



**SANS bridges the gap between atomic resolution (NMR and crystallography)** 

**and the light microscope**

# Typical lengthscales

NMR crystallograpy



right © Pearson Education, Inc., publishing as Benjamin Cumming:

# 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?**

# and study internal structure! lg  $\overline{c}$  $\overline{1}$  $\mathsf{o}$  $-1$  $\mathbf{0}$ 6 s. nm

SANS allows to go beyond the global shape

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





## 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?

- **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 58% H2O, 42% D2O 30% H2O, 70% D2O <sup>2</sup>O**

#### **Also possible for protein-protein complexes (deuteration)!**

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Artificial contrast using deuteration

Careful at **high D2O** 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*<sup>g</sup> , *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**



- multi-sample holder
- no radiation damage!

**Quartz cuvette (SANS)**

#### **Exposure times:**

- $\sim$  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 uniter plots obtained from chromatin particles (a) Particles in D.O (b) in H3O. Semi-log plot of interior interior in the interior interior interior in the straight line by variance-weighted least a

#### 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



<u>Figure 5</u>: Two possible kinds of structure for the chromatin<br>particle containing 140 base pairs of DEVA and eight histones.<br>(a) A spherical particle with overall diameter  $a_c = 53\hat{k}$ , derived<br>from  $R_1 = 41.1$ , in which

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

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# Scattering contrast of lipids: heterogeneity



H-lipid headgroup 'denser' D-lipid





A KcsA full length  $\ddot{\mathbf{0}}$  $0.0$  $\overline{a}$  $1E-3$  $1E-4$  $1E-5$  $0.00$  $0.15$  $0.20$  $0.25$  $0.30$  $0.10$ q [1/Å] A 0.025 KcsA full length  $0.020$  $0.015$ D(r)  $0.010$ 0.005  $0.000$  $\overline{20}$  $40$ 60 80  $100$  $120$ 140

r [Å]

## pH-induced transition in KcsA



concentration. On the basis of published detergent scattering<br>match points, we estimated the detergent scattering match<br>point for decyl- $\beta$ -D-mattopyramoside ( $C_2H_{2\text{-}}Q_{11}$ ) at  $\sim 22\%$ <br>D-O solvent concentration (24 ed neutron scattering. However, the fluctuation of so ng length of the DM detergent molecule must be stresse ñ This is due to the individual scattering match control the polar matterial content<br>matterial concentration for the hydrophobic decyls  $\sim 3\%$  D<sub>2</sub>O solvent<br>concentration for the hydrophobic decyl side chain. The over-<br>al

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

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# **EXAMPLE 4:**

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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# $(e)$

rRNA modifications and function

Dozens of modifications in structurally<br>and functionally important (and 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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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



**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 residues**



 $10^7$  $\overset{\scriptscriptstyle{10}^{\phantom{0}^{\phantom{0}^\dagger}}}{\mathsf{C}}$ 

 $\frac{1}{10}$  $10^{-1}$  $10<sup>1</sup>$  $10$ 

 $10^{\frac{1}{3}}$ 

A

uns

10  $10<sup>1</sup>$  $\overline{10}$  $10^{\circ}$ 

**Pyrococcus horizontal** 

d-SANS(

 $\frac{1}{200}$ 

 $\frac{1}{04}$ 

 $O(K^2)$ 

 $O(\text{A}^2)$ 

GFP (+36)

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**Kim, H.S.,** Martel, A., Girard, E., Moulin, M., Härtlein, M., Madern, D., Blackledge, M., Franzetti<sup>,</sup> B. and Gabel<sup>,</sup> 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.



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



And in the presence of the proteolytic core particle 20S?



# **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 D<sub>2</sub>O)!!
- "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 (Biology, Deuteration, Chemistry and Soft Matter) at ILL: getting the best from your neutron experiment!**





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

