

NMR-spectroscopy of proteins in solution

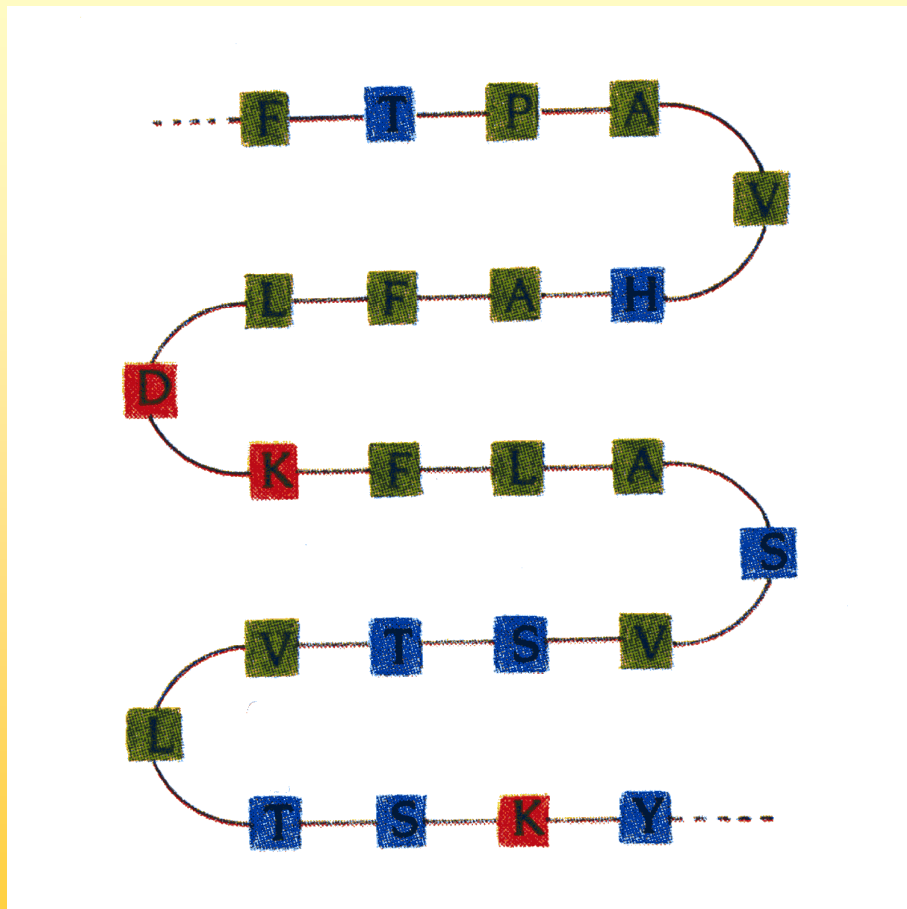
- spectra and assignment

Peter Schmieder



NMR-spectroscopy of proteins

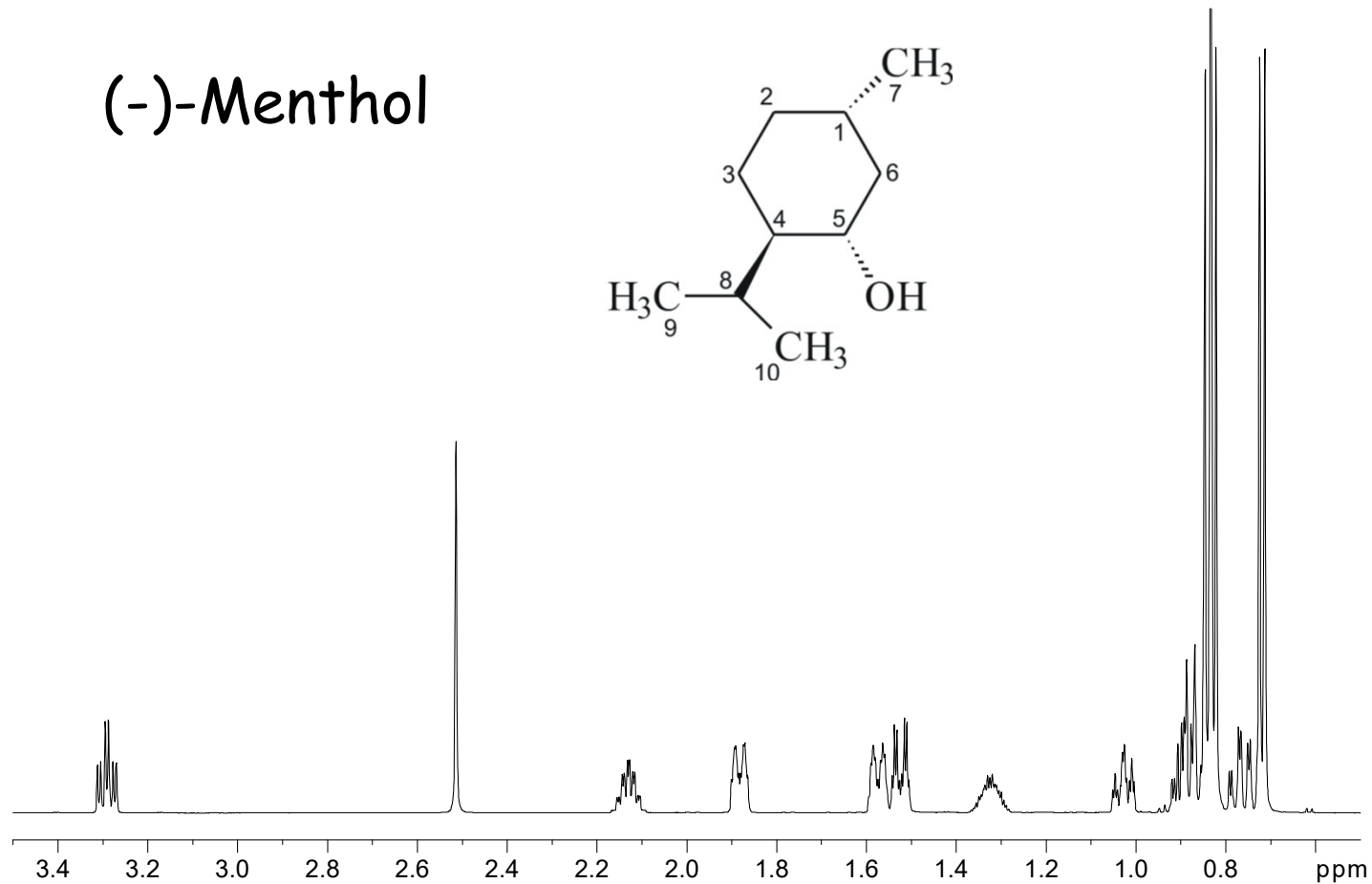
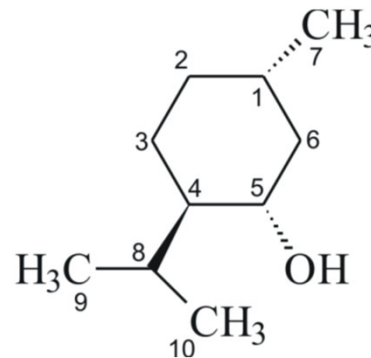
NMR-spectroscopy of proteins



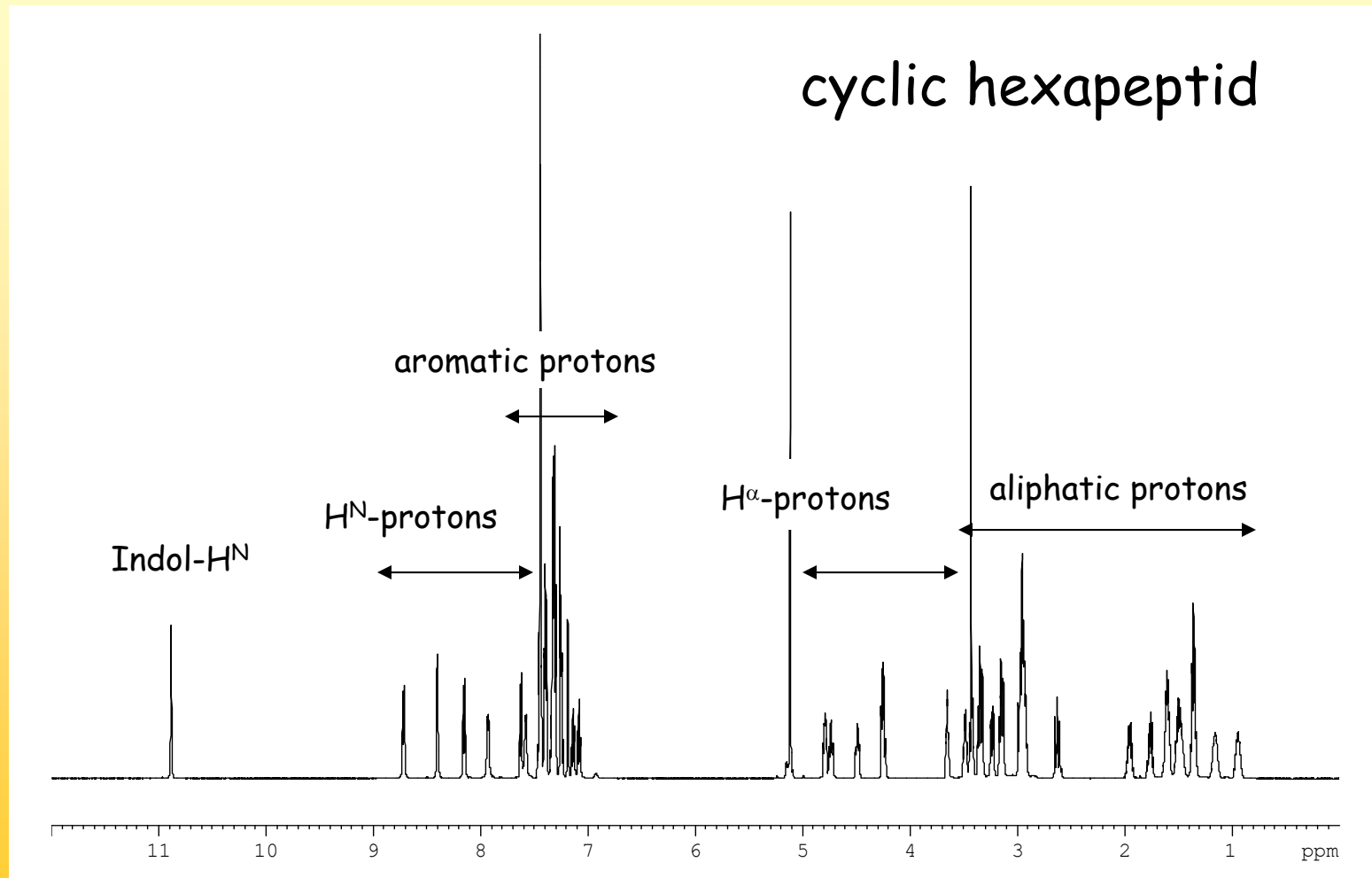
The main problem obtaining assignments of NMR-spectra of proteins result from the fact that a protein is a polymer, i.e. a repetition of identical units

NMR-spectroscopy of proteins

(-)-Menthol

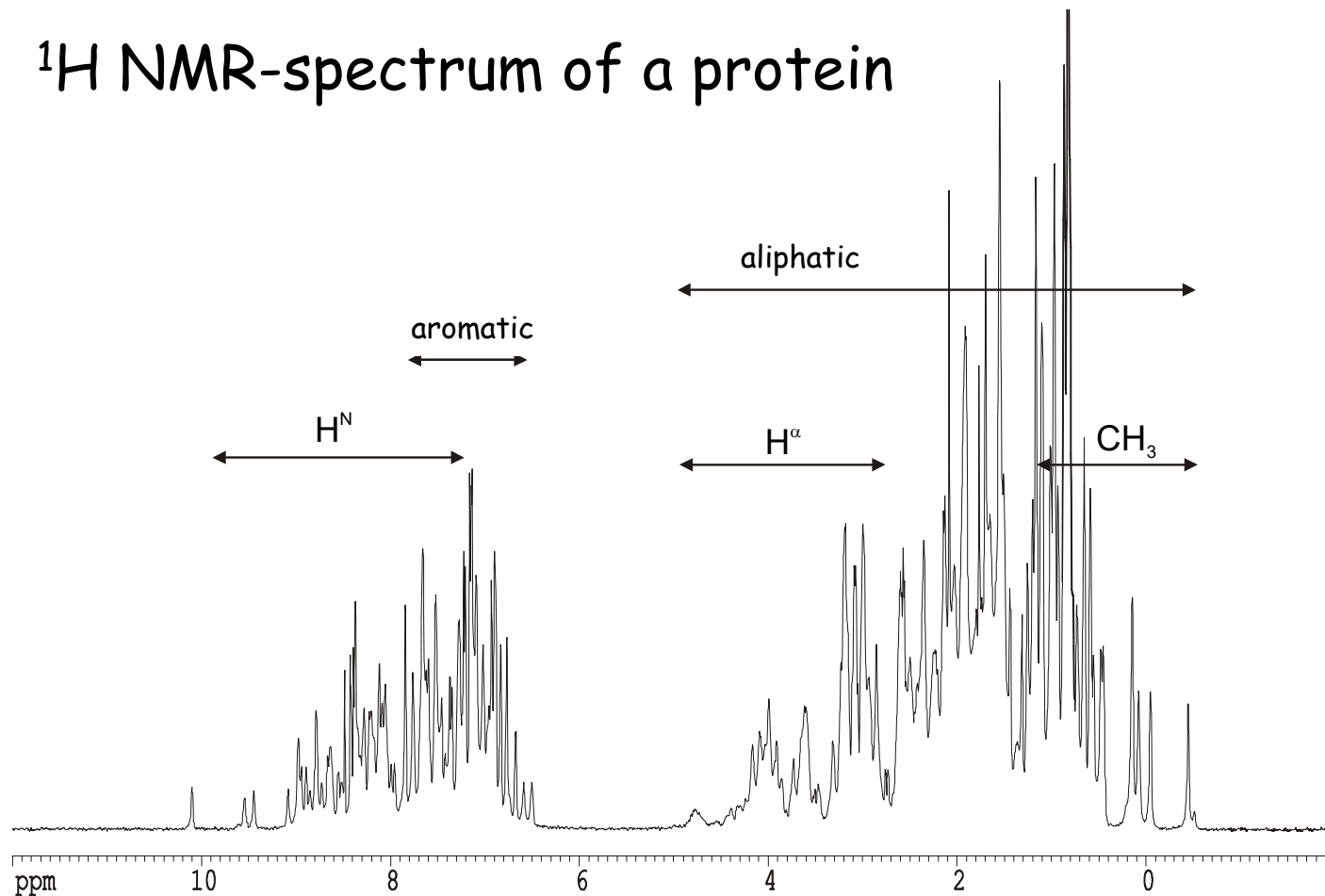


NMR-spectroscopy of proteins



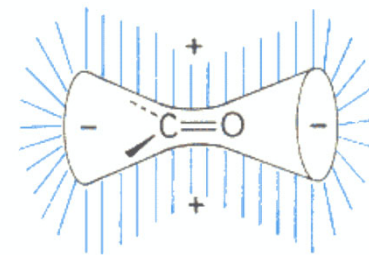
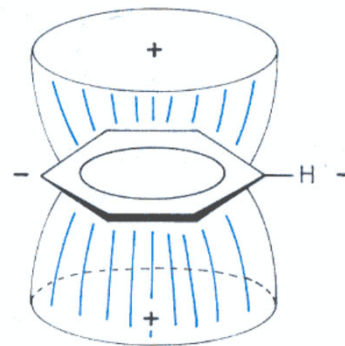
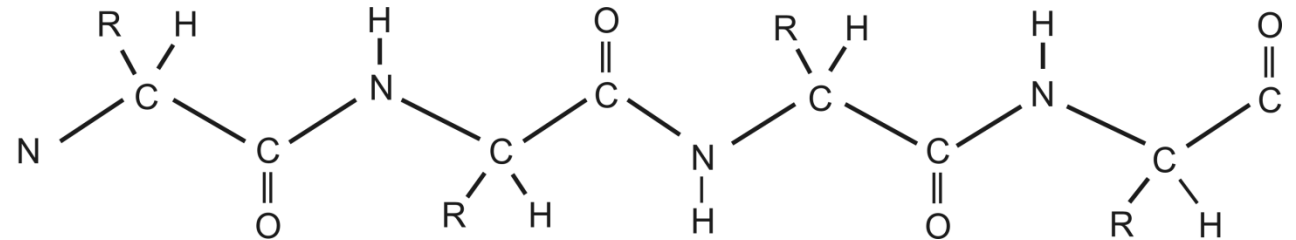
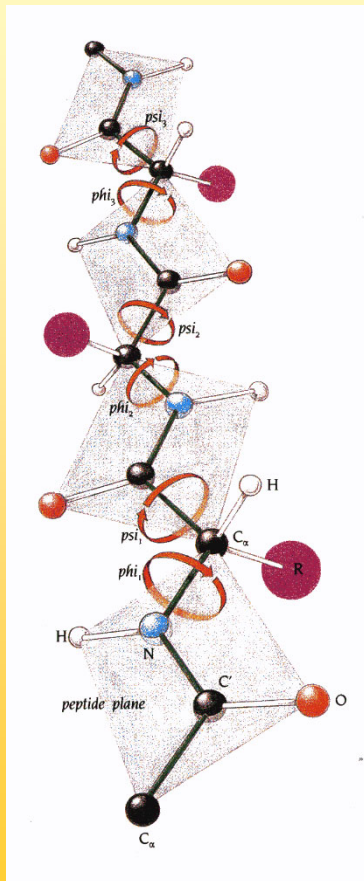
NMR-spectroscopy of proteins

^1H NMR-spectrum of a protein



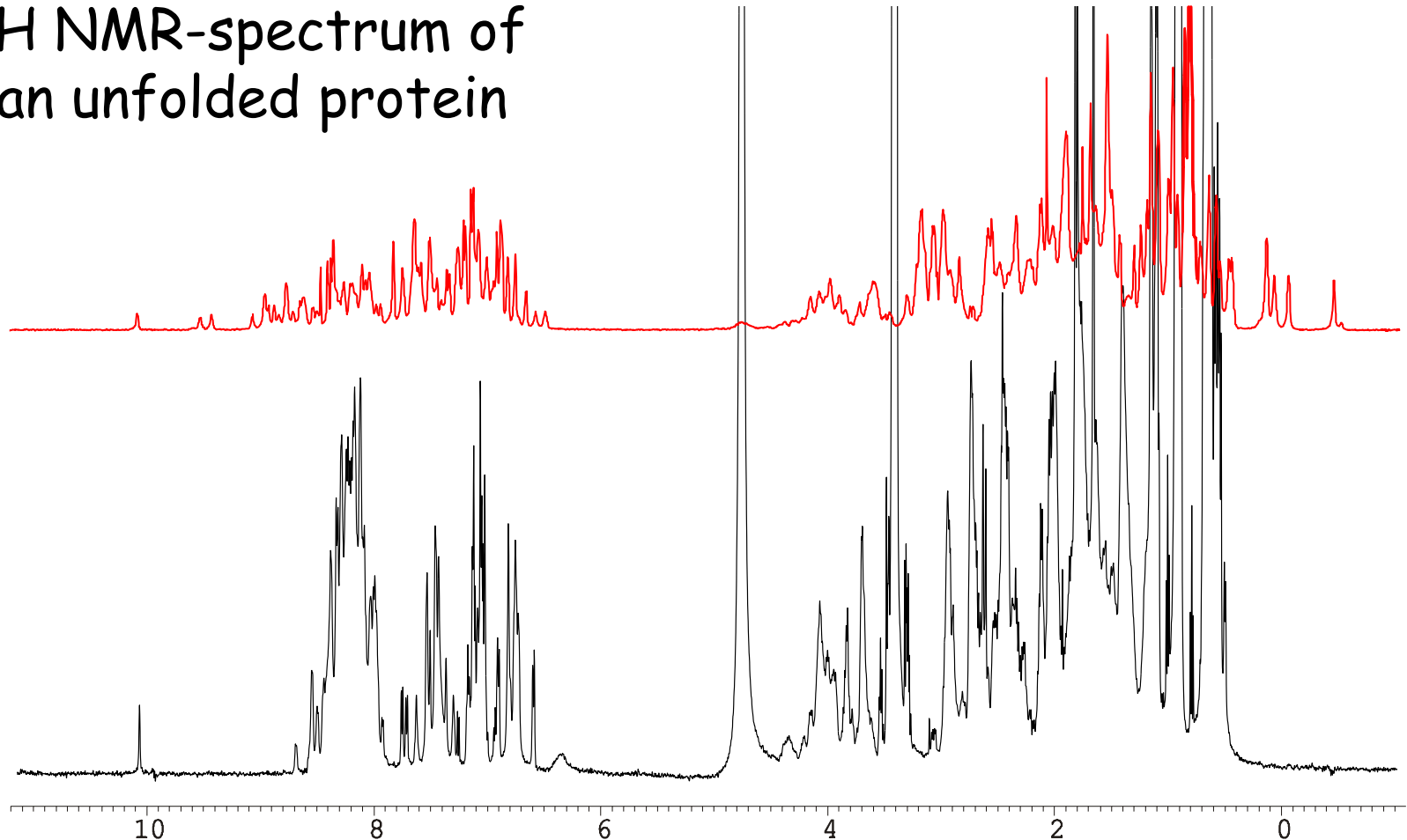
NMR-spectroscopy of proteins

Differences in the chemical shifts result from defined elements of secondary structure



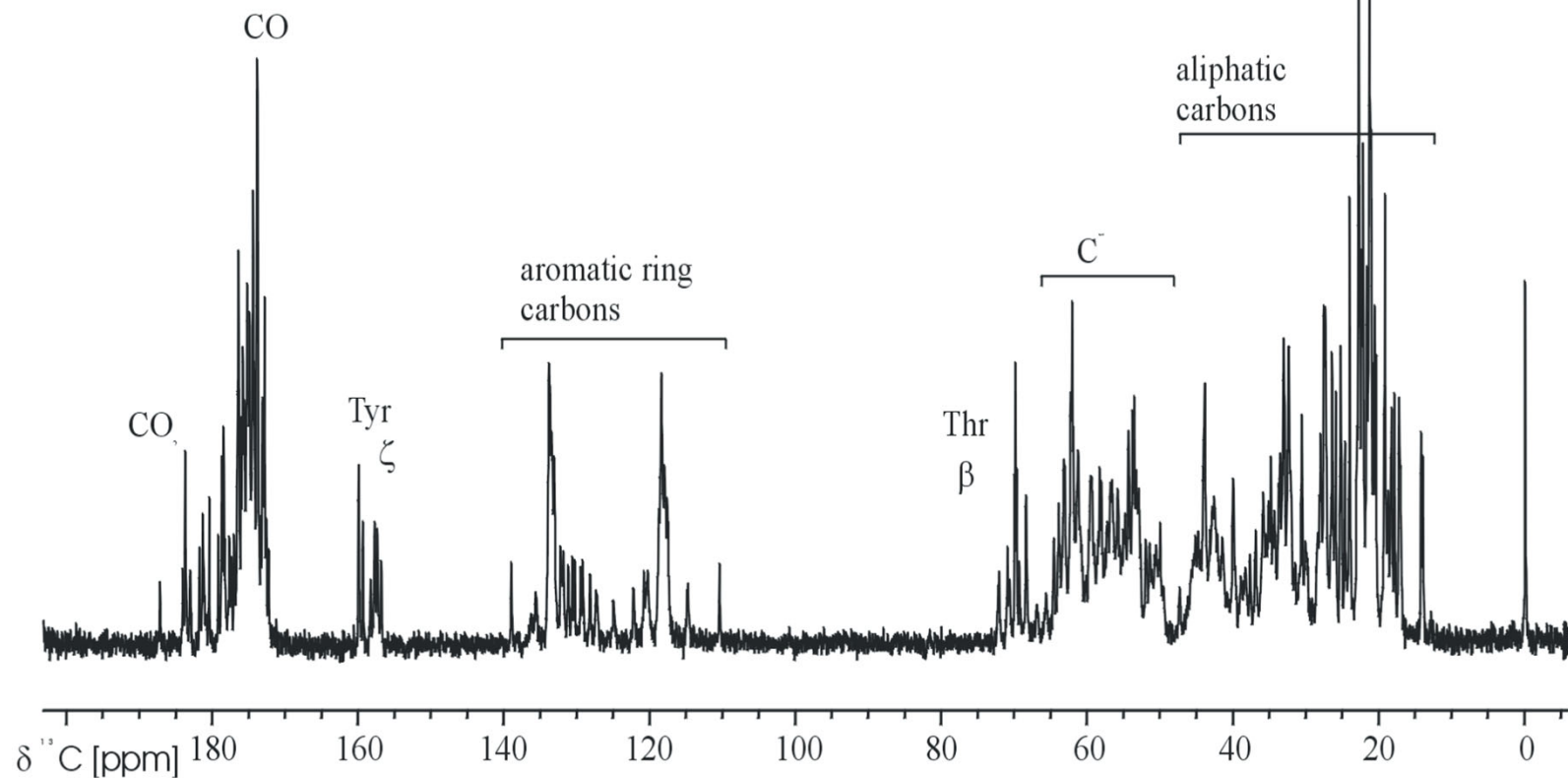
NMR-spectroscopy of proteins

^1H NMR-spectrum of
an unfolded protein



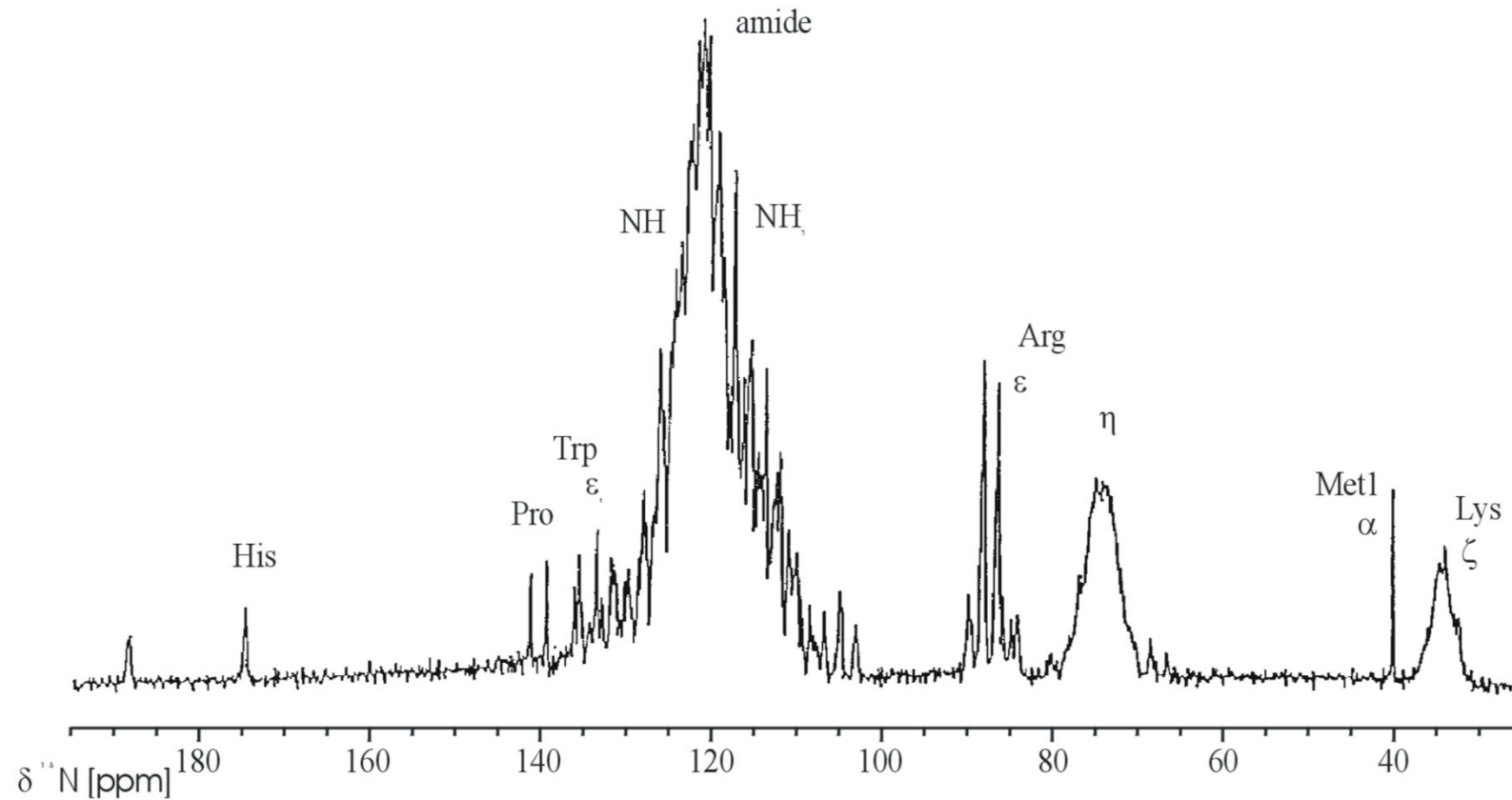
NMR-spectroscopy of proteins

^{13}C NMR-spectrum of a protein



NMR-spectroscopy of proteins

^{15}N NMR-spectrum of a protein



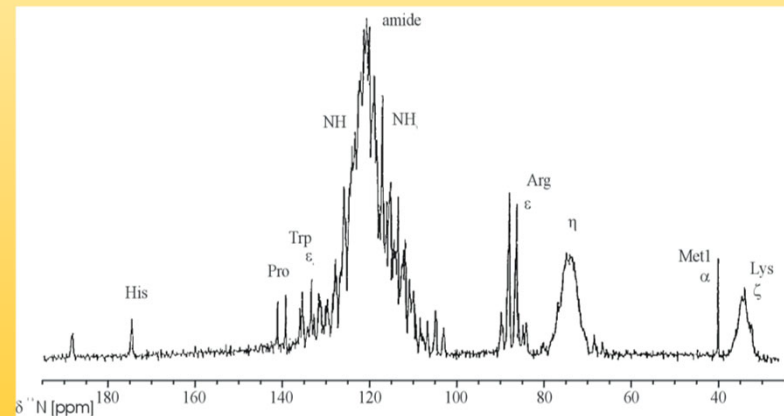
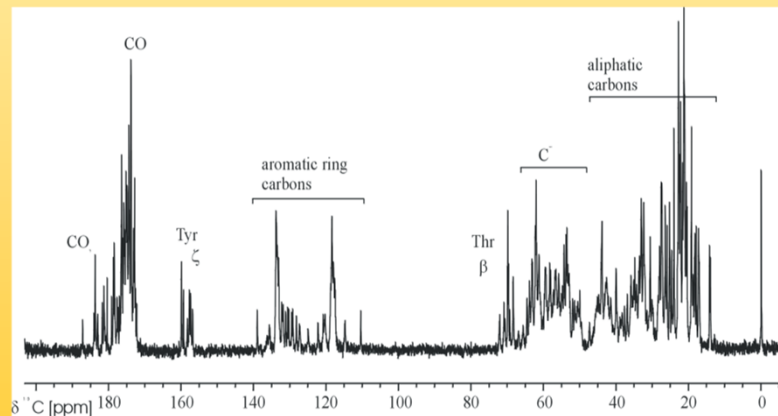
NMR-spectroscopy of proteins

NMR-spectra of proteins are thus not that much different from spectra of small molecules and show similar features (chemical shift, coupling etc.).

Because of the large number of nuclei contributing to the spectrum, the fact that they are polymers and their size, one-dimensional spectra are not sufficient and we need to use multidimensional ones.

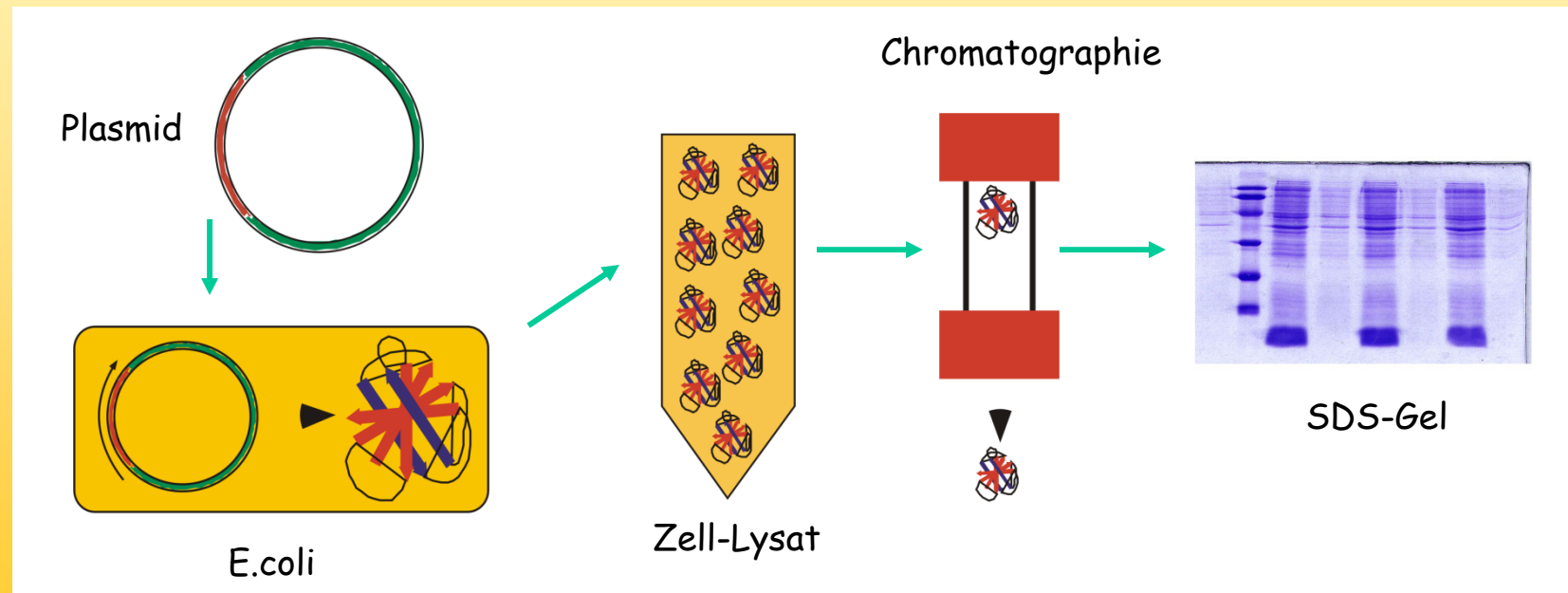
NMR-spectroscopy of proteins

In addition to the necessity to use multidimensional spectra the superior dispersion of the spectra of carbon and nitrogen make heteronuclear experiments the methods of choice for protein NMR spectroscopy:



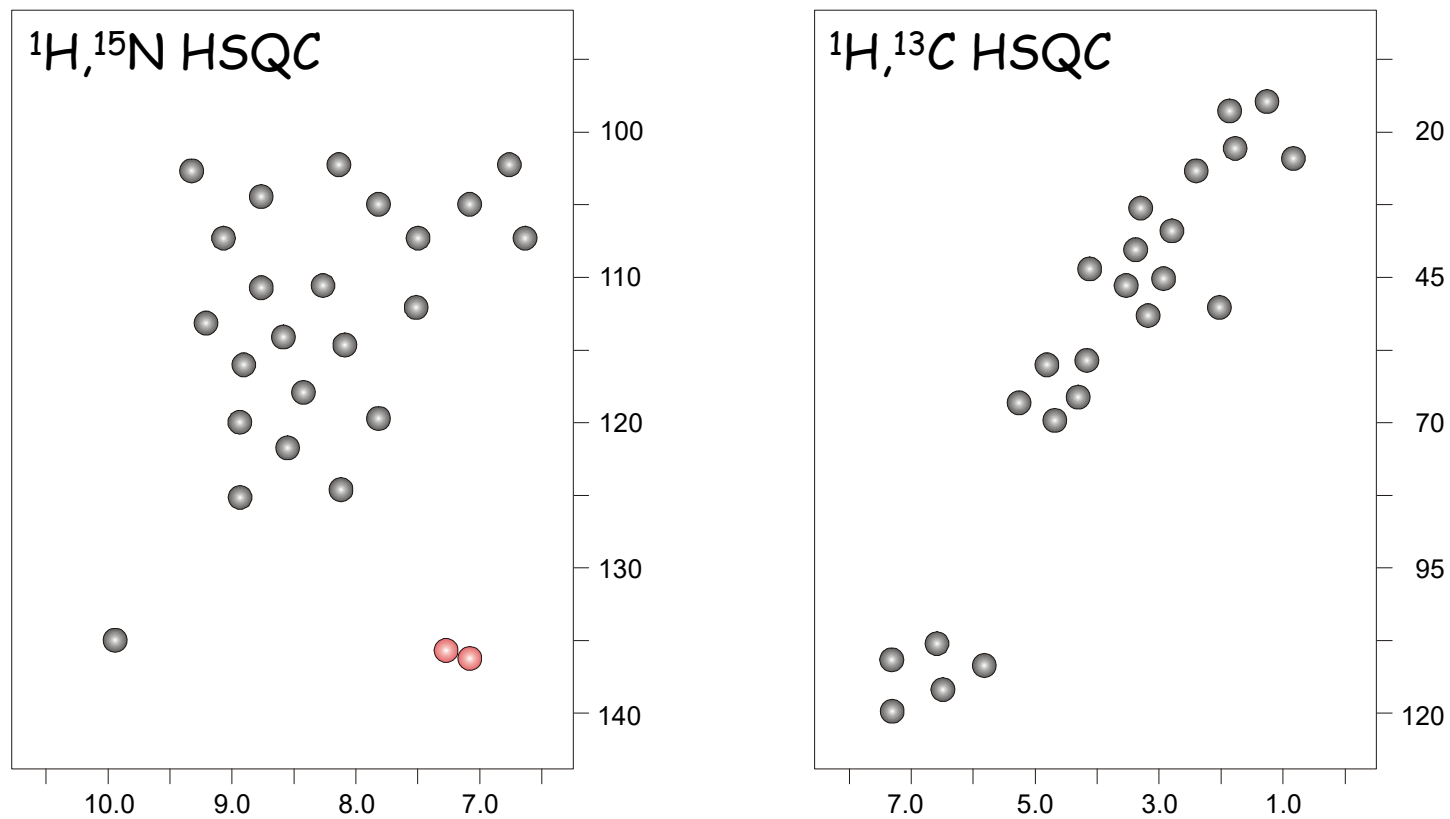
NMR-spectroscopy of proteins

Recording heteronuclear spectra using the heteronucleus in natural abundance ($^{13}\text{C}=1.1\%$ and $^{15}\text{N}=0.4\%$) is not realistic with proteins. The proteins need to be labeled when expressed.



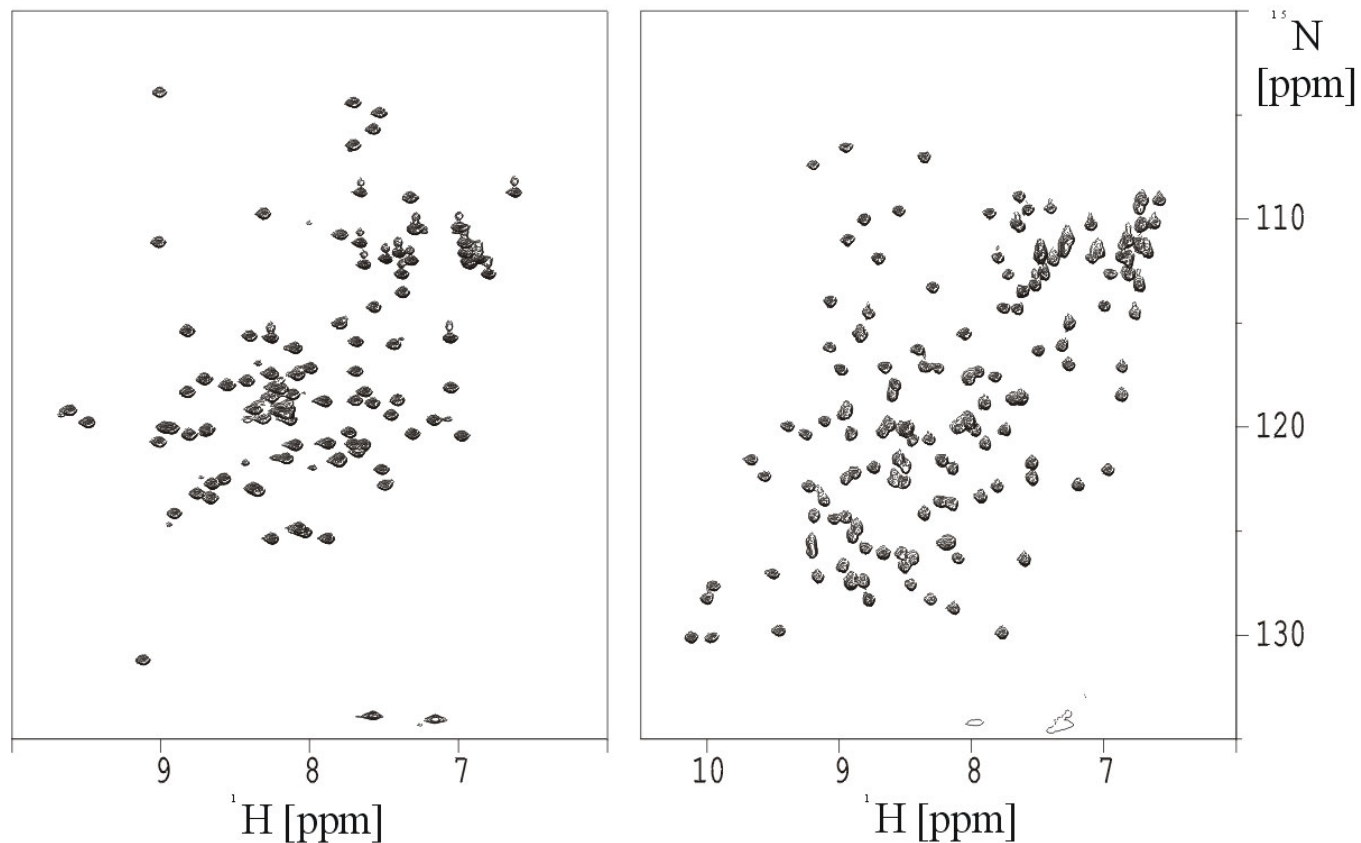
NMR-spectroscopy of proteins

$^1\text{H}, ^{15}\text{N}$ and $^1\text{H}, ^{13}\text{C}$ HSQC-spectra of proteins (schematic)



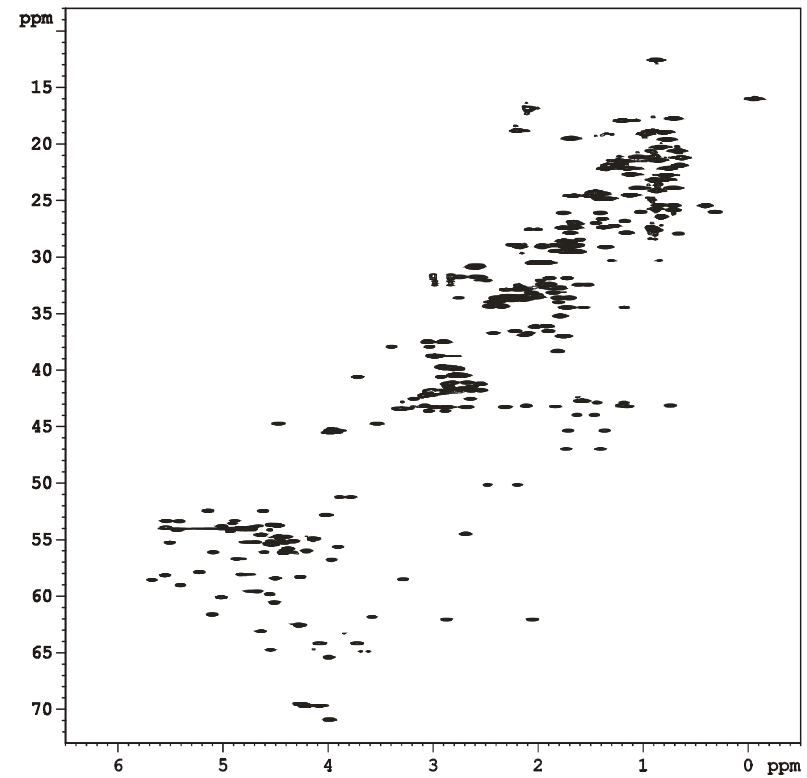
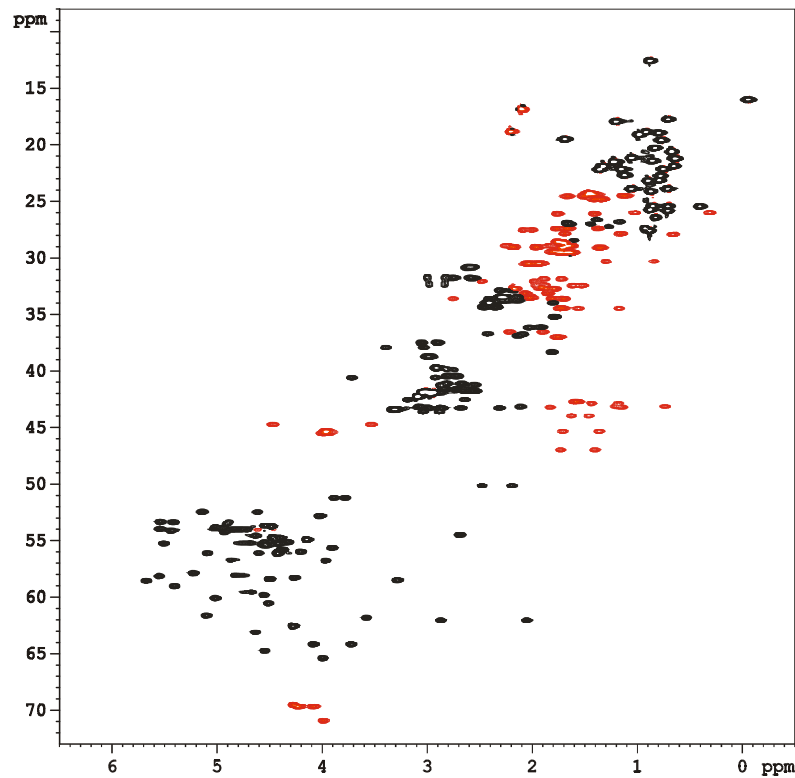
NMR-spectroscopy of proteins

^1H , ^{15}N HSQC-spectra are „fingerprints“ of proteins



NMR-spectroscopy of proteins

$^1\text{H}, ^{13}\text{C}$ HSQC-spectra of proteins

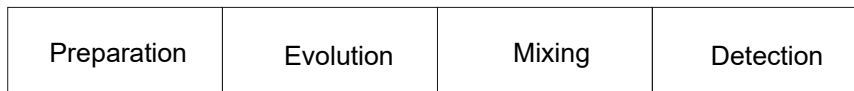


Multidimensional NMR-spectroscopy with more than two dimensions

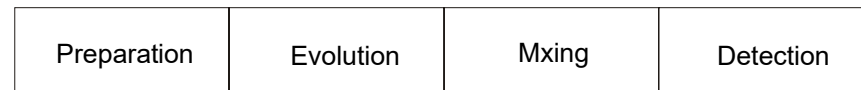
Multidimensional NMR-spectroscopy

Formally a 3D experiment is a combination of two
2D experiments

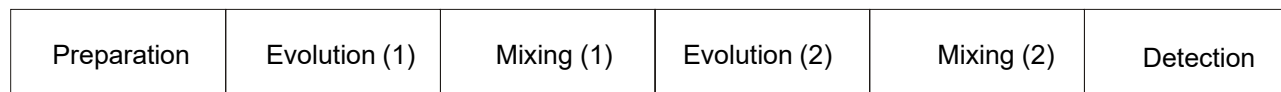
1. 2D-sequence



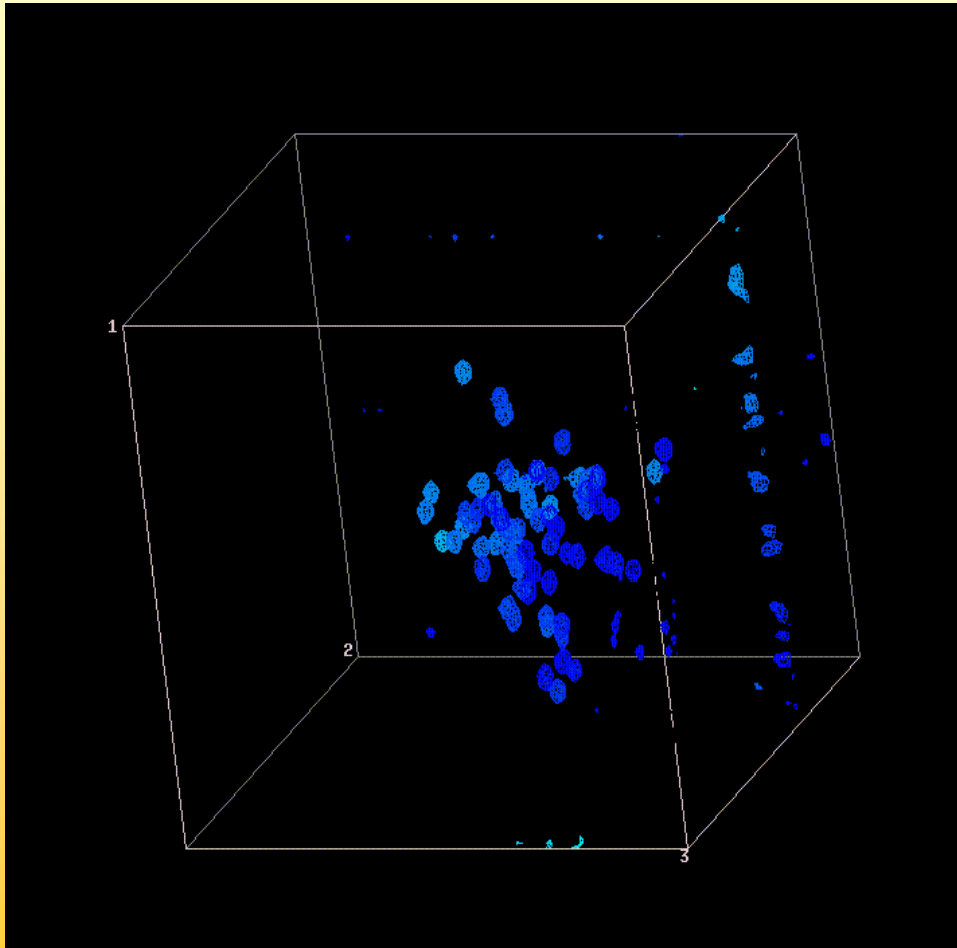
2. 2D-sequence



3D-sequence

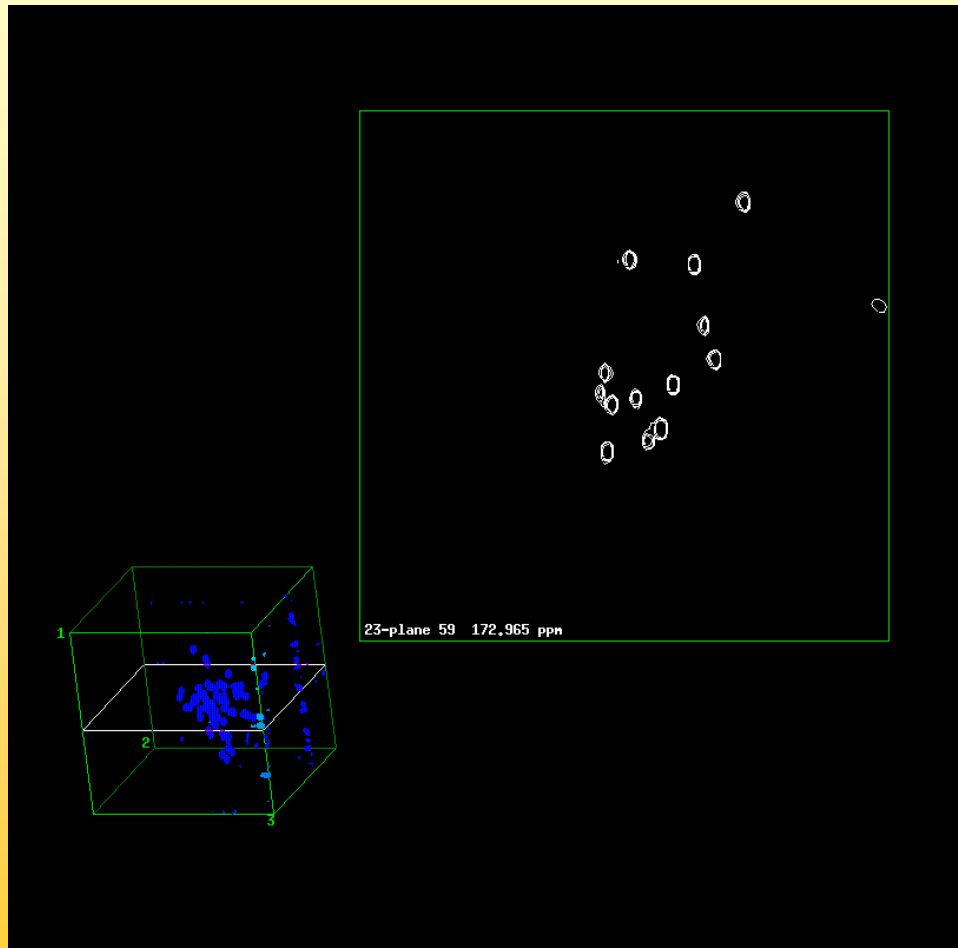


Multidimensional NMR-spectroscopy



The spectrum is then no longer a plane but a cuboid with chemical shift on all three axes. Intensity is then a „fourth dimension“

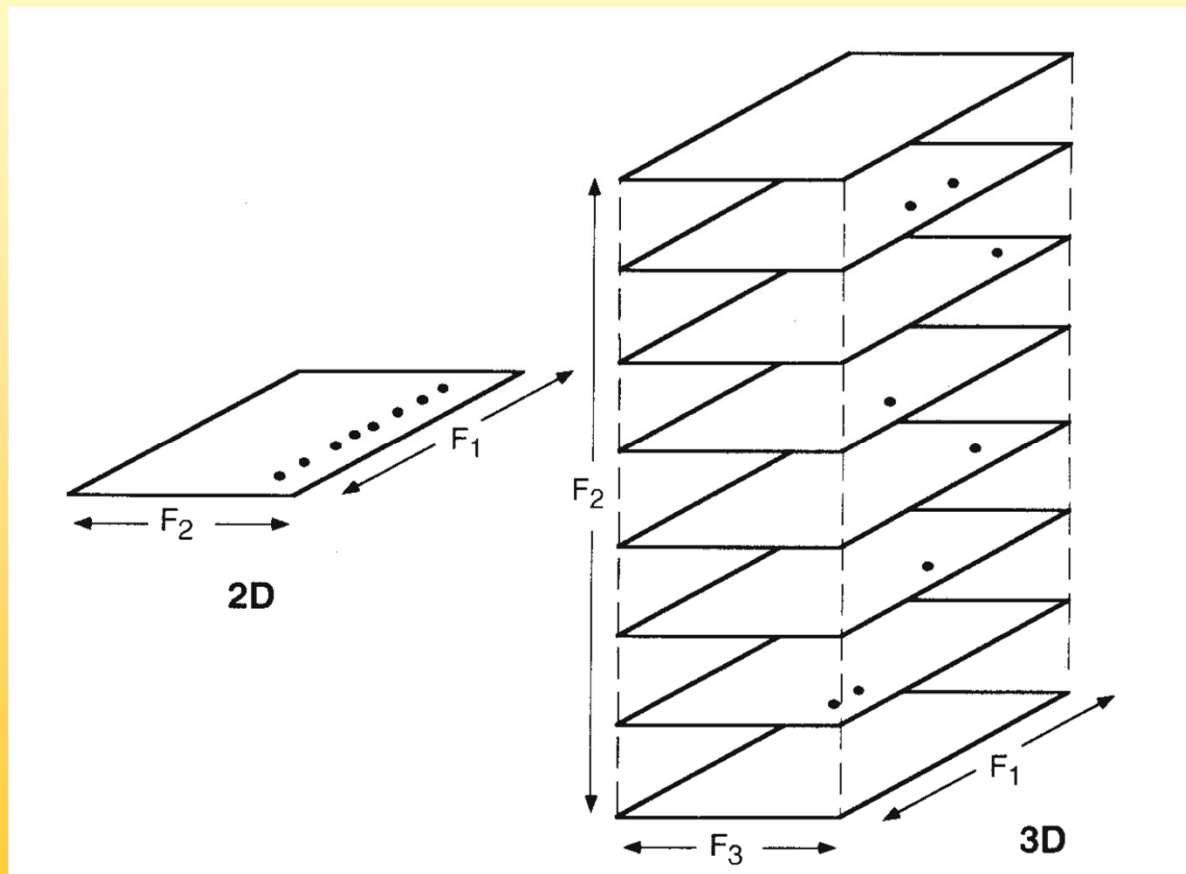
Multidimensional NMR-spectroscopy



Since the spectrum is difficult to analyze as a cuboid, planes are cut through at different chemical shifts of one axis resulting in a stack of 2D spectra

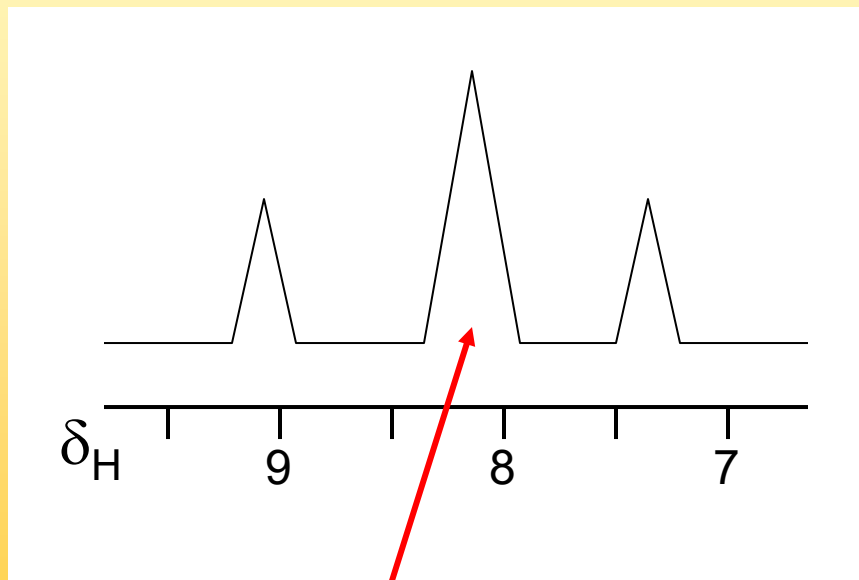
Multidimensional NMR-spectroscopy

3D-NMR



Multidimensional NMR-spectroscopy

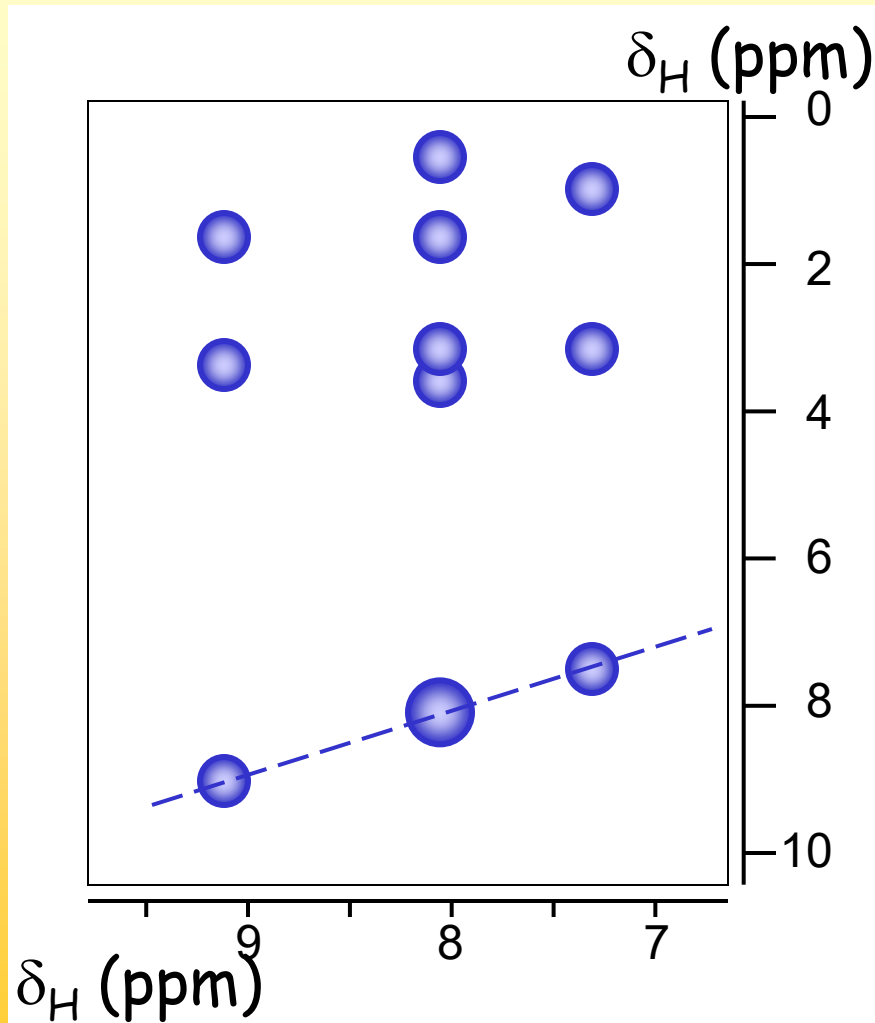
As a realistic example we look at an artificial region of amino protons, that are usually show overlap in the proton spectrum of a protein.



1D-¹H-spectrum:

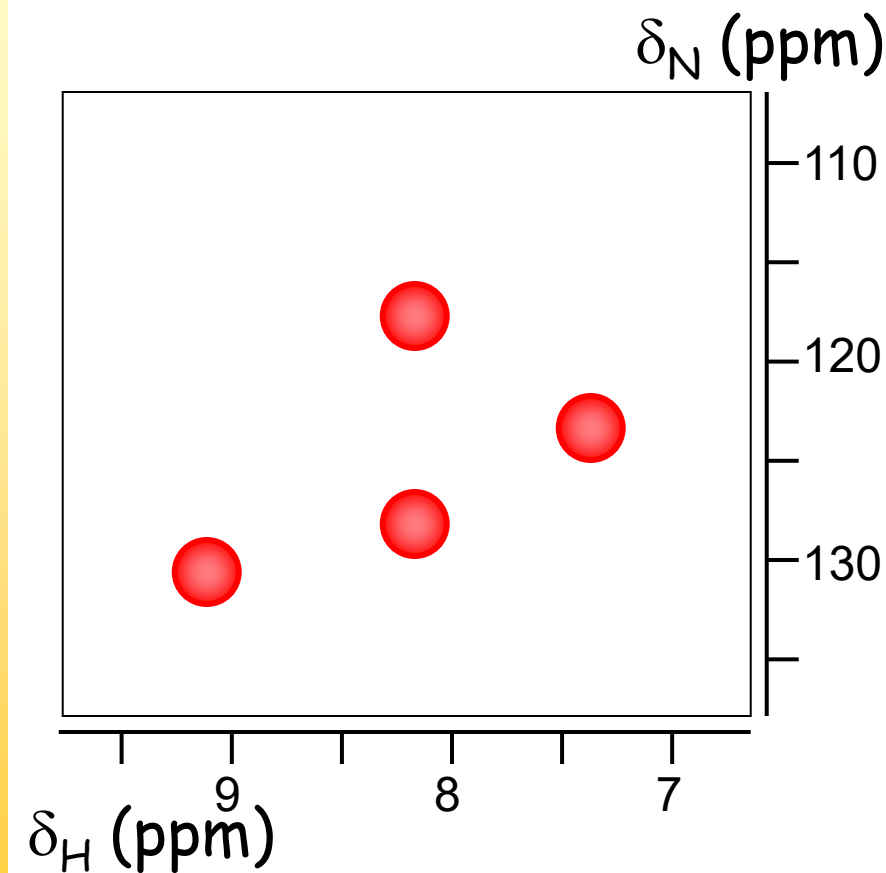
Region of the amino protons, 4 signals can be found, two of which overlap.

Multidimensional NMR-spectroscopy



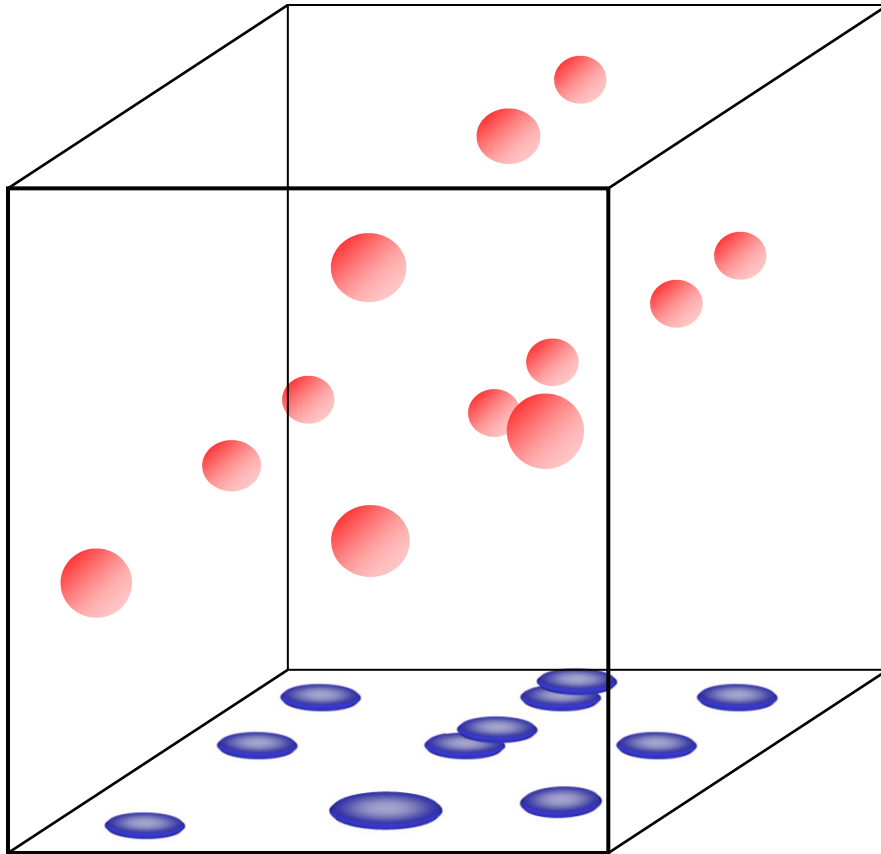
Homonuclear
 ^1H - ^1H spectrum:
 The overlap of the two
 amino protons prevent
 a distinction of the
 otherwise resolved
 protons in the aliphatic
 region

Multidimensional NMR-spectroscopy



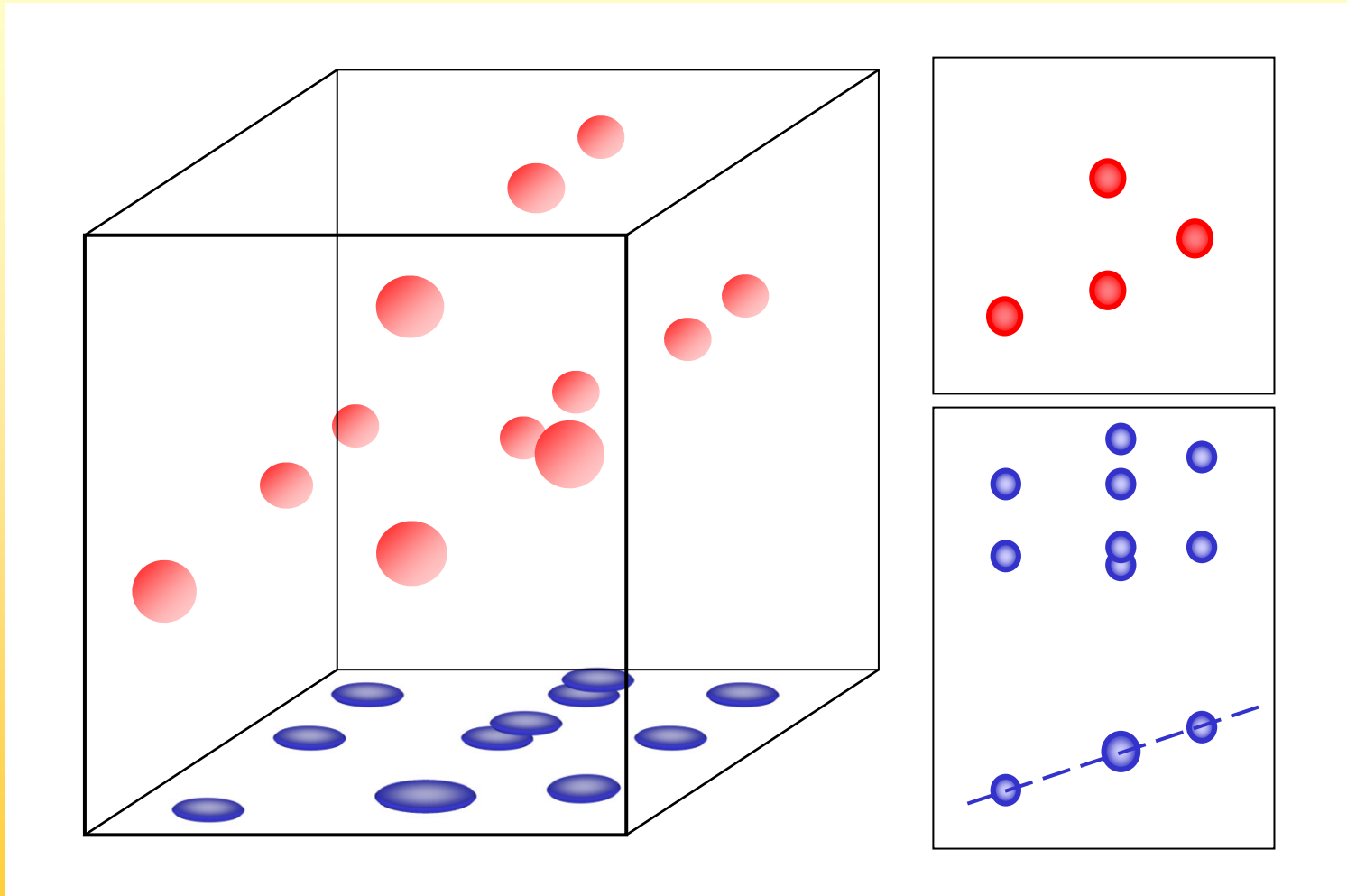
Heteronuclear
 ^1H - ^{15}N -spectrum:
The overlap of the
two amino protons is
resolved due to
different ^{15}N
chemical shifts.

Multidimensional NMR-spectroscopy



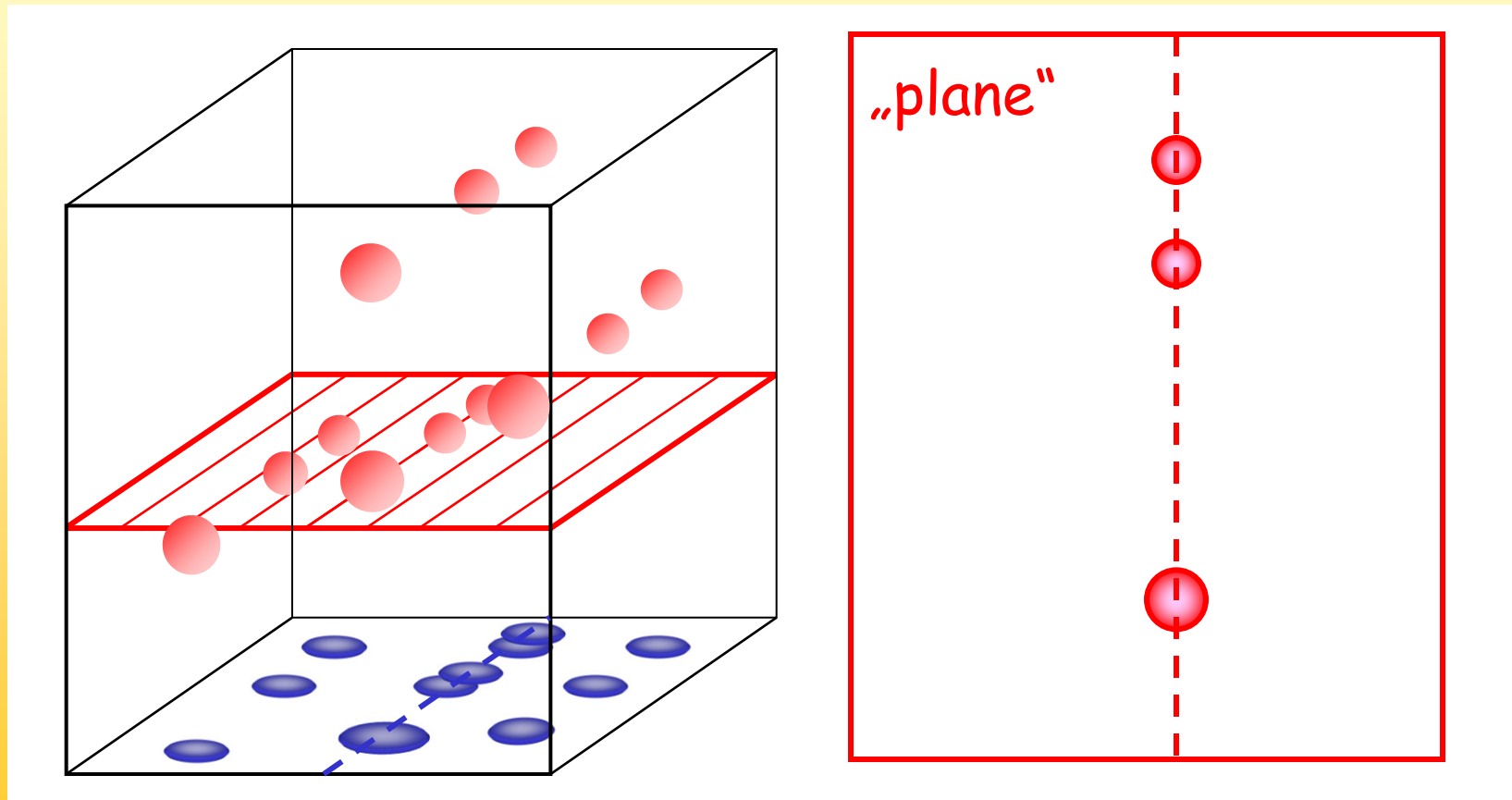
In an ^1H - ^1H - ^{15}N -3D-spectrum all overlap is resolved. Two of the faces of the cuboid correspond to the original 2D spectra.

Multidimensional NMR-spectroscopy



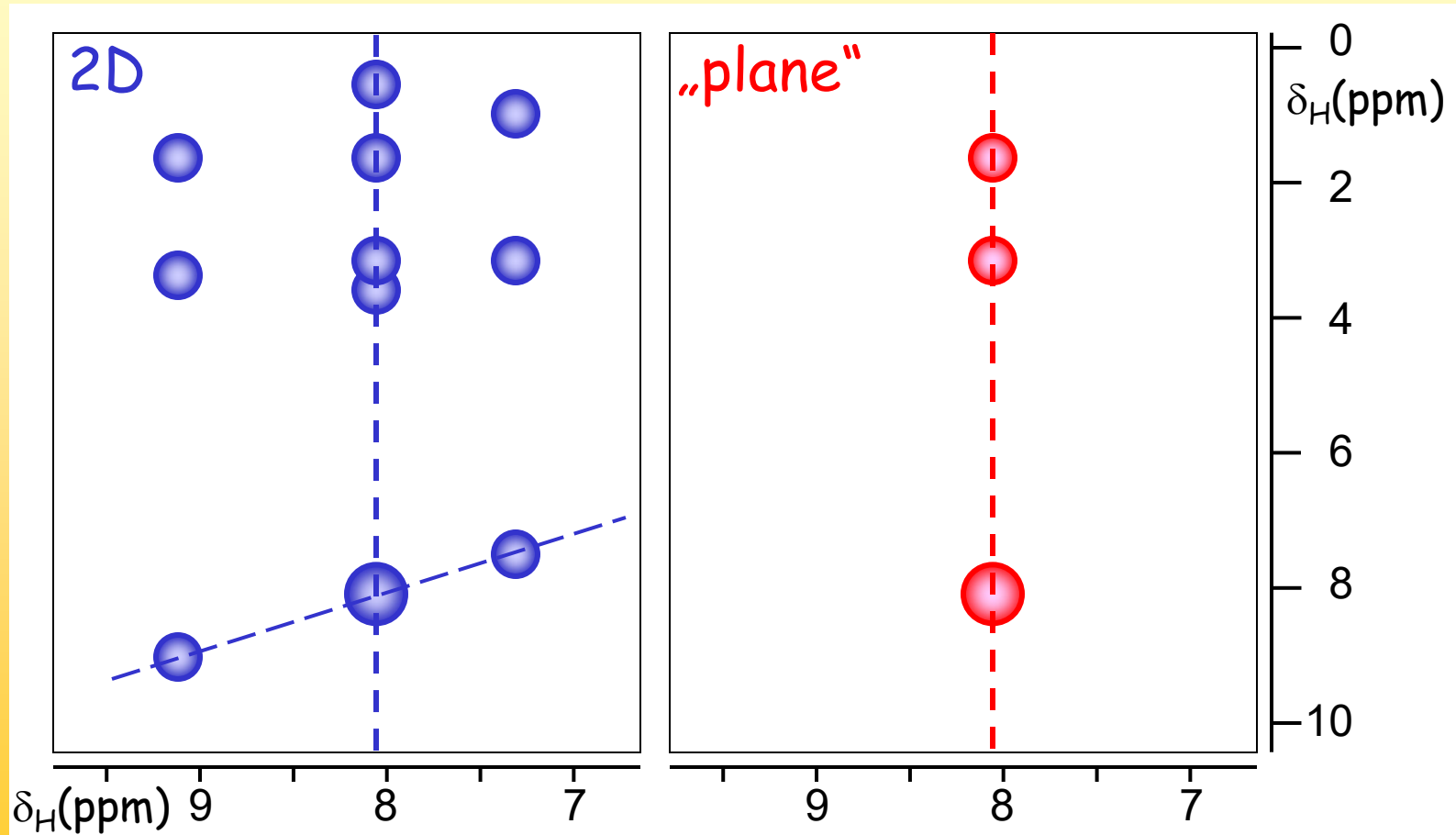
Multidimensional NMR-spectroscopy

Now we cut a „plane“ at a certain ^{15}N chemical shift out of the cuboid.

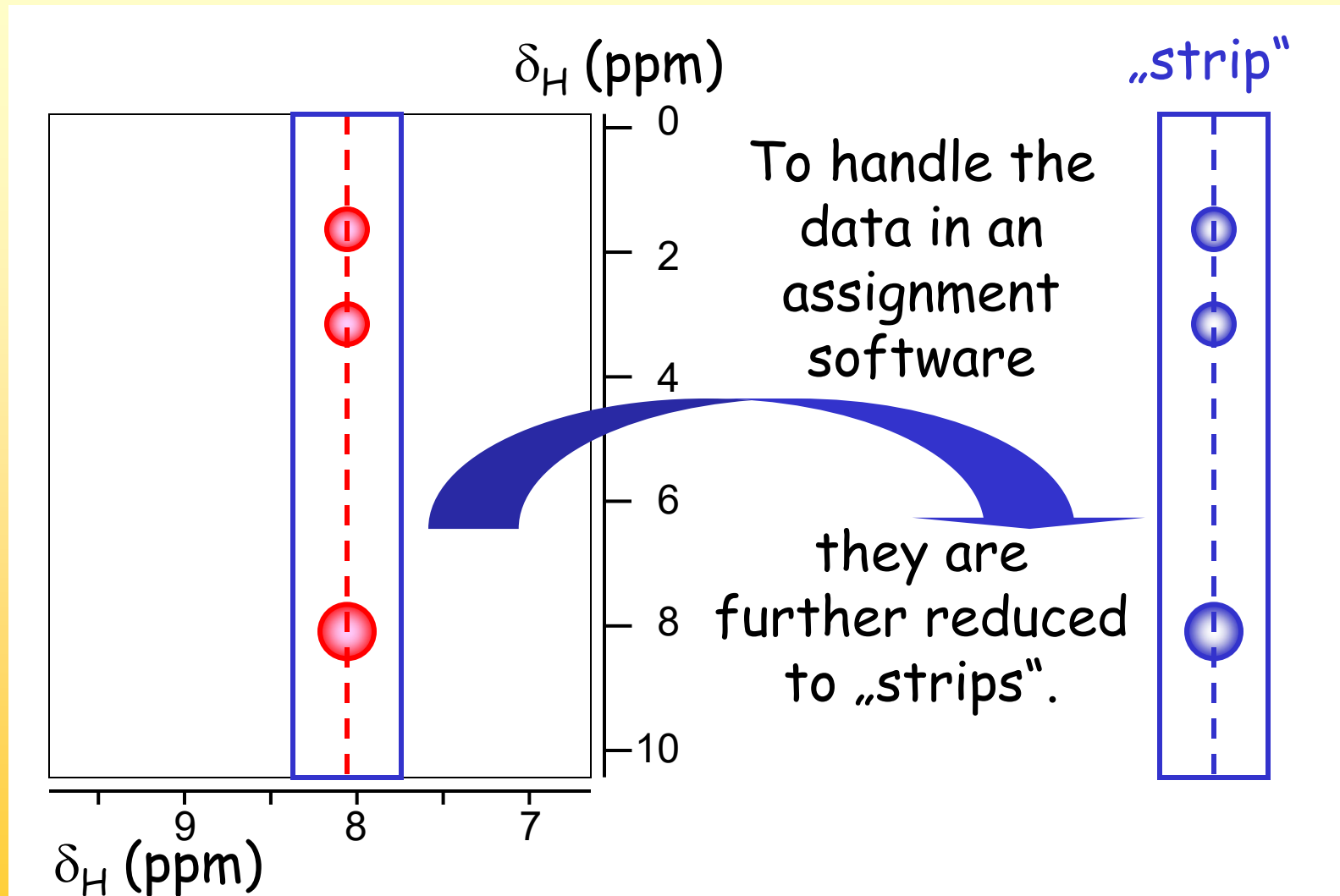


Multidimensional NMR-spectroscopy

The overlap has been removed from the ^1H - ^1H -2D spectrum

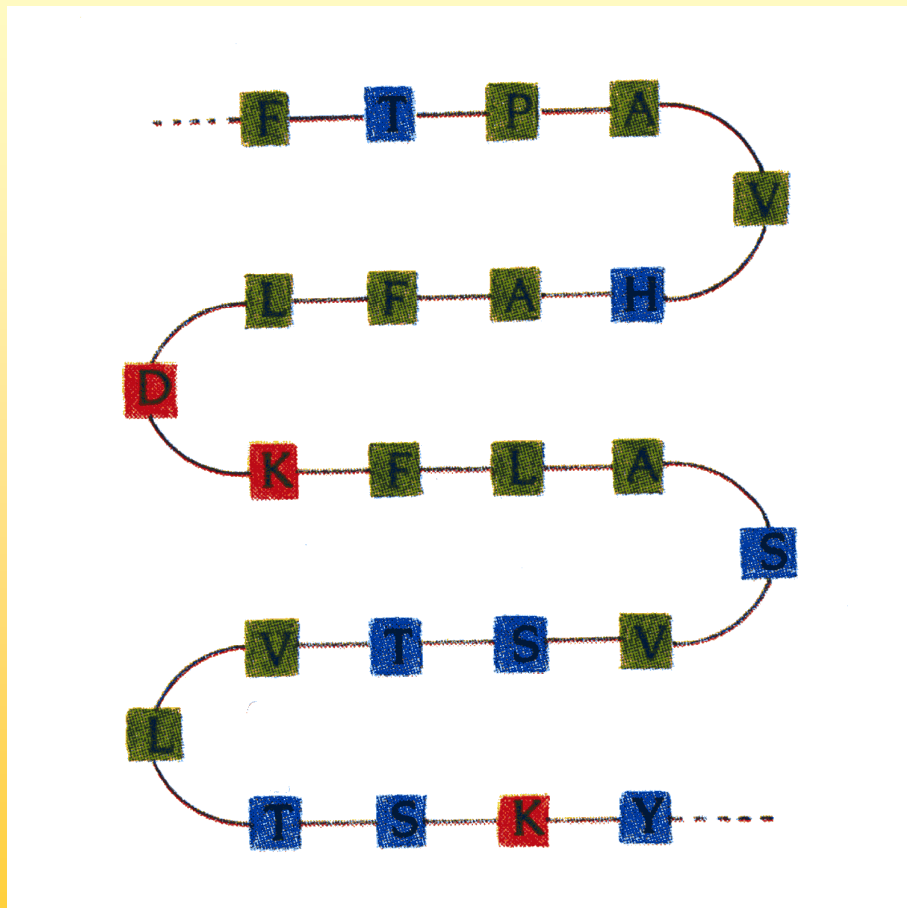


Multidimensional NMR-spectroscopy



Sequence specific assignment
of proteins using
tripel-resonance-techniques

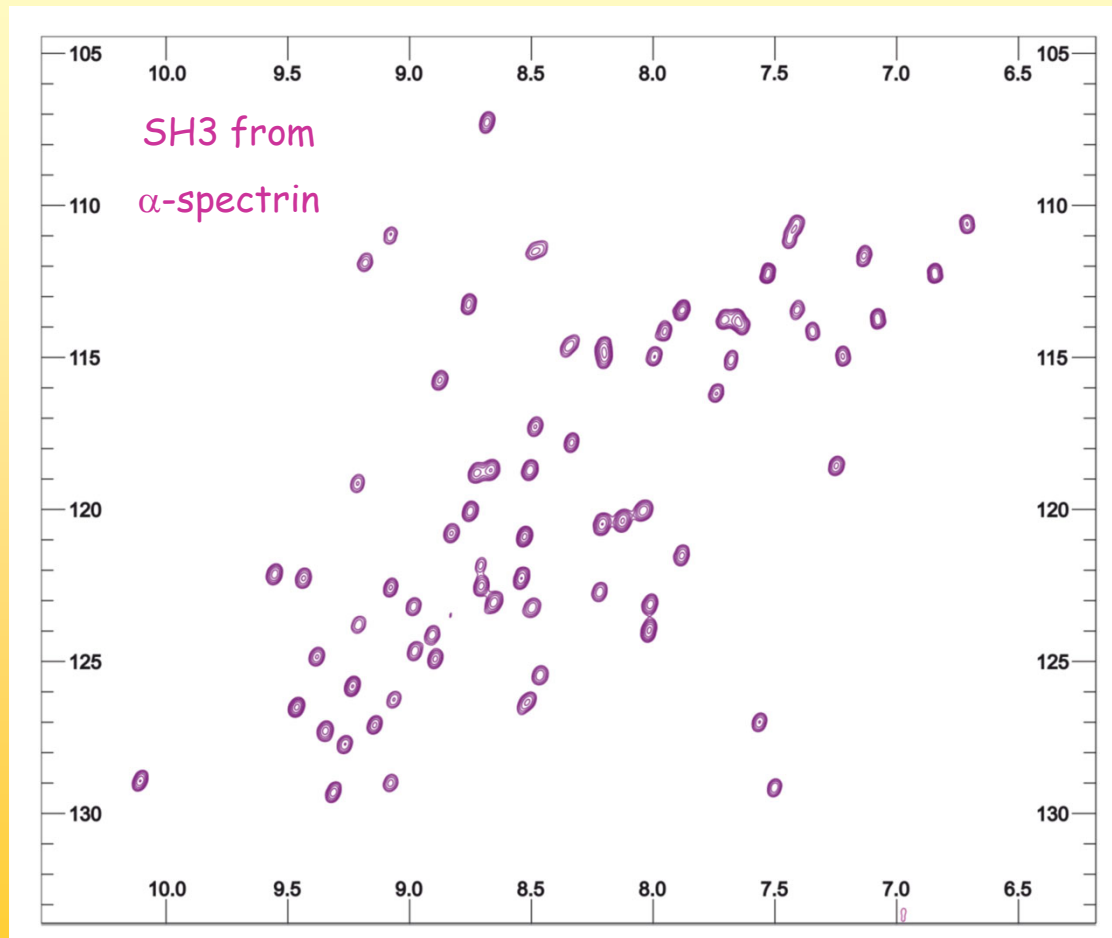
Sequence specific assignment of proteins



As with all other molecules the first step of an extraction of information is the assignment of resonances - but we have that „polymer-problem“.

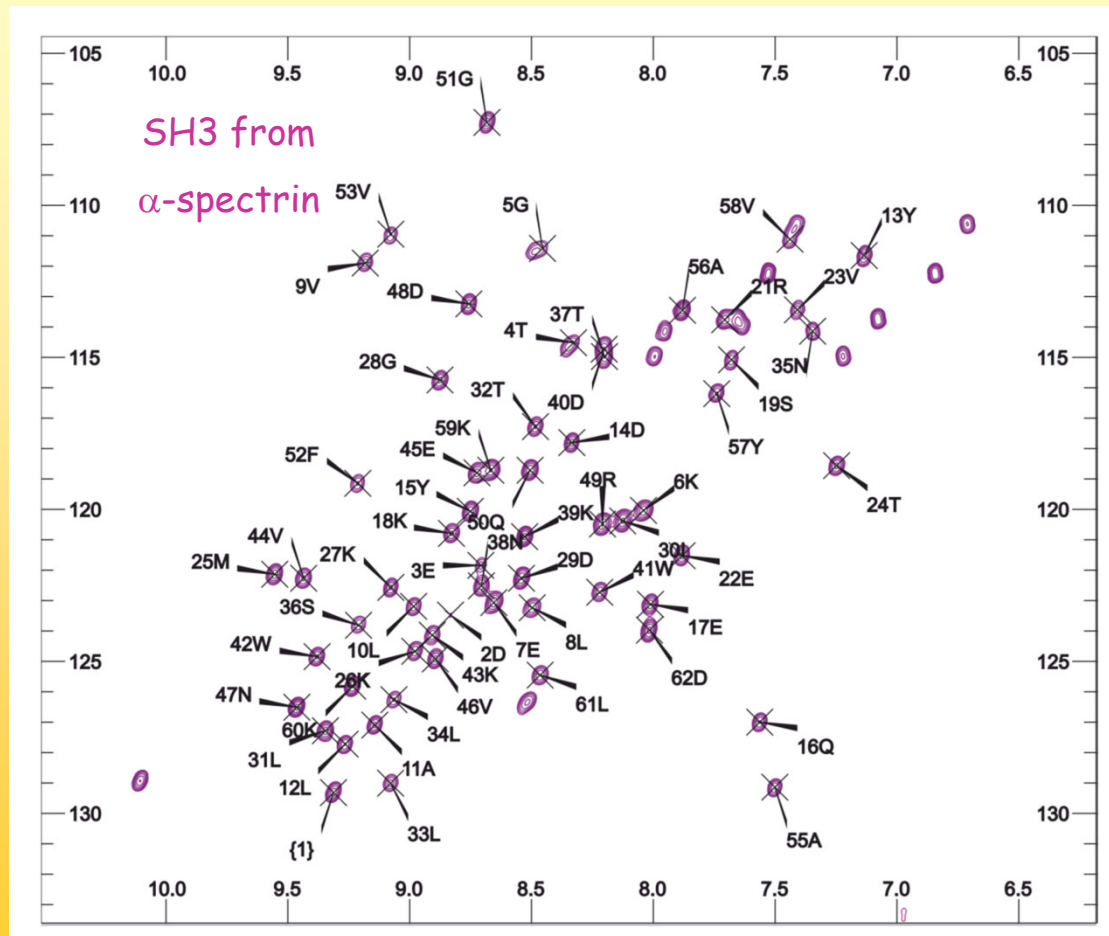
Sequence specific assignment of proteins

We need to get from here



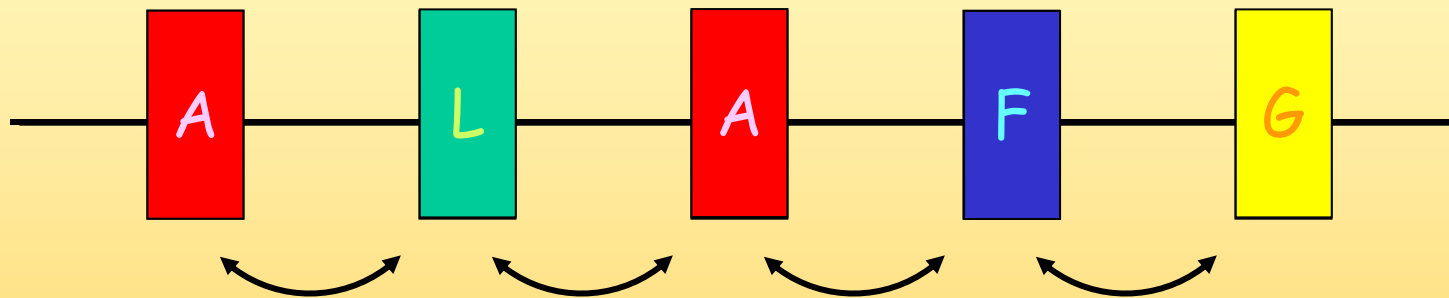
Sequence specific assignment of proteins

to here



Sequence specific assignment of proteins

The solution for the assignment problem is the
sequence-specific assignment



1. Which type of amino acid is present (color)
2. Which aa is next to which (neighborhood)
3. Comparison with the protein sequence to assign
4. The order of (1) and (2) is irrelevant

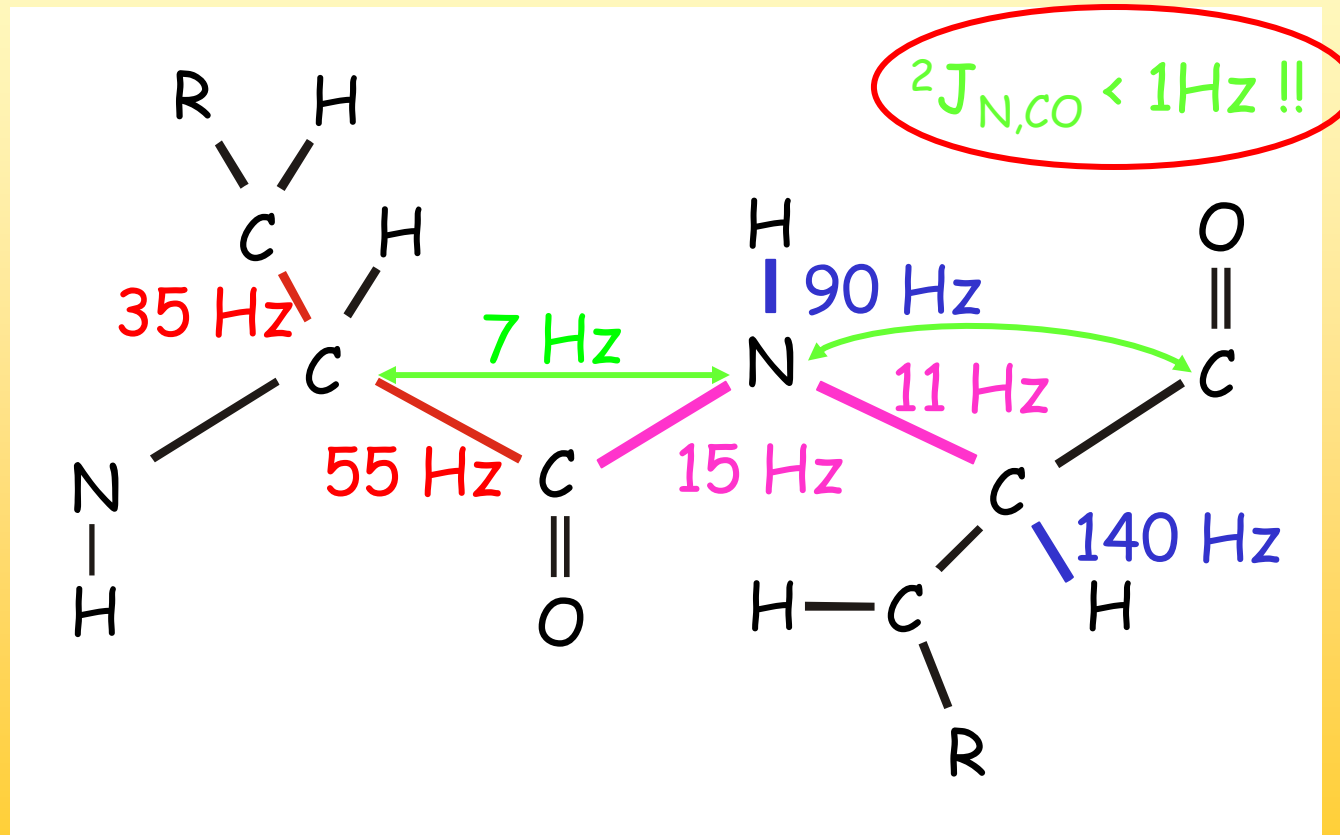
Sequence specific assignment of proteins

Besides the better dispersion of heteronuclear spectra that is utilized in 3D spectra a transfer of magnetization via scalar couplings between heteronuclei is more efficient for labeled proteins than a transfer via proton-proton couplings.

To record spectra all three nuclei (^1H , ^{13}C and ^{15}N) and the scalar couplings between them are therefor used. That's why the group of NMR experiments are called:
"Tripel-resonance-experiments"

Sequence specific assignment of proteins

J-couplings between heteronuclei in proteins



Sequence specific assignment of proteins

Because of the differences between the aliphatic and the carbonyl carbons in proteins both can be treated separately spectroscopically:

1. Chemical shift

$$\delta_{CO} \sim 170-180 \text{ ppm}$$

$$\delta_{C\alpha/\beta} \sim 10-70 \text{ ppm}$$

2. Scalar carbon-carbon coupling

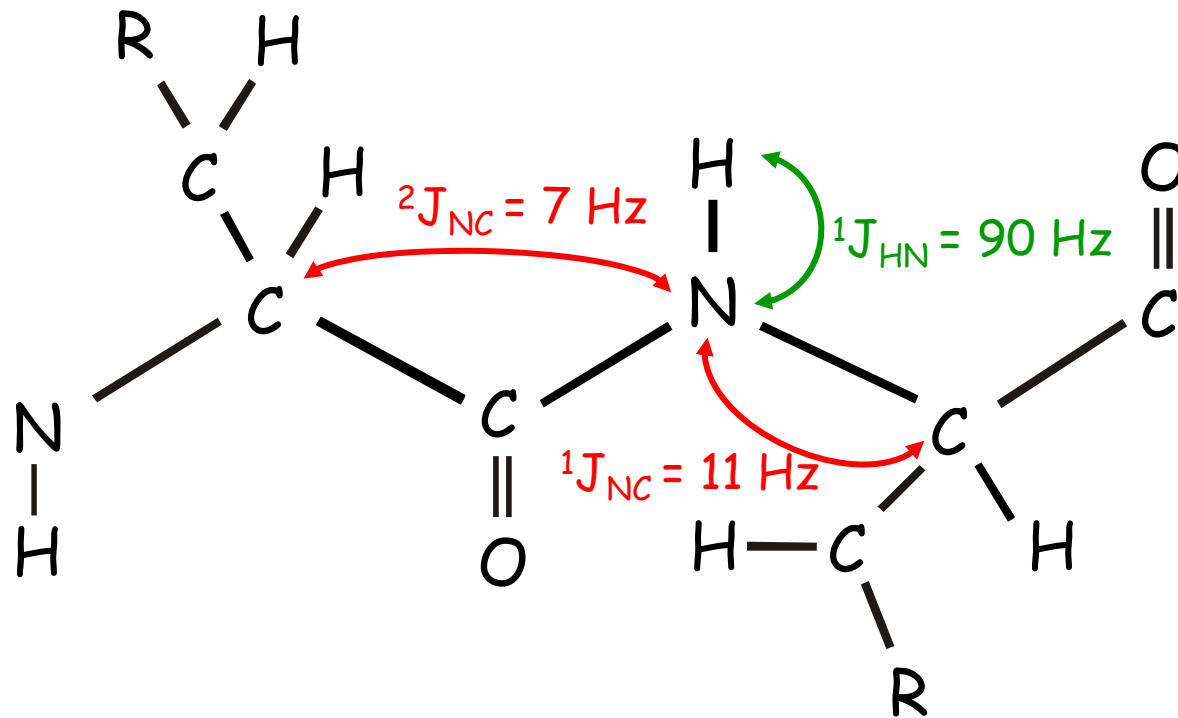
$$J(CO, C\alpha) \sim 55 \text{ Hz}$$

$$J(C, C) \sim 35 \text{ Hz}$$

3. Carbonyl carbons have no protons attached.
Carbonyl carbons are a "fourth" type of nucleus.

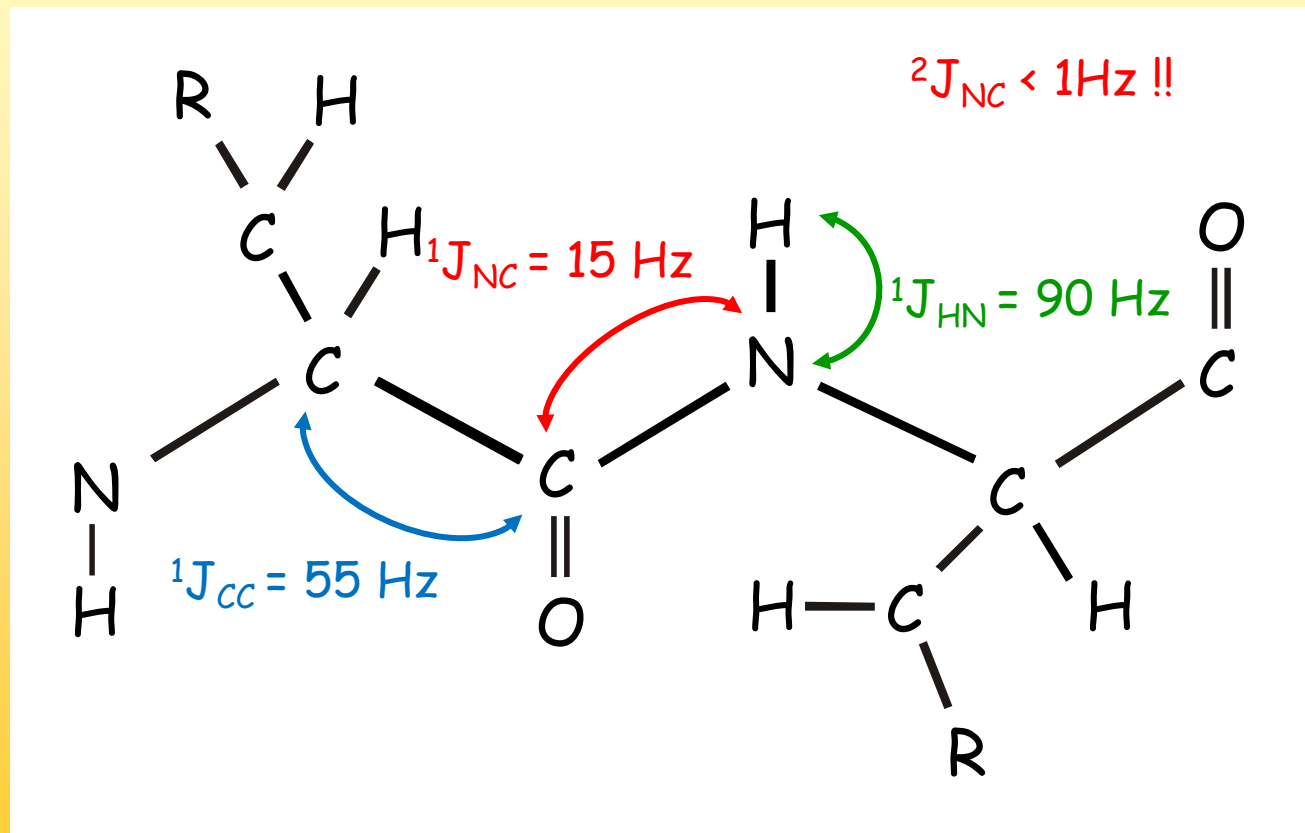
Sequence specific assignment of proteins

HNCA



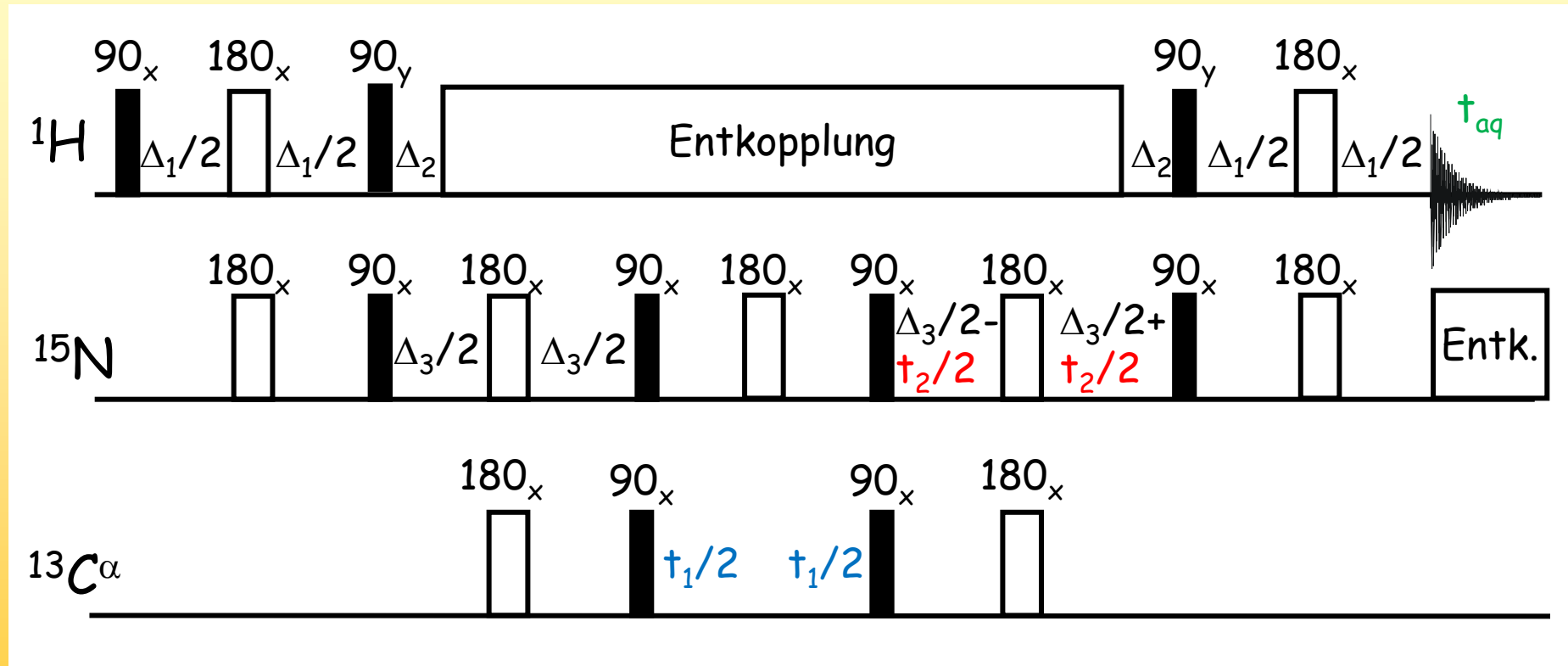
Sequence specific assignment of proteins

HN(CO)CA

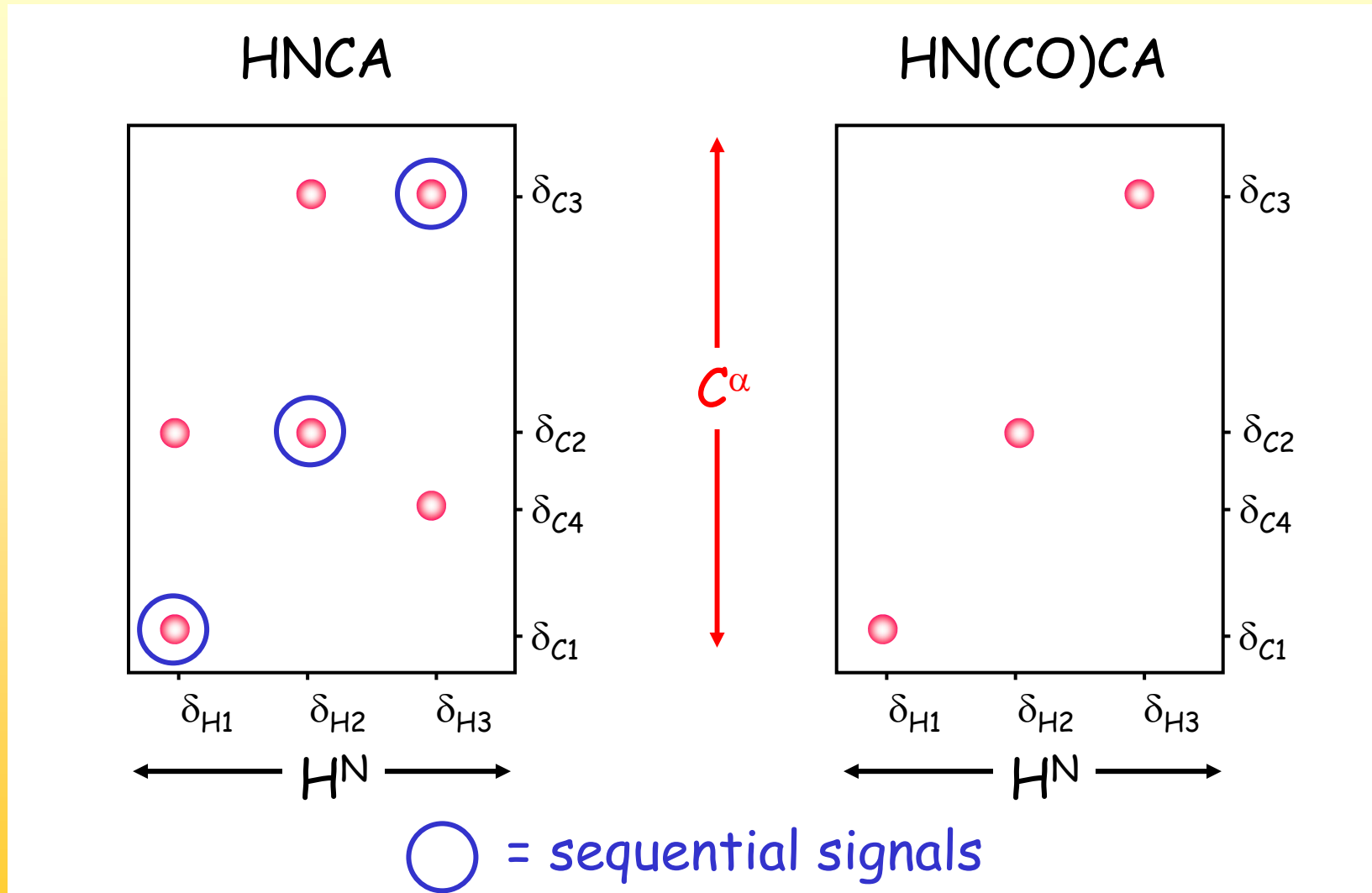


Sequence specific assignment of proteins

The pulse sequence of the HNCA

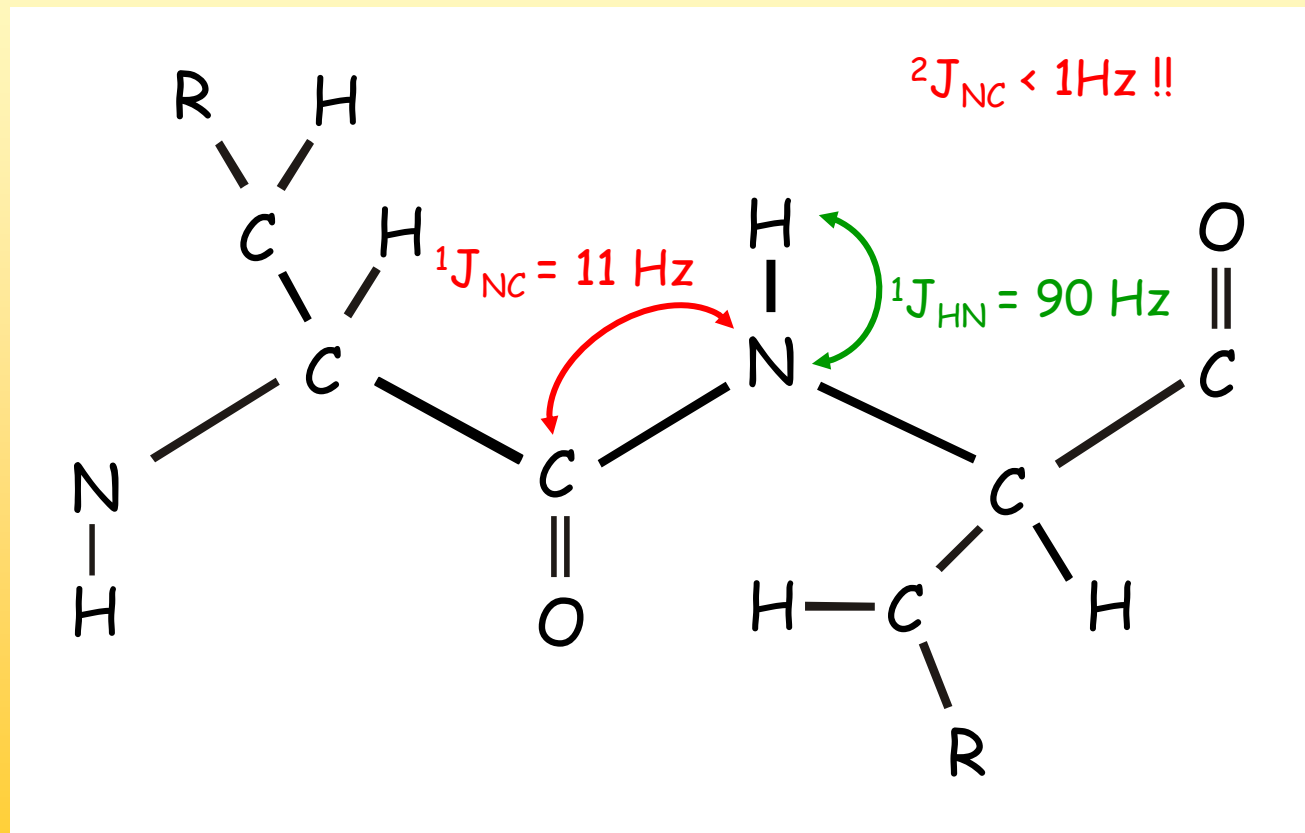


Sequence specific assignment of proteins



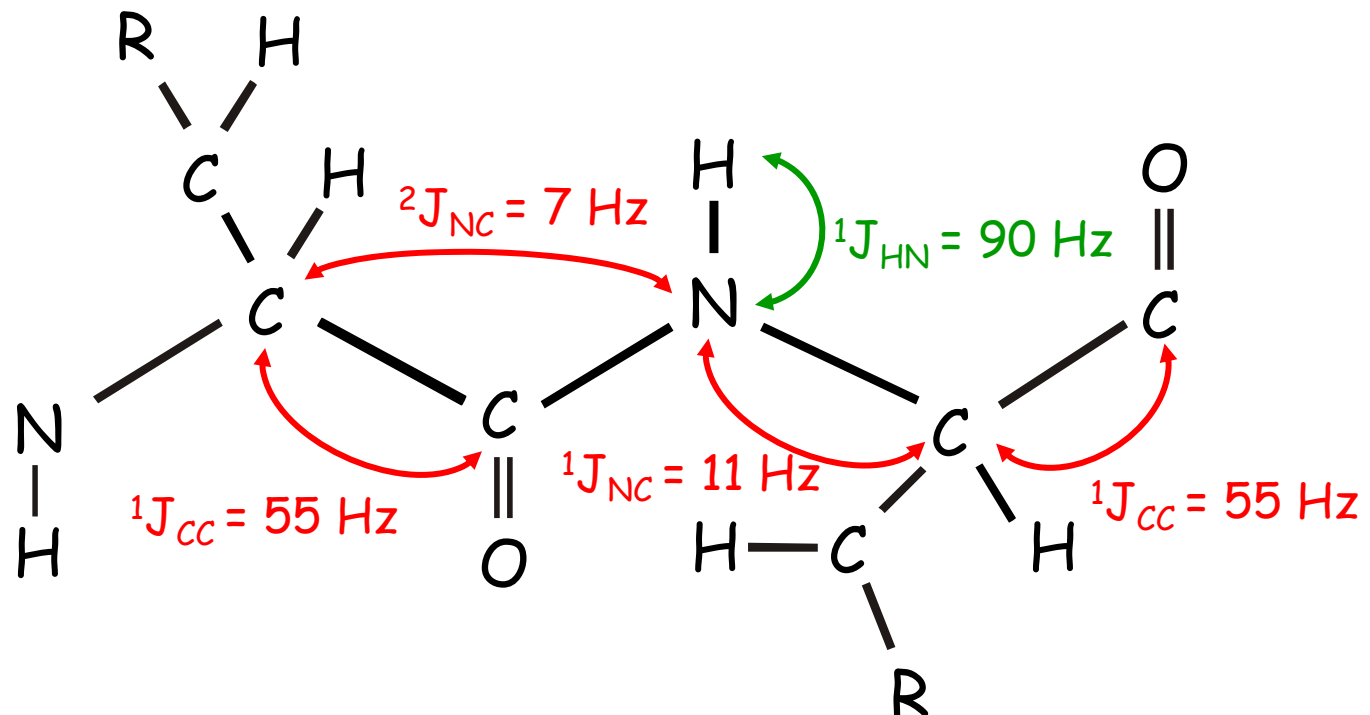
Sequence specific assignment of proteins

HNCO

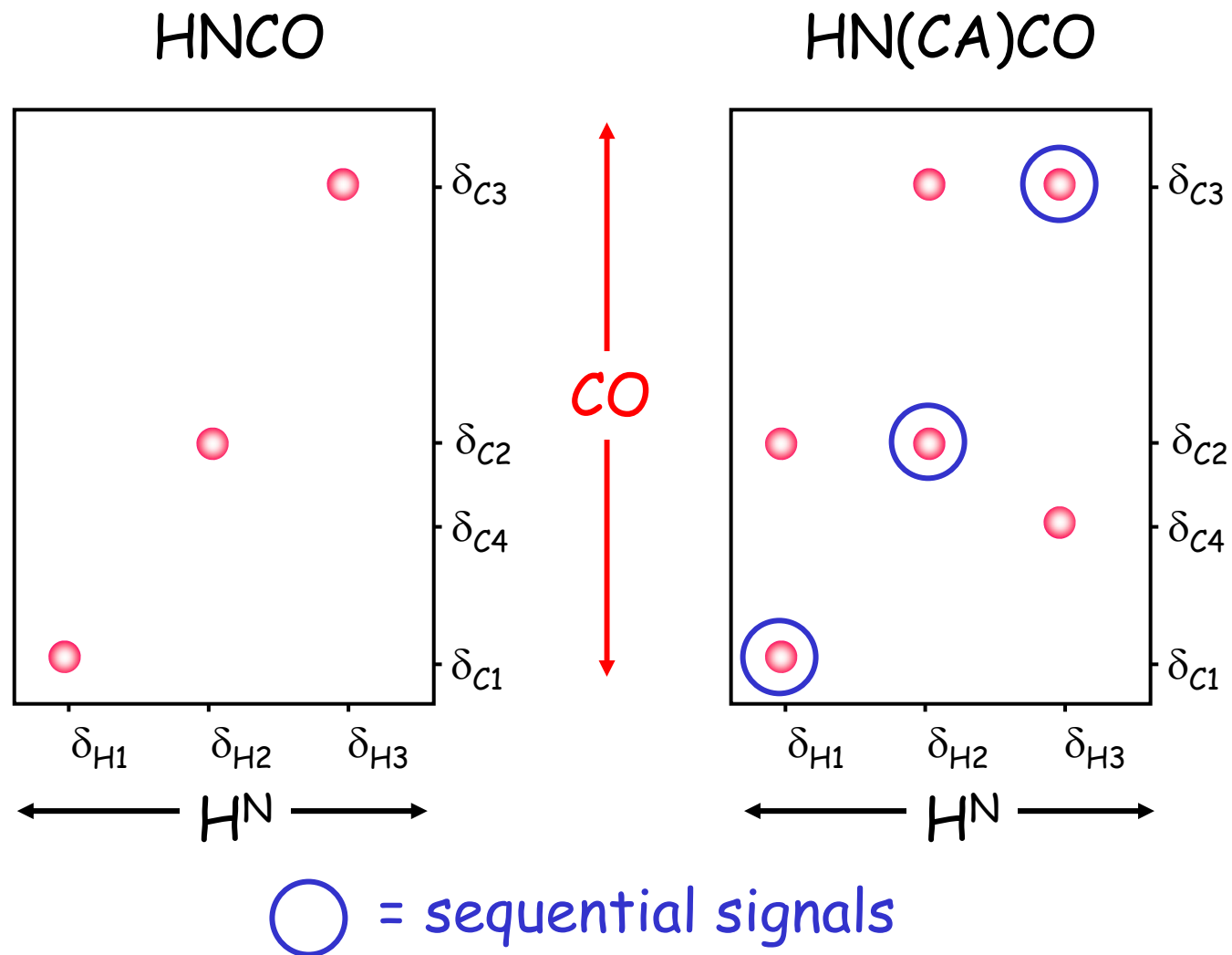


Sequence specific assignment of proteins

HN(CA)CO

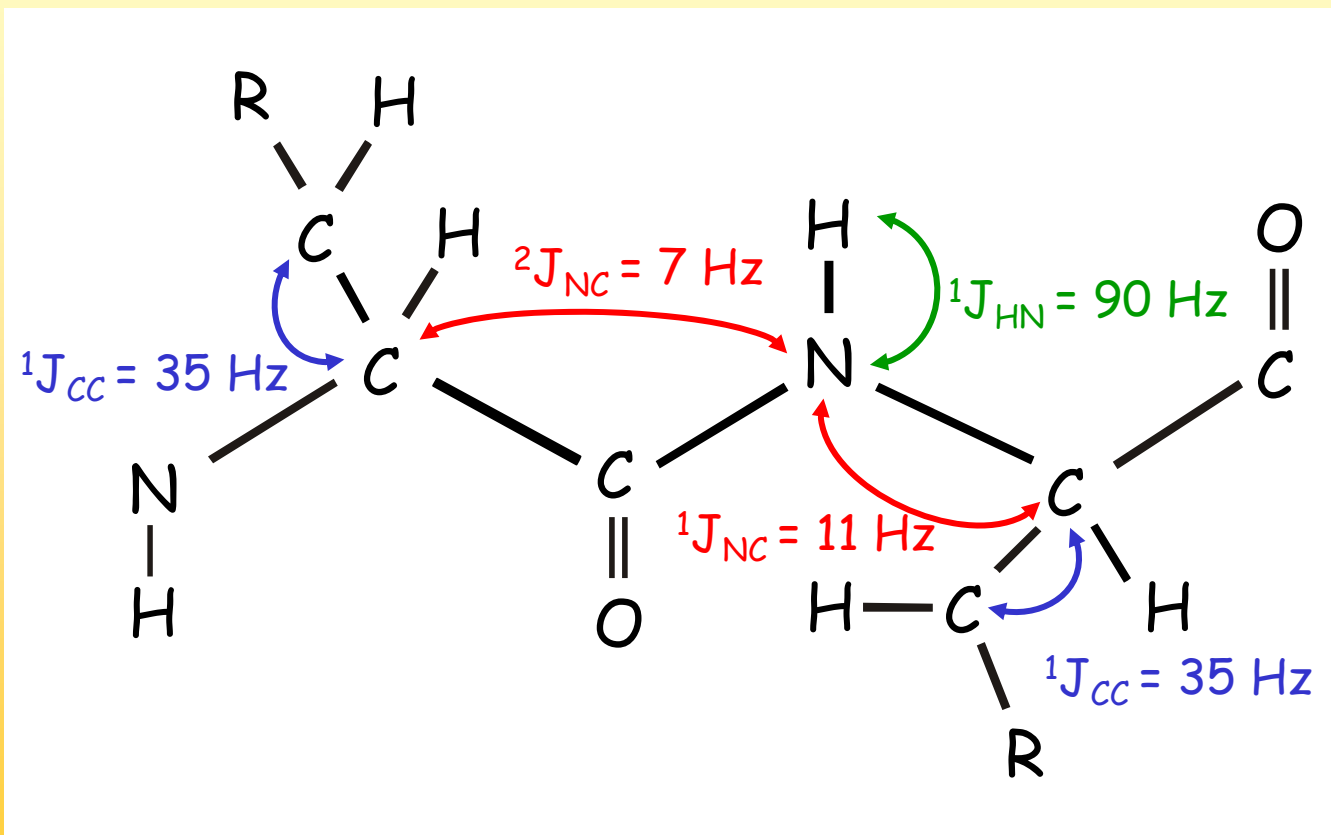


Sequence specific assignment of proteins



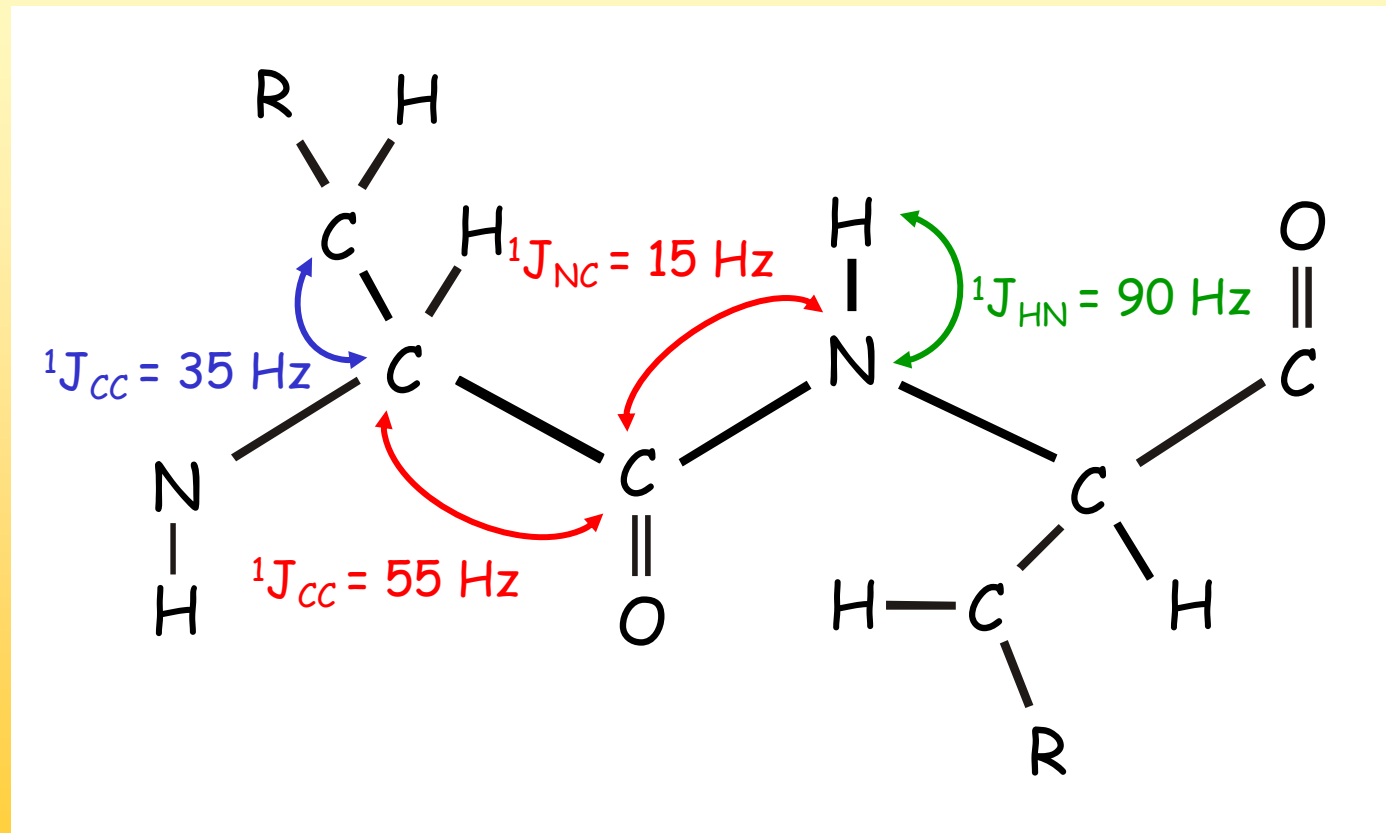
Sequence specific assignment of proteins

HNCACB



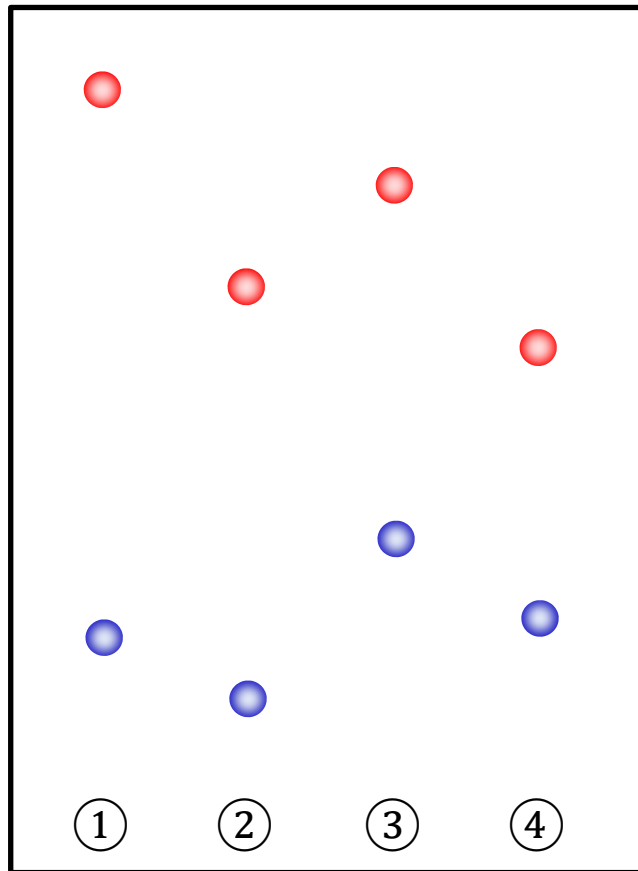
Sequence specific assignment of proteins

HN(CO)CACB

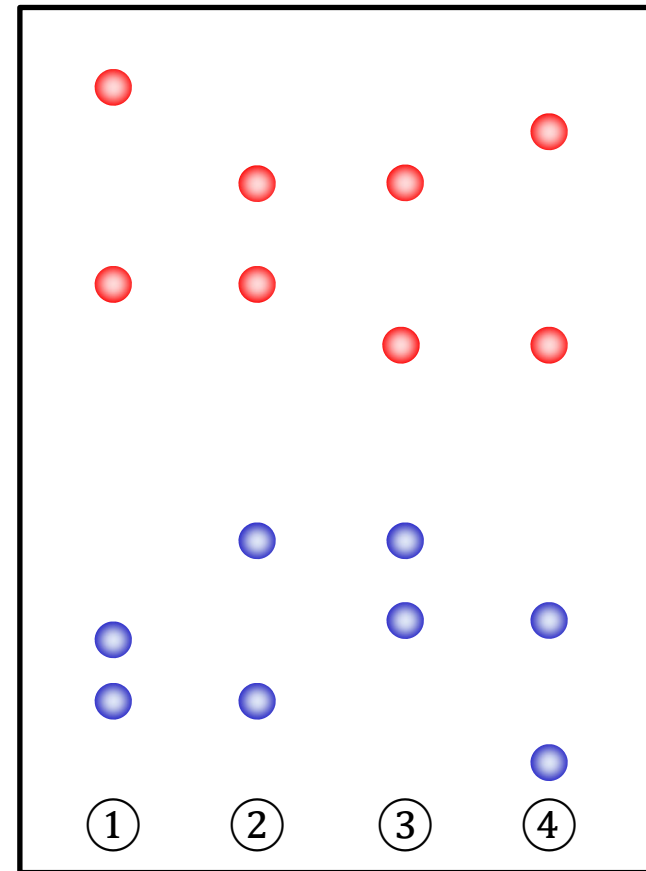


Sequence specific assignment of proteins

HN(CO)CACB

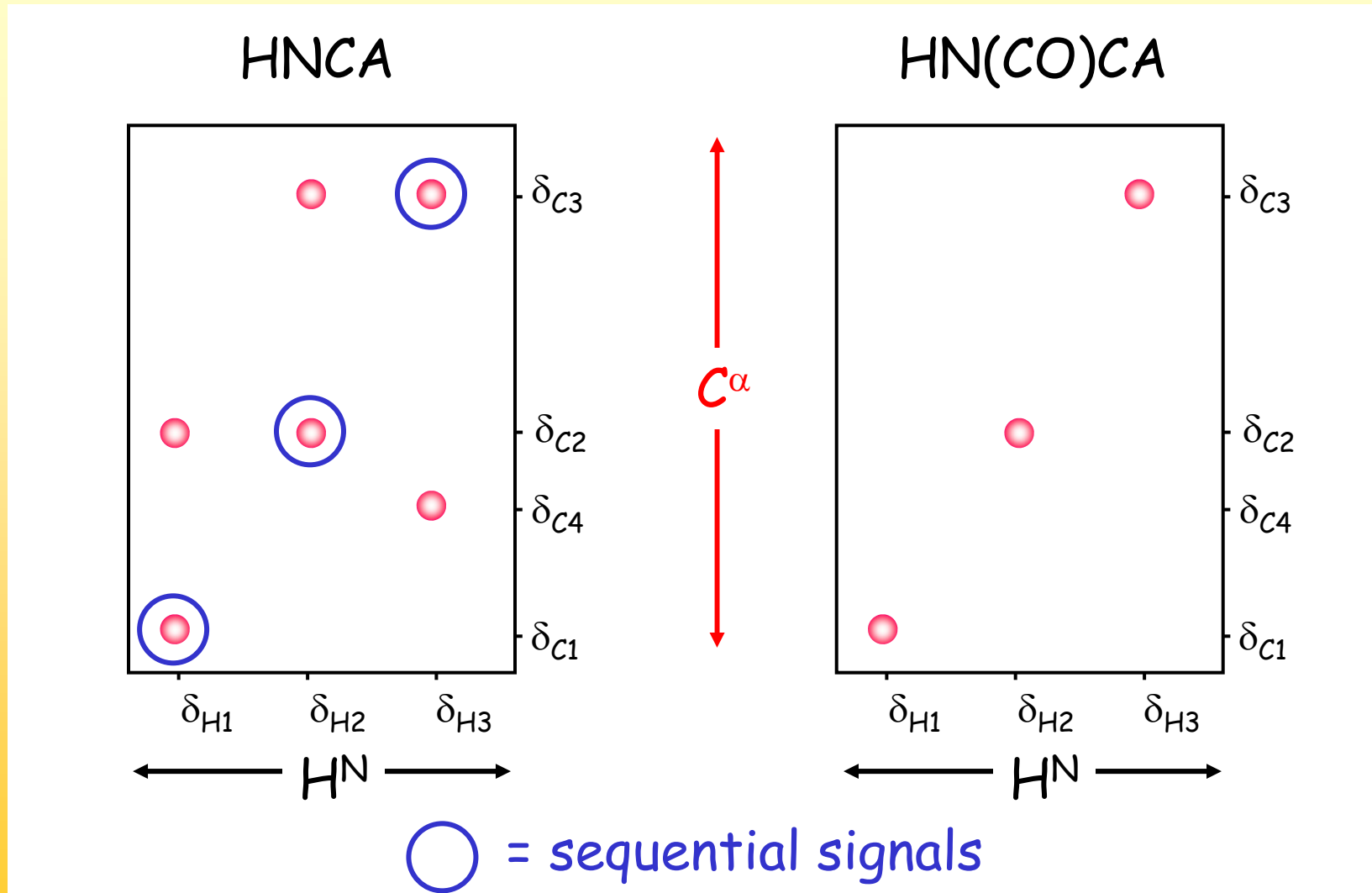


HNCACB

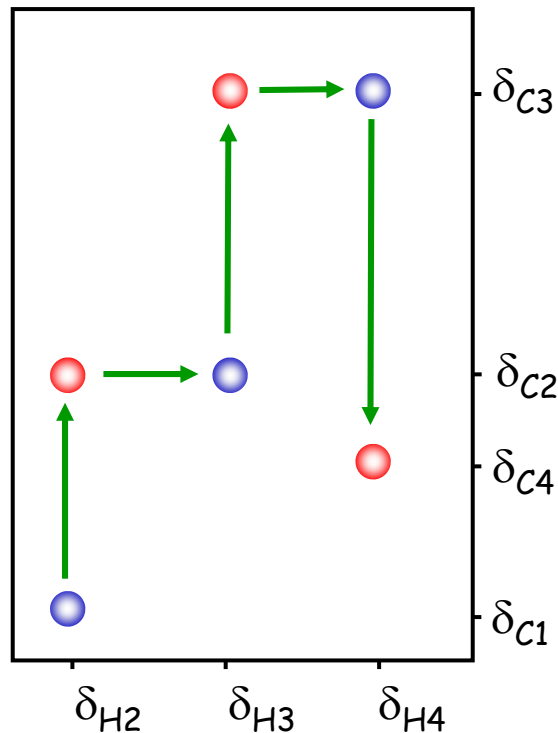
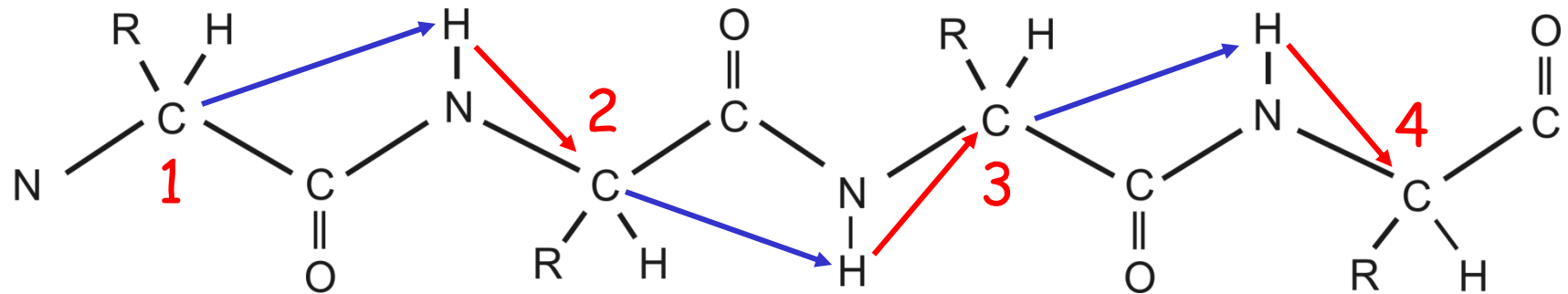


Vertical axis labels: C^{β} (red) and C^{α} (blue).

Sequence specific assignment of proteins



Sequence specific assignment of proteins



To do the assignment we do a „sequential walk“.

Blue arrows correspond to blue peaks, red arrows to red peaks.

The connection represented by the green arrows can be made by keeping the frequency (and thus the nucleus) constant

Sequence specific assignment of proteins

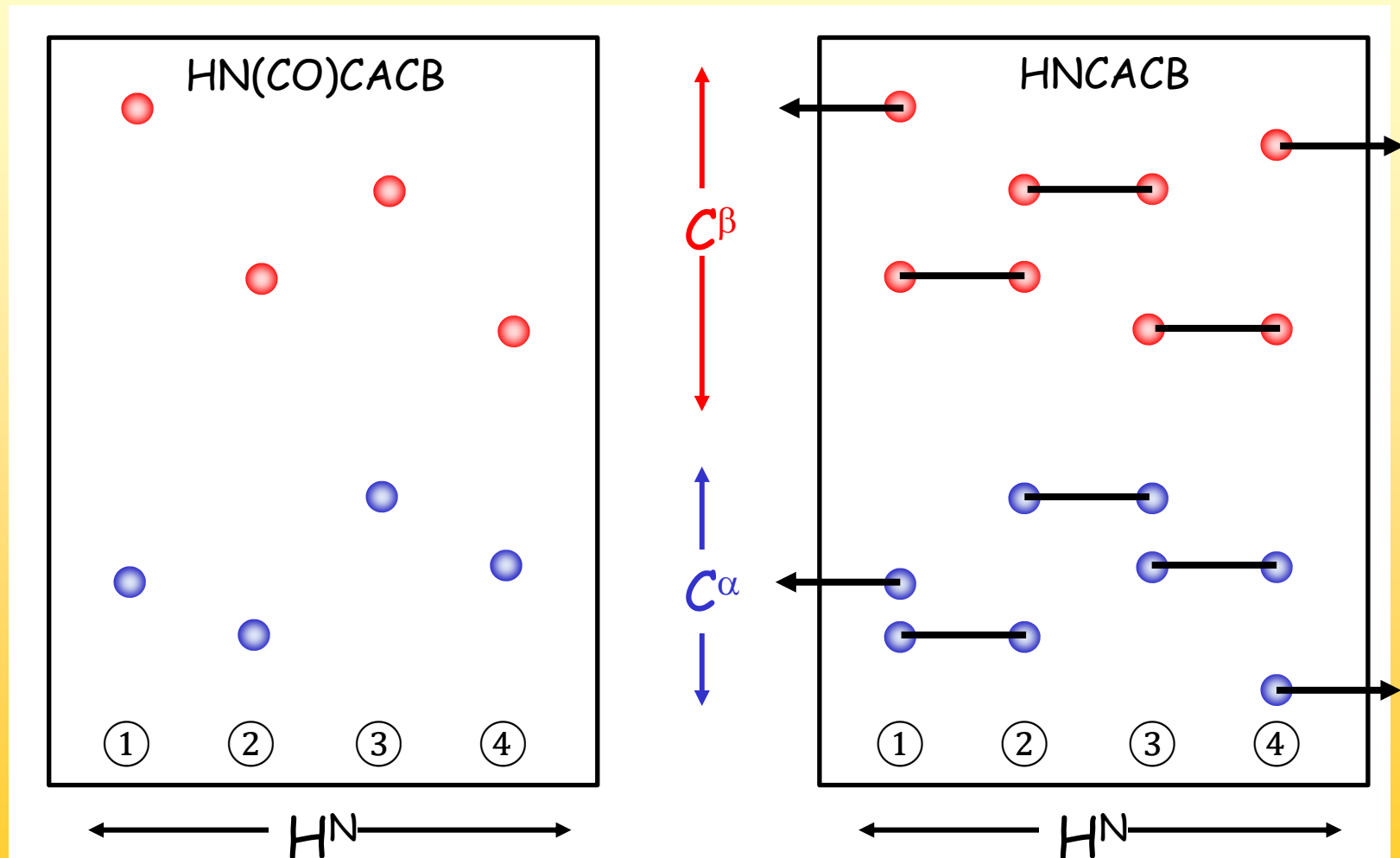
Using the HNCA/HN(CO)CA there can be the problem that two C^α shifts are identical.

Then it is possible to use the pair HNCO and HN(CA)CO as well in a similar fashion and it is unlikely that overlap occurs in both.

But it is even better to use C^α and C^β simultaneously in the HNCACB and HN(CO)CACB pair.

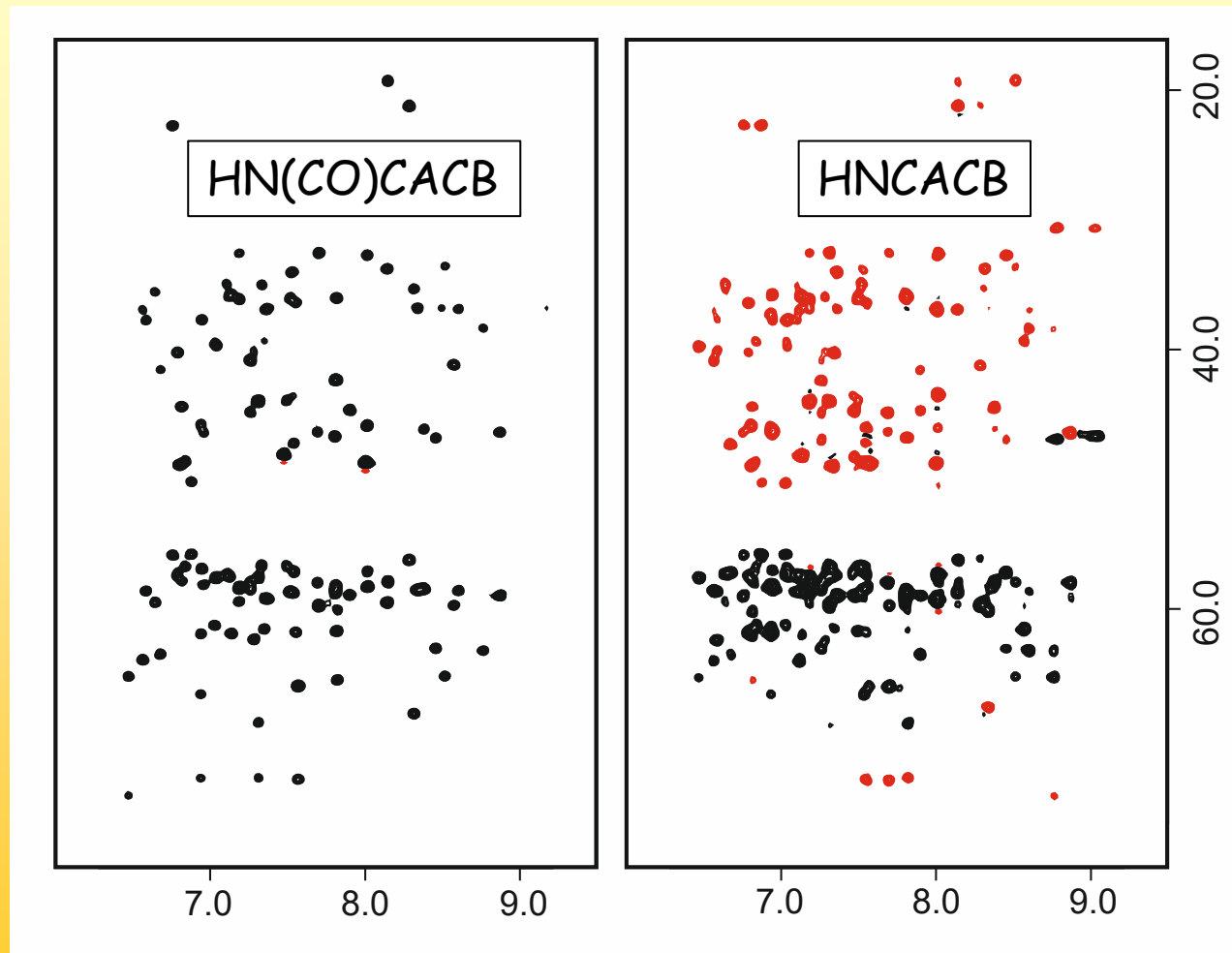
Sequence specific assignment of proteins

Sequential assignment



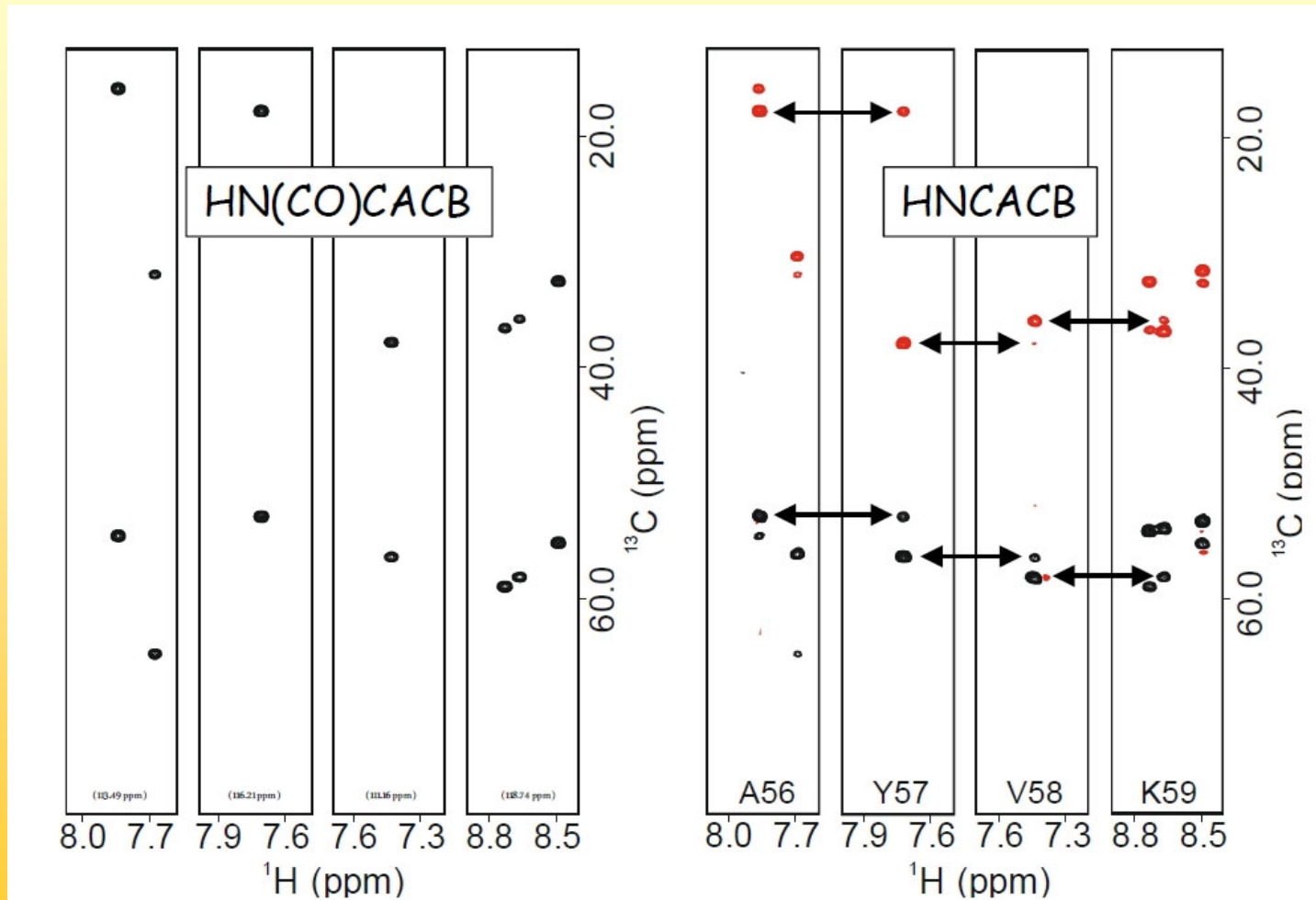
Sequence specific assignment of proteins

Sequentielle Zuordnung



Sequence specific assignment of proteins

Sequentielle Zuordnung



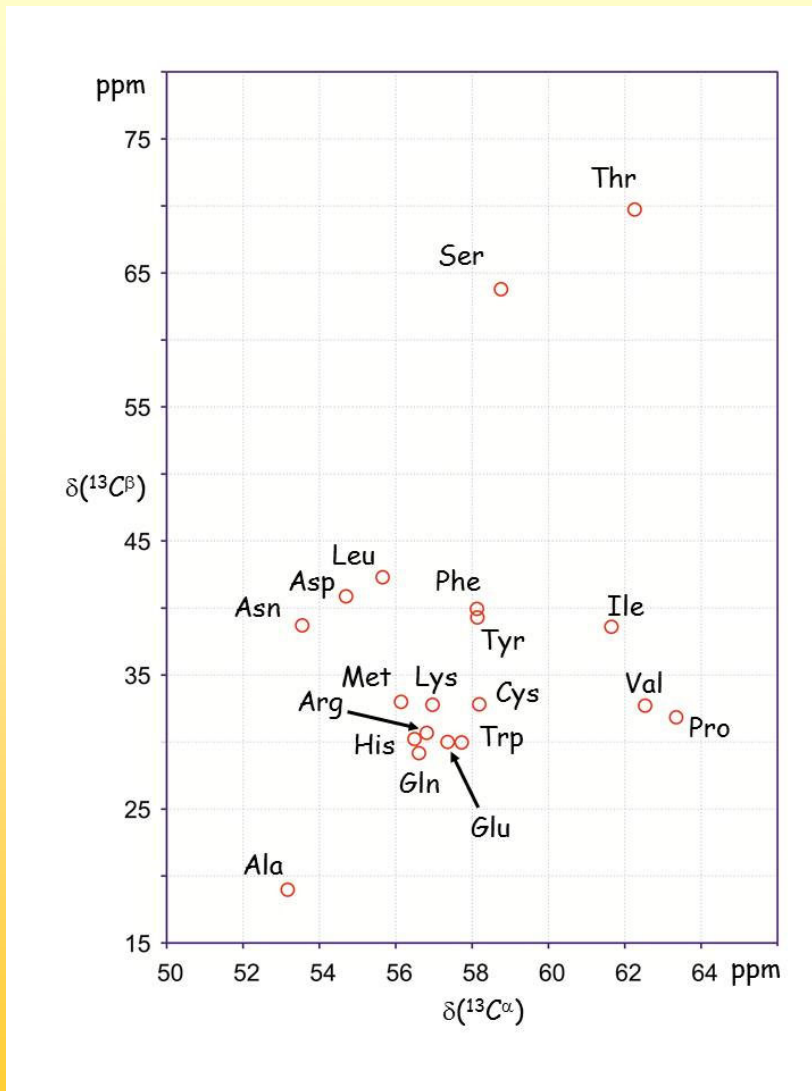
Sequence specific assignment of proteins

Using those pairs of triple resonance experiments it is possible to establish a sequential neighborhood.

To be able to compare the stretches of amino acids with the sequence in order to obtain the final assignment we need information on the type of amino acid.

That can be obtained from the chemical shifts of the C_α und C_β , that are available from the HNCACB or HN(CO)CACB

Sequence specific assignment of proteins



Amino acid „typing“

Ala	53,170	18,990
Arg	56,810	30,680
Asn	53,550	38,710
Asp	54,700	40,880
Cys	58,190	32,810
Gln	56,610	29,180
Glu	57,360	30,000
Gly	45,370	
His	56,490	30,220
Ile	61,650	38,610
Leu	55,660	42,300
Lys	56,970	32,790
Met	56,140	33,000
Phe	58,130	39,950
Pro	63,350	31,850
Ser	58,760	63,800
Thr	62,260	69,730
Trp	57,730	29,970
Tyr	58,140	39,300
Val	62,530	32,720

That's it

<https://www.schmieder-nmr.de/teaching/praktikum.htm>



LEIBNIZ-INSTITUT
FÜR MOLEKULARE
PHARMAKOLOGIE

Practical course „Molekulare Pharmakologie und
zelluläre Signaltransduktion : NMR-spectroscopy

Beerbaum/Schmieder
AG Solution-NMR