



SANS bridges the gap between atomic

resolution (NMR and crystallography)

and the light microscope

#### Typical lengthscales

NMR

crystallograpy



## Scattering basics: Huygens-Fresnel principle



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Reciprocal relationship between real space and the diffraction pattern





#### Information obtained by SANS:

- 1) Oligomeric state of macromolecules
- 2) Shape or conformation (globular, stick etc...)
- 3) Interaction of different macromolecules
- 4) Variation of points (1)-(3) as a function of pH, salt, ligands, T, p, ...
- 5) Contrast variation: visualisation of individual sub-units in situ

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#### Modeling techniques using SAS in structural biology



Putnam et al. (2007) Q. Rev. Biophys. 40(3), 191-285

What can SANS provide that is different from SAXS?

#### SANS allows to go beyond the global shape and study internal structure!



Often problematic to position/orient subunits in a larger complex using SAXS alone...

Internal structure: contrast variation and SANS!

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#### Idea of contrast variation



Vary scattering behaviour of (parts of) solutes with respect to solvent



Ideal solutions: no inter-particle effects, only form-factors

#### Scattering densities in biological samples

Atom Hydrogen Deuterium Carbon Nitrogen Oxygen Phosphorus Sulphur (a) Water	Nucleus <sup>1</sup> H <sup>2</sup> H (D) <sup>12</sup> C <sup>14</sup> N <sup>16</sup> O <sup>31</sup> P Mostly <sup>32</sup> S	$\begin{array}{cccc} b_{\rm coh} & f_{\rm X-ray} \left( \theta = 0^{\circ} \right) \\ \hline \left( 10^{-12}  {\rm cm} \right) & \left( 10^{-12}  {\rm cm} \right) \\ \hline \left( -0.3742 \right) & 0.28 \\ \hline 0.6671 & 0.28 \\ \hline 0.0651 & 1.69 \\ \hline 0.940 & 1.97 \\ \hline 0.5804 & 2.25 \\ \hline 0.517 & 4.23 \\ \hline 0.2847 & 4.5 \\ \hline \\ D & -0.168 & -0.00562 \\ \hline 0 & 1.90164 \\ \end{array} \right)$	(c) Nucleotides and Base Adenine { Guanine { Cytosine { Uracil 1 Thyminine 1	nucleic acid   Chemical composition H (   NA PNsC1008H11 3   NNA PNsC1003H11 3   NNA PNsC1008H11 4   NNA PNsC1008H11 4   NNA PNsC1008H11 3   NNA PNsC208H11 3   NNA PNsC208H11 3   NNA PNsC208H1 2   NNA PNsC208H10 2   NNA PNsC207H12 1	$\begin{array}{cccc} b_{\rm tot}  ({\rm H_2O}) & b_{\rm tot}  ({\rm D_2O}) \\ (10^{-13}  {\rm cm}) & (10^{-13}  {\rm cm}) \\ 11^{-23} & 14^{+35} \\ 10^{-65} & 12^{-73} \\ 11^{-81} & 15^{-98} \\ 11^{-23} & 14^{+35} \\ 9^{-26} & 12^{-39} \\ 8^{-68} & 10^{-77} \\ 9^{-28} & 11^{-36} \\ 8^{-61} & 9^{-65} \end{array}$	) b (deuterated) ) (10 <sup>-12</sup> cm) 22-68 22-10 23-26 22-68 20-72 20-14 19-69 21-11
(b) Amino acida Glycine Alanine Valine Leucine Isoleucine Phenylalanine Tyrpstophan Aspartic acid Glutamise Glutamise Glutamise Lysine Argaine Histidine Methionine Cysterine	and proteins   Chemical   Canposition   CaNOH5   CaNOH5   CaNOH5   CaNOH5   CaNOH5   CaNOH6   CaNOH5   CaNOH6   CaNOH6   CaNOH7   CaNOH8   CaNO4H8   CaNo5H8   CaNo8H8   CaNo8H8   CaNo8H8   CaNo8H8	$\begin{array}{c ccccc} b_{1et}\left(H_2O\right) & b_{1et}\left(D_2O\right) \\ I\left(\infty\right) & \left(10^{-12}\ cm\right) & \left(10^{-12}\ cm\right) \\ I\left(-1728 & 2.769 \\ 1.645 & 2.666 \\ 1.479 & 2.520 \\ 1.396 & 2.437 \\ 1.396 & 2.437 \\ 1.396 & 2.437 \\ 1.139 & 5.180 \\ 2.437 & 1.139 \\ 5.162 & 4.863 \\ 3.762 & 4.863 \\ 3.762 & 4.803 \\ 2.215 & 4.803 \\ 2.215 & 4.803 \\ 2.215 & 4.803 \\ 2.2142 & 4.224 \\ 1.365 & 6.580 \\ 3.373 & 6.497 \\ 3.373 & 6.497 \\ 3.374 & 5.521 \\ 1.586 & 5.752 \\ 3.465 & 9.751 \\ 3.465 & 6.521 \\ 1.764 & 2.805 \\ 1.764 & 2.805 \\ 1.930 & 4.013 \\ \end{array}$	biai (deuterated) (10 <sup>-13</sup> cm) pr.   (10 <sup>-13</sup> cm) pr.   4-85 6   6-852 9   10-854 14   12-850 16   13-51 20   14-09 20   16-45 23   8-010 14   7-431 12   9-704 13   11-70 16   15-12 17   11-73 16   11-14 7-137	A*) $\bar{l}$	$\overline{b} = \rho_{protein} = \sum_{j} \frac{b_{j}}{V}$ Example glycine in H <sub>2</sub> C .94+0.58+3*(-0.37)]10 <sup>-12</sup> cm/Å <sup>3</sup> = 2.68*10 <sup>10</sup> cm <sup>-2</sup>	<u>*</u> cm/66.4Å <sup>3</sup> 911.953



#### Destructive interference in SANS

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In practice, all biomacromolecules can be matched in SANS, i.e. made invisible!!! Not so easy with SAXS...



#### An analogon in optics: refractive index

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#### SAXS and contrast variation?

- Accessible range of solvent electron densities is limited
- Contrast agents (salt, sugar...) need to be added at high molarities and may not be inert to biomolecules
- Electron density of biomolecules cannot be modified globally

Mahieu, E. & Gabel, F. (2018). Acta Cryst. D74(Pt 8), 715-726.

#### Contrast variation in SANS: natural contrast

Proteins and RNAs have different proton densities









100% H<sub>2</sub>O

58% H<sub>2</sub>O, 42% D<sub>2</sub>O

30% H<sub>2</sub>O, 70% D<sub>2</sub>O

Also possible for protein-protein complexes (deuteration)!

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Artificial contrast using deuteration

Careful at high D<sub>2</sub>O levels in the solvent: favours oligomerisation/aggregation!



Measure I(0) experimentally (Guinier!) and trace the following expression as a function of H<sub>2</sub>O/D<sub>2</sub>O:



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#### Guinier approximation and radius of gyration



"See-saw analogy"

For a given molecular weight, a sphere has the smallest  $R_{\rm g}$ , *i.e.* it is the most compact object



Protein-protein complexes

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## Negative radii of gyration!?

- Scattering in forward direction, *I*(0), can be weak (or zero)
- Scattering can get stronger going to higher angles
- Result: "apparent" negative radius of gyration

# Can proteins be considered as homogeneous particles?



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# Summary: homogeneity of biomacromolecules in SANS experiments

- Proteins can be considered as homogeneous if the scattering density fluctuations are **not ordered** (*e.g.* core vs. outer shell) and/or if they occur on a **length-scale much smaller** than the overall protein dimension and/or the molecules of interest to be studied in a complex
- RNA and DNA are **more homogeneous** than proteins regarding the scattering density fluctuations
- Lipids are in general **heterogeneous** (head *vs.* tail) but can be made homogeneous using **deuteration**
- The approximation of homogeneous particles improves at smaller angles
- Careful with ab initio techniques in SANS!



# **Practical aspects**

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# SAXS vs SANS: some practical aspects

#### SANS vs SAXS instruments



BM29 (ESRF)

SAXS

- easy manipulation with pipettes
- multi-sample holder
- no radiation damage!

Quartz cuvette (SANS)

#### Exposure times:

- ~ 1-60 minutes (D22)
- ~ 1-10 seconds (BM29)

Including sample change:

- ~ 1-60 minutes (D22)
- ~ 2-3 minutes (BM29)

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Historical example 1: Chromatin



#### The chromatin structure

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#### Contrast variation: relative arrangement of DNA and protein



Figure 2: Examples of Guinier plots obtained from chromatin par-ticles (a) Particles in D<sub>0</sub> (b) in H<sub>30</sub>. Semi-log plot of intensity (1) verus h<sup>2</sup>, where h = 4 Hain $\theta_A$  and 26 is the angle of acattering. Each data set is fitted with a straight line by a variance-weighted least squares procedure. The data are arbit-rarily scaled with respect to I for convenience in plotting them on the same graph.

#### Relative topology of DNA and protein at low resolution before availability of high-resolution models!

Pardon et al. (1975) Nucl. Acids Res. 2(11) 2163-2176



Figure 5: Two possible kinds of structure for the chromatin particle containing 140 base pairs of DRA and eight histones. (a) A spherical particle with overall diameter  $a_{\rm g}=53$ , derived from  $R_{\rm g}=40.1$ , in which the inner protein core has radius  $a_{\rm g}=40$  from the experimental  $R_{\rm g}=30.6$ Å. The region occupying the DRA is shaded. (b) A cylindrical model in which two turns of helix with pitch 45Å and radius 37Å wound on an inner protein core of radius 27Å.

Historical example 2: **The ribosome** 

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#### The ribosome structure



## The "glassy" ribosome

Nierhaus et al. (1983) Proc. Natl. Acad. Sci. USA 80, 2889-2893

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The distance between two components of a complex can be extracted from the scattering curve

courtesy Roland May



3 distances define a triangle (a), another 3 a tetrahedron of undefined handedness (b). Each further 4 distances add another component in space (c).

courtesy Roland May

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#### Ribosome at low resolution

M.S. Capel, D.M. Engelman, B.R. Freeborn, M. Kjeldgaard, J.A. Langer, V. Ramakrishnan, D.G. Schindler, D.K. Schneider, B.P. Schoenborn, I.-Y. Sillers, S. Yabuki, P.B. Moore (1987) *Science* **238**, 1403-1406

Map of the 30S ribosomal subunit from E. coli. Each protein is represented by a sphere whose volume is the same as that of the protein. The maximum linear dimension of the array is about 190 Å.

courtesy Roland May

# **EXAMPLE 3:** Membrane proteins

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## Membrane proteins and lipids/detergents





Johs et al. (2006) J. Biol. Chem. 281, 19732-19739

#### Scattering contrast of lipids: heterogeneity



H-lipid headgroup 'denser'



D-lipid

(d) Lipids			h (deuterated)	
Phoenhyrylcholine	<i>b</i> (10 <sup>-12</sup> cm)	$\bar{b} (10^{-12} \text{ cm Å}^{-3})$	(10 <sup>-12</sup> cm)	$\bar{b} (10^{-12} \text{ cm Å}^{-3})$
CIL	-0.0833	-0.0031	2	0.0744
CH2 CH	-0.458	-0.0085	2.67	0.0495
C5H13NPO4	2.24	0.011	15.76	0.072

A KcsA full length 100% 0. 0.0 (b) 1E-3 1E-4 1E-0.00 0 10 0.15 0.20 0.25 0.30 q [1/Å] A 0.025 KcsA full length 0.020 0.015 D(r) 0.010 0.005 0.000 20 40 60 80 100 120 140 r [Å]

# 

concentration. On the basis of published detergent scattering match points, we estimated the detergent scattering match point for decyl-B-D-maltopyranoside (C<sub>2</sub>H<sub>4</sub>Q<sub>0</sub>)<sub>1</sub>) at -22% D<sub>2</sub>O solvent concentration (24), thereby minimizing the contribution of the colbatilities 1 detergent micelle to the obored neutron scattering. However, the fluctuation of scattering length of the DM detergent molecule must be stressed this is due to be individual contening mustle, points of the polar maltoside headgroup of -46% vs. ~3% D<sub>2</sub>O solvent concentration for the hydrophobic decyl side chain. The overall match point at 22% D<sub>2</sub>O solvent concentration, mini-

Zimmer et al. (2006) Biophys. J. 90, 1752-1766

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## **EXAMPLE 4:**

pH-induced transition in KcsA

Sophisticated approaches using SANS (SAXS) and NMR:

a « tour d'horizon » using a recent example: the BOX C/D complex



Gabel (2015) Small-angle neutron scattering for structural biology of protein-RNA complexes. Methods in Enzymology 558, 391-415.



#### Combining NMR with SAS

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rRNA modifications and function

Dozens of modifications in structurally and functionally important (and conserved) regions; their number increases with "complexity" of organism.

Single mutations can be tolerated, absence of all modifications is lethal.

Decatur, W.A. and Fournier, M.J. (2002) rRNA modifications and ribosome function TIBS 27(7), 344-351.



Number of modifications: bacteria < archaea < eukarya



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#### RNA modifications: snoRNPs, snoRNAs and box C/D



SANS (D22) and SAXS (BM29) data

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Relative positions of FIB proteins within the complex from SANS data



Important restraints for the atomic models!



Lapinaite, A., Simon, B., Skjaerven, L., Rakwalska-Bange, M., Gabel, F. and Carlomagno T. (2013) The structure of the box C/D enzyme reveals regulation of RNA methylation. *Nature* **502**(7472), 519-523.

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Large conformational change upon substrate binding!

#### The holo complex



RNA/RNA-binding pulls two NOP dimers apart, FIBs are in contact with RNA

Large conformational change upon substrate (RNA) binding to an elongated form (SAXS/SANS+ 257 PRE distance restraints)

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Proposed model for the sequential methylation and conformational changes



The structural model of the holo-enzyme, together with the NMR assays, suggests that methylation at the two sites occurs in a sequential, well-defined order!

Implications on folding pathways for ribosome...

# A novel mechanism for translational regulation in *Drosophila melanogaster*



Hennig J, Militti C, Popowicz G, Wang I, Sonntag M, Geerlof A, Gabel F, Gebauer F, and Sattler M (2014). Structural basis for the assembly of the SXL-UNR translation regulatory complex. *Nature* 515(7526), 287-290.

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## **Dosage compensation**

Human (male) karyotype





XIST gene silencing system in female mammals





Unequal proteins amounts from XX and XY pairs: needs compensation mechanisms

(Klinefelder syndrome in humans: XXY)

## Dosage compensation in *D. melanogaster*



- Up-regulated by "DCC" (Dosage compensation complex) constituted of 5 proteins and 2 non-coding RNAs
- Female-specific protein "SXL" (sex-lethal) silences the expression of a protein of the DCC complex in females by binding to its mRNA transcript and inhibiting its interaction with the ribosome





## **SANS-specific information**



Svergun DI, Richard S, Koch MH, Sayers Z, Kuprin S, Zaccai G. (1998) Protein hydration in solution: experimental observation by x-ray and neutron scattering. *Proc Natl Acad Sci U S A*. **95**(5):2267-2272.

Jochen Hub's talk



Residue-specific densities!	
Denser around acidic residue	s



10

Kim, H.S., Martel, A., Girard, E., Moulin, M., Härtlein, M., Madern, D., Blackledge, M., Franzetti B. and Gabel F. (2016) Solution scattering of X-rays and neutrons on supercharged proteins reveals residue-specific modifications of the hydration shell. *Biophys. J.* 110(10), 2185-2194.



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Ibrahim, Z., Martel, A., Moulin, M., Kim, H.S., Härtlein, M., Franzetti, B. and Gabel, F. (2017) Sci. Rep. 7, 40948.



Could this have been done by SAXS? (d SAXS curve predicted for protein mixture at t = 0 min 100 10 d-GFP (5 mg/mL) h-PAN (1.5 mg/mL) h-205 (23.5 mg/mL) 1 SAXS curve of interest 0.1 • d-GFP + h-PAN + h-20S h-20S (23.5 mg/mL) h-PAN (1.5 mg/mL) d-GFP (5 mg/mL) 0.01 0.30 q [Å<sup>-1</sup>] 0.05 0.10 0.15 0.20 0.25 Mahieu et al. (2020) EPJ Web Conf. 236, 03002

# Example 7: segmental labeling





Williams et al. (2016) PLoS ONE 11(4):e0154607

Sonntag et al. (2017) Angew. Chem. Intl. Ed. 56(32):9322-9325

More detailed insight into multi-domain proteins



Courtesy Anne Martel (ILL D22)

#### Practical aspects: doing SANS experiments

- use SAXS for homogeneous systems composed of a single body
- neutrons only possible at large facilities (no "home sources" for the moment!)
- request for measurement time is generally via an electronic proposal system
- deadlines are usually twice a year, beamtime is attributed about 6 months later
- BAG ("Block allocation group") systems allow more flexible access
- for continuation proposals, reports need to be submitted regularly
- experiments need to be prepared with great care (i.e. isotopic effect of D2O)!!
- "local contacts", often beamline responsibles, assist during experiments
- access (for non-industrial use) is in general free
- no maintenance, user friendly (software etc...)



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#### Summary

- 1) Low-resolution information in solution
- 2) Non-destructive technique, easy to use
- 3) Possibility to use contrast and focus on subsystems within complexes
- 4) Special applications: protein/DNA-RNA complexes, membrane systems
- 5) Doing biochemistry on samples in situ during measurement
- 6) Complementary information to SAXS

#### Literature

#### Basics (scattering, quantum mechanics):

- The Feynman lectures on Physics, Volume 3: Quantum mechanics (Addison Wesley, 2006)
- Cohen-Tannoudji et al.: Mécanique Quantique, Vol. 2. Chapter on diffusion. (Hermann, 1997)

#### General books on neutron scattering:

- Lovesey: Theory of Neutron Scattering from Condensed Matter (Clarendon, 1986)
- Geissler et al.: Structure and dynamics of biomolecules (Oxford University Press, 2000)

#### Books on small angle (neutron) scattering:

- Svergun: Structure Analysis by Small-Angle X-Ray and Neutron Scattering (Plenum, 1987)
- Guinier/Fournet: Small angle scattering of X-rays (John Wiley & Sons, 1955)
- Serdyuk, Zaccai, Zaccai: Methods in molecular biophysics (Cambridge University Press, 2007)

#### Reviews on SAXS/SANS:

- Jacrot, B. (1976) The Study of biological structures by neutron scattering from solution. *Rep. Prog. Phys.* 39, 911-953.
- Putnam et al. (2007) X-ray solution scattering (SAXS) combined with crystallography and computation: defining accurate macromolecular structures, conformations and assemblies in solution. *Q. Rev. Biophys.* 40(3):191-285.









The "BDCS" group (<u>Biology</u>, <u>Deuteration</u>, <u>Chemistry</u> and <u>Soft</u> Matter) at ILL: getting the best from your neutron experiment!





#### Joint SAXS/SANS contrast variation project: Postdoc position available (december call)!

