

Magnetic Resonance Imaging (MRI)



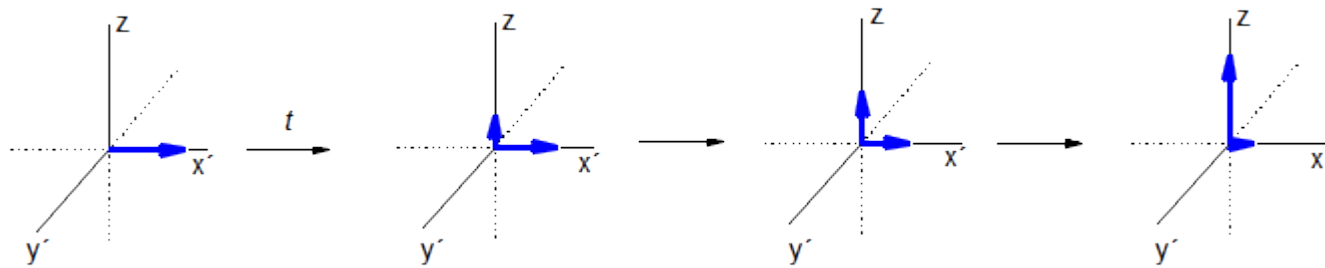
- Non-invasive and safe technique
- Great spatial resolution (μm scale)
- Outstanding diagnostic capability

MR sagittal image of human head

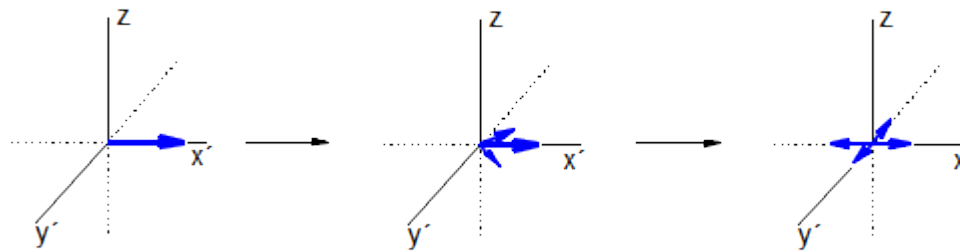
A MR-image represents a map of the intensity of
the $^1\text{H-NMR}$ signal of water protons

The contrast is mainly generated by difference
in the relaxation times (T_1 and T_2) of water protons

T_1 Relaxation (Spin-Lattice Relaxation)



T_2 Relaxation (Spin-Spin Relaxation)



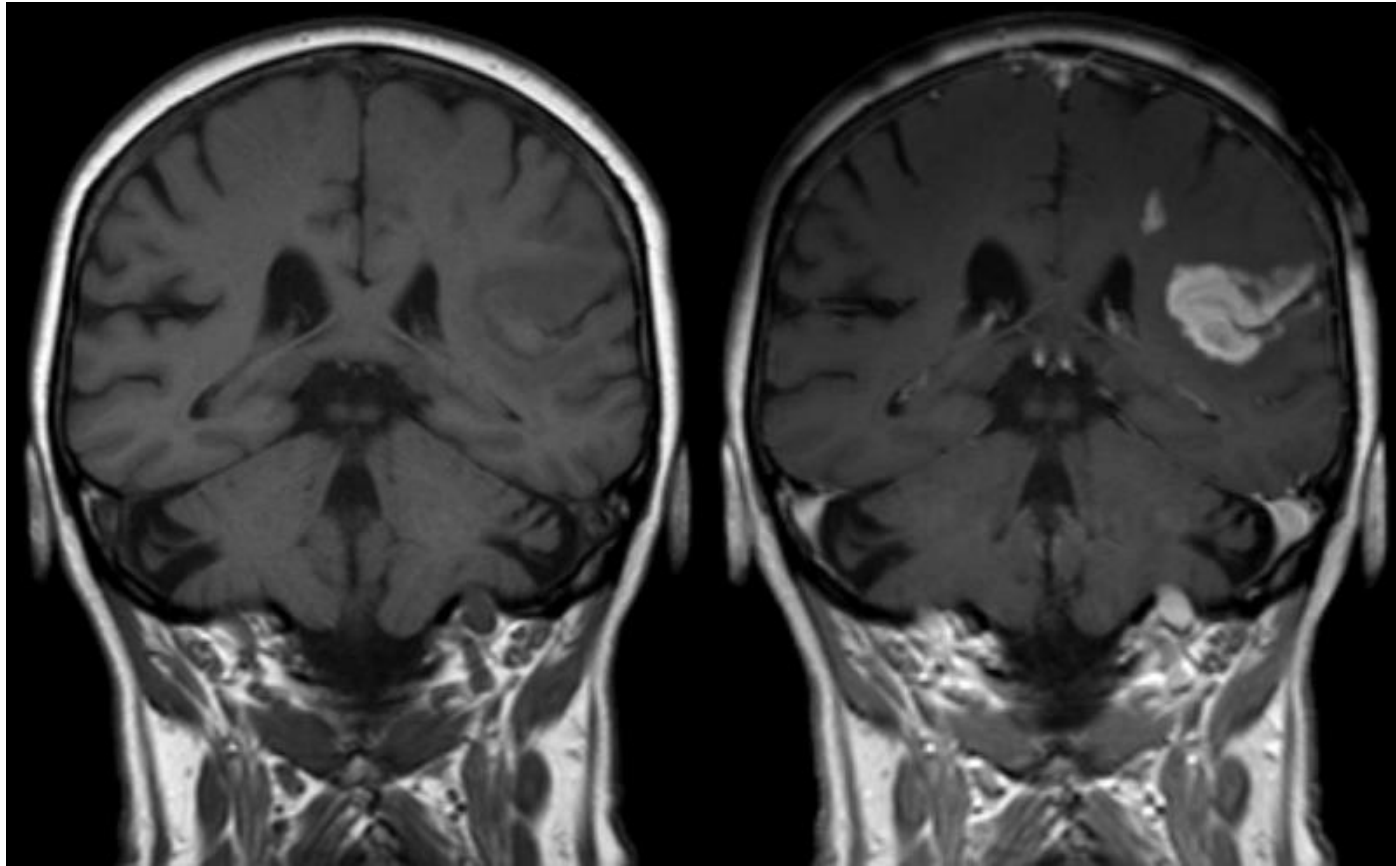
Contrast Agents (CA)

The purpose of a CA is **to reduce T_1 (parallel to B_0) or T_2 (perpendicular to B_0)** in order to obtain an hyper- or ipo-intense signal, respectively, in short times and with a better signal to noise ratio.

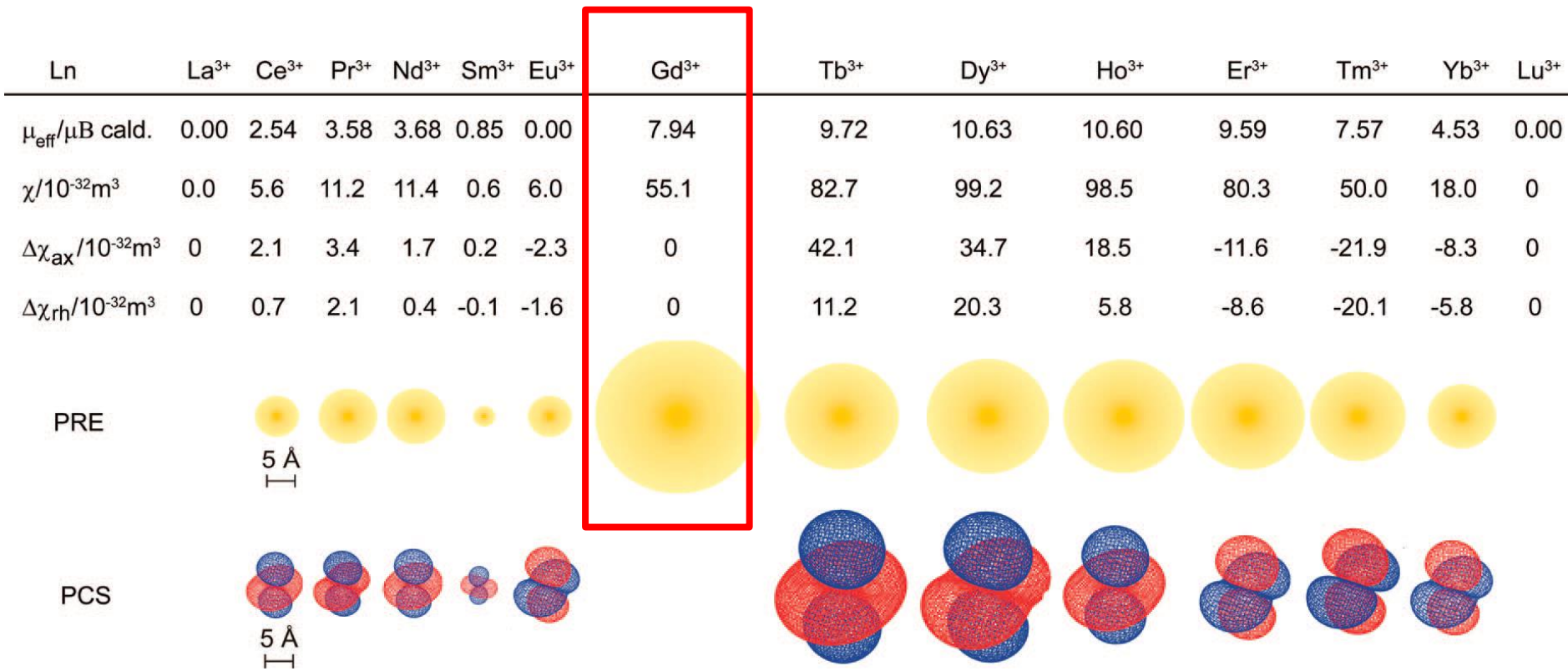
T_1 contrast agents (positive = hyper-intense signal): paramagnetic metal complexes Mn(II), **Gd(III)**

T_2 contrast agents (negative = ipo-intense signal): small super-paramagnetic iron oxide particles (SPIO) and ultra-small super-paramagnetic iron oxide (USPIO)

MRI CA's must have a catalytic (i.e. amplified) effect



Defect of the blood-brain barrier after stroke shown in MRI. T1-weighted images: left image = without; right image = with contrast medium administration



PRE = Paramagnetic Relaxation Enhancement

PCS = Pseudo-Contact Shift

il raggio della sfera gialla indica la distanza alla quale i segnali ¹H NMR subiscono un significativo accorciamento del tempo di rilassamento

>30 million MRI scans/year in USA
(>20 billion \$)

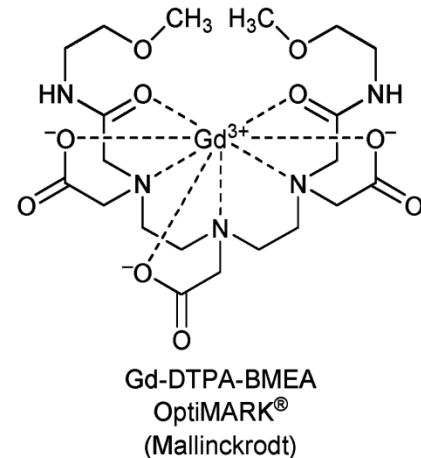
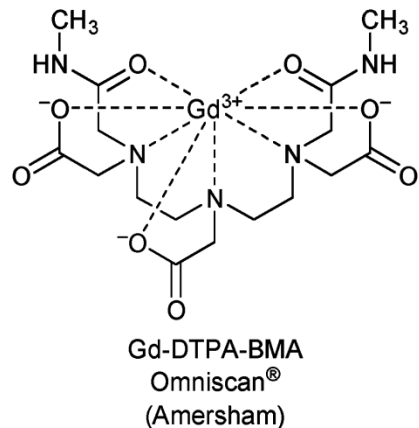
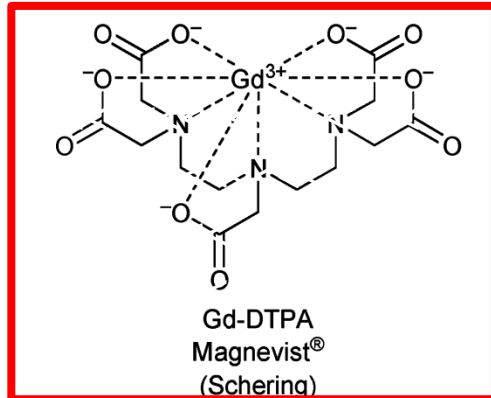
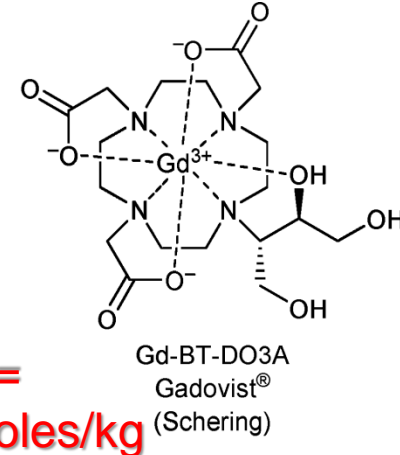
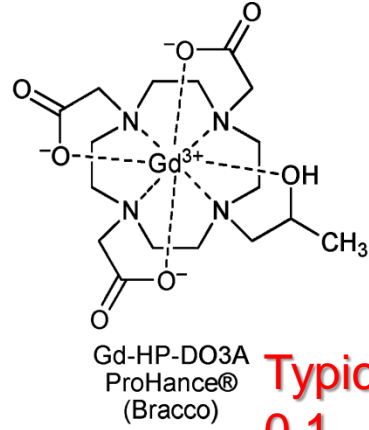
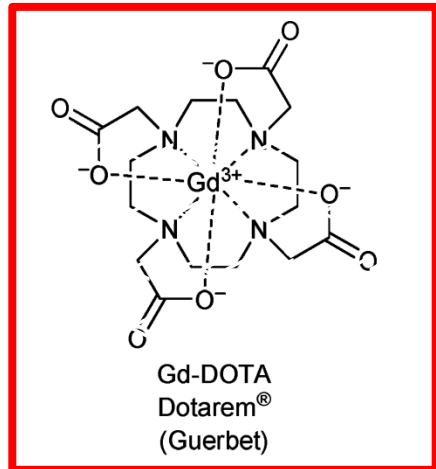
10 million MRI scans/year with Gd CA

9 commercially used Gd CA

The technique has a low sensitivity: **gram quantities** of Gd compounds are used in each scan. This causes toxicity problems (nephrogenic systemic fibrosis)

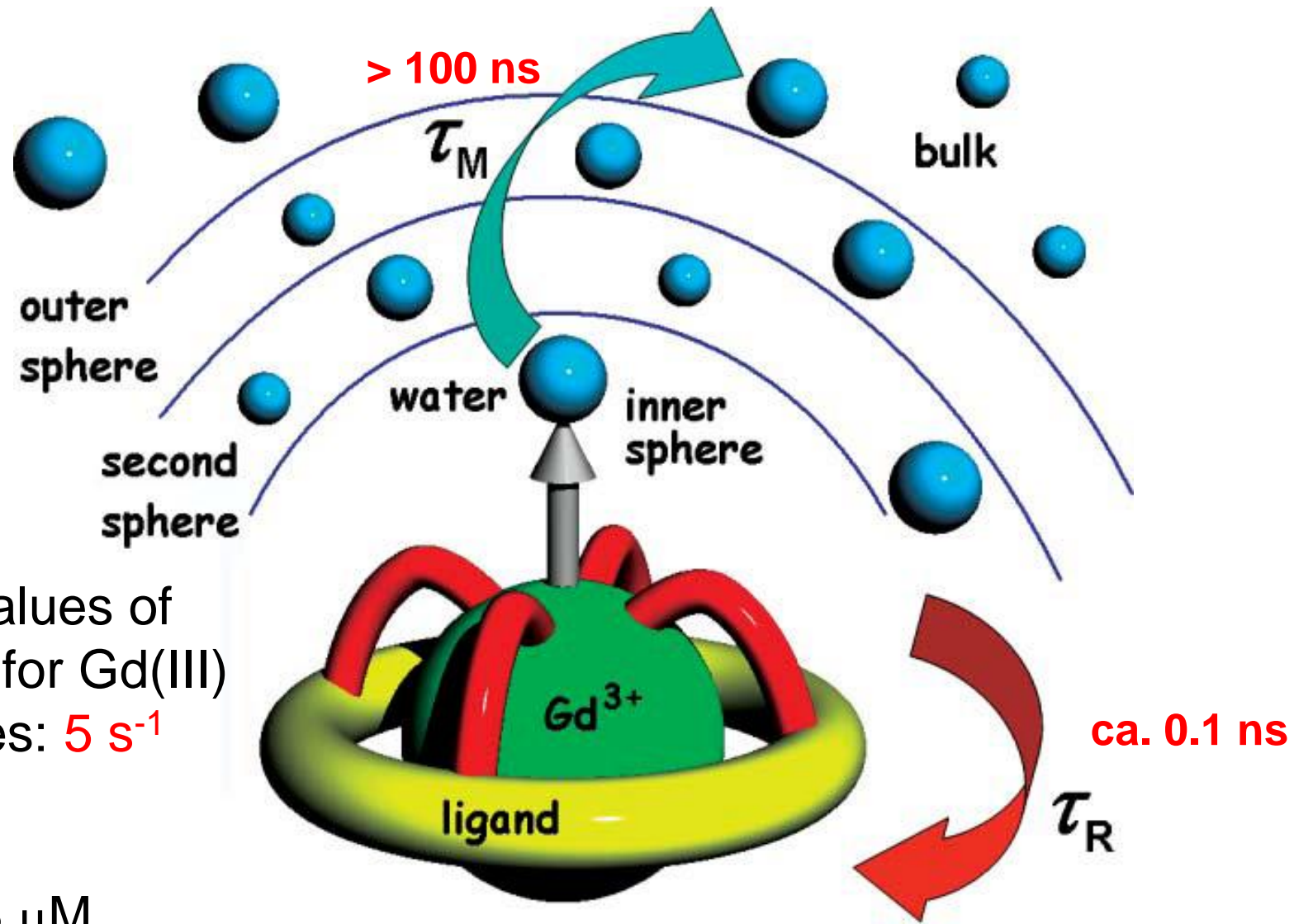
The Gd(III) ion is quite toxic ($LD_{50} = 0.2 \text{ mmol}\cdot\text{kg}^{-1}$)

Some commercial T₁ contrast agents (extracellular fluid CAs)

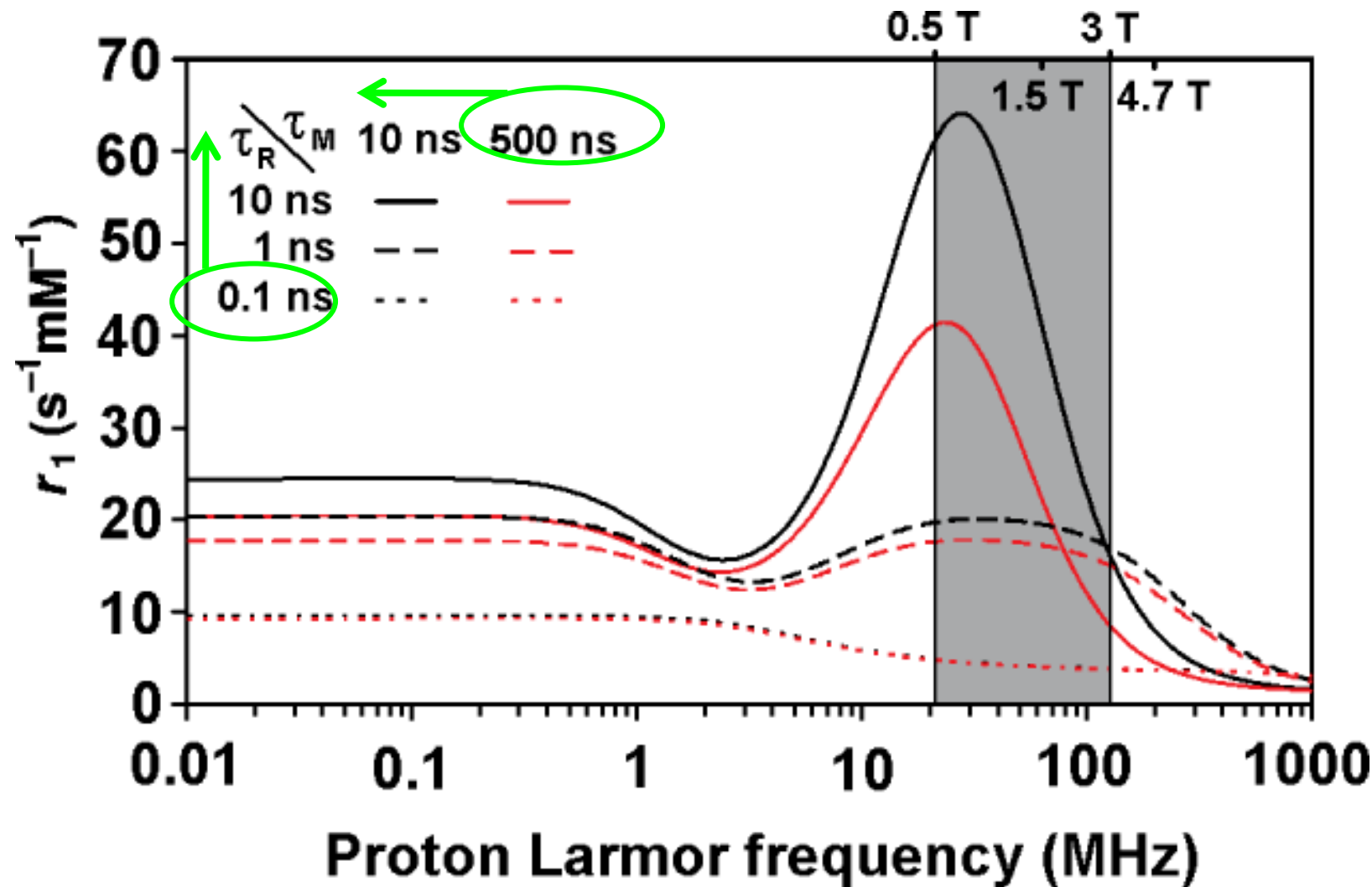


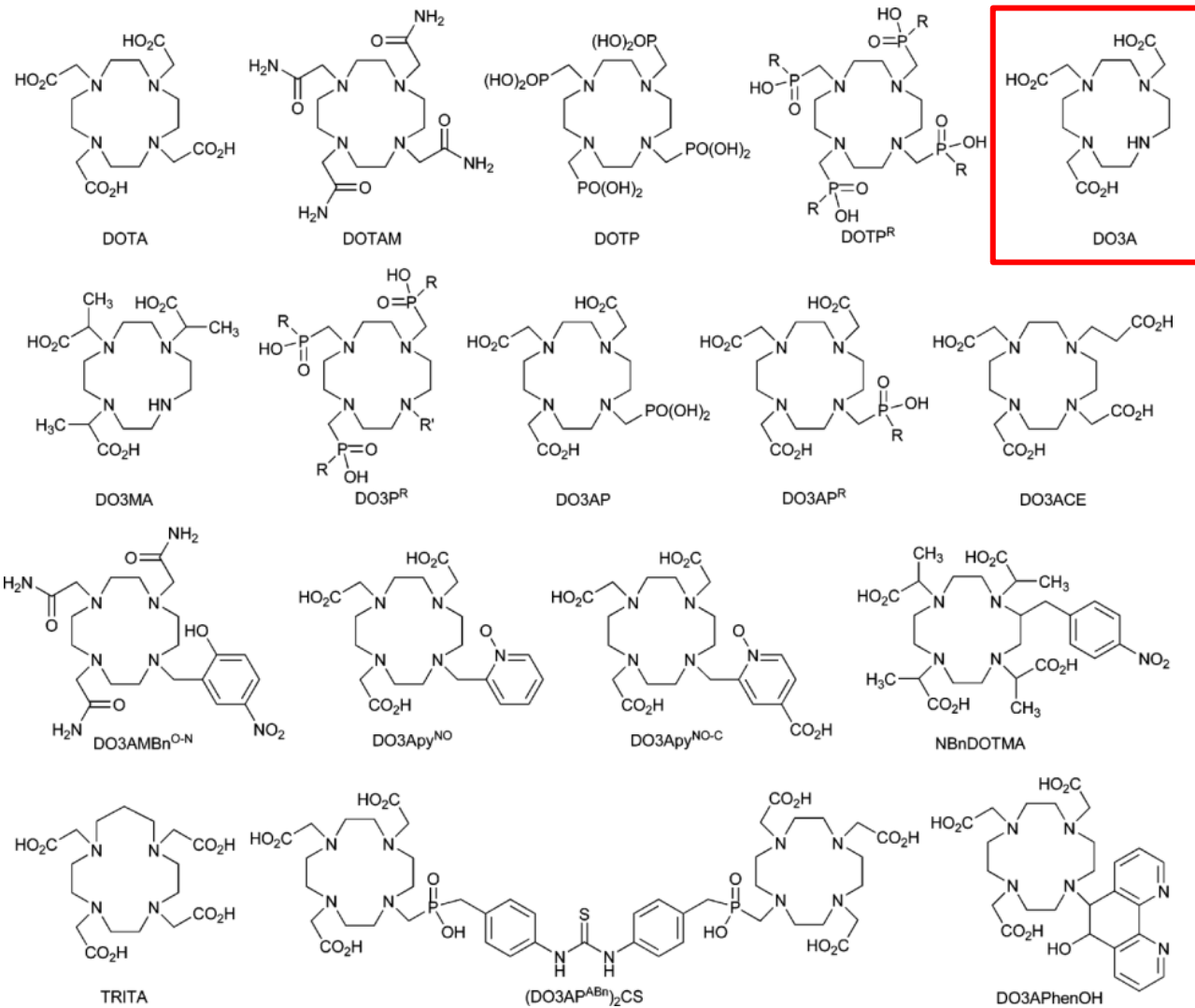
Typical dose =
0.1 – 0.3 mmoles/kg

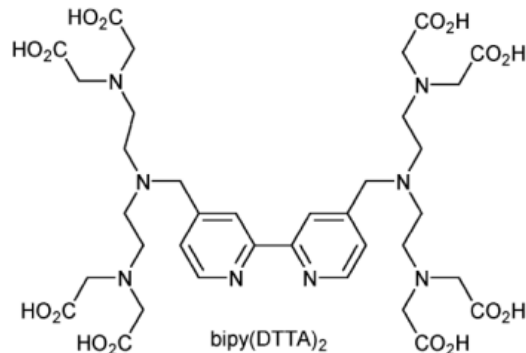
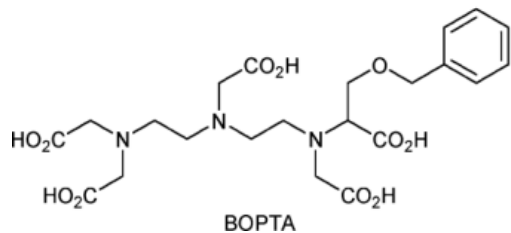
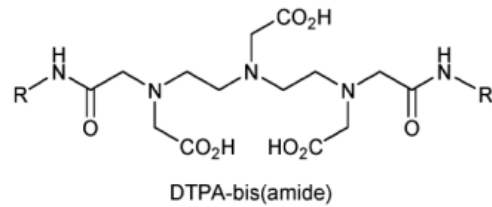
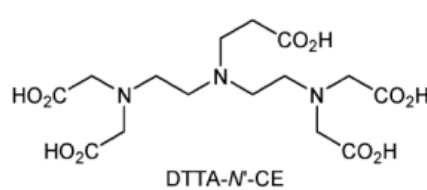
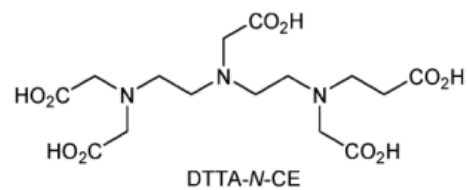
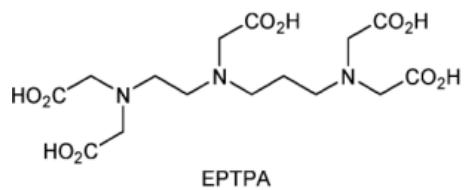
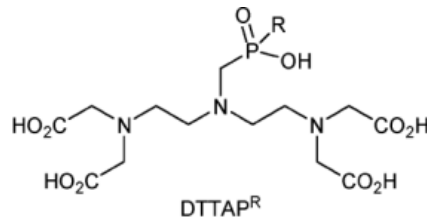
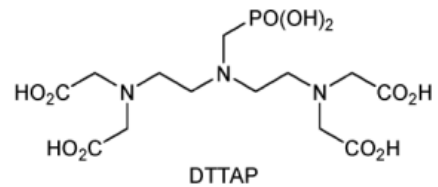
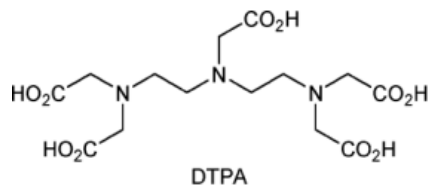
Parameters that affect Relaxivity



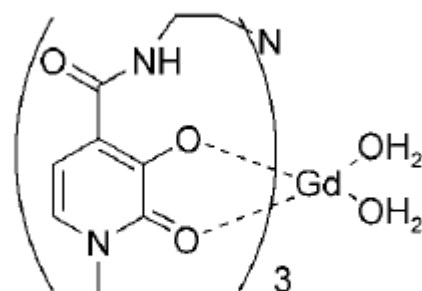
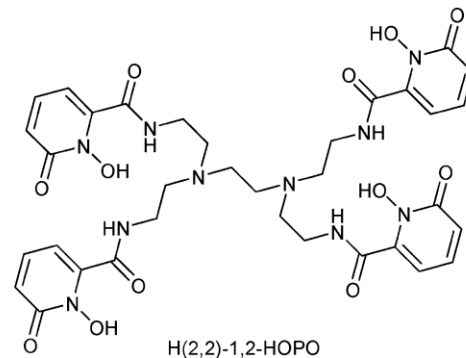
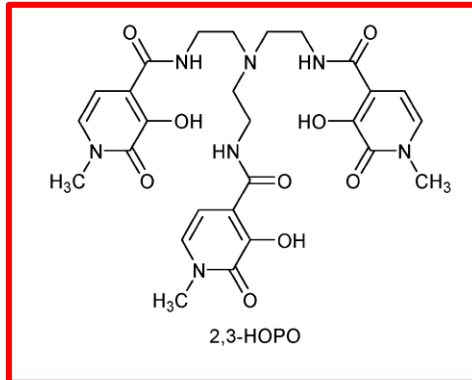
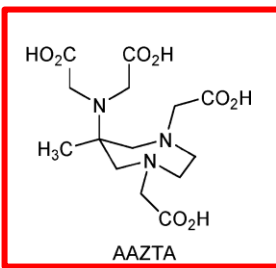
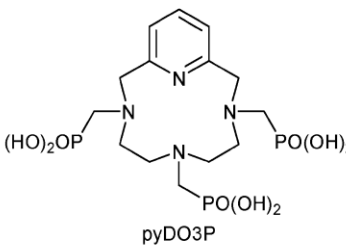
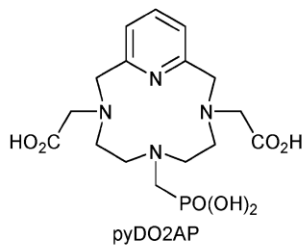
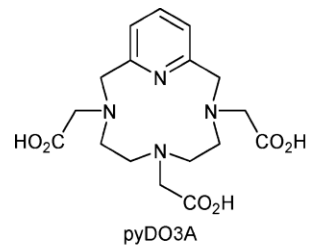
Teoria di Solomon-Bloembergen-Morgan (SBM Theory)





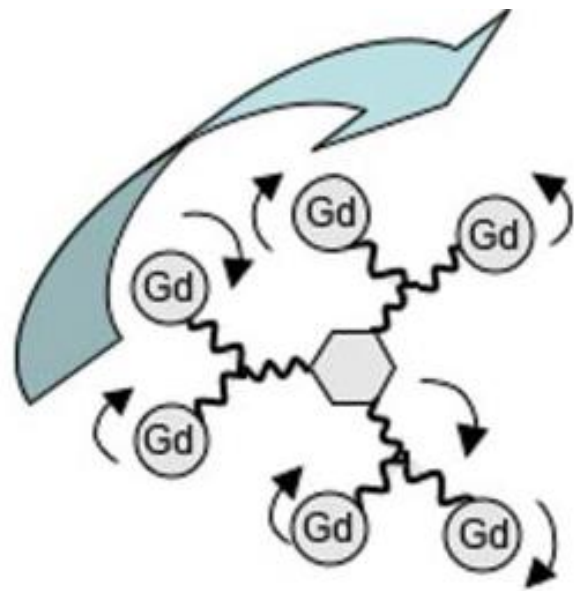
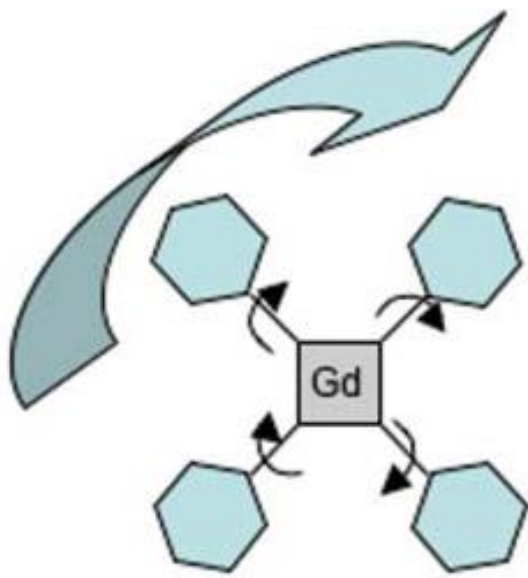


Nuovi leganti polidentati per CA di Gd(III)

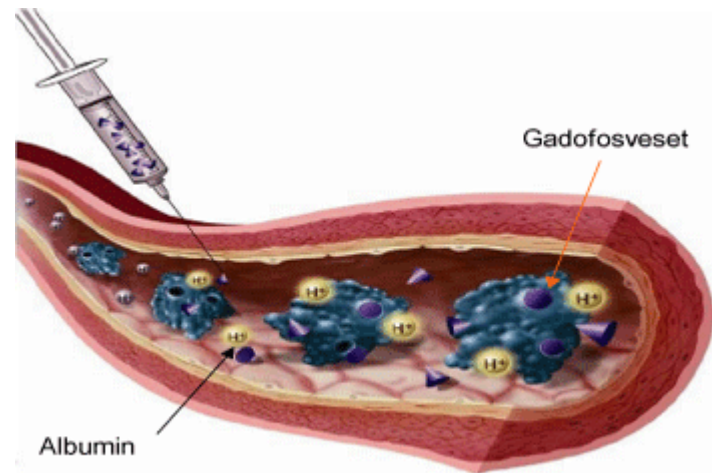
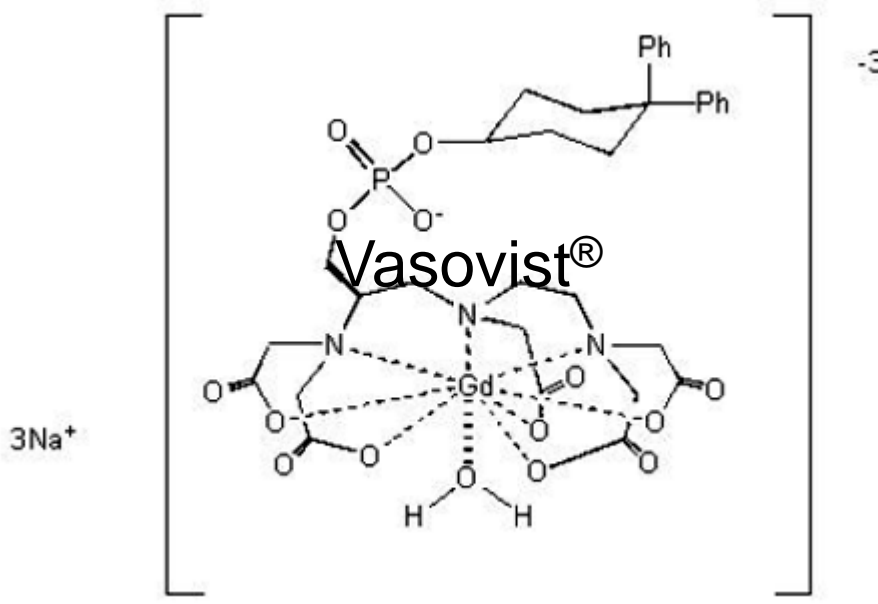


$[\text{Gd}\{\text{tren}(1\text{-Me-3,2-hopo})_3\}(\text{H}_2\text{O})_2]$

Strategie per aumentare τ_M



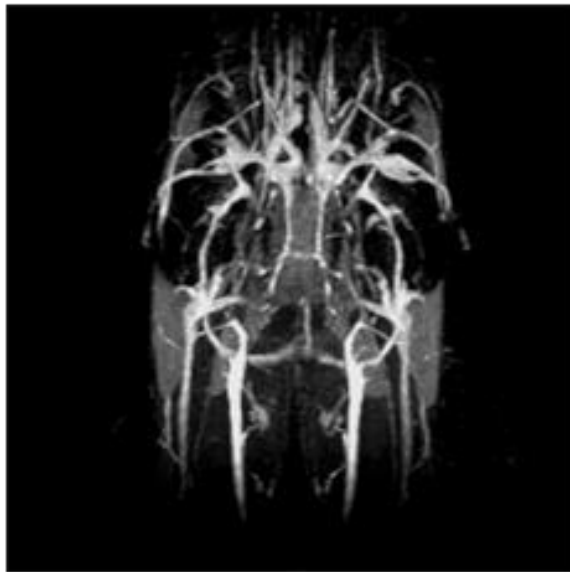
Blood pool contrast agents



Binding of the C.A.
to serum albumin
increases its
tumbling time (τ_R)



**5 min after
0.1 mmol/kg i.v.
of extracellular CA**



**5 min after
0.015 mmol/kg i.v.
of angiographic ca**

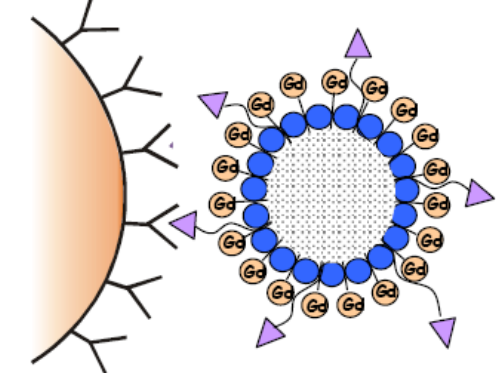
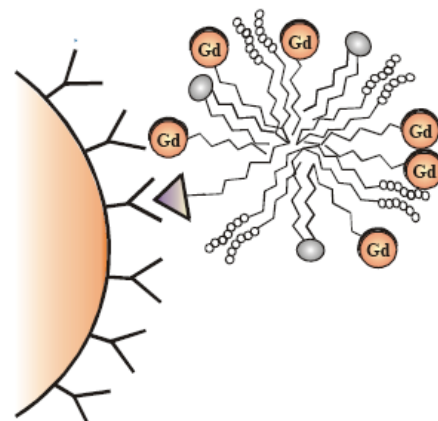
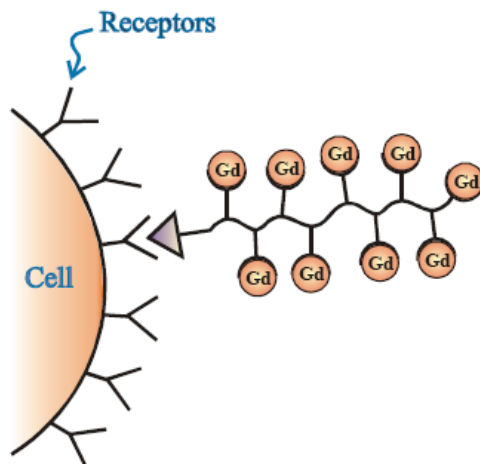
Towards molecular imaging with MRI

The very low concentration of the target requires the delivery of a high number, and possibly efficient, Gd(III) centres

$C > 125 \mu\text{M}$

Several strategies can be adopted

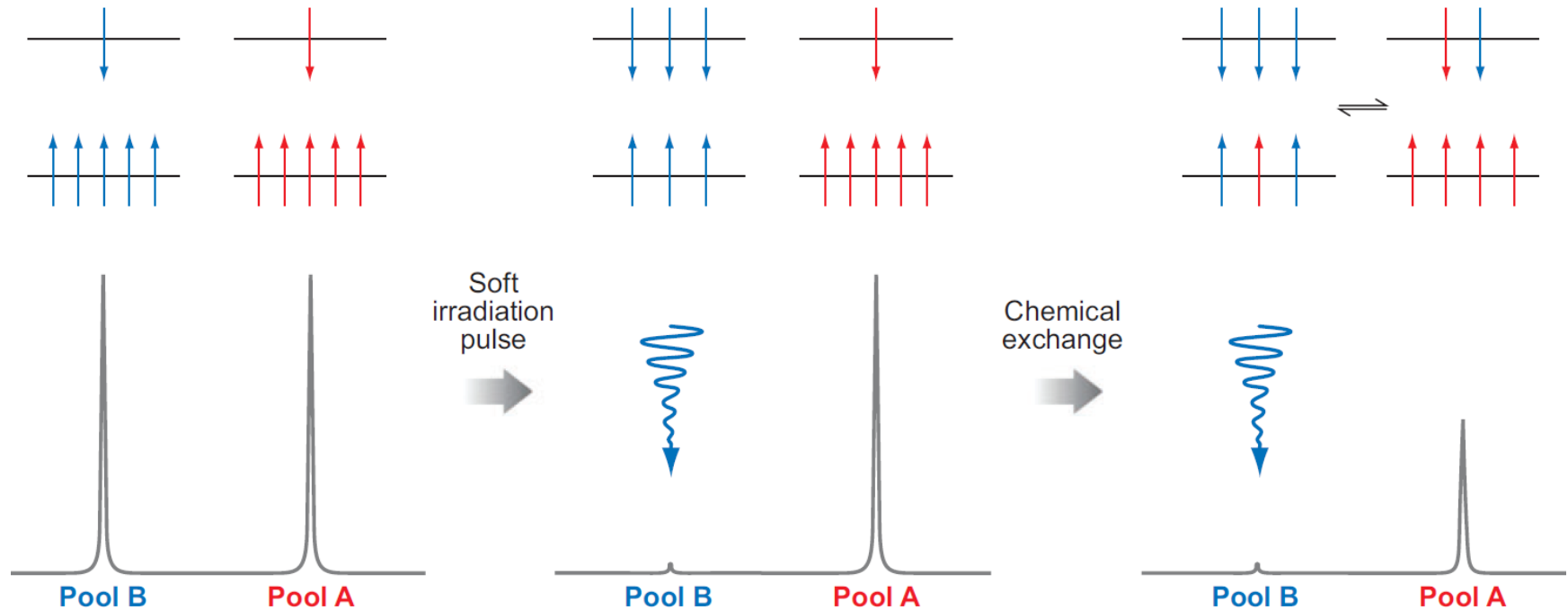
- Gd-chelates covalently or non-covalently linked to biocompatible polymer (proteins, polysaccharides, etc...)
- Self-assembling of complexes (e.g. micelles)
- Use of Gd-loaded nanoparticles (e.g. liposomes,...)



CEST Contrast Agents

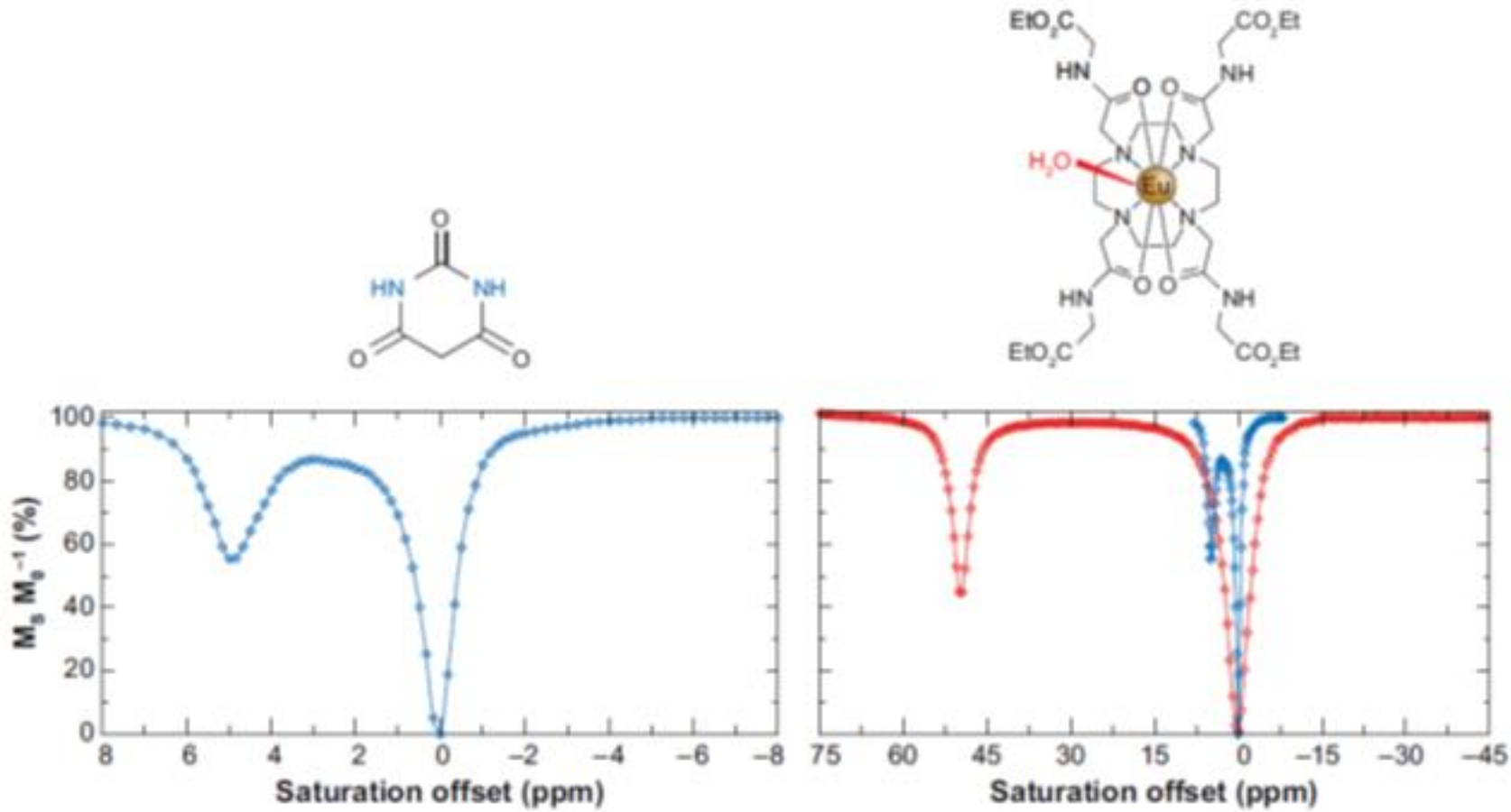
Chemical Exchange Saturation Transfer

composti mobili con protoni in scambio lento con l'acqua di *bulk*



$$k_{\text{CEST}} < \Delta\omega$$

CEST and PARACEST agents: saturation offset



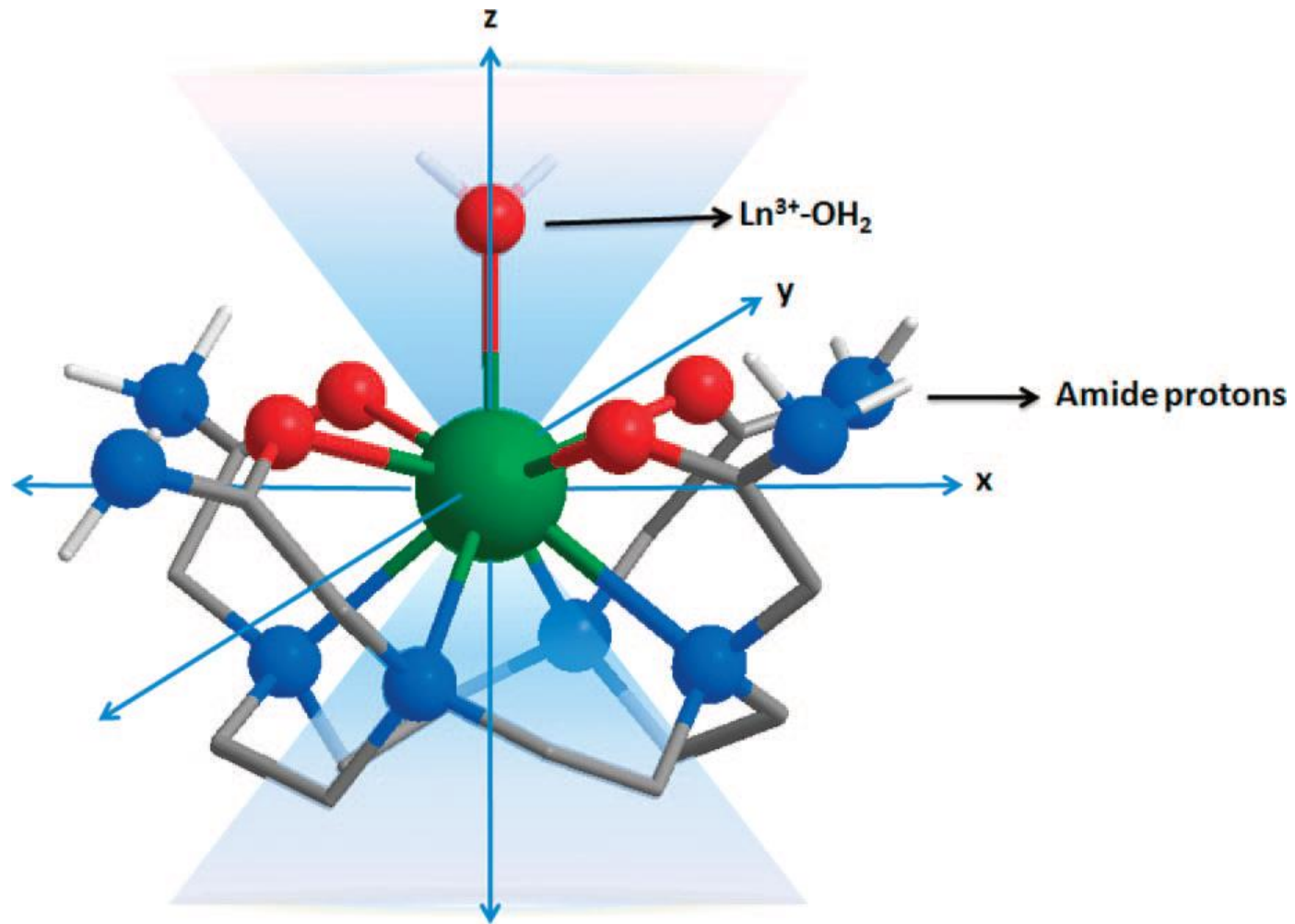
isotropo

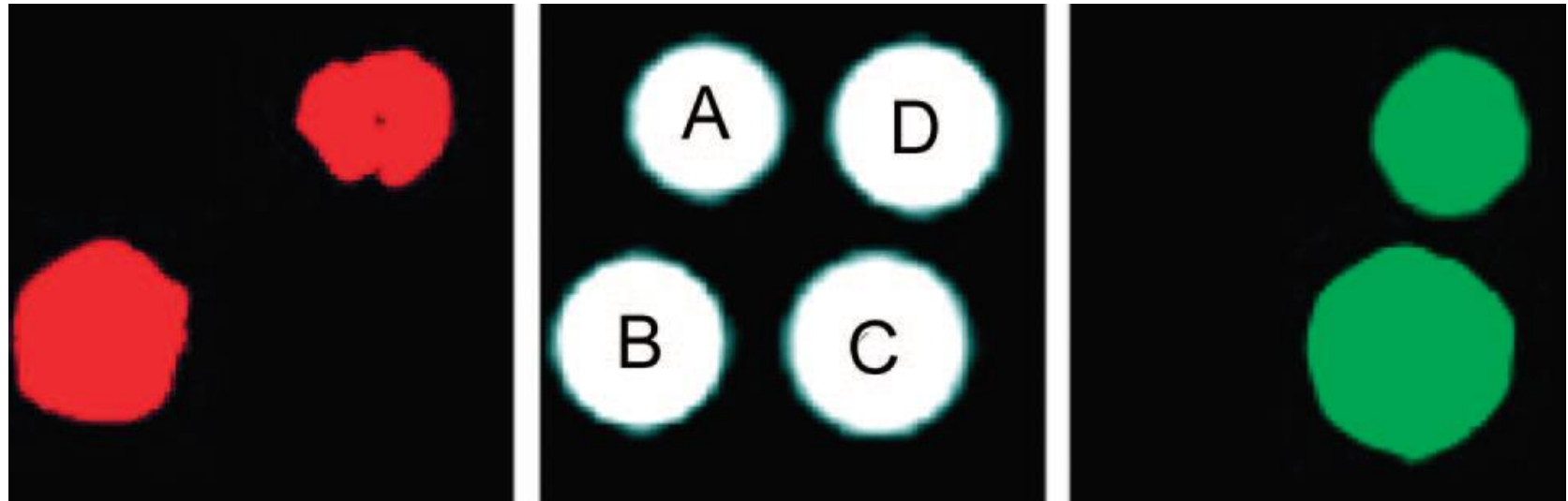
Ln	La ³⁺	Ce ³⁺	Pr ³⁺	Nd ³⁺	Sm ³⁺	Eu ³⁺	Gd ³⁺	Tb ³⁺	Dy ³⁺	Ho ³⁺	Er ³⁺	Tm ³⁺	Yb ³⁺	Lu ³⁺
μ_{eff}/μ_B cald.	0.00	2.54	3.58	3.68	0.85	0.00	7.94	9.72	10.63	10.60	9.59	7.57	4.53	0.00
$\chi/10^{-32}\text{m}^3$	0.0	5.6	11.2	11.4	0.6	6.0	55.1	82.7	99.2	98.5	80.3	50.0	18.0	0
$\Delta\chi_{\text{ax}}/10^{-32}\text{m}^3$	0	2.1	3.4	1.7	0.2	-2.3	0	42.1	34.7	18.5	-11.6	-21.9	-8.3	0
$\Delta\chi_{\text{rh}}/10^{-32}\text{m}^3$	0	0.7	2.1	0.4	-0.1	-1.6	0	11.2	20.3	5.8	-8.6	-20.1	-5.8	0
PRE														
PCS														

PRE = Paramagnetic Relaxation Enhancement

PCS = Pseudo-Contact Shift

determina la variazione di chemical shift indotta da ciascuno ione sui nuclei vicini e le iso-superfici rappresentano la grandezza e il segno del chemical shift



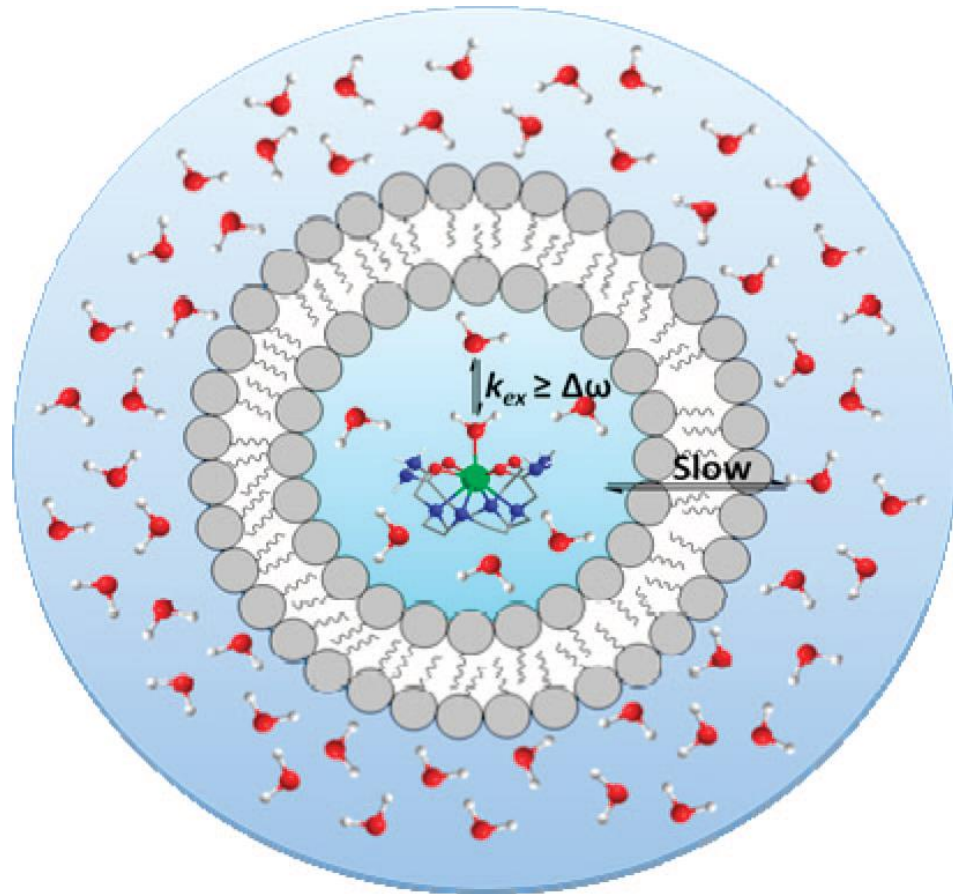
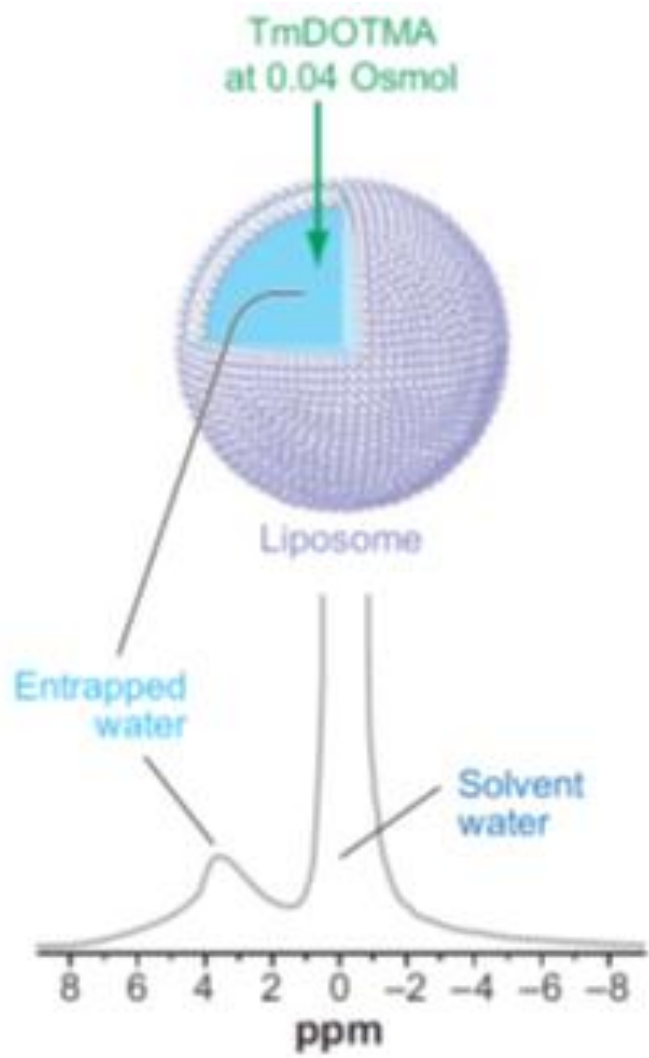


B = [Tb-DOTAMGly]⁻

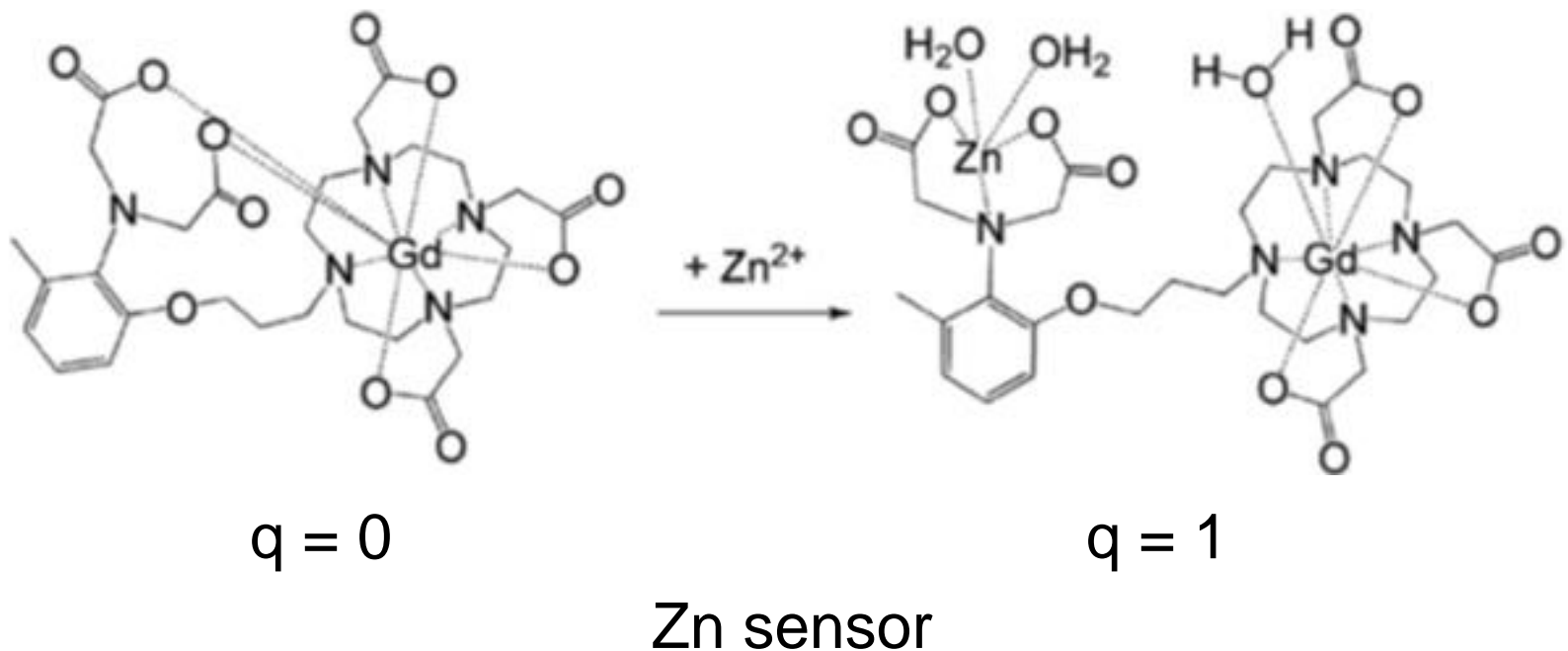
C = [Eu-DOTAMGly]⁻

D = [Tb-DOTAMGly]⁻ + [Eu-DOTAMGly]⁻

LipoCEST



Responsive (*smart*) CA



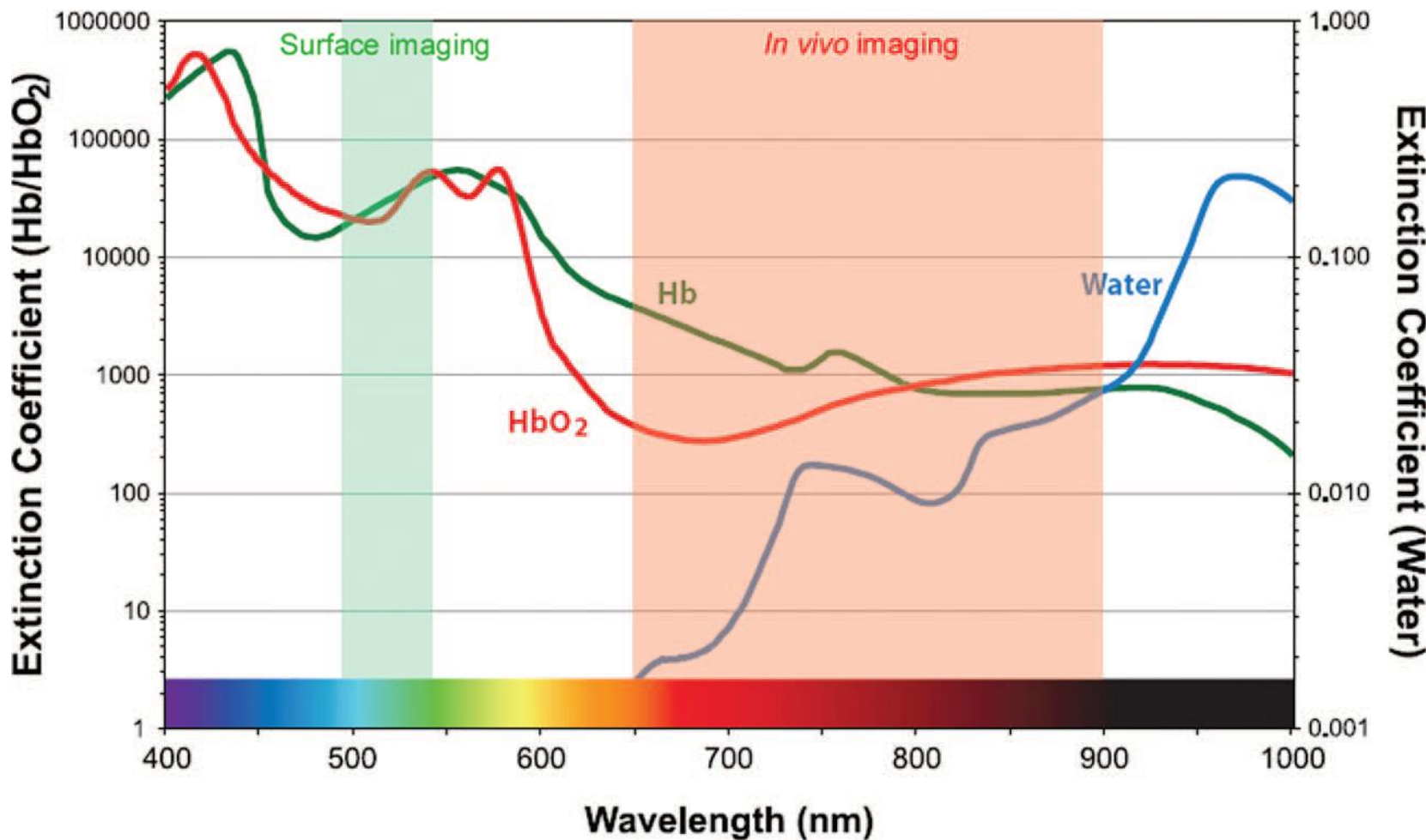
T_2 contrast agents super-paramagnetic iron oxide particles (SPIO) $d = 60 - 250 \text{ nm}$

Pre-Clinical Agent	Commercial Name	MR Target	Status
AMI-25	Ferumoxide, Feridex, Endoderm	Liver	Approved
OMP	Abdoscan	Bowel	Approved
AMI-121	Gastromark, Ferumoxsil, Lumirem	Bowel	Approved
SHU555A	Resovist	Liver	Approved (EU, Japan, Australia), Phase III (USA)
AMI-227	Combidex, Sinerem, Ferumoxtran	Lymph Node Metastases	Phase III
CODE 7228	Feraheme, Ferumoxytol	Vasculature	Phase II

Imaging ottico

- Sensibilità paragonabile a quella di SPECT e PET
- Possibilità di agenti *switchable*
- Possibilità di *time-resolved detection*
- **No quantificazione**

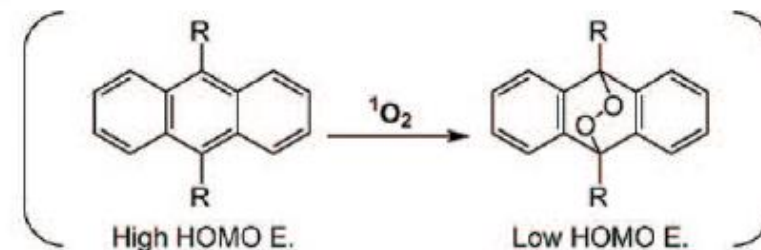
- Window
- Stokes shift
- Brightness
- Stability



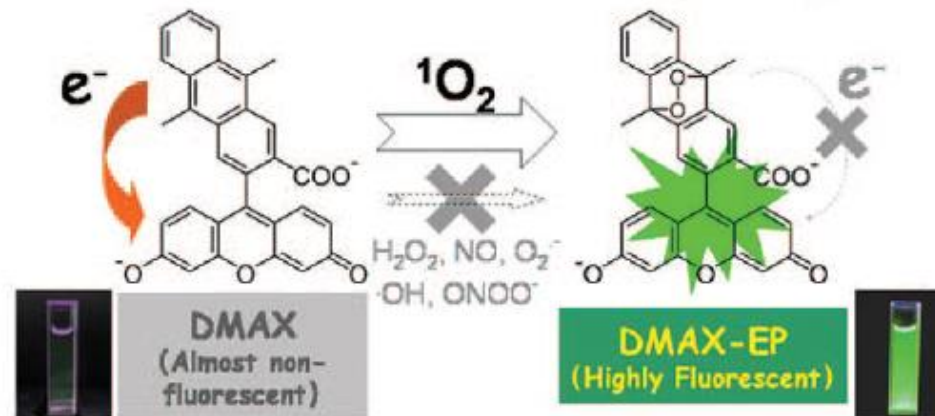
Esempio di switchable fluorescent probe sensore di ${}^1\text{O}_2$

(a) Singlet Oxygen Probes

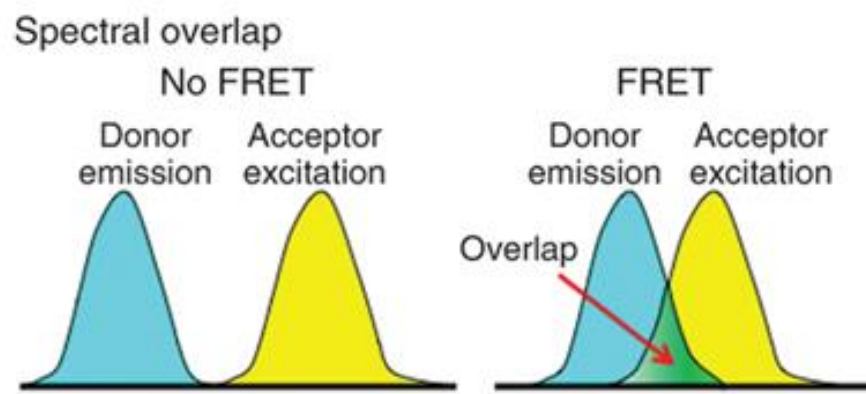
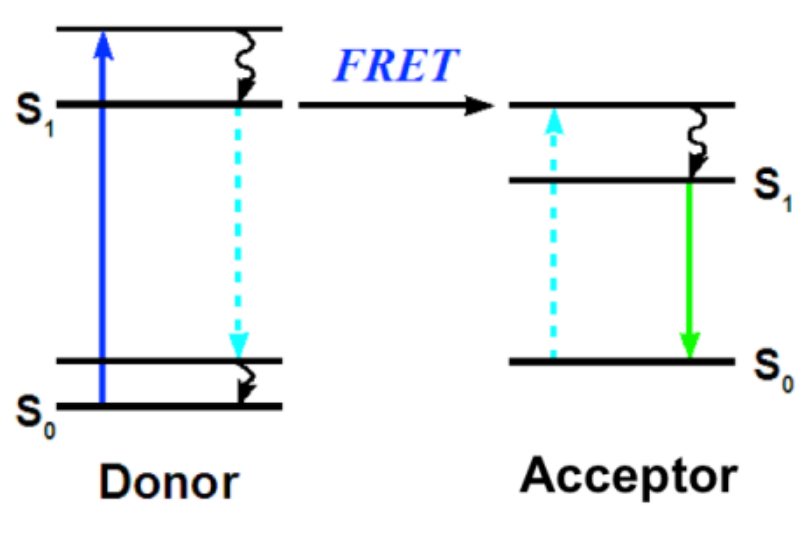
Key reaction: Endoperoxide formation



Reaction scheme for detection of singlet oxygen



FRET fluorescence – resonance energy transfer



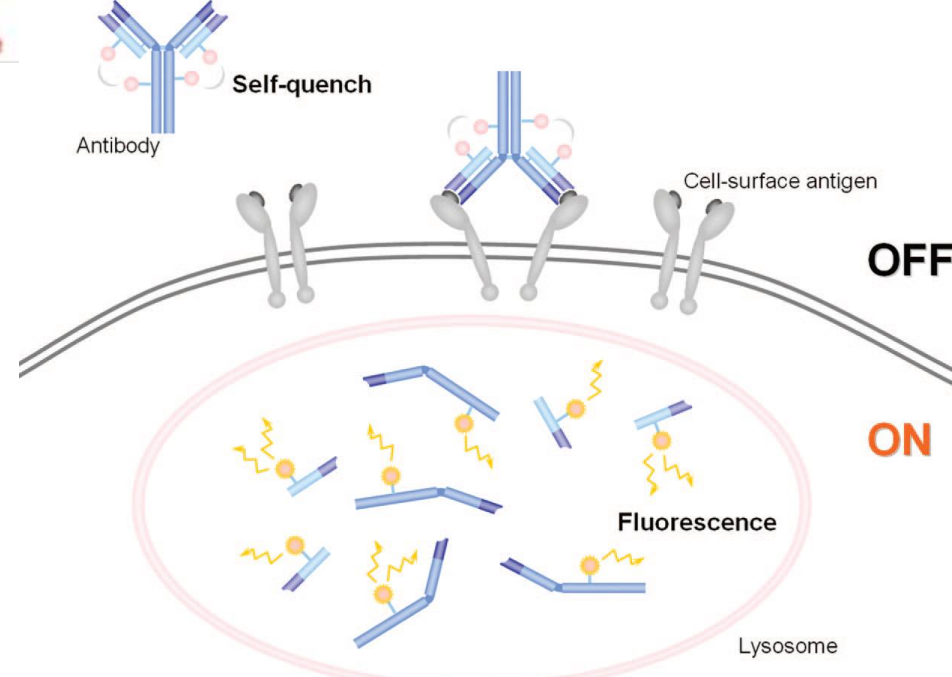
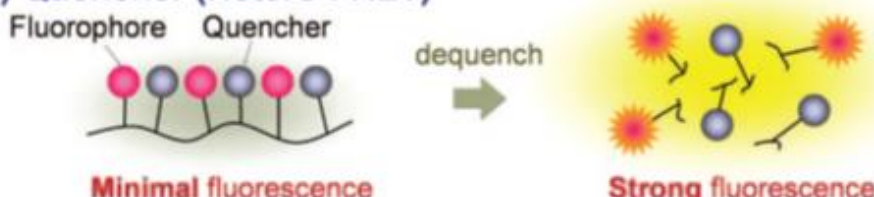
a) Self-quench (Homo-FRET)



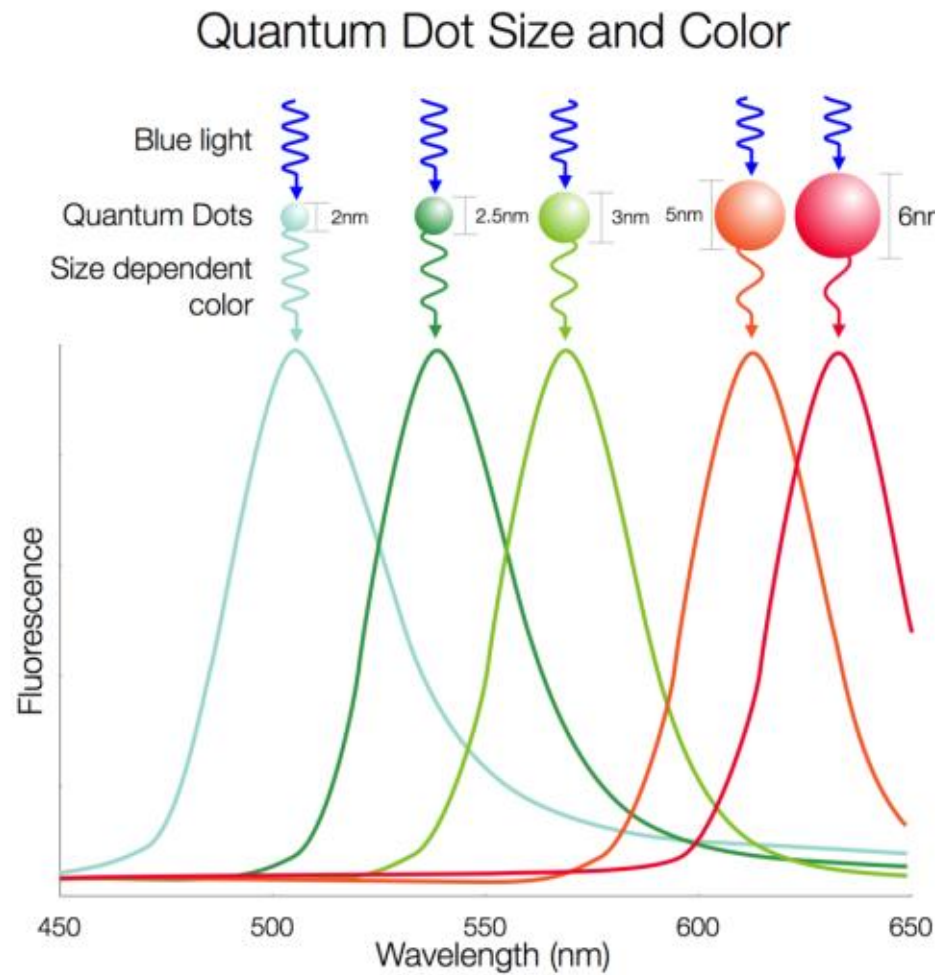
b) Fluorophore protein interaction



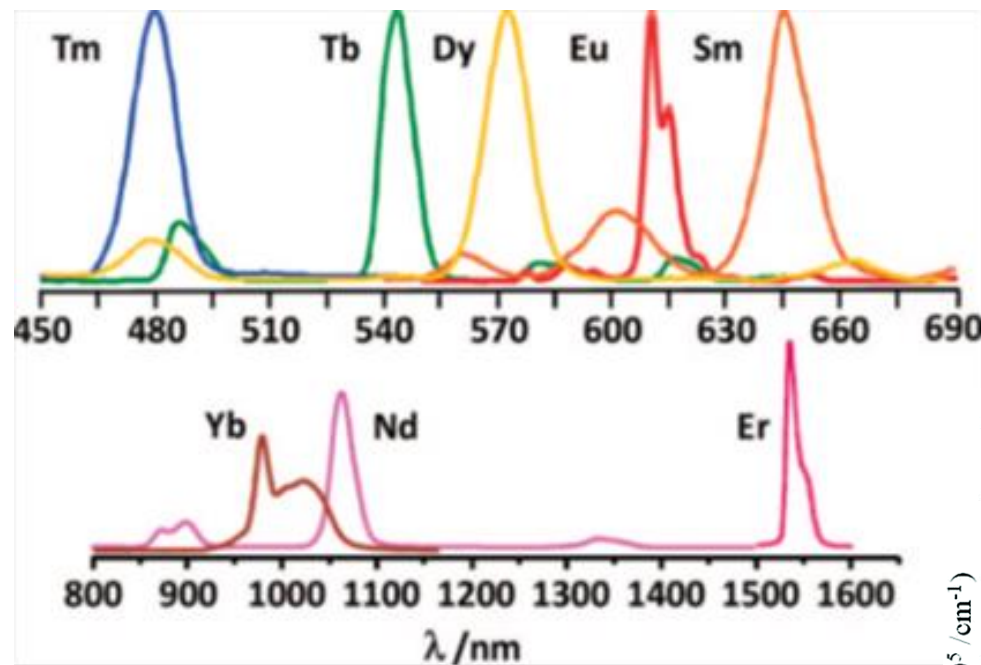
c) Quencher (Hetero-FRET)



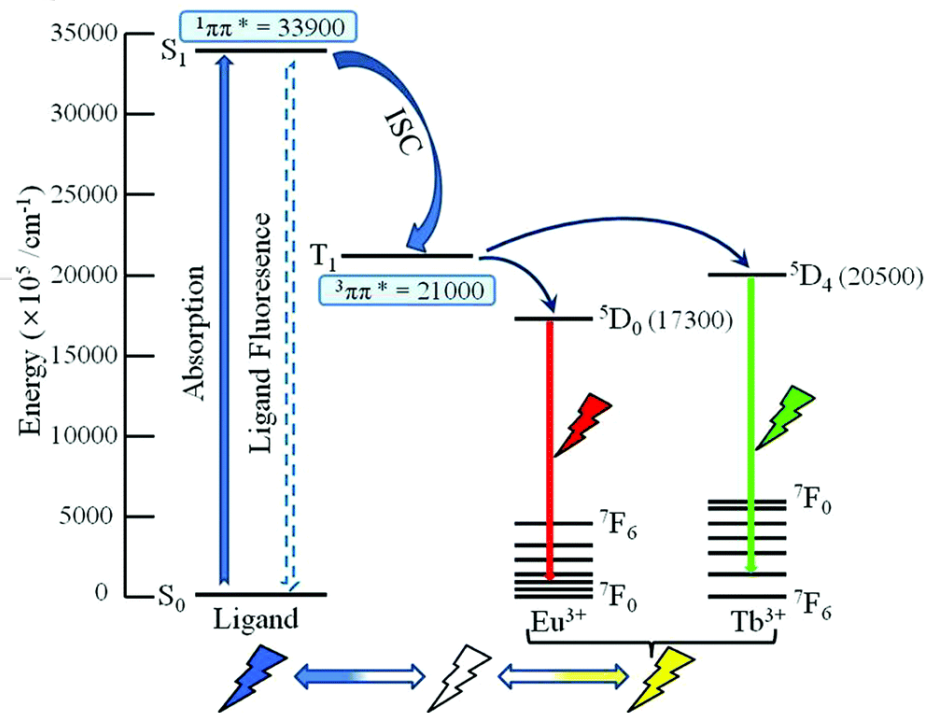
quantum dots (QD) nano-cristalli di semiconduttori (e.g. CdSe)



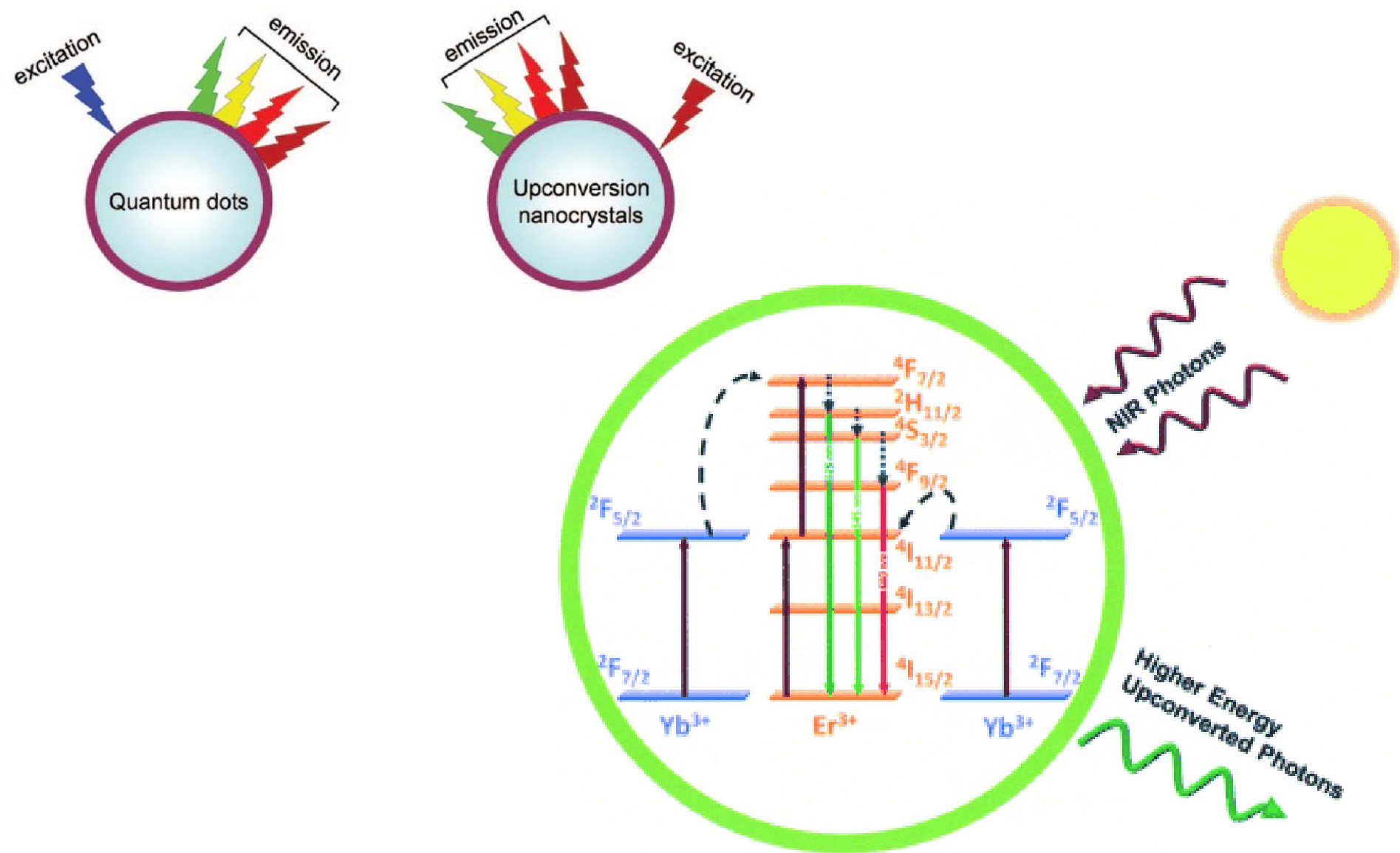
Complessi dei lantanidi



Effetto antenna dei leganti

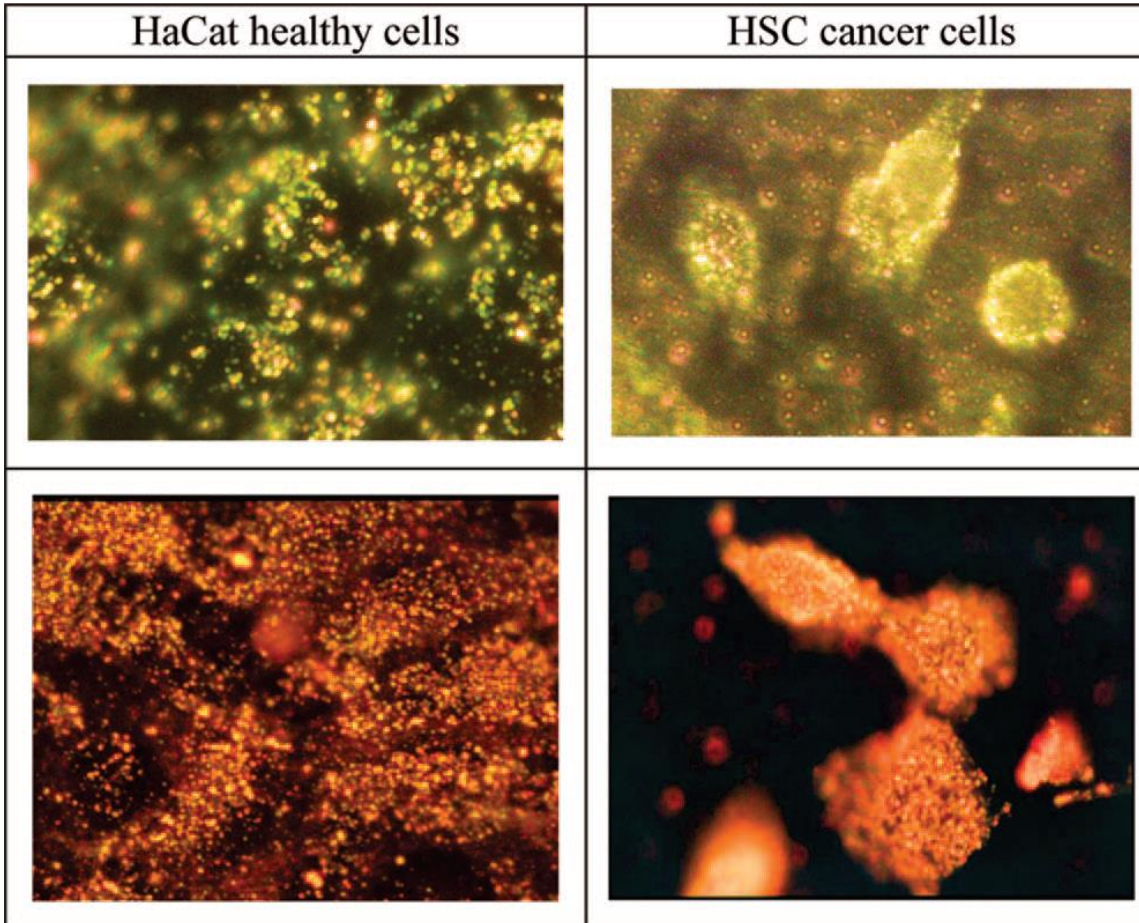


Upconverting QDs e LnNPs



Dark-field fluorescence imaging con AuNP

scattered
light



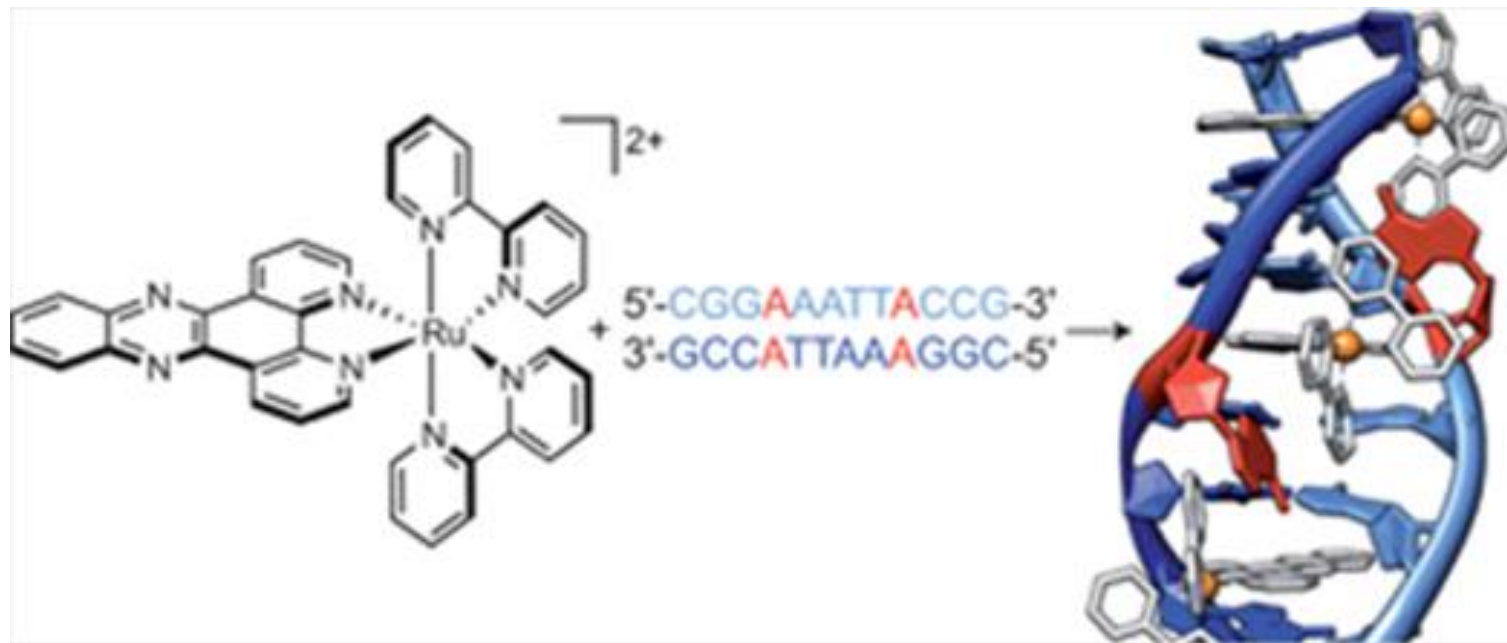
Au nanospheres

Au nanorods

AuNP coniugate a anticorpi anti-EGFR

EGFR = epidermal growth factor receptor, marcatore tumorale

Complessi polipiridilici di Ru(II) come *DNA light switch*



Sviluppi futuri

Multimodal imaging agents and theranostics

