MEP Introduction Report

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Abstract

MEP Introduction Report on the master thesis project of Isabel Postmes, supervised by Sonia Conesa Boj. My Master thesis will revolve around the automization of the electron energy loss spectrum (EELS) analysis, documented by the transmission electron microscope (TEM). The focus will be on the low-loss spectrum, where the influence of the zero loss peak (ZLP) is not negligible, and will include an error estimation, which makes it unique. The first step in this was done by Laurien Roest: the detection of the ZLP. I will focuss in my months in generilizing the program Laurien wrote, and I will expand it by including the Kramer-Kronig analysis and peak detection. The final goal is to be able to input an image of spectra, and have an automatic output of what pixels statisfy certain criteria.

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1 Planning

The first couple of weeks will be dedicated to getting acquainted with the subjects at hand, and Lauriens program. Then I will build the program step-by-step: starting by implementing the Fourier Deconvolution, continuing to subtracting the dielectric function with the Kramer-Kronig analysis and than working on the peak detection. Eventually I will implement all this into a program which takes a image of spectra, and gives a visual output and list of pixels depending on what characteristics are selected.

To test the final program, we will measure layered materials

2 Introduction

The supervisor on my project is Prof. Sonia Conesa Boj. Furthermore, I will be supervised in an unofficial capacity by dr. Juan Rojo of the Nikhef Institute in Amsterdam. The project is a continuation of the thesis project of Laurien Roest, whose paper can be found at

One of the features of the TEM is to obtain electron energy loss spectra of the reviewed sample. This spectrum documents the loss in kinetic energy of electrons travelling through a sample. [1]. Interpretation of these spectra can help in determining the composition and structure of the evaluated sample.

3 Background info

In the sections below, you will find a first overview of some of the critical background information.

3.1 TEM

The transmission electron microscope (TEM) projects a beam of electron trough an sample, to record the transmitte electrons on the opposite side of the sample to form an image. Due to the fact that the de Broglie wavelenghts of electrons (10^{-2}) are factors smaller than the wavelength of photons (10^{2}), mugh higher resolutions can be obtained than with light microscopy. The resolution of a TEM is mostly determined by the focusing power of the electron beam. For TEMs with monochromator, such as the TEM available at the Conesa Boj Laberatorium, resolutions of 0.1nm can be obtained, small enough to image single atoms.

3.1.1 EELS of the TEM

The electron energy loss spectrum (EELS) documents the loss in kinetic energy of electrons in their path through a specimen. Due to the quantification of energylevels at elementary level, this energy loss manifests in peaks in the spectrum. The broadness of the observed peaks mainly comes from the inherent imperfect electron source, which transmits electrons in a certain energy distribution focussed around the target energy. The analysis of these peaks gives inside in what energy levels are present in the sample, and with that gives inside in the composition and structure of the specimen.

3.1.2 Low loss spectrum

Much of the interesting information in the EELS is nested in the low loss part (¡50eV) of the spectrum. Here, ones finds info on bandgaps, plasmons and exitons among others. A significant problem in this part of the spectrum, is that the influence of the zero loss peak (ZLP) is non-negliglable. Older tactices to avoid this problem, are centered around fitting the ZLP in each individual spectrum, and subsequently substracting it. The most prominent problem with this approach is that there is no indication of the error in the estimation of the ZLP. Therefore, it can not be said with how significant the found peaks near the tail of the ZLP are. In her master thesis, Laurien Roest developed a method in which the ZLP is approximated by a neural network, which inherently results in a ZLP with error marges.

Since the ZLP is an indication of the distribution of the energies of the electrons produced by the electron source, the errors around the ZLP trackle trhough in all the rest of the spectrum. Therefore, in my project I will use the program developed by Laurien to estimate the ZLP, and use this in the detection and quantification of the peaks in the spectrum and the other variables to be extracted from the spectrum.

3.2 Dielectric function

The dielectric function, also called permitivity, is a measure for the polarisability of a material. From the dielectric function, a multiplitude of other characaristics can be calculated. Since the dielectric function can be extracted from the electron energy loss spectrum through Kramer-Kronig analysis, for an image of spectra, the dielectric function can be calculated for each pixel.

4 Calculations on the spectrum

4.1 Approximating continious Fourier transform with dicrete Fourier transform

Lets start with a small refresher on Fourier transforms, and evaluate how to approximate the continious Fourier transform with the discrete Fourier transform.

4.1.1 Original definitions

The continous Fourier transfrom (CFT) of function f(x) is defined as:

$$\mathcal{F}\left\{f(x)\right\} = F(\nu) = \int_{-\infty}^{\infty} e^{-i2\pi\nu x} f(x) dx. \tag{4.1}$$

The inverse of the CFT is given by:

$$\mathcal{F}^{-1}\{F(\nu)\} = f(x) = \int_{-\infty}^{\infty} e^{i2\pi\nu x} F(\nu) d\nu.$$
 (4.2)

The discrete Fourtier transform (DFT) of discrete function f[n] defined on $n \in \{0,...,N-1\}$ is given by:

DFT
$$\{f[n]\}\ = F[k] = \sum_{n=0}^{N-1} e^{-i2\pi kn} f[n], \forall k \in \{0, ..., N-1\}.$$
 (4.3)

The inverse of the CFT is given by:

$$DFT^{-1} \{F[k]\} = f[n] = \sum_{k=0}^{N-1} e^{i2\pi kn} F[k], \forall n \in \{0, ..., N-1\}.$$
 (4.4)

4.1.2 Discrete approximation CFT

If one approximates function f(x) by $f[x_0+n\Delta x], \forall n \in \{0,...,N-1\}$, and ν by $k\Delta \nu, \forall k \in \{0,...,N-1\}$, where:

$$\Delta x \Delta \nu = \frac{1}{N},\tag{4.5}$$

starting from eq. (4.1), one can obtain:

$$F(\nu) \approx F[k\Delta\nu] = \sum_{n=0}^{N-1} \Delta x f[x_0 + n\Delta x] \exp\left[-i2\pi k\Delta\nu(x_0 + n\Delta x)\right],$$

$$= \Delta x \exp\left[-i2\pi k\Delta\nu x_0\right] \sum_{n=0}^{N-1} \exp\left[-i2\pi nk/N\right] f[x_0 + n\Delta x],$$

$$= \Delta x \exp\left[-i2\pi k\Delta\nu x_0\right] \text{DFT} \left\{f[n]\right\}.$$

$$(4.6)$$

Similarly, we can reobtain the original function $f(x) \approx f[n\Delta x + x_0]$ from the approximation of $F(\nu) \approx F[k\Delta\nu]$, using eq. (4.2):

$$f(x) \approx f[n\Delta x + x_0] = \sum_{k=0}^{N-1} \Delta \nu F[k\Delta \nu] \exp\left[i2\pi k\Delta \nu (n\Delta x + x_0)\right]$$

$$= \Delta \nu \sum_{k=0}^{N-1} F[k\Delta \nu] \exp\left[i2\pi k\Delta \nu x_0\right] \exp\left[i2\pi kn/N\right].$$
(4.7)

Defining:

$$G[k\Delta\nu] = \exp\left[i2\pi k\Delta\nu x_0\right] F[k\Delta\nu],\tag{4.8}$$

we find:

$$f(x) \approx f[n\Delta x + x_0] = \Delta \nu \sum_{k=0}^{N-1} G[k\Delta \nu] \exp[i2\pi kn/N]$$

= $\Delta \nu \operatorname{DFT}^{-1} \{G[k\Delta \nu]\}$ (4.9)

4.1.3 Convolutions

Definition of concolution The concolution of two functions f(x) and g(x) is defined as te integral:

$$h(x) = \int_{-\infty}^{\infty} f(\bar{x})g(x - \bar{x})d\bar{x}.$$
 (4.10)

Assuming f(x) and g(x) have a limited domain, h(n) will also have a limited domain, given by:

$$f(x) = \begin{cases} f(x), & x_{0,f} \le x \le x_{1,f} \\ 0, & x < x_{0,f} \lor x > x_{1,f} \end{cases},$$

$$g(x) = \begin{cases} g(x), & x_{0,g} \le x \le x_{1,g} \\ 0, & x < x_{0,g} \lor x > x_{1,g} \end{cases},$$

$$h(x) = \begin{cases} h(x), & x_{0,f} + x_{0,g} \le x \le x_{1,f} + x_{1,g} \\ 0, & x < x_{0,f} + x_{0,f} \lor x > x_{1,f} + x_{1,g} \end{cases}.$$

$$(4.11)$$

This simplifies eq. (4.10) to:

$$h(x) = \int_{x_{0,f}}^{x_{1,f}} f(\bar{x})g(x - \bar{x})d\bar{x}.$$
 (4.12)

Futhermore, the Fourier transform of the convolution has the beautiful property that, for the limited domain functions, but this can be extended into infinity:

$$H(\nu) = \int_{x_{0,1}+x_{0,g}}^{x_{1,f}+x_{1,g}} h(x) \exp\left[-i2\pi\right] dx,$$

$$= \int_{x_{0,1}+x_{0,g}}^{x_{1,f}+x_{1,g}} \int_{x_{0,1}}^{x_{1,f}} f(\bar{x})g(x-\bar{x})d\bar{x} \exp\left[-i2\pi\nu x\right] dx,$$

$$= \int_{x_{0,1}}^{x_{1,f}} \int_{x_{0,1}+x_{0,g}}^{x_{1,f}+x_{1,g}} g(x-\bar{x}) \exp\left[-i2\pi\nu x\right] dx \quad f(\bar{x})d\bar{x}.$$
(4.13)

Defining $\hat{x} = x - \bar{x}$, evaluation the limits on the integrals and realising d(x) = dx, you obtain:

$$H(\nu) = \int_{x_{0,f}}^{x_{1,f}} \int_{x_{0,g}}^{x_{1,g}} g(\hat{x}) \exp\left[-i2\pi\nu(\hat{x} + \bar{x})\right] d\hat{x} \quad f(\bar{x}) d\bar{x},$$

$$= \int_{x_{0,g}}^{x_{1,g}} g(\hat{x}) \exp\left[-i2\pi\nu\hat{x}\right] d\hat{x} \int_{x_{0,f}}^{x_{1,f}} f(\bar{x}) \exp\left[-i2\pi\nu\bar{x}\right] d\bar{x},$$

$$= F(\nu)G(\nu). \tag{4.14}$$

Discretisation of the convolution and its CFT Again, we can approximate f(x), g(x) and h(n) by:

$$f(x) \approx f[n_f(\Delta x)_f + x_{0,f}], \forall n_f \in \{0, N_f - 1\}, (\Delta x)_f = (x_{1,f} - x_{0,f})/N_f,$$

$$g(x) \approx g[n_g(\Delta x)_g + x_{0,g}], \forall n_g \in \{0, N_g - 1\}, (\Delta x)_g = (x_{1,g} - x_{0,g})/N_g.$$
(4.15)

If we ensure that $(\Delta x)_f = (\Delta x)_g \equiv \Delta x$, we can approximate h(n):

$$h(x) \approx h[n_h \Delta x + x_{0,f} + x_{0,q}], \forall n_h \in \{0, N_f + N_q - 1\},$$
 (4.16)

where

$$h[n_h \Delta x + x_{0,f} + x_{0,g}] = \sum_{k=0}^{N_f - 1} f[k \Delta x + x_{0,f}] g[n \Delta x + x_{0,g} - k \Delta x] \Delta x.$$
 (4.17)

To discretisize the continious fourier transfrom, we once again need to define $\Delta \nu$, but mind the difference for $\Delta \nu$ for the discretisations of f(x) or g(x) and h(x):

$$(\Delta \nu)_f = \frac{1}{N_f \Delta x}$$

$$(\Delta \nu)_g = \frac{1}{N_g \Delta x}$$

$$(\Delta \nu)_h = \frac{1}{(N_f + N_g) \Delta x}$$

$$= \left((\Delta \nu)_f^{-1} + (\Delta \nu)_g^{-1} \right)^{-1}$$

$$(4.18)$$

Now the discrete approximation of the CFT of the convolution (given by eq. (4.13)) is given by:

$$H(\nu) \approx H[k(\Delta\nu)_h] = \sum_{n=0}^{N_f + N_g - 1} h[n\Delta x + x_{0,f} + x_{0,g}] \exp\left[-i2\pi k(\Delta\nu)_h(n\Delta x + x_{0,f} + x_{0,g})\right] \Delta x,$$

$$= \sum_{n=0}^{N_f + N_g - 1} \sum_{l=0}^{N_f - 1} f[l\Delta x + x_{0,f}]g[(n-l)\Delta x + x_{0,g}] \exp\left[-i2\pi k(\Delta\nu)_h(n\Delta x + x_{0,f} + x_{0,g})\right] (\Delta x)^2.$$

$$(4.19)$$

Defining m = n - l, and swapping summation order, you obtain:

$$H[k(\Delta\nu)_{h}] = (\Delta x)^{2} \exp[-2\pi i k(\Delta\nu)_{h} x_{0,f}] \sum_{l=0}^{N_{f}-1} f[l(\Delta\nu)_{h} + x_{0,f}]$$

$$\exp[-2\pi i k(\Delta\nu)_{h} x_{0,g}] \sum_{m=-l}^{N_{f}+N_{g}-l-1} g[m\Delta x + x_{0,g}] \exp[-2\pi i k(\Delta\nu)_{h} (m+l)\Delta x],$$
(4.20)

reevaluating the boundries on the second summation, inputing eq. (4.18) and shuffling above result leads to:

$$H[k(\Delta\nu)_{h}] = \exp[-2\pi i k(\Delta\nu)_{h} x_{0,f}] \sum_{l=0}^{N_{f}-1} f[l(\Delta\nu)_{h} + x_{0,f}] \exp[-2\pi i k l/(N_{f} + N_{g})]$$

$$\exp[-2\pi i k(\Delta\nu)_{h} x_{0,g}] \sum_{m=0}^{N_{g}-1} g[m\Delta x + x_{0,g}] \exp[-2\pi i k h m/(N_{f} + N_{g})].$$
(4.21)

Looking at the above equation, one almost recognizes a multiplication of two approximate CFT's (see eq. (4.6)). However, there is a significant difference, which means it is not a simple multiplication of the two CFT's: where in the CFT of the singular function, you see that $\Delta \nu = 1/N_f$ or $\Delta \nu = 1/N_g$ resectivily, whereas in eq. (4.21), $\Delta \nu = 1/(N_f + N_g)$, as given by eq. (4.18). Nor is it a simple factor with which the CFT's could be multiplied: the factor is present in the exponential within the summations. Each single exponential term in the summations need to raised to the power $(N_f + N_g)/N_f$ and $(N_f + N_g)/N_g$ respectively. THIS IS THE PROBLEM.

4.1.4 Other calculations on CFTs

For the deconvolution of the spectra, we will need the devision and logaritmic values of two CFT's.

4.2 Obtaining the single scattering distribution

4.2.1 Build-up of measured spectrum

When electrons go through the sample, the intensity of electrons that has no inelastic scattering is given by the zero-loss peak: $I_{ZLP}(E)$. The intensity of the electrons that do scatter, $I_{EEL}(E)$, is than dividable in the single scatter intensity, $I_1(E)$, the double scatter intensity, $I_2(E)$, the triple scatter intensity, $I_3(E)$, etc:

$$I(E) = I_{ZLP}(E) + I_{EEL}(E) = I_{ZLP}(E) + \sum_{n=0}^{\infty} I_n(E).$$
 (4.22)

MAYBE DISREGARD? The integrated intensity of each n-scattering spectrum N_n is this a logical choice of letter? depends on the total integrated intensity N, assuming independed scattering events, through the bionomal distribution:

$$N_n = \frac{N}{n!} \left(\frac{t}{\lambda}\right)^n \exp\left[-t/\lambda\right]. \tag{4.23}$$

Here t is the thickness of the sample, and λ is the mean free path of electrons in the sample. END DISREGARD

Since we know the zero-loss-peak due to Lau [2], the response function of the instrument, R(E), is easily deducted by:

$$R(E) = I_{ZLP}(E)/N_{ZLP}, \tag{4.24}$$

where N_{ZLP} is the integrated intensity of $I_{ZLP}(E)$.

Now we will show how the total recorded spectrum is build up from single-scattering distribution S(E), and the above defined response function R(E).

The spectrum recorded due to the single scattering events, $J^1(E)$, is these two distributions convoluted:

$$I_1(E) = R(E)^* S(E) \equiv \int_{-\infty}^{\infty} R(E - E') S(E') dE'.$$
 (4.25)

It can be easily seen, that as a double-scattering event, is a series of two single-scattering event, the double-scattering intensity is given by the self convolution of the single-scattering intensity, normalised to match eq. (4.23), and once again convoluted with the response function:

$$I_2(E) = R(E)^* S(E)^* S(E) / (2! N_{ZLP}).$$
 (4.26)

For higher order scattering spectra, this generalises to HOW TO WRITE DOWN A SUCCESSION OF CONVOLUTIONS?:

$$I_n(E) = R(E)^* S(E)^{*} S(E)^{n-1} / (n! N_{ZLP}^{n-1}).$$
(4.27)

The complete recorded spectrum, neglecting any surface plasmons, is than given by (analogous to eq. (4.22)):

$$I(E) = I_{ZLP}(E) + I^{1}(E) + I^{2}(E) + I^{3}(E) + \cdots$$

$$= I_{ZLP}(E)^{*} \left[\delta(E) + S(E)/N_{ZLP} + S(E)^{*}S(E) / \left(2!N_{ZLP}^{2} \right) + S(E)^{*}S(E)^{*}S(E) / \left(3!N_{ZLP}^{3} \right) + \cdots \right]$$

$$= I_{ZLP}(E)^{*} \left[\delta(E) + \sum_{n=1}^{\infty} R(E)^{*}S(E) \left[{}^{*}S(E) \right]^{n-1} / \left(n!N_{ZLP}^{n-1} \right) \right].$$
(4.28)

Since a convolution in energy domain translates to a multiplication in the frequency domain, it makes sense to take the Fourier transform (FT) of the above equation. Eq. (4.28) than becomes, using the taylor expansion of the exponential function:

$$i(\nu) = z(\nu) \left\{ 1 + s(\nu)/N_{ZLP} + [s(\nu)]^2 / \left(2! N_{ZLP}^2 \right) + [s(\nu)]^3 / \left(3! N_{ZLP}^3 \right) + \cdots \right\}$$

$$= z(\nu) \sum_{n=0}^{\infty} \frac{s(\nu)^n}{n! N_{ZLP}^n}$$

$$= z(\nu) \exp\left[s(\nu)/N_{ZLP} \right],$$
(4.29)

where $i(\nu)$ is the FT of the intensity spectrum, $z(\nu)$ is the FT of the zero-loss peak, and $s(\nu)$ is the FT of the single-scattering distribution.

The single scattering distribution can than be retrieved by rewriting eq. (4.29), and taking the inverse Fourier transform:

$$s(\nu) = N_{ZLP} \ln \left(\frac{i(\nu)}{z(\nu)} \right), \tag{4.30}$$

$$S(E) = \mathcal{F}^{-1} \left\{ s(\nu) \right\}$$

$$= \mathcal{F}^{-1} \left\{ N_{ZLP} \ln \left[\frac{i(\nu)}{z(\nu)} \right] \right\}$$

$$= \mathcal{F}^{-1} \left\{ N_{ZLP} \ln \left[\frac{\mathcal{F} \left\{ I(E) \right\}}{\mathcal{F} \left\{ I_{ZLP}(E) \right\}} \right] \right\}$$

$$(4.31)$$

However, eq. (4.36) only works for an "ideal" spectrum. Any noise on the spectrum will blow up, as

Discussion points What is the most official way to calculate the errors in R(E), from the errors in the ZLP? Just calculate for each ZLP and take the std of those, or can you use the error of the ZLP to calculate it at once? Because it comes back in the N_{ZLP} as well.

There are in NaN values in the $s(\nu)$ calculation in the python file, how best to handle those, just set them to 0?

Looking at the S(E) vs I_{ZLP} etc plot, there are some things that don't make sense

4.3 Kramer-Kronig relations

The Kramer-Kronig relations are two functions that relate the imaginary part of an complex function to the real part and vice versa. The relations hold as long as the complex function is analytic in the upper half-plane. The relations for function $\chi(\omega) = \chi_1(\omega) + i\chi_2(\omega)$, with ω a complex variable are given by [3]:

$$\chi_1(\omega) = \frac{1}{\pi} \mathcal{P} \int_{-\infty}^{\infty} \frac{\chi_2(\omega')}{\omega' - \omega} d\omega', \tag{4.32}$$

and:

$$\chi_2(\omega) = -\frac{1}{\pi} \mathcal{P} \int_{-\infty}^{\infty} \frac{\chi_1(\omega')}{\omega' - \omega} d\omega'. \tag{4.33}$$

Here \mathcal{P} denotes the Cauchy principal value of the integral. For causal functions, due to (anti)symmetries arrising from its causality, these can be rewritten to [3]:

$$\chi_1(\omega) = \frac{2}{\pi} \mathcal{P} \int_0^\infty \frac{\omega' \chi_2(\omega')}{\omega'^2 - \omega^2} d\omega', \tag{4.34}$$

and:

$$\chi_2(\omega) = -\frac{2}{\pi} \mathcal{P} \int_0^\infty \frac{\omega \chi_1(\omega')}{\omega'^2 - \omega^2} d\omega'. \tag{4.35}$$

Since the single scattering spectrum of a medium can be related to the imaginary part of the complex permittivity, the Kramer-Kronig relations can be used to retrieve energy dependence of the real permittivity [4].

4.3.1 Spectrum analysis

If one ignores the instrumental broadening, surface-mode scattering and the retardation effects, the single scattering spectrum is approached by the single scattering distribution, which in place can be obtained from the recorded energy loss spectrum by the Fourier log method. [4]

$$I_{1}(E) \approx S(E) = \frac{2N_{ZLP}t}{\pi a_{0}m_{0}v^{2}} \operatorname{Im} \left[\frac{-1}{\varepsilon(E)} \right] \int_{0}^{\beta} \frac{\theta d\theta}{\theta^{2} + \theta_{E}^{2}}$$

$$= \frac{N_{ZLP}t}{\pi a_{0}m_{0}v^{2}} \operatorname{Im} \left[\frac{-1}{\varepsilon(E)} \right] \ln \left[1 + \left(\frac{\beta}{\theta_{E}} \right)^{2} \right]$$

$$(4.36)$$

In this equation is $J^1(E)$ the single scattering distribution, S(E) the single scattering spectrum, N_{ZLP} the zero-loss intensity, t the sample thickness, v the velocity of the incoming electron, β the collection semi angle, α the angular divergence of the incoming beam, and θ_E the characteristic scattering angle for energy loss E. In this equation α is assumed small in comparison with β . If this is not the case, additional angular corrections are needed. Furthermore, θ_E is given by:

$$\theta_E = E/(\gamma m_0 v^2). \tag{4.37}$$

Furthermore, it should be noted that to retrieve Re $[1/\varepsilon(E)]$ from Im $[-1/\varepsilon(E)]$, equation (4.34) should be rewritten to [5]:

$$\operatorname{Re}\left[\frac{1}{\varepsilon(E)}\right] = 1 - \frac{2}{\pi} \mathcal{P} \int_0^\infty \operatorname{Im}\left[\frac{-1}{\varepsilon(E')}\right] \frac{E' dE'}{E'^2 - E^2}.$$
 (4.38)

QUESTIONs on my end Will we be working with J(E), and is there the need for a Fourier method, or is $J^1(E)$ provided? Are additional angular corrections needed? Where does the 1 come from in eq (4.38), is it due to the minus sign in the $\text{Im}[-1/\varepsilon]$?

4.3.2 Step 1: rescaling intensity

The first step of the K-K analysis is now to rewrite Eq. (4.36) to:

$$I_{1,ac}(E) = \frac{I_1(E)}{\ln\left[1 + \left(\frac{\beta}{\theta_E}\right)^2\right]} = \frac{N_{ZLP}t}{\pi a_0 m_0 v^2} \operatorname{Im}\left[\frac{-1}{\varepsilon(E)}\right]. \tag{4.39}$$

As θ_E scales linearly with E, see eq. (4.37), the intensity in on the left side of the equation above now relatively increases for high energy loss with respect to low energy loss. SOMETHING about aperture correction, is that relevant?

QUESTIONs on my end I assume β and v are known, and that we do not take a distribution for v?

4.3.3 Step 2: extrapolating

Since the upcoming integrals all extend to infinity, but the data acquisition is inherently up to a finite energy, the spectra need to be extrapolated. An often used form is AE^{-r} , where r=3 if you follow the Drude-model, or r can be deducted from experimental data.

4.3.4 Step 3: normalisation and retrieving $\operatorname{Im}\left[\frac{1}{\varepsilon(E)}\right]$

Taking E' = 0 in (4.38), one obtains:

$$1 - \operatorname{Re}\left[\frac{1}{\varepsilon(0)}\right] = \frac{2}{\pi} \int_0^\infty \operatorname{Im}\left[\frac{-1}{\varepsilon(E)}\right] \frac{dE}{E}.$$
 (4.40)

Now dividing both sides of Eq. (4.41) by the energy, and subsequently integrating them over energy results in a comparable integral:

$$\int_0^\infty I_{1,ac}(E) \frac{dE}{E} = \frac{N_{ZLP}t}{\pi a_0 m_0 v^2} \int_0^\infty \operatorname{Im} \left[\frac{-1}{\varepsilon(E)} \right] \frac{dE}{E}.$$
 (4.41)

Combining the two leads to:

$$\frac{\int_0^\infty I_{1,ac}(E) \frac{dE}{E}}{\frac{\pi}{2} (1 - \operatorname{Re} \left[\frac{1}{\varepsilon(0)} \right])} = \frac{N_{ZLP}t}{\pi a_0 m_0 v^2} \equiv K,$$
(4.42)

in which K is the proportionality constant, used to estimate the absolute thickness if the zero-loss integral and the indicent energy are known. This formula requires $\operatorname{Re}\left[\frac{1}{\varepsilon(0)}\right]$ to be known, as is the case in for example metals $\left(\operatorname{Re}\left[\frac{1}{\varepsilon_{metal}(0)}\right]\approx 0\right)$. If this is not the case, other options to estimate K will be discussed later on.

This value of K, which is constant over E, can than in turn be used to retrieve the function of $\operatorname{Im}\left[-\frac{1}{\varepsilon(E)}\right]$ from the observed single scattering energy distribution $J^1(E)$ with eq. (4.41).

need to add other estimations of K?

4.3.5 Step 4: retrieving $\operatorname{Re}\left[\frac{1}{\varepsilon(E)}\right]$

Having retrieved Im $\left[-\frac{1}{\varepsilon(E)}\right]$ from the steps above, one can now use eq. (4.38) to obtain Re $\left[\frac{1}{\varepsilon(E)}\right]$, where one must pay attention to avoid including E=E' in the discrete integral over the spectrum, as this is a singularity. To avoid this singularity in a discrete signal, a couple of approaches are possible:

- In the integral (for discrete signals: summation) in eq. (4.38), simply exclude the E=E' values.
- Shift the values of Re $\left[\frac{1}{\varepsilon(E)}\right]$ to values at $E_i'' = (E_i + E_{i+1})$, to make sure to avoid E'' = E' in the summation.
- The dielectric function in the energy domain relate to the dielectric response function $1/\varepsilon(t) \delta(t)$ through:

$$\operatorname{Re}\left[\frac{1}{\varepsilon(E)}\right] = \mathcal{C}\left\{\frac{1}{\varepsilon(t)} - \delta(t)\right\} = \mathcal{F}\{p(t)\},\tag{4.43}$$

and

$$\operatorname{Im}\left[\frac{-1}{\varepsilon(E)}\right] = \mathcal{S}\left\{\frac{1}{\varepsilon(t)} - \delta(t)\right\} = i\mathcal{F}\{q(t)\},\tag{4.44}$$

where p(t) and q(t) are the even and odd parts respectively of the dielectric response function, and C and S are the cosine and sine Fourier transforms respectively. Since the dielectric response function is a response function and therefor causal, it is 0 for t < 0. This results in:

$$p(t) = \operatorname{sgn}[q(t)]. \tag{4.45}$$

Combining all this means that one can also obtain $\operatorname{Re}\left[\frac{1}{\varepsilon(E)}\right]$ from $\operatorname{Im}\left[-\frac{1}{\varepsilon(E)}\right]$ by:

$$\operatorname{Re}\left[\frac{1}{\varepsilon(E)}\right] = \mathcal{C}\left\{\operatorname{sgn}\left[\mathcal{S}^{-1}\left\{\operatorname{Im}\left[\frac{-1}{\varepsilon(E)}\right]\right\}\right]\right\}. \tag{4.46}$$

4.3.6 Step 5: retrieving ε

The dielectric function can subsequently be obtained from:

$$\varepsilon(E) = \varepsilon_1(E) + i\varepsilon_2(E) = \frac{\operatorname{Re}[1/\varepsilon(E)] + i\operatorname{Im}[-1/\varepsilon(E)]}{\{\operatorname{Re}[1/\varepsilon(E)]\}^2 + \{\operatorname{Im}[-1/\varepsilon(E)]\}^2}.$$
(4.47)

References

- [1] R. F. Egerton, "Electron energy-loss spectroscopy in the tem," Reports on Progress in Physics, vol. 72, no. 1, p. 016502, 2008. DOI: 10.1088/0034-4885/72/1/016502.
- [2] Lau, "Amazing article,"
- [3] (2020). "Kramers-kronig relations," [Online]. Available: https://en.wikipedia.org/wiki/KramersKronig_relations.
- [4] R. F. Egerton, <u>Electron energy-loss spectroscopy in the electron microscope</u>. Springer, 2011.
- [5] M. Dapor, "Appendix g: The kramers-kronig relations and the sum rules," in <u>Transport of Energetic</u> Cham: Springer International Publishing, 2017, pp. 175–178, ISBN: 978-3-319-47492-2. DOI: 10.1007/978-3-319-47492-2_16. [Online]. Available: https://doi.org/10.1007/978-3-319-47492-2_16.