Inverse Adding-Doubling

(Version 3-12-0)

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§1 IAD (v 3-12-0) MAIN PROGRAM 1

1. Main Program.

Here is a relatively robust command line utility that shows how the iad and ad subroutines might be called. It suffers because it is written in CWEB and I used the macro expansion feature instead of creating separate functions. Oh well.

I create an empty file iad_main.h to simplify the Makefile $\langle iad_main.h \ 1 \rangle \equiv$

```
2. All the actual output for this web file goes into iad_main.c
```

```
\langle iad_main.c \rangle \equiv
   \langle Include files for main 3\rangle\langle print version function 20\rangle\langle print usage function 21\rangle\langle stringdup together
                    function 27\langle seconds elapsed function 28\langle print error legend function 26\langle print dot
                    function 30 \ \calculate coefficients function 22 \ \chi parse string into array function 29 \ \chi print
                    results header function 24\\Print results function 25\\int main(int argc, char **argv){
                    \langle \text{ Declare variables for } main \ 4 \rangle \langle \text{ Handle options } 5 \rangle Initialize\_Measure(\&m);
              \langle \text{Command-line changes to } m \ 18 \rangle Initialize\_Result(m, \&r); \langle \text{Command-line changes to } r \ 13 \rangle
              if (cl\_forward\_calc \neq \texttt{UNINITIALIZED}) {
                 ⟨ Calculate and Print the Forward Calculation 6⟩ return 0;
              \langle prepare file for reading 10\rangle
              if (process_command_line) {
                 \langle Count command-line measurements 19\rangle\langle Calculate and write optical properties 11\ranglereturn 0;
              if (Read\_Header(stdin, \&m, \&params) \equiv 0) {
                 start\_time = clock();
                 while (Read\_Data\_Line(stdin, \&m, params) \equiv 0)
                 \{\langle \text{Command-line changes to } m \text{ 18} \rangle \langle \text{Calculate and write optical properties } 11 \rangle \}
              if (cl\_verbosity > 0) fprintf(stderr, "\n");
              if (any\_error \land cl\_verbosity > 1) print\_error\_legend();
              return 0; }
```

2 MAIN PROGRAM IAD (v 3-12-0) §3

3. The first two defines are to stop Visual C++ from silly complaints

```
\langle Include files for main 3 \rangle \equiv
#define _CRT_SECURE_NO_WARNINGS
#define _CRT_NONSTDC_NO_WARNINGS
\#define NO_SLIDES 0
#define ONE_SLIDE_ON_TOP 1
\#define TWO_IDENTICAL_SLIDES 2
#define ONE_SLIDE_ON_BOTTOM 3
#define ONE_SLIDE_NEAR_SPHERE 4
#define ONE_SLIDE_NOT_NEAR_SPHERE 5
#define MR_IS_ONLY_RD 1
#define MT_IS_ONLY_TD 2
\#define NO_UNSCATTERED_LIGHT 3
#include <stdio.h>
#include <string.h>
#include <stdlib.h>
#include <time.h>
#include <math.h>
#include <ctype.h>
#include "ad_globl.h"
#include "ad_prime.h"
#include "iad_type.h"
#include "iad_pub.h"
#include "iad_io.h"
#include "iad_calc.h"
#include "iad_util.h"
#include "mygetopt.h"
#include "version.h"
#include "mc_lost.h"
#include "ad_frsnl.h"
 extern char *optarg;
 extern int optind;
```

```
4. \langle Declare variables for main \langle \rangle \equiv
           struct measure_type m;
           struct invert_type r;
           char *g\_out\_name = \Lambda;
           char c:
           long n_{-}photons = 100000;
           int MC_{-}iterations = 19:
           int any\_error = 0;
           int process\_command\_line = 0;
           int params = 0;
           int cl_quadrature_points = UNINITIALIZED;
           int cl\_verbosity = 2;
           double cl\_forward\_calc = UNINITIALIZED;
           double cl_{-}default_{-}a = \texttt{UNINITIALIZED}:
           double cl\_default\_g = UNINITIALIZED;
           double cl\_default\_b = \texttt{UNINITIALIZED};
           double cl\_default\_mua = \texttt{UNINITIALIZED};
           double cl\_default\_mus = \texttt{UNINITIALIZED};
           double cl\_tolerance = UNINITIALIZED;
           double cl\_slide\_OD = \texttt{UNINITIALIZED}:
           double cl\_cos\_angle = UNINITIALIZED;
           double cl\_beam\_d = \texttt{UNINITIALIZED};
           double cl\_sample\_d = \texttt{UNINITIALIZED};
           double cl\_sample\_n = \texttt{UNINITIALIZED};
           double cl\_slide\_d = UNINITIALIZED;
           double cl\_slide\_n = \texttt{UNINITIALIZED};
           double cl\_slides = \texttt{UNINITIALIZED};
           double cl\_default\_fr = \texttt{UNINITIALIZED};
           double cl_rstd_t = UNINITIALIZED;
           double cl\_rstd\_r = UNINITIALIZED;
           double cl\_rc\_fraction = UNINITIALIZED;
           double cl_{-}tc_{-}fraction = UNINITIALIZED;
           double cl\_search = UNINITIALIZED;
           double cl\_mus\theta = \texttt{UNINITIALIZED};
           double cl\_musp\theta = \texttt{UNINITIALIZED};
           double cl\_mus\theta\_pwr = \texttt{UNINITIALIZED};
           double cl\_mus0\_lambda = UNINITIALIZED:
           double cl_{-}UR1 = UNINITIALIZED;
           double cl_{-}UT1 = UNINITIALIZED;
           double cl_{-}Tc = UNINITIALIZED;
           double cl\_method = \texttt{UNINITIALIZED};
           double cl\_num\_spheres = UNINITIALIZED;
           \mathbf{double}\ cl\_sphere\_one[5] = \{ \mathtt{UNINITIALIZED}, \mathtt{UNINITIALIZE
                                  UNINITIALIZED \;
           \mathbf{double} \ \ \mathit{cl\_sphere\_two}[5] = \{ \mathtt{UNINITIALIZED}, \mathtt{UNINIT
                                   UNINITIALIZED \;
           clock_t \ start_time = clock();
           \mathbf{char}\ command\_line\_options[] = "?1:2:a:A:b:B:c:C:d:D:e:E:f:F:g:G:hi:n:N:M:o:p:q:r:R:s:S:t:T \setminus f(a) = f
                                   :u:vV:x:Xz";
```

4 MAIN PROGRAM IAD (v 3-12-0) $\S5$

```
use the my\_qetopt() to process options.
\langle Handle options 5\rangle \equiv
        while ((c = my\_getopt(argc, argv, command\_line\_options)) \neq EOF) {
              int n:
              char cc;
              \mathbf{switch}(c) {
               case '1': parse_string_into_array(optarg, cl_sphere_one, 5);
                       break:
               case '2': parse_string_into_array(optarg, cl_sphere_two, 5);
                       break:
               case 'a': cl\_default\_a = strtod(optarg, \Lambda);
                       break:
               case 'A': cl\_default\_mua = strtod(optarg, \Lambda);
                      break:
               case 'b': cl\_default\_b = strtod(optarg, \Lambda);
                       break:
               case 'B': cl\_beam\_d = strtod(optarg, \Lambda);
                       break;
               case 'c': cl\_rc\_fraction = strtod(optarq, \Lambda);
                      if (cl\_rc\_fraction < 0.0 \lor cl\_rc\_fraction > 1.0) {
                             fprintf(stderr, "required: \ 0 <= \ fraction \ of \ unscattered \ refl. \ in \ M_R_ <= \ 1 \ n");
                               exit(0);
                      break:
               case 'C': cl\_tc\_fraction = strtod(optarg, \Lambda);
                      if (cl\_tc\_fraction < 0.0 \lor cl\_tc\_fraction > 1.0) {
                             fprintf(stderr, "required: \_0 \subseteq = \_fraction \_of \_unscattered = trans. \_in \_M_T \subseteq 1 n");
                               exit(0);
                       break:
               case 'd': cl\_sample\_d = strtod(optarg, \Lambda);
                      break;
               case 'D': cl\_slide\_d = strtod(optarg, \Lambda);
                       break;
               case 'e': cl\_tolerance = strtod(optarq, \Lambda);
               case 'E': cl\_slide\_OD = strtod(optarg, \Lambda);
                       break;
               case 'f': cl\_default\_fr = strtod(optarg, \Lambda);
                      break:
               case 'F':
                                                               /* initial digit means this is mus is constant */
                      if (isdigit(optarg[0])) {
                               cl\_default\_mus = strtod(optarg, \Lambda);
                              break:
                                          /* should be a string like 'R 1000 1.2 -1.8' */
                       n = sscanf(optarg, "\c_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\c_{\c}\c_{\c}\c_{\c}\f_{\c}\f_{\c}\f_{\c}\c_{\c}\c_{\c}\f_{\c}\f_{\c}\f_{\c}\c_{\c}\c_{\c}\f_{\c}\f_{\c}\f_{\c}\c_{\c}\c_{\c}\f_{\c}\f_{\c}\f_{\c}\c_{\c}\c_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\f_{\c}\
                       if (n \neq 4 \lor (cc \neq P' \land cc \neq R')) {
                             fprintf(stderr, "Screwy\_argument\_for\_-F\_option.\_\_Try\_something\_like\n");
                             fprintf(stderr, " - F_1 1.0 - L_1 1.0 - L_2 1.0 + L_2 1.0 - L_2 
                             fprintf(stderr, " \_ -F_ \_'P_ \_500 \_ 1.0 \_ -1.3' \_ for_ \_mus_ = 1.0*(lambda/500)^(-1.3) \n");
                             fprintf(stderr, "_{\Box}-F_{\Box}, R_{\Box}500_{\Box}1.0_{\Box}-1.3, T_{\Box}) = 1.0*(lambda/500)^(-1.3) n");
                               exit(1);
```

```
§5 IAD (v 3-12-0)
```

```
if (cc \equiv 'R' \lor cc \equiv 'r') {
     cl\_musp\theta = cl\_mus\theta;
     cl\_mus\theta = {\tt UNINITIALIZED};
  break;
case 'g': cl\_default\_g = strtod(optarg, \Lambda);
  break:
case 'G':
  if (optarg[0] \equiv 0,0) cl\_slides = NO\_SLIDES;
  else if (optarg[0] \equiv '2') cl\_slides = TWO\_IDENTICAL\_SLIDES;
  else if (optarg[0] \equiv 't' \vee optarg[0] \equiv 'T') cl\_slides = ONE\_SLIDE\_ON\_TOP;
  else if (optarg[0] \equiv b' \lor optarg[0] \equiv B') cl\_slides = ONE\_SLIDE\_ON\_BOTTOM;
  else if (optarq[0] \equiv 'n' \lor optarq[0] \equiv 'N') cl\_slides = ONE\_SLIDE\_NEAR\_SPHERE:
  else if (optarg[0] \equiv 'f' \lor optarg[0] \equiv 'F') cl\_slides = ONE\_SLIDE\_NOT\_NEAR\_SPHERE;
  else {
    fprintf(stderr, "Argument_{\square}for_{\square}-G_{\square}option_{\square}must_{\square}be_{\square}\n");
    fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup}'t'_{\sqcup}---_{\sqcup}light_{\sqcup}always_{\sqcup}hits_{\sqcup}top_{\sqcup}slide_{\sqcup}first\n");
    fprintf(stderr, "_{\sqcup \sqcup \sqcup \sqcup} 'n'_{\sqcup} ---_{\sqcup} slide_{\sqcup} always_{\sqcup} closest_{\sqcup} to_{\sqcup} sphere \n");
    fprintf(stderr, "____'f'__---_slide_always_farthest_from_sphere\n");
     exit(1);
  break;
case 'i': cl\_cos\_angle = strtod(optarg, \Lambda);
  if (cl\_cos\_angle < 0 \lor cl\_cos\_angle > 90)
     fprintf(stderr, "Incident_langle_lmust_lbe_lbetween_l0_land_l90_ldegrees n");
  else cl\_cos\_angle = cos(cl\_cos\_angle * 3.1415926535/180.0);
case 'M': MC_{-iterations} = (int) strtod(optarg, \Lambda);
  break:
case 'n': cl\_sample\_n = strtod(optarg, \Lambda);
  break;
case 'N': cl\_slide\_n = strtod(optarg, \Lambda);
  break;
case 'o': g\_out\_name = strdup(optarg);
  break:
case 'p': n\_photons = (int) strtod(optarg, \Lambda);
  break;
case 'q': cl\_quadrature\_points = (int) \ strtod(optarg, \Lambda);
  if (cl\_quadrature\_points \% 4 \neq 0) {
    fprintf(stderr, "Number_lof_lquadrature_lpoints_must_lbe_la_lmultiple_lof_l4\n");
     exit(1);
  if ((cl\_cos\_angle \neq UNINITIALIZED) \land (cl\_quadrature\_points \% 12 \neq 0)) {
    fprintf(stderr, "Quadrature\_must\_be_12,_24,_36,..._for\_oblique\_incidence\n");
     exit(1);
  break;
case 'r': cl_{-}UR1 = strtod(optarq, \Lambda);
  process\_command\_line = 1;
  break;
```

6 Main program iad (v 3-12-0) §5

```
case 'R': cl_rstd_r = strtod(optarq, \Lambda);
       break:
     case 's': cl\_search = (int) \ strtod(optarq, \Lambda);
       break;
     case 'S': cl\_num\_spheres = (\mathbf{int}) \ strtod(optarg, \Lambda);
       break;
     case 't': cl_{-}UT1 = strtod(optarg, \Lambda);
       process\_command\_line = 1;
       break:
     case 'T': cl_rstd_t = strtod(optarg, \Lambda);
       break;
     case 'u': cl_{-}Tc = strtod(optarg, \Lambda);
       process\_command\_line = 1;
       break:
     case 'v': print_version();
     case 'V': cl\_verbosity = strtod(optarg, \Lambda);
     case 'x': Set\_Debugging((int) strtod(optarq, \Lambda));
       break:
     case 'X': cl\_method = \texttt{COMPARISON};
       break:
     case 'z': cl\_forward\_calc = 1;
       process\_command\_line = 1;
       break:
     default: fprintf(stderr, "unknown_option_',%c',\n",c); /* fall through */
     case 'h': case '?': print_usage();
       break;
     }
  argc -= optind;
  argv += optind;
This code is used in section 2.
```

6. We are doing a forward calculation. We still need to set the albedo and optical depth appropriately. Obviously when the -a switch is used then the albedo should be fixed as a constant equal to $cl_default_a$. The other cases are less clear. If scattering and absorption are both specified, then calculate the albedo using these values. If the scattering is not specified, then we assume that the sample is an unscattering sample and therefore the albedo is zero. On the other hand, if the scattering is specified and the absorption is not, then the albedo is set to one.

```
 \begin{split} &\langle \text{Calculate and Print the Forward Calculation } 6 \rangle \equiv \\ & \text{if } (\mathit{cl\_default\_a} \equiv \texttt{UNINITIALIZED}) \ \{ \\ & \text{if } (\mathit{cl\_default\_mus} \equiv \texttt{UNINITIALIZED}) \ \mathit{r.a} = 0; \\ & \text{else if } (\mathit{cl\_default\_mua} \equiv \texttt{UNINITIALIZED}) \ \mathit{r.a} = 1; \\ & \text{else } \mathit{r.a} = \mathit{cl\_default\_mus}/(\mathit{cl\_default\_mua} + \mathit{cl\_default\_mus}); \\ & \} \\ & \text{else } \mathit{r.a} = \mathit{cl\_default\_a}; \\ & \text{See also sections } 7, 8, \text{ and } 9. \\ & \text{This code is used in section } 2. \end{split}
```

 $\S7$ IAD (v 3-12-0) MAIN PROGRAM 7

7. This is slightly more tricky because there are four things that can affect the optical thickness — $cl_default_b$, the default mua, default mus and the thickness. If the sample thickness is unspecified, then the only reasonable thing to do is to assume that the sample is very thick. Otherwise, we use the sample thickness to calculate the optical thickness.

```
\langle Calculate and Print the Forward Calculation 6\rangle + \equiv
  if (cl\_default\_b \equiv \mathtt{UNINITIALIZED}) {
     if (cl\_sample\_d \equiv \mathtt{UNINITIALIZED}) r.b = \mathtt{HUGE\_VAL};
     else if (r.a \equiv 0) {
        if (cl\_default\_mua \equiv \texttt{UNINITIALIZED}) r.b = \texttt{HUGE\_VAL};
        else r.b = cl\_default\_mua * cl\_sample\_d;
     else {
        if (cl\_default\_mus \equiv \mathtt{UNINITIALIZED}) r.b = \mathtt{HUGE\_VAL};
        else r.b = cl\_default\_mus/r.a * cl\_sample\_d;
  }
  else r.b = cl\_default\_b;
8. The easiest case, use the default value or set it to zero
\langle Calculate and Print the Forward Calculation 6\rangle + \equiv
  if (cl\_default\_g \equiv \texttt{UNINITIALIZED}) \ r.g = 0;
  else r.g = cl\_default\_g;
9. \langle Calculate and Print the Forward Calculation _{6}\rangle +\equiv
  r.slab.a = r.a;
  r.slab.b = r.b;
  r.slab.g = r.g;
     double mu\_sp, mu\_a, m\_r, m\_t;
     Calculate\_MR\_MT(m, r, MC\_iterations, \&m\_r, \&m\_t);
     Calculate\_Mua\_Musp(m, r, \& mu\_sp, \& mu\_a);
     if (cl\_verbosity > 0) {
        Write\_Header(m, r, -1);
        print_results_header(stdout);
     print\_optical\_property\_result(stdout, m, r, m\_r, m\_t, mu\_a, mu\_sp, 0, 0);
```

8 Main Program IAD (v 3-12-0) §10

10. Make sure that the file is not named '-' and warn about too many files $\langle \text{ prepare file for reading } 10 \rangle \equiv$ if (argc > 1) { $fprintf(stderr, "Only_a_single_file_can_be_processed_at_a_time\n");$ $fprintf(stderr, "try_{\sqcup}'apply_{\sqcup}iad_{\sqcup}file1_{\sqcup}file2_{\sqcup}..._{\sqcup}fileN'\n");$ if $(argc \equiv 1 \land strcmp(argv[0], "-") \neq 0)$ { /* filename exists and != "-" */ int n; char *base_name, *rt_name; $base_name = strdup(argv[0]);$ $n = (\mathbf{int})(strlen(base_name) - strlen(".rxt"));$ if $(n > 0 \land strstr(base_name + n, ".rxt") \neq \Lambda)$ base_name $[n] = `\0';$ rt_name = strdup_together(base_name, ".rxt"); if $(freopen(argv[0], "r", stdin) \equiv \Lambda \land freopen(rt_name, "r", stdin) \equiv \Lambda)$ { $fprintf(stderr, "Could_not_open_either_', s'_or_', s'\setminus n", argv[0], rt_name);$ exit(1);if $(g_out_name \equiv \Lambda)$ $g_out_name = strdup_together(base_name, ".txt");$ $free(rt_name);$ $free(base_name);$ $process_command_line = 0;$ **if** $(g_out_name \neq \Lambda)$ { if $(freopen(g_out_name, "w", stdout) \equiv \Lambda)$ { $fprintf(stderr, "Could_not_open_file_', s'_for_output_n", g_out_name);$ exit(1); }

 $\S11$ IAD (v 3-12-0) MAIN PROGRAM 9

11. Need to explicitly reset *r.search* each time through the loop, because it will get altered by the calculation process. We want to be able to let different lines have different constraints. In particular consider the file *newton.tst*. In that file the first two rows contain three real measurements and the last two have the collimated transmission explicitly set to zero — in other words there are really only two measurements.

```
\langle Calculate and write optical properties |11\rangle \equiv
  { \langle Local Variables for Calculation 12 \rangle
  Initialize\_Result(m, \&r);
  \langle Command-line changes to r 13\rangle
  if (cl\_method \equiv \texttt{COMPARISON} \land m.d\_sphere\_r \neq 0 \land m.as\_r \equiv 0) {
     fprintf(stderr, "A_{\sqcup}dual-beam_{\sqcup}measurement_{\sqcup}is_{\sqcup}specified,_{\sqcup}but_{\sqcup}no_{\sqcup}port_{\sqcup}sizes.\n");
     fprintf(stderr, "You_might_mforsake_the_-X_moption_and_use_zero_spheres_(which_gives_n");
     fprintf(stderr, "the \_same \_result \_except \_lost \_light \_is \_not \_taken \_into \_account). \n");
     fprintf(stderr, "Alternatively, \_bite_\bot the_\bot bullet_\_and_\_enter_\bot your_\bot sphere_\bot parameters, \n");
     fprintf(stderr, "with_the_tknowledge_that_tonly_the_beam_tdiameter_and_sample_port\n");
     fprintf(stderr, "diameter_are_worth_obsessing_over.\n");
     exit(0);
  Write Header 14
  Inverse\_RT(m, \&r); if (r.error \equiv IAD\_NO\_ERROR)
     calculate\_coefficients(m, r, \&LR, \&LT, \&mu\_sp, \&mu\_a);
     (Improve result using Monte Carlo 15)
  }
  print_optical_property_result(stdout, m, r, LR, LT, mu_a, mu_sp, mc_iter, rt_total);
  if (Debug(DEBUG_LOST_LIGHT)) fprintf(stderr, "\n");
  else print_dot (start_time, r . error , mc_total, rt_total, 99, cl_verbosity, & any_error ); }
This code is used in section 2.
12.
\langle \text{Local Variables for Calculation } 12 \rangle \equiv
  static int rt_{-}total = 0;
  static int mc\_total = 0;
  int mc\_iter = 0;
  double ur1 = 0:
  double ut1 = 0;
  double uru = 0:
  double utu = 0;
  double mu_{-}a = 0;
  double mu\_sp = 0;
  double LR = 0;
  double LT = 0;
  rt_{-}total ++;
This code is used in section 11.
```

10 Main Program IAD (v 3-12-0) §13

```
13.
       \langle Command-line changes to r 13\rangle \equiv
  \textbf{if} \ (\textit{cl\_quadrature\_points} \neq \texttt{UNINITIALIZED}) \ \textit{r.method.quad\_pts} = \textit{cl\_quadrature\_points};
  else r.method.quad_pts = 8;
  if (cl\_default\_a \neq UNINITIALIZED) r.default\_a = cl\_default\_a;
  if (cl\_default\_mua \neq \texttt{UNINITIALIZED}) {
     r.default\_mua = cl\_default\_mua;
     if (cl\_sample\_d \neq UNINITIALIZED) r.default\_ba = cl\_default\_mua * cl\_sample\_d;
     else r.default_ba = cl_default_mua * m.slab_thickness;
  if (cl\_default\_b \neq UNINITIALIZED) r.default\_b = cl\_default\_b;
  if (cl\_default\_g \neq UNINITIALIZED) r.default\_g = cl\_default\_g;
  if (cl\_tolerance \neq \texttt{UNINITIALIZED}) {
     r.tolerance = cl\_tolerance;
     r.MC\_tolerance = cl\_tolerance;
  if (cl\_musp\theta \neq UNINITIALIZED)
     cl\_mus0 = (r.default\_g \neq UNINITIALIZED) ? cl\_musp0/(1 - r.default\_g) : cl\_musp0;
  if (cl\_mus0 \neq UNINITIALIZED \land m.lambda \neq 0)
     cl\_default\_mus = cl\_mus0 * pow(m.lambda/cl\_mus0\_lambda, cl\_mus0\_pwr);
  if (cl\_default\_mus \neq UNINITIALIZED) {
     r.default\_mus = cl\_default\_mus;
     if (cl\_sample\_d \neq UNINITIALIZED) r.default\_bs = cl\_default\_mus * cl\_sample\_d;
     else r.default_bs = cl_default_mus * m.slab_thickness;
  if (cl\_search \neq UNINITIALIZED) r.search = cl\_search;
This code is used in sections 2 and 11.
14. \langle \text{ Write Header } 14 \rangle \equiv
  if (rt\_total \equiv 1 \land cl\_verbosity > 0) {
      Write\_Header(m, r, params);
     if (MC\_iterations > 0) {
        if (n\_photons \ge 0)
           fprintf(stdout, "\#_{\sqcup\sqcup}Photons_{\sqcup}used_{\sqcup}to_{\sqcup}estimate_{\sqcup}lost_{\sqcup}light_{\sqcup}=_{\sqcup\sqcup\sqcup\sqcup}%ld\n", n_{\_}photons);
        else fprintf(stdout, "\#_{\sqcup\sqcup\sqcup\sqcup\sqcup} Time_{\sqcup}used_{\sqcup}to_{\sqcup}estimate_{\sqcup}lost_{\sqcup}light_{\sqcup}=_{\sqcup\sqcup\sqcup} %ld_{\sqcup}ms_{\square}", -n_{-}photons);
     else fprintf(stdout, "#_||Photons_used_to_estimate_lost_light_=||0\n");
     fprintf(stdout, "#\n");
     print_results_header(stdout);
This code is used in section 11.
```

§15 IAD (v 3-12-0) MAIN PROGRAM 11

Use Monte Carlo to figure out how much light leaks out. We use the sphere corrected values as the starting values and only do try Monte Carlo when spheres are used, the albedo unknown or non-zero, and there has been no error. The sphere parameters must be known because otherwise the beam size and the port size are unknown.

```
\langle \text{Improve result using Monte Carlo } 15 \rangle \equiv
            if (m.as\_r \neq 0 \land r.default\_a \neq 0 \land m.num\_spheres > 0) { double mu\_sp\_last = mu\_sp;
             double mu\_a\_last = mu\_a;
            if (Debug(DEBUG_LOST_LIGHT)) {
                          print\_results\_header(stderr);
                          print_optical_property_result(stderr, m, r, LR, LT, mu_a, mu_sp, mc_iter, rt_total);
             while (mc\_iter < MC\_iterations) { MC\_Lost(m, r, -1000, \&ur1, \&ut1, \&uru, \&utu, \&m.ur1\_lost, ur1, \&uru, \&
                                       \&m.ut1\_lost, \&m.uru\_lost, \&m.utu\_lost);
             mc\_total++;
             mc\_iter ++;
             Inverse\_RT(m, \&r);
             calculate\_coefficients(m, r, \&LR, \&LT, \&mu\_sp, \&mu\_a);
             if (fabs(mu\_a\_last-mu\_a)/(mu\_a+0.0001) < r.MC\_tolerance \land fabs(mu\_sp\_last-mu\_sp)/(mu\_sp+0.0001) < r.MC\_tolerance \land fabs(mu\_sp-0.0001) < r.MC\_tol
                                                    r.MC_tolerance) break;
             mu_-a_-last = mu_-a;
             mu\_sp\_last = mu\_sp;
             if (Debug(DEBUG_LOST_LIGHT))
                          print_optical_property_result(stderr, m, r, LR, LT, mu_a, mu_sp, mc_iter, rt_total);
             error \neq IAD\_NO\_ERROR ) break; }
```

12 Main Program iad (v 3-12-0) §16

```
16.
                       \langle \text{ Testing MC code } 16 \rangle \equiv
                   struct AD\_slab\_type s;
                   double ur1, ut1, uru, utu;
                   double adur1, adut1, aduru, adutu;
                   s.a = 0.0;
                   s.b = 0.5;
                   s.g = 0.0;
                   s.phase\_function = \texttt{HENYEY\_GREENSTEIN};
                   s.n_{-}slab = 1.0;
                   s.n_{-}top_{-}slide = 1.0;
                   s.n\_bottom\_slide = 1.0;
                   s.b\_top\_slide = 0;
                   s.b_bottom_slide = 0;
                  MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
                  RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                   fprintf(stderr, \n=\%5.4f_b=\%5.4f_g=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%
                                      s.n\_top\_slide);
                   fprintf(stderr, "
                   fprintf(stderr, "_{UU}AD_{UUU}MC_{UUUUUUUA}AD_{UUU}MC_{UUUUUUU}AD_{UUU}MC_{UUUUUUUUU}AD_{UUU}MC_{U} \n");
                   fprintf(stderr, "\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}", adur1, ur1, adut1, ut1);
                   fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", aduru, uru, adutu, utu);
                   s.b = 100.0;
                   s.n_{-}slab = 1.5;
                   s.n\_top\_slide);
                   MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
                   RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                   fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", adur1, ur1, adut1, ut1);
                   fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", aduru, uru, adutu, utu);
                   s.n_{-}slab = 2.0;
                   fprintf(stderr, "\n=\%5.4f_b=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=
                                      s.n\_top\_slide);
                   MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
                   RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                   fprintf(stderr, "\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}", adur1, ur1, adut1, ut1);
                   fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", aduru, uru, adutu, utu);
                   s.n_{-}slab = 1.5;
                   s.n_{-}top_{-}slide = 1.5;
                   s.n\_bottom\_slide = 1.5;
                   fprintf(stderr, \n=\%5.4f_b=\%5.4f_g=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%
                                      s.n\_top\_slide);
                   MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
                   RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                   fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", adur1, ur1, adut1, ut1);
                   fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", aduru, uru, adutu, utu);
                   s.n_{-}slab = 1.3;
                   s.n_{-}top_{-}slide = 1.5;
                   s.n\_bottom\_slide = 1.5;
                   s.n\_top\_slide);
                   MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
```

```
RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                         fprintf(stderr, "\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}\%5.4f_{"}", adur1, ur1, adut1, ut1);
                         fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", aduru, uru, adutu, utu);
                          s.a = 0.5;
                          s.b = 1.0;
                          s.n_{-}slab = 1.0;
                          s.n\_top\_slide = 1.0;
                          s.n\_bottom\_slide = 1.0;
                         fprintf(stderr, \n=\%5.4f_b=\%5.4f_g=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%
                                                    s.n\_top\_slide);
                         MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
                         RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                         \textit{fprintf} \, (\textit{stderr}, \texttt{"\%5.4f}_{\texttt{L}}\texttt{\%5.4f}_{\texttt{L}}\texttt{\%5.4f}_{\texttt{L}}\texttt{\%5.4f}_{\texttt{L}}\texttt{\%5.4f}_{\texttt{L}}\texttt{ULL}", \textit{adur1}, \textit{ur1}, \textit{adut1}, \textit{ut1});
                         fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", aduru, uru, adutu, utu);
                          s.g = 0.5;
                         fprintf(stderr, "\na=\%5.4f_b=\%5.4f_g=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n
                                                    s.n\_top\_slide);
                         MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
                        RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                         \textit{fprintf} \, (\textit{stderr}, \texttt{"\%5.4f}_{\texttt{L}} \texttt{\%5.4f}_{\texttt{L}} \texttt{\%5.4f}_{\texttt{L}} \texttt{\%5.4f}_{\texttt{L}} \texttt{\%5.4f}_{\texttt{L}} \texttt{ULL} \texttt{"}}, \textit{adur1}, \textit{ur1}, \textit{adut1}, \textit{ut1});
                         fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", aduru, uru, adutu, utu);
                         s.n_{-}slab = 1.5;
                          fprintf(stderr, "\n=\%5.4f_b=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=\%5.4f_n=
                                                    s.n\_top\_slide);
                          MC_RT(s, \&ur1, \&ut1, \&uru, \&utu);
                          RT(32, \&s, \&adur1, \&adut1, \&aduru, \&adutu);
                        fprintf(stderr, "\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}\%5.4f_{\square}", adur1, ur1, adut1, ut1);
                         \mathit{fprintf} \, (\mathit{stderr}, \verb"%5.4f_{\verb"\\"5.4f_{\verb"\\"5.4f_{\verb"\\"5.4f_{\verb"\\"}}} \verb"5.4f_{\verb"\\"1, aduru}, \mathit{uru}, \mathit{adutu}, \mathit{utu});
17. \langle old formatting 17\rangle \equiv
            if (cl\_verbosity > 0 \land count \% 100 \equiv 0) fprintf (stderr, "\n");
            if (cl\_verbosity > 0) printf (format2, m.m\_r, m.m\_t, m.m\_u, r.a, r.b, r.g, r.final\_distance);
             else printf("%9.5f\t%9.5f\t%9.5f\t%9.5f\n", r.a, r.b, r.g, r.final_distance);
```

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18. Stuff the command line arguments that should be constant over the entire inversion process into the measurement record and set up the result record to handle the arguments properly so that the optical properties can be determined.

```
\langle Command-line changes to m 18\rangle \equiv
  if (cl\_cos\_angle \neq \texttt{UNINITIALIZED}) {
     m.slab\_cos\_angle = cl\_cos\_angle;
     if (cl\_quadrature\_points \equiv UNINITIALIZED) cl\_quadrature\_points = 12;
     if (cl\_quadrature\_points \neq 12 * (cl\_quadrature\_points/12)) {
       fprintf (stderr,
             "If_{\cup}you_{\cup}use_{\cup}the_{\cup}-i_{\cup}option_{\cup}to_{\cup}specify_{\cup}an_{\cup}oblique_{\cup}incidence_{\cup}angle,_{\cup}then\n");
       fprintf(stderr, "the \ number \ of \ quadrature \ points \ must \ be \ a \ multiple \ of \ 12\ ");
        exit(0);
  if (cl\_sample\_n \neq UNINITIALIZED) m.slab\_index = cl\_sample\_n;
  if (cl\_slide\_n \neq UNINITIALIZED) {
     m.slab\_bottom\_slide\_index = cl\_slide\_n;
     m.slab\_top\_slide\_index = cl\_slide\_n;
  if (cl\_slide\_OD \neq \mathtt{UNINITIALIZED}) {
     m.slab\_bottom\_slide\_b = cl\_slide\_OD;
     m.slab\_top\_slide\_b = cl\_slide\_OD;
  if (cl\_sample\_d \neq UNINITIALIZED) m.slab\_thickness = cl\_sample\_d;
  if (cl\_beam\_d \neq UNINITIALIZED) m.d\_beam = cl\_beam\_d;
  if (cl\_slide\_d \neq UNINITIALIZED) {
     m.slab\_bottom\_slide\_thickness = cl\_slide\_d;
     m.slab\_top\_slide\_thickness = cl\_slide\_d;
  if (cl\_slides \equiv NO\_SLIDES) {
     m.slab\_bottom\_slide\_index = 1.0;
     m.slab\_bottom\_slide\_thickness = 0.0;
     m.slab\_top\_slide\_index = 1.0;
     m.slab\_top\_slide\_thickness = 0.0;
  if (cl\_slides \equiv \mathtt{ONE\_SLIDE\_ON\_TOP} \lor cl\_slides \equiv \mathtt{ONE\_SLIDE\_NEAR\_SPHERE}) {
     m.slab\_bottom\_slide\_index = 1.0;
     m.slab\_bottom\_slide\_thickness = 0.0;
  if (cl\_slides \equiv \mathtt{ONE\_SLIDE\_ON\_BOTTOM} \lor cl\_slides \equiv \mathtt{ONE\_SLIDE\_NOT\_NEAR\_SPHERE}) {
     m.slab\_top\_slide\_index = 1.0;
     m.slab\_top\_slide\_thickness = 0.0;
  if (cl\_slides \equiv \texttt{ONE\_SLIDE\_NEAR\_SPHERE} \lor cl\_slides \equiv \texttt{ONE\_SLIDE\_NOT\_NEAR\_SPHERE}) m.flip_sample = 1;
  else m.flip\_sample = 0;
  if (cl\_method \neq UNINITIALIZED) m.method = (int) cl\_method;
  if (cl\_rstd\_t \neq UNINITIALIZED) m.rstd\_t = cl\_rstd\_t;
  if (cl_rstd_r \neq UNINITIALIZED) m.rstd_r = cl_rstd_r;
  if (cl\_sphere\_one[4] \neq \texttt{UNINITIALIZED}) {
     double d_sample_r, d_entrance_r, d_detector_r;
     m.d\_sphere\_r = cl\_sphere\_one[0];
     d\_sample\_r = cl\_sphere\_one[1];
```

```
d_{-}entrance_{-}r = cl_{-}sphere_{-}one[2];
  d\_detector\_r = cl\_sphere\_one[3];
  m.rw_r = cl\_sphere\_one[4];
  m.as\_r = (d\_sample\_r/m.d\_sphere\_r/2) * (d\_sample\_r/m.d\_sphere\_r/2);
  m.ae\_r = (d\_entrance\_r/m.d\_sphere\_r/2) * (d\_entrance\_r/m.d\_sphere\_r/2);
  m.ad_r = (d\_detector\_r/m.d\_sphere\_r/2) * (d\_detector\_r/m.d\_sphere\_r/2);
  m.aw_r = 1.0 - m.as_r - m.ae_r - m.ad_r;
  m.d\_sphere\_t = m.d\_sphere\_r;
  m.as_t = m.as_r;
  m.ae_{-}t = m.ae_{-}r;
  m.ad_{-}t = m.ad_{-}r;
  m.aw_{-}t = m.aw_{-}r;
  m.rw_{-}t = m.rw_{-}r;
  if (cl\_num\_spheres \equiv UNINITIALIZED) m.num\_spheres = 1;
if (cl\_sphere\_two[4] \neq \texttt{UNINITIALIZED}) {
  double d_sample_t, d_entrance_t, d_detector_t;
  m.d\_sphere\_t = cl\_sphere\_two[0];
  d\_sample\_t = cl\_sphere\_two[1];
  d_{-}entrance_{-}t = cl_{-}sphere_{-}two[2];
  d\_detector\_t = cl\_sphere\_two[3];
  m.rw_t = cl\_sphere\_two[4];
  m.as_{-}t = (d\_sample\_t/m.d\_sphere\_t/2) * (d\_sample\_t/m.d\_sphere\_t/2);
  m.ae\_t = (d\_entrance\_t/m.d\_sphere\_t/2) * (d\_entrance\_t/m.d\_sphere\_t/2);
  m.ad\_t = (d\_detector\_t/m.d\_sphere\_t/2) * (d\_detector\_t/m.d\_sphere\_t/2);
  m.aw_{-}t = 1.0 - m.as_{-}t - m.ae_{-}t - m.ad_{-}t;
  if (cl\_num\_spheres \equiv UNINITIALIZED) m.num\_spheres = 2;
if (cl\_num\_spheres \neq UNINITIALIZED) {
  m.num\_spheres = (int) cl\_num\_spheres;
  if (m.num\_spheres > 0 \land m.method \equiv \texttt{UNKNOWN}) \ m.method = \texttt{SUBSTITUTION};
if (cl\_rc\_fraction \neq UNINITIALIZED) m.fraction\_of\_rc\_in\_mr = cl\_rc\_fraction;
if (cl\_tc\_fraction \neq UNINITIALIZED) m.fraction\_of\_tc\_in\_mt = cl\_tc\_fraction;
if (cl_{-}UR1 \neq UNINITIALIZED) m.m_{-}r = cl_{-}UR1;
if (cl_{-}UT1 \neq UNINITIALIZED) m.m_{-}t = cl_{-}UT1;
if (cl_{-}Tc \neq UNINITIALIZED) m.m_{-}u = cl_{-}Tc;
if (cl\_default\_fr \neq UNINITIALIZED) m.f\_r = cl\_default\_fr;
```

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put the values for command line reflection and transmission into the measurement record. \langle Count command-line measurements 19 $\rangle \equiv$ $m.num_measures = 3;$ if $(m.m_t \equiv 0)$ $m.num_measures ---;$ if $(m.m_u \equiv 0)$ $m.num_measures ---;$ $params = m.num_measures;$ if $(m.num_measures \equiv 3)$ { /* need to fill slab entries to calculate the optical thickness */ **struct** $AD_slab_type s;$ $s.n_slab = m.slab_index;$ $s.n_top_slide = m.slab_top_slide_index;$ $s.n_bottom_slide = m.slab_bottom_slide_index;$ $s.b_top_slide = m.slab_top_slide_b;$ $s.b_bottom_slide = m.slab_bottom_slide_b;$ $s.cos_angle = m.slab_cos_angle;$ $cl_default_b = What_Is_B(s, m.m_u);$ This code is used in section 2. **20.** $\langle \text{ print version function } 20 \rangle \equiv$ static void print_version(void) $fprintf(stderr, "iad_\%s\n", Version);$ fprintf(stderr, "Copyright_2020_Scott_Prahl,_scott.prahl@oit.edu\n"); $fprintf(stderr, "_{"} (see_Applied_Optics, _32:559-568, _1993) \n\n");$ $fprintf(stderr, "This \sqcup is \sqcup free \sqcup software; \sqcup see \sqcup the \sqcup source \sqcup for \sqcup copying \sqcup conditions. \n");$ fprintf(stderr, "There_is_no_warranty;_not_even_for_MERCHANTABILITY_or_FITNESS.\n"); $fprintf(stderr, "FOR_ A_ PARTICULAR_ PURPOSE. n");$ exit(0);

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17

```
21. \langle \text{ print usage function } 21 \rangle \equiv
       static void print_usage(void)
                fprintf(stderr, "iad_{\sqcup}%s\n\n", Version);
                fprintf(stderr, "iad_lfinds_loptical_lproperties_lfrom_lmeasurements\n\n");
                fprintf(stderr, "Usage: \sqcup iad \sqcup [options] \sqcup input \n');
                fprintf(stderr, "Options:\n");
                fprintf(stderr, "_{\sqcup\sqcup}-1_{\sqcup}, \#_{\sqcup}\#_{\sqcup}\#_{\sqcup}\#_{\sqcup}\#_{\sqcup})reflection_sphere_parameters_\n");
                fprintf(stderr, "_{\sqcup\sqcup}-2_{\sqcup}'\#_{\sqcup}\#_{\sqcup}\#_{\sqcup}\#_{\sqcup}"\#_{\sqcup}"transmission_{\sqcup}sphere_{\sqcup}parameters_{\sqcup}\n");
                fprintf(stderr, " \sqcup \sqcup \sqcup \sqcup \sqcup " sphere \sqcup d, \sqcup sample \sqcup d, \sqcup entrance \sqcup d, \sqcup detector \sqcup d, \sqcup wall \sqcup r ' \n");
                fprintf(stderr, "_{\cup \cup} - a_{\cup} \#_{\cup \cup \cup} use_{\cup} this_{\cup} albedo_{\cup} \n");
                fprintf(stderr, "_{\square\square} - A_{\square} \#_{\square\square \square \square} \text{use}_{\square} \text{this}_{\square} \text{absorption}_{\square} \text{coefficient}_{\square} \setminus n");
                fprintf(stderr, "_{\sqcup\sqcup}-b_{\sqcup}\#_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}use_{\sqcup}this_{\sqcup}optical_{\sqcup}thickness_{\sqcup}\n");
                fprintf(stderr, "_{\square\square} - B_{\square} \#_{\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square\square} beam_{\square} diameter_{\square} \n");
                fprintf(stderr, "_{\sqcup\sqcup} - c_{\sqcup} \#_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} fraction_{\sqcup} of_{\sqcup} unscattered_{\sqcup} refl_{\sqcup} in_{\sqcup} MR \ ";
                fprintf(stderr, "`` C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C " + C 
                fprintf(stderr, "uu-du#uuuuuuuuuuuthicknessuofusampleu\n");
                fprintf(stderr, "UU-eU#UUUUUUUUUUUUuerror_toleranceu(default_0.0001)_u n");
                fprintf(stderr, "UU-EU#UUUUUUUUUUUUUUOpticaludepthu(=mua*D)UforUslides\n");
                fprintf(stderr,
                                "uu-fu#uuuuuuuuuuuallowuaufractionu0.0-1.0uofulightutouhitusphereuwallufirst\n");
                fprintf(stderr, "uu-Fu#uuuuuuuuuuuuuuuseuthisuscatteringucoefficientu\n");
                fprintf(stderr, "_{L|L}-F_{L}'P_{L})ambda0_{L}mus0_{L}gamma'_{L|L|L}mus=mus0*(lambda/lambda0)^gamma'n");
                fprintf(stderr, "_{\sqcup \sqcup} - F_{\sqcup} \cdot R_{\sqcup} - B_{\sqcup} \cdot R_{\sqcup} - R_{\sqcup} R_
                fprintf(stderr, "_{\sqcup\sqcup}-g_{\sqcup}\#_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}scattering_{\sqcup}anisotropy_{\sqcup}(default_{\sqcup}0)_{\sqcup}\n");
                \mathit{fprintf}(\mathit{stderr}, \texttt{"uuuuuuuuuuuuuuu'0'uoru'2'uuuuuuuuuuuuuuuuuu---unumberuofuslides \n"});
               fprintf(stderr, "LULLULULULULULULULULUL't', (top)LorL'b', (bottom)L---LoneLslideL\
                                that is hit by light first \n");
               \mathit{fprintf}\,(\mathit{stderr}, \texttt{"}_{\verb"UUUUUUUUUUUUUUU'}, \texttt{n}, \texttt{`}_{\verb"U}(\texttt{near})_{\verb"UOT"}, \texttt{f}, \texttt{`}_{\verb"U}(\texttt{far})_{\verb"UUU} --- \texttt{UONE}_{\verb"Slide} \setminus \texttt{far})
                               position_relative_to_sphere\n");
                fprintf(stderr, "_{"} - h_{"} - h_{"});
                fprintf(stderr, "\_\_-i\_\#_\bot\_\_\_\_\_light\_is\_incident\_\_at\_\_this\_\_angle\_\_in\_\_degrees n");
                fprintf(stderr, "_{UU}-M_{U}\#_{UUUUUUUUUUUUUUUUUunumber_{U}} of_{U}Monte_{U}Carlo_{U}iterations \n");
                fprintf(stderr, "uu-nu#uuuuuuuuuuuuuspecifyuindexuofurefractionuofuslab\n");
                fprintf(stderr, "$_{$\sqcup\sqcup}$-N$_{$\sqcup\#}$_{$\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}$pecify$_{$\sqcup$index}_{$\sqcup$}$of$_{$\sqcup$refraction}_{$\sqcup$}of$_{$\sqcup$slides}");
                fprintf(stderr, "_{\sqcup\sqcup} - o_{\sqcup}filename_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} explicitly_specify_filename_{\sqcup}for_{\sqcup}output \");
                fprintf(stderr, "_{$\sqcup\sqcup} - p_{\sqcup}\#_{$\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}\#_{\sqcup}\#_{\sqcup}f_{\sqcup}Monte_{\sqcup}Carlo_{\sqcup}photons_{\sqcup}(default_{\sqcup}100000) \n");
                fprintf(stderr, "_{\Box\Box} - q_{\Box} \#_{\Box\Box\Box\Box\Box\Box\Box\Box\Box\Box\Box\Box\Box} number_{\Box} of_{\Box} quadrature_{\Box} points_{\Box} (default=8) n");
                fprintf(stderr, "_{"} - r_{"} + _{"} + _{"} - r_{"} + _{"} + _{"} - r_{"} + _{"} + _{"} - r_{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _{"} + _
                fprintf(stderr, "_{$\sqcup\sqcup} - R_{\sqcup} \#_{$\sqcup\sqcup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup} = \text{actual}_{$\sqcup$ reflectance}_{$\sqcup$} for_{$\sqcup} 100\%_{$\sqcup} measurement_{$\sqcup} \setminus n");
                fprintf(stderr, "_{\sqcup\sqcup} - S_{\sqcup} \#_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} number_{\sqcup} of_{\sqcup} spheres_{\sqcup} used \");
                fprintf(stderr, "_{$\sqcup\sqcup$}-t_{$\sqcup$}\#_{$\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}total_{$\sqcup$}transmission_{$\sqcup$}measurement\n");
                fprintf(stderr, "_{UU}-T_{U}+_{UUUUUUUUUUUUUUu} actual_{U} transmission_{U} for_{U} 100%_{U} measurement_{U} n");
                fprintf(stderr, "_{$\sqcup\sqcup} - u_{$\sqcup} \#_{$\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} unscattered_{$\sqcup$} transmission_{$\sqcup$} measurement \n");
                fprintf(stderr, "_{\sqcup\sqcup} - v_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} version_{\sqcup}information \");
                fprintf(stderr, "_{\square\square} - V_{\square}O_{\square\square\square\square\square\square\square\square\square\square\square} verbosity_{\square}low_{\square} - --_{\square}no_{\square}output_{\square}to_{\square}stderr n");
                fprintf(stderr, "_{\sqcup\sqcup} - V_{\sqcup} 2_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} verbosity_{\sqcup}high\n");
                fprintf(stderr, "_{\square\square} - x_{\square} \#_{\square\square\square\square\square\square\square\square\square\square\square\square\square\square} set_debugging_level\n");
```

18 Main Program iad (v 3-12-0) §21

```
fprintf(stderr, "_{$\sqcup\sqcup$}-X_{$\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}dual_{$\sqcup$}beam_{$\sqcup$}configuration\n");
     fprintf(stderr, "_{\sqcup\sqcup}-z_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}do_{\sqcup}forward_{\sqcup}calculation\n");
     fprintf(stderr, "Examples: \n");
     \mathit{fprintf} \, (\mathit{stderr}, \verb""") \verb"iad" file. \verb"rxt" \verb| uuuuuuuuuuuuu Results" will" \verb| be" put" \verb| in" file. \verb| txt \verb| n"");
     fprintf(stderr, "ulliadufileuluuuuuuuuuuuuuuuuuuuuuuusameuasuabove\n");
     fprintf(stderr, "``uuiad``u-c``u0.9``ufile.rxt``uuuuuuuAssume``uM_R``uincludes``u90\\\`uof``uuns\
          cattered reflectance \n");
     fprintf(stderr, "lliad_l-C_l0.8_lfile.rxt_lllllllllllllllllAssume_lM_T_lincludes_l80\%_lof_luns
          cattered<sub>□</sub>transmittance\n");
     fprintf(stderr, "\_u\_iad_u-e_u0.0001_ufile.rxt_uuuu\_Better_uconvergence_uto_uR_u&_uT_uvalues n");
     fprintf(stderr,
           "ULI iadu-ful. Oufile.rxtullullullAllulightuhitsureflectanceusphereuwallufirst\n");
     fprintf(stderr, "uliadu-oloutufile.rxtuluuuuuuCalculateduvaluesuinlout\n");
     fprintf(stderr, "lociad_l-r_l0.3_lociolociolociolocioloci_R_total=0.3, lobeinf, lofind_labedo\n");
     fprintf(stderr, ""uliadu-ru0.3u-tu0.4ululululululuR_total=0.3, uT_total=0.4, ufindua,b,g\n");
     fprintf(stderr, "\_\sqcup iad_\sqcup -r_\sqcup 0.3_\sqcup -t_\sqcup 0.4_\sqcup -n_\sqcup 1.5_\sqcup \sqcup R\_total = 0.3, \sqcup T\_total = 0.4, \sqcup n = 1.5, \sqcup find_\sqcup a, b n ");
     fprintf(stderr, "uuiadu-pu1000ufile.rxtuuuuuu00nlyu1000uphotons\n");
     fprintf(stderr, "uuiadu-pu-100ufile.rxtuuuuuuAllowuonlyu100msuperuiteration\n");
     fprintf(stderr, "_{\sqcup\sqcup}iad_{\sqcup}-q_{\sqcup}4_{\sqcup}file.rxt_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}Four_{\sqcup}quadrature_{\sqcup}points \n");
     fprintf(stderr, "uuiadu-MuOufile.rxtuuuuuuuuNouMCuuuu(iad)\n");
     fprintf(stderr, "uuiadu-Mu2ufile.rxtuuuuuuuuMCutwiceu(iadu->uMCu->uiadu->uMCu->uiad) \n");
     fprintf(stderr, "uuiadu-MuOu-quUufile.rxtuuuuuFastuanducrudeuconversion\n");
     fprintf (stderr,
           "uuiadu-Gutufile.rxtuuuuuuuu0neutopuslideuwithupropertiesufromufile.rxt\n");
     fprintf (stderr,
          "_{\sqcup\sqcup}iad_{\sqcup}-G_{\sqcup}b_{\sqcup}-N_{\sqcup}1.5_{\sqcup}-D_{\sqcup}1_{\sqcup}file_{\sqcup}Use_{\sqcup}1_{\sqcup}bottom_{\sqcup}slide_{\sqcup}with_{\sqcup}n=1.5_{\sqcup}and_{\sqcup}thickness=1\n");
     fprintf(stderr, "LLiad_l-x_LLL_Lfile.rxt_LLLLLLLLLLLLShow_lsphere_land_LMC_leffects\n");
     fprintf(stderr, "uuiadu-xuuu2ufile.rxtuuuuuuDEBUG_GRID\n");
     fprintf(stderr, "\_\_iad\_-x_\_\_4\_file.rxt_\_\_\_DEBUG\_ITERATIONS\n");
     fprintf(stderr, "uliad_-xulu_8 file.rxt_uuuuuuDEBUG_LOST_LIGHT\n");
     fprintf(stderr, "uuiadu-xuu16ufile.rxtuuuuuuDEBUG_SPHERE_EFFECTS\n");
     fprintf(stderr, "uuiadu-xuu32ufile.rxtuuuuuuDEBUG_BEST_GUESS\n");
     fprintf(stderr, "uuiadu-xuu64ufile.rxtuuuuuuDEBUG_EVERY_CALC\n");
     fprintf(stderr, "ulliad_l-x_l128_lfile.rxt_lullull_DEBUG_SEARCH\n");
     fprintf(stderr, "uuiadu-xu255ufile.rxtuuuuuuAlludebugginguoutput\n");
     fprintf(stderr,
          "_{\sqcup\sqcup} iad_{\sqcup} - X_{\sqcup} - i_{\sqcup} 8_{\sqcup} file.rxt_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} Dual_{\sqcup} beam_{\sqcup} spectrometer_{\sqcup} with_{\sqcup} 8_{\sqcup} degree_{\sqcup} incidence \n \n");
     fprintf(stderr,
           "_{\cup\cup}iad_{\cup}-z_{\cup}-a_{\cup}0.9_{\cup}-b_{\cup}1_{\cup}-i_{\cup}45_{\cup\cup}Forward_{\cup}calc_{\cup}assuming_{\cup}45_{\cup}degree_{\cup}incidence_{\setminus}n_{\setminus}n");
     fprintf(stderr, "`` Li Liapply Liad Lix.rxt Liy.rxt Li Li Li Li Li Li Process Limultiple Lifiles \n' ");
     fprintf(stderr, "Report_bugs_to_scott.prahl@oit.edu>\n\n");
     exit(0);
This code is used in section 2.
```

 $\S22$ IAD (v 3-12-0) MAIN PROGRAM 19

Just figure out the damn scattering and absorption \langle calculate coefficients function $22 \rangle \equiv$ static void $Calculate_Mua_Musp$ (struct $measure_type m$, struct $invert_type r$, double *musp, double if $(r.b \equiv \text{HUGE_VAL})$ { if $(r.a \equiv 0)$ { *musp = 0.0;*mua = 1.0;return; if $(r.default_mus \neq UNINITIALIZED)$ { $*musp = r.default_mus * (1 - r.g);$ $*mua = r.default_mus/r.a - r.default_mus;$ return; if $(r.default_mua \neq UNINITIALIZED)$ { $*musp = (r.default_mua/(1-r.a) - r.default_mua) * (1-r.g);$ $*mua = r.default_mua;$ return; *musp = 1.0 - r.g;*mua = (1.0 - r.a)/r.a;return; $*musp = r.a * r.b/m.slab_thickness * (1.0 - r.g);$ $*mua = (1 - r.a) * r.b/m.slab_thickness;$ See also section 23. This code is used in section 2. This can only be called immediately after *Invert_RT* You have been warned! Notice that *Calculate_Distance* does not pass any slab properties. \langle calculate coefficients function $22 \rangle + \equiv$ static void $calculate_coefficients$ (struct $measure_type$ m, struct $invert_type$ r, double *LR, double *LT, double *musp, double *mua) double delta; *LR = 0;*LT = 0;Calculate_Distance(LR, LT, & delta); $Calculate_Mua_Musp(m, r, musp, mua);$ }

20 Main Program iad (v 3-12-0) §24

```
\langle \text{ print results header function } 24 \rangle \equiv
         static void print_results_header(FILE *fp)
                   fprintf(fp, "\#_{UUUU} \land tMeasured_{U} \land t_{UUU} \land tMeasured_{U} \land t_{UUU} \land tEstimat \land t_{UUU} \land tMeasured_{U} \land t_{UUU} \land tEstimat \land t_{UUU} \land tMeasured_{U} \land tMeasured_{U
                                       ed\tEstimated\tEstimated");
                   if (Debug(DEBUG\_LOST\_LIGHT)) fprintf(fp,
                                                 "\tuuLostuuu\tuuLostuuu\tuuLostuuu\tuuLostuuu\tuuuMCuuuu\tuuuIADuuu\tuuErroruu");
                   fprintf(fp, "\n");
                  \mathit{fprintf} (\mathit{fp}, \texttt{"##wave} \land \texttt{tuuu} \texttt{M\_R} \texttt{uuu} \land \texttt{tuuu} \texttt{fit} \texttt{uuu} \land \texttt{tuuu} \texttt{M\_T} \texttt{uuu} \land \texttt{tuuu} \texttt{fit} \texttt{uuu} \land \texttt{tuuu} \texttt{mu\_au} \land \texttt{mu\_
                                       uu\tuumu_s'uu\tuuuuguuuu");
                   if (Debug(DEBUG_LOST_LIGHT)) fprintf(fp,
                                                 "\toooUR1ooo\toooURUooo\toooUT1ooo\toooUTUooo\toooo#oooo\toooo#oooo\tooStateoo");
                  fprintf(fp, "\n");
                  uu\tuu1/mmuuu\tuu[---]uu");
                   if (Debug(DEBUG_LOST_LIGHT)) fprintf(fp,
                                                 fprintf(fp, "\n");
This code is used in section 2.
25. When debugging lost light, it is handy to see how each iteration changes the calculated values for the
optical properties. We do that here if we are debugging, otherwise we just print a number or something to
keep the user from wondering what is going on.
\langle \text{ Print results function } 25 \rangle \equiv
          void print_optical_property_result (FILE *fp, struct measure_type m, struct invert_type r, double
                            LR, double LT, double mu_-a, double mu_-sp, int mc_-iter, int line ) {
          if (m.lambda \neq 0) fprintf (fp, "\%6.1f\t", m.lambda);
          else fprintf (fp, "\%6d\t", line);
          if (mu_a > 200) mu_a = 199.9999;
         if (mu\_sp > 1000) mu\_sp = 999.9999;
          fprintf(fp, "\% 9.4f \t\% 9.4f \t", m.m_r, LR);
          fprintf(fp, "\% 9.4f \t\% 9.4f \t", m.m_t, LT);
         fprintf(fp, "\% 9.4f\t", mu_a);
         fprintf(fp, "%_{\sqcup}9.4f\t", mu\_sp);
          fprintf(fp, "%\_9.4f\t", r.g);
         if (Debug(DEBUG_LOST_LIGHT)) {
                   fprintf(fp, "\% 9.4f \t\% 9.4f \t", m.ur1\_lost, m.uru\_lost);
                   fprintf(fp, "\% \cup 9.4f \t\% \cup 9.4f \t", m.ut1\_lost, m.utu\_lost);
                  \mathit{fprintf}\left(\mathit{fp}\,,\,\text{``} \sqcup \text{\ensuremath{\mbox{$\backslash$}}} 2d_{\sqcup \sqcup} \text{\ensuremath{\mbox{$\backslash$}}} \,,\, mc\_iter \right);
                  fprintf(fp, " \ \%4d \ ", r.iterations);
          fprintf(fp, "#_{\perp}\%c_{\perp}\n", what\_char(r.error));
         fflush(fp); \}
```

21

```
26.
        \langle \text{ print error legend function } 26 \rangle \equiv
   static void print_error_legend(void)
      fprintf(stderr, "_{"}==>_{"}Success_{"}");
      fprintf(stderr, "_{\sqcup\sqcup}0-9_{\sqcup}==>_{\sqcup}Monte_{\sqcup}Carlo_{\sqcup}Iteration\n");
      fprintf(stderr, "_{\sqcup\sqcup\sqcup}R_{\sqcup\sqcup}==>_{\sqcup}M_R_{\sqcup}is_{\sqcup}too_{\sqcup}big_{\sqcup\sqcup\sqcup}");
      fprintf(stderr, "_{\sqcup\sqcup\sqcup}r_{\sqcup\sqcup}==>_{\sqcup}M_R_{\sqcup}is_{\sqcup}too_{\sqcup}small\n");
      fprintf(stderr, "_{\sqcup\sqcup\sqcup}T_{\sqcup\sqcup}==>_{\sqcup}M_{\_}T_{\sqcup}is_{\sqcup}too_{\sqcup}big_{\sqcup\sqcup\sqcup}");
      fprintf(stderr, "_{\sqcup\sqcup\sqcup}t_{\sqcup\sqcup}==>_{\sqcup}M_{T_{\sqcup}}is_{\sqcup}too_{\sqcup}small\n");
      fprintf(stderr, "_{\sqcup\sqcup\sqcup}U_{\sqcup\sqcup}==>_{\sqcup}M_{\_}U_{\sqcup}is_{\sqcup}too_{\sqcup}big_{\sqcup\sqcup\sqcup}");
      fprintf(stderr, "_{\sqcup\sqcup\sqcup}u_{\sqcup\sqcup}==>_{\sqcup}M_{U_{\sqcup}}is_{\sqcup}too_{\sqcup}small\n");
      fprintf(stderr, "_{ \cup \cup \cup }!_{ \cup \cup }==>_{ \cup }M_R_{ \cup }+_{ \cup }M_T_{ \cup }>_{ \cup }1_{ \cup \cup \cup \cup }");
      fprintf(stderr, "_{\sqcup\sqcup\sqcup}+_{\sqcup\sqcup}==>_{\sqcup}Did_{\sqcup}not_{\sqcup}converge\\n\\n");
This code is used in section 2.
27. returns a new string consisting of s+t
\langle stringdup together function 27 \rangle \equiv
   static char *strdup\_together(\mathbf{char} *s, \mathbf{char} *t)
   {
      char *both;
      if (s \equiv \Lambda) {
         if (t \equiv \Lambda) return \Lambda;
         return strdup(t);
      if (t \equiv \Lambda) return strdup(s);
      both = malloc(strlen(s) + strlen(t) + 1);
      if (both \equiv \Lambda) fprintf(stderr, "Could_not_allocate_memory_for_both_strings.\n");
      strcpy(both, s);
      strcat(both, t);
      return both;
This code is used in section 2.
28. assume that start time has already been set
\langle seconds elapsed function 28\rangle \equiv
   static double seconds_elapsed(clock_t start_time)
      \mathbf{clock\_t}\ finish\_time = clock();
      return (double)(finish_time - start_time)/CLOCKS_PER_SEC;
This code is used in section 2.
```

22 Main Program iad (v 3-12-0) §29

29. given a string and an array, this fills the array with numbers from the string. The numbers should be separated by spaces.

Returns 0 upon successfully filling n entries, returns 1 for any error.

```
\langle parse string into array function 29 \rangle \equiv
  static int parse\_string\_into\_array(char *s, double *a, int n)
     \mathbf{char} \ *t, \ *last, \ *r;
     int i = 0;
     t = s;
     last = s + strlen(s);
     while (t < last) {
                               /* a space should mark the end of number */
       r = t;
       while (*r \neq ' \cup ' \land *r \neq ' \lor 0') r \leftrightarrow ;
       *r = '\0'; /* parse the number and save it */
       if (sscanf(t, "% if", \&(a[i])) \equiv 0) return 1;
       i++; /* are we done? */
       if (i \equiv n) return 0; /* move pointer just after last number */
       t = r + 1;
     return 1;
  }
```

```
30.
      \langle \text{ print dot function } 30 \rangle \equiv
  static char what_char(int err)
     if (err \equiv IAD\_NO\_ERROR) return '*';
    if (err = IAD_TOO_MANY_ITERATIONS) return '+';
    if (err \equiv IAD\_MR\_TOO\_BIG) return 'R';
    if (err \equiv IAD\_MR\_TOO\_SMALL) return 'r';
     if (err \equiv IAD\_MT\_TOO\_BIG) return 'T';
     if (err \equiv IAD\_MT\_TOO\_SMALL) return 't';
     if (err \equiv IAD\_MU\_TOO\_BIG) return 'U';
    if (err \equiv IAD\_MU\_TOO\_SMALL) return 'u';
     if (err \equiv IAD\_TOO\_MUCH\_LIGHT) return '!';
     return '?';
  static void print_dot(clock_t start_time, int err, int count, int points, int final, int verbosity, int
            *any\_error)
     static int counter = 0;
     counter ++;
    if (err \neq IAD\_NO\_ERROR) * any\_error = err;
    if (verbosity \equiv 0) return;
     if (final \equiv 99) fprintf(stderr, "%c", what_char(err));
     else {
       counter --;
       fprintf(stderr, "%1d\b", final \% 10);
     if (final \equiv 99) {
       if (counter \% 50 \equiv 0) {
          double rate = (seconds\_elapsed(start\_time)/points);
          fprintf(stderr, "_{\sqcup \sqcup} \%3d_{\sqcup}done_{\sqcup} (\%5.2f_{\sqcup}s/pt) \n", points, rate);
       else if (counter \% 10 \equiv 0) fprintf(stderr, "");
     fflush(stderr);
This code is used in section 2.
```

24 IAD TYPES IAD (v $_{3-12-0}$) §31

31. IAD Types. This file has no routines. It is responsible for creating the header file iad_type.h and nothing else. Altered 3/3/95 to change the version number below. Change June 95 to improve cross referencing using CTwill. Change August 97 to add root finding with known absorption

32. These are the various optical properties that can be found with this program. FIND_AUTO allows one to let the computer figure out what it should be looking for.

These determine what metric is used in the minimization process.

These give the two different types of illumination allowed.

Finally, for convenience I create a Boolean type.

```
\langle iad\_type.h \quad 32 \rangle \equiv
#undef FALSE
#undef TRUE
  ⟨ Preprocessor definitions ⟩
  ⟨Structs to export from IAD Types 35⟩
33.
#define FIND_A 0
#define FIND_B 1
#define FIND_AB 2
#define FIND_AG 3
#define FIND_AUTO 4
\#define FIND_BG 5
#define FIND_BaG 6
#define FIND_BsG 7
#define FIND_Ba 8
#define FIND_Bs 9
#define FIND_G 10
#define FIND_B_WITH_NO_ABSORPTION 11
#define FIND_B_WITH_NO_SCATTERING 12
\#define RELATIVE 0
#define ABSOLUTE 1
\#define COLLIMATED 0
\#define DIFFUSE 1
\#define FALSE 0
#define TRUE 1
#define IAD_MAX_ITERATIONS 500
```

 $\S34$ IAD (v 3-12-0) IAD TYPES 25

34. Need error codes for this silly program

```
#define IAD_NO_ERROR 0
#define IAD_TOO_MANY_ITERATIONS 1
#define IAD_AS_NOT_VALID 16
#define IAD_AE_NOT_VALID 17
#define IAD_AD_NOT_VALID 18
#define IAD_RW_NOT_VALID 19
#define IAD_RD_NOT_VALID 20
#define IAD_RSTD_NOT_VALID 21
#define IAD_GAMMA_NOT_VALID 22
#define IAD_F_NOT_VALID 23
#define IAD_BAD_PHASE_FUNCTION 24
#define IAD_QUAD_PTS_NOT_VALID 25
#define IAD_BAD_G_VALUE 26
#define IAD_TOO_MANY_LAYERS 27
#define IAD_MEMORY_ERROR 28
#define IAD_FILE_ERROR 29
#define IAD_EXCESSIVE_LIGHT_LOSS 30
#define IAD_RT_LT_MINIMUM 31
#define IAD_MR_TOO_SMALL 32
#define IAD_MR_TOO_BIG 33
#define IAD_MT_TOO_SMALL 34
#define IAD_MT_TOO_BIG 35
#define IAD_MU_TOO_SMALL 36
#define IAD_MU_TOO_BIG 37
#define IAD_TOO_MUCH_LIGHT 38
#define IAD_TSTD_NOT_VALID 39
#define UNINITIALIZED -99
#define DEBUG_A_LITTLE 1
#define DEBUG_GRID 2
#define DEBUG_ITERATIONS 4
#define DEBUG_LOST_LIGHT 8
#define DEBUG_SPHERE_EFFECTS 16
#define DEBUG_BEST_GUESS 32
#define DEBUG_EVERY_CALC 64
#define DEBUG_SEARCH 128
#define DEBUG_RD_ONLY 256
#define DEBUG_GRID_CALC 512
#define DEBUG ANY #FFFFFFF
\#define UNKNOWN 0
#define COMPARISON 1
\#define SUBSTITUTION 2
```

26 IAD TYPES IAD (v $_3$ -12-0) §35

35. The idea of the structure *measure_type* is collect all the information regarding a single measurement together in one spot. No information regarding how the inversion procedure is supposed to be done is contained in this structure, unlike in previous incarnations of this program.

```
\langle Structs to export from IAD Types 35\rangle \equiv
  typedef struct measure_type {
    double slab\_index;
    double slab_thickness;
    double slab_top_slide_index;
    double slab_top_slide_b;
    double slab_top_slide_thickness;
    double slab_bottom_slide_index;
    double slab_bottom_slide_b;
    {\bf double}\ slab\_bottom\_slide\_thickness;
    double slab\_cos\_angle;
    int num_spheres;
    int num_measures;
    int method;
    int flip_sample;
    double d\_beam;
    double fraction_of_rc_in_mr;
    double fraction_of_tc_in_mt;
    double m_{-}r, m_{-}t, m_{-}u;
    double lambda;
    double as_r, ad_r, ae_r, aw_r, rd_r, rw_r, rstd_r, f_r;
    double as_t, ad_t, ae_t, aw_t, rd_t, rw_t, rstd_t, f_t;
    double ur1_lost, uru_lost, ut1_lost, utu_lost;
    double d\_sphere\_r, d\_sphere\_t;
  } IAD_measure_type;
See also sections 36 and 37.
This code is used in section 32.
```

§36 IAD (v 3-12-0) IAD TYPES 27

36. This describes how the inversion process should proceed and also contains the results of that inversion process.

```
\langle Structs to export from IAD Types 35\rangle + \equiv
  typedef struct invert_type { double a;
                                                 /* the calculated albedo */
                /* the calculated optical depth */
  double b;
  double q;
                /* the calculated anisotropy */
  int found;
  int search;
  int metric;
  double tolerance;
  double MC_tolerance;
  double final_distance;
  int iterations; int error ;
  struct AD\_slab\_type slab;
  struct AD_method_type method;
  double default_a;
  double default_b;
  double default_g;
  double default_ba;
  double default_bs;
  double default_mua;
  double default_mus; } IAD_invert_type;
37. A few types that used to be enum's are now int's.
\langle Structs to export from IAD Types 35\rangle + \equiv
  typedef int search_type;
  typedef int boolean_type;
  typedef int illumination_type;
  typedef struct guess_t {
    double distance;
    double a;
    double b;
    double g;
  } guess_type;
  extern double FRACTION;
```

28 IAD PUBLIC IAD (v 3-12-0) §38

38. IAD Public.

This contains the routine $Inverse_RT$ that should generally be the basic entry point into this whole mess. Call this routine with the proper values and true happiness is bound to be yours.

Altered accuracy of the standard method of root finding from 0.001 to 0.00001. Note, it really doesn't help to change the method from ABSOLUTE to RELATIVE, but I did anyway. (3/3/95)

```
\langle iad_pub.c 38 \rangle \equiv
#include <stdio.h>
#include <math.h>
#include "nr_util.h"
#include "ad_globl.h"
#include "ad_frsnl.h"
#include "iad_type.h"
#include "iad_util.h"
#include "iad_calc.h"
#include "iad_find.h"
#include "iad_pub.h"
#include "iad_io.h"
#include "mc_lost.h"
  \langle \text{ Definition for } Inverse\_RT | 42 \rangle
   \langle \text{ Definition for } measure\_OK \mid 47 \rangle
   (Definition for determine_search 54)
   Definition for Initialize_Result 58
   Definition for Initialize_Measure 66
   \langle \text{ Definition for } ez\_Inverse\_RT | 64 \rangle
   (Definition for Spheres_Inverse_RT 68)
   (Definition for Spheres_Inverse_RT2 81)
   \langle \text{ Definition for } Calculate\_MR\_MT | 75 \rangle
   \langle \text{ Definition for } MinMax\_MR\_MT 79 \rangle
  \langle Definition for Calculate\_Minimum\_MR 77 \rangle
```

39. All the information that needs to be written to the header file iad_pub.h. This eliminates the need to maintain a set of header files as well.

```
 \begin{split} \langle \operatorname{iad\_pub.h} & 39 \rangle \equiv \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Inverse\_RT} \text{ } 41 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{measure\_OK} \text{ } 46 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{determine\_search} \text{ } 53 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Initialize\_Result} \text{ } 57 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Initialize\_Measure} \text{ } 63 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Initialize\_Measure} \text{ } 65 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Calculate\_MR\_MT} \text{ } 74 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Calculate\_Minimum\_MR} \text{ } 76 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Calculate\_Minimum\_MR} \text{ } 76 \rangle; \\ \langle \operatorname{Prototype} \text{ for } \operatorname{Spheres\_Inverse\_RT2} \text{ } 80 \rangle; \end{split}
```

40. Here is the header file needed to access one interesting routine in the libiad.so library.

```
⟨lib_iad.h 40⟩ ≡
  ⟨Prototype for ez_Inverse_RT 63⟩;
  ⟨Prototype for Spheres_Inverse_RT 67⟩;
  ⟨Prototype for Spheres_Inverse_RT2 80⟩;
```

 $\S41$ IAD (v 3-12-0) INVERSE RT 29

41. Inverse RT. $Inverse_RT$ is the main function in this whole package. You pass the variable m containing your experimentally measured values to the function $Inverse_RT$. It hopefully returns the optical properties in r that are appropriate for your experiment.

```
\langle Prototype for Inverse\_RT 41 \rangle \equiv
   void Inverse_RT(struct measure_type m, struct invert_type *r)
This code is used in sections 39 and 42.
42. \langle Definition for Inverse\_RT | 42 \rangle \equiv
   \langle Prototype for Inverse\_RT 41 \rangle
      if (0 \land Debug(DEBUG\_LOST\_LIGHT)) {
        fprintf(stderr, "** \bot Inverse_RT_{\sqcup}(%d_{\sqcup}spheres)_{\sqcup} ** \land ", m.num\_spheres);
        fprintf(stderr, "_{UUUUU}=_U%8.5f,_UMT_{UUUUUU}=_U%8.5f\\n", m.m_r, m.m_t);
        fprintf(stderr, "$\sqcup \sqcup \sqcup \sqcup UR1 \sqcup lost \sqcup = \sqcup \%8.5f, \sqcup UT1 \sqcup lost \sqcup = \sqcup \%8.5f \ ", m.ur1\_lost, m.ur1\_lost);
      r \rightarrow found = FALSE;
      if (r \rightarrow search \equiv FIND\_AUTO) r \rightarrow search = determine\_search(m, *r);
      if (r \rightarrow search \equiv FIND_B_WITH_NO_ABSORPTION) {
        r \rightarrow default_a = 1;
         r \rightarrow search = FIND_B;
      if (r \rightarrow search \equiv FIND_B_WITH_NO_SCATTERING) {
        r \rightarrow default_a = 0;
         r \rightarrow search = FIND_B;
      (Exit with bad input data 43)
      (Find the optical properties 44)
      if (r \neg final\_distance \leq r \neg tolerance) r \neg found = TRUE;
This code is used in section 38.
```

43. There is no sense going to all the trouble to try a multivariable minimization if the input data is bogus. So I wrote a single routine $measure_OK$ to do just this.

```
\langle Exit with bad input data 43\rangle \equiv r \rightarrow \mathbf{error} = measure\_OK(m,*r); if (r\rightarrow method.quad\_pts < 4) r \rightarrow \mathbf{error} = \mathtt{IAD\_QUAD\_PTS\_NOT\_VALID}; if (r \rightarrow \mathbf{error} \neq \mathtt{IAD\_NO\_ERROR}) return; This code is used in section 42.
```

30 INVERSE RT IAD (v 3-12-0) §44

44. Now I fob the real work off to the unconstrained minimization routines. Ultimately, I would like to replace all these by constrained minimization routines. Actually the first five already are constrained. The real work will be improving the last five because these are 2-D minimization routines.

```
\langle Find the optical properties 44 \rangle \equiv
  switch (r \rightarrow search) {
  case FIND_A: U_{-}Find_{-}A(m,r);
     break;
  case FIND_B: U_Find_B(m,r);
     break;
  case FIND_G: U_Find_G(m,r);
     break;
  case FIND\_Ba: U\_Find\_Ba(m,r);
     break;
  case FIND\_Bs: U\_Find\_Bs(m,r);
     break;
  case FIND_AB: U_Find_AB(m,r);
     break;
  case FIND_AG: U_Find_AG(m,r);
     break;
  case FIND_BG: U_Find_BG(m,r);
     break;
  case FIND\_BsG: U\_Find\_BsG(m,r);
  case FIND\_BaG: U\_Find\_BaG(m,r);
     break;
  if (r\text{-}iterations \equiv \texttt{IAD\_MAX\_ITERATIONS}) r \text{-} \mathbf{error} = \texttt{IAD\_TOO\_MANY\_ITERATIONS};
This code is used in section 42.
      Validation.
45.
      Now the question is — just what is bad data? Here's the prototype.
\langle \text{ Prototype for } measure\_OK \mid 46 \rangle \equiv
  int measure_OK(struct measure_type m, struct invert_type r)
This code is used in sections 39 and 47.
```

 $\S47$ IAD (v 3-12-0) VALIDATION 31

47. It would just be nice to stop computing with bad data. This does not work in practice because it turns out that there is often bogus data in a full wavelength scan. Often the reflectance is too low for short wavelengths and at long wavelengths the detector (photomultiplier tube) does not work worth a damn.

The two sphere checks are more complicated. For example, we can no longer categorically state that the transmittance is less than one or that the sum of the reflectance and transmittance is less than one. Instead we use the transmittance to bound the values for the reflectance — see the routine $MinMax_MR_MT$ below.

```
 \begin{tabular}{ll} $\langle \operatorname{Definition} \ & \operatorname{measure\_OK} \ 46 \end{tabular} = \\ & \langle \operatorname{Prototype} \ & \operatorname{for} \ & \operatorname{measure\_OK} \ 46 \end{tabular} = \\ & \langle \operatorname{Check} \ & \operatorname{MT} \ & \operatorname{for} \ & \operatorname{zero} \ & \operatorname{or} \ & \operatorname{one} \ & \operatorname{spheres} \ & 49 \end{tabular} \\ & \langle \operatorname{Check} \ & \operatorname{MR} \ & \operatorname{for} \ & \operatorname{zero} \ & \operatorname{or} \ & \operatorname{one} \ & \operatorname{spheres} \ & 48 \end{tabular} \\ & \rbrace \\ & \text{else} \ & \{ \ & \operatorname{int} \ & \operatorname{error} \ = \ & \operatorname{MinMax\_MR\_MT}(m,r); \ & \operatorname{if} \ & ( \ & \operatorname{error} \ \neq \ & \operatorname{IAD\_NO\_ERROR} \ ) \ & \operatorname{return} \ & \operatorname{error} \ ; \ \end{tabular} \ & \langle \operatorname{Check} \ & \operatorname{MU} \ & \operatorname{50} \end{tabular} \\ & \text{if} \ & (m.num\_spheres \ \neq 0) \ \ & \langle \operatorname{Check} \ & \operatorname{sphere} \ & \operatorname{parameters} \ & 51 \end{tabular} \\ & \rbrace \\ & \text{return} \ & \operatorname{IAD\_NO\_ERROR}; \ \end{tabular}
```

48. The reflectance is constrained by the index of refraction of the material and the transmission. The upper bound for the reflectance is just one minus the transmittance. The specular (unscattered) reflectance from the boundaries imposes minimum for the reflectance. Obviously, the reflected light cannot be less than that from the first boundary. This might be calculated by assuming an infinite layer thickness. But we can do better.

There is a definite bound on the minimum reflectance from a sample. If you have a sample with a given transmittance m_-t , the minimum reflectance possible is found by assuming that the sample does not scatter any light.

Knowledge of the indicies of refraction makes it a relatively simple matter to determine the optical thickness $b = mu_-a * d$ of the slab. The minimum reflection is obtained by including all the specular reflectances from all the surfaces.

If the default albedo has been specified as zero, then there is really no need to check MR because it is ignored.

32 Validation iad (v 3-12-0) §49

49. The transmittance is also constrained by the index of refraction of the material. The minimum transmittance is zero, but the maximum transmittance cannot exceed the total light passing through the sample when there is no scattering or absorption. This is calculated by assuming an infinitely thin (to eliminate any scattering or absorption effects).

There is a problem when spheres are present. The estimated values for the transmittance using Sp_mu_RT are not actually limiting cases. This will require a bit of fixing, but for now that test is omitted if the number of spheres is more than zero.

```
\langle Check MT for zero or one spheres 49 \rangle \equiv
        if (m.m_t < 0) return IAD_MT_TOO_SMALL;
        Sp\_mu\_RT\_Flip(m.flip\_sample, r.slab.n\_top\_slide, r.slab.n\_slab, r.slab.n\_bottom\_slide, r.slab.b\_top\_slide, 0,
                       r.slab.b_bottom_slide, r.slab.cos_angle, \&ru, \&tu);
        if (m.num\_spheres \equiv 0 \land m.m\_t > tu) {
               fprintf(stderr, "ntop=\%7.5f, \_nslab=\%7.5f, \_nbottom=\%7.5f \ ", r.slab.n\_top\_slide, r.slab.n\_slab, models = mo
                               r.slab.n_-bottom_-slide);
               fprintf(stderr, "tu_max=\%7.5f, um_t=\%7.5f, ut_std=\%7.5f n", tu, m.m_t, m.rstd_t);
               return IAD_MT_TOO_BIG;
This code is used in section 47.
50. The unscattered transmission is now always included in the total transmittance. Therefore the
unscattered transmittance must fall betwee zero and M_T
\langle \text{ Check MU 50} \rangle \equiv
       if (m.m_u < 0) return IAD_MU_TOO_SMALL;
        if (m.m_u > m.m_t) return IAD_MU_TOO_BIG;
This code is used in section 47.
```

51. Make sure that reflection sphere parameters are reasonable

```
 \begin{array}{l} \langle \, \text{Check sphere parameters 51} \, \rangle \equiv \\ \quad \text{if } \; (m.as\_r < 0 \lor m.as\_r \geq 0.2) \; \; \text{return IAD\_AS\_NOT\_VALID}; \\ \quad \text{if } \; (m.ad\_r < 0 \lor m.ad\_r \geq 0.2) \; \; \text{return IAD\_AD\_NOT\_VALID}; \\ \quad \text{if } \; (m.ae\_r < 0 \lor m.ae\_r \geq 0.2) \; \; \text{return IAD\_AE\_NOT\_VALID}; \\ \quad \text{if } \; (m.rw\_r < 0 \lor m.rw\_r > 1.0) \; \; \text{return IAD\_RW\_NOT\_VALID}; \\ \quad \text{if } \; (m.rd\_r < 0 \lor m.rd\_r > 1.0) \; \; \text{return IAD\_RD\_NOT\_VALID}; \\ \quad \text{if } \; (m.rstd\_r < 0 \lor m.rstd\_r > 1.0) \; \; \text{return IAD\_RSTD\_NOT\_VALID}; \\ \quad \text{if } \; (m.rstd\_t < 0 \lor m.rstd\_t > 1.0) \; \; \text{return IAD\_TSTD\_NOT\_VALID}; \\ \quad \text{if } \; (m.f\_r < 0 \lor m.f\_r > 1) \; \; \text{return IAD\_F\_NOT\_VALID}; \\ \quad \text{See also section 52}. \end{array}
```

This code is used in section 47.

52. Make sure that transmission sphere parameters are reasonable

```
⟨ Check sphere parameters 51⟩ +≡

if (m.as_-t < 0 \lor m.as_-t ≥ 0.2) return IAD_AS_NOT_VALID;

if (m.ad_-t < 0 \lor m.ad_-t ≥ 0.2) return IAD_AD_NOT_VALID;

if (m.ae_-t < 0 \lor m.ae_-t ≥ 0.2) return IAD_AE_NOT_VALID;

if (m.rw_-t < 0 \lor m.rw_-r > 1.0) return IAD_RW_NOT_VALID;

if (m.rd_-t < 0 \lor m.rd_-t > 1.0) return IAD_RD_NOT_VALID;

if (m.rstd_-t < 0 \lor m.rstd_-t > 1.0) return IAD_TSTD_NOT_VALID;

if (m.f_-t < 0 \lor m.f_-t > 1) return IAD_F_NOT_VALID;
```

 $\S53$ IAD (v 3-12-0) SEARCHING METHOD 33

53. Searching Method.

The original idea was that this routine would automatically determine what optical parameters could be figured out from the input data. This worked fine for a long while, but I discovered that often it was convenient to constrain the optical properties in various ways. Consequently, this routine got more and more complicated.

What should be done is to figure out whether the search will be 1D or 2D and split this routine into two parts.

It would be nice to enable the user to constrain two parameters, but the infrastructure is missing at this point.

```
\langle Prototype for determine\_search 53\rangle \equiv search_type determine\_search(struct measure_type m, struct invert_type r) This code is used in sections 39 and 54.
```

34 Searching method iad (v 3-12-0) §54

54. This routine is responsible for selecting the appropriate optical properties to determine.

```
\langle \text{ Definition for } determine\_search | 54 \rangle \equiv
  ⟨ Prototype for determine_search 53⟩
     double rt, tt, rd, td, tc, rc;
     int search = 0;
     int independent = m.num\_measures;
     if (Debug(DEBUG\_SEARCH)) {
       fprintf(stderr, "\n***_Determine_Search()\n");
       fprintf(stderr, "ullulustartinguwithu%dumeasurement(s)\n", m.num\_measures);
       fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup}m_r=\%.5f\n", m.m_r);
       fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup}m_t=\%.5f\n", m.m_t);
     Estimate\_RT(m, r, \&rt, \&tt, \&rd, \&rc, \&td, \&tc);
     if (m.m_{-}u \equiv 0 \land independent \equiv 3) {
       if (Debug(DEBUG\_SEARCH)) fprintf(stderr, "_{\sqcup \sqcup \sqcup \sqcup} no_{\sqcup} information_{\sqcup} in_{\sqcup} tc n");
       independent ---;
     if (rd \equiv 0 \land independent \equiv 2) {
       if (Debug(DEBUG\_SEARCH)) fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup}no_{\sqcup}information_{\sqcup}in_{\sqcup}rd\n");
       independent ---;
     if (td \equiv 0 \land independent \equiv 2) {
       independent ---;
     if (independent \equiv 1) {
       \langle \text{ One parameter search 55} \rangle
     else if (independent \equiv 2) {
       (Two parameter search 56)
           /* three real parameters with information! */
     else {
       search = FIND\_AG;
     if (Debuq(DEBUG\_SEARCH)) {
       fprintf(stderr, "uuuuindependentumeasurementsu=u%3d\n", independent);
       fprintf(stderr, "\____m_r=\%8.5f__m_t=\%8.5f__(rd__=_\%8.5f__td=\%8.5f) \n", m.m_r, m.m_t, rd, td);
       if (search \equiv FIND_B) fprintf(stderr, "_ \sqcup \sqcup \sqcup \sqcup search_ = \sqcup FIND_B \ ");
       \mathbf{if}\ (\mathit{search} \equiv \mathtt{FIND\_AG})\ \mathit{fprintf}(\mathit{stderr}, \texttt{"$$\sqcup\sqcup\sqcup\sqcup}\mathtt{search}_{\sqcup}\texttt{=}_{\sqcup}\mathtt{FIND\_AG}\texttt{\ensuremath{n"}});
        if (search \equiv FIND\_AUTO) \ fprintf(stderr, "\verb|u|| search = FIND\_AUTO \n"); \\
       if (search \equiv FIND\_BaG) fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup\sqcup}search_{\sqcup}=_{\sqcup}FIND\_BaG\n");
       if (search \equiv FIND\_BsG) fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup\sqcup}search_{\sqcup}=_{\sqcup}FIND\_BsG\n");
       if (search \equiv FIND\_Ba) fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup}search_{\sqcup}=_{\sqcup}FIND\_Ba\n");
       if (search \equiv FIND_G) fprintf(stderr, "_ \sqcup \sqcup \sqcup \sqcup search_ = \sqcup FIND_G \ ");
       if (search \equiv FIND_B_WITH_NO_ABSORPTION)
          fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup}search_{\sqcup}=_{\sqcup}FIND_B_WITH_NO_ABSORPTION\n");
```

55. The fastest inverse problems are those in which just one measurement is known. This corresponds to a simple one-dimensional minimization problem. The only complexity is deciding exactly what should be allowed to vary. The basic assumption is that the anisotropy has been specified or will be assumed to be zero.

If the anistropy is assumed known, then one other assumption will allow us to figure out the last parameter to solve for.

Ultimately, if no default values are given, then we look at the value of the total transmittance. If this is zero, then we assume that the optical thickness is infinite and solve for the albedo. Otherwise we will just make a stab at solving for the optical thickness assuming the albedo is one.

```
⟨One parameter search 55⟩ ≡

if (r.default_{-}a \neq \text{UNINITIALIZED}) {

if (r.default_{-}a \equiv 0) search = \text{FIND\_B\_WITH\_NO\_SCATTERING};

else if (r.default_{-}a \equiv 1) search = \text{FIND\_B\_WITH\_NO\_ABSORPTION};

else if (tt \equiv 0) search = \text{FIND\_G};

else search = \text{FIND\_B};
}

else if (r.default_{-}b \neq \text{UNINITIALIZED}) search = \text{FIND\_A};

else if (r.default_{-}bs \neq \text{UNINITIALIZED}) search = \text{FIND\_Ba};

else if (r.default_{-}ba \neq \text{UNINITIALIZED}) search = \text{FIND\_Bs};

else if (td \equiv 0) search = \text{FIND\_A};

else if (rd \equiv 0) search = \text{FIND\_B\_WITH\_NO\_SCATTERING};

else search = \text{FIND\_B\_WITH\_NO\_ABSORPTION};

This code is used in section 54.
```

36 Searching method iad (v 3-12-0) §56

56. If the absorption depth $\mu_a d$ is constrained return $FIND_BsG$. Recall that I use the bizarre mnemonic $bs = \mu_s d$ here and so this means that the program will search over various values of $\mu_s d$ and g.

If there are just two measurements then I assume that the anisotropy is not of interest and the only thing to calculate is the reduced albedo and optical thickness based on an assumed anisotropy.

```
\langle Two parameter search 56\rangle \equiv
  if (r.default_a \neq UNINITIALIZED) {
     if (r.default_a \equiv 0) search = FIND_B;
     else if (r.default_g \neq UNINITIALIZED) search = FIND_B;
     else search = FIND_BG;
  else if (r.default_b \neq UNINITIALIZED) {
     if (r.default\_g \neq UNINITIALIZED) search = FIND_A;
     \mathbf{else} \ \mathit{search} = \mathtt{FIND\_AG};
  else if (r.default_ba \neq UNINITIALIZED) {
    if (r.default\_q \neq UNINITIALIZED) search = FIND\_Bs;
     else search = FIND\_BsG;
  else if (r.default_bs \neq UNINITIALIZED) {
    if (r.default\_g \neq UNINITIALIZED) search = FIND\_Ba;
     else search = FIND\_BaG;
  else if (rt + tt > 1 \land 0 \land m.num\_spheres \neq 2) search = FIND_B_WITH_NO_ABSORPTION;
  else search = FIND\_AB;
This code is used in section 54.
```

57. This little routine just stuffs reasonable values into the structure we use to return the solution. This does not replace the values for $r.default_g$ nor for $r.method.quad_pts$. Presumably these have been set correctly elsewhere.

```
⟨ Prototype for Initialize_Result 57⟩ ≡
    void Initialize_Result(struct measure_type m, struct invert_type *r)
This code is used in sections 39 and 58.

58. ⟨ Definition for Initialize_Result 58⟩ ≡
    ⟨ Prototype for Initialize_Result 57⟩
    {
        ⟨ Fill r with reasonable values 59⟩
    }
This code is used in section 38.

59. Start with the optical properties.
⟨ Fill r with reasonable values 59⟩ ≡
    r→a = 0.0;
    r→b = 0.0;
    r→b = 0.0;
    r→g = 0.0;
See also sections 60, 61, and 62.
This code is used in section 58.
```

```
Continue with other useful stuff.
\langle \text{Fill } r \text{ with reasonable values } 59 \rangle + \equiv
   r \rightarrow found = FALSE;
   r \rightarrow tolerance = 0.0001;
   r \rightarrow MC\_tolerance = 0.01;
                                           /* percent */
   r \rightarrow search = FIND\_AUTO;
   r \rightarrow metric = RELATIVE;
   r \rightarrow final\_distance = 10;
   r \rightarrow iterations = 0; r \rightarrow error = IAD_NO_ERROR;
       The defaults might be handy
\langle \text{ Fill } r \text{ with reasonable values } 59 \rangle + \equiv
   r \rightarrow default_a = UNINITIALIZED;
   r \rightarrow default_b = UNINITIALIZED;
   r \rightarrow default_g = UNINITIALIZED;
   r \rightarrow default\_ba = UNINITIALIZED;
   r \rightarrow default\_bs = UNINITIALIZED;
   r \rightarrow default\_mua = UNINITIALIZED;
   r \rightarrow default\_mus = UNINITIALIZED;
      It is necessary to set up the slab correctly so, I stuff reasonable values into this record as well.
\langle \text{ Fill } r \text{ with reasonable values } 59 \rangle + \equiv
   r \rightarrow slab.a = 0.5;
   r \rightarrow slab.b = 1.0;
   r \rightarrow slab.g = 0;
   r \rightarrow slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};
   r \rightarrow slab . n\_slab = m.slab\_index;
   r \rightarrow slab.n\_top\_slide = m.slab\_top\_slide\_index;
   r \rightarrow slab.n\_bottom\_slide = m.slab\_bottom\_slide\_index;
   r \rightarrow slab.b\_top\_slide = m.slab\_top\_slide\_b;
   r \rightarrow slab.b\_bottom\_slide = m.slab\_bottom\_slide\_b;
   r \rightarrow slab.cos\_angle = m.slab\_cos\_angle;
   r \rightarrow method.a\_calc = 0.5;
   r \rightarrow method.b\_calc = 1;
   r \rightarrow method.g\_calc = 0.5;
   r \rightarrow method.quad\_pts = 8;
   r \rightarrow method.b_thinnest = 1.0/32.0;
```

63. EZ Inverse RT. *ez_Inverse_RT* is a simple interface to the main function *Inverse_RT* in this package. It eliminates the need for complicated data structures so that the command line interface (as well as those to Perl and Mathematica) will be simpler. This function assumes that the reflection and transmission include specular reflection and that the transmission also include unscattered transmission.

Other assumptions are that the top and bottom slides have the same index of refraction, that the illumination is collimated. Of course no sphere parameters are included.

```
\langle \text{Prototype for } ez\_Inverse\_RT \mid 63 \rangle \equiv 
void ez\_Inverse\_RT \mid \text{double } n, \text{double } nslide, \text{double UR1}, \text{double UT1}, \text{double } Tc, \text{double } *a, \text{double } *b, \text{double } *g, \text{ int } * \text{ error } )
This code is used in sections 39, 40, and 64.
```

38 EZ INVERSE RT IAD (v 3-12-0) §64

```
\langle \text{ Definition for } ez\_Inverse\_RT | 64 \rangle \equiv
64.
   \langle Prototype for ez\_Inverse\_RT 63 \rangle \{ struct measure\_type m; \}
        struct invert_type r;
        *a = 0;
        *b = \mathtt{HUGE\_VAL};
        *q = 0;
        Initialize\_Measure(\&m);
        m.slab\_index = n;
        m.slab\_top\_slide\_index = nslide;
        m.slab\_bottom\_slide\_index = nslide;
        m.slab\_cos\_angle = 1.0;
        m.num\_measures = 3;
        if (UT1 \equiv 0) m.num\_measures --;
        if (Tc \equiv 0) m.num_measures ---;
        m.m\_r = \mathtt{UR1};
        m.m_{-}t = \mathtt{UT1};
        m.m_u = Tc;
        Initialize\_Result(m, \&r);
        r.method.quad\_pts = 8;
        Inverse\_RT(m, \&r); *error = r . error ; if (r. error \equiv IAD\_NO\_ERROR)
        {
          *a = r.a;
          *b = r.b;
          *g = r.g;
This code is used in section 38.
     \langle \text{ Prototype for } Initialize\_Measure | 65 \rangle \equiv
  void Initialize_Measure(struct measure_type *m)
```

This code is used in sections 39 and 66.

```
\langle \text{ Definition for } Initialize\_Measure | 66 \rangle \equiv
⟨ Prototype for Initialize_Measure 65⟩
   double default\_sphere\_d = 8.0 * 25.4;
   double default\_sample\_d = 0.0 * 25.4;
   double default\_detector\_d = 0.1 * 25.4;
   double default\_entrance\_d = 0.5 * 25.4;
   double sphere = default\_sphere\_d * default\_sphere\_d;
   m \rightarrow slab\_index = 1.0;
   m \rightarrow slab\_top\_slide\_index = 1.0;
   m \rightarrow slab\_top\_slide\_b = 0.0;
   m \rightarrow slab\_top\_slide\_thickness = 0.0;
   m \rightarrow slab\_bottom\_slide\_index = 1.0;
   m \rightarrow slab\_bottom\_slide\_b = 0.0;
   m \rightarrow slab\_bottom\_slide\_thickness = 0.0;
   m \rightarrow slab\_thickness = 1.0:
   m \rightarrow slab\_cos\_angle = 1.0;
   m \rightarrow num\_spheres = 0;
   m \rightarrow num\_measures = 1;
   m \rightarrow method = UNKNOWN;
   m \rightarrow fraction\_of\_rc\_in\_mr = 1.0;
   m \rightarrow fraction\_of\_tc\_in\_mt = 1.0;
   m \rightarrow flip\_sample = 0;
   m \rightarrow m_{-}r = 0.0;
   m \rightarrow m_{-}t = 0.0;
   m \rightarrow m_{-}u = 0.0;
   m \rightarrow d\_sphere\_r = default\_sphere\_d;
   m \rightarrow as_r = default\_sample\_d * default\_sample\_d / sphere;
   m \rightarrow ad_r = default\_detector\_d * default\_detector\_d / sphere;
   m \rightarrow ae\_r = default\_entrance\_d * default\_entrance\_d/sphere;
   m \rightarrow aw_{-}r = 1.0 - m \rightarrow as_{-}r - m \rightarrow ad_{-}r - m \rightarrow ae_{-}r;
   m \rightarrow rd_{-}r = 0.0;
   m \rightarrow rw r = 1.0;
   m \rightarrow rstd_r = 1.0;
   m \rightarrow f_{-}r = 0.0;
   m \rightarrow d\_sphere\_t = default\_sphere\_d;
   m \rightarrow as_{-}t = m \rightarrow as_{-}r;
   m \rightarrow ad_{-}t = m \rightarrow ad_{-}r;
   m \rightarrow ae_{-}t = m \rightarrow ae_{-}r;
   m \rightarrow aw_{-}t = m \rightarrow aw_{-}r;
   m rd_t = 0.0;
   m \rightarrow rw_{-}t = 1.0;
   m \rightarrow rstd_-t = 1.0;
   m \rightarrow f_{-}t = 0.0;
   m \rightarrow lambda = 0.0;
   m \rightarrow d_- beam = 0.0;
   m \rightarrow ur1\_lost = 0;
   m \rightarrow uru\_lost = 0;
   m \rightarrow ut1\_lost = 0;
   m \rightarrow utu\_lost = 0;
```

This code is used in section 38.

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67. To avoid interfacing with C-structures it is necessary to pass the information as arrays. Here I have divided the experiment into (1) setup, (2) reflection sphere coefficients, (3) transmission sphere coefficients, (4) measurements, and (5) results.

 $\langle \text{Prototype for } Spheres_Inverse_RT | 67 \rangle \equiv$ **void** $Spheres_Inverse_RT \text{(double } *setup, \text{double } *analysis, \text{double } *sphere_r, \text{double } *sphere_t, \text{double } *results)$

This code is used in sections 40 and 68.

This code is used in section 38.

```
\langle \text{ Definition for } Spheres\_Inverse\_RT | 68 \rangle \equiv
\langle Prototype for Spheres\_Inverse\_RT \ 67 \rangle \{ struct measure\_type m; \}
     struct invert_type r;
     long num_photons;
     \mathbf{double} \ \mathit{ur1} \,, \ \mathit{ut1} \,, \ \mathit{uru} \,, \ \mathit{utu};
     int i, mc\_runs = 1;
      Initialize\_Measure(\&m);
      \langle \text{ handle setup } 69 \rangle
      \langle handle reflection sphere 72 \rangle
      (handle transmission sphere 73)
      \langle \text{ handle measurement } 71 \rangle
      Initialize\_Result(m, \&r);
      results[0] = 0;
      results[1] = 0;
      results[2] = 0;
      \langle \text{ handle analysis } 70 \rangle
      Inverse\_RT(m, \&r);
     for (i = 0; i < mc\_runs; i++) {
         MC\_Lost(m, r, num\_photons, \&ur1, \&ut1, \&uru, \&utu, \&m.ur1\_lost, \&m.ut1\_lost, \&m.uru\_lost,
              \&m.utu\_lost);
         Inverse\_RT(m, \&r);
     if (r \cdot \mathbf{error} \equiv \mathtt{IAD\_NO\_ERROR})
         results[0] = (1 - r.a) * r.b/m.slab\_thickness;
        results[1] = (r.a) * r.b/m.slab\_thickness;
        results[2] = r.g;
     }
      results[3] = r \cdot \mathbf{error}; }
```

This code is used in section 68.

These are in exactly the same order as the parameters in the .rxt header $\langle \text{ handle setup } 69 \rangle \equiv$ **double** *d_sample_r*, *d_entrance_r*, *d_detector_r*; **double** *d_sample_t*, *d_entrance_t*, *d_detector_t*; $m.slab_index = setup[0];$ $m.slab_top_slide_index = setup[1];$ $m.slab_thickness = setup[2];$ $m.slab_top_slide_thickness = setup[3];$ $m.d_beam = setup[4];$ $m.rstd_r = setup[5];$ $m.num_spheres = (int) setup[6];$ $m.d_sphere_r = setup[7];$ $d_sample_r = setup[8];$ $d_{-}entrance_{-}r = setup[9];$ $d_{-}detector_{-}r = setup[10];$ $m.rw_r = setup[11];$ $m.d_sphere_t = setup[12];$ $d_sample_t = setup[13];$ $d_{-}entrance_{-}t = setup[14];$ $d_{-}detector_{-}t = setup[15];$ $m.rw_{-}t = setup[16];$ $r.default_g = setup[17];$ $num_photons = (long) \ setup[18];$ $m.as_r = (d_sample_r/m.d_sphere_r) * (d_sample_r/m.d_sphere_r);$ $m.ae_r = (d_entrance_r/m.d_sphere_r) * (d_entrance_r/m.d_sphere_r);$ $m.ad_r = (d_detector_r/m.d_sphere_r) * (d_detector_r/m.d_sphere_r);$ $m.aw_r = 1.0 - m.as_r - m.ae_r - m.ad_r;$ $m.as_t = (d_sample_t/m.d_sphere_t) * (d_sample_t/m.d_sphere_t);$ $m.ae_t = (d_entrance_t/m.d_sphere_t) * (d_entrance_t/m.d_sphere_t);$ $m.ad_{-}t = (d_{-}detector_{-}t/m.d_{-}sphere_{-}t) * (d_{-}detector_{-}t/m.d_{-}sphere_{-}t);$ $m.aw_t = 1.0 - m.as_t - m.ae_t - m.ad_t;$ $m.slab_bottom_slide_index = m.slab_top_slide_index;$ $m.slab_bottom_slide_thickness = m.slab_top_slide_thickness;$ $fprintf(stderr, "****_lexecuting_lFIXME_l****/n");$ $m.slab_cos_angle = 1.0;$ /* FIXME */ This code is used in section 68. **70.** $\langle \text{ handle analysis } 70 \rangle \equiv$ $r.method.quad_pts = (int) \ analysis[0];$ $mc_runs = (\mathbf{int}) \ analysis[1];$

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```
71.
```

```
\langle \text{ handle measurement } 71 \rangle \equiv
  m.m_r = measurements[0];
  m.m_{-}t = measurements[1];
  m.m_{-}u = measurements[2];
  m.num\_measures = 3;
  if (m.m_t \equiv 0) m.num_measures --;
  if (m.m_u \equiv 0) m.num_measures ---;
This code is used in section 68.
72.
\langle handle reflection sphere 72\rangle \equiv
  m.as_r = sphere_r[0];
  m.ae_r = sphere_r[1];
  m.ad_r = sphere_r[2];
  m.rw_r = sphere_r[3];
  m.rd_r = sphere_r[4];
  m.rstd\_r = sphere\_r[5];
  m.f_r = sphere_r[7];
This code is used in section 68.
73.
\langle \text{ handle transmission sphere } 73 \rangle \equiv
  m.as_{-}t = sphere_{-}t[0];
  m.ae_{-}t = sphere_{-}t[1];
  m.ad_t = sphere_t[2];
  m.rw_t = sphere_t[3];
```

 $m.f_{-}t = sphere_{-}t[7];$ This code is used in section 68.

 $m.rd_{-}t = sphere_{-}t[4];$ $m.rstd_{-}t = sphere_{-}t[5];$

74. I needed a routine that would calculate the values of M_R and M_T without doing the whole inversion process. It seems odd that this does not exist yet.

The values for the lost light $m.uru_lost$ etc., should be calculated before calling this routine.

 $\langle \text{ Prototype for } Calculate_MR_MT | 74 \rangle \equiv$

void $Calculate_MR_MT$ (struct measure_type m, struct invert_type r, int $include_MC$, double $*M_R$, double $*M_T$)

This code is used in sections 39 and 75.

```
75.
      \langle \text{ Definition for } Calculate\_MR\_MT | 75 \rangle \equiv
  \langle Prototype for Calculate\_MR\_MT 74 \rangle
     double distance, ur1, ut1, uru, utu;
     struct measure_type old_mm;
     struct invert_type old_rr;
     if (include\_MC \land m.num\_spheres > 0) MC\_Lost(m, r, -2000, \&ur1, \&ut1, \&uru, \&utu, \&(m.ur1\_lost),
            &(m.ut1\_lost), &(m.uru\_lost), &(m.utu\_lost));
     Get\_Calc\_State(\&old\_mm,\&old\_rr);
     Set\_Calc\_State(m, r);
     Calculate\_Distance(M_R, M_T, \& distance);
     Set\_Calc\_State(old\_mm, old\_rr);
This code is used in section 38.
76. So, it turns out that the minimum measured M_R can be less than four percent for black glass! This is
because the sphere efficiency is much worse for the glass than for the white standard.
\langle \text{Prototype for } Calculate\_Minimum\_MR \ 76 \rangle \equiv
  void Calculate\_Minimum\_MR(struct measure_type m, struct invert_type r, double *mr, double
       *mt)
This code is used in sections 39 and 77.
77. \langle Definition for Calculate\_Minimum\_MR 77 \rangle \equiv
  \langle Prototype for Calculate\_Minimum\_MR 76 \rangle
    if (r.default_b \equiv \mathtt{UNINITIALIZED})
       if (r.slab.n\_slab > 1.0) r.slab.b = HUGE\_VAL;
       else r.slab.b = 1 \cdot 10^{-5};
     else r.slab.b = r.default_b;
     if (r.default_a \equiv UNINITIALIZED) r.slab.a = 0;
     else r.slab.a = r.default\_a;
     if (r.default\_g \equiv UNINITIALIZED) \ r.slab.g = 0.0;
     else r.slab.q = r.default_q;
     if (r.search \equiv FIND_G) \ r.slab.a = 0;
     r.a = r.slab.a;
     r.b = r.slab.b;
     r.g = r.slab.g;
     Calculate\_MR\_MT(m, r, 0, mr, mt);
     *mt = 0;
This code is used in section 38.
```

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The minimum possible value of MR for a given MT will be when the albedo is zero and the maximum value will be when the albedo is one. In the first case there will be no light loss and in the second we will assume that any light loss is neglible (to maximize MR).

The second case is perhaps over-simplified. Obviously for a fixed thickness as the albedo increases, the reflectance will increase. So how does $U_Find_B()$ work when the albedo is set to 1?

The problem is that to calculate these values one must know the optical thickness. Fortunately with the recent addition of constrained minimization, we can do exactly this.

The only thing that remains is to sort out the light lost effect.

```
\langle \text{ Prototype for } MinMax\_MR\_MT | 78 \rangle \equiv
  int MinMax_MR_MT (struct measure_type m, struct invert_type r)
This code is used in sections 39 and 79.
79. \langle \text{ Definition for } MinMax\_MR\_MT | 79 \rangle \equiv
  \langle Prototype for MinMax\_MR\_MT 78 \rangle
    double distance, measured_m_r, min_possible_m_r, max_possible_m_r, temp_m_t;
    if (m.m_r < 0) return IAD_MR_TOO_SMALL;
    if (m.m_r * m.rstd_r > 1) return IAD_MR_TOO_BIG;
    if (m.m_{-}t < 0) return IAD_MT_TOO_SMALL;
    if (m.m_t \equiv 0) return IAD_NO_ERROR;
    measured\_m\_r = m.m\_r;
    m.m_{-}r = 0;
    r.search = FIND_B;
    r.default_a = 0;
    U_{-}Find_{-}B(m, \&r);
    Calculate\_Distance(\&min\_possible\_m\_r, \&temp\_m\_t, \&temp\_m\_t, \&temp\_m\_t);
    if (measured_m_r < min_possible_m_r) return IAD_MR_TOO_SMALL;
    r.default_a = 1.0;
    U_Find_B(m, \&r);
    Calculate\_Distance(\&max\_possible\_m\_r, \&temp\_m\_t, \&distance);
    if (measured\_m\_r > max\_possible\_m\_r) return IAD_MR_TOO_BIG;
    return IAD_NO_ERROR;
This code is used in section 38.
     \langle Prototype for Spheres_Inverse_RT2 | 80 \rangle \equiv
  void Spheres_Inverse_RT2(double *sample, double *illumination, double *sphere_r, double
       **sphere_t, double **analysis, double *measurement, double **a, double **b, double **g)
This code is used in sections 39, 40, and 81.
```

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```
81.
      \langle \text{ Definition for } Spheres\_Inverse\_RT2 | 81 \rangle \equiv
  \langle Prototype for Spheres\_Inverse\_RT2 \ 80 \rangle \{  struct measure_type m;
       struct invert_type r;
       long num_photons;
       double ur1, ut1, uru, utu;
       int i, mc-runs = 1;
        Initialize\_Measure(\&m);
        \langle \text{ handle 2 sample 82} \rangle
        ⟨ handle2 illumination 83 ⟩
        \langle \text{ handle 2 reflection sphere } 84 \rangle
        \langle \text{ handle 2 transmission sphere } 85 \rangle
        (handle2 analysis 86)
        (handle2 measurement 87)
        Initialize\_Result(m, \&r);
        Inverse\_RT(m, \&r);
       for (i = 0; i < mc\_runs; i++) {
          MC\_Lost(m, r, num\_photons, \&ur1, \&ut1, \&uru, \&utu, \&m.ur1\_lost, \&m.ut1\_lost, \&m.uru\_lost,
               \&m.utu\_lost);
          Inverse\_RT(m, \&r);
       if (r \cdot error \equiv IAD_NO_ERROR)
          *a = r.a;
          *b = r.b;
          *g = r.g;
This code is used in section 38.
      Just move the values from the sample array into the right places
\langle \text{ handle 2 sample 82} \rangle \equiv
  m.slab\_index = sample[0];
  m.slab\_top\_slide\_index = sample[1];
  m.slab\_bottom\_slide\_index = sample[2];
  m.slab\_thickness = sample[3];
  m.slab\_top\_slide\_thickness = sample [4];
  m.slab\_bottom\_slide\_thickness = sample[5];
  m.slab\_top\_slide\_thickness = 0;
  m.slab\_bottom\_slide\_thickness = 0;
This code is used in section 81.
      Just move the values from the illumination array into the right places. Need to spend time to figure
out how to integrate items 2, 3, and 4
\langle \text{ handle 2 illumination } 83 \rangle \equiv
  m.d\_beam = illumination[0];
                                       /* m.lambda = illumination[1]; */
     /* m.specular-reflection-excluded = illumination[2]; */
                                                                         /* m.direct-transmission-excluded =
                                /* m.diffuse-illumination = illumination[4]; */
       illumination[3]; */
  m.num\_spheres = illumination [5];
This code is used in section 81.
```

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```
84.
\langle \text{ handle 2 reflection sphere } 84 \rangle \equiv
     double d_sample_r, d_entrance_r, d_detector_r;
     m.d\_sphere\_r = sphere\_r[0];
     d\_sample\_r = sphere\_r[1];
     d_{-}entrance_{-}r = sphere_{-}r[2];
     d\_detector\_r = sphere\_r[3];
     m.rw_r = sphere_r[4];
     m.rd_r = sphere_r[5];
     m.as\_r = (d\_sample\_r/m.d\_sphere\_r) * (d\_sample\_r/m.d\_sphere\_r);
     m.ae\_r = (d\_entrance\_r/m.d\_sphere\_r) * (d\_entrance\_r/m.d\_sphere\_r);
     m.ad_r = (d_detector_r/m.d_sphere_r) * (d_detector_r/m.d_sphere_r);
     m.aw_r = 1.0 - m.as_r - m.ae_r - m.ad_r;
This code is used in section 81.
85.
\langle \text{ handle 2 transmission sphere } 85 \rangle \equiv
     double d_sample_t, d_entrance_t, d_detector_t;
     m.d\_sphere\_t = sphere\_t[0];
     d\_sample\_t = sphere\_t[1];
     d_{-}entrance_{-}t = sphere_{-}t[2];
     d_{-}detector_{-}t = sphere_{-}t[3];
     m.rw_t = sphere_t[4];
     m.rd_t = sphere_t[5];
     m.as_t = (d\_sample\_t/m.d\_sphere\_t) * (d\_sample\_t/m.d\_sphere\_t);
     m.ae\_t = (d\_entrance\_t/m.d\_sphere\_t) * (d\_entrance\_t/m.d\_sphere\_t);
     m.ad_{-}t = (d_{-}detector_{-}t/m.d_{-}sphere_{-}t) * (d_{-}detector_{-}t/m.d_{-}sphere_{-}t);
     m.aw_{-}t = 1.0 - m.as_{-}t - m.ae_{-}t - m.ad_{-}t;
This code is used in section 81.
86.
\langle \text{ handle 2 analysis 86} \rangle \equiv
  r.method.quad\_pts = (int) \ analysis[0];
  mc\_runs = (\mathbf{int}) \ analysis[1];
  num\_photons = (\mathbf{long}) \ analysis[2];
This code is used in section 81.
87.
\langle \text{ handle 2 measurement } 87 \rangle \equiv
  m.rstd_r = measurement[0];
  m.m_{-}r = measurement[1];
  m.m_t = measurement[2];
  m.m_{-}u = measurement[3];
  m.num\_measures = 3;
  if (m.m_t \equiv 0) m.num_measures --;
  if (m.m_u \equiv 0) m.num_measures ---;
This code is used in section 81.
```

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88. IAD Input Output.

```
The special define below is to get Visual C to suppress silly warnings.
```

```
\langle iad_io.c 88 \rangle \equiv
#define _CRT_SECURE_NO_WARNINGS
#include <string.h>
#include <stdio.h>
#include <ctype.h>
#include <math.h>
#include "ad_globl.h"
#include "iad_type.h"
#include "iad_io.h"
#include "iad_pub.h"
#include "version.h"
   ⟨ Definition for skip_white 98 ⟩
   \langle \text{ Definition for } read\_number 100 \rangle
   \langle \text{ Definition for } check\_magic \ 102 \rangle
   ⟨ Definition for Read_Header 92 ⟩
   ⟨ Definition for Write_Header 104 ⟩
  ⟨ Definition for Read_Data_Line 96⟩
89. \langle iad_io.h 89 \rangle \equiv
  \langle Prototype for Read\_Header 91 \rangle;
  ⟨ Prototype for Write_Header 103⟩;
  \langle Prototype for Read\_Data\_Line 95 \rangle;
90. Reading the file header.
      \langle \text{ Prototype for } Read\_Header 91 \rangle \equiv
```

```
int Read_Header(FILE *fp, struct measure_type *m, int *params)
```

This code is used in sections 89 and 92.

92. Pretty straightforward stuff. The only thing that needs to be commented on is that only one slide thickness/index is specified in the file. This must be applied to both the top and bottom slides. Finally, to specify no slide, then either setting the slide index to 1.0 or the thickness to 0.0 should do the trick.

```
\langle \text{ Definition for } Read\_Header 92 \rangle \equiv
   ⟨ Prototype for Read_Header 91⟩
     double x;
     Initialize\_Measure(m);
     if (check\_magic(fp)) return 1;
     if (read_number(fp,&m¬slab_index)) return 1;
     if (read\_number(fp, \&m \neg slab\_top\_slide\_index)) return 1;
     if (read\_number(fp, \&m \neg slab\_thickness)) return 1;
     if (read_number(fp,&m¬slab_top_slide_thickness)) return 1;
     if (read\_number(fp,\&m\neg d\_beam)) return 1;
     if (m \rightarrow slab\_top\_slide\_thickness \equiv 0.0) m \rightarrow slab\_top\_slide\_index = 1.0;
     if (m \rightarrow slab\_top\_slide\_index \equiv 1.0) m \rightarrow slab\_top\_slide\_thickness = 0.0;
     if (m \rightarrow slab\_top\_slide\_index \equiv 0.0) {
        m \rightarrow slab\_top\_slide\_thickness = 0.0;
         m \rightarrow slab\_top\_slide\_index = 1.0;
     m \rightarrow slab\_bottom\_slide\_index = m \rightarrow slab\_top\_slide\_index;
     m \rightarrow slab\_bottom\_slide\_thickness = m \rightarrow slab\_top\_slide\_thickness;
     if (read\_number(fp, \&m \neg rstd\_r)) return 1;
     if (read\_number(fp, \&x)) return 1;
     m \rightarrow num\_spheres = (\mathbf{int}) x;
     m \rightarrow method = SUBSTITUTION;
     (Read coefficients for reflection sphere 93)
     \langle Read coefficients for transmission sphere 94\rangle
     if (read\_number(fp, \&x)) return 1;
     *params = (\mathbf{int}) x;
     m \rightarrow num\_measures = (*params \ge 3) ? 3 : *params;
     return 0;
This code is used in section 88.
      \langle Read coefficients for reflection sphere 93\rangle \equiv
93.
     double d_sample_r, d_entrance_r, d_detector_r;
     if (read\_number(fp, \&m \rightarrow d\_sphere\_r)) return 1;
     if (read_number(fp, &d_sample_r)) return 1;
     if (read\_number(fp, \&d\_entrance\_r)) return 1;
     if (read\_number(fp, \&d\_detector\_r)) return 1;
     if (read\_number(fp, \&m \neg rw\_r)) return 1;
     m \rightarrow as\_r = (d\_sample\_r/m \rightarrow d\_sphere\_r) * (d\_sample\_r/m \rightarrow d\_sphere\_r)/4.0;
     m \rightarrow ae\_r = (d\_entrance\_r/m \rightarrow d\_sphere\_r) * (d\_entrance\_r/m \rightarrow d\_sphere\_r)/4.0;
     m \rightarrow ad_r = (d_detector_r/m \rightarrow d_sphere_r) * (d_detector_r/m \rightarrow d_sphere_r)/4.0;
     m \rightarrow aw_r = 1.0 - m \rightarrow as_r - m \rightarrow ae_r - m \rightarrow ad_r;
This code is used in section 92.
```

```
94.
       \langle Read coefficients for transmission sphere 94 \rangle \equiv
     double d_sample_t, d_entrance_t, d_detector_t;
     if (read\_number(fp, \&m \rightarrow d\_sphere\_t)) return 1;
     if (read\_number(fp, \&d\_sample\_t)) return 1;
     if (read\_number(fp, \&d\_entrance\_t)) return 1;
     if (read\_number(fp, \&d\_detector\_t)) return 1;
     if (read\_number(fp, \&m \neg rw\_t)) return 1;
     m \rightarrow as\_t = (d\_sample\_t/m \rightarrow d\_sphere\_t) * (d\_sample\_t/m \rightarrow d\_sphere\_t)/4.0;
     m \rightarrow ae\_t = (d\_entrance\_t/m \rightarrow d\_sphere\_t) * (d\_entrance\_t/m \rightarrow d\_sphere\_t)/4.0;
     m \rightarrow ad_-t = (d_-detector_-t/m \rightarrow d_-sphere_-t) * (d_-detector_-t/m \rightarrow d_-sphere_-t)/4.0;
     m \rightarrow aw_{-}t = 1.0 - m \rightarrow as_{-}t - m \rightarrow ae_{-}t - m \rightarrow ad_{-}t;
This code is used in section 92.
95. Reading just one line of a data file.
  This reads a line of data based on the value of params.
  If the first number is greater than one then it is assumed to be the wavelength and is ignored. test on the
first value of the line.
  A non-zero value is returned upon a failure.
\langle Prototype for Read\_Data\_Line 95 \rangle \equiv
  int Read_Data_Line(FILE *fp, struct measure_type *m, int params)
This code is used in sections 89 and 96.
96. \langle \text{ Definition for } Read\_Data\_Line | 96 \rangle \equiv
   ⟨ Prototype for Read_Data_Line 95⟩
     if (read\_number(fp, \&m \neg m\_r)) return 1;
     if (m \rightarrow m_{-}r > 1) {
        m \rightarrow lambda = m \rightarrow m_r;
        if (read\_number(fp, \&m \neg m\_r)) return 1;
     if (params \equiv 1) return 0;
     if (read\_number(fp, \&m \neg m\_t)) return 1;
     if (params \equiv 2) return 0;
     if (read\_number(fp, \&m \neg m\_u)) return 1;
     if (params \equiv 3) return 0;
     if (read\_number(fp, \&m \neg rw\_r)) return 1;
     m \rightarrow rw_{-}t = m \rightarrow rw_{-}r;
     if (params \equiv 4) return 0;
     if (read\_number(fp, \&m \neg rw\_t)) return 1;
     if (params \equiv 5) return 0;
     if (read\_number(fp, \&m \neg rstd\_r)) return 1;
```

This code is used in section 88.

return 0;

if $(params \equiv 6)$ return 0;

if $(read_number(fp, \&m \neg rstd_t))$ return 1;

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97. Skip over white space and comments. It is assumed that # starts all comments and continues to the end of a line. This routine should work on files with nearly any line ending CR, LF, CRLF.

Failure is indicated by a non-zero return value.

```
\langle \text{ Prototype for } skip\_white 97 \rangle \equiv
  int skip_white(FILE *fp)
This code is used in section 98.
     \langle \text{ Definition for } skip\_white 98 \rangle \equiv
  ⟨ Prototype for skip_white 97⟩
     int c = fgetc(fp);
     while (\neg feof(fp)) {
        if (isspace(c)) c = fgetc(fp);
        else if (c \equiv "") do c = fgetc(fp); while (\neg feof(fp) \land c \neq "\n" \land c \neq "\");
        else break;
     if (feof(fp)) return 1;
     ungetc(c, fp);
     return 0;
This code is used in section 88.
99. Read a single number. Return 0 if there are no problems, otherwise return 1.
\langle \text{ Prototype for } read\_number 99 \rangle \equiv
  int read\_number(FILE *fp, double *x)
This code is used in section 100.
100. \langle \text{ Definition for } read\_number | 100 \rangle \equiv
  ⟨ Prototype for read_number 99⟩
     if (skip\_white(fp)) return 1;
     if (fscanf(fp, "%lf", x)) return 0;
     else return 1;
This code is used in section 88.
```

101. Ensure that the data file is actually in the right form. Return 0 if the file has the right starting characters. Return 1 if on a failure.

```
⟨ Prototype for check_magic 101 ⟩ ≡
int check_magic(FILE *fp)
This code is used in section 102.
```

```
102.
       \langle \text{ Definition for } check\_magic | 102 \rangle \equiv
  ⟨ Prototype for check_magic 101 ⟩
    char magic[] = "IAD1";
    int i, c;
    for (i = 0; i < 4; i++) {
       c = fgetc(fp);
       if (feof(fp) \lor c \neq magic[i]) {
         fprintf(stderr, "Sorry, \_but\_iad\_input\_files\_must\_begin\_with\_IAD1\n");
         fprintf(stderr, "_{\sqcup \sqcup \sqcup \sqcup \sqcup \sqcup \sqcup \sqcup \sqcup} as_{\sqcup} the_{\sqcup} first_{\sqcup} four_{\sqcup} characters_{\sqcup} of_{\sqcup} the_{\sqcup} file. \n");
         fprintf(stderr, "_{\sqcup \sqcup \sqcup \sqcup \sqcup \sqcup \sqcup \sqcup} Perhaps_{\sqcup} you_{\sqcup} are_{\sqcup} using_{\sqcup} an_{\sqcup} old_{\sqcup} iad_{\sqcup} format? \n");
         return 1;
    return 0;
This code is used in section 88.
       Formatting the header information.
\langle \text{ Prototype for } Write\_Header | 103 \rangle \equiv
  void Write_Header(struct measure_type m, struct invert_type r, int params)
This code is used in sections 89 and 104.
104. \langle \text{ Definition for } Write\_Header | 104 \rangle \equiv
  ⟨ Prototype for Write_Header 103⟩
     ⟨ Write slab info 105⟩
     ⟨ Write irradiation info 106⟩
     \langle \text{Write general sphere info } 107 \rangle
     Write first sphere info 108
     Write second sphere info 109
     \langle \text{Write measure and inversion info } 110 \rangle
This code is used in section 88.
105. \langle \text{Write slab info } 105 \rangle \equiv
  double xx;
  printf("\#_{\sqcup}Inverse_{\sqcup}Adding-Doubling_{\sqcup}\%s_{\sqcup}\n", Version);
  printf("#_{\sqcup}\n");
  printf("\#_{\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup}Sample_{\cup}index_{\cup}of_{\cup}refraction_{\cup}=_{\cup}\%7.4f\n", m.slab_{-}index);
  printf("\#_{\cup\cup\cup\cup\cup\cup\cup\cup\cup}Top_slide_index_of_refraction_=_\%7.4f\n", m.slab\_top\_slide\_index);
  This code is used in section 104.
106. \langle Write irradiation info 106 \rangle \equiv
  printf("#_{\sqcup}\n");
This code is used in section 104.
```

```
107. Write general sphere info 107 \geq
 printf("\#_{\sqcup\sqcup\sqcup\sqcup} Fraction_{\sqcup} unscattered_{\sqcup} refl._{\sqcup} in_{\sqcup} M_R_{\sqcup} = _{\sqcup} \%7.1 f_{\sqcup} \% \n", m. fraction_of_rc_in_mr * 100);
 printf("\#_{\sqcup\sqcup\sqcup} Fraction_{\sqcup} unscattered_{\sqcup} trans._{\sqcup} in_{\sqcup} M_{\_} T_{\sqcup} = _{\sqcup} \% 7.1 f_{\sqcup} \% \n", m. fraction_of_tc_in_mt * 100);
 printf("#_{\sqcup}\n");
This code is used in section 104.
108. (Write first sphere info 108) \equiv
 printf("#|Reflection|sphere\n");
 printf("\#_{\sqcup \sqcup \sqcup} \det \operatorname{etector}_{\sqcup} \operatorname{dameter}_{\sqcup = \sqcup} \%7.1 f_{\sqcup} \operatorname{mm}_{"}, 2*m.d\_sphere\_r * sqrt(m.ad\_r));
 printf("\#_{\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup}standard_reflectance_=\%7.1f_\%\n", m.rstd_r*100);
 printf("#_{UUUUUUUUUUUUUUUUUuuuudetector_reflectance_= \%7.1f_{\%}\n", m.rd_r*100);
 printf("#\n");
This code is used in section 104.
109. \langle \text{Write second sphere info } 109 \rangle \equiv
 printf("\#_{\square}Transmission_{\square}sphere\n");
 printf("\#_{\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup}standard_transmittance_=\%7.1f_\%\n", m.rstd_t * 100);
 printf("\#_{UUUUUUUUUUUUUUuuudetector_Ureflectance_U=_U\%7.1f_U\%\%n", m.rd_t*100);
This code is used in section 104.
```

```
110. \langle Write measure and inversion info | 110\rangle \equiv
  printf("#\n");
  switch (params) {
  case -1: printf("\#_{\square}No_{\square}M_{-}R_{\square}or_{\square}M_{-}T_{\square}--_{\square}forward_{\square}calculation.\n");
     break;
  case 1: printf("#□Just□M_R□was□measured");
     break:
  case 2: printf("\#_{\sqcup}M_{R_{\sqcup}}and_{\sqcup}M_{T_{\sqcup}}were_{\sqcup}measured");
     break:
  case 3: printf("#_M_R,_M_T,_and_M_U_were_measured");
     break;
  case 4: printf("\#_{\square}M_R,_{\square}M_T,_{\square}M_U,_{\square}and_{\square}r_w_{\square}were_{\square}measured");
     break:
  case 5: printf("#\luM_R,\luM_T,\luM_U,\lur_w,\luand\lut_w\luwere\lumeasured");
     break;
  case 6: printf("#\uM_R,\uM_T,\uM_U,\ur_w,\ut_w,\uand\ur_std\uwere\umeasured");
     break;
  case 7: printf("\#_{\sqcup}M_{\perp}R_{,\sqcup}M_{\perp}T_{,\sqcup}M_{\perp}U_{,\sqcup}r_{\perp}w_{,\sqcup}t_{\perp}w_{,\sqcup}r_{\perp}std_{\sqcup}and_{\sqcup}t_{\perp}std_{\sqcup}were_{\sqcup}measured");
     break;
  default: printf("#, Something, went, wrong,..., measures, should, be, 1, to, 5!\n");
     break;
  if (1 \leq params \wedge params \leq 7) {
     if (m.flip\_sample) printf("_{\sqcup}(sample_{\sqcup}flipped)_{\sqcup}");
     switch (m.method)  {
     case UNKNOWN: printf("usinguanunknownumethod.\n");
     case SUBSTITUTION: printf("usingutheusubstitutionu(single-beam)umethod.\n");
     case COMPARISON: printf("using_the_comparison_(dual-beam)_method.\n");
  switch (m.num\_spheres) {
  case 0: printf("#_No_sphere_corrections_were_used");
     break;
  case 1: printf("#⊔Single⊔sphere⊔corrections⊔were⊔used");
  case 2: printf("#_Double_sphere_corrections_were_used");
     break;
  printf("uwithulightuincidentuatu%dudegreesufromutheunormal",
        (int)(acos(m.slab\_cos\_angle)*57.2958));
  printf(".\n");
  switch (r.search) {
  case FIND_AB: printf("\#_{\square}The_{\square}inverse_{\square}routine_{\square}varied_{\square}the_{\square}albedo_{\square}and_{\square}optical_{\square}depth.\n");
     printf("#_{\sqcup}\n");
     xx = (r.default_g \neq UNINITIALIZED) ? r.default_g : 0;
     printf("\#_{\sqcup}Default_{\sqcup}single_{\sqcup}scattering_{\sqcup}anisotropy_{\sqcup}=_{\sqcup}\%7.3f_{\sqcup}\n", xx);
     break:
  case FIND_AG: printf("#UTheUinverseUroutineUvariedUtheUalbedoUandUanisotropy.\n");
     printf("#_{\sqcup}\n");
```

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```
if (r.default_b \neq UNINITIALIZED)
      else printf("#_{\sqcup}\n");
    break;
  case FIND_AUTO: printf("\#_{\square}The_{\square}inverse_{\square}routine_{\square}adapted_{\square}to_{\square}the_{\square}input_{\square}data.\n");
    printf("#_{\sqcup}\n");
    printf("#_{\sqcup}\n");
    break:
  case FIND_A: printf("#LTheLinverseLroutineLvariedLonlyLtheLalbedo.\n");
    printf("#_{\sqcup}\n");
    xx = (r.default_g \neq UNINITIALIZED) ? r.default_g : 0;
    printf("\#_{\square}Default_{\square}single_{\square}scattering_{\square}anisotropy_{\square}is_{\square}\%7.3f_{\square}", xx);
    xx = (r.default_b \neq UNINITIALIZED) ? r.default_b : HUGE_VAL;
    printf("_{\sqcup}and_{\sqcup}(mu_{t*d})_{\sqcup}=_{\sqcup}\%7.3g\n", xx);
    break;
  case FIND_B: printf("#\The\inverse\routine\varied\only\the\optical\depth.\n");
    printf("#_{\sqcup}\n");
    xx = (r.default_q \neq UNINITIALIZED) ? r.default_q : 0;
    printf("#_Default_single_scattering_anisotropy_is_%7.3f_", xx);
    if (r.default_a \neq UNINITIALIZED) printf("and_default_albedo_=_\%7.3g\n", r.default_a);
    else printf("\n");
    break;
  case FIND_Ba: printf("#_The_inverse_routine_varied_only_the_absorption.\n");
    printf("#_{\sqcup}\n");
    xx = (r.default\_bs \neq UNINITIALIZED) ? r.default\_bs : 0;
    break:
  case FIND_Bs: printf("#uTheuinverseuroutineuvarieduonlyutheuscattering.\n");
    printf("#_{\sqcup}\n");
    xx = (r.default_ba \neq UNINITIALIZED) ? r.default_ba : 0;
    break;
  default: printf("\#_{\!\!\!\perp} \n");
    printf("#_{\sqcup}\n");
    printf("#_{\sqcup}\n");
    break;
  printf("\#_{ \cup \cup} AD_{ \cup} tolerance_{ \cup} for_{ \cup} success_{ \cup} =_{ \cup} \%9.5f \n", r.tolerance);
  printf("\#_{\cup\cup\cup\cup\cup\cup}MC_{\cup}tolerance\_for_{\cup}mu_a_{\cup}and_{\cup}mu_s'_{\cup}=_{\cup}\%7.3f_{\cup}\%\%n", r.MC\_tolerance);
This code is used in section 104.
```

111. IAD Calculation.

```
\langle iad\_calc.c 111 \rangle \equiv
#include <math.h>
#include <string.h>
#include <stdio.h>
#include <stdlib.h>
#include "nr_util.h"
#include "nr_zbrent.h"
#include "ad_globl.h"
#include "ad_frsnl.h"
#include "ad_prime.h"
#include "iad_type.h"
#include "iad_util.h"
#include "iad_calc.h"
#define ABIT 1 \cdot 10^{-6}
#define A_COLUMN 1
#define B_COLUMN 2
#define G_COLUMN 3
#define URU_COLUMN 4
\#define UTU_COLUMN 5
#define UR1_COLUMN 6
#define UT1_COLUMN 7
#define REFLECTION_SPHERE 1
\#define TRANSMISSION_SPHERE 0
#define GRID_SIZE 101
#define T_TRUST_FACTOR 2
  static int CALCULATING_GRID = 1:
  static struct measure_type MM;
  static struct invert_type RR;
  static struct measure_type MGRID;
  static struct invert_type RGRID;
  static double **The\_Grid = \Lambda;
  static double GG_a:
  static double GG_{-}b;
  static double GG_{-q};
  static double GG_{-}bs;
  static double GG_{-}ba;
  static boolean_type The_Grid_Initialized = FALSE;
  static boolean_type The\_Grid\_Search = -1;
  ⟨ Definition for Set_Calc_State 127⟩
  \langle \text{ Definition for } Get\_Calc\_State | 129 \rangle
   (Definition for Same_Calc_State 131)
   (Prototype for Fill\_AB\_Grid\ 149);
   \langle \text{ Prototype for } Fill\_AG\_Grid \ 154 \rangle;
   \langle \text{ Definition for } RT\_Flip 147 \rangle
   Definition for Allocate_Grid 133 \
   Definition for Valid_Grid 137
   Definition for fill_grid_entry 148
   Definition for Fill_Grid 164
   Definition for Near_Grid_Points 145
   \langle \text{ Definition for } Fill\_AB\_Grid \ 150 \rangle
  \langle \text{ Definition for } Fill\_AG\_Grid \ 155 \rangle
```

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```
\langle \text{ Definition for } Fill\_BG\_Grid 158 \rangle
\langle \text{ Definition for } Fill\_BaG\_Grid \ 160 \rangle
\langle \text{ Definition for } Fill\_BsG\_Grid \ \ 162 \rangle
\langle \text{ Definition for } Grid\_ABG | 135 \rangle
Definition for Gain 116
\langle \text{ Definition for } Gain_{-}11 \text{ } 118 \rangle
\langle \text{ Definition for } Gain\_22 120 \rangle
(Definition for Two\_Sphere\_R 122)
 Definition for Two\_Sphere\_T 124\rangle
(Definition for Calculate_Distance_With_Corrections 170)
⟨ Definition for Calculate_Grid_Distance 168 ⟩
 Definition for Calculate_Distance 166
(Definition for abg\_distance 143)
\langle \text{ Definition for } Find\_AG\_fn \text{ 180} \rangle
\langle \text{ Definition for } Find\_AB\_fn \text{ 182} \rangle
 Definition for Find\_Ba\_fn 184\rangle
(Definition for Find_Bs_fn 186)
\langle \text{ Definition for } Find\_A\_fn \mid 188 \rangle
\langle \text{ Definition for } Find\_B\_fn 190 \rangle
 Definition for Find_-G_-fn 192\rangle
\langle \text{ Definition for } Find\_BG\_fn 194 \rangle
\langle \text{ Definition for } Find\_BaG\_fn \ \ 196 \rangle
 Definition for Find_{-}BsG_{-}fn 198 \rangle
 Definition for maxloss 200
```

112.

```
\langle iad\_calc.h \quad 112 \rangle \equiv
   \langle \text{ Prototype for } Gain \ 115 \rangle;
   \langle \text{Prototype for } Gain\_11 \text{ 117} \rangle;
   \langle \text{ Prototype for } Gain\_22 \text{ 119} \rangle;
    Prototype for Two\_Sphere\_R 121 \rangle;
   \langle Prototype for Two\_Sphere\_T 123 \rangle;
   \langle Prototype for Set_Calc_State 126 \rangle;
   \langle Prototype for Get\_Calc\_State 128 \rangle;
    Prototype for Same\_Calc\_State = 130:
    (Prototype for Valid\_Grid\ 136);
   \langle Prototype for Allocate\_Grid 132 \rangle;
   \langle \text{ Prototype for } Fill\_Grid \ 163 \rangle;
    Prototype for Near\_Grid\_Points 144;
    Prototype for Grid\_ABG 134\rangle;
   \langle \text{ Prototype for } Find\_AG\_fn \mid 179 \rangle;
    (Prototype for Find\_AB\_fn 181);
    Prototype for Find_Ba_fn = 183;
    (Prototype for Find_Bs_fn \ 185);
   \langle \text{ Prototype for } Find\_A\_fn \ 187 \rangle;
    (Prototype for Find_B = fn \ 189);
    Prototype for Find_{-}G_{-}fn 191 \;
   \langle \text{ Prototype for } Find\_BG\_fn \ 193 \rangle;
   \langle Prototype for Find\_BsG\_fn 197 \rangle;
   \langle Prototype for Find_BaG_fn 195 \rangle;
   \langle Prototype for Fill\_BG\_Grid 157 \rangle;
   \langle Prototype for Fill\_BsG\_Grid 161 \rangle;
   \langle Prototype for Fill\_BaG\_Grid 159 \rangle;
    Prototype for Calculate_Distance_With_Corrections 169;
   \langle Prototype for Calculate\_Distance 165 \rangle;
   \langle Prototype for Calculate\_Grid\_Distance 167 \rangle;
   \langle \text{ Prototype for } abg\_distance \ 142 \rangle;
    Prototype for maxloss 199;
   \langle Prototype for Max\_Light\_Loss 201 \rangle;
```

113. Initialization.

The functions in this file assume that the local variables MM and RR have been initialized appropriately. The variable MM contains all the information about how a particular experiment was done. The structure RR contains the data structure that is passed to the adding-doubling routines as well as the number of quadrature points.

history 6/8/94 changed error output to stderr.

114. Gain.

Assume that a sphere is illuminated with diffuse light having a power P. This light can reach all parts of sphere — specifically, light from this source is not blocked by a baffle. Multiple reflections in the sphere will increase the power falling on non-white areas in the sphere (e.g., the sample, detector, and entrance) To find the total we need to sum all the total of all incident light at a point. The first incidence is

$$P_w^{(1)} = a_w P, \qquad P_s^{(1)} = a_s P, \qquad P_d^{(1)} = a_d P$$

The light from the detector and sample is multiplied by $(1 - a_e)$ and not by a_w because the light from the detector (and sample) is not allowed to hit either the detector or sample. The second incidence on the wall is

$$P_w^{(2)} = a_w r_w P_w^{(1)} + (1 - a_e) r_d P_d^{(1)} + (1 - a_e) r_s P_s^{(1)}$$

The light that hits the walls after k bounces has the same form as above

$$P_w^{(k)} = a_w r_w P_w^{(k-1)} + (1 - a_e) r_d P_d^{(k-1)} + (1 - a_e) r_s P_s^{(k-1)}$$

Since the light falling on the sample and detector must come from the wall

$$P_s^{(k)} = a_s r_w P_w^{(k-1)}$$
 and $P_d^{(k)} = a_d r_w P_w^{(k-1)}$,

Therefore,

$$P_w^{(k)} = a_w r_w P_w^{(k-1)} + (1 - a_e) r_w (a_d r_d + a_s r_s) P_w^{(k-2)}$$

The total power falling on the walls is just

$$P_w = \sum_{k=1}^{\infty} P_w^{(k)} = \frac{a_w + (1 - a_e)(a_d r_d + a_s r_s)}{1 - a_w r_w - (1 - a_e)r_w(a_d r_d + a_s r_s)} P$$

The total power falling the detector is

$$P_d = a_d P + \sum_{k=2}^{\infty} a_d r_w P_w^{(k-1)} = a_d P + a_d r_w P_w$$

The gain $G(r_s)$ on the irradiance on the detector (relative to a black sphere),

$$G(r_s) \equiv \frac{P_d/A_d}{P/A}$$

in terms of the sphere parameters

$$G(r_s) = 1 + \frac{1}{a_w} \cdot \frac{a_w r_w + (1 - a_e) r_w (a_d r_d + a_s r_s)}{1 - a_w r_w - (1 - a_e) r_w (a_d r_d + a_s r_s)}$$

The gain for a detector in a transmission sphere is similar, but with primed parameters to designate a second potential sphere that is used. For a black sphere the gain G(0) = 1, which is easily verified by setting $r_w = 0$, $r_s = 0$, and $r_d = 0$. Conversely, when the sphere walls and sample are perfectly white, the irradiance at the entrance port, the sample port, and the detector port must increase so that the total power leaving via these ports is equal to the incident diffuse power P. Thus the gain should be the ratio of the sphere wall area over the area of the ports through which light leaves or $G(1) = A/(A_e + A_d)$ which follows immediately from the gain formula with $r_w = 1$, $r_s = 1$, and $r_d = 0$.

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115. The gain $G(r_s)$ on the irradiance on the detector (relative to a black sphere),

$$G(r_s) \equiv \frac{P_d/A_d}{P/A}$$

in terms of the sphere parameters

$$G(r_s) = 1 + \frac{a_w r_w + (1 - a_e) r_w (a_d r_d + a_s r_s)}{1 - a_w r_w - (1 - a_e) r_w (a_d r_d + a_s r_s)}$$

 $\langle \text{ Prototype for } Gain | 115 \rangle \equiv$

double $Gain(int sphere, struct measure_type m, double URU)$

This code is used in sections 112 and 116.

```
116. 〈 Definition for Gain \ 116〉 \equiv 〈 Prototype for Gain \ 115〉 { double G, \ tmp; if (sphere \equiv \texttt{REFLECTION\_SPHERE}) { tmp = m.rw\_r * (m.aw\_r + (1 - m.ae\_r) * (m.ad\_r * m.rd\_r + m.as\_r * \texttt{URU})); if (tmp \equiv 1.0) \ G = 1; else G = 1.0 + tmp/(1.0 - tmp); } else { tmp = m.rw\_t * (m.aw\_t + (1 - m.ae\_t) * (m.ad\_t * m.rd\_t + m.as\_t * \texttt{URU})); if (tmp \equiv 1.0) \ G = 1; else G = 1.0 + tmp/(1.0 - tmp); } return G; }
```

This code is used in section 111.

117. The gain for light on the detector in the first sphere for diffuse light starting in that same sphere is defined as

$$G_{1\to 1}(r_s, t_s) \equiv \frac{P_{1\to 1}(r_s, t_s)/A_d}{P/A}$$

then the full expression for the gain is

$$G_{1\to 1}(r_s, t_s) = \frac{G(r_s)}{1 - a_s a_s' r_w r_w' (1 - a_e) (1 - a_e') G(r_s) G'(r_s) t_s^2}$$

 $\langle \text{ Prototype for } Gain_{-}11 \text{ 117} \rangle \equiv$

double Gain_11 (struct measure_type m, double URU, double tdiffuse)

This code is used in sections 112 and 118.

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```
118. \langle Definition for Gain\_11 118\rangle \equiv \langle Prototype for Gain\_11 117\rangle {
    double G, GP, G11;
    G = Gain(REFLECTION_SPHERE, m, URU);
    GP = Gain(TRANSMISSION_SPHERE, m, URU);
    G11 = G/(1 - m.as\_r * m.as\_t * m.aw\_r * m.aw\_t * (1 - m.ae\_r) * (1 - m.ae\_t) * G * GP * tdiffuse * tdiffuse);
    return G11;
}
```

This code is used in section 111.

119. Similarly, when the light starts in the second sphere, the gain for light on the detector in the second sphere $G_{2\to 2}$ is found by switching all primed variables to unprimed. Thus $G_{2\to 1}(r_s,t_s)$ is

$$G_{2\to 2}(r_s, t_s) = \frac{G'(r_s)}{1 - a_s a_s' r_w r_w' (1 - a_e) (1 - a_e') G(r_s) G'(r_s) t_s^2}$$

 $\langle \text{ Prototype for } Gain_22 \text{ 119} \rangle \equiv$

double Gain_22 (struct measure_type m, double URU, double tdiffuse)

This code is used in sections 112 and 120.

```
120. \langle Definition for Gain\_22\ 120\rangle\equiv\langle Prototype for Gain\_22\ 119\rangle { double G, GP, G22; G=Gain(\texttt{REFLECTION\_SPHERE},m,\texttt{URU}); GP = Gain(\texttt{TRANSMISSION\_SPHERE},m,\texttt{URU}); G22 = GP/(1-m.as\_r*m.as\_t*m.aw\_r*m.aw\_t*(1-m.ae\_r)*(1-m.ae\_t)*G*GP*tdiffuse*tdiffuse); return G22; }
```

This code is used in section 111.

121. The reflected power for two spheres makes use of the formulas for Gain_11 above.

The light on the detector in the reflection (first) sphere arises from three sources: the fraction of light directly reflected off the sphere wall $fr_w^2(1-a_e)P$, the fraction of light reflected by the sample $(1-f)r_s^{\text{direct}}r_w^2(1-a_e)P$, and the light transmitted through the sample $(1-f)t_s^{\text{direct}}r_w'(1-a_e')P$,

$$\begin{split} R(r_s^{\text{direct}}, r_s, t_s^{\text{direct}}, t_s) &= G_{1 \to 1}(r_s, t_s) \cdot a_d (1 - a_e) r_w^2 f P \\ &+ G_{1 \to 1}(r_s, t_s) \cdot a_d (1 - a_e) r_w (1 - f) r_s^{\text{direct}} P \\ &+ G_{2 \to 1}(r_s, t_s) \cdot a_d (1 - a_e') r_w' (1 - f) t_s^{\text{direct}} P \end{split}$$

which simplifies slightly to

$$\begin{split} R(r_s^{\text{direct}}, r_s, t_s^{\text{direct}}, t_s) &= a_d (1 - a_e) r_w P \cdot G_{1 \rightarrow 1}(r_s, t_s) \\ &\times \left[(1 - f) r_s^{\text{direct}} + f r_w + (1 - f) a_s' (1 - a_e') r_w' t_s^{\text{direct}} t_s G'(r_s) \right] \end{split}$$

 $\langle \text{ Prototype for } Two_Sphere_R \ 121 \rangle \equiv$

double $Two_Sphere_R($ struct measure_type m, double UR1, double URU, double UT1, double UTU) This code is used in sections 112 and 122.

 $\{122 \text{ IAD (v } 3-12-0)\}$ GAIN 61

```
122. \langle Definition for Two\_Sphere\_R 122\rangle \equiv \langle Prototype for Two\_Sphere\_R 121\rangle \{ double x, GP; GP = Gain(TRANSMISSION\_SPHERE, m, URU); x = m.ad\_r * (1 - m.ae\_r) * m.rw\_r * <math>Gain\_11(m, URU, UTU); x *= (1 - m.f\_r) * UR1 + m.rw\_r * m.f\_r + (1 - m.f\_r) * m.as\_t * (1 - m.ae\_t) * m.rw\_t * UT1 * UTU * GP; return x; \} This code is used in section 111.
```

123. For the power on the detector in the transmission (second) sphere we have the same three sources. The only difference is that the subscripts on the gain terms now indicate that the light ends up in the second sphere

$$\begin{split} T(r_s^{\text{ direct}}, r_s, t_s^{\text{ direct}}, t_s) &= G_{1 \rightarrow 2}(r_s, t_s) \cdot a_d'(1 - a_e) r_w^2 f P \\ &+ G_{1 \rightarrow 2}(r_s, t_s) \cdot a_d'(1 - a_e) r_w (1 - f) r_s^{\text{ direct}} P \\ &+ G_{2 \rightarrow 2}(r_s, t_s) \cdot a_d'(1 - a_e') r_w'(1 - f) t_s^{\text{ direct}} P \end{split}$$

or

$$\begin{split} T(r_s^{\text{direct}}, r_s, t_s^{\text{direct}}, t_s) &= a_d'(1 - a_e') r_w' P \cdot G_{2 \to 2}(r_s, t_s) \\ &\times \left[(1 - f) t_s^{\text{direct}} + (1 - a_e) r_w a_s t_s (f r_w + (1 - f) r_s^{\text{direct}}) G(r_s) \right] \end{split}$$

 $\langle Prototype for Two_Sphere_T 123 \rangle \equiv$

This code is used in section 111.

double Two_Sphere_T (struct measure_type m, double UR1, double URU, double UT1, double UTU) This code is used in sections 112 and 124.

```
124. \langle Definition for Two\_Sphere\_T 124\rangle \equiv \langle Prototype for Two\_Sphere\_T 123\rangle \{ double x, G; G = Gain(\texttt{REFLECTION\_SPHERE}, m, \texttt{URU}); x = m.ad\_t * (1 - m.ae\_t) * m.rw\_t * Gain\_22(m, \texttt{URU}, \texttt{UTU}); x *= (1 - m.f\_r) * \texttt{UT1} + (1 - m.ae\_r) * m.rw\_r * m.as\_r * \texttt{UTU} * (m.f\_r * m.rw\_r + (1 - m.f\_r) * \texttt{UR1}) * G; return x; \}
```

125. Grid Routines. There is a long story associated with these routines. I spent a lot of time trying to find an empirical function to allow a guess at a starting value for the inversion routine. Basically nothing worked very well. There were too many special cases and what not. So I decided to calculate a whole bunch of reflection and transmission values and keep their associated optical properties linked nearby.

I did the very simplest thing. I just allocate a matrix that is five columns wide. Then I fill every row with a calculated set of optical properties and observables. The distribution of values that I use could certainly use some work, but they currently work.

SO... how does this thing work anyway? There are two possible grids one for calculations requiring the program to find the albedo and the optical depth (a and b) and one to find the albedo and anisotropy (a and g). These grids must be allocated and initialized before use.

62 GRID ROUTINES IAD (v 3-12-0) §126

126. This is a pretty important routine that should have some explanation. The reason that it exists, is that we need some 'out-of-band' information during the minimization process. Since the light transport calculation depends on all sorts of stuff (e.g., the sphere parameters) and the minimization routines just vary one or two parameters this information needs to be put somewhere.

I chose the global variables MM and RR to save things in.

The bottom line is that you cannot do a light transport calculation without calling this routine first.

```
void Set_Calc_State(struct measure_type m, struct invert_type r)
```

This code is used in sections 112 and 127.

This code is used in section 111.

This code is used in section 111.

```
127. ⟨ Definition for Set_Calc_State 127⟩ ≡
⟨ Prototype for Set_Calc_State 126⟩
{
    memcpy(&MM, &m, sizeof(struct measure_type));
    memcpy(&RR, &r, sizeof(struct invert_type));
    if (Debug(DEBUG_ITERATIONS) ∧ ¬CALCULATING_GRID) {
        fprintf(stderr, "UR1_loss=%g,_UT1_loss=%g,_UT1_loss=%g,_UT1_lost, m.ut1_lost, m.ut1_lost);
        fprintf(stderr, "URU_loss=%g,_UTU_loss=%g\n", m.uru_lost, m.utu_lost);
    }
}
```

128. The inverse of the previous routine. Note that you must have space for the parameters m and r already allocated.

```
⟨ Prototype for Get_Calc_State 128⟩ ≡
   void Get_Calc_State(struct measure_type *m, struct invert_type *r)
This code is used in sections 112 and 129.

129. ⟨ Definition for Get_Calc_State 129⟩ ≡
   ⟨ Prototype for Get_Calc_State 128⟩
   {
      memcpy(m, &MM, sizeof(struct measure_type));
      memcpy(r, &RR, sizeof(struct invert_type));
    }
```

130. The inverse of the previous routine. Note that you must have space for the parameters m and r already allocated.

```
\langle \text{Prototype for } Same\_Calc\_State \mid 130 \rangle \equiv  boolean_type Same\_Calc\_State (\text{struct measure\_type } m, \text{struct invert\_type } r) This code is used in sections 112 and 131.
```

```
GRID ROUTINES
```

```
§131 IAD (v 3-12-0)
```

```
131.
        \langle \text{ Definition for } Same\_Calc\_State | 131 \rangle \equiv
  \langle Prototype for Same\_Calc\_State 130 \rangle
     if (The\_Grid \equiv \Lambda) return FALSE;
     if (¬The_Grid_Initialized) return FALSE;
     if (r.search \neq RR.search) return FALSE;
     if (r.method.quad.pts \neq RR.method.quad.pts) return FALSE;
     if (r.slab.a \neq RR.slab.a) return FALSE;
     if (r.slab.b \neq RR.slab.b) return FALSE;
     if (r.slab.g \neq RR.slab.g) return FALSE;
     if (r.slab.phase\_function \neq RR.slab.phase\_function) return FALSE;
     if (r.slab.n\_slab \neq RR.slab.n\_slab) return FALSE;
     if (r.slab.n\_top\_slide \neq RR.slab.n\_top\_slide) return FALSE;
     if (r.slab.n\_bottom\_slide \neq RR.slab.n\_bottom\_slide) return FALSE;
     if (r.slab.b\_top\_slide \neq RR.slab.b\_top\_slide) return FALSE;
     if (r.slab.b\_bottom\_slide \neq RR.slab.b\_bottom\_slide) return FALSE;
     if (r.slab.cos\_angle \neq RR.slab.cos\_angle) return FALSE;
     if ((m.num\_measures \equiv 3) \land (m.m\_u \neq MGRID.m\_u)) return (FALSE);
     return TRUE;
This code is used in section 111.
        \langle Prototype for Allocate\_Grid 132 \rangle \equiv
  void Allocate_Grid(search_type s)
This code is used in sections 112 and 133.
133. \langle \text{ Definition for } Allocate\_Grid \ 133 \rangle \equiv
  ⟨ Prototype for Allocate_Grid 132⟩
     The\_Grid = dmatrix(0, GRID\_SIZE * GRID\_SIZE, 1, 7);
     if (The\_Grid \equiv \Lambda) \ AD\_error("unable\_to\_allocate\_the\_grid\_matrix");
     The\_Grid\_Initialized = FALSE;
This code is used in section 111.
134. This routine will return the a, b, and g values for a particular row in the grid.
\langle \text{ Prototype for } Grid\_ABG | 134 \rangle \equiv
  void Grid\_ABG(int i, int j, guess\_type *guess)
This code is used in sections 112 and 135.
```

64 GRID ROUTINES IAD (v 3-12-0) §135

```
135. \langle Definition for Grid\_ABG 135\rangle \equiv \langle Prototype for Grid\_ABG 134\rangle \{

if (0 \le i \land i < GRID\_SIZE \land 0 \le j \land j < GRID\_SIZE) \{

guess \neg a = The\_Grid [GRID\_SIZE * i + j] [A\_COLUMN];

guess \neg b = The\_Grid [GRID\_SIZE * i + j] [B\_COLUMN];

guess \neg distance = Calculate\_Grid\_Distance(i, j);
\}

else \{

guess \neg a = 0.5;

guess \neg b = 0.5;

guess \neg b = 0.5;

guess \neg distance = 999;
\}
\}

This code is used in section 111.
```

136. This routine is used to figure out if the current grid is valid. This can fail for several reasons. First the grid may not have been allocated. Or it may not have been initialized. The boundary conditions may have changed. The number or values of the sphere parameters may have changed. It is tedious, but straightforward to check these cases out.

If this routine returns true, then it is a pretty good bet that the values in the current grid can be used to guess the next starting set of values.

```
\langle \text{ Prototype for } Valid\_Grid \ 136 \rangle \equiv
  boolean_type Valid_Grid(struct measure_type m, search_type s)
This code is used in sections 112 and 137.
       \langle \text{ Definition for } Valid\_Grid 137 \rangle \equiv
  ⟨ Prototype for Valid_Grid 136⟩
     (Tests for invalid grid 138)
     return (TRUE);
This code is used in section 111.
       First check are to test if the grid has ever been filled
\langle Tests for invalid grid 138 \rangle \equiv
  if (The\_Grid \equiv \Lambda) {
     if (Debug(DEBUG\_GRID)) fprintf(stderr, "GRID: \bot Fill\_because \bot NULL \setminus n");
     return (FALSE);
  if (\neg The\_Grid\_Initialized) {
     if (Debug(DEBUG\_GRID)) fprintf(stderr, "GRID: \_Fill\_because\_not\_initialized \n");
     return (FALSE);
See also sections 139, 140, and 141.
This code is used in section 137.
```

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```
139. If the type of search has changed then report the grid as invalid
\langle Tests for invalid grid 138\rangle + \equiv
  if (The\_Grid\_Search \neq s) {
    if (Debug(DEBUG_GRID)) fprintf(stderr, "GRID: |Fill|because|search|type|changed\n");
    return (FALSE);
      Compare the m.m_u value only if there are three measurements
\langle Tests for invalid grid 138\rangle + \equiv
  if ((m.num\_measures \equiv 3) \land (m.m\_u \neq MGRID.m\_u)) {
    if (Debug(DEBUG\_GRID)) fprintf(stderr, "GRID: \botFill_ because_ unscattered_ light_ changed "");
    return (FALSE);
  }
141. Make sure that the boundary conditions have not changed.
\langle Tests for invalid grid 138\rangle + \equiv
  if (m.slab\_index \neq MGRID.slab\_index) {
    if (Debug(DEBUG\_GRID)) fprintf(stderr, "GRID: \_Fill\_slab\_refractive\_index\_changed\n");
    return (FALSE);
  if (m.slab\_cos\_angle \neq MGRID.slab\_cos\_angle) {
    if (Debug(DEBUG_GRID)) fprintf(stderr, "GRID: □Fill□incident□light□changed\n");
    return (FALSE);
  if (m.slab\_top\_slide\_index \neq MGRID.slab\_top\_slide\_index) {
    if (Debug(DEBUG\_GRID)) fprintf(stderr, "GRID: \_Fill\_top\_slide\_refractive\_index\_changed n");
    return (FALSE);
  if (m.slab\_bottom\_slide\_index \neq MGRID.slab\_bottom\_slide\_index) {
    if (Debug(DEBUG_GRID))
       fprintf(stderr, "GRID: □Fill □bottom □slide □refractive □index □changed \n");
    return (FALSE);
       Routine to just figure out the distance to a particular a, b, g point
\langle \text{ Prototype for } abg\_distance | 142 \rangle \equiv
  void abg\_distance(double \ a, double \ b, double \ g, guess\_type *guess)
This code is used in sections 112 and 143.
```

66 GRID ROUTINES IAD (v 3-12-0) $\S143$

```
\langle \text{ Definition for } abg\_distance | 143 \rangle \equiv
  \langle \text{ Prototype for } abg\_distance \ 142 \rangle
     double m_{-}r, m_{-}t, distance;
     struct measure_type old_mm;
     struct invert_type old_rr;
     Get\_Calc\_State(\&old\_mm, \&old\_rr);
     RR.slab.a = a;
     RR.slab.b = b;
     RR.slab.g = g;
     Calculate\_Distance(\&m\_r,\&m\_t,\&distance);
     Set\_Calc\_State(old\_mm, old\_rr);
     guess \neg a = a;
     guess \rightarrow b = b;
     guess \neg g = g;
     guess \neg distance = distance;
This code is used in section 111.
```

144. This just searches through the grid to find the minimum entry and returns the optical properties of that entry. The smallest, the next smallest, and the third smallest values are returned.

This has been rewritten to use Calculate_Distance_With_Corrections so that changes in sphere parameters won't necessitate recalculating the grid.

```
\langle \text{Prototype for } \textit{Near\_Grid\_Points } 144 \rangle \equiv  void \textit{Near\_Grid\_Points}(\textbf{double } r, \textbf{double } t, \textbf{search\_type } s, \textbf{int } *i\_min, \textbf{int } *j\_min) This code is used in sections 112 and 145.
```

```
145.
        \langle \text{ Definition for } Near\_Grid\_Points | 145 \rangle \equiv
  ⟨ Prototype for Near_Grid_Points 144⟩
     int i, j;
     double fval;
     double smallest = 10.0;
     struct measure_type old_mm;
     struct invert_type old_rr;
     Get\_Calc\_State(\&old\_mm,\&old\_rr);
     *i_{-}min = 0;
     *j_{-}min = 0;
     for (i = 0; i < GRID\_SIZE; i++) {
       for (j = 0; j < GRID\_SIZE; j \leftrightarrow) {
         CALCULATING\_GRID = 1;
         fval = Calculate\_Grid\_Distance(i, j);
         CALCULATING\_GRID = 0;
         if (fval < smallest) {
            *i\_min = i;
            *j_min = j;
            smallest = fval;
     Set\_Calc\_State(old\_mm, old\_rr);
This code is used in section 111.
```

146. Routine to incorporate flipping of sample if needed. This is pretty simple. The assumption is that flipping is handled relative to the reflection side of the sphere. Thus even when flipping is needed, the usual call to RT() will result in the correct values for the reflectances. The transmission values can then be calculated by swapping the top and bottom slides.

Technically, the value of slab should be **const** but it is not so that we don't pay a copying overhead whenever *flip* is false (the usual case).

```
\langle \text{Prototype for } RT\_Flip \mid 146 \rangle \equiv 
void RT\_Flip (\text{int } flip, \text{int } n, \text{struct } AD\_slab\_type *slab, \text{double *UR1}, \text{double *UT1}, \text{double *UTU})
This code is used in section 147.
```

68 GRID ROUTINES IAD (v 3-12-0) $\S147$

```
147. \langle Definition for RT_{-}Flip_{147}\rangle \equiv
   \langle \text{ Prototype for } RT\_Flip \ 146 \rangle
      double swap, correct\_UR1, correct\_URU;
      RT(n, slab, UR1, UT1, URU, UTU);
      if (flip) {
         correct_{-}UR1 = *UR1;
         correct_{-}URU = *URU;
         swap = slab \neg n\_top\_slide;
         slab \neg n\_top\_slide = slab \neg n\_bottom\_slide;
         slab \rightarrow n\_bottom\_slide = swap;
         swap = slab \rightarrow b\_top\_slide;
         slab \rightarrow b\_top\_slide = slab \rightarrow b\_bottom\_slide;
         slab \rightarrow b_-bottom\_slide = swap;
         RT(n, slab, UR1, UT1, URU, UTU);
         swap = slab \neg n\_top\_slide;
         slab \neg n\_top\_slide = slab \neg n\_bottom\_slide;
         slab \neg n\_bottom\_slide = swap;
         swap = slab \rightarrow b_top_slide;
         slab \rightarrow b\_top\_slide = slab \rightarrow b\_bottom\_slide;
         slab \rightarrow b\_bottom\_slide = swap;
         *UR1 = correct_UR1;
         *URU = correct_{-}URU;
```

This code is used in section 111.

```
148.
        Simple routine to put values into the grid
  Presumes that RR. slab is properly set up.
\langle \text{ Definition for } fill\_grid\_entry 148 \rangle \equiv
  static void fill_grid_entry(int i, int j)
     double ur1, ut1, uru, utu;
     if (RR.slab.b \le 1 \cdot 10^{-6}) RR.slab.b = 1 \cdot 10^{-6};
     if (Debug(DEBUG_EVERY_CALC)) {
       if (¬CALCULATING_GRID)
          fprintf(stderr, "a=\%8.5f_{\sqcup}b=\%10.5f_{\sqcup}g=\%8.5f_{\sqcup}", RR.slab.a, RR.slab.b, RR.slab.g);
          if (j \equiv 0) fprintf (stderr, ".");
          if (i + 1 \equiv GRID\_SIZE \land j \equiv 0) fprintf (stderr, "\n");
        }
     RT_Flip (MM.flip_sample, RR.method.quad_pts, &RR.slab, &ur1, &ut1, &utu, &utu);
     if (Debug(DEBUG\_EVERY\_CALC) \land \neg CALCULATING\_GRID)
        fprintf(stderr, "ur1=\%8.5f_ut1=\%8.5f_n", ur1, ut1);
     The\_Grid[GRID\_SIZE * i + j][A\_COLUMN] = RR.slab.a;
     The\_Grid[GRID\_SIZE * i + j][B\_COLUMN] = RR.slab.b;
     The\_Grid[GRID\_SIZE * i + j][G\_COLUMN] = RR.slab.g;
     The\_Grid[GRID\_SIZE*i+j][UR1\_COLUMN] = ur1;
     The\_Grid[GRID\_SIZE * i + j][UT1\_COLUMN] = ut1;
     The\_Grid[GRID\_SIZE * i + j][URU\_COLUMN] = uru;
     The\_Grid[GRID\_SIZE * i + j][UTU\_COLUMN] = utu;
     if (Debug(DEBUG_GRID_CALC)) {
       \mathit{fprintf}\,(\mathit{stderr}\,, \verb"+_{\sqcup}\nspace \verb"2d_{\sqcup}\nspace", i, j);
       fprintf(stderr, "\%10.5f_{\square}\%10.5f_{\square}\%10.5f_{\square}", RR.slab.a, RR.slab.b, RR.slab.g);
       fprintf(stderr, "\%10.5f_{\square}\%10.5f_{\square}| ", MM.m_r, uru);
       fprintf(stderr, "\%10.5f_{\square}\%10.5f_{\square}\n", MM.m_t, utu);
This code is used in section 111.
```

149. This routine fills the grid with a proper set of values. With a little work, this routine could be made much faster by (1) only generating the phase function matrix once, (2) Making only one pass through the array for each albedo value, i.e., using the matrix left over from b = 1 to generate the solution for b = 2. Unfortunately this would require a complete revision of the *Calculate_Distance* routine. Fortunately, this routine should only need to be calculated once at the beginning of each run.

```
\langle \text{Prototype for } Fill\_AB\_Grid \ 149 \rangle \equiv 
void Fill\_AB\_Grid (\text{struct measure\_type } m, \text{struct invert\_type } r)
This code is used in sections 111 and 150.
```

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```
150.
        \langle \text{ Definition for } Fill\_AB\_Grid \ 150 \rangle \equiv
   \langle Prototype for Fill\_AB\_Grid 149 \rangle
     int i, j;
     double a;
     double min_{-}b = -8;
                                  /* \exp(-10) is smallest thickness */
     double max_b = +8:
                                  /* \exp(+8) is greatest thickness */
     if (Debug(Debug(DEBUG_GRID))) fprintf(stderr, "Filling_AB_grid\n");
     if (The\_Grid \equiv \Lambda) Allocate\_Grid(r.search);
     \langle \text{ Zero GG } 156 \rangle
     Set\_Calc\_State(m,r);
     GG_{-}g = RR.slab.g;
     for (i = 0; i < GRID\_SIZE; i++) {
        double x = (\mathbf{double}) i/(\mathtt{GRID\_SIZE} - 1.0);
       RR.slab.b = exp(min\_b + (max\_b - min\_b) * x);
        for (j = 0; j < GRID\_SIZE; j \leftrightarrow) {
          ⟨Generate next albedo using j 152⟩
          fill\_grid\_entry(i, j);
     The\_Grid\_Initialized = TRUE;
     The\_Grid\_Search = FIND\_AB;
This code is used in section 111.
```

151. Now it seems that I must be a bit more subtle in choosing the range of albedos to use in the grid. Originally I just spaced them according to

$$a = 1 - \left[\frac{j-1}{n-1}\right]^3$$

where $1 \le j \le n$. Long ago it seems that I based things only on the square of the bracketed term, but I seem to remember that I was forced to change it from a square to a cube to get more global convergence.

So why am I rewriting this? Well, because it works very poorly for samples with small albedos. For example, when n = 11 then the values chosen for a are (1, .999, .992, .973, .936, .875, .784, .657, .488, .271, 0). Clearly very skewed towards high albedos.

I am considering a two part division. I'm not too sure how it should go. Let the first half be uniformly divided and the last half follow the cubic scheme given above. The list of values should then be (1, .996, .968, .892, 0.744, .5, .4, .3, .2, .1, 0).

Maybe it would be best if I just went back to a quadratic term. Who knows?

In the **if** statement below, note that it could read $j \geq k$ and still generate the same results.

```
 \begin{split} &\langle \, \text{Nonworking code 151} \, \rangle \equiv \\ &k = floor((\text{GRID\_SIZE} - 1)/2); \\ &\text{if } (j > k) \, \, \{ \\ &a = 0.5 * (1 - (j - k - 1)/(\text{GRID\_SIZE} - k - 1)); \\ &\text{RR.} slab.a = a; \\ &\} \\ &\text{else } \{ \\ &a = (j - 1.0)/(\text{GRID\_SIZE} - k - 1); \\ &\text{RR.} slab.a = 1.0 - a * a * a/2; \\ &\} \end{split}
```

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152. Well, the above code did not work well. So I futzed around and sort of empirically ended up using the very simple method below. The only real difference from the previous method what that the method is now quadratic and not cubic.

```
 \begin{split} &\langle \, \text{Generate next albedo using j } 152 \, \rangle \equiv \\ &a = (\mathbf{double}) \ j/(\mathtt{GRID\_SIZE} - 1.0); \\ &\mathbf{if} \ (a < 0.25) \ \mathtt{RR}.slab.a = 1.0 - a*a; \\ &\mathbf{else} \ \mathbf{if} \ (a > 0.75) \ \mathtt{RR}.slab.a = (1.0 - a)*(1.0 - a); \\ &\mathbf{else} \ \mathtt{RR}.slab.a = 1 - a; \end{split}  See also section 153.
```

§152

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This code is used in sections 150 and 155.

153. Well, the above code has gaps. Here is an attempt to eliminate the gaps

```
 \begin{split} &\langle \, \text{Generate next albedo using j 152} \, \rangle + \equiv \\ &a = (\mathbf{double}) \ j / (\texttt{GRID\_SIZE} - 1.0); \\ &\texttt{RR}.slab.a = (1.0 - a*a)*(1.0 - a) + (1.0 - a)*(1.0 - a)*a; \end{split}
```

154. This is quite similar to $Fill_AB_Grid$, with the exception of the little shuffle I do at the beginning to figure out the optical thickness to use. The problem is that the optical thickness may not be known. If it is known then the only way that we could have gotten here is if the user dictated FIND_AG and specified b and only provided two measurements. Otherwise, the user must have made three measurements and the optical depth can be figured out from $m.m_u$.

This routine could also be improved by not recalculating the anisotropy matrix for every point. But this would only end up being a minor performance enhancement if it were fixed.

```
\langle \text{ Prototype for } Fill\_AG\_Grid \ 154 \rangle \equiv
  void Fill_-AG_-Grid(struct measure_type m, struct invert_type r)
This code is used in sections 111 and 155.
155. \langle \text{ Definition for } Fill\_AG\_Grid \ 155 \rangle \equiv
   \langle \text{Prototype for } Fill\_AG\_Grid \ 154 \rangle
     int i, j;
     double a;
     if (Debug(Debug(DEBUG_GRID))) fprintf(stderr, "Filling_AG_grid\n");
     if (The\_Grid \equiv \Lambda) Allocate\_Grid(r.search);
     \langle \text{ Zero GG } 156 \rangle
     Set\_Calc\_State(m, r);
     GG_{-}b = r.slab.b;
     for (i = 0; i < GRID\_SIZE; i++) {
        RR.slab.g = 0.9999 * (2.0 * i/(GRID\_SIZE - 1.0) - 1.0);
        for (j = 0; j < GRID\_SIZE; j \leftrightarrow) {
           (Generate next albedo using j 152)
           fill\_grid\_entry(i, j);
      The\_Grid\_Initialized = TRUE;
      The\_Grid\_Search = FIND\_AG;
This code is used in section 111.
```

72 GRID ROUTINES IAD (v 3-12-0) $\S156$

```
156.  \langle \operatorname{Zero} \ \mathsf{GG} \ 156 \rangle \equiv \\ GG_{-}a = 0.0; \\ GG_{-}b = 0.0; \\ GG_{-}g = 0.0; \\ GG_{-}bs = 0.0; \\ GG_{-}ba = 0.0; \\ GG_{-}ba = 0.0; \\ \text{This code is used in sections 150, 155, 158, 160, and 162.}
```

157. This is quite similar to $Fill_AB_Grid$, with the exception of the that the albedo is held fixed while b and g are varied.

This routine could also be improved by not recalculating the anisotropy matrix for every point. But this would only end up being a minor performance enhancement if it were fixed.

```
\langle \text{ Prototype for } Fill\_BG\_Grid \ 157 \rangle \equiv
   void Fill_BG_Grid(struct measure_type m, struct invert_type r)
This code is used in sections 112 and 158.
       \langle \text{ Definition for } Fill\_BG\_Grid 158 \rangle \equiv
   \langle Prototype for Fill\_BG\_Grid 157 \rangle
     int i, j;
     if (The\_Grid \equiv \Lambda) Allocate\_Grid(r.search);
      \langle \text{ Zero GG } 156 \rangle
     if (Debug(Debug(DEBUG\_GRID))) fprintf(stderr, "Filling_{\square}BG_{\square}grid\n");
     Set\_Calc\_State(m, r);
     RR.slab.b = 1.0/32.0;
     RR.slab.a = RR.default_a;
     GG_{-}a = RR.slab.a;
     for (i = 0; i < GRID\_SIZE; i \leftrightarrow) {
        RR.slab.b *= 2;
        for (j = 0; j < GRID\_SIZE; j \leftrightarrow) {
           RR.slab.g = 0.9999 * (2.0 * j/(GRID\_SIZE - 1.0) - 1.0);
           fill\_grid\_entry(i, j);
        }
      The\_Grid\_Initialized = TRUE;
      The\_Grid\_Search = FIND\_BG;
This code is used in section 111.
```

159. This is quite similar to $Fill_BG_Grid$, with the exception of the that the $b_s = \mu_s d$ is held fixed. Here b and g are varied on the usual grid, but the albedo is forced to take whatever value is needed to ensure that the scattering constant remains fixed.

```
\langle \text{Prototype for } Fill\_BaG\_Grid \ 159 \rangle \equiv  void Fill\_BaG\_Grid (\text{struct measure\_type } m, \text{struct invert\_type } r) This code is used in sections 112 and 160.
```

```
§160
         IAD (v 3-12-0)
160.
        \langle \text{ Definition for } Fill\_BaG\_Grid \ 160 \rangle \equiv
  \langle Prototype for Fill\_BaG\_Grid 159 \rangle
     int i, j;
     double bs, ba;
     if (The\_Grid \equiv \Lambda) Allocate\_Grid(r.search);
     \langle \text{ Zero GG } 156 \rangle
     if (Debug(Debug(DEBUG_GRID))) fprintf(stderr, "Filling_BaG_grid\n");
     Set\_Calc\_State(m, r);
     ba = 1.0/32.0;
     bs = RR.default\_bs;
     GG_{-}bs = bs;
     for (i = 0; i < GRID\_SIZE; i++) {
       ba *= 2;
       ba = exp((double) i/(GRID\_SIZE - 1.0) * log(1024.0))/16.0;
       RR.slab.b = ba + bs;
       if (RR.slab.b > 0) RR.slab.a = bs/RR.slab.b;
       else RR.slab.a = 0;
       for (j = 0; j < GRID\_SIZE; j \leftrightarrow) {
          RR.slab.g = 0.9999 * (2.0 * j/(GRID\_SIZE - 1.0) - 1.0);
          fill\_grid\_entry(i, j);
     The\_Grid\_Initialized = \texttt{TRUE};
     The\_Grid\_Search = FIND\_BaG;
This code is used in section 111.
161. Very similar to the above routine. The value of b_a = \mu_a d is held constant.
\langle \text{ Prototype for } Fill\_BsG\_Grid \ 161 \rangle \equiv
  void Fill_BsG_Grid(struct measure_type m, struct invert_type r)
```

This code is used in sections 112 and 162.

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```
\langle \text{ Definition for } Fill\_BsG\_Grid \ 162 \rangle \equiv
   \langle Prototype for Fill\_BsG\_Grid 161 \rangle
     int i, j;
      double bs, ba;
     if (The\_Grid \equiv \Lambda) Allocate\_Grid(r.search);
      \langle \text{ Zero GG } 156 \rangle
      Set\_Calc\_State(m, r);
      bs = 1.0/32.0;
      ba = RR.default_ba;
      GG_{-}ba = ba;
       \mathbf{for} \ (i=0; \ i < \mathtt{GRID\_SIZE}; \ i +\!\!\!+\!\!\!) \ \big\{ 
        bs *= 2;
        RR.slab.b = ba + bs;
        if (RR.slab.b > 0) RR.slab.a = bs/RR.slab.b;
        else RR.slab.a = 0;
        for (j = 0; j < GRID\_SIZE; j \leftrightarrow) {
           RR.slab.g = 0.9999 * (2.0 * j/(GRID_SIZE - 1.0) - 1.0);
           fill\_grid\_entry(i, j);
      The\_Grid\_Initialized = TRUE;
      The\_Grid\_Search = FIND\_BsG;
This code is used in section 111.
163. \langle \text{ Prototype for } Fill\_Grid \ 163 \rangle \equiv
   void Fill\_Grid (struct measure_type m, struct invert_type r, int force\_new)
This code is used in sections 112 and 164.
```

```
164.
       \langle \text{ Definition for } Fill\_Grid \ 164 \rangle \equiv
  \langle \text{ Prototype for } Fill\_Grid \ 163 \rangle
    if (force\_new \lor \neg Same\_Calc\_State(m, r)) {
      switch (r.search) {
      case FIND_AB:
         if (Debug(DEBUG_SEARCH)) fprintf(stderr, "filling_AB_Grid\n");
         Fill\_AB\_Grid(m,r);
         break;
       case FIND_AG:
         if (Debug(DEBUG_SEARCH)) fprintf(stderr, "filling AGGGrid\n");
         Fill\_AG\_Grid(m,r);
         break;
      case FIND_BG:
         if (Debug(DEBUG_SEARCH)) fprintf(stderr, "filling_BG_Grid\n");
         Fill_BG_Grid(m,r);
         break;
      case FIND\_BaG:
         if (Debug(DEBUG_SEARCH)) fprintf(stderr, "filling_BaG_Grid\n");
         Fill\_BaG\_Grid(m,r);
         break;
      case FIND\_BsG:
         if (Debug(DEBUG_SEARCH)) fprintf(stderr, "filling_BsG_Grid\n");
         Fill\_BsG\_Grid(m,r);
         break;
      default: AD_error("Attempt to fill grid for unusual search case.");
    Get_Calc_State(&MGRID, &RGRID);
```

165. Calculating R and T.

This code is used in section 111.

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IAD (v 3-12-0)

Calculate_Distance returns the distance between the measured values in MM and the calculated values for the current guess at the optical properties. It assumes that the everything in the local variables MM and RR have been set appropriately.

```
⟨ Prototype for Calculate_Distance 165 ⟩ ≡ void Calculate_Distance (double *M_R, double *M_T, double *deviation)
This code is used in sections 112 and 166.
```

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```
166.
        \langle \text{ Definition for } Calculate\_Distance | 166 \rangle \equiv
  (Prototype for Calculate_Distance 165)
     double Rc, Tc, ur1, ut1, uru, utu;
     if (RR.slab.b \le 1 \cdot 10^{-6}) RR.slab.b = 1 \cdot 10^{-6};
     if (Debug(DEBUG_EVERY_CALC))
       fprintf(stderr, "a=\%8.5f_b=\%10.5f_g=\%8.5f_", RR.slab.a, RR.slab.b, RR.slab.g);
     RT_Flip(MM.flip\_sample, RR.method.quad\_pts, \&RR.slab, \&ur1, \&ut1, \&uru, \&utu);
     if (Debug(DEBUG_EVERY_CALC))
        fprintf(stderr, "ur1=\%8.5f_{\sqcup}ut1=\%8.5f_{\sqcup}(not_{\sqcup}M_{\perp}R_{\sqcup}and_{\sqcup}M_{\perp}T!)\n", ur1, ut1);
     Sp\_mu\_RT\_Flip (MM. flip\_sample, RR. slab .n\_top\_slide, RR. slab .n\_slab, RR. slab .n\_bottom\_slide,
          RR.slab.b\_top\_slide, RR.slab.b, RR.slab.b\_bottom\_slide, RR.slab.cos\_angle, & Rc, & Tc);
     if ((\neg \texttt{CALCULATING\_GRID} \land Debug(\texttt{DEBUG\_ITERATIONS})) \lor (\texttt{CALCULATING\_GRID} \land 
             Debug(DEBUG\_GRID\_CALC))) fprintf(stderr, "_____");
     Calculate_Distance_With_Corrections(ur1, ut1, Rc, Tc, uru, utu, M_R, M_T, deviation);
This code is used in section 111.
167. \langle \text{Prototype for } Calculate\_Grid\_Distance | 167 \rangle \equiv
  double Calculate\_Grid\_Distance(int i, int j)
This code is used in sections 112 and 168.
       \langle Definition for Calculate\_Grid\_Distance 168 \rangle \equiv
   ⟨ Prototype for Calculate_Grid_Distance 167⟩
     double ur1, ut1, uru, utu, Rc, Tc, b, dev, LR, LT;
     if (Debug(DEBUG\_GRID\_CALC)) fprintf(stderr, "g_{\parallel})/(2d_{\parallel})/(2d_{\parallel});
     b = The\_Grid[GRID\_SIZE * i + j][B\_COLUMN];
     ur1 = The\_Grid[GRID\_SIZE * i + j][UR1\_COLUMN];
     ut1 = The\_Grid[GRID\_SIZE * i + j][UT1\_COLUMN];
     uru = The\_Grid[GRID\_SIZE * i + j][URU\_COLUMN];
     utu = The\_Grid[GRID\_SIZE * i + j][UTU\_COLUMN];
     RR.slab.a = The\_Grid[GRID\_SIZE * i + j][A\_COLUMN];
     RR.slab.b = The\_Grid[GRID\_SIZE * i + j][B\_COLUMN];
     RR.slab.g = The\_Grid[GRID\_SIZE * i + j][G\_COLUMN];
     Sp\_mu\_RT\_Flip (MM. flip\_sample, RR. slab. n\_top\_slide, RR. slab. n\_slab, RR. slab. n\_bottom\_slide,
          RR.slab.b\_top\_slide, b, RR.slab.b\_bottom\_slide, RR.slab.cos\_angle, & Rc, & Tc);
     CALCULATING\_GRID = 1;
     Calculate_Distance_With_Corrections(ur1, ut1, Rc, Tc, uru, utu, &LR, &LT, &dev);
     CALCULATING\_GRID = 0;
     return dev;
This code is used in section 111.
```

CALCULATING R AND T 77

Rc and Tc refer to the unscattered (collimated) reflection and transmission.

The only tricky part is to remember that the we are trying to match the measured values. The measured values are affected by sphere parameters and light loss. Since the values UR1 and UT1 are for an infinite slab sample with no light loss, the light loss out the edges must be subtracted. It is these values that are used with the sphere formulas to convert the modified UR1 and UT1 to values for *M_R and *M_T.

This code is used in sections 112 and 170.

```
\langle Definition for Calculate\_Distance\_With\_Corrections 170 \rangle \equiv
  ⟨ Prototype for Calculate_Distance_With_Corrections 169⟩
     double R_direct, T_direct, R_diffuse, T_diffuse;
     R\_diffuse = URU - MM.uru\_lost;
     T_{-}diffuse = UTU - MM.utu_{-}lost;
     R\_direct = UR1 - MM.ur1\_lost - (1.0 - MM.fraction\_of\_rc\_in\_mr) * Rc;
     T\_direct = \mathtt{UT1} - \mathtt{MM}.ut1\_lost - (1.0 - \mathtt{MM}.fraction\_of\_tc\_in\_mt) * Tc;
     switch (MM. num_spheres) {
     case 0: (Calc M_R and M_T for no spheres 171)
       break:
     case 1: case -2:
       if (MM.method \equiv COMPARISON) (Calc M_R and M_T for dual beam sphere 173)
       else (Calc M_R and M_T for single beam sphere 172)
     case 2: (Calc M_R and M_T for two spheres 174)
       break:
     (Calculate the deviation 175)
     (Print diagnostics 178)
This code is used in section 111.
```

171. If no spheres were used in the measurement, then presumably the measured values are the reflection and transmission. Consequently, we just acertain what the irradiance was and whether the specular reflection ports were blocked and proceed accordingly. Note that blocking the ports does not have much meaning unless the light is collimated, and therefore the reflection and transmission is only modified for collimated irradiance.

```
\langle Calc M_R and M_T for no spheres 171\rangle \equiv *M_R = R-direct; *M_T = T-direct; This code is used in section 170.
```

172. The direct incident power is (1-f)P. The reflected power will be $(1-f)r_s^{\text{direct}}P$. Since baffles ensure that the light cannot reach the detector, we must bounce the light off the sphere walls to use to above gain formulas. The contribution will then be $(1-f)r_s^{\text{direct}}(1-a_e)r_wP$. The measured power will be

$$P_d = a_d(1 - a_e)r_w[(1 - f)r_s^{\text{direct}} + fr_w]P \cdot G(r_s)$$

Similarly the power falling on the detector measuring transmitted light is

$$P'_d = a'_d t_s^{\text{direct}} r'_w (1 - a'_e) P \cdot G'(r_s)$$

when the 'entrance' port in the transmission sphere is closed, $a_e'=0$. The normalized sphere measurements are

$$M_R = r_{\text{std}} \cdot \frac{R(r_s^{\text{direct}}, r_s) - R(0, 0)}{R(r_{\text{std}}, r_{\text{std}}) - R(0, 0)}$$

and

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$$M_T = t_{\text{std}} \cdot \frac{T(t_s^{\text{direct}}, r_s) - T(0, 0)}{T(t_{\text{std}}, r_{\text{std}}) - T(0, 0)}$$

```
 \left\{ \begin{array}{l} \text{ double $P$\_std$, $P$\_d$, $P$\_0$;} \\ \text{ double $P$\_std$, $P$\_d$, $GP$\_std$, $GP$\_std$, $GP$;} \\ \text{ G$\_0$ = $Gain$(REFLECTION\_SPHERE, MM, $0.0$);} \\ G=Gain$(REFLECTION\_SPHERE, MM, $R$\_diffuse$);} \\ G\_std=Gain$(REFLECTION\_SPHERE, MM, R$\_diffuse$);} \\ P\_d=G*(R\_direct*(1-MM.f\_r)+MM.f\_r*MM.rw\_r);} \\ P\_std=G*std*(MM.rstd\_r*(1-MM.f\_r)+MM.f\_r*MM.rw\_r);} \\ P\_0=G\_0*(MM.f\_r*MM.rw\_r);} \\ *M\_R=MM.rstd\_r*(P\_d-P\_0)/(P\_std-P\_0);} \\ \text{GP}=Gain$(TRANSMISSION\_SPHERE, MM, $R$\_diffuse$);} \\ GP\_std=Gain$(TRANSMISSION\_SPHERE, MM, 0.0);} \\ *M\_T=T\_direct*GP/GP\_std; \\ \right\}
```

This code is used in section 170.

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The dual beam case is different because the sphere efficiency is equivalent for measurement of light hitting the sample first or hitting the reference standard first. The dual beam measurement should report the ratio of these two reflectance measurements, thereby eliminating the need to calculate the gain completely. The same holds when no sample is present.

The normalized reflectance measurement (the difference between dual beam measurement for a port with the sample and with nothing) is

$$M_R = r_{\rm std} \cdot \frac{(1-f)r_s^{\rm direct} + fr_w}{(1-f')r_{\rm std} - f'r_w} - r_{\rm std} \cdot \frac{(1-f)(0) + fr_w}{(1-f')r_{\rm std} - f'r_w}$$

or

$$M_R = \frac{(1-f)r_s^{\text{direct}}}{(1-f') - f'r_w/r_{\text{std}}}$$

When f = f' = 1, then $M_R = 1$ no matter what the reflectance is. (Leave it in this form to avoid division by zero when f = 1.)

The normalized transmittance is simply t_s^{direct} .

When f = 0 then this result is essentially the same as the no spheres result (because no sphere corrections are needed). However if the number of spheres is zero, then no lost light calculations are made and therefore that is a potential error.

```
\langle \text{Calc M}_R \text{ and M}_T \text{ for dual beam sphere } 173 \rangle \equiv
      *M_R = (1 - MM.f_r) * R_direct/((1 - MM.f_r) + MM.f_r * MM.rw_r/MM.rstd_r);
      *M_T = T_direct;
```

This code is used in section 170.

174. When two integrating spheres are present then the double integrating sphere formulas are slightly more complicated.

I am not sure what it means when $rstd_{-}t$ is not unity.

The normalized sphere measurements for two spheres are

$$M_R = \frac{R(r_s^{\text{direct}}, r_s, t_s^{\text{direct}}, t_s) - R(0, 0, 0, 0)}{R(r_{\text{std}}, r_{\text{std}}, 0, 0) - R(0, 0, 0, 0)}$$

and

$$M_T = \frac{T(r_s^{\text{direct}}, r_s, t_s^{\text{direct}}, t_s) - T(0, 0, 0, 0)}{T(0, 0, 1, 1) - T(0, 0, 0, 0)}$$

Note that R_0 and T_0 will be zero unless one has explicitly set the fraction $m.f_r$ ore $m.f_r$ to be non-zero. $\langle \text{Calc M}_R \text{ and M}_T \text{ for two spheres } 174 \rangle \equiv$

```
{
  double R_0, T_0;
 R_0 = Two_Sphere_R(MM, 0, 0, 0, 0);
  T_0 = Two_Sphere_T(MM, 0, 0, 0, 0, 0);
  *M_R = MM.rstd_r * (Two_Sphere_R(MM, R_direct, R_diffuse, T_direct,
      T_{\text{-}}diffuse) - R_{\text{-}}0)/(Two_{\text{-}}Sphere_{\text{-}}R(MM, MM.rstd_{\text{-}}r, MM.rstd_{\text{-}}r, 0, 0) - R_{\text{-}}0);
  1) - T_0);
```

This code is used in section 170.

175. There are at least three things that need to be considered here. First, the number of measurements. Second, is the metric is relative or absolute. And third, is the albedo fixed at zero which means that the transmission measurement should be used instead of the reflection measurement.

```
⟨ Calculate the deviation 175⟩ ≡

if (RR.search ≡ FIND_A ∨ RR.search ≡ FIND_G ∨ RR.search ≡ FIND_B ∨ RR.search ≡ FIND_Bs ∨ RR.search ≡

FIND_Ba) {

⟨ One parameter deviation 176⟩

}

else {

⟨ Two parameter deviation 177⟩

}

This code is used in section 170.
```

176. This part was slightly tricky. The crux of the problem was to decide if the transmission or the reflection was trustworthy. After looking a bunches of measurements, I decided that the transmission measurement was almost always more reliable. So when there is just a single measurement known, then use the total transmission if it exists.

```
 \begin{split} &\langle \, \text{One parameter deviation 176} \, \rangle \equiv \\ & \quad \text{if } \, (\texttt{MM}.m\_t > 0) \, \, \{ \\ & \quad \text{if } \, (\texttt{RR}.metric \equiv \texttt{RELATIVE}) \, *dev = fabs(\texttt{MM}.m\_t - *\texttt{M\_T})/(\texttt{MM}.m\_t + \texttt{ABIT}); \\ & \quad \text{else } \, *dev = fabs(\texttt{MM}.m\_t - *\texttt{M\_T}); \\ & \quad \} \\ & \quad \text{else } \, \{ \\ & \quad \text{if } \, (\texttt{RR}.metric \equiv \texttt{RELATIVE}) \, *dev = fabs(\texttt{MM}.m\_r - *\texttt{M\_R})/(\texttt{MM}.m\_r + \texttt{ABIT}); \\ & \quad \text{else } \, *dev = fabs(\texttt{MM}.m\_r - *\texttt{M\_R}); \\ & \quad \} \end{split}  This code is used in section 175.
```

177. This stuff happens when we are doing two parameter searches. In these cases there should be information in both R and T. The distance should be calculated using the deviation from both. The albedo stuff might be able to be take out. We'll see.

```
 \begin{array}{l} \langle \, {\rm Two \; parameter \; deviation \; 177} \, \rangle \equiv \\ & \quad {\rm if \; (RR.} metric \equiv {\rm RELATIVE}) \; \left\{ \\ & \quad *dev = 0; \\ & \quad {\rm if \; (MM.} m_-t > {\rm ABIT}) \; *dev = {\rm T\_TRUST\_FACTOR} * fabs({\rm MM.} m_-t - *{\rm M\_T})/({\rm MM.} m_-t + {\rm ABIT}); \\ & \quad {\rm if \; (RR.} default_-a \neq 0) \; *dev \; += fabs({\rm MM.} m_-r - *{\rm M\_R})/({\rm MM.} m_-r + {\rm ABIT}); \\ & \quad {\rm else \; \{} \\ & \quad *dev = {\rm T\_TRUST\_FACTOR} * fabs({\rm MM.} m_-t - *{\rm M\_T}); \\ & \quad {\rm if \; (RR.} default_-a \neq 0) \; *dev \; += fabs({\rm MM.} m_-r - *{\rm M\_R}); \\ & \quad {\rm \}} \end{array}
```

This code is used in section 175.

183. $\langle \text{Prototype for } Find_Ba_fn \ 183 \rangle \equiv$ double $Find_Ba_fn(\text{double } x)$ This code is used in sections 112 and 184.

This is here so that I can figure out why the program is not converging. This is a little convoluted so that the global constants at the top of this file interact properly. $\langle \text{Print diagnostics } 178 \rangle \equiv$ if $((Debuq(DEBUG_ITERATIONS) \land \neg CALCULATING_GRID) \lor (Debuq(DEBUG_GRID_CALC) \land CALCULATING_GRID))$ static int once = 0; if $(once \equiv 0)$ { $fprintf(stderr, "\%10s_{\sqcup}\%10s_{\sqcup}\%10s_{\sqcup}\%10s_{\sqcup})\%10s_{\sqcup}\%10s_{\sqcup}\%10s_{\sqcup}\%10s_{\sqcup}\%10s_{\sqcap}, "a", "b", "g", "m_r", "fit", %fit", %fit"$ "m_t", "fit", "delta"); once = 1; $fprintf(stderr, "\%10.5f_{\square}\%10.5f_{\square}\%10.5f_{\square}", RR.slab.a, RR.slab.b, RR.slab.g);$ $fprintf(stderr, "\%10.5f_{\square}\%10.5f_{\square}|", MM.m_r, *M_R);$ $fprintf(stderr, "\%10.5f_{\square}\%10.5f_{\square}|", MM.m_{-}t, *M_T);$ $fprintf(stderr, "\%10.5f_{\square}\n", *dev);$ This code is used in section 170. $\langle \text{Prototype for } Find_AG_fn \mid 179 \rangle \equiv$ **double** $Find_AG_fn($ **double** x[])This code is used in sections 112 and 180. **180.** $\langle \text{ Definition for } Find_AG_fn \text{ 180} \rangle \equiv$ $\langle \text{ Prototype for } Find_AG_fn \ 179 \rangle$ **double** $m_{-}r$, $m_{-}t$, deviation; RR.slab.a = acalc2a(x[1]);RR.slab.g = gcalc2g(x[2]); $Calculate_Distance(\&m_r,\&m_t,\&deviation);$ return deviation; This code is used in section 111. **181.** $\langle \text{Prototype for } Find_AB_fn \ 181 \rangle \equiv$ **double** $Find_AB_fn($ **double** x[])This code is used in sections 112 and 182. **182.** $\langle \text{ Definition for } Find_AB_fn \text{ 182} \rangle \equiv$ $\langle \text{ Prototype for } Find_AB_fn \text{ 181} \rangle$ **double** $m_{-}r$, $m_{-}t$, deviation; RR.slab.a = acalc2a(x[1]);RR.slab.b = bcalc2b(x[2]); $Calculate_Distance(\&m_r,\&m_t,\&deviation);$ **return** deviation; This code is used in section 111.

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This is tricky only because the value in RR.slab.b is used to hold the value of bs or $d \cdot \mu_s$. It must be switched to the correct value for the optical thickness and then switched back at the end of the routine. $\langle \text{ Definition for } Find_Ba_fn \ 184 \rangle \equiv$ $\langle \text{ Prototype for } Find_Ba_fn \text{ 183} \rangle$ **double** $m_{-}r$, $m_{-}t$, deviation, ba, bs; bs = RR.slab.b;ba = bcalc2b(x);RR.slab.b = ba + bs;/* unswindle */ RR.slab.a = bs/(ba + bs); $Calculate_Distance(\&m_r,\&m_t,\&deviation);$ /* swindle */ RR.slab.b = bs;**return** deviation; This code is used in section 111. See the comments for the Find_Ba_fn routine above. Play the same trick but use ba. $\langle \text{ Prototype for } Find_Bs_fn \mid 185 \rangle \equiv$ **double** $Find_{-}Bs_{-}fn($ **double** x)This code is used in sections 112 and 186. **186.** $\langle \text{ Definition for } Find_Bs_fn \ 186 \rangle \equiv$ $\langle \text{ Prototype for } Find_Bs_fn \text{ 185} \rangle$ **double** $m_{-}r$, $m_{-}t$, deviation, ba, bs; ba = RR.slab.b;/* unswindle */ bs = bcalc2b(x);RR.slab.b = ba + bs;RR.slab.a = bs/(ba + bs); $Calculate_Distance(\&m_r,\&m_t,\&deviation);$ RR.slab.b = ba;/* swindle */ return deviation; This code is used in section 111. $\langle \text{Prototype for } Find_A_fn \mid 187 \rangle \equiv$ **double** $Find_A_fn($ **double** x)This code is used in sections 112 and 188. **188.** $\langle \text{ Definition for } Find_A_fn | 188 \rangle \equiv$ $\langle \text{ Prototype for } Find_A_fn \ 187 \rangle$ **double** $m_{-}r$, $m_{-}t$, deviation; RR.slab.a = acalc2a(x); $Calculate_Distance(\&m_r,\&m_t,\&deviation);$ **return** deviation;

This code is used in section 111.

```
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```

```
189.
         \langle \text{ Prototype for } Find\_B\_fn \text{ 189} \rangle \equiv
   double Find_B = fn(\mathbf{double} \ x)
This code is used in sections 112 and 190.
190. \langle \text{ Definition for } Find\_B\_fn \ 190 \rangle \equiv
   \langle \text{ Prototype for } Find\_B\_fn \ 189 \rangle
      double m_{-}r, m_{-}t, deviation;
     RR.slab.b = bcalc2b(x);
      Calculate\_Distance(\&m\_r,\&m\_t,\&deviation);
      return deviation;
This code is used in section 111.
191. \langle \text{Prototype for } Find_{-}G_{-}fn \mid 191 \rangle \equiv
   double Find_{-}G_{-}fn(double x)
This code is used in sections 112 and 192.
         \langle \text{ Definition for } Find_-G_-fn \mid 192 \rangle \equiv
   \langle \text{ Prototype for } Find\_G\_fn \ 191 \rangle
      double m_{-}r, m_{-}t, deviation;
      RR.slab.g = gcalc2g(x);
      Calculate\_Distance(\&m\_r,\&m\_t,\&deviation);
     return deviation;
This code is used in section 111.
        \langle \text{Prototype for } Find\_BG\_fn \ 193 \rangle \equiv
   double Find_BG_fn(double x[])
This code is used in sections 112 and 194.
194. \langle \text{ Definition for } Find\_BG\_fn \ 194 \rangle \equiv
   \langle \text{ Prototype for } Find\_BG\_fn \ 193 \rangle
      double m_{-}r, m_{-}t, deviation;
      RR.slab.b = bcalc2b(x[1]);
      RR.slab.g = gcalc2g(x[2]);
     RR.slab.a = RR.default_a;
      Calculate\_Distance(\&m\_r,\&m\_t,\&deviation);
     return deviation;
This code is used in section 111.
```

195. For this function the first term x[1] will contain the value of $\mu_s d$, the second term will contain the anisotropy. Of course the first term is in the bizarre calculation space and needs to be translated back into normal terms before use. We just at the scattering back on and voilá we have a useable value for the optical depth.

```
\langle Prototype for Find\_BaG\_fn\ 195 \rangle \equiv double Find\_BaG\_fn (double x[]) This code is used in sections 112 and 196.
```

```
196. \langle \text{ Definition for } Find\_BaG\_fn \ 196 \rangle \equiv
  \langle \text{ Prototype for } Find\_BaG\_fn \text{ 195} \rangle
     double m_{-}r, m_{-}t, deviation;
     RR.slab.b = bcalc2b(x[1]) + RR.default_bs;
     if (RR.slab.b \le 0) RR.slab.a = 0;
     else RR.slab.a = RR.default\_bs/RR.slab.b;
     RR.slab.g = gcalc2g(x[2]);
     Calculate\_Distance(\&m\_r,\&m\_t,\&deviation);
     return deviation;
This code is used in section 111.
       \langle \text{Prototype for } Find\_BsG\_fn \ 197 \rangle \equiv
  double Find_BsG_fn(double x[])
This code is used in sections 112 and 198.
198. \langle \text{ Definition for } Find\_BsG\_fn \ 198 \rangle \equiv
  \langle \text{ Prototype for } Find\_BsG\_fn \ 197 \rangle
     double m_{-}r, m_{-}t, deviation;
     RR.slab.b = bcalc2b(x[1]) + RR.default_ba;
     if (RR.slab.b \le 0) RR.slab.a = 0;
     else RR.slab.a = 1.0 - RR.default_ba/RR.slab.b;
     RR.slab.g = gcalc2g(x[2]);
     Calculate\_Distance(\&m\_r,\&m\_t,\&deviation);
     return deviation;
This code is used in section 111.
```

199. Routine to figure out if the light loss exceeds what is physically possible. Returns the descrepancy between the current values and the maximum possible values for the measurements $m_{-}r$ and $m_{-}t$.

```
\langle Prototype for maxloss 199\rangle \equiv double maxloss(double f)
This code is used in sections 112 and 200.
```

```
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```

```
200.
        \langle \text{ Definition for } maxloss | 200 \rangle \equiv
  \langle \text{ Prototype for } maxloss | 199 \rangle
     struct measure_type m_{-}old;
     struct invert_type r_old;
     double m_{-}r, m_{-}t, deviation;
     Get\_Calc\_State(\&m\_old,\&r\_old);
     RR.slab.a = 1.0;
     MM.ur1\_lost *= f;
     MM.ut1\_lost *= f;
     Calculate\_Distance(\&m\_r,\&m\_t,\&deviation);
     Set\_Calc\_State(m\_old, r\_old);
     deviation = ((MM.m_{-}r + MM.m_{-}t) - (m_{-}r + m_{-}t));
     return deviation;
This code is used in section 111.
      This checks the two light loss values ur1-loss and ut1-loss to see if they exceed what is physically
possible. If they do, then these values are replaced by a couple that are the maximum possible for the current
values in m and r.
\langle \text{ Prototype for } Max\_Light\_Loss \ 201 \rangle \equiv
  void Max.Light.Loss(struct measure_type m, struct invert_type r, double *ur1.loss, double
       *ut1\_loss)
This code is used in sections 112 and 202.
       \langle \text{ Definition for } Max\_Light\_Loss \ 202 \rangle \equiv
  \langle Prototype for Max\_Light\_Loss 201 \rangle
     struct measure_type m_{-}old;
     struct invert_type r_{-}old;
     *ur1\_loss = m.ur1\_lost;
     *ut1\_loss = m.ut1\_lost;
     if (Debug(DEBUG_LOST_LIGHT))
       fprintf(stderr, "\nlost_before_ur1=\%7.5f, ut1=\%7.5f \n", *ur1_loss, *ut1_loss);
     Get\_Calc\_State(\&m\_old,\&r\_old);
     Set\_Calc\_State(m, r);
     if (maxloss(1.0) * maxloss(0.0) < 0) {
       double frac;
       frac = zbrent(maxloss, 0.00, 1.0, 0.001);
       *ur1\_loss = m.ur1\_lost * frac;
       *ut1\_loss = m.ut1\_lost * frac;
     Set\_Calc\_State(m\_old, r\_old);
     if (Debug(DEBUG_LOST_LIGHT))
       fprintf(stderr, "lost_after_uur1=\%7.5f, ut1=\%7.5f n", *ur1_loss, *ut1_loss);
This code is used in section 111.
```

86 Calculating r and t iad (v 3-12-0) $\S 203$

```
203. this is currently unused
\langle \text{ Unused diffusion fragment } 203 \rangle \equiv
  static void DE_RT(int nfluxes, AD_slab_type slab, double *UR1, double *UT1, double *URU, double
     slabtypes;
     double rp, tp, rs, ts;
     s.f = slab.g * slab.g;
     s.gprime = slab.g/(1 + slab.g);
     s.aprime = (1 - s.f) * slab.a/(1 - slab.a * s.f);
     s.bprime = (1 - slab.a * s.f) * slab.b;
     s.boundary\_method = Egan;
     s.n_{-}top = slab.n_{-}slab;
     s.n_bottom = slab.n_slab;
     s.slide\_top = slab.n\_top\_slide;
     s.slide\_bottom = slab.n\_bottom\_slide;
     s.F0 = 1/pi;
     s.depth = 0.0;
     s.Exact\_coll\_flag = false;
    if (MM.illumination \equiv collimated) {
       compute_{-}R_{-}and_{-}T(\&s, 1.0, \&rp, \&rs, \&tp, \&ts);
       *UR1 = rp + rs;
       *UT1 = tp + ts;
       *URU = 0.0;
       *UTU = 0.0;
       return;
     quad\_Dif\_Calc\_R\_and\_T(\&s,\&rp,\&rs,\&tp,\&ts);
     *URU = rp + rs;
    *UTU = tp + ts;
     *UR1 = 0.0;
     *UT1 = 0.0;
```

 $\S204$ IAD (v 3-12-0) IAD FIND 87

204. IAD Find. March 1995. Incorporated the quick_quess algorithm for low albedos.

```
\langle iad\_find.c 204 \rangle \equiv
#include <math.h>
#include <stdio.h>
#include <stdlib.h>
#include "ad_globl.h"
#include "nr_util.h"
#include "nr_mnbrk.h"
#include "nr_brent.h"
#include "nr_amoeb.h"
#include "iad_type.h"
#include "iad_util.h"
#include "iad_calc.h"
#include "iad_find.h"
#include "iad_util.h"
#define NUMBER_OF_GUESSES 10
  guess_type guess[NUMBER_OF_GUESSES];
  int compare_quesses(const void *p1, const void *p2)
     \mathbf{guess\_type} *g1 = (\mathbf{guess\_type} *) p1;
     \mathbf{guess\_type} *g2 = (\mathbf{guess\_type} *) p2;
     if (g1 \neg distance < g2 \neg distance) return -1;
     else if (g1 \neg distance \equiv g2 \neg distance) return 0;
     else return 1;
  \langle \text{ Definition for } U\_Find\_Ba \text{ 218} \rangle
   \langle \text{ Definition for } U\_Find\_Bs \text{ 216} \rangle
   \langle \text{ Definition for } U\_Find\_A \text{ 220} \rangle
   \langle \text{ Definition for } U_{-}Find_{-}B \rangle
    Definition for U_Find_G = 222
   Definition for U_Find_AG 227
   \langle \text{ Definition for } U_F ind_A B | 207 \rangle
   \langle \text{ Definition for } U\_Find\_BG \text{ 232} \rangle
   Definition for U_Find_BaG 238
  \langle \text{ Definition for } U\_Find\_BsG \text{ 243} \rangle
```

205. All the information that needs to be written to the header file iad_find.h. This eliminates the need to maintain a set of header files as well.

```
 \begin{split} &\langle \text{iad\_find.h} \quad 205 \rangle \equiv \\ &\langle \text{Prototype for } \textit{U\_Find\_Ba} \text{ 217} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_Bs} \text{ 215} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_A} \text{ 219} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_B} \text{ 223} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_G} \text{ 221} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_AG} \text{ 226} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_AB} \text{ 206} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_BG} \text{ 231} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_BaG} \text{ 237} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_BaG} \text{ 237} \rangle; \\ &\langle \text{Prototype for } \textit{U\_Find\_BaG} \text{ 242} \rangle; \end{aligned}
```

88 FIXED ANISOTROPY IAD (v 3-12-0) $\S 206$

```
Fixed Anisotropy.
   This is the most common case.
\langle \text{ Prototype for } U_F ind_A B | 206 \rangle \equiv
   void U_Find_AB(struct measure_type m, struct invert_type *r)
This code is used in sections 205 and 207.
207. \langle \text{ Definition for } U\_Find\_AB | 207 \rangle \equiv
   \langle \, \text{Prototype for } \textit{U\_Find\_AB} \,\, \, 206 \, \rangle
     (Allocate local simplex variables 208)
     if (Debug(DEBUG\_SEARCH)) {
        fprintf(stderr, "In U_Find_AB");
        fprintf(stderr, "u(mu=\%6.4f)", r\rightarrow slab.cos\_angle);
        if (r - default_g \neq UNINITIALIZED) fprintf(stderr, "ulldefault_gu=u%8.5f", r-default_g);
        fprintf(stderr, "\n");
     r \rightarrow slab.g = (r \rightarrow default\_g \equiv \texttt{UNINITIALIZED}) ? 0 : r \rightarrow default\_g;
     Set\_Calc\_State(m, *r);
     \langle \text{ Get the initial } a, b, \text{ and } g \text{ 209} \rangle
      \langle Initialize the nodes of the a and b simplex 210\rangle
      \langle Evaluate the a and b simplex at the nodes 211\rangle
     amoeba(p, y, 2, r \rightarrow tolerance, Find\_AB\_fn, \&r \rightarrow iterations);
     \langle Choose the best node of the a and b simplex 212\rangle
      (Free simplex data structures 214)
      (Put final values in result 213)
This code is used in section 204.
208. To use the simplex algorithm, we need to vectors and a matrix.
\langle Allocate local simplex variables 208\rangle \equiv
  int i, i\_best, j\_best;
  double *x, *y, **p;
  x = dvector(1, 2);
  y = dvector(1,3);
  p = dmatrix(1, 3, 1, 2);
This code is used in sections 207, 227, 232, 238, and 243.
```

 $\S209$ IAD (v 3-12-0) FIXED ANISOTROPY 89

209. Just get the optimal optical properties to start the search process.

I had to add the line that tests to make sure the albedo is greater than 0.2 because the grid just does not work so well in this case. The problem is that for low albedos there is really very little information about the anisotropy available. This change was also made in the analogous code for a and b.

```
\langle \text{ Get the initial } a, b, \text{ and } g \text{ 209} \rangle \equiv
         /* double a3,b3,g3; */
     size_t \ count = NUMBER_OF_GUESSES;
                                                      /* distance to last result */
     abg\_distance(r \rightarrow slab.a, r \rightarrow slab.b, r \rightarrow slab.g, \&(guess[0]));
     if (\neg Valid\_Grid(m, r \rightarrow search)) Fill\_Grid(m, *r, 1);
                                                                      /* distance to nearest grid point */
     Near\_Grid\_Points(m.m\_r, m.m\_t, r \rightarrow search, \&i\_best, \&j\_best);
     Grid\_ABG(i\_best, j\_best, \&(guess[1]));
     Grid\_ABG(i\_best + 1, j\_best, \&(guess[2]));
     Grid\_ABG(i\_best - 1, j\_best, \&(guess[3]));
     Grid\_ABG(i\_best, j\_best + 1, \&(guess[4]));
     Grid\_ABG(i\_best, j\_best - 1, \&(guess[5]));
     Grid\_ABG(i\_best + 1, j\_best + 1, \&(guess[6]));
     Grid\_ABG(i\_best-1, j\_best-1, \&(guess[7]));
     Grid\_ABG(i\_best + 1, j\_best - 1, \&(guess[8]));
     Grid\_ABG(i\_best-1, j\_best+1, \&(quess[9]));
     qsort((void *) guess, count, sizeof(guess_type), compare_guesses);
     if (Debug(DEBUG_BEST_GUESS)) {
       int k;
        fprintf(stderr, "after\n");
       for (k = 0; k \le 6; k++) {
          fprintf(stderr, "%3d_{\sqcup\sqcup}", k);
          fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].a);
          fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].b);
          fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].g);
          fprintf(stderr, "%10.5f\n", guess[k].distance);
     }
```

This code is used in sections 207, 227, 232, 238, and 243.

90 FIXED ANISOTROPY IAD (v 3-12-0) §210

```
210. (Initialize the nodes of the a and b simplex 210) \equiv
     int k, kk;
    p[1][1] = a2acalc(quess[0].a);
    p[1][2] = b2bcalc(guess[0].b);
     for (k = 1; k < 7; k++) {
       if (guess[0].a \neq guess[k].a) break;
     p[2][1] = a2acalc(guess[k].a);
     p[2][2] = b2bcalc(quess[k].b);
     for (kk = 1; kk < 7; kk ++) {
       if (guess[0].b \neq guess[kk].b \land guess[k].b \neq guess[kk].b) break;
     p[3][1] = a2acalc(guess[kk].a);
     p[3][2] = b2bcalc(guess[kk].b);
     if (Debug(DEBUG_BEST_GUESS)) {
       fprintf(stderr, "guess<sub>□</sub>1");
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].a);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].g);
       fprintf(stderr, "%10.5f\n", guess[0].distance);
       fprintf(stderr, "guess_2");
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].a);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].b);
       fprintf(stderr, "\%10.5f_{\square}", guess[k].g);
       fprintf (stderr, \verb"%10.5f\n", guess[k].distance);
       fprintf(stderr, "guess_{\sqcup}3");
       fprintf(stderr, "%10.5f_{\sqcup}", guess[kk].a);
       fprintf(stderr, "%10.5f_{\sqcup}", guess[kk].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[kk].g);
       fprintf(stderr, "%10.5f\n", guess[kk].distance);
This code is used in section 207.
211. (Evaluate the a and b simplex at the nodes 211) \equiv
  for (i = 1; i \le 3; i++) {
    x[1] = p[i][1];
     x[2] = p[i][2];
     y[i] = Find\_AB\_fn(x);
This code is used in section 207.
```

```
212.
          \langle Choose the best node of the a and b simplex 212 \rangle \equiv
   r \rightarrow final\_distance = 10;
   for (i = 1; i < 3; i++) {
      if (y[i] < r \rightarrow final\_distance) {
         r \rightarrow slab.a = acalc2a(p[i][1]);
         r \rightarrow slab.b = bcalc2b(p[i][2]);
         r \rightarrow final\_distance = y[i];
   }
This code is used in section 207.
213. \langle \text{Put final values in result 213} \rangle \equiv
   r \rightarrow a = r \rightarrow slab.a;
   r \rightarrow b = r \rightarrow slab.b;
   r \rightarrow g = r \rightarrow slab.g;
   r \rightarrow found = (r \rightarrow tolerance \leq r \rightarrow final\_distance);
This code is used in sections 207, 216, 218, 220, 222, 224, 227, 232, 238, and 243.
214. Since we allocated these puppies, we got to get rid of them.
\langle Free simplex data structures 214\rangle \equiv
   free\_dvector(x, 1, 2);
   free\_dvector(y, 1, 3);
   free\_dmatrix(p, 1, 3, 1, 2);
This code is used in sections 207, 227, 232, 238, and 243.
```

215. Fixed Absorption and Anisotropy. Typically, this routine is called when the absorption coefficient is known, the anisotropy is known, and the physical thickness of the sample is known. This routine calculates the varies the scattering coefficient until the measurements are matched.

This was written for Ted Moffitt to analyze some intralipid data. We wanted to know what the scattering coefficient of the Intralipid was and made total transmission measurements through a sample with a fixed physical thickness. We did not make reflection measurements because the light source diverged too much, and we could not make reflection measurements easily.

In retrospect, we could have made URU measurements by illuminating the wall of the integrating sphere. However, these diffuse type of measurements are very difficult to make accurately.

This is tricky only because the value in slab.b is used to hold the value of ba or $d \cdot \mu_a$ when the $Find_Bs_fn$ is used.

```
\langle \text{Prototype for } U\_Find\_Bs \ 215 \rangle \equiv 
void U\_Find\_Bs (\text{struct measure\_type } m, \text{struct invert\_type } *r)
This code is used in sections 205 and 216.
```

IAD (v 3-12-0)

```
\langle \text{ Definition for } U\_Find\_Bs \ 216 \rangle \equiv
   \langle \text{ Prototype for } U_F ind_B s \text{ 215} \rangle
      double ax, bx, cx, fa, fb, fc, bs;
       \textbf{if} \ (Debug(\texttt{DEBUG\_SEARCH})) \ \{ \\
         fprintf(stderr, "In_U_Find_Bs");
         fprintf (stderr, "⊔(mu=%6.4f)", r→slab.cos_angle);
         if (r \rightarrow default\_ba \neq UNINITIALIZED) fprintf(stderr, " \sqcup \sqcup default\_ba \sqcup = \sqcup \%8.5f", r \rightarrow default\_ba);
         if (r \rightarrow default\_g \neq UNINITIALIZED) fprintf (stderr, "uudefault\_gu=u\%8.5f", r \rightarrow default\_g);
         fprintf(stderr, "\n");
      if (m.m_-t \equiv 0) {
         r \rightarrow slab.b = HUGE_VAL;
         U_{-}Find_{-}A(m,r);
         return;
      r \rightarrow slab.a = 0;
      r \rightarrow slab.g = (r \rightarrow default\_g \equiv UNINITIALIZED) ? 0 : r \rightarrow default\_g;
      r \rightarrow slab.b = (r \rightarrow default\_ba \equiv UNINITIALIZED)? HUGE_VAL: r \rightarrow default\_ba;
      Set\_Calc\_State(m, *r);
                                        /* store ba in RR.slab.b */
                                    /* first try for bs */
      ax = b2bcalc(0.1);
      bx = b2bcalc(1.0);
      mnbrak(\&ax,\&bx,\&cx,\&fa,\&fb,\&fc,Find_Bs_fn);
      r \rightarrow final\_distance = brent(ax, bx, cx, Find\_Bs\_fn, r \rightarrow tolerance, \&bs);
                                                                                                 /* recover true values */
      r \rightarrow slab.a = bcalc2b(bs)/(bcalc2b(bs) + r \rightarrow slab.b);
      r \rightarrow slab.b = bcalc2b(bs) + r \rightarrow slab.b;
      Set\_Calc\_State(m, *r);
      (Put final values in result 213)
This code is used in section 204.
```

217. Fixed Absorption and Scattering. Typically, this routine is called when the scattering coefficient is known, the anisotropy is known, and the physical thickness of the sample is known. This routine calculates the varies the absorption coefficient until the measurements are matched.

This is tricky only because the value in slab.b is used to hold the value of bs or $d \cdot \mu_s$ when the Find_Ba_fn is used.

```
\langle \text{ Prototype for } U_F ind_B a \text{ 217} \rangle \equiv
   void U_Find_Ba(struct measure\_type m, struct invert\_type *r)
This code is used in sections 205 and 218.
```

```
218.
         \langle \text{ Definition for } U_F ind_B a \text{ 218} \rangle \equiv
   \langle \text{ Prototype for } U_F ind_B a \text{ 217} \rangle
      double ax, bx, cx, fa, fb, fc, ba;
     if (Debug(DEBUG_SEARCH)) {
        fprintf(stderr, "In_U_Find_Bs");
        fprintf (stderr, "⊔(mu=%6.4f)", r→slab.cos_angle);
        if (r \rightarrow default\_bs \neq UNINITIALIZED) fprintf (stderr, "ulldefault\_bs = "%8.5f", r \rightarrow default\_bs);
        if (r \rightarrow default\_g \neq UNINITIALIZED) fprintf (stderr, "uudefault\_gu=u\%8.5f", r \rightarrow default\_g);
         fprintf(stderr, "\n");
      r \rightarrow slab.a = 0;
      r \rightarrow slab.g = (r \rightarrow default\_g \equiv UNINITIALIZED) ? 0 : r \rightarrow default\_g;
      r \rightarrow slab.b = (r \rightarrow default\_bs \equiv \texttt{UNINITIALIZED})? \texttt{HUGE\_VAL}: r \rightarrow default\_bs;
      if (m.m_t \equiv 0) {
        r\rightarrow slab.b = HUGE\_VAL;
         U_{-}Find_{-}A(m,r);
        return;
      Set\_Calc\_State(m, *r);
                                         /* store bs in RR.slab.b */
                                   /* first try for ba */
      ax = b2bcalc(0.1);
      bx = b2bcalc(1.0);
      mnbrak(\&ax,\&bx,\&cx,\&fa,\&fb,\&fc,Find_Ba\_fn);
      r \rightarrow final\_distance = brent(ax, bx, cx, Find\_Ba\_fn, r \rightarrow tolerance, \&ba);
                                                                                                 /* recover true values */
      r \rightarrow slab.a = (r \rightarrow slab.b)/(bcalc2b(ba) + r \rightarrow slab.b);
      r \rightarrow slab.b = bcalc2b(ba) + r \rightarrow slab.b;
                                                       /* actual value of b */
      Set\_Calc\_State(m, *r);
      (Put final values in result 213)
This code is used in section 204.
```

219. Fixed Optical Depth and Anisotropy. Typically, this routine is called when the optical thickness is assumed infinite. However, it may also be called when the optical thickness is assumed to be fixed at a particular value. Typically the only reasonable situation for this to occur is when the diffuse transmission is non-zero but the collimated transmission is zero. If this is the case then there is no information in the collimated transmission measurement and there is no sense even using it because the slab is not infinitely thick.

```
⟨ Prototype for U_Find_A 219⟩ ≡ void U_Find_A(struct measure_type m, struct invert_type *r)
This code is used in sections 205 and 220.
```

This code is used in sections 205 and 222.

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```
\langle \text{ Definition for } U_F ind_A \underline{220} \rangle \equiv
   \langle \text{ Prototype for } U\_Find\_A \text{ 219} \rangle
     double Rt, Tt, Rd, Rc, Td, Tc;
     if (Debug(DEBUG\_SEARCH)) {
        fprintf(stderr, "In U_Find_A");
        fprintf (stderr, "⊔(mu=%6.4f)", r→slab.cos_angle);
        if (r - default_b \neq UNINITIALIZED) fprintf(stderr, "uudefault_bu=u%8.5f", r- default_b);
        if (r \rightarrow default\_g \neq UNINITIALIZED) fprintf (stderr, "uudefault\_gu=u\%8.5f", r \rightarrow default\_g);
        fprintf(stderr, "\n");
     Estimate\_RT(m, *r, \&Rt, \&Tt, \&Rd, \&Rc, \&Td, \&Tc);
     r \rightarrow slab.g = (r \rightarrow default\_g \equiv UNINITIALIZED) ? 0 : r \rightarrow default\_g;
     r \rightarrow slab.b = (r \rightarrow default\_b \equiv UNINITIALIZED)? HUGE_VAL: r \rightarrow default\_b;
     r \rightarrow slab.a = 0.0;
     r \rightarrow final\_distance = 0.0;
     Set\_Calc\_State(m, *r);
     if (Rt > 0.99999) {
        r \rightarrow final\_distance = Find\_A\_fn(a2acalc(1.0));
        r \rightarrow slab.a = 1.0;
     else {
        double x, ax, bx, cx, fa, fb, fc;
        ax = a2acalc(0.3);
        bx = a2acalc(0.5);
        mnbrak(\&ax,\&bx,\&cx,\&fa,\&fb,\&fc,Find\_A\_fn);
        r \rightarrow final\_distance = brent(ax, bx, cx, Find\_A\_fn, r \rightarrow tolerance, \&x);
        r \rightarrow slab.a = acalc2a(x);
      (Put final values in result 213)
This code is used in section 204.
221. Fixed Optical Depth and Albedo.
\langle \text{ Prototype for } U_{-}Find_{-}G \text{ 221} \rangle \equiv
   void U_{-}Find_{-}G(struct measure_type m, struct invert_type *r)
```

```
222.
         \langle \text{ Definition for } U_F ind_G \underline{G} \underline{222} \rangle \equiv
   \langle \text{ Prototype for } U\_Find\_G \text{ 221} \rangle
     double Rt, Tt, Rd, Rc, Td, Tc;
     if (Debug(DEBUG\_SEARCH))  {
        fprintf(stderr, "In_U_Find_G");
        fprintf (stderr, "⊔(mu=%6.4f)", r→slab.cos_angle);
        if (r \rightarrow default_a \neq UNINITIALIZED) fprintf (stderr, "uudefault_au=u\%8.5f", r \rightarrow default_a);
        if (r \rightarrow default\_b \neq UNINITIALIZED) fprintf (stderr, "uudefault\_b_u=u\%8.5f", r \rightarrow default\_b);
        fprintf(stderr, "\n");
     Estimate\_RT(m, *r, \&Rt, \&Tt, \&Rd, \&Rc, \&Td, \&Tc);
     r \rightarrow slab.a = (r \rightarrow default\_a \equiv UNINITIALIZED) ? 0.5 : r \rightarrow default\_a;
     r \rightarrow slab.b = (r \rightarrow default\_b \equiv UNINITIALIZED)? HUGE_VAL: r \rightarrow default\_b;
     r \rightarrow slab.q = 0.0;
     r \rightarrow final\_distance = 0.0;
     Set\_Calc\_State(m, *r);
     if (Rd > 0.0) {
        double x, ax, bx, cx, fa, fb, fc;
         ax = g2gcalc(-0.99);
         bx = q2qcalc(0.99);
         mnbrak(\&ax,\&bx,\&cx,\&fa,\&fb,\&fc,Find\_G\_fn);
         r-final_distance = brent(ax, bx, cx, Find_G_fn, r-tolerance, &x);
        r \rightarrow slab.g = gcalc2g(x);
         Set\_Calc\_State(m, *r);
      (Put final values in result 213)
This code is used in section 204.
```

223. Fixed Anisotropy and Albedo. This routine can be called in three different situations: (1) the albedo is zero, (2) the albedo is one, or (3) the albedo is fixed at a default value. I calculate the individual reflections and transmissions to establish which of these cases we happen to have.

```
\langle \text{Prototype for } U\_Find\_B \text{ 223} \rangle \equiv 
void U\_Find\_B \text{(struct measure\_type } m, \text{struct invert\_type } *r)
This code is used in sections 205 and 224.
```

IAD (v 3-12-0)

```
\langle \text{ Definition for } U_F ind_B = 224 \rangle \equiv
  \langle \text{ Prototype for } U_F ind_B \ \underline{223} \rangle
     double Rt, Tt, Rd, Rc, Td, Tc;
     if (Debug(DEBUG_SEARCH)) {
        fprintf(stderr, "In_U_Find_B");
        fprintf (stderr, "⊔(mu=%6.4f)", r→slab.cos_angle);
        if (r \rightarrow default\_a \neq UNINITIALIZED) fprintf (stderr, "uudefault\_au=u\%8.5f", r \rightarrow default\_a);
        if (r - default_g \neq UNINITIALIZED) fprintf(stderr, "ulldefault_gu=u%8.5f", r-default_g);
        fprintf(stderr, "\n");
     Estimate\_RT(m, *r, \&Rt, \&Tt, \&Rd, \&Rc, \&Td, \&Tc);
     r \rightarrow slab.g = (r \rightarrow default\_g \equiv \texttt{UNINITIALIZED}) ? 0 : r \rightarrow default\_g;
     r \rightarrow slab.a = (r \rightarrow default\_a \equiv UNINITIALIZED) ? 0 : r \rightarrow default\_a;
     r \rightarrow slab.b = 0.5;
     r \rightarrow final\_distance = 0.0;
     Set\_Calc\_State(m, *r);
     \langle Iteratively solve for b \ 225 \rangle
     (Put final values in result 213)
     if (Debug(DEBUG_SEARCH)) {
        fprintf(stderr, "In_{I}U_Find_B_{I}final_{I}(a,b,g)_{I}=_{I}");
        fprintf(stderr, "(\%8.5f, \%8.5f, \%8.5f) \n", r \rightarrow a, r \rightarrow b, r \rightarrow g);
  }
This code is used in section 204.
225. This could be improved tremendously. I just don't want to mess with it at the moment.
\langle Iteratively solve for b = 225 \rangle \equiv
     double x, ax, bx, cx, fa, fb, fc;
     ax = b2bcalc(0.1);
     bx = b2bcalc(10);
     mnbrak(\&ax,\&bx,\&cx,\&fa,\&fb,\&fc,Find\_B\_fn);
     r-final_distance = brent(ax, bx, cx, Find_B-fn, r-tolerance, &x);
     r \rightarrow slab.b = bcalc2b(x);
     Set_{-}Calc_{-}State(m, *r);
This code is used in section 224.
226. Fixed Optical Depth.
  We can get here a couple of different ways.
  First there can be three real measurements, i.e., t_c is not zero, in this case we want to fix b based on the
t_c measurement.
  Second, we can get here if a default value for b has been set.
  Otherwise, we really should not be here. Just set b = 1 and calculate away.
\langle \text{ Prototype for } U_F ind_A G | 226 \rangle \equiv
  void U_Find_AG(struct measure_type m, struct invert_type *r)
This code is used in sections 205 and 227.
```

```
227.
          \langle \text{ Definition for } U\_Find\_AG \text{ 227} \rangle \equiv
   \langle \text{ Prototype for } U_{-}Find_{-}AG | 226 \rangle
      (Allocate local simplex variables 208)
      if (Debug(DEBUG\_SEARCH))  {
        fprintf(stderr, "In_U_Find_AG");
        fprintf (stderr, "⊔(mu=%6.4f)", r→slab.cos_angle);
         if (r \rightarrow default\_b \neq \texttt{UNINITIALIZED}) fprintf(stderr, "\_\_default\_b\_=\_\%8.5f", r \rightarrow default\_b);
         fprintf(stderr, "\n");
      if (m.num\_measures \equiv 3) r \rightarrow slab.b = What\_Is\_B(r \rightarrow slab, m.m\_u);
      else if (r \rightarrow default\_b \equiv UNINITIALIZED) r \rightarrow slab.b = 1;
      else r \rightarrow slab.b = r \rightarrow default\_b;
      Set_{-}Calc_{-}State(m, *r);
      \langle \text{ Get the initial } a, b, \text{ and } g \text{ 209} \rangle
      \langle Initialize the nodes of the a and g simplex 228\rangle
      \langle Evaluate the a and g simplex at the nodes 229\rangle
      amoeba(p, y, 2, r \rightarrow tolerance, Find\_AG\_fn, \&r \rightarrow iterations);
      \langle Choose the best node of the a and g simplex 230\rangle
      (Free simplex data structures 214)
      (Put final values in result 213)
```

This code is used in section 204.

98 FIXED OPTICAL DEPTH IAD (v 3-12-0) $\S 228$

```
(Initialize the nodes of the a and q simplex 228) \equiv
  {
    int k, kk;
    p[1][1] = a2acalc(quess[0].a);
    p[1][2] = g2gcalc(guess[0].g);
     for (k = 1; k < 7; k ++) {
       if (guess[0].a \neq guess[k].a) break;
    p[2][1] = a2acalc(guess[k].a);
    p[2][2] = g2qcalc(quess[k].q);
     for (kk = 1; kk < 7; kk ++) {
       if (guess[0].g \neq guess[kk].g \land guess[k].g \neq guess[kk].g) break;
    p[3][1] = a2acalc(guess[kk].a);
    p[3][2] = g2gcalc(guess[kk].g);
    if (Debug(DEBUG_BEST_GUESS)) {
       fprintf(stderr, "guess_1");
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].a);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].g);
       fprintf(stderr, "%10.5f\n", guess[0].distance);
       fprintf(stderr, "guess_2");
       fprintf(stderr, "\%10.5f_{11}", quess[k].a);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].g);
       fprintf (stderr, \verb"%10.5f\n", guess[k].distance);
       fprintf(stderr, "guess_{\sqcup}3");
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[kk].a);
       fprintf(stderr, "%10.5f", guess[kk].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[kk].g);
       fprintf(stderr, "%10.5f\n", guess[kk].distance);
This code is used in section 227.
229. (Evaluate the a and g simplex at the nodes 229) \equiv
  for (i = 1; i \le 3; i++) {
    x[1] = p[i][1];
    x[2] = p[i][2];
    y[i] = Find\_AG\_fn(x);
This code is used in section 227.
```

§230 IAD (v 3-12-0)

99

230. Here we find the node of the simplex that gave the best result and save that one. At the same time we save the whole simplex for later use if needed.

```
\langle Choose the best node of the a and g simplex 230\rangle \equiv
   r \rightarrow final\_distance = 10;
   for (i = 1; i \le 3; i ++) {
     if (y[i] < r \rightarrow final\_distance) {
         r \rightarrow slab.a = acalc2a(p[i][1]);
        r \rightarrow slab.g = gcalc2g(p[i][2]);
        r \rightarrow final\_distance = y[i];
This code is used in section 227.
231. Fixed Albedo. Here the optical depth and the anisotropy are varied (for a fixed albedo).
\langle \text{ Prototype for } U\_Find\_BG \text{ 231} \rangle \equiv
   void U_Find_BG(struct measure_type m, struct invert_type *r)
This code is used in sections 205 and 232.
        \langle \text{ Definition for } U\_Find\_BG \text{ 232} \rangle \equiv
   \langle \text{ Prototype for } U_F ind_B G \text{ 231} \rangle
      (Allocate local simplex variables 208)
      if (Debug(DEBUG\_SEARCH)) {
        fprintf(stderr, "In_U_Find_BG");
        fprintf(stderr, "u(mu=%6.4f)", r→slab.cos_angle);
        if (r \rightarrow default\_a \neq UNINITIALIZED) fprintf(stderr, "ulldefault\_al=u%8.5f", r \rightarrow default\_a);
        fprintf(stderr, "\n");
      r \rightarrow slab.a = (r \rightarrow default\_a \equiv UNINITIALIZED) ? 0 : r \rightarrow default\_a;
      Set\_Calc\_State(m, *r);
      \langle \text{ Get the initial } a, b, \text{ and } g \text{ 209} \rangle
      \langle Initialize the nodes of the b and g simplex 234\rangle
      \langle Evaluate the bq simplex at the nodes 235\rangle
      amoeba(p, y, 2, r \rightarrow tolerance, Find\_BG\_fn, \&r \rightarrow iterations);
      \langle Choose the best node of the b and g simplex 236\rangle
      ⟨Free simplex data structures 214⟩
      (Put final values in result 213)
This code is used in section 204.
```

233. A very simple start for variation of b and g. This should work fine for the cases in which the absorption or scattering are fixed.

100 FIXED ALBEDO IAD (v 3-12-0) $\S234$

```
234.
        \langle Initialize the nodes of the b and q simplex 234\rangle \equiv
  {
     int k, kk;
    p[1][1] = b2bcalc(quess[0].b);
    p[1][2] = g2gcalc(guess[0].g);
     for (k = 1; k < 7; k ++) {
       if (guess[0].b \neq guess[k].b) break;
     p[2][1] = b2bcalc(quess[k].b);
     p[2][2] = g2qcalc(quess[k],q);
     for (kk = 1; kk < 7; kk ++) {
       if (guess[0].g \neq guess[kk].g \land guess[k].g \neq guess[kk].g) break;
     p[3][1] = b2bcalc(guess[kk].b);
     p[3][2] = g2gcalc(guess[kk].g);
     if (Debug(DEBUG_BEST_GUESS)) {
       fprintf(stderr, "guess<sub>□</sub>1");
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].a);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[0].g);
       fprintf(stderr, "%10.5f\n", guess[0].distance);
       fprintf(stderr, "guess_2");
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].a);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[k].g);
       fprintf(stderr, "\%10.5f\n", guess[k].distance);
       fprintf(stderr, "guess_{\sqcup}3");
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[kk].a);
       fprintf(stderr, "%10.5f_{\sqcup}", guess[kk].b);
       fprintf(stderr, "\%10.5f_{\sqcup}", guess[kk].g);
       fprintf(stderr, "%10.5f\n", guess[kk].distance);
This code is used in section 232.
235. (Evaluate the bg simplex at the nodes 235) \equiv
  for (i = 1; i \le 3; i ++) {
    x[1] = p[i][1];
     x[2] = p[i][2];
     y[i] = Find\_BG\_fn(x);
This code is used in section 232.
```

 $\S236$ IAD (v 3-12-0) FIXED ALBEDO 101

236. Here we find the node of the simplex that gave the best result and save that one. At the same time we save the whole simplex for later use if needed.

```
 \langle \text{ Choose the best node of the } b \text{ and } g \text{ simplex } 236 \rangle \equiv r \neg final\_distance = 10; \\ \text{for } (i=1;\ i \leq 3;\ i++) \ \{\\ \text{if } (y[i] < r \neg final\_distance) \ \{\\ r \neg slab.b = bcalc2b(p[i][1]); \\ r \neg slab.g = gcalc2g(p[i][2]); \\ r \neg final\_distance = y[i]; \\ \}\\ \} \\ \text{This code is used in section } 232.
```

237. Fixed Scattering. Here I assume that a constant b_s ,

$$b_s = \mu_s d$$

where d is the physical thickness of the sample and μ_s is of course the absorption coefficient. This is just like U_-Find_-BG except that $b_a = \mu_a d$ is varied instead of b.

```
⟨Prototype for U_Find_BaG 237⟩ ≡ void U_Find_BaG(struct measure_type m, struct invert_type *r)
This code is used in sections 205 and 238.
238. ⟨Definition for U_Find_BaG 238⟩ ≡
⟨Prototype for U_Find_BaG 237⟩
{
⟨Allocate local simplex variables 208⟩
Set_Calc_State(m, *r);
```

 \langle Get the initial a, b, and $g 209 \rangle$ \langle Initialize the nodes of the ba and g simplex 239 \rangle \langle Evaluate the BaG simplex at the nodes 240 \rangle amoeba(p, y, 2, r-tolerance, $Find_BaG_fn, \&r$ -iterations); \langle Choose the best node of the ba and g simplex 241 \rangle \langle Free simplex data structures 214 \rangle \langle Put final values in result 213 \rangle

This code is used in section 204.

}

102 FIXED SCATTERING IAD (v 3-12-0) $\S 239$

```
239.
        (Initialize the nodes of the ba and q simplex 239) \equiv
  if (guess[0].b > r \rightarrow default\_bs) {
    p[1][1] = b2bcalc(guess[0].b - r \rightarrow default\_bs);
    p[2][1] = b2bcalc(2*(guess[0].b - r \rightarrow default\_bs));
    p[3][1] = p[1][1];
  else {
    p[1][1] = b2bcalc(0.0001);
    p[2][1] = b2bcalc(0.001);
    p[3][1] = p[1][1];
  p[1][2] = g2gcalc(guess[0].g);
  p[2][2] = p[1][2];
  p[3][2] = g2gcalc(0.9 * guess[0].g + 0.05);
This code is used in section 238.
240. (Evaluate the BaG simplex at the nodes 240) \equiv
  for (i = 1; i \le 3; i++) {
     x[1] = p[i][1];
     x[2] = p[i][2];
     y[i] = Find_BaG_fn(x);
This code is used in section 238.
```

241. Here we find the node of the simplex that gave the best result and save that one. At the same time we save the whole simplex for later use if needed.

```
 \langle \text{Choose the best node of the } ba \text{ and } g \text{ simplex } 241 \rangle \equiv r \neg final\_distance = 10; \\ \text{for } (i=1; i \leq 3; i++) \{\\ \text{if } (y[i] < r \neg final\_distance) \{\\ r \neg slab.b = bcalc2b(p[i][1]) + r \neg default\_bs; \\ r \neg slab.a = r \neg default\_bs/r \neg slab.b; \\ r \neg slab.g = gcalc2g(p[i][2]); \\ r \neg final\_distance = y[i]; \\ \} \\ \}
```

This code is used in section 238.

242. Fixed Absorption. Here I assume that a constant b_a ,

$$b_a = \mu_a d$$

where d is the physical thickness of the sample and μ_a is of course the absorption coefficient. This is just like U_-Find_-BG except that $b_s = \mu_s d$ is varied instead of b.

```
\langle \text{Prototype for } U\_Find\_BsG \ 242 \rangle \equiv 
void U\_Find\_BsG(\text{struct measure\_type } m, \text{struct invert\_type } *r)
This code is used in sections 205 and 243.
```

```
243.
         \langle \text{ Definition for } U_F ind_B sG | 243 \rangle \equiv
   \langle Prototype for U\_Find\_BsG 242 \rangle
      (Allocate local simplex variables 208)
     if (Debug(DEBUG\_SEARCH)) {
        fprintf(stderr, "In_U_Find_BsG");
        fprintf(stderr, "u(mu=%6.4f)", r→slab.cos_angle);
        if (r \rightarrow default\_ba \neq UNINITIALIZED) fprintf(stderr, "\lu default\_ba \lu = \lu %8.5f", r \rightarrow default\_ba);
        fprintf(stderr, "\n");
     Set\_Calc\_State(m, *r);
      \langle \text{ Get the initial } a, b, \text{ and } g \text{ 209} \rangle
      \langle Initialize the nodes of the bs and g simplex 244\rangle
     \langle \text{Evaluate the } BsG \text{ simplex at the nodes } 245 \rangle
     amoeba(p, y, 2, r \rightarrow tolerance, Find\_BsG\_fn, \&r \rightarrow iterations);
      \langle Choose the best node of the bs and g simplex 246\rangle
      (Free simplex data structures 214)
      (Put final values in result 213)
  }
This code is used in section 204.
244. (Initialize the nodes of the bs and g simplex 244) \equiv
  p[1][1] = b2bcalc(quess[0].b - r \rightarrow default\_ba);
  p[1][2] = g2gcalc(guess[0].g);
  p[2][1] = b2bcalc(2*guess[0].b - 2*r \rightarrow default\_ba);
  p[2][2] = p[1][2];
  p[3][1] = p[1][1];
  p[3][2] = g2gcalc(0.9 * guess[0].g + 0.05);
This code is used in section 243.
245. (Evaluate the BsG simplex at the nodes 245) \equiv
   for (i = 1; i \le 3; i++) {
     x[1] = p[i][1];
     x[2] = p[i][2];
     y[i] = Find_BsG_fn(x);
This code is used in section 243.
246. (Choose the best node of the bs and g simplex 246) \equiv
   r \rightarrow final\_distance = 10;
   for (i = 1; i \le 3; i++) {
     if (y[i] < r \rightarrow final\_distance) {
        r \rightarrow slab.b = bcalc2b(p[i][1]) + r \rightarrow default\_ba;
        r \rightarrow slab.a = 1 - r \rightarrow default\_ba/r \rightarrow slab.b;
        r \rightarrow slab.g = gcalc2g(p[i][2]);
        r \rightarrow final\_distance = y[i];
This code is used in section 243.
```

104 IAD UTILITIES IAD (v 3-12-0) $\S 247$

247. IAD Utilities.

March 1995. Reincluded quick_guess code.

```
\langle iad\_util.c 247 \rangle \equiv
#include <math.h>
#include <float.h>
#include <stdio.h>
#include "nr_util.h"
#include "ad_globl.h"
#include "ad_frsnl.h"
#include "ad_bound.h"
#include "iad_type.h"
#include "iad_calc.h"
#include "iad_pub.h"
#include "iad_util.h"
  unsigned long g_{-}util_{-}debugging = 0;
  ⟨ Preprocessor definitions ⟩
  \langle \text{ Definition for } What\_Is\_B \text{ 250} \rangle
   \langle \text{ Definition for } Estimate\_RT | 256 \rangle
   \langle \text{ Definition for } a2acalc | 262 \rangle
   (Definition for acalc2a 264)
   \langle \text{ Definition for } q2qcalc | 266 \rangle
   \langle Definition for gcalc2g 268\rangle
   Definition for b2bcalc 270 \rangle
   \langle \text{ Definition for } bcalc2b | 272 \rangle
   (Definition for twoprime 274)
   \langle \text{ Definition for } two unprime 276 \rangle
    Definition for abgg2ab 278\rangle
   (Definition for abgb2ag 280)
   (Definition for quick_quess 287)
   (Definition for Set_Debugging 300)
    Definition for Debug 302
   (Definition for Print_Invert_Type 304)
   ⟨ Definition for Print_Measure_Type 306⟩
```

 $\{248 \quad \text{IAD (v 3-12-0)} \quad \text{IAD UTILITIES} \quad 105$

```
248.
          \langle iad\_util.h \quad 248 \rangle \equiv
   \langle Prototype for What_Is_B \ 249 \rangle;
   \langle Prototype for Estimate\_RT \ 255 \rangle;
    Prototype for a2acalc \ 261;
     Prototype for acalc2a \ 263;
    (Prototype for g2gcalc \ 265);
    \langle \text{ Prototype for } gcalc2g \ 267 \rangle;
    (Prototype for b2bcalc 269);
    (Prototype for bcalc2b 271);
   \langle \text{ Prototype for } twoprime \ 273 \rangle;
   \langle Prototype for two unprime 275 \rangle;
    \langle Prototype for abgg2ab 277 \rangle;
    \langle \text{ Prototype for } abgb2ag 279 \rangle;
    \langle Prototype for quick_quess 286 \rangle;
   \langle Prototype for Set\_Debugging 299 \rangle;
    \langle Prototype for Debug 301 \rangle;
   \langle Prototype for Print_Invert_Type 303 \rangle;
   \langle Prototype for Print\_Measure\_Type 305 \rangle;
```

249. Finding optical thickness.

This routine figures out what the optical thickness of a slab based on the index of refraction of the slab and the amount of collimated light that gets through it.

It should be pointed out right here in the front that this routine does not work for diffuse irradiance, but then the whole concept of estimating the optical depth for diffuse irradiance is bogus anyway.

In version 1.3 changed all error output to *stderr*. Version 1.4 included cases involving absorption in the boundaries.

```
#define BIG_A_VALUE 999999.0

#define SMALL_A_VALUE 0.000001

⟨Prototype for What_Is_B 249⟩ ≡
double What_Is_B (struct AD_slab_type slab, double Tc)

This code is used in sections 248 and 250.

250. ⟨Definition for What_Is_B 250⟩ ≡
⟨Prototype for What_Is_B 249⟩

{
double r1, r2, t1, t2, mu_in_slab;
⟨Calculate specular reflection and transmission 251⟩
⟨Check for bad values of Tc 252⟩
⟨Solve if multiple internal reflections are not present 253⟩
⟨Find thickness when multiple internal reflections are present 254⟩
}

This code is used in section 247.
```

251. The first thing to do is to find the specular reflection for light interacting with the top and bottom airglass-sample interfaces. I make a simple check to ensure that the the indices are different before calculating the bottom reflection. Most of the time the $r1 \equiv r2$, but there are always those annoying special cases.

```
 \begin{array}{l} \langle \, \text{Calculate specular reflection and transmission } \, 251 \, \rangle \equiv \\ \, Absorbing\_Glass\_RT \, (1.0, slab.n\_top\_slide, slab.n\_slab, slab.cos\_angle, slab.b\_top\_slide, \&r1\,,\&t1 \, ); \\ \, mu\_in\_slab = \, Cos\_Snell \, (1.0, slab.cos\_angle, slab.n\_slab); \\ \, Absorbing\_Glass\_RT \, (slab.n\_slab, slab.n\_bottom\_slide, 1.0, mu\_in\_slab, slab.b\_bottom\_slide, \&r2\,,\&t2 \, ); \\ \, \text{This code is used in section } 250. \end{array}
```

Bad values for the unscattered transmission are those that are non-positive, those greater than one, and those greater than are possible in a non-absorbing medium, i.e.,

$$T_c > \frac{t_1 t_2}{1 - r_1 r_2}$$

Since this routine has no way to report errors, I just set the optical thickness to the natural values in these cases.

 $\langle\, {\rm Check} \ {\rm for} \ {\rm bad} \ {\rm values} \ {\rm of} \ Tc \ {\scriptstyle 252} \,\rangle \equiv$ if $(Tc \leq 0)$ return (HUGE_VAL); if $(Tc \ge t1 * t2/(1 - r1 * r2))$ return (0.001);

This code is used in section 250.

253. If either r1 or $r2 \equiv 0$ then things are very simple because the sample does not sustain multiple internal reflections and the unscattered transmission is

$$T_c = t_1 t_2 \exp(-b/\nu)$$

where b is the optical thickness and ν is slab.cos_angle. Clearly,

$$b = -\nu \ln \left(\frac{T_c}{t_1 t_2} \right)$$

 \langle Solve if multiple internal reflections are not present 253 $\rangle \equiv$ if $(r1 \equiv 0 \lor r2 \equiv 0)$ return $(-slab.cos_angle * log(Tc/t1/t2));$ This code is used in section 250.

254. Well I kept putting it off, but now comes the time to solve the following equation for b

$$T_c = \frac{t_1 t_2 \exp(-b)}{1 - r_1 r_2 \exp(-2b)}$$

We note immediately that this is a quadratic equation in $x = \exp(-b)$.

$$r_1 r_2 T_c x^2 + t_1 t_2 x - T_c = 0$$

Sufficient tests have been made above to ensure that none of the coefficients are exactly zero. However, it is clear that the leading quadratic term has a much smaller coefficient than the other two. Since r_1 and r_2 are typically about four percent the product is roughly 10^{-3} . The collimated transmission can be very small and this makes things even worse. A further complication is that we need to choose the only positive root.

Now the roots of $ax^2 + bx + c = 0$ can be found using the standard quadratic formula,

$$x = \frac{-b \pm \sqrt{b^2 - 4ac}}{2a}$$

This is very bad for small values of a. Instead I use

$$q = -\frac{1}{2} \left[b + \operatorname{sgn}(b) \sqrt{b^2 - 4ac} \right]$$

with the two roots

$$x = \frac{q}{a}$$
 and $x = \frac{c}{q}$

Substituting our coefficients

$$q = -\frac{1}{2} \left[t_1 t_2 + \sqrt{t_1^2 t_2^2 + 4r_1 r_2 T_c^2} \right]$$

With some algebra, this can be shown to be

$$q = -t_1 t_2 \left[1 + \frac{r_1 r_2 T_c^2}{t_1^2 t_2^2} + \cdots \right]$$

The only positive root is $x = -T_c/q$. Therefore

$$x = \frac{2T_c}{t_1 t_2 + \sqrt{t_1^2 t_2^2 + 4r_1 r_2 T_c^2}}$$

(Not very pretty, but straightforward enough.)

```
\langle Find thickness when multiple internal reflections are present 254 \rangle \equiv \{ double B; B = t1 * t2; return (-slab.cos\_angle * log(2 * Tc/(B + sqrt(B * B + 4 * Tc * Tc * r1 * r2)))); \}
```

This code is used in section 250.

108 ESTIMATING R AND T IAD (v 3-12-0) $\S255$

255. Estimating R and T.

In several places, it is useful to know an *estimate* for the values of the reflection and transmission of the sample based on the measurements. This routine provides such an estimate, but it currently ignores anything corrections that might be made for the integrating spheres.

Good values are only really obtainable when $num_measures \equiv 3$, otherwise we need to make pretty strong assumptions about the reflection and transmission values. If $num_measures < 3$, then we will assume that no collimated light makes it all the way through the sample. The specular reflection is then just that for a semi-infinite sample and Tc = 0. If $num_measures \equiv 1$, then Td is also set to zero.

```
rt
                   total reflection
                   primary or specular reflection
          rc
                   diffuse or scattered reflection
          rd
          tt
                   total transmission
                   primary or unscattered transmission
          tp
                   diffuse or scattered transmission
          td
\langle \text{ Prototype for } Estimate\_RT | 255 \rangle \equiv
  void Estimate\_RT(struct measure_type m, struct invert_type r, double *rt, double *tt, double
        *rd, double *rc, double *td, double *tc)
This code is used in sections 248 and 256.
256. \langle \text{ Definition for } Estimate\_RT | 256 \rangle \equiv
   \langle Prototype for Estimate\_RT \ 255 \rangle
     (Calculate the unscattered transmission and reflection 257)
     \langle Estimate the backscattered reflection 258\rangle
     (Estimate the scattered transmission 259)
```

257. If there are three measurements then the specular reflection can be calculated pretty well. If there are fewer then the unscattered transmission is assumed to be zero. This is not necessarily the case, but after all, this routine only makes estimates of the various reflection and transmission quantities.

If there are three measurements, the optical thickness of the sample is required. Of course if there are three measurements then the illumination must be collimated and we can call *What_Is_B* to find out the optical thickness. We pass this value to a routine in the fresnel.h unit and sit back and wait.

All the above is true if sphere corrections are not needed. Now, we just fob this off on another function.

 \langle Calculate the unscattered transmission and reflection 257 \rangle \equiv Calculate_Minimum_MR(m, r, rc, tc);

This code is used in section 256.

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258. Finding the diffuse reflection is now just a matter of checking whether V1% contains the specular reflection from the sample or not and then just adding or subtracting the specular reflection as appropriate.

This code is used in section 256.

259. The transmission values follow in much the same way as the diffuse reflection values — just subtract the specular transmission from the total transmission.

```
\langle Estimate the scattered transmission 259\rangle \equiv
  if (m.num\_measures \equiv 1) {
     *tt = 0.0;
     *td = 0.0;
  }
  else if (m.fraction_of_tc_in_mt) {
     *tt = m.m_{-}t;
     *td = *tt - *tc;
     if (*td < 0) {
        *tc = *tt;
        *td = 0;
     }
  else {
     *td = m.m_t;
     *tt = *td + *tc;
  if (Debug(DEBUG_SEARCH)) {
     fprintf(stderr, "_{""}tt_{""}tt_{"}=_{"}\%.5f\n",*tt);
     fprintf(stderr, "_{\sqcup\sqcup\sqcup\sqcup} est_{\sqcup}td_{\sqcup} = _{\sqcup}\%.5f n", *td);
This code is used in section 256.
```

260. Transforming properties. Routines to convert optical properties to calculation space and back.

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261. a2acalc is used for the albedo transformations according to

$$a_{calc} = \frac{2a - 1}{a(1 - a)}$$

Care is taken to avoid division by zero. Why was this function chosen? Well mostly because it maps the region between $[0,1] \to (-\infty, +\infty)$.

```
\langle \text{ Prototype for } a2acalc \text{ 261} \rangle \equiv
   double a2acalc (double a)
```

This code is used in sections 248 and 262.

```
262. \langle Definition for a2acalc \ 262 \rangle \equiv
   \langle \text{ Prototype for } a2acalc \text{ 261} \rangle
     if (a \le 0) return -BIG_A_VALUE;
     if (a \ge 1) return BIG_A_VALUE;
     return ((2*a-1)/a/(1-a));
```

This code is used in section 247.

263. acalc2a is used for the albedo transformations Now when we solve

$$a_c a l c = \frac{2a - 1}{a(1 - a)}$$

we obtain the quadratic equation

This code is used in section 247.

$$a_{calc}a^2 + (2 - a_{calc})a - 1 = 0$$

The only root of this equation between zero and one is

$$a = \frac{-2 + a_{calc} + \sqrt{a_{calc}^2 + 4}}{2a_{calc}}$$

I suppose that I should spend the time to recast this using the more appropriate numerical solutions of the quadratic equation, but this worked and I will leave it as it is for now.

```
\langle \text{ Prototype for } acalc2a \text{ 263} \rangle \equiv
  double acalc2a(double acalc)
This code is used in sections 248 and 264.
264. \langle Definition for acalc2a 264\rangle \equiv
   \langle \text{ Prototype for } acalc2a \text{ 263} \rangle
     if (acalc \equiv BIG_A_VALUE) return 1.0;
     else if (acalc \equiv -BIG_A_VALUE) return 0.0;
     else if (fabs(acalc) < SMALL_A_VALUE) return 0.5;
     else return ((-2 + acalc + sqrt(acalc * acalc + 4))/(2 * acalc));
```

265. *g2gcalc* is used for the anisotropy transformations according to

$$g_{calc} = \frac{g}{1 + |g|}$$

```
which maps (-1,1) \rightarrow (-\infty, +\infty).

\langle \text{Prototype for } g2gcalc | 265 \rangle \equiv \text{double } g2gcalc | (\text{double } g)

This code is used in sections 248 and 266.

266. \langle \text{Definition for } g2gcalc | 266 \rangle \equiv \langle \text{Prototype for } g2gcalc | 265 \rangle = \{ \text{if } (g \leq -1) \text{ return } (-\text{HUGE\_VAL}); \text{ if } (g \geq 1) \text{ return } (\text{HUGE\_VAL}); \text{ return } (g/(1 - fabs(g))); \}

This code is used in section 247.
```

267. gcalc2g is used for the anisotropy transformations it is the inverse of g2gcalc. The relation is

$$g = \frac{g_{calc}}{1 + |g_{calc}|}$$

```
double gcalc2g(double gcalc)
This code is used in sections 248 and 268.

268. ⟨Definition for gcalc2g 268⟩ ≡
⟨Prototype for gcalc2g 267⟩
{
   if (gcalc ≡ -HUGE_VAL) return -1.0;
   if (gcalc ≡ HUGE_VAL) return 1.0;
   return (gcalc/(1 + fabs(gcalc)));
}
This code is used in section 247.
```

 $\langle \text{ Prototype for } gcalc2q \text{ 267} \rangle \equiv$

269. b2bcalc is used for the optical depth transformations it is the inverse of bcalc2b. The relation is

$$b_{calc} = \ln(b)$$

The only caveats are to ensure that I don't take the logarithm of something big or non-positive.

```
\langle \text{ Prototype for } b2bcalc \text{ 269 } \rangle \equiv  double b2bcalc \text{ (double } b)
```

This code is used in sections 248 and 270.

```
270.
          \langle \text{ Definition for } b2bcalc | 270 \rangle \equiv
   \langle Prototype for b2bcalc 269 \rangle
      if (b \equiv \text{HUGE\_VAL}) return HUGE_VAL;
     if (b \le 0) return 0.0;
      return (log(b));
This code is used in section 247.
```

bcalc2b is used for the anisotropy transformations it is the inverse of b2bcalc. The relation is

$$b = \exp(b_{calc})$$

The only tricky part is to ensure that I don't exponentiate something big and get an overflow error. In ANSI C the maximum value for x such that 10^x is in the range of representable finite floating point numbers (for doubles) is given by DBL_MAX_10_EXP. Thus if we want to know if

$$e^{b_{calc}} > 10^x$$

or

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$$b_{calc} > x \ln(10) \approx 2.3x$$

and this is the criterion that I use.

```
\langle \text{ Prototype for } bcalc2b \text{ 271} \rangle \equiv
   double bcalc2b(double bcalc)
```

This code is used in sections 248 and 272.

```
272. \langle \text{ Definition for } bcalc2b | 272 \rangle \equiv
   \langle \text{ Prototype for } bcalc2b \text{ 271} \rangle
      if (bcalc \equiv HUGE_VAL) return HUGE_VAL;
     if (bcalc > 2.3 * DBL_MAX_10_EXP) return HUGE_VAL;
      return (exp(bcalc));
```

This code is used in section 247.

273. twoprime converts the true albedo a, optical depth b to the reduced albedo ap and reduced optical depth bp that correspond to g = 0.

```
\langle \text{ Prototype for } twoprime \ 273 \rangle \equiv
```

void $twoprime(\mathbf{double}\ a, \mathbf{double}\ b, \mathbf{double}\ g, \mathbf{double}\ *ap, \mathbf{double}\ *bp)$

This code is used in sections 248 and 274.

```
274. \langle Definition for two prime 274 \rangle \equiv
   \langle \text{ Prototype for } twoprime 273 \rangle
      if (a \equiv 1 \land g \equiv 1) *ap = 0.0;
      else *ap = (1 - g) * a/(1 - a * g);
      if (b \equiv \text{HUGE\_VAL}) *bp = \text{HUGE\_VAL};
      else *bp = (1 - a * g) * b;
```

275. two unprime converts the reduced albedo ap and reduced optical depth bp (for g=0) to the true albedo a and optical depth b for an anisotropy g.

```
 \begin{array}{l} \langle \operatorname{Prototype} \ \text{for} \ twounprime} \ \ 275 \rangle \equiv \\ \mathbf{void} \ twounprime} \ (\mathbf{double} \ ap, \mathbf{double} \ bp, \mathbf{double} \ g, \mathbf{double} \ *a, \mathbf{double} \ *b) \\ \text{This code is used in sections 248 and 276.} \\ \mathbf{276.} \quad \langle \operatorname{Definition} \ \text{for} \ twounprime} \ \ 276 \rangle \equiv \\ \langle \operatorname{Prototype} \ \text{for} \ twounprime} \ \ 275 \rangle \\ \{ \\ *a = ap/(1-g+ap*g); \\ \text{if} \ (bp \equiv \operatorname{HUGE\_VAL}) \ *b = \operatorname{HUGE\_VAL}; \\ \text{else} \ *b = (1+ap*g/(1-g))*bp; \\ \} \end{array}
```

277. abgg2ab assume a, b, g, and g1 are given this does the similarity translation that you would expect it should by converting it to the reduced optical properties and then transforming back using the new value of g

```
⟨Prototype for abgg2ab 277⟩ ≡
    void abgg2ab (double a1, double b1, double g1, double g2, double *a2, double *b2)
This code is used in sections 248 and 278.

278. ⟨Definition for abgg2ab 278⟩ ≡
    ⟨Prototype for abgg2ab 277⟩
{
    double a, b;
    twoprime(a1, b1, g1, &a, &b);
    twounprime(a, b, g2, a2, b2);
}
```

This code is used in section 247.

This code is used in section 247.

279. abgb2ag translates reduced optical properties to unreduced values assuming that the new optical thickness is given i.e., a1 and b1 are a' and b' for g=0. This routine then finds the appropriate anisotropy and albedo which correspond to an optical thickness b2.

If both b1 and b2 are zero then just assume g=0 for the unreduced values.

```
\langle \text{Prototype for } abgb2ag \ 279 \rangle \equiv  void abgb2ag (\text{double } a1, \text{double } b1, \text{double } b2, \text{double } *a2, \text{double } *g2) This code is used in sections 248 and 280.
```

```
280.
         \langle \text{ Definition for } abqb2aq 280 \rangle \equiv
   \langle \text{ Prototype for } abgb2ag 279 \rangle
     if (b1 \equiv 0 \lor b2 \equiv 0) {
        *a2 = a1;
        *g2 = 0;
     if (b2 < b1) b2 = b1;
     if (a1 \equiv 0) *a2 = 0.0;
     else {
        if (a1 \equiv 1) *a2 = 1.0;
        else {
           if (b1 \equiv 0 \lor b2 \equiv \text{HUGE\_VAL}) *a2 = a1;
           else *a2 = 1 + b1/b2 * (a1 - 1);
     if (*a2 \equiv 0 \lor b2 \equiv 0 \lor b2 \equiv \text{HUGE\_VAL}) *g2 = 0.5;
     else *g2 = (1 - b1/b2)/(*a2);
This code is used in section 247.
281. Guessing an inverse.
   This routine is not used anymore.
\langle \text{ Prototype for } slow\_guess | 281 \rangle \equiv
   void slow\_quess(struct measure_type m, struct invert_type *r, double *a, double *b, double *g)
This code is used in section 282.
        \langle \text{ Definition for } slow\_guess | 282 \rangle \equiv
   ⟨ Prototype for slow_guess 281 ⟩
     double fmin = 10.0;
     double fval;
     double *x;
     x = dvector(1, 2);
     switch (r \rightarrow search) {
     case FIND_A: \langle Slow guess for a alone 283\rangle
        break;
     case FIND_B: \langle Slow guess for b alone 284\rangle
        break:
     case FIND_AB: case FIND_AG: \langle Slow guess for a and b or a and g 285\rangle
        break;
     *a = r \rightarrow slab.a;
     *b = r \rightarrow slab.b;
     *g = r \rightarrow slab.g;
     free\_dvector(x, 1, 2);
```

```
283.
          \langle \text{Slow guess for } a \text{ alone } 283 \rangle \equiv
   r \rightarrow slab.b = HUGE\_VAL;
   r \rightarrow slab.g = r \rightarrow default\_g;
   Set\_Calc\_State(m, *r);
   for (r \rightarrow slab.a = 0.0; r \rightarrow slab.a \le 1.0; r \rightarrow slab.a += 0.1) {
      fval = Find\_A\_fn(a2acalc(r \rightarrow slab.a));
      if (fval < fmin) {
         r \rightarrow a = r \rightarrow slab.a;
         fmin = fval;
   }
   r \rightarrow slab.a = r \rightarrow a;
This code is used in section 282.
284. Presumably the only time that this will need to be called is when the albedo is fixed or is one. For
now, I'll just assume that it is one.
\langle Slow guess for b alone 284\rangle \equiv
   r \rightarrow slab.a = 1;
   r \rightarrow slab.g = r \rightarrow default_g;
   Set_{-}Calc_{-}State(m, *r);
   for (r \rightarrow slab.b = 1/32.0; r \rightarrow slab.b \le 32; r \rightarrow slab.b *= 2) {
      fval = Find_B fn(b2bcalc(r \rightarrow slab.b));
      if (fval < fmin) {
         r \rightarrow b = r \rightarrow slab.b;
         fmin = fval;
   r \rightarrow slab.b = r \rightarrow b;
This code is used in section 282.
285. (Slow guess for a and b or a and g 285) \equiv
      double min_{-}a, min_{-}b, min_{-}g;
      if (\neg Valid\_Grid(m, r \rightarrow search)) Fill\_Grid(m, *r);
      Near\_Grid\_Points(m.m\_r, m.m\_t, r \rightarrow search, \& min\_a, \& min\_b, \& min\_g);
      r \rightarrow slab.a = min_a;
      r \rightarrow slab.b = min_b;
      r \rightarrow slab.g = min_{g};
This code is used in section 282.
286. \langle \text{Prototype for } quick\_guess \ 286 \rangle \equiv
   void quick\_quess (struct measure_type m, struct invert_type r, double *a, double *b, double *q)
This code is used in sections 248 and 287.
```

§287

This code is used in sections 291, 295, and 296.

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```
287.
        \langle \text{ Definition for } quick\_quess | 287 \rangle \equiv
  ⟨ Prototype for quick_guess 286⟩
     double UR1, UT1, rd, td, tc, rc, bprime, aprime, alpha, beta, logr;
     Estimate\_RT(m, r, \&\mathtt{UR1}, \&\mathtt{UT1}, \&rd, \&rc, \&td, \&tc);
     (Estimate aprime 288)
     switch (m.num\_measures) {
     case 1: (Guess when only reflection is known 290)
       break;
     case 2: (Guess when reflection and transmission are known 291)
       break:
     case 3: (Guess when all three measurements are known 292)
       break;
     (Clean up guesses 297)
This code is used in section 247.
       \langle \text{ Estimate } aprime | 288 \rangle \equiv
  if (UT1 \equiv 1) aprime = 1.0;
  else if (rd/(1 - UT1) \ge 0.1)
     double tmp = (1 - rd - UT1)/(1 - UT1);
     aprime = 1 - 4.0/9.0 * tmp * tmp;
  else if (rd < 0.05 \land UT1 < 0.4) aprime = 1 - (1 - 10 * rd) * (1 - 10 * rd);
  else if (rd < 0.1 \land UT1 < 0.4) aprime = 0.5 + (rd - 0.05) * 4;
     double tmp = (1 - 4 * rd - UT1)/(1 - UT1);
     aprime = 1 - tmp * tmp;
This code is used in section 287.
289. \langle \text{ Estimate } bprime \ 289 \rangle \equiv
  if (rd < 0.01) {
     bprime = What\_Is\_B(r.slab, UT1);
     fprintf(stderr, "low_rd<0.01!_ut1=%f_aprime=%f_bprime=%f\n",UT1, aprime, bprime);</pre>
  else if (UT1 \le 0) bprime = HUGE_VAL;
  else if (UT1 > 0.1) bprime = 2 * exp(5 * (rd - UT1) * log(2.0));
  else {
     alpha = 1/log(0.05/1.0);
     beta = log(1.0)/log(0.05/1.0);
     logr = log(UR1);
     bprime = log(UT1) - beta * log(0.05) + beta * logr;
     bprime /= alpha * log(0.05) - alpha * logr - 1;
```

```
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```

```
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        IAD (v 3-12-0)
290.
\langle Guess when only reflection is known 290\rangle \equiv
  *g = r.default_g;
  *a = aprime/(1 - *q + aprime * (*q));
  *b = HUGE_VAL;
This code is used in section 287.
291. Guess when reflection and transmission are known 291 \ge 10^{-10}
  \langle \text{ Estimate } bprime 289 \rangle
  *g = r.default_g;
  *a = aprime/(1 - *g + aprime **g);
  *b = bprime/(1 - *a * *g);
This code is used in section 287.
292. (Guess when all three measurements are known 292) \equiv
  switch (r.search) {
  case FIND_A: (Guess when finding albedo 293)
  case FIND_B: (Guess when finding optical depth 294)
     break:
  case FIND_AB: (Guess when finding the albedo and optical depth 295)
  case FIND_AG: (Guess when finding anisotropy and albedo 296)
     break;
This code is used in section 287.
293.
\langle Guess when finding albedo 293\rangle \equiv
  *g = r.default_g;
  *a = aprime/(1 - *g + aprime **g);
  *b = What_Is_B(r.slab, m.m_u);
This code is used in section 292.
294.
\langle Guess when finding optical depth 294\rangle \equiv
  *g = r.default_g;
  *a = 0.0;
  *b = What_Is_B(r.slab, m.m_u);
This code is used in section 292.
295.
\langle Guess when finding the albedo and optical depth 295\rangle \equiv
  *g = r.default_g;
  if (*g \equiv 1) *a = 0.0;
  else *a = aprime/(1 - *g + aprime **g);
  (Estimate bprime 289)
  if (bprime \equiv HUGE\_VAL \lor *a **g \equiv 1) *b = HUGE\_VAL;
  else *b = bprime/(1 - *a * *g);
This code is used in section 292.
```

```
296.
\langle Guess when finding anisotropy and albedo 296\rangle \equiv
  *b = What_Is_B(r.slab, m.m_u);
  if (*b \equiv \text{HUGE\_VAL} \lor *b \equiv 0) {
     *a = aprime;
     *g = r.default_g;
  else {
     ⟨Estimate bprime 289⟩
     *a = 1 + bprime * (aprime - 1)/(*b);
     if (*a < 0.1) *g = 0.0;
     else *g = (1 - bprime/(*b))/(*a);
This code is used in section 292.
297.
\langle Clean up guesses 297\rangle \equiv
  if (*a < 0) *a = 0.0;
  if (*g < 0) *g = 0.0;
  else if (*g \ge 1) *g = 0.5;
This code is used in section 287.
298.
        Some debugging stuff.
         \langle Prototype for Set_Debugging 299 \rangle \equiv
  void Set_Debugging(unsigned long debug_level)
This code is used in sections 248 and 300.
300.
\langle \text{ Definition for } Set\_Debugging 300 \rangle \equiv
  \langle \text{ Prototype for } Set\_Debugging 299 \rangle
     g\_util\_debugging = debug\_level;
This code is used in section 247.
301.
\langle \text{ Prototype for } Debug | 301 \rangle \equiv
  int Debug(unsigned long mask)
This code is used in sections 248 and 302.
302.
\langle \text{ Definition for } Debug 302 \rangle \equiv
  \langle \text{ Prototype for } Debug 301 \rangle
     if (g_util_debugging & mask) return 1;
     else return 0;
```

```
SOME DEBUGGING STUFF
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```

```
§303
                                             IAD (v 3-12-0)
303.
\langle Prototype for Print_Invert_Type 303 \rangle \equiv
             void Print_Invert_Type(struct invert_type r)
This code is used in sections 248 and 304.
304.
\langle \text{ Definition for } Print\_Invert\_Type \ 304 \rangle \equiv
              ⟨ Prototype for Print_Invert_Type 303⟩
                          fprintf(stderr, "\n");
                         fprintf(stderr, "default_{\sqcup\sqcup}a=\%10.5f_{\sqcup\sqcup\sqcup\sqcup}b=\%10.5f_{\sqcup\sqcup\sqcup\sqcup}g=\%10.5f\\n", r.default_a, r.default_b, r.default_g);
                        fprintf(stderr, "slab_{\cup \cup \cup \cup \cup} a=\%10.5f_{\cup \cup \cup \cup} b=\%10.5f_{\cup \cup \cup \cup} g=\%10.5f_{\setminus n}", r.slab.a, r.slab.b, r.slab.g);
                        fprintf (stderr, \verb"nullulultop=%10.5flmid=%10.5flulbot=%10.5f\n", r.slab.n_top\_slide, r.slab.n_slab, n_top_slide, r.slab.n_top_slide, r.slab.n_t
                                                  r.slab.n\_bottom\_slide);
                        fprintf (stderr, \verb"thick_ultop=\%10.5f_ucos=\%10.5f_ulbot=\%10.5f_n", r.slab.b\_top\_slide, r.slab.cos\_angle, r.slab.b\_top\_slide, r.slab.cos\_angle, r.slab.b\_top\_slide, r
                                                 r.slab.b\_bottom\_slide);
                         fprintf(stderr, "search_= "\&d_quadrature_points_= "\&d\n", r.search, r.method.quad_pts);
This code is used in section 247.
305.
\langle Prototype for Print\_Measure\_Type 305 \rangle \equiv
              void Print_Measure_Type(struct measure_type m)
This code is used in sections 248 and 306.
```

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```
306.
```

```
\langle \text{ Definition for } Print\_Measure\_Type 306 \rangle \equiv
       (Prototype for Print_Measure_Type 305)
             fprintf(stderr, "\n");
              fprintf(stderr, "\#_{"} = "\%7.1f_{"} m\n", m.slab_thickness);
             m.slab\_top\_slide\_thickness);
             \mathit{fprintf} \, (\mathit{stderr}, \verb"#_UUUUUUUUUUUUUUUB \verb"ottom_Uslide_Uthickness_U=_U\%7.1 f_Umm \verb"n"}, \\
                            m.slab\_bottom\_slide\_thickness);
             fprintf(stderr, "\#_{$\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup}Sample_{$\sqcup$index}_{$\sqcup$fraction}_{$\sqcup$}%7.3f\n", m.slab\_index);
              fprintf(stderr, "\#_{\cup\cup\cup\cup\cup\cup\cup\cup\cup} Top_{\cup} slide_{\cup} index_{\cup} of_{\cup} refraction_{\cup} = _{\cup} %7.3f \ ", m.slab\_top\_slide\_index);
              fprintf(stderr, "\#_{\sqcup\sqcup\sqcup\sqcup\sqcup\sqcup} Bottom_{\sqcup} slide_{\sqcup} index_{\sqcup} of_{\sqcup} refraction_{\sqcup} = \cline{``locality}.3f\n", m.slab_bottom_slide_index);
             fprintf(stderr, "\#_{\sqcup\sqcup\sqcup\sqcup} Fraction_{\sqcup}unscattered_{\sqcup} light_{\sqcup}in_{\sqcup}M_R_{\sqcup}=_{\sqcup}\%.1f_{\sqcup}\%\%n",
                            m.fraction\_of\_rc\_in\_mr * 100);
              fprintf(stderr, "\#_{\sqcup\sqcup\sqcup\sqcup\sqcup} Fraction_{\sqcup}unscattered_{\sqcup} light_{\sqcup}in_{\sqcup}M_{T_{\sqcup}}=_{\sqcup}\%7.1f_{\sqcup}\%\n",
                             m.fraction\_of\_tc\_in\_mt * 100);
              fprintf(stderr, "#_{\sqcup}\n");
              fprintf(stderr, "\#_{\sqcup}Reflection_{\sqcup}sphere\n");
              fprintf(stderr, "\#_{\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup\cup} sample_{\cup} port_{\cup} diameter_{\cup} = _{\cup}\%7.1f_{\cup}mm\n",
                             2 * m.d\_sphere\_r * sqrt(m.as\_r);
              2*m.d\_sphere\_r*sqrt(m.ae\_r);
              2 * m.d\_sphere\_r * sgrt(m.ad\_r):
              fprintf(stderr, "#_{UUUUUUUUUUUUUUUUUUUdetector_lreflectance_l=l_%7.1f_l_%%\n", m.rd_r*100);
              fprintf(stderr, "area_r_as=\%10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}10.5f_{\coloredge}
                            m.ae_r, m.aw_r);
             fprintf(stderr, "refls_{\sqcup\sqcup}rd=\%10.5f_{\sqcup\sqcup}rw=\%10.5f_{\sqcup\sqcup\sqcup}rstd=\%10.5f_{\sqcup\sqcup\sqcup}f=\%10.5f \ , m.rd\_r, m.rw\_r, \ 
                             m.rstd_r, m.f_r);
             fprintf(stderr, "area_t|_as=\%10.5f_{|||}ad=\%10.5f_{|||||}ae=\%10.5f_{||||}av=\%10.5f_{|||}av=\%10.5f_{|||}, m.as_t, m.ad_t,
                            m.ae_{-t}, m.aw_{-t});
              fprintf(stderr, "refls_{\sqcup\sqcup} rd=\%10.5f_{\sqcup\sqcup} rw=\%10.5f_{\sqcup\sqcup} rstd=\%10.5f_{\sqcup\sqcup\sqcup} f=\%10.5f \ ".m.rd_t, m.rd_t, m.rw_t, m.rd_t, m.rw_t, m.rd_t, m.rd_t, m.rw_t, m.r
                             m.rstd_-t, m.f_-t);
             fprintf(stderr, "lost_{\sqcup\sqcup}ur1=\%10.5f_{\sqcup}ut1=\%10.5f_{\sqcup\sqcup\sqcup}uru=\%10.5f_{\sqcup\sqcup}utu=\%10.5f \ ".m.ur1\_lost,"
                            m.ut1\_lost, m.utu\_lost, m.utu\_lost);
```

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