

# C8953

## NMR structural analysis seminar

Information about classes + 1D  $^1\text{H}$ -NMR

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February 19, 2020

# Information about classes

## Credit:

- ▶ Max. 2 unexcused absences
- ▶ pass 2 tests (midterm and final)

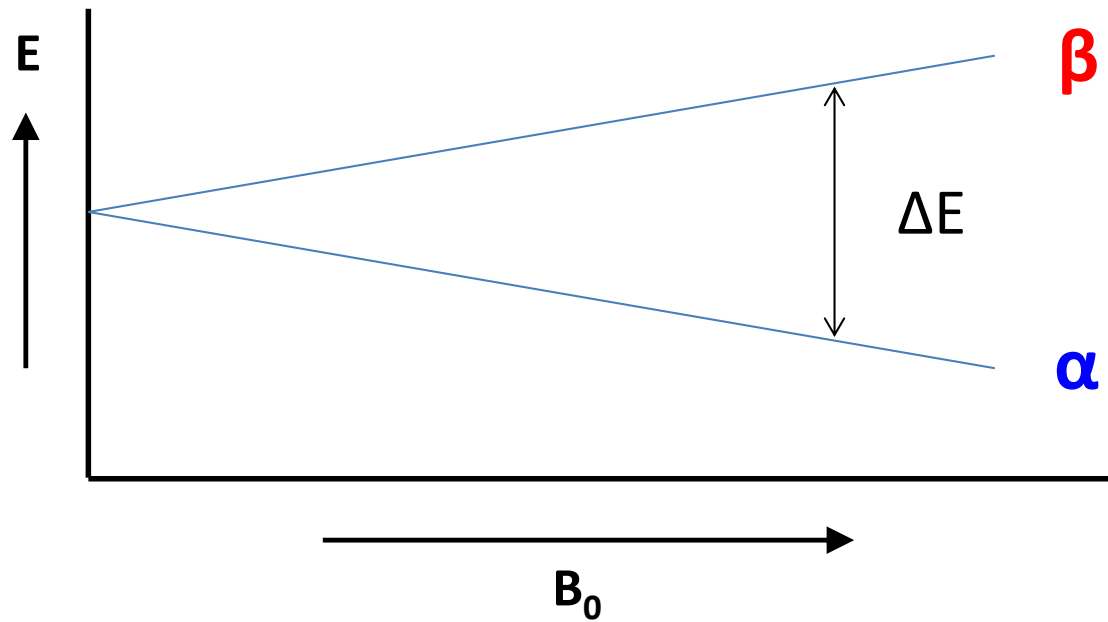
## Study materials:

<https://is.muni.cz/auth/el/1431/jaro2020/C8953/um>

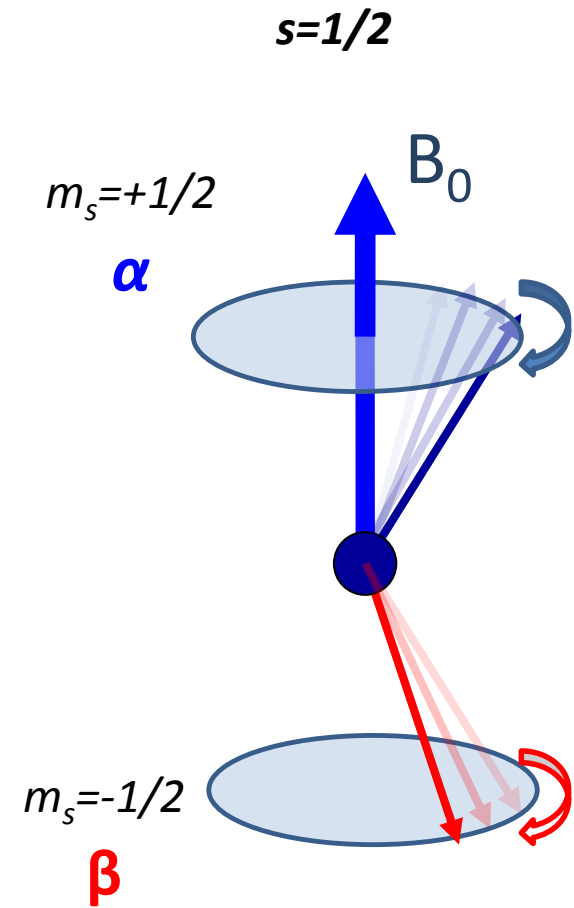
## E-tests:

<https://is.muni.cz/auth/el/1431/jaro2020/C8953/odp>

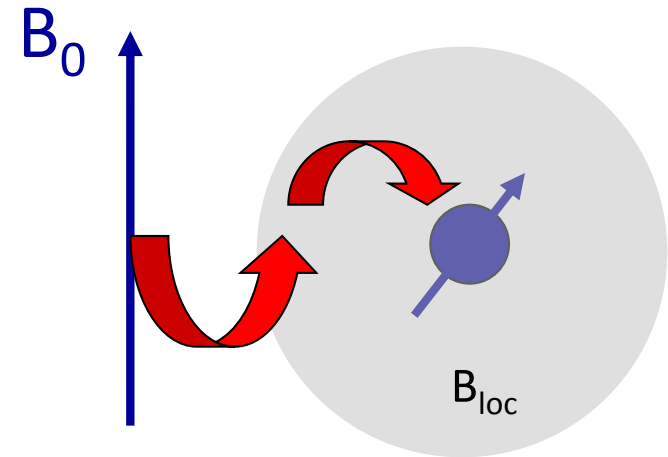
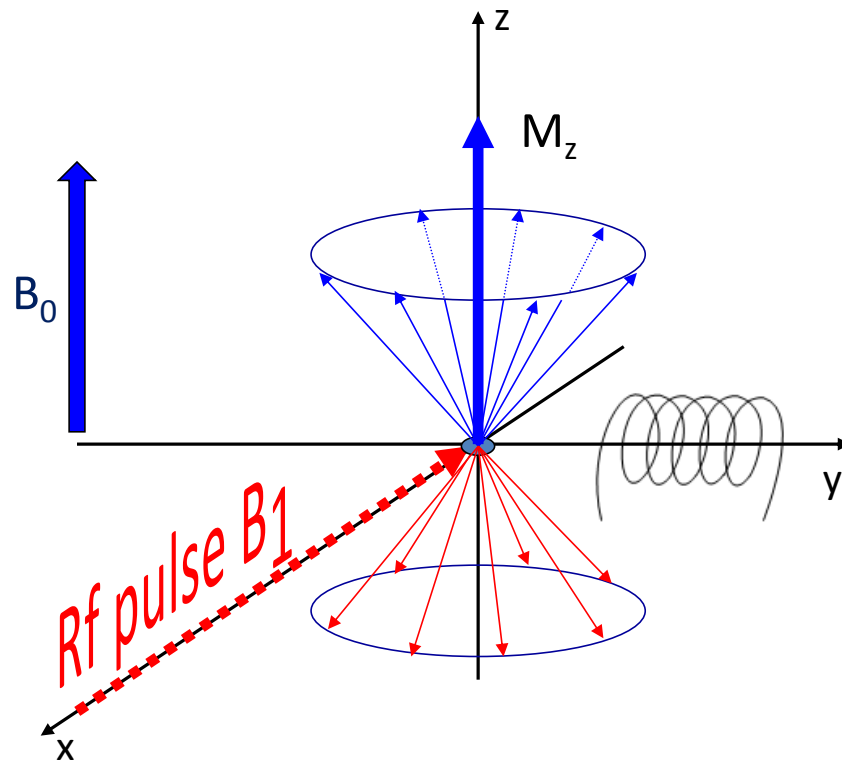
# Energy levels splitting



$$N_\alpha > N_\beta$$



# Behavior of nuclear spin after irradiation by RF pulse



$B_0$  induces local mag. field  $B_{loc}$ , which affects against  $B_0$

↓  
Nuclear shielding

Precession frequency:

Precession frequency affected by nuclear shielding:

Chemical shift:

Definition of the relative scale of the chemical shift:

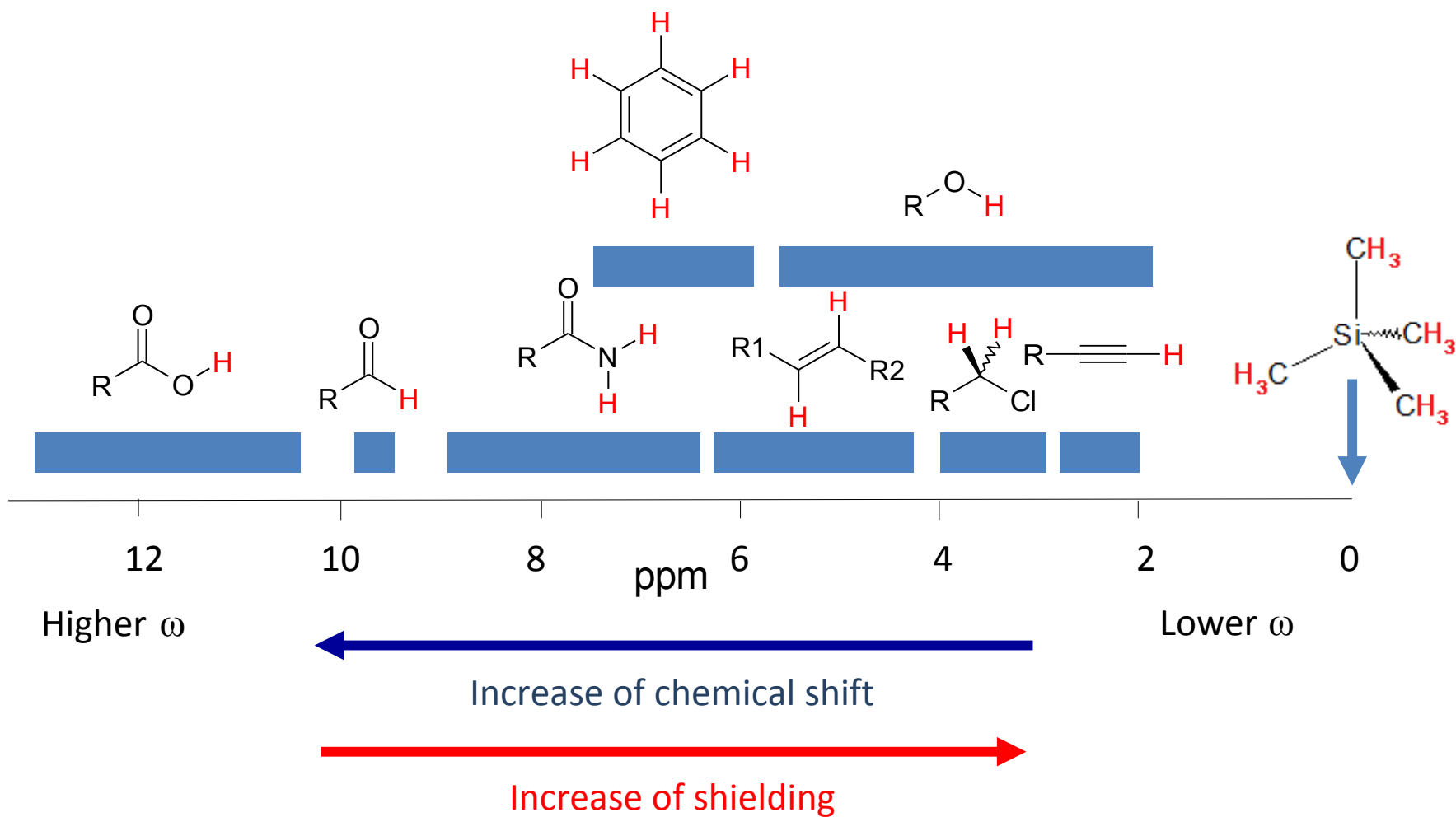
$$\omega = -\gamma B_0$$

$$\omega = -(1+\sigma)B_0$$

$$\delta = \omega - \omega_{ref}$$

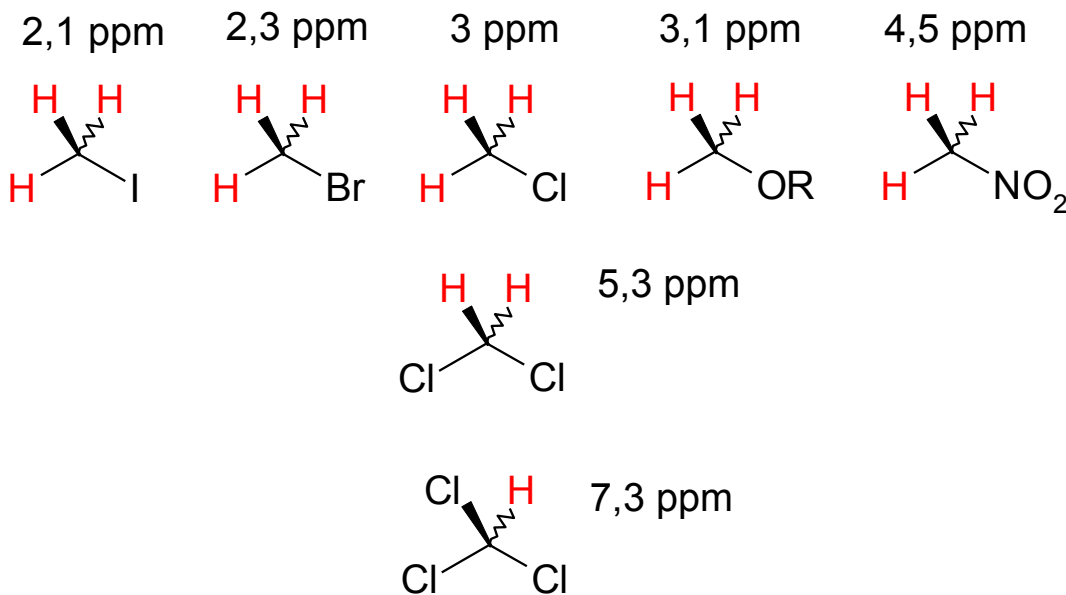
$$\delta = (\omega - \omega_{ref})/\omega_{ref} \cdot 10^6 \text{ ppm}$$

# Characteristic intervals of chemical shifts values



# Trends in chemical shifts

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



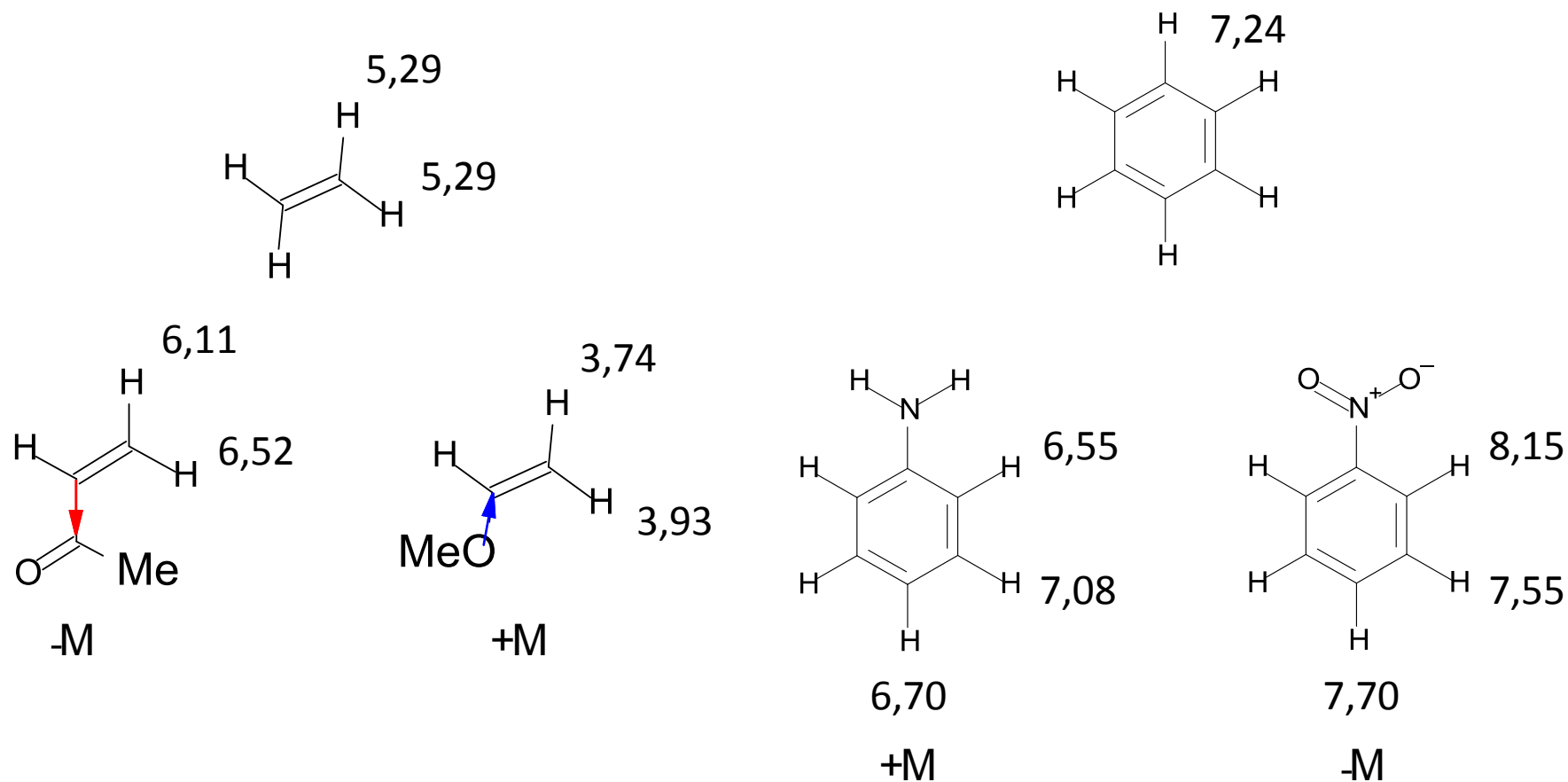
## Substituents with -I effect

$=N^+R_2 > -N^+R_3 > -NO_2 > -NR_2$   
 $-SO_2R > -SO_3 > -SOR > -SR$   
 $-F > -OR > -NR_2 > -CR_3$   
 $-F > -Cl > -Br > -I$   
 $\equiv N > =NR > -NR_2$   
 $-C\equiv CH > -CH=CH_2 > -CH_2-CH_3$

## Substituents with +I effects

$-N-R > -O- > S-$   
 $-C(CH_3)_3 > -CH(CH_3)_2 > -CH_2CH_3 > -CH_3$   
 metals

# Mesomeric effect



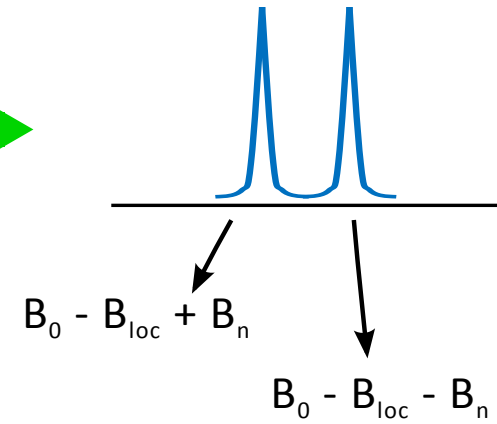
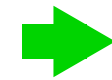
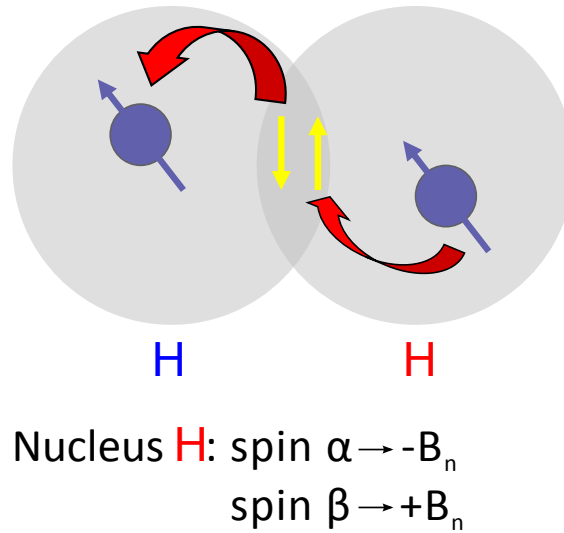
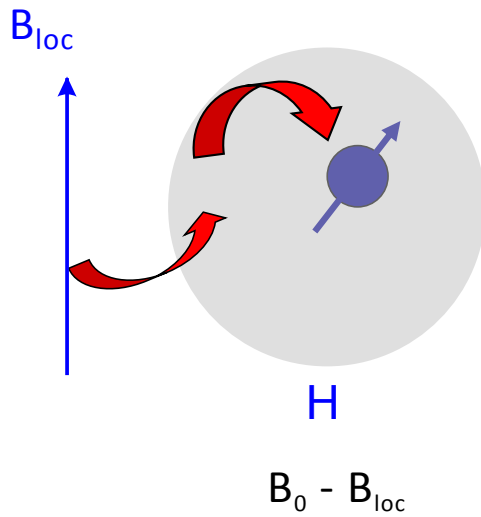
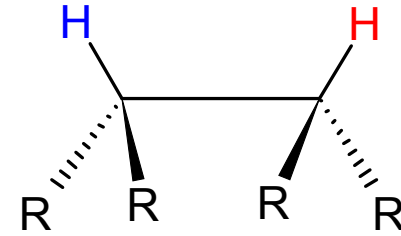
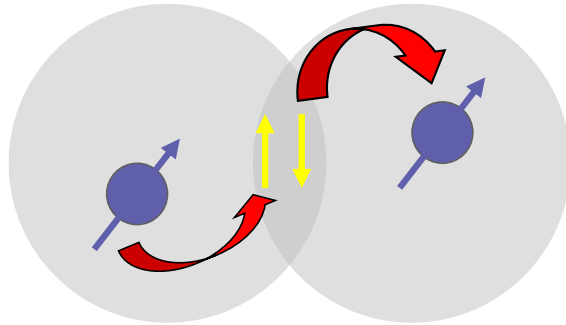
## Substituents with -M effects

-F, -Cl, -Br, -I, -OH, -OR, -NH<sub>2</sub>, -NHR, -NR<sub>2</sub>, -SH, -SR

## Substituents with +M effect

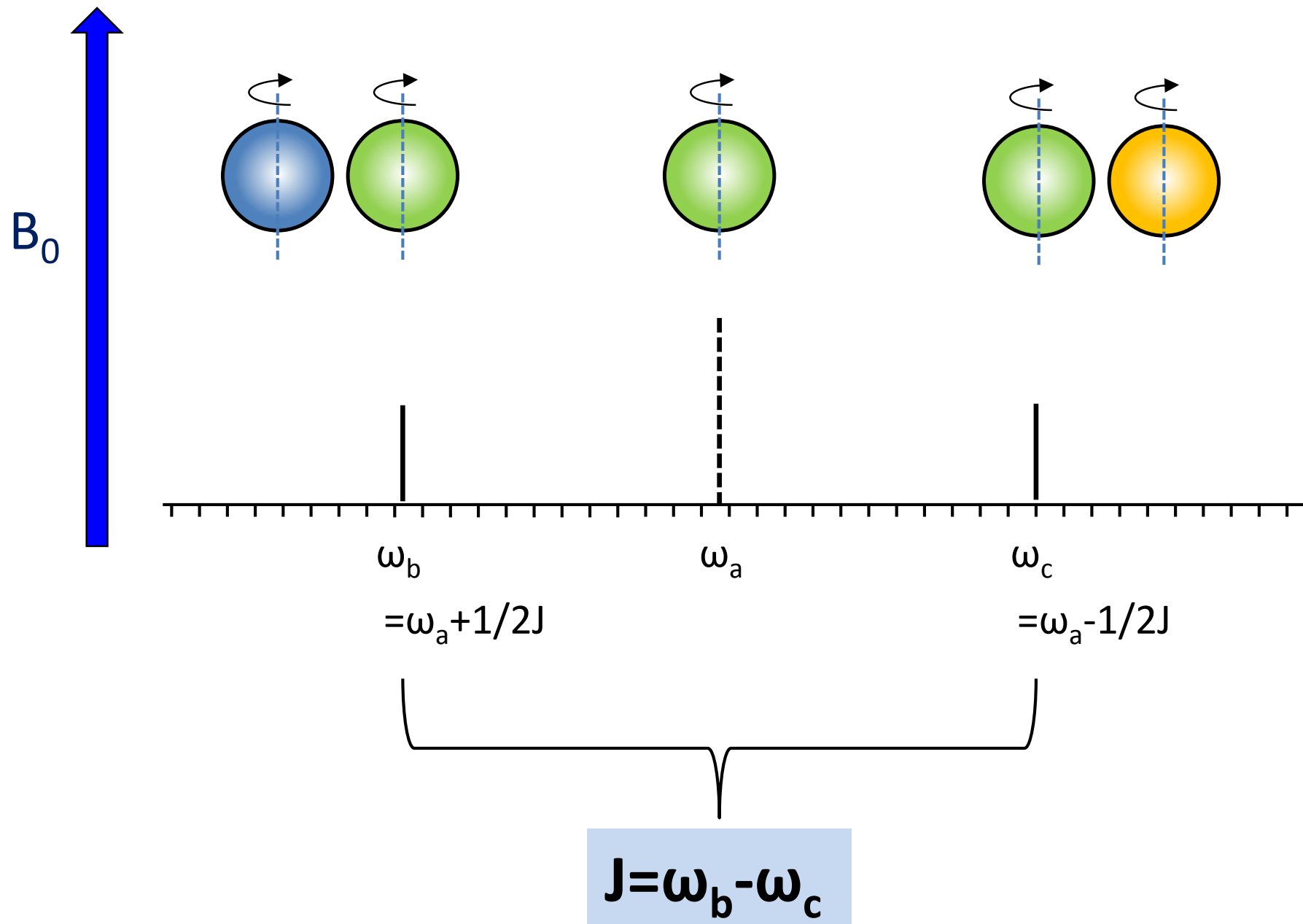
-CH=O, -RC=O, -C(OH)=O, -C(OR)=O, -C(NH<sub>2</sub>)=O, -NO<sub>2</sub>, -SO<sub>3</sub>H, -C≡N

# Spin-spin interaction, $J$ -coupling

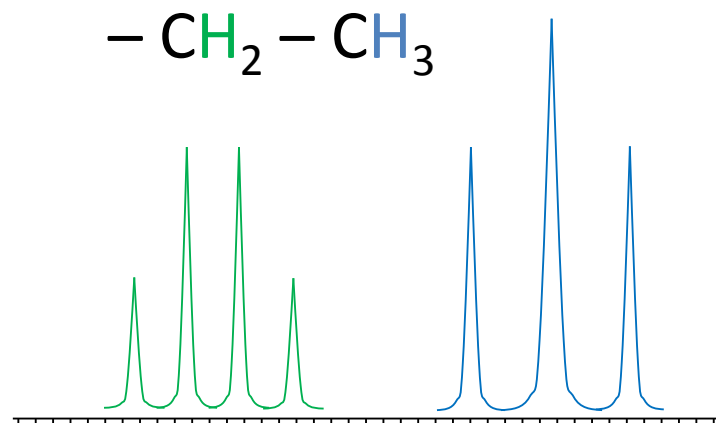




# Interaction constant $J$



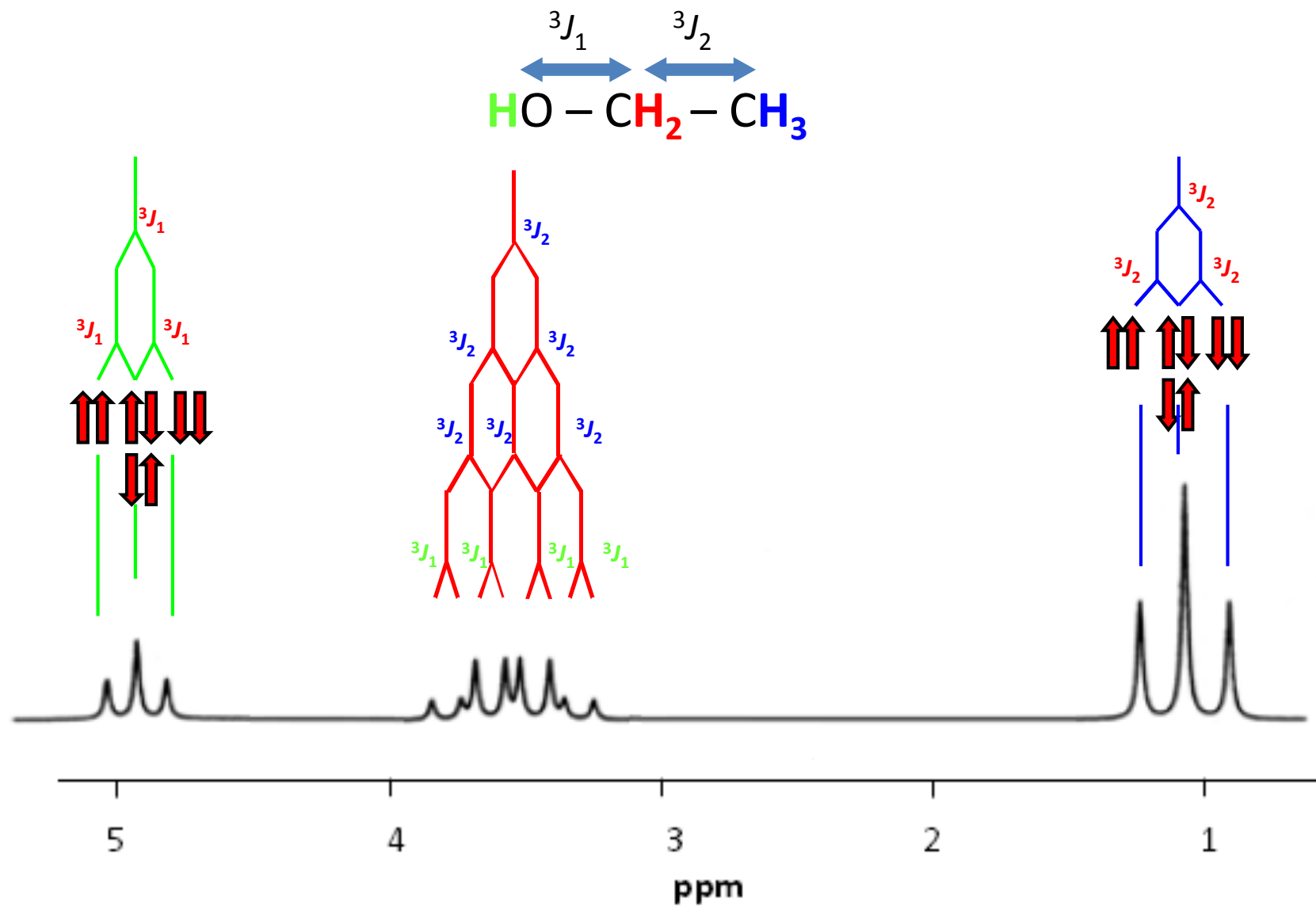
# Interaction constant $J$



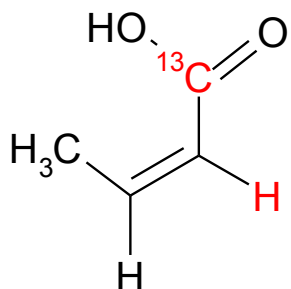
- ▶ Multiplicity of the nucleus I with the spin  $1/2$  is given by:  
 $m = n + 1$ ,  $n =$  number of interacting nuclei with nucleus I
- ▶ Intensity of lines in multiplet follows Pascal's triangle

			1			
			1	1		
		1	2	1		
		1	3	3	1	
	1	4	6	4	1	
1	5	10	10	5	1	

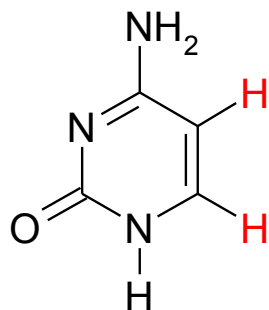
# 1D $^1\text{H}$ NMR spectrum



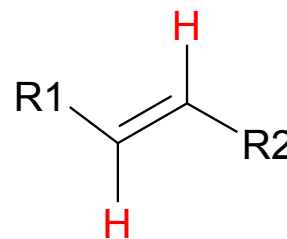
# Values of $J$ -constants - trends



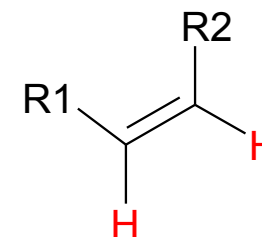
$${}^2J_{CH} = 3.1 \text{ Hz}$$



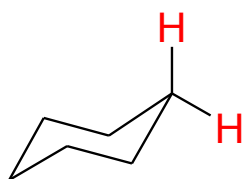
$${}^3J_{HH} = 12 \text{ Hz}$$



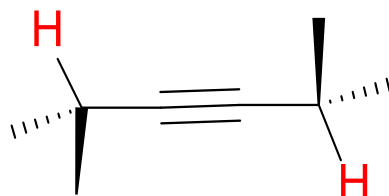
$${}^3J_{HH} = 13 - 18 \text{ Hz}$$



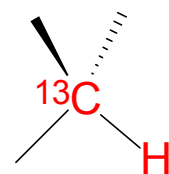
$${}^5J_{HH} = 7 - 12 \text{ Hz}$$



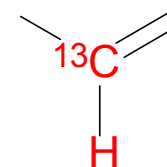
$${}^2J_{HH} = -12,5 \text{ Hz}$$



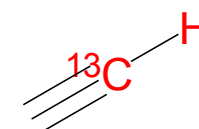
$${}^5J_{HH} = 2 - 3 \text{ Hz}$$



$${}^1J_{CH} = 125 \text{ Hz}$$

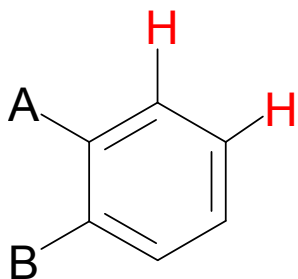


$${}^1J_{CH} = 160 \text{ Hz}$$

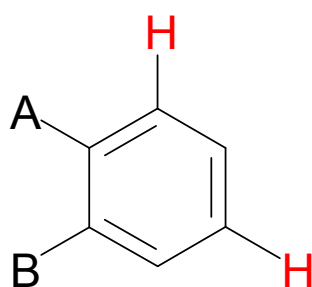


$${}^1J_{CH} = 250 \text{ Hz}$$

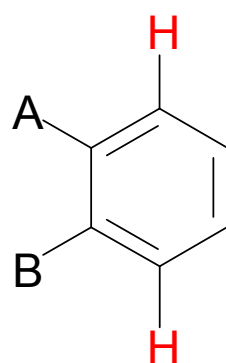
# Values of J-constants - trends



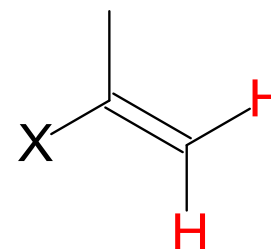
$${}^3J_{HH} = 7,5 \text{ Hz}$$



$${}^4J_{HH} = 1,5 \text{ Hz}$$



$${}^5J_{HH} = 0,7 \text{ Hz}$$



X=	Li	H	Cl	OMe	F
${}^2J_{HH}$ (Hz)	7,1	2,5	-1,4	-2,0	-3,2

# 1D $^1\text{H}$ NMR spectroscopy

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

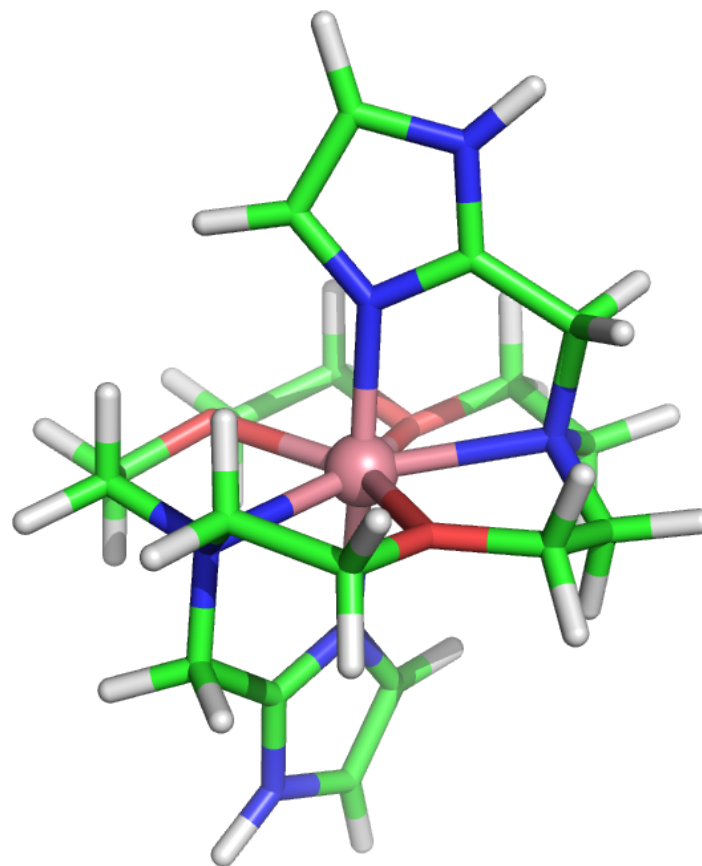
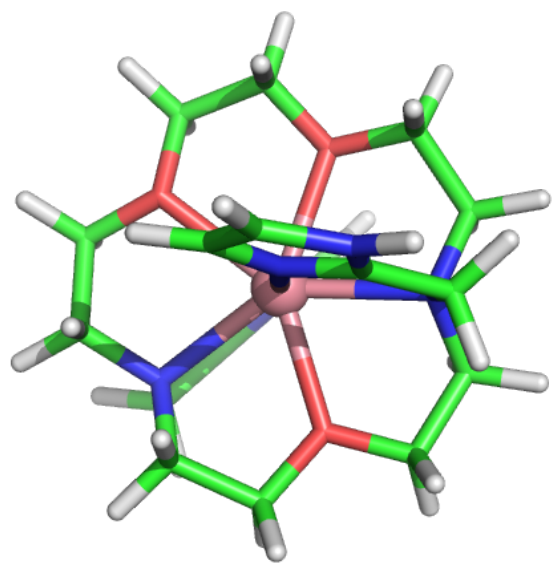
## We are looking for:

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

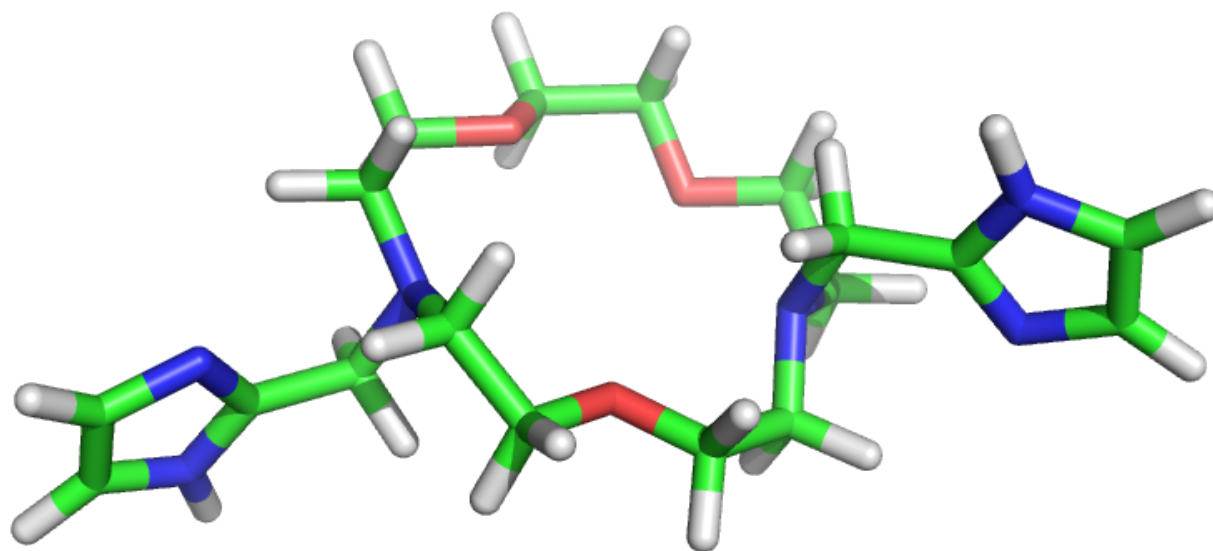
## We are considering:

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

# Determine number of signals in $^1\text{H}$ NMR spectrum

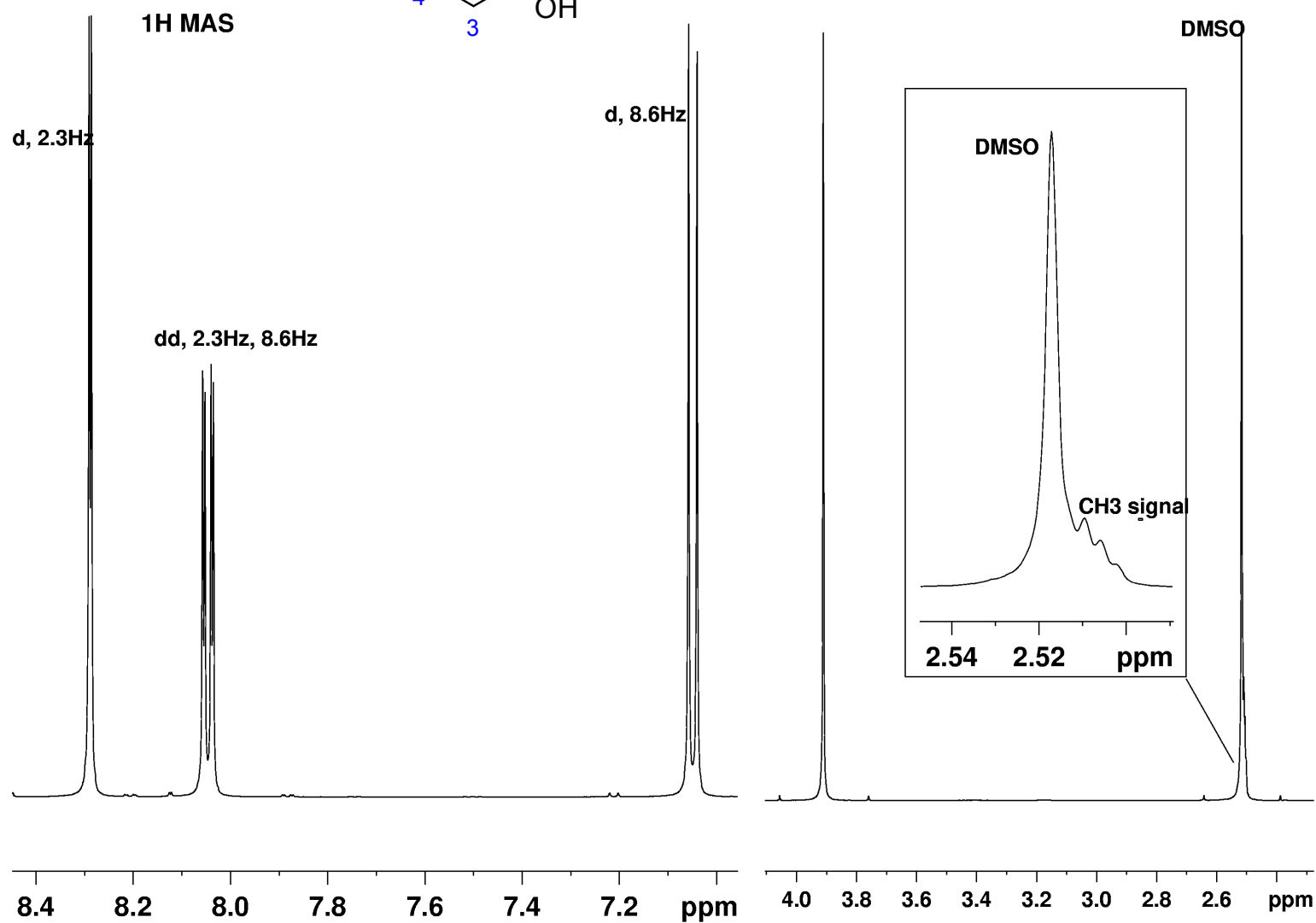
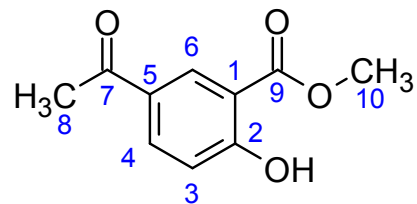


# Determine number of signals in $^1\text{H}$ NMR spectrum

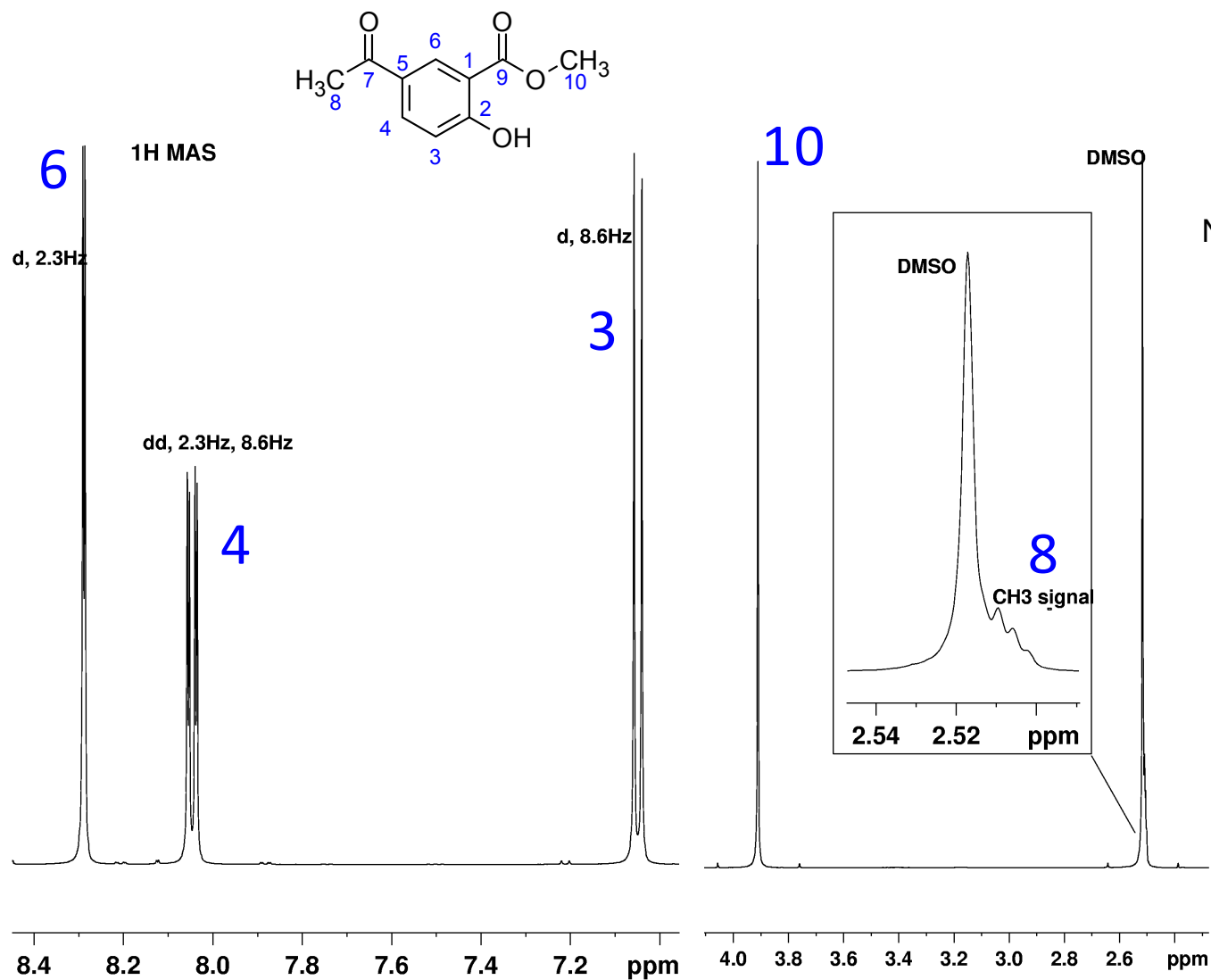




# 1D $^1\text{H}$ NMR spectrum of methyl-5-acetylsalicylate



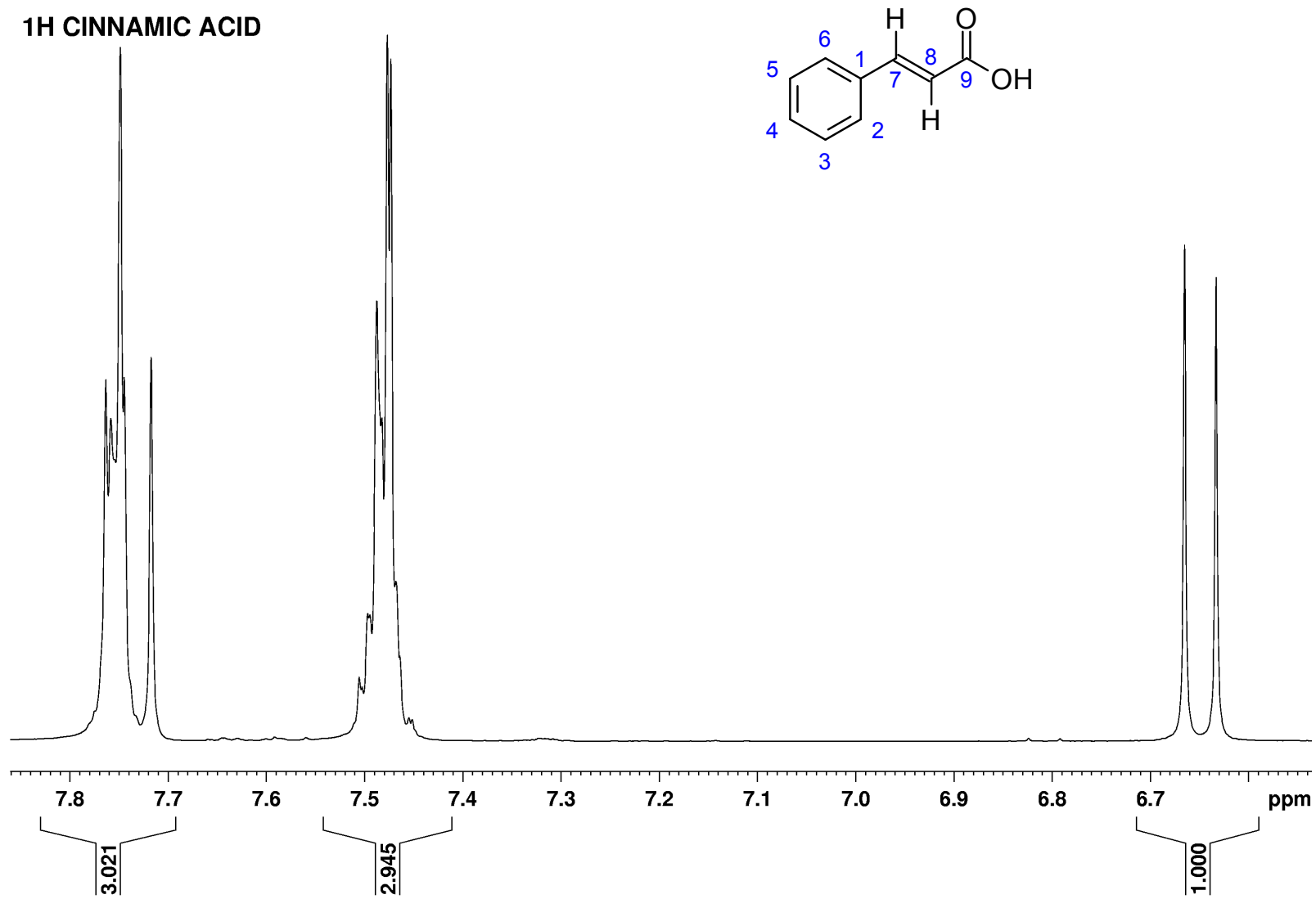
# 1D $^1\text{H}$ NMR spectrum of methyl-5-acetylsalicylate



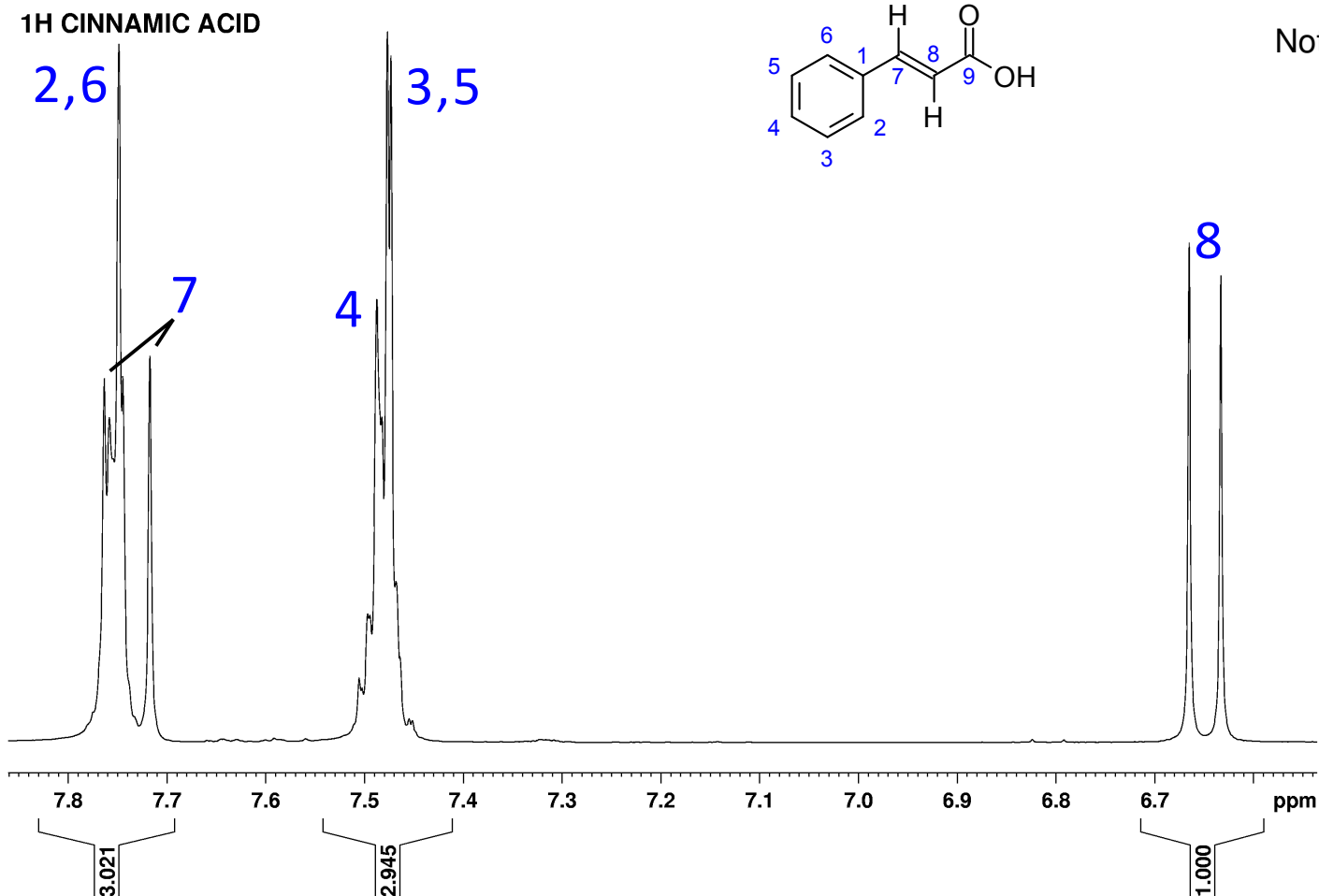
Notes:

- ▶ two singlets in the spectrum - two isolated groups in the structure -  $\text{CH}_3$  groups; **Met-8** neighboring carbonyl has lowest shift than ester **Met-10**
- ▶ doublet of doublets (cca 8.0 ppm) - proton signal splitted by two neighbors - **H-4**
- ▶ two doublets in interaction with H-4 - based on the  $J$ -interaction: doublet with larger  $J$ -constant belongs to close proton - **H-3**, smaller  $J$ -constant - more distant proton - **H-6**

# 1D $^1\text{H}$ NMR spectrum of cinnamic acid



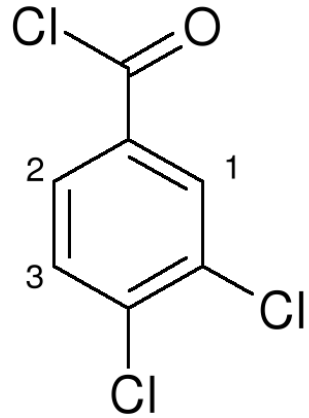
# 1D $^1\text{H}$ NMR spectrum of cinnamic acid



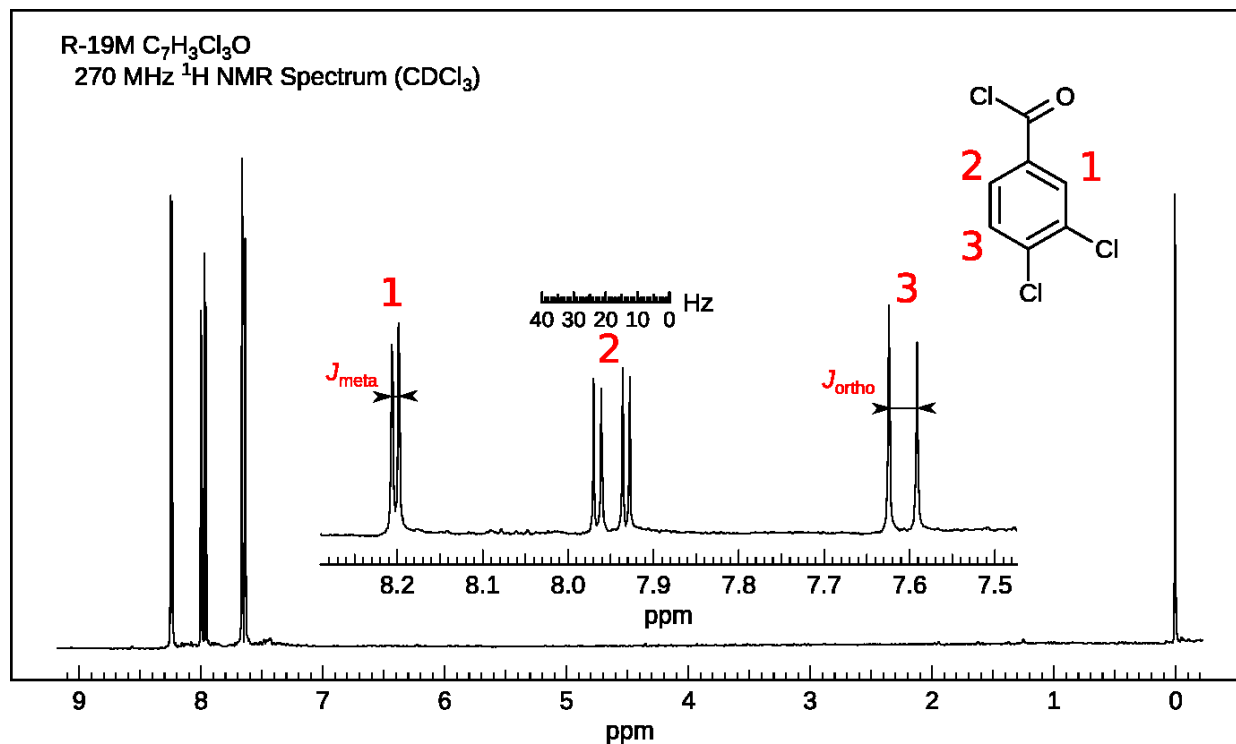
Notes:

- ▶ **H-8** - doublet with large coupling, in range of shifts of protons on double bond, integral = 1
- ▶ **H-7** - doublet with the same coupling like doublet H-8, deshielded due to -M effect of carboxyl and due to nearby aromatic ring
- ▶ more intensive signal between 7.7 and 7.8 ppm has integral: 3-1=2 protons - **H-2,6**, symmetrical, highest shift due to -M effect of substituent in *ortho* position on aromatic ring
- ▶ signal with integral = 3 around 7.5 ppm - less intensive signal - only one proton, highest shift due to -M effect of substituent in *para* position - **H-4**; more intensive signal with highest shift - **H-3,5**

Draw approximate 1D  $^1\text{H}$  NMR spectrum of the following compound



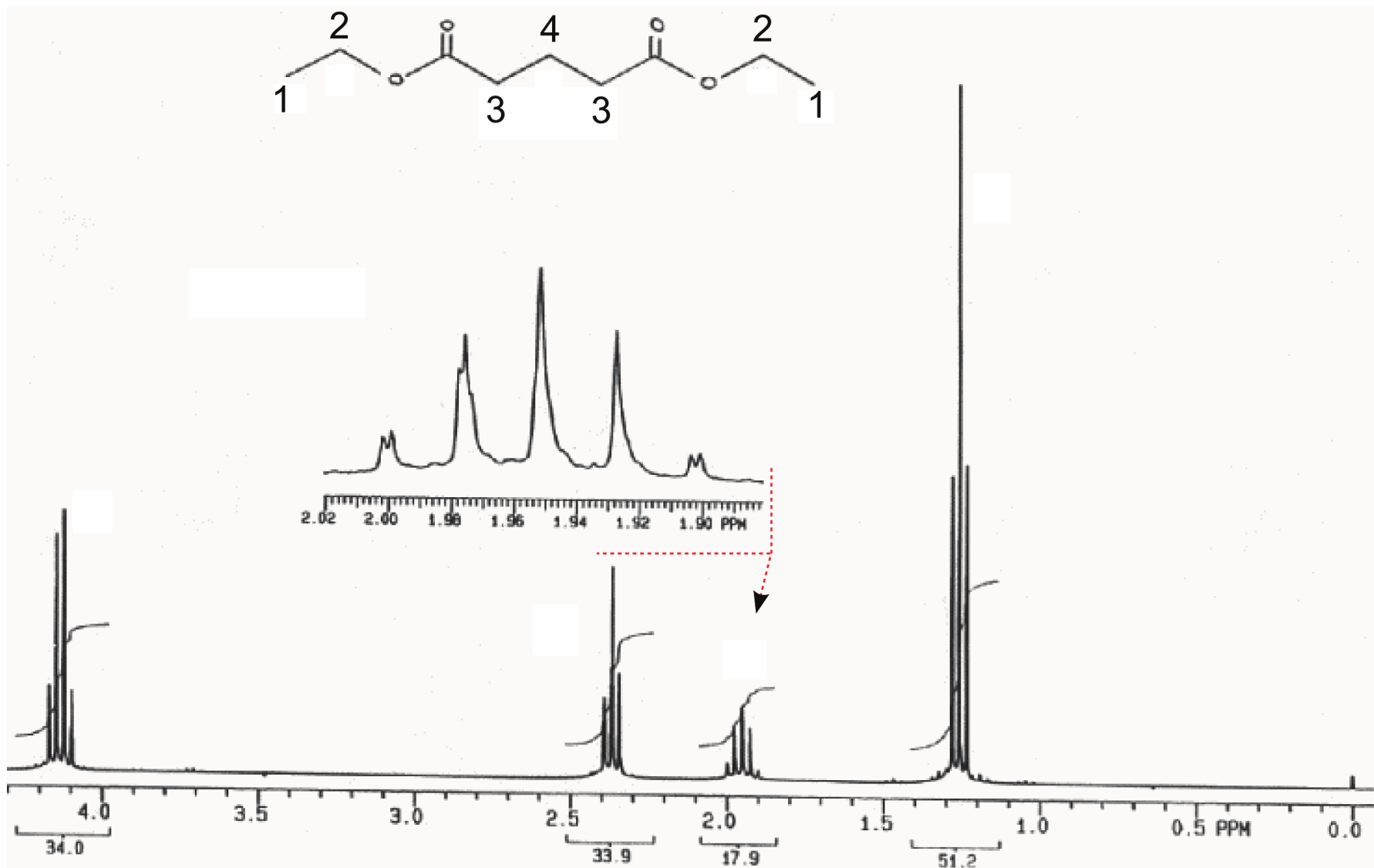
# Draw approximate 1D $^1\text{H}$ NMR spectrum of the following compound



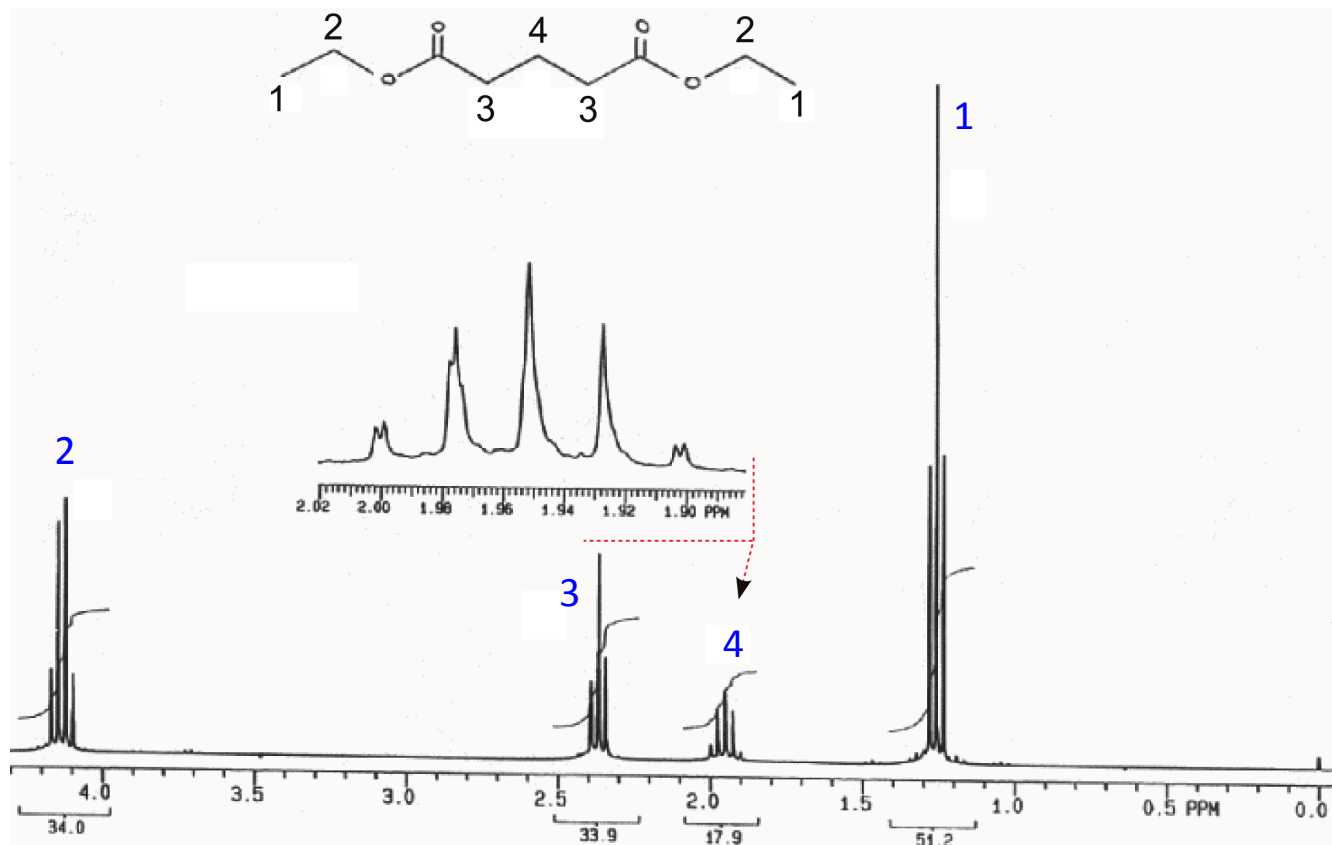
Notes:

- ▶ **H-2** - *meta*- and *ortho*- interaction with H-1 a H-3 - doublet of doublets
- ▶ **H-1** - only *meta*- interaction with H-2 - smaller coupling than **H-3** in *ortho*-interaction with H-2
- ▶ chemical shifts are result of overall effects of substituents on the aromatic ring

# 1D $^1\text{H}$ NMR spectrum of ethyl glutarate



# 1D $^1\text{H}$ spectrum of ethyl glutarate

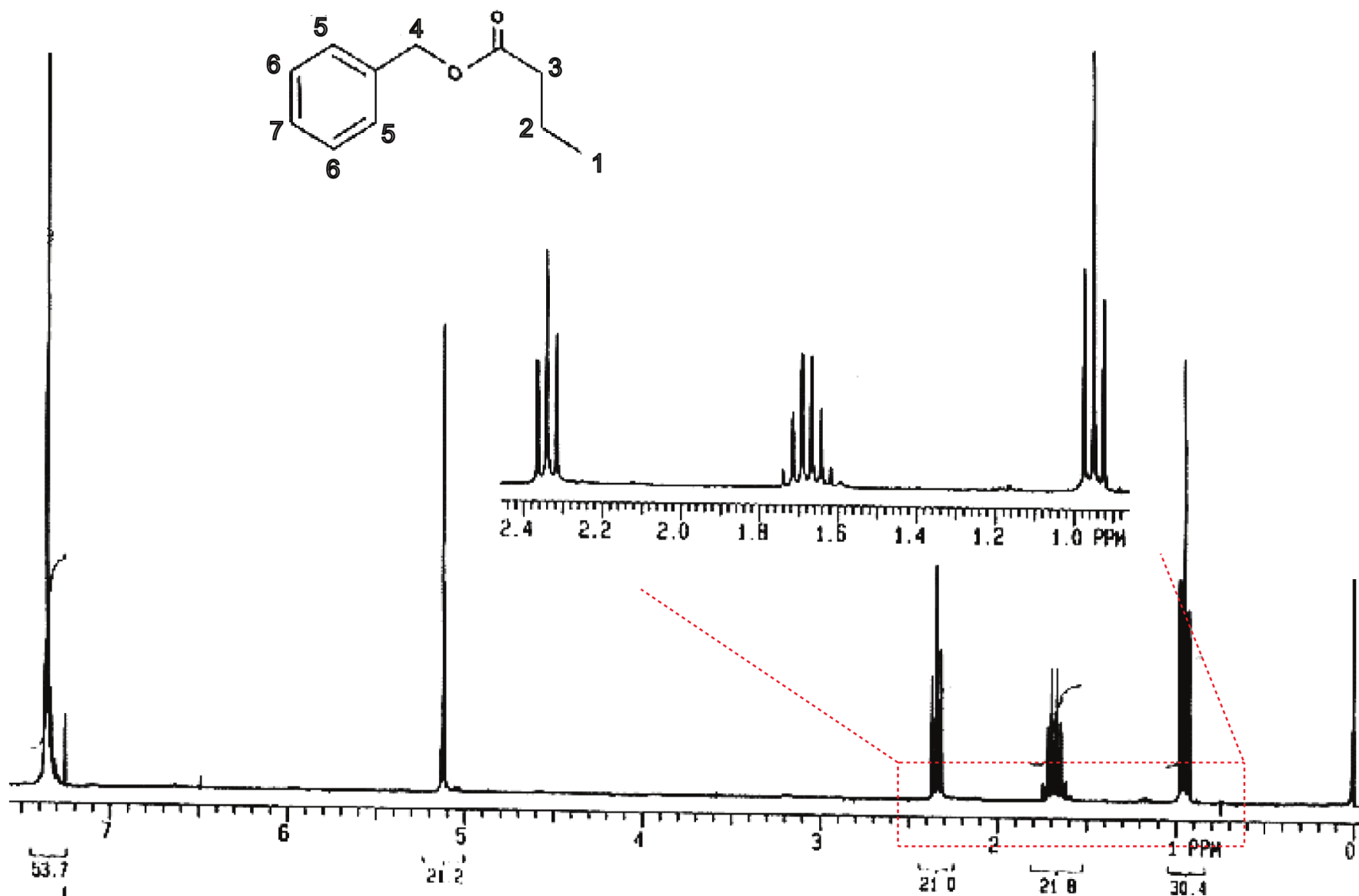


Notes:

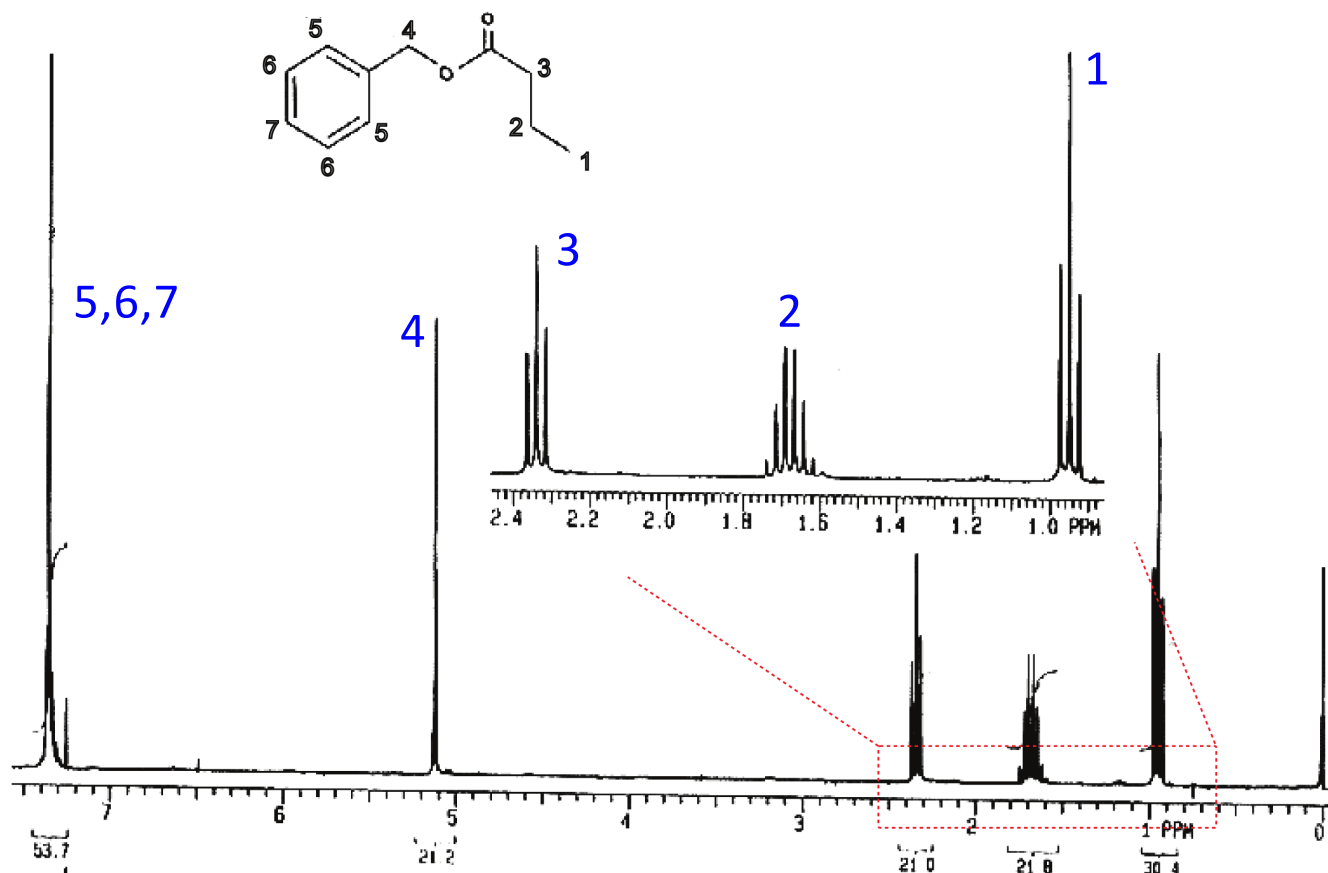
- ▶ symmetrical molecule - equivalent groups will give rise to only one signal
- ▶ splitting corresponds to  $n+1$  rule
- ▶ signal integrals correspond to number of protons in groups  
H-1:H-2:H-3:H-4 in the ratio 6:4:4:2
- ▶ chemical shifts depend on chemical environment of atoms



# 1D $^1\text{H}$ NMR spectrum of benzyl butyrate



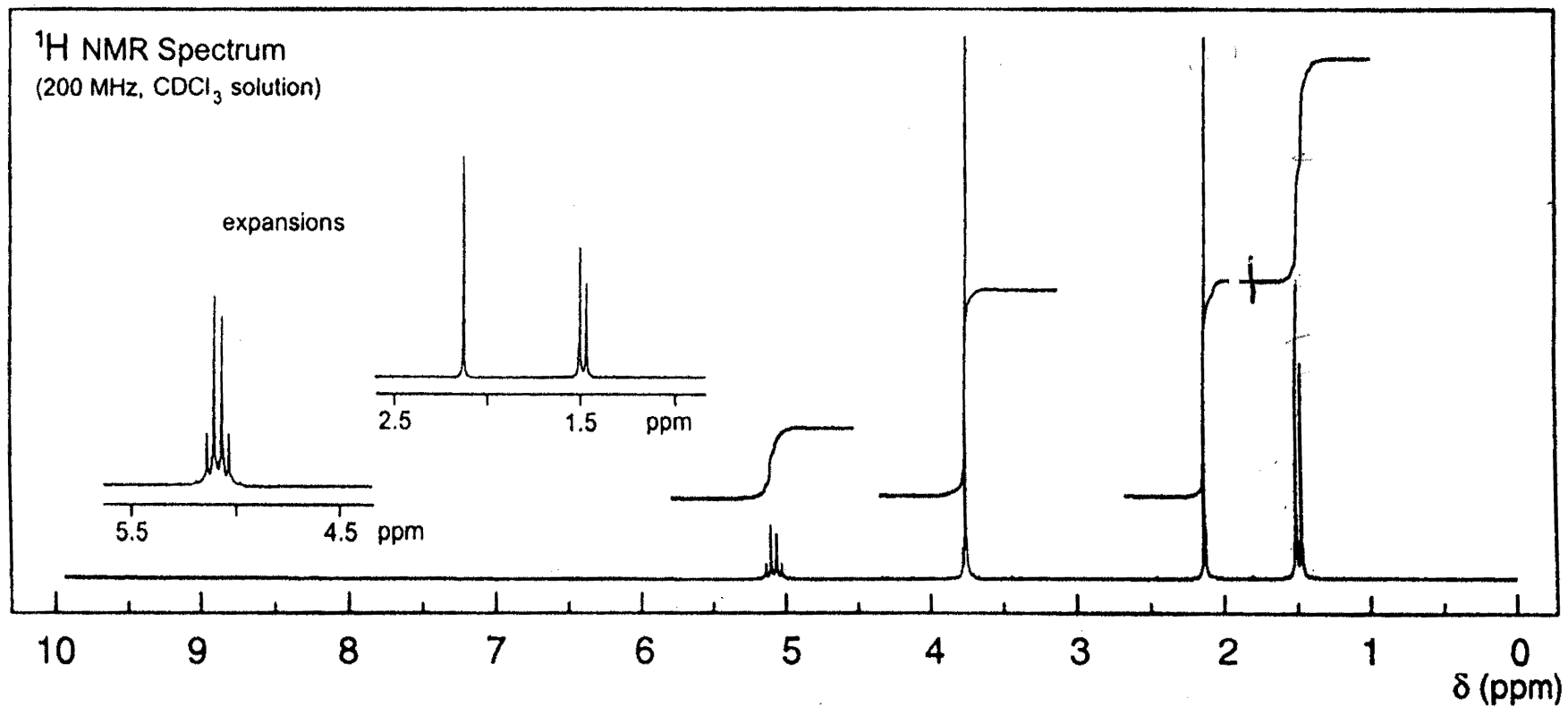
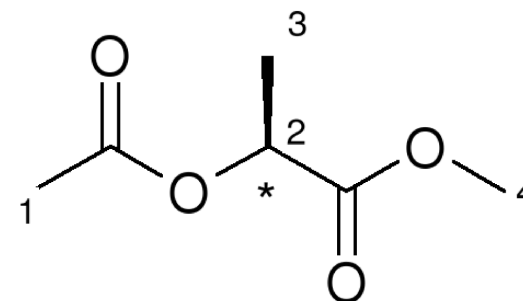
# 1D $^1\text{H}$ NMR spektrum benzylbutyrátu



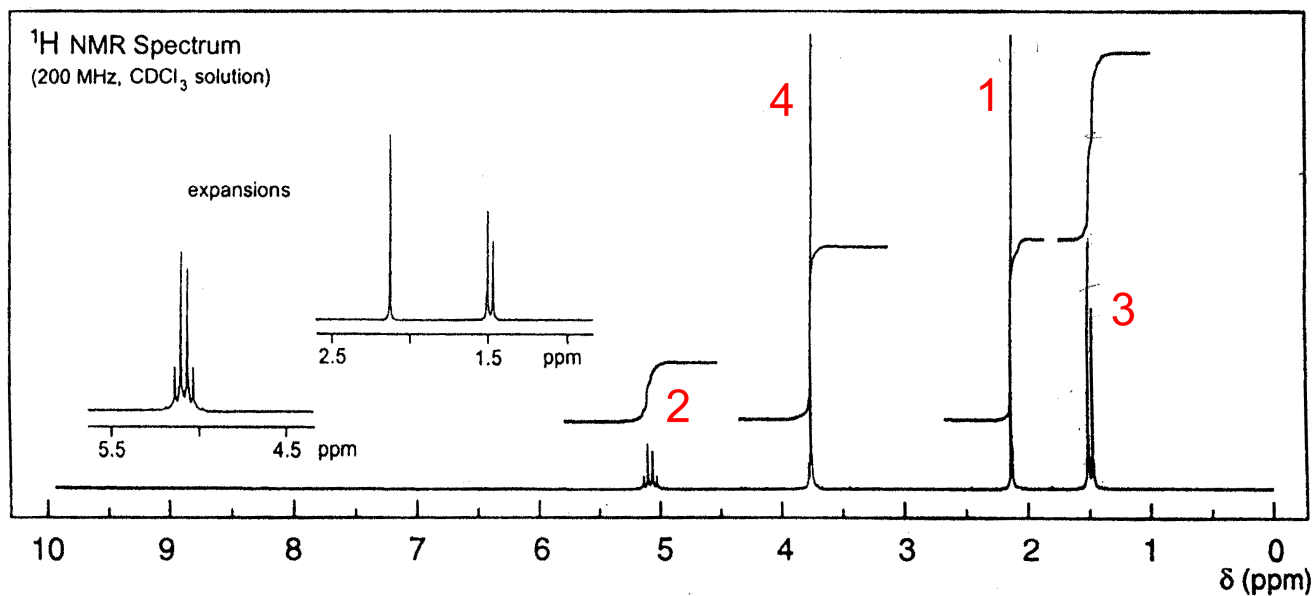
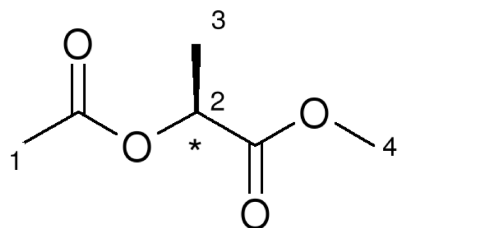
## Notes

- ▶ highest shift - aromatic ring, which is flexible - **H-5, H-6, H-7** are under one signal, integral value corresponds to five protons
- ▶ lowest shifts - alifatic chain H-1, H-2 a H-3: **H-1** - integral equals to three protons, splitted only by H-2 to triplet; **H-2** - integral equals to two protons, splitted by both H-1 and H-3 to triplet of quartets, which is due to similar  $J$ -coupling values fused into sextet; **H-3** - integral equals to two protons, splitted by H-2 to triplet
- ▶ **H-4** - integral equals to two protons, isolated signal - singlet, highest shift among all alifatic protons due to neighboring carboxyl and aromatic ring

# 1D $^1\text{H}$ NMR - methyl 2-acetoxy propanoate

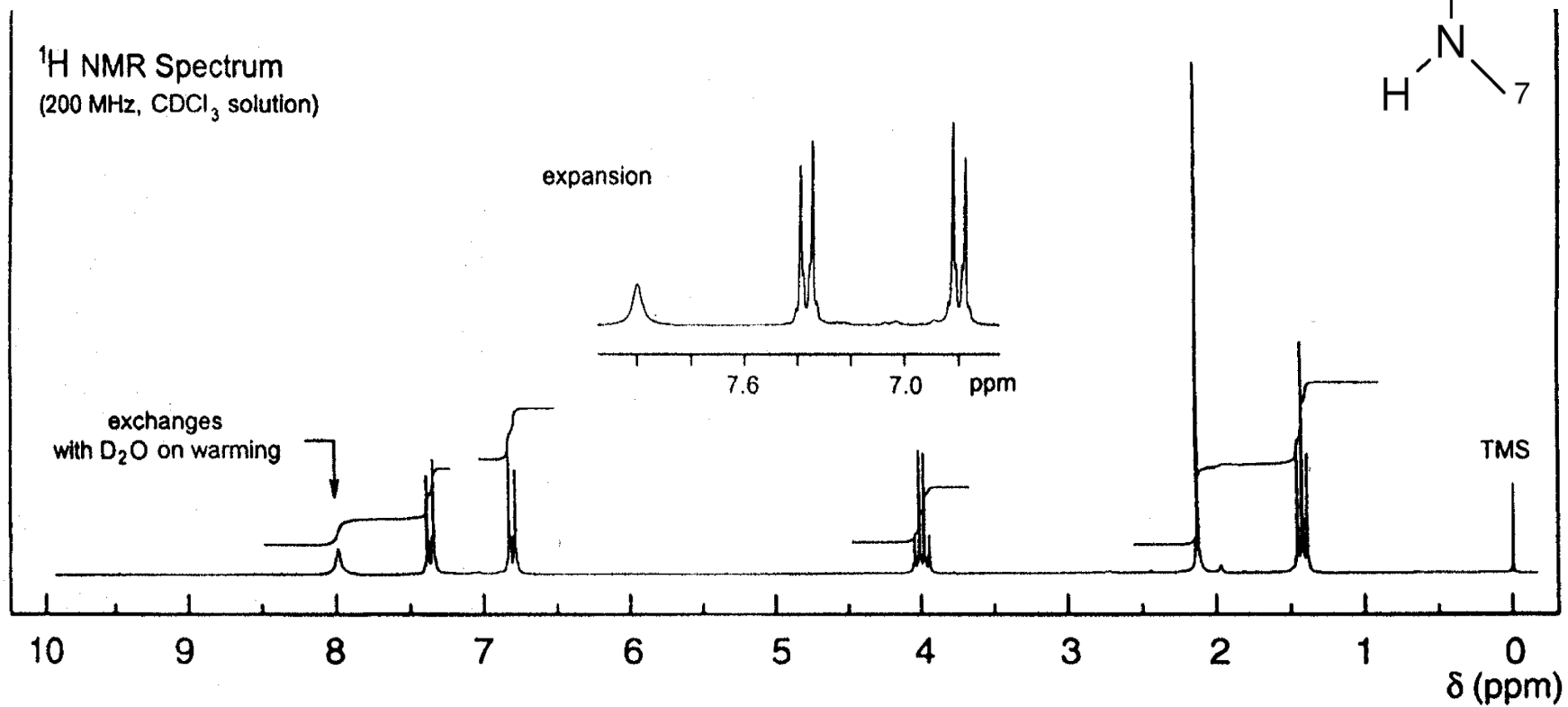
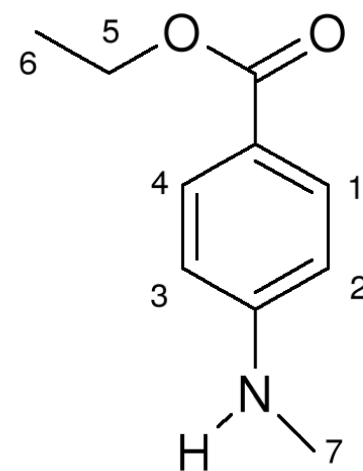


# 1D <sup>1</sup>H NMR - metyl 2-acetoxypropanoát

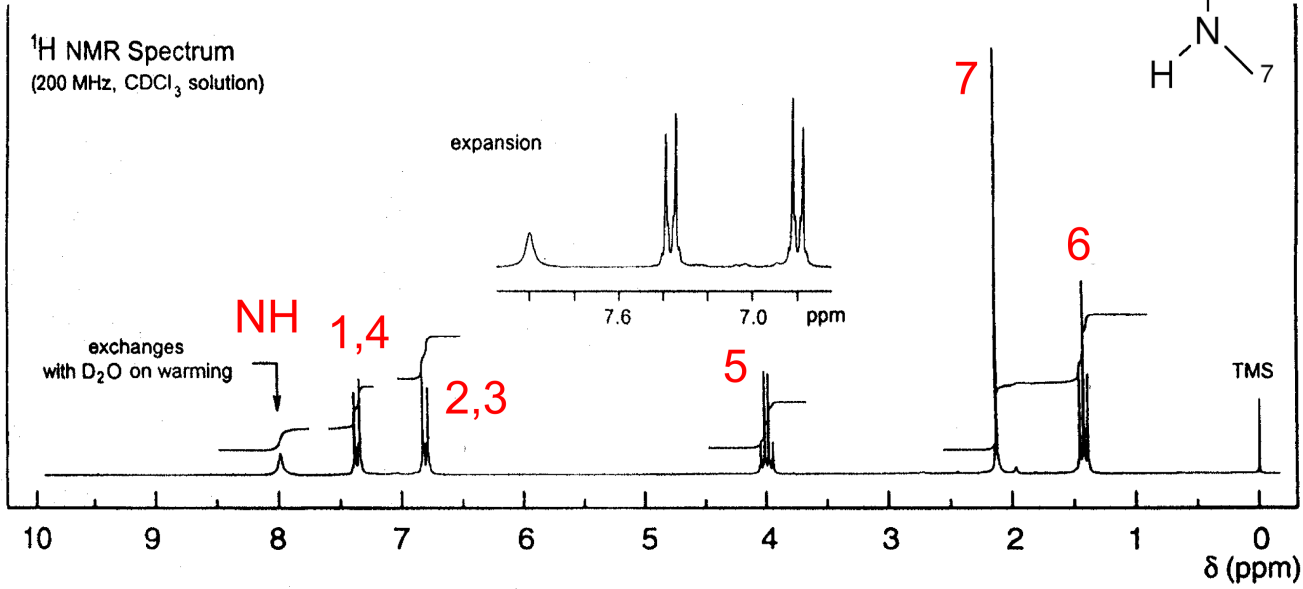
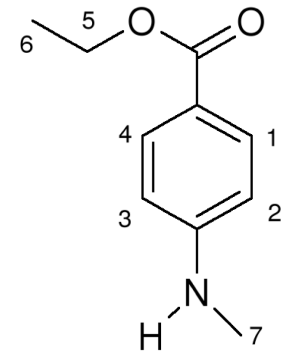


- ▶ lowest shift - methyl **H-3**, splitted due to neighboring stereogenic center C-2
- ▶ last splitted signal - **H-2** - splitted by H-3 to quartet, highest shift due to neighboring carboxyl groups
- ▶ two singlets - highest shift **H-4** next to carboxylic oxygen; lowest shift - **H-1** next to carboxylic carbon
- ▶ integrals corresponds to number of protons

# 1D $^1\text{H}$ NMR - ethyl 4-(methylamino)benzoate



# 1D <sup>1</sup>H NMR - etyl 4-(methylamino)benzoát



## Notes:

- ▶ highest shift - **NH**, least intensive and broad signal because there is dynamical exchange of the proton with the solvent, it is visible in the spectrum thanks to CDCl<sub>3</sub> used as solvent
- ▶ lowest shifts - alifatics - signal at 1.5 ppm splitted to triplet - methyl **H-6** splitted by H-5; singlet around 2 ppm - methyl **H-7** - isolated, highest shift than H-6 thanks to nitrogen; quartet at 4 ppm - **H-5** - splitted by methyl H-6, highest shift thanks to oxygen
- ▶ two doublets in aromatic region around 7 ppm - **H-1,4** a **H-2,3** - symmetrical, shifts are resulting from effects of both substituents on aromatic ring

# 1D $^1\text{H}$ NMR spectrum of cartilagineal

