeHgSR}$ ("Minamata toxine" southern Japan)



11. Biochemistry of the Lanthanides and Actinides

Lanthanides: Electron configuration



Cations

[Xe]	La ³⁺ Ce ³⁺ Pr ³⁺ Nd ³⁺ Pm ³⁺ Sm ³⁺ Eu ³⁺ Gd ³⁺ Tb ³⁺ Dy ³⁺ Ho ³⁺ Er ³⁺ Tm ³⁺ Yb ³⁺ Lu ³⁺												
	Ce ⁴⁺ Pr ⁴⁺ Nd ⁴⁺				Sm ²⁺ Eu ²⁺ Dy ⁴⁺						Tm ²⁺ Yb ²⁺		

4f 0 1 2 3 4 5 6 7 8 9 10 11 12 13 14

Electronic configuration

e.g. of Gd³⁺/Eu²⁺/Tb⁴⁺

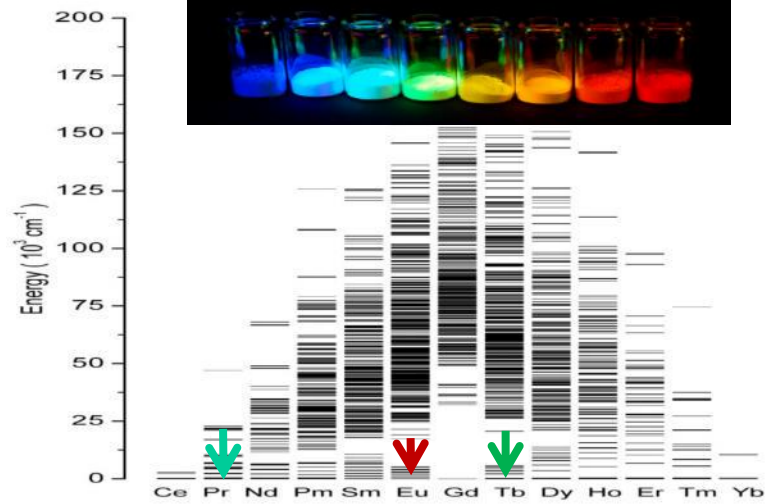
m _l	-3 -2 -1 0 1 2 3	-2 -1 0 1 2	0	-1 0 1																				
[Xe]	<table border="1"><tr><td>↑</td><td>↑</td><td>↑</td><td>↑</td><td>↑</td><td>↑</td><td>↑</td></tr></table>	↑	↑	↑	↑	↑	↑	↑	<table border="1"><tr><td></td><td></td><td></td><td></td><td></td><td></td></tr></table>							<table border="1"><tr><td></td><td></td><td></td><td></td></tr></table>					<table border="1"><tr><td></td><td></td><td></td></tr></table>			
↑	↑	↑	↑	↑	↑	↑																		

Ce³⁺ ... Yb³⁺, Pr⁴⁺, Nd⁴⁺, Tb⁴⁺, Dy⁴⁺, Sm²⁺, Eu²⁺, Tm²⁺

Ce³⁺ ... Yb³⁺

→ paramagnetic ions

→ ions with complex optical spectra



11. Biochemistry of the Lanthanides and Actinides

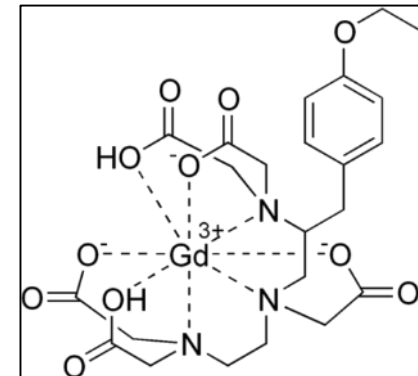
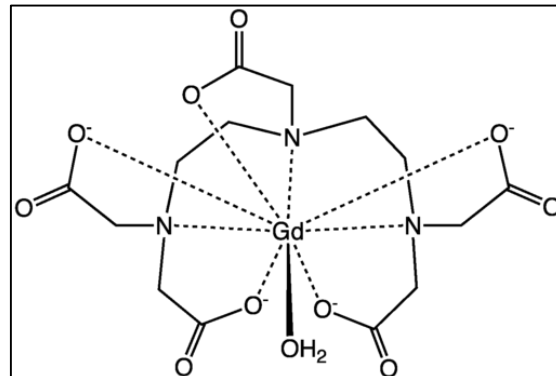
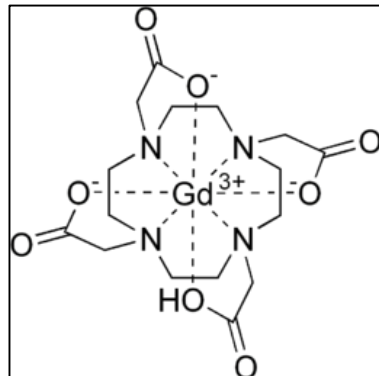
Lanthanides: MRT contrast enhancement agents – Eu^{2+} and Gd^{3+}

Gadolinium

Gd^{3+} [Xe]4f⁷ acutely toxic, only in complexed form quite safe
extremely strong paramagnet, long electronic relaxation time
Reduction of relaxation time of protons of those aqua ligands,
coordinating to Gd^{3+}

Gd^{3+} -complexes as contrast agents for MRI scans

- $[\text{Gd}(\text{DOTA})]^2-$ “Dotarem” DOTA = 1,4,7,10-tetra azacyclo dodecanetetraacetate
- $[\text{Gd}(\text{DTPA})(\text{H}_2\text{O})]^2-$ “Magnevist” DTPA = diethylene triamine pentaacetate
- $[\text{Gd}(\text{gadoxetic acid})]^2-$ “Primovist”



11. Biochemistry of the Lanthanides and Actinides

Lanthanides: Other than Gd³⁺

Trivalent ions

La³⁺

[Xe]4f⁰

La-citrate as additive in agriculture to improve feed conversion rate in livestock
Phosphate binder in hyperphosphatemia

Sm³⁺

[Xe]4f⁵

Antibacterial properties of samarium complexes
β-emitter ¹⁵³Sm-EDTMP accumulates in bone metastases and has a half-life of 46.3 hours

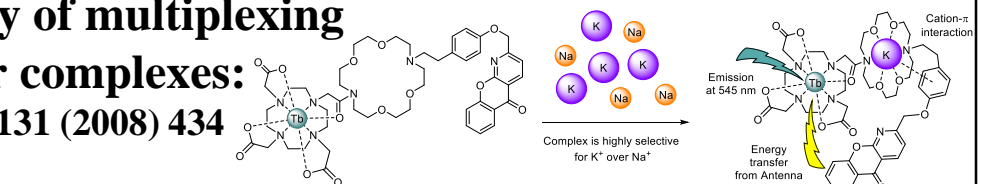
Tb³⁺

[Xe]4f⁸

Luminescent bioassays: Long-lived and bright PL, possibility of multiplexing

K⁺ sensor complexes:

Lit.: JACS 131 (2008) 434



Lu³⁺

[Xe]4f¹⁴

¹⁷⁷Lu as emitter of low-energy gamma radiation for imaging and medium energy β-particles for therapy
LuPO₄ nanoparticles doped with Pr³⁺ and/or Nd³⁺ as UV-C scintillator

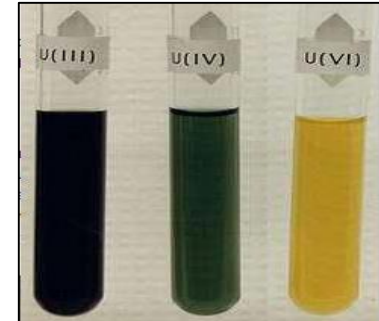
11. Biochemistry of the Lanthanides and Actinides

Actinides

Uranium

U^0	$[\text{Rn}]5\text{f}^36\text{d}^17\text{s}^2$
U^{3+}	$[\text{Rn}]5\text{f}^3$
U^{4+}	$[\text{Rn}]5\text{f}^2$
U^{6+}	$[\text{Rn}]$

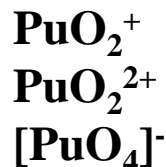
reacts with hot water: $\text{U} + 2 \text{H}_2\text{O} \rightarrow \text{UO}_2 + 2 \text{H}_2$
wide spread, each person comprise about 70 µg uranium
reducing
 UO_2^{2+} is the most stable ion in vivo or aqueous solution
 $\text{Na}_2\text{U}_2\text{O}_7$ “Yellow cake”



Plutonium

Pu^0	$[\text{Rn}]5\text{f}^67\text{s}^2$
Pu^{3+}	$[\text{Rn}]5\text{f}^5$
Pu^{4+}	$[\text{Rn}]5\text{f}^4$
Pu^{5+}	$[\text{Rn}]5\text{f}^3$
Pu^{6+}	$[\text{Rn}]5\text{f}^2$
Pu^{7+}	$[\text{Rn}]5\text{f}^1$

$[\text{Pu}(\text{H}_2\text{O})_n]^{3+}$
 $[\text{Pu}(\text{H}_2\text{O})_n]^{4+}$ similar ion charge density as Fe^{3+}
incorporation in iron-containing metalloenzymes
 PuO_2 in radionuclide batteries

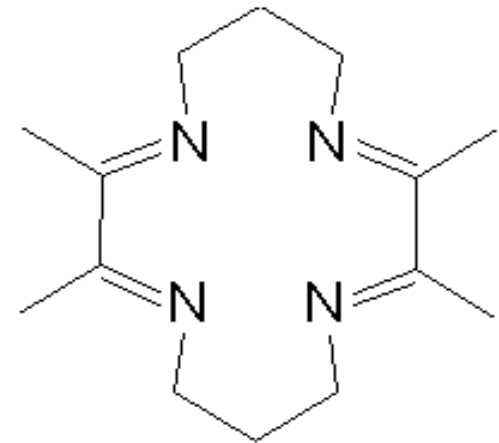


12. Modell Complexes

Aims of the Facilitation of Model Complexes

Structural Models

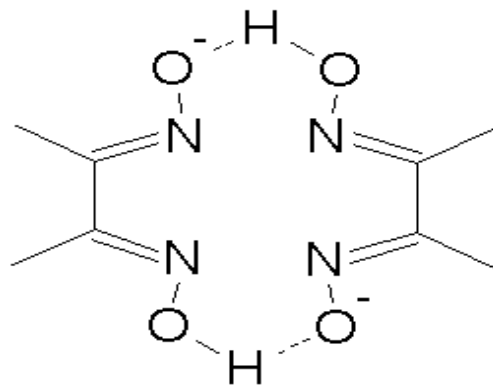
- Contribution to the structural elucidation of metalloenzymes
- Mimicking optical spectra and magnetic properties



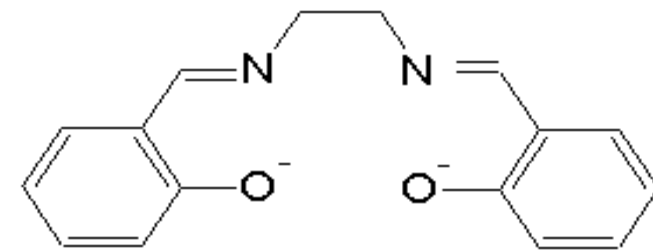
COSTA Ligand

Functional Models

- Elucidation of enzymatic reaction mechanisms
- Investigation of catalytic activity
- Application for lab syntheses
of small molecules



Bis(diacetyldioxim)



salen Ligand

13. Analytical Methods

Overview

1. Diffraction methods (→ clarification of 3D structure)
 - Problem: Crystallisation of proteins
 - Complex structures limit the resolution to only ca. 0.2 nm, i.e. identification of hydrogen atoms is impossible
2. Electron microscopy (→ 3D structure with intermediate resolution > 1 nm)
3. NMR-Spectroscopy (→ local structure and dynamic properties)
 - Problem: Complexity of proteins
4. X-ray absorption spectroscopy, e.g. EXAFS, XANES (→ local structure)
5. EPR-Spectroscopy (→ electronic properties of a species unpaired electrons)
6. Mößbauer Spectroscopy (→ identification of species with quadrupole moment)
7. Optical Spectroscopy (→ colour and electronic properties)
8. SQUID (→ characterisation of magnetic materials)
9. Cyclic voltammetry (→ characterisation of redox processes, e.g. electron transfers)
10. Vibrational spectroscopy, e.g. IR-, Raman-, Resonance-Raman-Spectroscopy (→ detection of functional groups)
 - Problem: Complexity of proteins

14. Applications Areas of Bioinorganic Chemistry

a) Commercial Production and Biotechnology

- Anaerobe bacterial decomposition in sewage treatment plants or sediments: Fe, Ni, Co
- Bacterial leaching (e.g. > 25% of global copper production): Fe, Cu, Au, U

b) Environmental Chemistry

- Agricultural trace element problems: nitrogen fixation (Fe, Mo, V)
- Environmental impact: Pb, Cd, Hg, As, Al, Cr
- Pollutant decomposition and detoxification, e.g. by peroxidases: Fe, Mn, V
- Phytoextraction/-leaching: Cr, Mn, Co, Ni, Cu, Ag, Au, Zn, Cd, Hg, In, Ga, Ge, Sn, As, Eu, Gd, Tb, Lu, ...

c) Pharmacy

- Diagnostics: Fe^{3+} , Gd^{3+} , Ba^{2+} , Tc^{3+} , Xe
- Therapeutics: Pt, Au, Li, B, Gd, Bi, As, Hg
- “Cis-platinum“, $\text{cis-PtCl}_2(\text{NH}_3)_2$, for the treatment of certain types of tumours
- Radio-iodine-therapy, e.g. in case of excessive thyroid function
- Metabolism through P-450-enzymes, metalloenzymes blocker: Fe, Zn

14. Applications Areas of Bioinorganic Chemistry

d) Biomaterials

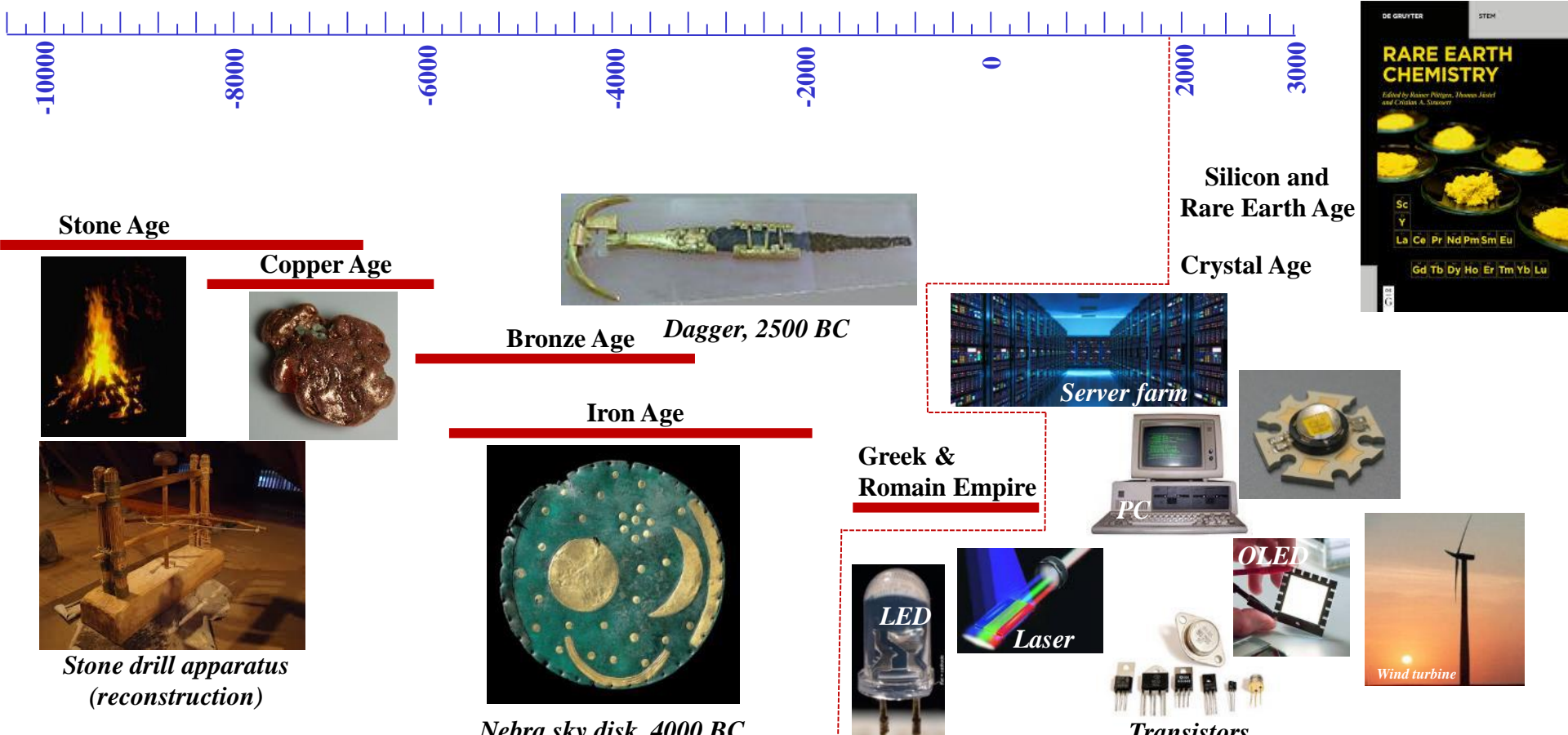
- Biocompatible (dental)implants
- Treatment of undesired demineralisation processes such as osteoporosis or caries: Ca^{2+} , PO_4^{3-} , F^-
- Biocompatible light guides

e) Inorganic Food Ingredients

- Deficiency symptoms → supplementation: Fe, Co, Zn, Se, ...
- Intoxications → complexation, e.g. during EDTA-therapy
- Precaution → iodine blockade through administration of KI, e.g. in case of a potential exposure to ^{131}I
- Food Design → TiO_2 nanoparticles

14. Applications Areas of Bioinorganic Chemistry

Environmental impact of heavy metals: Dissipation ↔ Toxicology



Source: Modified after Dr. Rytz, EOT

14. Applications Areas of Bioinorganic Chemistry

Phytoextraction/-leaching

Phytoextraction is to extract metal from soil substrates where plants capable of growing in high mineral environments

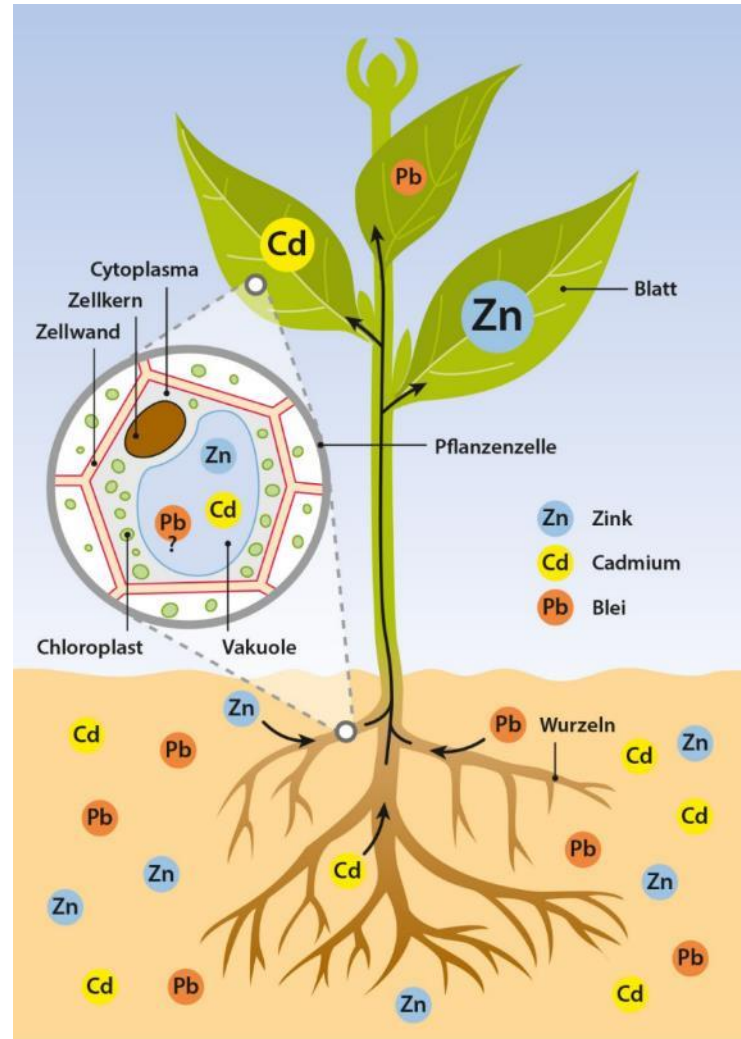
→ Zn, Cd, Hg, Pb,

(Lit.: Chaney et al., 1998)

Phytomining concerns extracting metals from soil substrates by harvesting specially selected hyper-accumulating plants

→ Ge, Ga, In, Sn,

(Lit.: Sheoran, S. Sheoran & Poonia, 2013)



Lit.: <https://motherboard.vice.com/de/article/phytomining>