Determination of structure of glycans and glycan interactions with proteins by NMR

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Why glycans are puzzling, why are they not obvious?





Open, pyranose and furanose forms of an aldose, showing the many equivalent OH groups. The OH groups can be modified by $-PO_4^{-3}$, $-SO_4^{-2}$, $-NH_2$, $-CO-CH_3$, $-NH-CO-CH_3$, -H.

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Structure determination of a glycan chain: wet organic chemistry for every step



Available online at www.sciencedirect.co ScienceDirect

Structural Biology

Solving the structural puzzle of bacterial glycome Roberta Marchetti, Rosa Ester Forgione, Ferran Nieto Fabregat, Cristina Di Carluccio, Antonio Molinaro and Alba Silipo

- Quali-quantitative analysis (OC, GC-MS, NMR)
- Absolute configuration (OC, GC-MS, NMR)
- Size of the ring (OC, GC-MS, NMR)
- Anomeric configuration (OC, NMR)
- Linkage analysis (OC, GC-MS, NMR)
- Monosaccharides sequence (OC, MALDI-MS, 2D NMR)
- Determination of non-carbohydrate appendages (OC, GC-MS, MALDI-MS, 2D NMR)



Meth. Enzymol., 2010; ACS Chem Biol, 2018; Curr. Opin. Struct. Biol., 2021; Carbohydr. Polym., 2021



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Carbohydrate Polymers

journal homepage: www.elsevier.com/locate/carbpol

Liquid-state NMR spectroscopy for complex carbohydrate structural analysis: A hitchhiker's guide

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Typical Applications of NMR:

Structural (chemical) elucidation

Natural product chemistry

Organic chemistry: Analytical tool of choice for synthetic chemists.

Study of dynamic processes

Reaction kinetics. Study of equilibrium (chemical or structural).

Structural (three-dimensional) studies Proteins Data (page)

DNA/RNA Polysaccharides Drug design

Structure Activity Relationships (SAR) by NMR

• Medicine - Magnetic Resonance Imaging (MRI)

NMR active nuclei



Nucleus (odd atomic number): ¹H, ¹³C, ¹⁵N, ¹⁹F, ³¹P

How does NMR work?

A spinning charge creates a magnetic moment, so these nuclei can be thought of as tiny magnets.



In presence of a magnetic field Magnetic moments precess and orient with or against the field

NMR Signal





$$\begin{aligned} & \mathbf{V}_{\text{PEAK}} - \mathbf{V}_{\text{REF}} \text{ (Hz)} \\ & \mathbf{\delta} \text{ (ppm)} = ----- = ppm \\ & \text{Freq of the nuclei (MHz)} \end{aligned}$$

• Depending on the *chemical environment* we have variations on the magnetic field that the nuclei feel, even for the same type of nuclei. It affects the local magnetic field.



¹H - ¹H Coupling

Signals do not appear as single lines, sometimes they appear as multiple lines. This is due to ¹H - ¹H coupling (also called spin-spin splitting **or J-coupling**).



A and B are scalarly coupled nuclei



The N + 1 Rule

If a signal is split by N equivalent protons, it is split into N + 1 peaks.



¹H NMR

☞ Number of signals (number of non-equivalent H)

Chemical shift (differences in chemical environment)

Splitting or Coupling (number of neighboring H)

☞ Integration (relative number of H at each signal)

- Chemical shift data tells us what kinds of protons we have.
- Integrals tells us the ratio of each kind of proton in our sample.
- ¹H ¹H coupling tells us about protons that are near other protons.



¹³C NMR

☞ Chemical shift is normally 0 to 220 ppm

Similar factors affect the chemical shifts in ¹³C as seen for H NMR

IF Long relaxation times (excited state to ground state) mean no integration

IF Number of peaks indicates the number of types of C



Types of NMR experiments



- COSY (Correlation Spectroscopy)
- -TOCSY (Total Correlation Spectroscopy)
- NOESY (Nuclear Overhauser effect spectroscopy)

 $^{1}\mathrm{H} - ^{1}\mathrm{H}$

 $^{1}H - ^{13}C$

- HSQC (Heteronuclear Single Quantum Coherence)
- HMBC (Heteronuclear Multiple Bond Coherence)

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DNA



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Structure Activity Relationships (SAR) by NMR

• Medicine - Magnetic Resonance Imaging (MRI)

Structure Determination

• Each observable NMR resonance needs to be assigned or associated with the atom .

- Molecular Formula
- Functional groups
- Carbon Connectivity
- Position of substitution on the carbon framework
 - Stereochemical properties.

NMR of carbohydrates:

Sugar assignment (COSY, TOCSY, HSQC, HMBC)

Anomeric configuration (³J_{H,H} coupling constants, NOE)

Ring size (¹³C chemical shift, HMBC)

Non carbohydrate substituents (Homo- and Heteronuclear NMR)

Sugar sequence: Linkage analysis and glycosylation pattern (HMBC, NOE)

¹H and ¹³C typical regions of carbohydrates:

The ¹H NMR Spectra can be <u>roughly</u> divided into the following regions:

Anomeric and Acylated Protons : 5.5-4.5 ppm. Ring Protons : 4.5-3 ppm Acetyl Groups, Methylene Protons: 3-2 ppm Methyl Groups: 0.8-2.0 ppm

The ¹³C NMR Spectra can be <u>roughly</u> divided into the following regions :

Anomeric Carbons Resonate Between 90-105 ppm Ring Carbons Between 60-85 ppm Nitrogen Bearing Carbons (In Amino Sugar) 50-60 ppm Acetyl Groups 20-25ppm Methylene Protons: 25-35 ppm Methyl Groups: 15-20ppm





Anomeric configuration

¹H NMR spectrum contains information on the configuration of glycosidic linkages







 β -linkage: $J_{1,2} < 2$ Hz



C

β-linkage: $J_{1,2} > 6$ Hz α-linkage: $J_{1,2} < 2$ Hz

α-linkage: ¹J_{C1,H1} 170-175 Hz β-linkage: ¹J_{C1,H1} 160-165 Hz



Anatomy of a 2D NMR Experiment



2D NMR - The Interferogram



A 2D data set can be thought of as a series of 1D . Each 1D file is different from the next by a change in t_1 .

Fourier transformation of each 1D in the t2 domain creates an interferogram.



Two Dimensional NMR

✓A 2D data set can be thought of as a series of 1D experiments collected with different timing.

✓ Fourier transformation of each 1D in the t2 domain creates an interferogram.

✓The ti domain is then Fourier transformed resulting in a 2D file with the frequency in each dimension.

✓ This 2D file will provide a map of all spin-to-spin correlations

✓ Each 2D experiment can provide either through bond (COSY type) or through space (NOESY type) correlation

COrrelation SpectroscopY (COSY)

In a 2D COSY spectrum, cross-peaks will exist where there is spin-spin coupling between nuclei.



Used to identify spins which are coupled to each other.

2D Experiments – COSY

The Power of 2D NMR: Resolving Overlapping Signals



2D Experiments – COSY



Sugar assignment



TOtal Correlation Spectroscopy (TOCSY) experiment

TOCSY Experiment

In general, the TOCSY mixing time determines the number of bonds over which signal can be Transferred, assuming that none of the coupling Constants = 0



•Cross peaks generated between all members of a coupled spin network



TOCSY

Cross peaks are generated between all members of a coupled spin network
Coherence transfer period occurs during a multi-pulse spin-lock period;
Length of spin-lock and J-coupling constants determine how far the spin coupling network will be probed


COSY and TOCSY – Sugar assignment



In Glucose, H1 and H2 protons are scalarly coupled, H1 and H3 are not. In COSY spectra → H1 and H2 correlation observed ; In TOCSY spectra → H1 and H3 observed

COSY and TOCSY – Sugar assignment



HSQC: Heteronuclear Single-Quantum Correlation



The spectrum contains a peak for each unique proton attached to the heteronucleus being considered.

The 2D HSQC experiment permits to obtain a 2D heteronuclear chemical shift correlation map between directly-bonded ¹H and X-heteronuclei (an atomic nucleus other than a proton), often ¹³C or ¹⁵N.

HSQC: Heteronuclear Single-Quantum Correlation

¹H–¹⁵N HSQC spectrum of a fragment of the protein NleG3-2. Each peak in the spectrum represents a bonded N-H pair, with its two coordinates corresponding to the chemical shifts of each of the H and N atoms. Some of the peaks are labeled with the amino acid residue that gives that signal



HMBC (Heteronuclear Multiple Bond Correlation)

2D HMBC experiment correlates chemical shifts of two types of nuclei **separated from each other with two or more chemical bonds.**



HSQC and HMBC of Menthol



Linkage analysis - Monosaccharide Sequence

Long range *inter*-residual correlations in the HMBC spectrum





Nuclear Overhauser Effect (NOE) Spectroscopy

The 2D spectrum will have chemical shifts in f1 and f2.

The cross peaks are for nuclei that are coupled dipolarly.



NOE contact: <u>C is close in space to spin A</u>

The NOE effect is the method for the elucidation of 3D structural features and stereochemistry



NOE and Distances Isolated spin pair aproximation (ISPA)



Intra-residue NOE contacts in monosaccharides: relative configuration of sugar residues

gluco, galacto configuration



α-linkage: H1/H2



β-linkage: H1/H3, H1/H5

manno configuration



α-linkage: no contact



β-linkage: H1/H2, H1/H3, H1/H5

Monosaccharide Sequence

Inter-residue NOE Glycosylation shift (HSQC spectrum) Inter-residual long range correlation (HMBC spectrum)

Inter-residue NOE contacts in saccharides (Linkage analysis)



 β -(1-3) linkage

α-(1-3)linkage



Sugar sequence - Linkage analysis

NOE in disaccharides may occur not only at the linkage protons but also at the neighbouring protons



....Saccharide conformation...

Sugar sequence - Linkage analysis

Sucrose NOESY and TOCSY



Application of various NMR techniques to carbohydrates

•HOMONUCLEAR (¹H-¹H) •HETERONUCLEAR (¹H-¹³C)



A BACTERIUM'S ENEMY ISN'T YOUR FRIEND

a Strong immune response against WTA

b Weak immune response against WTA





Nature 2019



MOLECULAR INTERACTION by NMR

The ligand-based approach

Representation of protein-ligand interactions

> STD NMR Protein TrNOESY Receptor WaterLOGSY Relaxation experiments Diffusion experiments Ligand Conformational Flexibility ➢ Other nuclei: ¹⁹F Paramagnetic tagging \succ Other methods The receptor based approach Isotope labelling Chemical shift perturbation mapping Paramagnetic tagging Other variations

Ligand observation





Two states equilibrium L_{free} L_{bound} Molar fractions

 $R_{Lobs} = L_f R_{Lf} + L_b R_{Lb}$ $\Delta R = L_b (R_{Lb} - R_{Lf})$

Experimental procedure: $L_0 >> R_0$; $L_0/R_0 > 10 - 100...$ $L_f >>> L_b$

Necessary condition: $|(R_{Lb} - R_{Lf})| >> 0$ R_{Lb} Strong dependency on molecular size NMR observable parameter R : NOE; Diffusion; Line Shape

Recapitulation

- K_{off} fast in the relaxation time scale, dissociation must occur before relaxation.
- K_{on} related to the efficacy of the interaction.
- Consider the molar fraction of ligand free and bound to the protein.
- Fast exchange in the chemical shift time scale
- L₀>>R₀ means excess of free ligand but since we are in conditions in which the exchange is high (rate) the system is dominated by the bound state
- K_{off} Dissociation before relaxation takes place; dissociation rate high since the molecule relax in the binding site

TRANSFERRED NOE

Information on the ligand bioactive conformation



During the mixing time *inter* and *intra*-molecular NOE effects build up
 Inter-molecular tr-NOE effects are visible, intermolecular trNOEs are usually much larger than intramolecular effects

The mixing time must be short enough so that the contribution of the free ligand is negligible and long enough to allow visualization of the signal in the spectrum.

Important notes

The molar ratio of ligand to receptor. It should be emphasized that the trNOESY experiment works well for ligands that have K_D in the range $10^{-3} - 10^{-6}$ M / mM- mM range

Small amount of purified receptor

Routinely used to probe ligand-receptor interaction

TRANSFERRED NOE and MOLECULAR MOTION



The bioactive conformation: Transfer NOESY



Is There Any Binding?Which Is The Ligand Bioactive Conformation?

Saturation Transfer Difference NMR Spectroscopy – STD NMR



At long irradiation times, the saturation is transferred to the bound ligand, first to the protons belonging to the ligand epitope, then to the rest of the ligand

(Meyer and Mayer, Peters, 1999, 2000)

Schematic representation of STD NMR method.



H close to receptor

- H far from receptor
- H furthest away from receptor

STD (Meyer and Mayer, Peters, 1999, 2000)

Key elements of protein-substrate binding

RECAPITULATION...

- The saturation is transferred first to the protons belonging to the ligand epitope, then to the rest of the ligand
- STD involves selective perturbation of protein-specific methyl proton resonances (0.5 p.p.m.-1 p.p.m.). This perturbation rapidly diffuses throughout the protein.
- This is usually done via a selective 180-degree pulse and results in a transfer of magnetization from the protein to any transiently bound ligands via the nuclear Overhauser effect (NOE).
- If the fragment binds to the target protein, the buildup of NOE that is transferred to the ligand results in enhanced signal corresponding to the resonances of that ligand in the STD spectrum.
- A number of factors affect the signal strength in an STD experiment, including protein size, duration of on-resonance irradiation, the frequency of irradiation, the dissociation constant of the ligand, the ligand/protein ratio and the field of the spectrometer.
- As such, it is useful to vary parameters such as ligand and protein concentration, the frequency of irradiation and the duration of irradiation

NMR Spectroscopy Techniques for Screening and Identifying Ligand Binding to Protein Receptors



 β -Gal-Ome and RCA120 (Ricinus communis agglutinin), 50 μ M and 1.2 mM, 1:40

M. Mayer, B. Meyer, J. Am. Chem. Soc. 2001;123(25):6108-17; Bernd Meyer, Thomas Peters, Angewandte Chemie International Edition, 42, 8, 864-890

NMR Spectroscopy Techniques for Screening and Identifying Ligand Binding to Protein Receptors

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In order to determine the magnitude of the STD effects, the intensity of the signal in the STD NMR spectrum are compared with the signal intensities of a reference spectrum (off-resonance). The STD signal with the highest intensity is set to 100% and the others are normalized to this signal.

M. Mayer, B. Meyer, J. Am. Chem. Soc. 2001;123(25):6108-17; Bernd Meyer, Thomas Peters, Angewandte Chemie International Edition, 42, 8, 864-890

Epitope mapping for sialyl LewisX bound to the the lectin Aleuria aurantia agglutinin (AAA)



STD TOCSY spectrum of sialyl LewisX in the presence of AAA (molar ratio 100:1): only the spin system of the I-fucose residue (F, yellow circle) is visible

1D and 2D STD spectra shows unambiguously that only the fucose interact with the protein

M. Mayer, B. Meyer, Angew.Chem. 1999, 111, 1902 – 1906 ;Angew.Chem.Int.Ed. 1999, 38, 1784 – 1788.

Water-LOGSY NMR



The resonances of non-binding compounds appear with opposite sign and tend to be weaker than those of the interacting ligands.

Experimentally, the first step is the selective water excitation; during the mixing time (to be optimized based on the size of the complex) the water magnetization that has migrated to the protein is transferred to the ligand via direct of relay processes.

Carr Purcell Meiboom Gill (CPMG) experiments

CPMG, is a relaxation-time-edited NMR experiment that exploits differences in transverse relaxation time (*T2*). *Proteins (and bound ligands)* have a small *T2 while free ligands have a large T2. Thus monitoring T2, binding* can be detected when the signal of the ligand decreases.



Carr, Y.H. & Purcell, M.E. Phys. Rev. 94, 630–638 (1954).; Hajduk, P.J., Olejniczak, E.T. & Fesik, S.W. J. Am. Chem. Soc. 119, 12257–12261 (1997).; E.H Mashalidis, P. Śledź, S. Lang, C. Abell, Nature Protocols, 8, 2309–2324 (2013)

Siglecs: <u>sialic-acid</u> binding immunoglobulin-like lectins

SIGLECS – SIALIC ACID AXIS



H-CD22- sialoglycans studies



ChemBioChem, 2019



*Kozmiński W, et al. J Magn Reson. 2000, 142, 294-299.

ChemBioChem, 2019
Siglecs-Glycan axis as attractive therapeutic target

Siglecs interaction with pathogens



Dynamic changes of structures and composition of microbial cell envelope glycans, their **camouflage with host selfantigens (molecular mimicry), their cross-talk with host immune Siglecs**, is central in GLYTUNES. The understanding of the role of sialic acid–Siglec interactions in the context of pathogen-associated immune suppression will pave the way to the **development of novel therapeutics glycomimetic to be used in clinical and diagnostic applications**

