C8953 NMR structural analysis seminar Information about classes + 1D ¹H-NMR

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Information about classes

Credit:

midterm test and final project

Study materials:

https://is.muni.cz/auth/el/1431/jaro2023/C8953/um

E-tests:

https://is.muni.cz/auth/el/1431/jaro2023/C8953/odp

Energy levels splitting



Behavior of nuclear spin after irradiation by RF pulse



Precession frequency:

Precession frequency affected by nuclear shielding: Chemical shift:

Definition of th relative scale of the chemical shift:



B₀ induces local mag. field B_{loc}, which affects against B₀ ↓ Nuclear shielding

$$\begin{split} & \omega = -\gamma B_{0} \\ & \omega = -(1+\sigma)B_{0} \\ & \delta = \omega - \omega_{ref} \\ & \delta = (\omega - \omega_{ref})/\omega_{ref}.10^{6} ppm \end{split}$$

Characteristic intervals of chemical shifts values



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Trends in chemical shifts

- Electronegativity, inductive and mesomeric effects of substituents
- Hybridisation
- Relative position towards the ring, double bond



Mesomeric effect



Substituents with -M effects

-F, -Cl, -Br, -I, -OH, -OR, -NH₂, -NHR, -NR₂, -SH, -SR

Substituents with +M effect

-CH=O, -RC=O, -C(OH)=O, -C(OR)=O, -C(NH₂)=O, -NO₂, -SO₃H, -C=N

Spin-spin interaction, J-coupling



Interaction constant J



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Interaction constant J



- Multiplicity of the nucleus I with the spin 1/2 is given by: m = n + 1, n = number of interacting nuclei with nucleus I
- Intensity of lines in multiplet follows Pascal's triangle



1D¹H NMR spectrum



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Values of *J*-constants - trends



 $^{2}J_{CH}$ = 3.1 Hz



 ${}^{3}J_{\rm HH}$ = 12 Hz



 ${}^{3}J_{HH} = 13 - 18 \text{ Hz}$ ${}^{5}J_{HH} = 7 - 12 \text{ Hz}$











 $^{2}J_{\rm HH}$ = -12,5 Hz

 ${}^{5}J_{HH}$ = 2 - 3 Hz ${}^{1}J_{CH}$ = 125 Hz ${}^{1}J_{CH}$ = 160 Hz ${}^{1}J_{CH}$ = 250 Hz

Values of *J*-constants - trends







X=	Li	Н	CI	OMe	F
² J _{HH} (Hz)	7,1	2,5	-1,4	-2,0	-3,2

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1D¹H NMR spectroscopy

- the fastest measuring, the highest sensitivity
- complicated interpretation in case of more complex systems

We are looking for:

- position of the signal (ppm)
- multiplicity $(^{2}J, ^{3}J, ^{4}J)$
- intensity (integral)
- halfwidth

We are considering:

- chemical/magnetic equivalence
- enantiotopicity/diastereotopicity
- averaging of signals (dynamics, chemical exchange)

1D¹H NMR spectrum of methyl-5-acetylsalicylate



1D¹H NMR spectrum of cinnamic acid



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Draw approximate 1D ¹H NMR spectrum of the following compound





1D¹H NMR spectrum of ethyl glutarate



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1D¹H NMR spectrum of benzyl butyrate



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1D ¹H NMR - ethyl 4-(methylamino)benzoate

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1D¹H NMR spectrum of cartilagineal



Next session:

1D ¹³C-NMR spectra

