

Introduction to solution NMR

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with thanks to Dr. Klaartje Houben

EMBO Global Exchange course, USP, Sao Paolo, Brazil

January 19th-26th, 2014



Universiteit Utrecht

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Solution NMR: 900-cryo, 750, 600-cryo, 600US, 2x500 MHz
Solid-state NMR: 800WB-DNP, 400WB-DNP, 700US, 500WB MHz
e-infrastructure: >1600 CPU cores + EGI grid (>90'000 CPU cores)

2014 : 950 MHz
2017: 1.2 GHz



and European infrastructure

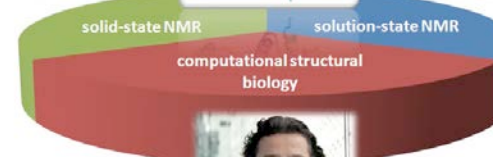


The NMR research group

Prof. Marc Baldus



Prof. Rolf Boelens



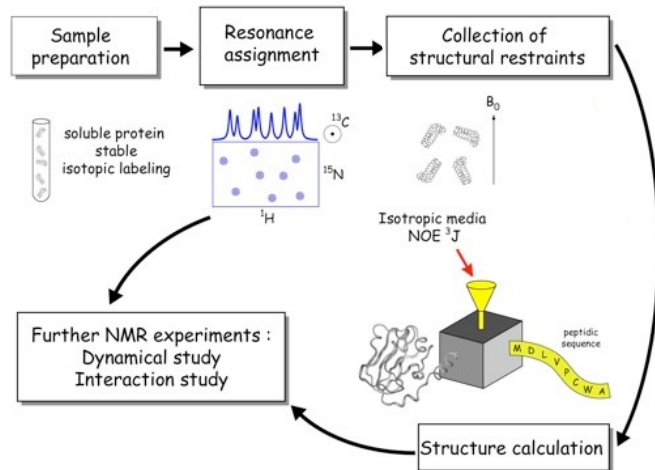
Prof. Alexandre Bonvin



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NMR 'journey'



Topics

- Why use NMR for structural biology...?
- The very basics
- Multidimensional NMR (intro)
- Resonance assignment (lecture Banci)
- Structure parameters & calculations (lecture Banci)
- NMR relaxation & dynamics



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Why use NMR.... ?

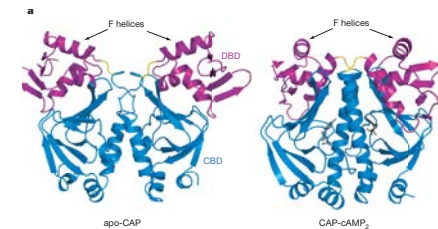


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NMR & Structural biology

DYNAMICS



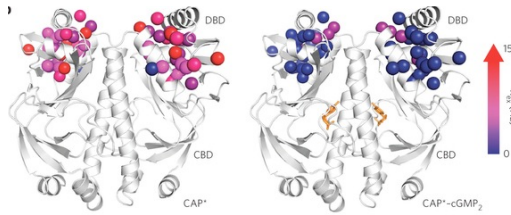
Dynamic activation of an allosteric regulatory protein Tzeng
S-R & Kalodimos *CG Nature* (2009)

NMR & Structural biology

DYNAMICS

• Allosteric regulation

- Dynamic interaction between ligand-binding & DNA binding site

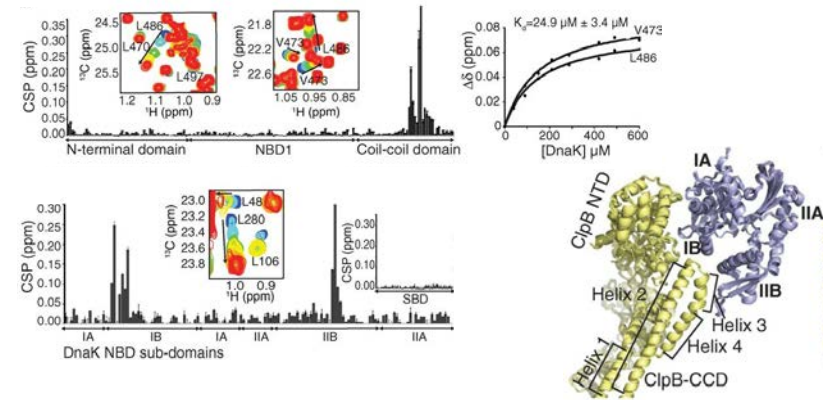


Dynamic activation of an allosteric regulatory protein Tzeng
S-R & Kalodimos CG Nature (2009)

NMR & Structural biology

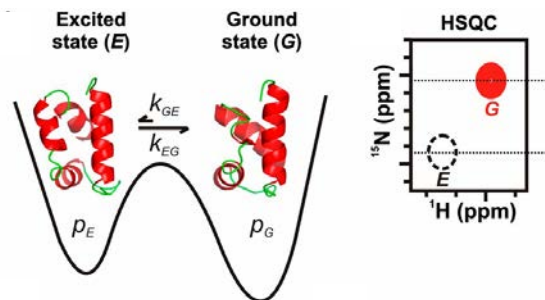
Biomolecular interactions

- Even weak and transient complexes can be studied



NMR & Structural biology

EXCITED STATES



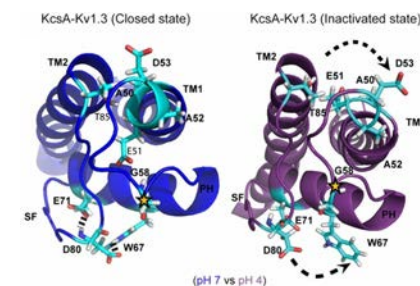
Shekhar & Kay PNAS 2013

NMR & Structural biology

MEMBRANE

• Native like environment

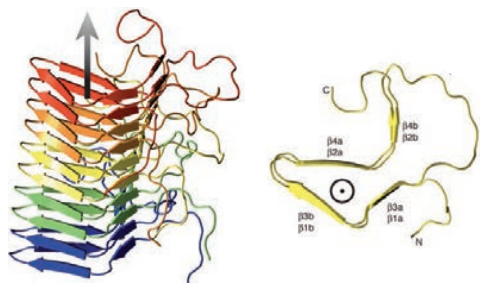
- Structural changes due to lipid environment



van der Cruisen, & Baldus PNAS 2013

NMR & Structural biology

AMYLOID FIBRILS

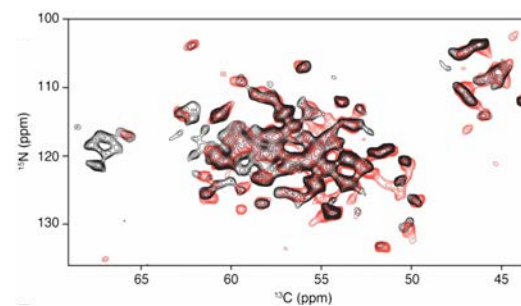


Amyloid Fibrils of the HET-s(218–289) Prion Form a β Solenoid with a Triangular Hydrophobic Core Wasmer C. *et al* Science (2008)

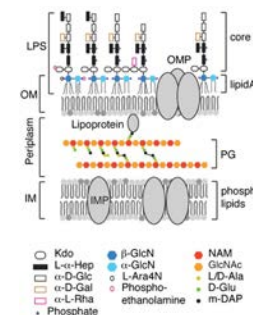
NMR & Structural biology

IN-CELL NMR

- Study proteins in their native cellular environment
 - Outermembrane protein in bacterial cell envelop



Renault M, & Baldus PNAS 2012



The NMR sample

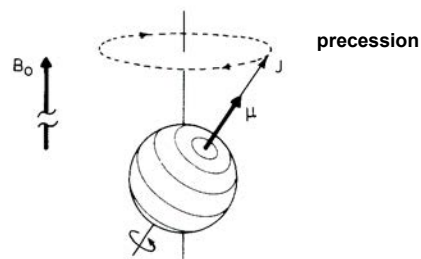
- isotope labeling
 - ^{15}N , ^{13}C , ^2H
 - selective labeling (e.g. only methyl groups)
 - recombinant expression in *E.coli*
- sample
 - pure, stable and high concentration
 - 500 μL of 0.5 mM solution $\rightarrow$ ~ 5 mg per sample
 - preferably low salt, low pH
 - no additives



The very basics of NMR



Nuclear spin



$$E = -\vec{\mu} \cdot \vec{B} = -\mu_z B_z$$

$$|\mu| = \gamma \hbar \sqrt{I(I+1)}$$

I = quantum number

$$\mu_z = \gamma \hbar m$$

$m = I, I-1, I-2, \dots, -I$ = allowed states

Nuclear spin

TABLE 1.1
Properties of Selected Nuclei^a

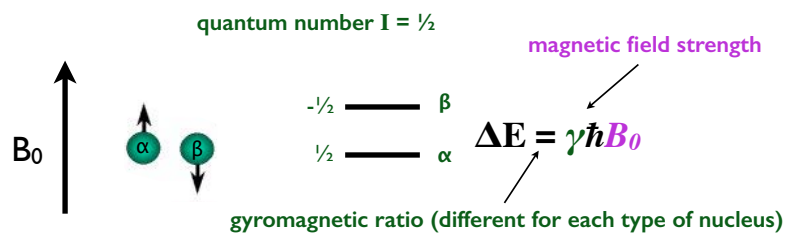
Nucleus	I	γ (rad · T ⁻¹ · s ⁻¹)	Natural abundance (%)
¹ H	$\frac{1}{2}$	2.6752×10^8	99.98
² H	1	4.107×10^7	0.02
¹³ C	$\frac{1}{2}$	6.728×10^7	1.11
¹⁴ N	1	1.934×10^7	99.64
¹⁵ N	$\frac{1}{2}$	-2.712×10^7	0.36
¹⁷ O	$\frac{5}{2}$	-3.628×10^7	0.04
¹⁹ F	$\frac{1}{2}$	2.5181×10^8	100.00
²³ Na	$\frac{3}{2}$	7.080×10^7	100.00
³¹ P	$\frac{1}{2}$	1.0841×10^8	100.00
¹¹³ Cd	$\frac{7}{2}$	5.934×10^7	12.26

^a The angular momentum quantum number, I , and the gyromagnetic ratio, γ , and natural isotopic abundance for nuclei of particular importance in biological NMR spectroscopy are shown.

Nuclear spin

• Nuclear magnetic resonance

- Only nuclei with non-zero spin quantum number are "magnets"
- Commonly used spins are spin $\frac{1}{2}$ nuclei: ¹H, ¹³C, ¹⁵N, ³¹P etc.

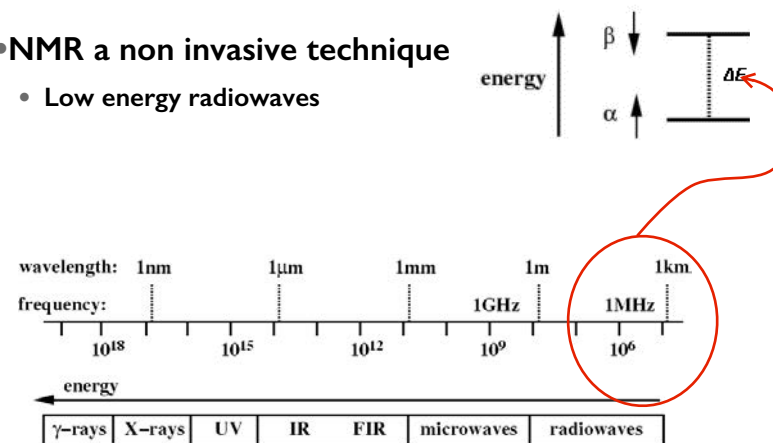


$$\text{Larmor frequency } \nu = (\gamma B_0)/2\pi$$

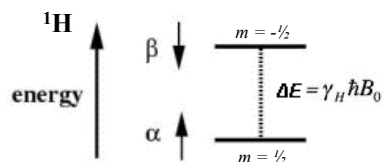
Nuclear spin & radiowaves

• NMR a non invasive technique

- Low energy radiowaves



Boltzman distribution

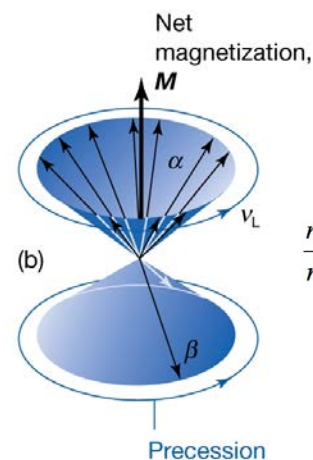


$$\frac{n_\beta}{n_\alpha} = \exp\left(-\frac{\Delta E}{k_B T}\right) = \exp\left(-\frac{\gamma_H \hbar B_0}{k_B T}\right) = 0.9999$$

Example

- 20.001 spins
- Only 1 more spin in lower energy state

Net magnetization



$$\frac{n_\beta}{n_\alpha} = \exp\left(-\frac{\Delta E}{k_B T}\right) = \exp\left(-\frac{\gamma_H \hbar B_0}{k_B T}\right) = 0.9999$$

Pulse

• Radio frequency pulses

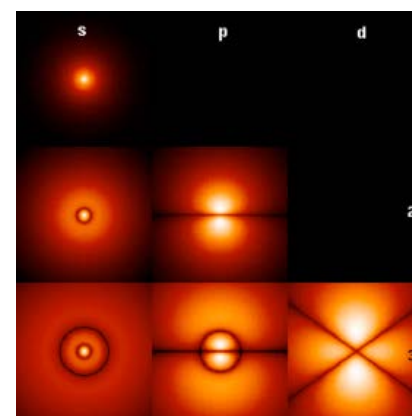
- Turn on an amplifier for a certain amount of time & certain amount of power (B_1 field)



only rotation
around B_1 is
observed

rotating frame: observe with frequency ν_0

Chemical shielding

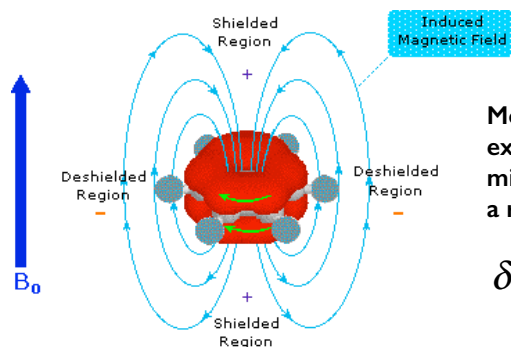


Local magnetic field is influenced by electronic environment
==> frequencies of nuclei will differ

Chemical shift

$$\nu = \frac{\gamma B_0}{2\pi} (1 - \sigma)$$

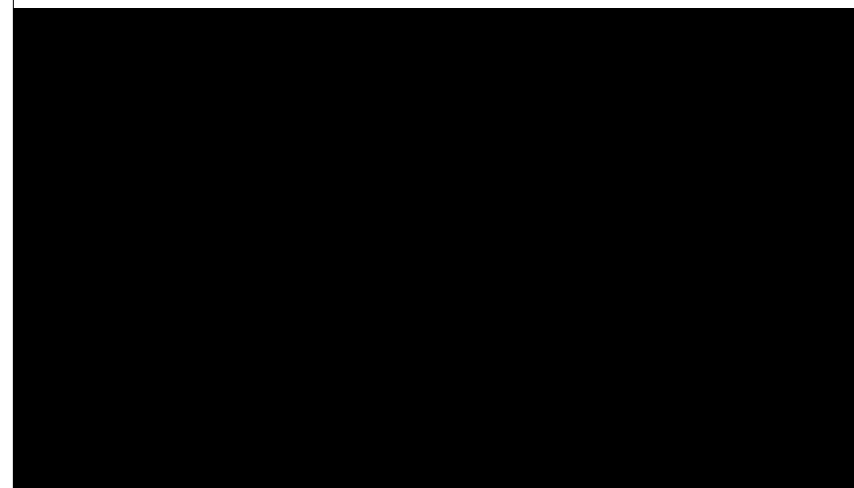
shielding constant



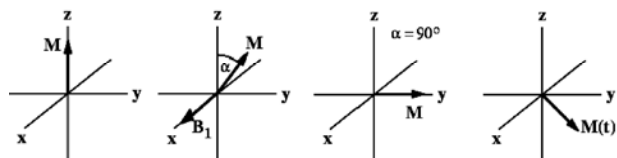
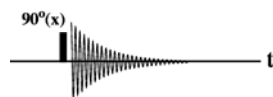
More conveniently expressed as part per million by comparison to a reference frequency:

$$\delta = 10^6 \frac{\nu - \nu_{ref}}{\nu_{ref}}$$

The spectrometer



Free induction decay (FID)



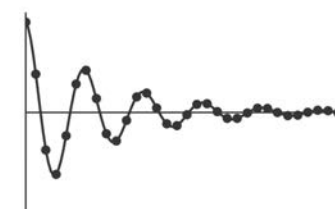
FID: analogue vs digital

(a)

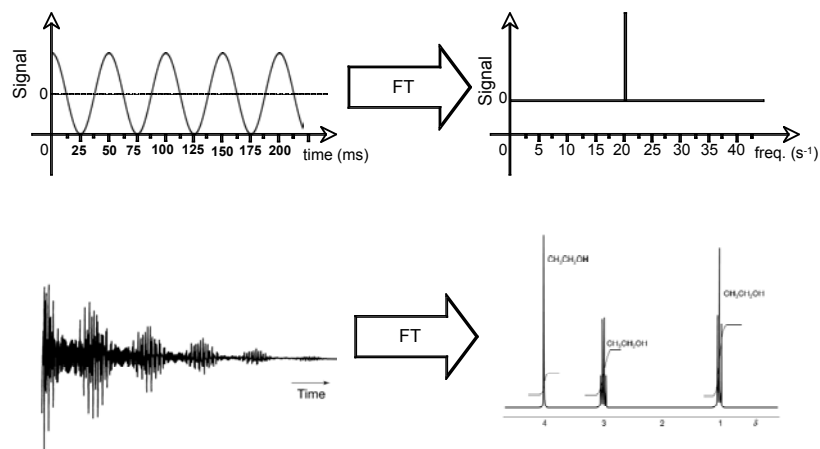


Free Induction Decay (FID)

(b)



Fourier Transform

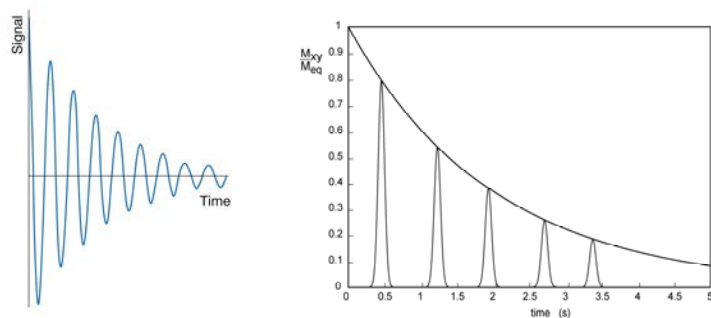


Relaxation

- **NMR Relaxation**
 - Restores Boltzmann equilibrium
- **T₂-relaxation (spin-spin)**
 - disappearance of transverse (x,y) magnetization
 - $1/T_2 \sim \text{signal line-width}$
- **T₁-relaxation (spin-lattice)**
 - build-up of longitudinal (z) magnetization
 - determines how long you should wait for the next experiment

Relaxation

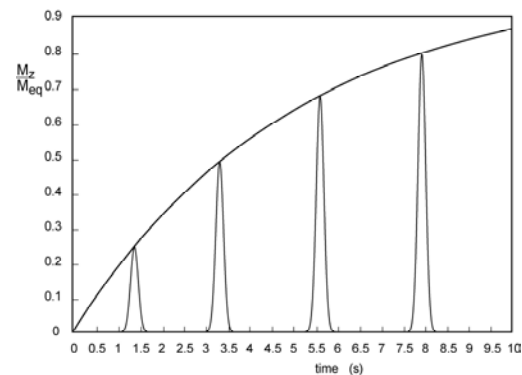
- **Restoring Boltzmann equilibrium**
 - T₂ relaxation: disappearance of transverse (x,y) magnetization



!! $1/T_2 \sim \text{signal line-width}$!!

Relaxation

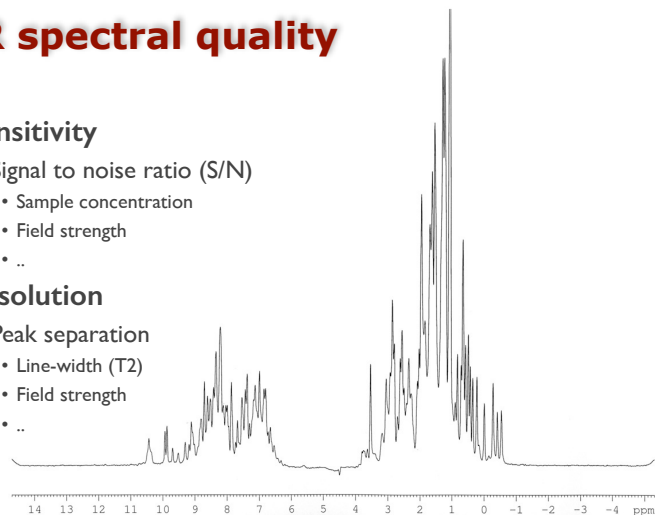
- **Restoring Boltzmann equilibrium**
 - T₁ relaxation: build-up of longitudinal (z) magnetization



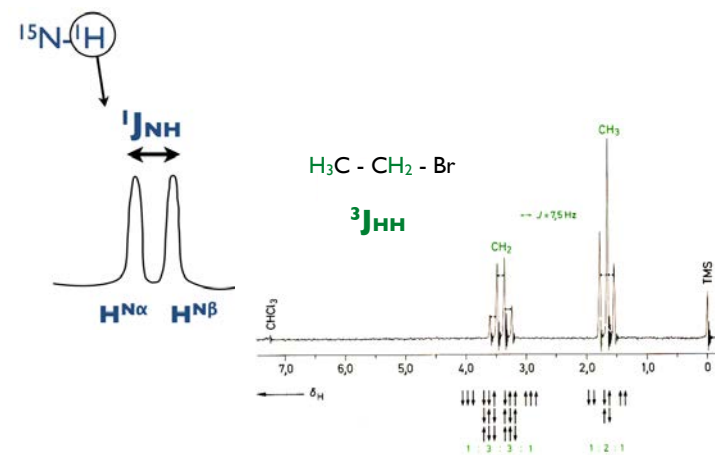
!! T₁ determines when to start the next experiment !!

NMR spectral quality

- **Sensitivity**
 - Signal to noise ratio (S/N)
 - Sample concentration
 - Field strength
 - ..
- **Resolution**
 - Peak separation
 - Line-width (T2)
 - Field strength
 - ..



Scalar coupling / J-coupling



Key concepts NMR

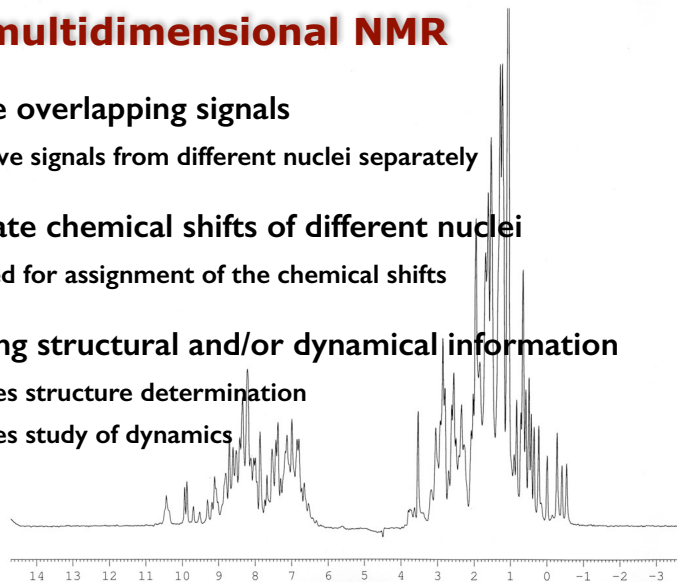
- **Nuclear magnetic resonance**
 - In a magnetic field magnetic nuclei will resonate with a specific frequency
- **FT-NMR**
 - Pulse, rotating frame, FID
- **Chemical shift**
 - Electronic environment influences local magnetic field -> frequency
- **NMR relaxation**
 - T_1 & T_2
- **J-coupling**

Multidimensional NMR

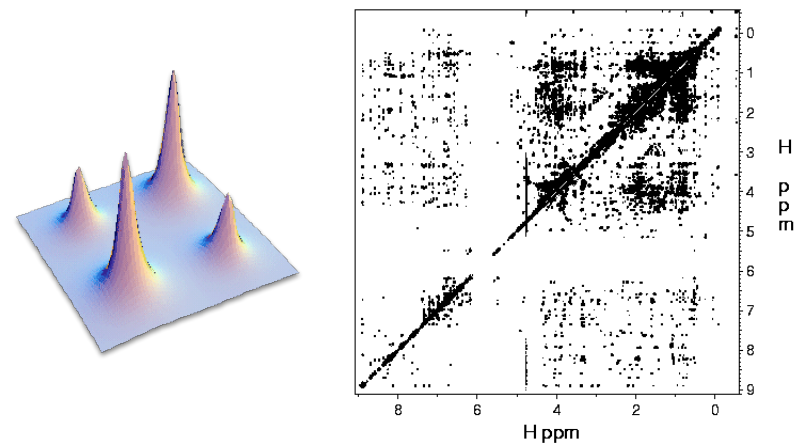


Why multidimensional NMR

- Resolve overlapping signals
 - observe signals from different nuclei separately
- Correlate chemical shifts of different nuclei
 - needed for assignment of the chemical shifts
- Encoding structural and/or dynamical information
 - enables structure determination
 - enables study of dynamics



2D NMR



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3D NMR

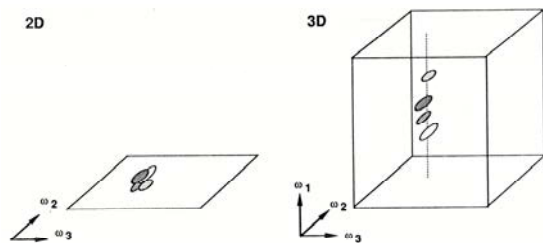
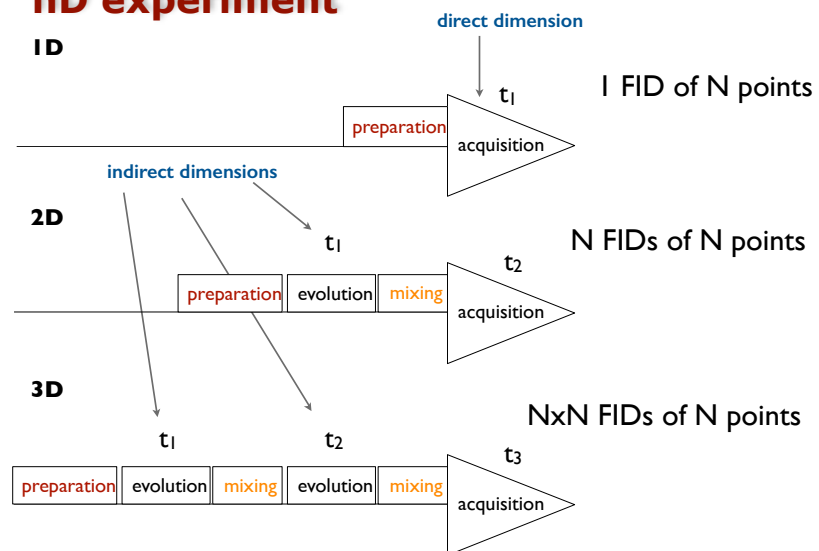


Figure 1. Illustration of the increase in resolution afforded by the increase in dimensionality. In the 2D spectrum, four cross-peaks overlap. By correlation with a third resonance frequency, each cross-peak obtains a different position along a line in the 3D spectrum, thus resolving the overlap problem.

(Unister, Ph.D. Thesis, 1991)

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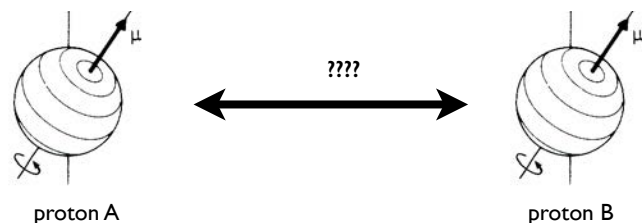
nD experiment



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Encoding information

- mixing/magnetization transfer

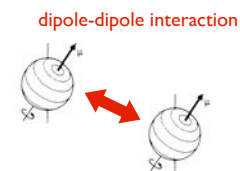


spin-spin interactions

Magnetization transfer

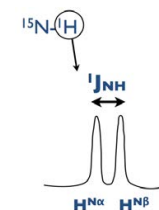
- **Magnetic dipole interaction (NOE)**

- Nuclear Overhauser Effect
- through space
- distance dependent ($1/r^6$)
- NOESY $\rightarrow$ distance restraints



- **J-coupling interaction**

- through 3-4 bonds max.
- chemical connectivities
- assignment
- also conformation dependent



homonuclear NMR

NOESY



magnetic dipole interaction
crosspeak intensity $\sim 1/r^6$
up to 5 Å

COSY



J-coupling interaction
transfer over one J-coupling, i.e.
max. 3-4 bonds

TOCSY

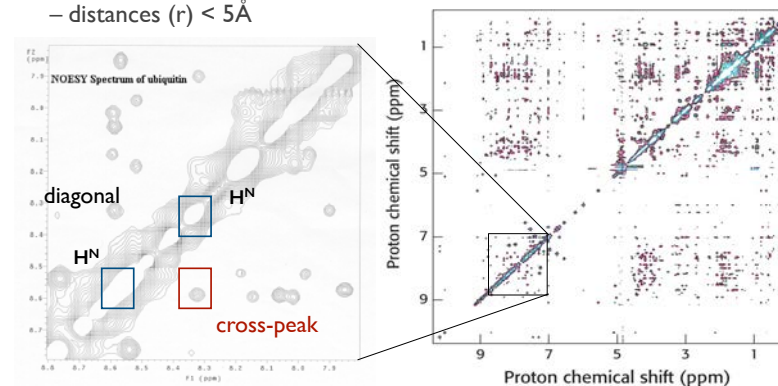


J-coupling interaction
transfer over several J-couplings, i.e. multiple steps
over max. 3-4 bonds

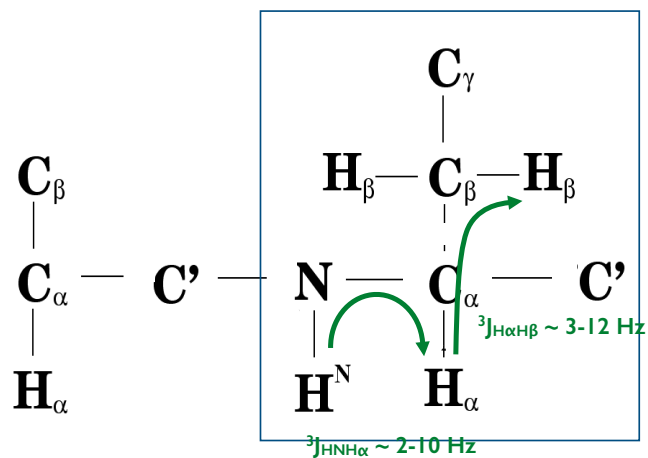
2D NOESY

- Uses dipolar interaction (NOE) to transfer magnetization between protons

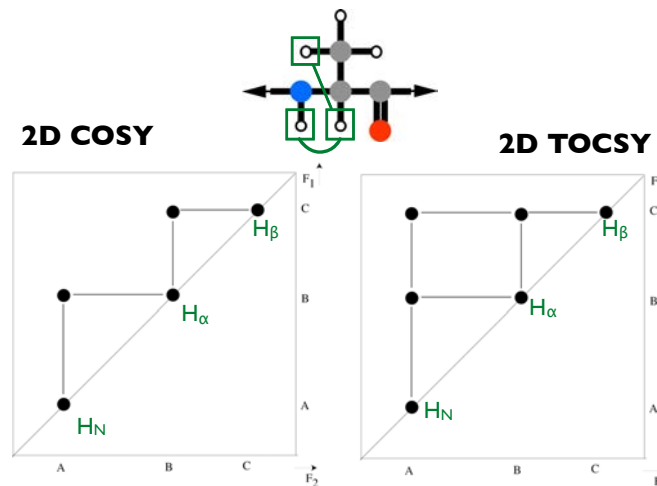
- cross-peak intensity $\sim 1/r^6$
- distances (r) $< 5 \text{ Å}$



Homonuclear scalar coupling

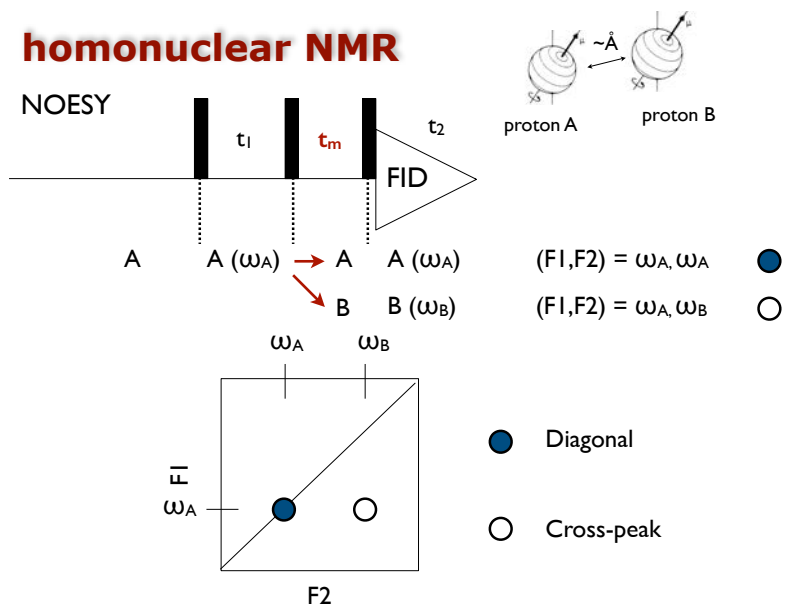


2D COSY & TOCSY

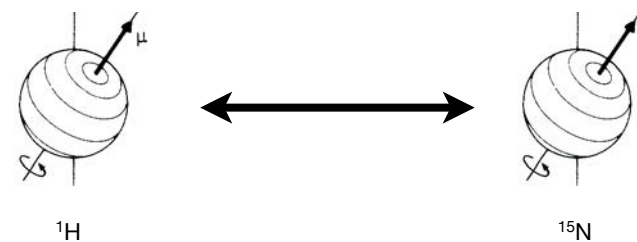


homonuclear NMR

NOESY

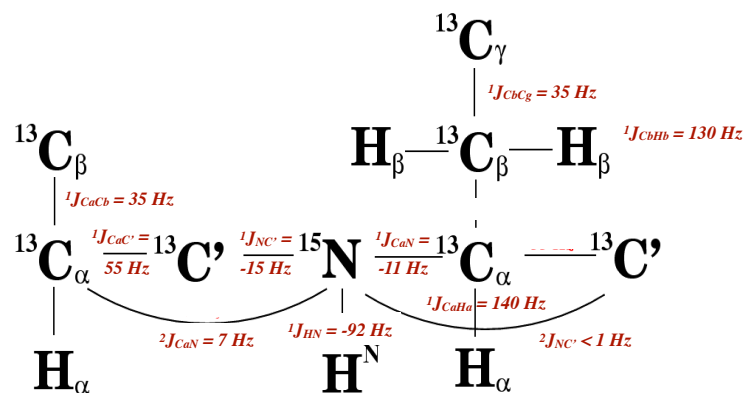


heteronuclear NMR



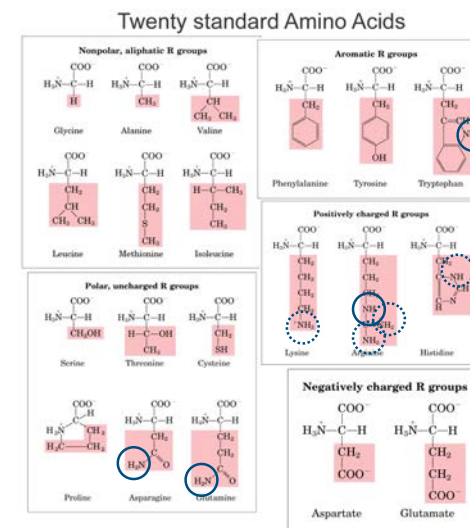
- measure frequencies of different nuclei; e.g. ^1H , ^{15}N , ^{13}C
- no diagonal peaks
- mixing not possible using NOE, only via J

J coupling constants

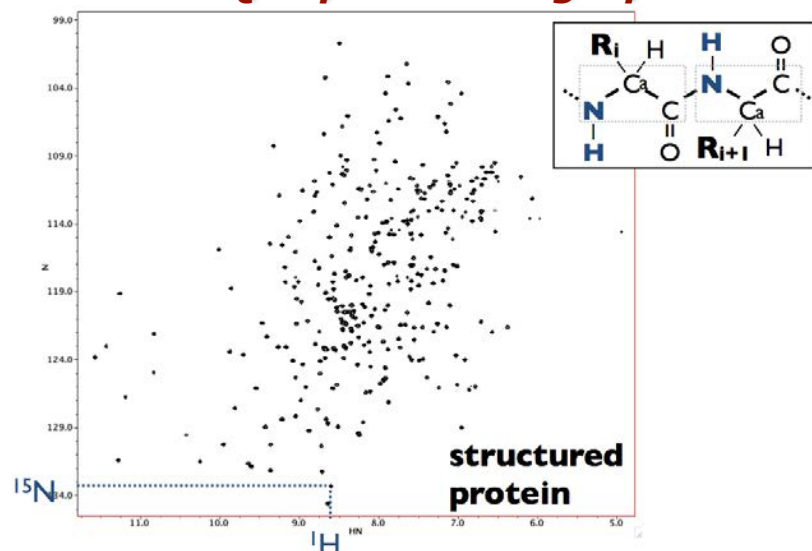


^{15}N HSQC

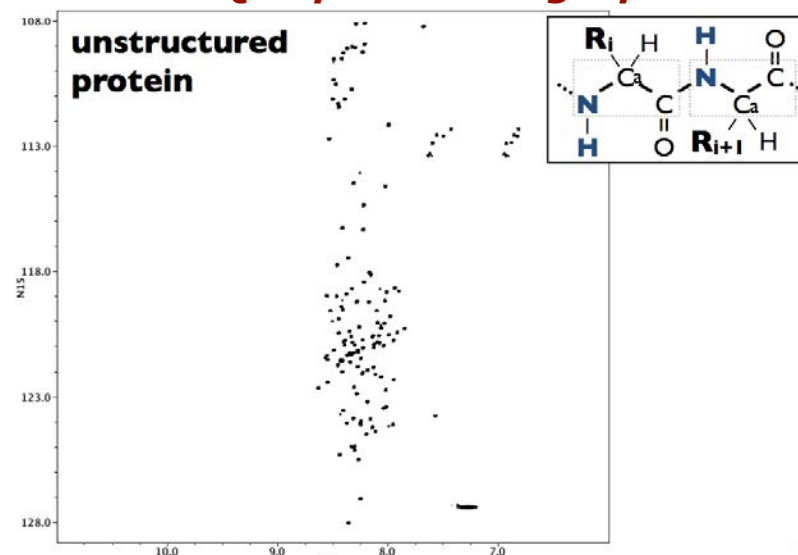
- Backbone HN
- Side-chain NH and NH_2



^1H - ^{15}N HSQC: 'protein fingerprint'



^1H - ^{15}N HSQC: 'protein fingerprint'



Key concepts multidimensional NMR

- Resolve overlapping signals
- Mixing/magnetization transfer
- NOESY, TOCSY, COSY
- HSQC
- 3D NOESY-HSQC, 3D TOCSY-HSQC
- Triple resonance

Relaxation & dynamics

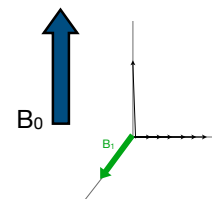
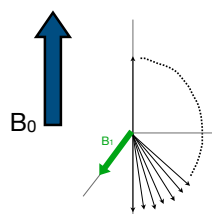


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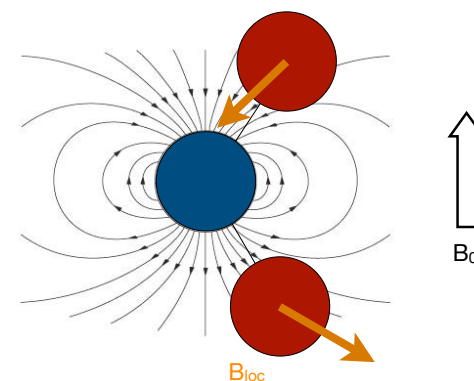
NMR relaxation

- Return to equilibrium
 - Spin-lattice relaxation
 - Longitudinal relaxation → **T1** relaxation
 - Return to z-axis
 - Spin-spin relaxation
 - Transversal relaxation → **T2** relaxation
 - Dephasing of magnetization in the x/y plane



Relaxation is caused by dynamics

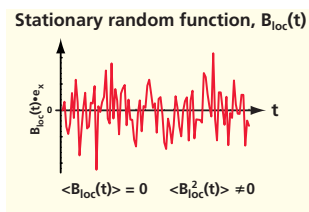
- Fluctuating magnetic fields
 - Overall tumbling and local motions cause the local magnetic fields to fluctuate in time



Relaxation is caused by dynamics

- **Fluctuating magnetic fields**

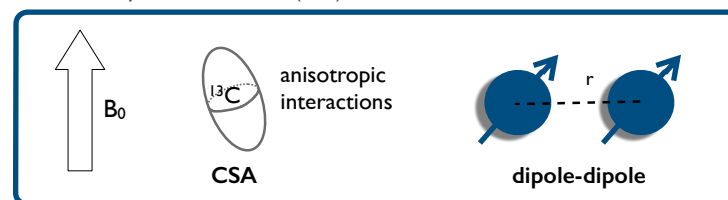
- Overall tumbling and local motions cause the local magnetic fields to fluctuate in time
- $B_{loc}(t)$ is thus time dependent
- If $B_{loc}(t)$ is fluctuating with frequency components near ω_0 then transitions may be induced that bring the spins back to equilibrium
- The efficiency of relaxation also depends on the amplitude of $B_{loc}(t)$



Local fluctuating magnetic fields

- $B_{loc}(t) = B_{loc}[iso] + B_{loc}(t)[aniso]$

- Isotropic part is not time dependent
 - chemical shift
 - J-coupling
- Only the anisotropic part is time dependent
 - chemical shift anisotropy (CSA)
 - dipolar interaction (DD)



Local fluctuating magnetic fields

- $B_{loc}(t) = B_{loc}[iso] + B_{loc}(t)[aniso]$

- Isotropic part is not time dependent
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 - chemical shift anisotropy (CSA)
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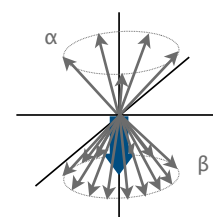
- Only $B_{loc}(t)[aniso]$ can cause relaxation

- Transverse fluctuating fields: $B_{loc}(t) \cdot e_x + B_{loc}(t) \cdot e_y$
- Longitudinal fluctuating fields: $B_{loc}(t) \cdot e_z$

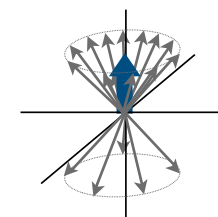
Components of the local field

- $B_{loc}(t) \cdot e_{xy}$

- Transverse fluctuating fields
- **Non-adiabatic:** exchange of energy between the spin-system and the lattice [environment]



transitions between states restore Boltzman equilibrium

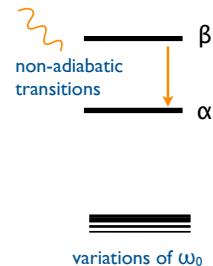


T_1 relaxation

Components of the local field

- $B_{loc}(t) \cdot e_{xy}$

- Transverse fluctuating fields
- **Non-adiabatic**: exchange of energy between the spin-system and the lattice [environment]

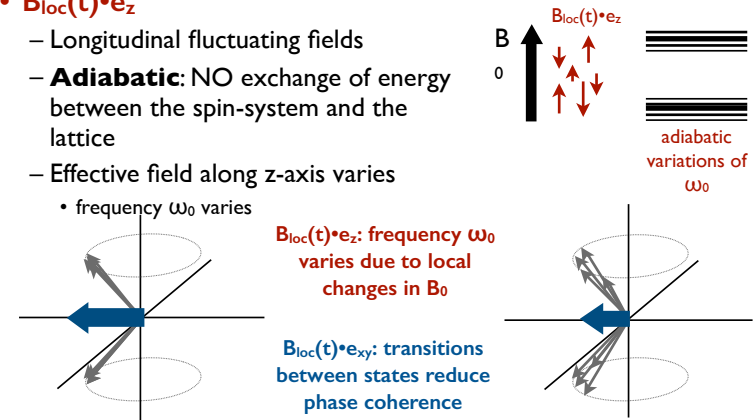


- Heisenberg's uncertainty relationship:
 - shorter lifetimes $\leftrightarrow$ broadening of energy levels

Components of the local field

- $B_{loc}(t) \cdot e_z$

- Longitudinal fluctuating fields
- **Adiabatic**: NO exchange of energy between the spin-system and the lattice
- Effective field along z-axis varies
 - frequency ω_0 varies

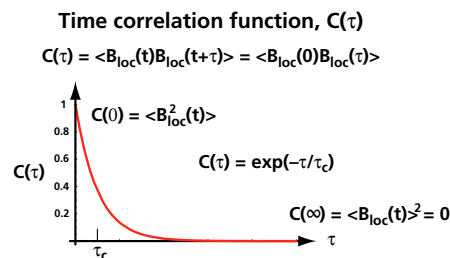
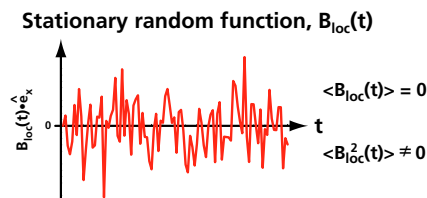


T_2 relaxation

Correlation function

- Describes the fluctuating magnetic fields

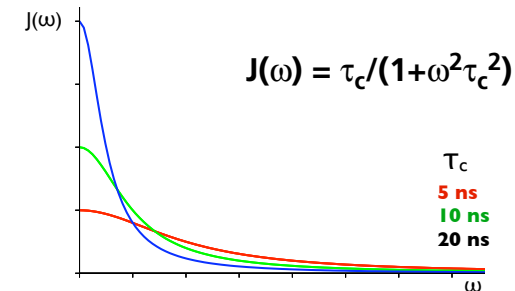
- correlation function $C(\tau)$ decays exponentially with a characteristic time τ_c



Spectral density function

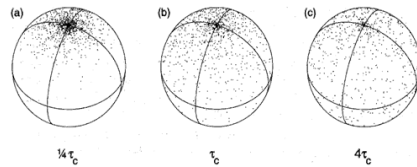
- Frequencies of the random fluctuating fields

- Spectral density function $J(\omega)$ is the Fourier transform of the correlation function $C(\tau)$
- $J(\omega)$ describes if a certain frequency can induce relaxation and whether it is efficient



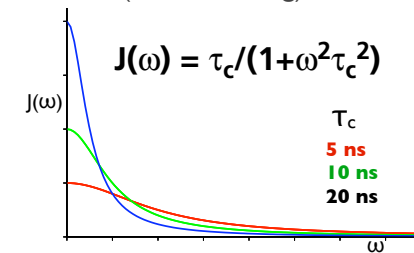
Link to rotational motions in liquids

- **Molecules in solution**
“tumble” (rotational diffusion combining rotations and collisions with other molecules)
- Can be characterized by a **rotational correlation time** τ_c
- τ_c is the time needed for the rms deflection of the molecules to be ~ 1 radian (60°)



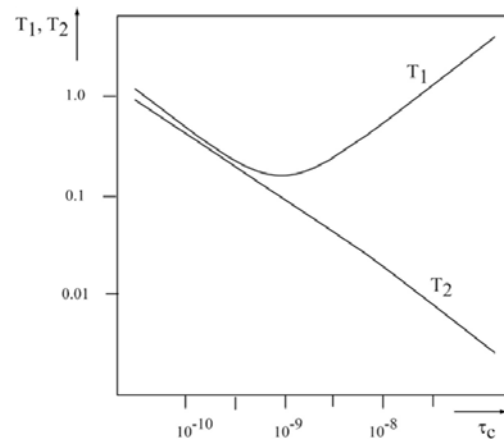
Link to rotational motions in liquids

- **Small molecules** (or high temperature):
 - smaller (shorter) correlation times (fast tumbling),
 - $J(\omega)$ extends to higher frequencies - spectrum is flatter
- **Large molecules** (or low temperature):
 - larger (longer) correlation times (slow tumbling)
 - $J(\omega)$ larger close to 0



Relaxation

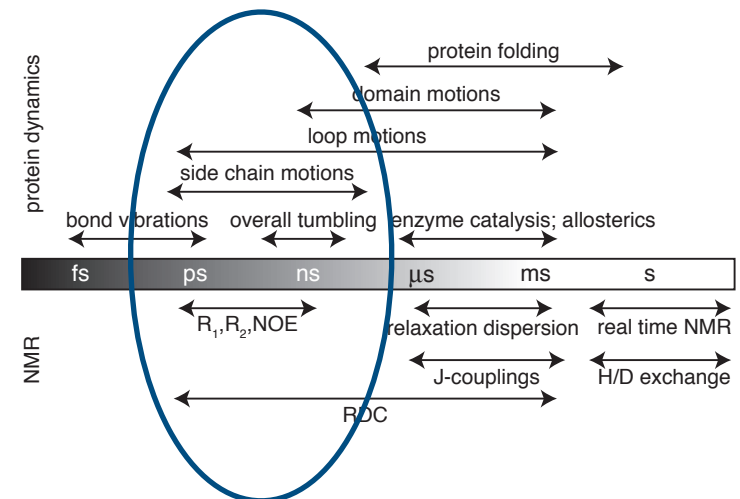
- relaxation time is related to rate of motion



$$R_1 = 1/T_1$$

$$R_2 = 1/T_2$$

NMR time scales



Protein backbone dynamics

- ^{15}N relaxation to describe ps-ns dynamics

- R_1 : longitudinal relaxation rate
- R_2 : transversal relaxation rate
- hetero-nuclear NOE: $\{^1\text{H}\}$ - ^{15}N

$$R_1 = d^2[J(\omega_{\text{H}} - \omega_{\text{N}}) + 3J(\omega_{\text{N}}) + 6J(\omega_{\text{H}} + \omega_{\text{N}})] + c^2J(\omega_{\text{N}})$$

$$R_2 = (d^2/2)[4J(0) + J(\omega_{\text{H}} - \omega_{\text{N}}) + 3J(\omega_{\text{N}}) + 6J(\omega_{\text{H}} + \omega_{\text{N}}) + 6J(\omega_{\text{H}})] + (c^2/6)[4J(0) + J(\omega_{\text{N}})]$$

$$\text{NOE} = 1 + [(\gamma_{\text{H}}/\gamma_{\text{N}})d^2\{6J(\omega_{\text{H}} + \omega_{\text{N}}) - J(\omega_{\text{H}} - \omega_{\text{N}})\}/R_1]$$

where $d^2 = (1/10) \gamma_{\text{H}}^2 \gamma_{\text{N}}^2 (\hbar/2\pi)^2 \langle r_{\text{NH}}^{-3} \rangle^2$ **dipole interaction**

and $c^2 = (2/15) \omega_{\text{N}}^2 \langle \sigma_{\parallel} \cdot \sigma_{\perp} \rangle^2$ **chemical shift anisotropy**

Protein backbone dynamics

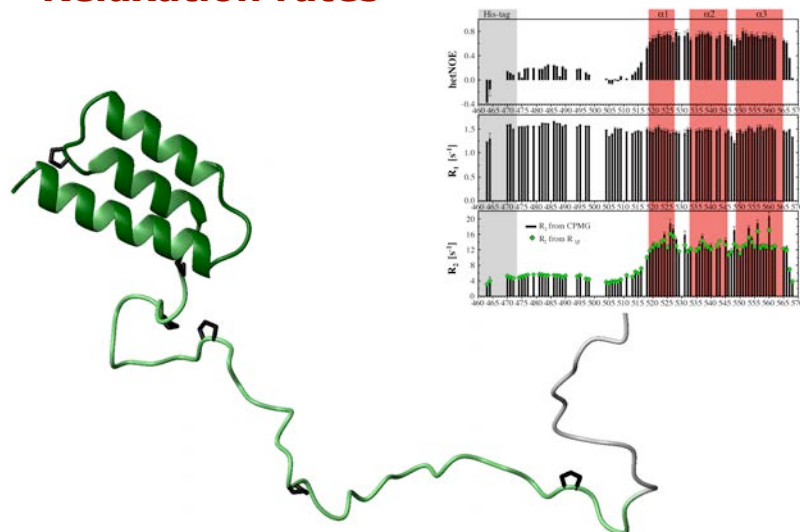
- ^{15}N relaxation to describe ps-ns dynamics

- R_1 : longitudinal relaxation rate
- R_2 : transversal relaxation rate
- hetero-nuclear NOE: $\{^1\text{H}\}$ - ^{15}N

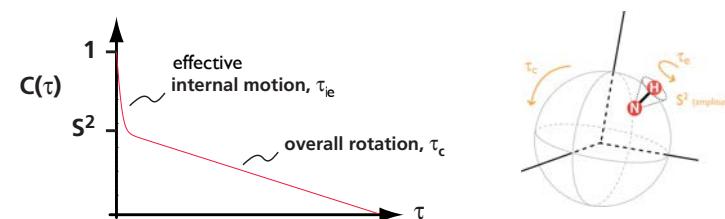
- Measured as a 2D ^1H - ^{15}N spectrum**

- R_1, R_2 : Repeat experiment several times with increasing relaxation-delay
- Fit the signal intensity as a function of the relaxation delay
 - $I_0 \exp(-Rt)$
- $\{^1\text{H}\}$ - ^{15}N NOE: Intensity ratio between saturated and non-saturated experiment

Relaxation rates



Lipari-Szabo MODELFREE

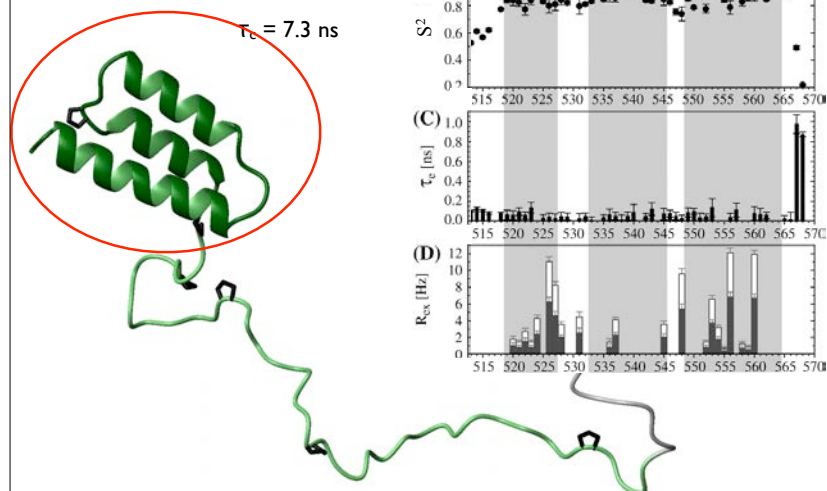


- Overall and local motion are considered to be **uncorrelated**
- S^2 = **order-parameter**

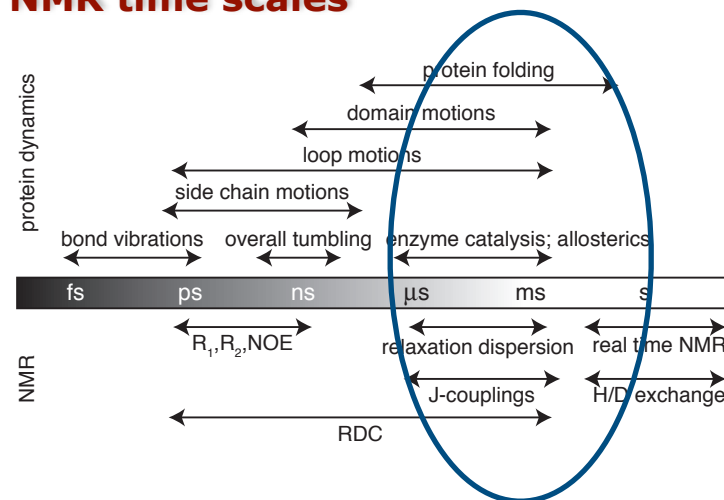
$$\left\{ \begin{array}{l} C_l(\tau=0) = 1 \\ C_l(\tau \rightarrow \infty) = S^2 \\ \int_0^\infty C_l(\tau) d\tau = \tau_c \end{array} \right\} \rightarrow C_l(\tau) = S^2 + (1 - S^2)e^{-\tau/\tau_c} \quad J(\omega) = \frac{2}{5} \left[\frac{S^2 \tau_c}{1 + \omega^2 \tau_c^2} + \frac{(1 - S^2) \tau}{1 + \omega^2 \tau^2} \right]$$

$$\tau^{-1} = \tau_c^{-1} + \tau_e^{-1}$$

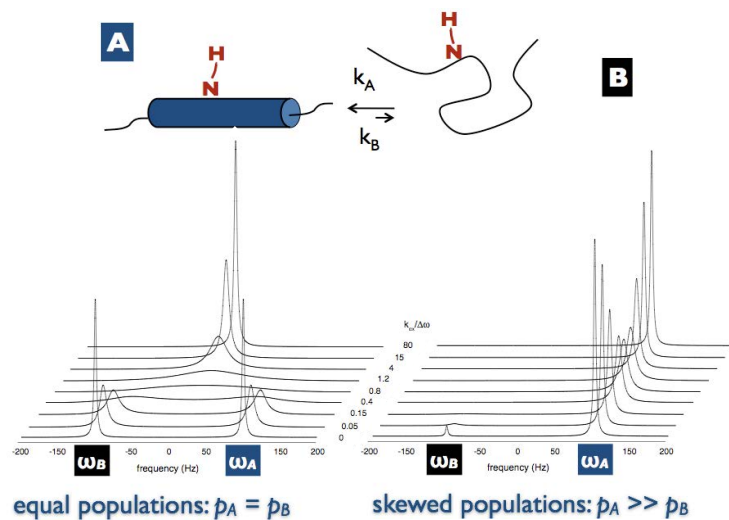
Modelfree analysis



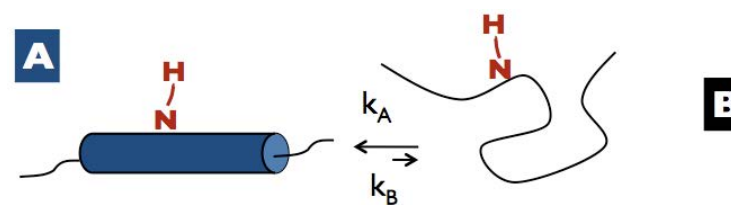
NMR time scales



Conformational exchange



Conformational exchange



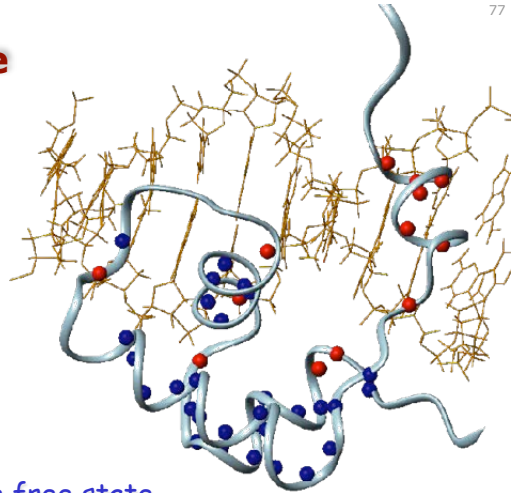
- Causes line-broadening of the signals

$$-R_{2,eff} = R_2 + R_{ex}$$

H/D exchange

Lac headpiece

Kalodimos et al. Science



- protected in the free state
- protected only in the DNA-bound state

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Key concepts relaxation

- time scales
- fluctuating magnetic fields
- correlation function, spectral density function
- molecular motions
- rotational correlation time (ns)
- fast time scale flexibility (ps-ns)
- slow time scale (μ s-ms): conformational exchange

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The End

Thank you for your attention!

