

20-nm-C-250-nm-ZnO-Silica-15kV-reflin-2e-1

John Minter

Started 2018-08-07, Last modified: 2018-08-11

Abstract

The objective was to measure the time required for this simulation using the **Zn L3-M4** transition to reach the REFLIN criterion of an uncertainty of $2.0e^{-1}$. This was a fairly weak transition (0.1140) compared to the stronger **Zn L3-M5** transition (1.0). Still, I want to see how the simulation scales with the value of the uncertainty criterion.

Contents

Set up for the analysis	1
Get the current simulation time	2
Retrieve the number of showers	2
Compute the total maximum intensity for each element	3
Retrieve the data	3
Compute the maximum total intensity for each element	4
Process the spectra data	5
Load the raw data	5
Plot the spectrum.	6
Write a spectrum file in MSA format	7

Set up for the analysis

Note: This simulation used the same MSIMPA parameters for all layers and a REFLIN parameter (2.E-1) to finish.

```
RSEED -10 1 [Seeds of the random-number generator]
REFLIN 30040800 1 2.E-1 [Zn L3-M4 IZ*1e6+S1*1e4+S2*1e2,dete,tol.]
NSIMSH 1.0e8 [Desired number of simulated showers]
TIME 1.0e6 [Allotted simulation time, in sec]
```

First load the libraries we need...

```
library(rpenepma)
library(dplyr)
library(pander)
panderOptions('table.split.table', Inf)
library(ggplot2)
```

Next, set the paths and constants that we need...

```
moved_data <- TRUE # set to TRUE when simu is done and .rda with data is moved...
e0 <- 15 # kV
int_lo_lim <- 1.0e-1
int_hi_lim <- 1.0e4
reflin_crit <- 2e-1
```

```

ori_sim_dir <- "C:/Users/jrminter/Documents/work/penepma16/20-nm-C-250-nm-ZnO-Silica-reflin2"
sim_dir      <- "/Users/jrminter/Desktop/test-plt"
sim_nam      <- "20-nm-C-250-nm-ZnO-Silica-reflin-2e-1"
res_fi       <- "./penepma-res.dat"
int_fi       <- "./pe-intens-01.dat"
spc_fi       <- "./pe-spect-01.dat"
out_ti       <- sprintf("%s-%gkV", sim_nam, e0)
msa_fi       <- sprintf("./%s-%gkV.msa", sim_nam, e0)

```

Get the current simulation time

```

if (moved_data==FALSE){
  # When we have not moved the sim measure the simulation time.
  # Note copying the files to another directory seems to change the time
  # Test a reload.
  delta_t <- penepma_measure_simulation_time(ori_sim_dir)
  # print(delta_t)
  save(delta_t, file="./delta_t.rda")
  rm(delta_t)
  load("./delta_t.rda")
  msg <- "Analyzed from simulation directory."
} else {
  # When we have moved the simulation, reload the delta_t.rda
  msg <- "Analyzed from data stored in delta_t.rda."
  load("./delta_t.rda")
}
print(msg)

```

```
[1] "Analyzed from data stored in delta_t.rda."
```

```
delta_t_time <- print(delta_t)
```

Time difference of 9.680407 hours

Sort summary: this is very short and noisy...

Retrieve the number of showers

We want retrieve the number of showers (trajectories) that our penepma16 simulation calculated.

```

options(scipen = -6) # force printing in exponential format
num_showers <- penepma_get_number_of_showers(res_fi)
cat(num_showers)

```

```
2.97344e+05
```

```
options(scipen = 3) # reset to default
```

Compute the total maximum intensity for each element

Retrieve the data

First retrieve the intensity data as a tibble and print a preview.

```
tib <- penepma_get_intensities(int-fi)
print(tib)
```

```
# A tibble: 23 x 14
  IZ S0 S1 keV P.mu P.se C.mu C.se B.mu
<int> <chr> <chr> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
1 6 K L2 0.277 8.85e-7 7.73e-7 1.70e-9 5.09e-9 2.66e-10
2 6 K L3 0.277 1.35e-6 9.59e-7 1.70e-9 5.09e-9 7.99e-10
3 8 K L2 0.525 2.11e-5 1.95e-5 3.57e-8 5.37e-8 8.52e- 9
4 8 K L3 0.525 3.06e-5 1.96e-5 5.60e-8 2.93e-8 9.32e- 9
5 30 L3 M1 0.884 2.77e-5 2.07e-6 2.72e-8 2.04e-8 2.09e- 8
6 30 L2 M1 0.906 1.40e-5 1.46e-6 1.70e-9 5.09e-9 1.15e- 8
7 30 L3 M2 0.932 6.79e-8 1.02e-7 0. 0. 0.
8 30 L3 M3 0.934 5.09e-8 8.82e-8 0. 0. 0.
9 30 L2 M3 0.957 6.79e-8 1.02e-7 0. 0. 0.
10 30 L3 M5 1.01 3.68e-5 2.37e-6 4.25e-8 2.55e-8 2.89e- 8
# ... with 13 more rows, and 5 more variables: B.se <dbl>, TF.mu <dbl>,
# TF.se <dbl>, Int.mu <dbl>, Int.se <dbl>
```

What features did we measure?

```
print(names(tib))
```

```
[1] "IZ" "S0" "S1" "keV" "P.mu" "P.se" "C.mu"
[8] "C.se" "B.mu" "B.se" "TF.mu" "TF.se" "Int.mu" "Int.se"
```

How many transitions did we measure?

```
print(nrow(tib))
```

```
[1] 23
```

Measure our test transition ("Zn L3-M4")

```
val <- penepma_get_total_intensity_z_transition(tib, 30, "L3", "M4")
print(val)
```

```
# A tibble: 1 x 7
  Symbol IZ S0 S1 Int.mu Int.se Int.snr
<chr> <int> <chr> <chr> <dbl> <dbl> <dbl>
1 Zn 30 L3 M4 0.00000383 0.000000764 5.01
```

And see how close we are to the REFLIN criterion

```
cur_ref_crit <- val$Int.se/val$Int.mu

rv <- c(cur_ref_crit, 100*reflin_crit/cur_ref_crit)
names(rv) <-c("value", "target %")
print(rv)
```

```
value target %
0.1995467 100.2271728
```

Let's estimate the remaining time

```
pct <- rv[2]/100.
names(pct) <- c()
tr <- delta_t*(1.0-pct)
print(tr)
```

Time difference of -0.02199125 hours

Let's store a .rda file with REFLIN info

```
tmp_v <- data.frame(as.character(sim_nam),
                    as.numeric(reflin_crit), as.character.POSIXt(delta_t_time),
                    as.numeric(num_showers), as.numeric(cur_ref_crit),
                    as.numeric(100*reflin_crit/cur_ref_crit))
names(tmp_v) <- c("name", "reflin", "time", "showers",
                  "value", "target %")

tmp_v <- as_tibble(tmp_v)
tmp_v$name <- as.character(tmp_v$name)
C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1_ro <- tmp_v
rm(tmp_v)

save(C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1_ro,
     file="./C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1_ro.rda")
rm(C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1_ro)
load("./C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1_ro.rda")
pander(C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1_ro)
```

name	reflin	time	showers	value	target %
20-nm-C-250-nm-ZnO-Silica-reflin-2e-1	0.2	9.680407 hours	297344	0.1995	100.2

Compute the maximum total intensity for each element

1. Compute the maximum total intensity for each element.
2. Stack each row of data together into a tibble.
3. Prepend the sample ID to the data.
4. Print a well-formatted a table using **pander**.
5. Save the final tibble so it can be r-loaded later

```
# 1
C_t <- penepma_get_max_total_intensity_z(tib, 6)
O_t <- penepma_get_max_total_intensity_z(tib, 8)
Zn_t <- penepma_get_max_total_intensity_z(tib, 30)
Si_t <- penepma_get_max_total_intensity_z(tib, 14)

# 2
tot_int <- dplyr::bind_rows(C_t, O_t, Zn_t, Si_t)

#3
C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1 <- prepend_sample_id_max_int_tib(tot_int, out_ti)

# 4
pander(C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1)
```

Sample.ID	Symbol	IZ	S0	S1	Int.mu	Int.se	Int.snr
20-nm-C-250-nm-ZnO-Silica-reflin-2e-1-15kV	C	6	K	L3	0.000001351	0.000000096	1.408
20-nm-C-250-nm-ZnO-Silica-reflin-2e-1-15kV	O	8	K	L3	0.00003065	0.0000196	1.564
20-nm-C-250-nm-ZnO-Silica-reflin-2e-1-15kV	Zn	30	L3	M5	0.00003687	0.00000237	15.56
20-nm-C-250-nm-ZnO-Silica-reflin-2e-1-15kV	Si	14	K	L3	0.00003387	0.0000337	1.005

```
# 5
save(C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1,
      file="C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1.rda")
# delete and reload be sure it worked...
rm(C_20_nm_ZnO_250_nm_SiO2_15kV_refli2n)
load("./C_20_nm_ZnO_250_nm_SiO2_15kV_reflin_2e_m1.rda")
```

Process the spectra data

Load the raw data

```
tib <- penepma_read_raw_data(spc-fi,min_intensity_clip=5.0e-10)
rownames(df) <- c()
```

Examine the start:

```
pander(head(tib))
```

keV	mu	se
0.01183	0.5	0.5
0.01549	0.5	0.5
0.01915	0.5	0.5
0.02281	0.5	0.5
0.02647	0.5	0.5
0.03013	0.5	0.5

Examine the end:

```
pander(tail(tib))
```

keV	mu	se
14.98	0.5	0.5
14.98	0.5	0.5
14.99	0.5	0.5
14.99	0.5	0.5
14.99	0.5	0.5
15	0.5	0.5

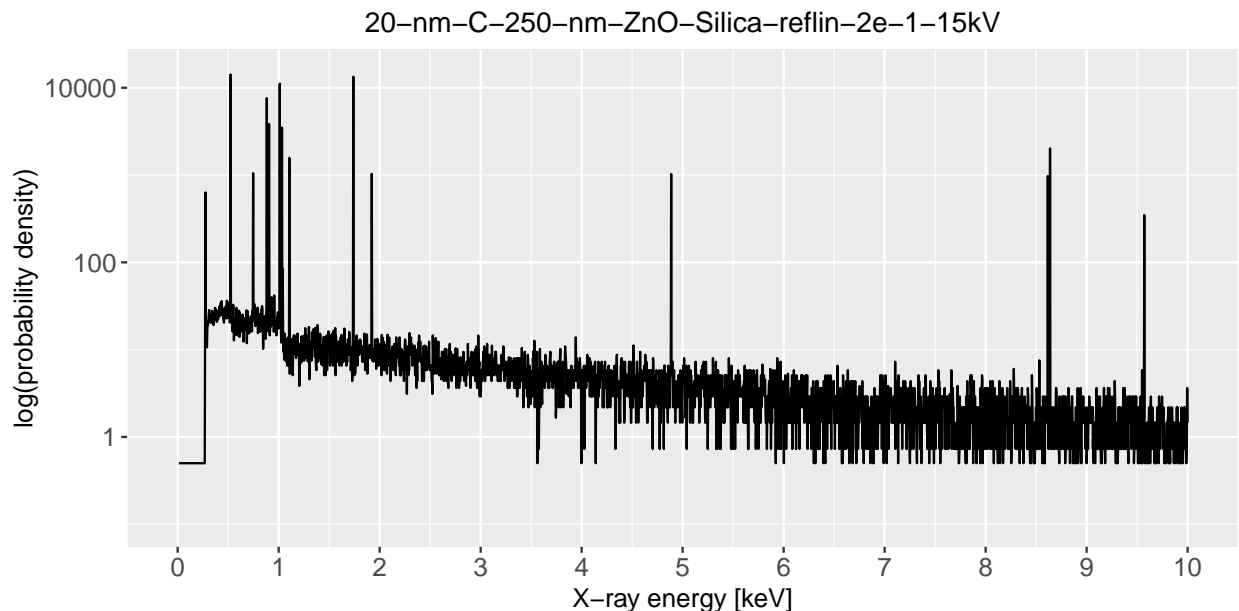
Plot the spectrum.

There is a very large dynamic range for both the **probability density** and the **uncertainty**. Penepma sets a lower limit for data at **1.0e-35**. Missing values are set to zero. We want to remove values from the dataframe that are below a useful limit. We do this below and plot a copy of the dataframe that is limited to the useful values.

```
m_t <- max(pretty(tib$mu))

plt <- ggplot(tib, aes(x = keV, y = mu)) +
  geom_line() +
  scale_x_continuous(breaks = seq(from = 0, to = e0-5, by = 1),
                    limits = c(0,e0-5)) +
  scale_y_log10(limits = c(int_lo_lim, m_t)) +
  xlab(label="X-ray energy [keV]") +
  ylab(label="log(probability density)" +
        # (1/(eV*sr*electron)) +
        ggtitle(out_ti) +
        theme(axis.text=element_text(size=12),
              axis.title=element_text(size=12),
              # center the title
              plot.title = element_text(hjust = 0.5))

print(plt)
```



And on a linear intensity scale...

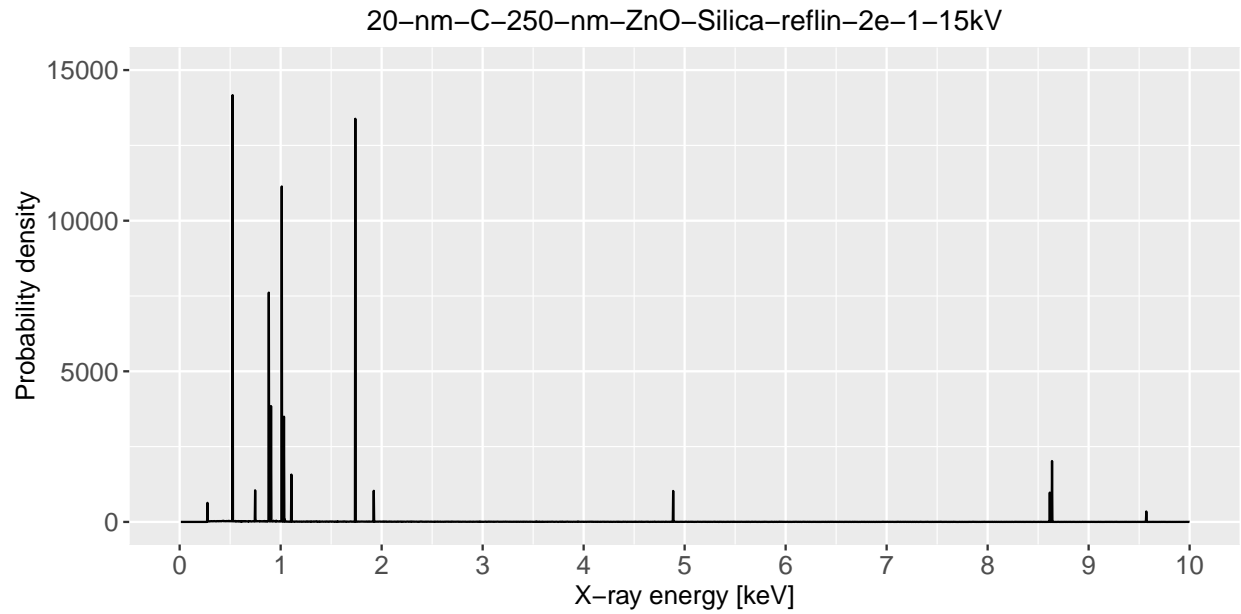
```
plt <- ggplot(tib, aes(x = keV, y = mu)) +
  geom_line() +
  scale_x_continuous(breaks = seq(from = 0, to = e0-5, by = 1),
                    limits = c(0,e0-5)) +
  scale_y_continuous(limits = c(int_lo_lim, m_t)) +
  xlab(label="X-ray energy [keV]") +
  ylab(label="Probability density" +
        # (1/(eV*sr*electron)) +
```

```

ggtitle(out_ti) +
theme(axis.text=element_text(size=12),
      axis.title=element_text(size=12),
      # center the title
      plot.title = element_text(hjust = 0.5))

print(plt)

```



Write a spectrum file in MSA format

```

penepma_to_msa(spc-fi, msa-fi,e0, out-ti)

```