

Frank Gabel: EMBO course Sep. 2024

Typical lengthscales

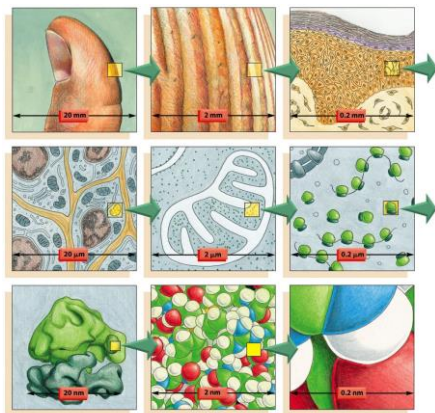
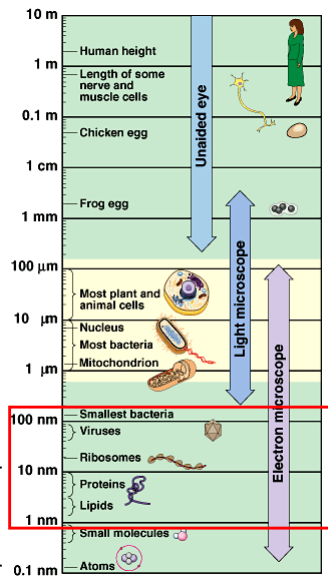


Figure 9-1 Molecular Biology of the Cell 5/e (© Garland Science 2008)

Small-Angle Neutron Scattering (SANS) in solution

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

NMR
crystallography



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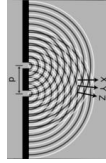
Scattering basics: Huygens-Fresnel principle



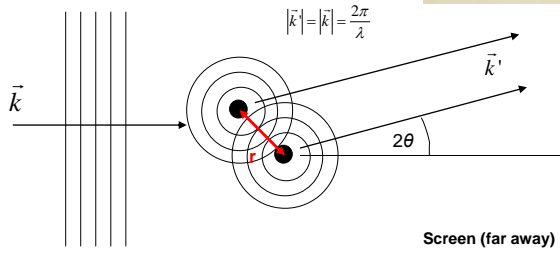
1629-1695



1788-1827



Incoming X-ray/neutron wave



Many scattering centers, FOURIER transform:

$$I(Q) = \left| \sum_j b_j e^{-i\vec{Q} \cdot \vec{r}_j} \right|^2$$

$$Q = |\vec{k}' - \vec{k}| = \frac{4\pi}{\lambda} \sin \theta$$

Reciprocal relationship between real space and the diffraction pattern

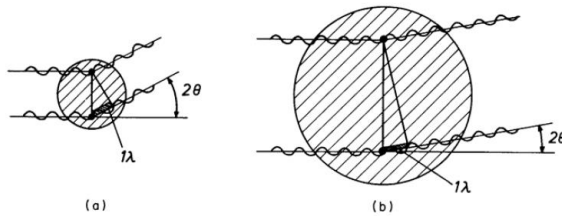


FIG. 1

Glatter and Kratky (1982)

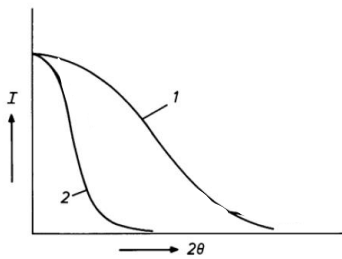
Many scattering centers, FOURIER transform:

$$I(Q) = \left\langle \left| \sum_j b_j e^{-i\vec{Q} \cdot \vec{r}_j} \right|^2 \right\rangle$$

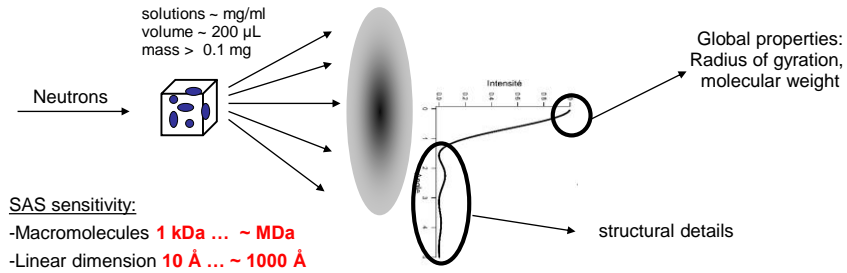
Oriental averaging

$$I(Q) = \sum_{i,j} b_i b_j \frac{\sin Q(r_i - r_j)}{Q(r_i - r_j)}$$

Debye equation (1915)



SANS sample conditions and information obtained

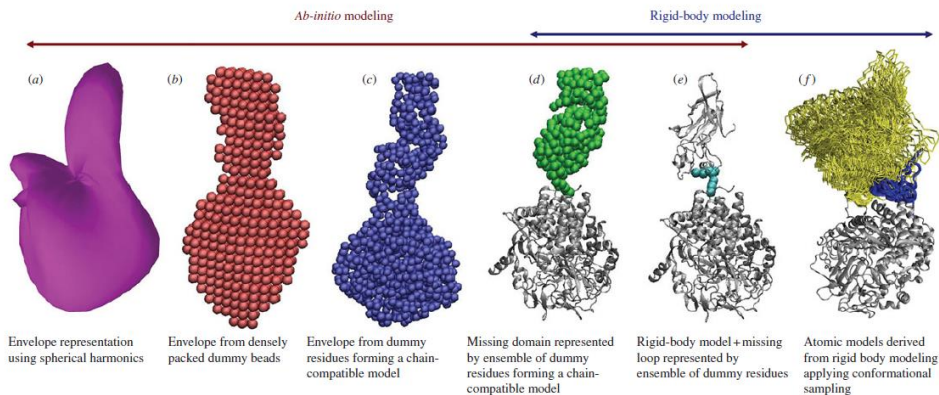


Information obtained by SANS:

- 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*

Modeling techniques using SAS in structural biology

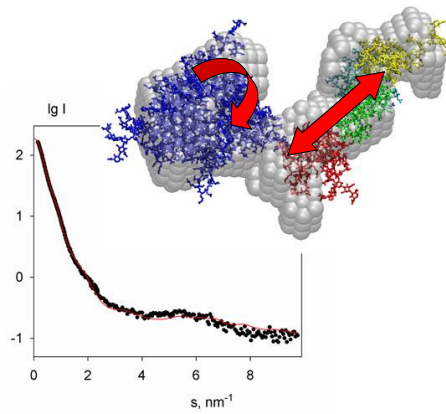
(SAXS lectures, SASscape tutorials,...)



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!**

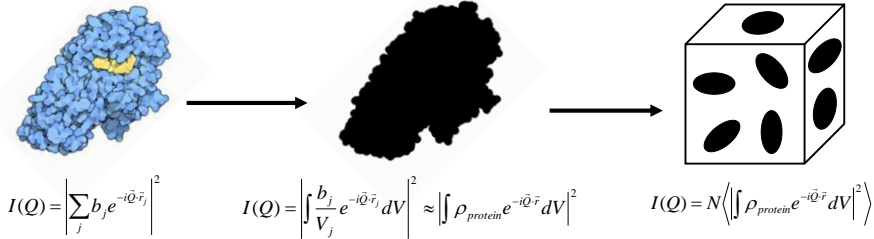
Idea of contrast variation



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

Concept of scattering density and contrast

In vacuo:



$$I(Q) = \left| \sum_j b_j e^{-i\vec{Q}\cdot\vec{r}_j} \right|^2$$

$$I(Q) = \left| \int \frac{b_j}{V_j} e^{-i\vec{Q}\cdot\vec{r}_j} dV \right|^2 \approx \left| \int \rho_{protein} e^{-i\vec{Q}\cdot\vec{r}} dV \right|^2$$

$$I(Q) = N \left\langle \left| \int \rho_{protein} e^{-i\vec{Q}\cdot\vec{r}} dV \right|^2 \right\rangle$$

In solution:

$$I(Q) = \left\langle \left| \int (\rho_{protein} - \rho_{solvent}) e^{-i\vec{Q}\cdot\vec{r}} dV \right|^2 \right\rangle$$

Continuum approximation:

$$\frac{b_j}{V_j} \approx \rho_{protein} = const$$

- How are scattering densities calculated?
- Under which conditions is the approximation valid?

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

Scattering densities in biological samples

Atom	Nucleus	b_{coh} (10^{-12} cm)	$f_{x-ray}(\theta=0^\circ)$ (10^{-12} cm)
Hydrogen	^1H	-0.3742	0.28
Deuterium	$^2\text{H (D)}$	0.6671	0.28
Carbon	^{12}C	0.6651	1.69
Nitrogen	^{14}N	0.940	1.97
Oxygen	^{16}O	0.5804	2.25
Phosphorus	^{31}P	0.517	4.23
Sulphur	Mostly ^{32}S	0.2847	4.5

(c) Nucleotides and nucleic acid					
Base	Chemical composition	H (ex)	b_{tot} (H_2O) (10^{-12} cm)	b_{tot} (D_2O) (10^{-12} cm)	b (deuterated) (10^{-12} cm)
Adenine	RNA $\text{PN}_5\text{C}_{10}\text{O}_4\text{H}_{11}$	3	11.23	14.35	22.68
	DNA $\text{PN}_5\text{C}_{10}\text{O}_2\text{H}_{11}$	2	10.65	12.73	22.10
Guanine	RNA $\text{PN}_5\text{C}_{10}\text{O}_4\text{H}_{11}$	4	11.81	15.98	23.26
	DNA $\text{PN}_5\text{C}_{10}\text{O}_2\text{H}_{11}$	3	11.23	14.35	22.68
Cytosine	RNA $\text{PN}_4\text{C}_9\text{O}_4\text{H}_{11}$	3	9.26	12.39	20.72
	DNA $\text{PN}_4\text{C}_9\text{O}_2\text{H}_{11}$	2	8.68	10.77	20.14
Uracil	RNA $\text{PN}_4\text{C}_8\text{O}_4\text{H}_{10}$	2	9.28	11.36	19.69
Thymine	DNA $\text{PN}_5\text{C}_{10}\text{O}_7\text{H}_{12}$	1	8.61	9.65	21.11

(a) Water					
		Σb	\bar{b} (10^{-12} cm \AA^{-3})		
H_2O		-0.168	-0.00562		
D_2O		1.915	+0.06404		

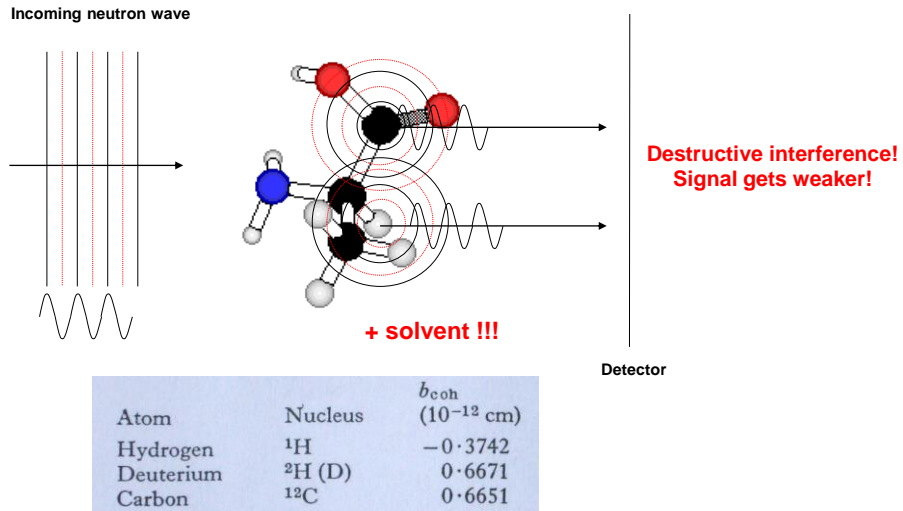
(b) Amino acids and proteins						
Amino acid	Chemical composition	H (ex)	b_{tot} (H_2O) (10^{-12} cm)	b_{tot} (D_2O) (10^{-12} cm)	b_{tot} (deuterated) (10^{-12} cm)	$V(\text{\AA}^3)$
Glycine	C_2NOH_3	1	1.728	2.769	4.85	66.4
Alanine	C_3NOH_5	1	1.645	2.686	6.852	91.5
Valine	C_6NOH_9	1	1.479	2.520	10.854	141.7
Leucine	$\text{C}_8\text{NOH}_{11}$	1	1.396	2.437	12.850	167.9
Isoleucine	$\text{C}_9\text{NOH}_{11}$	1	1.396	2.437	12.850	168.8
Phenylalanine	C_9NOH_9	1	4.139	5.180	13.51	203.4
Tyrosine	$\text{C}_9\text{NO}_2\text{H}_9$	2	4.719	6.802	14.09	203.6
Tryptophan	$\text{C}_{11}\text{NO}_2\text{H}_9$	2	6.035	8.118	16.45	237.6
Aspartic acid	$\text{C}_4\text{NO}_4\text{H}_4$	1	3.845	4.886	8.010	113.6
Glutamic acid	$\text{C}_5\text{NO}_4\text{H}_4$	1	3.762	4.803	10.01	140.6
Serine	$\text{C}_3\text{NO}_2\text{H}_5$	2	2.225	4.308	7.432	99.1
Threonine	$\text{C}_4\text{NO}_2\text{H}_7$	2	2.142	4.224	9.431	122.1
Asparagine	$\text{C}_4\text{N}_2\text{O}_4\text{H}_4$	3	3.456	6.580	9.704	135.2
Glutamine	$\text{C}_5\text{N}_2\text{O}_4\text{H}_4$	3	3.373	6.497	11.70	161.1
Lysine	$\text{C}_6\text{N}_2\text{O}_4\text{H}_8$	4	1.586	5.752	15.12	176.2
Arginine	$\text{C}_6\text{N}_4\text{O}_4\text{H}_8$	6	3.466	9.714	17.00	180.8
Histidine	$\text{C}_6\text{N}_3\text{O}_4\text{H}_6$	1.5	4.959	6.521	11.73	167.3
Methionine	$\text{C}_5\text{NOS}_2\text{H}_9$	1	1.764	2.805	11.14	170.8
Cysteine	$\text{C}_3\text{NOS}_2\text{H}_5$	2	1.930	4.013	7.137	105.6
Proline	C_5NOH_7	0	2.227	2.227	9.516	129.3

$$\bar{b} = \rho_{protein} = \sum_j \frac{b_j}{V}$$

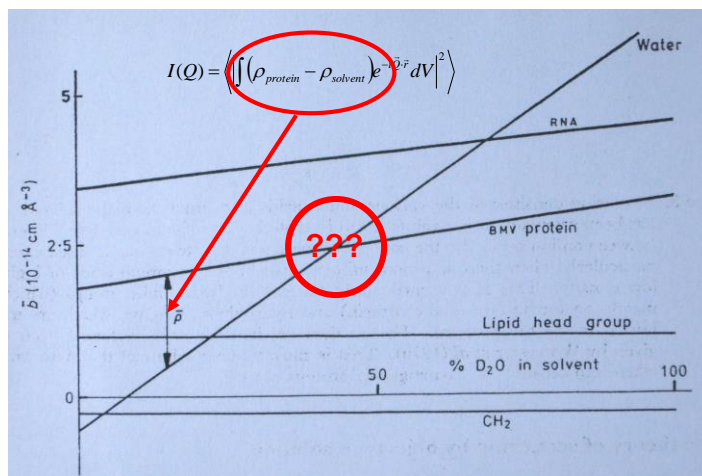
Example glycine in H_2O :
 $\rho = [2 \cdot 0.67 + 0.94 + 0.58 + 3 \cdot (-0.37)] 10^{-12} \text{cm} / 66.4 \text{\AA}^3$
 $= 2.68 \cdot 10^{-14} \text{cm} / \text{\AA}^3 = 2.68 \cdot 10^{10} \text{cm}^{-2}$

Jacrot, B. (1976) Rep. Prog. Phys. 39, 911-953.

Destructive interference in SANS

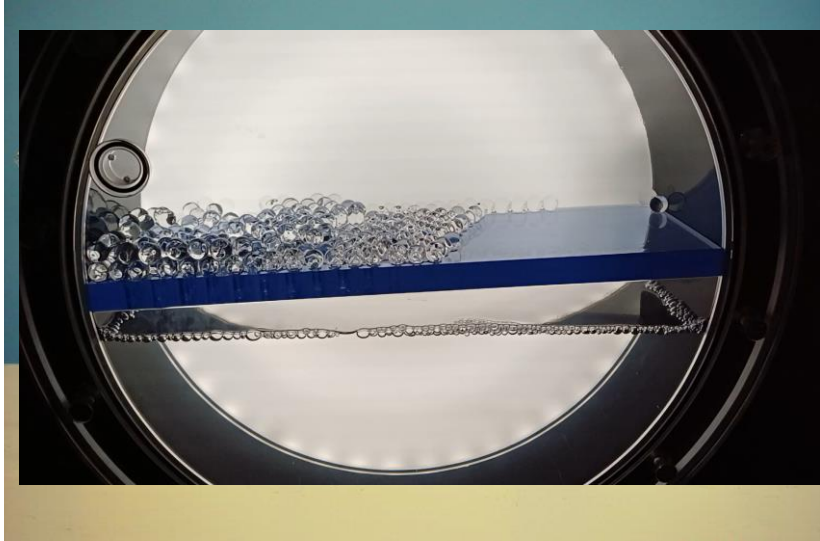


Natural Contrast

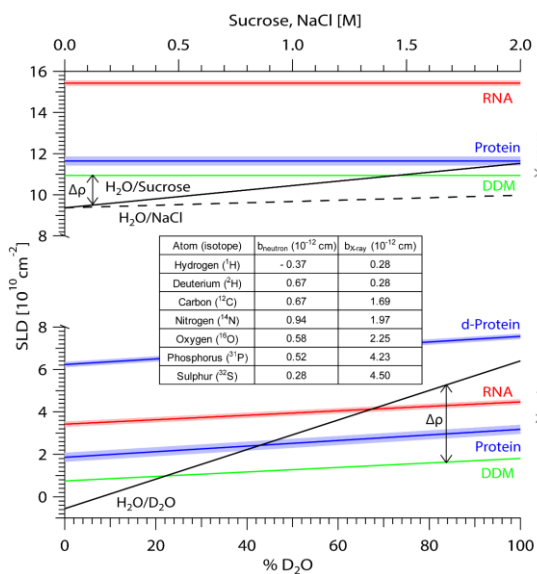


In practice, all biomacromolecules can be matched in SANS, i.e. made **invisible!!!**
Not so easy with SAXS...

An analogon in optics: refractive index



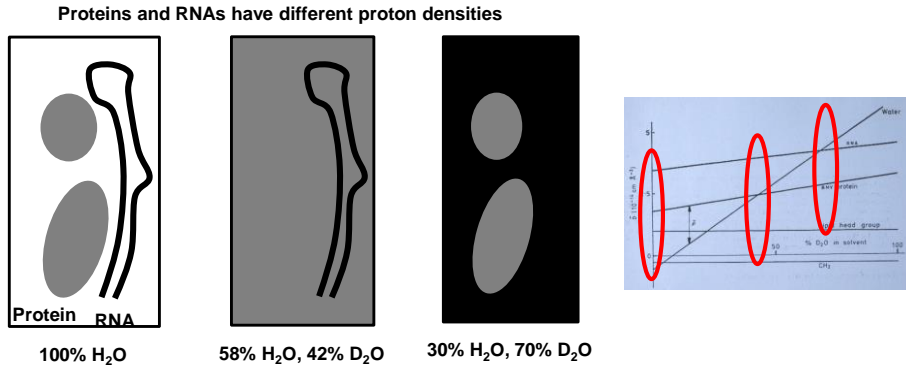
SAXS and contrast variation?



- 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**

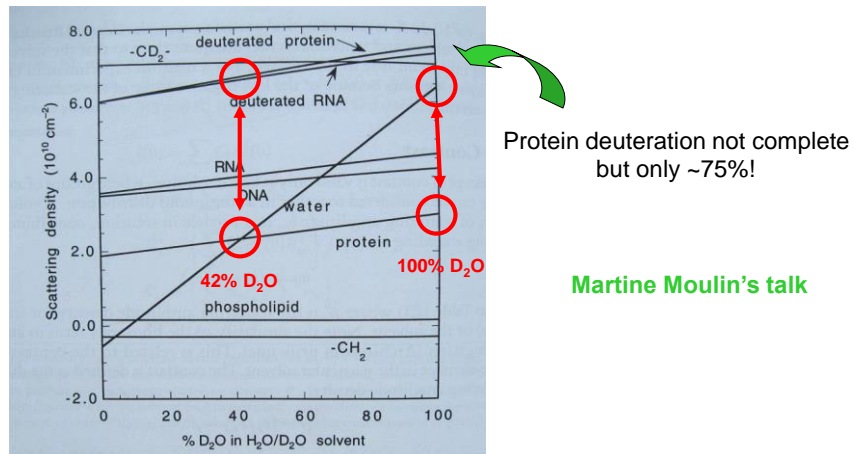
Mahieu, E. & Gabel, F. (2018). *Acta Cryst.* **D74**(Pt 8), 715-726.

Contrast variation in SANS: natural contrast



Also possible for protein-protein complexes (deuteration)!

Artificial contrast using deuteration



Careful at **high D₂O** levels in the solvent:
favours **oligomerisation/aggregation!**

Experimental determination of the zero contrast point (“contrast match point”)

$$I(Q) = \left\langle \left| \int (\rho_{\text{protein}} - \rho_{\text{solvent}}) e^{-i\vec{Q}\cdot\vec{r}} dV \right|^2 \right\rangle$$

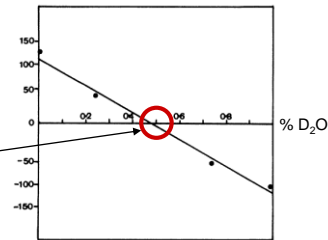
$$\rho_{\text{protein}} = \frac{\sum b_i}{V_{\text{prot}}}$$

$$\xrightarrow{Q=0} I(0) \sim C \cdot \left[\left(\frac{\sum b_i}{V_{\text{prot}}} - \rho_{\text{solvent}} \right) V_{\text{prot}} \right]^2$$

Contrast match-point: $I(0) = 0$

Measure $I(0)$ experimentally (Guinier!) and trace the following expression as a function of $\text{H}_2\text{O}/\text{D}_2\text{O}$:

$$\sqrt{\frac{I(0)}{C}} \sim (\rho_{\text{prot}} - \rho_{\text{solvent}}) V_{\text{prot}}$$



“Contrast match-point”

Guinier approximation and radius of gyration

$$I(Q) \approx I(0) \exp\left[-\frac{1}{3} R_g^2 Q^2\right] \quad R_g Q \leq 1 \dots 1.3$$

(from expansion of Debye equation)

$$\ln[I(Q)] \approx \ln[I(0)] - \frac{1}{3} R_g^2 Q^2$$

Radius of gyration:

$$R_g^2 = \frac{1}{M} \sum_i m_i r_i^2$$

Contrast x volume



“See-saw analogy”

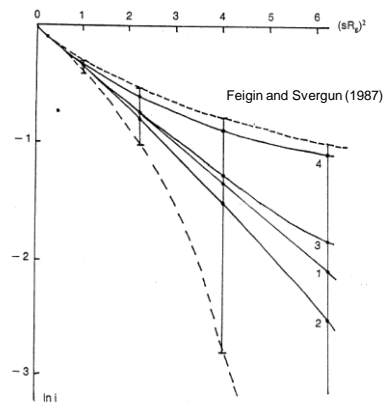
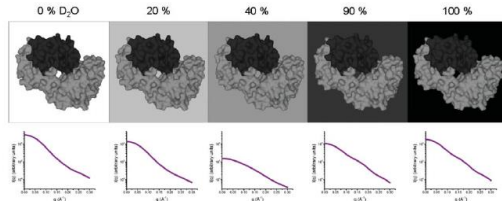


Figure 3.3. Accuracy of the Guinier law: (1) Guinier approximation with estimate (3.25); 2)-(4) correspond to scattering by a solid sphere, an infinitely thin disk, and an infinitely one rod.

For a given molecular weight, a sphere has the smallest R_g , i.e. it is the most compact object

Protein-protein complexes



Jacques and Trehella (2010)
Prot. Sci. **19**, 642-657

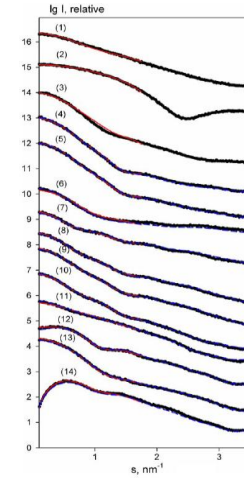
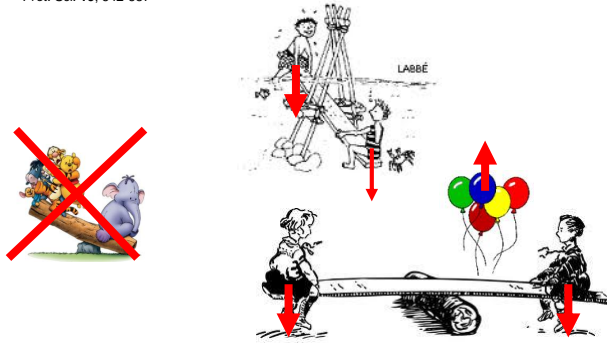
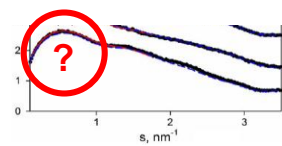


Fig. 5 Scattering profiles from the ternary complex. 1-4 are X-ray scattering curves of the DNA, theroctasin, polymerase and the entire complex, respectively; 5-9 are neutron scattering patterns from protonated complex in 0, 40, 25, 70 and 100% D₂O, respectively; 10-14 are neutron scattering patterns from the complex with perdeuterated theroctasin in 0, 40, 70, 100 and 55% D₂O, respectively. Notations of the simulated data and fit are as in Fig. 3. The patterns are displaced in logarithmic scale for better visualization.

Petoukhov and Svergun (2006)
Eur. Biophys. J. **35**, 567-576

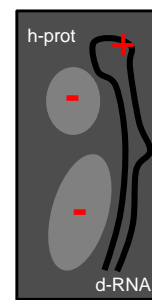
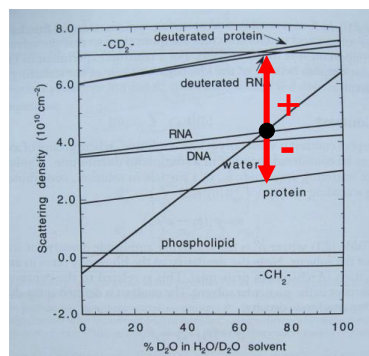
Negative radii of gyration!?



$$\ln[I(Q)] \approx \ln[I(0)] - \frac{1}{3} R_g^2 Q^2$$

$$R_g^2 = \frac{1}{M} \sum_i m_i r_i^2 = \frac{1}{M} \sum_i \Delta\rho_i V_i r_i^2$$

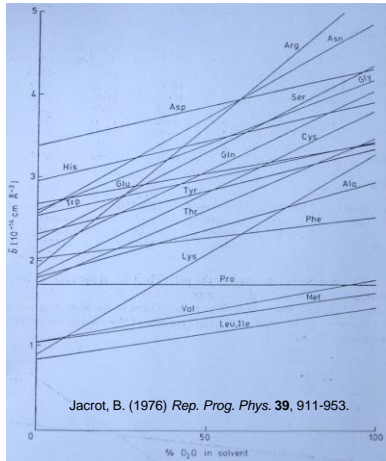
$$I(Q) = \left\langle \left| \sum_j b_j e^{-i\vec{Q}\cdot\vec{r}_j} \right|^2 \right\rangle$$



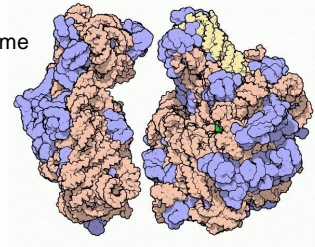
~ 70% D₂O

- 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

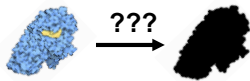
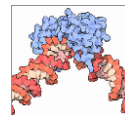
Can proteins be considered as homogeneous particles?



Ribosome



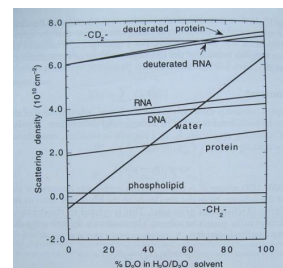
Transcription factor



Depends on the context and on the lengthscale studied!

Summary: homogeneity of biomacromolecules in SANS experiments

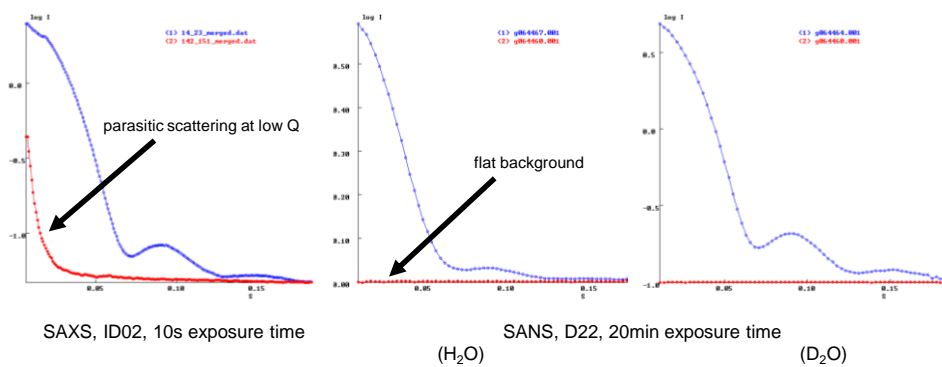
- Proteins can be considered as homogeneous if the scattering density fluctuations are **not ordered** (e.g. core vs. outer shell) and/or if they occur on a **length-scale much smaller** than the overall protein dimension and/or the molecules of interest to be studied in a complex
- RNA and DNA are **more 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

SAXS vs SANS: some practical aspects

	sample amount	flux	contrast
SAXS	~15 μ l @ 1-10 mg/ml	high	weak
SANS	~150 μ l @ 1-10 mg/ml	low	high (D_2O)



SANS vs SAXS instruments



Quartz cuvette (SANS)

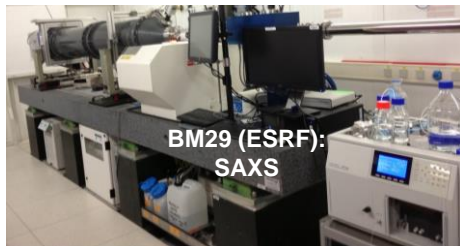
- easy manipulation with pipettes
- multi-sample holder
- no radiation damage!

Exposure times:

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

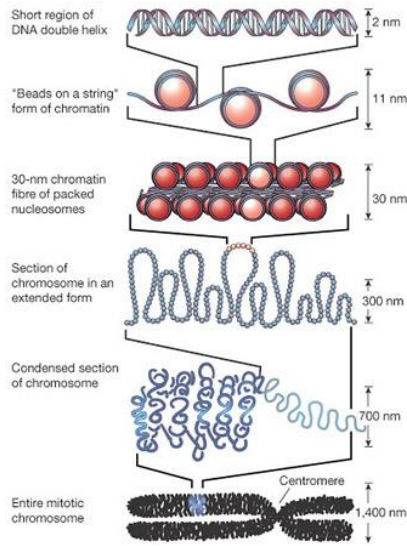
Including sample change:

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



Historical example 1: Chromatin

The chromatin structure



Contrast variation: relative arrangement of DNA and protein

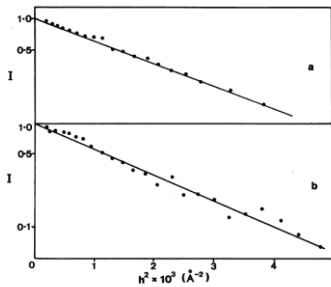


Figure 2: Examples of Guinier plots obtained from chromatin particles (a) Particles in D_2O (b) in H_2O . Semi-log plot of intensity (I) versus h^2 , where $h = 4\pi \sin \theta / \lambda$ and 2θ is the angle of scattering. Each data set is fitted with a straight line by a variance-weighted least squares procedure. The data are arbitrarily 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

TABLE 1

Percentage D_2O in Buffer	Particle Concentration mg/ml	Radius of Gyration (not corrected) \AA	Radius of Gyration (corrected for slit smearing) \AA
100	11	38.4 ± 0.3	39.1 ± 0.3
100	5	38.2 ± 0.3	
100	2.5	35.9 ± 1.1	
75	11	36.3 ± 0.9	36.8 ± 0.9
25	11	45.3 ± 6.0	49.9 ± 5.5
0	11	41.8 ± 0.7	43.0 ± 0.7

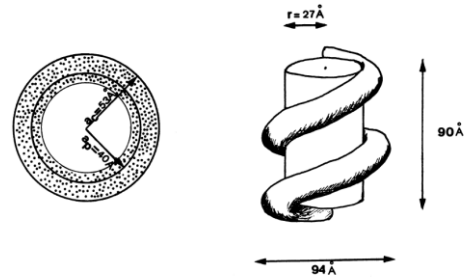
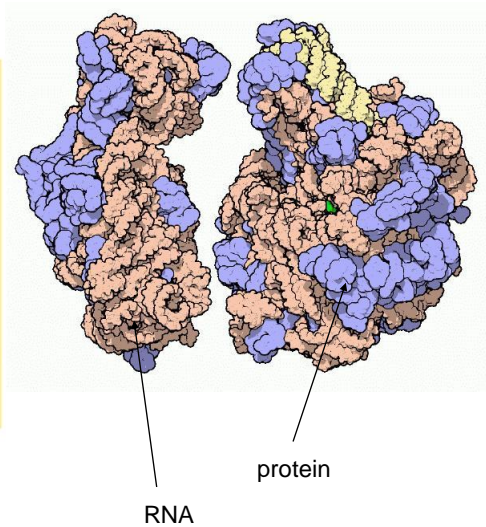
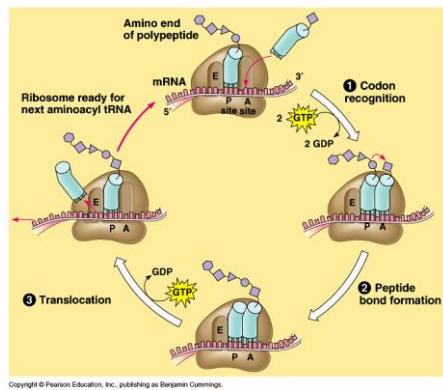


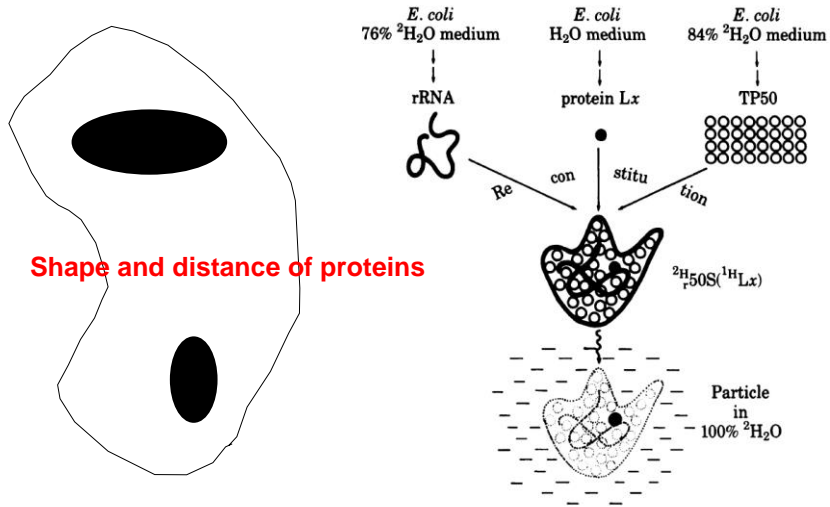
Figure 5: Two possible kinds of structure for the chromatin particle containing 140 base pairs of DNA and eight histones. (a) A spherical particle with overall diameter $a_o = 53 \text{\AA}$, derived from $R_g = 41.1$, in which the inner protein core has radius $a_p = 40$ from the experimental $R_g = 30.6 \text{\AA}$. The region occupying the DNA is shaded. (b) A cylindrical model in which two turns of helix with pitch 45\AA and radius 37\AA wound on an inner protein core of radius 27\AA .

Historical example 2: The ribosome

The ribosome structure

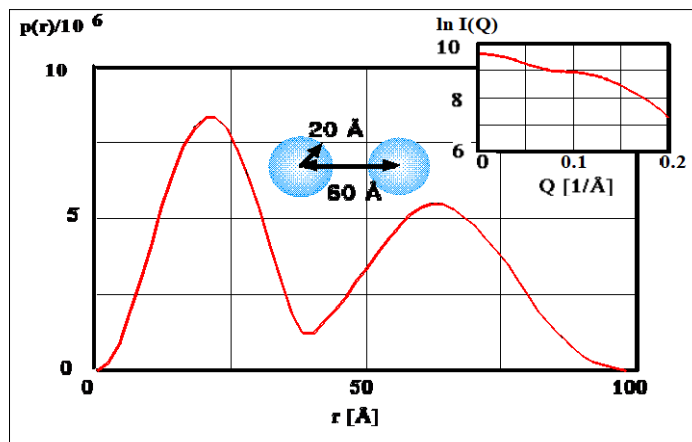


The “glassy” ribosome



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

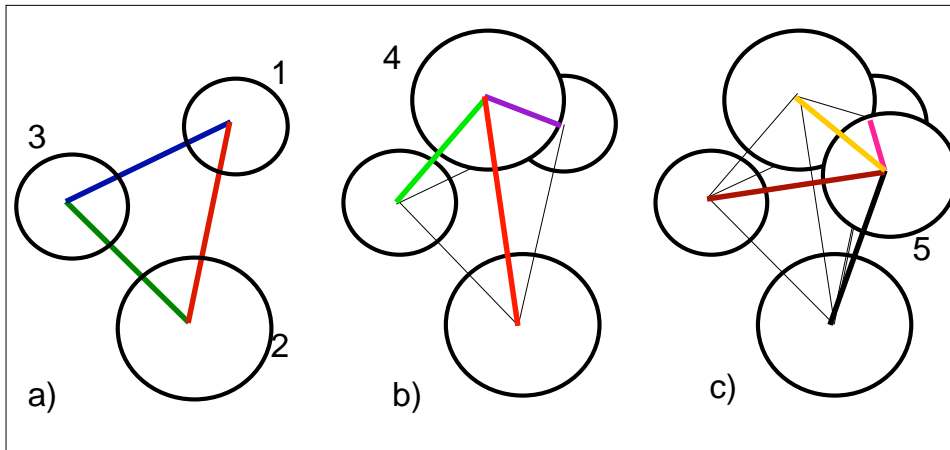
Triangulation



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

courtesy Roland May

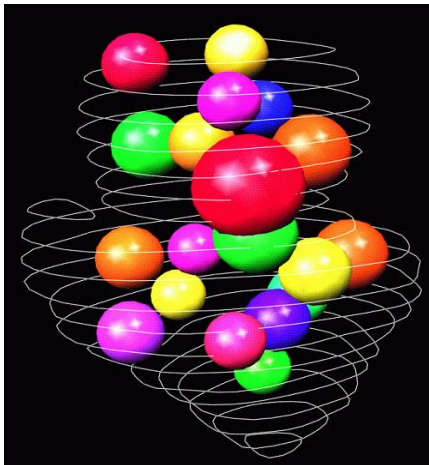
Triangulation



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

Ribosome at low resolution



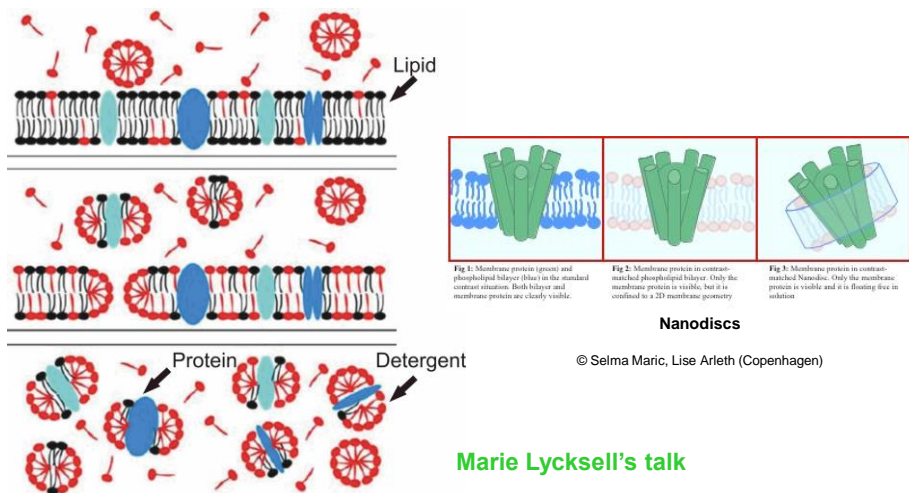
M.S. Capel, D.M. Engelman, B.R. Freeborn, M. Kjeldgaard, J.A. Langer, V. Ramakrishnan, D.G. Schindler, D.K. Schneider, B.P. Schoenborn, I.-Y. Sillers, S. Yabuki, P.B. Moore (1987) *Science* **238**, 1403-1406

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.

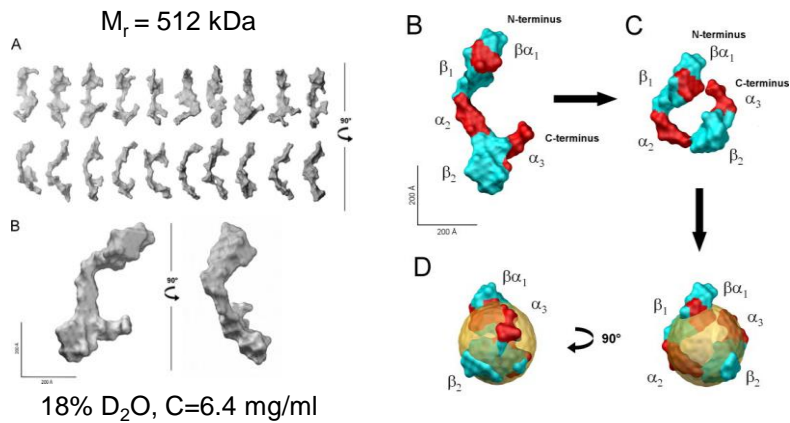
courtesy Roland May

EXAMPLE 3: Membrane proteins

Membrane proteins and lipids/detergents

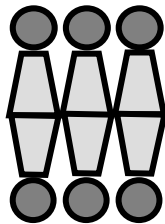


Modeling of a membrane protein: apolipoprotein B-100



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

Scattering contrast of lipids: heterogeneity



H-lipid headgroup 'denser'

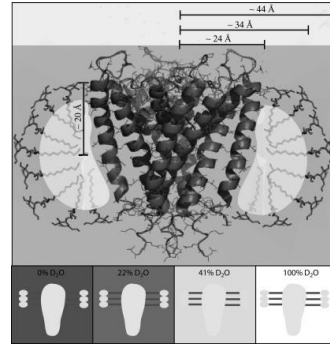
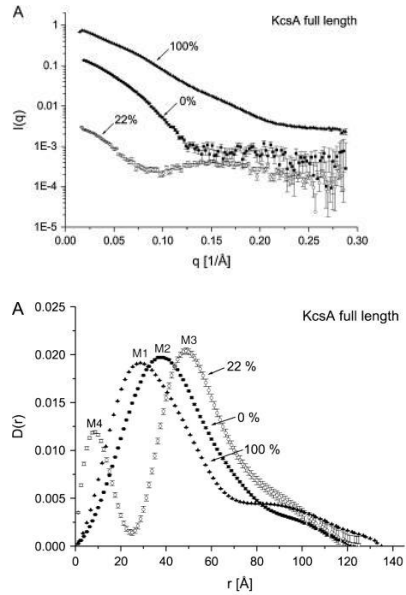


D-lipid

(d) Lipids

	b (10^{-12} cm)	\bar{b} (10^{-12} cm \AA^{-3})	b (deuterated) (10^{-12} cm)	\bar{b} (10^{-12} cm \AA^{-3})
Phosphorylcholine				
CH ₂	-0.0833	-0.0031	2	0.0744
CH ₃	-0.458	-0.0085	2.67	0.0495
C ₅ H ₁₃ NPO ₄	2.24	0.011	15.76	0.072

pH-induced transition in KcsA



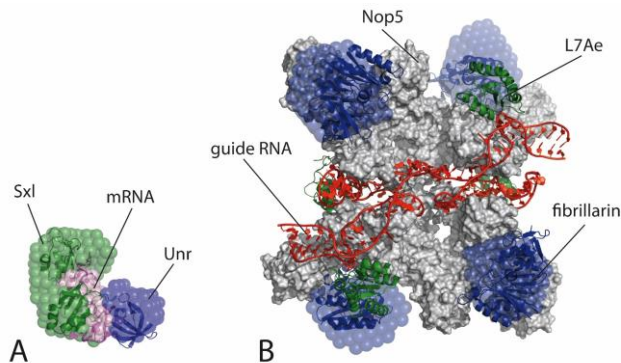
concentration. On the basis of published detergent scattering match points, we estimated the detergent scattering match point for decyl- β -D-maltopyranoside ($C_{22}H_{42}O_{11}$) at ~22% D_2O solvent concentration (24), thereby minimizing the contribution of the solubilizing detergent micelle to the observed neutron scattering. However, the fluctuation of scattering length of the DM detergent molecule must be stressed. This is due to the individual scattering match points of the polar maltoside headgroup of ~46% vs. ~3% D_2O solvent concentration for the hydrophobic decyl side chain. The overall match point at 22% D_2O solvent concentration, mini-

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

EXAMPLE 4:

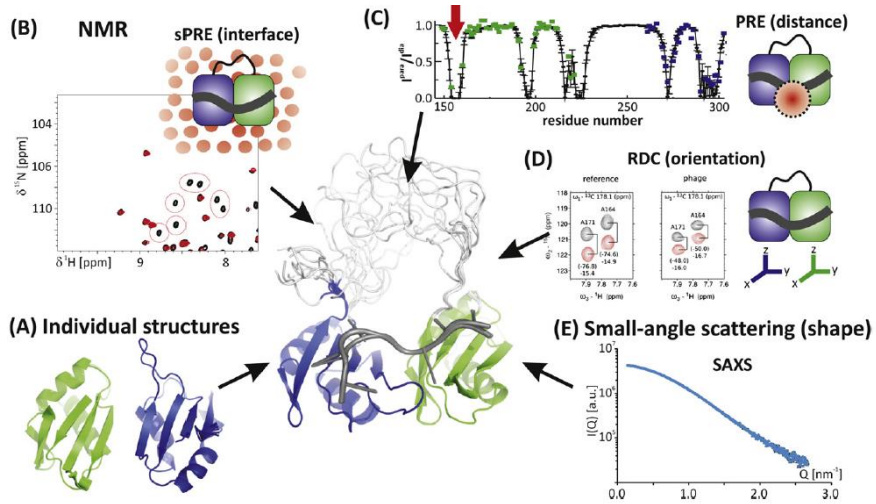
Sophisticated approaches using SANS (SAXS) and NMR:

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.

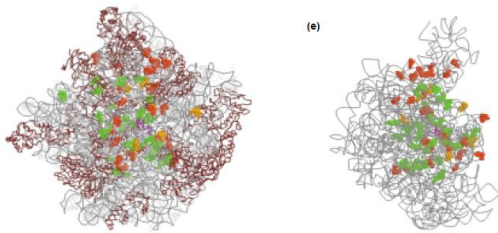
Combining NMR with SAS



Madl, T., Gabel, F. and Sattler, M. (2011) *J. Struct. Biol.* **173**, 472-482

Pau Bernado's talk

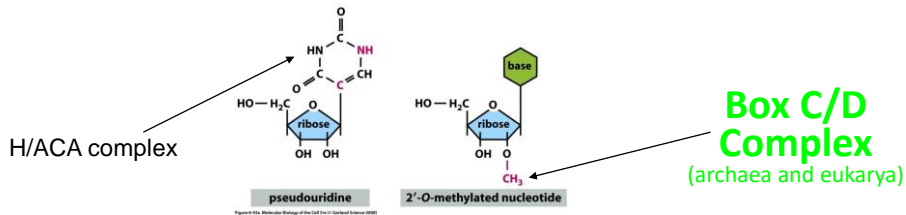
rRNA modifications and function



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.



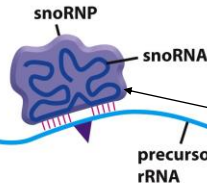
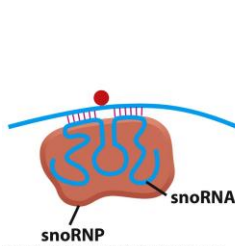
Number of modifications: bacteria < archaea < eukarya

RNA modifications: snoRNPs, snoRNAs and box C/D

snoRNP = "Small nucleolar Ribonucleo-Protein"

Only in archaea and eukaryotes,
not in bacteria

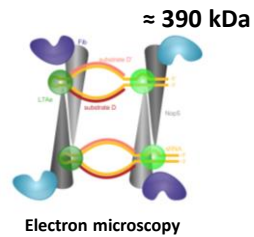
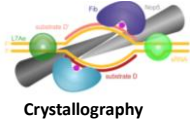
snoRNP = sRNP in archaea



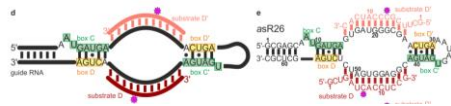
"Guide RNA"
(> 100 in humans!)

Figure 6-43b: Molecular Biology of the Cell 5/e (© Garland Science 2008)

Two different architectures
have been proposed:



- Where is the sRNA situated?
- What is mechanism of methylation?
- Why two asymmetric methylation sites?



Structural refinement strategy

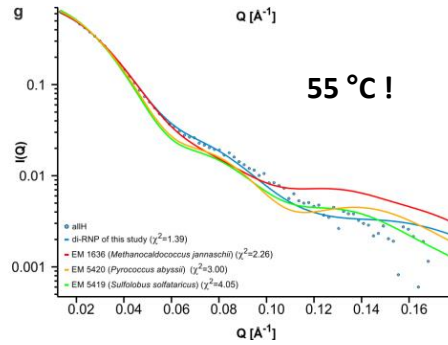
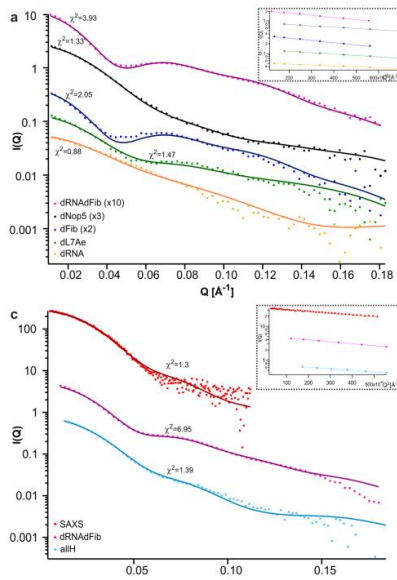
System from *Pyrococcus furiosus*

Rigid building blocks from crystal structures
+
flexible joints

Positions of spin labels

Methyl resonances CSPs of Fib and L7Ae
+
452 PRE distance restraints (apo-complex)
+
SAXS/SANS

SANS (D22) and SAXS (BM29) data

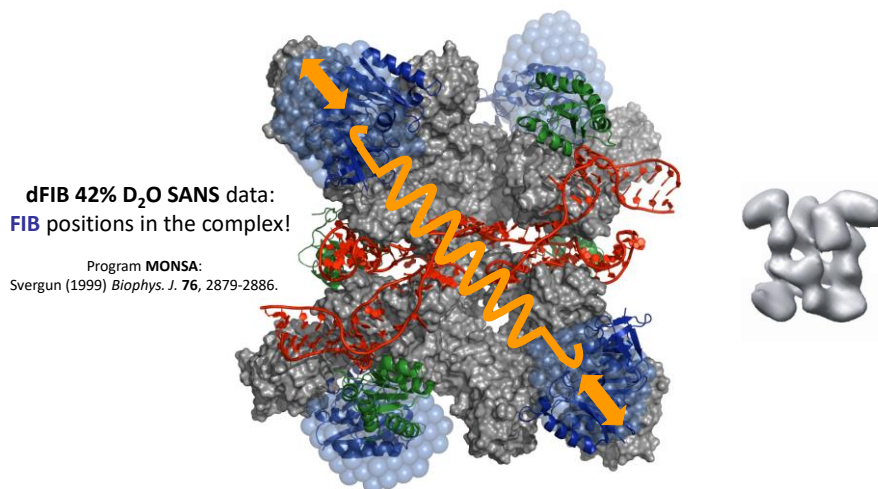


**A total of 26 SANS samples:
dFIB, dL7e, dRNA, dNOP5, dRNAdFIB
0, 42, 70% D₂O**

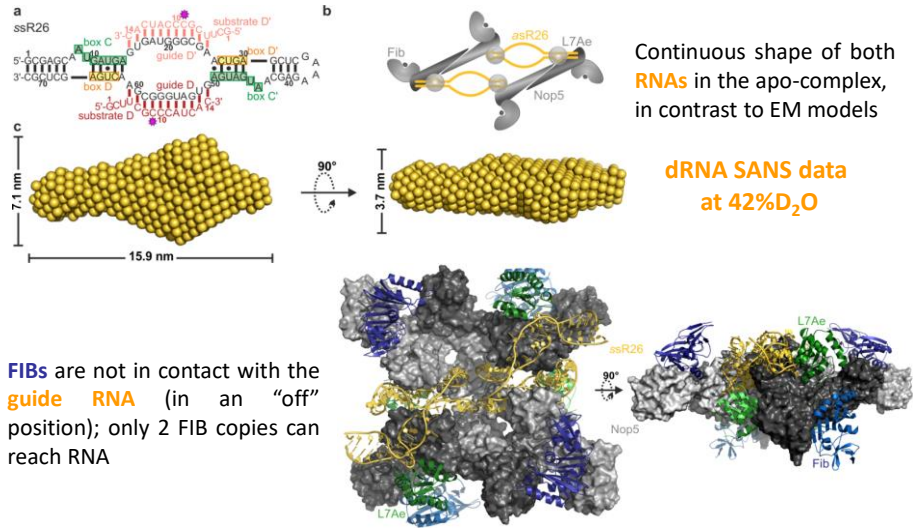
D22 (ILL) Local contact: **Anne Martel**

BM29 (ESRF) Local contact: **Petra Pernot**

Relative positions of FIB proteins within the complex from SANS data

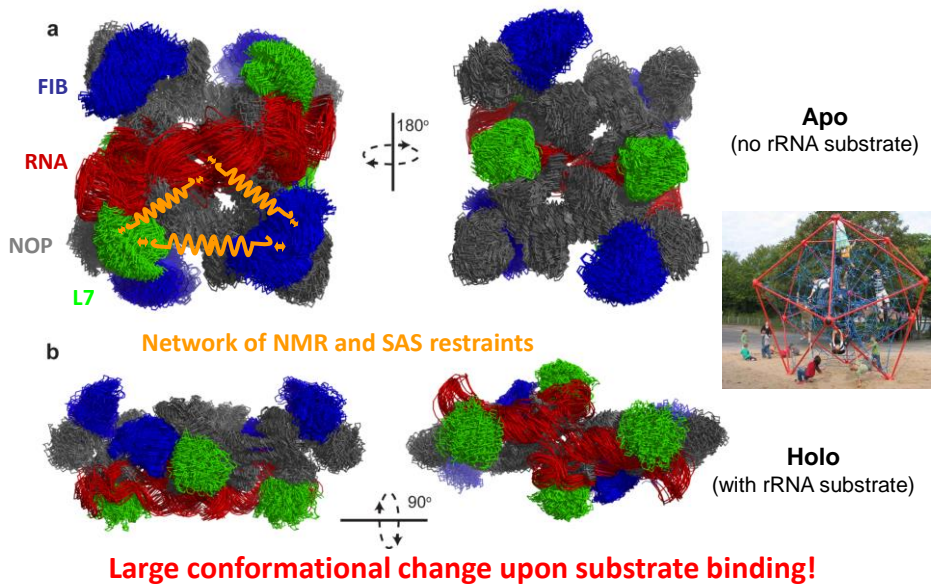


RNA shape within the complex from SANS data

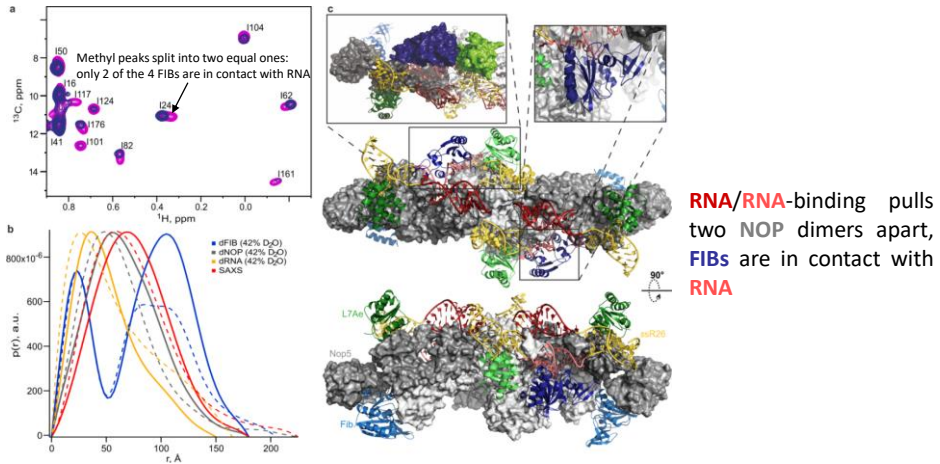


Lapinaite, A., Simon, B., Skjaerven, L., Rakwalska-Bange, M., Gabel, F. and Carlomagno T. (2013) The structure of the box C/D enzyme reveals regulation of RNA methylation. *Nature* **502**(7472), 519-523.

Family of refined structures

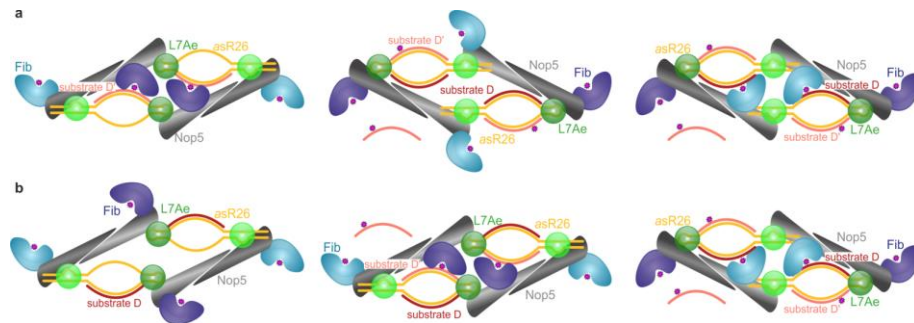


The holo complex



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

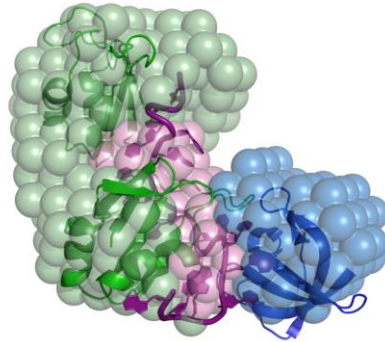
Proposed model for the sequential methylation and conformational changes



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...**

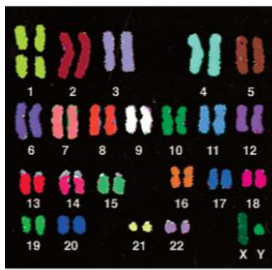
A novel mechanism for translational regulation in *Drosophila melanogaster*



Hennig J, Miliiti 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.

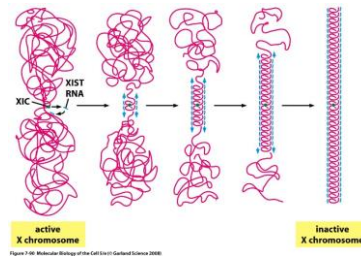
Dosage compensation

Human (male) karyotype



Females: XX

XIST gene silencing system in female mammals

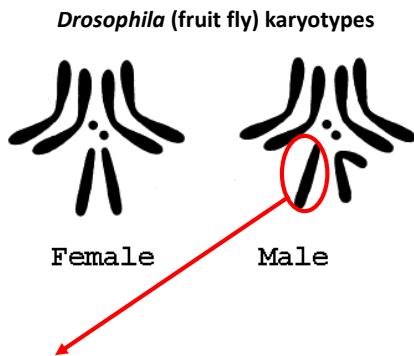


Unequal proteins amounts from XX and XY pairs: needs **compensation mechanisms**
(Klinefelter syndrome in humans: XXY)



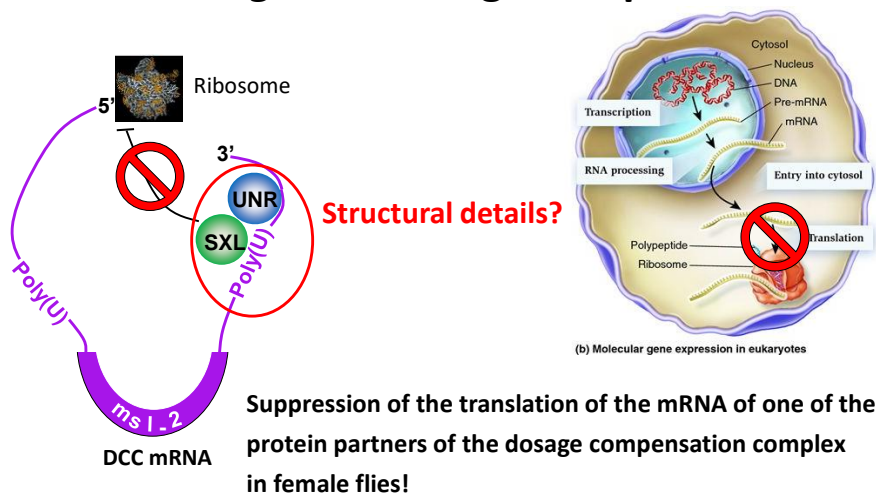
Calico cat

Dosage compensation in *D. melanogaster*



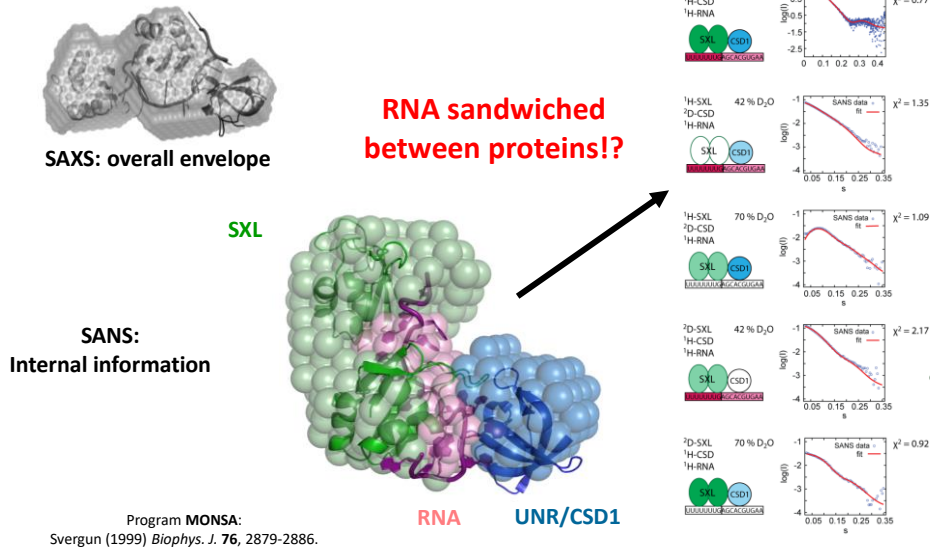
- **Up-regulated** by “DCC” (Dosage compensation complex) constituted of 5 proteins and 2 non-coding RNAs
- Female-specific protein “SXL” (sex-lethal) **silences the expression of a protein of the DCC complex** in females by binding to its mRNA transcript and inhibiting its interaction with the ribosome

Translational repression in *D. melanogaster* dosage compensation

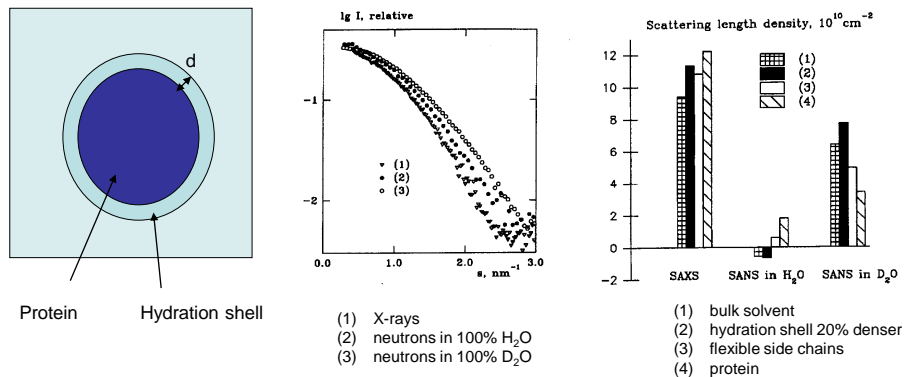


→ **Combined NMR/SAXS/SANS structural study**

SANS-specific information



Example 5: SANS provides more than shapes!



$$I(Q) = \left\langle \left| A_a(\vec{Q}) - \rho_s A_s(\vec{Q}) + \delta\rho_H A_H(\vec{Q}) \right|^2 \right\rangle_{\Omega}$$

$$\delta\rho_H \approx 0.1\rho_s$$

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

Influence of surface charge?

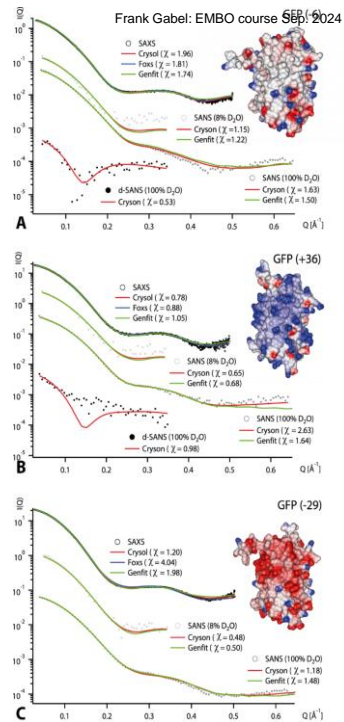
	GFP(-6)	GFP(-29)	GFP(+36)
SAXS			
HS $\Delta\rho / \rho_{bulk}$ (%)	5.00 ± 0.57	8.80 ± 0.62	4.13 ± 0.31
SANS (100 %)			
HS $\Delta\rho / \rho_{bulk}$ (%)	0.00 ± 0.00	2.65 ± 0.60	0.27 ± 0.47
SANS (8 %)			
HS $\Delta\rho / \rho_{bulk}$ (%)	(0.00±0.00)	(0.00±0.00)	(0.00±0.00)
d-SANS (100 %)			
HS $\Delta\rho / \rho_{bulk}$ (%)	0.95 ± 0.35	N.D.	0.00 ± 0.00

Residue-specific densities!

Denser around acidic residues

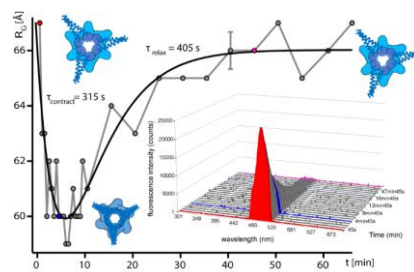
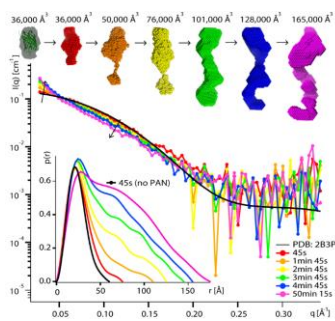


Kim, H.S., Martel, A., Girard, E., Moulin, M., Härtlein, M., Madern, D., Blackledge, M., Franzetti, B. and Gabel, F. (2016) Solution scattering of X-rays and neutrons on supercharged proteins reveals residue-specific modifications of the hydration shell. *Biophys. J.* 110(10), 2185-2194.

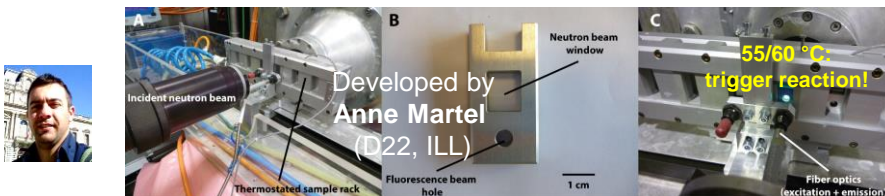


Example 6: time-resolved SANS

Frank Gabel: EMBO course Sep. 2024

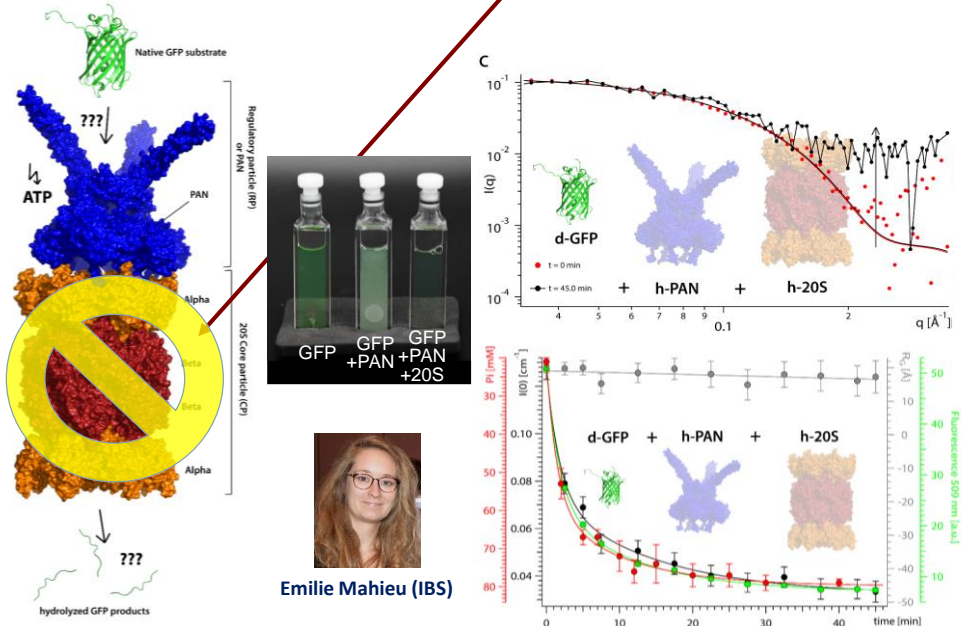


30 second frames!

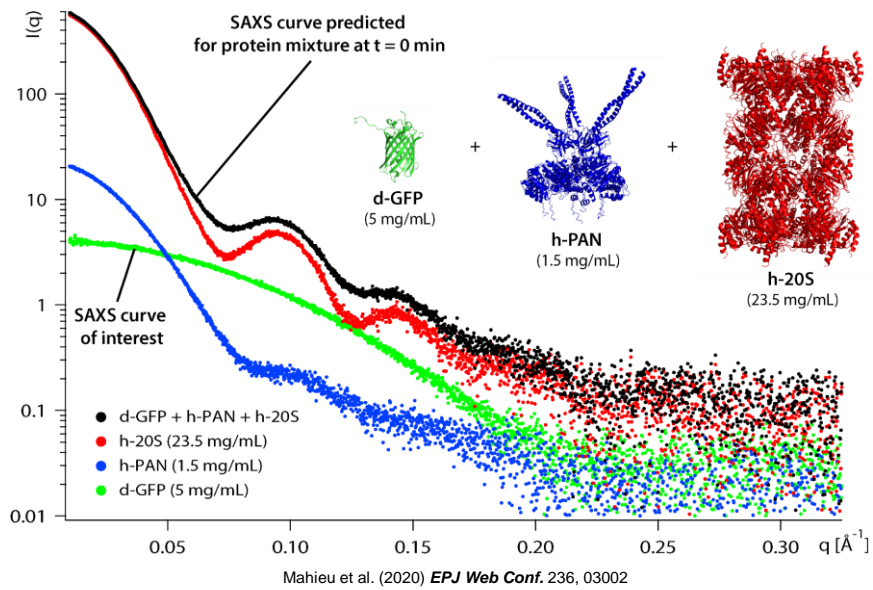


Ibrahim, Z., Martel, A., Moulin, M., Kim, H.S., Härtlein, M., Franzetti, B. and Gabel, F. (2017) *Sci. Rep.* 7, 40948.

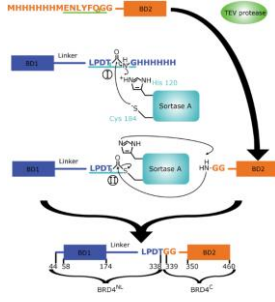
And in the presence of the proteolytic core particle 20S?



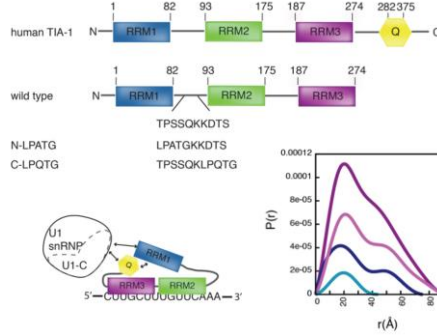
Could this have been done by SAXS?



Example 7: segmental labeling



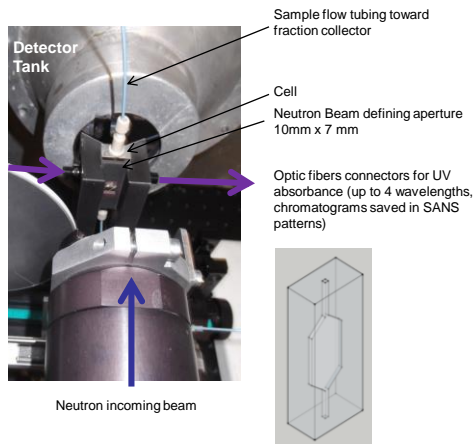
Williams et al. (2016) *PLoS ONE* 11(4):e0154607



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

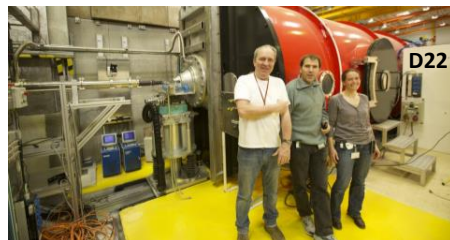
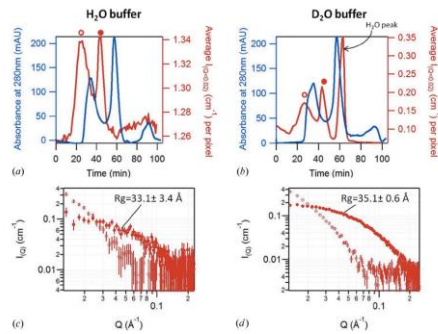
➔ More detailed insight into multi-domain proteins

Example 8: SEC-SANS



Cell: 8mm suprasil quartz + 1mm sample (T=82%)

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 responsables, **assist** during experiments
- access (for non-industrial use) is in general **free**
- **no maintenance**, user friendly (software etc...)



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:

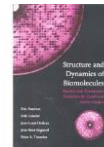
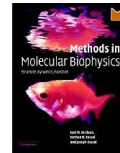
- Lovesey: Theory of Neutron Scattering from Condensed Matter (Clarendon, 1986)
- Geissler et al.: Structure and dynamics of biomolecules (Oxford University Press, 2000)

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!

Soft Matter Support Facilities (PSCM)
Infrastructure for ambitious large-scale soft matter research projects

The soft matter facilities at the ILL aim to offer a wide range of complementary techniques to the user community to allow improved sample preparation and beam-time optimisation.

Most studied topics

- Biomembranes and lipid assemblies
- Colloidal self-assembly (surfactants, polymers, micelles, etc.)
- Smart Coatings
- Structure, dynamics, and function of proteins

Our major equipment

Light scattering

- LS 3D static and dynamic light scattering
- ALV multiangle static and dynamic light scattering
- Corbourn portable light scattering
- Malvern particle size analyser

Solid and Liquid interfaces

- Beaglehole Picometer Light Ellipsometer
- Brewster Angle Microscope Nanofilm EMS
- 5x Langmuir troughs
- Kratos A11 Tensiometer and DS114 Drop shape analyser
- Q-Sense Quartz Crystal Microbalance E4

Calorimetry and viscometry

- 2 Differential scanning calorimeters (solids and liquids)
- 2 Densitometers and sound velocity meters

People: Leonardo Chiappisi, Martina Sandroni, Sandrine Verdon

Chemistry Lab Support Facilities
Enabling on-site sample handling and preparation

The laboratories are typically used for sample preparation, transfer to the sample holders, buffers preparation, substrates cleaning, sample dilution and deposition, sample hydration, dialysis, or thermal treatments before or during experiments. They are used by both soft and hard matter users.

Laboratory equipment

Besides standard equipment (fume hoods, scales, stirrers, pH-meters, ultrapure water,...) the laboratories are equipped with:

- Glovebox and glovebags under inert atmosphere
- Freeze dry
- Cold room and 80 °C freezer
- Centrifuge (standard large volume)
- Ovens and high-temperature furnaces (up to 1600 °C)
- Glove/booth fume hood for dry nanoparticles manipulation
- Schlenk line

Chemicals

A stock of standard chemicals is available. The needs for D₂O or specific chemicals are assessed before each experiment. The users can request chemicals to be ordered by ILL to be available onsite during the experiments. Costs will be invoiced after the experiment.

People: Martina Sandroni, Sandrine Verdon

ILL Lipid Lab (L-Lab)
Developing Advanced Models of Biological Membranes

Neutron scattering techniques are ideally suited for the study of lipid bilayers that are major components of cellular membranes. There's a dearth for biologically relevant deuterated lipids which is both expensive and difficult to synthesise them through chemical synthesis. Ideally, reconstituted microbial lipids extracted and purified from cells grown under deuterated conditions should work as model cellular membranes. The L-Lab has been successful in optimising methods to extract, purify and characterise deuterated lipids produced in yeast and bacteria.

People: Krishna Batchu

ILL Deuteration Lab (D-Lab)
Biomacromolecular Deuteration for Neutron Studies in Biology

The ILL D-Lab has been operating for more than 20 years as a dedicated platform to support the deuteration of biological molecules for neutron scattering experiments.

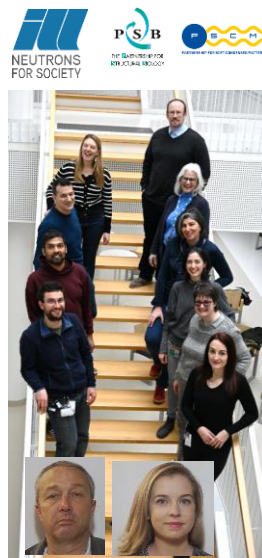
Activities of the D-Lab:

- Production of deuterated biomolecules for structural biology
- Development of a proposal system
- Method development and optimisation
- Biological organisation
- Isotope labelling strategies.

Capillary selections

- High cellulose, yeast and algae
- Purification of proteins and lipids by FPLC
- Chromatography of deuterated biomolecules.
- CIP
- HP
- Peptide synthesis, purification and characterisation.
- Protein crystallisation.

People: Valérie Laux, Juliette M. Devos, Martine Moulin



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



Gabel et al. (2019) *IUCr* 6, 521-525

water $\rho \approx 0.34 \text{ e}/\text{\AA}^3$

tail $\rho \approx 0.29 \text{ e}/\text{\AA}^3$

head $\rho \approx 0.52 \text{ e}/\text{\AA}^3$

