Nuclear magnetic resonance spectroscopy

II. ¹³C NMR

Reading: Pavia Chapter 6.1-6.5, 6.7, 6.11, 6.13

1. General

- more/better/additional structural information for larger compounds
- problems:
 - a) C isotopes and their natural abundance

¹H 99.95% ¹²C 98.9% ¹³C 1.1% nuclear spin I 0 $\frac{1}{2}$

b) gyromagnetic ratio

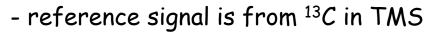
 \Rightarrow for B_0 1.41 T, $\omega(^1\text{H})$ 60 MHz $\omega(^{13}\text{C})$ 15 MHz

 \Rightarrow much lower E needed for transition

- \Rightarrow even less nuclei in excess in the lower level
- \Rightarrow smaller signal

- solutions: sum up more scans, use more sample, use a stronger field

2. Chemical shift

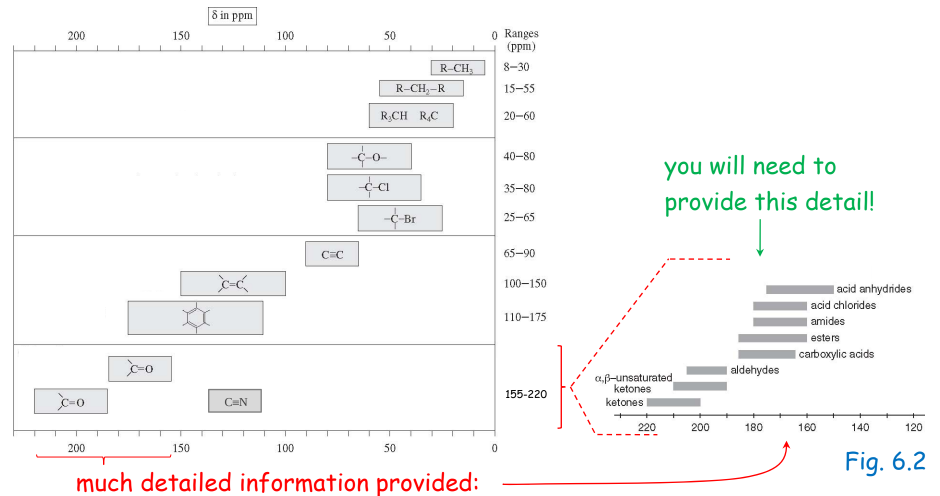


- δ range is \approx 200 ppm: advantage: - signals are spread out more

- more detailed information possible



Table 6.1



3. Estimation of $\delta^{13}C$

- from increment systems
 - for alkanes

 $\delta^{13}C$ (ppm) = -2.3 + increments for α , β ... C atoms and steric corrections see p. A-24 $C - C_{c} - C_{d} - C_{a} = -2.3 + 9.1 + 9.4 - 3.2.5 = 8.7 \text{ ppm (exp. 8.9 ppm) ok}$ $\delta^{13}C_{b} = -2.3 + 9.1 + 3.9.4 - 2.5 = 32.5 \text{ ppm}$ $\alpha \quad \beta \quad \gamma \quad -3.4 \text{ steric: 1°, attached to 4°}$ = 29.1 ppm (exp. 29.1 ppm) ok $\delta^{13}C_{c} = -2.3 + 4.9.1 + 9.4 = 43.5 \text{ ppm}$ α β - 3.1.5 - 8.4 steric: 4°, attached to 3.1° and 2° = 30.6 ppm (exp. 30.6 ppm) ok $\delta^{13}C_{d} = -2.3 + 2.9.1 + 3.9.4 = 44.1 \text{ ppm}$ α β - 7.5 steric: 2°, attached to 4° (and 1°) = 36.6 ppm (exp. 36.9 ppm) ok other substituents alkenes aromatic compounds

3. Estimation of $\delta^{13}C$ continued

- from increment systems
 - for substituted alkanes
 - for alkenes

see Table A8.3, add to values from Table A8.2!

(i.e., calculate from scratch!)

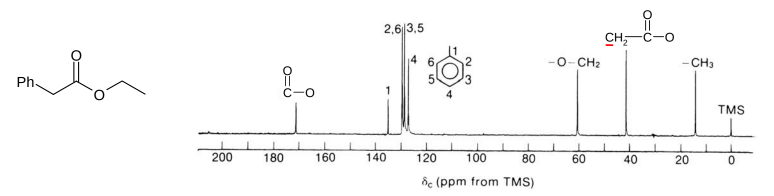
see Tables A8.5 and A8.6

- for substituted benzenes see Table A8.7

Work through these tables and examples on your own! You're (probably) ok if you can do the following two examples. Do not look at the answers before you have tried them!

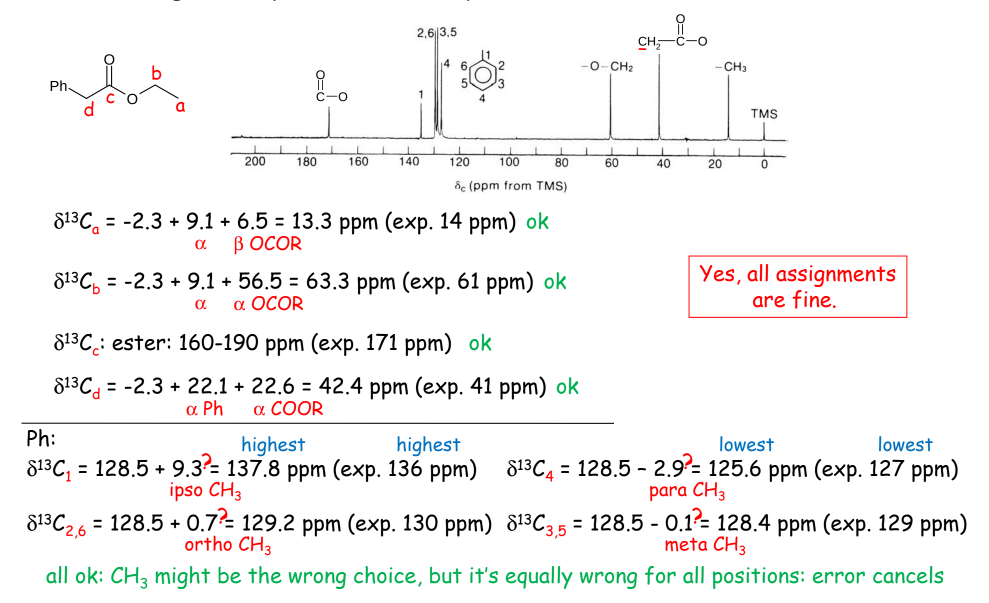
Example 1

Are the assignments provided on the spectrum correct? (Calculate/estimate $\delta^{13}C$!)



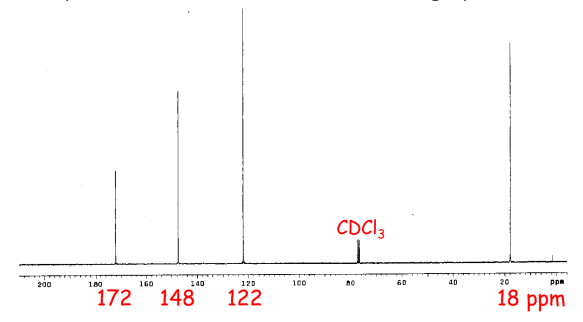
Example 1 answer

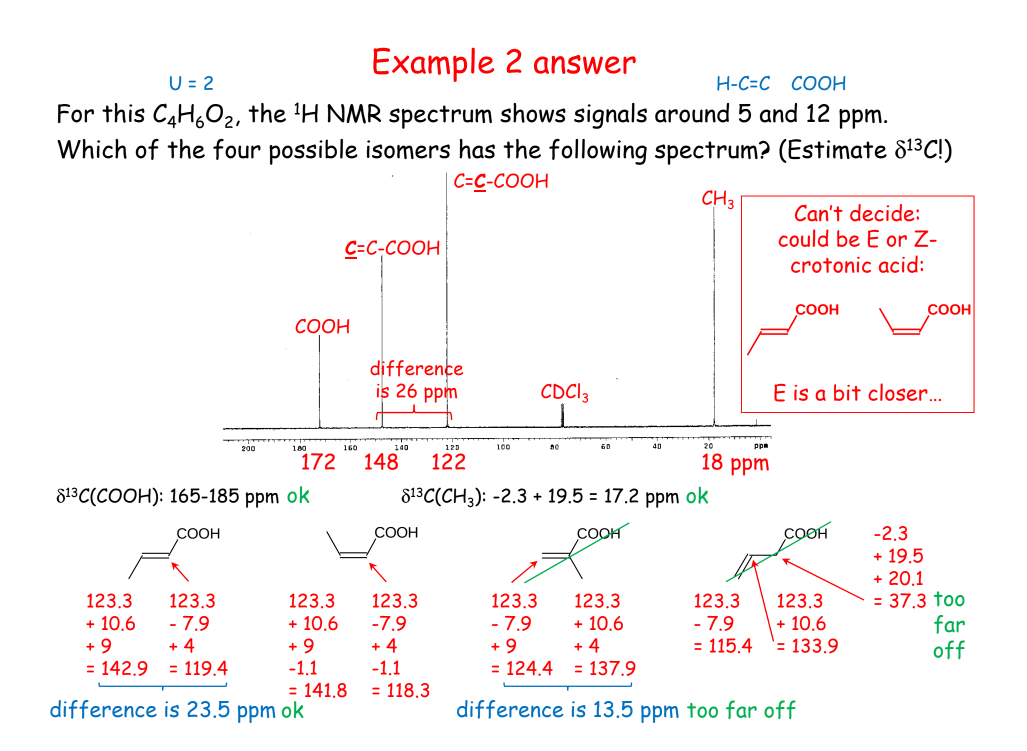
Are the assignments provided on the spectrum correct? (Calculate/estimate $\delta^{13}C$!)



Example 2

For this $C_4H_6O_2$, the ¹H NMR spectrum shows signals around 5 and 12 ppm. Which of the four possible isomers has the following spectrum? (Estimate $\delta^{13}C$!)





4. Spin-spin coupling

- as in ¹H NMR, but in principle two situations:

- ${}^{13}C$ coupling with ${}^{13}C$

 \Rightarrow rare event: probability of 2 ¹³C next to each other is very low

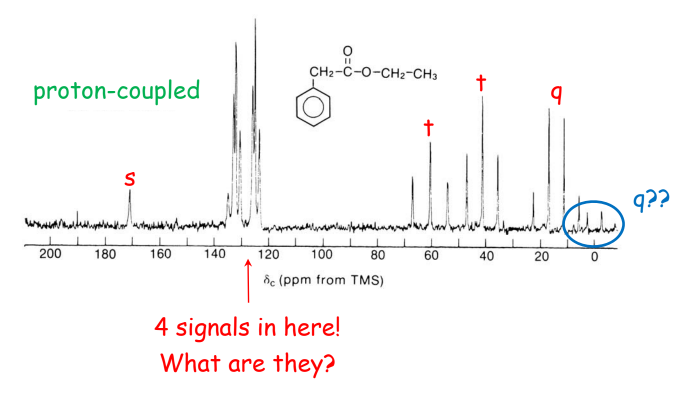
- $^{13}\mathrm{C}$ coupling with $^{1}\mathrm{H}$
 - \Rightarrow "heteronuclear" coupling
 - \Rightarrow follows the n+1 rule
 - \Rightarrow ¹J coupling:

$$-CH_3$$
 ¹³C signal split by ¹H into q

no proton: signal is not split, **s** (quaternary $C, R_2C=O, R-C=N...$)

4. Spin-spin coupling continued

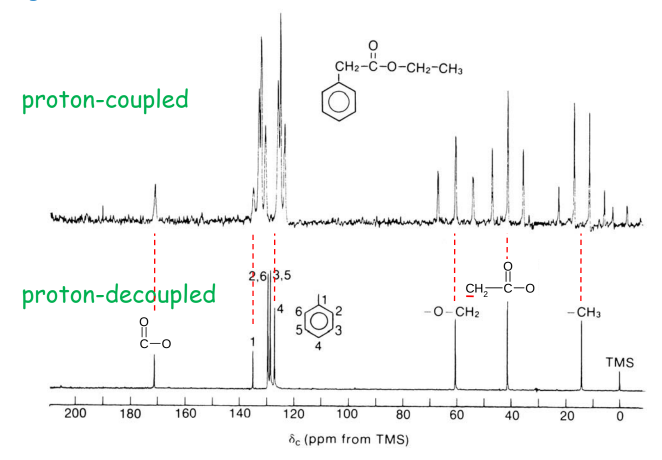
- ¹³C spectra that show ¹J(¹³C-¹H) coupling are called "proton-coupled" or "non-decoupled"



 \Rightarrow not often recorded, because of frequent overlap between multiplets, especially if the molecule contains many (sp³) C atoms

4. Spin-spin coupling continued

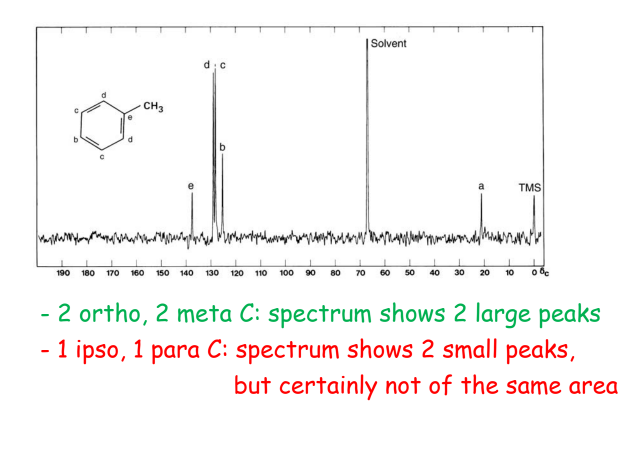
- ¹³C spectra that *do not* show ¹J(¹³C-¹H) coupling are called "proton-decoupled" \Rightarrow only singlets are observed

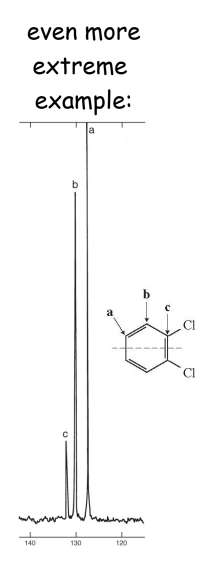


- advantage: simpler spectra
- disadvantage: important information on C-H connectivity is lost

5. Integration

- ¹³C spectra are not usually integrated





5. Integration continued

- two problems are the reason:

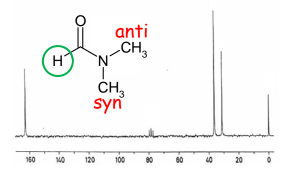
a) NOE (nuclear Overhauser enhancement)

- in proton-decoupled spectra
- intensity of a signal increases upon decoupling
 - signal for CH_3 grows most
 - signal for CH₂ grows less ...
 - C=O, ipso-C or quaternary C signal ends up being tiny
- reason lies in the decoupling experiment

Example application

NOE usage for peak assignment

- NOE also works "through space"





http://131.104.156.23/Lectures/CHEM_207/CHM_207_NMR.htm

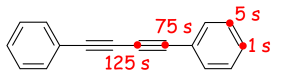
5. Integration continued

- two problems are the reason:

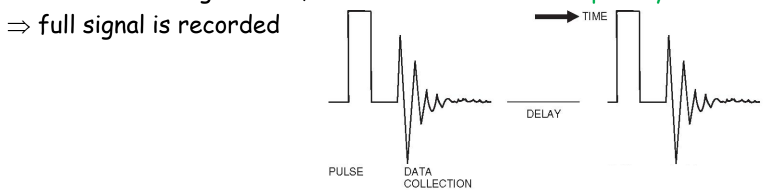
b) relaxation of a nucleus

- FT-NMR records the FID signal
 - \Rightarrow follows the excited nucleus as it relaxes to the ground state de-excitation is fast for ¹H

- variable for $^{13}\mbox{C}$, depending on the environment



- for a fast-relaxing nucleus, the FID is collected completely



- for a slow-relaxing nucleus, the FID is collected incompletely
 - \Rightarrow only partial signal is recorded

Why don't we just wait longer between pulses???