correct_saxs_Data

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1 Script for the Correction of differential scattering cross section in SAXS experiments (cooking book)

```
In [137]: import numpy as np
        import pandas as pd
        import matplotlib.pyplot as plt
```

URL for SAXS data of glassy carbon reference. There are several different onex available. The signal in high Q is nearly the same for all of them. Only in the low Q region they diverge from one another. The urls shown below correspond to the Glassy_Carbon_A, _B and _C samples from ANL. This script uses a Glassy carbon sample measured in another institute as reference.

Function to correct the thickness of the sample, by measuring an empty cuvette and one filled with water. For that two measurements with transmission and empty beam correspondingly have to be done with H2O and an empty cuvette. The mass attenuation coefficient for water in room temperature is $\frac{\mu}{\rho}=10.37\frac{cm^2}{g}$. The sample thickness d can then be calculated by using:

$$d = -\rho_{H_2O} \frac{\rho}{\mu} ln \left(\frac{T_{H_2O+cell}}{T_{cell}} \right)$$

, with T being the transmission factors for an empty cell and water and $\rho_{H_2O}=1\frac{g}{cm^2}.$

```
In [139]: # Übergabe:
    # - Transmissionfactor water+cell T_wc
    # - only cell T_c
    # - Anode type, most of the time copper

def calc_d(T_wc,T_c,typ='copper'):
```

```
if typ=='copper':
    MAC_w = 10.37 #cm²/g, for copper anode

rho_w = 1 #g/cm³

d = -np.log(T_wc/T_c)*1/(MAC_w*rho_w) #cm
return(d)
```

Function to calculate the transmission factor for intensity of sample S and empty beam intensity I_0 :

$$T_S = \frac{I_S}{I_0}$$

This Function calulates the Scaling or Correction Factor with a Glassy Carbon measurement done in your setup and scales it so it fits the intensity of the 'Glassy-Carbon-A' sample from ANL. You can chose whatever reference sample you like. They should turn out more or less the same. For consistency's sake, I will solely use the A sample. The formula for the SF calculation follows from Fan-Paper:

$$SF = \frac{\left(\frac{\delta \Sigma}{\delta \Omega}\right)_{st}}{\frac{(I_{st} - BG_{st})}{d_{st}T_{st+cell}}}$$

,with 'st' corresponding to the standard sample, in my case glassy carbon. BG corresponds to the background measurement and T the transmission factor for the GC measurements.

```
SF_vec = DSC_Ref/((DSC_GC)/(d_GC*T_GC))
return(SF_vec)
```

Finally the differential scattering crosssection for sample s will be correctly calculated using all previously determined parameters SF and d_s :

2 From here on follows an example

From here follows an example using sample TiS710 and data measured on our Anton Paar machine. First the intensities are extracted from .stat files. This process will not be discussed further, because it's simply parsing the files contained in the in 'dir' specified folder automatically. This step corresponds to point (1) in the cooking book. Calculate the transmission factors.

```
In [143]: import os

dir = '/home/zech/FAUbox/PhD/Auswertung/SAXS/test_data/'

l_EB_Stats = []

l_TR_Stats = []

table = pd.DataFrame(columns = ['TR', 'EB', 'I_TR', 'I_EB', 'T'])

pairs = pd.DataFrame(columns = ['typ', 'samp'])

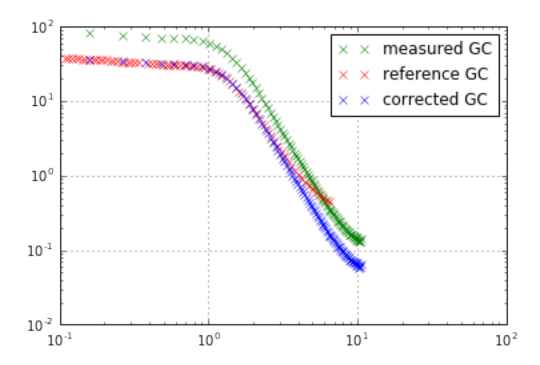
for iFile in os.listdir(dir):
    if os.path.isfile(os.path.join(dir, iFile)):
        if '.stat' in iFile:
        enum = iFile.split('_')[0]
        typ = iFile.split('_')[2]
        samp = iFile.split('_')[1]
        pairs.loc[str(enum)] = [str(typ), str(samp)]

for iFile in os.listdir(dir):
    if os.path.isfile(os.path.join(dir, iFile)):
```

```
for line in open(os.path.join(dir,iFile),'r'):
                           if 'Total intensity' in line:
                               inty = line.split()[3]
                               enum = iFile.split('_')[0]
                               samp = pairs.loc[str(enum), 'samp']
                               typ = pairs.loc[str(enum),'typ']
                               if str(samp) not in table.index:
                                   table.loc[str(samp)] = [0,0,0,0,0]
                               if typ == 'TR':
                                   table.loc[str(samp), 'TR'] = enum
                                   table.loc[str(samp),'I_TR'] = inty
                               elif typ == 'EB' or 'DB':
                                   table.loc[str(samp),'EB'] = enum
                                   table.loc[str(samp),'I_EB'] = inty
          for iInd in table.index:
              table.loc[iInd,'T'] = calc_T(float(table.loc[iInd,'I_TR']),float(table.loc[iInd,'I_TR'])
          table
Out[143]:
                   TR
                          EΒ
                                      I_TR
                                                    I_EB
                01744 01743 1.45425E+05 5.66515E+05 0.256701
          H20
          ЕC
                01746 01747 3.65022E+05 5.57136E+05 0.655176
                                                          0.535951
                01724 01723 3.75920E+05 7.01408E+05
          GC
          SGNR 01727 01728 2.11857E+05 7.99840E+05
                                                          0.264874
  In the next step the SF will be determined via the glassy carbon measurement. This is step (2).
In [144]: data_GC = np.loadtxt('/home/zech/FAUbox/PhD/Auswertung/SAXS/test_data/01")
          sample_GCA = np.loadtxt('/home/zech/FAUbox/PhD/Auswertung/SAXS/reference,
          plt.loglog(data_GC[:,0],data_GC[:,1],'xg', label='measured GC')
          plt.loglog(sample_GCA[:,0],sample_GCA[:,1],'xr',label='reference GC')
          data_BG_GC = np.zeros(shape=(len(data_GC[:,0])))
          SF = calc_SF(I_GC=data_GC[:,1],
                       BG_GC=data_BG_GC,
                        Q_GC=data_GC[:,0],
                        T_GC=table.loc['GC','T'],
                        d_{GC}=0.001)
          d_{GC}=0.001
          T_GC=table.loc['GC','T']
          plt.loglog(data_GC[:,0],data_GC[:,1]/(d_GC*T_GC)*SF,'xb',label='corrected
          plt.legend(loc='best')
          plt.grid()
          plt.show()
          print('calculated scaling factor SF = ' +str(SF))
```

#print(os.path.join(dir,iFile))

if '.stat' in iFile:



calculated scaling factor SF = 0.000243927104044

In step (3) the thickness of the samples will be calculated from H2O and empty cell measurements.

In the end of this procedure (step (4)) the differential scattering cross section of the sample will be calculated, by using the parameters determined in the steps above.

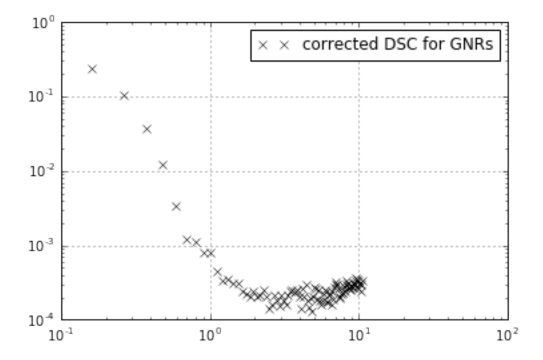
```
In [146]: T_sc = table.loc['SGNR','T']

    data_s = np.loadtxt('/home/zech/FAUbox/PhD/Auswertung/SAXS/test_data/0172
    data_H2O = np.loadtxt('/home/zech/FAUbox/PhD/Auswertung/SAXS/test_data/01

I_s = data_s[:,1]
    BG_s = data_H2O[:,1]
```

 $DSC_s = SF*(I_s-BG_s)/(d_s*T_sc)$

```
In [147]: plt.loglog(data_s[:,0],DSC_s,'xk',label='corrected DSC for GNRs');
     plt.legend()
     plt.grid()
     plt.show()
```



In []: