Radiomedicinské methody

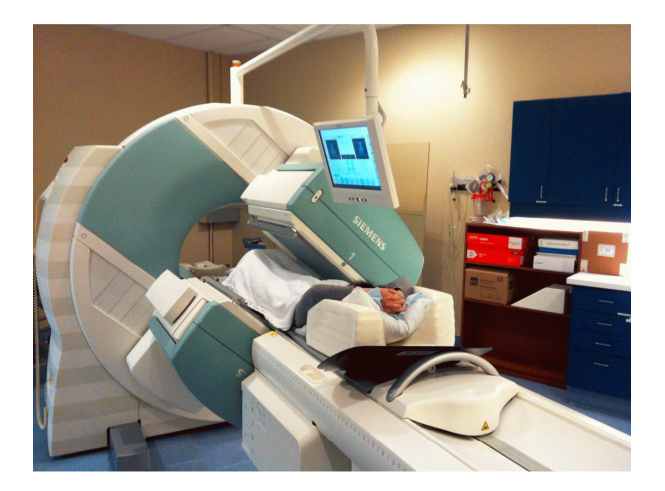
SPECT – single photon emission computer tomography

PET – **positron emission tomography**

Radiodiagnostické metody

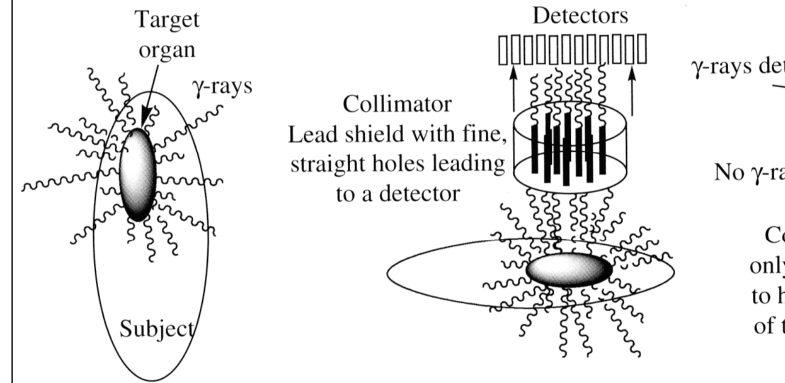
Jednofotonová emisní tomografie (SPECT)

- γ -emitující radioisotopy
 - rozlišení ~1 cm3





SPECT



View of the detectors through the Collimator γ-rays detected No γ-rays

> Collimator allows through only γ -rays travelling parallel to holes so creating an image of the radiation source in the detectors

Izotopy pro SPECT

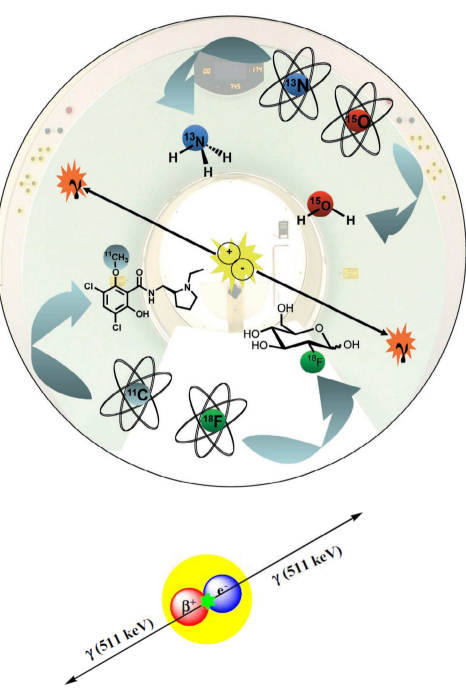
| Izotop | Přeměna | Poločas | Zdroj |
|-------------------|---------|---------|--|
| ^{99m} Tc | γ | 6 h | generátor, ⁹⁹ Mo(β ⁻) ^{99m} Tc, 66 h |
| ¹¹¹ In | γ | 68 h | cyklotron, ¹¹¹ Cd(p,n) ¹¹¹ In |
| ¹³¹ I | γ, β- | 8 d | reaktor |
| ¹⁵³ Sm | γ, β- | 46 h | reaktor |
| ¹⁶⁶ Ho | γ, β- | 26 h | reaktor |
| ¹⁷⁷ Lu | γ, β- | 6.7 d | reaktor |

PET



PET

- radioisotopes emitting positrons
 (β⁺-particles)
- annihilation with electrons
- two co-linear photons with an energy of 511 keV
- detection of both photons at the same time
- resolution about 1 mm³
- low energy \rightarrow better resolution



Izotopy pro PET

| Izotop | Přeměna | Poločas | Zdroj |
|-------------------|-----------|---------|---|
| ¹⁹ F | β+ | 110 min | cyklotron, ¹⁸ O(p,n) ¹⁸ F |
| ¹¹ C | β^+ | 20 min | cyklotron, $^{14}N(p,\alpha)^{11}C$ |
| ⁶¹ Cu | β^+ | 3.3 h | cyklotron, ⁶¹ Ni(p,n) ⁶¹ Cu |
| ⁶⁴ Cu | β^+ | 13 h | cyklotron, ⁶⁴ Ni(p,n) ⁶⁴ Cu |
| ⁶⁶ Ga | β^+ | 9.5 h | cyklotron, ${}^{63}Cu(\alpha,n\gamma){}^{66}Ga$ |
| ⁶⁸ Ga | β^+ | 68 min | generátor, ⁶⁸ Ge(β⁻) ⁶⁸ Ga, 288 d |
| ⁸⁶ Y | eta^+ | 15 h | cyklotron, ⁸⁶ Sr(p,n) ⁸⁶ Y |
| ¹¹⁰ In | eta^+ | 69 min | generátor, 110 Sn(β^{-}) 110 In, 4.11 d |
| | | | |

Každá minuta se počítá

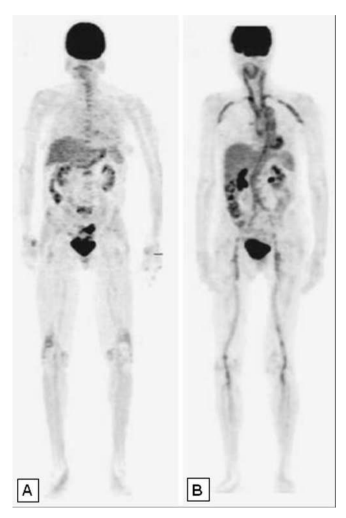
• Příprava izotopu

- Izolace izotopu
- Příprava radiofarmaka
- Separace radiofarmaka
- Analýza radiofarmaka
- Doprava k pacientovi
- Aplikace pacientovi
- Dosažení žádané biodistribuce
- Snímkování

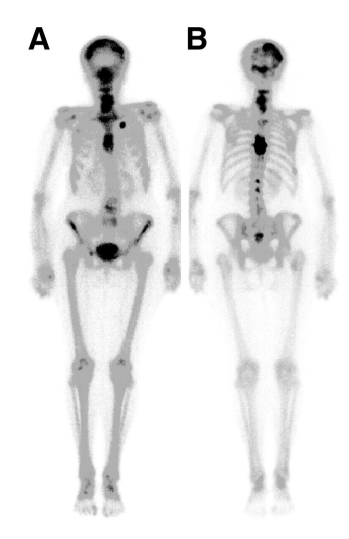


PET vs. SPECT

[¹⁸F]-FDG-PET

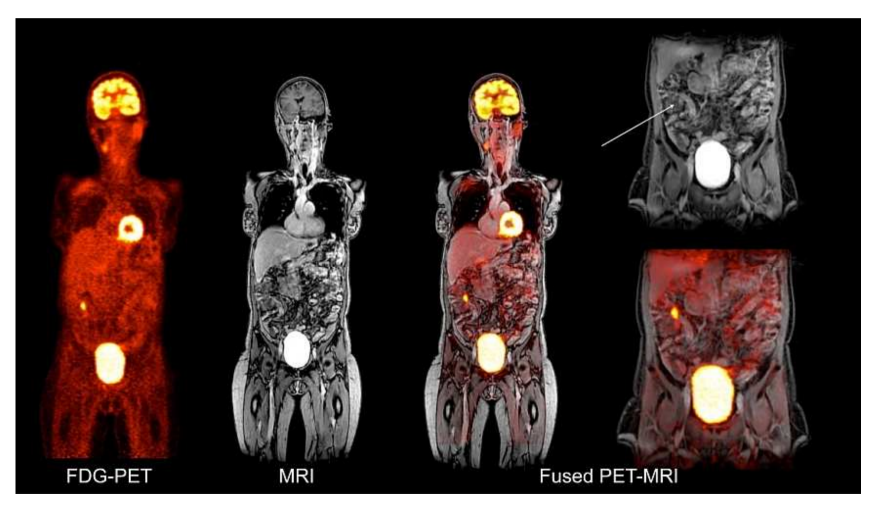


^{99m}Tc-MDP-SPECT

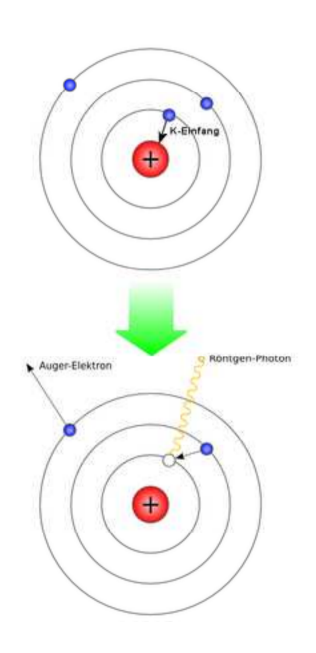


Fused images

- localization in tissues combined techniques with CT or MRI
- fused images PET/CT, PET/MRI, SPECT/CT, SPECT/MRI
- multimodal contrast agents



- Leksell gamma knife
 - focuses the radiation from external sources into tumour
- internal sources of radiation
 - short and defined radius of particles in tissue
 - α-emitters
 - β⁻-emitters
 - γ-emitters with low energy
 - emission of Auger electrons (EC isotopes)
- selective deposition in tumors half-lives in hours



| Isotope | Decay | Half-life | Source | Mean range in tissue |
|-------------------|-------|--------------|--|----------------------|
| ⁶⁴ Cu | β- | 12.8 h | cyclotron, ⁶⁴ Ni(p,n) ⁶⁴ Cu | 0.2 mm |
| ⁶⁷ Cu | β- | 62 h | cyclotron, ⁶⁷ Zn(n,p) ⁶⁷ Cu | |
| ⁶⁷ Ga | Auger | 3.26 d | cyclotron | |
| ⁸⁹ Sr | β- | 50.5 d | reactor | |
| ⁹⁰ Y | β- | 64 h | generator, 90 Sr(β^{-}) 90 Y | 3.9 mm |
| ¹¹¹ Ag | β- | 179 h | cyclotron | 1.1 mm |
| ¹⁴⁹ Pm | β- | 2.2 d | reactor | |
| ¹⁵³ Sm | β- | 1.9 d | reactor | |
| ¹⁶¹ Tb | β- | 166 h | reactor | 0.3 mm |
| ¹⁶⁶ Ho | β- | 1.1 d | reactor | |
| ¹⁷⁷ Lu | β- | 6.7 d | reactor | |
| ¹⁸⁶ Re | β- | 90 h | reactor | 1.1 mm |
| ¹⁸⁸ Re | β- | 17 h | generator, $^{188}W(\beta^{-})^{188}Re$ | 3.3 mm |
| ²¹² Pb | β-/α | 10.6 h/1.01h | reactor | 0.1 mm |

| lsotope | Half-life | Decay mode | Ē _β [MeV] (%) | E _γ [keV] (%) |
|-------------------|-----------|---|---|--------------------------|
| ⁴⁷ Sc | 3.35 days | β ⁻ , γ | 0.14 (68), 0.20 (32) | 159 (68) |
| ⁶⁷ Cu | 2.58 days | β ⁻ , γ 0.19 (20), 0.12 (57) | | 93 (16), 185 (49) |
| ⁹⁰ Y | 2.67 days | β ⁻ 0.93 (100) | | |
| ¹¹¹ Ag | 7.45 days | β ⁻ , γ | 0.36 (93) | 342 (6.7) |
| ¹⁹⁸ Au | 2.7 days | β ⁻ , γ | | |
| ¹⁹⁹ Au | 3.1 days | β ⁻ , γ | 0.13 (15), 0.08 (66) | 158 (37) |
| ²¹² Bi | 1.0 h | α | 6.1 (25) (α) | 727 (12) |
| | | β ⁻ , γ | 0.83 (48) (β ⁻) | |
| ²¹³ Bi | 46 min | α, γ | 5.8 (2) (α), 0.49 (65) (α), 0.32 (32) (α) | 440 (27) |
| ²²⁵ Ac | 10 days | α | 5.83 (51) (α) | |
| Lanthania | des | + | • | + |
| ¹⁴⁹ Pm | 2.21 days | 21 days β ⁻ , γ 0.37 (97) | | 286 (2.9) |
| ¹⁵³ Sm | 1.95 days | β ⁻ , γ | 0.23 (43), 0,20 (35) | 103 (28) |
| ¹⁶⁶ Dy | 3.40 days | β ⁻ , γ | 0.12 (91) | 82.5 (13) |
| ¹⁶⁶ Ho | 1.12 days | β ⁻ , γ | 0.69 (51), 0.65 (48) | 80.6 (6.2) |
| ¹⁷⁷ Lu | 6.71 days | β ⁻ , γ | 0.15 (79) | 208 (11) |

Common criteria for radiopharmaceuticals

- Selected molecule must be amenable to radiolabelling. Reaction must provide sufficient radiochemical yield, specific activity and must proceed in appropriate time, that means maximally 4, ideally less than 3 half-lives of radioisotope also depends on half-life itself. Reaction must proceed under reasonable conditions because of automation of procedure in the case of clinical production. Procedure including yield must be reliable and reproducible.
- **Biodistribution** of a radiopharmaceutical must be related to the physiological response to enable measuring functionality of biochemical process under investigation.
- **High affinity** to the target leading to high contrast of a PET image. Interaction between radiopharmaceutical and biomolecules in target tissue must be the major mechanism. Also high specificity for target molecule is essential because interaction with similar targets leads to interference with desired radioactive signal detected by a PET camera.
- The lipophilicity (defined as usual partition coefficient between *n*-octanol and water *log P*) that determines ability to cross cell membranes.
- Optimal passage of lipid bilayers requires $log P \sim 1.5 2$. Higher log P values result in nonspecific binding caused by hydrophobic interactions with lipids and proteins.
- Certain properties as passage across the cell membrane or other barriers like blood brain barrier (BBB). Besides mentioned lipophilicity, also active transport of compounds must be taken in account, e.g. dopamine, serotonin and amino acids.

Common criteria for radiopharmaceuticals

In general, a **low affinity to P-glycoprotein** (P-gp) is a desirable property for most radiopharmaceuticals. P-gp is an ATP-dependent efflux pump naturally expressed in BBB. It can be over-expressed in tumours. P-gp transports compounds that have high affinity for the pump out of the cell and then radiotracers that have high affinity for P-gp show little accumulation in tissues like brain and tumour.

Metabolism of a radiopharmaceutical is a very important point. Rapid metabolism is generally undesirable. Metabolites can then bind to other molecules or take part in other metabolic processes and result in non-specific accumulation of radioactivity. It is preferable to have the radioisotope in the part of molecule which reaches the target at first and after that is further metabolised. In some cases, metabolism of radiotracer is the mechanism underlying the accumulation of radioactivity in tissue.

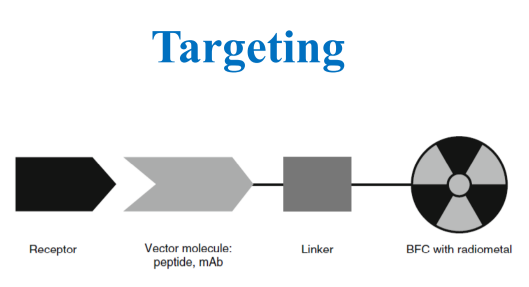
Clearance of non-specifically bound radioactivity by the time of measurement PET must be discriminated. This is relevant mainly for labelled macromolecules that slowly diffuse into cells and only small portion is bound to the target of interest. The unbound radiolabelled molecules must leave the cells again and be cleared from the circulation.

Mutagenicity and toxicity, despite the radiophramaceutical is prepared under non-carrier added (NCA) conditions and small mass is administrated to the patient, must be tested. This differs in different countries according to the law. Usually toxicity tests on rodent species and Ames test for mutagenicity are performed at dosages much higher than those applied in PET studies.

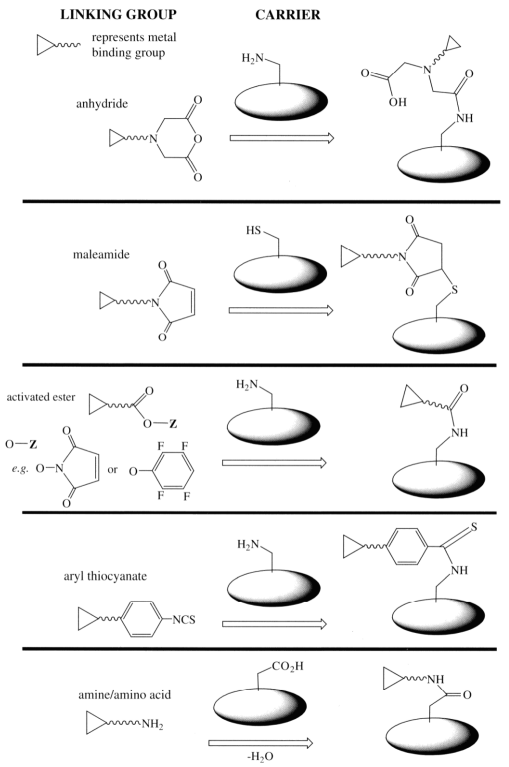
Preparation and administration

RCY – RadioChemical Yield CA – Carrier Added NCA – Non-Carrier Added





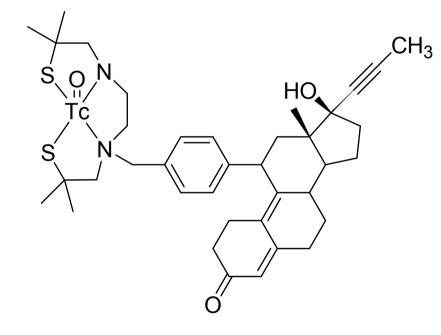
- vector selectivity for imaged tissue: peptide, oligosacharide, etc.
- chemically sensitive labelling at mild conditions

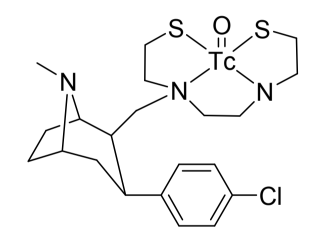


Targeting

• include a biologically active molecule covalently linked to the complex

• e.g.

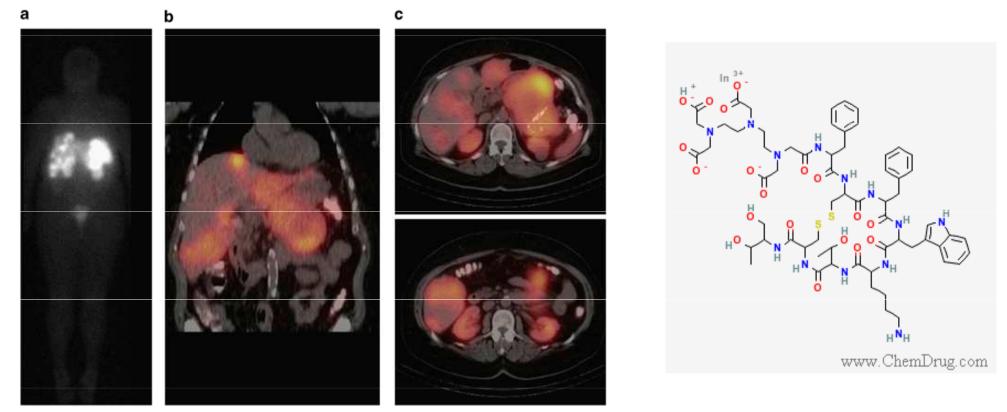




Progesterone receptor analogues (prostate cancer) Cocaine analogues (CNS diseases)

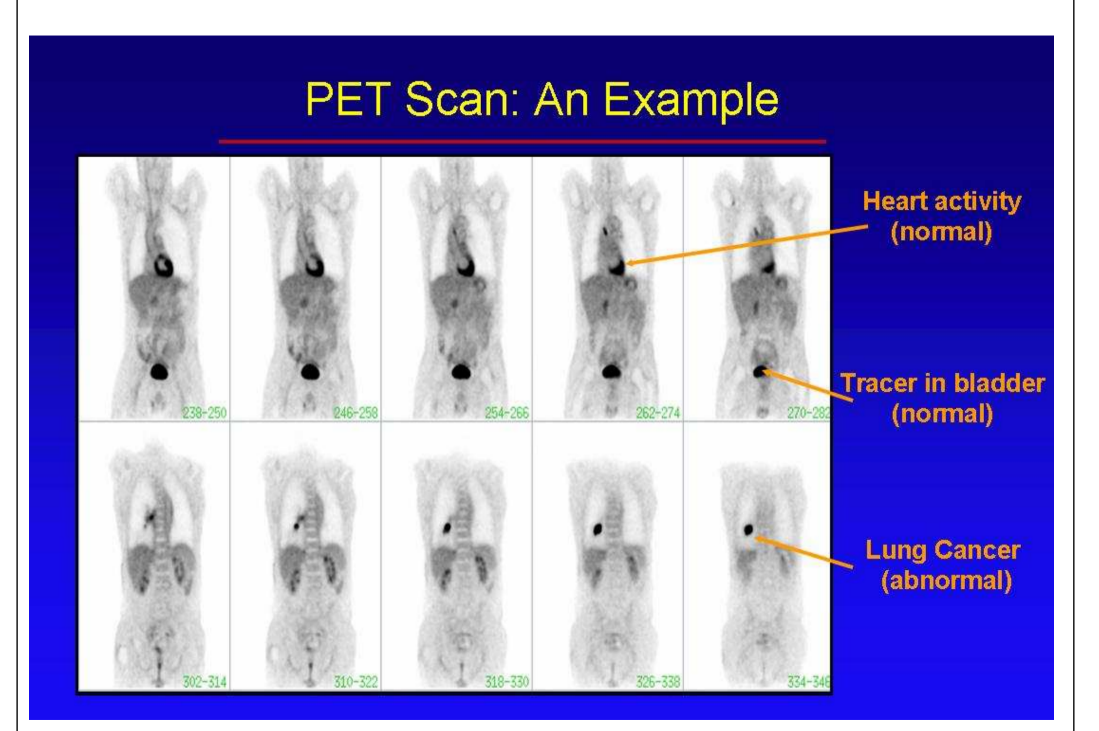
Targeting

Octreoscan – ¹¹¹**In** –**DTPA-Octreotide**

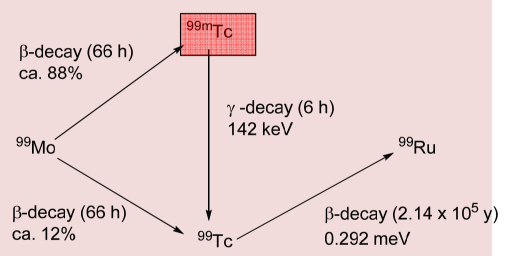


Octreoscan imaging for neuroendocrine tumors.

a) Coronal octreoscan image demonstrating radiotracer uptake in multiple liver metastases and a large pancreatic primary neuroendocrine carcinoma.
(b) Coronal octreoscan fusion images with single photon emission tomography (SPECT) providing increased anatomic detail.
(c) Axial octreoscan fusion images with SPECT.



- predicted by Mendeleev
- first isolated 1938
- 20 isotopes (7 relatively stable)
- used extensively (>90% of all diagnostic nuclear medicine)
- $t_{1/2} = 6$ h
- γ-ray emission energy of 141 keV
- versatile coordination chemistry
- multiple oxidation states
- easily generated from ⁹⁹Mo ($t_{1/2} = 66$ h)

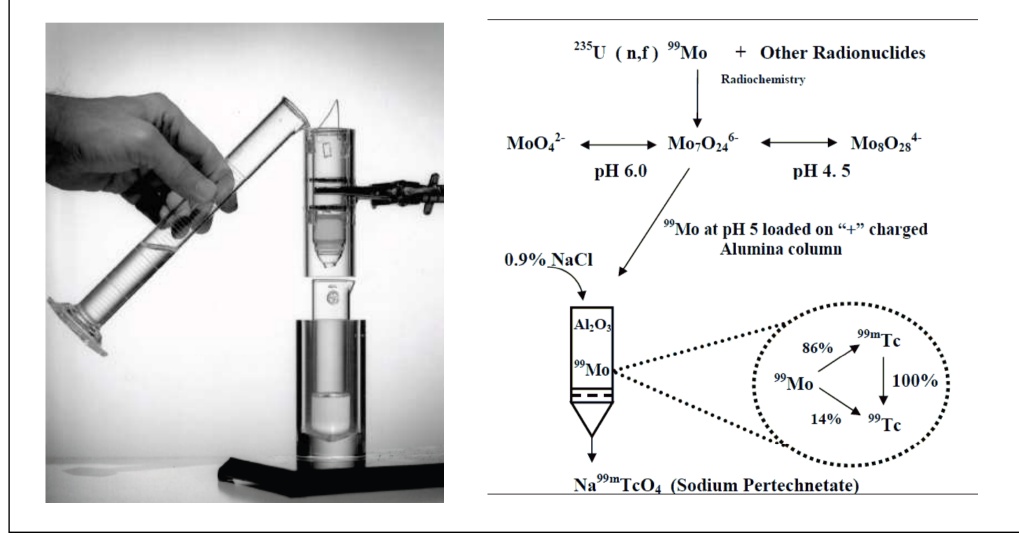


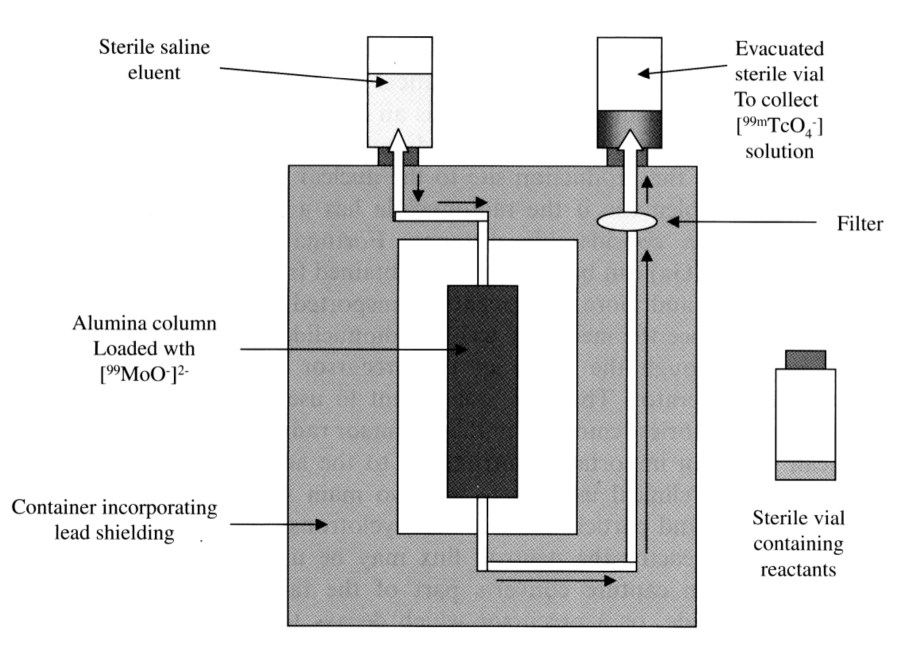
Rhenium

- •¹⁸⁶Re, $t_{1/2} = 90$ h, available from reactor
- ¹⁸⁸Re, $t_{1/2} = 17$ h, available from ¹⁸⁸W(β -)¹⁸⁸Re generator
- chemical properties similar to Terchnetium: diagnostic/therapeutic isotop-pair

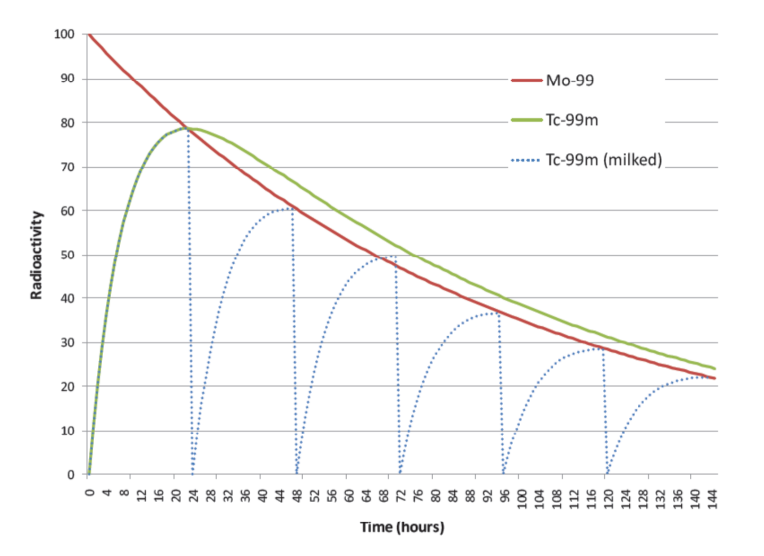
99Mo/99mTc Generator

- patented in 1958
- fission-produced ⁹⁹Mo is processed and purified to anionic molybdate
- loading on the positively charged alumina (Al_2O_3) column





Radionuklidový generátor



Počty elucí z generátoru

^{99m}Tc

Chemistry

- TcO_4^- most stable oxidation state, produced in generator, can not be complexed
- insoluble TcO₂
- reduction with ascorbic acid, FeCl₂, NaBH₄, Na₂S₂O₄, SnCl₂
 - oxidation states IV, $\rm V-$ oxocation technecyl
 - (disproportionation $V \rightarrow IV + VII$)
 - oxidation states I, II, III (oxidation \rightarrow IV)
- stabilization of oxidation states with ligands

Technetium kit

- reducing agent
- coordinating ligand
- antioxidants
- buffers
- lyophilized and sealed under inert atmosphere

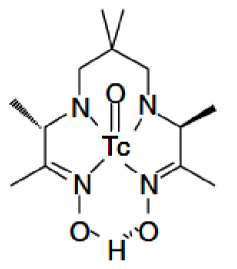
Pharmacology

- bio-distribution and targeting depends much on size and charge:
 - neutral brain
 - cationic hart
 - anionic bones and kidney
- so called technetium essential or first generation agent
- targeting of other organs requires designer ligands:
 - must traverse the blood brain barrier
 - moderately lipophilic
 - neutral charge

Neutral complexes

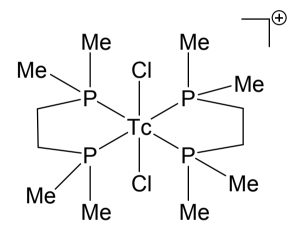
- brain imaging
- oxidation states IV, V

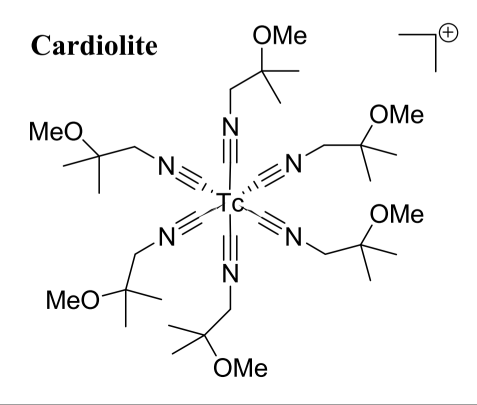
C₂H₅OOC COOC₂H₅ ٧Н S S

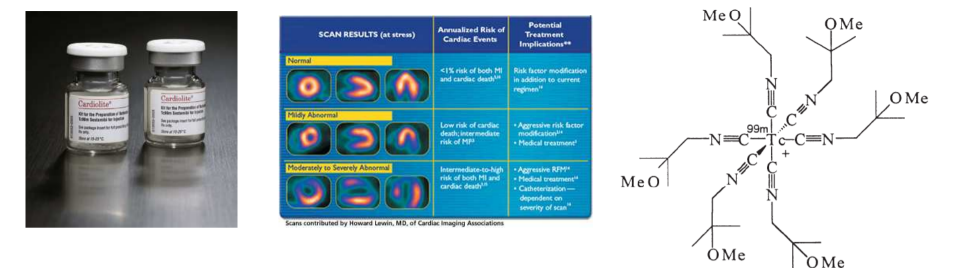


Cationic complexes

- hart imaging
- uptake via Na-K ATPase pump as K⁺ mimics







Cardiolite

Each 5 mL vial contains a sterile, non-pyrogenic, lyophilized mixture of:

- Tetrakis (2-methoxy isobutyl isonitrile) Copper (I) tetrafluoroborate 1.0 mg
- Sodium Citrate Dihydrate 2.6 mg
- L-Cysteine Hydrochloride Monohydrate 1.0 mg
- Mannitol 20 mg
- Stannous Chloride, Dihydrate, minimum (SnCl₂•2H₂O) 0.025 mg

Prior to lyophilization the pH is 5.3 to 5.9. The contents of the vial are lyophilized and stored under nitrogen.

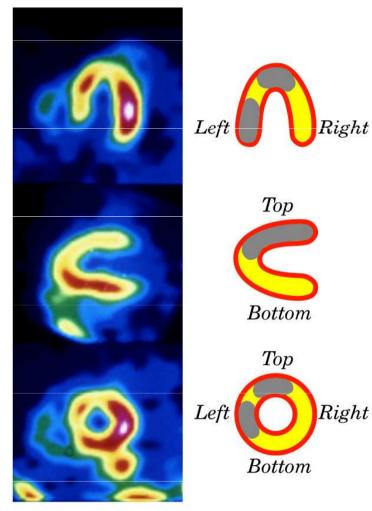
This drug is administered by intravenous injection for diagnostic use after reconstitution with sterile, non-pyrogenic, oxidant-free Sodium Pertechnetate ^{99m}Tc Injection. The pH of the reconstituted product is 5.5 (5.0 - 6.0). No bacteriostatic preservative is present.

The precise structure of the technetium complex is ^{99m}Tc[MIBI]⁶⁺ where MIBI is 2-methoxy isobutyl isonitrile.

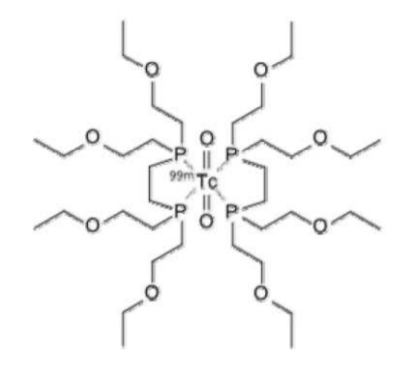
Over 40 million people have received cardiac scans using Cardiolite.

Scans of a human heart under stress taken with the ^{99m}Tc-based imaging agent Myoview[™]

Area with inadequate blood supply give less intense signals (the grey areas on the idealised images)



Imaging agents in action

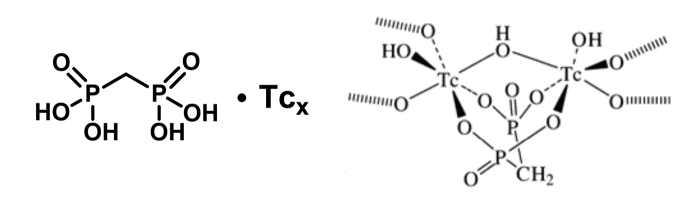


Myoview = tetrofosmin

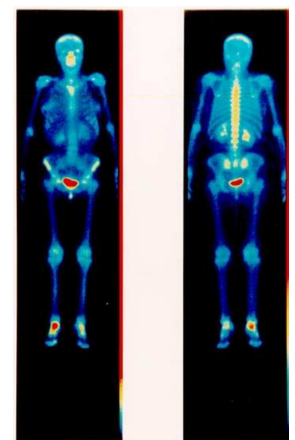
Heart scans courtesy of Amersham plc

Bone imaging

- hydroxyapatite principal mineral component of bones $Ca_{10}(PO_4)_6(OH)_2$
- phosphate (PO_4^{3-}) and pyrophosphate ($P_2O_7^{2-}$) bone seeking anions
- diphosphonates give improved performance



- absorption via calcium coordination to phosphonate
- stressed bone has higher calcium concentration
- main use to detect cancer metastasis into bone



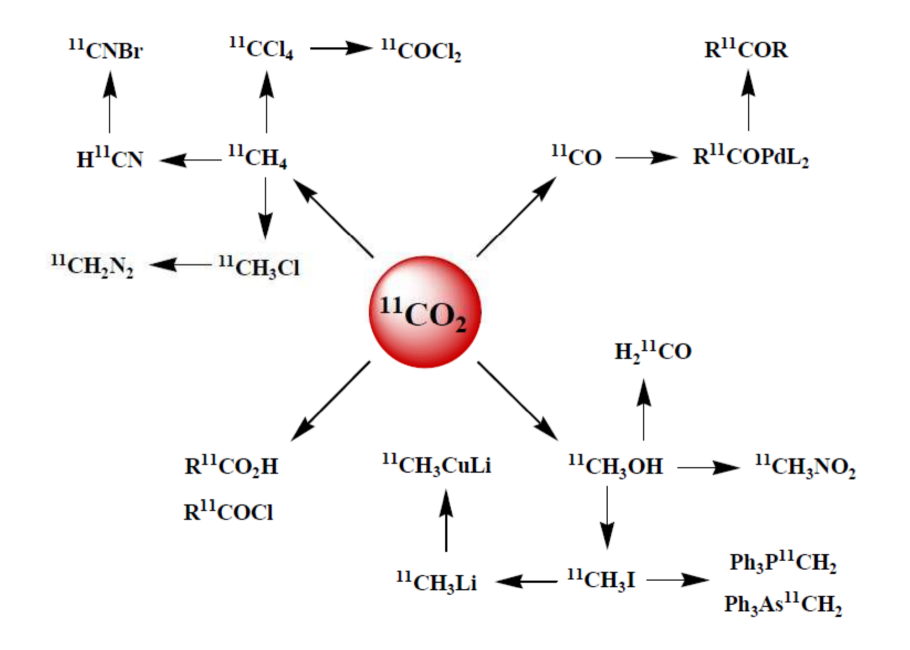
arthritic right ankle

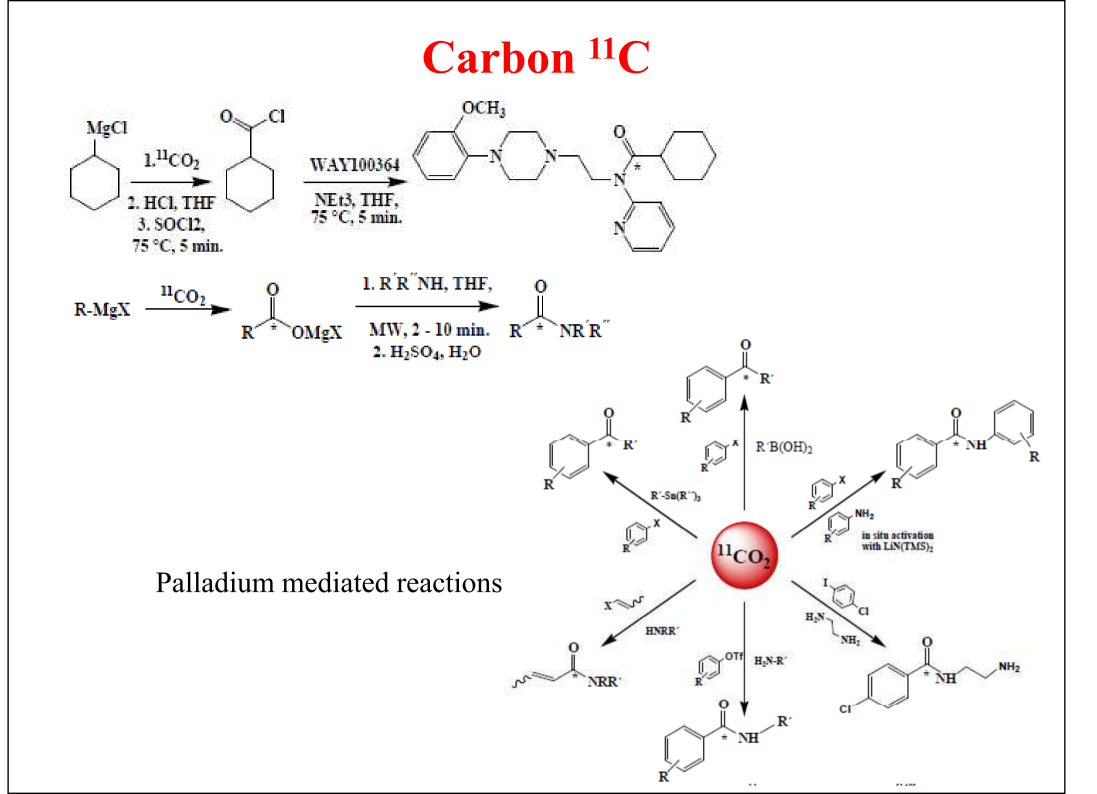
- half-life 20.40 min
- decay mode: 99.8 % $\beta^{\scriptscriptstyle +},$ 0.2 % EC
- max β^+ energy: 0.96 MeV
- range in tissue: 0.96 mm
- decay product: ¹¹B

Production

- \bullet cyclotron-generated: mainly produced by the proton bombardment of $^{14}\mathrm{N}$
- ${}^{14}N(p,\alpha){}^{11}C$ nuclear reaction
- target gas:

2% O_2 in $N_2 \rightarrow {}^{11}CO_2$ 5% H_2 in $N_2 \rightarrow {}^{11}CH_4$

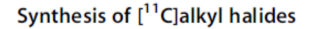


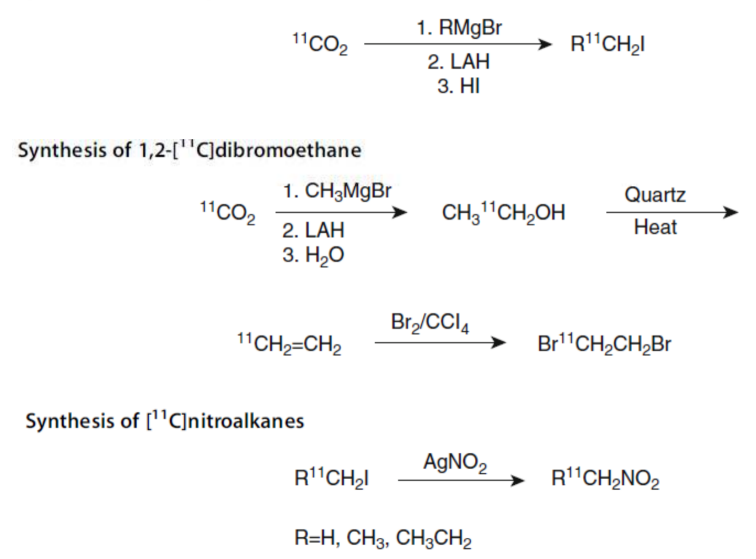


 $^{11}CH_{3}I$

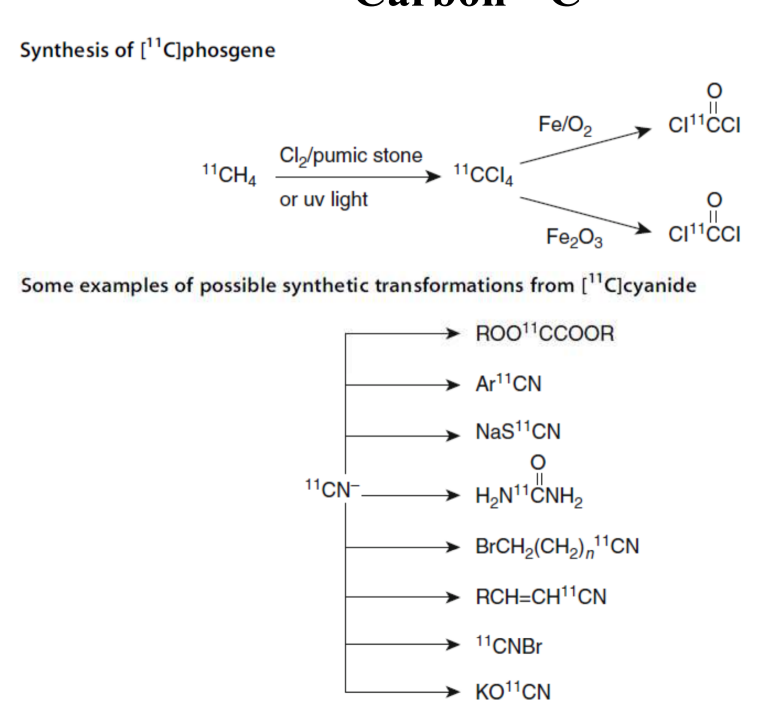
• methylation of N-, O-, S- compounds

¹¹CO₂
$$\xrightarrow{\text{Ni/H}_2}$$
 ¹¹CH₄ $\xrightarrow{\text{I}_2}$ ¹¹CH₃I $\xleftarrow{\text{HI}}$ ¹¹CH₃OH $\xleftarrow{\text{LAH}}$ ¹¹CO₂
 $\xrightarrow{\text{-IP(Ph)_3CH_3^+}}$
 $\xrightarrow{\text{-IP(Ph)_3CH$

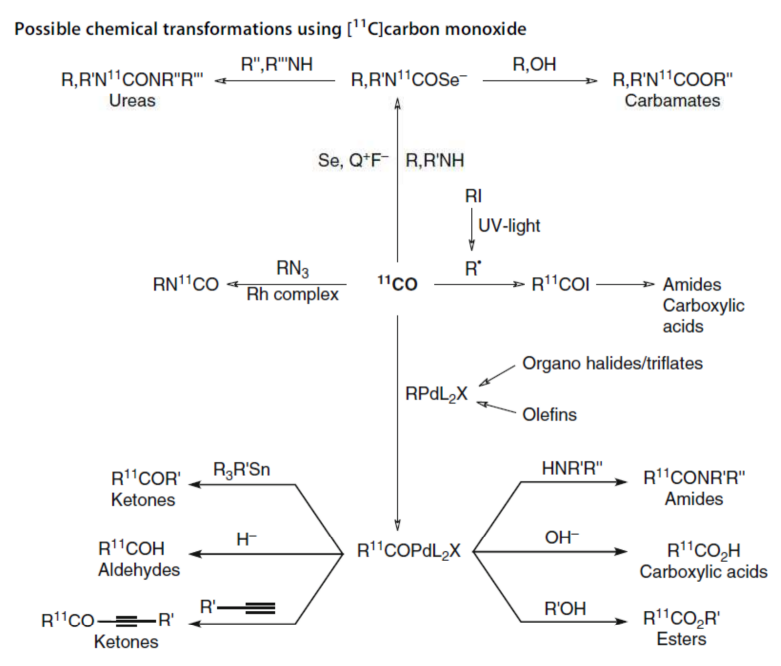




Carbon ¹¹C

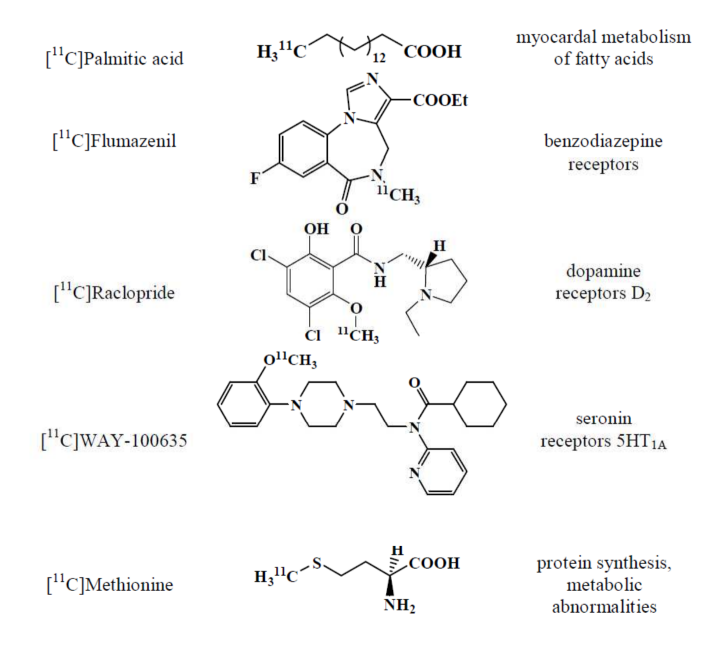


Carbon ¹¹C



X= halide or triflate

Carbon ¹¹C



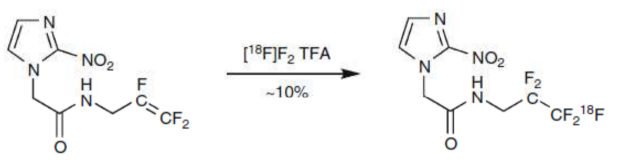
| half-life 109.7 min decay mode: 96.9 % β⁺, 3.1 % EC | | | |
|---|-----------------|----------------|-----------|
| max β⁺ energy: 0.63 MeV range in tissue: 0.54 mm | ¹⁷ F | β^+ | 64.5 s |
| • decay product: ¹⁸ O | ¹⁸ F | β^+ | 109.7 min |
| 18 | ²⁰ F | β^{-} | 11.0 s |
| ¹⁸ ₉ F (110 min.) 97 % β+ 3 % EC ¹⁸ ₈ O (stable) | ²¹ F | β¯ | 4.4 s |
| | ²² F | β ⁻ | 4.1 s |
| | ²³ F | β ⁻ | 2.2 s |
| | ²⁴ F | β ⁻ | 0.34 s |
| | ²⁵ F | β ⁻ | 59 ms |

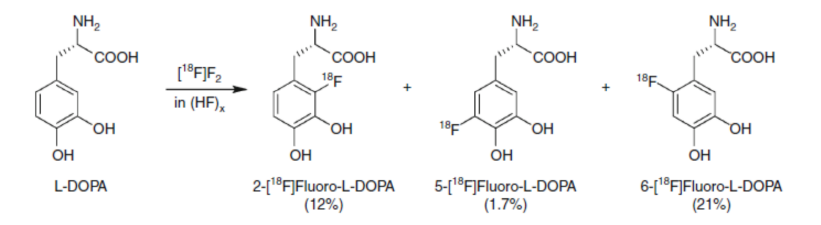
Production

| Product | Target | Beam energy (MeV) | Reaction | Specific activity |
|----------------------------------|---------------------------|-------------------|--|-------------------|
| [¹⁸ F]F ₂ | $0.1 \% F_2/^{20} Ne$ | 18 or 23 | 20 Ne(d, α) 18 F | 30 - 370 MBq/µmol |
| $[^{18}F]F_2$ | $0.1 \% F_2/^{18}O$ | 16 | ¹⁸ O(p,n) ¹⁸ F | 600 MBq/µmol |
| [¹⁸ F]HF | $15 \% H_2/^{20} Ne$ | 14 | 20 Ne(d, α) 18 F | 0.1 - 1 TBq/µmol |
| [¹⁸ F]F ⁻ | ${\rm H_{2}}^{18}{\rm O}$ | 15 | ¹⁸ O(p,n) ¹⁸ F | 0.01 - 7 TBq/µmol |
| [¹⁸ F]F ⁻ | H_2O | 36 | ${}^{6}\text{Li}(n,\alpha){}^{3}\text{H}/{}^{16}\text{O}({}^{3}\text{H},n){}^{18}\text{F}$ | 50 GBq/µmol |
| $[^{18}F]F$ | 2-fluoroaniline | 25 | ${}^{19}F(\gamma,n){}^{18}F$ | not published |

Electrophilic ¹⁸F-Fluorination

- reaction of highly polarized fluorine with an electron rich reactant, e.g., an aromatic system, an alkene, or a carbanion
- starting with $[^{18}F]F_2 50\%$ RCY (molecule compsition $^{18}F-^{19}F$)
- other fluorination agents [¹⁸F]XeF₂, [¹⁸F]AcOF





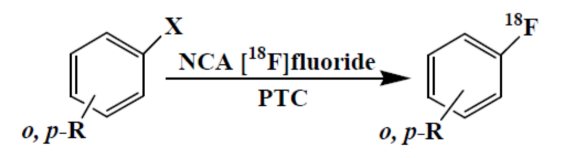
Nucleophilic ¹⁸F-Fluorination

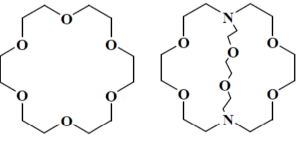
- [¹⁸F]fluoride
- protonation at low pH
- formation of ion pairs with cations decrease of reactivity
- \rightarrow phase transfer catalysis or use of crownethers and cryptands

Nucleophilic Aliphatic Fluorination

$$R-X+F^{-} \xrightarrow{} R-F+X^{-}$$

Nucleophilic Aromatic Fluorination



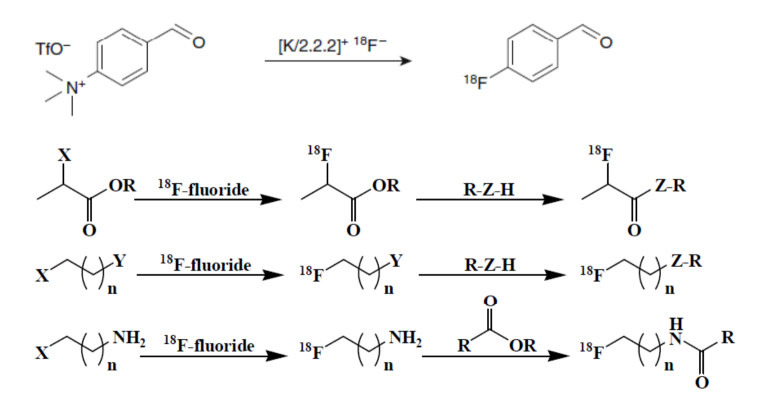


18-crown-6

Kryptofix 2.2.2.

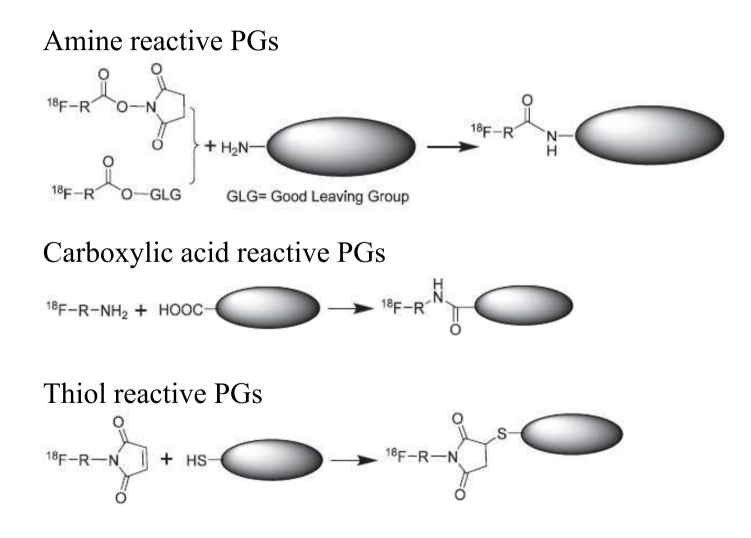
Prosthetic groups (PG)

- labelled compound containing reactive moiety
- fluoromethylation with [¹⁸F]FCH₂I or [¹⁸F]FCH₂Br
- fluoroethylation with $[^{18}F]FCH_2CH_2X$
- other precursors

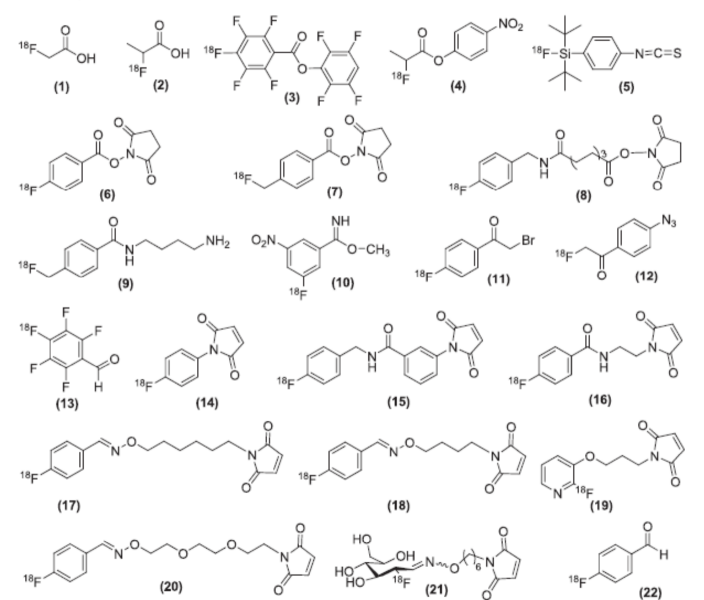


Prosthetic groups for biomolecules

• enable labelling of peptides and proteins



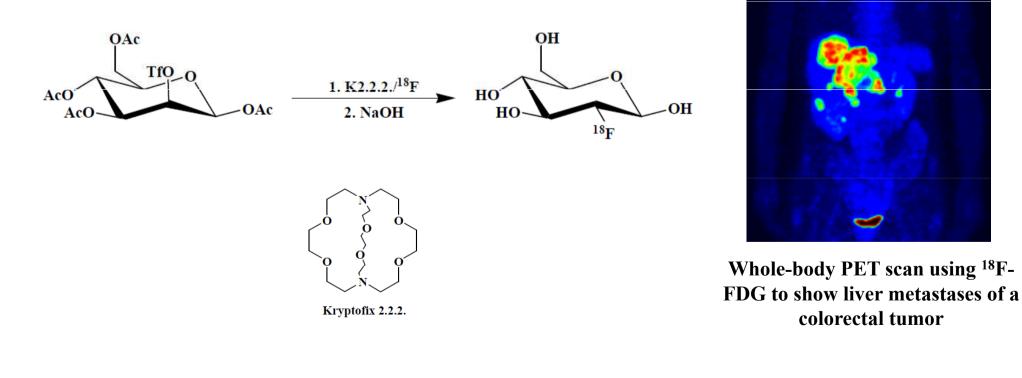
Prosthetic groups (PG)

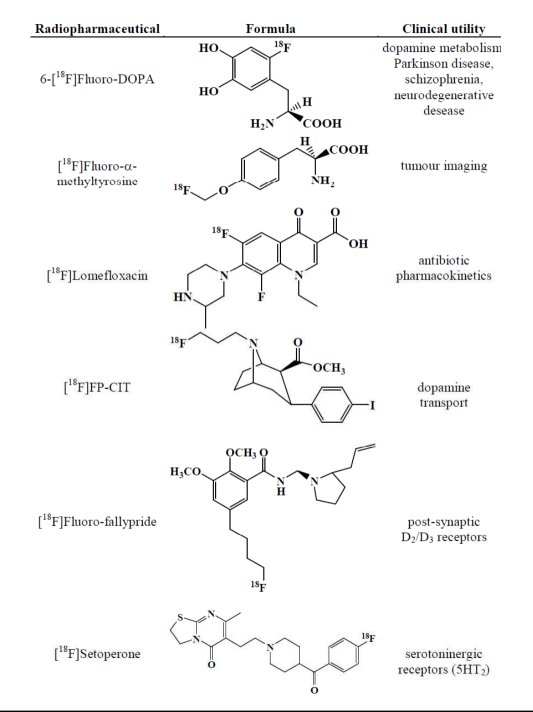


¹⁸F-FDG: [18F]fluoro-2-deoxy-D-glucose – vyrábí se cyklotronicky v MOÚ

colorectal tumor

- 1968, J. Pacák and M. Černý, Department of Organic Chemistry, UK
- the most common PET tracer
- distribution similar to glucose
- can not be metabolized
- accumulation in metabolically-active tissues





Iodine isotopes

| lsotope | Production | Modes of decay | Half-life | E_{γ}/E_{β}^{a} | Specific activity ^b | Application |
|----------------|--------------------|-------------------------|--------------|----------------------------|-----------------------------------|----------------------|
| ¹²³ | Cyclotron | EC | 13.2 h | 159/- | >600 ^c | Imaging |
| ¹²⁴ | Cyclotron | EC/β ⁺ (25%) | 4.18 days | 603/ 1,530 | >30 ^d | Imaging |
| ¹²⁵ | Nuclear reactor | EC | 59.4 days | 35/- | >90 | In vitro and therapy |
| ¹³¹ | Nuclear reactor | β- | 8.04 days | 364/606 | 110 | Imaging and therapy |

¹²³I:
124
Xe(p, 2n) 123 Cs — (β^+ , 0.1 h) $\rightarrow {}^{123}$ Xe — (β^+ , 2.08 h) $\rightarrow {}^{123}$ I

124
I: 124 Te(d, 2n)or(p, n)^{124}I

125
I : 124 Xe(n, γ) 125 Xe — (EC, 17 h) $\rightarrow ^{125}$ I

¹³¹I:
235
U(n, fission)¹³¹I, 99 Mo, 117 Pd, 137 Cs, ...

Metalic nuclides (non-Tc, non-Re)

| Metal | Ionic radius [pm],octahedral | Nuclides |
|--------------------------------------|------------------------------|--|
| Sc ³⁺ Ga ³⁺ | 75 | ⁴⁷ Sc |
| | 62 | ⁶⁶ Ga, ⁶⁷ Ga, ⁶⁸ Ga |
| Y ³⁺ | 90 (105) | ⁸⁶ Y, ⁹⁰ Y |
| ln ³⁺ | 80 | ¹¹¹ ln, ¹¹⁰ ln |
| Dy ³⁺ | 105 | ¹⁶⁶ Dy |
| Ho ³⁺ | 104 | ¹⁶⁶ Ho |
| Lu ³⁺ | 100 | ¹⁷⁷ Lu |
| Bi ³⁺ | 103 | ²¹² Bi, ²¹³ Bi |
| Ac ³⁺ | 126 | ²²⁵ Ac |
| Sm ³⁺ | 110 | ¹⁵³ Sm |
| Cu ²⁺ | 87 | ⁶⁷ Cu, ⁶⁴ Cu, ⁶² Cu, ⁶¹ Cu, ⁶⁰ Cu |
| Ag ⁺ | 129 | ¹¹¹ Ag |

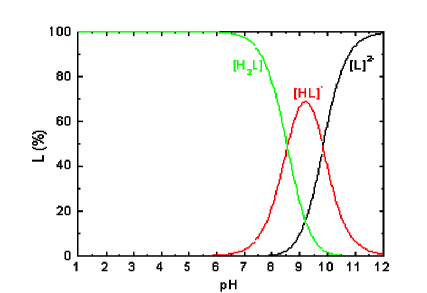
Complex stability

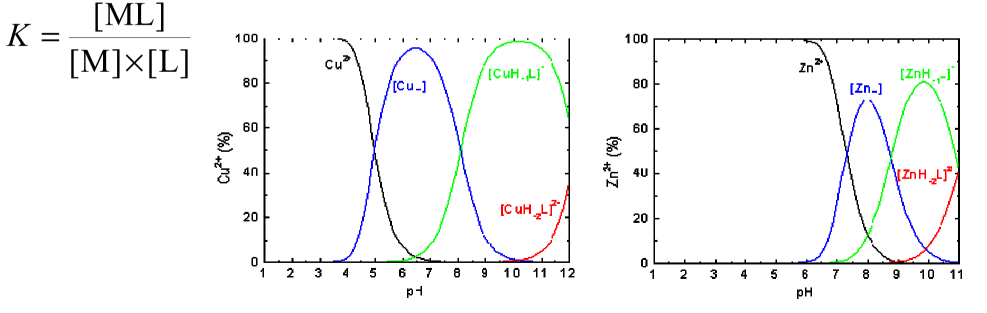
Thermodynamic stability

- proton vs. metal competition
- •ligand basicity

$$K_{a} = \frac{[H] \times [L]}{[HL]}$$

• stability constants





Complex kinetics

Formation kinetic

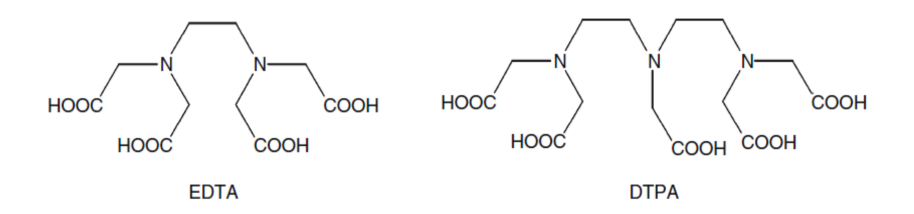
- chemistry high concentrations; NMR, UV-VIS
- radiochemistry low concentrations

Kinetic inertness

- *in vitro* experiments
 - transmetallation (Zn(II))
 - decomplexation in acidic solutions
 - incubation in plood plasma

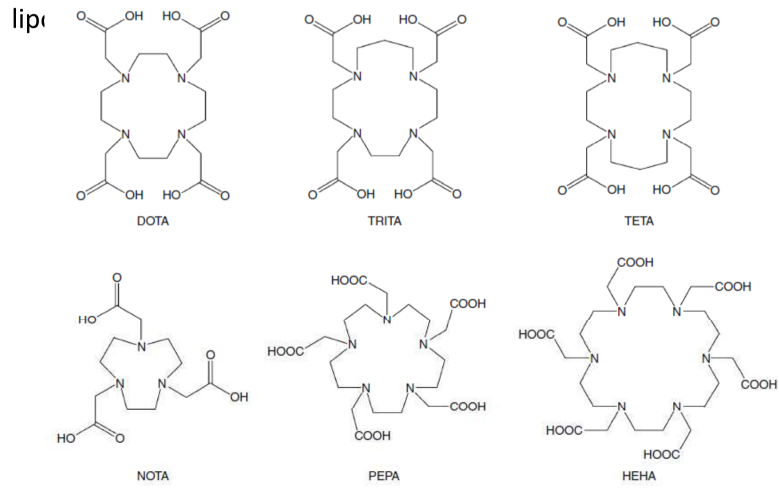
Open-chain ligands

- high thermodynamic stability
- kinetically labile
- fast complexation
- applied in large excess



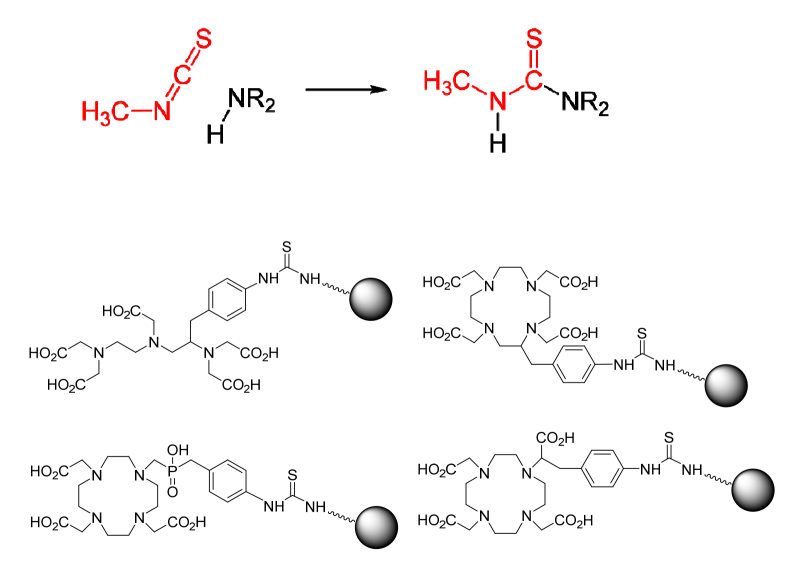
Macrocyclic ligands

- high thermodynamic stability
- kinetically inert
- slow complexation
- variation of pendant arms
 - carboxylates, alcohol, amine, phosphorus derivatives
 - changes in stability, inertness, complexation rate, charge,



Bifunctional chelators

Thiourea bond



Gallium ⁶⁸Ga

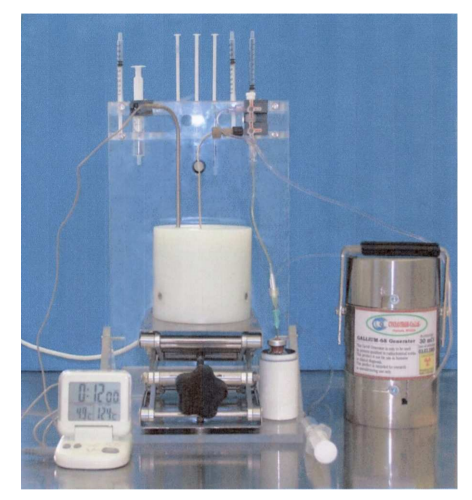
- half-life 67.6 min
- decay mode: 89 % $\beta^{\scriptscriptstyle +}$, 11 % EC
- max β^+ energy: 1.90 MeV
- range in tissue: 2.12 mm
- decay product: ⁶⁸Zn

Generator produced

- source ${}^{68}\text{Ge} (T_{1/2} = 271 \text{ d})$
- adsorbed on TiO_2 or SnO_2
- elution with HCl or citric acid

Chemistry

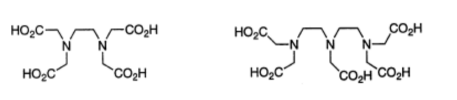
- trivalent
- hexacoordination
- hard metal ion
- precipitation of hydroxide at wide pH range
- formation of tetrahydroxido complex at high pH

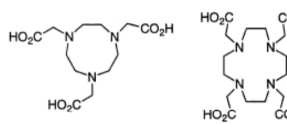


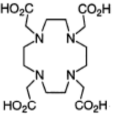
Gallium ⁶⁸Ga

Ligands for ⁶⁸Ga complexation

Aminocarboxylates

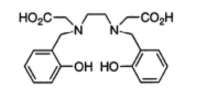


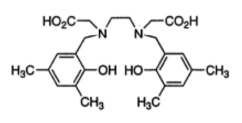


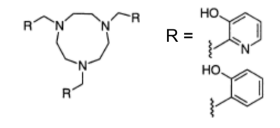


Ga

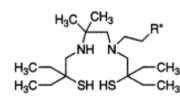
Hydroxybenzyl and hydroxypyridyl derivatives

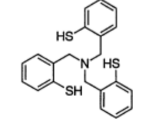


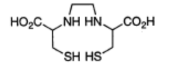


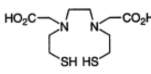


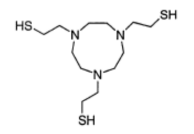
Thiol and amino-thiol chelates











Copper isotopes

| Isotope | Decay | Half-life |
|------------------|------------------------------------|-----------|
| ⁶⁰ Cu | β^+ | 24 min |
| ⁶¹ Cu | β^+ | 3.3 h |
| ⁶² Cu | β^+ | 9.74 min |
| ⁶⁴ Cu | $\beta^{+}(61\%), \beta^{-}(39\%)$ | 12.8 h |
| ⁶⁷ Cu | β- | 62 h |

| Source |
|--|
| cyclotron, ⁶⁰ Ni(p,n) ⁶⁰ Cu |
| cyclotron, ⁶¹ Ni(p,n) ⁶¹ Cu |
| generator, ${}^{62}Zn(\beta^{-}){}^{62}Cu$, 9.3 h |
| cyclotron, ⁶⁴ Ni(p,n) ⁶⁴ Cu |
| cyclotron, ⁶⁷ Zn(n,p) ⁶⁷ Cu |

- isotopes 60, 61, 62 and 64 PET
- isotopes 64 and 67 SPECT and therapy
- difficult production and isolation

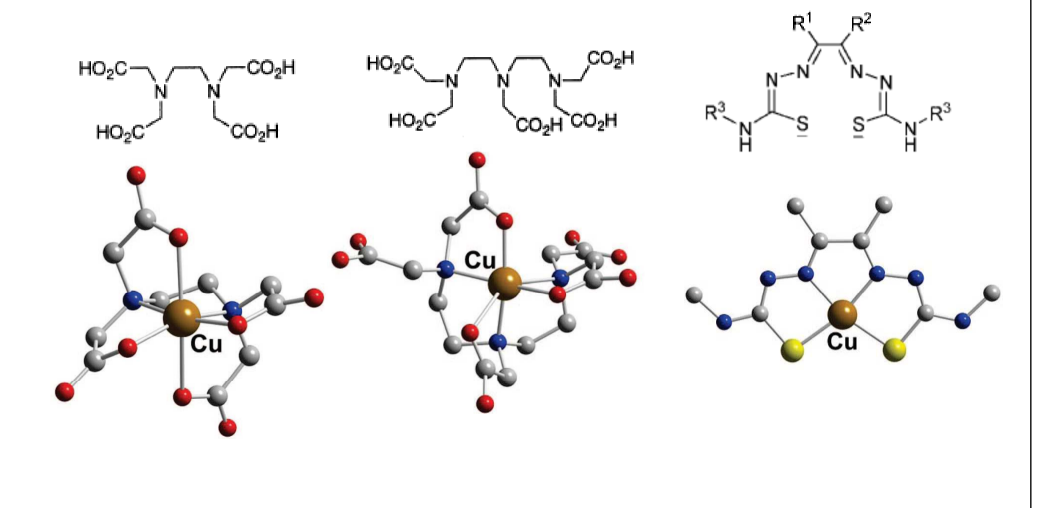
Chemistry

- divalent
- hexacoordination
- Jahn-Teller effect

Copper isotopes

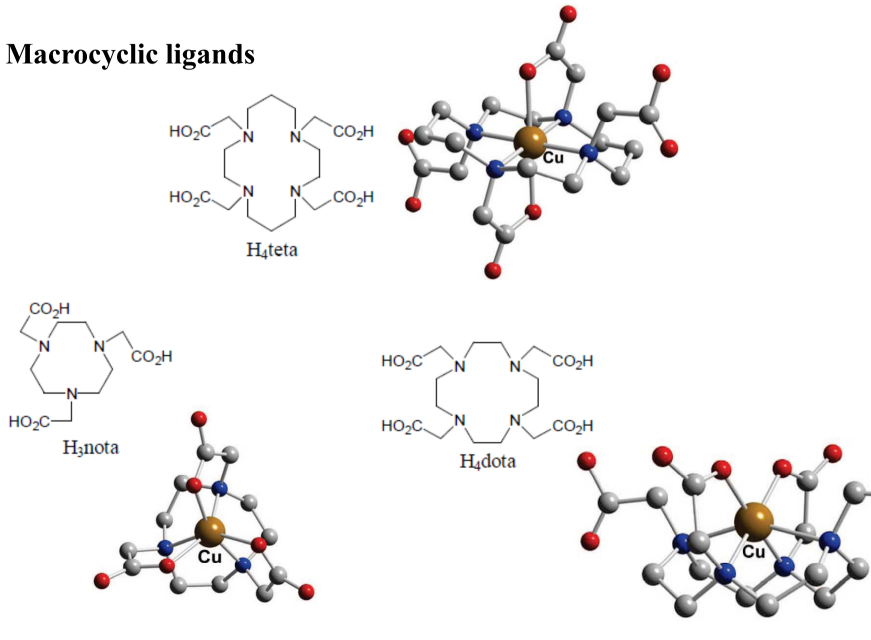
Ligands for copper complexation

Open-chain ligands



Copper isotopes

Ligands for copper complexation



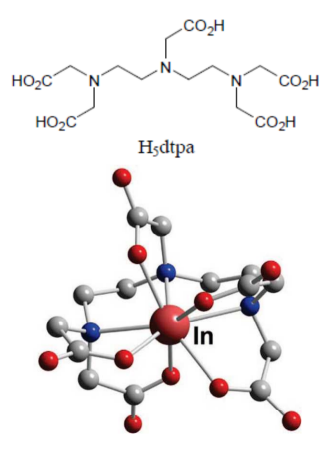
Sc, Y, In and Lanthanoides

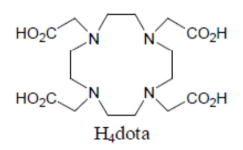
| | Decay | Half-life | Source |
|----------------------|-----------|-----------|--|
| Diagnotic isotopes | | | |
| ⁴⁴ Sc | eta^+ | 3.93 h | generator, ⁴⁴ Ti(EC) ⁴⁴ Sc, 59 y |
| | | | or cyclotron, ⁴⁴ Ca(p,n) ⁴⁴ Sc |
| ⁸⁶ Y | β^+ | 14.7 h | cyclotron, ⁸⁶ Sr(p,n) ⁸⁶ Y |
| ¹¹⁰ In | β^+ | 69.0 min | generator, ${}^{110}Sn(\beta^{-}){}^{110}In$, 4.11 d |
| ¹¹¹ In | γ | 68 h | cyclotron, ¹¹¹ Cd(p,n) ⁶⁴ In |
| ¹³⁴ La | β^+ | 6.70 min | generator, ${}^{134}Ce(\beta^{-}){}^{134}La$, 3.0 d |
| ¹⁴⁰ Pr | β^+ | 3.39 min | generator, 140 Nd(β -) 140 Pr, 3.3 d |
| Therapeutic isotopes | | | |
| ⁹⁰ Y | β- | 64 h | generator, 90 Sr(β^{-}) 90 Y, 28.8 y |
| ¹⁴⁹ Pm | β- | 2.2 d | reactor |
| ¹⁵³ Sm | β- | 1.9 d | reactor |
| ¹⁶¹ Tb | β- | 166 h | reactor |
| ¹⁶⁶ Ho | β- | 1.1 d | reactor |
| ¹⁷⁷ Lu | β- | 6.7 d | reactor |

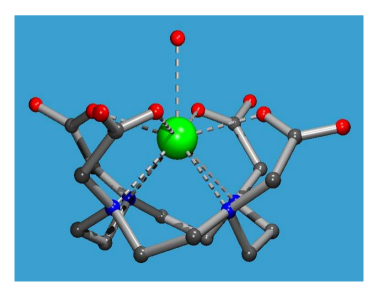
Sc, Y, In and Lanthanoides

Chemistry

- octa- or nona-coordination
- octadentate ligands
- hard metal ions







Bisphosphonate-containing dota-amides

- ligand synthesis and characterization
- chemical complexation study
- labelling (complexation of radionuclide)
 - pH, temperature, ligand concentration

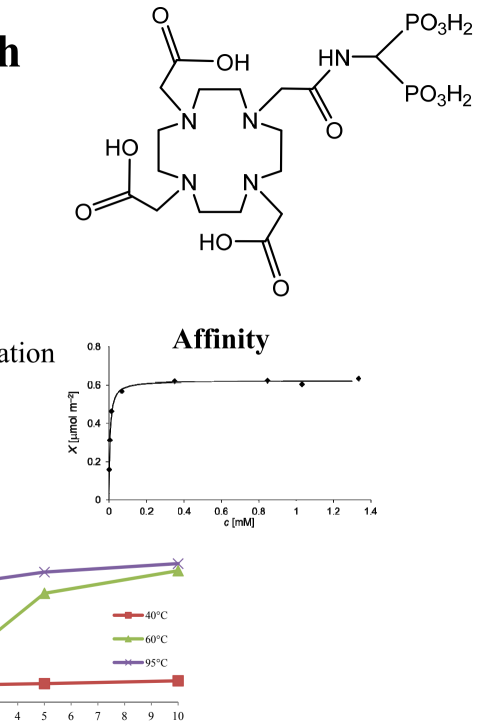
2

3

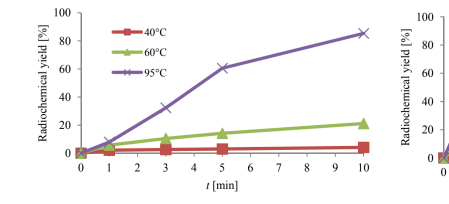
t [min]

1

• affinity and stability study



Labeling



 \cap

НΟ

07

·ОН

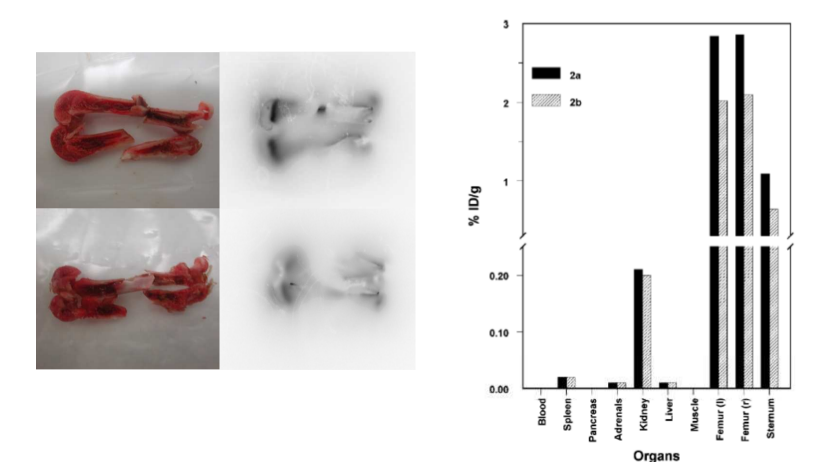
HO

PO₃H₂

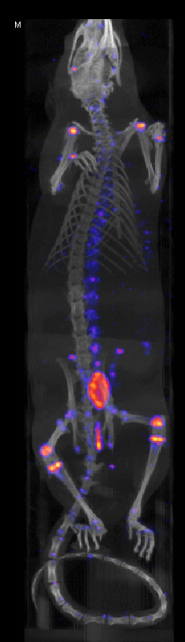
PO₃H₂

HN

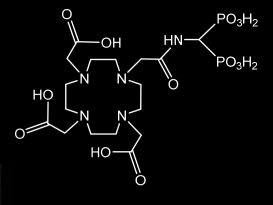
Biodistribution of ¹⁷⁷Lu-complexes in Lewis rat 24 h after injection



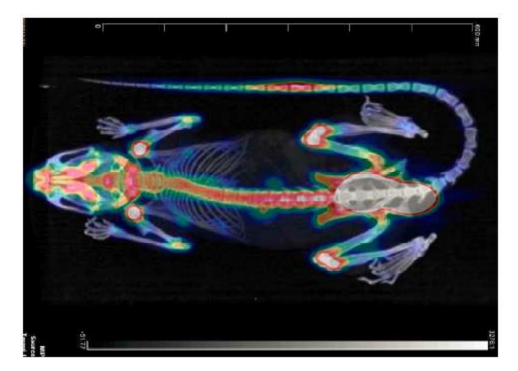
In vivo SPECT/CT visualization of ¹⁷⁷Lu-complex

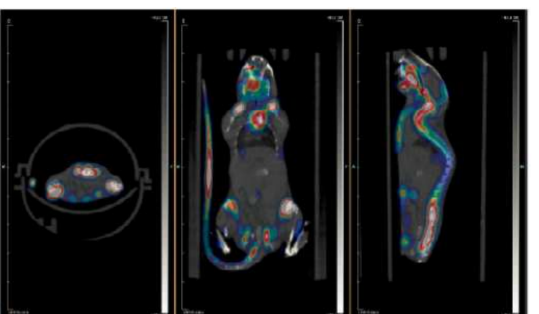


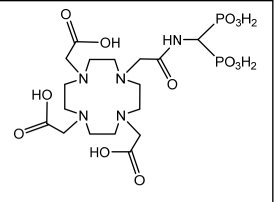




24 h *p.i*.







⁶⁴Cu complex

PO₃H₂

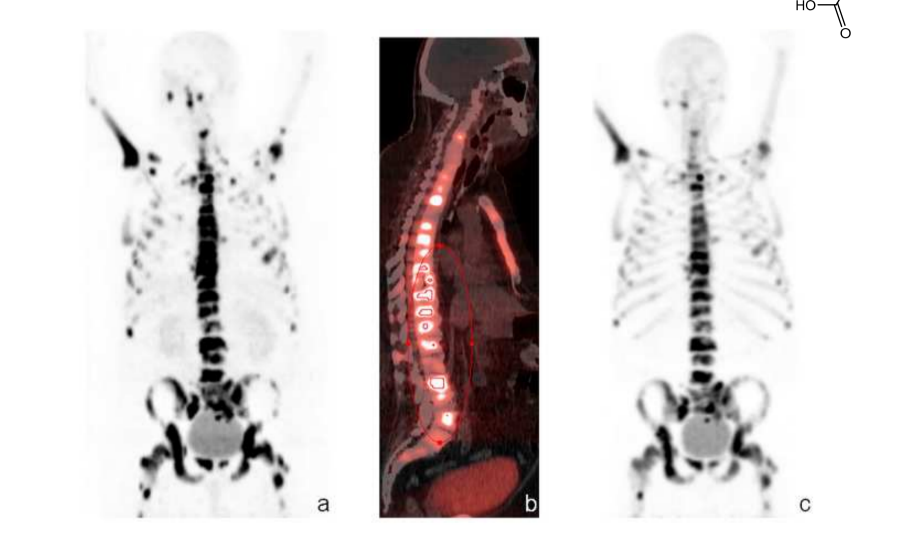
PO₃H₂

HN

OH

ΗΟ

PET/CT imaging of bone metastases with ⁶⁸Ga-complex



 (a) = coronal PET, (b) = sagittal PET/CT. For comparison (c) shows ¹⁸F-fluoride PET. University of Mainz, Zentral Klinik Bad Berka

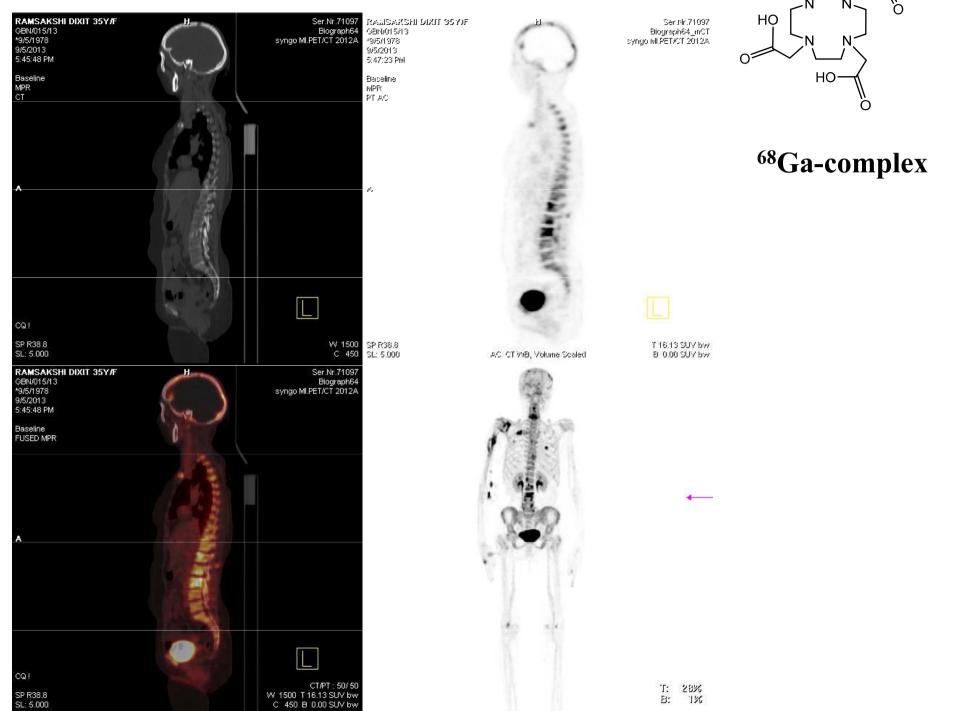
Ο

-OH

PO₃H₂

PO₃H₂

HN



Děkuji za pozornost!



