# Mass Spectrometry from observation to structural information

Marc-André Delsuc Renafobis - 2017











Emerging mass spectrometry (MS) methods for studying membrane protein complexes and their relationship to other biophysical techniques

from N.Barrera, C.Robinson Annu. Rev. Biochem. 2011. 80:247–71

### PLAN

- Principles of Mass Spectrometry
  - measuring m/z
  - ionisation methods
  - shape of a MS spectrum of a protein
  - fragmentation methods
- Examples of use in Structural Biology
  - Large multicomponent Complexes
  - Ligand binding
  - H-D exchange and other chemical labelling
  - Cross-Linking
  - Ionic Mobility
- 2D MS
  - ... teasing you ....

### Principles of Mass Spectrometry

- a charged molecule
- a fly in the vacuum
- the trajectory in inflected  $\overrightarrow{B}$   $\overrightarrow{E}$
- a detector senses the ion
- the sensing allows measuring the molecular mass





- We need ions in vacuum
- 3 fundamental steps
  - ionisation / separation / detection
- Electrostatic/electrodynamic interactions
  => we measure ONLY *m/z*
  - not just m
- m unit = 1 Dalton : 1atom-gram
  - definition 1/12 mass of <sup>12</sup>C atom
  - 1 Da = 1.66 10<sup>-27</sup> kg
- m/z unit : 1 Thomson = 1Da /  $1e^{-1}$

### Detection

- There is a large range of approaches for separating ions, in all cases:
  - the ion in vacuum flies in E and B fields
    - homogeneous or varying in space
    - static or varying in time
  - E and B field apply forces to the ion, proportionnal to the charge : z
  - the ion follows Newton law depend on the mass m
  - the displacement is dependent on the mass and the charge
  - only m/z can be measured
- different measurement methods
  - sector instruments
  - Time Of Flight
  - Quadrupole
  - Ion Trap
  - Orbitrap
  - Ionic Cyclotronic Resonance

### Equations of motion

• Electrostatic force  $\vec{F}_e = q\vec{E}$ 

• Lorentz force  $\vec{F}_L = q\vec{v} \wedge \vec{B}$ 

• Ponderomotive force  $\vec{F_p} = -\frac{q^2}{4m\omega^2}\nabla E^2$ 

• Newton law  $\vec{F} = m\vec{\gamma}$ 

### sector instrument

- Principle
  - simplest
    - deflection by a homogeneous, static E or B field
    - m/z is measured by the position of the impact
  - improved
    - deflection by a homogeneous, time varying E or B field
    - m/z is measured by the time of the impact at a given point
  - both field can be used to imp
  - not really used any more



### Paul trap

$$\phi_o = U + V \cos \Omega t$$

$$\phi_{r,z} = \frac{\phi_o}{r_o^2} (r^2 - 2z^2)$$



Wolfgang Paul - 1989 Physics Nobel Prize

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### lon trap

- unstable trajectory in a fluctuating E field
  - stable trajectories are trapped into a cell
  - some trajectories are unstable (depending on m/z)
  - sweeping the frequency ejects ion relative to m/z value
- in practical
  - Allows storing ions for some time
  - resolution is not very high





### Quadrupole

- resonant trajectory in a fluctuating E field
  - the frequency of the E determines m/z of the stable trajectory
  - sweeping the frequency
- improvement
  - longer quadrupoles
  - higher tensions
  - hexapoles



# Time Of Flight

- Principle
  - ions are accelerated by E field
  - ions are injected at a given time (MALDI)
  - m/z is measured by the time it takes to reach the detector



Zone d'accélération

Détecteur linéaire



Zone de vol, libre de champ

- improvements
  - The longer the path, the higher the resolution
  - refocalisation of different energies
    - Use of reflection chamber

### Orbitrap

- Stable ion orbits into a Electrostatic cell
  - static Eo
  - all ions are measured at the same time
  - orbit frequency depends on m/z
  - Fourier Transform gives frequency, thus m/z

### In practical

- very high resolution and sensitivity
- speed, sensitivity and resolution de
- requires very high vacuum
- patented by ThermoFisher

$$\frac{m}{z} \propto \sqrt{\frac{1}{f}}$$

Spectra obtained by Fourier Transform





**Supplementary Figure 1.** Schematic of the modified Exactive Plus instrument (ThermoFisher Scientific, Bremen, Germany) with HCD option.



Pening Trap







**FIGURE 7.** Incoherent ion cyclotron orbital motion (top left) is converted to coherent (and, therefore, detectable) motion (top right) by the application of a rotating electric field, which rotates in the same sense and at the ICR frequency of the ions of a given m/z value. The electronic circuitry is shown in the bottom diagram.

#### FOURIER TRANSFORM ION CYCLOTRON RESONANCE MASS SPECTROMETRY: A PRIMER

#### Alan G. Marshall, \*\* Christopher L. Hendrickson, and George S. Jackson\*

Center for Interdisciplinary Magnetic Resonance, National High Magnetic Field Laboratory, Florida State University, 1800 East Paul Dirac Dr., Tallahassee, FL 32310

Received 7 January 1998; revised 4 May 1998; accepted 6 May 1998

## FT-ICR

- Stable ion orbits into a magnetic field
  - static homogeneous Bo
  - all ions are measured at the same time
  - orbit frequency depends on m/z
  - Fourier Transform gives frequency, thus
- In practical
  - very high resolution and sensitivity
  - speed, sensitivity and resolution depend on value of Bo
  - requires very high vacuum
  - high Bo requires cryomagnet

$$\frac{m}{z} \propto \frac{1}{f}$$

Spectra obtained by Fourier Transform







### Chirp pulses



- broad-band spectra
  - ▶ 50kHz 1MHz
- Direct detection
  - no carrier
- Chirp pulses
  - t=0 not easy to defined
  - complex phase dependence
- Simulation needed
  - Lorentz + Newton

#### Evolution simulator

Swept pulse Frequency : 200.000-50.000 kHz sweep width : 150.000 kHz sweep steps : 1000 duration : 1.000 msec Epp : 1666.67 V/m approx excitation radius : 11.82 mm

resonant frequency 144151.41 Hz final radius : 7.56 mm



#### 700mina in a FT-ICR data-set

169





### Ionisation methods

• of course elephants can fly











John Fenn Koichi Tanaka Kurt Wüthrich Nobel prize in Chemistry 2002

### MALDI : Matrix Assisted Laser Desorption/Ionisation

### Matrix

absorbs light energy (UV laser)

- ionizes the molecule without breaking it
- Typically
  - cinnamic acid
  - but also

Ferrulic acid / Sinapic acid / DiHydroxy Benzoic acid / etc..









### ESI : Electro Spray Ionization





### ESI : Electro Spray Ionization





### ESI requirements

#### volatile buffers

so that no salt remains on the molecule of interest

- positively charged ammonium
- negatively charged
  - carbonate
  - formiate
  - acetate
- and that's about it !

#### see for instance

- http://www.rsc.org/suppdata/an/c1/c1an15123a/c1an15123a.pdf

### all together : a MS spectrometer

- Combine :
  - Usually some Chromatography
    - ► GC
    - ► HPLC
    - nanoLC
    - capillary electrophoresis
    - nothing => infusion
  - one source
    - ESI
    - MALDI
    - etc..
  - one measurement
    - trap / quadrupole
    - ► TOF
    - Orbitrap

- Hence
  - ESI-Orbitrap
  - Maldi-TOF
  - MALDI-LTQ
  - GC-TOF





• Whisky sample

### Resolution

- Resolution in MS
  - depends on several aspects
  - measured by the ratio R
- Isotopic patterns
  - monoisotopic mass / average mass
- Ionisation techniques
  - Multicharges patterns
    - difference MALDI / ESI
- Detection techniques
  - MS-MS < TOF LTQ Quad < Orbitrap FT-ICR</p>
    - 1.00010.000 100.000

 $\mathcal{M}$ R =

100.000 - 1.000.000

 $rac{m}{\Delta m}$ 

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### Resolution

- Resolution in MS
  - depends on several aspects
  - measured by the ratio R



- Detection techniques
  - Ion Trap < Quadrupole < TOF < Orbitrap < FT-ICR</li>
    1.000 5k 20k 50k 300k 1M 5M

### Isotopic pattern

- Each atom type displays an isotopic profile
  - list of isotope natural abundance for common atoms

isotopic ratio		+1	+2
н	99.99%	0.015%	
С	98.9%	1.1%	
N	99.63%	0.37%	
0	99.76%	0.038%	0.2%
Р	100%		
S	95.02%	0.75%	4.21%

http://www.sisweb.com/referenc/source/exactmaa.htm http://www.sisweb.com/mstools/isotope.htm
## Molecular Mass definition - Isotopic pattern

- For Cn peak at 12n+1 is proportionnal to n
- example here for linear alcanes C<sub>n</sub>H<sub>2n+2</sub>
  - monoisotopic mass / average mass



## Isotopic pattern

- Protein empirical formula
  - eg : C<sub>1</sub> H<sub>1.59</sub> N<sub>0.27</sub> O<sub>0.31</sub> S<sub>0.01</sub> (different expressions exist)
  - The aspect of the pattern depends on the resolution (and on the charge state)
  - here simulated for a 11kD protein
  - note how monoisotopic mass  $\neq$  average mass  $\neq$  top of the



#### C\_725 H\_1122 N\_194 O\_215 S\_10

16306.003396 0.0544783371 16307.006259 0.4815908078 16308.008995 2.1746869097 16309.011617 6.6792852462 16310.014141 15.6785827113 16311.016578 29.9699795375 16312.018937 48.5483260540 16313.021227 68.4889810809 16314.023455 85.8283420627 16315.025629 96.9892347469 16316.027753 100.000000000 16317.029832 94.9628506636 16318.031872 83.7024337398 16319.033877 68.9202526525 16320.035849 53.3017111235 16321.037793 38.8992040048 16322.039711 26.8962383101 16323.041605 17.6814074296 16324.043479\_11.0855202359 16325.045335 6.6465388834 16326.047174 3.8202539089 16327.048999 2.1095643109 16328.050811 1.1213685311 16329.052610 0.5748121676 16330.054400 0.2845871917 16331.056178 0.1362794827 16332.057943 0.0631972378 40 50

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100

80

60

40

monoisotopic

0

0

10

20

30

m/z

intensity



m/:

100

80

60

40

20

82 84 86 88 90 92

inter





P. Lössl, M. van de Waterbeemd & A. JR Heck

The EMBO Journal (2016) 35: 2634–2657

#### Prop to length $\Rightarrow$ to MW





# Charge/Mass dependence

- unfolded proteins
  - charge depends only on primary sequence and pH
- folded proteins
  - charge is only on the surface
  - Rayleigh model Z<sub>R</sub>
    - charges are in the droplet when sprayed
    - droplet evaporate by coulombic fission
    - protein is assumes folded ad spherical, charges on the protein are at the limit of the maximum coulombic density.



Number of observed mean charges of a number of globular proteins and protein complexes compared with the Raleigh limit model predicted charge. The number of observed charges is very close to the Raleigh limit on water droplets of the same size as the protein. All proteins were sprayed from 50 mM ammonium acetate at neutral pH. (O) Represents positive ion mode and (•) negative ion mode.

> $10kD \implies Z \sim 9+$   $100kD \implies Z \sim 28+$  $1MD \implies Z \sim 90+$







**FIGURE 22.** ESI FT-ICR mass spectra of chondroitinase I. Bottom: Heterodyne data for SWIFT-isolated ions, 1226 < m/z < 1273, with external ion accumulation (Senko et al., 1997), from 10 co-added time-domain signals; the peaks at m/z 1240 and 1254 correspond to an unidentified adduct of ~260 Da. Top: Mass scale-expansion showing unit mass resolution of the isotopic distribution of the z = 91 charge state. Data kindly provided by N. Kelleher and described in detail elsewhere (Kelleher et al., 1997).

## Charge depend on protein state



http://msr.dom.wustl.edu/tutorial-native-mass-spectrometry/

# Examples of use in Structural Biology

- Not proteomics!
- Structural Information
  - non covalent molecular interactions
    - Large multicomponent Complexes
    - Ligand binding
  - H-D exchange and other chemical labelling
  - Cross-Linking
  - Ionic Mobility
  - fragmentation



from E. Boeri Erba C. Petosa

*PROTEIN SCIENCE* 2015 VOL 24:1176-1192



## Denatured and native mass spectra of *H. pylori* urease.

(a) Denatured urease was electrosprayed from an aqueous 50% (vol/vol) acetonitrile containing 0.1 % (vol/vol) formic acid solution revealing individual charge distributions from the multiply charged  $\alpha$  (26.6 kDa, orange) and  $\beta$  (61.7 kDa, magenta) monomers of urease. (b) A mass spectrum of native urease electrosprayed from an aqueous ammonium acetate solution (bottom) displaying multiple ion signals that originate from multiple charged species of the  $\alpha_{12}\beta_{12}$  intact urease machinery with

a measured mass of 1,063.4  $\pm$  1.0 kDa. Insets are close-ups of the indicated regions. The cartoons are adapted from the X-ray structure of the intact  $\alpha_{12}\beta_{12}$  urease.

Albert J R Heck Nature Methods 5, 927 - 933 (2008) doi:10.1038/nmeth.1265

Native mass spectrometry: a bridge between interactomics and structural biology

## \_arge Complexes by MS /with Orbitrap



Orbitrap-based mass spectra of intact proteins and protein assemblies. (a–d) Native mass spectra of IgG antibody (a), bacteriophage HK97 capsid pentamers and hexamers (b), yeast 20S proteasome (c) and E. coli GroEL (d). Illustrative crystal structures are shown for each protein. Inset in a shows an enlargement of the 25+ charge state of IgG1.

Rose, R. J., Damoc, E., Denisov, E., Makarov, A. & Heck, A. J. R. High-sensitivity Orbitrap mass analysis of intact macromolecular assemblies. *Nat Meth* **9**, 1084– 1086 (2012).

# ESI & interactions in the vacuum

#### sequentially

- Ioss of water molecules
- Ioss of hydrophobic interactions

- Ioss of Van der Walls interactions
- weakening of electrostatic interactions

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K. Breuker and F. W. McLafferty, Proc. Natl. Acad. Sci. U. S. A., 2008, 105, 18145–18152

## Large Complexes

#### stabiliy collisions in source

Pi (mbar)



Pacholarz et al Chem. Soc. Rev., 2012, 41, 4335-4355



**Supplementary Figure 8**. Tandem mass spectrum using HCD activation of the 14-subunit GroEL precursor ions. This asymmetric charge/subunit dissociation pathway, as shown in the cartoon, is typical for gas-phase dissociation of non-covalently bound protein complexes by collisional activation. 13-subunit GroEL fragment ions are detected at m/z values up to and above 20,000 Th.

## Protein-Ligand

studying protein-ligand interactions in native mode



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## 185 - salivary protein

#### • IDP - low complexity

#### SPPGKPQGPPQQEGNKPQGPPPPG KPQGPPPAGGNPQQPQAPPAG KPQGPPPPQGGRPPRPAQGQQPPQ

• Interact selectively with polyphenols (proanthocyanidines)



#### **IB5 / TANNINS**



→ Formation de complexes IB5:EgCG avec différentes stoechiométries (de 1:1 à 1:5)

Francis Canon / Véronique Cheynier

#### **IB5 / TANNINS**

Impact de la structure des tanins sur l'interaction collision contre un gaz neutre depend de la vitesse d'accélération



	IB5-EgC	IB5-ECG	IB5-EgCG	IB5-B2	IB5- B2.3'OG
E <sub>50</sub> (eV)	67.5	105	120	150	180
E <sub>cm</sub> (eV)	0.26	0.4	0.45	0.55	0.65

Francis Canon / Véronique Cheynier

#### **IB5 / TANNINS**

#### Expériences de MS/MS: cas des complexes IB5:tanin 1:2

Exemple de l'hétérocomplexe IB5:(B2/ECG)



#### → Les deux tanins ne sont pas libérés en même temps

Francis Canon / Véronique Cheynier

## H / D exchange



S. R. Marcsisin and J. R. Engen Anal Bioanal Chem. 2010 June ; 397(3): 967–972. doi:10.1007/s00216-010-3556-4.

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Fig. 9 Scheme of on- and off-exchange approaches used in HDX experiments. Reproduced from A. Sinz, 2007<sup>5</sup> with permission from John Wiley and Sons.





Bruning JB, Chalmers MJ, Prasad S, et al.. Structure. 2007; 15(10):1258–1271.

View Article Online



**Fig. 12** Analytical strategy for analysing conformational changes in protein upon ligand binding by chemical cross linking and high resolution mass spectrometry. Reproduced from Muller and Sinz<sup>180</sup> with permission from Springer.





#### Pyruvate kinase (KPYM\_RABIT, P11974), PDB structure 2G50DSSPDHZL



A. Leitner, L.A.Joachimiak, P. Unverdorben, T.Walzthoeni, J.Frydman, F.Förster, and R.Aebersold

PNAS 111 26 9455–9460

vinculin forms hybrid complexes with components of the Arp2/3 actin polymerization complex











from N.Barrera, C.Robinson Annu. Rev. Biochem. 2011. 80:247–71

# Tandem MS : MS-MS

- Collision ΕI Photon Tandem MS : coupling ESI Surface MALDI the first MS select one m, a fragmentation is applied m/z 3 fragmen ionization detectior eparation eparation • the mass spectrum of the MS1 MS2 Precursor Product ion ion
- Several fragmentation techniques available Mostly :
  - CID : Collision Induced Dissociation collision with a neutral gaz : eg Argon
  - IRMPD : IR Multiple Photon Dissociation irradiation with a IR laser
  - ECD : Electron Capture Dissociation bombardment with e-
  - ETD : Electron Transfert Dissociation transfert of e- by collision with charged molecules

## Tandem MS : MS-MS

- Tandem MS : coupling two MS measure in series.
  - the first MS select one m/z
  - a fragmentation is applied to the parent peak
  - the mass spectrum of the gradients is determined
- Several fragementation technique available
  - Mostly :
  - CID : Collision Induced Dissociation
    - collision with a neutral gaz : eg Argon
    - SID HCD
  - ETD : Electron Transfer Dissociation
    - bombardment with e-
    - EDD ECD
  - IRMPD : IR Multiple Photon Dissociation
    - irradiation with a IR laser
    - BIRD

# MS-MS : several possible geometry

- coupling 2 MS plus a dissociation chamber
  - QqQ (Q3)
- but also
  - Q-TOF
  - Q-Orbitrap
  - TOF-TOF



### proteomics



Fig. 1 Comparison of top-down and bottom-up workflows.

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## Fragmentation

• fragmentation and identification from databases



P. Lössl, M. van de Waterbeemd & A. JR Heck

The EMBO Journal (2016) 35: 2634–2657

## Directly Obtaining Both Proteomics and Structural Information by Native Top-Down MS?

- Native MS for macromolecular complexes
- Top-down capability
  - High resolution
  - High mass accuracy
  - Multiple fragmentation techniques



solarıx 15-Tesla



From Huilin Li

#### Native Top-Down MS — Harvest the synergy between proteomics and native MS



Li, H. et al. J. Am. Soc. Mass Spectrom. 2014. 25, 2060-2068.

## All Information in One Experiment





Overall, 40% sequence coverage

Li, H. et al. J. Am. Soc. Mass Spectrom. 2014. 25, 2060-2068.
#### ECD Reveals Outer Surface Residues of Aldolase Tetramer (158 kDa)



PHSHPALTP	EQKKELSDIA	HRIVAPGKGI	LAADESTGSI	AKRLQSIGTE
NTEENRRFYR	QLLLTADDRV	NPCIGGVILF	HETLYQKADD	GRPFPQVIKS
KGGVVGIKVD	KGVVPLAGTN	GETTTQGLDG	LSERCAQYKK	DGADFAKWRC
VLKIGEHTPS	ALAIMENANV	LARYASICQQ	NGIV PIVEPE	I L P D G D <mark>H D L K</mark>
RCQYVTEKVL	A A V Y KA L S <mark>(</mark> D H	H I Y LEGTL <mark>L</mark> K	Р <mark>ММV</mark> ТРGНАС	TOKYSHEEIA
MAT VTALR	<u> γ Ρ ΡΑΥ ΤΙΘΥΤΙ</u> Ε	LEGGQSEEEA	SINLINA INKC	Ρ Ι[Ι[ΚΡΨ[Α][ Τ[F
SYGRALQABA	LKAWGGKKEN		RALANSLACQ	
			• •	•••

Li, H. et al. Anal. Chem. 2014. 86, 317-320

**Membrane proteins** 

**Revealing Structural Similarity upon Metal Binding** 



- Cu (His 46, 48, 63 and 120)
- Zn (His 63, 71, 80, and Asp 83)
- Disulfide bond (Cys57 and Cys146)



Green: Apo-WT SOD1; Cyan: Zn-SOD1; Purple: Cu,Zn-SOD1

**Revealing Structural Difference upon Metal Binding** 



#### Cu Binding Shields the Charged Residues (His46, His48)



Li, H. et al. Anal. Chem. 2017. 89, 2731-2738

#### Native Top-Down MS of β-Galactosidase Tetramer (465 kDa)



N-terminal tail in cyan

**ECD**: no c/z<sup>•</sup> ions (N- an C-termini are involved in interfaces)

**IRMPD**: 42% sequence coverage from the C-terminal, no PTMs observed

**CAD**: 12% sequence coverage from the N-Terminal; N-terminal is highly modified; at least four proteoforms



Li, H. et al. Nat. Chem. 2017. Under review

**Membrane proteins** 

#### **Reveals Empty Nanodisc Composition**

➤ Empty nanodisc is about 130~190 kDa

- > 2 scaffold proteins (22.45 kDa)
- > 125~214 DMPCs (677.5 Da)



I Campuzano, **H Li**, et al. *Anal. Chem.* **2016.** 88, 12427-12436.

#### **Release Membrane Protein Complex from Nanodisc**



2D-MS



## NMR - MS what's in common ?





NMR



Jean Baptiste Joseph Fourier (21 Mars 1768 – 16 Mai 1830)

 $f(t) : \mathbb{R} \to \mathbb{C}$  $F(\nu) : \mathbb{R} \to \mathbb{C}$ 

 $\mathcal{F}: f(t) \stackrel{\mathcal{F}}{\mapsto} F(\nu)$ 

$$F(\nu) = \int_{-\infty}^{\infty} f(t) e^{-2i\pi\nu t} dt$$

## Fourier Transform





#### Author's personal copy





- HI et H2 : aeux pnenomenes lles différents battements entre 2 oscillatet
- possible car retour à la situation initiale
- t1 : durée / t2 : date
- peu de contraintes sur le choix des valeurs de t1





M-A Delsuc Thèse d'État 1985 - Univ. Paris XI Volume 138, number 2,3

CHEMICAL PHYSICS LETTERS 1987



#### TWO-DIMENSIONAL FOURIER TRANSFORM ION CYCLOTRON RESONANCE MASS SPECTROMETRY

#### Peter PFÄNDLER, Geoffrey BODENHAUSEN

Institut de Chimie Organique, Université de Lausanne, Rue de la Barre 2, CH-1005 Lausanne, Switzerland

#### Jacques RAPIN, Raymond HOURIET and Tino GÄUMANN

Institut de Chimie Physique, Ecole Polytechnique Fédérale, CH-1015 Lausanne, Switzerland





#### 1988

P Pfaendler, G Bodenhausen, J Rapin, M Walser, T Gaümann

Broad-band two-dimensional Fourier transform ion cyclotron resonance.

JAm Chem Soc (1988) vol. 110 (17) 5625-5628



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Pfaendler, P., Bodenhausen, G., Rapin, J., Walser, M. E., & Gäumann, T. (1988). *J.Am.Chem.Soc.*, **110**, 5625-5628. van Agthoven, M. A., Delsuc, M.-A., Bodenhausen, G. & Rolando, C. (2013) *Anal Bioanal Chem* **405**, 51–61.

#### Principle of 2D FT-ICR

author's personal copy



#### exploration of the 2D map



#### Author's personal copy



### <u>Top-down proteomics</u>



#### Calmoduline

- 2D FT-ICR IRMPD
  - ▶ 512 x 4M = 2 Gpoints
  - > 20 min. acquisition
  - ▶ R1 ~ 180
  - ▶ R2 ~ 420.000



Floris, F., van Agthoven, M. A., Chiron, L., Soulby, A., Wootton, C. A., Lam, P. Y., Barrow, M.P., Delsuc, M-A., O'Connor, P. (2016). *J. Am. Soc. MS*, *27*(9), 1531–1538

#### <u>Top-down proteomics</u>



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#### Top-down proteomics

## Calmoduline 2D fragment ion scan at 14+



### Bottom-up proteomics

• Calmoduline

- 2D FT-ICR IRMPD
  - ▶ 4096 x 512k = 2 Gpoint
  - ▶ 50 min. acquisition
  - ▶ R1 ~ 1200
  - ▶ R2 ~ 60.000



#### om-un nrotec

Collage protein

Université de Strasbou

- compléx protein
  - heavyly transformed: hydroxy-proline / hydroxy-lysine
  - packed helices and dense Hbound network
  - Bovin  $\widehat{E}$  Collagen Type1:  $\alpha$ 1 and  $\alpha$ 2 chains (~2x 1400 aa)
  - tryptig digest

Blank regions are low complexity, with repetitic of X-Y-Gly pattern

a1-[1] MFSFVDLRLL LLLAATALLT HGQEEGQEEG QEEDIPPVTC VQNGLRYHD DNGNVLC DDVICDELKD CPNAKVPTDE EGPKGDTGPR GPRGPAGPPG RDGTPGOPGL POLSYGYDEK STGISVPGPM GPSGPRGLPG PPGAPGPOGF OGPPGEPGEP GASGPMGPRG PPGPPGKNGD DGEAGKPGRP GERGPPGPOG ARGLPGTAGL PGMKGHRGFS GLDGAKGDAG PAGPKGEPGS PGENGAPGQM GPRGLPGERG RPGAPGPAGA RGNDGATGAA GPPGPTGPAG PPGFPGAVGA KGEGGPQGPR GSEGPQGVRG EPGPPGPAGA AGPAGNPGAD GQPGAKGANG APGIAGAPGF PGARGPSGPQ GPSGPPGPKG NSGEPGAPGS KGDTGAKGEP GPTGIQGPPG PAGEEGKRGA RGEPGPAGLP GPPGERGGPG SRGFPGADGV AGPKGPAGER GAPGPAGPKG SPGEAGRPGE AGLPGAKGLT GSPGSPGPDG KTGPPGPAGQ DGRPGPPGPP GARGQAGVMG FPGPKGAAGE PGKAGERGVP GPPGAVGPAG KDGEAGAQGP PGPAGPAGER GEQGPAGSPG FQGLPGPAGP PGEAGKPGEQ GVPGDLGAPG PSGARGERGF PGERGVQGPP GPAGPRGANG APGNDGAKGD AGAPGAPGSQ GAPGLQGMPG ERGAAGLPGP KGDRGDAGPK GADGAPGKDG VRGLTGPIGP PGPAGAPGDK GEAGPSGPAG PTGARGAPGD RGEPGPPGPA GFAGPPGADG QPGAKGEPGD AGAKGDAGPP GPAGPAGPPG PIGNVGAPGP KGARGSAGPP GATGFPGAAG RVGPPGPSGN AGPPGPPGPA GKEGSKGPRG ETGPAGRPGE VGPPGPPGPA GEKGAPGADG PAGAPGTPGP 1000 QGIAGQRGVV GLPGQRGERG FPGLPGPSGE PGKQGPSGAS GERGPPGPMG 1050 PPGLAGPPGE SGREGAPGAE GSPGRDGSPG AKGDRGETGP AGPPGAPGAP GAPGPVGPAG KSGDRGETGP AGPAGPIGPV GARGPAGPQG PRGDKGETGE 1100 OGDRGIKGHR GFSGLOGPPG PPGSPGEOGP SGASGPAGPR GPPGSAGSPG 1150 KDGLNGLPGP IGPPGPRGRT GDAGPAGPPG PPGPPGPPGP PSGGYDLSFL 1200 POPPOEKAHD GGRYYRADDA NVVRDRDLEV DTTLKSLSOO IENIRSPEGS 1250 1300 RKNPARTCRD LKMCHSDWKS GEYWIDPNOG CNLDAIKVFC 1350 PTOPSVAOKN WYISKNPKEK RHVWYGESMT GGFOFEYGGO GSDPADVAIC 1400 ONTTYHCK NSVAYMDOOT GNLKKALLLO GSNETEIRAF 1450 HTGA WGKTVIEYKT TKTSRLPIID VAPLDVGAPD QEFGFDVGPA CFL

50

100

150

200

250

300

350

400

450

500

550

600

650

700

750

800

850

900

950

	a	2	-1	
		-		

MLSFVDTRTL LLLAVTSCLA TCQSLQEATA RKGPSGDRGP RGERGPPGP GRDGDDGIPG PPGPPGPPGP PGLGGNFAAO FDAKGGGPGP 100 GASGAPGPQG FQGPPGEPGE PGQTGPAGAR GPPGPPGKAG EDGHPGKPGF 150 PGERGVVGPQ GARGFPGTPG LPGFKGIRGH NGLDGLKGQP GAPGVKGEPG 200 APGENGTPGO TGARGLPGER GRVGAPGPAG ARGSDGSVGP VGPAGPIGSA 250 GPPGFPGAPG PKGELGPVGN PGPAGPAGPR GEVGLPGLSG PVGPPGNPGA 300 NGLPGAKGAA GLPGVAGAPG LPGPRGIPGP VGAAGATGAR GLVGEPGPAG 350 SKGESGNKGE PGAVGQPGPP GPSGEEGKRG STGEIGPAGP PGPPGLRGNP 400 450 GSRGLPGADG RAGVMGPAGS RGATGPAGVR GPNGDSGRPG EPGLMGPRGF PGSPGNIGPA GKEGPVGLPG IDGRPGPIGP AGARGEPGNI GFPGPKGPSG 500 550 DPGKAGEKGH AGLAGARGAP GPDGNNGAQG PPGLQGVQGG KGEQGPAGPP GFQGLPGPAG TAGEAGKPGE RGIPGEFGLP GPAGARGERG PPGESGAAGP 600 TGPIGSRGPS GPPGPDGNKG EPGVVGAPGT AGPSGPSGLP GERGAAGIPG 650 GKGEKGETGL RGDIGSPGRD GARGAPGAIG APGPAGANGD RGEAGPAGPA 700 GPAGPRGSPG ERGEVGPAGP NGFAGPAGAA GQPGAKGERG TKGPKGENGP 750 VGPTGPVGAA GPSGPNGPPG PAGSRGDGGP PGATGFPGAA GRTGPPGPSG 800 ISGPPGPPGP AGKEGLRGPR GDQGPVGRSG ETGASGPPGF VGEKGPSGEP 850 GTAGPPGTPG PQGLLGAPGF LGLPGSRGER GLPGVAGSVG EPGPLGIAGP 900 PGARGPPGNV GNPGVNGAPG EAGRDGNPGN DGPPGRDGQP GHKGERGYPG 950 NAGPVGAAGA PGPQGPVGPV GKHGNRGEPG PAGAVGPAGA VGPRGPSGPQ 1050 GIRGDKGEPG DKGPRGLPGL KGHNGLQGLP GLAGHHGDQG APGAVGPAGP RGPAGPSGPA GKDGRIGOPG AVGPAGIRGS OGSOGPAGPP GPPGPPGPG 1100 PSGGGYEFGF DGDFYRADOP RSPTSLRPKD YEVDATLKSL NNOIETLLTH 1150 1200 1250 1300 1350 EGNSRFTYTV LVDGCSKKTN EWOKTITEYK TNKPSRLPIL DIAPLDIGGA 1364 DOEIRLNIGP VCF

Simon, H., van Agthoven, M., Lam, P. Y., Floris, F., Chiron, L., Delsuc, M.-A., Rolando, C., Barrow, M., O'Connor, P. (2016). Analyst, 141, 157-165

# Bottom-up proteomics

- Collagen protein
  - 2D FT-ICR IRMPD

    - ▶ 60 min. acquisi<sup>§</sup>ion
    - ▶ R1 ~ 630
    - ▶ **R2 ~ 13.000** (*m/z*=850)



Simon, H., van Agthoven, M., Lam, P. Y., Floris, F., Chiron, L., Delsuc, M.-A., Rolando, C., Barrow, M., O'Connor, P. (2016). Analyst, 141, 157–165

#### Bottom-up proteomics

- Collagen protein
  - comparing CAD, ECD, IRMPD, 2D IRMPD

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### <u>Bottom-up proteomics</u>

- yeast cells
  - cell extract
  - tryptic digest
  - no separation



- 2D FT-ICR IRMPD
  - ▶ 4096 x 256k = 1 Gpoint
  - processed to 4k x 1M = 4 Gpoints
  - ▶ 50 min. acquisition
  - ▶ R1 ~ 1500
  - ▶ R2 ~ 60.000

# Parent Excitation limited to $m/z = 330 \dots 700$





<sup>&</sup>gt;100k? potential PSM



## NUS: substance P, 2D spectrum overview

Non-Uniform Sampling / non-Fourier Analysis



The overall aspect of the 2D spectrum is preserved.





Classic

NUS 32k div 16

## NUS (32k, 1/16) 2D zoom on doubly charged



*in preparation* 

a triglyceride mixture



1. ligand binding

a. Recent developments in protein-ligand affinity mass spectrometry (http://europepmc.org/articles/PMC3043251)

b. Mass spectrometry-based approaches to protein-ligand interactions (<u>http://www.tandfonline.com/doi/abs/10.1586/14789450.2.4.475?</u>

src=recsys&journalCode=ieru20)

c. Mass spectrometry based tools to investigate protein–ligand interactions for drug discovery (<u>http://pubs.rsc.org/en/Content/ArticleLanding/2012/CS/</u> <u>C2CS35035A#ldivAbstract</u>)

2. Kd measurements - with all the caveat on ESI, etc..

a. Converting Solution Macromolecular Thermodynamic Properties into Gas-Phase Mass Spectrometry Observations (<u>http://www.cell.com/cell-chemical-biology/fulltext/S1074-5521(02)00221-1</u>)

b. Sizing Up Protein–Ligand Complexes: The Rise of Structural Mass Spectrometry Approaches in the Pharmaceutical Sciences (<u>http://www.annualreviews.org/</u><u>doi/full/10.1146/annurev-anchem-061516-045414#f1</u>)</u>

c. A General Mass Spectrometry-Based Assay for the Quantitation of Protein–Ligand Binding Interactions in Solution (<u>http://pubs.acs.org/doi/abs/10.1021/ja026574g</u>)

3. multi molecular complexes

a. The diverse and expanding role of mass spectrometry in structural and molecular biology (<u>http://onlinelibrary.wiley.com/doi/10.15252/embj.201694818/full</u>) b. Advances in the Mass Spectrometry of Membrane Proteins: From Individual Proteins to Intact Complexes (<u>http://www.annualreviews.org/doi/abs/10.1146/</u><u>annurev-biochem-062309-093307?journalCode=biochem</u>)</u>

c. The emerging role of native mass spectrometry in characterizing the structure and dynamics of macromolecular complexes (<u>http://onlinelibrary.wiley.com/doi/10.1002/pro.2661/pdf</u>)

d. Mass spectrometry guided structural biology (http://www.sciencedirect.com/science/article/pii/S0959440X16301440)

4. HDX

a. Hydrogen exchange mass spectrometry for studying protein structure and dynamics (<u>http://pubs.rsc.org/en/content/articlelanding/2011/cs/c0cs00113a#!</u> divAbstract)

b. Differential hydrogen/deuterium exchange mass spectrometry analysis of protein-ligand interactions (http://europepmc.org/articles/PMC3113475)

c. Hydrogen exchange mass spectrometry: what is it and what can it tell us? (https://link.springer.com/article/10.1007/s00216-010-3556-4?no-access=true)

5. cross-linking

a. Chemical cross-linking and mass spectrometry to map three-dimensional protein structures and protein-protein interactions (<u>http://onlinelibrary.wiley.com/doi/10.1002/mas.20082/abstract</u>)

b. Probing Native Protein Structures by Chemical Cross-linking, Mass Spectrometry, and Bioinformatics (http://www.mcponline.org/content/9/8/1634.full)

c. Chemical cross-linking/mass spectrometry targeting acidic residues in proteins and protein complexes (http://www.pnas.org/content/111/26/9455.long)

d. Chemical cross-linking and native mass spectrometry: A fruitful combination for structural biology (<u>http://onlinelibrary.wiley.com/doi/10.1002/pro.2696/</u> full#references)

6. fragmentation

a. Top-down mass spectrometry: Recent developments, applications and perspectives (<u>http://pubs.rsc.org/en/content/articlelanding/2011/an/c1an15286f#!</u> divAbstract)

b. Native mass spectrometry of photosynthetic pigment-protein complexes (http://www.sciencedirect.com/science/article/pii/S0014579313000197)

c. 193 nm Ultraviolet Photodissociation Mass Spectrometry of Tetrameric Protein Complexes Provides Insight into Quaternary and Secondary Protein Topology (http://pubs.acs.org/doi/abs/10.1021/jacs.6b03905)

acknowledgment to Huilin Li

## Thank you



