

What Do We Mean By ABSOLUTE?

- There are 4 Absolute Methods of Measuring MW
- 1) Membrane **Osmometry** (Number Average MW)
- 2) Light Scattering (Weight Average MW)
- 3) Sedimentation Equilibrium (Ultracentrifugation) (z-average MW)
 - 4) Mass spectroscopy
- NO Reference to standards of mass
- NO assumptions of molecular model/conformation
- ALL parameters measured directly from 1st principles
- **Refractive indices**
- geometries of cell and detector
- wavelength
- concentrations
- detector response
- *temperature*
- dn/dc

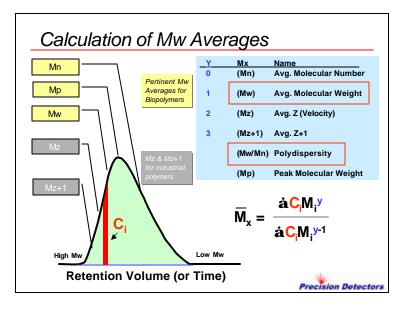
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Goals for this unit:

- Theory 101 just the basics Know why sky is blue, sunset burnt orange
- Rayleigh Scattering (Lord Rayleigh ~1871) / Rayleigh Ratio
- How does LS yield an "absolute" molecular weight? LS vs. RI
- What is polydispersity? How is it defined?
- Types of "Molecular Weight Averages"
- LS Instrument / Practical Considerations
- Static vs. Dynamic vs. X-ray Scattering (info from each)

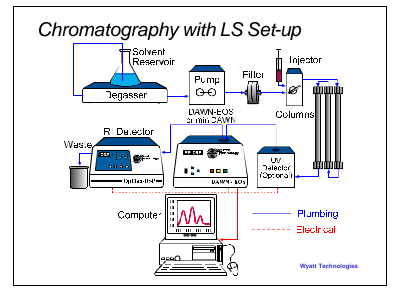


Abstract

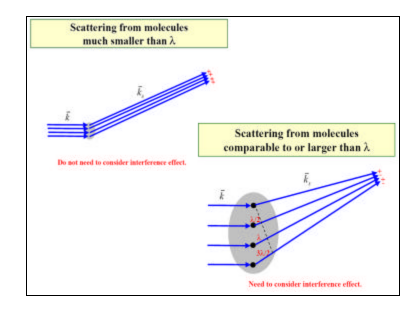
Recent advances in genomics and proteomics have produced a proliferation of new proteins requiring characterization. Mass spectrometry is ideally suited for identification and primary structural purposes but is not suited for determining well conformational structures in solution. As these molecules are expressed in cell culture, purified and then formulated, rigorous production processes must be carefully evaluated to minimize impact on the protein structure and its long-term shelf life. Obtaining a conformational stability profile of protein or antibodies can help weed out bad drug candidates from good ones as environmental factors can change their tertiary and quaternary structure. Environmental factors include pH, ionic strength, temperature, and excipient composition.

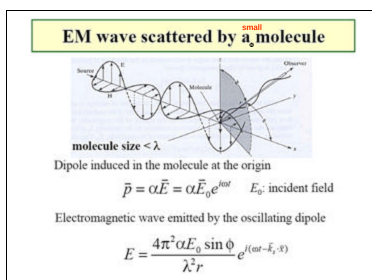
Precision Detectors

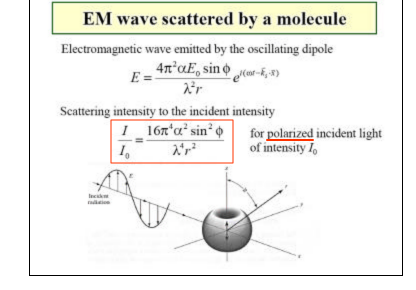
HPLC is used in flow injection mode with a detector array composed of laser light scattering (static and dynamic modes) and a concentration source detector (RI or UV). This configuration determines the average molecular weight and average hydrodynamic radius with run times as short as 1 minute. Alternatively, a SEC guard column can be used isolate analyte from excipients (eliminating blank runs) with run times under 3 minutes.

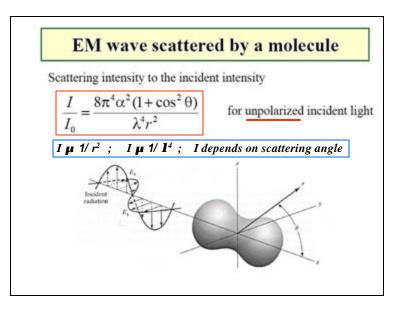


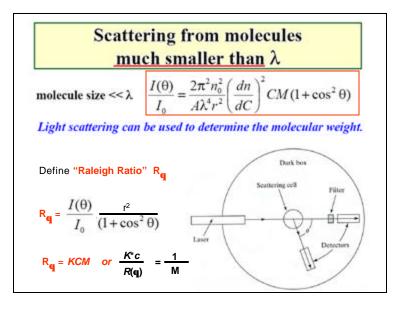












Basic Light Scattering Principles

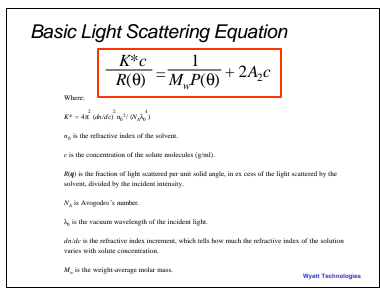
• The amount of **light scattered** is directly proportional to the product of the **molar mass** and the **molecular concentration**

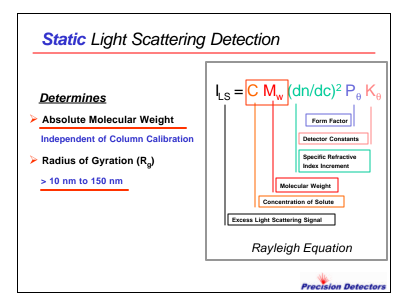
 $I_{LS} = C M_w (dn/dc)^2 P_{\theta} K_{\theta}$

• The variation of scattered light with scattering angle is proportional to the average size of the scattering molecules.

$$\frac{K^*c}{R(\mathbf{q})} = \frac{1}{M} \left[1 + \frac{16p^2}{3l^2} < r_g^2 > \sin^2(\mathbf{q}/2) + \cdots \right] + 2A_2c$$

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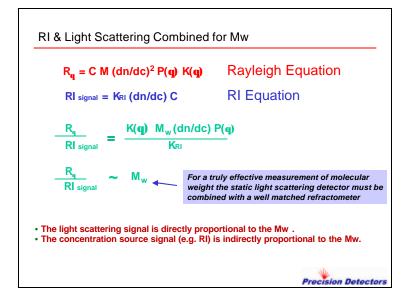


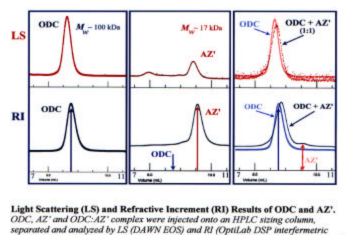
Accuracy of Molecular Masses of Test Proteins Determined by Light Scattering

Protein	Mass From Structure	Light Scattering*	Apparent Error	
	[Da]	[Da]	[%]	
Carbonic anhydrase	29,023	29,800	+2.7	
Alcohol dehydrogenase	145,980	149,000	+1.4	
β-Amylase	224,340	228,000	+1.6	
Apoferritin	476,316	484,400	+1.7	
Thyroglobulin	669,000	679,000	+1.5	
Ornithine decarboxylase	990,684	978,000	-1.3	
Octopus Hemocyanin	3,440,000	3,450,000	+0.3	

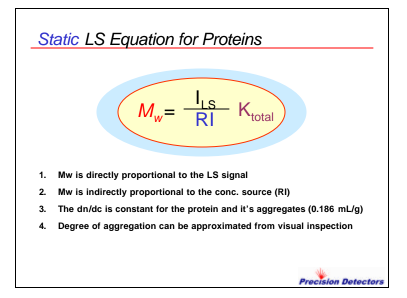
*DAWN detector model-F, 0.19 was used as dn/dc value for all the proteins Adapted from "Assembly of the Gigantic Hemoglobin of the Earthworm Lumbricus terrestris by A. Riggs et.al. In J. Bio. Chem., Vol. 271, No. 47, pp 30007-30021, 1996.

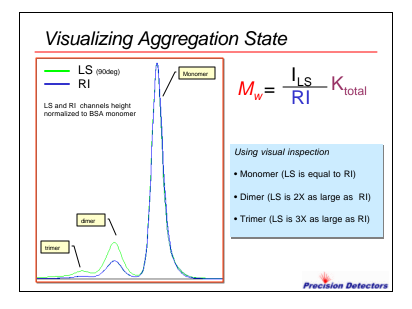


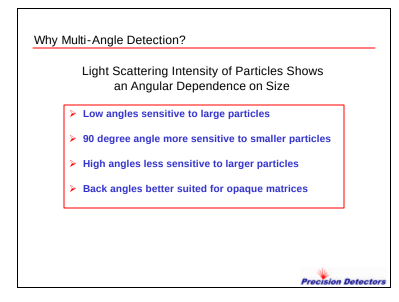


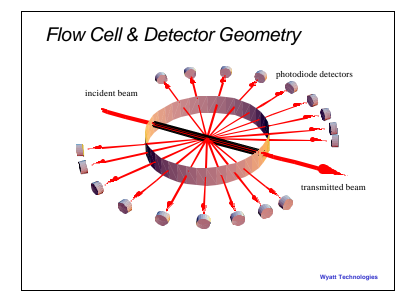


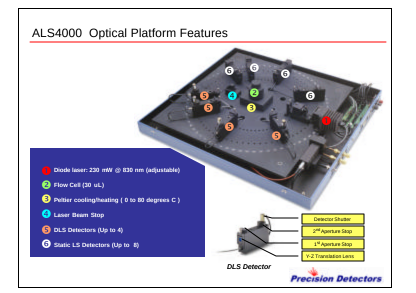
ODC, AZ and ODC:AZ complex were injected onto an HPLC sizing column, separated and analyzed by LS (DAWN EOS) and RI (OptiLab DSP interfermetric refractometer). The results shown are for 7 to 11 minutes of elution volume. Frame 1 is for ODC, frame 2 for AZ, and frame 3 for the ODC:AZ mixture at a 1:1 subunit ratio with the ODC trace from frame 1 superimposed for reference.

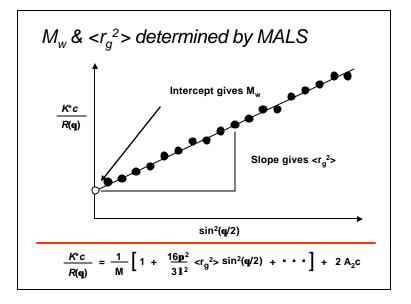


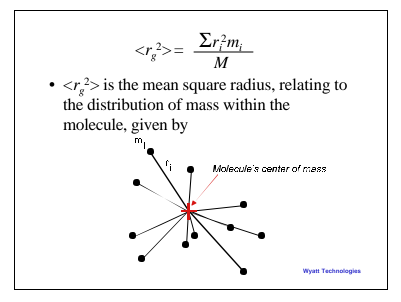


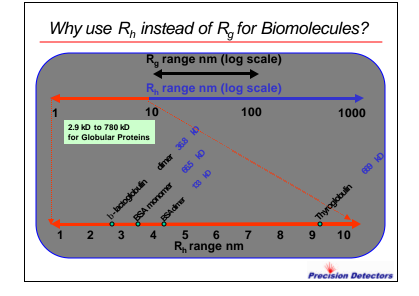


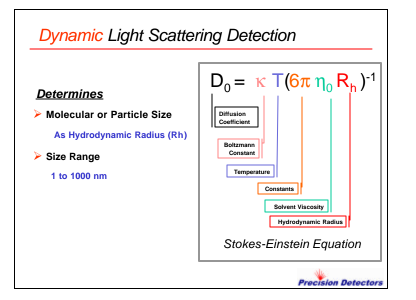


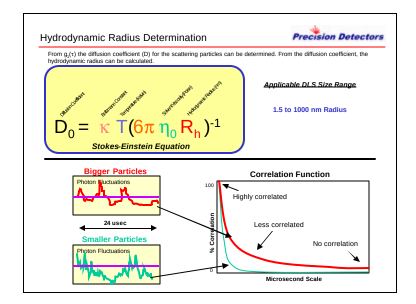


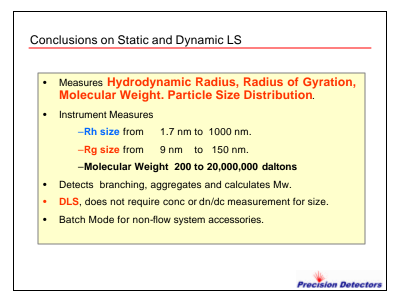


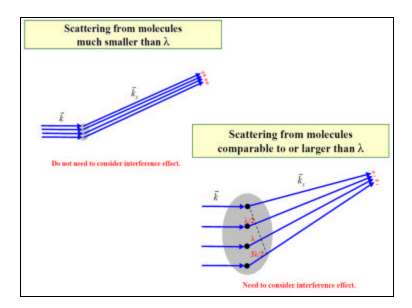


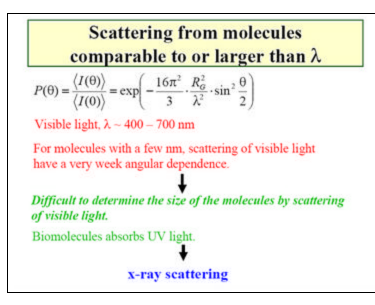


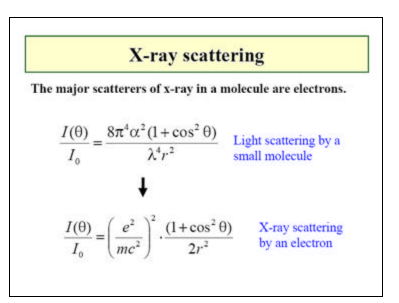


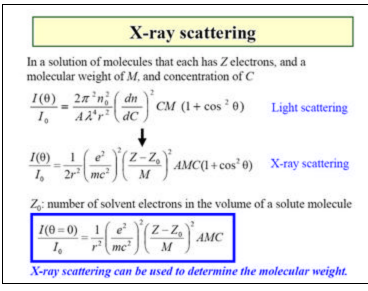


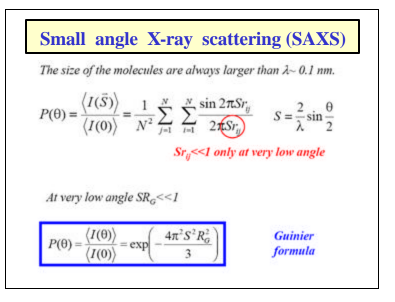








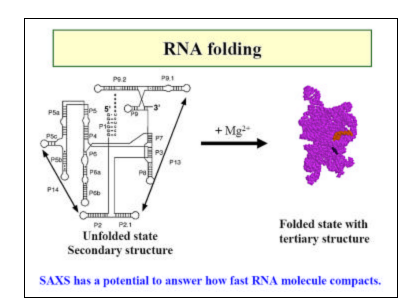


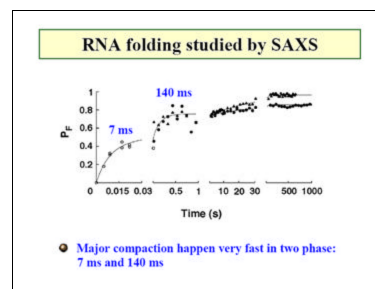


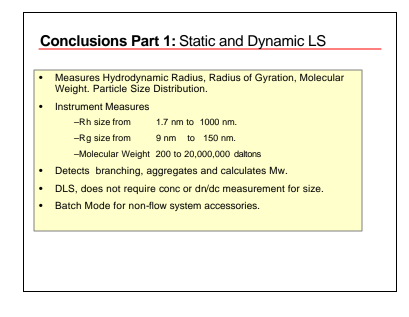
Molecular weight and size measured by Light scattering or SAXS

Material	M_w	$R_G (nm)$
Ribonuclease	12,700	1.48
a-Lactalbumin	13,500	1.45
Lysozyme	13,600	1.43
B-Lactoglobulin	36,000	
	36,700	2.17
Serum albumin	70,000	2.98
Myosin	493,000	46.8
Turnip yellow mosaic virus		10.4
Tobacco mosaic virus	39×10^{6}	92.4

Values in *italic* are from low angle X-ray scattering.







•Static Light Scattering •Provides absolute Mw in Solution.. •Sensitive Aggregation Detection •Oynamic Light Scattering •Provides Rh Sizing down to 1 nm for Biomolecules. •"On-the-fly Determinations" •Oives more insight on structure of material •Can be added to any existing static LS system from any manufacturer •Flow Injection Analysis is Ideal for Rapid Screening for Avg. Mw and Size

