

- Free radicals in solids, liquids or gases
- Paramagnetic ions of transition metals (up to 5 or 9 unpaired e-)
- localized defects in solids (point defects)

 systems with more than one unpaired e- : triplet systems

biradicals

 systems with conduction e- (semiconductors and metals)

Organic Radicals

- very popolar in EPR
- one unpaired e-
- the effect of the spin orbit coupling sul g_e is very small (in the case of p orbitals)
- the paramagnetism can be considered as exclusively due to e- spin (ESR)
- "free" nomenclature of 1800 to adress groups of atoms with one unpaired e- that can migrate from one molecule to another

Organic Radicals

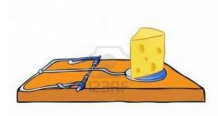
- The first radical to be proved is the triphenylmethyl (trytil)
- The are distinguished as σ or π
- The π with an extended π systems are usually both **thermodinamically** more stable
- and have a longer lifetime (kinetics): very important for the ESR study

Organic Radicals

- Neutral radicals (from homolysis)
- Radicalcations
- Radicalanions
- Biradicals (triplet state)
- also from photoexcitation (ESR proof of the phosphorescence mechanism)
- The stable radicals are used as spin probes and spin labels

Nitroxide Radical

- amongst the most diffuse persistent radicals in ESR
- RR' NO·
- derivatives of nitric oxide NO \cdot
- preparation: by oxidation of the parent secondary amines or oximes by H₂O₂, peroxides, Ag(I), H₂O₂ and Ti(III)
- adducts of transient radicals to nitroso compounds or nitrones (spin trap)

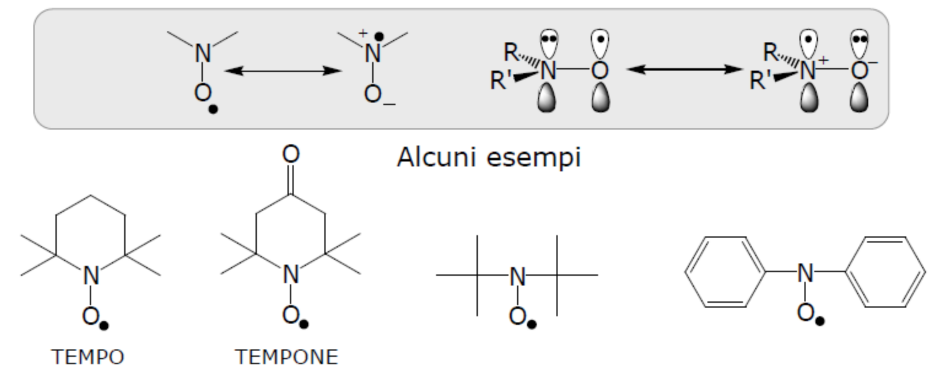


 $\begin{array}{ll} \text{spin trap} & \text{sp} \\ \text{R}^{\cdot} + \text{R}^{\prime} \text{N}=O \rightarrow & \text{RR} \end{array}$

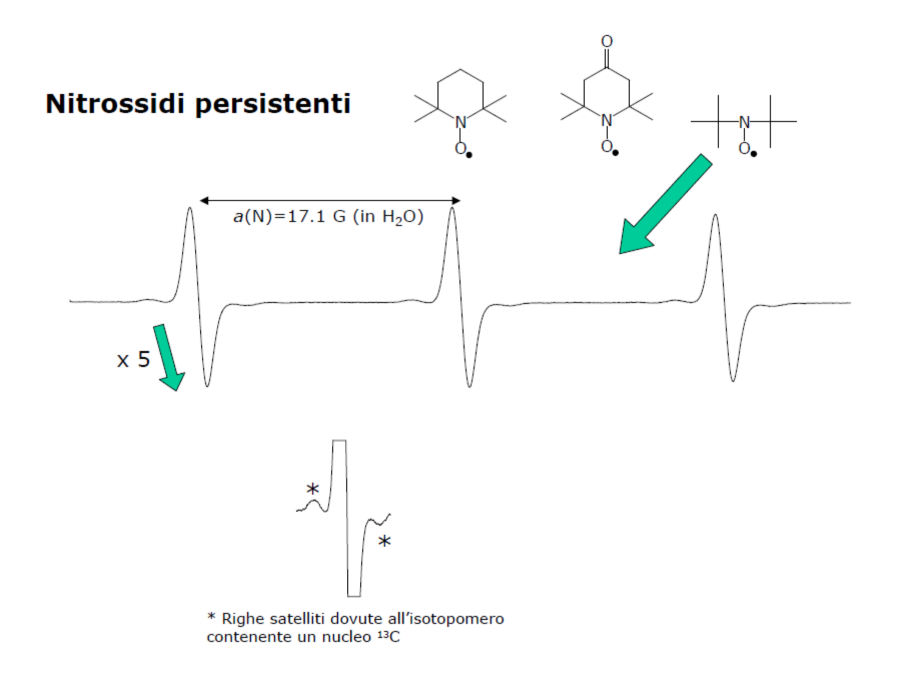
spin adduct RR' N-O·

Nitroxide Radicals

persistent (from months till years), temperature stable, soluble in water and many organic solvents

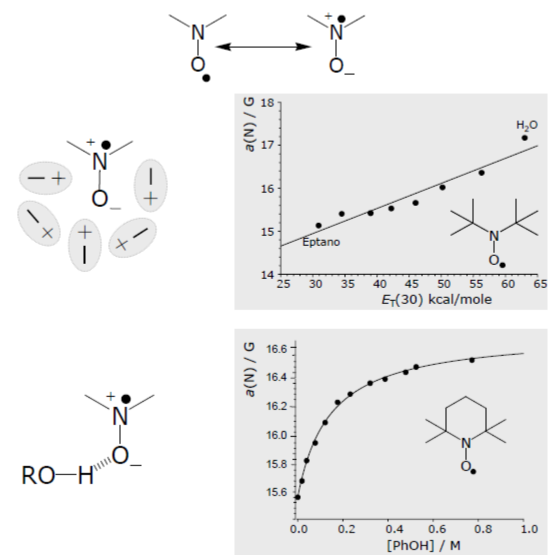


also **TEMPOLO**

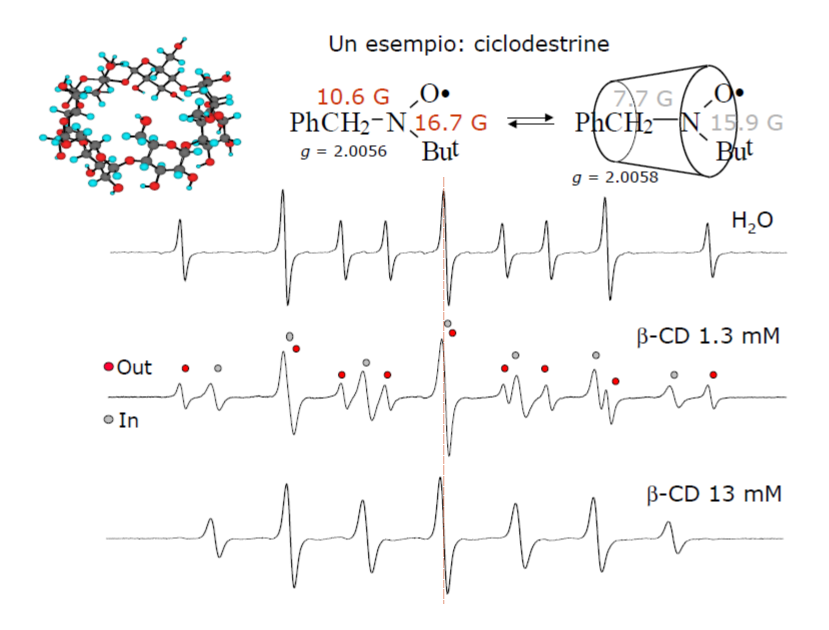


Spin probe

Dipendenza di a(N) dall'ambiente circostante



Spin probe



Origin of Polarity Dependence of Nitroxides

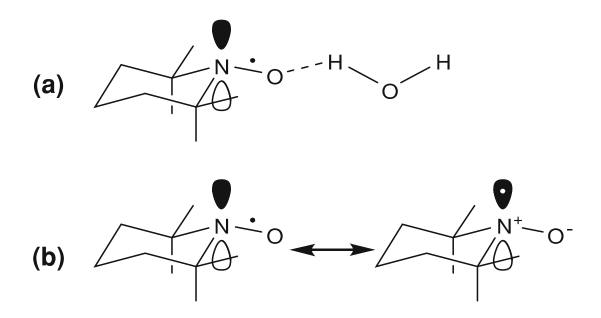


Fig. 2.14 Origin of the polarity and proticity dependence of nitroxide spectra. **a** Hydrogen bonding leads to a decrease of the spin-orbit coupling. **b** A stabilization of the zwitterionic resonance structure causes an increase of the spin density at the nitrogen atom. The combined effects lead to a decrease of g_{iso} and to an increase of a_{iso}

Zeeman Interaction

ESR spectroscopy is based on the interaction of the magnetic moment of e- with the magnetic field

$$\alpha$$
 — $E_{\alpha} = 1/2g_{e}\mu_{B}B$

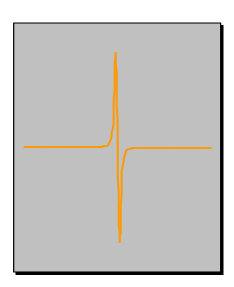
 $\Delta E = hv = g_e \mu_B B$

$$B = -1/2g_e\mu_B B$$

e-gyromagnetic ratio

 $\gamma_e = \nu/B = g_e \mu_B / h = 2.8024 \cdot 10^{10} \text{ s}^{-1}\text{T}^{-1}$

considering the g_e of the free e-



Electron-Nuclear Magnetic Interaction

Hyperfine Interactions

Hyperfine interaction for the hydrogen atom

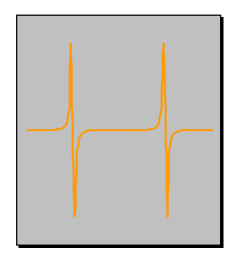
a(H)= 506.8 G

selction rules

 $\Delta M_s = \pm 1$

 $\Delta M_I = 0$

Fermi contact mechanism



m_l=+1/2 m_l=-1/2

B-a/2 B+a/2