

• Form Factors & Structure factors

This is list of available form factor and structure factors in Irena package. This is current as of release of 2.52.

Content:

[Form & Structure factors parameters](#)

[Form Factor description](#)

[Spheroid](#)

[Integrated_Spheroid](#)

[Cylinder](#)

[CylinderAR](#)

removed in v2.62: Algebraic_Globules

removed in v2.62: Algebraic_Rods

removed in v2.62: Algebraic_Disks

[Unified_Sphere](#)

[Unified_Rod](#)

[Unified_RodAR](#)

[Unified_Disk](#)

[CoreShell](#)

[CoreShellCylinder](#)

[CoreShellPrecipitate](#)

[Fractal_Aggregate](#)

[CoreShellShell](#)

[SphereWHSLocMonoSq](#)

[Janus_CoreShell_Micelle_1](#)

[Janus_CoreShell_Micelle_2](#)

[Janus_CoreShell_Micelle_3](#)

[User](#)

[Testing and using Form factors in users own code](#)

[Structure factors description](#)

[Interferences](#)

[HardSpheres](#)

[SquareWell](#)

[StickyHardSpheres](#)

[HayerPenfoldMSA](#)

[Interprecipitate](#)

[Calling the library and use](#)

Form & Structure factors parameters

<u>Form Factor</u>	<u>Parameter(s)</u>
spheroid	AspectRatio = ParticlePar1
Integrated_Spheroid	AspectRatio=ParticlePar1
Cylinders	Length=ParticlePar1
CylindersAR	AspectRatio=ParticlePar1
Unified_Disc	thickness = ParticlePar1

Unified_Rod length = ParticlePar1
Unified_RodAR AspectRatio = ParticlePar1
Unified_Sphere none needed
User uses user provided functions.
 There are two user provided functions necessary -
 F(q,R,par1,par2,par3,par4,par5)
 and V(R,par1,par2,par3,par4,par5)
 the names for these need to be provided in strings...
 the input is q and R in angstroms

Tube length=ParticlePar1 //length in A
 WallThickness=ParticlePar2 //in A
 CoreRho =ParticlePar3
 //rho [10^10 cm-2] (not delta rho squared!!!) of core material
 ShellRho =ParticlePar4
 //rho [10^10 cm-2] (not delta rho squared!!!) of shell material
 SolventRho =ParticlePar5
 //rho [10^10 cm-2] (not delta rho squared!!!) of surrounding
 medium (air=0)

CoreShell CoreShellThickness=ParticlePar1 //skin thickness in Angstroms
 CoreRho =ParticlePar2
 //rho [10^10 cm-2] (not delta rho squared!!!) of core material
 ShellRho =ParticlePar3
 //rho [10^10 cm-2] (not delta rho squared!!!) of shell material
 SolventRho =ParticlePar4
 //rho [10^10 cm-2] (not delta rho squared!!!) of surrounding
 medium (air=0)

CoreShellPrecipitate CoreShellThickness is calculated for each size, so the average contrast of
 the core+shell is same as contrast of the solvent.
 CoreRho =ParticlePar2
 //rho [10^10 cm-2] (not delta rho squared!!!) of core material
 ShellRho =ParticlePar3
 //rho [10^10 cm-2] (not delta rho squared!!!) of shell material
 SolventRho =ParticlePar4
 //rho [10^10 cm-2] (not delta rho squared!!!) of surrounding
 medium (air=0)

Fractal aggregate FractalRadiusOfPriPart=ParticlePar1 //radius of primary particle
 FractalDimension=ParticlePar2 //Fractal dimension

SphereWHSLocMonoSq Distance for HS model as ratio to part radius
 Hard Spheres model fraction

Janus CoreShell Micelle 1 //particle size is total size of the particle
 (R0 in the figure in description)
 Shell_Thickness=ParticlePar1//shell thickness
 SolventRho=ParticlePar2 // rho for solvent
 CoreRho=ParticlePar3 // rho for core material
 Shell1Rho=ParticlePar4 // rho for shell 1
 Shell2Rho=particlePar5 // rho for shell 2

Janus CoreShell Micelle 2 //particle size here is shell thickness!!!
 Core_Size=ParticlePar1 // Core radius A
 SolventRho=ParticlePar2 // rho for solvent
 CoreRho=ParticlePar3 // rho for core material
 Shell1Rho=ParticlePar4 // rho for shell 1
 Shell2Rho=particlePar5 // rho for shell 2

Janus CoreShell Micelle 3 //particle size here is core radius
 Shell_Thickness=ParticlePar1 // Shell Thickness A

```

SolventRho=ParticlePar2      // rho for solvent
CoreRho=ParticlePar3    // rho for core material
Shell1Rho=ParticlePar4 // rho for shell 1
Shell2Rho=particlePar5 // rho for shell 2

```

Structure factors included

Interferences reference: Beaucage, G. (1995). J Appl Crystallogr **28**, 717-728.

Par1: ETA (center-to-center distance)

Par2: Pack (number of particles In nearest neighbor sphere)

Hard Spheres reference: Percus-Yevick model, PERCUS,YEVICK PHYS. REV. **110** 1 (1958),
THIELE J. CHEM PHYS. **39** 474 (1968), WERTHEIM PHYS. REV. LETT. **47** 1462 (1981)

Par1: Radius [A]

Par2: Volume fraction (fraction)

Square Well reference: SHARMA,SHARMA, PHYSICA **89A**,(1977),212, NOTE - depths >1.5kT
and volume fractions > 0.08 give UNPHYSICAL RESULTS when compared to Monte Carlo
simulations

Par1: Radius [A]

Par2: Volume fraction [fraction]

Par3: Well depth e/kT, dimensionless, positive values are attractive

Par4: Well width, multiples of diameters

Sticky hard spheres no reference given in NIST macros \AA

Par1: Radius [A]

Par2: Vol. fraction

Par3: Perturbation parameter (0.1)

Par4: Stickiness, tau

Hayer Penfold MSA no reference given in NIST macros \AA

Par1: Radius [A]

Par2: Charges

Par3: Volume fraction

Par4: Temperature in Kelvin

Par5: Monovalent salt concentration (M)

Par6: dielectric constant of solvent

Interprecipitate SF Formula 6 in APPLIED PHYSICS LETTERS **93**, 161904 (2008) \AA

Par1: Distance L [A]

Par2: Sigma (root-mean-square deviation (ordering factor))

Important comment for Core-shell and Core shell cylinder (and Unified tube)

The volume definition for Core-shell objects is matter of discussion. Heated at times and I suspect that the appropriate answer depends on the case when and how the FF is used. Therefore from version 2.26 Irena macros include option which needs to be set - both Core shell and Core shell cylinder will share common parameter (this parameter is global for all cases of calls to core shell form factors or their volumes) of volume definition.

The options are: whole particle, core, and shell.

Note: Unified tube is using as volume the volume of shell. It is how it is defined at this time and it is meant for cases like Carbon nanotubes, when this is appropriate. To match with core shell cylinder us "shell" as volume

Form Factor description

```
*****
*****
```

Spheroid

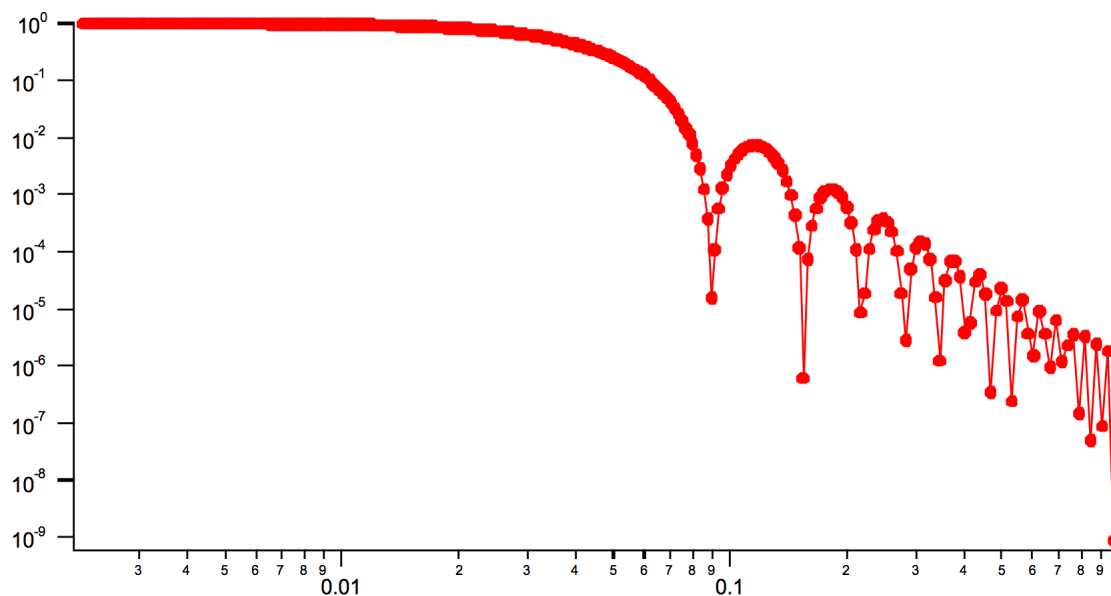
uses sphere form factor for aspect ratio between 0.99 and 1.01:

$$F^2 = 3/(QR^3) * (\sin(QR) - (QR * \cos(QR)))$$

$$\text{volume : } V = ((4/3) * \pi * \text{radius}^3)$$

This calculation approximates integral over R as rectangle (compare with Integrated spheroid).

graph for R = 50A



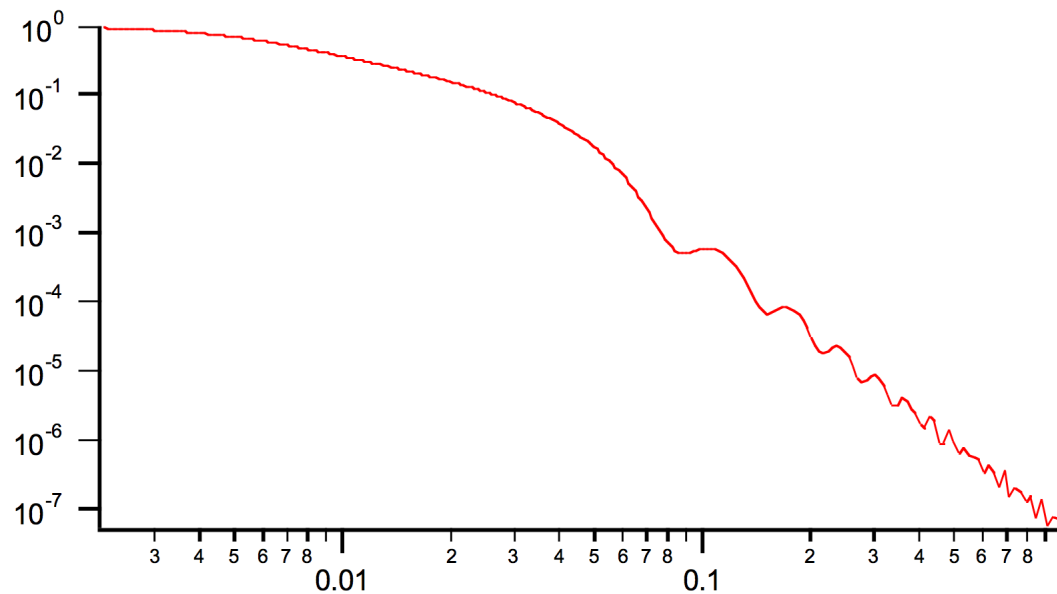
For aspect ratios smaller than 0.99 and larger than 1.01 uses standard form factor for spheroid:

$$F = \text{Integral of } (3/(QR^3)) * (\sin(QR) - (QR * \cos(QR)))$$

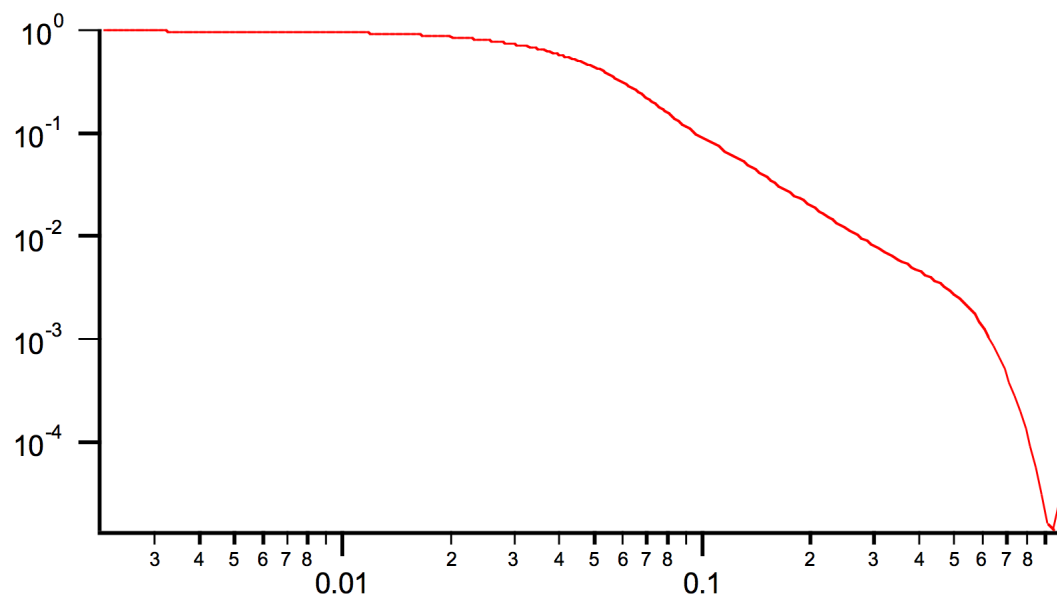
$$\text{where } QR = Q\text{value} * \text{radius} * \sqrt{1 + ((AR^2) - 1) * \cos^2(\theta)}$$

over of $\cos(\theta) = 0$ to 1. This is numerically calculated using 50 points (step in $\cos(\theta) = 0.02$). Following graphs are examples:

AR = 10



AR=0.1



Integrated_Spheroid

same code as in the spheroid, but in this case the code integrates over the width of the R bin.

Note, the bin star and end points are calcualted linearly (even for log-binned data) as half way distance:

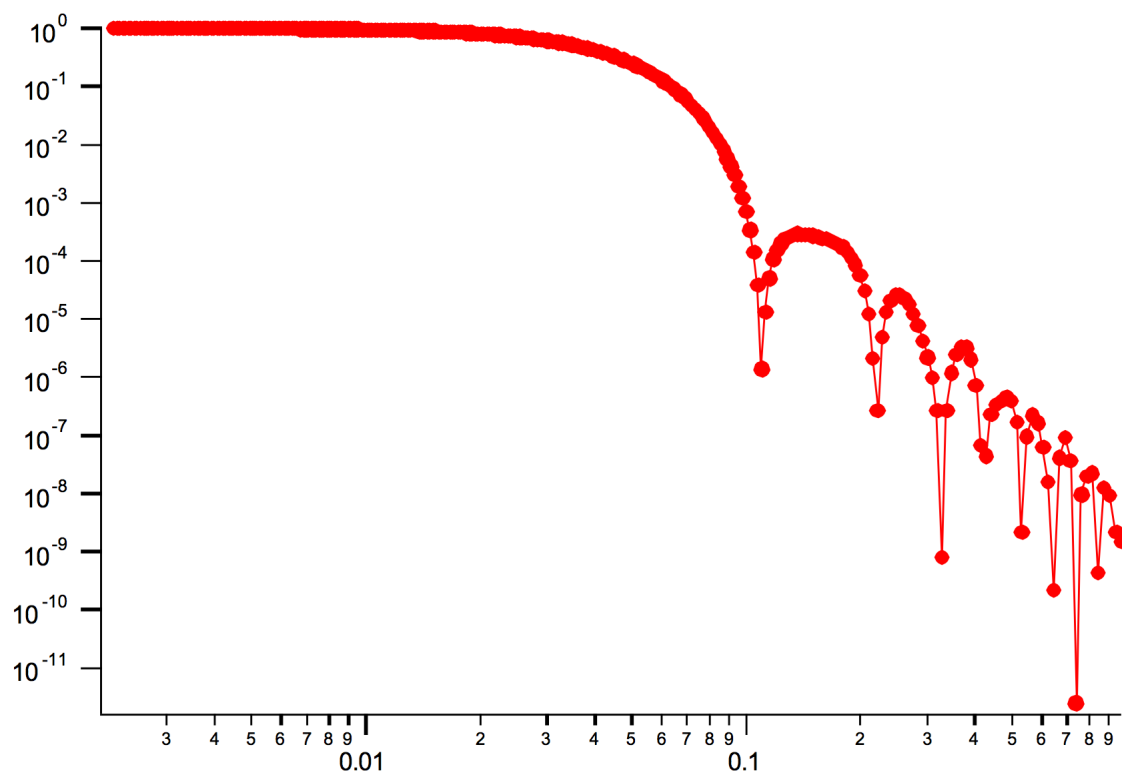
$$R_{\text{start}} = (R_n + R_{n-1})/2$$

$$R_{\text{end}} = (R_n + R_{n+1})/2$$

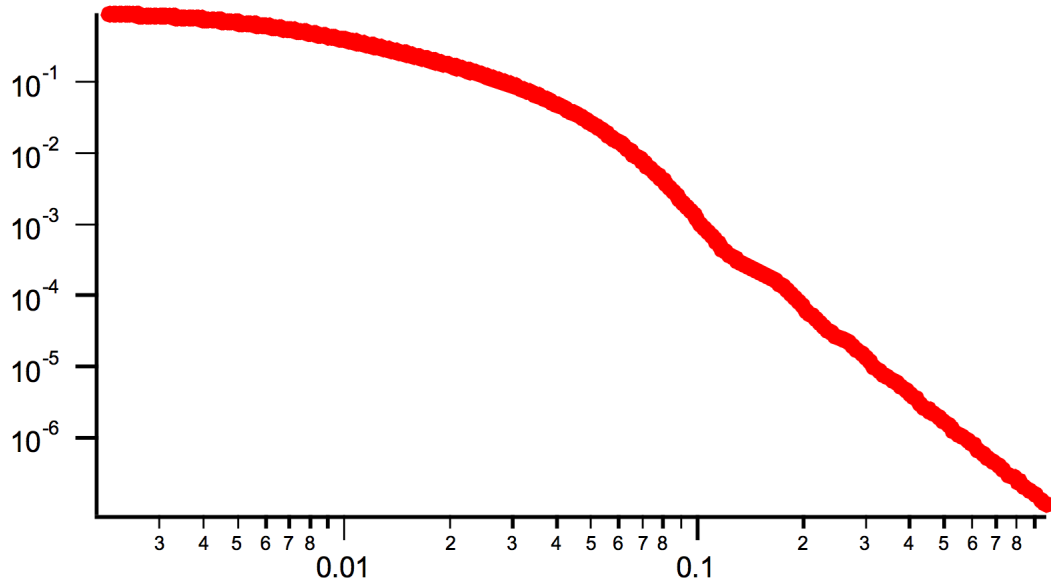
Uses adaptive steps to integrate bessel function oscillations of the form factor over the width of the bin in R - note, the averaging is done including the volume of particles involved. This code is quite convoluted and time consuming. Its only reasonable use is for cases with wide bins in radius (R), when this removes some of the bessel function oscillations.

Examples with R width 40A, average size 50A (that means R varies from 30 to 70A). Note that the bessel function oscillations are somewhat smooth out. With wider bins in R these oscillations may disappear all together.

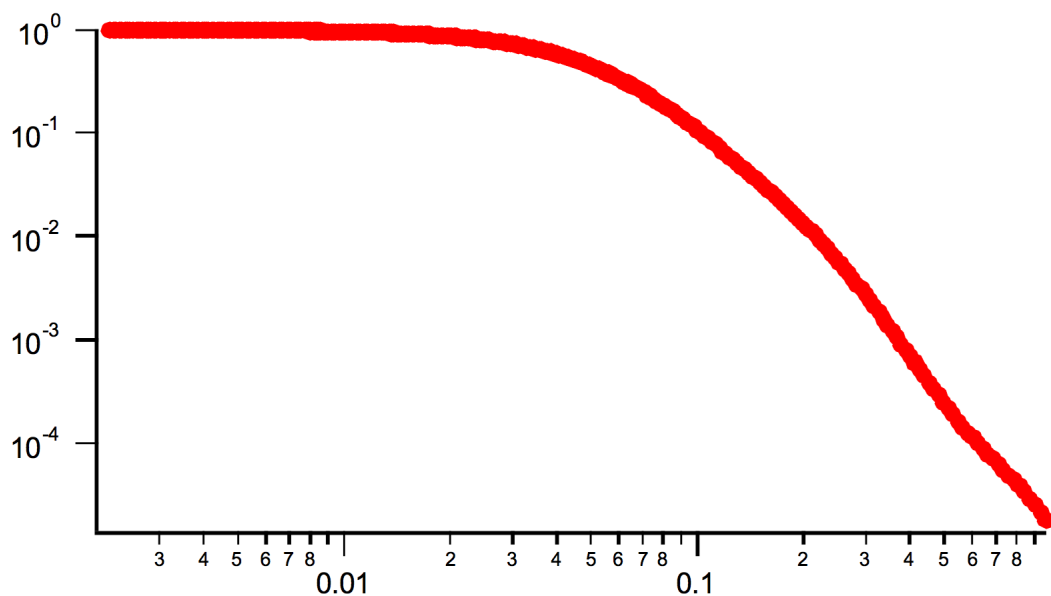
AR = 1 (sphere)



AR=10 (Spheroid)



AR=0.1 (spheroid)



```
*****
*****
```

Cylinder**CylinderAR**

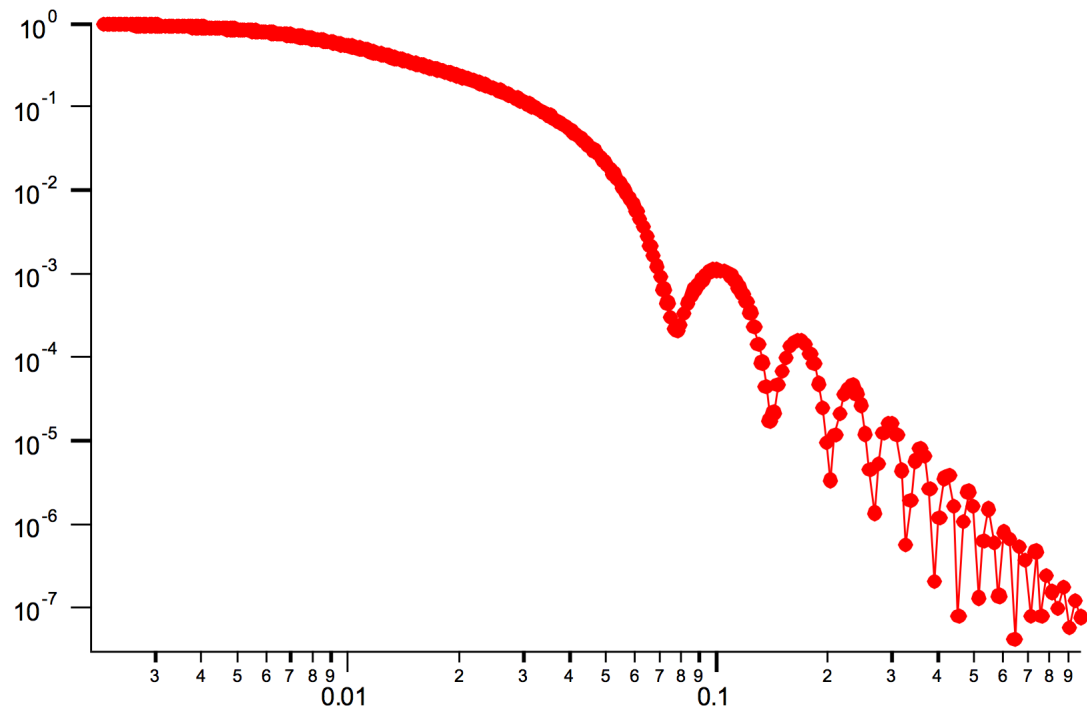
The code uses the following code to calculate form factor for cylinder. Note, that also this code is doing the same integration as integrated spheroid above (see 2).

Form factor = integral over (F_t) for $\Alpha = 0$ to $\pi/2$, F_t is below:
 $\text{LargeBes} = \sin(0.5 * \text{Qvalue} * \text{length} * \cos(\Alpha)) / \text{LargeBesArg}$

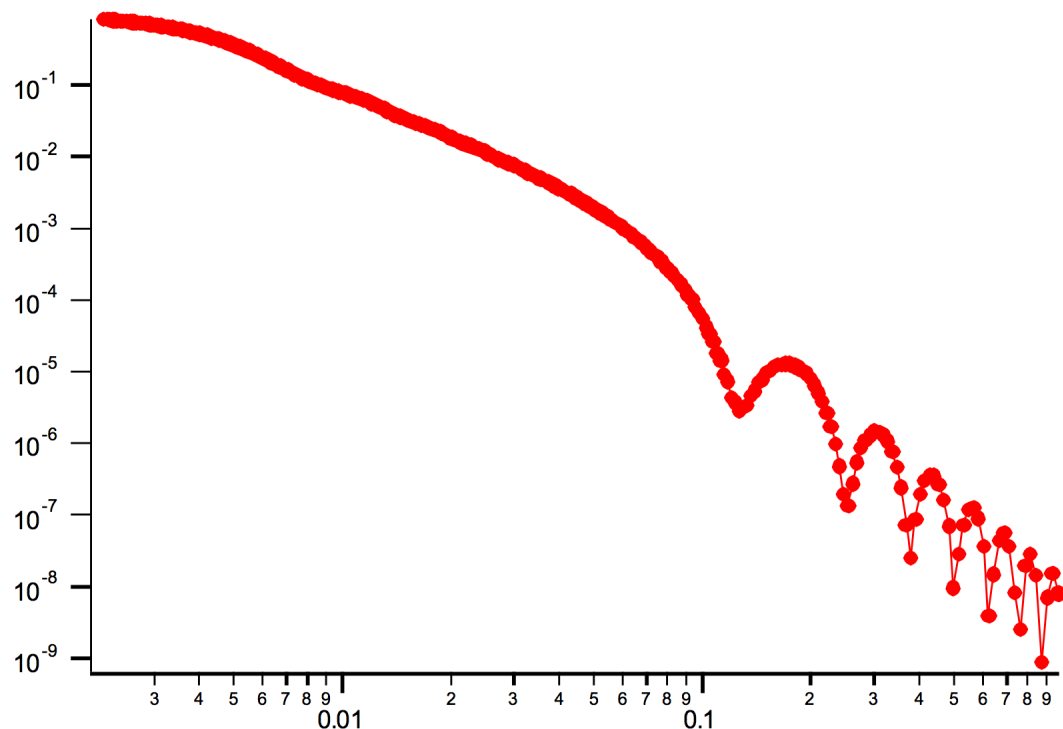
```
SmallBessDivided=BessJ(1, Qvalue*radius*Sin(Alpha))/Qvalue*radius*Sin(Alpha)
Ft = LargeBes*SmallBessDivided
```

Examples

Cylinder with length 500A and radius 50A.



Disk (cylinder) with radius 500A and length 50A.



```
*****
*****
```

removed in v2.52: **Algebraic_Globules**

removed in v2.52 - use Unified FF approximation (it is much better)

```
*****
*****
```

removed in v2.52: **Algebraic_Rods**

removed in v2.52 - use Unified FF approximation (it is much better)

```
*****
*****
```

removed in v2.52: **Algebraic_Disks**

removed in v2.52 - use Unified FF approximation (it is much better)

```
*****
*****
```

Unified_Sphere

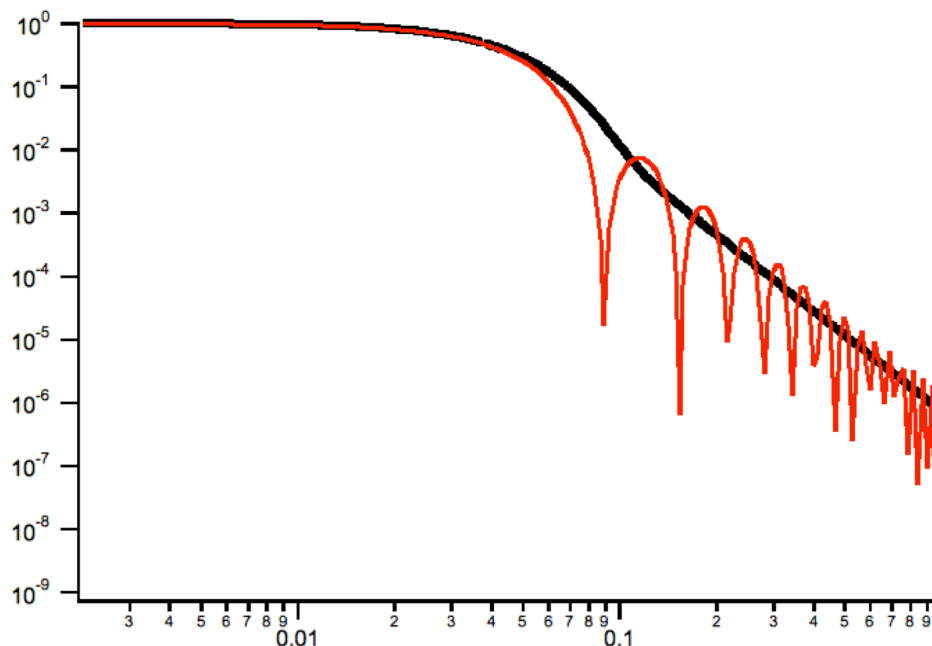
This is formula from Unified fit model by Greg Beaucage (see Unified tool and

documentation for it). The parameters are calculated from the code in the manual for each different shape. Specific formulas for these shapes were provided by Dale Schaefer...

This is the code:

```
G1=1
P1=4
Rg1=sqrt(3/5)*radius
B1=1.62*G1/Rg1^4
QstarVector=qvalue/(erf(qvalue*Rg1/sqrt(6)))^3
F^2 = G1*exp(-qvalue^2*Rg1^2/3)+(B1/QstarVector^P1)
```

Example for R=50A compared with the spheroid with aspect ratio =1



Unified_Rod

Unified_RodAR

This is formula from Unified fit model by Greg Beaucage (see Unified tool and documentation for it). The parameters are calculated from the code in the manual for each different shape. Specific formulas for these shapes were provided by Dale Schaefer...

This is the code:

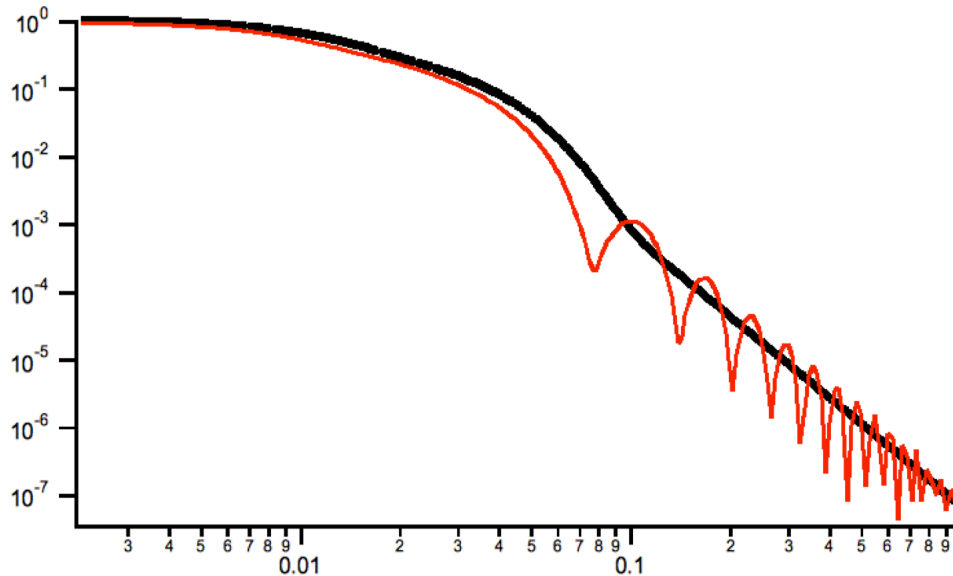
```
G2 =1
Rg2=sqrt(Radius^2/2+Length^2/12)
B2=G2*pi/length
P2=1
Rg1=sqrt(3)*Radius/2
```

```

RgC02=Rg1
G1=2*G2*Radius/(3*Length)
B1=4*G2*(Length+Radius)/(Radius^3*Length^2)
P1=4
QstarVector=qvalue/(erf(qvalue*Rg2/sqrt(6)))^3
A=G2*exp(-qvalue^2*Rg2^2/3)+(B2/QstarVector^P2) * exp(-RGC02^2 * qvalue^2)
QstarVector=qvalue/(erf(qvalue*Rg1/sqrt(6)))^3
F^2 = A + G1*exp(-qvalue^2*Rg1^2/3)+(B1/QstarVector^P1)

```

Example for R=50A and length 500A compared with the cylinder



Unified_Disk

This is formula from Unified fit model by Greg Beauchage (see Unified tool and documentation for it). The parameters are calculated from the code in the manual for each different shape. Specific formulas for these shapes were provided by Dale Schaefer...

This is the code:

```

G2=1
Rg2=sqrt(Radius^2/2+thickness^2/12)
B2=G2*2/(radius^2)//dws guess
P2=2
Rg1=sqrt(3)*thickness/2// Kratky and glatter = Thickness/2
RgC02=1.1*Rg1
G1=2*G2*thickness^2/(3*radius^2)
B1=4*G2*(thickness+Radius)/(Radius^3*thickness^2)//same as rod
P1=4
QstarVector=Q/(erf(Q*Rg2/sqrt(6)))^3

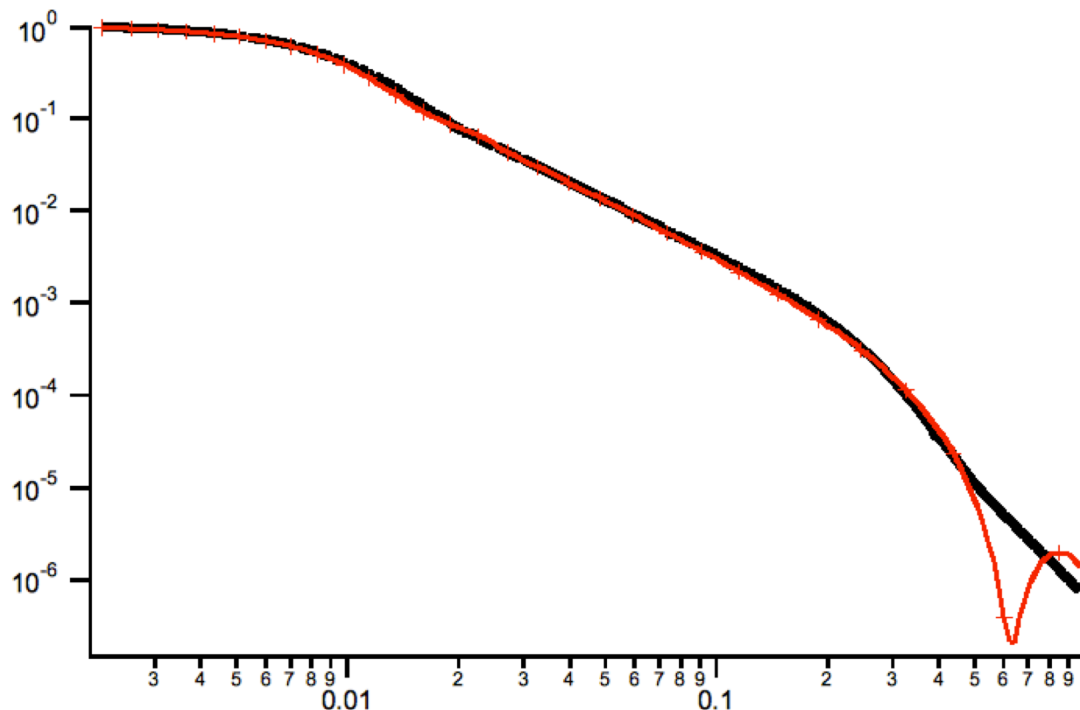
```

```

A=G2*exp(-Q^2*Rg2^2/3)+(B2/QstarVector^P2) * exp(-RGC02^2 * Q^2/3)
QstarVector=Q/(erf(Q*Rg1/sqrt(6)))^3
F^2 = A + G1*exp(-Q^2*Rg1^2/3)+(B1/QstarVector^P1)

```

Example for R=250Å and thickness 10Å compared with the cylinder



```

*****
*****
*****
CoreShell
```

One thing to remember: the total radius of this particle is core radius + shell thickness... Note, this form factor calculation also includes integration over the width of bin in radii (same as integrated spheroid and cylinder).

Note: Input contrast for core/shell/solvant is rho in [10^{10} cm^{-2}] - this is very important to keep in mind.

Note, that there is volume definition choice you need to do: Whole particle, core, or shell, as appropriate for given problem. This volume definition is used for all volume calculations for this particle. It is global parameter for all core shell cylinder or core shell calls in the WHOLE EXPERIMENT.

Code (heavily simplified!):

```
Contrast = CoreRho - ShellRho
```

```

//core
Result1=(3/(Q*R)^3)*(sin(Q*R)-(Q*R*cos(Q*R))) * Contrast * (4/3 * pi * R^3)

//Now add the shell (skin) , thickness Rshell
r = R+Rshell
Contrast = ShellRho - SolventRho
Result2 = (3/(Q*r)^3)*(sin(Q*r)-(Q*r*cos(Q*r))) * Contrast * (4/3 * pi * r^3)

//summ them together and normalize by the total particle volume
F^2 =( result1 + result2 )^2 / Volume

```

Volume definition depends on the setting of above discussed global parameter and is either:

Whole particle volume = $4/3 * \pi * (R+r)^3$

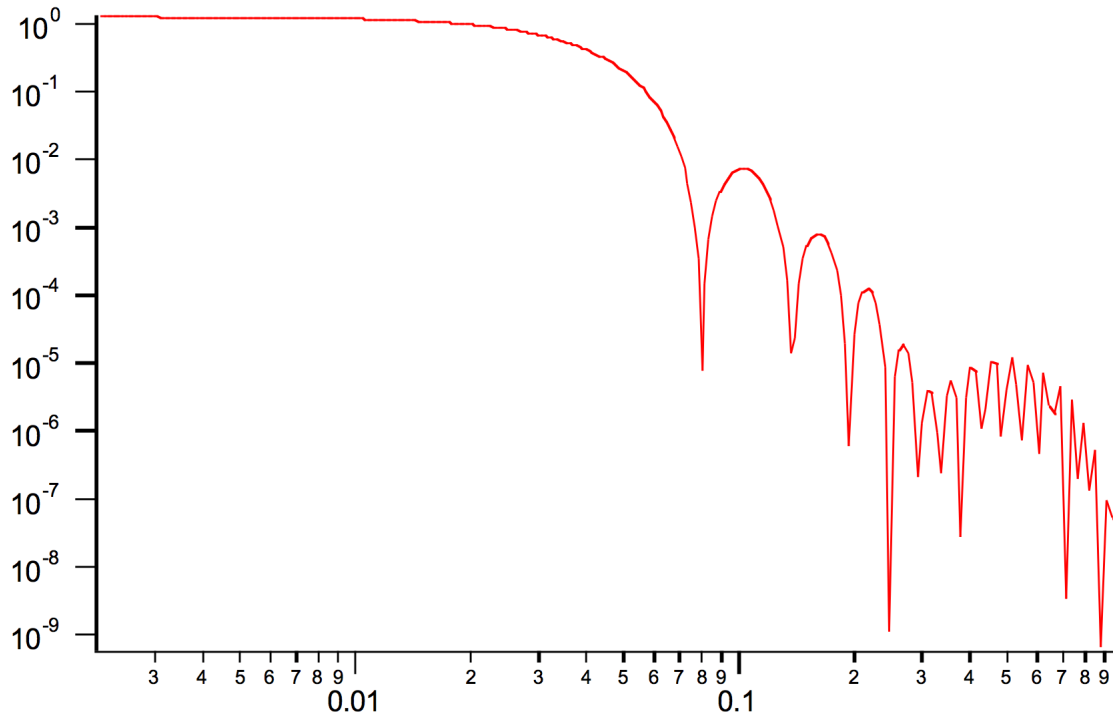
Core volume = $4/3 * \pi * R^3$

Shell volume = $4/3 * \pi * r^3$

Make sure your choice is appropriate

Note, that to my surprise these calculations (copied from NIST Form factors) do not normalize correctly to 1 at low q. The reason is that the weighting is done by volume and contrast. I'll need to look into this again and in detail...

Example, Radius 50A, skin thickness 10A, contrast ratio 0.6



```
*****
*****
```

CoreShellPrecipitate

This is unique form factor, which - even for dilute system - results in "diffraction peak" type scattering. It is a very special case, when coreshell particle is formed from matrix and as it is formed, the core chemistry/contrast deviates from matrix chemistry/contrast. If the diffusion in the matrix is not fast enough, the chemistry around the particle changes, which results in contrast changing in the other direction. Therefore one can end with coreshell particle which has higher-than-solvent contrast core surrounded by lower-than-solvent contrast shell (or the other way). With average contrast same as matrix. In such case at low-qs the particle "disappears" since we are probing material on larger length scales, and on average at those length scales the contrast is the same. For example of this type of precipitation see:

Imhoff, S.D., et al., Kinetic transition in the growth of Al nanocrystals in Al-Sm alloys. *Journal of Applied Physics*, 2012. 111(6): p. 063525-9.

Remember, that by basic nature of this logic, the contrast of the core/shell needs to be one larger and the other smaller than solvent contrast. Also, they probably need to be pretty close together. If this is not correct, the code would create negative shell thicknesses and abort. Do not do it, it is not very physical...

The particle volume is always volume of the core. I think no other logic makes too much sense.

Code uses regular coreshell form factor (see above). For each size the shell thickness is calculated so the average contrast of the particle matches the contrast of the solvent.

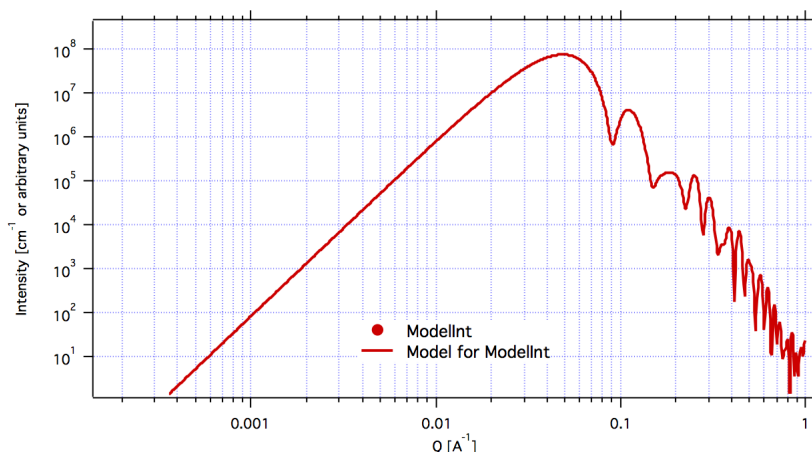
First we calculate:

`ShellVolume = CoreVolume*(SolventContrast - CoreContrast) / (ShellContrast - SolventContrast)`

Then we calculate the shell thickness for known ShellVolume and known core radius.

`Core volume = 4/3 * pi * R^3`

Example, Radius 50A, Core contrast 110, Shell Contrast 85, Solvent Contrast 90; note, this internally resolves to shell thickness of 35.5A.



```
*****
*****
```

CoreShellCylinder

Note, this form factor calculation also includes integration over the width of bin in radii (same as integrated spheroid and cylinder).

This code has been developed some time ago and I am not sure about it's function...

Code which is being used is direct copy of NIST Core shell cylinder.

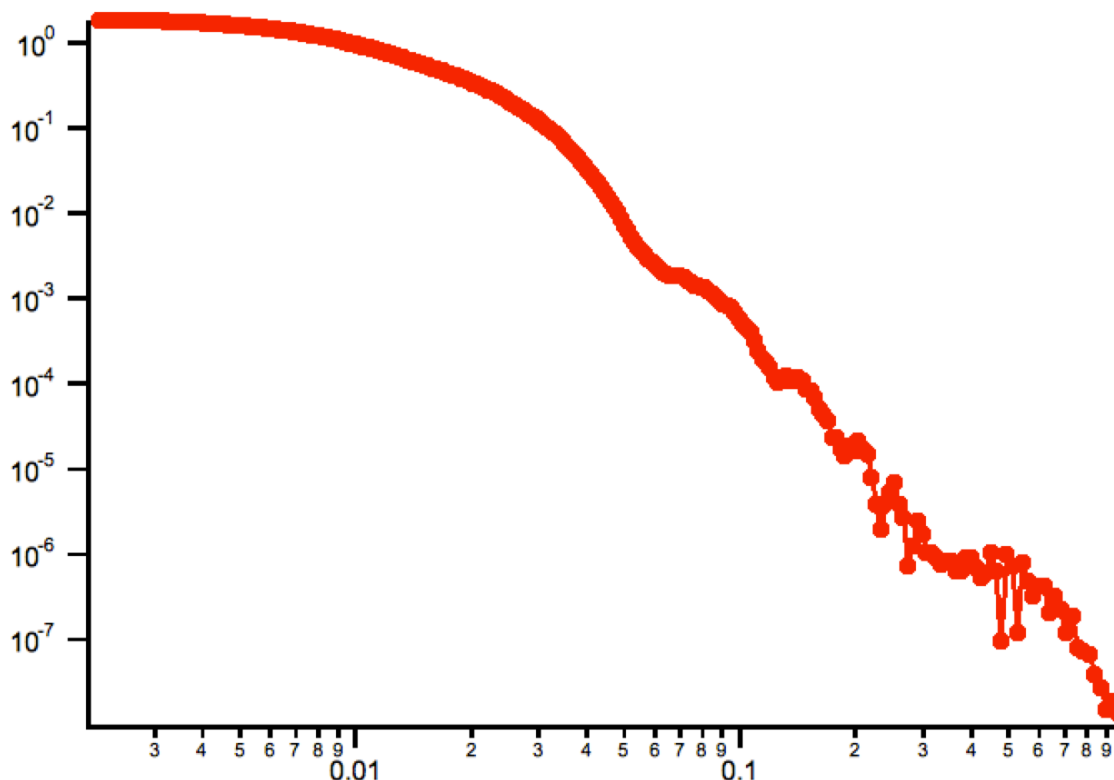
Note, that there is volume definition choice you need to do: Whole particle, core, or shell, as appropriate for given problem. This volume definition is used for all volume calculations for this particle. It is global parameter for all core shell cylinder or core shell calls in the WHOLE EXPERIMENT \AA .

Volume definition depends on the setting of above discussed global parameter and is either:

Whole particle volume = $\pi * (R+r)^2 * (L+2*r)$

Core volume = $\pi * R^2 * L$

Shell volume = $\pi * r^2 * L$



```
*****
*****
```

Fractal Aggregate

This form factor was requested by Dale Schaefer and I cannot very well guarantee its functionality....

code:

```
f = IR1T_CalcSphereFormFactor(Qw[p],(2*Param1))
//calculates the F(Q,r) part fo formula
//this is same as for sphere of diameter = 2*Param1
// (= radius of primary particle, which is hard sphere)
//fractal part is next
F^2 =f^2 * IR1T_CalculateFractAggSQPoints(Qw[p],currentR,Param1, Param2)
```

where

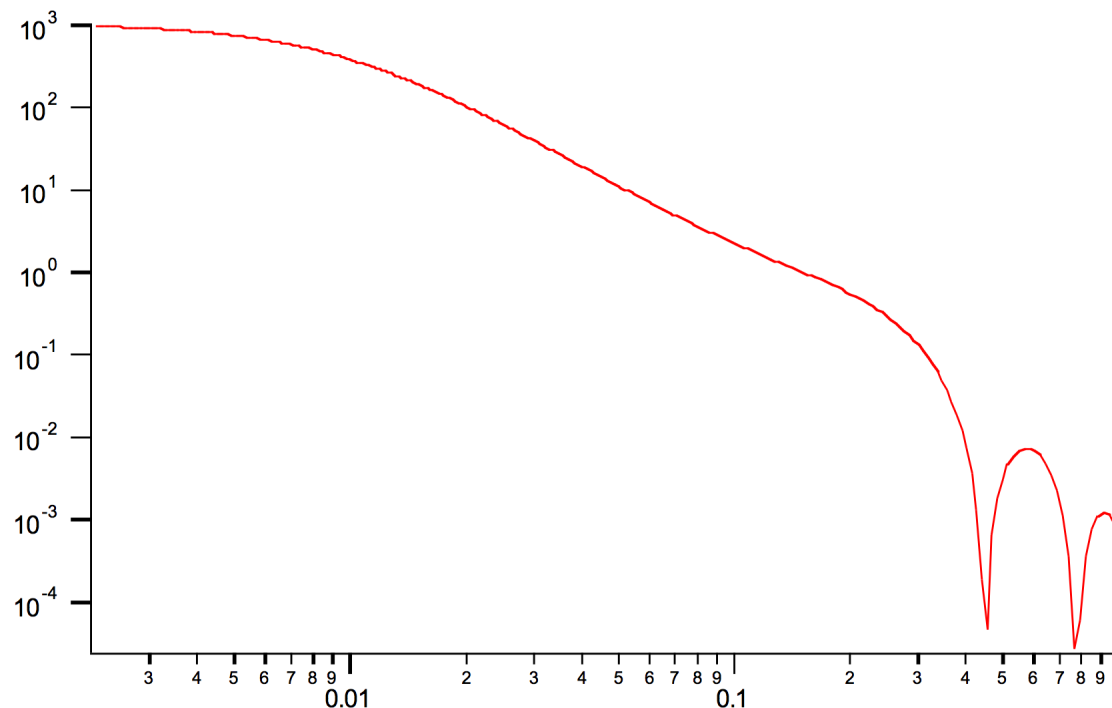
```
IR1T_CalculateFractAggSQPoints(Qvalue,R,r0, D) is
QR=Qvalue*R
part1=1
part2=(qR*r0/R)^-D
part3=D*(exp(gammln(D-1)))
part5= (1+(qR)^-2)^((D-1)/2)
part4=abs(sin((D-1)*atan(qR)))
return (part1+part2*part3*part4/part5)
```

Note, that parameters are :

Param1 - radius of primary particle

param2 - fractal dimension of the fractal particles

Example for R=100A, radius of primary particle 10 A and fractal dimension 2.5.



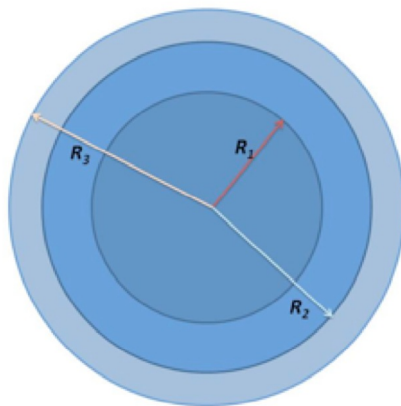
Comment: Note, that this is not scaled correctly at all... I have no idea why - apparently this formula is either wrongly coded or plainly does not behave right.

```
*****  
*****  
*****  
*****  
*****  
*****
```

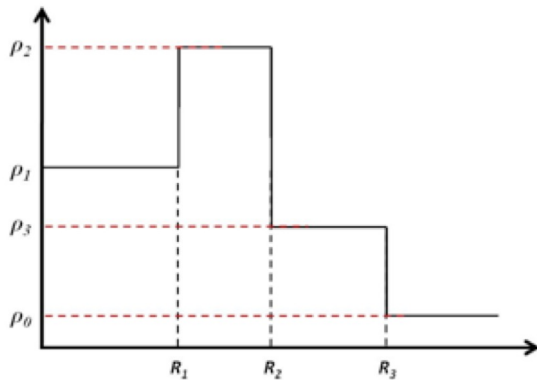
CoreShellShell

This form factor has been provided by Fan Zhang, many thanks to him.

Description of the model:



Scattering Length Density:



List of Model Parameters:

R_1 : core radius

R_2 : outer radius of the first shell

R_3 : outer radius of the second shell

ρ_0 : scattering length density of the matrix

ρ_1 : scattering length density of the core

ρ_2 : scattering length density of the first shell

ρ_3 : scattering length density of the second shell

First-order Bessel function of the first kind is defined as

$$J_1(x) = \frac{\sin(x) - x \cos(x)}{x^2}$$

Volume is defined as

$$V_i = \frac{4\pi}{3} R_i^3$$

Form factor of the core-shell-shell structure is:

$$F(q) = \frac{3V_1}{qR_1} (\rho_1 - \rho_2) J_1(qR_1) + \frac{3V_2}{qR_2} (\rho_2 - \rho_3) J_1(qR_2) + \frac{3V_3}{qR_3} (\rho_3 - \rho_0) J_1(qR_3)$$

Volume definition depends on the setting of global parameter described in Core-shell form factor

and is either:

$$\text{Whole particle volume} = 4/3 * \pi * (R+r)^3$$

$$\text{Core volume} = 4/3 * \pi * R^3$$

$$\text{Shell volume} = 4/3 * \pi * r^3$$

Where shell thickness "r" is sum of the two shell thicknesses (R_3-R_1).

Make sure your choice is appropriate

```
*****
*****
*****
*****
```

SphereWHSLocMonoSq

This is form factor combined with structure factor – Based on Jan Skov Pedersen J. Appl. Cryst paper : J. Appl. Cryst. (1994) 27, 595-608. The model is locally mono dispersed system, therefore locally one can use spheres Form factor combined with structure factor.

For each bin here the code calculates $F(Q,R)^2 * S(Q,D,\phi)$, where $D \sim R$ via input parameter. Phi is simply fraction of Percus Yevic structure factor.

The result is different than multiplying dilute system by Structure factor – that assumes that the distance for Structure factor is the same for all sizes. In this case the ratio of distance to size of particle is the same. We assume here that the phi is the same for all sizes.

Suffise to say, that using this form factor with another structure factor is meaningless and garbage will be produced.

```
*****
*****
*****
```

Janus CoreShell Micelle 1

Janus CoreShell Micelle 2

Janus CoreShell Micelle 3

This is form factor based on manuscript:

T. Futterer, G. A. Vliegenthart, and P. R. Lang, "Particle Scattering Factor of janus Micelles", Macromolecules 2004, 37, 8407-8413.

The Form factor follows formula 3 of this manuscript

$$\begin{aligned}
 b_J^2(\mathbf{Q}, \theta) = & \frac{4\pi^2}{Q^2(1-\mu^2)V_n} \left[((\rho_B + \rho_C) \int_0^{R_o} \cos(kz) F(R_o, z, \mu) dz \right. \\
 & + (\Delta_{BA} + \Delta_{CA}) \int_0^{R_i} \cos(kz) F(R_i, z, \mu) dz)^2 \\
 & + ((\rho_B - \rho_C) \int_0^{R_o} \sin(kz) F(R_o, z, \mu) dz \\
 & \left. + (\Delta_{BA} - \Delta_{CA}) \int_0^{R_i} \sin(kz) F(R_i, z, \mu) dz)^2 \right]
 \end{aligned}$$

where $Q=|\mathbf{Q}|$, $\mu=\cos\theta$, $k=Q\mu$, $\Delta_{BA}=\rho_A-\rho_B$, $\Delta_{CA}=\rho_A-\rho_C$ and the functions $F(R_j, z, \mu)$ are given by

$F(R_j, z, \mu) = \sqrt{R_j^2 - z^2} J_1(Q\sqrt{1-\mu^2} \sqrt{R_j^2 - z^2})$, with J_1 the Bessel function of first order. For this

structure the normalization constant is $V_n = \frac{2}{3}\pi((\rho_B + \rho_C) \cdot (R_o^3 - R_i^3) + 2\rho_A \cdot R_i^3)$.

which describes scattering from the particle on the left of the Figure 1 from their manuscript (below).

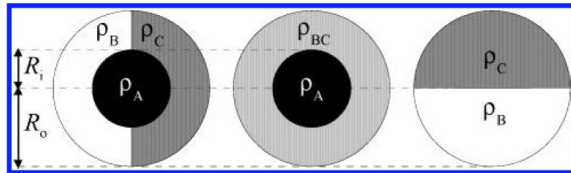
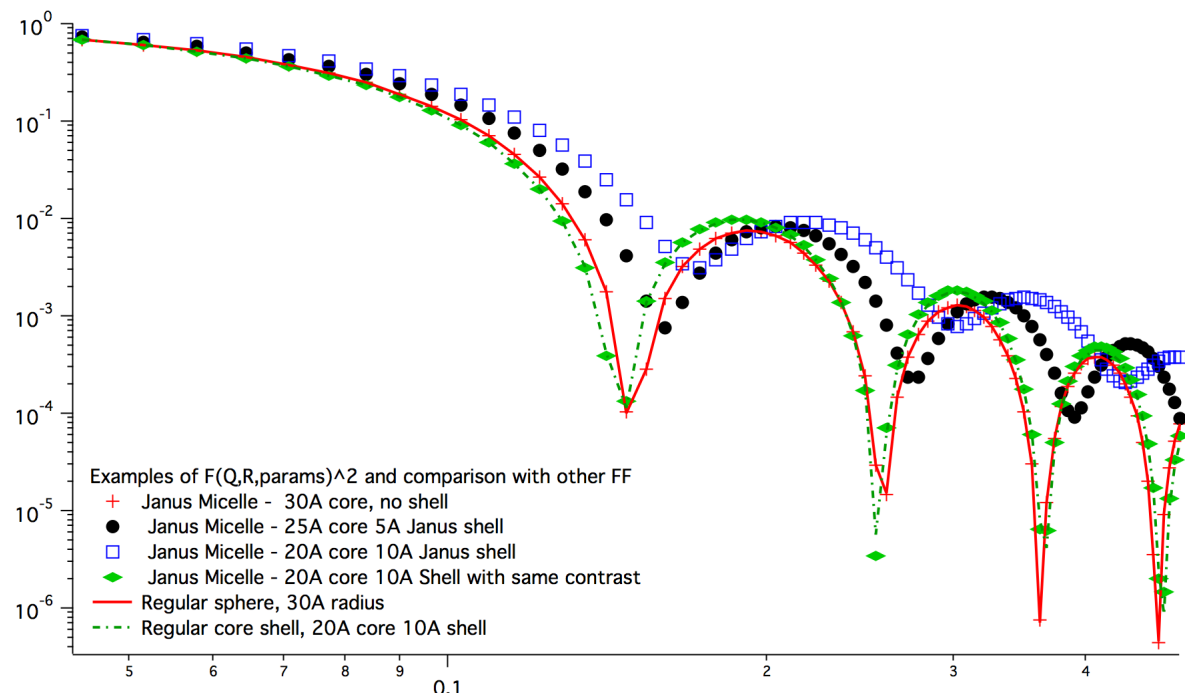


Figure 1. Sketch of competing particle structures. From left to right: Janus micelle, core–shell particle, and simplified Janus micelle.

Example of results:



Note: the results in the above graph are scaled to $F^2(Q=0) = 1$. Since the formula included scattering length densities, normalization by the volume does not result in $F^2(Q=0) = 1$. This may result in unexpected problems with absolute calibration.

This FF is implemented twice...

"Janus CoreShell Micelle 1" ... particle size is total size of the particle (R_0 in the figure in description), parameters:

Shell_Thickness=ParticlePar1	//shell thickness A
CoreRho=ParticlePar2	// rho for core material
Shell1Rho=ParticlePar3	// rho for shell 1 material
Shell2Rho=particlePar4	// rho for shell 2 material
SolventRho=ParticlePar5	// rho for solvent material

"Janus CoreShell Micelle 2" ... particle size here is shell thickness!!! This may be _very_ confusing!!!!, parameters:

Core_Size=ParticlePar1	// Core radius A
CoreRho=ParticlePar2	// rho for core material
Shell1Rho=ParticlePar3	// rho for shell 1 material
Shell2Rho=particlePar4	// rho for shell 2 material
SolventRho=ParticlePar5	// rho for solvent material

"Janus CoreShell Micelle 3" ... particle size is radius of the core (R_i in the figure in description), parameters:

Shell_Thickness=ParticlePar1	//shell thickness A
CoreRho=ParticlePar2	// rho for core material
Shell1Rho=ParticlePar3	// rho for shell 1 material
Shell2Rho=particlePar4	// rho for shell 2 material
SolventRho=ParticlePar5	// rho for solvent material

The reason for the two implementations is, that in usual implementation the shell thickness is fixed while the particle size has size distribution - but this is possible ONLY if core has distribution of sizes. This may be incorrect, as someone can have monodispersed cores, but distribution of shell thicknesses.

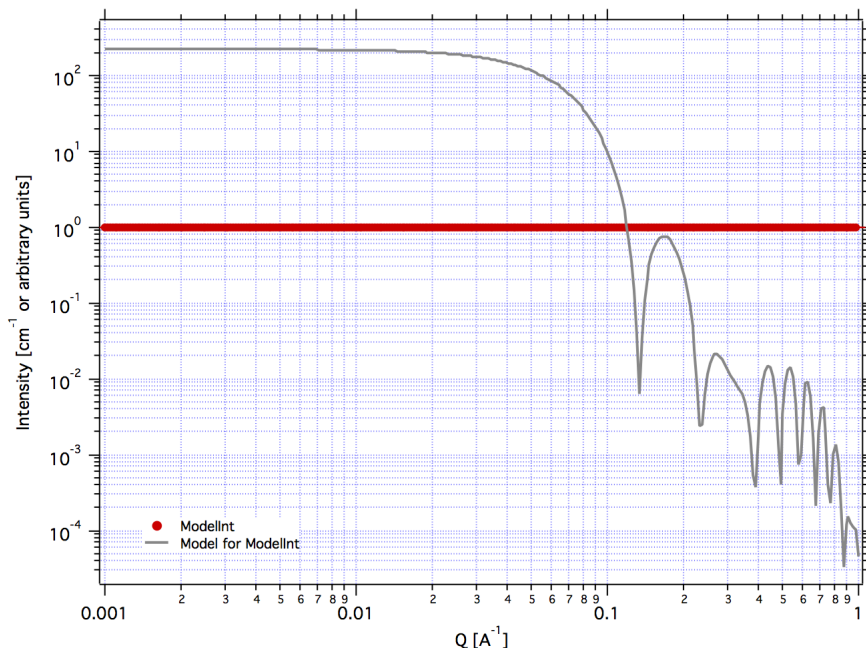
Note, that the "Janus CoreShell Micelle 2 and 3" will not work with some of the tools in Irena as all assume size represents total size (core+shell). Be warned, results will be difficult to present meaningfully! You are on your own...

Model comparison:

Core (Au):	131.5	10^{10} cm^{-1}
Shell 1(Al ₂ O ₃)	34.95	10^{10}
Shell 2(ZrO ₂)	46.27	10^{10}
Solvant (H ₂ O)	9.42	10^{10}
volume = 0.05		

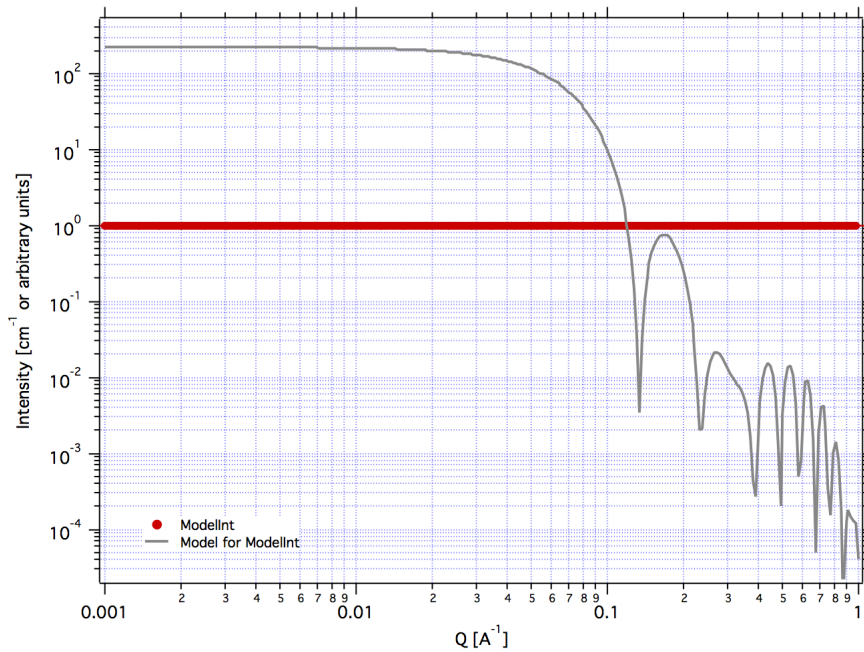
Janus CoreShell Micelle 1:

Mean radius 40A, width 0.3A (Gauss), Shell thickness 10A,

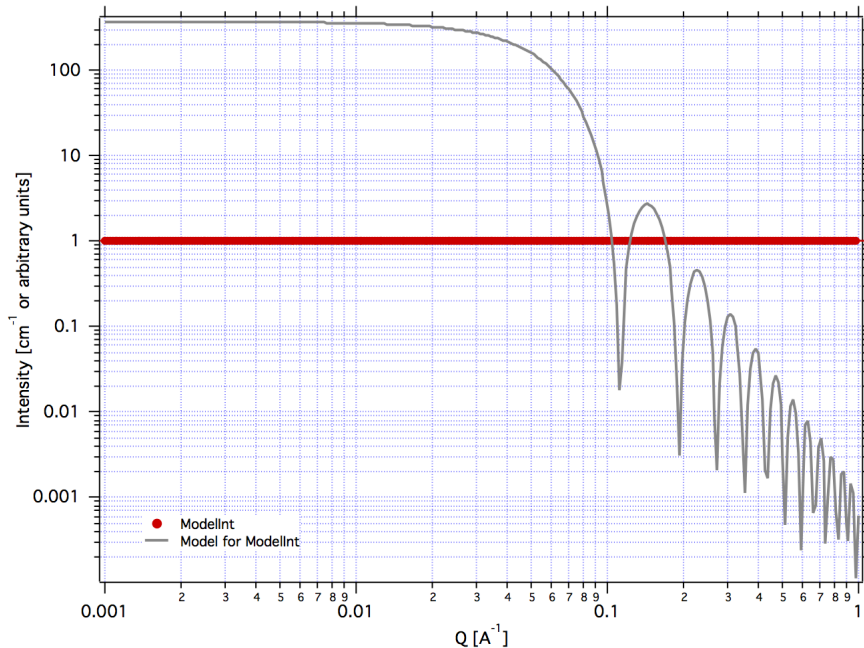


Janus CoreShell Micelle 2:

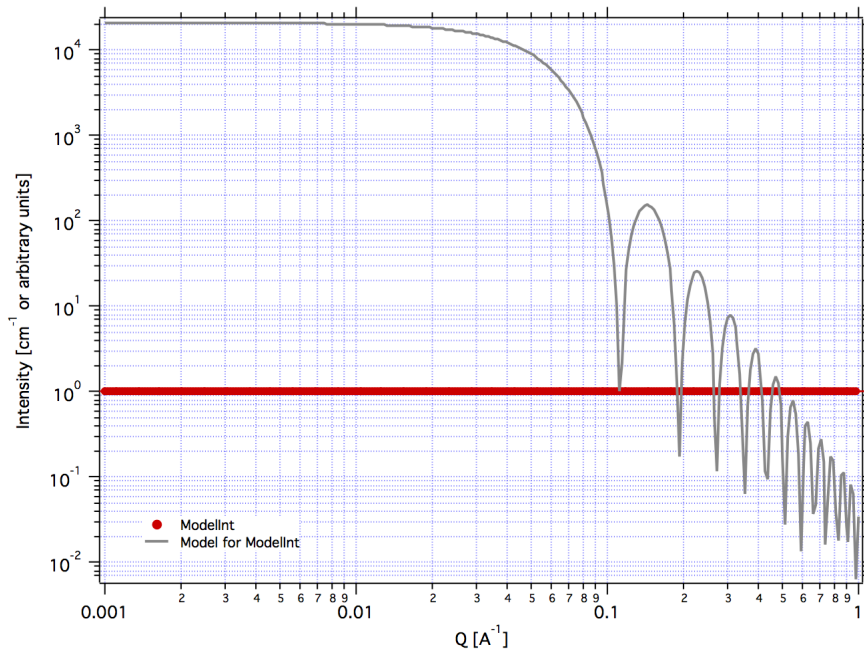
Core radius 30A, Mean radius 40A, width 0.3A (Gauss) :



Janus CoreShell Micelle 1:
Pseudo sphere (shell thickness = 0), Radius = 40 Å,

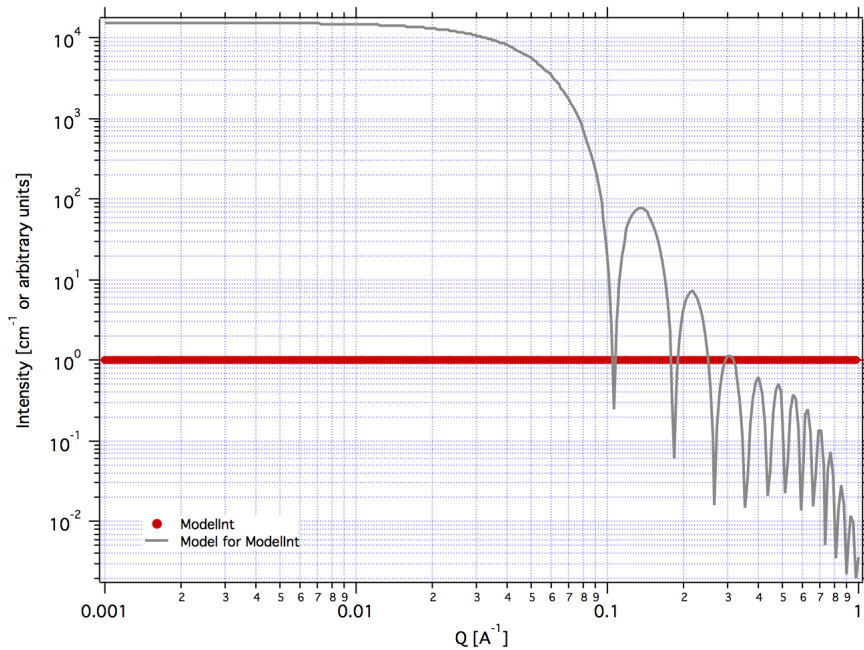


Real sphere, contrast 14903.5 (Au-water):

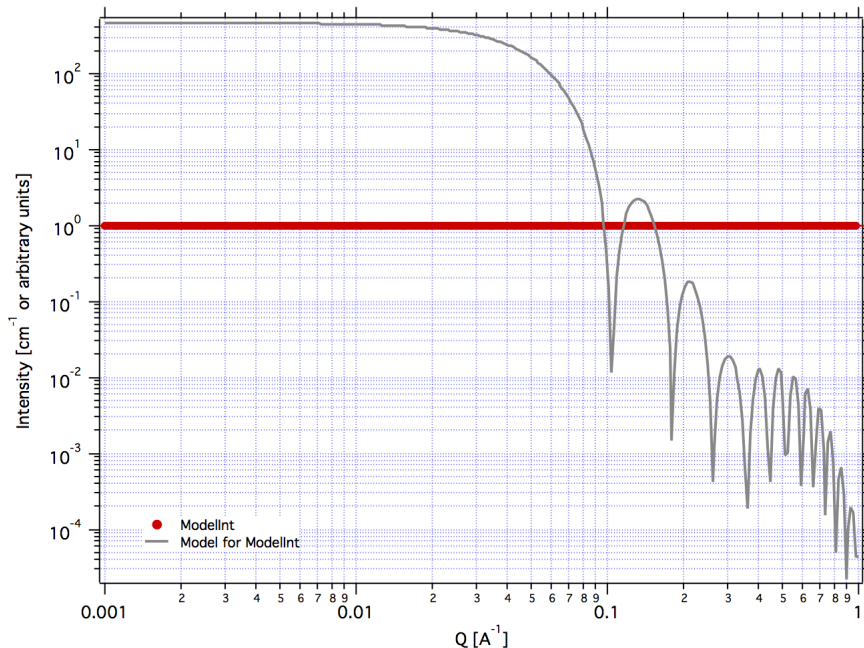


Note the suspicious difference in calibrations. See note above about my suspicion on the problem here...

Real core shell system (pick shell contrast 34.95). Use "Whole particle" as volume.



Janus CoreShell Micelle 1, fake the core shell with same contrast (34.95) for both shells. Recall that the total size of the CoreShell in Irena is radius of core ("Radius") + shell thickness; while for Janus CoreShell Micelle 1 it is just Radius (see figure).



The difference in absolute intensity here is surely related to different assumptions on volume of particle.

User

To use "User" form factor you will need to supply two functions:

- 1. Form factor itself
 - 2. Volume of particle function

Both have to be supplied. Use of form factors which would include volume scaling within is possible, but MUCH more challenging due to other parts of code. If you really insist on doing so, contact me and I will create rules and explanation.

Both functions must work with radius in Angstroems and Q in inverse Angstroems.

Both have to declare following parameters, in following order:

Form factor: Q, radius, par1,par2,par3,par4,par5

Volume : radius, par1,par2,par3,par4,par5

These function are not required to use these 5 user parameters, but they have to declare them.

Examples for sphere:

```

Function IR1T_ExampleSphereFFPoints(Q, radius, par1,par2,par3,par4,par5) //Sphere
  variable Q, radius, par1,par2,par3,par4,par5
  variable QR=Q*radius
  return (3/(QR*QR*QR))*(sin(QR)-(QR*cos(QR)))
end

```

```

Function IR1T_ExampleSphereVolume(radius, par1,par2,par3,par4,par5)           //returns
  variable radius, par1,par2,par3,par4,par5

  return ((4/3)*pi*radius*radius*radius)
end

*****
*****
```

Testing and using Form factors in users own code

To verify that the form factor works for you and to use the form factor if your own functions use following process and functions:

1. Generate Q wave with Qs for which the data are to be calculated
2. Generate intensity wave (will be redimesnioned as necessary, so the only thing is, it should be double precision).
3. Generate distributipon of radii wave - if you want to use single R, create wave with single point
4. decide what you want to calculate:

F^2	powerFct=0
V*F^2	powerFct=1
V^2 * F^2	powerFct=2

5. Run following command:

```
IR1T_GenerateGMatrix(R_FF,Q_wave,R_dist,powerFct,"form factor name",param1,param2,param3,param4,param5,
  "", "")
```

This function will return R_intensity, which is generally matrix with dimensions numpoints(Q_vector) x numpoints(R_dist), if R_dist has 1 point only, returned is wave (vector) as expected and reasonable...

The param1 - param5 are form factor parameters, as desribed in chapter 1, the "" at the end are for user form factor functions (there go the strings with names of user form factor and volume function).

"form factor name" is name from list in chapter 1.

6. Create log-log plot of the data if R_dist has single point. If R_dist has more point, well, you have to pull out the right column of data you need to plot.

Note, that if the IR1T_GenerateGMatrix function returns wave of NaN values if unknown name of form factor is passed in.

Example of code:

```
make/N=100 Q_wave
Q_wave=0.001+p/100
  //will create 100 points wave with values 0.001 to 1) values
```

```

Make/O/D R_FF
    //makes some place for form factor
make R_dist
R_dist=50
//or
//make/N=3 R_dist
//R_dist={10,50,100}
    //creates R distribution and sets values
IR1T_GenerateGMatrix(R_FF,Q_wave,R_dist,powerFct,"form factor name",param1,param2,param3,param4,param5,
    "", "")
    //Note, above lines belong on one line together!
    // replace powerFct with 0, 1, or 2!
// replace "form factor name" with name of form factor you want to use
Display R_FF vs Q_wave
ModifyGraph log=1
    //creates log-log graph of

```


Structure factors description

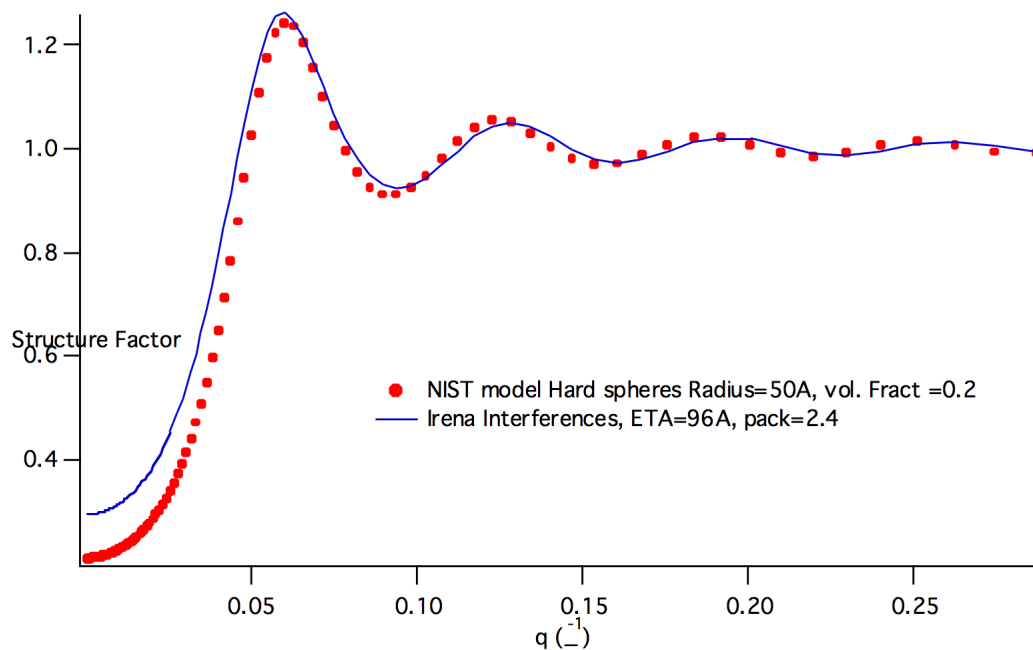
This is list of library of structure factors. These structure factors enable to deal with limited S(Q) effects in Irena package. The functionality is provided by library, which can be called by any other user code. The library provides also GUI for setting the user parameters. In principle, further structure factors can be added if they have less than 5 parameters.

Interferences

This is original structure factor in Irena package. It has been provided as part of Unified fit model by Gregg Beaucage and is listed in his publication: Beaucage, G. (1995). Chapter 9 in ÅgHybrid Organic-Inorganic CompositesÅh, ACS symposium Series 585, edited by J. E. Mark, C. Y-C. Lee, and P. A. Bianconi, 207th National Meeting of the American Chemical Society, San Diego, CA, March 13-17, 1994. American Chemical Society, Washington, DC 1995. Pg. 97 – 111.

$$S(Q) = \frac{1}{1 + k * \frac{3 * (\sin(Q\xi) - Q\xi \cos(Q\xi))}{(Q\xi)^3}}$$

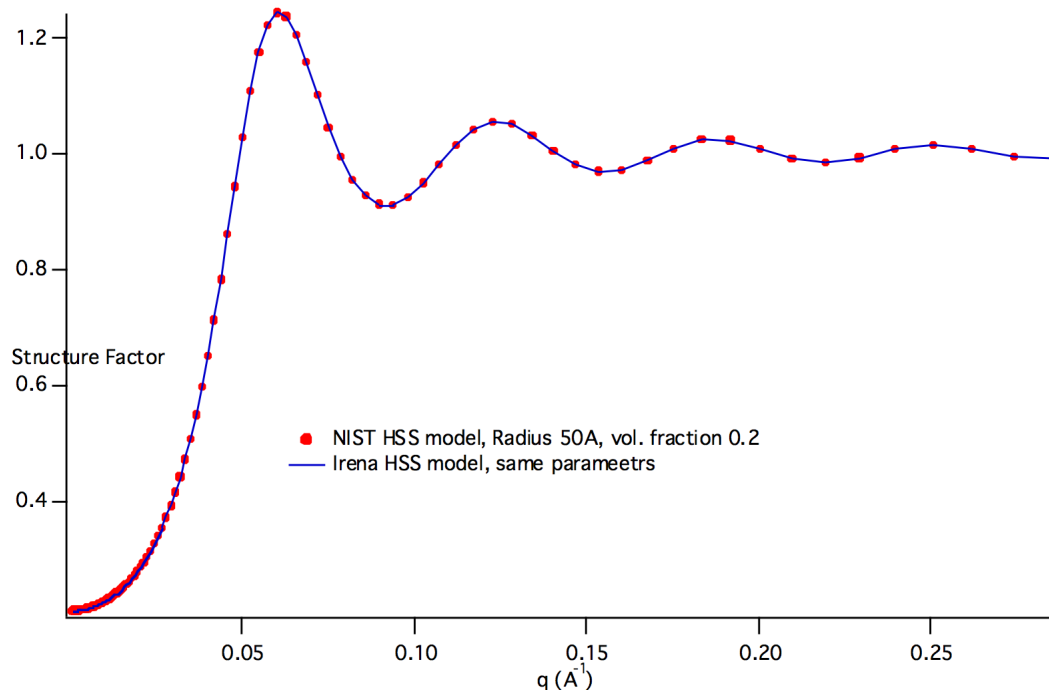
Note, that this model is, for most practical purposes, close to Hard spheres model with different definition of the parameters k (ÅgpackÅh) and f Å (ÅgETAÅh). Modeling II extends the capabilities by including three more structure factors using code available from NIST Igor package (ref). Included are now: Hard spheres, Square Well, and Sticky Hard Spheres, which can be used in addition to interferences model above and no structure factor (dilute limit).



HardSpheres

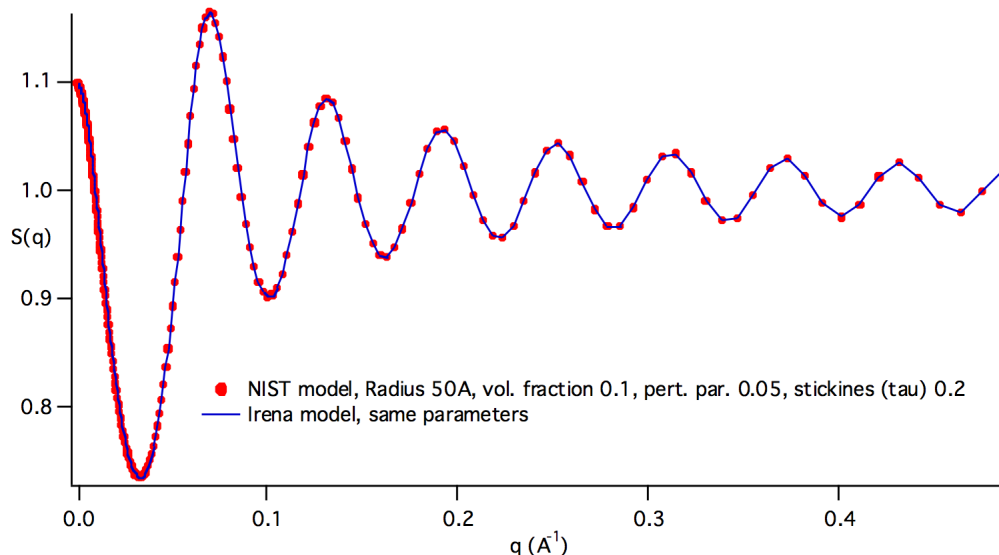
The code for this structure factor has been copied from NIST SAS macros (Kline, S. R. (2006). J Appl Crystallogr **39**, 895-900). Please, give them credit when using this structure factor. (http://www.ncnr.nist.gov/programs/sans/data/data_anal.html)

This is graph of NIST model and Irena implementation.



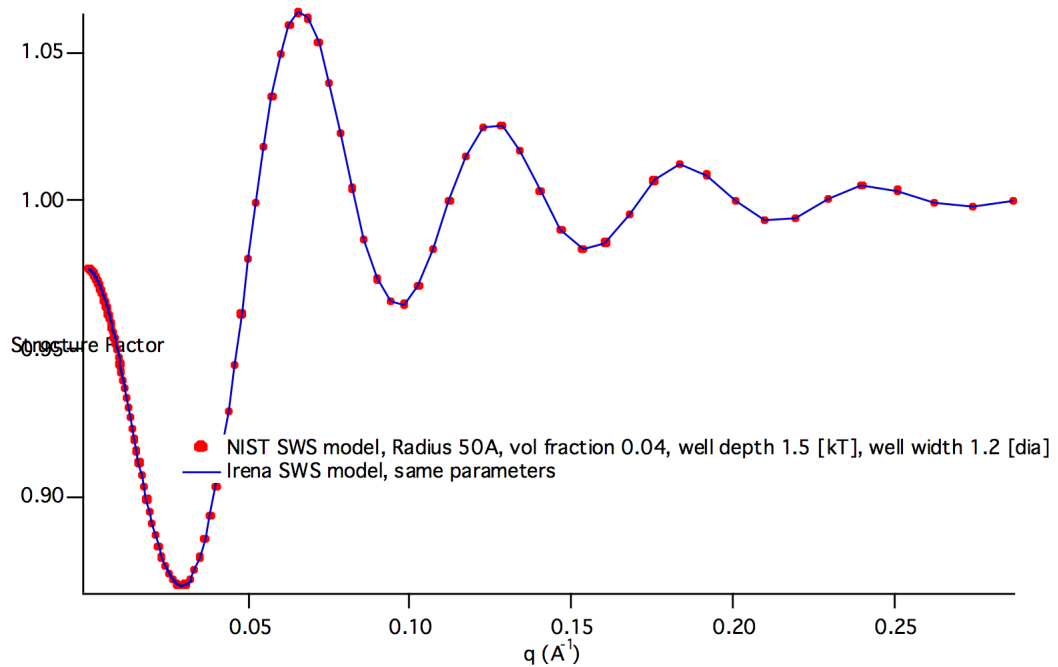
StickyHardSpheres

The code for this structure factor has been copied from NIST SAS macros (Kline, S. R. (2006). J Appl Crystallogr **39**, 895-900). Please, give them credit when using this structure factor. (http://www.ncnr.nist.gov/programs/sans/data/data_anal.html) Åc



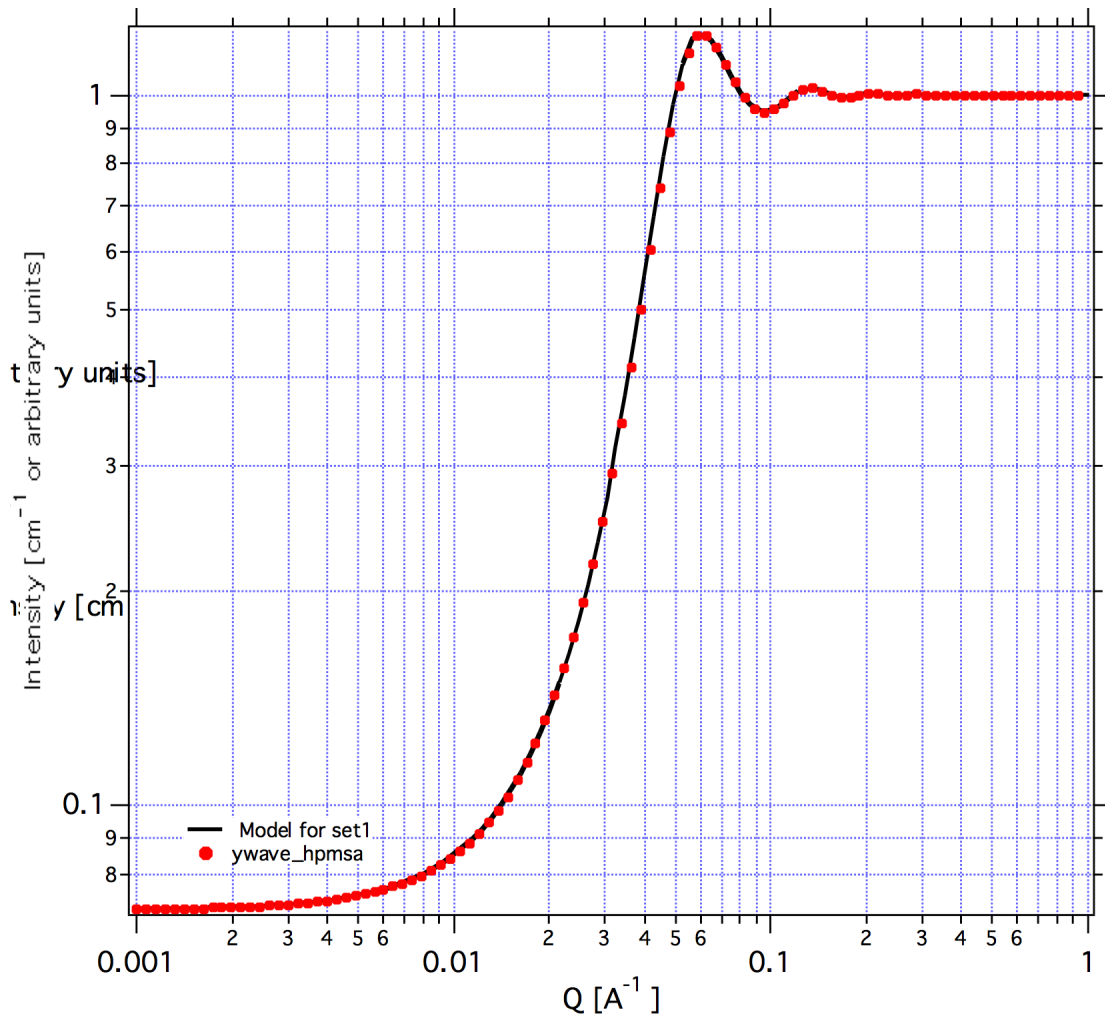
SquareWell

The code for this structure factor has been copied from NIST SAS macros (Kline, S. R. (2006). J Appl Crystallogr **39**, 895-900). Please, give them credit when using this structure factor. (http://www.ncnr.nist.gov/programs/sans/data/data_anal.html) Åc



HayerPenfoldMSA

The code for this structure factor has been copied from NIST SAS macros (Kline, S. R. (2006). For any details on the use of these, please see the original code and description from NIST data analysis package (http://www.ncnr.nist.gov/programs/sans/data/data_anal.html). Please, give them credit when using this structure factor.



This is graph fro standard NIST set of parameters for both Irena package (black line) and NIST package (red dots).

Both assume ONLY structure factor (Form factor is set to 1). The parameters were:

Diameter (A) 41.5 NOTE: Irena uses here radius, which is converted to diameter inside the structure factor.

This is to keep consistency with other structure factors.

Charge 19

Volume Fraction 0.0192

Temperature(K) 298

monovalent salt conc. (M) 0

dielectric constant of solvent 78

Units are mentioned in the help for each filed on the Structure factor panel (you may have to enable help on Mac, it is shown always on PC in the bottom left corner of the Igor window).

Important note: this is comment from original NIST code.

// *** NOTE **** THIS CALCULATION REQUIRES THAT THE NUMBER

OF

// Q-VALUES AT WHICH THE S(Q) IS CALCULATED BE

// A POWER OF 2

!!!!!! this is at this time NOT enforced in Irena implementation...

```
// I am not sure if this is really problem or not.
// How do I find out? Users need to test this for me and if necessary, I
// need to try it out.
// in my testing there was NO problem with the results when the number
// of q pointds was arbitrary number of points...
```


InterPrecipitate

The code for this structure factor has been created on user request for study of precipitation in metals. It is based on formula 6 from APPLIED PHYSICS LETTERS 93, 161904 (2008), Study of nanoprecipitates in a nickel-based superalloy using small-angle neutron scattering and transmission electron microscopy by : E-Wen Huang, Peter K. Liaw, Lionel Porcar, Yun Liu, Yee-Lang Liu, Ji-Jung Kai, and Wei-Ren Chen. This manuscript refers for this formula to paper by R. Giordano, A. Grasso, and J. Teixeira, Phys. Rev. A 43, 6894 (1991). I did not look up original reference, so check it youself to make sure theformula is OKÅc

Structure factor has two parameters - L distance and sigma - root-mean-square deviation (ordering factor):

The interprecipitate structure factor¹³ $S(Q)$ is expressed as a function of Q , L , and σ

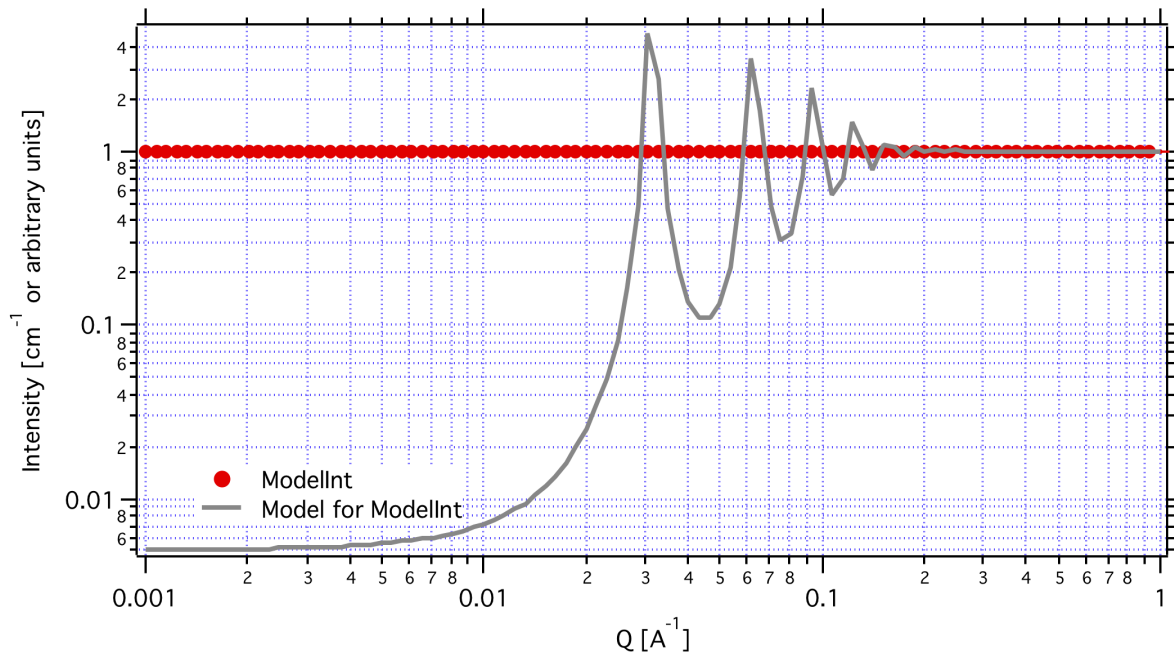
$$S(Q, L, \sigma) = 2 \left\{ \frac{1 - \exp[-(Q^2 \sigma^2)/4] \cos(QL)}{1 - 2 \exp[-(Q^2 \sigma^2)/4] \cos(QL) + \exp[-(Q^2 \sigma^2)/2]} \right\} - 1. \quad (6)$$

In Igor code this is programmed:

```
top = 1 - exp(-(Q^2*sigma^2)/4)*cos(Q*L)
bot = 1-2*exp(-(Q^2 * sigma^2)/4)* cos(Q*L) + exp(-(Q^2*sigma^2)/2)

S(Q,L,sigma) = 2*(top/bot) - 1
```

This is model of the SF for L=200 and Sigma=20 (Sigma/L=10). I have no way of testing this so this formula has not been checked against any data.



Calling the library and use

Users can use built in library in their own code using following calls:

1. initialize by calling: IR2S_InitStructureFactors()
this is where the list of known structure factors is:
SVAR

ListOfStructureFactors=root:Packages:StructureFactorCalc>ListOfStructureFactors

2. use by calling:
IR2S_CalcStructureFactor(SFname,Qvalue,Param1,Param2,Param3,Param4,Param5,Param6)

$I(Q) = I(Q, \text{dilute limit}) *$
IR2S_CalcStructureFactor(SFname,Qvalue,Param1,Param2,Param3,Param4,Param5,Param6)

```
//Dilute
system;Interferences;HardSpheres;SquareWell;StickyHardSpheres;HayterPenfoldMSA
```

3. Get panel by calling:
IR2S_MakeSFParamPanel(TitleStr,SFStr,P1Str,FitP1Str,LowP1Str,HighP1Str,P2Str,FitP2Str,LowP2Str,HighP2Str,P3Str,FitP3Str,LowP3Str,HighP3Str,P4Str,FitP4Str,LowP4Str,HighP4Str,P5Str,FitP5Str,LowP5Str,HighP5Str,P6Str,FitP6Str,LowP6Str,HighP6Str,SFUserSFformula)
to disallow fitting of parameters, simply set FitP1Str="" etc.
then do not have to set low and high limits ...

Structure factors package...
IR2_OldInterferences

this is roughly hard spheres (close to Percus-

Yevick model, not exactly), the ETA = 2* radius and Phi = 8 * vol. fraction for PC model.

IR2_HardSphereStruct	this is Percus-Yevick model
IR2_StickyHS_Struct	this is sticky hard spheres
IR2_SquareWellStruct	this is Square well
IR2_HayterPenfoldMSA	this is HayterPenfoldMSA
IR2_InterPrecipitateSF	this is InterPrecipitate