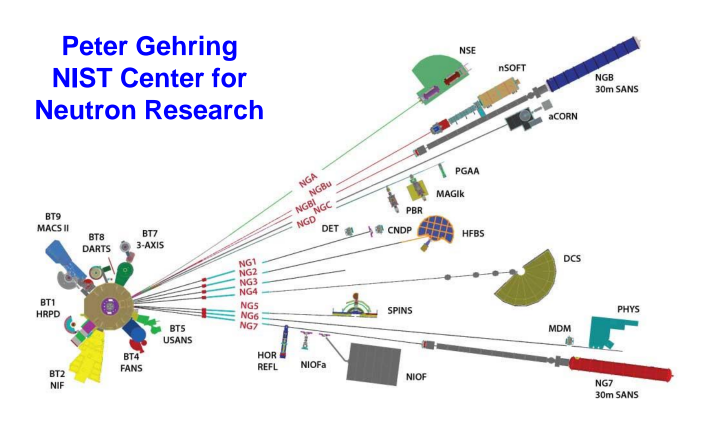


# Choosing the Right Spectrometer











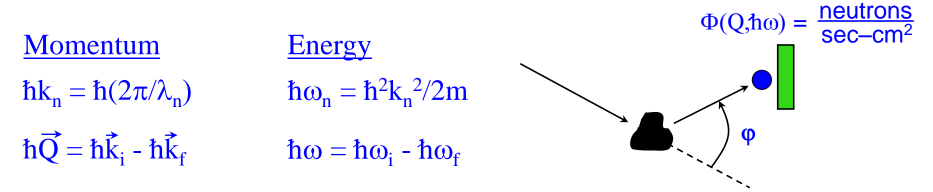




## Review: Main Messages of the Week



(1) Neutron scattering experiments measure the <u>flux</u> of neutrons scattered by a sample into a detector as a function of the <u>change</u> in neutron wave vector  $(\vec{Q})$  and energy  $(\hbar\omega)$ .



(2) The expressions for the scattered neutron flux  $\Phi$  depend on the positions and motions of atomic nuclei or unpaired electron spins.

$$\Phi = \mathbf{F}\{\vec{r}_{i}(t), \, \vec{r}_{j}(t), \, \vec{S}_{i}(t), \, \vec{S}_{j}(t)\}$$



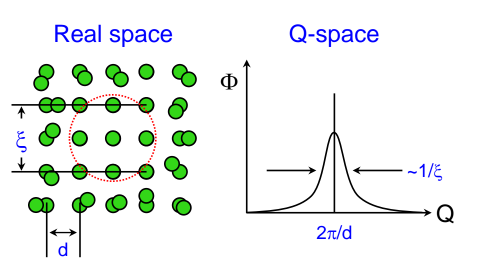
Φ provides information about <u>all</u> of these quantities!

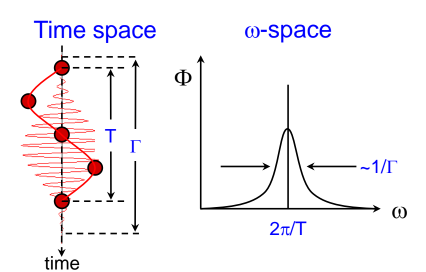
## Review: Main Messages of the Week



(3) The scattered neutron flux  $\Phi(\vec{Q},\hbar\omega)$  is proportional to the <u>space</u>  $(\vec{r})$  and <u>time</u> (t) Fourier transform of the <u>probability</u>  $G(\vec{r},t)$  of finding one or two atoms separated by a particular distance at a particular time.

$$\Phi \propto \frac{\partial^2 \sigma}{\partial \Omega \partial \omega} \propto \iint e^{i(\vec{Q}\cdot\vec{r}-\omega t)} G(\vec{r},t) d^3 \vec{r} dt$$





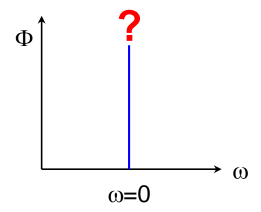
## Pop Quiz!



Question:

Can one measure elastic scattering from a liquid?



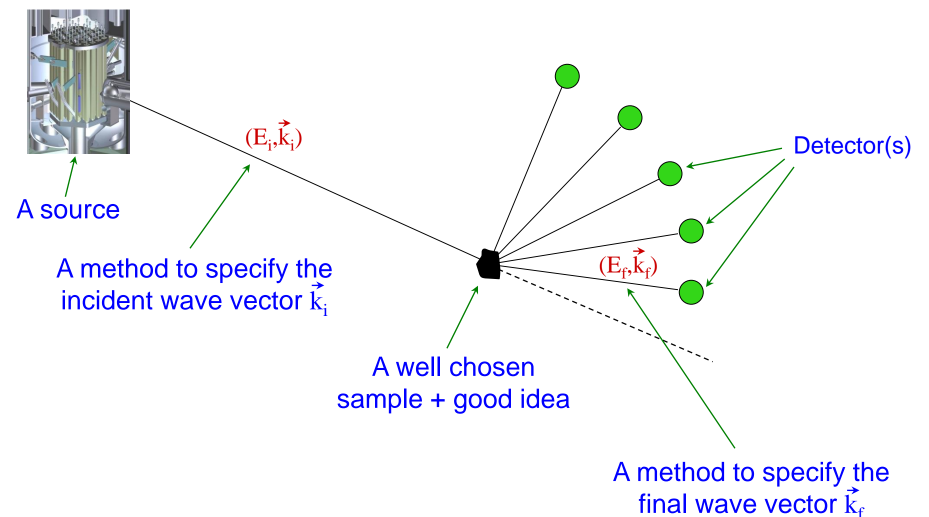


Why? Why not?

Hint: What is the correlation in time of one atom in a liquid with another atom a distance r away?

## Basics Elements of a Neutron Inelastic Scattering Experiment



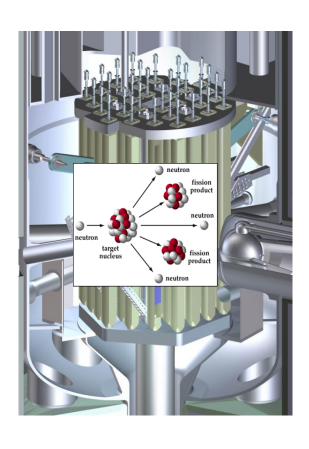


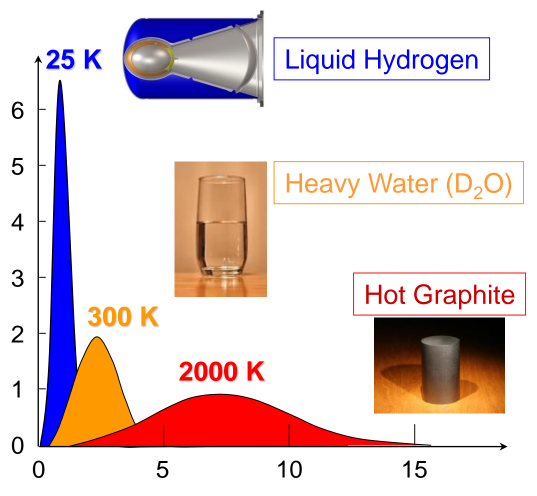
## Neutron Source: Moderation

Maxwellian Distribution

$$\Phi \sim v^3 e^{(-mv^2/2k_BT)}$$







"Fast" neutrons: v = 20,000 km/sec

Neutron velocity *v* (km/sec)

# Methods of Specifying and Measuring $\vec{k}_i$ and $\vec{k}_f$

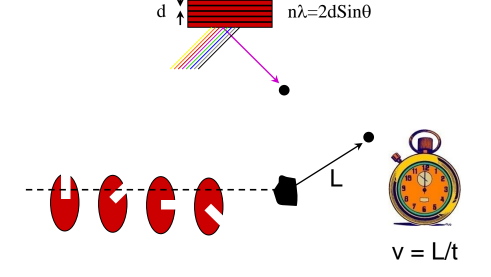


1. Bragg Diffraction

BT7, MACS, HFBS

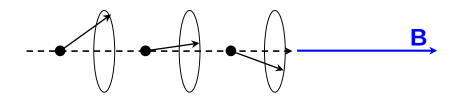
2. Time-of-Flight (TOF)

DCS, HFBS

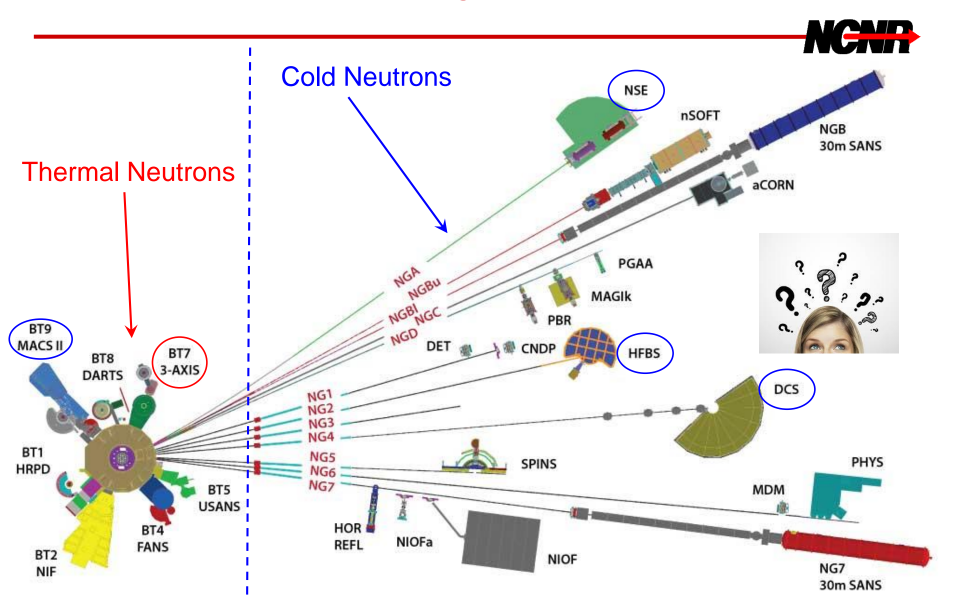


3. Larmor Precession

**NSE** 



## The NCNR Menagerie of Instruments



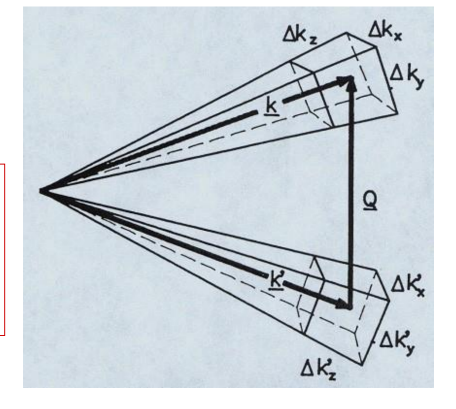
## Why So Many Different Spectrometers?

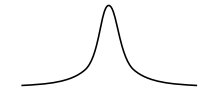


Because neutron scattering is an <u>intensity-limited</u> technique. Thus detector coverage and resolution MUST be tailored to the science.

Uncertainties in the neutron wavelength and direction imply  $\mathbf{Q}$  and  $\hbar\omega$  can only be defined with a finite precision.

The total signal in a scattering experiment is proportional to the resolution volume → better resolution leads to lower count rates! Choose carefully ...





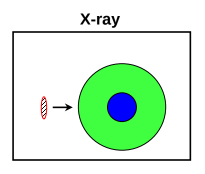


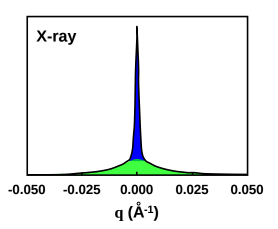


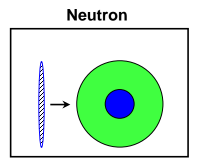
### **Q-Resolution Matters!**

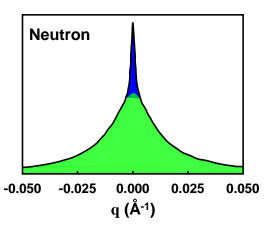


The "right" resolution depends on what you want to study.





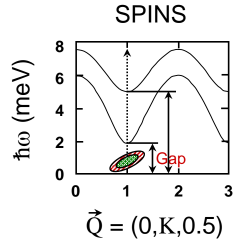


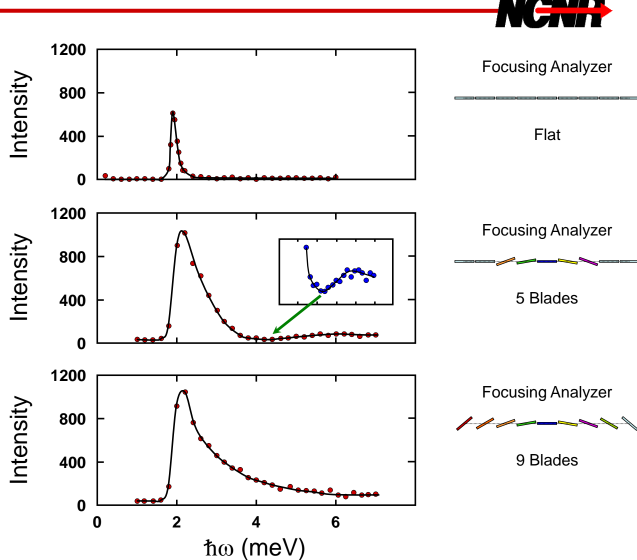


### ħω-Resolution Matters!



Another example ...





## How do I Choose the Right Spectrometer?



#### Two basic considerations:

- 1. What are the time scales ( $\hbar\omega$ ) of interest?
- 2. What are the length scales (Q) of interest?

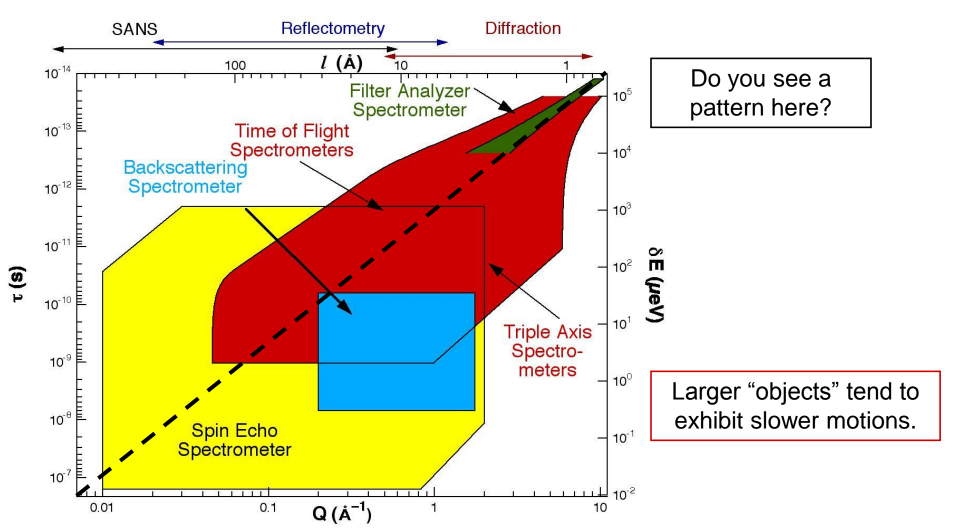
(Some spectrometers overlap → the choice may boil down to one of resolution)

#### Two additional considerations:

- 1. What energy resolution ( $\Delta\hbar\omega$ ) is required?
- 2. What momentum resolution ( $\Delta Q$ ) is required?

## Different Spectrometers Cover Different Regions of Phase Space





#### Rules of Thumb



1. What are the energies  $(\hbar\omega)$ , i.e. time scales  $(\Delta t \sim 1/\omega)$ , of interest?

 $\hbar\omega > 10-20 \text{ meV}$  - use a thermal triple-axis spectrometer like BT7.

 $\hbar\omega$  < 20-30  $\mu eV$  - use HFBS or NSE.

In between - use MACS or DCS or a cold-neutron triple-axis spectrometer like SPINS.

2. Make sure that the length scales L of the relevant motions lie within the range of the spectrometer. For example, consider the HFBS. ( $\mathbf{Q} \sim 2\pi/\mathbf{L}$ )

$$\mathbf{Q}_{\text{min}} = 0.25 \, \text{Å}^{-1} \rightarrow \mathbf{L}_{\text{max}} \sim 25 \, \text{Å}$$

$$\mathbf{Q}_{\text{max}} = 1.75 \, \text{Å}^{-1} \rightarrow \mathbf{L}_{\text{min}} \sim 3.5 \, \text{Å}$$

REMEMBER - **Q**<sub>min</sub> and **Q**<sub>max</sub> are <u>inversely</u> proportional to the incident neutron wavelength

#### More Rules of Thumb



Is your sample polycrystalline or amorphous?

Does ONLY the magnitude (not the direction) of **Q** matter?

Is the expected **Q**-dependence of the scattering weak?

This often means that you want to look at a large region of  $\mathbf{Q}$ - $\hbar\omega$  space, or that you can sum the data over a large region of  $\mathbf{Q}$ - $\hbar\omega$  space.

YES? Consider instruments with large analyzer areas.

NO? Consider using BT7, SPINS, or NSE.

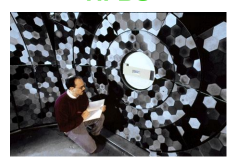
**MACS** 



**DCS** 



**HFBS** 



BT7



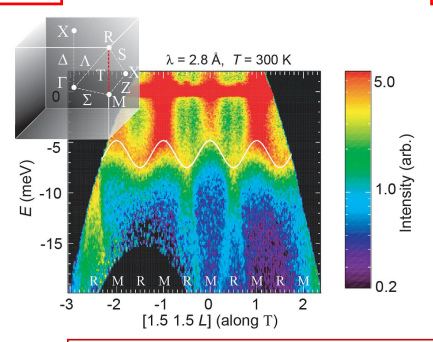
## Example: DCS versus BT7

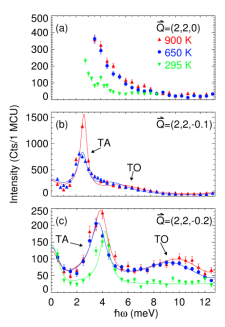


DCS Broad surveys in  $\mathbf{Q}$ - $\mathbf{\omega}$ 

BT7

Limited regions in  $\mathbf{Q}$ - $\mathbf{\omega}$ 





Rules of Thumb: (think carefully before violating)

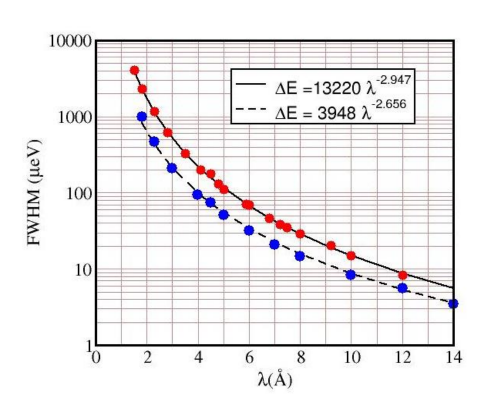
DCS, MACS – systems requiring resolution < 400  $\mu eV$ 

BT7 – single crystals

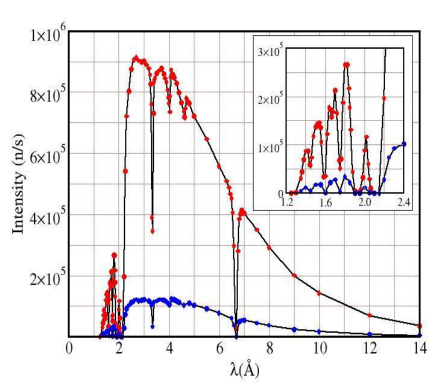
## Things to Consider When Choosing DCS







#### I(E)



Quantities varied

- wavelength  $\lambda$
- chopper slot widths W

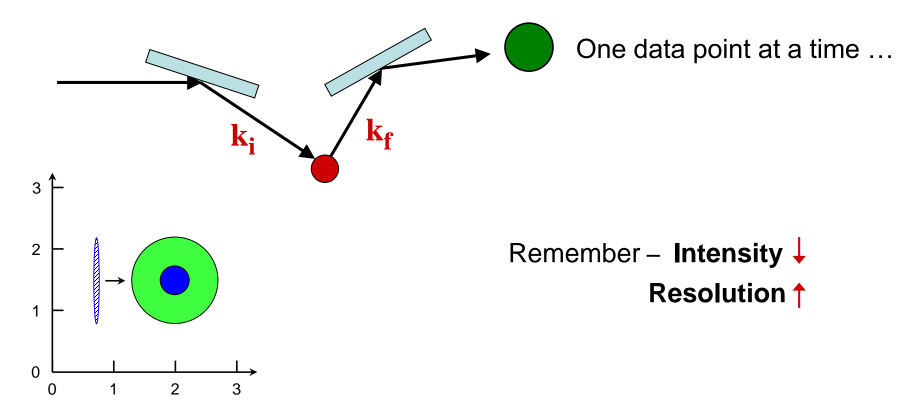
Remember – Intensity ↓
Resolution ↑

## Things to Consider When Choosing BT7



Triple axis spectrometers are typically used when either -

- (1) the *direction* of **Q** is important or
- (2) the interesting region of  $\mathbf{Q}$ - $\mathbf{\omega}$  space is of *limited extent*.



## Things to Consider When Choosing HFBS

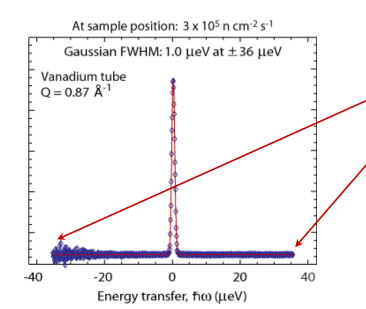


$$0.25 \, \text{Å}^{-1} < \mathbf{Q} < 1.75 \, \text{Å}^{-1}$$

Do the length scales of interest lie within this Q-range?

$$\delta \mathbf{Q} < 0.1 - 0.2 \, \text{Å}^{-1}$$

Can you live with such coarse Q-resolution?



Do the features of interest lie within this  $h\omega$ -range?

Do you really require such good energy resolution  $\delta E \sim 1 \mu eV$  (or perhaps even better resolution)?

## Things to Consider When Choosing NSE



If the  $h\omega$ -resolution of backscattering is "not good enough," or if you are only interested in a "limited" region of  $\mathbf{Q}$ -space (typically small  $\mathbf{Q}$ ) ...

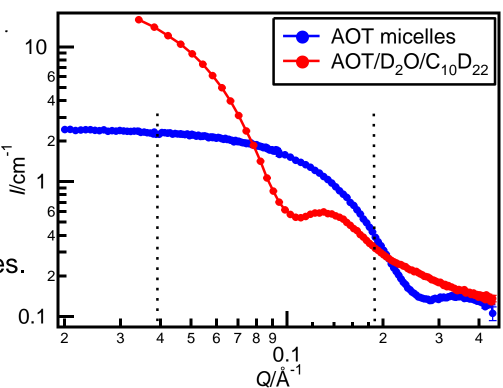
... then use NSE (low Q, long times)

These cases typically involve coherent scattering, which tends to peak near

Q ~  $\frac{2\pi}{\text{relevant length scale}}$ 

Remember – slower motions usually imply longer length scales.

Many atoms moving together→ Coherent scattering



## General Sample "Design"



## Know as much about your sample as possible!! (Beamtime costs ~ \$5000/day!!)

Other considerations:

What's the structure (in a general sense)?

Are there any phase transitions (or a glass transition)?

What isotopes are present?

Supplementary data from other measurements ...

Magnetization vs T

Muon spin relaxation

X-ray data

## General Sample "Design"



Try to avoid isotopes that are strongly absorbing.

<sup>6</sup>Li <sup>10</sup>B <sup>113</sup>Cd <sup>157</sup>Gd

For a complete listing go to

http://www.ncnr.nist.gov/resources/n-lengths

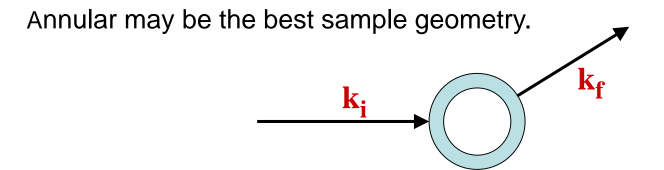
## Sample "Design" for Triple-Axis Spectrometers



Single crystals yield the most information.

Increase the intensity by increasing the amount of sample.

If you have a <u>powder</u>, use a cylindrical container (rather than flat plate).



## Sample "Design" for DCS and HFBS



Increase the intensity by increasing the amount of sample

→ Fill the beam with sample

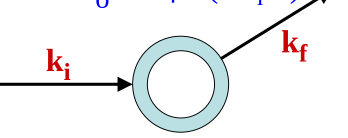
The maximum beam size is usually given in the instrument description:

DCS: 3 cm x 10 cm (or 1.5 cm x 10 cm)

Backscattering: 3 cm x 3 cm

If your sample is a powder, use cylindrical samples (rather than a flat plate). Remember - for incoherent, quasielastic scattering the transmission of the beam should be ~90%.  $I/I_0 = \exp{-(n\sigma_T D)}$ 

Often annular is the best sample geometry (if your sample is absorbing).



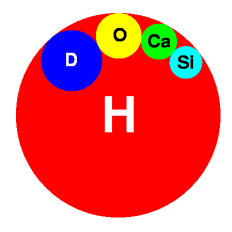
### Sample "Design" for DCS and HFBS



Does the sample contain H?

Remember: Neutrons LOVE H!!

Create a sample where the "interesting" portions are <u>hydrogenated</u> and the "uninteresting" portions are <u>deuterated</u>.



## Sample "Design" for NSE



Create a sample where the "interesting" portions of the sample have a <u>different SLD</u> than the "uninteresting" portions

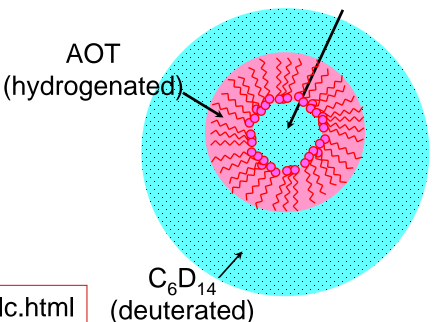
Typically this means <u>deuterating</u> the major phase in order to reduce the incoherent background

D<sub>2</sub>O (deuterated)

SLD core 6.4×10<sup>-6</sup> Å<sup>-2</sup>

SLD shell 10.0×10<sup>-6</sup> Å<sup>-2</sup>

SLD solvent 6.1×10<sup>-6</sup> Å<sup>-2</sup>



http://www.ncnr.nist.gov/resources/sldcalc.html

## Sample "Design" for NSE



Increase the intensity by increasing the amount of sample

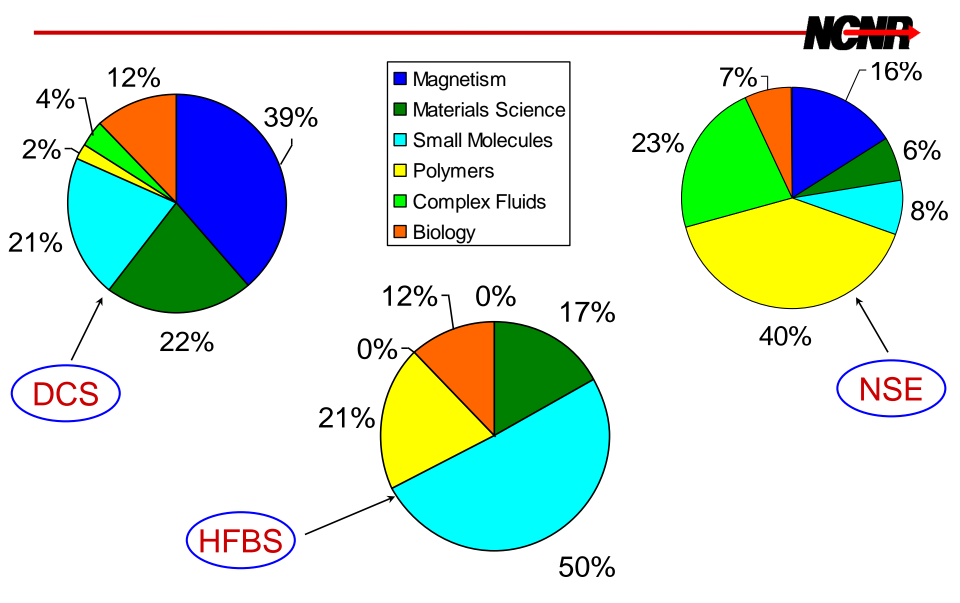
→ Fill the beam with sample

Typically use flat plate for powder samples (because most NSE experiments occur at small angles)



Rule of thumb - the transmission should be ~70%

## Typical Distributions of Science by Instrument



## Applying for Beam Time



Access to the neutron scattering instruments that you've used over the past week is merit-based. Open to all qualified users, but subject to an anonymous peer-review of proposals.

Calls for proposals are issued about twice/yr.



Next deadline for new proposals: June 16, 2015



Further information on submitting proposals:

http://www.ncnr.nist.gov/programs/CHRNS/CHRNS\_prop.html

#### Some Summer School Success Stories



2001 2003

Jae-Ho Chung University Prof.





Vicky Garcia-Sakai ISIS Staff Scientist

1999





1997

William Ratcliff NCNR Staff Physicist Rob Dimeo NCNR Director

## Acknowledgements



#### Organizers – Yun Liu and Yamali Hernandez

Administrative staff Experiment teams Invited speakers



**Scatter Well!**