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Chapter 1

Introduction

The utilities to SHADOW are a collection of programs written to perform a specific task on either the input or output. They are designed to give the user more information about the results in either tabular or graphic form. Currently the graphics package used is TopDrawer. SHADOW utilities are run through a shell program. Once inside the shell, the utilities are divided into six categories. They are:

- Preprocessors
- Postprocessors
- Macros
- Optical properties
- Graphics
- General

When the shell is invoked, a menu comes up with the first level of options. The up and down arrow keys (↑, ↓) move the cursor to the desired option within a list, <RETURN> moves to the list. Selecting an option moves one level down to the list of utilities that fall in that category. Some utilities may fall in more than one category, in which case they will be found in both. Each category will list the utility followed by a one line description of what the program does. At this point the user can either run the utility (R), get help on the utility (H), quit (Q), or return to the main menu (<RETURN>).
Chapter 2

Preprocessors

This section of the Utilities Manual will cover those utilities that need to be run before SHADOW. The programs generate files, some of them unformatted, that are used as input to SHADOW in various places such as source optimization, crystal diffraction, reflectivities of coatings and/or multilayers, and surface distortions. In this way SHADOW can trace more specific systems with more realistic results.
**Purpose:** Create a file which can be used by SHADOW in the computation of the reflectivity of a single crystal.

**Input:** Interactive from terminal.

**Output:** Formatted files (user specified).

**Description:**

In order to ray trace an optical system containing a crystal, SHADOW needs some preliminary data. These data are prepared by BRAGG. SHADOW can handle the so-called symmetric Bragg case where the crystal planes are parallel to the surface, as well as the asymmetric case. Within the context of the symmetric case, the local reflectivity calculated by the Darwin-Prins formalism is:

\[
R = \left| \sqrt{F_H/F_{\tilde{H}}} \left[ \eta \pm \sqrt{\eta^2 - 1} \right] \right|^2
\]  

(2.1)

where,

\[
\eta = \frac{-\Delta \theta \sin(2\theta_B) + \Gamma F_0}{|P| \Gamma (F_HF_{\tilde{H}})}
\]  

(2.2)

\(\theta\) is the glancing angle, \(\theta_B\) the Bragg angle, \(\Gamma\) a constant, \(P\) a polarization factor, and \(F_H\) the structure factor relative to the scattering vector \(H\) (\(F_{\tilde{H}}\) is in the \(\tilde{H}\) direction).

The structure factor \(F_H\) needs more explanation.

\[
F_H = \sum_{j=1}^{H} f_j e^{i\tilde{r}_j \cdot \tilde{H}}
\]  

(2.3)

It is the cell structure factor, and as such depends on:

1. \(\tilde{H}\), the scattering vector
2. \(\tilde{r}_j\), the location of atoms within a unit cell in units of \(a\) (lattice count).
3. \(f\), the atomic scattering factors of the individual atom.

\(H\) is simply the vector defined by \((H,K,L)\), where \(H\), \(K\), and \(L\) are the indices of the crystal plane.

Four types of crystal structure are supported. Let say there are atom A and atom B making up the basis within each cubic unit cell, then the locations of each atom for the crystal
structures are:

<table>
<thead>
<tr>
<th>Structure</th>
<th>A</th>
<th>B</th>
<th>Lattice</th>
</tr>
</thead>
<tbody>
<tr>
<td>ZincBlende</td>
<td>(0,0,0)</td>
<td>(1/4,1/4,1/4)</td>
<td>fcc</td>
</tr>
<tr>
<td>Rocksalt</td>
<td>(0,0,0)</td>
<td>(1/2,1/2,1/2)</td>
<td>fcc</td>
</tr>
<tr>
<td>Simple fcc</td>
<td>(0,0,0)</td>
<td></td>
<td>fcc</td>
</tr>
<tr>
<td>CsCl</td>
<td>(0,0,0)</td>
<td>(1/2,1/2,1/2)</td>
<td>cubic</td>
</tr>
</tbody>
</table>

Of course atom A can be the same as atom B. In that case, the ZincBlende reduces to the diamond structure; rocksalt to simple cubic; and CsCl to simple bcc. All together they cover the geometry of most commonly found crystals.

The scattering factor $f$ depends on the atomic properties. This quantity is complex, and is written as:

$$ f = f_0 + f' + i f'' \tag{2.4} $$

$f_0$ is the non-dispersive portion of the real part, while $f'$ and $f''$ are the dispersive part. $f'$ and $f''$ are functions of photon energy for each element, and they are readily obtained from our optical constant library (see PREREFL about the library). On the other hand, we don’t have a database for $f_0$ and it must be supplied by the user. $f_0$ is a function of the perpendicular component of the photon momentum, i.e., $\left(\frac{2\pi}{\lambda}\sin \theta\right)$. At the Bragg peak, $k_\perp$ is exactly $\frac{\pi}{\lambda}$. Since we are working near the Bragg peak, BRAGG asks the user to enter the values of $f_0 \left(\frac{\sin \theta}{\lambda}\right)$ at the neighborhood where $\sin \theta$ is approximately $\frac{1}{2d}$. Three points at that neighborhood are required in order to do a quadratic fitting.

A good listing of all $f_0$ values is found in the International Tables of X-Ray Crystallography, Vol. III, sect.2.2, table 2.2A, p.72. A shorter version, adapted from X-ray Diffraction, by B.E.Warren, Addison-Wesley, is listed in the SHADOW PRIMER, sect.6.4, p.84.

The example we use here is GaAs(111) crystal. It is the same one as in SHADOW PRIMER. Here we concentrate on the BRAGG part of the modelling, while the user can refer to PRIMER for the SHADOW part. A description on how to run a two crystals case is given there. GaAs has ZincBlende structure with $a = 5.65$ Å. Note two capitalized letters must be used for each element (Gallium as GA, Tungsten as WW). Then the user is asked to input (3) values of $f_0$ for each atom. Since the program now knows the 2d spacing, it will type out the value of $\frac{\sin \theta}{\lambda}$ near the Bragg peak in order to assist the user in selecting the correct $f_0 \left(\frac{\sin \theta}{\lambda}\right)$. After that, $f'$ and $f''$ are simply defined by the photon energy range. A Debye-Waller factor $\exp^{-M}$ (between 0 and 1) is introduced to account for thermal vibrations. To neglect the finite temperature effect, enter the value of 1 for the Debye-Waller factor. At this point, all the parameters are defined for SHADOW and they are stored in
a formatted file (GAAS.PAR). Rocking curves can also be generated by BRAGG for the user’s convenience.

As an example, we pick 11160 eV. BRAGG outputs some useful data. Note \( \theta \) (graz) is almost the same as Bragg angle, the difference being that the latter is corrected for refraction of the radiation at the crystal surface. The last two items printed are Real(Ss) and Real(Sp), they are the widths of the rocking curves in the two polarizations (s-perpendicular, p-parallel to the plane of incidence). Thus the user can choose the angular range of the rocking curves he wants to generate (60 microradian cover more than twice the width).

These data are stored in ROCK_CURVE.PAR, while the rocking curves themselves are stored in ROCK_CURVE.S and ROCK_CURVE.P. Graphs of both files are shown below.

The file GAAS.SHA is the input to SHADOW. It is formatted so that the user may edit the data for \( f' \) and \( f'' \). The data format is explained in the example. However, we strongly advise not to edit the file unless the data have been determined to be inaccurate e.g. for the description near an absorption edge.

**Examples:**

The first example will be for a symmetrical case, after which we will run the asymmetrical case.

```$```
```
$ BRAGG

All crystal structures are refered to a cubic unit cell.

Bravais lattice type :
0 for ZincBlende
1 for Rocksalt
2 for simple FCC
3 for CsCl structure

Then ?
0 /* select ZincBlende */

Lattice constant (Aangs) ?
5.65 /* lattice constant */

Index of crystal plane of reflection H,K,L :
1,1,1 /* indices */

The ZINCBLENDE structure is defined by atom A located at (0,0,0) and atom B at (1/4,1/4,1/4) of the fcc lattice.

Enter 2-letters (capitalized) atomic symbol for atom A :
```
Enter 2-letters (capitalized) atomic symbol for atom B :
AS  \* Arsenic *

Atomic scattering factor is defined by \( f_0 + f' + if'' \), where
\( f_0 = f_0(SIN(\theta)/\Lambda) \) is the non-dispersive part
\( f', f'' \) (\( \Lambda \)) are the dispersive part.

We need \( f_0 \) at 3 different values of \( SIN(\theta)/\Lambda \), which
should cover the range of interest and center around :
\( SIN(\theta)/\Lambda = 0.1532788325282192 \) ratio.

Please enter 1) \( SIN(\theta)/\Lambda \), 2) \( f_0 \),
For atom A, first set :  \* Ga *
\[ 0.14, 27.162 \]  \* The International Tables for X-ray
, second set :  \* Crystallography lists \( f_0 \) as a
\[ 0.15, 26.783 \]  \* function of \( SIN(\theta)/\Lambda \) for
, third set :  \* each element and some ions.
\[ 0.16, 26.406 \]  
For atom B, first set :  \* As! Here the user needs to supply \( f_0 \)
\[ 0.14, 28.742 \]  \* centered around 0.15327..., and
, second set :  \* within some limits that SHADOW can
\[ 0.15, 28.307 \]  \* interpolate for \( f_0 \).
, third set :  
\[ 0.16, 27.877 \]  

\( f', f'' \) is furnished from optical constant library within ...
minimum photon energy (eV) :
\[ 11100 \]  \* lower range boundary *
maximum photon energy (eV) :
\[ 11200 \]  \* upper range boundary *
energy step (eV) :
10
Do you want to include crystal absorption [1/0] ?
1  \* always *
Temperature (Debye-Waller) factor :
0.979
Output file name (for SHADOW) :
GAAS.SHA  \* try to use a meaningful name *
Do you want to generate a rocking curve [1/0] ?
1 \* for examination *

... at what energy (eV) ? \* following only for rocking curve *
11160

So far, we are working with:
Lambda = 1.110978494623656 Angstroms
Theta (graz) = 9.804650841199757 degrees
Bragg angle = 9.807153296220226 degrees
Structure factor = (142.6426218594671,11.03752285669588)
refraction index = 1 - delta - i*beta :
delta = 7.3270650274570581E-06
beta = 4.8385681505652676E-07
Absorption coeff = 547.2944878508624 cm^{-1}

Do you want an asymmetrical cut [1/0] ?
0 \* symmetrical this time *
1/2 width of Rock Curve  s-pol = : 25.54182356671078 microradians
1/2 width of Rock Curve  p-pol = : 24.06047395262200 microradians

+/- how many microradians :
60 \* width of calculation *

How many points :
200
Do you want to use sec [1/0] ?
0
Do you want the R.C. not centered [1/0] ?
0
Do you want to try another energy [1/0] ?
0
$ DIR/SINCE

Directory CXRL$DUA0:[SHADOW.XRAYOP.DOCUMENT.UTIL.BRAGG]

GAAS.SHA;1 4 19-SEP-1989 13:13:12.77 (RWED,RWED,RE,)
ROCK_CURVE.P;1 20 19-SEP-1989 13:13:25.78 (RWED,RWED,RE,)
ROCK_CURVE.PAR;1 3 19-SEP-1989 13:14:57.71 (RWED,RWED,RE,)
ROCK_CURVE.S;1 20 19-SEP-1989 13:13:25.64 (RWED,RWED,RE,)

Total of 4 files, 47 blocks.
$
At the end of the file GAAS.SHA is a table of:
eV  f'(Ga)  f''(Ga)  f'(As)  f''(As)
from the optical constant library. The users can edit this part if they have more accurate
values, especially near an absorption edge.

\$TY\ GAAS.SHA

\begin{verbatim}
0 1562378464.990918 3.2620290209213856E-08
31 33 0.9790000000000000
(4.0000000000000000,-6.8654264713317982E-17)
(4.0000000000000000,6.8654264713317982E-17)
(8.4736075498395345E-17,-4.0000000000000000)
(8.4736075498395345E-17,4.0000000000000000)
32.678000000000011 -40.800000000000000 10.0000000000000364
35.356999999999986 -50.7499999999999727 24.9999999999999272

11
11100.000000000000 -1.895109680624937 3.414937245198015
-2.436850441775594 0.5607204097485073
11110.000000000000 -1.8801062553822577 3.410127243375178
-2.449320912049193 0.5598036778163483
11120.000000000000 -1.8651028301395772 3.405317241552341
-2.461791382322792 0.558869458841894
11130.000000000000 -1.850099404896898 3.400507239729505
-2.474261852596391 0.5579702139520304
11140.000000000000 -1.835095979654219 3.3956972379066668
-2.486732322689990 0.5570534820198715
11150.000000000000 -1.820092554411540 3.390887236083831
-2.499202793143589 0.5561367500877125
11160.000000000000 -1.805089129168860 3.386077234260994
-2.511673263417189 0.5552200181555536
11170.000000000000 -1.790085703926181 3.381267232438158
-2.52414373690788 0.5543032862233946
11180.000000000000 -1.775082278683501 3.376457230615321
-2.53661420364387 0.5533865542912357
11190.000000000000 -1.760078853440822 3.371647228792484
-2.549084674237986 0.5524698223590767
11200.000000000000 -1.745075428198142 3.366837226969648
-2.56155144511585 0.5515530904269178
\end{verbatim}
$ ! ROCK\_CURVE.PAR lists the parameters used to generate the rocking
$ ! curve at 11600 eV.
$ !
$ !
$ ! TY ROCK\_CURVE.PAR

ZincBlende structure:
For atom A, \( fo + f' + if'' = (24.8507850869210, 3.386077234260994) \)
\( B, = (25.65378599830614, 0.555220018155536) \)
Lattice constant = 5.650000000000000 Angstroms
d-spacing = 3.262029020921386 Angstroms
Photon energy = 11160.00000000000 eV
Lambda = 1.11097849623656 Angstroms
SIN(theta)/Lambda = 0.1532788325282192 ratio.
refraction index = 1 - delta - i*beta :
delta = 7.327065274570581E-06
beta = 4.8385681505652676E-07
Absorption coeff = 547.2944878508624 cm\(^{-1}\)
Temperature factor = 0.979000000000000
Structure factor F(000) = (238.7329504296558, 15.76518900966619)
Structure factor F(hkl) = (142.6426218594671, 11.0375285669588)
The width of the Rock Curve is
1/2 width for s-pol = : 25.54182356671078 microradians
1/2 width for p-pol = : 24.06047395262200 microradians
Theta (graz) = 9.804650841199757 degrees
Bragg angle (corr) = 9.807153296220226 degrees
$ !
$ ! TY ROCK\_CURVE.S

The following example was run for an asymmetrical\index{crystal, asymmetric}
cut silicon crystal\index{crystal} with the
cut at seven degrees between the face and Bragg planes. We run between 7700 and 7800 eV, and generate a rocking curve at 7750 eV. Notice for this case you get a rocking curve for the incidence and reflection at the crystal. The parameter file that acts as input to SHADOW does not specify symmetric or asymmetric. This information is input to SHADOW when tracing, so once a file is set up for a crystal, one does not have to rerun BRAGG for the other case of the same crystal unless new rocking curves are desired.

All crystal structures are refered to a cubic unit cell.

Bravais lattice type:
0 for ZincBlende
1 for Rocksalt
2 for simple FCC
3 for CsCl structure

Then?

Lattice constant (Angs)?
5.43

Index of crystal plane of reflection H,K,L:
1,1,1

The ZINCBLENDE structure is defined by atom A located at (0,0,0) and atom B at (1/4,1/4,1/4) of the fcc lattice.

Enter 2-letters (capitalized) atomic symbol for atom A:
SI
Enter 2-letters (capitalized) atomic symbol for atom B:
Atomic scattering factor is defined by \( f_0 + f' + if'' \), where
\( f_0 = f_0(\sin(\theta)/\Lambda) \) is the non-dispersive part
\( f', f'' \) (\( \Lambda \)) are the dispersive part.

We need \( f_0 \) at 3 different values of \( \sin(\theta)/\Lambda \), which should cover the range of interest and center around:
\( \sin(\theta)/\Lambda = 0.1594890246380182 \) ratio.

Please enter 1) \( \sin(\theta)/\Lambda \), 2) \( f_0 \),

Figure 2.1: BRAGG: Rocking curve for S- and P- polarization
For atom A, first set:
0.1, 12.16
, second set:
0.2, 9.67
, third set:
0.3, 8.22
For atom B, first set:
0.1, 12.16
, second set:
0.2, 9.67
, third set:
0.3, 8.22

f', f" is furnished from optical constant library within ...
minimum photon energy (eV): 7700
maximum photon energy (eV): 7800
energy step (eV): 10
Do you want to include crystal absorption [1/0]? 1
Temperature (Debye-Waller) factor:
1.0
Output file name (for SHADOW):
SI.PAR
Do you want to generate a rocking curve [1/0]? 1
... at what energy (eV)? 7750

So far, we are working with:
Lambda = 1.599809032258065 Angstroms
Theta (graz) = 14.78259099147192 degrees
Bragg angle = 14.78449376730854 degrees
Structure factor = (61.29837293588675, 1.976536272774998)
refraction index = 1 - delta - i*beta:
delta = 8.1919893381993125E-06
beta = 2.0040488614427989E-07
Absorption coeff = 157.4164179247638 cm⁻¹
Do you want an asymmetrical cut [1/0] ?
1
Angle (deg) between face and bragg planes (CW) =
7
The width of the Rock Curve in function of incident angle is
1/2 width for s-pol = : 10.76098572573549 microradians
1/2 width for p-pol = : 9.359850847544060 microradians
The width of the Rock Curve in function of reflected angle is
1/2 width for s-pol = : 29.48906494122322 microradians
1/2 width for p-pol = : 25.64943923522630 microradians
Asymmetric factor b= : 2.740368372638810

+/- how many microradians:
80
How many points:
150
Do you want to use sec [1/0] ?
1
Do you want the R.C. not centered [1/0] ?
1
Do you want to try another energy [1/0] ?
0

$ TY SI.PAR

0  1760078747.302978  3.1350119616996679E-08
14   14 1.0000000000000000
(4.0000000000000000, -6.8654264713317982E-17)
(4.0000000000000000,  6.8654264713317982E-17)
(8.4736075498395345E-17, -4.0000000000000000)
(8.4736075498395345E-17,  4.0000000000000000)
15.690000000000000  -40.500000000000000  51.999999999999999
15.690000000000000  -40.500000000000000  51.999999999999999
11
7700.0000000000000  0.2846972178807969  0.3536787673872427
0.2846972178807969  0.3536787673872427
7710.0000000000000  0.2843014434140994  0.3528241239968134
0.2843014434140994  0.3528241239968134
7720.0000000000000  0.2839056689474015  0.3519694806063841
0.2839056689474015  0.3519694806063841
TY ROCK_CURVE.PAR

ZincBlende structure:

For atom A, \( f_o + f' + i f'' = (10.83612379466686, 0.3494055504350961) \)

B, \( = (10.83612379466686, 0.3494055504350961) \)

Lattice constant = 5.430000000000000 Angstroms

d-spacing = 3.13501196169968 Angstroms

Photon energy = 7750.000000000000 eV

\( \Lambda = 1.599809032258065 \) Angstroms

\( \sin(\theta)/\Lambda = 0.1594890246380182 \) ratio.

Refractive index = 1 - \( \delta - i \beta \):

\( \delta = 8.191989338193125E-06 \)

\( \beta = 2.0040488614427989E-07 \)

Absorption coeff = 157.4164179247638 cm\(^{-1}\)

Temperature factor = 1.000000000000000

Structure factor \( F(000) = (114.2617467643785, 2.795244403480769) \)

Structure factor \( F(hkl) = (61.29837293588675, 1.976536272774998) \)

Width of Rock Curve in function of incident angle:

1/2 width for s-pol = : 10.76098572573549 microrad

1/2 width for p-pol = : 9.359850847544060 microrad

Incident Bragg angle = 21.78259099147192 degrees

Incident corrected angle = 21.78388955458310 degrees

Width of Rock Curve in function of reflected angle:

1/2 width for s-pol = : 29.48906494122322 microrad
Figure 2.2: BRAGG: Asymmetric Crystal cut

1/2 width for p-pol = : 25.64943923522630 microrad
Reflected Bragg angle = 7.782590991471924 degrees
Reflected corrected angle = 7.786149532751679 degrees
Asymmetric factor b= : 2.740368372638810
Angle between face and bragg planes (CW)= 7.000000000000000 degrees

The following plot shows the rocking curves for incident and reflected light. The S-polarization is denoted by a solid line, while the P-polarization is simply plotted points. We show the pertinent information when running TRACE.
$ go trace

Ray Tracing Selected. Begin procedure.

Are we dealing with a crystal [ Y/N ] ?
1
File containing crystal parameters ?
si.par
Is the crystal asymmetric [ Y/N ] ?
1
Enter the angle formed by the crystal planes with the crystal surface. Use a positive value if the plane normal lies along the +Y direction (focusing), negative otherwise.
Planes angle [ deg ] ?
7
Automatic Tuning of Crystal [ Y/N ] ?
1
You must then supply the wavelength (or photon energy) at which you want the crystal tuned.
Energy, in eV, [ 0 ] or wavelength, in Angs., [ 1 ] ?
0
Photon Energy ?
7750
Purpose: To set up an ERG /Grasshopper type monochromator at a user specified photon energy, given an existing system.

Input: A set of STARTxx files describing the optical system at zero order. The user will also need to input the following parameters:
- $M_0$–exit slit distance
- $M_1$–codling slit distance
- codling slit incidence angle
- grating radius
- grating incidence angle
- lines per cm
- photon energy or wavelength the calculations are desired at the maximum slit-to-slit distance.

Output: A set of STARTxx files describing the optical system at a user specified photon energy. The user may give them different names from the original system. Also writes the system parameters to a text file or terminal (TT:) if the user chooses.

Description:

The ERG /Grasshopper monochromators are fairly complex to setup because of the correlated rotations of grating, codling slit, and condensing mirror. The user typically knows the beamline parameters at zero order. To avoid annoying calculations, this utility will set up the parameters for an ERG /Grasshopper type monochromator at a given photon energy, given the system at zero order. It will compute zero order slit distance, carriage movement, and codling slit tilt. It will then modify the system to include these parameters, “tuning” the monochromator to the specified wavelength or energy. Notice that no other parameters are modified (e.g. reflectivity switches, sizes, errors, etc.) so that the operation is indeed exactly equivalent to retuning the monochromator.

Examples:

Below is the output from SYSINFO on a 2m Grasshopper monochromator at zero order. The output from SYSPLOT for two and three dimensional plots follows.

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
************** SYSTEM DESCRIPTION **************
System at zero order
2m Grasshopper -- ERGSET Example

Input file specified:

Optical Element: Creation Time:
1 CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET]END.01;5 8-JAN-1990 15:38
2 CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET]END.02;5 8-JAN-1990 15:38
3 CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET]END.03;5 8-JAN-1990 15:39
4 CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET]END.04;5 8-JAN-1990 15:39
5 CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET]END.05;5 8-JAN-1990 15:39

Optical Element # 1 System Number:

MIRROR SPHERICAL DIM CHECK COMPUTED REFLEC. OFF

Orientation 270.0000000000000000 deg.
Source Plane 230.0000000000000000
Incidence Ang. 88.0000000000000000 deg.
Reflection Ang. 88.0000000000000000 deg.
Image Plane 108.4500000000000000

Optical Element # 2 System Number:

MIRROR ELLIPTICAL DIM CHECK COMPUTED REFLEC. OFF

Orientation 90.0000000000000002 deg.
Source Plane 108.4500000000000000
Incidence Ang. 88.5000000000000002 deg.
Reflection Ang. 88.5000000000000002 deg.
Image Plane 22.5000000000000000

Optical Element # 3 System Number:

MIRROR CODLING SLIT DIM CHECK COMPUTED REFLEC. OFF
Orientation  180.0000000000000 deg.
Source Plane  22.50000000000000
Incidence Ang.  88.00000000000000 deg.
Reflection Ang.  88.00000000000000 deg.
Image Plane  3.48950000000000

Optical Element #  4  System Number:

GRATING  SPHERICAL  DIM CHECK  EXTERNAL  REFLEC. OFF

Orientation  180.0000000000000 deg.
Source Plane  3.48950000000000
Incidence Ang.  88.00000000000000 deg.
Reflection Ang.  88.00000000000000 deg.
Image Plane  6.979899340500188

Optical Element #  5  System Number:

MIRROR  ELLIPTICAL  DIM CHECK  COMPUTED  REFLEC. OFF

Orientation  180.0000000000000 deg.
Source Plane  100.0000000000000
Incidence Ang.  88.00000000000000 deg.
Reflection Ang.  88.00000000000000 deg.
Image Plane  100.0000000000000

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

OPT. Elem #  X =  Y =  Z =

0  0.000000000000000E+00  0.000000000000000E+00  0.000000000000000E+00
1  0.000000000000000E+00  230.00000000  0.000000000000000E+00
1’  7.5650895776  338.18582125  0.29010291601E-1
2  15.130179155  446.37164250  0.58020583202E-15
2’  16.697548841  468.78607328  1.1775590155
Three dimensional and two dimensional plots of system at zero order. Now run ERGSET on the zero order START files.

$ ERGSET
Definition of ERG at ZERO Order Position.
Distance M0-exit slit ? 238.9
M1-Codling slit distance ? 45
Codling Slit Incidence Angle ? 88
Grating Radius ? 200
Grating Incidence Angle ? 87.8
Lines per CM ? 12000
Enter [0] to use eV, [1] Angs. 0
Then ? 40
Order to be used ? -1
Maximum slit-to-slit distance : 611.5
Do you want to save the inputs [Y/N] ? y
File describing M1 ? START.02
Codling Slit ? START.03
Grating ? START.04

START.04

read OK.
Enter new file name for grating. To use the same, just type <ret>
Figure 2.3: Three dimensional before ERGSET
Figure 2.4: Two Dimensional before ERGSET—Side view
Figure 2.5: Two Dimensional before ERGSET – Top view
Then ? 40EV.04
START.03

read OK.
Enter new file name for Codling Slit. To use the same, just type <ret>
Then ? 40EV.03
Enter new file name for M1. To use the same, just type <ret>
Then ? 40EV.02
File name ? 40EV.PAR
$
$ type 40EV.PAR
238.96300000000000 ! Distance M0-exit slit
45.00000000000000 ! Codling slit distance
88.00000000000000 ! Codling Slit Incidence Angle
200.00000000000000 ! Grating Radius
88.00000000000000 ! Grating Incidence Angle
13.95979868100039 ! Zero order slit distance
47.19515409835923 ! Carriage movement
6.902390810838892 ! Codling slit tilt
start.02
! M1 file
start.03
! Codling Slit file
start.04
! Grating file

40EV.PAR contains the fixed parameters of the system that we entered when we ran ERGSET. It also contains the parameters it computed for our Grasshopper at 40 eV – the zero order slit distance, carriage movement, and codling slit tilt.
The output from SYSINFO for the new system is shown below.
### Optical Element: Creation Time:
- Input file specified:

```
+---------------------------------------------------------------------+
  # Optical Element: Creation Time:                                  
  1  CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET.40EV]END.01;1 29-DEC-1989 15:  
  2  CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET.40EV]END.02;1 29-DEC-1989 15:  
  3  CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET.40EV]END.03;1 29-DEC-1989 15:  
  4  CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET.40EV]END.04;1 29-DEC-1989 15:  
  5  CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.ERGSET.40EV]END.05;1 29-DEC-1989 15:  
+---------------------------------------------------------------------+
```

#### Optical Element # 1: System Number:
**MIRROR SPHERICAL**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orientation</td>
<td>270.0000000000000000</td>
<td>deg.</td>
</tr>
<tr>
<td>Source Plane</td>
<td>230.0000000000000000</td>
<td></td>
</tr>
<tr>
<td>Incidence Ang.</td>
<td>88.0000000000000000</td>
<td>deg.</td>
</tr>
<tr>
<td>Reflection Ang.</td>
<td>88.0000000000000000</td>
<td>deg.</td>
</tr>
<tr>
<td>Image Plane</td>
<td>108.45000000000000</td>
<td></td>
</tr>
</tbody>
</table>

#### Optical Element # 2: System Number:
**MIRROR ELLIPTICAL**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orientation</td>
<td>90.0000000000000000</td>
<td>deg.</td>
</tr>
<tr>
<td>Source Plane</td>
<td>61.25484590164077</td>
<td></td>
</tr>
<tr>
<td>Incidence Ang.</td>
<td>88.5000000000000000</td>
<td>deg.</td>
</tr>
<tr>
<td>Reflection Ang.</td>
<td>88.5000000000000000</td>
<td>deg.</td>
</tr>
<tr>
<td>Image Plane</td>
<td>25.00000000000000</td>
<td></td>
</tr>
</tbody>
</table>

#### Optical Element # 3: System Number:
**MIRROR CODLING SLIT**

<table>
<thead>
<tr>
<th>Dimension</th>
<th>Value</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orientation</td>
<td>180.0000000000000000</td>
<td>deg.</td>
</tr>
<tr>
<td>Source Plane</td>
<td>20.0000000000000000</td>
<td></td>
</tr>
<tr>
<td>Incidence Ang.</td>
<td>81.09760918916109</td>
<td>deg.</td>
</tr>
<tr>
<td>Reflection Ang.</td>
<td>81.09760918916109</td>
<td>deg.</td>
</tr>
</tbody>
</table>
### Optical Element # 4 System Number:

**Grating Spherical**

- **Orientation**: 180.0000000000000 deg.
- **Source Plane**: 3.489949670250097
- **Incidence Ang.**: 88.0000000000000 deg.
- **Reflection Ang.**: 74.19521837832219 deg.
- **Image Plane**: 54.47210960704609

### Optical Element # 5 System Number:

**Mirror Elliptical**

- **Orientation**: 180.0000000000000 deg.
- **Source Plane**: 100.0000000000000
- **Incidence Ang.**: 88.00000000000000 deg.
- **Reflection Ang.**: 88.00000000000000 deg.
- **Image Plane**: 100.0000000000000

### Optical System Configuration

**Laboratory Reference Frame**

<table>
<thead>
<tr>
<th>OPT. Elem #</th>
<th>X =</th>
<th>Y =</th>
<th>Z =</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.000000000000000E+00</td>
<td>0.000000000000000E+00</td>
<td>0.000000000000000E+00</td>
</tr>
<tr>
<td>1</td>
<td>0.000000000000000E+00</td>
<td>230.0000000000000</td>
<td>0.000000000000000E+00</td>
</tr>
<tr>
<td>1'</td>
<td>7.5650895776</td>
<td>338.18582125</td>
<td>0.29010291601E-15</td>
</tr>
<tr>
<td>2</td>
<td>11.838011627</td>
<td>399.29145343</td>
<td>0.45395915774E-15</td>
</tr>
<tr>
<td>2'</td>
<td>13.579533500</td>
<td>424.19637651</td>
<td>1.3083989061</td>
</tr>
<tr>
<td>3</td>
<td>14.972750999</td>
<td>444.12031498</td>
<td>2.3551180309</td>
</tr>
<tr>
<td>3'</td>
<td>15.208115658</td>
<td>447.48618642</td>
<td>1.4633436043</td>
</tr>
</tbody>
</table>
The new system is plotted below in two and three dimensions.
Figure 2.6: Three dimensional after tuning with ERGSET
Figure 2.7: Two dimensional side view after tuning with ERGSET
Figure 2.8: Two dimensional top view after tuning with ERGSET
HISTO3

Purpose: To create a file needed by SHADOW for optimized sources.
Input: A file generated by REFLAG or SHADOWIT that has modified the source by flagging the rays that will be lost in the system.
Output: A file that has partitioned the useful volume of phase space into bins. This file will serve as input to the SOURCE option of SHADOW and will be a guide to the 'optimization' of the source.

Description:

HISTO3 defines the volume of phase space in which the rays from the source will not be lost. The steps involved in this matching of the source and system are as follows:

1. Setup source, system and trace as usual.

2. If the number of rays lost in the system is significant, it may be advisable to “optimize” the source. Since SHADOW keeps track of all rays, it is in principle possible to work back toward the source and tag all the rays that are lost somewhere along the line. This can be accomplished with REFLAG (for a single file) or SHADOWIT (for multiple files).

3. The modified source ray file from the previous step is used as input to HISTO3 and is used to define the discrete chunks of phase space which are good or bad. The histograms are given the value 1 if one or more rays that were generated at the source have ‘good’ flags at the check point in the system. The histograms are created in the phase spaces of \((X,X')\), \((Y,Y')\), \((Z,Z')\). Thus the three coordinates are treated independently (which is not always their true behavior) since the phase space is six-dimensional.

4. The output from HISTO3 is then used as input to the source generation program by selecting the optimization. After the filename has been given to SOURCE, one can exit and let the matching begin. SOURCE will reject any rays that have been generated in the zero space of the histogram. The number of rays accepted and the total number are displayed at regular intervals.

Since HISTO3 separates good and bad phase space into discrete intervals, it is not guaranteed that all of the rays will be good after one run of the optimization procedure. The above routine may be repeated on the “optimized source” to increase the efficiency further,
but with diminishing returns. By increasing the number of bins, one achieves a finer resolu-
tion in phase space at the cost of additional memory and computing time.

**Examples:**

Here we will consider a typical synchrotron radiation source incident upon a toroidal mirror
with an exit slit at the continuation plane, where the image will be formed. The slit has been
defined such that it will block a substantial number of rays. Below are the descriptions of
the source and system.

```
+----------------------------------------------------------------+
| *************************************************************** |
| **SOURCE DESCRIPTION**                                        |
| Optimized source                                              |
| Optimized for exit slit (Star.01)                             |
| +----------------------------------------------------------------+
| Input file specified:                                         |
| end.00                                                        |
| Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.HISTO3]END.00;2|
| Creation Date :10-APR-1989 16:05                             |
| +----------------------------------------------------------------+
| Random Source.                                               |
| Generated total 1000 rays.                                   |
| Source assumed TRIDIMENSIONAL.                                |
| Source Spatial Characteristics: GAUSSIAN                      |
| Sigma X : 0.100000000  Sigma Z : 0.100000000E-01             |
| Depth: SYNCHROTRON SOURCE.                                    |
| +----------------------------------------------------------------+
| Source Emission Characteristics                              |
| Distribution Type: SYNCHROTRON                               |
| Distribution Limits. +X : 0.100000000E-01 -X: 0.100000000E-01 rad|
| +Z : 0.500000000E-02 -Z: 0.500000000E-02 rad                  |
| Magnetic Radius = 4.81000000 m.  Beam Energy = 1.5000 GeV.  |
| Beam Emittances. EPSI_X: 0.000000000E+00 EPSI_Z: 0.000000000E+00|
| Distance from Waist. X: 0.000000000E+00 Z: 0.000000000E+00    |
| Polarization Used: SR TOTAL                                   |
| +----------------------------------------------------------------+
| Source Photon Energy Distribution: SINGLE LINE                |
| Photon Energy: 1000.0 eV, or 12.399 Angs.                     |
| +----------------------------------------------------------------+
| *************************************************************** |
| END                                                            |
```
Torroidal Mirror With Exit Slit.

Central Axis parameters:
- Source Plane Distance: 10000.00000000000
- Image Plane: 5000.000000000000
- Incidence Angle: 88.99999999999998
- Reflection/Diffraction Angle: 88.99999999999998

Mirror parameters:
- Major Radius: 381874.9072807532
- Minor Radius: 116.3493762485566
- Source of this O.E. moved: NO
- Mirror at pole position (no mov.): YES
The first plot shows how a large portion of the beam is blocked by the exit slit.
We will now use the lost rays from the exit slit (STAR.01) to modify the good/lost flag in the
source using REFLAG.

$ reflag
  File to be modified [ e.g. BEGIN.DAT ] ? begin.dat
  File to use as modifier [ e.g. SCREEN.0503 ] ? star.01
  Output file ? opt.dat

Now HISTO3 will divide the phase space up into good and lost bins.

$ histo3
  Input file ? opt.dat
  Output ? opt.hist
  Found 527 good points out of 1000
  Here we are.
  X max is 0.2972638002614275
  X min is -0.3260431318536972
  Y max is 24.30820832467235
  Y min is -24.00455259924489
  Z max is 2.6589882743400154E-02
  Z min is -3.3074220129884787E-02
  X prime max is 5.0536091497003565E-03
  X prime min is -4.9907608257961093E-03
  Y prime max is 0.999999987862705
  Y prime min is 0.9999871499645169
  Z prime max is 5.8449901106416506E-04
  Z prime min is -5.4485118546760329E-04
  How many bins in X  [ default 25 ] ?
  How many bins in X’ ?
  How many bins in Y  ?
  How many bins in Y’ ?
  How many bins in Z  ?
  How many bins in Z’ ?
Now the file generated by HISTO3 is used as the optimization file in SOURCE. The easiest way to perform this is to enter the menu mode of SOURCE and go into the 'Number of random rays' submenu by using the right arrow. The optimization of source should be changed to 'YES', and the file name is that of the output from HISTO3. Upon exit of SOURCE the number of rays generated vs the number accepted is displayed:

```
Phase space boundaries file read successfully.
500 rays have been rejected so far.
208 accepted.
Generated 250 rays out of 1000
1000 rays have been rejected so far.
393 accepted.
    500
1500 rays have been rejected so far.
595 accepted.
    750
2000 rays have been rejected so far.
778 accepted.
2500 rays have been rejected so far.
963 accepted.
    1000
Exit from SOURCE
SOURCE => Source has been successfully generated.
```

SOURCE procedure completed.

Now the system can be retraced in the BATCH mode since the parameters have not changed. We can then examine the exit slit of the optimized system using PLOTXY. Note that 995 out of the 1000 rays have been accepted instead of 527 for the first run. Let's optimize the TGM system with respect to the first mirror. Most of the rays lost are at the edge of the mirror. HISTO3 cannot optimize at the exit slit of a monochromator, because the lost rays are a function of energy, not phase space. In the previous example we plotted the progress against a limiting aperture. In this case we will look at a MINMAX file of the mirror in question to see how the limits change:

```
$ minmax
Input file ?
mirr.01
```
Figure 2.9: HISTO3: Before source optimization
Figure 2.10: HISTO3: After source optimization
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ?
2
Comment ?
First Mirror of TGM without optimization
***********
File examined is : D13:[XRAYOP.DOCUMENT.TGM]MIRR.01;29
Was created : 9-MAY-1989 11:16
***********
Found 927 good points out of 1000
Here we are.
X max is  2.233099
X min is -2.203895
Y max is  4.030588
Y min is -5.178271
Z max is  0.1620322
Z min is  5.9245693E-05
X prime max is  1.2591288E-02
X prime min is -1.3506259E-02
Y prime max is  0.9964082
Y prime min is  0.9960193
Z prime max is  8.9137740E-02
Z prime min is  8.3950877E-02

We pay most attention to the X, Y limits since these define the mirror’s edge.

$ reflag
File to be modified [ e.g. BEGIN.DAT ] ?
begin.dat
File to use as modifier [ e.g. SCREEN.0503 ] ?
mirr.01
Output file ?
reflag.opt

$ histo3
Input file ?
reflag.opt
Output ?
histo.opt
Found 927 good points out of 1000
Here we are.
X max is 0.2020543762143582
X min is -0.1922062231865444
Y max is 2.489119530789779
Y min is -2.421645912872256
Z max is 3.3317030366455779E-02
Z min is -3.4645052849801248E-02
X prime max is 1.1941889145443042E-02
X prime min is -1.1634579482826180E-02
Y prime max is 0.9999999788492193
Y prime min is 0.9999280937975692
Z prime max is 1.9853600893657853E-03
Z prime min is -1.9938319051862862E-03
How many bins in X [ default 25 ] ?
How many bins in X' ?
How many bins in Y ?
How many bins in Y' ?
How many bins in Z ?
How many bins in Z' ?

The user must change the source parameters to include the optimization file as explained in the SOURCE manual. The following run is made with the file histo.opt as the optimization file.

$ go source
SOURCE selected. Begin procedure.

This procedure generates a SOURCE for SHADOW.
Mode selected [ ? <ret> for HELP ] ? : batch
File containing source description [ START.00 ] ?: synch.00
Phase space boundaries file read successfully.
Generated 250 rays out of 1000
500
750
1000

Exit from SOURCE
SOURCE => Source has been successfully generated.

Now we'll run MINMAX on the mirror file again to see how the limits have changed. Note that the improvement was slight, but this is where the phase space dependent rays were being lost.

$ minmax
Input file ?
mirr.01
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ?
2
Comment ?
First mirror of TGM after optimization
***********
File examined is :
D13:[XRAYOP.DOCUMENT.TGM]MIRR.01;30
Was created :
10-MAY-1989 12:47
***********
Found 973 good points out of 1000

Here we are.

X max is 2.083563
X min is -2.086486
Y max is 4.180661
Y min is -4.992939
Z max is 0.1412461
Z min is 5.4842703E-06
X prime max is 1.1868485E-02
X prime min is -1.1789099E-02
Y prime max is 0.9964100
Y prime min is 0.9960120
Z prime max is 8.9194998E-02
Z prime min is 8.3885789E-02
Purpose: To create a file which can be used by SHADOW to calculate reflectivity of mirrors.

Input: Interactive from terminal.


Description:

PREREFL is a program to compute the complex dielectric function of a given material as a function of photon energy. The results are stored in an unformatted file ready to be read by SHADOW for calculating reflectivity of mirrors (multilayers coatings are handled differently by PREMLAYER and crystals by BRAGG). The user must run PREREFL first if he is going to include reflectivity when calling SHADOW later on. (Another case would be absorbing screens along the optical system. The user will still run PREREFL first, specifying the absorbing material, then pass the output file to SHADOW.)

PREREFL looks up an optical constant library which contains all the elements and covers the range 100 eV to 100,000 eV. The lower energy range (100-10,000 eV) is from B. Henke’s compilation, while the higher range (10,000-100,000 eV) is calculated by Cromer and Liberman’s method. Anything outside the range will create fatal errors. Naturally, the photon energy range should cover the range used in generating the source rays. Some consideration should be given to whether the range includes absorption edges, when choosing the number of points.

In the example, we define an elemental (gold) and a compound (SiC) coatings, from 100 to 500 eV in step of 2 eV. Note silicon must be typed in as SI, and carbon as CC.

Examples:

```bash
\ \ First let us define a gold mirror.

$ PREREFL
Element [0] or compound [1] ? 0
Density [ g/cm3 ] ? 19.3
Enter 2-letters (capitalized) atomic symbol ? AU \ \ Must be CAPITALIZED \
```
Enter starting photon energy, end and step
100
500
2
Output file?
AU.REFL
$
$ DIR AU.REFL

Directory D13:[XRAYOP.DOCUMENT.PREREFL]

AU.REFL;1  7  22-FEB-1989 21:58  (RWED,RWED,RE,RE)

Total of 1 file, 7 blocks.
$

\ Now we define another mirror made of SiC.

$ PREREFL
Element [0] or compound [1] ?
1
Density [ g/cm3 ] ?
3.21
The program is setup to compute optical constants of
compounds with up to 5 elements.
How many atomic species ?
2
H2O would be: HH,2 and OO,1. Then ?
Enter 2-letters (capitalized) atomic symbol and formula index for : 1
SI
1
Enter 2-letters (capitalized) atomic symbol and formula index for : 2
CC \ Must be 2-letters \ 1
Molecular weight is : 40.10000  Number of molecules/cm3:  4.8206819E+22
Enter starting photon energy, end and step
100
500
2
Output file?
SIC.REFL
$ DIR *.REFL

Directory D13:[XRAYOP.DOCUMENT.PREREFL]

AU.REFL;1 7 22-FEB-1989 21:58 (RWED,RWED,RE,RE)
SIC.REFL;1 7 22-FEB-1989 22:03 (RWED,RWED,RE,RE)

Total of 2 files, 14 blocks.

Either of these files can now be specified to SHADOW for reflectivity calculation.

The followings are typical responses to SHADOW:

Reflectivity of Surface. SHADOW may solve the Fresnel equations locally.
Available options:'No reflectivity .......... 0
Full polarization dependence ............ 1
No " .......... 2
Reflectivity mode [ 0,1,2 ] ?
1 " Either 1 or 2 selects reflectivity mode \\

Optical constants from file ( for multi-line source ) ...... 0
Optical constants from keyboard (single-line source ) ...... 1
Define multilayer from file .......... 2
Optical Constant Source: [file=0,tt:=1], multilayer [2] ?
0
File with optical constants ?
AU.REFL
Purpose: To generate a spline file used by SHADOW to simulate distortions in the surface of an optical element.

Input: A file containing the array size for the distortion and the surface mesh.

Output: A file readable by SHADOW containing the spline for the distortion.

Description:

This utility takes a file with a surface mesh array and computes the bi-cubic spline for the surface. Normally this is interpreted as an “error surface”, although one could use weird shapes by combining suitable error surfaces and plane or other mirrors. It creates a file that can be read by SHADOW under “Distorted Surface?”. The system will then be traced as if this defect was present on the mirror. This utility is particularly useful in conjunction with MAKE_PWR and PWR_DENS in modelling the effects of heating on a mirror surface. PRESURFACE takes a file in the format of:

\[
N_x, N_y, \ y(1), \ldots, y(N_y), \ x(1), z(1,1), \ldots, z(1,N_y), \\
\vdots, \\
x(N_x), z(N_x,1), \ldots, z(N_x, N_y)
\]

where \(N_x\) and \(N_y\) are the size of the surface mesh.

Examples:

To start, we will use a program generated error of the form \(\delta z = re^{-r^2/\sigma^2}\) and run that through PRESURFACE. PRESURFACE will then create a binary file for SHADOW. The second example is more complex, but a better demonstration of the usefulness of the program. We look at the distortions to the image of the TGM caused by heat loading on the mirror. This is done by taking the output from PWR_DENS/MAKE_PWR and submitting it to the finite element program ANSYS for analysis. ANSYS then returns a file in the form needed by PRESURFACE so we can give the information to SHADOW.

In the first example, the test program creates a gaussian error file in the form given above for an error with a height of 500 Å. This is run through PRESURFACE and traced through SHADOW as shown below.
Source used in PRESURFACE Example

Input file specified: end.00
Full file specification: CXRL$DUA0:[XRAYOP.DOCUMENT.UTIL.PRESURFACE]END.00;1
Creation Date: 7-SEP-1989 16:12

Grid Source.
Generated total 3721 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: RECTANGULAR
Source Width: 40.0000000 and Height: 5.0000000

Source Emission Characteristics
Distribution Type: UNIFORM
Distribution Limits. +X: 0.000000000E+00 -X: 0.000000000E+00 rad
+Z: 0.000000000E+00 -Z: 0.000000000E+00 rad

PRESURFACE Example
Distorted surface using test program
Surface figure was defined as: PLANE
Cylindrical figure          NO
Element type                REFLECTOR
Reflectivity                OFF
Mirror dimensions           UNLIMITED

Central Axis parameters :
Source Plane Distance       100.00000000000000
Image Plane                 100.00000000000000
Incidence Angle             0.0000000000000000E+00
Reflection/Diffraction Angle 0.0000000000000000E+00

Mirror parameters          COMPUTED
Same configuration as Central Axis NO
Objective focus at         0.0000000000000000E+00
Image focus at             0.0000000000000000E+00
Incidence angle            0.0000000000000000E+00
Parameters used follow:
  Plane mirror
Source of this O.E. moved   NO
Mirror at pole position ( no mov. ) YES

The plots shown below are the cross-sections at the image plane after the mirror. The first is without distortion, the second, after the 500 Å gaussian error was applied. We also show the output from running SURPLOT on the mirror surface to show the distortions in a clearer manner.
Figure 2.11: PRESURFACE: No distortion
Figure 2.12: PRESURFACE: With gaussian error
Figure 2.13: PRESURFACE: Surface generated with SURPLOT
Purpose: To create a file which has the parameters that SHADOW needs to calculate reflectivity of multilayers.
Input: Interactive from terminal.

Description:
PRE_MLAYER operates along the same lines as PREREFL. It lets the user specify the multilayer parameters prior to running SHADOW. The necessary complex refractive indices are obtained from an optical constant library (see PREREFL), within the user specified photon energy range. These are stored together with other parameters in a format understandable by SHADOW, where both the incident angle and photon energy might vary from ray to ray. Naturally, the photon energy range used here should contain that of the source rays. Valid energies are from 100 to 100,000 eV. Each layer material could be a compound or simply an element. For simplicity, elemental materials are implemented as the case of compounds with only one constituent.

In the example, we defined a 20 layer tungsten-carbon (W-C) multilayers deposited on silicon substrate. The geometrical structure is as follows:

<table>
<thead>
<tr>
<th>Layer</th>
<th>Material</th>
<th>Thickness (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20th</td>
<td>C</td>
<td>14.59</td>
</tr>
<tr>
<td></td>
<td>W</td>
<td>7.74</td>
</tr>
<tr>
<td>19th</td>
<td>C</td>
<td>14.59</td>
</tr>
<tr>
<td></td>
<td>W</td>
<td>7.74</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>

1SHADOW uses a multilayer reflectivity code developed by J. H. Underwood.
Note TWO capitalized letters must be used to specify each element (silicon as SI, tungsten as WW, and so on). Also note the thicknesses are entered upward starting from the substrate as:

\[
\begin{align*}
7.71,14.56 \\
7.72,14.57 \\
7.74,14.59 \\
-1,-1
\end{align*}
\]

The (-1,-1) means the 4th to 20th layers have the same thicknesses as the last entered layer (3rd layer). Of course, if the thicknesses are different from layer to layer, all twenty pairs must be entered individually in ascending order. If they all have the same thicknesses, then just enter:

\[
\begin{align*}
7.71,14.56 \\
-1,-1
\end{align*}
\]

For the reflectivity of these multilayers as a function of angle, at a fixed incident wavelength of 18 Å, refer to the example under MLAYER.

**Examples:**

```
$ PRE_MLAYER
Name of output file :
CW.REFL
Photon energy (eV) from :
500
    to :
```
2000
****************************************************************** \ Si substrate \\
Specify the substrate material:
Density [ g/cm3 ] ?
2.32
How many atomic species (up to 5) ?
1
H2O would be: HH,2 and OO,1. Then ?
Enter 2-letters (capitalized) atomic symbol and formula index for:
SI
1
Formula:
SI( 1)

Molecular weight is : 28.10000  Number of molecules/cm3: 4.9719812E+22  Atoms/Cm3 of each species:
SI
0.49720E+23
Density 2.320000 g/cm3
****************************************************************** \ W layers \\
Right above the substrate is the even layer material.
Specify the even layer material:
Density [ g/cm3 ] ?
19.3
How many atomic species (up to 5) ?
1
H2O would be: HH,2 and OO,1. Then ?
Enter 2-letters (capitalized) atomic symbol and formula index for:
WW
1
Formula:
WW( 1)

Molecular weight is : 184.0000  Number of molecules/cm3: 6.316570E+22  Atoms/Cm3 of each species:
WW
0.63167E+23
Density 19.30000 g/cm3
****************************************************************** \ C layers \\
Odd layer material is on top of the even layer.
Specify the odd layer material:
Density [ g/cm^3 ]?
2.26
How many atomic species (up to 5)?
1
H2O would be: HH,2 and OO,1. Then?
Enter 2-letters (capitalized) atomic symbol and formula index for:
1 CC
Formula:
CC(1)

Molecular weight is: 12.00000 Number of molecules/cm^3: 1.1341619E+23
Atoms/Cm^3 of each species:
  CC
    0.11342E+24 Density 2.260000 g/cm^3

Thicknesses

No. of layer pairs:
20
Now specify the thickness (angstroms) of each pair starting from the substrate surface. Type two -1 whenever you want the remaining layers to assume the thicknesses of the previous one.

Even and odd thicknesses of layer 1
7.71,14.56

Even and odd thicknesses of layer 2
7.72,14.57

Even and odd thicknesses of layer 3
7.74,14.59

Even and odd thicknesses of layer 4
-1,-1

$ $ DIR CW.REFL

Directory D13:[XRAYOP.DOCUMENT.PRE_MLAYER]

CW.REFL;1  7 23-FEB-1989 20:37 (RWED,RWED,RE,RE)

Total of 1 file, 7 blocks.
Now CW.REFL can be used by SHADOW to compute the multilayers reflectivity.

The followings are typical responses to SHADOW:

Reflectivity of Surface. SHADOW may solve the Fresnel equations locally.
Available options:
- 'No reflectivity' ............ 0
- Full polarization dependence ............ 1
- No polarization dependence ............ 2
Reflectivity mode [ 0,1,2 ] ?
1 \ Either 1 or 2 selects reflectivity mode \n
Optical constants from file (for multi-line source) ...... 0
Optical constants from keyboard (single-line source) ...... 1
Define multilayer from file ...... 2
Optical Constant Source: [file=0,tt:=1], multilayer [2] ?
2
File with optical constants ?
CW.REFL
Purpose: To examine the output from MAKE_ID in a formatted fashion.

Input: CDF100.DAT type unformatted file created by MAKE_ID.

Output: A listing to the terminal of the following information contained in CDF100.DAT:
- ENERGY
- THETA
- PHI
- ENERGY CDF
- THETA CDF
- PHI CDF
- POLARIZATION

Description:

This utility may be used after having run MAKE_ID to create an undulator source. MAKE_ID writes two unformatted files called CDF100.DAT and RN100.DAT. The first contains the integrated probability distribution function for the undulator specified. The second contains the unintegrated version (i.e., the probability density function). READCDF will translate these files and display them on the screen, listing them in the following order: $\omega, \theta, \phi, \omega CDF, \theta CDF, \phi CDF,$ and polarization.

Examples:

Shown here is an abbreviated log run of READCDF. Since we have chosen an array of size (7,31,31), the number of $\phi$ values is quite high, 6727. Therefore we have indicated where part of the numbers have been taken out for brevity.

```
$ run util:readcdf
CDF file name : CDF100.DAT
7       31       31
ENERGY
166.6000000000000   166.6514536735605   166.7083895768990
166.7634291358226   166.8087044526493   166.8511310282644
166.9000000000000
THETA
```
0.0000000000000000E+00 1.4791657137678076E-05 2.251941231713891E-05 2.7687702415863914E-05 3.2313066897178147E-05 3.6571308379987740E-05
::


PHI
0.0000000000000000E+00 5.2365113547351804E-02 0.1047198074795338 0.1570796332030884 0.2617993877992018
::

0.9931394837863106 1.052217027334260 1.12161870390235

ENERGY CDF
0.0000000000000000E+00 1597452437754297. 3352259513027127.
5052544494091979. 6617314861817053. 7933820056896882.
94838647996998803.

THETA CDF
0.0000000000000000E+00 1137879769018200. 2302194120683513.
3445473695689648. 4583600464381745. 5719824268258108.
::

3.2037618174895696E+16 3.3120834622079286E+16 3.4135948814895818E+16
3.523969259666078E+16 3.636833680816223E+16 3.674552875051814E+16
3.6769491832312990E+16

PHI CDF
0.0000000000000000E+00 2.6945087860844903E+23 5.3884814185598183E+23
8.0827181134233465E+23 1.0776957448910698E+24 1.3471196810804287E+24
::

4.1471861771090057E+19 4.3182396273454665E+19 4.4909831649742257E+19
4.6637392633570821E+19 4.8364869209620913E+19 5.0091263816753908E+19
5.1815261815261810E+19

POLARIZATION
1.0000000000000000
1.0000000000000000
1.0000000000000000
<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.80</td>
<td>0.67</td>
<td>0.52</td>
</tr>
<tr>
<td>0.33</td>
<td>6.23</td>
<td>0.43</td>
</tr>
<tr>
<td>0.71</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
REFLAG

**Purpose:**
Modifies the good/lost ray flag (column 10) in one BOF (object) according to that of another BOF (control). This utility is typically used to study which rays will be lost from a given source and to define masks for scattered light.

**Input:**
Two Binary Output Files (Usually BEGIN.DAT and another ray file like STAR.xx).

**Output:**
A modified version of the first input file.

**Description:**
The program reads in both files and copies the status flag of the control file to the object file, ray by ray. A typical application is for example, in answering the question “what fraction of the source is the system accepting?” or “how much of a condenser mirror are we really using?”. In the first case the object file would be BEGIN.DAT, in the second MIRR.XX, while the control file could be for example the final image STAR.YY.

Another application is that of reducing scattered light. Let’s suppose that a beam overfills a mirror thus creating stray rays that may have harmful effects. One can then use REFLAG to see which part of the incoming beam (e.g. at a suitable screen) will miss the mirror and thus design a suitable absorbing mesh that will effectively eliminate stray radiation.

**Examples:**

First the output from the source and single element system that we’ll use for our example.

```
+----------------------------------------------------------------------------------------------------------+
|                          SOURCE DESCRIPTION                                               |
+----------------------------------------------------------------------------------------------------------+
| REFLAG Example                     |
| Source used in REFLAG before optimization                                                               |
+----------------------------------------------------------------------------------------------------------+
| Input file specified:               |
| end.00                               |
| Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.REFLAG]END.00;1                                    |
| Creation Date :17-MAY-1989 16:26                                                                |
| Random Source.                                                                      |
| Generated total 2000 rays.                                                              |
| Source assumed TRIDIMENSIONAL.                                                            |
| Source Spatial Characteristics: GAUSSIAN                                                  |
+----------------------------------------------------------------------------------------------------------+
```
Sigma X : 0.100000000 Sigma Z : 0.10000000E-01
Depth: SYNCHROTRON SOURCE.

Source Emission Characteristics
Distribution Type: SYNCHROTRON
Distribution Limits. +X : 0.100000000E-01 -X: 0.100000000E-01 rad
+Z : 0.500000000E-02 -Z: 0.500000000E-02 rad
Magnetic Radius = 4.81000000 m. Beam Energy = 1.5000 GeV.
Beam Emittances. EPSI_X: 0.000000000E+00 EPSI_Z: 0.000000000E+00
Distance from Waist. X: 0.000000000E+00 Z: 0.000000000E+00
Polarization Used: SR TOTAL

Source Photon Energy Distribution: SINGLE LINE
Photon Energy: 1000.0 eV, or 12.399 Angs.

REFLAG Example
Mirror used in REFLAG example
Input file specified:end.01
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.REFLAG]END.01;1
Creation Date :17-MAY-1989 16:26

Surface figure was defined as: TOROIDAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity OFF
Mirror dimensions UNLIMITED
Central Axis parameters :
Source Plane Distance 10000.00000000000
Image Plane 5000.00000000000
Incidence Angle 88.99999999999998
Reflection/Diffraction Angle 88.99999999999998
Mirror parameters: COMPU TED

Same configuration as Central Axis: YES

Objective focus at: 10000.0000000000
Image focus at: 5000.0000000000
Incidence angle: 88.9999999999999

Parameters used follow:
   Major Radius: 381874.9072807532
   Minor Radius: 116.3493762485566

Source of this O.E. moved: NO
Mirror at pole position (no mov.): YES

$ reflag

File to be modified [e.g. BEGIN.DAT] ? begin.dat
File to use as modifier [e.g. SCREEN.0503] ? star.01
Output file ? refl.dat

$ histo3
Input file ? refl.dat
Output file ? new.dat

Found 550 good points out of 1000

Here we are.

X max is 0.2730171936559060
X min is -0.2595255943453094
Y max is 24.25826988410192
Y min is -24.24719610097799
Z max is 3.1774282219930756E-02
Z min is -4.2464660404542004E-02
X prime max is 5.0432633039571831E-03
X prime min is -5.041184507958304E-03
Y prime max is 0.99999999990291811
Y prime min is 0.9999872410237632
Z prime max is 6.1683691834940080E-04
Z prime min is -5.8033590292263016E-04

How many bins in X [default 25] ?
How many bins in X’ ?
How many bins in Y ?
How many bins in Y’ ?
How many bins in $Z$  
How many bins in $Z'$  

The output file from HISTO3 is used to optimize the source. To perform this optimization, the user should enter the source generation program by typing: `$GO SOURCE` and selecting MENU mode.

1. LOAD SOURCE
2. SEL SOURCE
3. enter the submenu for "Number of Random Rays" by placing the cursor next to the '1000' in the rightmost column and pressing the right arrow key.
4. Change 'optimization of source' from NO to YES
5. Enter the name of the file that was generated by HISTO3
6. Exit the menus by pressing the left arrow twice
7. SAVE SOURCE
8. EXIT

The source will now generate only those rays that fit into the bounds defined by HISTO3:

Shown below is the exit from SOURCE after optimization.

```
Phase space boundaries file read successfully.
   500 rays have been rejected so far.
      46 accepted.
  1000 rays have been rejected so far.
       82 accepted.
  1500 rays have been rejected so far.
      115 accepted.
 2000 rays have been rejected so far.
      157 accepted.
 2500 rays have been rejected so far.
      197 accepted.
 3000 rays have been rejected so far.
      233 accepted.
Generated 250 rays out of 1000
   3500 rays have been rejected so far.
```
280 accepted.
4000 rays have been rejected so far.
321 accepted.
4500 rays have been rejected so far.
364 accepted.
5000 rays have been rejected so far.
403 accepted.
5500 rays have been rejected so far.
446 accepted.
6000 rays have been rejected so far.
489 accepted.
500
6500 rays have been rejected so far.
529 accepted.
7000 rays have been rejected so far.
574 accepted.
7500 rays have been rejected so far.
620 accepted.
8000 rays have been rejected so far.
667 accepted.
8500 rays have been rejected so far.
714 accepted.
750
9000 rays have been rejected so far.
750 accepted.
9500 rays have been rejected so far.
793 accepted.
10000 rays have been rejected so far.
847 accepted.
10500 rays have been rejected so far.
898 accepted.
11000 rays have been rejected so far.
946 accepted.
11500 rays have been rejected so far.
982 accepted.
1000
Exit from SOURCE
SOURCE => Source has been succesfully generated.

SOURCE procedure completed.
Shadow:: exit

Exit to DCL
Chapter 3

Postprocessors

After running SHADOW, it is necessary to have programs to analyze the results and put them in a more understandable form. The Utilities in this section do summarization and ray analysis. They are broken down into four sections: reporting utilities, graphical analysis, non-graphical analysis, and specialized post-processors.
Purpose: Reads an ERF (exit record file - e.g. END.xx) associated with an optical element and displays information (including parameters computed during a trace) about that OE in an intelligible format.

Input: An ERF associated with an optical element.

Output: TT: or a user named and labeled text file.

Description:

Takes the information from an exit record file and puts it into a readable format. The following data is included in the output file:

- Exit Record File Data: Name, Creation Date
- Surface Type
- Element Type
- Reflectivity Mode
- Mirror Dimensions
- Source/Image plane Distances
- Incidence/Reflection Angles
- Mirror Parameters
- Focal positions in Object and Image Space

Examples:

Shown below is the output from three runs through MIRINFO. The first run is the first mirror in the two mirror example from the primer.

$ mirinfo

MINFO> Input file MUST be an ERF (Exit Record File) type from SHADOW
MININFO> Please input filename:
end.01
MININFO> File read correctly.
MININFO> Title ?
Two Mirror Example: First Mirror
MININFO> Comment ?
Original position from primer example
MININFO> Output file ?
mirr1.inf
MININFO> Prepare output to file : mirr1.inf

+-----------------------------------------------------------------------------+
|                           MIRROR DESCRIPTION                             |
+-----------------------------------------------------------------------------+
Two Mirror Example: First Mirror
Original position from primer example
+-----------------------------------------------------------------------------+
Input file specified:end.01
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.MIRINFO]END.01;1
Creation Date :10-APR-1989 15:03
+-----------------------------------------------------------------------------+
Surface figure was defined as: ELLIPTICAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity OFF
Mirror dimensions UNLIMITED
+-----------------------------------------------------------------------------+
Central Axis parameters :
Source Plane Distance 2000.000000000000
Image Plane 500.00000000000000
Incidence Angle 88.00000000000000
Reflection/Diffraction Angle 88.00000000000000

Mirror parameters COMPUTED
Same configuration as Central Axis YES
Objective focus at 2000.000000000000
Image focus at 500.00000000000000
Incidence angle 88.00000000000000
Parameters used follow:

- Semi-major axis: 1250.000000000000
- Semi-minor axis: 34.89949670250097
- Semi-focal-length: 0.000000000000000E+00
- Eccentricity: 0.000000000000000E+00

Source of this O.E. moved: NO
Mirror at pole position (no mov.): YES

Using the same two mirror example, we move the first mirror in the mirror reference frame.
Notice the new information MIRINFO gives:

- Surface figure was defined as: ELLIPTICAL
- Cylindrical figure: NO
- Element type: REFLECTOR
- Reflectivity: OFF
- Mirror dimensions: UNLIMITED

Central Axis parameters:
- Source Plane Distance: 2000.000000000000
- Image Plane: 500.0000000000000
- Incidence Angle: 88.0000000000000
- Reflection/Diffraction Angle: 88.0000000000000

Mirror parameters: COMPUTED
Same configuration as Central Axis: YES
Objective focus at 2000.000000000000
Image focus at 500.000000000000
Incidence angle 88.00000000000000
Parameters used follow:
   Semi-major axis 1250.00000000000000
   Semi-minor axis 34.89949670250097
   Semi-focal-length 0.00000000000000E+00
   Eccentricity 0.00000000000000E+00

Source of this O.E. moved NO
Mirror moved from pole. Parameters:
   Displacement along X: 0.00000000000000E+00
   Y: 0.00000000000000E+00
   Z: 100.00000000000000
   Rotation around X: 0.00000000000000E+00
   Y: 0.00000000000000E+00
   Z: 0.00000000000000E+00

The final example we show is the first crystal example from the primer. Compare the information in this example with that of earlier examples.
Bragg Reflection from GAAS.PAR
Reflectivity ON coefficients from: GAAS.PAR
Polarization dependence YES
Mirror dimensions UNLIMITED

Central Axis parameters:
Source Plane Distance 5000.000000000000
Image Plane 149.8956000000000
Incidence Angle 80.19284670377978
Reflection/Diffractive Angle 80.19284670377978

Mirror parameters COMPUTED
Same configuration as Central Axis NO
Objective focus at 0.000000000000000E+00
Image focus at 0.000000000000000E+00
Incidence angle 0.000000000000000E+00
Parameters used follow:
Plane mirror
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

******************************************************************************
******************************************************************************
**Purpose:** Reads a source ERF and creates a file with all relevant source information in an intelligible format.

**Input:** Any source Exit Record File: e.g END.00

**Output:** A user defined and labeled file (or TT:) with a description of source.

**Description:**

Given the source Exit Record File, SOURCINFO will tell the user all of the information regarding creation of the source:

- Type (Random or Grid)
- Number of rays generated
- Source Depth Type
- Spacial Characteristics
- Emission Characteristics (angle distribution)
- Photon Energy Distribution

The user can put a title along with a comment at the top of the file to help identify the source information.

**Examples:**

We'll run two examples. The first will be the source used in the first crystal example in section 6.4 of the primer.

```
$ sourcinfo
```

```
------------------ SOURCINFO --------------------
v3.2.01 Feb 1989

INPUT> File containing source specs ?
end.00
```
INPUT> Title ?
Primer Crystal 1 Example
INPUT> Comment ?
Source used in sec. 6.4 of primer, crystal example
INPUT> Output file-name ?
xtal.inf
SYSINF> Prepare output to file : xtal.inf
SOURCE_INFO> File
end.00
read correctly.
$
$ type xtal.inf

+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
************** SOURCE DESCRIPTION **************
Primer Crystal 1 Example
Source used in sec. 6.4 of primer, crystal example
+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified:
end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.SOURCINFO]END.00;1
Creation Date :26-MAR-1987 14:22
+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Random Source.
Generated total 1000 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: POINT
+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Source Emission Characteristics
Distribution Type: LAMBERTIAN
Distribution Limits. +X : 0.000000000E+00 -X: 0.000000000E+00 rad
+Z : 0.600000000E-04 -Z: 0.600000000E-04 rad
+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Source Photon Energy Distribution: SINGLE LINE
Photon Energy: 11160. eV, or 1.1110 Angs.
+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
END
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Second example: We'll just show the output of the file synch.inf, a model of the synchrotron source Aladdin.

```
+++
*********** SOURCE DESCRIPTION ***********
+++
Synchrotron Source
Sourcinfo Example 2
+++
Input file specified:
end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.SOURCINFO]END.00;1
Creation Date :10-APR-1989 16:38
+++
Random Source.
Generated total 1000 rays.
Source assumed TRIDIMENSIONAL.
Source Spatial Characteristics: GAUSSIAN
Sigma X : 0.675000000 Sigma Z : 0.350000000
Depth: SYNCHROTRON SOURCE.
+++
Source Emission Characteristics
Distribution Type: SYNCHROTRON
Distribution Limits. +X : 0.100000000E-01 -X: 0.100000000E-01 rad
+Z : 0.500000000E-02 -Z: 0.500000000E-02 rad
Magnetic Radius = 2.08330000 m. Beam Energy = 1.0000 GeV.
Beam Emittances. EPSI_X: 0.000000000E+00 EPSI_Z: 0.000000000E+00
Distance from Waist. X: 0.000000000E+00 Z: 0.000000000E+00
Polarization Used: SR TOTAL
+++
Source Photon Energy Distribution: BOX DISTR.
From Photon Energy: 400.000000 or 30.9963000 Angs.
to Photon Energy: 5000.000000 or 2.47970400 Angs.
Angular difference in phase is 90.000
Degree of polarization is 0.99462
Source points have INCOHERENT phase.
+++
END
+++```

SOURCINFO
**Purpose:** Creates a user friendly simplified description of an optical system from the exit record files after a trace has been completed.

**Input:** A file containing a listing of the ERF’s for all the optical elements in the system. Also the optical axis file is requested as input. A description on how to create the ERF list is given below.

**Output:** A user defined text file that describes the optical system.

**Description:**

Gives a description of the optical system from the exit record files. Each optical element is labeled in the following format:

- **TYPE** (e.g. GRATING)
- **SURFACE** (e.g. TOROIDAL)
- **SIZE** ( (UN) LIMITED)
- **PARAMETERS** ( (not) COMPUTED)
- **REFLECTIVITY** (OFF/FULL/NO POL)

- Orientation Ang. (Degrees)
- Source Plane (Distance from OE to source: arbitrary units)
- Incidence Ang. (Degrees)
- Reflection Ang. (Degrees)
- Image Plane (Distance from OE to continuation plane)

Following the description of each OE, The Optical System Configuration is given in the Laboratory Reference Frame. This involves giving the position in X, Y, Z space of each of the optical elements and of their continuation planes:
Examples:

Below we show two runs through SYSINFO and the respective files generated. The first example is the two mirror example from the primer. The second example is the first crystal example from the primer.

$ sysinfo

------------------------ SYSINFO ------------------------

vs. 2.02 June 1989

Default filenames [ Y/N ] ? y
How many OE's ? 2
INPUT> Title ? SYSINFO Example
INPUT> Comment ? Two Mirror System from Primer Chapter 6
INPUT> Output file-name ? sysi.twom
SYSINF> Prepare output to file : sysi.twom

0
SYSINF> File
END.01

    read correctly.

0
SYSINF> File
END.02

    read correctly.
$ ty sysi.twom

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
SYSINFO

************** SYSTEM DESCRIPTION **************

SYSINFO Example
Two Mirror System from Primer Chapter 6

Input file specified:

# Optical Element: Creation Time:
1 D13:[XRAYOP.PRIMER.CHAPTER6.TWOMIRR]END.01;2 5-MAY-1987 14:
2 D13:[XRAYOP.PRIMER.CHAPTER6.TWOMIRR]END.02;2 5-MAY-1987 14:

Optical Element # 1 System Number:

MIRROR ELLIPTICAL UNLIMITED COMPUTED REFLEC. OFF

Orientation 0.0000000000000000E+00 deg.
Source Plane 2000.0000000000000
Incidence Ang. 88.00000000000000 deg.
Reflection Ang. 88.00000000000000 deg.
Image Plane 500.0000000000000

----------------

Optical Element # 2 System Number:

MIRROR ELLIPTICAL UNLIMITED COMPUTED REFLEC. OFF

Orientation 180.00000000000000 deg.
Source Plane 1000.0000000000000
Incidence Ang. 88.00000000000000 deg.
Reflection Ang. 88.00000000000000 deg.
Image Plane 500.0000000000000

----------------

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

OPT. Elem # X = Y = Z =
0 0.00000000000000E+00 0.00000000000000E+00 0.00000000000000E+00
$ sysinfo

-------------------- SYSINFO ----------------------
vs. 2.02 June 1989

Default filenames [ Y/N ] ? y
How many OE’s ? 2
INPUT> Title ? SYSINFO Example
INPUT> Comment ? Crystal Example 1 from Primer Chapter 6
INPUT> Output file-name ? sysi.xtal
SYSINF> Prepare output to file : sysi.xtal
  0
SYSINF> File
END.01
read correctly.
  0
SYSINF> File
END.02
read correctly.
$ ty sysi.xtal

$ sysinfo

-------------------- SYSINFO ----------------------
vs. 2.02 June 1989

Default filenames [ Y/N ] ? y
How many OE’s ? 2
INPUT> Title ? SYSINFO Example
INPUT> Comment ? Crystal Example 1 from Primer Chapter 6
INPUT> Output file-name ? sysi.xtal
SYSINF> Prepare output to file : sysi.xtal
  0
SYSINF> File
END.01
read correctly.
  0
SYSINF> File
END.02
read correctly.
$ ty sysi.xtal
Input file specified:

Optical Element: Creation Time:
1  D13:[XRAYOP.PRIMER.CHAPTER6.XTAL1]END.01;1  26-MAR-1987 14:
2  D13:[XRAYOP.PRIMER.CHAPTER6.XTAL1]END.02;1  26-MAR-1987 14:

Optical Element #  1  System Number:

<table>
<thead>
<tr>
<th>BRAGG</th>
<th>PLANE</th>
<th>UNLIMITED</th>
<th>COMPUTED</th>
<th>REFLEC. OFF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orientation</td>
<td>0.0000000000000000E+00 deg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Source Plane</td>
<td>5000.000000000000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incidence Ang.</td>
<td>80.19284670377978 deg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reflection Ang.</td>
<td>80.19284670377978 deg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image Plane</td>
<td>149.8956000000000</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

-------------------------

Optical Element #  2  System Number:

<table>
<thead>
<tr>
<th>BRAGG</th>
<th>PLANE</th>
<th>UNLIMITED</th>
<th>COMPUTED</th>
<th>REFLEC. OFF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orientation</td>
<td>180.0000000000000 deg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Source Plane</td>
<td>149.8956000000000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Incidence Ang.</td>
<td>80.19284670377978 deg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reflection Ang.</td>
<td>80.19284670377978 deg.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Image Plane</td>
<td>5000.0000000000000</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

-------------------------

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

<table>
<thead>
<tr>
<th>OPT. Elem #</th>
<th>X =</th>
<th>Y =</th>
<th>Z =</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.0000000000000000E+00</td>
<td>0.0000000000000000E+00</td>
<td>0.0000000000000000E+00</td>
</tr>
<tr>
<td>1</td>
<td>0.0000000000000000E+00</td>
<td>5000.000000000000</td>
<td>0.0000000000000000E+00</td>
</tr>
<tr>
<td>1’</td>
<td>0.0000000000000000E+00</td>
<td>5141.1977073</td>
<td>50.317972551</td>
</tr>
<tr>
<td>2</td>
<td>0.0000000000000000E+00</td>
<td>5282.3954145</td>
<td>100.63594510</td>
</tr>
<tr>
<td>2’</td>
<td>0.73966525681E-13</td>
<td>10282.395415</td>
<td>100.63594510</td>
</tr>
</tbody>
</table>
**Purpose:** Translates a BOF (e.g. BEGIN.DAT, STAR.xx) into a formatted file. Each of the twelve to eighteen rows (depending on source definition) for each ray is included in the output file.

**Input:** Any binary output file.

**Output:** A user-named formatted file or the terminal (TT).

**Description:**
This utility translates a binary output file into a formatted file. The total number of rays in the BOF is given to the user who then chooses how many to translate. The output is in double precision and is displayed in the following format.

- \( X \) \( Y \) \( Z \) Spatial coordinates
- \( X' \) \( Y' \) \( Z' \) Direction
- \( A_x \) \( A_y \) \( A_z \) EM vector (s-)
- FF q kk Flag, Wavenumber, ray index
- opd \( F_s \) \( F_p \) Optical path length, phases
- \( A_{p_x} \) \( A_{p_y} \) \( A_{p_z} \) EM Vector (p-)

As usual, the number of rows depends on the source definition (i.e. if opd is selected in source but polarization was not, then the number of rows is 13). TRANSLATE tells the user how many rows for each ray will be translated.

**Examples:**
Here we will use the synchrotron source set up in SOURCINFO, shown below:

```
+++
********** SOURCE DESCRIPTION **********
Synchrotron Source
Translate Example
+++...
```

Input file specified:
Random Source.
Generated total 1000 rays.
Source assumed TRIDIMENSIONAL.
Source Spatial Characteristics: GAUSSIAN
Sigma X : 0.6750000000 Sigma Z : 0.3500000000
Depth: SYNCHROTRON SOURCE.
Source Emission Characteristics
Distribution Type: SYNCHROTRON
Distribution Limits. +X : 0.1000000000E-01 -X: 0.1000000000E-01 rad
+Z : 0.5000000000E-02 -Z: 0.5000000000E-02 rad
Magnetic Radius = 2.08330000 m. Beam Energy = 1.0000 GeV.
Beam Emittancies. EPSI_X: 0.0000000000E+00 EPSI_Z: 0.0000000000E+00
Distance from Waist. X: 0.0000000000E+00 Z: 0.0000000000E+00
Polarization Used: SR TOTAL
Source Photon Energy Distribution: BOX DISTR.
From Photon Energy: 400.000000 or 30.9963000 Angs.
to Photon Energy: 5000.000000 or 2.47970400 Angs.
Angular difference in phase is 90.000
Degree of polarization is 0.99462
Source points have INCOHERENT phase.

Now let’s run BEGIN.DAT through TRANSLATE. Notice the non-zero values for depth and electric field, and compare them to the output from SOURCINFO above.

$ translate
File for input ? BEGIN.DAT
and for output?
trans.out
Read 1000 rays.
Each ray has 18 entries.
how many rays to translate?
5
All done.
$ dir/since

Directory D13:[XRAYOP.DOCUMENT.UTIL.TRANSLATE]

TRANS.OUT;1 5 13-APR-1989 12:10 (RWED,RWED,RE,RE)

Total of 1 files, 5 blocks.
$
$ type trans.out

-1.390714334133603 -9.879931896167709 5.0067780073250970E-02
-4.7392262218636672E-03 0.9999887652748167 9.5178598615866550E-05
0.9996444645105199 4.7375944414025118E-03 4.5092266573523865E-07
1.0000000000000000 95734809.56938891 1.0000000000000000E+00
0.0000000000000000E+00 3.074243931170887 4.645040257965784
0.0000000000000000E+00 -2.4974458683843627E-06 2.623927906675882E-02

2.7511145072241483E-02 -14.98043983256070 6.6027240850530335E-02
-7.1906338941880916E-03 0.9999739618211993 6.0865738670193557E-04
0.895468443927048 6.4391510223206930E-03 3.9193888627151111E-06
1.0000000000000000 46186641.30730766 2.0000000000000000E+00
0.0000000000000000E+00 2.071735804317716 3.642532131112613
-4.7124511123640432E-23 -2.7090727399034242E-04 0.4450783412567988

0.1530800437340783 -14.92438730181352 -0.3691629931304233
-7.1641626783434953E-03 0.9999742780491507 -3.4353049060675017E-04
0.9788676621087926 7.0129467309467256E-03 -2.4092230009967500E-06
1.0000000000000000 30557276.28737217 3.0000000000000000E+00
0.0000000000000000E+00 5.645537282077025 4.074740955282128
1.0819493662900201E-23 7.021065927267415E-05 0.2043744448811408

0.7045787542230621 -13.36189239266564 -0.566560849177220
-6.4158488038517787E-03 0.9999794153140405 7.636959785068862E-05
<table>
<thead>
<tr>
<th>X</th>
<th>Y</th>
<th>Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9997970251734600</td>
<td>6.4146785547840900E-03</td>
<td>4.8989651834470764E-07</td>
</tr>
<tr>
<td>1.0000000000000000</td>
<td>186852493.9196818</td>
<td>4.0000000000000000E00</td>
</tr>
<tr>
<td>0.0000000000000000E+00</td>
<td>1.763037151036863</td>
<td>3.33333477831759</td>
</tr>
<tr>
<td>0.0000000000000000E+00</td>
<td>-1.4585901185907957E-06</td>
<td>1.9098700242978440E-02</td>
</tr>
<tr>
<td>-0.6917105883993459</td>
<td>-14.81614060873230</td>
<td>0.1114545705002670</td>
</tr>
<tr>
<td>-7.1093210068994241E-03</td>
<td>0.9999747192975676</td>
<td>-1.3535350638976451E-04</td>
</tr>
<tr>
<td>0.9993560424606797</td>
<td>7.1049223932968824E-03</td>
<td>-9.617004710232756E-07</td>
</tr>
<tr>
<td>1.0000000000000000</td>
<td>21913642.01124334</td>
<td>5.0000000000000000E00</td>
</tr>
<tr>
<td>0.0000000000000000E+00</td>
<td>2.059520134068875</td>
<td>0.4887238072739785</td>
</tr>
<tr>
<td>0.0000000000000000E+00</td>
<td>4.7606793400995139E-06</td>
<td>3.5171301532988853E-02</td>
</tr>
</tbody>
</table>
**Purpose:**
To create a histogram of the frequency distribution of ray coordinates, from a selected column in an ASCII file (e.g., output from PREPLOT or after processing with FILEX).

**Input:**
Any column formatted file.

**Output:**
TT: or a user defined file.

**Description:**
This is a specialized version of HISTO1 and uses the same algorithm. H2KOL will create the histogram from a chosen column in a formatted two column file.

**Examples:**
To demonstrate H2KOL, we'll use a synchrotron source with an energy from 50 - 5000 eV. The output from SOURCINFO is shown. We run BEGIN.DAT through PREPLOT and extract columns 6 (angle distribution in Z) and 11 (photon energy).
Before we create the histogram, we should use FMAX to find the extrema in the data so we can make a reasonable estimate of the center and width of the histogram.

```
+---------------------------------------------------------------------+
<table>
<thead>
<tr>
<th>SOURCE DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>H2KOL Example</td>
</tr>
<tr>
<td>Synchrotron Source</td>
</tr>
<tr>
<td>----------------------------------</td>
</tr>
<tr>
<td>Input file specified:</td>
</tr>
<tr>
<td>end.00</td>
</tr>
<tr>
<td>Full file Specification :CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.H2KOL]END.00;2</td>
</tr>
<tr>
<td>Creation Date :28-DEC-1989 11:05</td>
</tr>
<tr>
<td>Random Source.</td>
</tr>
<tr>
<td>Generated total 5000 rays.</td>
</tr>
<tr>
<td>Source assumed TRIDIMENSIONAL.</td>
</tr>
<tr>
<td>Source Spatial Characteristics: GAUSSIAN</td>
</tr>
<tr>
<td>Sigma X : 0.600000000E-01 Sigma Z : 0.100000000E-01</td>
</tr>
<tr>
<td>Depth: SYNCHROTRON SOURCE.</td>
</tr>
</tbody>
</table>
+---------------------------------------------------------------------+```
Source Emission Characteristics
Distribution Type: SYNNCHROTRON
Distribution Limits. +X : 0.120000000E-01 -X: 0.120000000E-01 rad
+Z : 0.200000000E-02 -Z: 0.200000000E-02 rad
Magnetic Radius = 2.08330000 m. Beam Energy = 0.80000 GeV.
Beam Emittances. EPSI_X: 0.000000000E+00 EPSI_Z: 0.000000000E+00
Distance from Waist. X: 0.000000000E+00 Z: 0.000000000E+00
Polarization Used: SR TOTAL

Source Photon Energy Distribution: BOX DISTR.
From Photon Energy: 50.0000000 or 247.970400 Angs.
to Photon Energy: 5000.00000 or 2.47970400 Angs.
Angular difference in phase is 90.000
Degree of polarization is 0.62859
Source points have INCOHERENT phase.

*************** END ***************

$ dir
Directory CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.H2KOL]
BEGIN.DAT;1 1410 21-DEC-1989 13:19:30.57 (RWED,RWED,RE,)
END.00;1 7 21-DEC-1989 13:19:35.16 (RWED,RWED,RE,)
H2KOL.ABS;3 1 11-JUL-1989 16:22:53.90 (RWED,RWED,RWED,RE)
H2KOL.DSC;5 1 2-JUN-1989 10:19:00.35 (RWED,RWED,RWED,RE)
H2KOL.EXA;9 9 2-JUN-1989 10:20:01.53 (RWED,RWED,RWED,RE)
H2KOL.INP;1 1 24-JUN-1988 12:17:52.00 (RWED,RWED,RWED,RE)
H2KOL.LOG;1 1 21-DEC-1989 13:21:26.40 (RWED,RWED,RE,)
H2KOL.OUT;2 1 21-FEB-1989 14:25:32.78 (RWED,RWED,RWED,RE)
PHOTONS.DAT;1 313 21-DEC-1989 13:20:46.28 (RWED,RWED,RE,)
SOU.INF;1 5 21-DEC-1989 13:20:05.54 (RWED,RWED,RE,)
START.00;3 7 21-DEC-1989 13:17:55.70 (RWED,RWED,RE,)

Total of 11 files, 1755 blocks.
$ fmax
File-name ?
photons.dat
Read 5000 points.
Found:
XMIN  -1.9855129E-03  . Set at : -2.4330688E-03
XMAX  1.9927619E-03     2.4403178E-03
YMIN  50.12449    -357.5591
YMAX  3673.979      4081.663
Good luck!
$
$h2kol
Input file ? photons.dat
Column to analyze     [ 1 or 2 ] ? 1
Center    ? 0
Width     ? 5e-03
Number of bins      [ max=100 ] ? 31
Normalization kind. Enter :
0 for no normalization
1 to normalize to 1    [ at the peak ]
2 to the total counts   [ divide by N ]
3 use the other col. to fill in the bins
<?, 0
Read
Read 5000
Output options :
[ 0 ] store histogram in a file
[ 1 ] plot histogram on screen
[ 2 ] both
Then 0
$ dir/since=-00:02

Directory CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.H2KOL]

H2KOL.DAT;1 4 21-DEC-1989 13:25:17.00 (RWED,RWED,RE,)

Total of 1 file, 4 blocks.

The histogram can be normalized or scaled in four ways. The first is not to normalize the plot. The value of each bin is the actual number of points that fall in that bin. The second normalization, to normalize to 1, divides the actual counts by the greatest number of counts to fall in a bin. Thus the maximum value is 1. To normalize to the total counts divides each actual number of points by the total number of points plotted (N). The last type of normalization is most useful when trying to show intensity or amplitude. The phrase “use
the other col. to fill in the bins” means to weight each count by the value of the other column. For our example, this means we are plotting power distribution in angle.
Figure 3.1: H2KOL: No Normalization – ray distribution in angle
Figure 3.2: H2KOL: Normalize to 1
Figure 3.3: H2KOL: Normalize to total counts
Figure 3.4: H2KOL: Weighted Normalization – power distribution in angle
Purpose: To plot a two or three dimensional diagram of the rays near the position of a component of an optical system corresponding to a binary output file. Extends the use of FOCNEW.

Input: Any Binary Output File.

Output: A plottable file in two or three dimensions that can be used as input to a graphics routine such as TopDrawer.

Description:
Creates a file suitable for plotting the rays near a continuation plane. The utility asks the user how far in front of and how far beyond the plane the rays should be plotted. The distances are not absolute so the quantity before the plane should be negative. The paths are found by extrapolating:

\[ X(d + \Delta) = X(d) + \Delta \cdot X'(d) \]

where \( d \) is the location of the continuation plane and \( \Delta \) is the distance away from the plane along the optical axis.

Examples:
Here we will make a three dimensional plot in the vicinity of the focus in the tangential plane for the example run in FOCNEW. The source has been modified to a conical grid source angle distribution with only twenty rays for the purpose of clarity. From the information we obtained in FOCNEW, we can set the limits to five units on either side of the tangential and sagittal focii.

$ plotfoc
Input file ? star.01
Output file ? pf.dat
Read 20 points.
Distance before plane [ <0 ] : -5
after [ >0 ] : 15
Two dimension [ 0 ] or three [ 1 ] ? 1
Only good rays [ 0 ] or all [ 1 ] ? 1
OK. Here we go.
The following TopDrawer command file was used with the data from PLOTFOC to generate the plot shown below. The outlines of the tangential and sagittal planes were drawn for the sake of clarity.
Figure 3.5: PLOTFOC: Tangential and Sagittal focii
Figure 3.6: PLOTFOC: Sagittal plane focus
Figure 3.7: PLOTFOC: Tangential plane focus
Purpose: To generate four plots on the same screen of \((X,X'), (Z,Z'), (X,Z), (X',Z')\) of a ray file, thus providing a single view of the phase space.

Input: A BOF file. (such as STAR.xx or MIRR.xx)

Output: Four plots of specified file to terminal.

Description:

This utility will generate four plots on the terminal of a single ray file. The plots, clockwise from the top left, are of \(X\)-phase space \((X,X')\), \(Z'-X'\) space, \(Z\)-phase space \((Z',Z)\), and the \(X-Z\) plane. A TopDrawer command file is not written. If plotting only losses at a given OE, the flag value of those rays must be entered. This is simply the number of the optical element multiplied by 11000.

This program is very useful to have a general overview of the beam properties at any point along the optical axis and to verify, for example, the existence of astigmatism.

Examples:

We'll run this example on the STAR.01 file of the system used in FOCNEW.

```
$ run util:plotspace
PREPLOT> Input file ? star.01
Read 1000 rays.
Each ray has 12 entries.
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays.
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
PREPLOT> Then ?
0
***********
Found 1000 good points out of 1000
Read 1000 good rays. Plot all ?
> 1
```
We see that the sagittal direction ($X$) is not at focus while $Z$ is. The meaning of the plots is the following:

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>3.</td>
<td>4.</td>
</tr>
<tr>
<td>$X$ Phase space ($X,X'$)</td>
<td>Far field image ($Z',X'$)</td>
</tr>
<tr>
<td>1.</td>
<td>2.</td>
</tr>
<tr>
<td>Physical image ($X,Z$)</td>
<td>$Z$ Phase space ($Z',Z$)</td>
</tr>
</tbody>
</table>

where 1. is the beam cross section, 2. and 3. are the phase spaces, 4. is the image detected at large distances (far field). Notice also that the shared boundaries between the plots correspond to the same variables.

Next, we’ll run PLOTSPACE on the STAR.03 file from the TGM example. This will show us $X$ phase space, $Z$ phase space, the $X$-$Z$ plane, and the far field plot of $X'$ vs. $Z'$.

```
$ run util:plotspace
PREPLOT> Input file ? STAR.03
Read 1000 rays.
Each ray has 12 entries.
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays .
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
PREPLOT> Then ?
0
***********
Found 865 good points out of 1000
```
Figure 3.8: PLOTSPACE example one
Read 865 good rays. Plot all?
> y
0
$
Figure 3.9: PLOTSPACE of TGM final image
### PLOTXY

<table>
<thead>
<tr>
<th>Purpose:</th>
<th>General plotting utility used with SHADOW.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Input:</td>
<td>Unformatted file generated by SHADOW (BEGIN.DAT, STAR.xx, MIRR.xx, SCREEN.xxyy etc.)</td>
</tr>
<tr>
<td>Output:</td>
<td>Plot to screen and to file if desired.</td>
</tr>
</tbody>
</table>

#### Description:

PLOTXY is the basic graphics utility for SHADOW. It will plot any combination of “columns” from the BOF that SHADOW writes, i.e., BEGIN.DAT, MIRR.xx, STAR.xx, SCREEN.xxyy etc. The figure below shows all possible options and, where applicable, the additional information required.
(contour plot)

Number of bins in X

Number of bins in Y

Save TD command file?

Don't save command file

Write TD.FIL command file

Terminal type

0: VT240  1: TEK40XX  2: HIREZ  3: TEK4107

Plotting Begins
Examples:

We'll show the contour plot of the footprint on the first mirror of the two mirror example from the primer.

```
$ dir

Directory D13:[XRAYOP.DOCUMENT.UTIL.PLOTXY]

MIRR.01;2     282  5-MAY-1987 14:55:09.61 (RWED,RWED,RE,RE)

Total of 1 file, 282 blocks.
$

$ plotxy

PLOT> Input file? mirr.01 /* Use mirror BOF file */
PLOT> Options --- Enter
PLOT> 0 for excluding the losses
PLOT> 1 for including only the losses
PLOT> 2 for including all the rays.
PLOT> Then ? 0 /* no losses */
PLOT> Comment for plot [ 80 char ] ? /* a title, if you will */
First Mirror footprint

PLOT> File read OK. Full specifications:
    D13:[XRAYOP.DOCUMENT.UTIL.PLOTXY]MIRR.01;2
    Was created : 5-MAY-1987 14:55

PLOT> Found 1000 good points out of 1000
PLOT> The following columns are defined for each ray :
      1) the regular columns [1-12]
      2) optical path [13]
      3) phase angle of As [14], Ap [15], and the Ap vector [16-18]

Col Par Minimum: Maximum: Center: St. Dev.:

1  X  -12.123  12.010  0.65111E-01  5.7198
Options. You may plot any two rows from the above list versus each other. You may also plot any of them versus the ray numerical aperture. N.A. -- enter 20.

Rows to use for plot:
- for horizontal axis ? 1  /* X */
- for vertical axis  ? 2  /* Y */

Scaling options. Enter
- 0 For automatic scaling
- 1 For cartesian scaling
- 2 For external limits
Then ? 2  /* choose external limits */
Enter limits.
- Horizontal min.: -50
- Horizontal max.: 50
- Vertical min. : -150
- Vertical max. : 100

Plotting options:
- 0 For scattered plot
- 1 For connected plot
- 2 For contour plot
Then ? 2  /* contour plot */
Number of bins [Nx by Ny] to prepare the contours. /* odd number */
- Nx : 21
- Ny : 21

Number of contours : 11  /* number of lines or levels */
Degree of polynomial used for joining [0=default] : 0
Included reflectivity ? 0  /* don’t include reflectivity here */
Data normalization :
0  For no normalization
1  For normalized to 1
2  For normalized to total counts
PLOT> Then ? 2
   /* normalize to total counts */
Smoothing [ Y/N ] ? 1
   /* choose smoothing */
Save grid file [ Y/N ] ? 0
PLOT> Hairline at [ 0,0 ] ? 0
PLOT> Overlay a mirror/slit ? 0
PLOT> Ready for histograms. Enter:
   -1  to skip
   0  for same limits as plot
   1  3*stdev
   2  external
PLOT> ? 0
PLOT> Do you want to save the TD command file ? 1
   /* save the command file */

$ dir/since
   /* show the command file */

Directory D13:[XRAYOP.DOCUMENT.UTIL.PLOTXY]

TD.FIL;1 37  2-AUG-1989 10:09:14.99  (RWED,RWED,RE,RE)

Total of 1 file, 37 blocks.
$

The command file TD.FIL is plottable using TDTEK and UGTEK.
Figure 3.10: PLOTXY output
Purpose: Creates a two or three column formatted file from a binary output file. The user specifies which two or three columns of the BOF to use for output. The generated file is in a form suitable for input to a graphics routine like TopDrawer or an analysis program that requires formatted input like FMAX or FILEX. Also, a quick alternative to PLOTXY.

Input: Any Binary output file.
Output: A user specified file (use TT to direct output to terminal). The file is formatted and in single precision.

Description:
PREPLOT generates a two or three column formatted file from a BOF input file. Any of the standard 12 rows (13 or 18 if optical path or polarization is selected when source is defined) may be selected in addition to “rows” 20-23 which are defined during PREPLOT execution (see example). The output can then be used as input to a graphics, histogram, or analysis program that requires formatted input. For example, if one wanted to create a table of optical path length vs. angle from the optical axis at the 2nd image plane, then rows 21 and 13 would be selected from the file STAR.02. Note: output is in single precision.

Examples:
Let’s use the STAR.01 file from sec 2.4 of the primer (spherical mirror).

$ preplot
PREPLOT> Input file ?
star.01
Read 1000 rays.
Each ray has 12 entries.
PREPLOT> How many columns to write out ?
2
Row [1-12] : the individual column
Row [20] : R = SQRT(X**2 + Y**2 + Z**2)
Row [21] : angle from the Y-axis
Row [22] : the magnitude of A vector
Row [23] : A**2
PREPLOT> Row # 1 :
1
PREPLOT>  2 :
4
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays.
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
PREPLOT> Then ?
0
***********
Found 1000 good points out of 1000
Output options :
[ 0 ] store rays in a file
[ 1 ] plot directly on screen
[ 2 ] both
Then ?
0
PREPLOT> Output file ?
pre.dat

Let’s look at the first 15 lines of pre.dat

$ ty pre.dat
-0.2780961  1.8455550E-02
-0.2831062  1.8694960E-02
 0.3307338 -2.2033621E-02
-0.3311352  2.2213560E-02
-9.4173871E-02  6.2023830E-03
 0.2767766  -1.8385222E-02
 0.3668378  -2.4258027E-02
 0.1166269  -7.7635958E-03
-0.3735043  2.5044683E-02
-0.3660054  2.4130033E-02
PREPLOT

Figure 3.11: PREPLOT: Energy vs. Z for TGM exit slit

Dispersion of TGM on white (49.8 - 50.2eV) source

-9.1828540E-02 6.0481941E-03
0.3607364 -2.4109263E-02
0.3719847 -2.5014305E-02
-3.2931246E-02 2.2053795E-03
0.1509321 -1.0015748E-02
Purpose: Creates a two column formatted file from two binary output files. The user specifies which row of each ray file to use for output. With this utility it is possible to study the correlation between different optical systems, or between different sections of the same OS.

Input: Any two binary output files.

Output: A user-specified formatted two column file or TT.

Description:
This utility allows the user to create a two column file from two different ray files. Any one of the standard twelve to eighteen columns from the first ray file may be tabulated against any column from the second file. The user may then use this data to plot the correlation between various optical systems or between different elements of the same system. For example, the distance off axis in the x-direction at a mirror may be correlated with the optical path length at, say, a image plane further along in the system.

Examples:
One example of PREPLOT2 would be to use it to study the correlation between footprint width on the grating of a TGM and the dispersion at the exit slit. Using our TGM setup at an energy of 59 eV, we run PREPLOT2 using column 1 at the grating (X) and column 3 at the exit slit (Z).

$ dir

Directory CXRL$DUA0:[SHADOW.ZDOCUMENT.UTIL.PREPLOT2]

MIRR.02;3  282 28-DEC-1989 10:23:10.19  (RWED,RWED,RE,) <<<
STAR.02;3  282 28-DEC-1989 10:23:13.49  (RWED,RWED,RE,) <<<
SYNCH.00;126  7 28-DEC-1989 10:19:23.61  (RWED,RWED,RE,)
SYSTEMFILE.DAT;2  1 28-DEC-1989 10:22:48.04  (RWED,RWED,RE,)
TGMEG.01;47  12 28-DEC-1989 10:22:46.61  (RWED,RWED,RE,)
TGMEG.02;47  12 28-DEC-1989 10:22:47.11  (RWED,RWED,RE,)
TGMEG.03;48  12 28-DEC-1989 10:22:47.67  (RWED,RWED,RE,)

Total of 7 files, 608 blocks.
$ preplot2
All the losses will be determined by the SECOND file.
First file to be prepared?
mirr.02 /* grating footprint file */
Second file to be prepared?
star.02 /* exit slit (image) file */
Columns to write out from file
mirr.02
1 /* X -- width of footprint */
Columns to write out from file
star.02
3 /* Z -- height of image at slit */
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays.
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
? 2 /* plot all rays to show full dispersion */
***********
Found 876 good points out of 1000
Output options:
[ 0 ] store rays in a file
[ 1 ] plot directly on screen
[ 2 ] both
Then ? 0
Output file? aberr
$

The output is shown in figure 3.12.
TGM Aberrations

Figure 3.12: PREPLOT2 output
Purpose: Creates a plottable three dimensional file of a mirror surface. It can be used to plot out the “real” shape of a mirror.
Input: A Binary Output File (for example MIRR.xx).
Output: A three dimensional plottable file to be used as input to TDFILE.

Description:

SURPLOT takes a binary output file (BOF) and creates a plottable file that can be put into TopDrawer to create a “real” picture of the mirror surface. It must be used with a rectangular, GRID/GRID source, with the grid in X and Z, and only a 1 x 1 grid in phase space. The source should be about the size of the surface you wish to plot, and should have no depth and zero divergence. This assures that the rays are parallel when they hit the mirror’s surface. The system must consist of only the optical element you wish to plot because after the first element the rays will no longer be parallel.

Examples:

We’ll show two examples of mirror surfaces, one elliptical and one toroidal. Below is the output from SOURCINFO on the source used, as well as the MIRINFO for each example.

+++
*********** SOURCE DESCRIPTION ***********
GRID/GRID Source
Example of source used for SURPLOT
+++
Input file specified:
end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.SURPLOT]END.00;2
Creation Date :13-APR-1989 13:48
+++
Grid Source.
Generated total 225 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: RECTANGULAR
Source Width: 8.00000000 and Height: 18.0000000
Source Emission Characteristics
Distribution Type: UNIFORM
Distribution Limits. +X : 0.000000000E+00 -X: 0.000000000E+00 rad
+Z : 0.000000000E+00 -Z: 0.000000000E+00 rad

Elliptical Mirror SURPLOT Example
Elliptical mirror with shape exaggerated.
Input file specified: end.01
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.SURPLOT]END.01;6
Creation Date :14-APR-1989 15:40

Surface figure was defined as: ELLIPTICAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity OFF
Mirror dimensions (elliptical):
  Major Axis Minor axis
  -9.0000 -6.0000

Central Axis parameters:
Source Plane Distance 100.0000000000000
Image Plane 100.0000000000000
Incidence Angle 0.000000000000000E+00
Reflection/Diffractive Angle 0.000000000000000E+00

Mirror parameters EXTERNAL
Parameters used follow:
  Semi-major axis 50.0000000000000
  Semi-minor axis 20.0000000000000
  Semi-focal-length 0.000000000000000E+00
Eccentricity 0.0000000000000000E+00
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Now we'll show a log of the run through SURPLOT and the output after running it through TopDrawer.

$ surplot
SURPLOT> File ?
mirr.01
Mesh size must be Nx by Ny.
>> Nx ?
15
>> Ny ?
15
READ 225 POINTS.
$ dir/since
Directory D13:[XRAYOP.DOCUMENT.UTIL.SURPLOT]
SURPLOT.DAT;7 66 11-APR-1989 14:44 (RWED,RWED,RE,RE)
Total of 1 files, 66 blocks.
$ type sur.com
SET THREE

SET ORDER X Y Z
SET LIMITS X -7 7 y -7 7 z -7 7
SET AXIS OFF
@SURPLOT.DAT
END
The plot can be generated using TDTEK and UGTEK with the use of a simple command file in TDTEK, shown above. For best results, the size of the source should not exceed the size of the mirror. The limits in the command file may have to be adjusted slightly to fit the entire mirror.

+++++ MIRROR DESCRIPTION ********

Toroidal Mirror
Second example of SURPLOT
+++++ Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.SURPLOT]END.01;3
Creation Date :13-APR-1989 13:48
+++++

Surface figure was defined as: TOROIDAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity OFF
Mirror dimensions ( elliptical ) :
   Major Axis Minor axis
     -9.0000  -6.0000
+++++

Central Axis parameters :
Source Plane Distance  100.0000000000000
Image Plane  100.0000000000000
Incidence Angle  0.0000000000000000E+00
Reflection/Diffraction Angle  0.0000000000000000E+00

Mirror parameters EXTERNAL
Parameters used follow:
   Major Radius  15.0000000000000
   Minor Radius  5.0000000000000
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES
+++++
Figure 3.13: SURPLOT: Elliptical Mirror

Elliptical Mirror Surface
A better illustration may be the toroidal grating from the TGM. It is shown below, having been traced with a grid source and run through SURPLOT. The shape is exaggerated due to the scaling, X -3 to 3, Y -50 to 50 and Z 0 to 1. The user can also add extra lines to the plot, to better define the mirror shape.
Figure 3.14: SURPLOT: Toroidal Mirror Surface
Grating Surface of 6m TGM

Figure 3.15: SURPLOT: TGM Grating
Purpose: Creates a plottable file of all the rays in an optical system in the laboratory reference frame. Either a two or three dimensional plot can be made.

Input: All of the following are required for input to SYSPLOT:

<table>
<thead>
<tr>
<th>File</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optical Axis File</td>
<td>OPTAX.nn</td>
</tr>
<tr>
<td>Source Binary Output File</td>
<td>BEGIN.DAT</td>
</tr>
<tr>
<td>Final Image Binary Output File</td>
<td>STAR.nn</td>
</tr>
<tr>
<td></td>
<td>(There are nn OEs in the OS)</td>
</tr>
<tr>
<td>All OE Binary Output Files</td>
<td>MIRR.01-MIRR.nn</td>
</tr>
</tbody>
</table>

Output: A plottable file SYSPLOT.DAT that can be used as input to TopDrawer.

Description:

SYSPLOT will create a plottable file that shows the path of each ray with respect to the physical optical system. Two dimensional projections of $X$ vs. $Y$ or $Z$ vs. $Y$ can be implemented by selecting columns 2,1 or 2,3 respectively when prompted for horizontal and vertical columns. The output file SYSPLOT.DAT should be examined manually for extrema which will be used to set limits on the plot (see examples below). The output from TDTEK is then used as input to the routine UGTEK to generate the plot.

Examples:

We’ll run SYSPLOT on the two mirror example from section 6.1 of the primer. The output from SYSINFO is shown first so one can compare the system to the plot. Ideally one would use a grid/grid source to generate the info for SYSPLOT since the rays would be regularly spaced. We won’t do that here to be consistent with data in the Primer.

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
************** SYSTEM DESCRIPTION **************
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
System shown by SYSPLOT

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified:TT:
Optical Element # 1  System Number:

MIRROR  ELLIPTICAL  UNLIMITED  COMPUTED  REFLEC. OFF

Orientation  0.000000000000000E+00  deg.
Source Plane  2000.000000000000
Incidence Ang.  88.00000000000000  deg.
Reflection Ang.  88.00000000000000  deg.
Image Plane  500.00000000000000

Optical Element # 2  System Number:

MIRROR  ELLIPTICAL  UNLIMITED  COMPUTED  REFLEC. OFF

Orientation  180.00000000000000  deg.
Source Plane  1000.000000000000
Incidence Ang.  88.00000000000000  deg.
Reflection Ang.  88.00000000000000  deg.
Image Plane  500.00000000000000

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

OPT. Elem #  X =  Y =  Z =
0  0.0000000000000E+00  0.0000000000000E+00  0.0000000000000E+00
1  0.0000000000000E+00  2000.000000000  0.0000000000000E+00
1′  0.0000000000000E+00  2498.7820251  34.878236872

***************  END  ***************
$ sysplot
File [ 0 ] or tt [ 1 ] ?
1
Bidimensional [ 0 ] or 3d plot [ 1 ] ?
1
How many mirrors ?
2
How many rays [ suggest no more than 200 (500 max) ] ?
20
Optaxis file ?
optax.02
Source file and (final) image file ?
begin.dat
star.02
Mirror N. 1 ?
mirr.01
Mirror N. 2 ?
mirr.02
Plot all the losses too [ Y/N ] ?
y
$ dir/since

Directory D13:[XRAYOP.DOCUMENT.UTIL.SYSPLOT]

SYSPLOT.DAT;5 8 11-APR-1989 13:29 (RWED,RWED,RE,)
SYSPLOT.STR;2 1 11-APR-1989 13:29 (RWED,RWED,RE,)

Total of 2 files, 9 blocks.

SYSPLOT creates the two files SYSPLOT.DAT and SYSPLOT.STR. SYSPLOT.DAT can now be plotted using TopDrawer. The limits were chosen by typing the file SYSPLOT.DAT and looking for the extrema. The simplest way to plot the system is to create a command file like the following:

```
set three
set order x y z
```
To generate a plot on your terminal, use the utilities TDTEK and UGTEK. This is done by first running TDTEK as follows:

```
$ TDTEK
TOP DRAWER (10/10/80) NOW EXECUTING

@SYSPLOT.COM

<ctrl>O (suppresses output)
```

where SYSPLOT.COM is the command file shown above. Then you must put your terminal in graphics mode to run UGTEK on the file created by TDTEK called SAVE.DAT. This is shown in the following:

```
$UGTEK
UG element file: SAVE.DAT
Plot Which Pictures (default=all): <RET>
Enter Picture Scale Factor (default=1.0): <RET>
```
Two Mirror System in 3-D

Figure 3.16: SYSPLoT: 3-D system
Figure 3.17: SYSPLOT: Top view
Figure 3.18: SYSPLLOT: Side view
Purpose: To generate a connected X-Y plot from a grid RAY file.

Input: A grid RAY file that has been processed by TRANSLATE and COL_2, or any other two column file.

Output: A plottable file suitable for TopDrawer output.

Description:

This utility will create a TopDrawer file, that when plotted will be a connected plot. It requires as input a two column formatted file. If one wishes to plot a ray file from SHADOW, it must be a grid file and it must be run through PREPLOT before being suitable for TDCONN. The maximum number of data points it can handle is 5000. The program assumes that the M\{x,y\} records describes a grid, \{N_x,N_y\}. Thus a first pass is made by joining on \{x\}, then a second by joining on the other variable. In this way a connected plot is created. Very useful to show for example image distortions.

Examples:

To illustrate this utility, we will use a grid source with conical angle distribution traced through a toroidal mirror. This is the same system used in the SOURCE section of the User's Guide under Conical Angle Distribution. Here we show how to obtain the plot using PREPLOT and TDCONN instead of PLOTXY. The source and mirror are described, then we will show a log of how to get the results, and finally the plot.

+---------------------------------------------------------------------------+  
| TDCONN Example                                                            |
| Conical Source Angle Distribution-Grid source                              |
| +---------------------------------------------------------------------------+  
| Input file specified:                                                     |
| end.00                                                                     |
| Full file Specification :D13:[XRAYOP.DOCUMENTSOURCE.CONICSAD]END.00;2      |
| Creation Date :27-MAR-1989 16:35                                           |
| Grid Source.                                                              |
| Generated total 160 rays.                                                 |
| Source assumed BIDIMENSIONAL (flat).                                      |
| Source Spatial Characteristics: POINT                                       |  

Source Emission Characteristics
Distribution Type: CONICAL
Cone Outer Aperture : 0.500000000E-02 Inner Aperture : 0.000000000E+00

Source of this O.E. moved NO
Mirror at pole position ( no mov. )  YES

$ preplot
PREPLOT> Input file ? star.01  \* Takes BOF and writes text file *
Read 160 rays.  \* Maximum number of points allowed
Each ray has 12 entries.  \* by TDCONN is 5000.
PREPLOT> How many columns to write out ? 2
Row [1-12] : the individual column
Row [20] : \( R = \sqrt{X^2 + Y^2 + Z^2} \)
Row [21] : angle from the Y-axis
Row [22] : the magnitude of A vector
Row [23] : \( A^2 \)
PREPLOT> Row # 1 : 1  \* Here we show the x phase space *
PREPLOT> 2 : 4
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays .
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
PREPLOT> Then ? 0
***********
Found 160 good points out of 160
Output options :
[ 0 ] store rays in a file
[ 1 ] plot directly on screen
[ 2 ] both
Then ? 0
PREPLOT> Output file ? conic14.dat
$ r nutil:tdconn
TDCONN> Input file : conic14.dat  \* Read text file to create
Read 160 from conic14.dat  \* plottable file. *

TDCONN> Grid size [ Nx by Ny ].  \* Nx times Ny must equal
TDCONN> Nx : 10
TDCONN> Ny : 16

/* N x times Ny must equal the total number of points */
Figure 3.19: TDCONN Example – X phase space plot

Xmax = 9.9340184E-03 Xin = -9.9340184E-03
Ymax = 4.9745785E-03 Ymin = -4.9745785E-03
TDCONN> Limits to use on the plot:
TDCONN> Xmin : -1.5e-02
TDCONN> Xmax : 1.5e-02
TDCONN> Ymin : -1e-02
TDCONN> Ymax : 1e-02
TDCONN> Output file : conic14.td

/* Plottable directly using
$ */
/* TD_NEW */
Purpose: To determine the distribution of the values in a given column of a ray file. The data are binned in a histogram with conditions specified by the user.

Input: Any BOF (e.g. BEGIN.DAT, STAR.xx).

Output: A plottable file named histo1.dat or TT: or both.

Description:

This utility will create a one-dimensional histogram from any column of a BOF. The extrema are given to the user, who must decide what the mean (center) and width of the histogram should be. Options are included to ignore the lost rays, to include reflectivity, and to direct output to the terminal or a file. If the output is in a file, the histogram may be plotted using the graphics routine TopDrawer as in the following example:

TDTEK
@ histo1.dat // <CTRL> O may be toggled to supress output
histo // The histogram is now plotted
end

Examples:

Here we will examine the system set up in the primer section 6.2. We have a rectangular source that models an entrance slit with very narrow vertical divergence. The optical element is a spherical diffraction grating with the slits placed on the Rowland circle. We have an energy range from 198 to 202 eV, and will examine how the exit slit determines the output spectrum. For this, we will need to know the photon energy distribution, i.e., the histogram of the number of rays in function of the photon energy. The source and system descriptions follow.

+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++Source Incident upon Rowland Circle Mount GratingHISTO1 Example+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++Input file specified:
end.00
Full file Specification :DISK4:[RUNKLE.UG.UTIL.HISTO1]END.00;3
Creation Date :25-APR-1989 17:44
++-------------------------------
Random Source.
Generated total 5000 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: RECTANGULAR
Source Width: 2.00000000 and Height: 0.200000000E-01
++-------------------------------
Source Emission Characteristics
Distribution Type: UNIFORM
Distribution Limits. +X : 0.500000000E-02 -X: 0.500000000E-02 rad
+Z : 0.100000000E-01 -Z: 0.100000000E-01 rad
++-------------------------------
Source Photon Energy Distribution: BOX DISTR
From Photon Energy 198.000000 or 62.6187879 Angs.
To Photon Energy 202.000000 or 61.3788119 Angs.
++-------------------------------
++-------------------------------
******************** MIRROR DESCRIPTION ********************
++-------------------------------
Rowland Circle Mount Diffraction Grating
HISTO1 Example
++-------------------------------
Input file specified: end.01
Full file Specification :DISK4:[RUNKLE.UG.UTIL.HISTO1]END.01;7
Creation Date :25-APR-1989 17:55
++-------------------------------
Surface figure was defined as: SPHERICAL
Cylindrical figure NO
Element type REFLECTOR
Element type GRATING
Order choosen ( inside are < 0 ) -1.000000000000000
Automatic Tuning NO
Constant ruling [ lines/cm ] 12000.00000000000
We have added an exit slit with dimensions 10 mm X 3.5 mm at the continuation plane. We will create a histogram of the photon energy of the rays passing through the slit. The spectrum is dependent on the slit width, the divergence in the vertical, and on aberrations in the spherical mirror.

```bash
$Hist01
File for analysis ? star.01
Column to analyze ? 11
Options : [0] Angstroms
          [1] eV
          [2] cm⁻¹
<??> 1
Read : 5000 rays
Maximum : 201.9990
Minimum : 198.0021
Distribution center ? 200
   width ? 1
Number of bins (odd, please) ?
   25
Flag checks. Enter :
0 to exclude lost rays
1 to include lost rays too
```
2 to use only lost rays
<?> 0
Normalization kind. Enter:
0 for no normalization
1 to normalize to 1
2 area normalized to 1
<?>
0
Include reflectivity? 0

Output options
[ 0 ] store histogram in a file
[ 1 ] plot histogram on screen
[ 2 ] both
Then? 0
Another run [ Y/N ]? 0
Return to DCL
$

The output file can be used as input to TopDrawer by using the following commands in TDTEK:

@Hist01.dat
Histo
End

The results are shown below.

Often the histogram assumes a shape close to that of a gaussian curve. This could be the case with the energy profile of a monochromator, or the distribution of rays across an image plane. We will plot the distribution in the vertical direction of rays in the final image plane for the TGM system. The procedure will be to obtain the mean and variance of the vertical distribution using VARIANCE on a PREPLOT file that has been generated from the final image ray file (STAR.03). The same data can be obtained from PLOTXY. These statistics will be used to generate a gaussian curve with the utility TGAUSS. Then HIST01 will be run on the STAR.03 file using the mean and interval that the gaussian curve was generated with. Finally, both the histogram and superimposed gaussian will be plotted with TopDrawer.
Figure 3.20: HISTO1 example
$ preplot
PREPLOT> Input file ?
TGM:star.03
Read 1000 rays.
Each ray has 12 entries.
PREPLOT> How many columns to write out ?
2
Row [1-12] : the individual column
Row [20] : R = SQRT(X**2 + Y**2 + Z**2)
Row [21] : angle from the Y-axis
Row [22] : the magnitude of A vector
Row [23] : A**2
PREPLOT> Row # 1 :
1
PREPLOT> 2 :
3
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays.
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
PREPLOT> Then ?
0
***********
Found 927 good points out of 1000
Output options :
[ 0 ] store rays in a file
[ 1 ] plot directly on screen
[ 2 ] both
Then ?
0
PREPLOT> Output file ?
pre.dat

$ variance
Log file ?
var.dat
File for analysis? 
pre.dat
Read 927 from pre.dat
Column to analyze? 2

Results from column 2
Mean value: -1.0513511961653615E-02
Variance: 1.4434200420233979E-04
Standard deviation: 1.2014241723984905E-02

Enter:
0 to exit
1 new file
2 new column (same file)
3 best fit (plottable)
?
0

FILEX must be used on the TGAUSS data file to add the mean value to the data, otherwise the curve would be centered on zero. It is the spherical aberration (notice tail) that gives an apparent shift.

$ filex
FILEX>
TG = TG 
FILEX> read 50 records.
FILEX> Successful completion
FILEX>
TG = TG - %1.037e-2
FILEX> read 50 records.
FILEX> Successful completion
FILEX>
TG = TG 
FILEX> read 50 records.
FILEX> Successful completion
FILEX>
FILEX>
$ histo1
File for analysis ?
TGM:star.03
Column to analyze ?
3
Read : 1000 rays
Maximum : 1.0586338E-02
Minimum : -7.2909199E-02
Distribution center ?
-1.037e-2
    width ?
.08
Number of bins (odd, please) ?
25
Flag checks. Enter :
0 to exclude lost rays
1 to include lost rays too
2 to use only lost rays
< ?>
0
Normalization kind. Enter :
0 for no normalization
1 to normalize to 1
2 area normalized to 1
< ?>
1
Include reflectivity ?
0
Output options :
[ 0 ] store histogram in a file
[ 1 ] plot histogram on screen
[ 2 ] both
Then ?
0
Another run [ Y/N ] ?
0
Return to DCL
Figure 3.21: HISTO1: Histogram with superimposed gaussian curve

The following TopDrawer commands will superimpose the TGAUSS curve onto the histogram:

@histo1.dat
histo.dat
@TG.dat
join 1
end
HISTO2

Purpose: To create a two dimensional histogram from any two of the twelve to eighteen columns of a ray file.

Input: Any BOF.

Output: A formatted plottable file named MESH.DAT which can be used as input to a graphics routine like TopDrawer.

Description:

This utility enables the user to create a two dimensional histogram from any two columns of a ray file. The input file should first be run through the utility MINMAX to find the extrema (only for the first 6 columns) since HISTO2 asks for minimum and maximum values for each dimension. Like HISTO1, options for treating lost rays, reflectivity, and normalization are included. The output file MESH.PLT can be plotted using TopDrawer:

    TDTEK
    MESH.PLT  // <CTRL> O may be used to supress output
    histo     // histogram is plotted here
    end       // Exits TopDrawer

We do not have yet a good 3-d plotting utility, so the user will have to use the (limited) resources of TopDrawer or find another package. The NCAR package has powerful contouring capabilities and requires a different file format.

Examples:

Here will make two histograms using the TGM system. We will look at the ray distributions on the first mirror, and at the final image plane. Since HISTO2 asks for the limits, we will run MINMAX on each ray file to determine the appropriate intervals. The output is then used in TopDrawer. Note how the first histogram fills up the space of the mirror, which has a rectangular shape (4 X 9.5). The Final image also has a shape similar to that of the exit slit.

    $ minmax
    Input file ?
    tgm:mirr.01
    Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ?
0
Comment ?
TGM example - first mirror footprint
***********
File examined is :
D13:[XRAYOP.DOCUMENT.TGM]MIRR.01;36
Was created :
15-MAY-1989 07:32
***********
Found 973 good points out of 1000
Here we are.
X max is 1.993178
X min is -1.999294
Y max is 4.180661
Y min is -4.735539
Z max is 0.1313010
Z min is 5.4842703E-06
X prime max is 1.1284630E-02
X prime min is -1.1375175E-02
Y prime max is 0.9963850
Y prime min is 0.9960120
Z prime max is 8.9194998E-02
Z prime min is 8.4335230E-02

$ histo2
File for analysis ?
tgm:mirr.01
Data ready. Read 1000 points.
Columns for x-axis and y-axis?
1
2
X min, X max and number of bins :
-5
5
Y min, Y max and number of bins :
-5
5
25
Do you want reflectivity too (Y=1,N=0) ?
0
Data normalization. Enter:
0 for no normalization
1 to normalize to 1
2 to normalize to total counts
1
There were 27 lost out of 1000 rays.

$ minmax
Input file ?
tgm:star.03
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ?
0
Comment ?
TGM example - Final image
*************
File examined is :
D13:[XRAYOP.DOCUMENT.TGM]STAR.03;36
Was created :
15-MAY-1989 07:32
*************
Found 927 good points out of 1000
Here we are.
X max is 9.4501123E-02
$ histo2
File for analysis ?
tgm:star.03
Data ready. Read 1000 points.
Columns for x-axis and y-axis?
1
3
X min, X max and number of bins :
-.15
.15
25
Y min, Y max and number of bins :
-.15
.15
25
Do you want reflectivity too (Y=1,N=0) ?
0
Data normalization. Enter:
0for no normalization
1to normalize to 1
2to normalize to total counts
1
There were 73 lost out of 1000 rays.
Computations done.
File for TOP DRAWER [ 0 ] or NCAR [ 1 ] ?
All done.

The output files from HISTO2 are named MESH.PLT. These files can be plotted by issuing the following commands to TopDrawer:

@MESH.PLT  (or a renamed file)
HISTO
END
Photon counts on first mirror of TGM

Figure 3.22: HISTO2: First mirror
Histogram of final image plane of TGM

Figure 3.23: HISTO2: Final image
**INTENS**

**Purpose:** Computes the power reflected, absorbed or transmitted ray by ray.

**Input:** One or two RAY files such as ST AR.xx, MIRR.xx, SCREEN.xxyy

**Output:** To a user-specified file of \( \{x, y\} \) or \( \{x, y, z(x, y)\} \) format.

**Description:**

The program works by computing the total energy carried by a RAY file after a reflection by a mirror or transmission by a screen. This can be compared with that of the incoming beam and the local reflectivity or absorbed power can then be computed. The program creates a 2 or 3 columns file of the type \( \{x, y, z(x, y)\} \) where \( (x, y) \) can be any two columns and \( z \) is the power. If the rescaling option is selected, then the beam is considered to be a sample of the source (user-specified) power. The output can be plotted out using TopDrawer or further analyzed.

**Examples:**

For the purpose of demonstration, we use an elliptical mirror to focus a point source. Since the source covers an energy range of 500-2000 eV, the reflectivity of the mirror (Au-coated) will be a function of both the photon energy and the angle. For the narrow divergence (20 mrad) we used, the reflected power shows up mainly depending on the energy. The way the total power is subdivided among rays depends on how the source was generated. In general, SHADOW assumes that the photons are generated by sampling \( N(h, \omega) \), unless otherwise specified (as in the SR source).

\[
W = \sum_{i=1}^{N} n_i \hbar \omega_i
\]

In the second case, the photons are generated by sampling \( P(h, \omega) \) so that each ray carries the same amount of power:

\[
W = \sum_{i=1}^{N} W_i
\]

To make it more realistic, we assign 100 Watt to the 5000 rays in BEGIN.DAT; that translates into 0.02 Watt per ray, and it is easy to see that the reflected and absorbed power of each ray do add up to 0.02 Watt. Has this not been done, then INTENS simply assumes the intensity of each ray is 1 at the source. The user has the option of writing out the intensity of each ray versus one or two other columns, thus making a bi- or tri-dimensional plot. This is very similar to PREPLOT.
$type source.fil  

Example file  
Source  

Input file specified:
start.00  
Full file Specification :D13:[XRAYOP.DOCUMENT.INTENS]START.00;1  
Creation Date :25-FEB-1989 17:09  
Random Source.  
Generated total 5000 rays.  
Source assumed BIDIMENSIONAL (flat).  
Source Spatial Characteristics: POINT  
Source Emission Characteristics  
Distribution Type: UNIFORM  
Distribution Limits. +X : 0.100000000E-02 -X: 0.100000000E-02 rad  
+Z : 0.100000000E-01 -Z: 0.100000000E-01 rad  
Source Photon Energy Distribution: BOX DISTR.  
From Photon Energy: 500.000000 or 24.7970400 Angs.  
Angular difference in phase is 0.00000E+00  
Degree of polarization is 0.00000E+00  
Source points have COHERENT phase.  

$  
$  
$type mirror.fil  

$
Input file specified: end.01
Full file Specification :D13:[XRAYOP.DOCUMENT.INTENS]END.01;2
Creation Date : 25-FEB-1989 17:21

Surface figure was defined as: ELLIPTICAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity ON coefficients from: AUF1F2
Polarization dependence YES
Mirror dimensions UNLIMITED

Central Axis parameters:
Source Plane Distance 1000.000000000000
Image Plane 1000.000000000000
Incidence Angle 84.00000000000001
Reflection/Diffraction Angle 84.00000000000001

Mirror parameters COMPUTED
Same configuration as Central Axis YES
Objective focus at 1000.000000000000
Image focus at 1000.000000000000
Incidence angle 84.00000000000001
Parameters used follow:
  Semi-major axis 1000.000000000000
  Semi-minor axis 104.5284632676535
  Semi-focal-length 0.0000000000000000E+00
  Eccentricity 0.0000000000000000E+00
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

$ type system.fil

$
Input file specified:

# Optical Element: Creation Time:
1  D13:[XRAYOP.DOCUMENT.UTIL.INTENS]END.01;2  25-FEB-1989 17:21

Optical Element #  1  System Number:
MIRROR ELLIPTICAL UNLIMITED COMPUTED REFLEC. ON

Orientation  0.000000000000000E+00 deg.
Source Plane  1000.000000000000
Incidence Ang.  84.00000000000001 deg.
Reflection Ang.  84.00000000000001 deg.
Image Plane  1000.000000000000

----------------

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

OPT. Elem #  X =  Y =  Z =
0  0.00000000000E+00  0.00000000000E+00  0.00000000000E+00
1  0.00000000000E+00  1000.0000000  0.00000000000E+00
1'  0.00000000000E+00  1978.1476007  207.91169082

END

$ dir
Directory D13:[XRAYOP.DOCUMENT.INTENS]
INTENS

File for Intensity calculations ? MIRR.01
Read 5000 rays.
Each ray has 18 entries.
Total intensity is: 118.2835503036797

Normalized intensity is: 2.3656710060735936E-02

Options:
Intensity transmitted/reflected ........... 0
Intensity absorbed ........................ 1
Local reflectivity/transmission .......... 2

Spectrum type:
[0] # photons/sec.
[1] Watt

Normalize to source power [ Y/N ] ? 1
Total source power emitted ? 100
File-name for [x,y] file ? FIRST
Bi [ 0 ] or Tri-dimensional plot [ 1 ] ? 0

Total of 17 files, 4293 blocks.

$ INTENS

/* We perform now several runs of INTENS to demonstrate all the features. */

$
Row for plot x-axis ? 11 /* Reflected power vs. hv */
Enter:
0 cm-1
1 eV
2 angst
Then ? 1
Another run [1/0] ? 1 /* End of first plot */
File for Intensity calculations ? MIRR.01
Read 5000rays.
Each ray has 18 entries.
Total intensity is : 118.2835503036797
Normalized intensity is : 2.3656710060735936E-02
Options:
Intensity transmitted/reflected ........ 0 /* Same as above, except the
Intensity absorbed .................... 1 absorbed power is used. */
Local reflectivity/transmission ........ 2
Then ? 1
Spectrum type :
[0] # photons/sec.
[1] Watt
Then ? 1
Normalize to source power [ Y/N ] ? 1
Total source power emitted ? 100
Input I0 file : BEGIN.DAT
File-name for [x,y] file ? SECOND
Bi [ 0 ] or Tri-dimensional plot [ 1 ] ? 0
Row for plot x-axis ? 11
Enter:
0 cm-1
1 eV
2 angst
Then ? 1
Another run [1/0] ? 1 /* End of second run */
File for Intensity calculations ? MIRR.01 /* this time we use a different
Read 1000rays. /* example. Here we have a
Each ray has 18 entries. /* single energy source at
Total intensity is : 707.5300660592716 /* 2000 eV incident on a
Normalized intensity is : 0.7075300660592716 /* spherical mirror at 89 deg.
Options:
Intensity transmitted/reflected ........ 0
Intensity absorbed ..................... 1
Local reflectivity/transmission .... 2
Then ? 1
Spectrum type :
   [0] # photons/sec.
   [1] Watt
Then ? 1
Normalize to source power [ Y/N ] ? 1
Total source power emitted ? 100
Input I0 file : BEGIN.DAT
File-name for [x,y] file ? ABSORB
Bi [ 0 ] or Tri-dimensional plot [ 1 ] ? 0
Row for plot x-axis ? 2
Another run [1/0] ? 0
$ 
$ 
$ dir/since=-00:05

Directory D13:[XRAYOP.DOCUMENT.INTENS]

FIRST.DAT;1 313 25-FEB-1989 17:28 (RE,RWED,RWED,RE)
SECOND.DAT;1 313 25-FEB-1989 17:29 (RE,RWED,RWED,RE)

Total of 2 files, 616 blocks.

These files are plotted in Figures 3.24 and 3.25

$ 
$ type first.dat /* Units are eV and Watt. */
  916.4160  4.8564727E-04
  941.6163  4.3134825E-04
  1500.448  5.0780447E-05
  1441.797  6.1421124E-05
  1961.732  1.2236715E-05
  1883.999  1.5970762E-05
^Y

$ type second.dat
  916.4160  1.9514352E-02
  941.6163  1.9568652E-02
The last data file (ABSORB.DAT) is plotted in figure 3.26. Following that we ran H2KOL on the data file to generate a histogram. Note that the total power is divided among the bins, so the vertical scale is Watts/bin.
Figure 3.25: INTENS: Absorbed power vs. energy (second.dat)
Figure 3.26: INTENS: Absorbed power vs. mirror coordinate (Z) (absorb.dat)
Figure 3.27: INTENS: H2KOL output from ABSORB.DAT
Purpose: To find the extrema of the phase space occupied by a beam described by a RAY file. In other words, finds the extrema of each coordinate \((X,Y,Z)\) and direction cosines \((X',Y',Z')\).

Input: Binary output file (e.g. STAR.xx)

Output: TT and formatted file MINMAX.DAT

Description:
Given a ray file it will find the absolute minima and maxima for the three coordinates in position and momentum space. Options to include the lost rays in the analysis are provided. A comment is requested so the output file will have a title. The user may wish to rename this file since all output files will be named MINMAX.DAT. Very useful for quick verification of results.

Examples:
The first example will be using the spherical mirror example from the primer. After that we’ll run MINMAX on the STAR.03 and the MIRR.01 files from the TGM example.

```
$ minmax
Input file ?
star.01
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ?
0
Comment ?
Simple Spherical Mirror (Primer sec 2.4)
***********
File examined is :
D13:[XRAYOP.PRIMER.CHAPTER2]STAR.01;1
Was created :
4-MAY-1987 16:37
***********
```
Found 1000 good points out of 1000
Here we are.
X max is 0.4081510
X min is -0.4102450
Y max is 1.8064706E-15
Y min is -1.6632746E-15
Z max is 9.7605148E-03
Z min is 8.7015678E-07
X prime max is 2.7311902E-02
X prime min is -2.7292412E-02
Y prime max is 0.9999999
Y prime min is 0.9996073
Z prime max is 7.4768672E-03
Z prime min is -7.6602534E-03

First we’ll show the log of the run, then we’ll type the output file to show it’s contents.

$ minmax
Input file ? star.03
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ? 0
Comment ?
Image plane after tgm refocus mirror
***********
File examined is :
D13:[XRAYOP.DOCUMENT.UTIL.MINMAX]STAR.03;25
Was created :
4-MAY-1989 10:40
***********
Found 865 good points out of 1000
Here we are.
X max is 0.1150454
X min is -0.1366614
Y max is 2.7868333E-15

First we’ll show the log of the run, then we’ll type the output file to show it’s contents.

$ minmax
Input file ? star.03
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ? 0
Comment ?
Image plane after tgm refocus mirror
***********
File examined is :
D13:[XRAYOP.DOCUMENT.UTIL.MINMAX]STAR.03;25
Was created :
4-MAY-1989 10:40
***********
Found 865 good points out of 1000
Here we are.
X max is 0.1150454
X min is -0.1366614
Y max is 2.7868333E-15
Y min is -4.3796347E-15
Z max is 1.1263924E-02
Z min is -5.2220654E-02
X prime max is 2.3804422E-02
X prime min is -2.4356563E-02
Y prime max is 0.9999999
Y prime min is 0.9996949
Z prime max is 7.3660947E-03
Z prime min is -1.0705367E-02
$
$
$ type minmax.star
Searching file:
D13:[XRAYOP.DOCUMENT.UTIL.MINMAX]STAR.03;25
4-MAY-1989 10:40

Image plane after tgm refocus mirror

Total rays : 1000
Lost rays : 135
EXCLUDING LOSSES
X max is 0.115045405924
X min is -0.136661350727
Y max is 0.278683326416E-14
Y min is -0.437963467576E-14
Z max is 0.112639237195E-01
Z min is -0.522206537426E-01
X prime max is 0.238044224679E-01
X prime min is -0.243565626442E-01
Y prime max is 0.9999999880791
Y prime min is 0.999694883823
Z prime max is 0.736609473825E-02
Z prime min is -0.107053667307E-01
$
$
$ minmax
Input file ? mirr.01
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ? 0
Comment ?
First mirror of tgm

File examined is:
D13:[XRAYOP.DOCUMENT.UTIL.MINMAX]MIRR.01;25
Was created:
  4-MAY-1989 10:40

***********
Found 927 good points out of 1000
Here we are.
X max is  1.998689
X min is  -1.995066
Y max is   4.030588
Y min is  -4.720492
Z max is  0.1304751
Z min is  5.9245693E-05
X prime max is 1.1206643E-02
X prime min is -1.1470584E-02
Y prime max is  0.9963953
Y prime min is  0.9960193
Z prime max is  8.9137740E-02
Z prime min is  8.4174052E-02
$
$ type minmax.mirr
Searching file:
D13:[XRAYOP.DOCUMENT.UTIL.MINMAX]MIRR.01;25
  4-MAY-1989 10:40

First mirror of tgm

Total rays :     1000
Lost rays :       73
EXCLUDING LOSSES
X max is  1.99868941307
X min is  -1.99506604671
Y max is   4.03058767319
Y min is  -4.72049236298
Z max is  0.130475103855
Z min is  0.592456926825E-04
X prime max is 0.112066427246E-01
X prime min is -0.114705841988E-01
Y prime max is 0.996395349503
Y prime min is 0.996019303799
Z prime max is 0.891377404332E-01
Z prime min is 0.841740518808E-01

$
PSPREAD

Purpose: Creates a histogram of the point spread function at an image plane.

Input: A ray file corresponding to the location of the desired point spread analysis, along with an interval and variable with respect to which the point spread is computed.

Output: A file (PSPREAD.DAT) containing data for the point spread histogram and/or a direct screen histogram.

Description:
The point spread function for an optical system is the two dimensional analog of the impulse response function for a linear system. PSPREAD computes a point spread histogram for any plane in the system in which a ray file exists. In order to get a true point spread the system must be traced with a point source in the XZ plane having no depth and a conical angle distribution. The histogram bins are created with respect to one of three variables: constant radius, where each bin represents an equal step outward from the user defined center; constant area, where each bin represents an equal area from the user defined center; or constant volume. The histogram counts may be made with respect to the number of photons or, in the case where reflectivity is used, the number of photons weighted by the electric field intensity. PSPREAD may be used to determine the spread for extended sources, which would be the convolution of the source shape with the point spread if the system was shift invariant.

Examples:
The first example we show is the point spread of a uniform source before it has been traced through an optical system. This shows the ideal point spread when calculated with respect to constant area.

Uniform Source
PSPREAD Example
Input file specified:end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.PSPREAD]END.00;1
Creation Date : 7-JUL-1989 09:31
Random Source.
Generated total 1000 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: ELLIPTICAL
Source Width: $0.100000000$ and Height: $0.100000000$
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Source Emission Characteristics
Distribution Type: LAMBERTIAN
Distribution Limits. +X: $0.200000000E-01$ -X: $0.200000000E-01$ rad
+Z: $0.200000000E-01$ -Z: $0.200000000E-01$ rad
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
*************** E N D ***************
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++

Below is a plot of the cross section of the source. We’ll now run PSPREAD and show the histogram it produces.

$ 
$ pspread
File for analysis ? begin.dat /* source file */
Read : 1000 rays
Maximum radius : 9.992726050813235E-02
Minimum radius : 3.9308915161286589E-03
Histogram is done for absolute radius /* negative values are meaningless
...from : 0 /* here. Work from zero. */
...to : 0.15
Number of bins (odd, please) ? 31 /* must be odd. 31 or 25 work well */
Generate the histogram at :
0 constant radius
1 constant area
2 constant volume
Then ? 1 /* choose constant area */
Flag checks. Enter :
0 to exclude lost rays
1 to include lost rays too
2 to use only lost rays
<?> 0 /* no lost rays */
Include reflectivity ? 0 /* no reflectivity yet */
Figure 3.28: PSPREAD: Source cross section
Output options:
[ 0 ] store histogram in a file
[ 1 ] plot histogram on screen
[ 2 ] both
Then ? 0
Another run [ Y/N ] ?n
Return to DCL
$ dir/since=-00:01
 /* information is output to PSPREAD.DAT */

Directory D13:[XRAYOP.DOCUMENT.UTIL.PSPREAD]

PSPREAD.DAT;2 5 10-JUL-1989 10:32:43.39 (RWED,RWED,RE,RE)

Total of 1 file, 5 blocks.
$

The histogram is shown below.

The next example will run PSPREAD on the STAR.02 file created with the Schwarzchild source and system. The source and system files are shown below.

```
+-------------------------------------------------------------------------------------------------+
************* SOURCE DESCRIPTION *************
Schwarzchild Source
PSPREAD Example
+-------------------------------------------------------------------------------------------------+
Input file specified:
end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.PSPREAD.MODEL]END.00;4
Creation Date : 7-JUL-1989 10:00
+-------------------------------------------------------------------------------------------------+
Random Source.
Generated total 5000 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: POINT
+-------------------------------------------------------------------------------------------------+
Source Emission Characteristics
Distribution Type: CONICAL
```
Figure 3.29: PSPREAD output
Cone Outer Aperture : 0.100000000E-01 Inner Aperture : 0.500000000E-02
Source Photon Energy Distribution: SINGLE LINE
Photon Energy: 281.78 eV, or 44.000 Angs.

Swarcchild system
PSPREAD Example

Optical Element # 1 System Number:
MIRROR SPHERICAL UNLIMITED EXTERNAL REFLEC. OFF
Orientation 0.00000000000000000E+00 deg.
Source Plane 51.210000000000000
Incidence Ang. 0.00000000000000000E+00 deg.
Reflection Ang. 0.00000000000000000E+00 deg.
Image Plane 2.500000000000000

Optical Element # 2 System Number:
MIRROR SPHERICAL UNLIMITED EXTERNAL REFLEC. OFF
Orientation 0.00000000000000000E+00 deg.
Source Plane 2.701370000000000
Incidence Ang. 0.00000000000000000E+00 deg.
Reflection Ang. 0.0000000000000000E+00 deg.
Image Plane 11.02186800000000

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

<table>
<thead>
<tr>
<th>OPT. Elem #</th>
<th>X =</th>
<th>Y =</th>
<th>Z =</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.000000000000000E+00</td>
<td>0.000000000000000E+00</td>
<td>0.000000000000000E+00</td>
</tr>
<tr>
<td>1</td>
<td>0.000000000000000E+00</td>
<td>51.210000000</td>
<td>0.000000000000000E+00</td>
</tr>
<tr>
<td>1'</td>
<td>0.000000000000000E+00</td>
<td>48.710000000</td>
<td>0.28605943631E-02</td>
</tr>
<tr>
<td>2</td>
<td>0.000000000000000E+00</td>
<td>46.008630000</td>
<td>0.59516038809E-16</td>
</tr>
<tr>
<td>2'</td>
<td>0.000000000000000E+00</td>
<td>57.030498000</td>
<td>-0.19271670896E-02</td>
</tr>
</tbody>
</table>

Shown below is the PLOTXY output of the second image file.
The STAR.02 file was run through PSPREAD. The run is shown below, followed by the histogram created with PSPREAD.DAT.

```
$ pspread
File for analysis? star.02  /* image file */
Read: 5000 rays
Maximum radius: 2.7598328716075391E-06
Minimum radius: 2.138662054996055E-10
Histogram is done for absolute radius
  ...from: 0
  ...to: 4.0E-06
Number of bins (odd, please)? 31
Generate the histogram at:
  0 constant radius
  1 constant area
  2 constant volume
```
Figure 3.30: PSPREAD: Second Image
Notice how the PSPREAD graph shows a constant density in the radius, while the histograms from PLOTXY show a peaked structure (because of the binning procedure).
Figure 3.31: PSPREAD output of STAR.02
Purpose: To generate a two-dimensional histogram containing the power density using an output file from SHADOW.

Input: Source file (i.e., BEGIN.DAT) and file for analysis (i.e., MIRR.xx etc).

Output: A file containing the power density grid in one of the following formats: TopDrawer, NCAR, generic format, or ANSYS, suitable for further processing.

Description:

PWR_DENS calculates the power at a given point in an optical system. It can be power transmitted/reflected, power absorbed, local reflectivity/transmission, or incoming power. The program reads a source file (BEGIN.DAT) and for all options except the power transmitted/reflected, the binary output file of the previous element. (For the first element in a system this would also be BEGIN.DAT.)

SHADOW stores the amplitudes of all the rays as they are traced through the system. The MIRR.xx file for an element stores the amplitude of the i-th ray “after” reflection. The incoming intensity is calculated from the source file for the element in question, the previous STAR file or BEGIN.DAT for the first element. If the amplitude, $A_s$ and $A_p$, is represented by “columns” 7, 8, 9 and 13, 14, 15 respectively, the intensity “after” reflection can be written as:

$$A_s^2 + A_p^2 = I_{after}(x, y)$$

(3.1)

A binning operation is done so that the reflected intensity is:

$$I_{after}(x, y) = \sum_{j=1}^{M} [A_{js}^2 + A_{jp}^2]$$

(3.2)

Along the same lines, we can write the absorbed intensity as:

$$I_{absorbed} = \sum_{j=1}^{M} [I_{j, reflected} - (I_{j, incoming})]$$

(3.3)

The local reflectivity is then:

$$r_s(x, y) = \frac{I_{after}}{I_{before}}$$

(3.4)

For this utility to give meaningful results, the source must have full polarization and reflectivities must be turned on for the elements in question.
For cleaner statistics, the utility MAKE_PWR will take the user through PWR_DENS and create a file that can be submitted as a batch job to run SHADOW iteratively as many times as desired.

**Examples:**

```
$ pwr_dens
Interactive [0] or namelist mode [1] : 0
This program needs a file containing the source spectrum computed within the limits used. Or you can specify the total power.
Source file [0] or total power [1] : 1
Total power in Watt : 100

Please specify now the file containing the SOURCE array (e.g., BEGIN.DAT)
Then ? BEGIN.DAT
Read 5000 rays. The file has 18 columns.
Computing renormalizing factor.

All preliminaries completed.
Ready to proceed with density analysis.

File for analysis ? MIRR.01
Data ready. Read 5000 points.
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ? 0
***********
File examined is :

Was created :

***********
Found 4717 good points out of 5000
Found:
Column 1: X max is 1.999296941008725
```
1: X min is -1.999883817471559
2: Y max is 3.568274926989652
2: Y min is -4.166031351586952
3: Z max is 0.1300002931936204
3: Z min is 3.5289946631600344E-07

for a photon energy range:
101.1613479791329 to 2959.045637676470

I need to define the two-dimensional histogram for the power density. Please specify:
Columns for x-axis and y-axis?
1
2

Limits to use for the histogram.
To use scaling, enter:
  automatic 0
  external 1

Then? 0

Number of bins in X [ max 51 ] ? 31
  Y [ max 51 ] ? 31

Optical Properties options.

To select enter

Power transmitted/reflected ........ 0
Power absorbed ....................... 1
Local reflectivity/transmission .... 2
Incoming Power ....................... 3

Then ?
1

File to use for Io ? BEGIN.DAT

The program need to know how many iterations have been already performed. Enter [ 0 ] if this is the first time.
No. of time this power density case has run : 0
How many times will you be looping through ? 1

There were 283 lost out of 5000 rays.

Begin verification.
Total power in file: 72.12355434039542
Compare with: 100.0000000000000

Computations done. Ready to store output data.
The files contain the power density at the nodes but for the case of ANSYS, that stores total power.

File for:
  TOP DRAWER [ 0 ]
  NCAR [ 1 ]
  Generic [ 2 ] ** to be used for loop
  For ANSYS [ 3 ]

2

Output file ? MIRR1
All done.

$ DIR/SINCE

Directory D1: [SHADOW.XRAYOP.DOCUMENT.TGM]

MIRR1.DAT;1 102 7-AUG-1989 10:38:55.61 (RWED,RWED,RE,RE)
PWR_DENS_PAR.DAT;1 3 7-AUG-1989 10:38:58.14 (RWED,RWED,RE,RE)

Total of 2 files, 105 blocks.

$ TY MIRR1.DAT

-1.9354 -4.0413 0.00000000E+00 0
-1.9354 -3.7918 0.35433976E-01 1
-1.9354 -3.5423 0.00000000E+00 0
-1.9354 -3.2928 0.35721396 4
-1.9354 -3.0433 0.55205592E-01 2
-1.9354 -2.7938 2.8768778 15
-1.9354 -2.5443 4.1098812 12
-1.9354 -2.2948 4.6196552 12
-1.9354 -2.0453 5.7607246 14
-1.9354 -1.7958 3.5439385 11

$
**Purpose:**
To combine two RAY files in a single one in order to merge two sources or two images.

**Input:**
Two Binary RAY files, e.g., STAR.01;0, STAR.01;2

**Output:**
A binary Ray file.

**Description:**
The program reads two ray files and combines them by appending the second to the first in anew file. The counter flag is reset to take in account the addition of rays. If the total number of rays exceeds 5000 an error message is typed and no output file produced. This program is designed to create “special” sources.

**Examples:**

As an example, let’s create two source files with a circular and rectangular shape. They are illustrated in Fig. 3.32 and fig. 3.33. After running COMB_RAY we obtain the source shown in Fig. 3.34. This program can be used to generate quite complicated sources; one could also combine STAR files with other STAR or BEGIN files for interesting systems.
Figure 3.32: COMB_RAY: Source one
Figure 3.33: COMB_RAY: Source two
Figure 3.34: COMB_RAY: Combined source
**Purpose:** To calculate the diffraction image from an aperture using the Fresnel-Kirchoff integral.

**Input:** Interactive from terminal and a SCREENxxxx.DAT file.

**Output:** Formatted 2-columns file.

**Description:**

Ordinarily, SHADOW is a ray-tracing program using geometrical optics. But it also has the capability of working in the physical optics regime. This is accomplished by properly defining the source and using FFRESNEL. It can be shown that the diffracted ("real") image can be computed by applying the Fresnel-Kirchoff Integral at the exit pupil of an optical system where the information on the illumination function has been obtained by geometrical optics. The placement of the pupil is arbitrary (as long as proper scaling is applied) since a shift leads to a phase change only. It is particularly useful for well corrected systems where most of the geometrical aberrations have been cancelled out and the image is diffraction limited. Such a case can be found in the Schwarzschild objective which we will be using as example here.

FFRESNEL is basically a numerical implementation of the full Fresnel-Kirchoff integral. It integrates over all the rays across a suitably located aperture. The principle is discussed in B. Lai, K. Chapman and F. Cerrina, *SHADOW: New Developments*, Nucl. Instr. and Meth. A266, 544-549 (1988). Briefly, the FK integral can be written as:

\[
A'(\vec{r}') = \frac{iu}{\lambda} \int_S A(\vec{r}) e^{ikR} \frac{d\vec{r}}{R}
\]  

(3.5)

where \( R = |\vec{r} - \vec{r}'| \) and \( r(\vec{r}') \) is the position vector at the image (exit pupil) plane. \( A(A') \) is the electric field at the same positions. SHADOW knows exactly \( A(\vec{r}) \) at the exit pupil while the mapping performed by the ray tracing operation has transformed the integral from a Riemann to a Stieltjes type. In order to do that, it must know for each ray the amplitude and phase of the electric field vector \( \vec{A} \). In general, we separate \( \vec{A} \) into two components:

\[
\vec{A} = \vec{A}_s e^{i\theta_s} + \vec{A}_p e^{i\theta_p}
\]  

(3.6)

where \( s \) and \( p \) are the perpendicular and parallel component (respect to the plane of incidence). At the source, they are along the X and Z direction. Thus the \( \vec{A} \) vector is defined
initially at SOURCE by:

\[
\text{degree of polarization} = \frac{\tilde{A}_x}{\tilde{A}_x + \tilde{A}_z}
\]
\[
\text{phase difference} = \theta_z - \theta_x
\]  

(3.7)  

(3.8)

From then on, both the phase and amplitude may be modified upon reflecting off a surface (be it a grating, crystal, multilayers or reflective coatings), and these are all tracked by SHADOW ray by ray. In addition, the user also has the option of storing the optical path distance (OPD) of each ray from the source. This fully specifies (with 18 columns) all the rays anywhere along the optical system. With this information, FFRESNEL can easily evaluate the integral.

A very important point is that, since we are dealing with interference pattern, good cancellation cannot be achieved if we just use a random source due to finite sampling. Thus a grid source with coherent phase should be used if FFRESNEL is to give meaningful results. The type of source to use requires careful analysis. For example, a SR source is essentially fully incoherent so that the final image will be a convolution of the diffracted image (computed from a point source) with the geometrical optics image (computed from the extended source).

The first example shows the diffraction of a very simple system, a spherical mirror with the point source located at the center of curvature. The geometrical image is clearly a point while the diffracted image is given by an Airy pattern of width depending on the radiation numerical aperture. The first minimum is indeed located at

\[
\Delta = \frac{\lambda}{2 \text{N.A.}}
\]  

(3.9)

In the second example, a conical source is used to match the acceptance of the Schwarzschild objective. We place 30 points along the radius and only 20 points around the circle since the system is spherically symmetric. This system then images the point source to 11.02 cm after the second mirror. MINMAX tells us that the geometrical spot is about 276˚A which demonstrates the quality of the image. So we define an aperture, using the screen option in SHADOW, 1 cm after the second mirror and use FFRESNEL to calculate the diffraction image 10.02 cm from the screen. At 44 Åwavelength, the diffraction image size is certainly comparable to that of the geometric image. Note that we do not specify the photon energy in SOURCE and input it as parameter in FFRESNEL for flexibility. The user can do it the other way if he wants to use a chromatic source.

Examples:
The first example we will show is a simple case of a point source at the focal point of a spherical mirror. We run the example with three different numerical apertures, 0.1, 0.2, and 0.4. The diffraction patterns are plotted together following the example.

```
Spherical mirror source example
FFRESNEL Example
Input file specified:
en.d0
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.FFRESNEL.EXAMPLE]END.00;
Creation Date :25-JUL-1989 16:12
Grid Source.
Generated total 2601 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: POINT
Source Emission Characteristics
Distribution Type: CONICAL
Cone Outer Aperture : 0.200000000 Inner Aperture : 0.000000000E+00
Source Photon Energy Distribution: SINGLE LINE
Photon Energy: 2.4797 eV, or 5000.0 Angs.
Angular difference in phase is 0.00000E+00 /* full specification of rays
Degree of polarization is 0.50000 /* neccesary. */
Source points have COHERENT phase.
```

```
END
```

```
Spherical mirror
FFRESNEL example
Input file specified: end.01
```
A screen was defined 10 cm after the mirror for the purposes of running FFRESNEL.

$ ffresnel

Define integration limits [ cm ].
From -5E-04
to 5E-04
No. of points 201 /* odd number works best. */
Integration direction [ 0 = x, 90 = z ] ? 0
Distance from plane [ cm ] ? 90 /* distance of screen from image */
Ray input file ? screen.0101 /* input SCREEN.xxyy file */
[ 0 ] to discard losses, [ 1 ] to keep them. 0
Do you want to override the wavelength in the file ? 0
Files read OK.
Conversion to \[ \text{cm} \] factor (*) ? 1.0
Include the S- and P- phase shifts in the computations ?1
$ \text{dir/since=-00:02}$

Directory D13: [XRAYOP.DOCUMENT.UTIL.FFRESNEL.EXAMPLE]

FFPAR.DAT;23 1 11-JUL-1989 11:20:03.11 (RWED,RWED,RE,RE)
FFRESNEL.DAT;16 20 11-JUL-1989 11:20:03.81 (RWED,RWED,RE,RE)

Total of 2 files, 21 blocks.
$

The plot follows.
Next the source was offset by 0.5 units to test its affect on the diffraction pattern. This
offsets the image the same distance in the opposite direction, so our integration will no
longer be around zero. The output from SOURCINFO is the same as the previous example.
The numerical aperture is 0.2. The mirror description follows:

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
******************** MIRROR DESCRIPTION ********************
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Spherical Mirror--Offset
FFRESNEL Example
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified:end.01
Full file Specification :D1: [XRAYOP.DOCUMENT.UTIL.FFRESNEL.EXAMPLE]END.01
Creation Date : 2-AUG-1989 16:32
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Surface figure was defined as: SPHERICAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity OFF
Mirror dimensions UNLIMITED
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Central Axis parameters :
Source Plane Distance 100.0000000000000
Image Plane 100.0000000000000
Figure 3.35: FFRESNEL output with N.A. 0.1, 0.2, 0.4
Incidence Angle 0.0000000000000000E+00
Reflection/Diffraction Angle 0.0000000000000000E+00

Mirror parameters EXTERNAL
Parameters used follow:
Spherical Radius 100.000000
Source of this O.E. moved YES

In SOURCE reference frame:
Source Movement X: 0.0000000000000000E+00
Y: 0.0000000000000000E+00
Z: 0.5000000000000000E+00 <<< offset source 0.5 cm *\nSource rotat X: 0.0000000000000000E+00
Y: 0.0000000000000000E+00
Z: 0.0000000000000000E+00

In MIRROR reference frame:
Source distance 100.0000000000000
rotation 0.0000000000000000E+00
Incidence angle 0.0000000000000000E+00
Source offset X: 0.0000000000000000E+00
Y: 0.0000000000000000E+00
Z: 0.0000000000000000E+00
Mirror at pole position ( no mov. ) YES

Define integration limits [ cm ].
From 0.495 /* offset integration as well */
to 0.505
No. of points 201
Integration direction [ 0 = x, 90 = z ] ? 90 /* integrate in Z since */
/* offset in Z */
Distance from plane [ cm ] ? 90
Ray input file ? SCREEN.0101
[ 0 ] to discard losses, [ 1 ] to keep them. 0
Do you want to override the wavelength in the file ? 0
Files read OK.
Conversion to [ cm ] factor (*) ? 1.0
Include the S- and P- phase shifts in the computations ?
$ DIR/SINCE

Directory D1:[SHADOW.XRAYOP.DOCUMENT.UTIL.FFRESNEL.EXAMPLE]

FFPAR.DAT;1 1 2-AUG-1989 16:44:52.84 (RWED,RWED,RE,RE)
FFRESNEL.DAT;1 20 2-AUG-1989 16:44:53.24 (RWED,RWED,RE,RE)

Total of 2 files, 21 blocks.
$ TY FFPAR.DAT
From 0.4950000000000000 to 0.5050000000000000
Step 5.0000000000000010E-05
Wavelength 0.0000000000000000E+00 angs.
Image located at 90.00000000000000
Integration direction 90.00000000000000
Scale factor [ units ] * 1.0000000000000000 = [ cm ]
Losses: NOT INCLUDED
S- and P- phase shifts : INCLUDED
Files used:
SCREEN.0101

The offset plot superimposed on the first example with N.A.=0.2 follows.
The next example is more complex. It involves the Schwarzschild source and system. First
SHADOW was run to define the source and the Schwarzschild objective. The output from
SOURCINFO and SYSINFO follows.

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
************** SOURCE DESCRIPTION **************
Source for FFRESNEL

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified:
end.00
Full file Specification :D1:[XRAYOP.DOCUMENT.UTIL.FFRESNEL]END.00;2
Figure 3.36: FFRESNEL: Offset example
Grid Source. "* grid source a must *
Generated total 600 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: POINT
Source Emission Characteristics
Distribution Type: CONICAL
Cone Outer Aperture: 0.10000000E-01 Inner Aperture: 0.50000000E-02
Source Photon Energy Distribution: SINGLE LINE
Photon Energy: 281.78 eV, or 44.000 Angs.
Angular difference in phase is 0.00000E+00 "* full specification of rays
Degree of polarization is 0.50000 "* needed OPD + A vector
Source points have COHERENT phase. "* No random phase
*******************************************************************************
*************** E N D ***************
*******************************************************************************
\\ Now define the first mirror (convex). \\
\\ Then define the second mirror (concave). \\
\\ Following the log from the screen definition is the system output from
SYSINFO. \\
\\ Define a screen as our aperture. \\
:\n:\n\n Any screens in this OE [ Y/N ] ? 1
How many in this OE [ total ] ? 1
Screen N. 1
Is this screen before mirror [ Y/N ] ? 0
Distance from mirror [ absolute ] ? 1.0
Is Screen Carrying an Aperture Stop [ Y/N ] ? 0
Include absorption [ Y/N ] ? 0
:\n:\n*******************************************************************************
*************** S Y S T E M       D E S C R I P T I O N ***************
*******************************************************************************
FFRESNEL Example

Input file specified:

# Optical Element: Creation Time:
1  D13:[XRAYOP.DOCUMENT.UTIL.FFRESNEL]END.01;2  3-JUL-1989 14 225
2  D13:[XRAYOP.DOCUMENT.UTIL.FFRESNEL]END.02;2  3-JUL-1989 14 225

Optical Element # 1  System Number:

MIRROR SPHERICAL UNLIMITED EXTERNAL REFLEC. ON

Orientation 0.0000000000000000E+00 deg.
Source Plane 51.210000000000000
Incidence Ang. 0.0000000000000000E+00 deg.
Reflection Ang. 0.0000000000000000E+00 deg.
Image Plane 2.500000000000000

Optical Element # 2  System Number:

MIRROR SPHERICAL UNLIMITED EXTERNAL REFLEC. ON

Orientation 0.0000000000000000E+00 deg.
Source Plane 2.701370000000000
Incidence Ang. 0.0000000000000000E+00 deg.
Reflection Ang. 0.0000000000000000E+00 deg.
Image Plane 11.021868000000000

SCREENS: 1 defined.
BEFORE Mirror at 1.000000000000000

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

OPT. Elem # X = Y = Z =


\ \ First check the geometric image size. \ \\

$ 
$ MINMAX 
Input file ? STAR.02 
Options --- Enter 
0 for excluding the losses 
1 for including losses at a particular O.E. 
2 for including all the losses. 
Then ? 0 
Comment ? Geometrical Image 
********* 
File examined is : 
D13:[XRAYOP.DOCUMENT.UTIL.FFRESNEL]STAR.02;2 
Was created : 
3-JUL-1989 14:32 
******* 
Found 600 good points out of 600 
Here we are. 
X max is 2.7609133E-06 
X min is -2.7609133E-06 
Y max is 2.2204460E-16 
Y min is 0.0000000E+00 
Z max is 2.7609133E-06 
Z min is -2.7609133E-06 
X prime max is 0.2000885 
X prime min is -0.2000885 
Y prime max is 0.9949863
Y prime min is 0.9797778
Z prime max is 0.2000885
Z prime min is -0.2000885

$ DIR SCREEN*  
  * The aperture file *

Directory D13:[XRAYOP.DOCUMENT.UTIL.FFRESNEL]

SCREEN.0201;2  170  3-JUL-1989 14:32:26.41 (RWED,RWED,RE,RE)
Total of 1 file, 170 blocks.

$ FFRESNEL  
  * calculate the diffraction image *

Define integration limits [ cm ].
From -500E-8  
  * -500 to 500 Angstroms *
  to 500E-8
No. of points 101
Integration direction [ 0 = x, 90 = z ] ? 0  
  * along the horizontal *
Distance from plane [ cm ] ? 10.021868
Ray input file ? SCREEN.0201
[ 0 ] to discard losses, [ 1 ] to keep them. 0
Do you want to override the wavelength in the file ? 0
Files read OK.
Conversion to [ cm ] factor (*) ? 1.0
Include the S- and P- phase shifts in the computations ?1

$ DIR FF*

Directory D13:[XRAYOP.DOCUMENT.UTIL.FFRESNEL]

FFPAR.DAT;2  1  3-JUL-1989 15:00:20.70 (RWED,RWED,RE,RE)
FFRESNEL.DAT;2 10  3-JUL-1989 15:00:21.49 (RWED,RE,RE,RE)
Total of 2 files, 11 blocks.

$ TY FFPAR.DAT  
  * stores parameters for FFRESNEL *
From -5.0000000000000000E-06 to 5.00000000000000000E-06
Step  1.0000000000000000E-07
Wavelength  0.0000000000000000E+00 angs.
Image located at  10.02186800000000
Integration direction  0.0000000000000000E+00
Scale factor [ units ] *  1.0000000000000000  = [ cm ]
Losses: NOT INCLUDED
S- and P- phase shifts : INCLUDED
Files used:
SCREEN.0201
$
$ TY FFRESNEL.DAT      /* the diffraction image */
-5.0000000000000000E-06  1973087.868071882
-4.9000000000000000E-06  1695296.044110004
-4.8000000000000000E-06  1316561.473098089
   :                      :            :            :
   :                      :            :
 4.8000000000000000E-06  1316561.466332203
 4.9000000000000001E-06  1695296.058811655
 5.0000000000000000E-06  1973087.881998670
$
Figure 3.37: FFRESNEL: Schwarzschild example
Purpose: To search a BOF for the location of the minimum beam waist (i.e. the best gaussian focus in both the sagittal (x) and tangential (z) directions).

Input: STAR.xx

Output: A file FOCUS.DAT containing the analysis results plus an optional user defined plottable file that tabulates waist size vs. distance along optical axis. Analysis results are also displayed on terminal.

Description:

Locates the waist of the beam by minimizing the variance in real space with respect to the optical axis in the sagittal, tangential planes and for the circle of least confusion. If \( x_i \) is the coordinate of the \( i^{th} \) ray (in the sagittal plane), and \( v_i \) is its direction vector then the variance will be:

\[
\sigma(t) = \frac{1}{N} \sum_{i=1}^{N} (x_i + v_i t)^2
\]

Expanding the sum:

\[
\sigma(t) = \frac{1}{N} \sum_{i=1}^{N} (x_i^2 + 2x_i v_i + v_i^2 t^2)
\]

For many purposes, the focus can be defined as the position where the beam spread is minimum. Thus we have to find the minimum of the above expression, which is just a parabola, with respect to \( t \). Setting the derivative equal to zero and solving for \( t \) we have,

\[
t = \frac{-\sum_{i=1}^{N} x_i v_i}{\sum_{i=1}^{N} v_i^2}
\]

The expressions for the tangential focus and circle of least confusion are similar. The user may also choose a center of the distribution other than the optical axis.

This utility is very important in the study of monochromators and spectrographs, since it allows the user to verify the real location of the focal position.

Note: One must be careful in using FOCNEW and verify with PLOTXY that the calculations are correct. This is because with strongly asymmetric or multimodel distribution the standard deviation is not a good measure of the image “size”.

Examples:
Here we will have a spherical mirror create an astigmatic image by having a point source incident upon it at 20 degrees. SHADOW computes the curvature of the mirror to satisfy the focal condition for the tangential plane. The sagittal focus occurs further away from the mirror, and the circle of least confusion is located between the two. Below are the source and mirror descriptions.

++++++++++ Source Description ++++++++++

Point source
Focnew Example

Input file specified:
end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.FOCNEW]END.00;1
Creation Date :11-APR-1989 10:56
Random Source.
Generated total 1000 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: POINT
Source Emission Characteristics
Distribution Type: CONICAL
Cone Outer Aperture : 0.500000000E-01 Inner Aperture : 0.100000000E-01
++++++++++ End +++++++++

++++++++++ Mirror Description ++++++++++

Spherical Mirror Illuminated at 20 degrees incidence
Focnew Example to Generate Astigmatic Image

Input file specified: end.01
Surface figure was defined as: SPHERICAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity OFF
Mirror dimensions UNLIMITED

Central Axis parameters:
Source Plane Distance 30.00000000000000
Image Plane 30.00000000000000
Incidence Angle 20.00000000000000
Reflection/Diffractive Angle 20.00000000000000

Mirror parameters COMPUTED
Same configuration as Central Axis YES
Objective focus at 30.00000000000000
Image focus at 30.00000000000000
Incidence angle 20.00000000000000

Parameters used follow:
Spherical Radius 31.9253332
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

We will have FOCNEW create files for the sagittal and tangential focii as well as the circle of least confusion. These files will then be used as input to TopDrawer to plot the results.

$ focnew
Input file ? star.01
We have 1000 good points out of 1000
Row Par Minimum: Maximum: Center: St. Dev.:

1 X -0.35104 0.35104 0.36242E-17 0.17898
2 Y -0.74221E-15 0.59533E-15 -0.29388E-16 0.37404E-15
3 Z -0.50159E-03 0.56424E-02 0.14659E-02 0.20068E-02
4 X' -0.38286E-01 0.38286E-01 -0.45285E-18 0.19524E-01
5 Y' 0.99870 0.99997 0.99948 0.37334E-03
6 Z' -0.49071E-01 0.50888E-01 0.76300E-04 0.25491E-01

Options:
center at origin [ 0 ]
center at center of gravity [ 1 ]
external [ 2 ]
Then ? 0

Searching file:
star.01
D13:[XRAYOP.DOCUMENT.UTIL.FOCNEW]STAR.01;5 11-APR-1989 10:56

Center at :Origin
X = 0.0000000000000000E+00 Z = 0.0000000000000000E+00

............. SAGITTAL ............
X coefficients : 0.3815973746E-03 -0.3496278593E-02 0.3203366904E-01
Center : 0.3624200855E-17 Average versor : 0.2795689524E-18
Sagittal focus at : 9.162218680
Waist size at best focus (rms) : 0.0000000000000000E+00
Waist size at origin : 0.1789795213

............. TANGENTIAL ............
Z coefficients : 0.6510081993E-03 -0.5464547342E-05 0.6176084572E-05
Center : 0.1465870279E-02 Average versor : 0.7687155566E-04
Tangential focus at : 0.8393976217E-02
Waist size at best focus (rms) : 0.2475927158E-02
Waist size at origin : 0.2485172946E-02

............. LEAST CONFUSION ............
T coefficients : 0.1032605574E-02 -0.3501743141E-02 0.3203984512E-01
Center : 0.1465870279E-02 Average versor : 0.7687155566E-04
Circle of least confusion : 3.391172031
Waist size at best focus (rms) : 0.1420029286
Waist size at origin : 0.1789967741
All done. File out data.
Do you want a plottable file ? 1
Enter :
1 for X
2 for Z
3 for T
4 for all
Then ? 4
Please specify Limits [ YMIN, YMAX, STEP ] ?
Ymin ? -5
Ymax ? 15
Ystep ? 0.5
File root [ 7 letters max ] ? foc.dat
Enter :
0 for another plot
1 to restart
2 to exit
Then ? 2
$
$

Shown below are the Phase space projections in X and Z showing lack of focus in the sagittal plane and focus in the tangential plane, since the points are uncorrelated.

Notice how the beam is smallest at the slit position.

A more complex case may be that of the toroidal grating monochromator (TGM). This mount is based on the use of a Toroidal grating rotated around its surface with fixed slits located at p and q respectively. A Rowland Circle mount would give the best image when the Rowland circle condition is satisfied, but the quality degrades quickly if the grating is rotated. In the TGM mount the quality of the “best” focus is sacrificed so that a wider range with “acceptable” resolution is maintained.

Typically, the entrance slit is located “inward” relative to the rowland circle position at zero order (asymmetric mount) so that its image moves outward. When scanning the wavelength the change in angle of incidence brings the image back closer to the grating until it is located at the exit slit; this is the first “sweet spot” of the TGM. Then the image moves even closer to a minimum distance and the process reverses, with the image moving away through the second “sweet spot” of the spectrum.

Clearly if one could move the exit slit to follow the “best” focus, the performance of the monochromator would be improved. This has originated many designs based on movable exit slits. FOCNEW is particularly important because it allows one to locate the position
Figure 3.38: FOCNEW: X phase space
Figure 3.39: FOCNEW: Z phase space
Figure 3.40: FOCNEW: Full beam size
Figure 3.41: FOCNEW: Sagittal beam size
Figure 3.42: FOCNEW: Tangential beam size
of this best focus. It is also useful to verify that the optical system we have just traced is indeed performing as expected.

With the TGM, let’s look at how the tangential focus at the final image plane is dependent upon the beam energy. We first examined the focus at 50eV, the energy to which the grating was tuned. Note that the center of gravity was used, because at different energies, the ray bundle will be displaced from the origin. Then the source was recolored to 50.1eV, the system was retraced, and FOCNEW was run in the same manner, again using the center of gravity for the distribution center. The FOCNEW file FOCUS.DAT for each run is displayed below, along with a plot around the tangential focal points.

```
FOCNEW

Searching file:
tgm:star.03
D13:[XRAYOP.DOCUMENT.TGM]STAR.03;29  9-MAY-1989 11:1

Center at: Baricenter
X = 0.0000000000000000E+00  Z = 0.0000000000000000E+00

Working with 865 "good" rays out of 1000

.............. S A G I T T A L ..............
X coefficients: 0.1279945686E-03  0.2849121654E-03  0.1589516586E-02
Center: 0.3572783631E-02  Average versor: 0.2159759878E-03
Sagittal focus at: -2.232813118
Waist size at best focus (rms): 0.3064103890E-01
Waist size at origin: 0.3970833417E-01

.............. T A N G E N T I A L ............
Z coefficients: 0.1814695449E-04  0.5456865017E-04  0.2534060565E-03
Center: 0.1037995525E-01  Average versor: 0.1706396676E-02
Tangential focus at: -2.419161809
Waist size at best focus (rms): 0.7516725477E-02
Waist size at origin: 0.1206907559E-01

.............. LEAST CONFUSION ............
T coefficients: 0.1461415231E-03  0.3394808156E-03  0.1842922642E-02
Center: 0.1097762515E-01  Average versor: 0.1720010245E-02
Circle of least confusion: -2.252641249
Waist size at best focus (rms): 0.3155704656E-01
Waist size at origin: 0.4150198054E-01
```
Searching file:
tgm:star.03;28
D13:[XRAYOP.DOCUMENT.TGM]STAR.03;28 9-MAY-1989 11:10

Center at : Baricenter
X = 0.0000000000000000E+00  Z = 0.0000000000000000E+00

Working with 865 "good" rays out of 1000

.............. S A G I T T A L ..............
X coefficients : 0.1300886749E-03 0.2994988597E-03 0.1629528588E-02
Center : 0.3521177601E-02 Average versor : 0.2175865878E-03
Sagittal focus at : -2.308996827
Waist size at best focus (rms) : 0.3039438956E-01
Waist size at origin : 0.4021355364E-01

.............. T A N G E N T I A L .............
Z coefficients : 0.1549933837E-04 0.3405311760E-04 0.2902239386E-03
Center : 0.1251010550E-01 Average versor : 0.4472218222E-04
Tangential focus at : -2.233454297
Waist size at best focus (rms) : 0.7511028279E-02
Waist size at origin : 0.1156378826E-01

.............. LEAST CONFUSION ..............
T coefficients : 0.1455880133E-03 0.3335519773E-03 0.1919752527E-02
Center : 0.1299620835E-01 Average versor : 0.2213508688E-03
Circle of least confusion : -2.300952861
Waist size at best focus (rms) : 0.3130995822E-01
Waist size at origin : 0.4184317263E-01
Figure 3.43: TGM example with FOCNEW
### Purpose:
To modify the photon energies of each ray in a source array file according to a user defined distribution. Only random photon energy distributions are allowed.

### Input:
A ray Binary Output File.

### Output:
A ray Binary Output File.

### Description:
This utility will create a ray file with a distribution of energies. Such a modification is useful in studying the effects of a nonmonochromatic source. For example, one may wish to examine how power is absorbed in a mirror as a function of wavelength or one could determine how different energies are diffracted in a given crystal. A typical application can be found when one runs first a monochromatic source through a system dispersing, say, in $Z$. PREPLOT can be used to extract $\text{FIL1} = \{h\omega_0, z\}$. Then RECOLOR is used to modify $h\omega_0$ to the range $h\omega_0 \pm \Delta h\omega$. After running SHADOW, one creates $\text{FIL2} = \{h\omega, z\}$. Then using FILEX:

$$\text{DISP} = \text{FIL2} - \text{FIL1}$$

and plot DISP. The graph will show directly $\Delta(h\omega)$ around $h\omega_0$. Notice that for extended sources the values of $Z$ in FIL1 will not be all the same.

RECOLOR will change the photon energy distribution to one of the following:

1. **Monochromatic:** All rays have the same energy.

2. **Multi Lines:** Discrete energies, each specified by the user (not necessarily equispaced), are assigned to rays at random.

3. **Uniform Distribution:** Any value between user specified limits may be assigned at random to the ray energy.

The energies can be expressed in eV or as a function of wavelength.

### Examples:
We will examine an ERG monochromator with a monochromatic point source having a much larger vertical divergence than horizontal. Then we will use RECOLOR to create a white source with the energy distribution centered on that of the tuned grating. The beam will be dispersed in the tangential plane as a function of energy. The dispersion may be
examined by finding the difference between the Z coordinates for the monochromatic and white rays and plotting this vs energy. Below are descriptions of the source and system.

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
************** SOURCE DESCRIPTION **************
Point source to be incident upon ERG
Recolor Example
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified:
end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.RECOLOR]END.00;2
Creation Date :13-APR-1989 09:26
Random Source.
Generated total 1000 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: POINT
Source Emission Characteristics
Distribution Type: LAMBERTIAN
Distribution Limits. +X : 0.100000000E-03 -X: 0.100000000E-03 rad
+Z : 0.200000000E-01 -Z: 0.200000000E-01 rad
Source Photon Energy Distribution: SINGLE LINE
Photon Energy: 400.00 eV, or 30.996 Angs.
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
******************** MIRROR DESCRIPTION ********************
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
ERG auto tuned to order -1
RECOLOR EXAMPLE
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified: end.01
Full file Specification : D13:[XRAYOP.DOCUMENT.UTIL.RECOLOR] END.01;2
Creation Date : 13-APR-1989 09:27

Surface figure was defined as: SPHERICAL
Cylindrical figure NO
Element type REFLECTOR
Element type GRATING
Order choosen ( inside area < 0 ) -1.000000000000000
Automatic Tuning YES
Mount ERG
Grating tuned at [ eV ] 400.0000000000000
Constant ruling [ lines/cm ] 12000.00000000000
Reflectivity OFF
Mirror dimensions UNLIMITED

Central Axis parameters :
Source Plane Distance 174.5000000000000
Image Plane 464.7235755591278
Incidence Angle 88.00000000000000
Reflection/Diffraction Angle 84.66696278217028

Mirror parameters EXTERNAL
Parameters used follow:
Spherical Radius 5000.00000
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

Now we will create a file with the energy and Z coordinate of each ray in the image plane.

PREPLOT> Input file ?
400ev.img
Read 1000 rays.
Each ray has 12 entries.
PREPLOT> How many columns to write out ?
Now we will create the white light source in the interval 395 to 405 eV. The system is then retraced with the new source.

$ RECOLOR
File for input ? BEGIN.DAT
and for output ? BEGIN.DAT
Data ready. Read 1000 points.
Are you going to specify energies in eV [ 0 ] or wavelengths in angstroms [1] ? 0
Available options :
After rerunning TRACE in the BATCH mode on the system, we create the file of Z coordinate vs energy on the new image file with the aid of PREPLOT.

$ preplot
PREPLOT> Input file ?
delev.img
Read 1000 rays.
Each ray has 12 entries.
PREPLOT> How many columns to write out ?
2
Row [1-12] : the individual column
Row [20] : R = SQRT(X**2 + Y**2 + Z**2)
Row [21] : angle from the Y-axis
Row [22] : the magnitude of A vector
Row [23] : A**2
PREPLOT> Row # 1 : 11
PREPLOT> 2 : 3
Option: Angstroms [ 0 ]
Electronvolts [ 1 ]
Cm-1 [ 2 ]
PREPLOT> Then ? 1
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays.
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
PREPLOT> Then ? 0
***********
Found 1000 good points out of 1000
Output options :
[ 0 ] store rays in a file
[ 1 ] plot directly on screen
[ 2 ] both
Then ? 0
PREPLOT> Output file ? delev.dat

Use FILEX to get the difference between the Z coordinates.

$ filex
FILEX >
disper = delev - 400ev
FILEX > read 1000 records.
FILEX > Successful completion
FILEX >

The file disper.dat can now be plotted using TDTEK or TDFILE and UGTEK. Notice the variance in the dispersion away from 400eV. This is a result of the spherical aberration of the grating since the divergence in the tangential plane is significant.
Figure 3.44: RECOLOR: Image at 400 eV
Figure 3.45: RECOLOR: Image from 395-405 eV
Figure 3.46: RECOLOR: Dispersion in tangential plane from 395-405 eV
RETRACE

Purpose: To create a new BOF (e.g. STAR.xx) from an existing BOF at a different position along the optical axis by continuing the trajectory of the rays. Notice that the distance is arbitrary so that for example it is possible to study virtual images.

Input: Any BOF

Output: A BOF

Description:

This utility enables the user to look at the 'image' formed by the rays at some distance along the optical axis from the location of the continuation plane. The rays are assumed to be undeflected between the continuation plane and the desired location. Thus the new position can be calculated as follows:

\[ X(t) = X(0) + X' t \]

Here \( X' \) is assumed constant, of course.

RETRACE should be used in conjunction with FOCNEW and PLOTFOC, for example, to look at the image at the circle of least confusion without having to go back into SHADOW to modify the image plane and retrace the entire system.

Examples:

In this example we will examine the astigmatic image that was generated in Section 2.4 of the Primer. Recall that FOCNEW told us that the sagittal focus and the circle of least confusion were about 9.151 and 3.281 units away from the tangential focus. We may plot out the cross sections of the beam or use other utilities to analyze the new ray files. In order to create a BOF for the beam at the position of the sagittal focus and circle of least confusion without retracing the system, we can let RETRACE do the work for us:

```
$ retrace
RETRACE> Ray input file ? star.01
RETRACE> Ray output file ? ret.dat
RETRACE> Distance ? 9.151
Read 1000 rays.
```
Another run [ Y/N ] ? y
RETRACE> Ray input file ? star.01
RETRACE> Ray output file ? ret2.dat
RETRACE> Distance ? 3.281
Read 1000 rays.
Another run [ Y/N ] ? 0

Now that we have the ray files for the desired positions, we can plot out the images of the beam. Notice that neither coordinate is in focus, but that the optical size is smaller.
Figure 3.48: RETRACE: Tangential focus
Figure 3.49: RETRACE: Circle of least confusion
**Purpose:** To combine up to four tracings from SHADOW in a single sectored output.

**Input:** A BEGIN/STAR file and the corresponding START.xx file, up to four MIRR.xx files, one for each sector.

**Output:** A STAR.xx file containing the image from the sectored surface and a MIRRS.xx file containing the combined MIRR file.

**Description:**
This utility will combine up to four separate tracings from SHADOW to create a “compound” mirror otherwise not possible with SHADOW. Each tracing is done separately by the user, using the same source or previous optical element. Then the utility is run. The utility reads in a source file and its corresponding namelist file (i.e. BEGIN.DAT and START.00 or STAR.02 and START.02, etc.). It then reads in up to four mirror files (MIRR.xx). Using these files, the program computes the optical path to the mirror. The intercepts associated with shorter optical paths are then put into a single continuation file. The program writes two binary files; MIRRS.00 and STAR.00, the mirror file and continuation plane file respectively. These files can be plotted with PLOTXY, like any other mirror or image file.

**Examples:**
The following example shows how one would trace a flashlight composed of two cylindrical parabolical mirrors with the cylinder axes 90 degrees apart. First we show the output from MIRINFO on each of the mirrors we start with.

```
+===============================================================================================================
*                         MIRROR DESCRIPTION                          *
+===============================================================================================================
SECTOR Example - First Mirror
0 deg orientation
+===============================================================================================================
Input file specified:end.01
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.SECTOR.M1]END.01;5
Creation Date :12-JUN-1989 13:35
+===============================================================================================================
Surface figure was defined as: PARABOLICAL
```
Cylindrical figure: YES  << note angle
Cylinder axis angle from X-axis: 0.0000000000000000E+00
Element type: REFLECTOR
Reflectivity: OFF
Mirror dimensions: UNLIMITED

Central Axis parameters:
- Source Plane Distance: 1.0000000000000000E+00
- Image Plane: 5.0000000000000000E+00
- Incidence Angle: 0.0000000000000000E+00
- Reflection/Diffraction Angle: 0.0000000000000000E+00

Mirror parameters: COMPUTED
- Same configuration as Central Axis: YES
- Objective focus at: 1.0000000000000000E+00
- Image focus at: 5.0000000000000000E+00
- Incidence angle: 0.0000000000000000E+00

Parameters used follow:
- Parabola Param.: 2.0000000000000000E+00
- Source of this O.E. moved: NO
- Mirror at pole position (no mov.): YES

-----------------------------------------------
************** MIRROR DESCRIPTION **************
-----------------------------------------------
SECTOR Example - Second Mirror
0 deg orientation

Input file specified: end.01
Full file Specification: D13:[XRAYOP.DOCUMENT.UTIL.SECTOR.M2]END.01;5
Creation Date: 12-JUN-1989 13:39

Surface figure was defined as: PARABOLICAL
Cylindrical figure: YES  << note angle of
Cylinder axis angle from X-axis: 90.000000000000002
Element type: REFLECTOR
Reflectivity OFF
Mirror dimensions UNLIMITED

Central Axis parameters:
Source Plane Distance 1.000000000000000
Image Plane 5.000000000000000
Incidence Angle 0.000000000000000
Reflection/Diffraction Angle 0.000000000000000

Mirror parameters COMPUTED
Same configuration as Central Axis YES
Objective focus at 1.000000000000000
Image focus at 5.000000000000000
Incidence angle 0.000000000000000
Parameters used follow:
Parabola Param. 2.000000000000000
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

Using the renamed mirror files from the above mirrors, we'll now run SECTOR using the same source used in tracing of each of the separate mirrors.

$ run util:sector
Source file ? BEGIN.DAT /* or the previous STAR.xx file */
Parameter file [ START.xx ] ? START.00 /* or START.xx file corresponding */
How many sectors [ max 4 ] ? 2 /* to previous optical element */
Call to SETSOUR
Exit from SETSOUR
Call to IMREF
Exit from IMREF
Call to OPTAXIS
Exit from OPTAXIS
Read 1000 from
BEGIN.DAT

Sector # 1
File name ? MIRR1.01  /* MIRR.xx file of first sector */
Sector # 2
File name ? MIRR2.01  /* MIRR.xx file of second sector */
Call to RESTART
Exit from RESTART
Call to IMAGE
Exit from IMAGE
$ DIR/SINCE=-00:05

Directory D13:[XRAYOP.DOCUMENT.UTIL SECTOR]

<table>
<thead>
<tr>
<th>File Name</th>
<th>Blocks</th>
<th>Date/Time</th>
<th>Permissions</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIRRS.00;3</td>
<td>188</td>
<td>8-JUN-1989 16:18:59.20</td>
<td>(RWED,RWED,RE,RE)</td>
</tr>
<tr>
<td>OPTAX.00;3</td>
<td>0</td>
<td>8-JUN-1989 16:18:37.54</td>
<td>(RWED,RWED,RE,RE)</td>
</tr>
<tr>
<td>STAR.00;3</td>
<td>188</td>
<td>8-JUN-1989 16:19:02.36</td>
<td>(RWED,RWED,RE,RE)</td>
</tr>
</tbody>
</table>

Total of 3 files, 376 blocks.
$ /* Both MIRRS.00 and STAR.00 are binary files plottable using PLOTXY */
  /* giving the results of the combined mirror */

The image and footprint plots are shown below as well as plots of the resulting system generates by SYSPLOT.
For comparison, we also show the results of tracing a full parabolical mirror.

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
******************** MIRROR DESCRIPTION ********************
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Sector Example - Full Parabolical Mirror w/o SECTOR
0 deg orientation
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified:end.01
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL SECTOR.PARAB]END.01;1
Creation Date :12-JUN-1989 14:16
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++

Surface figure was defined as: PARABOLICAL
Cylindrical figure: NO
Element type: REFLECTOR
Reflectivity: OFF
Mirror dimensions: UNLIMITED

++++++++++++++++++
Figure 3.50: SECTOR: PLOTSPACE output of SECTORed mirror
Figure 3.51: SECTOR: System in 3-D
Figure 3.52: SECTOR: Side view of resulting system
Figure 3.53: SECTOR: Top view
Central Axis parameters:
Source Plane Distance 1.0000000000000000
Image Plane 5.0000000000000000
Incidence Angle 0.0000000000000000E+00
Reflection/Diffraction Angle 0.0000000000000000E+00

Mirror parameters COMPUTED
Same configuration as Central Axis YES
Objective focus at 1.0000000000000000
Image focus at 5.0000000000000000
Incidence angle 0.0000000000000000E+00

Parameters used follow:
Parabola Param. 2.0000000000000000
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Figure 3.54: SECTOR: PLOTSPACE of full parabolical mirror
Figure 3.55: SECTOR: System in 3-D

Parabolical Mirror
Figure 3.56: SECTOR: Top view
Figure 3.57: SECTOR: Side view
Purpose: To study the resolving power (resolution) of monochromators.

Input: The ray files where the entrance and exit slits will be positioned, along with the maximum and minimum slit sizes, and number of steps for each slit.

Output: A file containing energy and lost ray statistics for each combination of entrance and exit slit width.

Description:

SLITS will aid the user in determining the performance of a monochromator i.e., its resolution. The program answers the basic question of “what is the resolution of the real system for given slit settings?”.

The program is based on the following algorithm: Given two ray files located at the entrance and exit slit positions (e.g. STAR.xx or SCREEN.xxyy), the program sets the width of the entrance slit and then computes the number of rays that are transmitted for a given exit slit setting. From that, the transmitted photon energy distribution (i.e., bandwidth or resolution) is easily computed. The setting of the exit slit is then modified, and the process is repeated to show the trends in the resolution. After the computations have been performed for all exit slit settings, the entrance slit is modified, and the entire process is repeated.

The entrance slit should be set to an opening that brackets the source image. Ideally, the optical system must be designed to relay an image of the source consistent with the required resolution. For example, let us consider a grating in a Rowland geometry. A line source will have a resolution limited by the spherical aberration of the grating. This means that the overall resolution will not be affected too much by the entrance slit size. The image will be (figuratively speaking):

$$\Delta_{\text{image}} = \Delta_{\text{source}} + \Delta_{\text{aberr}}$$

SLITS allows one to easily assess these terms by changing $\Delta_{\text{source}}$ and studying the effects on $\Delta_{\text{image}}$.

The following steps must be taken by the user before running SLITS:

1. All limiting apertures between the desired entrance and exit slit should be turned off in the system. This includes slits at continuation planes and screens.

2. The source should have a continuous energy range. The extent of the range may be varied according to the number of rays lost at the exit slit: if more than 70 percent of rays are lost for the widest slit use a narrower distribution. The energy range may be easily changed with the utility RECOLOR.
3. The system must be retraced after the slits are removed and for each redefinition of the source energy.

The output from SLITS is arranged in rows. Each row presents the statistics for each combination of slit openings. The entries are in the order: Entrance slit width, Exit slit width, Minimum energy passing through exit slit, Maximum transmitted energy, Mean transmitted energy, standard deviation of transmitted energy, number of transmitted rays, and average intensity of the electric field (maximum intensity = 1).

Perhaps the most useful information is the standard deviation and the number of good rays, for these provide luminosity and resolution of the monochromator. The slit widths are varied in increments specified by the user. One of the slits may be held at a constant width to determine the effect of the other slit on the beam as demonstrated in the TGM example below.

After running SLITS, make sure to restore the slits and screens in the system.

**Examples:**

To start let's run SLITS on a Rowland circle grating found in a 2m Grasshopper. The system is described below. We run SLITS for six different entrance slit widths through the same range of exit slit widths. We then plot exit slit vs. the standard deviation on energy to show the resolution of the system.

```
+------------------------------------------------------------------------------------------+
** SOURCE DESCRIPTION **
+------------------------------------------------------------------------------------------+
Source used in SLITS
+------------------------------------------------------------------------------------------+
Input file specified: end.00
Full file Specification :CXRL$DUA0:[XRAYOP.DOCUMENT.UTIL.SLITS.ROWLAND]END.00;
Creation Date :25-AUG-1989 16:30
Random Source.
Generated total 5000 rays.
Source assumed TRIDIMENSIONAL.
Source Spatial Characteristics: GAUSSIAN
 Sigma X : 0.600000000E-01 Sigma Z : 0.100000000E-01
Depth: SYNCHROTRON SOURCE.
Source Emission Characteristics
Distribution Type: SYNCHROTRON
+------------------------------------------------------------------------------------------+
```
Distribution Limits. +X : 0.120000000E-01 -X: 0.120000000E-01 rad
+Z : 0.200000000E-02 -Z: 0.200000000E-02 rad
Magnetic Radius = 2.08330000 m. Beam Energy = 0.80000 GeV.
Beam Emittances. EPSI_X: 0.000000000E+00 EPSI_Z: 0.000000000E+00
Distance from Waist. X: 0.000000000E+00 Z: 0.000000000E+00
Polarization Used: SR TOTAL

Source Photon Energy Distribution: BOX DISTR.
From Photon Energy: 450.000000 or 27.5522667 Angs.
to Photon Energy: 550.000000 or 22.5427636 Angs.

Rowland mount grating used in SLITS Example

Surface figure was defined as: SPHERICAL
Cylindrical figure NO
Element type REFLECTOR
Element type GRATING
Order choosen ( inside are < 0 ) -1.000000000000000
Automatic Tuning YES
Mount Const. INCIDENCE
Grating tuned at [ eV ] 500.0000000000000
Constant ruling [ lines/cm ] 12000.0000000000000
Reflectivity OFF
Mirror dimensions UNLIMITED

Central Axis parameters:
Source Plane Distance 6.978000000000000
Image Plane 25.320000000000000
Incidence Angle 88.00000000000000
Reflection/Diffractive Angle 85.14710385779864

Mirror parameters EXTERNAL
Parameters used follow:
Spherical Radius 200.000000
Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

$ slits
Entrance slit file : begin.dat
Exit slit file : star.01
Units : [ 0 ] eV
[ 1 ] Angstroms
[ 2 ] cm⁻¹

Then ? 0
So far we are working with :
Total number of rays = 5000
Number of rays transmitted = 5000
Intensity of transmitted rays = 5000.000000000000

Maximum entrance slit width : 0.4
Minimum entrance slit width : 0.4
No. of points : 1
Maximum exit slit width : 0.3
Minimum exit slit width : 0.001
No. of points : 101
Output file : slits.dat

$ type slits.dat

<table>
<thead>
<tr>
<th>EnS</th>
<th>ExS</th>
<th>Min NRG</th>
<th>Max NRG</th>
<th>Mean NRG</th>
<th>std. dev</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.40000E+00</td>
<td>0.30000E+00</td>
<td>0.45001E+03</td>
<td>0.54997E+03</td>
<td>0.49729E+03</td>
<td>0.28377E+02</td>
</tr>
</tbody>
</table>

# good rays Avg reflectivity *
4994 0.10000E+01

0.40000E+00 0.29701E+00 0.45001E+03 0.54997E+03 0.49729E+03 0.28377E+02
4994 0.10000E+01
Let's now look at the TGM to determine appropriate values for the entrance and exit slits. The slits are located at the image continuation planes of the first and second OE. We will run SLITS two times: first to determine the width of the entrance slit that begins to limit the divergence of the incoming rays; then with the threshold value of the entrance slit we will look at the resolution (standard deviation of the energy) as the exit slit width varies.

```
$ slits
Entrance slit file :
tgm:star.01
Exit slit file :
tgm:star.02
Units : [ 0 ] eV
       [ 1 ] Angstroms
       [ 2 ] cm-1

Then ?
0
So far we are working with :

  Total number of rays = 1000
  Number of rays transmitted = 927
  Intensity of transmitted rays = 450.9344767072463

Maximum entrance slit width :
.5
Minimum entrance slit width :
.025
No. of points :
20
Maximum exit slit width :
1
Minimum exit slit width :
1
No. of points :
1
```
Figure 3.58: SLITS: Resolution of Rowland circle grating
Notice the saturation of the resolution for large entrance slit sizes.
Output file: entrance.dat

$ 
$ ty entrance.dat

<table>
<thead>
<tr>
<th>x</th>
<th>y</th>
<th>w1</th>
<th>w2</th>
<th>w3</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5000000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.4750000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.4500000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.4250000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.4000000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3750000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3500000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3250000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.3000000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2750000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2500000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2250000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.2000000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.1750000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.1500000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.1250000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.1000000</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00345</td>
</tr>
<tr>
<td>0.2290445</td>
<td>927</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.5000003E-02</td>
<td>1.000000</td>
<td>49.60062</td>
<td>50.39836</td>
<td>50.00284</td>
</tr>
<tr>
<td>0.2289153</td>
<td>925</td>
<td>0.4864450</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Notice that the rays start to be limited by the slit at a width of 0.075cm. We will then use 0.1 cm as our entrance slit width. Any source that has a larger divergence will then be limited to that of the current source (4 mrad in the z direction).

$\text{slits}$

Entrance slit file:

tgm:star.01

Exit slit file:

tgm:star.02

Units: [ 0 ] eV
[ 1 ] Angstroms
[ 2 ] cm⁻¹

Then?

0

So far we are working with:

Total number of rays = 1000
Number of rays transmitted = 927
Intensity of transmitted rays = 450.9344767072463

Maximum entrance slit width:
.1
Minimum entrance slit width:
.1
No. of points:
1
Maximum exit slit width:
1
Minimum exit slit width:
.05
No. of points:
20
Output file:
exit.dat

Now we can look at the energy range for the various exit slit widths:
Figure 3.59: TGM SLITS example
Chapter 4

Macros

The programs are defined as macros because they are command files rather than straight executables. They generally call other programs. The functions of the programs in this category are not necessarily similar; MAKEJD is basically a preprocessor, MAKE_PWR is a postprocessor.
**Purpose:**
To prepare the files describing an insertion device for use by SHADOW. Includes wigglers and transverse undulators.

**Input:**
From terminal.

**Output:**
Files for SHADOW to use to define the source.

**Description:**

MAKE_ID generates the insertion device parameters for both the wiggler source and the undulator source. For both sources, the single electron trajectory is generated. The program then splits to compute the photon radiation depending on the insertion device desired.

**Undulator:**
Since no simple scaling relations exist for an undulator, the photon distribution is function of \((h\omega, \theta, \phi)\) where \(\theta\) and \(\phi\) are the polar and azimuthal angles respectively referred to the SOURCE Y-axis (forward direction). Thus the distribution must be recomputed every time the photon energy range or the polar angle range is changed (\(\phi\) is always taken from 0 to \(2\pi\)), even though the physical parameters of the undulator are the same.

This process makes the undulator calculation very computer time-consuming, so we limit the \((h\omega, \theta, \phi)\) array to a maximum size of \((51,31,31)\). If the user is only looking at the peak of a harmonic, \(\approx 5\) array points in \(\omega\) is sufficient (at least 2 points are needed for the interpolation scheme to work). For a good description of an entire harmonic, \(\approx 10\) to 20 points should be used to cover the wider energy range. Finally to cover more than one harmonic, the full 51 points should be used. For the angular part, we recommend using the \((31,31)\) points whenever possible. This is especially true for the \(\theta\) array because it has more structures, while the features are usually not as sharp in the azimuthal \(\phi\) direction.

For the Undulator Radiation part, the user can choose between entering the parameters from terminal (TT:) or a namelist file. At the end of each run a namelist file called UP-HOT.PAR is always created; this can be used (ITER must be reset to 0 though, see Appendix in User's Guide on User's Program) by the user if he needs to run the program again. For the first pass, one must use the interactive process. We provide a general purpose program to compute the undulator radiation distribution, which is sufficient for most cases; but the user can use his own program if he finds our program insufficient for his needs (see Appendix in User's Guide on User's Program).

To make better use of the array \((h\omega, \theta, \phi)\), the variables are not equally spaced; rather they are put where there are more photons. This involves computing the angular “rings” at each energy for the first pass, and what we call the optimization for the latter passes. At
the end of the first pass, we know the approximate photon distribution. The optimization process then comes in: it finds out where there are higher emission of photons according to the existing distribution, relocates the array points in \((h, \theta, \phi)\), and then recomputes the entire photon distribution for the new array using either the radiation program we provided or the User's Program. The upper and lower limits on the three variables are of course unchanged. This procedure can be repeated as many times as needed, we do it four times for the example. The angular patterns as a result of each optimization are shown here as X'-Z' plots. We note that the overall features are fairly evident even before optimization, while the plot after the last optimization looks more realistic. So the last distribution (stored in CDF100.DAT;5) should be used in subsequent ray tracing. A warning here about artifact: a higher amount of rays may be generated between the rings due to the linear interpolation scheme used\(^1\).

The essential output of MAKE_ID is in CDF100.DAT (CDF100.DAT;1 is before optimization, CDF100.DAT;5 after fourth optimization). It contains the integrated probability distribution function (CDF) ready as input to SOURCE. An unintegrated version is found in RN100.DAT:*\(^*\). They are both unformatted files and use exactly the same format. For SOURCE, we select a zero emittance source. Since the photon energy and the angle range are already pre-determined by MAKE_ID, there is no need to specify them in SOURCE.

Wiggler:
For insertion devices with large K, SHADOW uses the wiggler source. It basically assumes incoherent emission processes, thus the wiggler can be approximated by bending magnets of changing magnetic radius. Then the task is divided into computing: a) electron trajectory, b) photon radiation. They are both computed by MAKE_ID which must be called before running SOURCE.

The correction field factor is used to adjust the end magnetic fields so there is smooth transition in and out of the wiggler. Typically, 101 points is sufficient to describe the trajectory of each period. The output of this first part is in WIGGLER.PAR and WIGGLER.TRAJ. The first file simply stores the parameters for the trajectory, while the second file stores the actual trajectory in the following format:

\[
\begin{array}{ccc}
X & Y & Z \\
\beta_x & \beta_y & 1/R
\end{array}
\]

where \((X,Y,Z)\) is the usual SOURCE reference frame, \(\beta=v/c\), \(R\) the instantaneous radius of curvature. They are both formatted files and metric units (meter) are used.

The second part uses the trajectory file (WIGGLER.TRAJ) from the first part and computes the radiation at each point, within the photon energy range specified by the user. The output

\(^1\)This part of the code is under intensive development and interested users should contact us for the latest version.
put file (RADIAT.DAT) then stores the CDF of photon emission along the wiggler in binary form ready for input to SOURCE. Note that the highest critical energy \( \lambda_c \) is determined by the smallest radius of curvature.

**Examples:**

In the example below, we take an undulator of \( \lambda = 3.5 \text{cm} \), \( K=1.87 \), \( E=1.3 \text{GeV} \) (as suitable for an undulator on SSRL). The correction field factor modifies the magnetic fields at the two ends so that the electron trajectory transits smoothly in and out of the undulator. 101 points are used for each period. That fully defines the trajectory, whose parameters are stored in the file SSRL.PAR and the trajectory itself in SSRL.TRAJ. An optional plottable file SSRL.PLOT is also created for demonstration, which is a formatted 5 columns file in \((X,\beta_x,Y,1-\beta_y,t)\) where \((X,Y,Z)\) is the usual SOURCE coordinates, \( \beta \) is \( v/c \), and \( t \) the time. The computation uses metric units and are equally spaced in the \( Y \)-axis.

In this example we are centered at the peak of the first harmonic, so 7 points in energy are enough. From the parameters of the trajectory file we note that the first harmonic is at 166.81 eV, and so we take the range from 166.6 to 166.9 eV which is still narrower than the width for a 100 periods undulator. For \( \theta \) we go from 0 to 1.0 mrad. Now the calculation is fully defined which involves 6727 (7x31x31) points, and the user can tell our program to update on the screen whenever it finishes every 20 or 50 points.

\[
\$
\]

\[
\$
\]

**Type of Insertion Device.**

Enter:

for wiggler (large \( K \)) [ 1 ]

for undulator (small \( K \)) [ 2 ]

Then ? 2

**Define Insertion Device parameters:**

wavelength of insertion device (m) : 0.035

deflection parameter \( K \) : 1.87

correction field factor (=1) : 1.0
electron energy (GeV) : 1.3
Enter the number of points to be used in the trajectory calculation.
( max = 1001, suggested 101 ) : 101

Two files will be created. One will contain a record of the parameters used in the calculation, the other the trajectory itself. The names of the files can be specified by the user, e.g., MYFILE.PAR and MYFILE.TRAJ.

Output files specification:

Name for parameter file: SSRL.PAR
Name for UNDULATOR trajectory file: SSRL.TRAJ
Do you want a plottable file [ Y/N ]? Y
Name for plottable file: SSRL.PLOT

Calculation Completed. File out results.

Files:
SSRL.TRAJ
SSRL.PAR
SSRL.PLOT

Written to disk.

All Done. Trajectory computed and stored on Disk.
************ UNDULATOR RADIATION ************

Parameters from:
User interactive process    (0)
NAMELIST file              (1)
Choice : 0
Computation done by:
    supplied program     (0)
    user’s program       (1)
Choice : 0

---------------------------------------------------------------------------

Define Radiation Computational parameters:

Polar coordinate is used here.
Number of points in energy (51 max) : 7
    theta (31 max) : 31
    phi (31 max) : 31

---------------------------------------------------------------------------

Enter name of trajectory file from EPATH (input ) : SSRL.TRAJ

---------------------------------------------------------------------------

Undulator case.
Trajectory computed by EPATH with following parameters:

Number of points : 101
Undulator wavelength : 3.500000000000000E-02 meters
Deflection parameter : 1.870000000000000
Peak magnetic field : 0.5722597698020819 tesla
Electron energy : 1.300000000000000 GeV
Gamma : 2543.735230444036
******
Fundamental energy : 166.8162940850640 eV
Fundamental wavelength : 74.33307164977071 Angstrom

Parameters for Radiation Computation.
Please note value of first harmonic from above table.

Enter number of periods: 100
  spectrum starting energy [ eV ] : 166.6
  final [ eV ] : 166.9
  electron current [ A ] : 1.0
Maximum angle between radiation and undulator axis [ mrad ]: 1.0

All undulator parameters defined.
How many times of optimization ? 4
How often do you want a report on calculations ?
E.G., 20, 50,... ? 4500

\\ First pass (before optimization): \\

Begin calculations.

Phi / Horz: Theta / Vert : Energy: CPU Time: % Completed:
  0.209E-03  0.917E-03  167.  290.  66.89

Spectra Computations completed.

Total CPU time used so far: 435.5199794769287
CPU time per point: 6.4742081087695661E-02
Calculation completed.

Begin computation of total power.
Total Power emitted in the specified angles is:
0.1967898468476837 Watts.

Preliminary calculations completed.
Number of optimization finished : 0

Do you want to write out spectra ? 1
Name of file for storing spectra: RN100.DAT
Do you want to create a SHADOW file ? 1
Name of (binary) file for SHADOW: CDF100.DAT
File name for parameter info : UNDULATOR.PAR

Files:
UNDULATOR.PAR
RN100.DAT
CDF100.DAT
written to disk.

\ First optimization : \ 

Begin calculations.
Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.921E-03</td>
<td>167.</td>
<td>280.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.804E-03</td>
<td>167.</td>
<td>283.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.804E-03</td>
<td>167.</td>
<td>283.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

Spectra Computations completed.

Total CPU time used so far: 425.2399902343750
CPU time per point: 6.3213912625891927E-02
Calculation completed.

Begin computation of total power.
Total Power emitted in the specified angles is: 0.2482396424982191 Watts.

Preliminary calculations completed.

Number of optimization finished : 1

Do you want to write out spectra ? 1
Name of file for storing spectra: RN100.DAT
Do you want to create a SHADOW file ? 1
Name of (binary) file for SHADOW: CDF100.DAT
File name for parameter info : UNDULATOR.PAR

Files:
UNDULATOR.PAR
RN100.DAT
CDF100.DAT
written to disk.

\\ Second optimization : \\

Begin calculations.
<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.921E-03</td>
<td>167.</td>
<td>292.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.661E-03</td>
<td>167.</td>
<td>285.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.661E-03</td>
<td>167.</td>
<td>284.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

Spectra Computations completed.

Total CPU time used so far: 428.3701171875000
CPU time per point: 6.3679220631410733E-02

Calculation completed.

Begin computation of total power.
Total Power emitted in the specified angles is:
0.2960115135632488 Watts.

Preliminary calculations completed.

Number of optimization finished: 2

Do you want to write out spectra? 1
Name of file for storing spectra: RN100.DAT
Do you want to create a SHADOW file? 1
Name of (binary) file for SHADOW: CDF100.DAT
File name for parameter info: UNDULATOR.PAR

Files:
UNDULATOR.PAR
RN100.DAT
CDF100.DAT
written to disk.

\ Third optimization : \
Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.690E-03</td>
<td>167.</td>
<td>286.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

---

Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.653E-03</td>
<td>167.</td>
<td>285.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

---

Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.653E-03</td>
<td>167.</td>
<td>286.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

---

Spectra Computations completed.
Total CPU time used so far: 430.8698730468750
CPU time per point: 6.4050821026739259E-02

Calculation completed.

Begin computation of total power.
Total Power emitted in the specified angles is:
0.2668414494602727 Watts.
Preliminary calculations completed.

Number of optimization finished : 3

Do you want to write out spectra? 1
Name of file for storing spectra: RN100.DAT
Do you want to create a SHADOW file? 1
Name of (binary) file for SHADOW: CDF100.DAT
File name for parameter info: UNDULATOR.PAR

Files:
UNDULATOR.PAR
RN100.DAT
CDF100.DAT
written to disk.

Fourth optimization:
Begin calculations.

<table>
<thead>
<tr>
<th>Phi / Horz:</th>
<th>Theta / Vert:</th>
<th>Energy:</th>
<th>CPU Time:</th>
<th>% Completed:</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.209E-03</td>
<td>0.921E-03</td>
<td>167.</td>
<td>283.</td>
<td>66.89</td>
</tr>
</tbody>
</table>

Spectra Computations completed.
Total CPU time used so far: 430.2602539062500
CPU time per point: 6.3960198291400327E-02

Calculation completed.

Begin computation of total power.
Total Power emitted in the specified angles is:
0.2533485171223082 Watts.

Preliminary calculations completed.

Number of optimization finished : 4

Do you want to write out spectra ? 1
Name of file for storing spectra: RN100.DAT
Do you want to create a SHADOW file ? 1
Name of (binary) file for SHADOW: CDF100.DAT
File name for parameter info : UNDULATOR.PAR

Files:
UNDULATOR.PAR
RN100.DAT
CDF100.DAT
written to disk.

FORTRAN STOP
$ DIR

Directory DEVEL: [MAXIMUM.FRANCO.FRINGLE.DEMO]
CDF100.DAT;5  324  13-MAR-1989  23:31  (RE,RWED,RE,RE)
CDF100.DAT;4  324  13-MAR-1989  23:03  (RE,RWED,RE,RE)
CDF100.DAT;3  324  13-MAR-1989  22:33  (RE,RWED,RE,RE)
CDF100.DAT;2  324  13-MAR-1989  21:59  (RE,RWED,RE,RE)
CDF100.DAT;1  324  13-MAR-1989  21:33  (RE,RWED,RE,RE)
RN100.DAT;5  324  13-MAR-1989  23:31  (RE,RWED,RE,RE)
RN100.DAT;4  324  13-MAR-1989  23:03  (RE,RWED,RE,RE)
RN100.DAT;3  324  13-MAR-1989  22:33  (RE,RWED,RE,RE)
RN100.DAT;2  324  13-MAR-1989  21:59  (RE,RWED,RE,RE)
RN100.DAT;1  324  13-MAR-1989  21:33  (RE,RWED,RE,RE)
SSRL.PAR;1    2  13-MAR-1989  21:20  (RE,RWED,RE,RE)
SSRL.PLOT;1   21  13-MAR-1989  21:21 (RE,RWED,RE,RE)
SSRL.TRAJ;1   16  13-MAR-1989  21:21 (RE,RWED,RE,RE)
UNDULATOR.PAR;5 4  13-MAR-1989  23:31  (RE,RWED,RE,RE)
UNDULATOR.PAR;4 4  13-MAR-1989  23:03  (RE,RWED,RE,RE)
UNDULATOR.PAR;3 4  13-MAR-1989  22:33  (RE,RWED,RE,RE)
UNDULATOR.PAR;2 4  13-MAR-1989  21:59  (RE,RWED,RE,RE)
UNDULATOR.PAR;1 4  13-MAR-1989  21:33  (RE,RWED,RE,RE)

Total of 20 files, 3621 blocks.
$ 
\% SSRL.PAR is the parameters file from trajectory calculation. \%
$
$ TY SSRL.PAR
Parameters used for run creating
UNDULATOR case. Trajectory stored in:
SSRL.TRAJ
and in (formatted file):
SSRL.PLOT
The UNDULATOR case uses 1 period only.
------- INPUT ---------
Insertion device Wavel. [ m ] = 3.5000000000000000E-02
Beam Energy [ GeV ] = 1.3000000000000000
K = 1.8700000000000000
Field correction factor = 1.0000000000000000
------- OUTPUT ---------
Gamma = 2543.735230444036
Peak Magnetic field [ Tesla ] = 0.5722597698020819
Fundamental [ Ev ] = 166.8162940850640
Fundamental [ Angstroms ] = 74.33307164977071
Equivalent SR C.W. [ Angs ] = 19.23237966486749
C.E. [ eV ] = 644.6690537546344

SSRL.PLOT is the plottable file of the trajectory. The first 3 points are shown.

UNDULATOR.PAR is the parameter file from the radiation part.

Trajectory computed by EPATH with following parameters:
Number of points : 101
Wavenlen. (und) : 3.50000000000000E-02 meters
Fundamental wvl : 74.33307164977071 angstroms
Fund. energy : 166.8162940850640 eV
K is : 1.870000000000000
Gamma : 2543.735230444036
Beta0 : 0.999999992272728 C units
Field B0 : 0.5722597698020819 tesla
Electron energy : 1.300000000000000 GeV

Read 101 trajectory records from SSRL.TRAJ
Number periods used in ERAD: 100
Total power radiated in the limits [ W ]: 1.8898209057174996E-04
Working with constant dE (units eV).
Energy interval = 0.0000000000000000E+00
Limits: 166.6000000000000 166.9000000000000
Step : 0.0000000000000000E+00 Number of points: 7

POLAR ANGLES CHOSEN
Azimuthal angle (units rad).
Limits: 0.0000000000000000E+00 1.5707963267948966E-03
Step : 0.0000000000000000E+00 Number of points: 31

Polar angle (units mrad).
Limits: 0.0000000000000000E+00 1.0000000000000000
Step : 0.0000000000000000E+00 Number of points: 31
Spectra written into (binary) file:
RN100.DAT
in units: PHOTONS/SEC/eV/RAD**2

$\\ UPHOT.PAR is a namelist file generated at the end of each optimization.\\$

$\$\$ TY UPHOT.PAR
$PARAIN
NCOMP = 100,
RCURR = 1.0000000000000000,
ICOMP = 1,
BPASS = 0.0000000000000000E+00,
IANGLE = 1,
IAPERTURE = 2,
IEXTERNAL = 0,
Shown below are the plots from the first pass and the subsequent optimizations.
Figure 4.1: MAKE_ID: Before optimization
Figure 4.2: MAKE_ID: After first pass
Figure 4.3: MAKE_ID: After second pass
Figure 4.4: MAKE_ID: After third pass
Figure 4.5: MAKE_ID: After fourth pass
Wiggler Example:
This example illustrates how to generate a wiggler source with the following parameters: 
N=10 periods, \( \lambda = 10 \) cm, \( K = 14.94 \), \( E = 1 \) GeV. For the SOURCE part, we use finite emittance to demonstrate its effects which can be seen at both the top view and cross view. The electrons come to a waist at the middle of the wiggler, as we have specified in SOURCE. Most photons are clearly generated near turning points, thus giving rise to the two loops seen when viewing the wiggler head on. Note the entire wiggler is offset to the positive x-axis. The photon energy range, on the other hand, is taken from MAKE_ID (1-100 eV).

\[
\]

$ MAKE_ID

-----------------------------------------------------------------------------
Type of Insertion Device.
Enter:
for wiggler (large K) [ 1 ]
for undulator (small K) [ 2 ]
Then ? 1

-----------------------------------------------------------------------------
Define Insertion Device parameters:

Enter number of periods : 10
    wavelength of insertion device (m) : 0.1
    deflection parameter K : 14.94
definition field factor (=1) : 1.0
electron energy (GeV) : 1.0
Enter the number of points to be used in the trajectory calculation.
    ( max = 1001, suggested 101 ) : 101

-----------------------------------------------------------------------------
Two files will be created. One will contain a record of the parameters used in the calculation, the other the trajectory itself. The names of the files can be specified by the user, e.g., MYFILE.PAR and MYFILE.TRJ.

Output files specification:
Name for parameter file: WIGGLER.PAR
Name for WIGGLER trajectory file: WIGGLER.TRAJ

Trajectory Calculations begins.
Calculation Completed. File out results.

Files:
WIGGLER.TRAJ

************************* WIGGLER RADIATION *************************
Name of input file: WIGGLER.TRAJ
Read 1001 points from input file.
Beam energy (GeV) = 1.000000000174087
Radius of curvature (max.) = 6.2266522507136881E+16 m
(min.) = 2.084481700350993 m
Critical Energy (max.) = 1063.821612058376 eV
(min.) = 3.5613305408526866E-14 eV
Initial photon energy [ eV ] : 1
Final photon energy [ eV ] : 100
Name of output file: RADIAT.DAT
Total no.of photons = 1.7399069816581998E+16

Directory DEVEL:[MAXIMUM.FRANCO.FRIDGE.WIGGLER]
\$ TY WIGGLER.PAR \ Parameter file. \ 
Parameters used for run creating WIGGLER case. Trajectory stored in: WIGGLER.TRAJ

Number of periods = 10

--- INPUT ---
Insertion device Wavel. [ m ] = 0.1000000000000000
Beam Energy [ GeV ] = 1.000000000000000
K = 14.94000000000000
Field correction factor = 1.000000000000000

--- OUTPUT ---
Gamma = 1956.719408033874
Peak Magnetic field [ Tesla ] = 1.600185121255940
Fundamental [ Ev ] = 0.8432615847877075
Fundamental [ Angstroms ] = 14704.76986532713
Equivalent SR C.W. [ Angs ] = 11.62365513397686
C.E. [ eV ] = 1066.662754279258

\$ \ WIGGLER.TRAJ is the actual trajectory. The first three points are listed.

\$ TY WIGGLER.TRAJ

0.0000000000000000E+00 -0.5000000000000000 0.0000000000000000E+00
0.0000000000000000E+00 0.9999998694090963 0.4999999999999999E+00
2.3978852546284019E-07 -0.4999999999999999E+00 0.0000000000000000E+00
4.794199083475524E-04 0.9999997544873504 0.4787889134551865
9.5820947756009422E-07 -0.4980000000000000E+00 0.0000000000000000E+00
9.5694776534338392E-04 0.9999994115344188 0.4759527038051818
Figure 4.6: MAKE_ID: Cross view
Figure 4.7: MAKE_ID: Top view
Purpose: To generate a batch command file for power density calculations on a given optical element or elements.

Input: Mirror file and BEGIN file for element interested in.

Output: A batch file that when submitted generates two data files and two namelist files for each iteration.

Description:
When computing a two-dimensional power distribution it is essential to accumulate a good statistics in order to have a clean result. This program creates a batch file that, when submitted, runs SHADOW iteratively for a specified number of times; at the same time, the power absorbed is accumulated in a file. At the last iteration a file suitable for input to ANSYS is generated. The batch file purges after every run, leaving the two latest versions. The file before the ANSYS file can be used as input to TopDrawer as well.

Examples:
We have turned on the reflectivities and put a gold coating on the first mirror and the grating of the 6m TGM. We run MAKE_PWR to generate a command file that can be submitted in batch mode. When submitted, the intermediate files are in a generic form of X, Y, power, and counts per bin. The data file generated after the last loop is in a form that can be input to ANSYS for further power analysis. It may be useful to use the LOG option when submitting the batch file so if the job does not terminate correctly, you have some indication of the reason.

$ sd [.xrayop.document.tgm]
$ MAKE_PWR

Name for the batch job [ myjob.com ] ? : MYJOB1.COM /*command file */

How many elements do you want to analyze ? : 2 /* analyze the mirror
* and grating */

Defining element: 2 /* grating */

Interactive [0] or namelist mode [1] : 0 /* first time must be interactive */
This program needs a file containing the source spectrum computed within the limits used. Or you can specify the total power.
Source file [0] or total power [1] : 1 /* choose total power */
Total power in Watt : 100

Please specify now the file containing the SOURCE array (e.g., BEGIN.DAT)
Then ? BEGIN.DAT
Read 5000 rays. The file has 18 columns.
Computing renormalizing factor.

All preliminaries completed.
Ready to proceed with density analysis.

File for analysis ?
MIRR.02 /* grating file */
Data ready. Read 5000 points.
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ? 0
***********
File examined is :
Was created :

***********
Found 4483 good points out of 5000
Found:
Column 1: X max is 2.247921404888549
1: X min is -2.248152741557152
2: Y max is 3.733963121236766
2: Y min is -3.241366038463902
3: Z max is 5.5920629967978197E-02
3: Z min is 9.874196393864412E-07
for a photon energy range:
101.1074845359784 to 2971.487995770237
I need to define the two-dimensional histogram for the power density. Please specify:
Columns for x-axis and y-axis?
/* calculate the power on the surface
 * of the mirror ($X,Y$)
 */

Limits to use for the histogram.
To use scaling, enter:
   automatic 0
   external 1
Then? 0
Number of bins in X [ max 51 ] ? 31 /* odd number of bins works best */
   Y [ max 51 ] ? 31

Optical Properties options.
To select enter

Power transmitted/reflected ........ 0
Power absorbed .......................... 1 <<<<
Local reflectivity/transmission .... 2
Incoming Power .......................... 3

Then?
1 /* choose power absorbed */

File to use for Io?
   /* source file, i.e. BEGIN.DAT */
BEGIN.DAT

The program need to know how many iterations have been already performed. Enter
   [ 0 ] if this is the first time.
No. of time this power density case has run : 0
How many times will you be looping through? 50 /* this number will be entered
   again, remember it! */

There were 517 lost out of 5000 rays.

Begin verification.
Total power in file: 83.98999144696705
Compare with: 100.00000000000000
Computations done. Ready to store output data.
The files contain the power density at the nodes but
for the case of ANSYS, that stores total power.
File for:
   TOP DRAWER [ 0 ]
   NCAR [ 1 ]
   Generic [ 2 ] ** to be used for loop
   For ANSYS [ 3 ]
2 /* for running more than once, choose generic */
Output file? GRATING  \* file name, no extension *\nAll done.

Output file name for this element [ ele1.pwr, ele2.pwr, ... ] ? : ELE2G

Defining element: 1  \* first mirror *\n
Interactive [0] or namelist mode [1] : 0
This program needs a file containing the source spectrum computed within the
limits used. Or you can specify the total power.
Source file [0] or total power [1] : 1
Total power in Watt : 100

Please specify now the file containing the SOURCE array (e.g., BEGIN.DAT)
Then ? BEGIN.DAT
Read 5000 rays. The file has 18 columns.
Computing renormalizing factor.

All preliminaries completed.
Ready to proceed with density analysis.

File for analysis?
MIRR.01  \* mirror file *\nData ready. Read 5000 points.
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for including all the losses.
Then ? 0
***********
File examined is:

Was created:

***********
Found 4696 good points out of 5000
Found:
Column 1: X max is 1.999925668739772
   1: X min is -1.998543900746248
   2: Y max is 3.943993346718717
MAKE_PWR

2: Y min is -4.268345358794168
3: Z max is 0.1313206504810491
3: Z min is 6.9529741764107200E-07
for a photon energy range:
101.0305579596815 to 2971.487995770237

I need to define the two-dimensional histogram for the power density. Please specify:
Columns for x-axis and y-axis?
1
2

Limits to use for the histogram.
To use scaling, enter:
automatic 0
external 1
Then? 0

Number of bins in X [ max 51 ] ? 31
Y [ max 51 ] ? 31

Optical Properties options.

To select enter

Power transmitted/reflected ....... 0
Power absorbed ................... 1 <<<
Local reflectivity/transmission .... 2
Incoming Power ................... 3

Then?
1

File to use for Io ? BEGIN.DAT

The program need to know how many iterations have been already performed. Enter [ 0 ] if this is the first time.
No. of time this power density case has run : 0 /* still 0, hasn’t finished */

How many times will you be looping through ? 50 /* same as before */

There were 304 lost out of 5000 rays.

Begin verification.
Total power in file: 72.89175060410695
Compare with: 100.0000000000000

Computations done. Ready to store output data.
The files contain the power density at the nodes but for the case of ANSYS, that stores total power.
File for:
TOP DRAWER [ 0 ]
NCAR [ 1 ]
Generic [ 2 ] ** to be used for loop
For ANSYS [ 3 ]
2
Output file ? MIRROR
All done.

Output file name for this element [ ele1.pwr, ele2.pwr, ... ] ? : ELE1M

Input completed. Prepare batch file.
Job File name ? MYJOB1.COM /* same job name as above */
Command used to define SHADOW environment [ setsha, setup, ... , no $ sign] ? 
@[SHADOW]SETUP
Default directory for calculations ? [SHADOW.XRAYOP.DOCUMENT.TGM]
How many OE’s in this system ? 2 /* # of OE’s for calculations */
How many loops ? 50 /* same loop number again */
Read 2 files.

Batch file ready. Check for the file:
%SHOW-S-NOTRAN, no translation for logical name JOBFILE
in your default directory.
You may use the command:

$submit/notify/noprinter myjob.com

Directory D1:[SHADOW.XRAYOP.DOCUMENT.TGM]

ELE1M.DAT;1 3 31-JUL-1989 10:38:58.42 (RWED,RWED,RE,RE)
ELE2G.DAT;1 3 31-JUL-1989 10:37:28.48 (RWED,RWED,RE,RE)
Total of 6 files, 213 blocks.

$ TY MYJOB1.COM                        " job file to be submitted "
$SET DEFAULT [SHADOW.XRAYOP.DOCUMENT.TGM]
$ASSIGN SYS$COMMAND TT:
$@[SHADOW]SETUP
$YES_TYPE == "SECOND"
$ASSIGN SYS$COMMAND TT:
$PWR_DENS === RUN UTIL:PWR_DENS
$ASSIGN SYS$COMMAND TT:
$SEED === RUN UTIL:SEED
$I = 0
$AGAIN:
$GO
SOURCE
BATCH
START.00                " if you have renamed the namelist
TRACE                 " files for your source and system
BATCH                 " you must edit the command file
0                    " and replace the defaults with
START.01             " the correct names or rename your
0                    " namelist files to START.xx    "
START.02
0
%EXIT
EXIT
$ASSIGN/USER_MODE ELE2G PWR_DENS_PAR
$PWR_DENS
1
$ASSIGN/USER_MODE ELE1M PWR_DENS_PAR
$PWR_DENS
1
$PURGE/KEEP=2           " only two latest versions kept "
$SEED
START.00              " make sure to change name of namelist
After the batch job was completed, the latest generic data files were run through a contouring program. The results were plotted with TopDrawer. The plots show the outline of the mirror and grating and how the power is absorbed on each.
Figure 4.8: MAKE_PWR: Mirror
Figure 4.9: MAKE_PWR: Grating
Chapter 5

Optics Utilities

The optical utilities are mainly standalone programs that perform optical calculations that may be of use when running SHADOW. They provide information about element/compound characteristics, grating information, multilayer calculations, synchrotron spectra and the resulting transmission through a system. They do not generate any files that can be input to SHADOW itself, however similar utilities for some exist in Preprocessors. Again we have broken them down into several sections for convenience: Optical properties, sources, and geometrical optics.
Purpose: Computes several properties of a given material (over a specified photon energy range) that are important for physical optics calculations.

Input:

From terminal:

1. Material Density
2. Material formula
3. Photon range

Output:

ABSOR_COEFF.DAT
CROSS_SECTION.DAT
DIELEC_FUNC.DAT
ELECTRON_YIELD.DAT
ENERGY_LOSS.DAT
REFLECTIVITY.DAT
REFRACT_INDEX.DAT
TRANSMISSION.DAT

Description:

The program is based on the compilation of atomic scattering factor by B.Henke, with extension to 10,000 by G.Tirsell. We have extended it to 100,000 eV by using the tables from Cromer and Liberman. The division is then:


The program requires no user input beside the knowledge of the material atomic composition and its density. The formulae used are described in detail in B.Henke’s article in Low Energy X-ray Diagnostics and are valid for the x-ray region.
The program can compute the optical properties of materials containing up to five atomic species. Notice that when specifying the formula the atomic symbols must be capitalized and two-letters long (e.g. SI for silicon and WW for tungsten). Another input parameter needed by the program is the density, $\rho$, in g/cm$^3$ of the material.

Finally the program prompts for “Full printout at terminal”. If the user answers yes, then all the results are printed to the terminal beside being saved in a file. This is useful only if one is interested in the value at one or two photon energies.

The files are saved with fixed file names. It is a good idea to rename them immediately (e.g. REFLECTIVITY.DAT $\rightarrow$ REFLEC.AU,89) to keep things orderly.

The following properties are computed over the specified photon energy range:

- **Dielectric Function**: The complex dielectric function is computed and stored in a file called DIELEC_FUNC.DAT. The units are in $\text{farad/cm}$ and the program stores $\alpha, \beta$ where:
  \[ \tilde{\varepsilon} = 1 - \alpha + i\beta \]  
  (5.1)

- **Absorption Coefficient**: Units are in $\text{cm}^{-1}$ and the result is stored in the file ABSOR_COEFF.DAT.

- **Transmission**: The transmission through a film of thickness $d$ (as specified by the user):
  \[ T(h\omega) = e^{-\mu(h\omega)d} \]  
  (5.2)

  and is stored in the file TRANSMISSION.DAT.

- **Reflectivity**: The $s$– and $p$–components of the reflectivity $R$ are computed from the Fresnel equations and the dielectric function as obtained above. The file includes average, $s$– and $p$– reflectivities.

- **Refraction Index**: This value is again obtained from the dielectric function above ($n^2 = \varepsilon$); only the real part is stored. The complex refractive index is of the form: $1 - \delta - i\beta$. The output is formatted (ANSI).

- **Cross Section**: for photoelectric effect. Obtained directly from $f_1, f_2$ in $\text{cm}^2$. It is the basic “ingredient” of the absorption coefficient $\mu$.

- **Electron Yield**: as defined by B.Henke. In essence, the primary electron yield is proportional to the square of the electric field at the surface of the material.

- **Energy Loss**: The rate at which energetic electrons lose energy when travelling through a medium is proportional to the imaginary part of the dielectric function.
This is obtained directly from the linear medium response theory in the limit of small $q$ (Raether, 1960),

$$R(\Delta E) = -\text{Im} \frac{1}{\epsilon}$$  \hspace{1cm} (5.3)

The files are all in a two or three columns tabular format and can be plotted out using TopDrawer. It may be a good idea to run CLIP on the files if an error is obtained when running TopDrawer because of possible underflows.

**Examples:**

```
$ abrefc
Full printout at Terminal ?
n \(* suppresses extra output *\)
Thickness (microns) ?
1.0 \(* needed for transmission *\)
The program is setup to compute optical constants of
compounds with up to 5 elements.
How many atomic species ?
2 \(* choose a compound *\)
H2O would be: HH,2 and OO,1. Then ?
Enter 2-letters (capitalized) atomic symbol and formula index for : 1
SI \(* silicon *\)
1
Enter 2-letters (capitalized) atomic symbol and formula index for : 2
CC \(* carbon *\)
1
Density ?
3.44 \(* g/cm^3 *\)
Enter starting photon energy,end and step
200,5000,20 \(* computation range *\)
Incidence angle ?
88 \(* for reflectivity *\)
Molecular weight is :  40.10000
Atoms/Cm3
5.1660890E+22 5.1660890E+22 0.000000E+00 0.000000E+00 0.000000E+00
$ dir

Directory D13:[XRAYOP.DOCUMENT.UG.ABREFC]
```
Total of 9 files, 149 blocks.

$ $ $ $ type absor_coeff.dat   \ units are in cm$^{-1}$

```
200.0000  179319.5
220.0000  148879.4
240.0000  124486.7
260.0000  104718.1
280.0000   88981.34
300.0000   125351.2
320.0000   107683.3
340.0000    93240.84
360.0000    81363.08
380.0000    71451.71
400.0000   63132.95
420.0000   56095.28
440.0000   50092.01
460.0000   44927.46
480.0000   40454.81
500.0000   36563.99
520.0000   33177.92
540.0000   30197.32
560.0000   27553.70

.....     ....
```

$ $ $ $ t cross_section.dat   \ cm$^2$

```
200.0000  3.4710877E-18
220.0000  2.8818582E-18
```
<table>
<thead>
<tr>
<th>photon</th>
<th>real</th>
<th>imaginary</th>
</tr>
</thead>
<tbody>
<tr>
<td>energy</td>
<td>part</td>
<td>part</td>
</tr>
<tr>
<td>200.0000</td>
<td>2.3322932E-02</td>
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<td>1.3353705E-02</td>
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<td>1.0406492E-02</td>
<td>6.2709148E-03</td>
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<tr>
<td>300.0000</td>
<td>1.1901873E-02</td>
<td>8.2451245E-03</td>
</tr>
<tr>
<td>320.0000</td>
<td>1.1594478E-02</td>
<td>6.6403067E-03</td>
</tr>
<tr>
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<td>1.0792673E-02</td>
<td>5.4114950E-03</td>
</tr>
<tr>
<td>360.0000</td>
<td>9.9309748E-03</td>
<td>4.4597946E-03</td>
</tr>
<tr>
<td>380.0000</td>
<td>9.1089029E-03</td>
<td>3.7103852E-03</td>
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<td>8.3512999E-03</td>
<td>3.1144836E-03</td>
</tr>
<tr>
<td>420.0000</td>
<td>7.6613128E-03</td>
<td>2.6355239E-03</td>
</tr>
<tr>
<td>440.0000</td>
<td>7.0428853E-03</td>
<td>2.2464965E-03</td>
</tr>
<tr>
<td>460.0000</td>
<td>6.4875023E-03</td>
<td>1.9272763E-03</td>
</tr>
<tr>
<td>480.0000</td>
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</tr>
<tr>
<td>500.0000</td>
<td>5.5409130E-03</td>
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$ type dielec_func.dat

$ type refrac_index.dat

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<tr>
<td>260.0000</td>
<td>2.4264235E-18</td>
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<tr>
<td>280.0000</td>
<td>2.0844250E-18</td>
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<tr>
<td>300.0000</td>
<td>1.8048632E-18</td>
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<td>1.5749454E-18</td>
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<tr>
<td>340.0000</td>
<td>1.3830910E-18</td>
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<td>360.0000</td>
<td>1.2206473E-18</td>
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<tr>
<td>400.0000</td>
<td>9.6963125E-19</td>
</tr>
<tr>
<td>420.0000</td>
<td>8.6966089E-19</td>
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<td>7.6613128E-03</td>
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<tr>
<td>460.0000</td>
<td>6.4875023E-03</td>
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<td>480.0000</td>
<td>5.9888614E-03</td>
</tr>
<tr>
<td>500.0000</td>
<td>5.5409130E-03</td>
</tr>
</tbody>
</table>

$ type dielec_func.dat

$ type refrac_index.dat
<table>
<thead>
<tr>
<th>energy</th>
<th>part</th>
</tr>
</thead>
<tbody>
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<td>7.5257206E-03</td>
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<td>280.0000</td>
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<td>300.0000</td>
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<td>320.0000</td>
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</tr>
<tr>
<td>500.0000</td>
<td>2.7704565E-03</td>
</tr>
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</table>

```
$ type reflectivity.dat
```

<table>
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<th>photon</th>
<th>r</th>
<th>r_s</th>
<th>r_p</th>
</tr>
</thead>
<tbody>
<tr>
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<td></td>
<td></td>
</tr>
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<tr>
<td>220.0000</td>
<td>0.7695174</td>
<td>0.7719697</td>
<td>0.7670650</td>
</tr>
<tr>
<td>240.0000</td>
<td>0.7701561</td>
<td>0.7722075</td>
<td>0.7681047</td>
</tr>
<tr>
<td>260.0000</td>
<td>0.7644928</td>
<td>0.7662324</td>
<td>0.7627532</td>
</tr>
<tr>
<td>280.0000</td>
<td>0.6992385</td>
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<td>0.6977182</td>
</tr>
<tr>
<td>300.0000</td>
<td>0.6961607</td>
<td>0.6979848</td>
<td>0.6943366</td>
</tr>
<tr>
<td>320.0000</td>
<td>0.7215253</td>
<td>0.7230968</td>
<td>0.7199538</td>
</tr>
<tr>
<td>340.0000</td>
<td>0.7340940</td>
<td>0.7354625</td>
<td>0.7327254</td>
</tr>
<tr>
<td>360.0000</td>
<td>0.7424233</td>
<td>0.7436262</td>
<td>0.7412204</td>
</tr>
<tr>
<td>380.0000</td>
<td>0.7487475</td>
<td>0.7498126</td>
<td>0.7476823</td>
</tr>
<tr>
<td>400.0000</td>
<td>0.7537164</td>
<td>0.7546662</td>
<td>0.7527666</td>
</tr>
<tr>
<td>420.0000</td>
<td>0.7576135</td>
<td>0.7584658</td>
<td>0.7567613</td>
</tr>
<tr>
<td>440.0000</td>
<td>0.7609170</td>
<td>0.7616860</td>
<td>0.7601480</td>
</tr>
<tr>
<td>460.0000</td>
<td>0.7636897</td>
<td>0.7643870</td>
<td>0.7629924</td>
</tr>
<tr>
<td>480.0000</td>
<td>0.7660290</td>
<td>0.7666641</td>
<td>0.7653939</td>
</tr>
</tbody>
</table>

......
$\text{type transmission.dat}$

<table>
<thead>
<tr>
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<th>transmission</th>
</tr>
</thead>
<tbody>
<tr>
<td>200.0000</td>
<td>1.6302495E-08</td>
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<td>220.0000</td>
<td>3.4217771E-07</td>
</tr>
<tr>
<td>240.0000</td>
<td>3.9229426E-06</td>
</tr>
<tr>
<td>260.0000</td>
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<tr>
<td>420.0000</td>
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<tr>
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<td>6.6762352E-03</td>
</tr>
<tr>
<td>460.0000</td>
<td>1.1189881E-02</td>
</tr>
</tbody>
</table>

$\text{type energy_loss.dat}$

<table>
<thead>
<tr>
<th>photon energy</th>
<th>Im (1/epsilon)</th>
</tr>
</thead>
<tbody>
<tr>
<td>200.0000</td>
<td>1.8541420E-02</td>
</tr>
<tr>
<td>220.0000</td>
<td>1.3918069E-02</td>
</tr>
<tr>
<td>240.0000</td>
<td>1.0608352E-02</td>
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<tr>
<td>260.0000</td>
<td>8.1918687E-03</td>
</tr>
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</tr>
<tr>
<td>420.0000</td>
<td>2.6763568E-03</td>
</tr>
</tbody>
</table>
The files are plotted in the following two figures.
Figure 5.2: ABREFC output continued
**Purpose:** Computes the reflectivity cutoff angle for a given element.

**Input:** From the Keyboard.

**Output:** To two \{x,y\} files, CUTOFFDEG.DAT (degrees, reflectivity) and CUTOFF.DAT (radians, reflectivity).

**Description:**

The program is based on the optical constants described in ABREFC. At each photon the cutoff wavelength, i.e., the extrapolation to zero is computed. The formula is defined by B. Henke in *Low Energy X-ray Diagnostics*. Notice that for an absorbing system this wavelength is not well defined as it is for a dielectric.

**Examples:**

We will consider the case of a gold mirror.

```
$CUTOFF
Enter 2-letters (capitalized) atomic symbol ?
AU
Density ?
19.7
Enter starting photon energy, end and step
200,5000,25
$
$ dir *.dat

Directory D13:[XRAYOP.DOCUMENT.CUTOFF]

CUTOFF.DAT;2 13 25-FEB-1989 13:02 (RE,RWED,RWED,RE)
CUTOFFDEG.DAT;2 13 25-FEB-1989 13:02 (RE,RWED,RWED,RE)

Total of 2 files, 26 blocks.
$
$```

```
Figure 5.3: CUTOFF example using gold

TopDrawer was then used to generate the plot displayed here.
Purpose: Computes the absorbed X-ray dose in a given material vs. depth.

Input:
- Incoming power data file (as generated by TRANSMIT)
- Absorption coefficient file (as generated by ABREFC)

Output:
A file $z$, dose with the value of the absorbed dose within the material.

Description:
The program relies on two preprocessors, TRANSMIT and ABREFC to generate respectively the incoming power density file and the absorption coefficient. Beer's law is used to compute the local absorbed power. Particularly useful for radiation damage studies. No geometrical information is used so that the implicit geometry is that of the x-rays incoming normally onto the sample.

Examples:
In order to demonstrate the program, we will prepare a system consisting of a thin Be window followed by an X-ray mask; an organic material (a resist) is located behind the mask and absorbs the radiation. We will compute the power density absorbed in function of the depth in this material. The absorbed dose is given by:

$$ D(z) = \int_{E_1}^{E_2} I(z, h\omega)\mu(h\omega) \, d\, h\omega $$  \hspace{1cm} (5.4)

$$ I(z, h\omega) = I_o e^{-\mu(h\omega)z} $$  \hspace{1cm} (5.5)

where $\mu$ is the absorption coefficient. It is necessary to compute the incoming power density file and the absorption coefficient of the material under study. TRANSMIT generates a file that represents the Watts of radiation incident on a given area, hence the units are W/cm$^2$. Since $\mu$ is in [cm$^{-1}$], the dose is obtained in W/cm$^3$. If one is interested in a monochromatic or narrow band source, then create with EDT or another program, a file of the same format as TRANSMIT, i.e., \{photon energy, power\}.

Note: care must be taken that the different files be coherent for the photon energy range and energy steps.

$TRANSMIT$
Do you want to generate a source file ?: y
Machine name ?
Aladdin /* obviously */
Title ?
Demo for Dose Session
Machine Radius [ m ] and Beam Energy [ GeV ] ?
2.0833
1.0
Current [ A ] and milliradians of orbit (horizontal) ?
.100
5
Minimum photon energy, maximum, step :
200,2500,25 /* calculation range */
Spectrum type:
dE = constant 0 [ Photons/sec/eV ]
dL = constant 1 [ Photons/sec/Angs ]
dL/L = constant 2 [ Photons/sec/%bandwidth ]
dE/E = constant 3 [ Photons/sec/%bandwidth ]
Power dL=const 4 [ Watt/Angs ]
Power dE=const 5 [ Watt/eV ] /* use this */
Then ? 5
Do you want to specify the vertical acceptance ? 0
Read 1001 records.
+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Spectrum completed
+++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
: DATA file roots for local properties ?
> loc
: CUMULATIVE DATA file roots ?
> tot
Do you want a conversion to Watts/cm^2 ? [Y/N] ? > y /* not neccessary */
Please enter the vertically scanned height (including overshoot) in CM.
> 10
The program will assume that you have used 1 mrad of orbit. If you have specified 1 mrad when creating the source, simply enter 1. If not, enter the number of milliradians you used.
Number of milliradians used: 5
Please enter the horizontal dimension of the exposed field [cm].
> 2.0 /* we will use an area of 20 cm^2 */
Distance from the tangent point [m] ?
> 5 /* located at 5 m from the source */
Total area is: 20.00000
All set.
Read 93 records.
Initial power is: 0.1030330

Currently at Element N.  0

: Enter a negative number to exit, zero to cancel; OE ?
> 1
: Enter [ 0 ] for pure element, [ 1 ] for compound
> 0
Enter 2-letters (capitalized) atomic symbol ?
BE
Density ?
1.98
Mirror [ 1 ] or filter [ 0 ] ?
0
Thickness (microns) ?
15
OUT > Power density [W/cm^2] is 3.0929089E-02
Comment for this OE > Op First filter

Currently at Element N.  1

: Enter a negative number to exit, zero to cancel; OE ?
> 2
: Enter [ 0 ] for pure element, [ 1 ] for compound
> 1
Density [ g/cm3 ] ?
4.6
Mirror [ 1 ] or filter [ 0 ] ?
0
Thickness (microns) ?
2
The program is setup to compute optical constants of compounds with up to 5 elements.
How many atomic species ?
3
H2O would be: HH,2 and OO,1. Then ?
Enter 2-letters (capitalized) atomic symbol and formula index for : 1
BB
4
Enter 2-letters (capitalized) atomic symbol and formula index for : 2
NN
1
Enter 2-letters (capitalized) atomic symbol and formula index for : 3
HH
1
OUT > Power density [W/cm^2] is 2.0523597E-02
Comment for this OE > This could be an X-ray mask membrane

Currently at Element N. 2

: Enter a negative number to exit, zero to cancel; OE ?
> -2

$ dir *.dat

Directory D13:[XRAYOP.DOCUMENT.DOSE]

LOC1.DAT;1 6 25-FEB-1989 13:25 (RE,RWED,RWED,RE)
LOC2.DAT;1 6 25-FEB-1989 13:26 (RE,RWED,RWED,RE)
MACHINE.DAT;1 3 25-FEB-1989 13:24 (RE,RWED,RWED,RE)
SRCOMP.DAT;1 6 25-FEB-1989 13:24 (RE,RWED,RWED,RE)
TOT1.DAT;1 6 25-FEB-1989 13:26 (RE,RWED,RWED,RE)
TOT2.DAT;1 6 25-FEB-1989 13:26 (RE,RWED,RWED,RE)
TRANSMIT.DAT;1 4 25-FEB-1989 13:24 (RE,RWED,RWED,RE)
TRANSMIT.LOG;1 6 25-FEB-1989 13:26 (RE,RWED,RWED,RE)

Total of 8 files, 43 blocks.

$ $ /* verify that the limits are OK */
$ fmax
File-name ?
tot1.dat
Read 93 points.
Found:
XMIN 200.0000 . Set at : -58.75000
Good luck!

$ \* compute absorption coefficient *$

$ ABREFC$

Full printout at Terminal? 0

Thickness (microns)? 2

The program is setup to compute optical constants of compounds with up to 5 elements.

How many atomic species? 3

H2O would be: HH,2 and OO,1. Then?

Enter 2-letters (capitalized) atomic symbol and formula index for: 1 CC

7

Enter 2-letters (capitalized) atomic symbol and formula index for: 2 OO

6

Enter 2-letters (capitalized) atomic symbol and formula index for: 3 HH

11

Density? 2.03

Enter starting photon energy, end and step 200,2500,25

Incidence angle? 0

Molecular weight is: 191.1100

Atoms/Cm3

4.477364E+22 3.8380595E+22 7.0364425E+22 0.0000000E+00 0.0000000E+00

$ dir abs* \* this file contains the abs. coeff. in cm-1 *$

Directory D13:[XRAYOP.DOCUMENT.DOSE]
Total of 1 file, 6 blocks.

$ dose
Power density file ? tot2.dat
Absorption Coefficient File ? absor_coeff.dat
Output file ? dose abs_dose
Files read OK.
Thickness (MICRONS) ? 2  /* film thickness */
Step size (MICRONS) ? 0.05 2  /* calculation step -- 41 points */
Area over which power is spread [ cm2 ] ? 20
All calculations completed

$ type abs_dose.dat

$ dir .dat;0

Directory D13:[XRAYOP.DOCUMENT.DOSE]

ABS_COEFF.DAT;1 6 25-FEB-1989 13:29 (RE,RWED,RWED,RE)
ABS_DOSE.DAT;1 7 25-FEB-1989 13:30 (RE,RWED,RWED,RE) <<<<<<
CROSS_SECTION.DAT;1 6 25-FEB-1989 13:29 (RE,RWED,RWED,RE)
DIELEC_FUNC.DAT;1 9 25-FEB-1989 13:29 (RE,RWED,RWED,RE)
ELECTRON_YIELD.DAT;1 6 25-FEB-1989 13:29 (RE,RWED,RWED,RE)
ENERGY_LOSS.DAT;1 6 25-FEB-1989 13:29 (RE,RWED,RWED,RE)
Total of 16 files, 101 blocks.

\* The session of Transmit is recorded in TRANSMIT.LOG *\n
$ type transmit.log

Demo Session

Parameters of Aladdin

Magnetic Radius : 2.083300 meters
Beam energy : 1.000000 GeV
Gamma : 1957.000
Magnetic Field : 1.600695 tesla

Critical ENERGY : 1064.883 eV
WAVELENGTH: 11.64308 Angstroms
Power/Amp : 42.48068 kilowatts/amp
Total Power : 4.248068 kilowatts
at current of : 0.1000000 Amperes.
Total Power/mrad : 0.6761010 watts/mrad

For unlimited vertical acceptance,
power accepted : 3.380505 watts
hori. acceptance : 5.000000 mrad
Photon energy range used:

\[
\begin{align*}
\text{start} & : 200.0000 \\
\text{end} & : 2500.000 \\
\text{step} & : 25.00000
\end{align*}
\]

Results are in Watts/cm\(^2\).

Power in source file [W] = 2.575826
Full power in W/cm\(^2\) = 0.1030330

Exposure field is defined by:

\[
\begin{align*}
\text{Horizontal [cm]} & = 2.000000 \\
\text{Vertical [cm]} & = 10.00000 \\
\text{Area [cm]} & = 20.00000 \\
\text{Milliradians} & = 4.000000 \\
\text{Distance [m]} & = 5.000000
\end{align*}
\]

Optical Element : 1
Defined FILTER 15.000 microns thick
Working with \( Z = 4 \) Symbol: BE
Atomic Weight: 9.010000 and density 1.980000 g/cm\(^3\)

OE # 1
First filter
OUT > Power density [W/cm\(^2\)] is 3.0929089E-02

Optical Element : 2
Defined FILTER 2.0000 microns thick

Formula: BB(4) NN(1) HH(1)

Molecular weight is: 58.21000 Number of molecules/cm³: 4.7589159E23

Atoms/Cm³ of each species:

<p>| | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>BB</td>
<td>NN</td>
<td>HH</td>
</tr>
<tr>
<td>0.19036E+24</td>
<td>0.47589E+23</td>
<td>0.47589E+23</td>
</tr>
</tbody>
</table>

Density 4.600000 g/cm³

---------------------------

OE # 2
This could be an X-ray mask membrane

OUT > Power density [W/cm²] is 2.0523597E-02

$
$
$
$
$
Figure 5.4: DOSE: Graphical output from TRANSMIT
Figure 5.5: DOSE: Dose seen by photo-resist
MLAYER

Purpose: A standalone program to compute the reflectivity of multilayers. (J.H. Underwood)
Input: Multilayers defined either from terminal or by a formatted file.
Output: Formatted (8 columns) file.

Description:

MLAYER is not really a utility to SHADOW but rather, a standalone program written by J. H. Underwood of the Center for X-ray Optics. It can calculate the reflectivity of multilayer as a function of a) incident angle or b) incident wavelength, leaving the other one fixed. MLAYER also assumes that a narrow wavelength range will be used, so that a set of constant refractive indices (supplied by the user) is sufficient. Thus a possible sequence of events may be:

1. Run MLAYER to get an idea of how the multilayers perform
2. Run PRE_MLAYER to set up the multilayers
3. Run SHADOW for a complete simulation.

MLAYER has the following options:

<table>
<thead>
<tr>
<th>OPTIONS</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>fixed wavelength, scan angle, periodic layers</td>
</tr>
<tr>
<td>2</td>
<td>fixed angle, scan wavelength, periodic layers</td>
</tr>
<tr>
<td>3</td>
<td>fixed wavelength, scan angle, individual layers defined</td>
</tr>
<tr>
<td>4</td>
<td>fixed angle, scan wavelength, individual layers defined</td>
</tr>
<tr>
<td>5</td>
<td>fixed angle, scan wavevector, individual layers defined</td>
</tr>
</tbody>
</table>

Whether the multilayers are periodic or not depends on whether the absorbing and non-absorbing materials all have the same thicknesses for all layers. Of course, the user can always define the thicknesses one by one.

For the example here we use the same W-C multilayers as in PRE_MLAYER (see PRE_MLAYER).

We use option 3 to scan the glancing angle from 0 to 90 degrees, at a fixed wavelength of 18 Å. Then the user can direct the program to read all the input parameters from a file or type them in from the terminal. For the first pass, we use the latter mode. We proceed to define the complex refractive indices of the substrate and the multilayers by δ and β:

\[ n = (1 - \delta) - i \times (\beta) \]
The $\delta$'s and $\beta$'s can in turn be obtained at the energy you are interested in from ABREFC. We put the wavelength at 18 Å and the initial angle 0 degrees. Using a step size of 0.5 and 181 points, this will take us up to 90 degrees. Had we chosen to scan the wavelength instead (option 4), the same step size and number of points will scan from 18 to 108 Å (but then the refractive indices are not nearly constant). After that, we type in the individual thickness of the layers. This process fully specifies the multilayers, and the parameters can be saved in a file (CW.PAR). In future run, the user can direct MLAYER to read from the file, saving the labor of re-defining the multilayers.

The output of this calculation is saved in a formatted file (CW.OUT). There are eight columns and they have the following meanings:

<table>
<thead>
<tr>
<th>COLUMN</th>
<th>Meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$\lambda$ (or $\delta$, whichever is varied)</td>
</tr>
<tr>
<td>2</td>
<td>S reflectivity</td>
</tr>
<tr>
<td>3</td>
<td>P reflectivity</td>
</tr>
<tr>
<td>4</td>
<td>Average reflectivity</td>
</tr>
<tr>
<td>5</td>
<td>S phase shift</td>
</tr>
<tr>
<td>6</td>
<td>P phase shift</td>
</tr>
<tr>
<td>7</td>
<td>S (electric field)$^2$</td>
</tr>
<tr>
<td>8</td>
<td>P (electric field)$^2$</td>
</tr>
</tbody>
</table>

where S is the perpendicular component and P the parallel component (to the plane of incidence). Phase shift is in units of $\pi$.

Examples:

```
\ For the geometry, refer to the example in PRE_MLAYER. \n
$  
$ MLAYER
option 1, 2, 3, 4, or 5? (0 = help, 6 = exit)
0  
  \ Help menu \n
1 = fixed wavelength, scan angle, periodic layers
2 = fixed angle, scan wavelength, periodic layers
3 = fixed wavelength, scan angle, individual layers defined
4 = fixed angle, scan wavelength, individual layers defined
5 = fixed angle, scan wavevector, individual layers defined
option 1, 2, 3, 4, or 5? (0 = help, 6 = exit)
3
use existing input file? (cr = yes)
```
N
input multilayer parameters
substrate material is    . (cr = no change)    \ Substrate \\
SI
delta =    0.00000,   beta =    0.00000(, = no change)
1.0E-3,1.0E-4
odd layer material is    . (cr = no change)    \ Odd layer \\
W
delta =    0.00000,   beta =    0.00000(, = no change)
1.1E-2,1.25E-2
even layer material is    . (cr = no change)    \ Even layer \\
C
delta =    0.00000,   beta =    0.00000(, = no change)
1.3E-3,1.86E-4
lambda(begin) = 0.00000   theta(begin) = 0.00000(, = no change)
18.0,0.0                   \ wavelength, angle \\
step size = 0.00000   number of steps = 0(, = no change)
0.5,181
number of layer pairs = 0(, = no change)
20
for layer pair  1,
odd thickness = 0.00000    even thickness = 0.00000 (, = no change)
7.71,14.56
for layer pair  2,
odd thickness = 0.00000    even thickness = 0.00000 (, = no change)
7.72,14.57
for layer pair  3,
odd thickness = 0.00000    even thickness = 0.00000 (, = no change)
7.74,14.59
for layer pair  4,
odd thickness = 0.00000    even thickness = 0.00000 (, = no change)
7.74,14.59
for layer pair  5,
odd thickness = 0.00000    even thickness = 0.00000 (, = no change)
7.74,14.59
::     ::     ::     ::     ::     ::     ::
for layer pair 19,
    odd thickness = 0.00000 even thickness = 0.00000 (, = no change)
7.74,14.59
for layer pair 20,
    odd thickness = 0.00000 even thickness = 0.00000 (, = no change)
7.74,14.59
save layer parms in which file? (cr = don’t save)
CW.PAR
name of output file:
CW.OUT
option 1, 2, 3, 4, or 5? (0 = help, 6 = exit)
6
FORTRAN STOP
$
$ DIR CW

Directory D13:[XRAYOP.DOCUMENT.MLAYER]

CW.OUT; 1   36 23-FEB-1989 23:13 (RWED,RWED,RE,RE)
CW.PARCN; 1  2 23-FEB-1989 23:13 (RWED,RWED,RE,RE)

Total of 2 files, 38 blocks.

\ CW.PARCN can be edited or simply reused for input to MLAYER. \n
$
$ TY CW.PARCN
SI    \ Substrate
  0.00100  0.00010 \ delta, beta
W     \ Odd layer
   0.01100  0.01250 \ delta, beta
C     \ Even layer
  0.00130  0.00019 \ delta, beta
18.00000  0.00000 \ wavelength, angle
  0.50000  181 \ step size, no. of points
 20    \ no. of layers
  7.71000  14.56000 \ thickness (odd, even)
  7.72000  14.57000
  7.74000  14.59000
  7.74000  14.59000
\[ \text{CW.OUT is the output we are interested in.} \]

\$  
$ \text{TY CW.OUT}  
\text{/* the file CW.OUT contains the eight values on a single line, however to display the values, we have continued them on the next line. */}  
\[ 
\begin{array}{ccccccc}
90.000000 & 0.000025 & 0.000025 & 0.000025 & 1.988236 & 0.988236 \\
1.009956 & 1.009956 \\
89.500000 & 0.000025 & 0.000025 & 0.000025 & 1.988777 & 0.988776 \\
1.009947 & 1.009945 \\
89.000000 & 0.000024 & 0.000024 & 0.000024 & 1.990377 & 0.990375 \\
1.009917 & 1.009911 \\
88.500000 & 0.000024 & 0.000024 & 0.000024 & 1.992974 & 0.992969 \\
1.009863 & 1.009849 \\
\end{array} \]

\[ 
\begin{array}{ccccccc}
: & : & : & : & : & : \\
: & : & : & : & : & : \\
1.500000 & 0.703394 & 0.696951 & 0.700172 & 0.892246 & 0.895151 \\
0.121218 & 3.276859 \\
\end{array} \]
Now let us graph the average reflectivity (col. 4) vs the angle (col. 1) using TOPDRAWER.

$ TDTEK
TOP DRAWER (10/10/80) NOW EXECUTING

MOD NOECHO
1 MOD NOECHO

SET ORDER X DUMMY DUMMY Y DUMMY DUMMY DUMMY DUMMY DUMMY
2 SET ORDER X DUMMY DUMMY Y DUMMY DUMMY DUMMY DUMMY

@CW.OUT
READING FROM INDIRECT COMMAND FILE: CW.OUT
3 90.000000 0.000025 0.000025 0.000025 1.988236
   0.988236 1.009956 1.009956
END OF INDIRECT COMMAND FILE
JOIN 1
   184 JOIN 1
END
   185 END
Figure 5.6: MLAYER: S-reflectivity vs. angle
Figure 5.7: MLAYER: P-reflectivity vs. angle
Figure 5.8: MLAYER: Average reflectivity
Purpose: To compute the transmission in terms of energy vs. power of an x-ray beamline from the universal SR curves (C. K. Green. BNL Internal Report. Upton, 1976.) with reflectivities and transmission coefficients as obtained from the Fresnel equations and a comprehensive optical constants database.

Input: Machine parameters, photon energy, and system information.

Output: Formatted ASCII files containing the source spectra \{photon energy, power\}, and for each element a file with local properties and cumulative properties, and a log file of the session.

Description:

The TRANSMIT code was developed for the modelling of x-ray lithography systems – hence the terminology. It is however of wide application and the program performs calculations for mirrors, filters, masks and photo resists consisting of single elements or compounds of up to five atomic species.

The program is file oriented. A user interface creates a set of files describing the calculations to be performed and then invokes a computational engine to compute the actual physical properties. The program requires a set of startup files with a generic machine description for the source, a generic x-ray beamline, and a resist. These files can then be created and modified in a menu driven interface to represent specific machines and beamlines. New startup files are then saved for future use. They can be named by the user.

The program outputs several files. The first is a data file containing the source spectra in the form \{photon energy, power\}. For each optical element, two files are written when the LOGGING option is on. One contains the local properties for that element, the other contains the cumulative properties. The final file that can be written is a log file of the session. Whether or not this file is written is dependent on the value given to the LOGGING flag in the main menu. The log file contains all the information found in a summary to the screen, plus a simple ASCII graph of the flux after each element. It is recommended that the user make all the changes and additions to his system, then turn the logging flag on and run TRANSMIT. This way only the latest version of the system is written to the log file. Otherwise, if the flag is turned on when starting, every time a change is made and saved and TRANSMIT is run, a log of the entire system is written. This can lead to a very large file containing duplicate information; from another point of view, it will contain a log of the entire session.
TRANSMIT is run by a menu driven shell program. The shell allows the user to enter a source and system, generate the flux characteristics, and then modify any parameter without having to reenter the entire source and system. The same source or system can also be used with other runs, providing the startup files have been saved. TRANSMIT provides information on the spectral power distribution generated by a source and modified by an optical system. The program provides information on the total power along the beamline, making it quite useful for the estimation of power loadings. The units of the files created by TRANSMIT (e.g. Watts) may also be scaled to a given surface, thus giving W/cm².

The program can model a specific storage ring when the parameters of the machine are input. The characteristics of the user’s machine are scaled to the universal SR curve based on formulas calculated by C. K. Green. The source spectra is then calculated and stored. The user can then specify optical elements in a beamline or system. An optical element can consist of mirrors, filters, a mask or a photo resist. The user must input the materials used (element or compound), their densities, and other variables depending on the type of optical element (i.e. thickness for a filter). The reflectivity/transmission characteristics for various polarizations are calculated using an optical constants library shared with SHADOW. Meaningful energies are from 100 eV to 100,000 eV. If energies below 100 eV are entered, the program will assume an energy of 100 eV. If both the minimum and maximum energy are chosen to be below 100 eV, the program will abort. The information is output in the form of energy or wavelength vs. power.

**Examples:**

The following example uses the SR source Aladdin at 800 MeV to generate the power profiles. The x-ray beamline contains three mirrors at 88, 89, 89 degrees angle of incidence with a beryllium window and silicon carrier/gold absorber mask. The photoresist is modelled by 1 μm of PMMA.

Moving around in the shell is accomplished with the up/down arrows and the return key. The arrow keys move you vertically within a menu. The return key selects an item for modification and moves from the main menu down to the sub-menus, and returns to the main menu. Each screen shown below will have a description below it of what is being done and how to move to the next screen.
The program is started by typing TRANSMIT at the DCL level. The screen displays a prompt for system name. For first time users, this will be a generic system. The system name is input in the main menu at the EXIT & SAVE option. At this point, hitting <RETURN> will take you to the main menu.
This is the generic startup menu. It contains basic machine parameters for a source, but does not have any kind of beamline set up. Options, Machine, Geometry, Beamline, Mask, and Resist are all sub-menus where parameters can be entered. These will be shown. The Logging flag should be OFF for now, turn it on after the system is completely set up, then move the highlight bar to Transmit and hit <RETURN> to generate a log file of the final system. Overview will give a summary of the system to the terminal. Quit will exit without saving any parameters. The last option, EXIT & SAVE, is where the user's system name should be entered after setting up the rest of the parameters. Using the arrow keys, position the highlighted bar so it is on Options, then hit <RETURN>. 
This is the Options menu. It allows the user to give meaningful names to the data files that TRANSMIT writes and to give a default system or set of files to read. The title is just a comment to the user, it has no bearing on the program. This is where the first time user should name the Option file. That way the parameters displayed in this menu will be saved in a file as specified in the Option Name field. If changes have been made, choose the Exit option, otherwise choose Quit by moving the arrow keys and hitting <RETURN> to return to the main menu. Enter the Machine sub-menu by highlighting the option and again hitting <RETURN>. 
The machine menu sets up the basic x-ray source. Here we model the synchrotron called Aladdin. To enter the specific parameters interested in, move the arrow to highlight the parameter and hit return, then enter the correct data. Under Spectrum type, there are five options. Highlighting and hitting return will list the available spectra. Create Source will generate the source with the given parameters, and among other things, calculate the critical energy, wavelength, and total power/mrad. Exit will also save the parameters and generate the source. Return to the main menu. At this point we notice that next to Machine name is the name of the machine we have generated. This name can be used in future runs if the parameters are desired. Next move down to Geometry and enter that menu.
The geometry menu refers to the acceptance of the system in terms of mrads. This is usually specific to location in relation to the source. The vertical acceptance can either be specified or assumed infinite. If the user desires a conversion to Watts/cm², he will need also to enter the scan height, horizontal field, and distance from tangent. Update will perform the calculations and remain in this menu so that the user can see the results immediately, if desired, Exit will also run the calculations, but will return you to the main menu when complete.
The next menu is the Beamline menu. At this point the user can set up any combination of mirrors and filters knowing the element or compound they are made of, the material density, the formula, and either the incidence angle of the light or the thickness of the filter. Add optical element adds a new element on the end of the beamline. Delete optical element asks which element you wish to delete. Run Transmit recalculates the power. Quit returns to the main menu without saving any changes or updating any numbers. Exit saves, updates, and returns to the main menu.
The Mask menu appears much like the beamline menu. The elements of the mask are entered in the same fashion; the only difference being the user must distinguish between a carrier and an absorber.
The final step in an x-ray beamline is the photo resist. The Resist menu is similar in setup to the beamline and mask menus. Here we demonstrate a compound, PMMA, with molecular formula $C_{5}H_{8}O_{2}$. 
We return to the main menu and notice all the system names and titles appear here. This is where if the user had existing beamlines or masks etc., and wished to use them, he could load the specific system by name. If the user wishes to get a hardcopy of the entire system, this is the time to turn Logging on, run TRANSMIT, and turn it back off. The Overview option will give the same information as is in the log file, only to the screen. A sample run through the Overview option is shown in figures 5.9 – 5.10. To save the source, beamline, mask, and resist so that they can all be loaded together at another time, move down to the Exit & Save option and name the startup file.

When we exit the program we get an assortment of files similar to the following:

```
Directory D1:[CWELNAK.TEMP]

101LOC.DAT;1 11 /* local properties at first mirror */
101TOT.DAT;1 11 /* cumulative properties at first */
```
Figure 5.9: TRANSMIT: Overview screen

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnetic Field</td>
<td>2.883388 meters</td>
</tr>
<tr>
<td>Beam Energy</td>
<td>0.880000 GeV</td>
</tr>
<tr>
<td>Name</td>
<td>1565.609</td>
</tr>
<tr>
<td>Magnetic Field</td>
<td>1.260556 tesla</td>
</tr>
<tr>
<td>Critical Energy</td>
<td>545.2201 eV</td>
</tr>
<tr>
<td>Length</td>
<td>22.74039 Å</td>
</tr>
<tr>
<td>Power/amp</td>
<td>17.4000 kWs/amp</td>
</tr>
<tr>
<td>Total Power</td>
<td>1.740000 kW</td>
</tr>
<tr>
<td>Total Current</td>
<td>0.160000 Amp</td>
</tr>
<tr>
<td>Torque/degree</td>
<td>0.2769311 w/mr</td>
</tr>
<tr>
<td>Acceptance</td>
<td>8.309732</td>
</tr>
<tr>
<td>Total Acceptance</td>
<td>30.00000 mrad</td>
</tr>
</tbody>
</table>

Page 1
Press a key to continue.
Figure 5.10: TRANSMIT: Overview screen

<table>
<thead>
<tr>
<th>Compound</th>
<th>Filter/Mirror</th>
<th>Density</th>
<th>Thick/Angle</th>
<th>Watts</th>
</tr>
</thead>
<tbody>
<tr>
<td>11 H1</td>
<td>Mirror</td>
<td>15.38</td>
<td>88.00</td>
<td>1.02</td>
</tr>
<tr>
<td>21 H2</td>
<td>Mirror</td>
<td>15.38</td>
<td>88.00</td>
<td>1.51</td>
</tr>
<tr>
<td>31 H3</td>
<td>Mirror</td>
<td>19.38</td>
<td>89.00</td>
<td>1.19</td>
</tr>
<tr>
<td>41 BE</td>
<td>Filter</td>
<td>1.850</td>
<td>15.00</td>
<td>0.274</td>
</tr>
<tr>
<td>51 BE</td>
<td>Filter</td>
<td>1.786E-03, 2.000E-06</td>
<td>0.245</td>
<td></td>
</tr>
</tbody>
</table>
Figure 5.11: TRANSMIT: Overview screen

<table>
<thead>
<tr>
<th>Compound</th>
<th>Carrier Density</th>
<th>Thick Angle</th>
<th>Watts</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-51</td>
<td>2.330</td>
<td>2.000</td>
<td>0.160</td>
</tr>
<tr>
<td>25-81</td>
<td>19.30</td>
<td>0.580</td>
<td>0.149E-81</td>
</tr>
</tbody>
</table>

Optical Contrast: 19.72
Figure 5.12: TRANSMIT: Overview screen

<table>
<thead>
<tr>
<th>Compound</th>
<th>Resist</th>
<th>Density</th>
<th>Thick/</th>
<th>Watts</th>
</tr>
</thead>
<tbody>
<tr>
<td>T-PMN</td>
<td>1.000</td>
<td>1.000</td>
<td>5.13E-3</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Page 4
Press a key to continue
Figure 5.13: TRANSMIT: Overview screen
Figure 5.14: TRANSMIT: Overview screen
Figure 5.15: TRANSMIT: Overview screen
<table>
<thead>
<tr>
<th>File Name</th>
<th>Blocks</th>
</tr>
</thead>
<tbody>
<tr>
<td>102LOC.DAT</td>
<td>1</td>
</tr>
<tr>
<td>102TOT.DAT</td>
<td>1</td>
</tr>
<tr>
<td>103LOC.DAT</td>
<td>1</td>
</tr>
<tr>
<td>103TOT.DAT</td>
<td>1</td>
</tr>
<tr>
<td>104LOC.DAT</td>
<td>1</td>
</tr>
<tr>
<td>104TOT.DAT</td>
<td>1</td>
</tr>
<tr>
<td>105LOC.DAT</td>
<td>1</td>
</tr>
<tr>
<td>105TOT.DAT</td>
<td>1</td>
</tr>
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</tr>
<tr>
<td>202LOC.DAT</td>
<td>1</td>
</tr>
<tr>
<td>202TOT.DAT</td>
<td>1</td>
</tr>
<tr>
<td>301LOC.DAT</td>
<td>1</td>
</tr>
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<td>301TOT.DAT</td>
<td>1</td>
</tr>
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</tr>
<tr>
<td>401TOT.DAT</td>
<td>1</td>
</tr>
<tr>
<td>BOOTSTRAP.DAT</td>
<td>19</td>
</tr>
<tr>
<td>G-ES2.DAT</td>
<td>41</td>
</tr>
<tr>
<td>M-ALADDIN.DAT</td>
<td>42</td>
</tr>
<tr>
<td>MACHINE.DAT</td>
<td>4</td>
</tr>
<tr>
<td>O-CLW.DAT</td>
<td>14</td>
</tr>
<tr>
<td>S-EXAMPLE.DAT</td>
<td>22</td>
</tr>
<tr>
<td>S-PMMA.DAT</td>
<td>32</td>
</tr>
<tr>
<td>S-SIMASK.DAT</td>
<td>31</td>
</tr>
<tr>
<td>SRCOMP.DAT</td>
<td>4</td>
</tr>
<tr>
<td>STARTUP.DAT</td>
<td>19</td>
</tr>
<tr>
<td>TRANS.DAT</td>
<td>1</td>
</tr>
<tr>
<td>TRANSMIT.LOG</td>
<td>4</td>
</tr>
</tbody>
</table>

Total of 30 files, 291 blocks.

Next we'll show plots generated by TopDrawer of some of the various data files generated.
Figure 5.16: TRANSMIT: Source spectra – SRCOMP.DAT
Figure 5.17: TRANSMIT: Source spectra and after each of the three mirrors
Figure 5.18: TRANSMIT: Source, third mirror, beryllium window, and helium pathway
Figure 5.19: TRANSMIT: After complete beamline, after Si carrier, after Au absorber
Purpose: To generate the synchrotron spectra for a given machine.
Input: Machine parameters and energy range interested in.
Output: File containing photon energy, $\theta$, power in TopDrawer format if desired.

Description:

SYNC_SPECTRA takes the parameters of the user's machine and scales them to the universal curves using the formula from C. K. Green. From this information, it generates a spectra in the desired form at the desired photon energy. The results can be scaled by several factors including to the critical energy, to the beam energy, logarithmically, and changing $\theta$ by a factor of $\gamma$.

Examples:

The following example uses the SR machine Aladdin for its parameters. The output file is plotted using TopDrawer for $\theta$ vs. power.

```
$ r sync_spectra
Machine radius [ m ] : 2.083 /* Machine parameters */
   energy [ GeV ] : 0.8 /*
current [ A ] : 0.1 /*
Subtended angle [ mrad ] : 30 /* geometry at port */
Critical energy = 545.2986288035259 /* calculated by SYNC_SPECTRA */
Options:
dE = const 0 [ phot/sec/mrad(V)/eV ]
dL = const 1 [ phot/sec/mrad(V)/Angs ]
dL/E = const 2 [ phot/sec/mrad(V)/%bandwidth ]
dE/E = const 3 [ phot/sec/mrad(V)/%bandwidth ]
Power dL = const 4 [ watts/mrad(V)/angs ]
Power dE = const 5 [ watts/mrad(V)/eV ]
Then ? 5 /* here choose power in watts/mrad */
Maximum Photon energy [wavelength]: 2200 /* "white" source */
Minimum : 600
Number of points in energy [wavelength]: 17
Log Scale [ Y/N ] ? 0
```
Maximum vertical angle [ mrad ] : 1.25
Minimum : -1.25
No. of points in THETA : 250 /* vertical angle of above */
Polarization [1,2,3] : 3 /* parallel, perpendicular, total */
Scale energy to critical energy [ Y/N ] ? 0
Scale flux to beam energy [ Y/N ] ? 0
Multiply theta by gamma [ Y/N ] ? 0
File for 3d Top Drawer ? 0
File for Top Drawer (with embedded JOIN)? 1 /* 2d plot */
Output file name: spectd.dat
$ dir/since=-00:02

Directory CXRL$DUA0:[SHADOW.XRAYOP.DOCUMENT.UTIL.SYNC_SPECTRA]

SPECTD.DAT;1 399 27-SEP-1989 15:27:29.91 (RWED,RWED,RE,)

Total of 1 file, 399 blocks.
$ ty spectd.dat
  photon  theta  power
  energy

  :  :  :
  :  :  :
 600.0000 -1.019076  0.0000000E+00
 600.0000 -1.009036  0.0000000E+00
 600.0000 -0.9989960  0.0000000E+00
 600.0000 -0.9889558  0.0000000E+00
 600.0000 -0.9789157  0.0000000E+00
 600.0000 -0.9688755  0.0000000E+00
 600.0000 -0.9588354  0.0000000E+00
 600.0000 -0.9487952  0.0000000E+00
 600.0000 -0.9387550  0.0000000E+00
 600.0000 -0.9287149  9.8043092E-05
 600.0000 -0.9186747  1.1077573E-04
 600.0000 -0.9086345  1.3059203E-04
 600.0000 -0.8985944  1.4686001E-04
 600.0000 -0.8885542  1.6397580E-04
 600.0000 -0.8785141  1.8351113E-04
 600.0000 -0.8684739  2.0489228E-04
 600.0000 -0.8584337  2.2823217E-04
A plot of $\theta$ vs. power follows. Notice the decrease of the FWHM with the photon energy.
**Purpose:** To compute some useful angles for diffraction gratings studies.

**Input:** From keyboard.

**Output:** To the terminal and a file DIFFRAC.DAT

### Description:

The program recognizes the following cases:

1. **Blazed grating.** The grating is set at an angle relative to the incoming radiation that satisfies the grating equation for that wavelength and the specular reflection from the groove facet. Double grating systems can be configured efficiently in this mount.

2. **Constant Incidence Angle.** The user specify the incidence angle and the program computes the diffraction angles that satisfies the grating equation. Rowland Circle monochromators belong to this class.

3. **Constant Included Angle.** The deflection angle of the radiation (included angle) is kept constant and the incidence angle is computed to satisfy the diffraction condition. Typical examples of this case several varieties of Toroidal Grating Monochromators.

### Examples:

We will consider each of the three cases that the program can handle. The data will also be stored in the file DIFFRAC.DAT. All the cases are based on the grating equation:

\[
\sin(\alpha) - \sin(\beta) = \frac{m \lambda}{d}
\]  

(5.6)

where \(d\) is the grating spacing. The difference between the three cases is explained in the following figure. If the wavelength is too long to satisfy the grating equation, an error message will be issued.

```bash
$RUN UTIL:GRATING_SET
File out the results ? 1 yes
Options:
   Diffraction angles for given blaze..... [ 0 ]
   fixed incidence angle..... [ 1 ]
   fixed included angle..... [ 2 ]
```
Figure 5.21: GRATING_SET: Three diffraction cases available
Choice ? 1
Do you want to use eV [ 0 ] or Angstroms [ 1 ] ?
Choice ? 0
Photon Energy ? 500
Grating Line Density [ Lines / mm ] ? 1200
Order [ negatives are inside ] ? -1
Incidence Angle [ degrees ]? 88
Diffraction angle BETA: 85.14710385779866
Options:
Diffraction angles for given blaze..... [ 0 ]
fixed incidence angle..... [ 1 ]
fixed included angle..... [ 2 ]

Choice ? 0
Do you want to use eV [ 0 ] or Angstroms [ 1 ] ?
Choice ? 0
Photon Energy ? 800
Grating Line Density [ Lines / mm ] ? 1200
Order [ negatives are inside ] ? -1
Blaze angle [ degrees ]: 1
Incidence angle ALPHA: 87.94575321306993
Diffraction angle BETA: 85.94575321306993
Options:
Diffraction angles for given blaze..... [ 0 ]
fixed incidence angle..... [ 1 ]
fixed included angle..... [ 2 ]

Choice ? 2
Do you want to use eV [ 0 ] or Angstroms [ 1 ] ?
Choice ? 0
Photon Energy ? 250
Grating Line Density [ Lines / mm ] ? 1200
Order [ negatives are inside ] ? -1
Included Angle [ degs ]? 170
Incidence Angle [ degs ]: 86.95655527500508
Diffraction Angle [ degs ]: 83.04344472499492
Rotation Angle [ degs ]: 1.956555275005078
Options:
Diffraction angles for given blaze..... [ 0 ]
fixed incidence angle..... [ 1 ]
fixed included angle..... [ 2 ]

Choice ?
^Y
$
$

The following example was run using the TGM example. The six meter TGM has a fixed included angle of 160 degrees.

$ run util:grating_set
File out the results ? 1
Options:
  Diffraction angles for given blaze..... [ 0 ]
    fixed incidence angle..... [ 1 ]
    fixed included angle..... [ 2 ]

Choice ? 2
Do you want to use eV [ 0 ] or Angstroms [ 1 ] ?
Choice ? 0
Photon Energy ? 50
Grating Line Density [ Lines / mm ] ? 1200
Order [ negatives are inside ] ? -1
Included Angle [ degs ]? 160
Incidence Angle [ degs ]: 84.91514459719740
Diffraction Angle [ degs ]: 75.08485540280260
Rotation Angle [ degs ]: 4.915144597197396
Options:
  Diffraction angles for given blaze..... [ 0 ]
    fixed incidence angle..... [ 1 ]
    fixed included angle..... [ 2 ]

Choice ? 2
Do you want to use eV [ 0 ] or Angstroms [ 1 ] ?
Choice ? 0
Photon Energy ? 60
Grating Line Density [ Lines / mm ] ? 1200
Order [ negatives are inside ] ? -1
Included Angle [ degs ]? 160
Incidence Angle [ degs ]: 84.09441581524222
Diffraction Angle [ degs ]: 75.90558418475778
Rotation Angle [ degs ]: 4.09441581524222

Options:
Diffraction angles for given blaze..... [ 0 ]
fixed incidence angle..... [ 1 ]
fixed included angle..... [ 2 ]

Choice ?
<CTRL>Y
$ dir/since

Directory D13:[XRAYOP.DOCUMENT.UTIL.GRATING]

DIFFRAC.DAT;3 3 10-MAY-1989 14:56:22.01 (RWED, RWED, RE, RE)

Total of 1 file, 3 blocks.
$
$ type diffrac.dat

-------------------------------------------
-------------------------------------------
Fixed Included Angle.
-------------------------------------------
-------------------------------------------
-------------------------------------------
-------------------------------------------

Photon Energy [ eV ]: 50.000000000000000
Wavelength [ Angs ]: 247.970400000000000
Line density [ mm-1 ]: 1200.0000000000000
Order : -1.000000000000000
Included Angle [ degs ]: 1.396263401595464

Incidence Angle [ degs ]: 84.91514459719740
Diffraction Angle [ degs ]: 75.0848540280260
Rotation angle [ degs ]: 4.915144597197396

-------------------------------------------
end

-------------------------------------------
-------------------------------------------
-------------------------------------------
Fixed Included Angle.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Photon Energy [ eV ]</td>
<td>60.00000000000000</td>
</tr>
<tr>
<td>Wavelength [ Angs ]</td>
<td>206.64200000000000</td>
</tr>
<tr>
<td>Line density [ mm⁻¹ ]</td>
<td>1200.00000000000000</td>
</tr>
<tr>
<td>Order</td>
<td>-1.0000000000000000</td>
</tr>
<tr>
<td>Included Angle [ degs ]</td>
<td>1.396263401595464</td>
</tr>
<tr>
<td>Incidence Angle [ degs ]</td>
<td>84.09441581524222</td>
</tr>
<tr>
<td>Diffraction Angle [ degs ]</td>
<td>75.90558418475778</td>
</tr>
<tr>
<td>Rotation angle [ degs ]</td>
<td>4.09441581524222</td>
</tr>
</tbody>
</table>

end
Purpose: Computes the major and minor radii for a toroidal mirror given the setup parameters.

Input: The distances to the tangential and sagittal object and image planes, and the angle of incidence with respect to the mirror normal.

Output: A log file containing the input parameters and the computed focal lengths and radii of curvature for the tangential and sagittal planes.

Description:

If a toroidal mirror is to be used in a system where the object and image distances for the tangential and sagittal planes as well as the angle of incidence are known, TORUS may be used to compute the parameters of the mirror surface without setting up and tracing the system. The object and image distances need not be coincident for the tangential and sagittal planes. If they are coincident, TORUS computes the parameters that minimize astigmatism for the optical element. The focal length is computed in the conventional method:

$$\frac{1}{p} + \frac{1}{q} = \frac{1}{f}$$

where $p$ and $q$ are the object and image distances and $f$ is the focal length. The radii of curvature for the tangential and sagittal planes may be computed from the relations

$$\frac{1}{p} + \frac{1}{q} = \frac{2}{R \cos \theta}$$

and

$$\frac{1}{p} + \frac{1}{q} = \frac{2 \cos \theta}{r}$$

where $R$ and $r$ are the radii for the tangential and sagittal planes respectively, and $\theta$ is the angle of incidence. Notice that \{R,r\} are the “optical” radii of the mirrors ($R$ is, for example, the maximum radius). Sometimes other definitions are used in the equations defining the surface so that there may be an apparent difference of $r$ or $2r$ with the values of $R$ used by different authors.

Examples:

In this example we will have TORUS compute the radii for a toroidal mirror that has object and image distances both 100 cm from the mirror, and is illuminated at 85 degrees incidence. The results will be compared to a SHADOW run where a similar system is traced.
with a point source having a conical angle distribution of 10 mrad. Note that we wish to have the tangential and sagittal focal planes coincide. Let's run TORUS first:

$ run util:torus
Name to use for log file ? tor.par
Tangential Object: 100
    Image: 100
Sagittal Object: 100
    Image: 100
Incidence Angle: 85

Tangential.
Focal length = 50.00000000000000
Optical Radius = 1147.371324566986  \* R_0
Sagittal.
Focal length = 50.00000000000000
Radius = 8.715574274765817
Continue ? 0

After tracing the system described above, we check out the computed mirror parameters using MIRINFO:\index{MIRINFO}:

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
******************** MIRROR DESCRIPTION ********************
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Toroidal Mirror
TORUS example
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified:end.01
Full file Specification :DISK4:[RUNKLE.UG.UTIL.TORUS]END.01;1
Creation Date : 1-MAY-1989 14:31
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Surface figure was defined as: TOROIDAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity OFF
Mirror dimensions UNLIMITED
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Central Axis parameters :
Source Plane Distance 100.0000000000000000
Image Plane 100.0000000000000000
Incidence Angle 84.9999999999999999
Reflection/Diffraction Angle 84.9999999999999999

Mirror parameters COMPUTED
Same configuration as Central Axis YES
Objective focus at 100.0000000000000000
Image focus at 100.0000000000000000
Incidence angle 84.9999999999999999
Parameters used follow:
    Major Radius 1138.655750292220 * R_1 *
    Minor Radius 8.715574274765816 * rho *
Source of this O.E. moved NO
Mirror at pole position (no mov.) YES

Notice that the radii computed by TORUS and SHADOW are in agreement. Indeed:

\[ R_1 = R_0 - \rho \]

We can also look at the positions of the sagittal and tangential focii with respect to the final continuation plane by using FOCNEW:

---
Searching file: star.01
DISK4:[RUNKLE.UG.UTIL.TORUS]STAR.01;1 1-MAY-1989 14:3
---
Center at: Origin
X = 0.0000000000000000E+00 Z = 0.0000000000000000E+00
---
Working with 500 "good" rays out of 500
.............. S A G I T T A L ..............
X coefficients: 0.2539159622E-04 -0.5607838486E-05 0.2861789653E-05
Center: 0.8078084052E-04 Average versor: 0.1794262007E-03
Sagittal focus at: 0.2208541140
Waist size at best focus (rms): 0.1274078276E-02
Waist size at origin : \(0.1691682492\times10^{-2}\)

............... T A N G E N T I A L ............

Z coefficients : \(0.2398763062\times10^{-4} - 0.2063942800\times10^{-4} + 0.2015928704\times10^{-4}\)
Center : \(0.1149163952\times10^{-3}\) Average versor : \(0.5822703881\times10^{-4}\)
Tangential focus at : \(0.8604196190\)
Waist size at best focus (rms) : \(0.1549425142\times10^{-2}\)
Waist size at origin : \(0.4489909469\times10^{-2}\)

............... LEAST CONFUSION ............

T coefficients : \(0.4937922683\times10^{-4} - 0.2624726648\times10^{-4} + 0.2302107669\times10^{-4}\)
Center : \(0.1404682245\times10^{-3}\) Average versor : \(0.1886376144\times10^{-3}\)
Circle of least confusion : \(0.5315447034\)
Waist size at best focus (rms) : \(0.3011557938\times10^{-2}\)
Waist size at origin : \(0.4798028417\times10^{-2}\)

In practice it is quite difficult to eliminate astigmatism, but the toroidal mirror performed reasonably well here, considering the grazing angle of incidence.
Chapter 6

Graphics

This chapter covers the core level graphical utilities used with SHADOW and its utilities. They are a subset of TopDrawer a graphics program written at Stanford University.
Purpose: General graphics program used in plotting much of the output from SHADOW and its utilities.

Input: TopDrawer commands or a command file containing TopDrawer commands and reference to a data file.

Output: Terminal (TT:) and TD metafile with default name SAVE.DAT.

Description:

TDTEK is a graphics program that is part of TopDrawer. The output to the terminal can be suppressed by hitting <CTRL>O, as shown in specific examples. For the purpose of SHADOW and its utilities, example command files are given in the example for that utility. For more information on commands and other functions of TopDrawer, see the TopDrawer manual.

Notice that TDTEK writes to the virtual device UGDEVICE, which is often assigned to the terminal ($ASSIGN TT: UGDEVICE) but can be assigned to another terminal if the system implementation allows it ($ASSIGN RRE7: UGDEVICE). In this way text I/O will be directed to the standard terminal while the graphics will go to RRE7:. This is the best way to run TDTEK, otherwise text and graphics get hopelessly mingled. 1 TDTEK is quite useful for a “quick and dirty” examination of data files.

Examples:

Here we will use the example shown in SYSPLOT. We will wait to hit <CTRL>O, to demonstrate the output to the terminal.

```
$ type sysplot.com
set three
set order x y z
set limits x -50 50 y -100 4100 z -20 120
set ticks off
plot axes at 0 0 0
@sysplot.dat
end
```

1 New graphics packages are being developed.
$  
$ TDTEK  
TOP DRAWER (10/10/80) NOW EXECUTING  

@SYSPLOT.COM  
READING FROM INDIRECT COMMAND FILE: SYSPLOT.COM
1  set three
  
2  set order x y z
  
3  set limits x -50 50 y -100 4100 z -20 120
  
4  set ticks off
  
5  plot axes at 0 0 0

<CTRL>O \ \ suppresses output to screen
$ \ \ It’s probably a good idea to
$ \ \ hit <CTRL>O sooner than shown
$ dir/since \ \ if not running in graphics mode.

Directory D13:[XRAYOP.DOCUMENT.UTIL.TDTEK]

SAVE.DAT;1    11  5-MAY-1989 09:16:29.40 (RWED,RWED,RE,RE)
TD.FIL;1      1   5-MAY-1989 09:16:20.09 (RWED,RWED,RE,RE)

Total of 2 files, 12 blocks.
$  
$ type td.fil
@SYSPLOT.COM  
$
**Purpose:** Non interactive version of TopDrawer.

**Input:** A set of TopDrawer commands or a TopDrawer command file.

**Output:** A TopDrawer log file (TD.FIL) that contains the commands for the non-interactive session, and a file (SAVE.DAT) to be input to the Unified Graphics utilities for graphics output.

### Description:

TDFILE controls a non-interactive session of TopDrawer. The procedures for running TopDrawer through TDFILE are the same as those in TDTEK except that there is no direct graphics output. Advantages of running in this mode include an absence of clutter from a mixture of text and graphics and creating graphics files on terminals without graphics capability to be plotted elsewhere, as it may happen in a batch run. The graphics file SAVE.DAT may be used as input to UGTEK or a similar Unified Graphics utility for video display or hard copy.
Purpose: General purpose graphics program.
Input: TopDrawer command file or commands and data file.
Output: Plot to terminal.

Description:
This utility calls a general graphics program that processes a TopDrawer command file and plots the information to the screen. The terminal must be in graphics mode before calling TD_NEW. The command file input would be the same as for TDTEK. Instead of writing the information to a TD metafile, it is plotted to the screen directly.

Examples:
As with UGTEK, the transition between text and graphics mode cannot be shown well. We will again use the files SYSPLOT.COM and SYSPLOT.DAT to illustrate this utility.
To get output to an LN03 laser printer is fairly simple with TD_NEW. One need not put the terminal graphics mode. Simply make the first line typed or in the command file SET DEVICE LASER. This generates a file with the default name LN03.PLOT that can be printed using a standard print command with no set format.

```
$ TYPE SYSPLOT.COM
  set three
  set order x y z
  set limits x -50 50 y -100 4100 z -20 120
  set ticks off
  plot axes at 0 0 0
  @sysplot.dat
  end
$ $ DTEKON \ terminal in graphics mode

$ TD_NEW
  Top Drawer 08/11/82
  @SYSPLOT.COM
```
\ at this point the plot is being drawn to the screen. \ hitting <return> will get you back to the DCL prompt \ when finished plotting.

FORTRAN STOP
$
$
$ DTEKOFF \ terminal returns to text mode.
$
$
Purpose: Plots a TopDrawer metafile to the screen.
Input: A TopDrawer metafile (i.e. SAVE.DAT from TDTEK or TDFILE).
Output: A plot on the screen.

**Description:**

The command UGTEK will plot a TopDrawer metafile to the screen. The terminal must be in graphics mode for the plot to be drawn to the screen. A TD metafile is created by running either TDTEK or TDFILE on a TopDrawer command file. Its default name is SAVE.DAT. If there is more than one picture in a SAVE.DAT file, UGTEK gives you the opportunity to specify which pictures to plot, defaulting to plotting all (sequentially), and also gives you the chance to reduce the size of the picture.

**Examples:**

The page does not show well the transition from text to graphics mode, and consequently the output of UGTEK. However, we will show how to run through UGTEK, the result being a plot on the screen.

```
$ DTEKON \ puts the terminal in graphics mode. Clears screen for plot.
$ UGTEK

UG element file: SAVE.DAT
Plot which pictures (default=all)?
Enter picture scale factor (default=1.0):

At this point the screen clears again to make space for the plot. Hitting <RETURN> will return you to the DCL command level when the plot has finished.

$ DTEKOFF \ Returns the terminal to text mode.
```
<table>
<thead>
<tr>
<th>Purpose:</th>
<th>To plot a TopDrawer metafile on an HP 7470 plotter in high quality.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Input:</td>
<td>TopDrawer metafile such as SAVE.DAT output from TDTEK or TDFILE.</td>
</tr>
<tr>
<td>Output:</td>
<td>High quality plot of TopDrawer metafile on HP plotter.</td>
</tr>
</tbody>
</table>

**Description:**

The command UGHP7470 will plot a TopDrawer metafile to an HP plotter in high quality, suitable for publication. A TD metafile is created by running either TDTEK or TDFILE on a TopDrawer command file. Its default name is SAVE.DAT. If there is more than one picture in a SAVE.DAT file, UGHP7470 gives you the opportunity to specify which pictures to plot, defaulting to plotting all, and also gives you the chance to reduce the size of the picture. Notice that the program expects the plotter to be connected between the computer and the terminal with the “Y” cable. Alternatively, the file can be captured in a log session or (if using a PC emulator) on a local media for later plotting.
Figure 6.1: UGHP7470: Plotter/terminal setup
| **Purpose:** | To plot a TopDrawer metafile on an HP 7470 plotter in draft quality. |
| **Input:**  | TopDrawer metafile such as SAVE.DAT output from TDTEK or TDFILE. |
| **Output:** | Draft quality plot of TopDrawer metafile on HP plotter. |

**Description:**

The command UGHP7470S will plot a TopDrawer metafile to an HP 7470 plotter in draft quality. A TD metafile is created by running either TDTEK or TDFILE on a TopDrawer command file. Its default name is SAVE.DAT. If there is more than one picture in a SAVE.DAT file, UGHP7470S gives you the opportunity to specify which pictures to plot, defaulting to plotting all, and also gives you the chance to reduce the size of the picture. The same setup as in UGHP7470 applies here as well.
Chapter 7

General Utilities

The utilities in this chapter are more convenience utilities than SHADOW utilities. They perform tasks that, though rather simple, take multiple steps and are easier used in program form.
**Purpose:**
Clips the second column in a two column formatted file. CLIP gives the user the extrema for the second column, and asks for upper and lower clipping values. If any data is outside the specified range it is set equal to the clipping bound.

**Input:**
Any two column formatted file (e.g. output from PREPLOT).

**Output:**
A two column formatted file (similar to input file).

**Description:**
This utility clips the second column of a two column formatted file using the following rules:

\[
\begin{align*}
\text{IF } Y > \text{MAX} \text{ THEN } Y:=\text{MAX} \\
\text{IF } Y < \text{MIN} \text{ THEN } Y:=\text{MIN}
\end{align*}
\]

The input file is scanned and the extrema are displayed. Then the user can enter the desired values for MAX and MIN. This program is useful for cleaning up files that have some rays that fall far from the mean. Then the clipped file can be plotted without overscaling to accommodate the stray rays. Also, the graphics program TopDrawer crashes if the range is too wide. TD converts first to integers then checks for limits; in this step a very small number \(10^{-30}\) will cause an overflow while zero values are handled correctly. This may happen when computing absorption or transmission data near an edge. In this case, CLIP to zero.

**Examples:**
We'll run CLIP on the first 15 rows in the file PRE.DAT (See PREPLOT document for details on how PRE.DAT was generated).

```
$ clip
Input file ? pre.dat
Output file ? clip.dat
CLIP> Read 15 records.
Enter 0 to clip at zero only, 1 to other limits
1
Found:
Ymax = 2.5044683E-02
```
Ymin = -2.5014305E-02
CLIP> Enter Ymin,Ymax
CLIP> Ymax = 2.0e-2
CLIP> Ymin = -2.0e-2
CLIP> Do you want IF (y,lt.1.0e-10) y = 0 [ y=1,n=0 ] ? 1

Here's a before and after look at the data.

Before:

-0.2780961 1.8455550E-02
-0.2831062 1.8694960E-02
0.3307338 -2.2033621E-02
-0.3311352 2.2213560E-02
-9.4173871E-02 6.2023830E-03
0.2767766 -1.8385222E-02
0.3686378 -2.4258027E-02
0.1166269 -7.7635958E-03
-0.3735043 2.5044683E-02
-0.3660054 2.4130033E-02
-9.1828540E-02 6.0481941E-03
0.3607364 -2.4109263E-02
0.3719847 -2.5014305E-02
-3.2931246E-02 2.2053795E-03
0.1509321 -1.0015748E-02

After:

-0.2780961 1.8455550E-02
-0.2831062 1.8694960E-02
0.3307338 -2.0000000E-02
-0.3311352 2.0000000E-02
-9.4173871E-02 6.2023830E-03
0.2767766 -1.8385222E-02
Another example is the transmission of a 25 μm beryllium window in the energy range 100-1000 eV. We run ABREFC to get the TRANSMISSION.DAT file, then run that through CLIP.

```
$ $ abrefc
Full printout at Terminal ?
0
Thickness (microns) ?
25
The program is setup to compute optical constants of compounds with up to 5 elements.
How many atomic species ?
1
H2O would be: HH,2 and OO,1. Then ?
Enter 2-letters (capitalized) atomic symbol and formula index for :
1
BE /* beryllium */
1
Density ?
1.96 /* g/cm^3 */
Enter starting photon energy,end and step
100
1000
50
Incidence angle ?
0
Molecular weight is : 9.010000
```
Atoms/Cm3
1.3100236E+23 0.0000000E+00 0.0000000E+00 0.0000000E+00 0.0000000E+00
$ dir/since

Directory D13:[XRAYOP.DOCUMENT.UTIL.CLIP]

ABSOR_COEFF.DAT;1 2 13-JUL-1989 15:06:36.28 (RWED,RWED,RE,RE)
CROSS_SECTION.DAT;1 2 13-JUL-1989 15:06:38.09 (RWED,RWED,RE,RE)
DIELEC_FUNC.DAT;1 2 13-JUL-1989 15:06:36.91 (RWED,RWED,RE,RE)
ELECTRON_YIELD.DAT;1 2 13-JUL-1989 15:06:37.48 (RWED,RWED,RE,RE)
ENERGY_LOSS.DAT;1 2 13-JUL-1989 15:06:37.20 (RWED,RWED,RE,RE)
REFLECTIVITY.DAT;1 3 13-JUL-1989 15:06:35.97 (RWED,RWED,RE,RE)
REFRAC_INDEX.DAT;1 2 13-JUL-1989 15:06:37.75 (RWED,RWED,RE,RE)
TRANSMISSION.DAT;1 2 13-JUL-1989 15:06:36.53 (RWED,RWED,RE,RE) <<<

Total of 9 files, 17 blocks.

$ ty transmission.dat
original file. Notice the very small
values that would trip up a program
" like TopDrawer.

... crash site

$ clip
Input file ? transmission.dat
Output file ? trans_clip.dat
CLIP> Read 19 records.
Enter 0 to clip at zero only, 1 to other limits
1
Found:
Ymax = 5.3043921E-02
Ymin = 0.0000000E+00
CLIP> Enter Ymin,Ymax
CLIP> Ymax = 6e-02
CLIP> Ymin = 0
CLIP> Do you want IF (y,lt.1.0e-10) y = 0 [ y=1,n=0 ] ?1 /* get rid of 1E-
$ */ values. */
$ type trans_clip.dat  /* after */

100.0000 0.0000000E+00
150.0000 0.0000000E+00
200.0000 0.0000000E+00
250.0000 0.0000000E+00
300.0000 0.0000000E+00
350.0000 0.0000000E+00
400.0000 0.0000000E+00
450.0000 0.0000000E+00
500.0000 2.6327970E-10
550.0000 4.6298648E-08
600.0000 1.9180404E-06
650.0000 2.9635275E-05
700.0000 2.2219188E-04
750.0000 1.7964149E-02
800.0000 3.2535858E-02
850.0000 5.3043921E-02
900.0000 5.3043921E-02
950.0000 5.3043921E-02
1000.000 5.3043921E-02

$
Purpose: Extract any two columns from an ASCII, N column file.
Input: A file with N columns.
Output: A file with any two columns extracted from the input file.

Description:

This utility reads in a formatted ASCII file with a known number of columns, and allows the user to choose any two columns to be written to another file in the order specified by the user. This utility will not work on binary output files such as MIRR.xx or STAR.xx without first running them through TRANSLATE.

Examples:

Two examples follow. The first example demonstrates the most common type of file this utility works on, a simple file with multiple columns, where each line is a column. The second example demonstrates how the utility works on a translated ray file.

```
$ type example.lis
  1  101  102  103  104
  2  201  202  203  204
  3  301  302  303  304
  4  401  402  403  404
    :    :    :    :    :
    :    :    :    :    :
101  111  222  333  444
$
$ col_2
input file : example.lis
How many columns in the input file : 5
output file : col2.out
Column 1 : 2
Column 2 : 4
$
$ type col2.out
  /* column 2
   column 4 */
For this example we'll use the synchrotron source file BEGIN.DAT created for use with the TGM examples. First we'll run TRANSLATE to format the file. Then we'll run COL_2 to extract columns 4 and 6, the angle distribution. For this example we'll only use 20 rays.

$ translate
File for input ? begin.dat
and for output ? tbegin.dat
Read 1000 rays.
Each ray has 12 entries.
how many rays to translate ? 20
All done.
$

$ col_2
input file : tbegin.dat
How many columns in the input file : 12
output file : col.out
Column 1 : 4
Column 2 : 6
$

$ dir/since

Directory D13:[XRAYOP.DOCUMENT.UTIL.COL_2]

<table>
<thead>
<tr>
<th>File_name</th>
<th>Num</th>
<th>Date</th>
<th>Time</th>
<th>Format</th>
</tr>
</thead>
<tbody>
<tr>
<td>COL.OUT</td>
<td>1</td>
<td>5-MAY-1989</td>
<td>14:24:28.07</td>
<td>(RWED, RWED, RE, RE)</td>
</tr>
<tr>
<td>TBEGIN.DAT</td>
<td>1</td>
<td>5-MAY-1989</td>
<td>14:14:02.81</td>
<td>(RWED, RWED, RE, RE)</td>
</tr>
</tbody>
</table>
Total of 2 files, 14 blocks.

```
$ type col.out
7.1207349033216965E-03 4.2598658113385647E-04
-2.6712548929609998E-03 -5.3321707887277658E-04
-4.245040397124077561E-03 8.9222914618205597E-04
-1.5401936936191181E-03 1.4590310682059307E-03
1.0545386278912323E-02 5.3498849279228146E-04
1.5371153788922542E-03 -2.9422052535095319E-04
-9.0826729006043025E-04 -4.2770848179471997E-04
6.4885397124077561E-03 2.6606883484927588E-04
-9.0210832491599969E-03 1.1547317216979092E-03
4.5896466841608658E-03 1.1607701177610969E-03
6.0678659520798649E-03 7.6186301014691257E-04
-7.8908855034182015E-04 4.8762170237351560E-04
-2.0309429492565774E-03 -4.3717432212851493E-05
1.4943430919349669E-03 -3.8288106268715352E-04
-7.6212021143264583E-03 -4.8180567614367067E-05
-8.3078372735220177E-03 5.8870256818982272E-04
9.5664992971012362E-03 -2.8970588545932720E-04
-1.7791916484991329E-03 6.1577849756395542E-04
3.702018414177026E-03 -1.9409225197437758E-04
-2.5605721601357427E-04 -5.8362582323245487E-04
$```

`* now compare the above list with the entries in columns 4 and 6 from *\`
`* the first two records of tbegin.dat *\`

```
$ type tbegin.dat
5.0982915276736750E-04 1.483496817385351 -1.3608385926677304E-02
7.1207349033216965E-03 0.9999745565112491 4.2598658113385647E-04
0.9999746472458372 -7.1207342572083670E-03 -3.0334144220366471E-06
1.0000000000000000 2533844.889220482 1.0000000000000000
-1.892307584387773E-02 0.5565551450115362 -5.9508659084241021E-03
-2.6712548929609998E-03 0.9999962900315399 -5.3321707887277658E-04
0.9999964321922838 2.6712545132120622E-03 -1.4243638128054778E-06
1.0000000000000000 2533844.889220482 2.0000000000000000
: : :
: : 
: : 
```

**Purpose:** Prepares a file with a listing of the default directory sorted by date.

**Input:** None.

**Output:** A file DIRTIME.LIS containing the sorted directory.

**Description:**

The utility writes a file of the default directory to disk and then sorts it according to the date, using VMS $DIRECTORY and $SORT commands. Strangely enough, VMS does not have a switch to change the sorting of the directory listing. It is sometimes important to see when files were created relative to others. This simple utility fills the gap.

**Examples:**

```
$ show def
   BEAMROOT:[XRAYOP.DOCUMENT]
$ DIRTIME
Sorted listing ready in DIRTIME.LIS.
$
```

```
$ type dirtime.lis

Directory D13:[XRAYOP.DOCUMENT]

\* name size date *

UG.DIR;1 2 16-JUN-1988 23:
PAGE.TEX;2 2 8-JUL-1988 22:
DIRTEX.DAT;2 2 10-JUL-1988 11:
DIRTEMP.DAT;9 1 11-JUL-1988 15:
DIRTRY.DAT;1 1 11-JUL-1988 15:
DOC_PREP.FOR;28 11 11-JUL-1988 16:
DOC_PREP.EXE;19 14 11-JUL-1988 17:
DOCPROG.DAT;6 4 2-AUG-1988 11:
INTRODUCTION.DIR;1 1 6-DEC-1988 10:
```
<table>
<thead>
<tr>
<th>Directory</th>
<th>ID</th>
<th>Date</th>
<th>Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>SOURCE.DIR</td>
<td>1</td>
<td>12-DEC-1988</td>
<td>15:57</td>
</tr>
<tr>
<td>CLIP.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:46</td>
</tr>
<tr>
<td>HISTO2.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>MIRINFO.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>FMAX.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>FILEX.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>MINMAX.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>H2KOL.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>HISTO1.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>FOCNEW.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>MERGEROW.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>HISTO3.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>DOSE.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:48</td>
</tr>
<tr>
<td>PLOTXY.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:53</td>
</tr>
<tr>
<td>PLOTFOC.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:53</td>
</tr>
<tr>
<td>PREPLOT.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:54</td>
</tr>
<tr>
<td>REFFLAG.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:54</td>
</tr>
<tr>
<td>RECOLOR.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:54</td>
</tr>
<tr>
<td>PREPLOT2.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:54</td>
</tr>
<tr>
<td>RETRACE.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:57</td>
</tr>
<tr>
<td>SURPLOT.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:58</td>
</tr>
<tr>
<td>REVPRINT.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:58</td>
</tr>
<tr>
<td>SCAN.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:58</td>
</tr>
<tr>
<td>SOURCINFO.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>12:58</td>
</tr>
<tr>
<td>SYSINFO.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>13:00</td>
</tr>
<tr>
<td>COP.COM</td>
<td>7</td>
<td>30-JAN-1989</td>
<td>13:00</td>
</tr>
<tr>
<td>TRANSLATE.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>13:01</td>
</tr>
<tr>
<td>VARIANCE.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>13:01</td>
</tr>
<tr>
<td>SYSPLOT.DIR</td>
<td>1</td>
<td>30-JAN-1989</td>
<td>13:01</td>
</tr>
<tr>
<td>DIR.LST</td>
<td>4</td>
<td>30-JAN-1989</td>
<td>13:07</td>
</tr>
<tr>
<td>TRACE.DIR</td>
<td>1</td>
<td>17-FEB-1989</td>
<td>15:56</td>
</tr>
<tr>
<td>PREREFL.DIR</td>
<td>1</td>
<td>22-FEB-1989</td>
<td>21:52</td>
</tr>
<tr>
<td>PRE_MLAYER.DIR</td>
<td>1</td>
<td>23-FEB-1989</td>
<td>20:33</td>
</tr>
<tr>
<td>MLAYER.DIR</td>
<td>1</td>
<td>23-FEB-1989</td>
<td>21:56</td>
</tr>
<tr>
<td>ABREFC.DIR</td>
<td>1</td>
<td>24-FEB-1989</td>
<td>21:02</td>
</tr>
<tr>
<td>COMB_RAY.DIR</td>
<td>1</td>
<td>24-FEB-1989</td>
<td>22:42</td>
</tr>
<tr>
<td>CUTOFF.DIR</td>
<td>1</td>
<td>25-FEB-1989</td>
<td>12:53</td>
</tr>
<tr>
<td>DIRTIME.DIR</td>
<td>1</td>
<td>25-FEB-1989</td>
<td>13:08</td>
</tr>
</tbody>
</table>
Total of 48 files.
FILEX

Purpose: To perform simple algebraic operations on the second column of a two column formatted file and to plot out the results. A list of legal operations is given to the user upon request.

Input: Any two column ASCII files such as those generated by PREPLOT. One or two files may be used as input.

Output: A two column ASCII file.

Description:

It is often desirable to perform simple algebraic operations among files, such as division or multiplication without having to write every time a program ex-novo. The operation may be performed between two files or a file and a scalar. FILEX implements a logic as described below. This guide to FILEX can be displayed by typing ? after the prompt FILEX>.

PROGRAM FILEX

This program is a file manipulation utility. Operations are performed on two files having the format of an (x,y) table and the same number of elements. Command format is:

file-out = file1 /oper/ file2

where /oper/ is an operation symbol, like *, +, -, /.

If file2 is written as %123, then 123 will be used as a factor for the Y (second) column.

The following symbols operates only on one file. They are

& Sum of X, Y # Computes 1/Y
^ Derivative of Y ~ SQRT (Y)
@ Integral of second column " Interchange X and Y
\ Inverts the sequential order ( 1 --> N, N --> 1 )
| Replace Y with X*Y
[ Mean and st. dev. of X weighed by Y
To plot out a file, type: %Plot = file-name

If the extension of the files in the command line is *.dat, then the extension may be omitted. Also, the file-out may have the same name as one of the files on which the operation is being performed. This helps keep the directory clean when several operations are to be applied to a file.

Examples:
Add X, Y: file.out = file1 + file2
Multiply Y by 200: file.out = file1 * %200
Subtract Y2 from Y1: file.out = file1 - file2

Note: Because of the simple parser used, correct spacing of the '=' and operations between filenames is critical for proper generation of output file.

Examples:
Here we will examine the reflectivity of a gold coated mirror as a function of grazing angle. The mirror will be illuminated by a 100 points between 0 and 12 degrees. PREPLOT will be used to create a two column file with the intensity vs divergence in radians in the image plane. We will use FILEX to convert the radians to degrees and to add the grazing angle of the central ray to the angle of all the rays. FILEX will also be used to plot out the results.

The source and system descriptions follow:

++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
************** SOURCE DESCRIPTION **************
++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++++
Input file specified: end.00
Full file Specification :D13:[XRAYOP.DOCUMENT.UTIL.FILEX]END.00;4
Creation Date :14-APR-1989 14:44
Mixed Type Source. Spatial: RANDOM, directions: GRID
Generated total 100 rays.
Source assumed BIDIMENSIONAL (flat).
Source Spatial Characteristics: POINT
Source Emission Characteristics
Distribution Type: LAMBERTIAN
Distribution Limits. +X : 0.000000000E+00  -X: 0.000000000E+00 rad
+Z : 0.100000000  -Z: 0.100000000 rad

Source Photon Energy Distribution: SINGLE LINE
Photon Energy: 1000.0 eV, or 12.399 Angs.
Angular difference in phase is 0.00000E+00
Degree of polarization is 0.50000
Source points have INCOHERENT phase.

Gold coated spherical mirror
Filex example
Input file specified:end.01
Full file Specification :D13:\[XRAYOP.DOCUMENT.UTIL.FILEX\]END.01;3
Creation Date :14-APR-1989 14:46

Surface figure was defined as: PLANE
Cylindrical figure NO
Element type REFLECTOR
Reflectivity ON coefficients from: aufl.opt
Polarization dependence YES
Mirror dimensions UNLIMITED

Central Axis parameters :
Source Plane Distance 10.00000000000000
Image Plane: 10.00000000000000
Incidence Angle: 84.00000000000001
Reflection/Diffractive Angle: 84.00000000000001

Mirror parameters: COMPUTED
Same configuration as Central Axis: NO
Objective focus at: 0.0000000000000000E+00
Image focus at: 0.0000000000000000E+00
Incidence angle: 0.0000000000000000E+00

Parameters used follow:
   Plane mirror
Source of this O.E. moved: NO
Mirror at pole position (no mov.): YES

$ preplot
PREPLOT> Input file? star.01
Read 100 rays.
Each ray has 18 entries.
PREPLOT> How many columns to write out? 2
Row [1-18]: the individual column
Row [20]: \( R = \sqrt{X^2 + Y^2 + Z^2} \)
Row [21]: angle from the Y-axis
Row [22]: the magnitude of A vector
Row [23]: A**2
PREPLOT> Row # 1: 23 /* intensity vs. */
PREPLOT> 2: 6 /* Z' */
Options --- Enter
0 for excluding the losses
1 for including losses at a particular O.E.
2 for plotting all the rays.
3 for plotting ONLY the losses (all of them)
4 for plotting ONLY the losses at a given O.E.
PREPLOT> Then? 0
**********
Found 100 good points out of 100
Output options:
[ 0 ] store rays in a file
[ 1 ] plot directly on screen
[ 2 ] both
Then ? 0
PREPLOT> Output file ? reflec2.dat

$ filex
FILEX >  reflec = reflec * %57.30  /* converts radians to degrees */
FILEX >  read    100 records.
FILEX >  Succesful completion
FILEX >  reflec = reflec + %6   /* Adds grazing angle of central */
         /* ray to all rays */
FILEX >  read    100 records.
FILEX >  Succesful completion
FILEX >  reflec = reflec "  /* Switches columns for plotting */
FILEX >  read    100 records.
FILEX >  Succesful completion
FILEX >  %plot = reflec2   /* Writes to plottable file */
Figure 7.1: FILEX example – grazing angle vs. reflectivity
Purpose: Scans an ASCII two column file and returns the extrema.
Input: Any formatted two column file (e.g. output from the utility PREPLOT is suitable for input to FMAX).
Output: TT:

Description:
FMAX accepts as input a formatted two column file (e.g. output from PREPLOT) and returns the extrema found in the two columns. The first column lists the absolute extrema. The second column lists extrema with a slightly larger variance than the absolute extrema so that these values can be used to set, e.g., limits of a plot. The program is similar to MINMAX and useful to verify if data are reasonable, and to figure out plotting limits.

Examples:
We’ll use the formatted file generated from PREPLOT using the STAR.01 file from the primer sec 2.4 (Refer to PREPLOT example).

```
$ fmax
File-name ?
pre.dat
Read       1000 points.
Found :
XMIN  -0.4102450  . Set at : -0.5023146
XMAX   0.4081510   0.5002205
YMIN  -2.7292412E-02 -3.3435397E-02
YMAX   2.7311902E-02  3.3454888E-02
Good luck!
```
LOCPRI

Purpose: To insert a Form Feed every 60 lines and prepare a nicely formatted file. Can directly drive a local printer connected to the printer port of a terminal.

Input: A formatted/text file.

Output: A nicely printable formatted file or printout.

Description:

This utility will take a file and format it to print “nice” pages. The output can be written to a file or sent directly to a printer if one is connected to the terminal on the “printer” port.

Examples:

For an example we’ll run LOCPRI on the file REFLEC2.DAT used in H2KOL. Here we write the formatted output to a file.

```bash
$ run util:locpri
Input file ? reflec2.dat
Output file ? lreflec.dat
$ ty lreflec.dat
```

reflec2.dat

<p>| | | |</p>
<table>
<thead>
<tr>
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Purpose: Takes two files of similar format and combines them into one file by taking alternate rows from each input file.
Input: Any two text files of similar format, especially column files (e.g. output from TRANSLATE or PREPLOT).
Output: A single file with alternating rows from the two input files.

Description:
This utility makes it easy to compare the rows of two files like those that are output from TRANSLATE or PREPLOT or similar column formatted files. Two input files are requested and are combined row by row with the labels 1: and 2: at the left of each row to designate which file it belongs to.

Examples:
Let's translate a couple of BOFs and then merge them.

```
$ translate star.01
File for input ?
star.01
and for output ?
star.dat
Read 1000 rays.
Each ray has 12 entries.
how many rays to translate ?
2
All done.

$ translate
File for input ?
mirr.01
and for output ?
mirr.dat
Read 1000 rays.
```
Each ray has 12 entries.
how many rays to translate?
2
All done.

$ run util:merge
First file?
star.dat
Second file?
mirr.dat
Output file?
mrg.dat
End of file found.

We’ll type the input files and the merged file:

$ ty star.dat
-0.278096064037422 1.0318894176630788E-16 4.3164806578071226E-03
 1.8455549611618461E-02 0.9998271538721697 -2.2483479289981678E-03
 0.9989241080626123 -1.853447430375961E-02 -4.2509994043719599E-02
 1.000000000000000 0.0000000000000000E+00 1.000000000000000

-0.2831061672594486 1.7334495384241366E-15 4.6566385577880755E-03
 1.8694959705118020E-02 0.9998115448711023 -5.2319426682332429E-03
 0.9988783014611897 -1.8904690404164647E-02 -4.341372537624238E-02
 1.000000000000000 0.0000000000000000E+00 2.000000000000000

$ ty mirr.dat
-1.016798475029074 -9.5158502246510235E-02 -3.1955982989010609E-02
 1.8455549611618461E-02 0.9387612182741329 0.9387612182741329
 0.9989241080626123 3.3607164154692846E-02 -3.1955982989010609E-02
 1.000000000000000 0.0000000000000000E+00 1.000000000000000
Notice how much easier it is to compare ray values.

```
$ty mrg.dat
1: -0.2780960640374226 1.0318894176630788E-16 4.3164806578071226E-03
2: -1.016798475029074 -9.5158502246510235E-02 1.4295016022463081E-02
1: 1.8455549611618461E-02 0.9998271538721697 -2.2483479289981678E-03
2: 1.8455549611618461E-02 0.3440737824260938 0.9387612182741329
1: 0.9989241080626123 -1.8534474303775961E-02 -4.2509994043719599E-02
2: 0.9989241080626123 3.3607164154692846E-02 -3.1955982989010609E-02
1: 1.0000000000000000 0.0000000000000000E+00 1.0000000000000000
2: 1.0000000000000000 0.0000000000000000E+00 1.0000000000000000
1:
2:
1: -0.2831061672594486 1.7334495384241366E-15 4.6566385577880755E-03
2: -1.032199854951956 -0.2224876977694570 1.5282134990743490E-02
1: 1.8694959705118020E-02 0.3468721057931864 0.9377261011106732
0.9988783014611897 3.4329772451938084E-02 -3.2612966645288010E-02
1.0000000000000000 0.0000000000000000E+00 2.0000000000000000
2: 1.0000000000000000 0.0000000000000000E+00 2.0000000000000000
1:
2:
```
**Purpose:** Arithmetic double-precision calculator.

**Input:** Numbers and functions suitable for calculator input.

**Output:** The results of the calculations.

### Description:

Simulates a double-precision calculator, like an HP. Old and crude, but useful at 3:00 AM when you have lost your HP. Functions are chosen by entering the appropriate number. The inverse of a function may be used by entering a negative of the number. For example, to take the square root, one enters 7, to square a number, one enters -7.

### Examples:

A simple example using this utility follows.

```
$ run util:pp
This program simulates an RPN calculator, like an HP. It works in double precision. For the moment, the stack is 10 registers. The trig functions work in radians, but there is an easy conversion key. The "display" is the lowest register in the stack. Enter the number correspondent to the function. Use the negative code for the inverse function. Press RETURN when ready to go

- 10 clear all
0 enter X
1 SIN , 2 COS , 3 TAN
4 LN , 5 LOG , 6 - not used yet-
7 SQRT
8 to degrees
9 +9 CHS, -9 1/x
10 EXIT
11 (+) , 12 (+) , 13 (-) , 14 (/)
```
15 write x to register
16 read in x from register
17 exchange the two lowest stack regs.
18 step up (down) the stack
19 E (G) Format
20 print the registers and stack
Register 11 contains conversion eV-angstroms, 12 PI.

Display = 0.0000000000000000E+00

- 10 clear all
0 enter X
1 SIN, 2 COS, 3 TAN
4 LN, 5 LOG, 6 - not used yet-
7 SQRT
8 to degrees
9 +9 CHS, -9 1/x
10 EXIT
11 (*) , 12 (+), 13 (-), 14 (/)
15 write x to register
16 read in x from register
17 exchange the two lowest stack regs.
18 step up (down) the stack
19 E (G) Format
20 print the registers and stack
Register 11 contains conversion eV-angstroms, 12 PI.

Display = 25.00000000000000

- 10 clear all
0 enter X
1 SIN, 2 COS, 3 TAN
4 LN , 5 LOG, 6 - not used yet-
7 SQRT
8 to degrees
9 +9 CHS, -9 1/x
10 EXIT
11 (*) , 12 (+) , 13 (-) , 14 (/)
15 write x to register
16 read in x from register
17 exchange the two lowest stack regs.
18 step up (down) the stack
19 E (G) Format
20 print the registers and stack
Register 11 contains conversion eV-angstroms, 12 PI.

--------------------------------- Display = 5.0000000000000000 ---------------------------------

-10

- 10 clear all
0 enter X
1 SIN , 2 COS , 3 TAN
4 LN , 5 LOG, 6 - not used yet-
7 SQRT
8 to degrees
9 +9 CHS, -9 1/x
10 EXIT
11 (*) , 12 (+) , 13 (-) , 14 (/)
15 write x to register
16 read in x from register
17 exchange the two lowest stack regs.
18 step up (down) the stack
19 E (G) Format
20 print the registers and stack
Register 11 contains conversion eV-angstroms, 12 PI.

--------------------------------- Display = 0.0000000000000000E+00 ---------------------------------
Purpose: Displays a formatted file in reverse order line by line.
Input: A formatted file of 10,000 lines or less.
Output: TT:

Description:
Displays a file in reverse order line by line. REVPRINT can be useful when the data are in a large file and the last entries are to be examined. A typical case could be to examine a long .LOG file to find the reason for a crash.

Examples:
We'll run REVPRINT on a short PREPLOT output file Here's the file before REVPRINT:

```
$ ty pre.dat
-0.2780961 1.8455550E-02
-0.2831062 1.8694960E-02
0.3307338 -2.2033621E-02
-0.3311352 2.2213560E-02
-9.4173871E-02 6.2023830E-03
0.2767766 -1.8385222E-02
0.3686378 -2.4258027E-02
0.1166269 -7.7635958E-03
-0.3735043 2.5044683E-02
-0.3660054 2.4130033E-02
-9.1828540E-02 6.0481941E-03
0.3607364 -2.4109263E-02
0.3719847 -2.5014305E-02
-3.2931246E-02 2.2053795E-03
0.1509321 -1.0015748E-02
```

After:
$ revprint
Input file ?
pre.dat
    0.1509321     -1.0015748E-02
   -3.2931246E-02    2.2053795E-03
     0.3719847     -2.5014305E-02
     0.3607364     -2.4109263E-02
   -9.1828540E-02    6.0481941E-03
  -0.3660054     2.4130033E-02
  -0.3735043     2.5044683E-02
     0.1166269    -7.7635958E-03
     0.3686378    -2.4258027E-02
     0.2767766   -1.8385222E-02
   -9.4173871E-02    6.2023830E-03
  -0.3311352     2.2213560E-02
     0.3307338    -2.2033621E-02
  -0.2831062     1.8694960E-02
  -0.2780961     1.8455550E-02
**Purpose:**
Creates a two column formatted file of a discrete gaussian function.

**Input:**
The standard deviation and interval for the gaussian function, and the number of points on the interval for which the function is to be computed.

**Output:**
A user named two column file that contains the value of the gaussian function for each point on the interval.

**Description:**
TGAUSS may be used to compare a randomly generated distribution to an actual gaussian with the same parameters. It could also be useful for comparing with the spectrum from the output of a narrow band device such as a monochromator. For demonstrations, see the example in HISTO1. If the normalization is to be to the area, then the integral over the interval is equal to one. If normalization is to one, the value of the gaussian at the mean (zero) is one and the other values are weighted accordingly.

**Examples:**
To start out, let's make a plot of a gaussian function with a four s standard deviation width on either side of the mean. We'll normalize to area and use TopDrawer to plot out the results.

```
$ run util:tgauss
Output file ? Gauss.dat
Enter Mean, Stnd. Dev., Interval Width, Number of Points
 0
 1
 8
100
Normalize to area [ 1 ] or to 1 [ 0 ]?
1
$
```
Figure 7.2: TGAUSS Example
Purpose: Prints a file, one page at a time, forward or backwards.
Input: A file to be displayed on the terminal.
Output: The contents of the file printed one page at a time.

Description:

TYPEOUT allows scrolling through a file one page at a time in either direction. Maximum file size is 1000 lines.

Examples:

The following run was done using the file reflec2.dat used in H2KOL.

```
$ typeout
Input file ? reflec2.dat

11.98835424000000  7.004043999999999E-04
11.86860270000000  7.315066800000000E-04
11.74888608000000  7.643602400000000E-04
11.62903906170000  7.990854800000000E-04
11.50911554790000  8.358124800000000E-04
11.38911909120000  8.746822700000000E-04
11.26911909000000  9.158772000000000E-04
11.14891267800000  9.594743599999999E-04
11.02870615950000  1.005742500000000E-03
10.90843260000000  1.054847500000000E-03
10.78809371850000  1.107025000000000E-03
10.66769083290000  1.162439500000000E-03
10.54722560490000  1.221415000000000E-03
10.42669975350000  1.284195000000000E-03
10.30611494040000  1.351092100000000E-03
10.18547254080000  1.422433800000000E-03
10.06477421640000  1.498582400000000E-03
  9.94021227800000  1.579935900000000E-03
  9.82321529400000  1.666931900000000E-03
```
9.702358592400000 1.7600510000000000E-03

Cursor at page: 1; Page printed: 1
<ret>, +n, -n
<ret>

9.581451925200000 1.8598255000000000E-03
9.460497240600000 1.9668394000000000E-03
9.339496200300000 2.0817409000000000E-03
9.218450351400000 2.2052468000000000E-03
9.097361183700000 2.3381521000000000E-03
8.976230416200000 2.4813407000000000E-03
8.855096533000000 2.6357956000000000E-03
8.733850556700000 2.8026134000000000E-03
8.612604443000000 2.9830188000000000E-03
8.491323207000000 3.1783830000000000E-03
8.370085638000000 3.3902437000000000E-03
8.248661775300000 3.6203295000000000E-03
8.127284847000000 3.8705876000000000E-03
8.005879096800000 4.1432166000000000E-03
7.884446472900000 4.4407044000000000E-03
7.762988350500000 4.7658742000000000E-03
7.641506391300000 5.1219380000000000E-03
7.520002371600000 5.5125575000000000E-03
7.398477239000000 5.9419232000000000E-03
7.276934224500000 6.4148409000000000E-03

Cursor at page: 2; Page printed: 2
<ret>, +n, -n
-1

11.988354240000000 7.0040339999999999E-04
11.868660270000000 7.3150668000000000E-04
11.748886080000000 7.6436024000000000E-04
11.629039061700000 7.9908548000000000E-04
11.509115547900000 8.3581248000000000E-04
11.389119091200000 8.7468227000000000E-04
11.269050894900000 9.1584772000000000E-04
11.148912678000000 9.5947435999999999E-04
11.028706159500000 1.0057425000000000E-03
<table>
<thead>
<tr>
<th>Value 1</th>
<th>Value 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>10.908432600000000</td>
<td>1.0548475000000000E-03</td>
</tr>
<tr>
<td>10.788093718500000</td>
<td>1.1070025000000000E-03</td>
</tr>
<tr>
<td>10.667690832900000</td>
<td>1.1624395000000000E-03</td>
</tr>
<tr>
<td>10.547225604900000</td>
<td>1.2214115000000000E-03</td>
</tr>
<tr>
<td>10.426699753500000</td>
<td>1.2841950000000000E-03</td>
</tr>
<tr>
<td>10.306149404000000</td>
<td>1.3510921000000000E-03</td>
</tr>
<tr>
<td>10.185472540800000</td>
<td>1.4224338000000000E-03</td>
</tr>
<tr>
<td>10.064774216400000</td>
<td>1.4985824000000000E-03</td>
</tr>
<tr>
<td>9.944021227800000</td>
<td>1.5799359000000000E-03</td>
</tr>
<tr>
<td>9.823215294000000</td>
<td>1.6669319000000000E-03</td>
</tr>
<tr>
<td>9.702358592400000</td>
<td>1.7600515000000000E-03</td>
</tr>
</tbody>
</table>

2.297641407600000 | 0.5318510000000000 |
2.176784706000000 | 0.5574403000000000 |
2.055978772200000 | 0.5825618000000000 |
1.935225783600000 | 0.6072831000000000 |
1.814527459200000 | 0.6316712000000000 |
1.693885059600000 | 0.6557913000000000 |
1.573300246500000 | 0.6797072000000000 |
1.452774395100000 | 0.7034808000000000 |
1.332309167100000 | 0.7271726000000000 |
1.211906281500000 | 0.7508422000000000 |
1.091567400000000 | 0.7745487000000000 |
0.971293840500000 | 0.7983509000000000 |
0.851087322000000 | 0.8223084000000000 |
0.730949105100000 | 0.8464814000000000 |
0.610880908800000 | 0.8709326000000000 |
0.490884521000000 | 0.8957261000000000 |
0.370960938300000 | 0.9209299000000000 |
0.251139200000000 | 0.9466155000000000 |
0.131339730000000 | 0.9728594000000000 |
1.164579999999949E-02 | 0.9997437000000000 |

Cursor at page: 2; Page printed: 1
<ret>, +n, -n
+2

Cursor at page: 5; Page printed: 5
<ret>, +n, -n

$
VARIANCE

Purpose: Analyzes a formatted two column file (such as output from PREPLOT) and returns the mean, variance and standard deviation for the selected column. A plottable file of the distribution function is optional.

Input: Any formatted two column file.

Output: TT and a user-named log file. A plottable file is optional.

Description:

Gives parameters for the statistical distribution of a column from a formatted file: mean, variance, standard deviation. The results are saved in a log file named by the user. A best fit plottable file is also optional: the user specifies the interval, mean and number of points to plot in the interval. It is useful to define the interval as the difference between the extrema of the data, which can be found using FMAX before running VARIANCE as in the example below.

Examples:

// If we want a plottable file, we should find the extrema using FMAX.

// We’re using the spherical mirror example (pri sec 2.4).

$ fmax
File-name ?
pre.dat
Read 1000 points.
Found :
XMIN -0.4102450 . Set at : -0.5023146
XMAX 0.4081510 0.5002205
YMIN -2.7292412E-02 -3.3435397E-02
YMAX 2.7311902E-02 3.3454888E-02
Good luck!
$ variance
Log file ?
var.dat
File for analysis ?
[.ug.variance]pre.dat
Read 1000 from
[.ug.variance]pre.dat
Column to analyze ?
1

Results from column 1
Mean value: -8.2237018150000001E-03
Variance : 5.5424558469452820E-02
Standard deviation : 0.2354242096077904

Enter:
0 to exit
1 new file
2 new column (same file)
3 best fit (plottable)
?
3
Output file name ?
var.plt
Enter interval width, number of points
Interval width ?
1
Interval center ?
0
Number of points ?
30
Area = 1 ?
1
Enter:
0 to exit
1 new file
2 new column (same file)
3 best fit (plottable)
? 0
Figure 7.3: VARIANCE best fit example
Appendix A

Toroidal Grating Monochromator

Throughout the examples in this manual we have referred to a 6 meter toroidal grating monochromator (TGM). The monochromator modelled is a six-meter installed at the Synchrotron Radiation Center on the ring Aladdin. The system consists of an elliptical entrance mirror, an entrance slit, a toroidal grating, an exit slit and an elliptical refocus mirror. The two mirrors and the grating are gold coated. It has three holographic gratings, low, medium and high energy. The grating we have used for all our examples is the medium energy grating.

We follow with the output from SYSINFO and MIRINFO for each element. The screens defined are there to show the location of valves or future filters.

```
+---------------------------------------------------------------------+
|                         S Y S T E M   D E S C R I P T I O N                     |
+---------------------------------------------------------------------+
| 6-m TGM                                                              |
+---------------------------------------------------------------------+

Input file specified:
+---------------------------------------------------------------------+
| #   Optical Element:  Creation Time:  |
| 1   CXRL$DUA0:[SHADOW.XRAYOP.DOCUMENT.TGM]END.01;56  1-NOV-1989 10: |
| 2   CXRL$DUA0:[SHADOW.XRAYOP.DOCUMENT.TGM]END.02;55  1-NOV-1989 10: |
| 3   CXRL$DUA0:[SHADOW.XRAYOP.DOCUMENT.TGM]END.03;5  1-NOV-1989 10: |
+---------------------------------------------------------------------+

Optical Element #  1  System Number:
<table>
<thead>
<tr>
<th>Orientation</th>
<th>Source Plane</th>
<th>Incidence Ang.</th>
<th>Reflection Ang.</th>
<th>Image Plane</th>
</tr>
</thead>
<tbody>
<tr>
<td>180.0000000000000</td>
<td>180.0000000000000</td>
<td>84.99999999999999</td>
<td>84.99999999999999</td>
<td>180.0000000000000</td>
</tr>
</tbody>
</table>

Optical Element # 2 System Number:

<table>
<thead>
<tr>
<th>Orientation</th>
<th>Source Plane</th>
<th>Incidence Ang.</th>
<th>Reflection Ang.</th>
<th>Image Plane</th>
</tr>
</thead>
<tbody>
<tr>
<td>180.0000000000000</td>
<td>211.1000000000000</td>
<td>80.00000000000002</td>
<td>80.00000000000002</td>
<td>388.8000000000000</td>
</tr>
</tbody>
</table>

SCREENS: 4 defined.
AFTER Mirror at 190.0000000000000
Type :SLIT ELLIPTICAL
AFTER Mirror at 76.0000000000000
BEFORE Mirror at 329.0000000000000
Type :SLIT ELLIPTICAL
BEFORE Mirror at 388.7900000000000

Optical Element # 3 System Number:

<table>
<thead>
<tr>
<th>Orientation</th>
<th>Source Plane</th>
<th>Incidence Ang.</th>
<th>Reflection Ang.</th>
<th>Image Plane</th>
</tr>
</thead>
<tbody>
<tr>
<td>180.0000000000000</td>
<td>240.0000000000000</td>
<td>80.00000000000002</td>
<td>80.00000000000002</td>
<td>80.00000000000000</td>
</tr>
</tbody>
</table>

OPTICAL SYSTEM CONFIGURATION
Laboratory Reference Frame.

<table>
<thead>
<tr>
<th>OPT. Elem #</th>
<th>X =</th>
<th>Y =</th>
<th>Z =</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.0000000000000E+00</td>
<td>0.0000000000000E+00</td>
<td>0.0000000000000E+00</td>
</tr>
<tr>
<td>1</td>
<td>0.0000000000000E+00</td>
<td>180.00000000</td>
<td>0.0000000000000E+00</td>
</tr>
<tr>
<td>1'</td>
<td>0.1377432066E-14</td>
<td>357.2659554</td>
<td>-31.25671980</td>
</tr>
<tr>
<td>2</td>
<td>0.29928779894E-14</td>
<td>565.15831220</td>
<td>-67.913802286</td>
</tr>
<tr>
<td>2'</td>
<td>-0.5842516932E-14</td>
<td>948.0516657</td>
<td>-0.39939080863</td>
</tr>
<tr>
<td>3</td>
<td>-0.11296520633E-13</td>
<td>1184.4054273</td>
<td>41.276171831</td>
</tr>
<tr>
<td>3'</td>
<td>-0.94785309865E-14</td>
<td>1263.1900475</td>
<td>27.384317618</td>
</tr>
</tbody>
</table>

To show the system layout, we ran the beamline through SYSPLOT.

Surface figure was defined as: ELLIPTICAL
Cylindrical figure NO
Element type REFLECTOR
Reflectivity ON coefficients from: AU.REF
Polarization dependence YES
Mirror dimensions (rectangular):
X plus    X minus    Y plus    Y minus
2.0000   -2.0000    4.7500   -4.7500
Figure A.1: TGM Layout – side view
Figure A.2: TGM Layout – top view
Central Axis parameters:
Source Plane Distance 180.0000000000000
Image Plane 180.0000000000000
Incidence Angle 84.99999999999999
Reflection/Diffraction Angle 84.99999999999999

Mirror parameters
Same configuration as Central Axis YES
Objective focus at 180.0000000000000
Image focus at 180.0000000000000
Incidence angle 84.99999999999999

Parameters used follow:
Semi-major axis 180.0000000000000
Semi-minor axis 15.68803369457847
Semi-focal-length 0.0000000000000000E+00
Eccentricity 0.0000000000000000E+00

Source of this O.E. moved NO
Mirror at pole position ( no mov. ) YES

Grating
Orientation 180 deg

Surface figure was defined as: TOROIDAL
Cylindrical figure NO
Element type REFLECTOR
Element type GRATING
Order chosen ( inside are < 0 ) -1.0000000000000000
Automatic Tuning YES
Mount SEYA / TGM
Grating tuned at \[ \text{eV} \] 59.00000000000000
Holographic grating. Recording Wavelength: 4879.860000000000

<table>
<thead>
<tr>
<th>Input Slit Dist.</th>
<th>Exit Slit Dist.</th>
<th>Input Slit Angle</th>
<th>Exit Slit Angle</th>
</tr>
</thead>
<tbody>
<tr>
<td>919.300000</td>
<td>619.000000</td>
<td>10.3600000</td>
<td>-12.8100000</td>
</tr>
</tbody>
</table>

Input Slit rotation angle 0.000000000000000E+00
Output Slit rotation angle 0.000000000000000E+00
Spherical / Spherical
Spherical / Spherical
Real / Real
Reflectivity ON coefficients from: AU.REF
Polarization dependence YES

Mirror dimensions (rectangular):

<table>
<thead>
<tr>
<th>X plus</th>
<th>X minus</th>
<th>Y plus</th>
<th>Y minus</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.2500</td>
<td>-2.2500</td>
<td>6.7500</td>
<td>-6.7500</td>
</tr>
</tbody>
</table>

Central Axis parameters:

<table>
<thead>
<tr>
<th>Source Plane Distance</th>
<th>211.10000000000000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Image Plane</td>
<td>388.80000000000000</td>
</tr>
<tr>
<td>Incidence Angle</td>
<td>80.00000000000002</td>
</tr>
<tr>
<td>Reflection/Diffraction Angle</td>
<td>80.00000000000002</td>
</tr>
</tbody>
</table>

Mirror parameters EXTERNAL

Parameters used follow:

<table>
<thead>
<tr>
<th>Major Radius</th>
<th>1877.550000000000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minor Radius</td>
<td>47.05000000000000</td>
</tr>
</tbody>
</table>

Source of this O.E. moved NO

Mirror moved from pole. Parameters:

<table>
<thead>
<tr>
<th>Displacement along X:</th>
<th>0.0000000000000000E+00</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y:</td>
<td>0.0000000000000000E+00</td>
</tr>
<tr>
<td>Z:</td>
<td>0.0000000000000000E+00</td>
</tr>
</tbody>
</table>

Rotation around X: 2.853995871245999

<table>
<thead>
<tr>
<th>Y:</th>
<th>0.0000000000000000E+00</th>
</tr>
</thead>
<tbody>
<tr>
<td>Z:</td>
<td>0.0000000000000000E+00</td>
</tr>
</tbody>
</table>

******************** MIRROR DESCRIPTION ********************
Refocus Mirror
Orientation 180 deg.

Input file specified: end.03
Full file Specification: CXRL$DUA0: [SHADOW.XRAYOP.DOCUMENT.TGM] END.03; 3
Creation Date: 24-OCT-1989 15:15

Surface figure was defined as: ELLIPTICAL
Cylindrical figure: NO
Element type: REFLECTOR
Reflectivity: OFF
Mirror dimensions: UNLIMITED

Central Axis parameters:
Source Plane Distance: 240.000000000000000
Image Plane: 80.000000000000000
Incidence Angle: 80.00000000000002
Reflection/Diffraction Angle: 80.00000000000002

Mirror parameters: COMPUTED
Same configuration as Central Axis: YES
Objective focus at: 240.000000000000000
Image focus at: 80.00000000000002
Incidence angle: 80.00000000000002

Parameters used follow:
- Semi-major axis: 160.000000000000000
- Semi-minor axis: 24.06139730886965
- Semi-focal-length: 0.0000000000000000E+00
- Eccentricity: 0.0000000000000000E+00

Source of this O.E. moved: NO
Mirror at pole position (no mov.): YES
Appendix B

Glossary

*Active Surface*  
A surface where the rays are reflected/refracted.

*Aligned System*  
An optical system where nothing has been "moved" when running SHADOW.

*Aperture*  
A slit or hole at a screen that limits the latitude of the beam.

*BATCH mode*  
Input mode that takes parameter files (START.xx, SYSTEM-FILE.DAT) generated by a previous input session and supplies them to SHADOW for a re-run. You will simply be prompted for these parameter file names.

*Beam*  
A collection of rays, pretty much like in flashlight beam. The rays can be random or ordered (grid). The beam is stored in the BOF at different locations.

*Beamline*  
The hardware implementation of an optical system for XUV, used to relay with or without monochromatization the beam from source to sample. Optical System.

*Bins*  
Divisions of a histogram that determine the resolution of ray statistics; A greater number of bins yields higher resolution but less reliable statistics, i.e. a smaller signal/noise ratio.

*Blaze*  
The blaze condition proper applies only to triangular grooves, when the light is incident and diffraction angles, relative to the grating normal, are such that the ray is exactly "reflected" by the facet. In other words, the incoming ray forms with the facet normal the same angle as the diffracted beam. It is the condition of maximum efficiency of the grating. Lamellar and sinusoidal gratings have both high-efficiency angles, but for other reasons (see *Gratings*).
**BOF**

Binary Output File – the large binary files generated by the run of SHADOW. They contain the full beam description at some location along the optical system. Use TRANSLATE to type/print them.

**Central Ray**

The ray generated by SOURCE with \(x = y = z = 0\), \(x' = z' = 0\), and \(y' = 1\). It is the ray propagating exactly along the source Y axis. It coincides with the optical axis, in the case of an aligned system.

**Collimated**

The condition of a beam in which all rays are parallel, i.e., the object/image is at infinity.

**Column**

Each ray has up to eighteen values defining it in terms of \(X\), \(Y\), \(Z\), \(X'\), \(Y'\), \(Z'\) etc.; they form the columns containing \(X\), \(Y\), etc.

**Continuation Plane**

An intermediate plane perpendicular to the \(Y\) axis that acts as a delimiter between two optical elements, or as the image plane at the end of the optical system. Continuation planes may or may not coincide with the beam focus. The beam information for the XXth continuation plane is located in the BOF STAR.XX.

**Cylindrical Figure**

A figure originated by the translation of a plane curve perpendicular to the plane. If the plane curve is an ellipse, we speak of elliptical cylinder; if it is a circle, of circular cylinder; etc.

**DCL**

DIGITAL Command Language - The command language for the VAX/VMS operating system.

**DEFINE**

(OE, SCREEN) This MENU mode command is used declare existence of OEs and SCREENs in the System. The elements are numbered starting from the source, which is OE 0. Elements can be inserted and deleted from the system, since the pattern for numbering the OEs of the OS follows that of a stack. Similarly, SCREENS are placed within each OE.

**DELETE**

(OE, SCREEN) Menu mode command used to remove either an OE from the SYSTEM list or a screen from an OE. The system is renumbered down.
**Diffraction Grating**

An element whose surface produces a strongly peaked autocorrelation power series. The relative height of the first, second, etc. peak represents the relative efficiency of the first, second, ... diffraction order. As the ACF depends on the coherent spacing for the peak position, the diffraction angles are determined solely by the grating pitch. The height of the peaks represents the Fourier components of the surface at each groove, so that the efficiency depends on the shape of the grooves. The condition of blaze is usually the most efficient.

**Diffraction Limit**

The observed width of a (geometrically) point-like image. It is due to the finite wavelength of the radiation used to reconstruct the image and to the finite convergence angle. Typically:

\[ \delta = \frac{\lambda}{2 \text{N.A.}} \]

**Diffraction Limited Optics**

An optical system where the dimensions of the image are larger than those of the geometrical image and defined by the radiation employed. For a focusing system, the diffraction limited focus is given by \(\frac{\lambda}{2 \text{N.A.}}\), N.A. being the numerical aperture of the system. Aberrations and figure errors mainly modify the intensity distribution within the limits predicted by the diffraction limit.

**Diffraction Order**

See Order.

**Element**

The part of an optical system that encloses an active surface.

**ERF**

Exit Record File - Contains information about a source or optical element after the rays have been traced through the system. Information such as computed parameters, and positions in the X,Y,Z and Laboratory Reference Frames are given. SHADOW gives the ERFs the default name END.XX.

**EXIT**

Exit from SHADOW command level to VMS DCL level, restoring the terminal status to the standard one.

**External**

Pertaining to the definition of an optical element; the parameters are supplied by the user rather than being computed by SHADOW.

**Figure**

The mathematical function describing the surface of a mirror or of a lens. We speak of elliptical figure, etc.
**Figure Limited Optics**

Optical systems where the dimensions of the image are dominated by the figure errors. At grazing angle, slope errors are the most serious. Common problems with high brightness sources (small dimensions).

**Flag**

Tags a ray if it was not transmitted through an OE or screen. The Flag is 1 if the ray passed and < 0 if it was lost. The Flag is the tenth column in the BOFs; it may assume other values as well.

**Footprint**

The intensity pattern of a beam on the surface of an optical element.

**Fresnel Equations**

Determine the reflectivity of a medium for the two polarization directions as a function of incident and transmitted angles:

\[
R_\perp = \frac{\sin^2(\theta_i - \theta_t)}{\sin^2(\theta_i + \theta_t)}, \quad R_\parallel = \frac{\tan^2(\theta_i - \theta_t)}{\tan^2(\theta_i + \theta_t)}
\]

where \(\theta_i\) and \(\theta_t\) are the angles of incidence and transmission respectively, and \(R_\perp\) and \(R_\parallel\) correspond to the reflectivities in the perpendicular (s-polarization) and parallel (p-polarization) directions with respect to the plane of incidence.

**Geometrical Optics**

The study of an optical system purely on the basis of geometrical relations between rays and surfaces.

**GOTO**

MENU mode command that allows the user to set the pointer to a particular OE. Useful for forgotten screens or if you’re in the (good) habit of DEFINing all OEs at once and then using the SELECT command to input parameters.

**Grating Horizon**

The wavelength such that, for a particular incidence angle, the diffracted first order would be at \(-90^\circ\), i.e., diffracted back and tangent to the surface. Longer wavelengths would be diffracted ‘inside’ the surface, so that the grating behaves like a normal mirror for wavelengths beyond the horizon.
**Grating Mount**
The particular optical system configuration chosen to implement an instrument. A mount is specified by the geometry of the grating element and by focal conditions. Also used, more loosely, to indicate a type of scanning action, like in “TGM mount”: this simply means a mount with Constant Included angle and rotation achieved by simple rotation of the grating.

**Grazing Angle**
An Incidence or reflection angle in which the rays are nearly parallel to the optical surface.

**Groove Shape**
Refers to a grating’s grooves. They can be generally divided in lamellar, triangular or sinusoidal.

**HELP**
Evokes the help facility at the SHADOW command level and provides some on-line description of the local commands and their purposes.

**HELPSHA**
Invokes SHADOW help facility at DCL command level.

**Image Distance**
The distance along the Y axis from an optical element to the next continuation plane, which may or may not be the location of a real image or focus point.

**Incidence Angle**
The angle that an incident ray makes with respect to the normal of the optical surface in question.

**Input Modes**
Means to enter the parameters characterizing source and OE. The three input modes are PROMPT, MENU and BATCH.

**LIST**
Menu mode command that gives a list of the currently defined OEs and SCREENs in an OS and their relative numbering.

**LOAD**
(SYSTEM, SOURCE, OE) MENU mode command that invokes the LOAD menu. This command LOADs an already existing SYSTEM, OE or SOURCE by reading existing parameter file(s) into memory. This is useful for major or minor modifications. The parameters requested are the directory and file-name root describing the location of the information you wish the menu to access. Use “[ ]” or nothing to specify the current directory; the default file-root name is START unless you’ve renamed it. When you have defined the files, hit the ← to return to the “Ready::” prompt.
Marginal Rays

The rays on the periphery (margin) of the beam. They are normally the ones suffering the most aberration.

MENU

This command is used to define, examine and modify parameters defining source and/or optical elements. The order of entry is entirely free. The system displays a list of variables with their current values; each item is accessed using the cursor’s arrows. No calculations are performed and only START files are generated.

MENU mode

One of the three input modes. The program for MENU is subdivided into a COMMAND interpreter and a set of screen oriented menus. It is the most powerful method. A list of parameters are presented on the screen and these can be randomly accessed. This method offers the flexibility to make a single change thus bypassing the lengthy process of the PROMPT mode. It allows also the possibility of going back and forth between the parameters specifying an OE or source. On the other hand, one runs the risk of leaving out a single parameter, and spending a sleepless night wondering why a plane mirror magnified the source. MENU has its own command level and the prompt is “Ready::”. The possible commands are: DEFINE, DELETE, GOTO, LIST, LOAD, SAVE, SELECT, SPAWN.

Momentum Space

The momentum \( \hat{p} \) is the conjugate variable to the coordinate \( \hat{r} \). In general,

\[
\hat{p} = \frac{2\pi}{\lambda} \{a_x, a_y, a_z\}
\]

where \( a \)'s are the cartesian cosines

\[
a_i = \frac{\hat{p} \cdot \hat{x}_j}{|\hat{p}|} \]

Rows 4,5,6 store the \( a \)'s of the ray, while \( |\hat{p}| = \frac{2\pi}{\lambda} \) is stored in row 11. The coordinates in momentum space for a ray are in columns 4, 5, and 6 in any BOF. Also called reciprocal space.
Monochromator
An optical instrument capable of transmitting a very narrow bandpass of a white spectrum. Normally consisting of one or more gratings/crystals in a particular mount. Notice that the monochromator is an optical device whose function is simply to define the bandpass: it does not necessarily refocus the diffracted beam. It is normally teamed with condensing and refocusing optics.

Namelist
Text file that contains the input parameters for the source or optical system. These files (named by SHADOW as START.XX) are the only files that are needed, besides the files created by the preprocessors, to regenerate the beam data for the entire system.

Normal
A vector which is perpendicular to the surface in question. Usually located at the geometric center of an optical surface.

Numerical Aperture
The sine of the angle formed by a ray with the optical axis.

Optical Axis
The ray connecting the poles of the optical elements. It connects the origin points of the consecutive OE's. Notice that even when the mirror are “moved”, the optical axis remains the same. Same as central ray, if the source has not been “moved” in SHADOW.

Optical Efficiency
The ratio of the number of photons departing to the number incident on an optical element (or system).

Optical Element or O.E.
The part of the optical system encompassing an active surface. It is delimited by two planes (source/object and image, respectively) located along the optical axis at some convenient location.

Optical Path
The optical path of a ray is given by:

$$T_{BA} = \int_{a}^{b} n(s)ds$$

where $n(s)$ is the index of refraction as a function of position, and $a$ and $b$ are the initial and terminal points of the ray. The focal condition states that all the optical paths are the same, independent of angle.

Optical system
The collection of optical elements under study.
Orders
The labels tagged to a beam when diffracted from a grating. The European convention defines negative the orders diffracted inside (inside order) the zero-order reflection beam, positive those laying outside. The American convention is the opposite. Most XUV systems work in the first order and the XUV community follows the European convention.

Phase Space
The union of Real and Momentum spaces. Phase space projections may aid in interpreting the character of the beam at a given point in the system by determining the correlation between two of the components such as $X$ and $X'$. The volume of phase space occupied by a beam is always conserved.

Physical Optics
The study of an optical system that explicitly includes physical aspects, such as reflectivity, diffractions, etc.

Preprocessor
Any utility that generates data that is used for SOURCE generation or defining an element of the optical system, e.g., PREREF is used to determine the reflection and transmission coefficients for a given Atomic Number.

PROMPT mode
Input mode in SHADOW that asks the user for parameters to cover the full specification of a source or OE. Useful when setting up a new system.

Ray
The path described by a single photon. It is specified by 12-18 double precision real numbers. They are described by a starting point, a direction, an electric field, a phase, a polarization degree, an energy, a status flag.

Ray Index
Index used to number the rays in order of their generation: the first ray is given Ray Index 1, the second 2, etc... The Ray index is the 12th column in the BOFs.

Real Space
The ordinary geometrical space. The real space coordinates for a ray are in columns 1, 2, and 3 in the BOFs.

Reflection Angle
The angle that a reflected or refracted ray that is coincident with the $Y$ axis makes with respect to the normal of the optical surface in question.
**Roughness**
The incoherent part of a surface height autocorrelation power series. It causes a diffuse scattering of radiation away from the geometrical image. Although it does not affect the image sharpness, it decreases the system MTF. The roughness becomes less important at glancing angles, due to the roll-off in the power spectrum and to the increase in the photon horizontal wavevector at glancing angles.

**Sagittal**
A direction perpendicular to the incidence plane (s-polarization).

**SAVE**
(SYSTEM, SOURCE, OE) MENU mode command that invokes the SAVE menu. The parameters requested are the directory and the file-name root, i.e., where and under what name do you want the data stored. This is the command to use to write the definition of OE, SOURCE or entire SYSTEM to disk. SAVE SYSTEM also creates SYSTEM-FILE.DAT; when in doubt, always use SAVE SYSTEM.

NOTE: You *MUST* issue the SAVE command on exit from the MENU or else the data is lost. SHADOW will warn you if you have modified a file and not saved it.

**Seed**
An odd eight-digit number used as input to a pseudo-random number generator for creating source photon distributions.

**SELECT**
(SOURCE, OE, SCREEN) MENU mode command that SELECTs which OE or SCREEN you wish to characterize. This applies also for the SOURCE. After issuing the SELECT OE (or SCREEN) command, you are prompted for the element number (screen/slit number). This positions the pointer to the chosen location. Note, an OE or screen must be DEFINEd before SELECTed, though SELECT SOURCE automatically verifies its existence. Before issuing the SELECT SCREEN command, you must be in a defined (i.e., existing) OE.

**Shape**
The shape of the mirror outline or contour: a spherical mirror can be round or rectangular. The same is true for any mirror figure.

**Sigma**
Usually refers to the standard deviation of the distribution function in question.
**SOURCE**

Used to define the source parameters and generate the source rays. After entering the command the user will be prompted to choose an input mode during which the physical description is set up.

*Source Depth*

The distribution of the source points in the \( Y \) direction.

*Source Distance*

The distance along the \( Y \) axis from the source or previous continuation plane to the next optical element in the system.

*Source Limited optics*

An optical system where the dimensions of the image are dominated by the aberrations caused by the large extent of the source, either in plane or in angle. Diffraction and figure errors have a small effect.

**SPAWN**

MENU mode command that spawns the process, giving full access to DCL without interrupting the menu session, by creating a new VMS session. The program will resume where it left when you log off from the sub-process.

Note: Be sure to “log-off” from all sub-processes when done.

*Tangential*

A direction laying in the incidence plane (p-polarization).

**TRACE**

Used to define the optical elements and the optical system. On completion of the OS definition, TRACE computes the path of each ray through the OS. A source must exist before issuing the command TRACE.

*Virtual Image*

Rays emerging from an OE or OS in image space are diverging, so image appears to exist in object space.

*Virtual Object*

Rays entering an OE or OS from object space are converging, so object appears to exist in image space.

*Wavenumber*

The wavenumber for a photon is given by \( K = \frac{2\pi}{\lambda} \), where \( \lambda \) is the wavelength. (see *Momentum*.)

*Zero order*

The beam reflected by a grating without scattering, whereby it acts like a simple mirror.