# The Adding-Doubling Program

 $({\rm Version}\ 3\text{-}11\text{-}5)$ 

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AD GLOBAL VARIABLES

AD Global Variables. Global Routines and Variables. Changed version to reflect bug fix in the Fresnel routine section.

Revised in May 1995 to allow slides to absorb and various modifications to improve the way that the file looks.

```
Revision May 1996 to remove uninitialized tfluence
  Revision May 1998 to improve wrarray.
\langle ad\_globl.c 1 \rangle \equiv
#include <math.h>
#include <stdio.h>
#include <stdlib.h>
#include "ad_globl.h"
#include "ad_frsnl.h"
   (Global variables for adding-doubling 11)
   Definition for Zero\_Layer 16 \rangle
   (Definition for AD_{-error} 14)
   \langle \text{ Definition for } URU\_and\_UR1 | 22 \rangle
   (Definition for URU\_and\_UR1\_Cone 18)
    Definition for URU_and_URx_Cone 20
   \langle \text{ Definition for } UFU\_and\_UF1 \text{ 24} \rangle
   (Definition for wrmatrix 26)
   (Definition for wrarray 28)
2. \langle ad\_globl.h 2 \rangle \equiv
  ⟨ Preprocessor definitions ⟩
   (Types to export from AD Globals 9)
   (External variables to export from AD Globals 12)
   \langle Prototype for Zero\_Layer 15 \rangle;
   \langle \text{ Prototype for } AD\_error \ 13 \rangle;
    Prototype for URU_and_UR1 21);
   \langle Prototype for URU\_and\_UR1\_Cone 17 \rangle;
   \langle Prototype for URU\_and\_URx\_Cone 19 \rangle;
   \langle Prototype for UFU_and_UF1 \ 23 \rangle;
    Prototype for wrmatrix 25;
   \langle \text{ Prototype for } wrarray | 27 \rangle;
```

#### 3. Constants.

This is Version 2.0.0 of the adding-doubling code. (The inverse adding-doubling code may have a different version number.)

4. The number of quadrature points determines how accurately the integrals are performed. Larger numbers of quadrature points lead to more accurate solutions. Fewer points yield much faster computations since the computation time is proportional to  $n^3$  or  $n^2 \ln n$  because an  $n \times n$  matrix must be inverted.

For most practical purposes four quadrature points is plenty. However, if you need very accurate reflection and transmission values, then increase the number of quadrature points. For example, if you want to verify a Monte Carlo implementation, then just crank the number up to 16 or 32 and you are almost certain to get 5 significant digits in your answer.

The number of quadrature points does not need to be a power of 2, but it should be an even number. If it isn't then somewhere in the bowels of this program it will get changed. Finally, if you are unsure of how accurate a solution is, then increase the number of quadrature points and repeat the algorithm.

There is no intrinsic reason that the maximum number of quadrature points is limited to 128. If you have enough memory then this number can be increased. But if you have read the stuff above, my feeling is, why bother?

```
#define MAX_QUAD_PTS 128
#define DEFAULT_QUAD_PTS 4
```

5. The two permissible phase functions are isotropic and Henyey-Greenstein.

```
#define ISOTROPIC 0
#define HENYEY_GREENSTEIN 1
```

**6.** The last two constants are related to the details of how the initial adding-doubling layer is generated. It is very unlikely that these will ever be used by anyone.

```
#define DIAMOND 0
#define INFINITESIMAL_GENERATOR 1
```

7. This last define is so that intermediate values can be generated during the calculation of the initial layer matrices. It is named after Martin Hammer who requested it.

```
#define MARTIN_HAMMER 1
```

8. And finally something for whether the light is conical or oblique

```
#define CONE 1 #define OBLIQUE 0
```

#### 9. Types.

#endif

This code is used in section 2.

The fundamental structure for an adding-doubling calculation keeps all the details of the optical properties of the sample together. The sample is bounded by a glass slide above and below. The glass slides have indicies of refraction  $n\_top\_slide$  and  $n\_bottom\_slide$ . The glass slides may absorb light, in which case  $b\_top\_slide$  or  $b\_bottom\_slide$  may be non-zero.

The albedo of the slab is denoted a, the optical thickness of the slab by  $b = (\mu_a + \mu_s)d$ , and the average cosine of the phase function by g. The phase function of the slab is restricted to just isotropic and Henyey-Greenstein phase functions at the moment.

```
\langle \text{Types to export from AD Globals } 9 \rangle \equiv
  typedef struct AD_slab_type {
    double a:
    double b;
    double g;
    int phase_function;
    double n\_slab;
    double n\_top\_slide;
    double n\_bottom\_slide;
    double b\_top\_slide;
    double b\_bottom\_slide;
    double cos_angle;
  } slab_type;
See also section 10.
This code is used in section 2.
    \langle \text{ Types to export from AD Globals } 9 \rangle + \equiv
  typedef struct AD_method_type {
    int quad_pts;
    double a\_calc, b\_calc, g\_calc, b\_thinnest;
  } method_type;
     The Martin_Hammer variable only exists to print internal results when testing. Its only a integer and
doesn't take up much space so here it is.
\langle Global variables for adding-doubling 11 \rangle \equiv
#define AD_GLOBAL_SOURCE
  double angle[MAX_QUAD_PTS + 1];
  double weight[MAX_QUAD_PTS + 1];
  double twoaw [MAX_QUAD_PTS + 1];
  int Martin\_Hammer = 0;
This code is used in section 1.
      \langle External variables to export from AD Globals 12\rangle \equiv
12.
#ifndef AD_GLOBAL_SOURCE
  extern double angle[MAX_QUAD_PTS + 1];
  extern double weight[MAX_QUAD_PTS + 1];
  extern double twoaw[MAX_QUAD_PTS + 1];
  extern int Martin_Hammer;
```

13. Global routines. My standard error handler

```
\langle \text{ Prototype for } AD\_error \ 13 \rangle \equiv
   void AD_error(char error_text[])
This code is used in sections 2 and 14.
14. \langle Definition for AD_{-error} 14\rangle \equiv
   \langle \text{ Prototype for } AD\_error \ 13 \rangle
     fprintf(stderr, "Adding-Doubling_error\n");
     fprintf(stderr, "%s\n", error\_text);
     fprintf(stderr, "...now_exiting_to_system...\n");
     exit(1);
This code is used in section 1.
15. \langle \text{Prototype for } Zero\_Layer \ 15 \rangle \equiv
   void Zero\_Layer(int n, double **r, double **t)
This code is used in sections 2 and 16.
     \langle \text{ Definition for } Zero\_Layer | 16 \rangle \equiv
   ⟨ Prototype for Zero_Layer 15⟩
     int i, j;
     for (i = 1; i \le n; i++)
        for (j = 1; j \le n; j ++) {
           t[i][j] = 0.0;
           r[i][j] = 0.0;
     \hat{\mathbf{for}}\ (i=1;\ i \leq n;\ i++)\ t[i][i] = 1/twoaw[i];
This code is used in section 1.
```

17. Figure out the reflection for collimated irradiance returning within a cone whose cosine is mu. Note that mu is defined on the air side of the slab and that mu is the cosine of the angle that the cone makes with the normal to the slab,

$$UR1 \equiv \int_{\mu}^{1} R(\nu', 1) 2\nu' \, d\nu'$$

Similarly for irradiance characterized by diffuse light within a cone one can calculate the amount of reflectance returing within that cone as

URU 
$$\equiv n^2 \int_{\mu}^{1} \int_{0}^{1} R(\nu', \nu'') 2\nu' \, d\nu' 2\nu'' \, d\nu''$$

where,  $n^2$  term is to account for the  $n^2$  law of radiance.

```
\langle\, {\rm Prototype} \mbox{ for } \mbox{\it URU\_and\_UR1\_Cone} \mbox{\ \ } 17 \,\rangle \equiv
```

void  $URU\_and\_UR1\_Cone$  (int n, double  $n\_slab$ , double mu, double \*\*R, double \*URU, double \*UR1) This code is used in sections 2 and 18.

```
\langle \text{ Definition for } URU\_and\_UR1\_Cone \text{ 18} \rangle \equiv
  \langle Prototype for URU\_and\_UR1\_Cone 17 \rangle
     int i, j, last_{-}j;
     double mu\_slab;
     double temp = 0.0;
     if (n\_slab \equiv 1) mu\_slab = mu;
     else mu\_slab = sqrt(n\_slab * n\_slab - 1 + mu * mu)/n\_slab;
     last_{-}j = 1;
     while (angle[last_{-}j] \leq mu\_slab) last_{-}j \leftrightarrow ;
     *URU = 0.0;
     for (i = 1; i \le n; i++) {
       temp = 0.0;
       *URU += temp * twoaw[i];
     *UR1 = temp;
     *URU *= n_{-}slab * n_{-}slab/(1 - mu * mu);
This code is used in section 1.
```

19. Figure out the reflection for oblique irradiance returning from a layer Note that mu is the cosine of the angle that the cone makes with the normal to the slab in air,

$$\text{URx} = \int_{\mu}^{1} R(\nu', \mu) 2\nu' \, d\nu'$$

For diffuse irradiance over the cone, the total flux back URU is somewhat arbitrarily chosen as the that flux returning in the same cone. Specifically as

$$URU = n^2 \int_{\mu}^{1} \int_{\mu}^{1} R(\nu', \nu'') 2\nu' \, d\nu' 2\nu'' \, d\nu''$$

where,  $n^2$  term is to account for the  $n^2$  law of radiance. (If you want the total flux returning within a cone for uniform diffuse illumination then use  $URU_{-}$  and  $UR1_{-}$  Cone.)

```
\langle \text{Prototype for } URU\_and\_URx\_Cone \ 19 \rangle \equiv
```

void  $URU_{-}$  and  $URx_{-}$  Cone (int n, double  $n_{-}$  slab, double mu, double \*\*R, double \*URU, double \*URX) This code is used in sections 2 and 20.

\*URU \*=  $n_{-}slab * n_{-}slab$ ;

This code is used in section 1.

```
6
     \langle \text{ Definition for } URU\_and\_URx\_Cone \ 20 \rangle \equiv
  \langle Prototype for URU\_and\_URx\_Cone 19 \rangle
     int i, j, cone\_index;
     double mu_slab, urx, delta, closest_delta;
     double degrees = 180.0/3.1415926535;
     mu\_slab = sqrt(n\_slab * n\_slab - 1 + mu * mu)/n\_slab;
     closest\_delta = 1;
     cone\_index = n;
     for (i = n; i \ge 1; i--) {
       delta = fabs(angle[i] - mu\_slab);
       if (delta < closest\_delta) {
          closest\_delta = delta;
          cone\_index = i;
       }
     if (fabs(angle[cone\_index] - mu\_slab) > 1 \cdot 10^{-5}) {
       fprintf(stderr, "Something is wrong with the quadrature n");
       fprintf(stderr, "theta_i_= \%5.2f_degrees_or_", acos(mu) * degrees);
       fprintf(stderr, "cos(theta_i)_= \%8.5f\n", mu);
       fprintf(stderr, "theta_t|=|.%5.2f_|)degrees_||or_||", acos(mu\_slab)*degrees);
       fprintf(stderr, "cos(theta_t)_{\sqcup} = \ \%8.5f\n", mu\_slab);
       fprintf(stderr, "\_index\_\_degrees\_cosine\n");
       for (i = n; i \ge 1; i --) {
         fprintf(stderr, "$\_\%5d_{\lumbda\lumbda}\%5.2f_{\lumbda}", i, acos(angle[i])*degrees);
         fprintf(stderr, "$\bot \%8.5f \n", angle[i]);
       fprintf(stderr, "Closest_uquadrature_uangle_uis_ui=\%5d_u", cone_index);
       fprintf(stderr, "or_{\sqcup}cos(theta) = \%8.5f \n", angle[cone_index]);
       fprintf(stderr, "Assuming \_normal \_incidence \n");
     *URU = 0.0;
     for (i = 1; i < n; i ++) {
       urx = 0.0;
       {\bf for}\ (j=1;\ j\le n;\ j+\!\!+)\ urx\ +\!\!=R[i][j]*twoaw[j];
       *URU += urx * twoaw[i];
       if (i \equiv cone\_index) * URx = urx;
```

21. Just add up all the angles up to the critical angle. This is a commonly used convenience function to easily calculate UR1 and URU. We select the entire range of angles by passing  $\cos(\pi/2) = 0$  to the  $URU_{-}and_{-}UR1_{-}Cone$  routine.

```
\langle \text{ Prototype for } URU\_and\_UR1 \text{ 21} \rangle \equiv
  void URU_and_{-}UR1 (int n, double n_slab, double **R, double *URU, double *UR1)
This code is used in sections 2 and 22.
```

This code is used in sections 2 and 26.

```
22.
      \langle \text{ Definition for } URU\_and\_UR1 | 22 \rangle \equiv
   \langle \text{ Prototype for } URU\_and\_UR1 \text{ 21} \rangle
      URU_{-}and_{-}UR1_{-}Cone(n, n_{-}slab, 0.0, R, URU, UR1);
This code is used in section 1.
23. \langle \text{Prototype for } UFU\_and\_UF1 \text{ 23} \rangle \equiv
   void UFU_and_UF1 (int n, double n\_slab, double **Lup, double **Ldown, double *UFU, double
        *UF1)
This code is used in sections 2 and 24.
24. \langle Definition for UFU_and_UF1 \ 24 \rangle \equiv
   \langle Prototype for UFU\_and\_UF1 \ 23 \rangle
     int i, j;
     double temp = 0.0;
     *UFU = 0.0;
     for (j = 1; j \le n; j++) {
        temp = 0.0;
        for (i = 1; i \le n; i++) temp += (Lup[i][j] + Ldown[i][j]) * 2 * weight[i];
        *UFU += twoaw[j] * temp;
     *UF1 = temp * n\_slab * n\_slab;
     *UFU *= n_{-}slab * n_{-}slab/2;
This code is used in section 1.
25. \langle \text{Prototype for } wrmatrix \ 25 \rangle \equiv
  void wrmatrix(int n, double **a)
```

This code is used in sections 2 and 28.

```
26. \langle \text{ Definition for } wrmatrix | 26 \rangle \equiv
  ⟨ Prototype for wrmatrix 25⟩
    int i, j;
     double tflux, flux;
     printf("%9.5f", 0.0);
     for (i = 1; i \le n; i++) printf("%9.5f", angle[i]);
     tflux = 0.0;
     for (i = 1; i \le n; i++) {
       printf ("%9.5f", angle[i]);
       for (j = 1; j \le n; j ++)
         if ((a[i][j] > 10) \lor (a[i][j] < -10)) printf("____*****");
          else printf("\%9.5f", a[i][j]);
       flux = 0.0;
       for (j = 1; j \le n; j++)
         if ((a[i][j] < 10) \land (a[i][j] > -10)) flux += a[i][j] * twoaw[j];
       printf("\%9.5f\n", flux);
       tflux += flux * twoaw[i];
    printf("\%9s","flux_{\sqcup\sqcup\sqcup}");
    for (i = 1; i \le n; i++) {
       flux = 0.0;
       for (j = 1; j \le n; j ++)
         if ((a[j][i] < 10) \land (a[j][i] > -10)) flux += a[j][i] * twoaw[j];
       printf ("%9.5f", flux);
     printf("\%9.5f\n", tflux);
     for (i = 1; i \le (n + 2); i++) printf("********");
     printf("\n\n");
This code is used in section 1.
27. \langle \text{ Prototype for } wrarray | 27 \rangle \equiv
  void wrarray(int n, double *a)
```

```
28.
      \langle \text{ Definition for } wrarray | 28 \rangle \equiv
  \langle \text{ Prototype for } wrarray | 27 \rangle
    int i;
     double sum;
     for (i = 1; i \le n; i++) printf("%9.5f", angle[i]);
     printf("%9s\n", "\_angles");
     sum = 0.0;
     for (i = 1; i \le n; i++) {
       if (a[i] > 10 \lor a[i] < -10) printf("____*****");
       else printf("\%9.5f", a[i]);
       if (a[i] < 10 \land a[i] < -10) sum += a[i];
     printf("%9.5f", sum);
     printf ("%9s\n", "☐ (natural)");
     sum = 0.0;
     for (i = 1; i \le n; i++) {
       if (a[i] > 10 \lor a[i] < -10) printf("____*****");
       else printf("\%9.5f", a[i]/twoaw[i]);
       if (a[i] < 10 \land a[i] < -10) sum += a[i];
     printf("%9.5f", sum);
     printf("%9s\n", "*2aw");
    for (i = 1; i \le (n + 2); i ++) printf("********");
     printf("\n\n");
This code is used in section 1.
29. Just print out an array without mucking
\langle \text{ Prototype for } swrarray | 29 \rangle \equiv
  void swrarray(int n, double *a)
This code is used in section 30.
30. \langle \text{ Definition for } swrarray | 30 \rangle \equiv
  (Prototype for swrarray 29)
     int i:
     double sum;
     for (i = 1; i \le n; i ++) printf("%9.5f", angle[i]);
     printf("%9s\n", "*2aw");
     sum = 0.0;
     for (i = 1; i < n; i ++) {
       if (a[i] > 10 \lor a[i] < -10) printf ("____*****");
       else printf("\%9.5f", a[i]/twoaw[i]);
       if (a[i] < 10 \land a[i] < -10) sum += a[i];
     printf("%9.5f\n", sum);
    for (i = 1; i \le (n + 2); i ++) printf("*********");
     printf("\n\n");
```

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AD Prime. This has the rather stupid name prime because I was at a loss for another. Currently this is poorly commented. The fluence routine has not even been checked. There may or may not be errors associated with the  $n^2$ -law in there. It just needs to be checked.

```
\langle ad\_prime.c 31 \rangle \equiv
#include <math.h>
#include <float.h>
#include <stdio.h>
#include "nr_util.h"
#include "ad_globl.h"
#include "ad_bound.h"
#include "ad_start.h"
#include "ad_doubl.h"
#include "ad_prime.h"
#include "ad_matrx.h"
#include "ad_cone.h"
   \langle \text{ Definition for } RT\_Matrices 35 \rangle
   \langle \text{ Definition for RT } 37 \rangle
    Definition for ez_RT 52
    \langle \text{ Definition for } RTabs | \mathbf{56} \rangle
   ⟨ Definition for Flux_Fluence 66 ⟩
   \langle \text{ Definition for } ez\_RT\_unscattered 54 \rangle
32. \langle ad_prime.h \quad 32 \rangle \equiv
   ⟨ Preprocessor definitions ⟩
   \langle \text{ Prototype for } RT\_Matrices 34 \rangle;
   \langle \text{ Prototype for RT } 36 \rangle;
   \langle \text{ Prototype for } ez\_RT \text{ 51} \rangle;
   \langle \text{ Prototype for } RTabs | 55 \rangle;
    \langle Prototype for Flux_Fluence 65 \rangle;
   \langle Prototype for ez\_RT\_unscattered 53 \rangle;
33. \langle \text{lib\_ad.h } 33 \rangle \equiv
   \langle \text{ Prototype for } ez\_RT \text{ 51} \rangle;
   \langle \text{Prototype for } ez\_RT\_unscattered 53 \rangle;
```

34. R and T Matrix routines. This section contains the routine to calculate the reflection and transmission matrix for a scattering and absorbing slab. Basically you just need to set the number of quadrature points method-quad\_pts and the optical properties (the albedo, anisotropy, optical thickness, and choice of phase function) in slab. Call this routine and get back matrices filled with cool numbers.

```
\langle \text{ Prototype for } RT\_Matrices \ 34 \rangle \equiv
  void RT\_Matrices (int n, struct AD_slab_type *slab, struct AD_method_type *method, double
       **R, double **T)
```

This code is used in sections 32 and 35.

```
35.
       \langle \text{ Definition for } RT\_Matrices | 35 \rangle \equiv
  \langle \text{ Prototype for } RT\_Matrices 34 \rangle
     double d;
     if (n < 3) method-quad_pts = DEFAULT_QUAD_PTS;
     else if (n > MAX_QUAD_PTS) method \neg quad_pts = MAX_QUAD_PTS;
     else if ((n \& 1) \equiv 1) method-quad_pts = n/2 * 2;
     else method \neg quad\_pts = n;
     Choose_Method(slab, method);
     if (slab \rightarrow b < 0) {
        Zero\_Layer(n, R, T);
        return;
     n = method \neg quad\_pts;
     Init\_Layer(*slab, *method, R, T);
     if (slab \rightarrow b \equiv HUGE\_VAL) d = 1.0;
                                                 /* Ignored ... just set it something. */
     else d = method \neg b\_thinnest * slab \neg b/method \neg b\_calc;
     Double\_Until(n, R, T, d, slab \rightarrow b);
This code is used in section 31.
```

## 36. Total reflection and transmission.

RT is the top level routine for accessing the adding-doubling algorithm. By passing the optical paramters characteristic of the slab, this routine will do what it must to return the total reflection and transmission for collimated and diffuse irradiance.

This routine has three different components based on if zero, one, or two boundary layers must be included. If the index of refraction of the slab and the top and bottom slides are all one, then no boundaries need to be included. If the top and bottom slides are identical, then some simplifications can be made and some time saved as a consequence. If the top and bottom slides are different, then the full red carpet treatment is required.

Since the calculation time increases for each of these cases we test for matched boundaries first. If the boundaries are matched then don't bother with boundaries for the top and bottom. Just calculate the integrated reflection and transmission. Similarly, if the top and bottom slides are similar, then quickly calculate these.

```
\langle \text{Prototype for RT 36} \rangle \equiv \text{void RT(int } n, \text{struct AD\_slab\_type } *slab, \text{double *UR1, double *UT1, double *URU, double *UTU)}
This code is used in sections 32 and 37.
```

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```
37.
      \langle \text{ Definition for RT } 37 \rangle \equiv
  \langle \text{ Prototype for RT } 36 \rangle
     (Declare variables for RT 38)
     if (slab \rightarrow cos\_angle \neq 1.0) {
        RT_{-}Cone(n, slab, OBLIQUE, UR1, UT1, URU, UTU);
       return:
     (Validate input parameters 39)
     (Allocate and calculate R and T for homogeneous slab 40)
     if (slab \rightarrow b \equiv 0) {
        Sp_{-}RT(n,*slab,UR1,UT1,URU,UTU);
     else if (slab \neg n\_slab \equiv 1 \land slab \neg n\_top\_slide \equiv 1 \land slab \neg n\_bottom\_slide \equiv 1 \land slab \neg b\_top\_slide \equiv
             0 \land slab \neg b\_bottom\_slide \equiv 0) {
        \langle \text{ Do slab with no boundaries 41} \rangle
     else if (slab \neg n\_top\_slide \equiv slab \neg n\_bottom\_slide \land slab \neg b\_top\_slide \equiv 0 \land slab \neg b\_bottom\_slide \equiv 0) {
        (Allocate and generate top boundary 42)
        (Do slab with matched top and bottom boundaries 43)
        (Free top boundary 44)
     else {
        (Allocate and generate top boundary 42)
        (Allocate and generate bottom boundary 45)
        (Allocate misc matrices 46)
        (Do slab with mismatched boundaries 47)
        (Free misc matrices 48)
        (Free bottom boundary 49)
        (Free top boundary 44)
     \langle \text{ Free R and T } 50 \rangle
This code is used in section 31.
38. \langle \text{ Declare variables for RT } 38 \rangle \equiv
  double **R, **T, **R2, **T2;
  double *R01, *R10, *T01, *T10;
  double *R23, *R32, *T23, *T32;
  double **R02, **R20, **T02, **T20;
  double **R03, **R30, **T03, **T30;
  double **atemp, **btemp;
  struct AD_method_type method;
  *UR1 = -1:
  *URU = -1;
  *UT1 = -1;
  *UTU = -1;
This code is used in section 37.
```

```
ξ39
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39.
\langle \text{ Validate input parameters } 39 \rangle \equiv
  if (slab \neg n\_slab < 0) return;
  if (slab \rightarrow n\_top\_slide < 0) return;
  if (slab \neg n\_bottom\_slide < 0) return;
  if (slab \neg a < 0 \lor slab \neg a > 1) return;
  if (slab \rightarrow g < -1 \lor slab \rightarrow g > 1) return;
  if (slab \rightarrow b < 0) return;
This code is used in section 37.
40. Find the R and T for a homogeneous slab without boundaries
\langle Allocate and calculate R and T for homogeneous slab 40 \rangle \equiv
  R = dmatrix(1, n, 1, n);
  T = dmatrix(1, n, 1, n);
  RT\_Matrices(n, slab, \& method, R, T);
This code is used in sections 37 and 56.
41. \langle Do slab with no boundaries \langle 41 \rangle \equiv
   URU_{-}and_{-}UR1(n, slab \rightarrow n_{-}slab, R, URU, UR1);
   URU_{-}and_{-}UR1 (n, slab \rightarrow n_{-}slab, T, UTU, UT1);
This code is used in section 37.
42. \langle Allocate and generate top boundary \langle \langle \rangle
  RO1 = dvector(1, n);
  R10 = dvector(1, n);
  T01 = dvector(1, n);
  T10 = dvector(1, n);
  Init\_Boundary (*slab, method. quad\_pts, R01, R10, T01, T10, T0P_BOUNDARY);
This code is used in sections 37 and 60.
43. (Do slab with matched top and bottom boundaries 43) \equiv
  atemp = dmatrix(1, n, 1, n);
  btemp = dmatrix(1, n, 1, n);
  R2 = dmatrix(1, n, 1, n);
  T2 = dmatrix(1, n, 1, n);
  Add\_Slides(n, R01, R10, T01, T10, R, T, R2, T2, atemp, btemp);
  URU_{-}and_{-}UR1 (n, slab \rightarrow n_{-}slab, R2, URU, UR1);
  URU_{-}and_{-}UR1 (n, slab \rightarrow n_{-}slab, T2, UTU, UT1);
  free\_dmatrix(atemp, 1, n, 1, n);
  free\_dmatrix(btemp, 1, n, 1, n);
  free\_dmatrix(R2, 1, n, 1, n);
  free\_dmatrix(T2, 1, n, 1, n);
This code is used in section 37.
44. \langle Free top boundary 44 \rangle \equiv
  free\_dvector(RO1, 1, n);
  free\_dvector(R10, 1, n);
```

 $free\_dvector(\texttt{T01},1,n); \\ free\_dvector(\texttt{T10},1,n); \\ This code is used in sections 37 and 56.$ 

```
45.
      \langle Allocate and generate bottom boundary 45\rangle \equiv
  R23 = dvector(1, n);
  \texttt{R32} = dvector(1, n);
  T23 = dvector(1, n);
  T32 = dvector(1, n);
  Init_Boundary (*slab, method.quad_pts, R23, R32, T23, T32, BOTTOM_BOUNDARY);
This code is used in sections 37 and 61.
46. \langle Allocate misc matrices |46\rangle \equiv
  R02 = dmatrix(1, n, 1, n);
  R20 = dmatrix(1, n, 1, n);
  T02 = dmatrix(1, n, 1, n);
  T20 = dmatrix(1, n, 1, n);
  R03 = dmatrix(1, n, 1, n);
  R30 = dmatrix(1, n, 1, n);
  T03 = dmatrix(1, n, 1, n);
  T30 = dmatrix(1, n, 1, n);
  atemp = dmatrix(1, n, 1, n);
  btemp = dmatrix(1, n, 1, n);
This code is used in sections 37 and 56.
47. \langle Do slab with mismatched boundaries 47\rangle \equiv
  Add_{-}Top(n, R01, R10, T01, T10, R, R, T, T, R02, R20, T02, T20, atemp, btemp);
  Add\_Bottom(n, RO2, R20, TO2, T20, R23, R32, T23, T32, RO3, R30, T03, T30, atemp, btemp);
  URU_{-}and_{-}UR1 (n, slab \rightarrow n_{-}slab, RO3, URU, UR1);
  Transpose\_Matrix(n, T03);
  URU_{-}and_{-}UR1 (n, slab \rightarrow n_{-}slab, TO3, UTU, UT1);
This code is used in section 37.
48. \langle Free misc matrices 48 \rangle \equiv
  free\_dmatrix(RO2, 1, n, 1, n);
  free\_dmatrix(R20, 1, n, 1, n);
  free\_dmatrix(T02, 1, n, 1, n);
  free\_dmatrix(T20, 1, n, 1, n);
  free\_dmatrix(RO3, 1, n, 1, n);
  free\_dmatrix(R30, 1, n, 1, n);
  free\_dmatrix(TO3, 1, n, 1, n);
  free\_dmatrix(T30, 1, n, 1, n);
  free\_dmatrix(atemp, 1, n, 1, n);
  free\_dmatrix(btemp, 1, n, 1, n);
This code is used in sections 37 and 56.
49. \langle Free bottom boundary \langle 49\rangle \equiv
  free\_dvector(R23, 1, n);
  free\_dvector(R32, 1, n);
  free\_dvector(T23, 1, n);
  free\_dvector(T32, 1, n);
This code is used in sections 37 and 56.
```

```
50. \langle Free R and T 50\rangle \equiv free\_dmatrix(R,1,n,1,n); free\_dmatrix(T,1,n,1,n); This code is used in sections 37 and 56.
```

#### 51. Simple interfaces for Perl, Python, or Mathematica.

 $ez\_RT$  is a top level routine for accessing the adding-doubling algorithm. This routine was originally created so that I could make a Perl .xs module. Since I did not know how to mess around with passing structures, I changed the interface to avoid using structures.

```
\langle \text{ Prototype for } ez\_RT \text{ 51} \rangle \equiv
```

This code is used in sections 32, 33, and 52.

```
52. \langle Definition for ez_RT 52\rangle \equiv \langle Prototype for ez_RT 51\rangle {

\mathbf{struct\ AD\_slab\_type\ } slab;

slab.n\_slab = nslab;

slab.n\_top\_slide = ntopslide;

slab.n\_bottom\_slide = nbottomslide;

slab.b\_top\_slide = 0;

slab.b\_bottom\_slide = 0;

slab.b\_bottom\_slide = 0;

slab.b = b;

slab.b = b;

slab.g = g;

slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};

slab.cos\_angle = 1.0;

\texttt{RT}(n, \&slab, \texttt{UR1}, \texttt{UT1}, \texttt{URU}, \texttt{UTU});
}
```

#### 53. Unscattered reflection and transmission.

ez\_RT\_unscattered is a top level routine for accessing the adding-doubling algorithm. This routine was created so that I could make a Perl module. Since I did not know how to mess around with passing structures, I changed the interface to avoid using structures.

```
\langle \text{ Prototype for } ez\_RT\_unscattered 53 \rangle \equiv
```

 $\begin{array}{l} \textbf{void} \ \ ez\_RT\_unscattered(\textbf{int} \ n, \textbf{double} \ nslab, \textbf{double} \ ntopslide, \textbf{double} \ nbottomslide, \textbf{double} \ a, \textbf{double} \ b, \textbf{double} \ \ *\texttt{UR1}, \textbf{double} \ \ *\texttt{UT1}, \textbf{double} \ \ *\texttt{URU}, \textbf{double} \ \ *\texttt{UTU}) \end{array}$ 

This code is used in sections 32, 33, and 54.

This code is used in section 31.

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```
54.
      \langle \text{ Definition for } ez\_RT\_unscattered 54 \rangle \equiv
  \langle Prototype for ez\_RT\_unscattered 53 \rangle
     struct AD_slab_type slab;
     slab.n\_slab = nslab;
     slab.n\_top\_slide = ntopslide;
     slab.n\_bottom\_slide = nbottomslide;
     slab.b\_top\_slide = 0;
     slab.b\_bottom\_slide = 0;
     slab.a = a:
     slab.b = b;
     slab.g = g;
     slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};
     slab.cos\_angle = 1.0;
     Sp\_RT(n, slab, UR1, UT1, URU, UTU);
```

This code is used in section 31.

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#### Including absorbing slides.

The idea is to create a function that includes absorption in the top and bottom slides. This is done by creating two extra layers, finding the full reflection and transmission matrices for these layers and adding them to the slab. Of course this only works when all the indices of refraction are the same. Yikes!

This routine returns UR1 and UT1 for light incident from the top of the slab. The values for light incident from the bottom will be different when the slides on the top and bottom are different. Caveat emptor!

```
\langle \text{ Prototype for } RTabs | 55 \rangle \equiv
```

void RTabs (int n, struct AD\_slab\_type \*slab, double \*UR1, double \*UT1, double \*URU, double \*UTU)

This code is used in sections 32 and 56.

```
\langle \text{ Definition for } RTabs | 56 \rangle \equiv
  \langle \text{ Prototype for } RTabs | 55 \rangle
     \langle \text{ Declare variables for } RTabs 57 \rangle
     double **Rtop, **Ttop, **Rbottom, **Tbottom;
     struct AD_slab_type slab1;
     double btop, bbottom;
     \langle Allocate and calculate R and T for homogeneous slab 40\rangle
     (Allocate and calculate top absorbing slide 58)
      (Allocate and calculate bottom absorbing slide 59)
      Allocate misc matrices 46
      Allocate and calculate top non-absorbing boundary 60
      Allocate and calculate bottom non-absorbing boundary 61
      Add all the stuff together 62
      Free misc matrices 48
      Free bottom boundary 49
     (Free top boundary 44)
      \langle \text{Free R and T 50} \rangle
     \langle Free matrices for the top and bottom absorbing slides 63\rangle
This code is used in section 31.
```

```
57.
      \langle \text{ Declare variables for } RTabs | 57 \rangle \equiv
  double **R, **T;
  double *R01, *R10, *T01, *T10;
  double *R23, *R32, *T23, *T32;
  double **R02, **R20, **T02, **T20;
  double **R03, **R30, **T03, **T30;
  double **atemp, **btemp;
  struct AD_method_type method;
This code is used in section 56.
58. \langle Allocate and calculate top absorbing slide 58 \rangle \equiv
  slab1.b = slab \rightarrow b\_top\_slide;
  slab1.cos\_angle = slab\neg cos\_angle;
  slab1.a = 0;
  slab1.g = 0;
  slab1.phase\_function = \texttt{HENYEY\_GREENSTEIN};
  slab1.n\_slab = slab \rightarrow n\_slab;
  slab1.n\_top\_slide = 1.0;
  slab1.n\_bottom\_slide = 1.0;
  slab1.b_{-}top_{-}slide = 0.0;
  slab1.b_bottom_slide = 0.0;
  Rtop = dmatrix(1, n, 1, n);
  Ttop = dmatrix(1, n, 1, n);
  RT\_Matrices(n, \&slab1, \&method, Rtop, Ttop);
This code is used in section 56.
59. \langle Allocate and calculate bottom absorbing slide 59\rangle \equiv
  slab1.b = slab \rightarrow b\_bottom\_slide;
  slab1.cos\_angle = slab \neg cos\_angle;
  Rbottom = dmatrix(1, n, 1, n);
  Tbottom = dmatrix(1, n, 1, n);
  RT\_Matrices(n, \&slab1, \&method, Rbottom, Tbottom);
This code is used in section 56.
60.
\langle Allocate and calculate top non-absorbing boundary 60 \rangle \equiv
  btop = slab \rightarrow b\_top\_slide;
  slab \rightarrow b_- top_- slide = 0;
  (Allocate and generate top boundary 42)
  slab \rightarrow b\_top\_slide = btop;
This code is used in section 56.
61.
\langle Allocate and calculate bottom non-absorbing boundary 61\rangle \equiv
  bbottom = slab \rightarrow b\_bottom\_slide;
  slab \rightarrow b_-bottom\_slide = 0;
  (Allocate and generate bottom boundary 45)
  slab \rightarrow b\_bottom\_slide = bbottom;
This code is used in section 56.
```

```
18 62.
```

```
\langle \text{ Add all the stuff together } 62 \rangle \equiv
  Add(n, Rtop, Rtop, Ttop, Ttop, R, R, T, T, RO2, R20, T02, T20);
  Add(n, RO2, R20, TO2, T20, Rbottom, Rbottom, Tbottom, Tbottom, RO3, R30, TO3, T30);
  Add_{-}Top(n, R01, R10, T01, T10, R03, R30, T03, T30, R02, R20, T02, T20, atemp, btemp);
  Add\_Bottom(n, RO2, R20, TO2, T20, R23, R32, T23, T32, RO3, R30, T03, T30, atemp, btemp);
   URU_{-}and_{-}UR1 (n, slab \rightarrow n_{-}slab, RO3, URU, UR1);
  Transpose\_Matrix(n, T03);
   URU_{-}and_{-}UR1 (n, slab \rightarrow n_{-}slab, TO3, UTU, UT1);
This code is used in section 56.
63.
\langle Free matrices for the top and bottom absorbing slides 63\rangle \equiv
  free\_dmatrix(Rtop, 1, n, 1, n);
  free\_dmatrix(Ttop, 1, n, 1, n);
  free\_dmatrix(Rbottom, 1, n, 1, n);
  free\_dmatrix(Tbottom, 1, n, 1, n);
This code is used in section 56.
```

#### 64. Flux and Fluence.

This code is used in section 31.

Calculates the flux and fluence at various depths between the optical depths zmin and zmax for a slab. The number of values is intervals + 1 times...i.e. it calculates at zmin, zmin + (zmax - zmin)/intervals, ..., zmax

The fluence and fluxes at 0 and *slab.b* are calculated just inside the boundary, i.e. beneath any existing glass slide or just below a mismatched boundary.

This routine could be improved dramatically. I just have not had the need so far.

This has not been adequately tested.

```
#define MAX_FLUENCE_INTERVALS 200
     \langle \text{ Prototype for } Flux\_Fluence | 65 \rangle \equiv
  void Flux_Fluence(int n, struct AD_slab_type *slab, double zmin, double zmax, int
       intervals, double *UF1\_array, double *UFU\_array, double *flux\_up, double *flux\_down)
This code is used in sections 32 and 66.
66. \langle Definition for Flux_Fluence 66 \rangle \equiv
  \langle Prototype for Flux_Fluence 65 \rangle
     ⟨ Declare variables for Flux_Fluence 67⟩
     if (intervals > MAX_FLUENCE_INTERVALS)
       AD_{error}("too_{lmany_{ll}}intervals_{ll}requested._{ll_{ll}}increase_{ll}the_{ll}const_{ll}max_{ll}fluence_{ln}tervals_{ll});
     (Find the 02 matrix for the slab above all layers 68)
     (Find the 46 matrix for the slab below all layers 69)
     (Allocate intermediate matrices 70)
     for (i = 0; i \leq intervals; i++) {
       (Find radiance at each depth 71)
       (Calculate Fluence and Flux 72)
     (Free all those intermediate matrices 73)
```

```
67.
      \langle \text{ Declare variables for } Flux\_Fluence | 67 \rangle \equiv
  double *R01, *R10, *T01, *T10;
  double *R56, *R65, *T56, *T65;
  double **R12, **T12;
  double **R23, **T23;
  double **R34, **T34;
  double **R45, **T45;
  double **R02, **R20, **T02, **T20;
  double **R46, **R64, **T46, **T64;
  double **R03, **R30, **T03, **T30;
  double **R36, **R63, **T36, **T63;
  double **Lup, **Ldown;
  double **a, **b;
  double flx_{-}down, flx_{-}up, UFU, UF1;
  double slab_thickness;
  struct AD_method_type method;
  int i, j;
This code is used in section 66.
68.
\langle Find the 02 matrix for the slab above all layers 68\rangle \equiv
  slab\_thickness = slab \neg b;
                               /* save it for later */
  slab \rightarrow b = zmin;
  R12 = dmatrix(1, n, 1, n);
  T12 = dmatrix(1, n, 1, n);
  RT\_Matrices(n, slab, \& method, R12, T12);
  RO1 = dvector(1, n);
  R10 = dvector(1, n);
  T01 = dvector(1, n);
  T10 = dvector(1, n);
  Init_Boundary (*slab, method.quad_pts, RO1, R10, T01, T10, TOP_BOUNDARY);
  R20 = dmatrix(1, n, 1, n);
  T20 = dmatrix(1, n, 1, n);
  R02 = dmatrix(1, n, 1, n);
  T02 = dmatrix(1, n, 1, n);
  a = dmatrix(1, n, 1, n);
  b = dmatrix(1, n, 1, n);
  Add_{-}Top(n, R01, R10, T01, T10, R12, R12, T12, T12, R02, R20, T02, T20, a, b);
  free\_dmatrix(R12, 1, n, 1, n);
  free\_dmatrix(T12, 1, n, 1, n);
  free\_dvector(RO1, 1, n);
  free\_dvector(R10, 1, n);
  free\_dvector(TO1, 1, n);
  free\_dvector(T10, 1, n);
This code is used in section 66.
```

```
69. \langle Find the 46 matrix for the slab below all layers 69\rangle \equiv
  slab \rightarrow b = slab\_thickness - zmax;
  R45 = dmatrix(1, n, 1, n);
  T45 = dmatrix(1, n, 1, n);
  RT-Matrices (n, slab, \& method, R45, T45);
  R56 = dvector(1, n);
  R65 = dvector(1, n);
  T56 = dvector(1, n);
  T65 = dvector(1, n);
  Init_Boundary(*slab, method.quad_pts, R56, R65, T56, T65, BOTTOM_BOUNDARY);
  R46 = dmatrix(1, n, 1, n);
  T46 = dmatrix(1, n, 1, n);
  R64 = dmatrix(1, n, 1, n);
  T64 = dmatrix(1, n, 1, n);
  Add_Bottom(n, R45, R45, T45, T45, R56, R65, T56, T65, R46, R64, T46, T64, a, b);
  free\_dmatrix(R45, 1, n, 1, n);
  free\_dmatrix(T45, 1, n, 1, n);
  free\_dvector(R56, 1, n);
  free\_dvector(R65, 1, n);
  free\_dvector(T56, 1, n);
  free\_dvector(T65, 1, n);
  free\_dmatrix(a, 1, n, 1, n);
  free\_dmatrix(b, 1, n, 1, n);
This code is used in section 66.
70. \langle Allocate intermediate matrices \langle 70 \rangle \equiv
  R23 = dmatrix(1, n, 1, n);
  T23 = dmatrix(1, n, 1, n);
  R03 = dmatrix(1, n, 1, n);
  T03 = dmatrix(1, n, 1, n);
  R30 = dmatrix(1, n, 1, n);
  T30 = dmatrix(1, n, 1, n);
  R34 = dmatrix(1, n, 1, n);
  T34 = dmatrix(1, n, 1, n);
  R63 = dmatrix(1, n, 1, n);
  T63 = dmatrix(1, n, 1, n);
  R36 = dmatrix(1, n, 1, n);
  T36 = dmatrix(1, n, 1, n);
  Lup = dmatrix(1, n, 1, n);
  Ldown = dmatrix(1, n, 1, n);
This code is used in section 66.
71. \langle Find radiance at each depth 71 \rangle \equiv
  slab \rightarrow b = (zmax - zmin)/intervals * i;
  RT\_Matrices(n, slab, \& method, R23, T23);
  Add(n, R02, R20, T02, T20, R23, R23, T23, T23, R03, R30, T03, T30);
  slab \rightarrow b = (zmax - zmin) - slab \rightarrow b;
  RT-Matrices (n, slab, \& method, R34, T34);
  Add(n, R34, R34, T34, T34, R46, R64, T46, T64, R36, R63, T36, T63);
  Between(n, R03, R30, T03, T30, R36, R63, T36, T63, Lup, Ldown);
This code is used in section 66.
```

This code is used in section 66.

 $free\_dmatrix(Lup, 1, n, 1, n);$  $free\_dmatrix(Ldown, 1, n, 1, n);$  **74. AD Layers.** This file provides routines to obtain reflection and transmission values for normal illumination of several multiple scattering and absorbing layers.

```
\langle ad\_layers.c 74 \rangle \equiv
#include <math.h>
#include <float.h>
#include "nr_util.h"
#include "ad_globl.h"
#include "ad_bound.h"
#include "ad_doubl.h"
#include "ad_prime.h"
#include "ad_matrx.h"
#include "ad_prime.h"
  \langle \text{ Definition for } RT\_Layers\_All | 77 \rangle
  \langle \text{ Definition for } RT\_Layers 88 \rangle
75. \langle ad_{layers.h} | 75 \rangle \equiv
  (Preprocessor definitions)
   \langle \text{ Prototype for } RT\_Layers 87 \rangle;
  \langle Prototype for RT\_Layers\_All 76 \rangle;
```

 $\langle \text{ Prototype for } RT\_Layers\_All | 76 \rangle \equiv$ 

**76. RT Layers.** Sometimes you just need to know the total reflection and transmission from a target consisting of multiple layers. This is the routine for you. It adds a bunch of scattering and absorbing layers together which have the same index of refraction together. The top and bottom are possibly bounded by glass slides. This is not particularly fast, but it should get the job done.

nlayers specifies the number of different layers (not including possible glass slides above and below the composite sample. The optical properties are passed in three zero-based arrays of doubles. For example a[1] is the albedo of the second layer.

```
void RT_Layers_All(int n, double nslab, double ntopslide, double nbottomslide, int nlayers, double
       a[], double b[], double g[], double *dUR1, double *dUT1, double *dURU, double
       *dUTU, double *uUR1, double *uUT1, double *uURU, double *uUTU)
This code is used in sections 75 and 77.
    \langle \text{ Definition for } RT\_Layers\_All | 77 \rangle \equiv
  \langle \text{ Prototype for } RT\_Layers\_All | 76 \rangle
     \langle Declare variables for RT\_Layers 79\rangle
     ⟨Validate layer properties 78⟩
     (Allocate slab memory 80)
     (Initialize slab structure 82)
     (Initialize composite layer 83)
     (Allocate and generate top and bottom boundaries 81)
     (Add all composite layers together 84)
     (Add top and bottom boundaries 85)
     Free memory for RT-Layers 86
This code is used in section 74.
```

**78.** 

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```
\langle Validate layer properties 78 \rangle \equiv
  if (nlayers < 1) return;
  if (nslab < 0) return;
  if (ntopslide < 0) return;
  if (nbottomslide < 0) return;
  for (i = 0; i < nlayers; i++) {
    if (a[i] < 0 \lor a[i] > 1) return;
    if (b[i] < 0) return;
    if (g[i] < -1 \lor g[i] > 1) return;
This code is used in section 77.
79. \langle Declare variables for RT_Layers 79\rangle \equiv
  struct AD_slab_type slab;
  struct AD_method_type method;
  double *R01, *R10, *T01, *T10;
  double *R34, *R43, *T34, *T43;
  double **R12, **R21, **T12, **T21;
  double **R23, **R32, **T23, **T32;
  double **R13, **R31, **T13, **T31;
  double **atemp, **btemp;
  int i;
  *dUR1 = -1;
  *dUT1 = -1;
  *dURU = -1;
  *dUTU = -1;
  *uUR1 = -1;
  *uUT1 = -1;
  *uURU = -1:
  *uUTU = -1;
This code is used in section 77.
80. \langle Allocate slab memory 80 \rangle \equiv
  R12 = dmatrix(1, n, 1, n);
  R21 = dmatrix(1, n, 1, n);
  T12 = dmatrix(1, n, 1, n);
  T21 = dmatrix(1, n, 1, n);
  R23 = dmatrix(1, n, 1, n);
  R32 = dmatrix(1, n, 1, n);
  T23 = dmatrix(1, n, 1, n);
  T32 = dmatrix(1, n, 1, n);
  R13 = dmatrix(1, n, 1, n);
  R31 = dmatrix(1, n, 1, n);
  T13 = dmatrix(1, n, 1, n);
  T31 = dmatrix(1, n, 1, n);
  atemp = dmatrix(1, n, 1, n);
  btemp = dmatrix(1, n, 1, n);
This code is used in section 77.
```

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Create the matrices needed for the top and bottom. This needs to be done after a call to  $RT\_matrices()$ so that quadrature angles are chosen.

```
\langle Allocate and generate top and bottom boundaries 81 \rangle \equiv
  RO1 = dvector(1, n);
  R10 = dvector(1, n);
  T01 = dvector(1, n);
  T10 = dvector(1, n);
  Init\_Boundary(slab, n, RO1, R10, T01, T10, TOP\_BOUNDARY);
  R34 = dvector(1, n);
  R43 = dvector(1, n);
  T34 = dvector(1, n);
  T43 = dvector(1, n);
  Init\_Boundary(slab, n, R34, R43, T34, T43, BOTTOM\_BOUNDARY);
This code is used in section 77.
```

We set this to be a clear layer so that the composite layer will be created properly. The index of refraction of the slab is important so that the quadrature angles will be chosen correctly.

```
\langle \text{Initialize slab structure } 82 \rangle \equiv
  slab.n_{-}slab = nslab;
  slab.n\_top\_slide = ntopslide;
  slab.n\_bottom\_slide = nbottomslide;
  slab.b\_top\_slide = 0;
  slab.b\_bottom\_slide = 0;
  slab.a = 0.0;
  slab.b = 0.0;
  slab.q = 0.0;
  slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};
  slab.cos\_angle = 1.0;
This code is used in section 77.
```

83. The composite layer initially has 0% reflection and 100% transmission. We fob the details on how this layer is created to the RT\_Matrices which goes to the trouble to initialize method and call Zero\_Layer for us. Finally, since this optical problem is not reversible (illumination from below gives a different answer), we need to initialize the upward matrices as well. This simplifies the code when adding successive layers.

```
\langle \text{Initialize composite layer 83} \rangle \equiv
  RT\_Matrices(n, \&slab, \&method, R23, T23);
  Copy\_Matrix(n, R23, R32);
  Copy\_Matrix(n, T23, T32);
This code is used in section 77.
```

84. Now add the layers together. Since the composite layer has been initialized to be a clear layer, we can just add layers to it. We start from the bottom. Find the transport matrices for this layer. Add this layer to the top of the composite layer. This is repeated for each of the layers.

85. The only confusing part about this piece of code is that the layer numbering gets all messed up. The composite layer is in the 23 matrices. This gets added to the top 01 boundary and should be labeled the 03 matrix. Instead I use the already allocated 13 matrices. This layer is then added to the bottom 34 matrices and should result in 04 matrices, but once again I use the 23 matrices. Finally, the total reflectances and transmittances are calculated, so that all the remains is to free the allocated memory! Not so hard after all.

```
 \langle \mbox{Add top and bottom boundaries } 85 \rangle \equiv \\ \mbox{$Add\_Top(n, R01, R10, T01, T10, R23, R32, T23, T32, R13, R31, T13, T31, atemp, btemp)$; } \\ \mbox{$Add\_Bottom(n, R13, R31, T13, T31, R34, R43, T34, T43, R23, R32, T23, T32, atemp, btemp)$; } \\ \mbox{$URU\_and\_UR1(n, slab.n\_slab, R23, $dURU, dUR1)$; } \\ \mbox{$URU\_and\_UR1(n, slab.n\_slab, R32, $uURU, uUR1)$; } \\ \mbox{$Transpose\_Matrix}(n, T23)$; } \\ \mbox{$Transpose\_Matrix}(n, T32)$; } \\ \mbox{$URU\_and\_UR1(n, slab.n\_slab, T23, $dUTU, dUT1)$; } \\ \mbox{$URU\_and\_UR1(n, slab.n\_slab, T32, $uUTU, uUT1)$; } \\ \mbox{This code is used in section } 77.
```

This code is used in section 74.

```
86.
               \langle \text{ Free memory for } RT\_Layers 86 \rangle \equiv
      free\_dvector(RO1, 1, n);
     free\_dvector(R10, 1, n);
      free\_dvector(TO1, 1, n);
      free\_dvector(T10, 1, n);
      free\_dmatrix(R12, 1, n, 1, n);
     free\_dmatrix(R21, 1, n, 1, n);
      free\_dmatrix(T12, 1, n, 1, n);
     free\_dmatrix(T21, 1, n, 1, n);
     free\_dmatrix(R23, 1, n, 1, n);
      free\_dmatrix(R32, 1, n, 1, n);
      free\_dmatrix(T23, 1, n, 1, n);
      free\_dmatrix(T32, 1, n, 1, n);
      free\_dmatrix(R13, 1, n, 1, n);
      free\_dmatrix(R31, 1, n, 1, n);
      free\_dmatrix(T13, 1, n, 1, n);
      free\_dmatrix(T31, 1, n, 1, n);
      free\_dmatrix(atemp, 1, n, 1, n);
      free\_dmatrix(btemp, 1, n, 1, n);
      free\_dvector(R34, 1, n);
      free\_dvector(R43, 1, n);
     free\_dvector(T34, 1, n);
      free\_dvector(T43, 1, n);
This code is used in section 77.
87. This just returns the reflection and transmission for light travelling downwards. This is most often
what is desired.
\langle \text{ Prototype for } RT\_Layers 87 \rangle \equiv
      void RT\_Layers (int n, double nslab, double ntopslide, double nbottomslide, int nlayers, double
                  a[], double b[], double g[], double *UR1, double *UT1, double *URU, double *UTU)
This code is used in sections 75 and 88.
            \langle \text{ Definition for } RT\_Layers 88 \rangle \equiv
      \langle \text{ Prototype for } RT\_Layers 87 \rangle
            double uUR1, uUT1, uURU, uUTU;
            RT\_Layers\_All(n, nslab, ntopslide, nbottomslide, nlayers, a, b, g, 39uR1, UT1, URU, UTU, &uUR1, &uUT1, &u
                        \&uURU, \&uUTU);
```

AD CONE

ξ89

**AD Cone.** This file provides routines to obtain reflection and transmission values returning within a cone assuming normal illumination.

```
\langle ad\_cone.c 89 \rangle \equiv
#include <math.h>
#include <float.h>
#include <stdio.h>
#include "nr_util.h"
#include "ad_globl.h"
#include "ad_matrx.h"
#include "ad_bound.h"
#include "ad_doubl.h"
#include "ad_start.h"
   \langle \text{ Definition for } RT\_Cone 93 \rangle
    Definition for ez_RT_Cone = 103
   \langle \text{ Definition for } ez\_RT\_Oblique \ 105 \rangle
90. \langle ad\_cone.h 90 \rangle \equiv
   (Preprocessor definitions)
   \langle \text{ Prototype for } RT\_Cone 92 \rangle;
   \langle \text{ Prototype for } ez\_RT\_Cone \ 102 \rangle;
   \langle \text{ Prototype for } ez\_RT\_Oblique \ 104 \rangle;
91. \langle ad\_cone\_ez.h \quad 91 \rangle \equiv
   ⟨ Preprocessor definitions ⟩
   \langle \text{ Prototype for } ez\_RT\_Cone \ 102 \rangle;
   \langle \text{ Prototype for } ez\_RT\_Oblique \ 104 \rangle;
```

Sometimes you just need to know the total reflection and transmission from a target within a specified cone of angles. For example, you might want to test a Monte Carlo implementation of fiber illumination. The way that this works is to divide the integration over angles into two or three pieces. A separate quadrature is done over each integration range. For example if  $\nu_{\rm cone}$  is the cosine of the cone angle and there are no index of refraction changes that need to accounted for, then

$$\int_0^1 A(\nu, \nu') B(\nu', \nu'') d\nu' = \int_0^{\nu_{\text{cone}}} A(\nu, \nu') B(\nu', \nu'') d\nu' + \int_{\nu_{\text{cone}}}^1 A(\nu, \nu') B(\nu', \nu'') d\nu'.$$

otherwise one needs to include the critical angle as a special point in the integration and the integration becomes

$$\begin{split} \int_{0}^{1} A(\nu, \nu') B(\nu', \nu'') \, d\nu' &= \int_{0}^{\nu_{\text{crit}}} A(\nu, \nu') B(\nu', \nu'') \, d\nu' \\ &+ \int_{\nu_{\text{crit}}}^{\nu_{\text{cone}}} A(\nu, \nu') B(\nu', \nu'') \, d\nu' + \int_{\nu_{\text{cone}}}^{1} A(\nu, \nu') B(\nu', \nu'') \, d\nu'. \end{split}$$

Radau quadrature is chosen for the integration range from  $\nu_{\text{cone}}$  to 1. The other two use Gaussian quadrature.

```
\langle \text{ Prototype for } RT_{-}Cone 92 \rangle \equiv
```

void RT\_Cone(int n, struct AD\_slab\_type \*slab, int use\_cone, double \*UR1, double \*UT1, double \*URU, double \*UTU)

This code is used in sections 90 and 93.

```
\langle \text{ Definition for } RT\_Cone 93 \rangle \equiv
   \langle \text{ Prototype for } RT\_Cone 92 \rangle
      \langle RT\_Cone \text{ Declare variables } 94 \rangle
      \langle RT_{-}Cone \text{ Check inputs } 95 \rangle
      \langle RT\_Cone \text{ Allocate slab memory } 97 \rangle
      \langle RT\_Cone \text{ Initialize homogeneous layer 98} \rangle
      \langle RT\_Cone \text{ Allocate and generate top and bottom boundaries 99} \rangle
      \langle RT\_Cone \text{ Add top and bottom boundaries } 100 \rangle
      \langle RT\_Cone \text{ Free memory } 101 \rangle
  }
This code is used in section 89.
94. \langle RT\_Cone \text{ Declare variables 94} \rangle \equiv
   struct AD_method_type method;
   double *R01, *R10, *T01, *T10;
   double *R23, *R32, *T23, *T32;
   double **R12, **T12;
   double **R02, **T02, **T20, **R20;
   double **R03, **T03, **T30, **R30;
   double **atemp, **btemp;
   double d;
   *UR1 = -1;
   *URU = -1;
   *UT1 = -1;
   *UTU = -1;
This code is used in section 93.
95.
\langle RT\_Cone \text{ Check inputs } 95 \rangle \equiv
  if (slab \rightarrow n\_slab < 0) return;
  if (slab \rightarrow n_- top_- slide < 0) return;
  if (slab \neg n\_bottom\_slide < 0) return;
  if (slab \neg a < 0 \lor slab \neg a > 1) return;
  if (slab \rightarrow g < -1 \lor slab \rightarrow g > 1) return;
  if (slab \rightarrow b < 0) return;
  if (slab \neg cos\_angle < 0 \lor slab \neg cos\_angle > 1) return;
See also section 96.
This code is used in section 93.
```

**96.** The number of quadrature points must be fixed before starting to allocate memory. We want the number of points to be at least twelve so that each of the three integrals will have four quadrature points.

```
\langle RT\_Cone \text{ Check inputs 95} \rangle +\equiv n = 12 * (n/12);

if (n < 12) n = 12;

method.quad\_pts = n;
```

```
97. \langle RT\_Cone \text{ Allocate slab memory } 97 \rangle \equiv
  R12 = dmatrix(1, n, 1, n);
  T12 = dmatrix(1, n, 1, n);
  R02 = dmatrix(1, n, 1, n);
  T02 = dmatrix(1, n, 1, n);
  R20 = dmatrix(1, n, 1, n);
  T20 = dmatrix(1, n, 1, n);
  R03 = dmatrix(1, n, 1, n);
  T03 = dmatrix(1, n, 1, n);
  R30 = dmatrix(1, n, 1, n);
  \mathsf{T30} = dmatrix(1, n, 1, n);
  atemp = dmatrix(1, n, 1, n);
  btemp = dmatrix(1, n, 1, n);
This code is used in section 93.
```

The homogeneous layer initially has 0% reflection and 100% transmission. We cannot fob the details on how this layer is created to RT\_Matrices because we need to (1) set the quadrature angles to a multiple of three, and (2) explicitly make a call to Choose\_Cone\_Method so that the quadrature angles will get chosen appropriately.

This code is directly lifted from the RT\_Matrices routine.

```
\langle RT\_Cone \text{ Initialize homogeneous layer } 98 \rangle \equiv
  Choose\_Cone\_Method(slab, \&method);
  if (slab \rightarrow b \leq 0) {
     Zero\_Layer(n, R12, T12);
     return;
  n = method.quad_pts;
  Init\_Layer(*slab, method, R12, T12);
  d = 1.0;
  if (slab \rightarrow b \neq HUGE\_VAL) d = method.b\_thinnest * slab \rightarrow b / method.b\_calc;
  Double\_Until(n, R12, T12, d, slab \rightarrow b);
```

This code is used in section 93.

99. Create the matrices needed for the top and bottom

```
\langle RT\_Cone \text{ Allocate and generate top and bottom boundaries 99} \rangle \equiv
  RO1 = dvector(1, n);
  R10 = dvector(1, n);
  T01 = dvector(1, n);
  T10 = dvector(1, n);
  Init\_Boundary(*slab, n, R01, R10, T01, T10, T0P\_BOUNDARY);
  R23 = dvector(1, n);
  R32 = dvector(1, n);
  T23 = dvector(1, n);
  T32 = dvector(1, n);
  Init\_Boundary(*slab, n, R23, R32, T23, T32, BOTTOM\_BOUNDARY);
This code is used in section 93.
```

100. Here the layer numbering is pretty consistent. The top slide is 01, the scattering layer is 12, and the bottom slide is 23. Light going from the top of the slide to the bottom of the scattering layer is 02 and similarly light going all the way through is 03.

The only tricky part is that the definitions of UR1 and URU have changed from their usual definitions. When  $use\_cone \equiv OBLIQUE$  then UR1 refers to the light reflected back into the specified cone for normal irradiance and URU is for light reflected back into the cone for light incident uniformly at all angles within that cone. Otherwise, assume that the incidence is oblique. UR1 then refers to the total amount of light reflected back for light incident only at the cone angle.

```
\langle RT\_Cone \text{ Add top and bottom boundaries } 100 \rangle \equiv
  Add_{-}Top(n, R01, R10, T01, T10, R12, R12, T12, T12, R02, R20, T02, T20, atemp, btemp);
  Add\_Bottom(n, RO2, R20, TO2, T20, R23, R32, T23, T32, RO3, R30, T03, T30, atemp, btemp);
  if (use\_cone \equiv CONE) {
     URU\_and\_UR1\_Cone(n, slab \rightarrow n\_slab, slab \rightarrow cos\_angle, RO3, URU, UR1);
      Transpose\_Matrix(n, T03);
     URU\_and\_UR1\_Cone(n, slab \rightarrow n\_slab, slab \rightarrow cos\_angle, T03, UTU, UT1);
  }
  else {
     if (use\_cone \neq OBLIQUE)
        fprintf(stderr, "Unknown_{\sqcup}type_{\sqcup}for_{\sqcup}use\_cone._{\sqcup \sqcup}Assuming_{\sqcup}oblique_{\sqcup}incidence.\n");
     URU\_and\_URx\_Cone(n, slab \rightarrow n\_slab, slab \rightarrow cos\_angle, RO3, URU, UR1);
     Transpose\_Matrix(n, T03);
     URU\_and\_URx\_Cone(n, slab \rightarrow n\_slab, slab \rightarrow cos\_angle, T03, UTU, UT1);
  }
This code is used in section 93.
101. \langle RT\_Cone \text{ Free memory } 101 \rangle \equiv
  free\_dvector(RO1, 1, n);
  free\_dvector(R10, 1, n);
  free\_dvector(TO1, 1, n);
  free\_dvector(T10, 1, n);
  free\_dmatrix(R12, 1, n, 1, n);
  free\_dmatrix(T12, 1, n, 1, n);
  free\_dmatrix(RO3, 1, n, 1, n);
  free\_dmatrix(R30, 1, n, 1, n);
  free\_dmatrix(T03, 1, n, 1, n);
  free\_dmatrix(T30, 1, n, 1, n);
  free\_dmatrix(RO2, 1, n, 1, n);
  free\_dmatrix(R20, 1, n, 1, n);
  free\_dmatrix(T02, 1, n, 1, n);
  free\_dmatrix(T20, 1, n, 1, n);
  free\_dmatrix(atemp, 1, n, 1, n);
  free\_dmatrix(btemp, 1, n, 1, n);
  free\_dvector(R32, 1, n);
  free\_dvector(R23, 1, n);
  free\_dvector(T32, 1, n);
  free\_dvector(T23, 1, n);
This code is used in section 93.
```

This code is used in section 89.

```
102.
                Simple wrapper that avoids data structures
\langle \text{ Prototype for } ez\_RT\_Cone \ 102 \rangle \equiv
     void ez_RT_Cone(int n, double nslab, double ntopslide, double nbottomslide, double a, double
               b, double q, double cos_cone_angle, double *UR1, double *UT1, double *URU, double *UTU)
This code is used in sections 90, 91, and 103.
103. \langle \text{ Definition for } ez\_RT\_Cone \ 103 \rangle \equiv
     \langle \text{ Prototype for } ez\_RT\_Cone \ 102 \rangle
          struct AD_slab_type slab;
          slab.n_{-}slab = nslab;
          slab.n\_top\_slide = ntopslide;
          slab.n\_bottom\_slide = nbottomslide;
          slab.b\_top\_slide = 0;
          slab.b\_bottom\_slide = 0;
          slab.a = a;
          slab.b = b;
          slab.g = g;
          slab.cos\_angle = cos\_cone\_angle;
          slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};
          RT_{-}Cone(n, \&slab, CONE, UR1, UT1, URU, UTU);
This code is used in section 89.
                This routine calculates reflection and transmission for oblique incidence. URx and UTx are the total
light reflected and transmitted for light incident at at cos_oblique_angle. URU and UTU are the same thing for
diffuse incident light.
\langle \text{ Prototype for } ez\_RT\_Oblique \ 104 \rangle \equiv
     void ez_RT_Oblique(int n, double nslab, double ntopslide, double nbottomslide, double a, double normalistic norm
                b, double q, double cos\_oblique\_angle, double *URx, double *UTx, double *URU, double *UTU)
This code is used in sections 90, 91, and 105.
105. \langle \text{ Definition for } ez\_RT\_Oblique | 105 \rangle \equiv
      \langle Prototype for ez\_RT\_Oblique 104 \rangle
          struct AD_slab_type slab;
          slab.n\_slab = nslab;
          slab.n\_top\_slide = ntopslide;
          slab.n\_bottom\_slide = nbottomslide;
          slab.b\_top\_slide = 0;
          slab.b\_bottom\_slide = 0;
          slab.a = a;
          slab.b = b;
          slab.g = g;
          slab.cos\_angle = cos\_oblique\_angle;
          slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};
          RT_{-}Cone(n, \&slab, OBLIQUE, URx, UTx, URU, UTU);
```

32

106. AD Start. This has the routines for forming the initial matrix to start off an adding-doubling calculation.

Added printing of intermediate results for Martin Hammer.

```
#include <math.h>
#include <float.h>
#include <stdio.h>
#include "ad_frsnl.h"
#include "ad_globl.h"
#include "ad_matrx.h"
#include "ad_phase.h"
#include "ad_radau.h"
#include "ad_start.h"
#include "nr_gaulg.h"
#include "nr_util.h"
  \langle Definition for Get\_Start\_Depth 110 \rangle
  (Definition for Quadrature 113)
   (Definition for Choose_Method 115)
   Definition for Choose_Cone_Method 117
   \langle \text{ Definition for } Get\_IGI\_Layer 127 \rangle
  ⟨ Definition for Get_Diamond_Layer 128⟩
  ⟨ Definition for Init_Layer 140⟩
107. \langle ad_start.h | 107 \rangle \equiv
  \langle Prototype for Get\_Start\_Depth 109 \rangle;
  ⟨ Prototype for Choose_Method 114⟩;
  ⟨ Prototype for Choose_Cone_Method 116⟩;
  \langle Prototype for Init\_Layer 139 \rangle;
  \langle Prototype for Quadrature 112 \rangle;
```

#### 108. Basic routines.

This file contains the three procedures which must be called before any doubling may take place. They should be called in the following order:

```
Choose_Method — to fill the method record Quadrature — to calculate the quad angles and weights code to initialize angle, weight, and twoaw Init\_Layer — to calculate the thin layer R and T Double\_Until — to obtain R and T for the desired thickness
```

BASIC ROUTINES

109. Get\_Start\_Depth selects the best minimum starting thickness to start the doubling process. The criterion is based on an assessment of the (1) round-off error, (2) the angular initialization error, and (3) the thickness initialization error. Wiscombe concluded that an optimal starting thickness depends on the smallest quadrature angle, and recommends that when either the infinitesimal generator or diamond initialization methods are used then the initial thickness is optimal when type 2 and 3 errors are comparable, or when

```
d \approx \mu
```

Note that round-off is important when the starting thickness is less than  $1 \cdot 10^{-4}$  for diamond initialization and less than  $1 \cdot 10^{-8}$  for infinitesimal generator initialization assuming about 14 significant digits of accuracy.

Since the final thickness is determined by repeated doubling, the starting thickness is found by dividing by 2 until the starting thickness is less than  $\mu$ . Also we make checks for a layer with zero thickness and one that infinitely thick.

```
⟨ Prototype for Get_Start_Depth 109⟩ ≡
   double Get_Start_Depth(double mu, double d)
This code is used in sections 107 and 110.

110. ⟨ Definition for Get_Start_Depth 110⟩ ≡
   ⟨ Prototype for Get_Start_Depth 109⟩
   {
      if (d ≤ 0) return 0.0;
      if (d ≡ HUGE_VAL) return (mu/2.0);
      while (d > mu) d /= 2;
      return d;
   }
This code is used in section 106.
```

#### 111. Quadrature.

112. This returns the quadrature angles using Radau quadrature over the interval 0 to 1 if there is no critical angle for total internal reflection in the slab. If there is a critical angle whose cosine is  $\mu_c$  then Radau quadrature points are chosen from 0 to  $\mu_c$  and Radau quadrature points over the interval  $\mu_c$  to 1.

```
\langle Prototype for Quadrature 112\rangle \equiv void Quadrature (int n, double n_slab, double *x, double *w) This code is used in sections 107 and 113.
```

```
\langle \text{ Definition for } Quadrature | 113 \rangle \equiv
  \langle \text{ Prototype for } \textit{Quadrature } 112 \rangle
     int i, nby2;
     double *x1, *w1;
     double mu\_c;
     if (n\_slab \equiv 1) {
        Radau(0.0, 1.0, x, w, n);
        return;
     mu\_c = Cos\_Critical\_Angle(n\_slab, 1.0);
     nby2 = n/2;
     gauleg(0.0, mu\_c, x, w, nby2);
     x1 = dvector(1, nby2);
     w1 = dvector(1, nby2);
     Radau(mu_{-}c, 1.0, x1, w1, nby2);
     for (i = 1; i \le nby2; i++) {
       x[nby2 + i] = x1[i];
        w[nby2 + i] = w1[i];
     free\_dvector(x1, 1, nby2);
     free\_dvector(w1, 1, nby2);
This code is used in section 106.
```

Choose\_Method fills the method structure with correct values for a\_calc, b\_calc, q\_calc, and b\_thinnest based on the delta-M method. Furthermore, the quadrature angles and weights are also calculated. Before calling this routines method.quad\_pts must be set to some multiple of 2. If this routine is not called then it is up to you to

- 1. to fill the method record appropriately
- 2. call Quadrature
- 3. fill global arrays angle, weight, and twoaw
- 4. determine the thickness of the thinnest layer

 $\langle Prototype for Choose\_Method 114 \rangle \equiv$ 

void Choose\_Method(struct AD\_slab\_type \*slab, struct AD\_method\_type \*method)

This code is used in sections 107 and 115.

```
115.
        \langle \text{ Definition for } Choose\_Method | 115 \rangle \equiv
   ⟨ Prototype for Choose_Method 114⟩
     double af;
     int i, n;
     if (0 < slab \neg cos\_angle \land slab \neg cos\_angle < 1) {
        Choose\_Cone\_Method(slab, method);
        return;
     n = method \neg quad\_pts;
     af = pow(slab \rightarrow g, n) * slab \rightarrow a;
     method \neg a\_calc = (slab \neg a - af)/(1 - af);
     method \neg b\_calc = (1 - af) * slab \neg b;
     method \neg g\_calc = slab \neg g;
     Quadrature(n, slab \rightarrow n\_slab, angle, weight);
     for (i = 1; i < n; i++) twoaw[i] = 2 * angle[i] * weight[i];
     method \neg b\_thinnest = Get\_Start\_Depth(angle[1], method \neg b\_calc);
This code is used in section 106.
        Choose_Cone_Method adds the ability to specify a specific quadrature angle so that accurate estimates
of the reflection and transmission might be made for when the light returning in a particular cone is of interest.
This code mimicks the usual Choose_Method above, and in fact explicitly uses it for a couple of special cases.
\langle Prototype for Choose\_Cone\_Method 116 \rangle \equiv
  void Choose_Cone_Method(struct AD_slab_type *slab, struct AD_method_type *method)
This code is used in sections 107 and 117.
117. \langle Definition for Choose\_Cone\_Method 117 \rangle \equiv
   ⟨ Prototype for Choose_Cone_Method 116⟩
     double af, *angle1, *weight1, cos_crit_angle, mu;
     int i, n, nby2, nby3;
     n = method \neg quad\_pts;
     af = pow(slab \rightarrow g, n) * slab \rightarrow a;
     method \neg a\_calc = (slab \neg a - af)/(1 - af);
     method \rightarrow b\_calc = (1 - af) * slab \rightarrow b;
     method \neg g\_calc = slab \neg g;
     (Special case when cosine is zero 120)
     (Special case when no index of refraction change 121)
```

This code is used in section 106.

118.  $\langle \text{ print angles } 118 \rangle \equiv$  This code is used in sections 120, 121, and 124.

(Gaussian quadrature from 0 to the critical angle 122)

Radau quadrature from the cone angle to 1 124

(Radau quadrature from the critical angle to the cone angle 123)

```
119. \langle \text{debug print angles } 119 \rangle \equiv \{ \\ printf("****Cone\_Angle_{\Box\Box\Box\Box\Box\Box\Box\Box\Box\Box}= \%6.2f_{\Box}degrees, \_Cosine() = \%6.4f \", \\ acos(slab \neg cos\_angle) * 180.0/3.14159, slab \neg cos\_angle); \\ double sum = 0; \\ for (i = 1; i \leq n; i++) \{ \\ sum += twoaw[i]; \\ printf("\%02d\_theta = \%6.2f\_cos(theta) = \%6.4f\_w = \%6.4f\_2aw = \%6.4f \", i, \\ acos(angle[i])/3.1415926 * 180.0, angle[i], weight[i], twoaw[i]); \\ \} \\ printf("twoaw\_sum\_= \%8.4f \", sum); \\ \}
```

**120.** When the cone angle is zero or ninety degrees then we can just use the standard method for choosing the quadrature points.

```
 \langle \, \text{Special case when cosine is zero } \, 120 \, \rangle \equiv \\ \quad \text{if } \, (slab \neg cos\_angle \equiv 0 \lor slab \neg cos\_angle \equiv 1) \, \, \{ \\ \quad Choose\_Method(slab, method); \\ \quad \langle \, \text{print angles } \, 118 \, \rangle \\ \quad \text{return;} \\ \quad \}
```

This code is used in section 117.

121. When there is no index of refraction change, there is no critical angle to worry about. Since we want the cone angle to be included as one of our angles, we use Radau quadrature. That way both the cone angle and perpendicular angles are included.

```
\langle Special case when no index of refraction change 121 \rangle \equiv
  if (slab \neg n\_slab \equiv 1 \land slab \neg n\_top\_slide \equiv 1 \land slab \neg n\_bottom\_slide \equiv 1) {
     nby2 = n/2;
     Radau(0.0, slab \rightarrow cos\_angle, angle, weight, nby2);
     angle 1 = dvector(1, nby 2);
     weight1 = dvector(1, nby2);
     Radau(slab \rightarrow cos\_angle, 1.0, angle1, weight1, nby2);
     for (i = 1; i \le nby2; i++) {
        angle[nby2 + i] = angle1[i];
        weight[nby2 + i] = weight1[i];
     free\_dvector(angle1, 1, nby2);
     free\_dvector(weight1, 1, nby2);
     for (i = 1; i \le n; i ++) twoaw[i] = 2 * angle[i] * weight[i];
     method \neg b\_thinnest = Get\_Start\_Depth(angle[1], method \neg b\_calc);
     (print angles 118)
     return;
```

This code is used in section 117.

122. Now we need to include three angles, the critical angle, the cone angle, and perpendicular. Now the important angles are the ones in the slab. So we calculate the cosine of the critical angle in the slab and cosine of the cone angle in the slab.

The critical angle will always be greater than the cone angle in the slab and therefore the cosine of the critical angle will always be less than the cosine of the cone angle. Thus we will integrate from zero to the cosine of the critical angle (using Gaussian quadrature to avoid either endpoint) then from the critical angle to the cone angle (using Radau quadrature so that the cosine angle will be included) and finally from the cone angle to 1 (again using Radau quadrature so that 1 will be included).

```
\langle Gaussian quadrature from 0 to the critical angle |122\rangle \equiv
  cos\_crit\_angle = Cos\_Critical\_Angle(slab \rightarrow n\_slab, 1.0);
  nby\beta = n/3;
  gauleg(0.0, cos\_crit\_angle, angle, weight, nby3);
This code is used in section 117.
       \langle Radau quadrature from the critical angle to the cone angle 123\rangle
  mu = sqrt(slab \neg n\_slab * slab \neg n\_slab - 1 + slab \neg cos\_angle * slab \neg cos\_angle)/slab \neg n\_slab;
  angle1 = dvector(1, nby3);
  weight1 = dvector(1, nby3);
  Radau(cos_crit_angle, mu, angle1, weight1, nby3);
  for (i = 1; i \le nby3; i++) {
     angle[nby\beta + i] = angle1[i];
     weight[nby3 + i] = weight1[i];
  }
This code is used in section 117.
124. \langle Radau quadrature from the cone angle to 1 124\rangle \equiv
  Radau(mu, 1.0, angle1, weight1, nby3);
  for (i = 1; i \le nby3; i++) {
     angle[nby3 * 2 + i] = angle1[i];
     weight[nby3 * 2 + i] = weight1[i];
  free\_dvector(angle1, 1, nby3);
  free\_dvector(weight1, 1, nby3);
  for (i = 1; i \le n; i++) twoaw [i] = 2 * angle[i] * weight[i];
  method \neg b\_thinnest = Get\_Start\_Depth(angle[1], method \neg b\_calc);
  (print angles 118)
This code is used in section 117.
```

### 125. Initialization.

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The basic idea behind diamond initialization is to rewrite the time-independent, one-dimensional, azimuthally averaged, radiative transport equation

$$\nu \frac{\partial L(\tau, \nu)}{\partial \tau} + L(\tau, \nu) = \frac{a}{2} \int_{-1}^{1} h(\nu, \nu') L(\tau, \nu') \, d\nu'$$

in a discrete form as

$$\pm \nu_i \frac{\partial L(\tau, \pm \nu_i)}{\partial \tau} + L(\tau, \pm \nu_i) = \frac{a}{2} \sum_{i=1}^{M} w_j \left[ h(\nu_i, \nu_j) L(\tau, \pm \nu_i) + h(\nu_i, -\nu_j) L(\tau, \mp \nu_i) \right]$$

When this equation is integrated over a thin layer from  $\tau_0^*$  to  $\tau_1^*$  then get

$$\pm \nu_i [L(\tau_1^*, \pm \nu_i) - L(\tau_0^*, \pm \nu_i)] + dL_{1/2}(\pm \nu_i)$$

$$= \frac{a}{2} \sum_{i=1}^M w_j d \left[ h(\nu_i, \nu_j) L_{1/2}(\pm \nu_i) + h(\nu_i, -\nu_j) L_{1/2}(\mp \nu_i) \right]$$

where  $d = \tau_1^* - \tau_0^*$ . The integrated radiance  $L_{1/2}(\nu)$  is

$$L_{1/2}(\nu) \equiv \frac{1}{\Delta \tau^*} \int_{\tau_0^*}^{\tau_1^*} L(\tau, \nu) d\tau$$

Exactly how this integral is approximated determines the type of initialization. Wiscombe evaluated a number of initialization methods and found two that were useful. These are the infinitesimal generator and the diamond methods. The infinitesimal generator initialization makes the approximation

$$L_{1/2}(-\nu) = L(\tau_1^*, -\nu)$$
  $L_{1/2}(\nu) = L(\tau_0^*, \nu)$ 

and the diamond initialization assumes

$$L_{1/2}(\nu) = \frac{1}{2} [L(\tau_0^*, \nu) + L(\tau_1^*, \nu)]$$

# 126. Infinitesmal Generator Initialization.

127.  $Get\_IGI\_Layer$  generates the starting matrix with the inifinitesimal generator method. The accuracy is O(d) and assumes that the average irradiance upwards is equal to that travelling downwards at the top and the average radiance upwards equals that moving upwards from the bottom.

$$L_{1/2}(-\nu) = L(\tau_1^*, -\nu)$$
  $L_{1/2}(\nu) = L(\tau_0^*, \nu)$ 

After manipulation, Wiscombe obtains these basic formulas for the infinitesimal generator method,

$$R = \hat{R}d \qquad \qquad T = I - \hat{T}d$$

where d is the optical thickness of the layer and I is the identity matrix. The values for  $\hat{R}$  and  $\hat{T}$  are given by

$$\hat{R} = \frac{a}{2} M^{-1} h^{+-} W \qquad \qquad \hat{T} = M^{-1} (I - \frac{a}{2} h^{++} W)$$

where M and W are diagonal matrices composed of the quadrature angles and their corresponding weights. Therefore

$$\hat{R}_{ij} = \frac{a}{2\mu_i} h_{ij}^{+-} w_j \qquad \hat{T}_{ij} = \frac{\delta_{ij}}{\mu_i} - \frac{a}{2\mu_i} h_{ij}^{++} w_j$$

and

$$R_{ij} = \frac{ad}{2\mu_i} h_{ij}^{+-} w_j$$
  $T_{ij} = \frac{ad}{2\mu_i} h_{ij}^{++} + \delta_{ij} \left[ 1 - \frac{d}{\mu_i} \right]$ 

This would be fine, but the way that the reflection and transmission matrices are set-up requires that each we multiply each matrix on the right by  $1/(2\mu_i w_i)$ . Putting things together we get

$$R_{ij} = \frac{ad}{4\mu_i \mu_j} h_{ij}^{+-}$$

and

$$T_{ij} = \frac{ad}{4\mu_i \mu_j} h_{ij}^{++} + \frac{\delta_{ij}}{2\mu_i w_i} \left[ 1 - \frac{d}{\mu_i} \right]$$

```
 \langle \text{ Definition for } \textit{Get\_IGI\_Layer} \ | \ \textbf{127} \rangle \equiv \\ \textbf{static void } \textit{Get\_IGI\_Layer} \ (\textbf{struct AD\_method\_type} \ \textit{method}, \textbf{double} \ **h, \textbf{double} \ **R, \textbf{double} \ **T) \ \\ \{ \textbf{int } i, j, n; \\ \textbf{double } a, c, d, \textit{temp}; \\ a = \textit{method.a\_calc}; \\ d = \textit{method.b\_thinnest}; \\ n = \textit{method.quad\_pts}; \\ \textbf{for } (j = 1; j \leq n; j + ) \ \\ \textit{temp} = a * d/4 / angle [j]; \\ \textbf{for } (i = 1; i \leq n; i + +) \ \\ c = \textit{temp} / angle [i]; \\ R[i][j] = c * h[i][-j]; \\ T[i][j] = c * h[i][j]; \\ \} \\ T[j][j] + = (1 - d / angle [j]) / \textit{twoaw}[j]; \\ \}
```

This code is used in section 106.

### 128. Diamond Initialization.

It should be noted up front that the implementation contained herein is somewhat cryptic. Much of the complexity comes from using the tricks in the appendix A of Wiscombe's paper ("On initialization, error and flux conservation in the doubling method.") After spending a whole day tracking down a small error in the calculation of the reflection matrix, I will spend a few moments trying to improve the documentation for this whole section. It should be apparent that this is no substitute for reading the paper.

The advantage of the diamond initialization method is that its accuracy is of the order of the square of the optical thickness  $O(d^2)$ . This means that much thicker starting layers and retain good starting accuracy. This reduces the number of doubling steps that are required. However, if the layer thickness is too thin then the accuracy gets much worse because errors in the numerical precision start to affect the results.

Get\_Diamond\_Layer generates the starting matrix with the diamond method. This implies that the integral can be replaced by a simple average of the radiances at the top and bottom of the layer,

$$L_{1/2}(\nu) = \frac{1}{2}[L(\tau_0^*, \nu) + L(\tau_1^*, \nu)]$$

```
 \langle \text{ Definition for } \textit{Get\_Diamond\_Layer} \ 128 \rangle \equiv \\ \textbf{static void } \textit{Get\_Diamond\_Layer}(\textbf{struct AD\_method\_type} \ \textit{method}, \textbf{double} \ **h, \textbf{double} \ **R, \textbf{double} \ **T) \\ \\ \{ \\ \langle \text{Local variables and initialization 136} \rangle \\ \langle \text{Find } r \text{ and } t \ 129 \rangle \\ \langle \text{Find } C = r/(1+t) \ 130 \rangle \\ \langle \text{Find } G = 0.5(1+t-Cr) \ 131 \rangle \\ \langle \text{print } r, t, \text{ and } g \text{ for Martin Hammer 132} \rangle \\ \langle \text{Calculate } R \text{ and } T \ 133 \rangle \\ \langle \text{Free up memory 137} \rangle \\ \}
```

This code is used in section 106.

129. This diamond initialization method uses the same  $\hat{R}$  and  $\hat{T}$  as was used for infinitesimal generator method. However, we want to form the r and t

$$r = \frac{d}{2}\hat{R} \qquad \qquad t = \frac{d}{2}\hat{T}$$

Recall that

$$\hat{R}_{ij} = \frac{a}{2\mu_i} h_{ij}^{+-} w_j \qquad \qquad \hat{T}_{ij} = \frac{\delta_{ij}}{\mu_i} - \frac{a}{2\mu_i} h_{ij}^{++} w_j$$

therefore

$$r_{ij} = \frac{adw_j}{4\mu_i} h_{ij}^{+-}$$
  $t_{ij} = \delta_{ij} \frac{d}{2\mu_i} - \frac{adw_j}{4\mu_i} h_{ij}^{++}$ 

If you happen to be wondering why right multiplication by  $1/(2\mu_j w_j)$  is not needed, you would be a thinking sort of person. Division by  $1/(2\mu_j w_j)$  is not needed until the final values for R and T are formed.

```
 \begin{split} \langle \, \text{Find } r \, \text{ and } t \, \, 129 \, \rangle &\equiv \\ \mathbf{for} \, \left( j = 1; \, \, j \leq n; \, \, j + + \right) \, \left\{ \\ temp &= a * d * weight[j]/4; \\ \mathbf{for} \, \left( i = 1; \, \, i \leq n; \, \, i + + \right) \, \left\{ \\ c &= temp/angle[i]; \\ R[i][j] &= c * h[i][-j]; \\ T[i][j] &= -c * h[i][j]; \\ \right\} \\ T[j][j] &+ = d/(2 * angle[j]); \\ \left\} \end{split}
```

This code is used in section 128.

130. Wiscombe points out (in Appendix A), that the matrix inversions can be avoided by noting that if we want C from the combination

$$C = r(I+t)^{-1}$$

then one needs only solve the system

$$(I+t)^T C^T = r^T$$

for C. This is done in the routine  $Left\_Inverse\_Multiply$ . We just need to create A = I + T and fire it off to  $Left\_Inverse\_Multiply$ . Actually, Wiscome goes on to suggest a faster method that takes advantage of the column oriented structure of storage on the computer. Since we are using the Numerical Recipes scheme, I don't think that his refinement will prove faster because it involves more multiplications and divisions. (Actually, that improvement was exactly what the bug in the program was. I included the required multiplications and voilá! It worked.)

This code is used in section 128.

### **131.** Here the matrix

$$G = \frac{1}{2}(I + t - Cr)$$

is formed.  $\langle \operatorname{Find} G = 0.5(1+t-Cr) \ \ 131 \rangle \equiv \\ \operatorname{Matrix\_Multiply}(n,C,R,G); \\ \mathbf{for} \ (i=1; \ i \leq n; \ i++) \ \{ \\ \mathbf{for} \ (j=1; \ j \leq n; \ j++) \ \ G[i][j] = (T[i][j]-G[i][j])/2; \\ G[i][i] \ += 0.5; \\ \}$ 

This code is used in section 128.

132. To print intermediate results for Chapter 4 of AJ's book, then it is necessary to print things from within  $Get\_Diamond\_Layer$ . Martin Hammer requested that I provide these results. Since this is the only time that they are of interest, they are only printed when both the compiler define MARTIN\_HAMMER is defined, and when the variable  $Martin\_Hammer \neq 0$ .

```
\langle \text{ print } r, t, \text{ and } g \text{ for Martin Hammer } 132 \rangle \equiv
#ifdef MARTIN_HAMMER
  {
     double **Ginv, **G2;
     if (Martin\_Hammer \neq 0) {
        printf("A_{\sqcup}from_{\sqcup}equation_{\sqcup}5.55\n");
        wrmatrix(n,T);
        printf("B_{\square}from_{\square}equation_{\square}5.55\n");
        wrmatrix(n, R);
        Ginv = dmatrix(1, n, 1, n);
        \texttt{G2} = dmatrix(1, n, 1, n);
        for (i = 1; i \le n; i++) {
          for (j = 1; j \le n; j++) {
             G2[i][j] = G[i][j] * 2.0;
        Matrix\_Inverse(n, G2, Ginv);
        printf("Inverse_lof_lG_lfrom_lequation_l5.56\n");
        wrmatrix(n, G2);
        printf("G_{\sqcup}from_{\sqcup}equation_{\sqcup}5.56\n");
        wrmatrix(n, Ginv);
        free\_matrix(Ginv, 1, n, 1, n);
        free\_matrix(G2, 1, n, 1, n);
  }
#endif
```

This code is used in section 128.

133. Now we get the part that I really don't understand. However, I know that this works. There are a couple of confusing transposes and bizarre incorporation of twoaw, but everything hangs together. Now since the single layer matrices R and T are the solutions to the systems of equations

$$GR = C$$
  $G(t+I) = I$ 

```
We do the little shuffle and only find the LU decomposition of G once and use it to find both R and T+1.
```

```
\langle \text{ Calculate } R \text{ and } T \text{ 133} \rangle \equiv
   Transpose\_Matrix(n, G);
   Decomp(n, G, \& condition, ipvt);
   \textbf{if} \ (\textit{condition} \equiv 1 \cdot 10^{32}) \ \textit{AD\_error}(\texttt{"Singular} \sqcup \texttt{Matrix} \sqcup \ldots \sqcup \texttt{failed} \sqcup \texttt{in} \sqcup \texttt{diamond\_init} \setminus \texttt{n"});\\
   for (i = 1; i \le n; i++) {
      \langle Solve for row of R 134\rangle
      \langle Solve for row of T 135\rangle
\#\mathbf{ifdef} MARTIN_HAMMER
   {
      double **T2, **Ginv;
      if (Martin\_Hammer \equiv 5) {
         T2 = dmatrix(1, n, 1, n);
         Ginv = dmatrix(1, n, 1, n);
         Copy\_Matrix(n, T, T2);
         for (i = 1; i \le n; i++) {
            T2[i][i] += 1/twoaw[i];
         for (i = 1; i \le n; i++) {
            for (j = 1; j \le n; j ++) {
               T2[i][j] *= twoaw[j] * 0.5;
         }
         printf("G=(T-1)/2 \cup from \cup equation \cup 5.55 \setminus n");
         wrmatrix(n, T2);
         Matrix\_Inverse(n, T2, Ginv);
         printf("1/G\n");
         wrmatrix(n, Ginv);
         free\_matrix(T2, 1, n, 1, n);
         free\_matrix(Ginv, 1, n, 1, n);
   }
#endif
```

This code is used in section 128.

134 We use the decomposed form of C to find I

**134.** We use the decomposed form of G to find R. Since G is now the LU decomposition of  $G^T$ , we must pass rows of the C to Solve and get rows back. Note the finess with

$$\operatorname{work}_{j} = C_{ji} \frac{a_{j} w_{j}}{a_{i} w_{i}}$$

To get everything in the right place. This is discussed in Wiscombe's appendix. Finally, we dutifully put these values back in R and divide by  $1/(2\mu_j w_j)$  so that R will be symmetric and have the proper form.

```
 \begin{split} &\langle \text{ Solve for row of } R \ 134 \rangle \equiv \\ & \textbf{for } (j=1; \ j \leq n; \ j++) \ work[j] = C[j][i] * twoaw[j]/twoaw[i]; \\ & Solve(n,G,work,ipvt); \\ & \textbf{for } (j=1; \ j \leq n; \ j++) \ R[i][j] = work[j]/twoaw[j]; \end{split}  This code is used in section 133.
```

135. We again use the decomposed form of G to find T. This is much simpler since we only need to pass rows of the identity matrix back and forth. We again carefully put these values back in T and divide by  $1/(2\mu_j w_j)$  so that T is properly formed. Oh yes, we can't forget to subtract the identity matrix!

```
 \langle \text{Solve for row of } T \mid 135 \rangle \equiv \\ \text{for } (j=1; \ j \leq n; \ j++) \quad work[j] = 0; \\ work[i] = 1.0; \\ Solve(n,G,work,ipvt); \\ \text{for } (j=1; \ j \leq n; \ j++) \quad T[i][j] = work[j]/twoaw[j]; \\ T[i][i] = 1.0/twoaw[i]; \quad /* \text{ Subtract Identity Matrix */}  This code is used in section 133.
```

This code is used in section 128.

136. Pretty standard stuff here. Allocate memory and print a warning if the thickness is too small.

```
\langle \text{Local variables and initialization } 136 \rangle \equiv
  int i, j, n;
  double **A, **G, **C;
  double a, c, d, temp;
  double *work:
  double condition;
  int *ipvt;
  d = method.b_thinnest;
  a = method.a\_calc;
  n = method.quad.pts;
  A = dmatrix(1, n, 1, n);
  G = dmatrix(1, n, 1, n);
  C = dmatrix(1, n, 1, n);
  work = dvector(1, n);
  ipvt = ivector(1, n);
  if (d < 1 \cdot 10^{-4}) AD_{error}("****_LRoundoff_Lerror_Lis_La_Lproblem--Use_LIGI_Lmethod n");
```

```
137.
```

```
\langle \text{ Free up memory } 137 \rangle \equiv \\ free\_dvector(work, 1, n); \\ free\_ivector(ipvt, 1, n); \\ free\_dmatrix(A, 1, n, 1, n); \\ free\_dmatrix(G, 1, n, 1, n); \\ free\_dmatrix(C, 1, n, 1, n); \\ \text{This code is used in section } 128.
```

# 138. Layer Initialization.

139.  $Init\_Layer$  returns reflection and transmission matrices for a thin layer. Space must previously been allocated for R and T.

```
\langle \text{Prototype for } \textit{Init\_Layer } 139 \rangle \equiv 
void \textit{Init\_Layer}(\text{struct AD\_slab\_type } slab, \text{struct AD\_method\_type } method, \text{double } **R, \text{double } **T)
```

This code is used in sections 107 and 140.

This code is used in section 106.

141. AD Double. This has the routines needed to add layers together in various combinations.

```
\langle ad\_doubl.c 141 \rangle \equiv
#include <math.h>
#include <float.h>
#include "nr_util.h"
#include "ad_matrx.h"
#include "ad_globl.h"
#include "ad_doubl.h"
  ⟨ Definition for Star_Multiply 161⟩
   ⟨ Definition for Star_One_Minus 162 ⟩
   \langle Definition for Basic\_Add\_Layers 143 \rangle
   (Definition for Basic_Add_Layers_With_Sources 144)
   \langle \text{ Definition for } Add | 147 \rangle
    Definition for Add_-With\_Sources 149\rangle
   Definition for Add_Homogeneous 151
   (Definition for Double_Once 153)
   (Definition for Double_Until 155)
   (Definition for Double_Until_Infinite 157)
  \langle \text{ Definition for } Between 159 \rangle
        \langle ad_doubl.h \quad 142 \rangle \equiv
142.
   \langle \text{ Prototype for } Add \ 146 \rangle;
   \langle Prototype for Add_With_Sources 148 \rangle;
   \langle Prototype for Add\_Homogeneous 150 \rangle;
   \langle Prototype for Double\_Once 152 \rangle;
   ⟨ Prototype for Double_Until 154⟩;
   ⟨ Prototype for Double_Until_Infinite 156⟩;
  \langle \text{ Prototype for } Between | 158 \rangle;
```

# 143. Basic Routine to Add Layers Without Sources.

The basic equations for the adding-doubling method (neglecting sources) are

$$\begin{split} \mathbf{T}^{02} &= \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \mathbf{R}^{12})^{-1} \mathbf{T}^{01} \\ \mathbf{R}^{20} &= \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \mathbf{R}^{12})^{-1} \mathbf{R}^{10} \mathbf{T}^{21} + \mathbf{R}^{21} \\ \mathbf{T}^{20} &= \mathbf{T}^{10} (\mathbf{E} - \mathbf{R}^{12} \mathbf{R}^{10})^{-1} \mathbf{T}^{21} \\ \mathbf{R}^{02} &= \mathbf{T}^{10} (\mathbf{E} - \mathbf{R}^{12} \mathbf{R}^{10})^{-1} \mathbf{R}^{12} \mathbf{T}^{01} + \mathbf{R}^{01} \end{split}$$

Upon examination it is clear that the two sets of equations have the same form. Therefore if I implement the first two equations, then the second set can be obtained by suitable switching of the parameters. Furthermore, these equations assume some of the multiplications are star multiplications. Explicitly,

$$\mathbf{T}^{02} = \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1} \mathbf{T}^{01}$$

and

$$\mathbf{R}^{20} = \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1} \mathbf{R}^{10} \star \mathbf{T}^{21} + \mathbf{R}^{21}$$

where the identity matrix E is then

$$\mathbf{E}^{ij} = \frac{1}{2\mu_i w_i} \delta_{ij}$$

where  $\delta_{ij}$  is the usual Kronecker delta. It is noteworthy that if say  $R^{10} \equiv 0$ , then  $\mathbf{E}^{-1} \equiv \mathbf{c}$  and so

$$\mathbf{T}^{02} = \mathbf{T}^{12}\mathbf{c}\mathbf{T}^{01} = \mathbf{T}^{12}\star\mathbf{T}^{01}$$

One goal of this routine was to make it efficient and easy to use. It is possible to call this routine with the same pointer for all the different reflection matrices and the pointer for the transmission matrices may be the same also. (The reflection and transmission pointers may need to be distinct. The temporary memory pointers a and b must be distinct from each other and distinct from the reflection and transmission matrices.)

Note: it should be possible to eliminate the need for the matrix b if  $Inverse\_Multiply$  could be called with an argument list like  $Inverse\_Multiply(n, A, B, A)$ . A quick glance at the code suggests that this would just force the allocation of the matrix into the  $Inverse\_Multiply$  routine and no net gain would result.

This code is used in section 141.

# 144. Basic Routine to Add Layers With Sources.

The adding-doubling equations including source terms  $\mathbf{J}$  are identical to those given above for the reflection and transmission. The only difference is that the source terms must be kept track of separately according to

$$\mathbf{J}_{+}^{02} = \mathbf{J}_{+}^{12} + \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \mathbf{R}^{12})^{-1} (\mathbf{J}_{+}^{01} + \mathbf{R}^{10} \mathbf{J}_{-}^{21})$$

and

$$\mathbf{J}_{+}^{20} = \mathbf{J}_{-}^{10} + \mathbf{T}^{10} (\mathbf{E} - \mathbf{R}^{12} \mathbf{R}^{10})^{-1} (\mathbf{J}_{-}^{21} + \mathbf{R}^{12} \mathbf{J}_{+}^{01})$$

where the + subscript indicates the downward direction and - indicates the upward direction. Note that these subscripts are not needed. Thus we have

$$\mathbf{J}^{02} = \mathbf{J}^{12} + \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \mathbf{R}^{12})^{-1} (\mathbf{J}^{01} + \mathbf{R}^{10} \mathbf{J}^{21})$$

and

$$\mathbf{J}^{20} = \mathbf{J}^{10} + \mathbf{T}^{10} (\mathbf{E} - \mathbf{R}^{12} \mathbf{R}^{10})^{-1} (\mathbf{J}^{21} + \mathbf{R}^{12} \mathbf{J}^{01})$$

Again, it is apparent that clever switching of the arguments requires that only one set of equations needs to be calculated. These equations assume some of the multiplications are star multiplications. Explicitly,

$$\mathbf{J}^{02} = \mathbf{J}^{12} + \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1} (\mathbf{J}^{01} + \mathbf{R}^{10} \star \mathbf{J}^{21})$$

```
\langle Definition for Basic\_Add\_Layers\_With\_Sources 144 \rangle \equiv
   static void Basic_Add_Layers_With_Sources(int n, double **R10, double **T01, double **R12, double
                 **R21, double **T12, double **T21, double **R20, double **T02, double **J01, double
                **J12, double **J21, double **J02, double **a, double **b)
                                                        /* a = \mathbf{R}^{10} \star \mathbf{R}^{12} */
      Star\_Multiply(n, R10, R12, a);
                                               /* a = \mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12} */
      Star\_One\_Minus(n, a);
                                                            /* b = \mathbf{T}^{12} (\dot{\mathbf{E}} - \mathbf{R}^{10} \mathbf{R}^{12})^{-1} */
      Left\_Inverse\_Multiply(n, a, T12, b);
                                                       //* a = \mathbf{T}^{12} (\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1} \mathbf{R}^{10} */
      Matrix\_Multiply(n, b, R10, a);
                                                    /*'a = \mathbf{T}^{12}(\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1}\mathbf{R}^{10} \star \mathbf{T}^{21} */
      Star_Multiply(n, a, T21, a);
      Matrix\_Sum(n, R21, a, R20);
      Copy_{-}Matrix(n, TO1, a);
      Matrix_Multiply(n, b, a, T02);
                                                  /* \ a = \mathbf{R}^{10} \star \mathbf{J}^{21} * / 
/* \ a = \mathbf{J}^{01} + \mathbf{R}^{10} \star \mathbf{J}^{21} * / 
      Star\_Multiply(n, R10, J21, a);
      Matrix\_Sum(n, JO1, a, a);
                                                        /* J02 = \mathbf{T}^{12}(\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1}(\mathbf{J}^{01} + \mathbf{R}^{10} \star \mathbf{J}^{21}) */
      Matrix\_Multiply(n, b, a, J02):
      Matrix\_Sum(n, J02, J12, J02);
```

# This code is used in section 141.

# 146.

145.

```
\langle \text{ Prototype for } Add | 146 \rangle \equiv
```

 $\begin{array}{l} \textbf{void} \ \textit{Add} (\textbf{int} \ \textit{n}, \textbf{double} \ **\text{R01}, \textbf{double} \ **\text{R10}, \textbf{double} \ **\text{T10}, \textbf{double} \ **\text{R12}, \textbf{double} \ **\text{R12}, \textbf{double} \ **\text{R21}, \textbf{double} \ **\text{R21}, \textbf{double} \ **\text{R20}, \textbf{double} \ **\text{R20}, \textbf{double} \ **\text{T02}, \textbf{double} \ **\text{T20}) \\ \end{array}$ 

This code is used in sections 142 and 147.

Higher level routines.

147. Add returns the reflection and transmission matrices for two different layers added together. These matrices do not have to be homogeneous. The output matrices R20, R02, T20, and T02 should be distinct from the input matrices.

```
\langle \text{ Definition for } Add | 147 \rangle \equiv
  \langle Prototype for Add 146\rangle
     \langle Allocate memory for a and b 163\rangle
     Basic\_Add\_Layers(n, R10, T01, R12, R21, T12, T21, R20, T02, a, b);
     Basic\_Add\_Layers(n, R12, T21, R10, R01, T10, T01, R02, T20, a, b);
     \langle Free Memory for a and b 164\rangle
This code is used in section 141.
148.
\langle Prototype for Add_With_Sources 148 \rangle \equiv
  void Add_With_Sources(int n, double **R01, double **R10, double **T11, double **T10, double
        **J01, double **J10, double **R12, double **R21, double **T12, double **T21, double
        **J12, double **J21, double **R02, double **R20, double **T02, double **T20, double
        **J02, double **J20)
This code is used in sections 142 and 149.
       Add_With_Sources returns the reflection and transmission matrices for two different layers added
together. These matrices do not have to be homogeneous. The output matrices R20, R02, T20, T02, J20,
and J02 should be distinct from the input matrices.
\langle \text{ Definition for } Add\_With\_Sources \ 149 \rangle \equiv
  \langle Prototype for Add_With_Sources 148 \rangle
     \langle Allocate memory for a and b 163\rangle
     Basic_Add_Layers_With_Sources (n, R10, T01, R12, R21, T12, T21, R20, T02, J01, J12, J21, J02, a, b);
     Basic_Add_Layers_With_Sources (n, R12, T21, R10, R01, T10, T01, R02, T20, J21, J10, J01, J20, a, b);
     \langle Free Memory for a and b 164\rangle
This code is used in section 141.
150.
\langle \text{ Prototype for } Add\_Homogeneous | 150 \rangle \equiv
  void Add_Homogeneous(int n, double **R01, double **T01, double **R12, double **T12, double
        **R02, double **T02)
This code is used in sections 142 and 151.
151.
\langle \text{ Definition for } Add\_Homogeneous | 151 \rangle \equiv
  \langle Prototype for Add_Homogeneous 150 \rangle
     \langle Allocate memory for a and b 163\rangle
     Basic\_Add\_Layers(n, R01, T01, R12, R12, T12, T12, R02, T02, a, b);
     \langle Free Memory for a and b 164\rangle
This code is used in section 141.
```

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This just adds a layer to itself. Couldn't Basic\_Add\_Layers be used? It would mean that there would be no restriction on the use of variables — i.e., R could be used as both a factor and as a result.

```
\langle Prototype for Double\_Once 152 \rangle \equiv
  void Double\_Once(int n, double **R, double **T)
This code is used in sections 142 and 153.
153.
\langle \text{ Definition for } Double\_Once | 153 \rangle \equiv
  ⟨ Prototype for Double_Once 152⟩
     \langle Allocate memory for a and b 163\rangle
     Basic\_Add\_Layers(n, R, T, R, R, T, T, R, T, a, b);
     \langle Free Memory for a and b 164\rangle
  }
This code is used in section 141.
```

154. Double\_Until and Double\_Until\_Infinite are the only ones that really take advantage of the external allocation of memory from the routine. I was kind of careful to make sure that this routine terminates if bad start and end values are given i.e., end  $\neq$  start  $\cdot 2^k$ . Futhermore, it should work correctly if the target thickness is infinite. I suppose that I could put some error warnings in...but right now I don't want to take the time.

```
\langle Prototype for Double\_Until 154 \rangle \equiv
  void Double_Until(int n, double **r, double **t, double start, double end)
This code is used in sections 142 and 155.
155.
\langle \text{ Definition for } Double\_Until \ 155 \rangle \equiv
   ⟨ Prototype for Double_Until 154⟩
     if (end \equiv \texttt{HUGE\_VAL}) {
        Double\_Until\_Infinite(n, r, t);
        return;
        \langle Allocate memory for a and b 163\rangle
        while (fabs(end - start) > 0.00001 \land end > start) {
           Basic\_Add\_Layers(n, r, t, r, r, t, t, r, t, a, b);
           start *= 2;
        \langle Free Memory for a and b 164\rangle
This code is used in section 141.
```

156. Double\_Until\_Infinite continues doubling until the thickness of the slab is essentially infinite. Originally I had defined infinite as a diffuse transmission less than  $10^{-6}$ . However, when the albedo is unity, then this is kind of impractical and I changed the definition of infinity to be that the diffuse transmission changes by less than one part in  $10^{-6}$  after one doubling step. The more I think about this, the less sense it makes....

```
\langle Prototype for Double\_Until\_Infinite 156 \rangle \equiv
  void Double_Until_Infinite(int n, double **r, double **t)
This code is used in sections 142 and 157.
```

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### 157.

```
 \langle \text{ Definition for } \textit{Double\_Until\_Infinite } 157 \rangle \equiv \\ \langle \text{ Prototype for } \textit{Double\_Until\_Infinite } 156 \rangle \\ \{ \\ \textbf{double } \textit{oldutu}, \text{ UTU, UT1;} \\ \langle \text{ Allocate memory for } a \text{ and } b \text{ 163} \rangle \\ \text{UTU} = 0.0; \\ \textbf{do } \{ \\ \textit{oldutu} = \text{UTU;} \\ \textit{Basic\_Add\_Layers}(n,r,t,r,t,t,r,t,a,b); \\ \textit{URU\_and\_UR1}(n,1.0,t,\&\text{UTU},\&\text{UT1}); \\ \} \textbf{ while } (\textit{fabs}(\text{UTU}-\textit{oldutu}) \geq 0.000001); \\ \langle \text{ Free Memory for } a \text{ and } b \text{ 164} \rangle \\ \}
```

This code is used in section 141.

### 158. Internal Radiance.

Between finds the radiance between two slabs. This equation for the upward radiance at the interface between two layers is

$$\mathbf{L}_{-} = (\mathbf{E} - \mathbf{R}^{12} \star \mathbf{R}^{10})^{-1} (\mathbf{R}^{12} \star \mathbf{T}^{01} \star \mathbf{L}_{+}^{0} + \mathbf{T}^{21} \star \mathbf{L}_{-}^{2})$$

where  $\mathbf{L}_{+}^{0}$  is the downward radiance on the top layer and  $\mathbf{L}_{-}^{2}$  is the upward radiance on the bottom layer. The equation for the downward mid-layer radiance can be obtained similarly using

$$\mathbf{L}_{+} = (\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1} (\mathbf{T}^{01} \star \mathbf{L}_{+}^{0} + \mathbf{R}^{10} \star \mathbf{T}^{21} \star \mathbf{L}_{-}^{2})$$

Now assume that  $\mathbf{L}_{-}^{2}$  is zero. Then the matrix

$$\mathbf{L}_{-} = (\mathbf{E} - \mathbf{R}^{12} \star \mathbf{R}^{10})^{-1} \mathbf{R}^{12} \star \mathbf{T}^{01}$$

can be used to find the downward fluence by simply star multiplying with the downward irradiance. Similarly,

$$\mathbf{L}_{+} = (\mathbf{E} - \mathbf{R}^{10} \star \mathbf{R}^{12})^{-1} \mathbf{T}^{01}$$

```
\langle \text{Prototype for } Between | 158 \rangle \equiv 
void Between(\text{int } n, \text{double } **R01, \text{double } **R10, \text{double } **T01, \text{double } **T10, \text{double } **R12, \text{double } **R21, \text{double } **T12, \text{double } **T21, \text{double } **Lup, \text{double } **Ldown)
```

This code is used in sections 142 and 159.

```
159.
        \langle \text{ Definition for } Between | 159 \rangle \equiv
  ⟨ Prototype for Between 158⟩
     \langle Allocate memory for a and b 163\rangle
     Star_Multiply(n, R10, R12, a);
     Star\_One\_Minus(n, a);
     Right\_Inverse\_Multiply(n, a, TO1, Ldown);
     Star\_Multiply(n, R12, R10, a);
     Star\_One\_Minus(n, a);
     Right\_Inverse\_Multiply(n, a, R12, b);
     Star\_Multiply(n, b, TO1, Lup);
     \langle Free Memory for a and b 164\rangle
```

This code is used in section 141.

#### 160. Utility routines.

Star matrix multiplication  $A \star B$  is defined to directly correspond to an integration, i.e. 161.

$$A \star B = \int_0^1 A(\mu, \mu') B(\mu', \mu'') 2\mu d\mu$$

then

$$A\star B = \sum_j A^{ij} 2\mu_j w_j B^{jk}$$

where  $\mu_i$  is the jth quadrature angle and  $w_i$  is its corresponding weight. It is sometimes useful to consider these matrix "star multiplications" as normal matrix multiplications which include a diagonal matrix c

$$\mathbf{c}_{ij} = 2\mu_i w_i \delta_{ij}$$

Thus a matrix star multiplication may be written

$$A \star B = A \mathbf{c} B$$

where the multiplications on the RHS of the above equation are usual matrix multiplications.

Since the routine Matrix Multiply that multiplies the matrices A and B to get C, allows A and C to be coincident. I first find C = Ac and then do  $C = C \cdot B$ . This allows us to avoid allocating a temporary matrix. A may occupy the same memory as C, but B and C must be distinct.

```
\langle \text{ Definition for } Star\_Multiply | 161 \rangle \equiv
  static void Star_Multiply (int n, double **A, double **B, double **C)
     Right\_Diagonal\_Multiply(n, A, twoaw, C);
     Matrix\_Multiply(n, C, B, C);
This code is used in section 141.
```

 $free\_dmatrix(a, 1, n, 1, n);$  $free\_dmatrix(b, 1, n, 1, n);$ 

This code is used in sections 147, 149, 151, 153, 155, 157,and 159.

This subtracts the matrix A from the unit matrix for star multiplication.  $\langle \text{ Definition for } Star\_One\_Minus | 162 \rangle \equiv$ static void Star\_One\_Minus(int n, double \*\*A) int i, j;  ${\bf for}\ (i=1;\ i\le n;\ i\!+\!\!+)\ \{$ for  $(j = 1; j \le n; j++) A[i][j] *= -1;$ A[i][i] += 1.0/twoaw[i];} This code is used in section 141. **163.**  $\langle$  Allocate memory for a and b 163 $\rangle \equiv$ **double** \*\*a, \*\*b; a = dmatrix(1, n, 1, n);b = dmatrix(1, n, 1, n);This code is used in sections 147, 149, 151, 153, 155, 157, and 159. 164.  $\langle$  Free Memory for a and b 164 $\rangle$   $\equiv$ 

# AD Boundary.

This section has routines associated with incorporating boundary conditions into the adding-doubling algorithm.

```
\langle ad\_bound.c \quad 165 \rangle \equiv
#include <math.h>
#include <stdio.h>
#include "ad_globl.h"
#include "ad_bound.h"
#include "ad_frsnl.h"
#include "ad_matrx.h"
#include "nr_util.h"
   \langle \text{ Prototype for } A\_Add\_Slide \ 174 \rangle;
   \langle \text{ Prototype for } B\_Add\_Slide \ 176 \rangle;
    Definition for Init_Boundary 169
    Definition for Boundary_RT 172
    \langle \text{ Definition for } Add\_Top 180 \rangle
    Definition for Add\_Bottom 182
    Definition for A\_Add\_Slide 175
    \langle \text{ Definition for } B\_Add\_Slide 177 \rangle
   \langle \text{ Definition for } Add\_Slides | 184 \rangle
   \langle \text{ Definition for } Sp\_RT | 186 \rangle
        \langle ad\_bound.h \quad 166 \rangle \equiv
   ⟨ Preprocessor definitions ⟩
   \langle Prototype for Init\_Boundary 168 \rangle;
   \langle Prototype for Boundary\_RT | 171 \rangle;
   \langle \text{ Prototype for } Add\_Top \ 179 \rangle;
   \langle Prototype for Add\_Bottom 181 \rangle;
    \langle Prototype for Add\_Slides 183 \rangle;
   \langle \text{ Prototype for } Sp\_RT \mid 185 \rangle;
```

#### 167. Boundary Initialization.

 $Init\_Boundary$  creates reflection and transmission matrices to simulate a boundary. If boundary  $\equiv$ TOP\_BOUNDARY then the arrays returned are for the top surface and the labels are as expected i.e. T01 is the reflection for light from air passing to the slab. Otherwise the calculations are made for the bottom surface and the labels are backwards i.e.  $T01 \equiv T32$  and  $T10 \equiv T23$ , where 0 is the first air slide surface, 1 is the slide/slab surface, 2 is the second slide/slab surface, and 3 is the bottom slide/air surface

```
\#define TOP_BOUNDARY 0
#define BOTTOM_BOUNDARY 1
\langle Prototype for Init\_Boundary 168 \rangle \equiv
  void Init_Boundary(struct AD_slab_type slab, int n,
  double *R01, double *R10, double *T01, double *T10,
  char boundary)
This code is used in sections 166 and 169.
```

This code is used in section 165.

170. Boundary\_RT computes the diagonal matrix (represented as an array) that characterizes reflection and transmission at an air (0), absorbing glass (1), slab (2) boundary. The reflection matrix is the same entering or exiting the slab. The transmission matrices should differ by a factor of  $(n_{\text{slab}}/n_{\text{outside}})^4$ , due to  $n^2$  law of radiance, but there is some inconsistency in the program and if I use this principle then regular calculations for R and T don't work and the fluence calculations still don't work. So punted and took all that code out.

The important point that must be remembered is that all the angles in this program assume that the angles are those actually in the sample. This allows angles greater that the critical angle to be used. Everything is fine as long as the index of refraction of the incident medium is 1.0. If this is not the case then the angle inside the medium must be figured out.

```
\langle \text{ Prototype for } Boundary\_RT | 171 \rangle \equiv
  void Boundary_RT(double n_-i, double n_-g, double n_-t, int n, double b,
  double *R, double *T)
This code is used in sections 166 and 172.
        \langle \text{ Definition for } Boundary\_RT | 172 \rangle \equiv
  \langle Prototype for Boundary\_RT 171 \rangle
     int i:
     double refl, trans;
     double mu;
     for (i = 1; i < n; i ++) {
        if (n_{-i} \equiv 1.0) mu = Cos\_Snell(n_{-t}, angle[i], n_{-i});
        else mu = angle[i];
        Absorbing\_Glass\_RT(n\_i, n\_g, n\_t, mu, b, \&refl, \&trans);
        R[i] = refl * twoaw[i];
        T[i] = trans;
  }
```

This code is used in section 165.

### 173. Boundary incorporation algorithms.

The next two routines  $A\_Add\_Slide$  and  $B\_Add\_Slide$  are modifications of the full addition algorithms for dissimilar layers. They are optimized to take advantage of the diagonal nature of the boundary matrices. There are two algorithms below to facilitate adding slides below and above the sample.

174.  $A\_Add\_Slide$  computes the resulting R20 and T02 matrices for a glass slide on top of an inhomogeneous layer characterized by R12, R21, T12, T21. It is ok if R21  $\equiv$  R12 and T12  $\equiv$  T21. But I do not think that it is required by this routine. The result matrices R20 and T02 should be independent of the input matrices None of the input matrices are changed

The critical quantites are

$$T_{02} = T_{12}(E - R_{10}R_{12})^{-1}T_{01}$$

and

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$$R_{20} = T_{12}(E - R_{10}R_{12})^{-1}R_{10}T_{21} + R_{21}$$

```
\langle \text{ Prototype for } A\_Add\_Slide \ 174 \rangle \equiv
  static void A_Add_Slide(int n,double **R12,double **R21,double **T12,double **T12,
  double *R10, double *T01, double **R20, double **T02,
  double **atemp, double **btemp)
This code is used in sections 165 and 175.
       \langle \text{ Definition for } A\_Add\_Slide | 175 \rangle \equiv
  \langle \text{ Prototype for } A\_Add\_Slide \ 174 \rangle
     double **ctemp;
     ctemp = R20;
     Left\_Diagonal\_Multiply(n, R10, R12, atemp);
     One\_Minus(n, atemp);
     Left\_Inverse\_Multiply(n, atemp, T12, ctemp);
     Right\_Diagonal\_Multiply(n, ctemp, T01, T02);
     Right\_Diagonal\_Multiply(n, ctemp, R10, btemp);
     Matrix\_Multiply(n, btemp, T21, atemp);
     Matrix\_Sum(n, R21, atemp, R20);
```

This code is used in section 165.

176.  $B\_Add\_Slide$  computes the resulting R02 and T20 matrices for a glass slide on top of an inhomogeneous layer characterized by R12, R21, T12, T21. It is ok if R21  $\equiv$  R12 and T12  $\equiv$  T21. But I do not think that it is required by this routine. The result matrices R02 and T20 should be independent of the input matrices None of the input matrices are changed

The critical equations are

$$T_{20} = T_{10}(E - R_{12}R_{10})^{-1}T_{21}$$

and

$$R_{02} = T_{10}(E - R_{12}R_{10})^{-1}R_{12}T_{01} + R_{01}$$

```
⟨ Prototype for B_Add_Slide 176⟩ ≡
    static void B_Add_Slide(int n, double **R12, double **T21,
    double *R01, double *R10, double *T01, double *T10,
    double **R02, double **T20,
    double **atemp, double **btemp)
```

This code is used in sections 165 and 177.

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```
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177.
        \langle \text{ Definition for } B\_Add\_Slide | 177 \rangle \equiv
  \langle \text{ Prototype for } B\_Add\_Slide \ 176 \rangle
     double **ctemp;
     int i;
     ctemp = R02;
     Right\_Diagonal\_Multiply(n, R12, R10, atemp);
     One\_Minus(n, atemp);
     Diagonal\_To\_Matrix(n, T10, btemp);
     Left\_Inverse\_Multiply(n, atemp, btemp, ctemp);
     Matrix\_Multiply(n, ctemp, T21, T20);
     Matrix\_Multiply(n, ctemp, R12, btemp);
     Right\_Diagonal\_Multiply(n, btemp, TO1, RO2);
     for (i = 1; i \le n; i ++) RO2[i][i] += RO1[i]/twoaw[i]/twoaw[i];
```

This code is used in section 165.

# Routines to incorporate slides.

Add\_Top calculates the reflection and transmission matrices for a slab with a boundary placed on 179. top of it.

```
size of matrix
                  R01, R10, T01, T10
                                            R, T for slide assuming 0=air and 1=slab
                  R12, R21, T12, T21
                                            R, T for slab assuming 1=slide and 2=?
                  R02, R20, T02, T20
                                            calc R, T for both assuming 0=air and 2=?
                  atemp, btemp
                                             previously allocated temporary storage matrices
\langle \text{ Prototype for } Add\_Top \ 179 \rangle \equiv
  void Add_Top(int n, double *R01, double *R10, double *T01, double *T10,
  double **R12, double **R21, double **T12, double **T21,
  double **R02, double **R20, double **T02, double **T20,
  double **atemp, double **btemp)
This code is used in sections 166 and 180.
180.
\langle \text{ Definition for } Add\_Top | 180 \rangle \equiv
  \langle \text{ Prototype for } Add\_Top 179 \rangle
     A\_Add\_Slide(n, R12, R21, T12, T21, R10, T01, R20, T02, atemp, btemp);
     B\_Add\_Slide(n, R12, T21, R01, R10, T01, T10, R02, T20, atemp, btemp);
```

This code is used in section 165.

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181. Add\_Bottom calculates the reflection and transmission matrices for a slab with a boundary placed beneath it

```
size of matrix
           R01, R10, T01, T10
                                      R, T for slab assuming 0=slab top and 1=slab bottom
           R12, R21, T12, T21
                                      R, T for slide assuming 1=slab bottom and 2=slide bottom
           R02, R20, T02, T20
                                      calc R, T for both assuming 0=slab top and 2=slide bottom
           atemp, btemp
                                      previously allocated temporary storage matrices
\langle \text{ Prototype for } Add\_Bottom | 181 \rangle \equiv
  void Add\_Bottom(int n, double **R01, double **R10, double **T11, double **T10,
  double *R12, double *R21, double *T12, double *T21,
  double **R02, double **R20, double **T02, double **T20,
  double **atemp, double **btemp)
This code is used in sections 166 and 182.
182.
\langle \text{ Definition for } Add\_Bottom | 182 \rangle \equiv
  \langle \text{ Prototype for } Add\_Bottom \ 181 \rangle
    A\_Add\_Slide(n, R10, R01, T10, T01, R12, T21, R02, T20, atemp, btemp);
    B\_Add\_Slide(n, R10, T01, R21, R12, T21, T12, R20, T02, atemp, btemp);
This code is used in section 165.
```

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183. Including identical slides.  $Add\_Slides$  is optimized for a slab with equal boundaries on each side.  $Add\_Slides$  calculates the reflection and transmission matrices for a slab with the same boundary placed above and below it. It is assumed that the slab is homogeneous. in this case the resulting R and T matrices are independent of direction. There are no constraints on R01, R10, T01, and T10. The handles for R and T cannot be equal to those for  $R\_total$  and  $T\_total$ .

n size of matrix R01, R10, T01, T10 R, T for slide assuming 0=air and 1=slab R, T R\_total,  $T_t$ total R, T for all 3 with top = bottom boundary atemp, btemp temporary storage matrices

If equal boundary conditions exist on both sides of the slab then, by symmetry, the transmission and reflection operator for light travelling from the top to the bottom are equal to those for light propagating from the bottom to the top. Consequently only one set need be calculated. This leads to a faster method for calculating the reflection and transmission for a slab with equal boundary conditions on each side. Let the top boundary be layer 01, the medium layer 12, and the bottom layer 23. The boundary conditions on each side are equal:  $R_{01} = R_{32}$ ,  $R_{10} = R_{23}$ ,  $T_{01} = T_{32}$ , and  $T_{10} = T_{23}$ . For example the light reflected from layer 01 (travelling from boundary 0 to boundary 1) will equal the amount of light reflected from layer 32, since there is no physical difference between the two cases. The switch in the numbering arises from the fact that light passes from the medium to the outside at the top surface by going from 1 to 0, and from 2 to 3 on the bottom surface. The reflection and transmission for the slab with boundary conditions are  $R_{30}$  and  $T_{03}$  respectively. These are given by

$$T_{02} = T_{12}(E - R_{10}R_{12})^{-1}T_{01}$$

$$R_{20} = T_{12}(E - R_{10}R_{12})^{-1}R_{10}T_{21} + R_{21}$$

$$T_{03} = T_{10}(E - R_{20}R_{10})^{-1}T_{02}$$

and

and

and

$$R_{30} = T_{10}(E - R_{20}R_{10})^{-1}R_{20}T_{01} + R_{01}$$

Further increases in efficiency may be made by exploiting the diagonal nature of the reflection and transmission operators for an interface, since most matrix/matrix multiplications above become vector/matrix multiplications.

```
\langle Prototype for Add\_Slides\ 183\rangle \equiv void Add\_Slides\ (int\ n, double\ *R01, double\ *R10, double\ *T01, double\ *T10, double\ **R, double\ **T, double\ **R\_total, double\ **T\_total, double\ **atemp, double\ **btemp)
This code is used in sections 166 and 184.
```

```
184.
```

```
\langle Definition for Add\_Slides 184\rangle \equiv
  \langle \text{ Prototype for } Add\_Slides \text{ 183} \rangle
    int i;
     double **R12, **R21, **T12, **T21;
    double temp;
    R12 = R:
    R21 = R;
    T21 = T;
    T12 = T;
     Left\_Diagonal\_Multiply(n, R10, R12, atemp);
     One\_Minus(n, atemp);
     Left\_Inverse\_Multiply(n, atemp, T12, T\_total);
     Right\_Diagonal\_Multiply(n, T\_total, R10, btemp);
     Matrix\_Multiply(n, btemp, T21, R\_total);
     Matrix\_Sum(n, R\_total, R21, R\_total);
     Right\_Diagonal\_Multiply(n, R\_total, R10, atemp);
     One\_Minus(n, atemp);
     Matrix\_Inverse(n, atemp, btemp);
     Left\_Diagonal\_Multiply(n, T10, btemp, atemp);
     Matrix\_Multiply(n, atemp, T\_total, btemp);
     Right\_Diagonal\_Multiply(n, btemp, TO1, T\_total);
     Matrix\_Multiply(n, atemp, R\_total, btemp);
     Right\_Diagonal\_Multiply(n, btemp, TO1, R\_total);
     for (i = 1; i \le n; i ++) {
       temp = twoaw[i];
       R_{-}total[i][i] += RO1[i]/(temp * temp);
  }
```

This code is used in section 165.

# 185. Specular R and T.

 $Sp\_RT$  calculates the specular reflection and transmission for light incident on a slide-slab-slide sandwich. The sample is characterized by the record slab. The total unscattered reflection and transmission for oblique irradiance (urx and utx) together with their companions uru and utu for diffuse irradiance. The cosine of the incident angle is specified by  $slab.cos\_angle$ .

The way that this routine calculates the diffuse unscattered quantities based on the global quadrature angles previously set-up. Consequently, these estimates are not exact. In fact if n=4 then only two quadrature points will actually be used to figure out the diffuse reflection and transmission (assuming mismatched boundaries).

This algorithm is pretty simple. Since the quadrature angles are all chosen assuming points **inside** the medium, I must calculate the corresponding angle for light entering from the outside. If the the cosine of this angle is greater than zero then the angle does not correspond to a direction in which light is totally internally reflected. For this ray, I find the unscattered that would be reflected or transmitted from the slab. I multiply this by the quadrature angle and weight twoaw[i] to get the total diffuse reflectance and transmittance.

Oh, yes. The mysterious multiplication by a factor of  $n\_slab * n\_slab$  is required to account for the  $n^2$ -law of radiance.

```
\langle \text{ Prototype for } Sp\_RT \mid 185 \rangle \equiv
  void Sp_RT (int n, struct AD_slab_type slab, double *ur1, double *ut1, double *uru, double *utu)
This code is used in sections 166 and 186.
       \langle \text{ Definition for } Sp\_RT | 186 \rangle \equiv
  \langle \text{ Prototype for } Sp\_RT \text{ 185} \rangle
     double mu\_outside, r, t;
     int i;
     *uru = 0;
     *utu = 0;
     for (i = 1; i \le n; i ++) {
        mu\_outside = Cos\_Snell(slab.n\_slab, angle[i], 1.0);
        if (mu\_outside \neq 0) {
           Sp\_mu\_RT(slab.n\_top\_slide, slab.n\_slab, slab.n\_bottom\_slide, slab.b\_top\_slide, slab.b,
                slab.b\_bottom\_slide, mu\_outside, \&r, \&t);
          *uru += twoaw[i]*r;
           *utu += twoaw[i] *t;
        }
     Sp\_mu\_RT(slab.n\_top\_slide, slab.n\_slab, slab.n\_bottom\_slide, slab.b\_top\_slide, slab.b, slab.b\_bottom\_slide,
           slab.cos\_angle, ur1, ut1);
     *uru *= slab.n\_slab * slab.n\_slab;
     *utu *= slab.n\_slab * slab.n\_slab;
```

This code is used in section 165.

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**187. AD Fresnel.** This is a part of the core suite of files for the adding-doubling program. Not surprisingly, this program includes routines to calculate Fresnel reflection.

```
\langle ad_frsnl.c 187 \rangle \equiv
#include <math.h>
#include <float.h>
#include <stdio.h>
#include "ad_frsnl.h"
   \langle \text{ Prototype for } Fresnel \ 194 \rangle;
    \langle \text{ Prototype for R1 } 208 \rangle;
    Definition for Cos\_Critical\_Angle 190
    (Definition for Cos_Snell 192)
    \langle \text{ Definition for } Fresnel 195 \rangle
    (Definition for Glass 197)
     Definition for Absorbing\_Glass\_RT 199\rangle
     Definition for R1 209
    \langle \text{ Definition for } Sp\_mu\_RT \text{ 204} \rangle
    \langle \text{ Definition for } Sp\_mu\_RT\_Flip 202 \rangle
   \langle \text{ Definition for } Diffuse\_Glass\_R \text{ 211} \rangle
188. \langle ad\_frsnl.h 188 \rangle \equiv
    \langle Prototype for Cos\_Critical\_Angle 189 \rangle;
    \langle Prototype for Cos_Snell 191 \rangle;
    \langle Prototype for Absorbing\_Glass\_RT 198 \rangle;
    \langle \text{ Prototype for } Sp\_mu\_RT \text{ 203} \rangle;
    \langle \text{ Prototype for } Sp\_mu\_RT\_Flip \ 201 \rangle;
    \langle Prototype for Diffuse\_Glass\_R 210 \rangle;
   \langle \text{ Prototype for } Glass \ 196 \rangle;
```

# 189. The critical angle.

 $Cos\_Critical\_Angle$  calculates the cosine of the critical angle. If there is no critical angle then 0.0 is returned (i.e.,  $cos(\pi/2)$ ). Note that no trigonmetric functions are required. Recalling Snell's law

$$n_i \sin \theta_i = n_t \sin \theta_t$$

To find the critical angle, let  $\theta_t = \pi/2$  and then

$$\theta_c = \sin^{-1} \frac{n_t}{n_i}$$

The cosine of this angle is then

$$\cos \theta_c = \cos \left( \sin^{-1} \frac{n_t}{n_i} \right) = \frac{\sqrt{n_i^2 - n_t^2}}{n_i}$$

or more simply

$$\cos \theta_c = \sqrt{1 - n^2}$$

```
where n = n_t/n_i.

\langle \text{Prototype for } Cos\_Critical\_Angle | 189 \rangle \equiv 
double Cos\_Critical\_Angle (\textbf{double } ni, \textbf{double } nt)
This code is used in sections 188 and 190.
```

```
190. \langle \text{Definition for } Cos\_Critical\_Angle \ 190 \rangle \equiv \langle \text{Prototype for } Cos\_Critical\_Angle \ 189 \rangle  {
    double x;
    if (nt \geq ni) return 0.0;
    else {
        x = nt/ni;
        x = sqrt(1.0 - x * x);
        return x;
    }
}
```

This code is used in section 187.

# 191. Snell's Law.

 $Cos\_Snell$  returns the cosine of the angle that the light propagates through a medium given the cosine of the angle of incidence and the indices of refraction. Let the cosine of the angle of incidence be  $\mu_t$ , the transmitted cosine as  $\mu_t$ , the index of refraction of the incident material  $n_i$  and that of the transmitted material be  $n_t$ .

Snell's law states

$$n_i \sin \theta_i = n_t \sin \theta_t$$

but if the angles are expressed as cosines,  $\mu_i = \cos \theta_i$  then

$$n_i \sin(\cos^{-1} \mu_i) = n_t \sin(\cos^{-1} \mu_t)$$

Solving for  $\mu_t$  yields

$$\mu_t = \cos\{\sin^{-1}[(n_i/n_t)\sin(\cos^{-1}\mu_i)]\}$$

which is pretty ugly. However, note that  $\sin(\cos^{-1}\mu) = \sqrt{1-\mu^2}$  and the above becomes

$$\mu_t = \sqrt{1 - (n_i/n_t)^2 (1 - \mu_i^2)}$$

and no trigonmetric calls are necessary. Hooray!

A few final notes. I check to make sure that the index of refraction of changes before calculating a bunch of stuff. This routine should not be passed incident angles greater than the critical angle, but I shall program defensively and test to make sure that the argument of the sqrt function is non-negative. If it is, then I return  $\mu_t = 0$  i.e.,  $\theta_t = 90^{\circ}$ .

I also pretest for the common but trivial case of normal incidence.

```
\langle \text{Prototype for } Cos\_Snell \ 191 \rangle \equiv  double Cos\_Snell(double n\_i, double mu\_i, double n\_t)
```

This code is used in sections 188 and 192.

```
192. \langle \text{Definition for } Cos\_Snell \ 192 \rangle \equiv \langle \text{Prototype for } Cos\_Snell \ 191 \rangle
{
    double temp;
    if (mu\_i \equiv 1.0) return 1.0;
    if (n\_i \equiv n\_t) return mu\_i;
    temp = n\_i/n\_t;
    temp = n\_i/n\_t;
    temp = 1.0 - temp * temp * (1.0 - mu\_i * mu\_i);
    if (temp < 0) return 0.0;
    else return (sqrt(temp));
    }

This code is used in section 187.
```

# 193. Fresnel Reflection.

Fresnel calculates the specular reflection for light incident at an angle  $\theta_i$  from the normal (having a cosine equal to  $\mu_i$ ) in a medium with index of refraction  $n_-i$  onto a medium with index of refraction  $n_-t$ .

The usual way to calculate the total reflection for unpolarized light is to use the Fresnel formula

$$R = \frac{1}{2} \left[ \frac{\sin^2(\theta_i - \theta_t)}{\sin^2(\theta_i + \theta_t)} + \frac{\tan^2(\theta_i - \theta_t)}{\tan^2(\theta_i + \theta_t)} \right]$$

where  $\theta_i$  and  $\theta_t$  represent the angle (from normal) that light is incident and the angle at which light is transmitted. There are several problems with calculating the reflection using this formula. First, if the angle of incidence is zero, then the formula results in division by zero. Furthermore, if the angle of incidence is near zero, then the formula is the ratio of two small numbers and the results can be inaccurate. Second, if the angle of incidence exceeds the critical angle, then the calculation of  $\theta_t$  results in an attempt to find the arcsine of a quantity greater than one. Third, all calculations in this program are based on the cosine of the angle. This routine forces the calling routine to find  $\theta_i = \cos^{-1} \mu$ . Fourth, the routine also gives problems when the critical angle is exceeded.

Closer inspection reveals that this is the wrong formulation to use. The formulas that should be used for parallel and perpendicular polarization are

$$R_{\parallel} = \left[ \frac{n_t \cos \theta_i - n_i \cos \theta_t}{n_t \cos \theta_i + n_i \cos \theta_t} \right]^2, \qquad R_{\perp} = \left[ \frac{n_i \cos \theta_i - n_t \cos \theta_t}{n_i \cos \theta_i + n_t \cos \theta_t} \right]^2.$$

The formula for unpolarized light, written in terms of  $\mu_i = \cos \theta_i$  and  $\mu_t = \cos \theta_t$  is

$$R = \frac{1}{2} \left[ \frac{n_t \mu_i - n_i \mu_t}{n_t \mu_i + n_i \mu_t} \right]^2 + \frac{1}{2} \left[ \frac{n_i \mu_i - n_t \mu_t}{n_i \mu_i + n_t \mu_t} \right]^2$$

This formula has the advantage that no trig routines need to be called and that the case of normal irradiance does not cause division by zero. Near normal incidence remains numerically well-conditioned. In the routine below, I test for matched boundaries and normal incidence to eliminate unnecessary calculations. I also test for total internal reflection to avoid possible division by zero. I also find the ratio of the indices of refraction to avoid an extra multiplication and several intermediate variables.

```
194.
        \langle \text{ Prototype for } Fresnel \ 194 \rangle \equiv
  static double Fresnel (double n_{-i}, double n_{-t}, double mu_{-i})
This code is used in sections 187 and 195.
       \langle \text{ Definition for } Fresnel | 195 \rangle \equiv
   (Prototype for Fresnel 194)
     double mu_t, ratio, temp, temp1;
     if (n_{-}i \equiv n_{-}t) return 0.0;
     if (mu_{-}i \equiv 1.0) {
        temp = (n_{-i} - n_{-t})/(n_{-i} + n_{-t});
        return (temp * temp);
     if (mu_{-}i \equiv 0.0) return 1.0;
     mu_{-}t = Cos_{-}Snell(n_{-}i, mu_{-}i, n_{-}t);
     if (mu_{-}t \equiv 0.0) return 1.0;
     ratio = n_{-}i/n_{-}t;
     temp = ratio * mu_-t;
     temp1 = (mu\_i - temp)/(mu\_i + temp);
     temp = ratio * mu_i;
     temp = (mu\_t - temp)/(mu\_t + temp);
     return ((temp1 * temp1 + temp * temp)/2);
  }
This code is used in section 187.
```

# 196. Reflection from a glass slide.

Glass calculates the total specular reflection (i.e., including multiple internal reflections) based on the indices of refraction of the incident medium  $n_{-i}$ , the glass  $n_{-g}$ , and medium into which the light is transmitted  $n_{-t}$  for light incident at an angle from the normal having cosine  $mu_{-i}$ .

In many tissue optics problems, the sample is constrained by a piece of glass creating an air-glass-tissue sequence. The adding-doubling formalism can calculate the effect that the layer of glass will have on the radiative transport properties by including a layer for the glass-tissue interface and a layer for the air-glass interface. However, it is simpler to find net effect of the glass slide and include only one layer for the glass boundary.

The first time I implemented this routine, I did not include multiple internal reflections. After running test cases, it soon became apparent that the percentage errors were way too big for media with little absorption and scattering. It is not hard to find the result for the reflection from a non-absorbing glass layer (equation A2.21 in my dissertation) in which multiple reflections are properly accounted for

$$r_g = \frac{r_1 + r_2 - 2r_1r_2}{1 - r_1r_2}$$

Here  $r_1$  is the reflection at the air-glass interface and  $r_2$  is the reflection at the glass-sample interface.

There is one pitfall in calculating  $r_g$ . When the angle of incidence exceeds the critical angle then the formula above causes division by zero. If this is the case then  $r_1 = 1$  and can easily be tested for.

To eliminate unnecessary computation, I check to make sure that it really is necessary to call the Fresnel routine twice. It is noteworthy that the formula for  $r_g$  works correctly if the the first boundary is not totally reflecting but the second one is. Note that  $\mu_g$  gets calculated twice (once in the first call to Fresnel and once directly).

```
\langle \text{ Prototype for } Glass | 196 \rangle \equiv  double Glass(double n_{-}i,double n_{-}g,double n_{-}t,double mu_{-}i) This code is used in sections 188 and 197.
```

```
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```

```
197. \langle \text{ Definition for } Glass | 197 \rangle \equiv \langle \text{ Prototype for } Glass | 196 \rangle {

double r1, r2, mu\_g, temp;

if (n\_i \equiv n\_g) return (Fresnel(n\_g, n\_t, mu\_i));

r1 = Fresnel(n\_i, n\_g, mu\_i);

if (r1 \geq 1.0 \vee n\_g \equiv n\_t) return r1;

mu\_g = Cos\_Snell(n\_i, mu\_i, n\_g);

r2 = Fresnel(n\_g, n\_t, mu\_g);

temp = r1 * r2;

temp = (r1 + r2 - 2 * temp)/(1 - temp);

return temp;

}
```

This code is used in section 187.

# 198. Reflection from an absorbing slide.

Absorbing\_Glass\_RT calculates the total specular reflection and transmission (i.e., including multiple internal reflections) based on the indices of refraction of the incident medium  $n_-i$ , the glass  $n_-g$ , and medium into which the light is transmitted  $n_-t$  for light incident at an angle from the normal having cosine  $mu_-i$ . The optical thickness of the glass  $b = \mu_a d$  is measured normal to the glass.

This routine was generated to help solve a problem with the inverse adding-doubling program associated with samples with low absorbances. A particular situation arises when the slides have significant absorption relative to the sample absorption. Anyway, it is not hard to extend the result for non-absorbing slides to the absorbing case

$$r = \frac{r_1 + (1 - 2r_1)r_2 \exp(-2b/\mu_g)}{1 - r_1 r_2 \exp(-2b/\mu_g)}$$

Here  $r_1$  is the reflection at the sample-glass interface and  $r_2$  is the reflection at the glass-air interface and  $\mu_g$  is the cosine of the angle inside the glass. Note that if  $b \neq 0$  then the reflection depends on the order of the indices of refraction, otherwise  $n_{-i}$  and  $n_{-t}$  can be switched and the result should be the same.

The corresponding result for transmission is

$$t = \frac{(1 - r_1)(1 - r_2)\exp(-b/\mu_g)}{1 - r_1r_2\exp(-2b/\mu_g)}$$

There are two potential pitfalls in the calculation. The first is when the angle of incidence exceeds the critical angle then the formula above causes division by zero. If this is the case, *Fresnel* will return  $r_1 = 1$  and this routine responds appropriately. The second case is when the optical thickness of the slide is too large.

I don't worry too much about optimal coding, because this routine does not get called all that often and also because *Fresnel* is pretty good at avoiding unnecessary computations. At worst this routine just has a couple of extra function calls and a few extra multiplications.

I also check to make sure that the exponent is not too small.

```
\langle \text{Prototype for } Absorbing\_Glass\_RT \text{ 198} \rangle \equiv 
void Absorbing\_Glass\_RT \text{ (double } n\_i, \text{ double } n\_g, \text{ double } n\_t, \text{ double } mu\_i, \text{ double } b, \text{ double } *r, \text{ double } *t \rangle
```

This code is used in sections 188 and 199.

```
199.
        \langle Definition for Absorbing\_Glass\_RT 199 \rangle \equiv
  \langle Prototype for Absorbing\_Glass\_RT 198 \rangle
     double r1, r2, mu_{-}g, expo, denom;
     *t = 0:
     *r = Fresnel(n_{-i}, n_{-q}, mu_{-i});
     if (*r \ge 1.0 \lor b \equiv \texttt{HUGE\_VAL} \lor mu\_i \equiv 0.0) return;
     mu\_g = Cos\_Snell(n\_i, mu\_i, n\_g);
     r1 = *r;
     r2 = Fresnel(n_{-q}, n_{-t}, mu_{-q});
     if (b \equiv 0.0) {
       *r = (r1 + r2 - 2.0 * r1 * r2)/(1 - r1 * r2);
        *t = 1.0 - (*r);
     else {
       expo = -b/mu_{-}q;
       if (2 * expo \leq DBL_MIN_10_EXP * 2.3025851) return;
        expo = exp(expo);
        denom = 1.0 - r1 * r2 * expo * expo;
       *r = (r1 + (1.0 - 2.0 * r1) * r2 * expo * expo)/denom;
       *t = (1.0 - r1) * (1.0 - r2) * expo/denom;
  }
This code is used in section 187.
```

# 200. Unscattered refl and trans for a sample.

**201.**  $Sp\_mu\_RT\_Flip$  finds the reflectance to incorporate flipping of the sample. This is needed when the sample is flipped between measurements.

```
\langle \text{Prototype for } Sp\_mu\_RT\_Flip | 201 \rangle \equiv 
void Sp\_mu\_RT\_Flip (\text{int } flip, \text{double } n\_top, \text{double } n\_slab, \text{double } n\_bottom, \text{double } tau\_top, \text{double } tau\_slab, \text{double } tau\_bottom, \text{double } mu, \text{double } *r, \text{double } *t)
```

This code is used in sections 188 and 202.

```
202. \langle \text{Definition for } Sp\_mu\_RT\_Flip \ 202 \rangle \equiv \langle \text{Prototype for } Sp\_mu\_RT\_Flip \ 201 \rangle  {  Sp\_mu\_RT (n\_top, n\_slab, n\_bottom, tau\_top, tau\_slab, tau\_bottom, mu, r, t);  if (flip \land n\_top \neq n\_bottom \land tau\_top \neq tau\_bottom) {  double \ correct\_r = *r;   Sp\_mu\_RT (n\_bottom, n\_slab, n\_top, tau\_bottom, tau\_slab, tau\_top, mu, r, t);   *r = correct\_r;  } } } }  This code is used in section 187.
```

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 $Sp_{-}mu_{-}RT$  calculates the unscattered reflection and transmission (i.e., specular) through a glass-slabglass sandwich. Light is incident at an angle having a cosine mu from air onto a possibly absorbing glass plate with index  $n_{-}top$  on a sample with index  $n_{-}slab$  resting on another possibly absorbing glass plate with index  $n\_bottom$  and then exiting into air again.

```
The optical thickness of the slab is tau\_slab.
\langle \text{ Prototype for } Sp\_mu\_RT \text{ 203} \rangle \equiv
   void Sp\_mu\_RT (double n\_top, double n\_slab, double n\_bottom, double tau\_top, double
        tau\_slab, double tau\_bottom, double mu, double *r, double *t)
This code is used in sections 188 and 204.
204. \langle \text{ Definition for } Sp\_mu\_RT | 204 \rangle \equiv
   \langle \text{ Prototype for } Sp\_mu\_RT \text{ 203} \rangle
     double r_{-}top, r_{-}bottom, t_{-}top, t_{-}bottom, mu_{-}slab, beer, denom, temp, mu_{-}in_{-}slab;
     *r = 0;
     *t = 0;
     Absorbing\_Glass\_RT(1.0, n\_top, n\_slab, mu, tau\_top, \&r\_top, \&t\_top);
     mu\_in\_slab = Cos\_Snell(1.0, mu, n\_slab);
     Absorbing\_Glass\_RT(n\_slab, n\_bottom, 1.0, mu\_in\_slab, tau\_bottom, \&r\_bottom, \&t\_bottom);
     ⟨ Calculate beer 206⟩
      \langle \text{ Calculate } r \text{ and } t \text{ 207} \rangle
```

This code is used in section 187.

- 205. Nothing tricky here except a check to make sure that the reflection for the top is not equal to that on the bottom before calculating it again. I also drop out of the routine if the top surface is totally reflecting.
- 206. I am careful here not to cause an underflow error and to avoid division by zero.

It turns out that I found a small error in this code fragment. Basically I misunderstood what one of the values in float.h represented. This version is now correct

```
\langle \text{ Calculate } beer 206 \rangle \equiv
  mu\_slab = Cos\_Snell(1.0, mu, n\_slab);
  if (mu\_slab \equiv 0) beer = 0.0;
  else if (tau\_slab \equiv HUGE\_VAL) beer = 0.0;
     temp = -tau\_slab/mu\_slab;
     if (2 * temp \le DBL_MIN_10_EXP * 2.3025851) beer = 0.0;
     else beer = exp(temp);
This code is used in section 204.
```

207. If  $r_{\text{top}}$  is the reflection for the top and  $r_{\text{bottom}}$  is that for the bottom surface then the total reflection will be

UNSCATTERED REFL AND TRANS FOR A SAMPLE

$$r = r_{\text{top}} + \frac{r_{\text{bottom}} t_{\text{top}}^2 \exp(-2\tau/\mu)}{1 - r_{\text{top}} r_{\text{bottom}} \exp(-2\tau/\mu)}$$

and the transmission is

$$t = \frac{t_{\text{top}}t_{\text{bottom}} \exp(-\tau/\mu)}{1 - r_{\text{top}}r_{\text{bottom}} \exp(-2\tau/\mu)}$$

where  $\mu$  is the angle inside the slab and  $\tau$  is the optical thickness of the slab.

I have already calculated the reflections and the exponential attenuation, so I can just plug into the formula after making sure that it is really necessary. The denominator cannot be zero since I know  $r\_top < 1$  and that  $r\_bottom$  and beer are less than or equal to one.

The bug that was fixed was in the calculated reflection. I omitted a  $r_{\text{bottom}}$  in the numerator of the fraction used to calculate the reflection.

```
 \begin{split} &\langle \, \text{Calculate } r \, \, \text{and} \, \, t \, \, 207 \, \rangle \equiv \\ & \quad \text{if} \, \left( \, beer \equiv 0.0 \right) \, \left\{ \\ & \quad *r = r\_top; \\ & \quad \rbrace \\ & \quad \text{else} \, \left\{ \\ & \quad temp = t\_top * beer; \\ & \quad denom = 1 - r\_top * r\_bottom * beer * beer; \\ & \quad *r = r\_top + r\_bottom * temp * temp / denom; \\ & \quad *t = t\_bottom * temp / denom; \\ & \quad \rbrace \end{aligned}
```

This code is used in section 204.

# 208. Total diffuse reflection.

R1 calculates the first moment of the Fresnel reflectance using the analytic solution of Walsh. The integral of the first moment of the Fresnel reflection  $(R_1)$  has been found analytically by Walsh, [see Ryde 1931]

$$R_1 = \frac{1}{2} + \frac{(m-1)(3m+1)}{6(m+1)^2} + \left[ \frac{m^2(m^2-1)^2}{(m^2+1)^3} \right] \log \left( \frac{m-1}{m+1} \right) - \frac{2m^3(m^2+2m-1)}{(m^2+1)(m^4-1)} + \left[ \frac{8m^4(m^4+1)}{(m^2+1)(m^4-1)^2} \right] \log m$$

where Walsh's parameter  $m = n_t/n_i$ . This equation is only valid when  $n_i < n_t$ . If  $n_i > n_t$  then using (see Egan and Hilgeman 1973),

$$\frac{1 - R_1(n_i/n_t)}{n_t^2} = \frac{1 - R_1(n_t/n_i)}{n_i^2}$$

or

$$R(1/m) = 1 - m^2[1 - R(m)]$$

```
\langle \text{Prototype for R1 208} \rangle \equiv  static double R1(double ni, double nt)
```

This code is used in sections 187 and 209.

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```
209. \langle Definition for R1 \frac{209}{} \rangle \equiv
  ⟨Prototype for R1 208⟩
    double m, m2, m4, mm1, mp1, r, temp;
    if (ni \equiv nt) return 0.0;
    if (ni < nt) m = nt/ni;
    else m = ni/nt;
    m2 = m * m;
    m4 = m2 * m2;
    mm1 = m - 1;
    mp1 = m + 1;
    temp = (m2 - 1)/(m2 + 1);
    r = 0.5 + mm1 * (3 * m + 1)/6/mp1/mp1;
    r += m2 * temp * temp/(m2 + 1) * log(mm1/mp1);
    r = 2 * m * m2 * (m2 + 2 * m - 1)/(m2 + 1)/(m4 - 1);
    r += 8 * m4 * (m4 + 1)/(m2 + 1)/(m4 - 1)/(m4 - 1) * log(m);
    if (ni < nt) return r;
    else return (1 - (1 - r)/m2);
This code is used in section 187.
210. Diffusion reflection from a glass slide.
  Diffuse_Glass_R returns the total diffuse specular reflection from the air-glass-tissue interface
\langle Prototype for Diffuse\_Glass\_R 210 \rangle \equiv
  double Diffuse_Glass_R(double nair, double nslide, double nslab)
This code is used in sections 188 and 211.
211. \langle \text{ Definition for } Diffuse\_Glass\_R \ 211 \rangle \equiv
  \langle Prototype for Diffuse\_Glass\_R 210 \rangle
    double rairglass, rglasstissue, rtemp;
    rairglass = R1(nair, nslide);
    rglasstissue = R1(nslide, nslab);
    rtemp = rairglass * rglasstissue;
    if (rtemp \ge 1) return 1.0;
    else return ((rairglass + rglasstissue - 2 * rtemp)/(1 - rtemp));
This code is used in section 187.
```

## 212. AD Matrix.

This is a part of the core suite of files for the adding-doubling program. Not surprisingly, this program includes routines to manipulate matrices. These routines require that the matrices be stored using the allocation scheme outline in  $Numerical\ Recipes$  by Press  $et\ al.$  I have spent some time optimizing the matrix multiplication routine  $Matrix\_Multiply$  because roughly half the time in any adding-doubling calculation is spent doing matrix multiplication. Lastly, I should mention that all the routines assume a square matrix of size n by n.

213. In this module I collect up information that needs to be written to the header file ad\_matrx.h so that other source files that want to make use of the function defined here will have the necessary declarations available.

## 214. Simple Matrix Routines.

```
215. Copy\_Matrix replaces the matrix B by A \langle Prototype for Copy\_Matrix\ _{215}\rangle \equiv void Copy\_Matrix(\mathbf{int}\ n, \mathbf{double}\ **A, \mathbf{double}\ **B) This code is used in sections 213 and 216.
```

```
\langle \text{ Definition for } Copy\_Matrix \ 216 \rangle \equiv
  ⟨ Prototype for Copy_Matrix 215⟩
     double *a\_ptr, *b\_ptr, *a\_last;
     a\_last = \&A[n][n];
     a_{-}ptr = \&A[1][1];
     b_{-}ptr = \&B[1][1];
     while (a_{-}ptr \le a_{-}last) *b_{-}ptr ++ = *a_{-}ptr ++;
This code is used in section 212.
217. One_Minus replaces the matrix A by 1-A
\langle \text{ Prototype for } One\_Minus \ 217 \rangle \equiv
  void One\_Minus(int n, double **A)
This code is used in sections 213 and 218.
218. \langle Definition for One\_Minus \ 218 \rangle \equiv
  \langle \text{ Prototype for } One\_Minus 217 \rangle
     int i, j;
     for (i = 1; i \le n; i++) {
        for (j = 1; j \le n; j++) A[i][j] *= -1;
        A[i][i] += 1.0;
This code is used in section 212.
        Transpose_Matrix transposes a matrix.
\langle Prototype for Transpose\_Matrix 219 \rangle \equiv
  void Transpose_Matrix(int n, double **a)
This code is used in sections 213 and 220.
220. \langle \text{ Definition for } Transpose\_Matrix \underline{220} \rangle \equiv
  \langle Prototype for Transpose\_Matrix 219 \rangle
     int i, j;
     double swap;
     for (i = 1; i \le n; i++) {
        for (j = i + 1; j \le n; j ++) {
           swap = a[i][j];
          a[i][j] = a[j][i];
          a[j][i] = swap;
This code is used in section 212.
```

**225.** Left\_Diagonal\_Multiply multiplies the diagonal matrix a by the matrix B, puts the result in C. B and C can be the same matrix

```
\langle \text{Prototype for } \textit{Left\_Diagonal\_Multiply } 225 \rangle \equiv  void \textit{Left\_Diagonal\_Multiply}(\text{int } n, \text{double } *A, \text{double } **B, \text{double } **C) This code is used in sections 213 and 226.
```

```
226. \langle Definition for Left\_Diagonal\_Multiply\ 226 \rangle \equiv \langle Prototype for Left\_Diagonal\_Multiply\ 225 \rangle {

int i,\ j;

for (i=1;\ i\leq n;\ i++)

for (j=1;\ j\leq n;\ j++)\ C[i][j]=A[i]*B[i][j];
}
```

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227. Matrix\_Sum adds the two matrices A and B, puts the result in C The matrices need not be distinct  $\langle \text{ Prototype for } Matrix\_Sum \ \underline{227} \rangle \equiv$ 

```
void Matrix\_Sum(int n, double **A, double **B, double **C)
This code is used in sections 213 and 228.
        \langle \text{ Definition for } Matrix\_Sum \ 228 \rangle \equiv
   \langle \text{ Prototype for } Matrix\_Sum \ 227 \rangle
```

for  $(j = 1; j \le n; j ++)$  C[i][j] = A[i][j] + B[i][j];

This code is used in section 212.

for  $(i = 1; i \le n; i++)$ 

int i, j;

229. Matrix Multiplication. This is the crux of this whole unit at present. Most of the time in the adding-doubling algorithm is spent doing matrix multiplication and this implementation has been optimized using pointers.

Matrix Multiply multiplies the two matrices A and B and puts the result in C. The following routine requires that C does not occupy the same space as B, but it can be coincident with A. There is no inherent reason that A, B, and C must all be  $n \times n$  matrices. However, all the matrices in the adding-doubling method are square and I did not want to pass three separate dimensions to this routine.

The usual way matrix multiplication uses an algorithm something similar to:

```
\langle \text{ unused fragment one } 229 \rangle \equiv
  for (i = 1; i \le n; i++) {
     for (j = 1; j \le n; j ++) {
        C[i][j] = 0.0;
        for (k = 1; k \le n; k++) C[i][j] += A[i][k] * B[k][j];
  }
```

This has the unfortunate problem that the innermost loop indexes successive columns of A and successive rows of B. Because indexing successive rows requires something other than a unit increment of the matrix pointer, a different algorithm is used. In this case,

```
\langle \text{ unused fragment two } 230 \rangle \equiv
   for (i = 1; i \le n; i++)
      for (j = 1; j \le n; j ++) C[i][j] = 0.0;
   for (i = 1; i \le n; i++) {
      for (k = 1; k \le n; k++) {
         {\bf for} \ (j=1; \ j \leq n; \ j + +) \ C[i][j] \ + = A[i][k] * B[k][j]; 
   }
```

Explictly using pointers to the entries in the salient matrices makes this routine roughly 20% faster than when the above implementation is used. Profiling of the code indicates that roughly 45% of the time spent in an adding-doubling calculation is spent in this one routine. Therefore even a modest 20% increase will translate to a ten percent improvement in performance.

Finally, the algorithm can be improved to allow the pointers to A and C to be the same. This is sufficient to allow us to avoid allocating an extra matrix here and there. It can easily be adapted to work with "star" multiplication by premultiplying using  $Right\_Diagonal\_Multiply$ . The drawbacks are that a vector D must be allocated on each call. It is also necessary to copy the data from the vector D to the output matrix C.

```
232.
         \langle \text{Prototype for } Matrix\_Multiply \ 232 \rangle \equiv
   void Matrix\_Multiply(int n, double **A, double **B, double **C)
This code is used in sections 213 and 233.
        \langle \text{ Definition for } Matrix\_Multiply 233 \rangle \equiv
   ⟨ Prototype for Matrix_Multiply 232⟩
      (Local variables for Matrix_Multiply 234)
      (Do awkward cases 235)
      (Allocate memory for D 236)
      \langle Initialization for Matrix_Multiply 237 \rangle
      \langle Multiplying A \text{ and } B \text{ 240} \rangle
      \langle Free memory for D 241\rangle
This code is used in section 212.
234. \langle \text{Local variables for } Matrix\_Multiply 234 \rangle \equiv
   double *a\_ptr, *a\_start;
   double *b\_start, *b\_last;
   double *c\_start, *c\_very\_last, *c\_ptr;
   double *D;
   double *d\_start, *d\_last;
   register double t, *d_ptr, *b_ptr;
   ptrdiff_t row;
This code is used in section 233.
235. \langle \text{ Do awkward cases } 235 \rangle \equiv
   if (n \leq 0) {
     AD_{-}error("Non-positive_{\sqcup}dimension_{\sqcup}passed_{\sqcup}to_{\sqcup}Matrix_{\perp}Multiply");
   else if (n \equiv 1) {
     C[1][1] = A[1][1] * B[1][1];
     return;
This code is used in section 233.
```

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I need a temporary vector equal to the row length of C to hold intermediate calculations. This will allow A and C to point to the same matrix and still yield the correct results.

```
\langle Allocate memory for D 236\rangle \equiv
   D = dvector(1, n);
This code is used in section 233.
```

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237. During the initialization phase, I need to know how far it is from one row to the next row. Because of the peculiar way that Numerical Recipes allocates the matrices, this may and probably is not equal to n. The number of entries is found explicitly by subtracting a pointer to the first entry in row one from the first entry in row two. This assumes that the size of the matrix is at least two. To make this routine bulletproof, this would need to be changed—but I do not think it is really necessary.

```
\langle \text{Initialization for } Matrix\_Multiply 237 \rangle \equiv
   a\_start = \&A[1][1];
   b_{-}last = \&B[n][1];
   row = \&A[2][1] - a\_start;
   c\_very\_last = \&C[n][n];
   d\_start = \&D[1];
   d_{-}last = \&D[n];
This code is used in section 233.
```

There may be a better way of doing this, but I bet it would depend on specific knowlege about how zero is stored in the computer.

```
\langle \text{Zero } D | \mathbf{238} \rangle \equiv
   d_{-}ptr = d_{-}start;
   while (d_{-}ptr \le d_{-}last) *d_{-}ptr ++ = 0.0;
This code is used in section 240.
```

**239.** Copy the contents of D to C. This could potentially be sped up using memmove() but I just want it to work for now.

```
\langle \text{Copy } D \text{ into } C \text{ 239} \rangle \equiv
   d_{-}ptr = d_{-}start;
   c_{-}ptr = c_{-}start;
   while (d_{-}ptr \le d_{-}last) *c_{-}ptr ++ = *d_{-}ptr ++;
This code is used in section 240.
```

**240.** Here is the heart of the routine. The first row of C is filled completely, then the routine goes on to the second row and so on. The inner loop is responsible for multiplying A[i][k] (represented by  $t = *a\_ptr$ ) by every element in row i and adding it to the appropriate element in row i of C.

```
\langle Multiplying A \text{ and } B \text{ 240} \rangle \equiv
   for (c\_start = \&C[1][1]; c\_start \le c\_very\_last; c\_start += row) {
      a_{-}ptr = a_{-}start;
      \langle \text{ Zero } D | \mathbf{238} \rangle
      for (b\_start = \&B[1][1]; b\_start \le b\_last; b\_start += row) {
         t = *a_ptr ++;
         b_{-}ptr = b_{-}start;
         d_{-}ptr = d_{-}start;
         while (d_{-}ptr \leq d_{-}last) *d_{-}ptr +++ = t *(*b_{-}ptr +++);
      \langle \text{ Copy } D \text{ into } C \text{ 239} \rangle
      a\_start += row;
This code is used in section 233.
241. Dump the memory that was allocated.
\langle Free memory for D 241\rangle \equiv
   free\_dvector(D, 1, n);
This code is used in section 233.
242.
         Matrix Decomposition.
243.
          \langle \text{ Prototype for } Decomp \ 243 \rangle \equiv
   void Decomp(\mathbf{int}\ n, \mathbf{double}\ **A, \mathbf{double}\ *condition, \mathbf{int}\ *ipvt)
This code is used in sections 213 and 244.
```

**244.** Decomp decomposes a double matrix by Gaussian elimination and estimates the condition of the matrix.

Use solve to compute solutions to linear systems

On input n is the order of the matrix and A is the matrix to be triangularized.

On output A contains an upper triangular matrix U and a permuted version of a lower triangular matrix I - L so that (permutation matrix)\*A=L\*U. condition is an estimate of the condition of A. For the linear system AX = B, changes in A and B may cause changes condition times as large in X. If condition+1.0 = condition, A is singular to working precision. condition is set to  $1.0 \cdot 10^{32}$  if exact singularity is detected. ipvt is the pivot vector ipvt(k) is the index of the kth pivot row  $ipvt(n) = (-1)^{(number of interchanges)}$ 

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This code is used in section 247.

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**245.** This should probably be fixed to compute the inverse of a non-zero 1by 1 matrix.  $\langle \text{ Do } n \equiv 1 \text{ case } 245 \rangle \equiv$ 

```
ipvt[n] = 1;
  if (n \equiv 1) {
     if (A[1][1] \equiv 0) {
        AD_{-error}("1_{\sqcup}X_{\sqcup}1_{\sqcup}Matrix_{\sqcup}is_{\sqcup}Singular_{\sqcup}---_{\sqcup}i.e._{\sqcup}zero");
        return;
     }
  }
This code is used in section 244.
246. \langle Compute 1-norm of A 246\rangle \equiv
  anorm = 0.0;
  for (j = 1; j \le n; j ++) {
     t = 0.0:
     for (i = 1; i \le n; i++) t += fabs(A[i][j]);
     if (t > anorm) anorm = t;
This code is used in section 244.
247. \langle Gaussian elimination with partial pivoting 247 \rangle \equiv
  for (k = 1; k < n; k ++) {
     (Find pivot 248)
     (Compute multipliers 249)
     (Interchange and eliminate by columns 250)
This code is used in section 244.
248. \langle \text{ Find pivot } 248 \rangle \equiv
  m=k:
  for (i = k + 1; i \le n; i++)
     if (fabs(A[i][k]) > fabs(A[m][k])) m = i;
  ipvt[k] = m;
  if (m \neq k) ipvt[n] *= -1;
  t = A[m][k];
  A[m][k] = A[k][k];
  A[k][k] = t; /* skip step if pivot is zero */
  if (t \equiv 0) continue;
This code is used in section 247.
249. \langle Compute multipliers 249 \rangle \equiv
  for (i = k + 1; i \le n; i++) A[i][k] /= -t;
```

```
250.
        \langle Interchange and eliminate by columns 250 \rangle \equiv
  for (j = k + 1; j \le n; j++) {
     t = A[m][j];
     A[m][j] = A[k][j];
     A[k][j] = t;
     if (t \equiv 0) continue;
      {\bf for}\ (i=k+1;\ i\le n;\ i+\!\!\!+)\ A[i][j]\ +=A[i][k]*t;
This code is used in section 247.
251. \langle Check for singularity 251 \rangle \equiv
  *condition = 1.0;
  for (k = 1; k \le n; k ++) {
     if (A[k][k] \equiv 0.0) {
        *condition = 1 \cdot 10^{32};
        return;
  }
This code is used in section 244.
```

252. Solving systems of equations.

```
253.
```

```
\langle Prototype for Solve\ 253 \rangle \equiv void Solve\ (int\ n, double\ **A, double\ *B, int\ *ipvt) This code is used in sections 213 and 254.
```

**254.** This procedure finds the solution of the linear system AX = B Don't use if Decomp has found a singularity

On input n is the order of matrix, A is the triangularized matrix obtained form Decomp. B is the right hand side vector and ipvt is the pivot vector obtained from Decomp

On output B is the solution vector X.

```
⟨ Definition for Solve 254⟩ ≡
⟨ Prototype for Solve 253⟩
{
    int i, k, m;
    double t;
⟨ Forward elimination 255⟩
⟨ Back substitution 256⟩
}

This code is used in section 212.

255. ⟨ Forward elimination 255⟩ ≡

for (k = 1; k < n; k++) {
    m = ipvt[k];
    t = B[m];
    B[m] = B[k];
    B[m] = B[k];
    B[k] = t;
    for (i = k + 1; i \le n; i++) B[i] += A[i][k] * t;
}
```

This code is used in sections 213 and 260.

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```
256. \langle \text{ Back substitution } 256 \rangle \equiv
  for (k = n; k > 1; k --) {
     B[k] /= A[k][k];
     t = -B[k];
     for (i = 1; i < k; i++) B[i] += A[i][k] * t;
  B[1] /= A[1][1];
This code is used in section 254.
257. Finds the inverse of the matrix A (of order n) and stores the answer in Ainv.
\langle Prototype for Matrix_Inverse \ 257 \rangle \equiv
  void Matrix\_Inverse(int n, double **A, double **Ainv)
This code is used in sections 213 and 258.
258. \langle Definition for Matrix\_Inverse 258\rangle \equiv
  ⟨ Prototype for Matrix_Inverse 257⟩
     int *ipvt;
     int i, j;
     double *work:
     double condition;
     ipvt = ivector(1, n);
     Decomp(n, A, \& condition, ipvt);
     if (condition \equiv (condition + 1) \lor condition \equiv 1 \cdot 10^{32}) {
       free\_ivector(ipvt, 1, n);
       AD\_error("Singular \sqcup Matrix \sqcup \ldots \sqcup failed \sqcup in \sqcup Inverse\_Multiply \setminus n");
     work = dvector(1, n);
     for (i = 1; i \le n; i ++) {
       for (j = 1; j \le n; j++) work[j] = 0.0;
        work[i] = 1.0;
       Solve(n, A, work, ipvt);
       for (j = 1; j \le n; j++) Ainv[j][i] = work[j];
     free\_dvector(work, 1, n);
     free\_ivector(ipvt, 1, n);
This code is used in section 212.
       \langle Prototype for Left_Inverse\_Multiply 259 \rangle \equiv
  void Left\_Inverse\_Multiply (int n, double **D, double **C, double **A)
```

**260.** Left\_Inverse\_Multiply computes  $\mathbf{A} = \mathbf{C} \cdot \mathbf{D}^{-1}$  where A, C and D are all n by n matrices. This is faster than inverting and then multiplying by a factor of six. Space for A should be allocated before calling this routine.

```
\langle \text{ Definition for } Left\_Inverse\_Multiply 260 \rangle \equiv
  ⟨ Prototype for Left_Inverse_Multiply 259⟩
     int *ipvt;
     int i, j;
     double *work;
     double condition:
     Transpose\_Matrix(n, D);
     ipvt = ivector(1, n);
                                              /* Check for singular result */
     Decomp(n, D, \&condition, ipvt);
     if (condition \equiv (condition + 1) \lor condition \equiv 1 \cdot 10^{32}) {
       free\_ivector(ipvt, 1, n);
       AD_{-error}("Singular_{\square}Matrix_{\square}..._{\square}failed_{\square}in_{\square}Left_{Inverse\_Multiply}n");
     work = dvector(1, n);
                                      /* Cycle through all the row in C */
     for (i = 1; i < n; i++) {
       for (j = 1; j \le n; j ++)
                                       /* put a row of C into work */
                                 /* and avoid a Transpose Matrix */
          work[j] = C[i][j];
       Solve(n, D, work, ipvt);
                                        /* Again avoiding a Transpose Matrix */
       for (j = 1; j \le n; j ++)
          A[i][j] = work[j];
                                  /* stuff the results into a row of A */
     free\_dvector(work, 1, n);
     free\_ivector(ipvt, 1, n);
This code is used in section 212.
261. \langle Prototype for Right_Inverse_Multiply 261 \rangle \equiv
  void Right\_Inverse\_Multiply (int n, double **D, double **C, double **A)
This code is used in sections 213 and 262.
```

Right\_Inverse\_Multiply computes  $\mathbf{A} = \mathbf{D}^{-1} \cdot \mathbf{C}$  where A, C and D are all n by n matrices. This is faster than inverting and then multiplying by a factor of six. Space for A should be allocated before calling this routine.

```
\langle \text{ Definition for } Right\_Inverse\_Multiply 262 \rangle \equiv
  ⟨ Prototype for Right_Inverse_Multiply 261⟩
     int *ipvt;
     int i, j;
     double *work;
     double condition:
     ipvt = ivector(1, n);
                                            /* Check for singular result */
     Decomp(n, D, \& condition, ipvt);
     if (condition \equiv (condition + 1) \lor condition \equiv 1 \cdot 10^{32}) {
       free\_ivector(ipvt, 1, n);
       AD_error("Singular_Matrix_..._failed_in_Right_Inverse_Multiply\n");
     work = dvector(1, n);
                                     /* Cycle through all the rows */
     for (i = 1; i \le n; i++) {
       for (j = 1; j \le n; j++)
                                      /* put a column of C into work */
          work[j] = C[j][i];
       Solve(n, D, work, ipvt);
       for (j = 1; j \le n; j ++)
                                      /* stuff the results into a column of A */
         A[j][i] = work[j];
     free\_dvector(work, 1, n);
     free\_ivector(ipvt, 1, n);
This code is used in section 212.
```

### 263. AD Radau Quadrature.

This global variable is needed because the degree of the Legendre Polynomial must be known. The routine Radau stores the correct value in this.

```
#define NSLICES 512
#define EPS 1 \cdot 10^{-16}
\langle ad_radau.c 263 \rangle \equiv
   ⟨ Preprocessor definitions ⟩
   # include "ad_globl.h" # include "ad_radau.h" # include "nr_rtsaf.h" # include "nr_util.h"
   {\bf include} \ "nr\_zbrak.h" \ {\bf static} \ {\bf int} \ {\it local\_n\_size};
      \langle Prototype for Pn\_and\_Pnm1 \ 274 \rangle;
   \langle \text{ Prototype for } Pnd | 276 \rangle;
   \langle \text{ Prototype for } phi | 282 \rangle;
   ⟨ Prototype for phi_and_phiprime 278⟩;
   \langle \text{ Definition for } Pn\_and\_Pnm1 275 \rangle
   \langle \text{ Definition for } Pnd 277 \rangle
   \langle \text{ Definition for } phi | 283 \rangle
   \langle \text{ Definition for } phi\_and\_phiprime 279 \rangle
   ⟨ Definition for Radau 267⟩
264. \langle ad_radau.h 264 \rangle \equiv
   \langle \text{ Prototype for } Radau \text{ 266} \rangle;
```

#### 265. Introduction.

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The adding-doubling method is based on numerical integration of functions using quadrature,

$$\int_0^1 f(\nu, \nu') \, d\nu' = \sum_{k=1}^N w_k f(x_k)$$

The values of the quadrature points  $x_k$  and the weights  $w_k$  are chosen in such a way that the integral is evaluated exactly for a polynomial of order 2N-1 (or possibly 2N-2 depending on the quadrature method). Using N quadrature points (Gaussian) is equivalent to the spherical harmonic method of order  $P_{N-1}$ , i.e. four quadrature points corresponds to the  $P_3$  method. The specific choice of quadrature methods for samples with mismatched boundaries is described in the next section.

Total internal reflection causes problems by changing the effective range of integration. Usually, adding-doubling integrals range from 0 to 1, since the angle varies from  $\frac{\pi}{2}$  to 0 and therefore the cosine varies from 0 to 1. The integrations are calculated using numerical quadrature, and the quadrature angles are optimized for this range. If the cosine of the critical angle is denoted by  $\nu_c$  for a boundary layer with total internal reflection, then the effective range of integration is reduced to  $\nu_c$  to 1 (because the rest of the integration range is now zero). To maintain integration accuracy, the integral is broken into two parts and each is evaluated by quadrature over the specified subrange,

$$\int_0^1 A(\nu, \nu') B(\nu', \nu'') d\nu' = \int_0^{\nu_c} A(\nu, \nu') B(\nu', \nu'') d\nu' + \int_{\nu_c}^1 A(\nu, \nu') B(\nu', \nu'') d\nu'.$$

Here  $A(\nu, \nu')$  and  $B(\nu, \nu')$  represent reflection or transmission functions, and clearly if either is identically zero for values of  $\nu$  less than  $\nu_c$ , the integration range is reduced. The calculations in this paper used Gaussian quadrature for the range from 0 to  $\nu_c$ , thereby avoiding calculations at both endpoints (in particular, the angle  $\nu = 0$  is avoided, which may cause division by zero). Radau quadrature is used for the range from  $\nu_c$  to 1, so  $\nu = 1$  could be specified as a quadrature point. Each part of the integration range gets half of the quadrature points; when no critical angle exists, Radau quadrature is used over the entire range.

Radau quadrature requires finding the n roots of the following equation

$$P_{n-1}(x_i) + \frac{x_i - 1}{n} P'_{n-1}(x_i) = 0$$

Here  $P_n(x)$  is the *n*th Legendre polynomial of order zero and  $P'_{n-1}(x_i)$  is the first derivative of the n-1 Legendre polynomial. These roots are the required quadrature points for the integration range -1 to 1. The *n*th integration angle  $\nu_n$  corresponds with  $x_n = -1$  (normal incidence).

**266.** Radau. Radau calculates the n quadrature points  $x_i$  and weights  $w_i$ .

 $\langle \text{ Prototype for } Radau \text{ 266} \rangle \equiv$ 

void Radau(double x1, double x2, double \*x, double \*w, int n)

This code is used in sections 264 and 267.

```
267. \langle Definition for Radau\ 267 \rangle \equiv \langle Prototype for Radau\ 266 \rangle {
 x[n] = -1.0;
 w[n] = 2.0/(n*n);
 switch\ (n)\ \{
 case\ 2:\ \langle \text{Values for } n \equiv 2\ 285 \rangle
 case\ 4:\ \langle \text{Values for } n \equiv 4\ 286 \rangle
 case\ 8:\ \langle \text{Values for } n \equiv 8\ 287 \rangle
 case\ 16:\ \langle \text{Values for } n \equiv 16\ 288 \rangle
 default:\ \langle \text{Values for arbitrary } n\ 269 \rangle
 \}
 \langle \text{Scale values } 268 \rangle
 \}
This code is used in section 263.
```

**268.** The code to scale values is easy. Radau quadrature is defined over the range -1 to 1. Here we just linearly scale the width of each interval and weight as appropriate. To modify for the range  $\nu_c$  to 1 the following relations are needed to find the necessary integration angles  $\nu_i$  and weights  $w_i$ 

$$\nu_i = \frac{1 + \nu_c - (1 - \nu_c)x_i}{2}$$

and

$$w_i = \frac{1 - \nu_c}{(1 - x_i)\sqrt{P'_{n-1}(x_i)}}$$

```
 \langle \, \text{Scale values} \, \, 268 \, \rangle \equiv \\ \{ \\ \mathbf{double} \, \, xm, \, \, xl; \\ \mathbf{int} \, \, i; \\ xm = (x2 + x1) * 0.5; \\ xl = (x2 - x1) * 0.5; \\ \mathbf{for} \, \, (i = 1; \, \, i \leq n; \, \, i + +) \, \, \{ \\ x[i] = xm - xl * x[i]; \\ w[i] = xl * w[i]; \\ \} \\ \}
```

```
269. Here is the method for finding Radau quadrature points for non-tabulated values.
\langle \text{ Values for arbitrary } n \text{ 269} \rangle \equiv
     int i, nb, ndiv;
     double z;
    double *xb1, *xb2;
     (Allocate memory for Radau 270)
     (Bracket roots 271)
     (Find roots and weights 272)
     ⟨ Free memory for Radau 273 ⟩
     break;
This code is used in section 267.
270. \langle Allocate memory for Radau 270 \rangle \equiv
  xb1 = dvector(1, NSLICES);
  xb2 = dvector(1, NSLICES);
This code is used in section 269.
271. Bracket n-1 roots, double ndiv if not enough roots are found.
\langle \text{ Bracket roots } 271 \rangle \equiv
  local\_n\_size = n;
  if (2 * n > NSLICES) ndiv = NSLICES;
  else ndiv = 2 * n;
  do {
     nb = n - 1;
     zbrak(phi, -1.0, 1.0, ndiv, xb1, xb2, \&nb);
    ndiv *= 2;
  } while (nb < n - 1 \land ndiv \leq \texttt{NSLICES});
  if (nb < n-1) AD\_error("Cannot find enough roots for Radau quadrature");
This code is used in section 269.
272. Find the roots with an accuracy EPS and store them in the array x. Put them in backwards so that
x[n] = -1 is in the correct spot.
\langle Find roots and weights 272 \rangle \equiv
  for (i = 1; i < n; i++) {
     double tmp;
     z = rtsafe(phi\_and\_phiprime, xb1[i], xb2[i], EPS);
     x[n-i]=z;
     tmp = Pnd(n-1,z);
     w[n-i] = 1/((1-z)*tmp*tmp);
This code is used in section 269.
273. \langle Free memory for Radau 273 \rangle \equiv
  free\_dvector(xb1, 1, NSLICES);
  free\_dvector(xb2, 1, NSLICES);
This code is used in section 269.
```

**276.** To calculate the weights for the quadrature points we need to evaluate the first derivative of the Legendre polynomial. To do this we use a recurrence relation given by H. H. Michels, in "Abscissas and weigh coefficients for Lobatto quadrature," *Math Comp*, **17**, 237-244 (1963).

```
\langle Prototype for Pnd\ 276 \rangle \equiv static double Pnd (int n, double x) This code is used in sections 263 and 277.
```

```
277. \langle \text{ Definition for } Pnd | 277 \rangle \equiv
  \langle Prototype for Pnd 276\rangle
     \mathbf{double}\ p,\ pminus,\ pplus;
     int i;
    if (x > 1.0) {
       x = 1;
     else if (x < -1.0) {
       x = -1;
     pminus = 0;
     p = 1;
     if (n \le 0) return pminus;
     for (i = 1; i < n; i ++) {
       pplus = ((2 * i + 1) * x * p - (i + 1) * pminus)/i;
       pminus = p;
       p = pplus;
     return p;
```

$$\phi_{n-1}(x) = \frac{P_{n-1}(x) + P_n(x)}{1+x}$$

we need to find the derivative. This is

$$\phi'_{n-1}(x) = \frac{P'_{n-1}(x) + P'_n(x)}{1+x} - \frac{P_{n-1}(x) + P_n(x)}{(1+x)^2}$$

Now we can use our recurrence relation

$$(1 - x^2)P'_{n-1}(x) = nxP_{n-1}(x) - nP_n(x)$$

To eliminate the derivative terms in the above equation to get

$$\phi'_{n-1} = \frac{(nx+x-1)P_{n-1}(x) + (nx+2x-n-1)P_n(x) - (n+1)P_{n+1}(x)}{(1-x)(1+x)^2}$$

The higher order Legendre Polynomial can be eliminated using

$$(n+1)P_{n+1}(x) = (2n+1)xP_n(x) - nP_{n-1}(x)$$

to get

$$\phi'_{n-1}(x) = \frac{(nx+x+n-1)P_{n-1}(x) + (-nx+x-n-1)P_n(x)}{(1-x)(1+x)^2}$$

And therefore we just call the routine that will return  $P_n(x)$  and  $P_{n-1}(x)$  and multiply by the appropriate factors to obtain both terms.

The only problem is when x = 1 or x = -1. Then we get this spurious division by zero. So we special case these and evaluate them elsewhere.

 $\langle \text{ Prototype for } phi\_and\_phiprime \ 278 \rangle \equiv$ 

static void  $phi\_and\_phiprime$  (double x, double \*phi, double \*phiprime)

This code is used in sections 263 and 279.

```
279. \langle \text{Definition for } phi\_and\_phiprime \ 279 \rangle \equiv \langle \text{Prototype for } phi\_and\_phiprime \ 278 \rangle

{
    double Pn, Pnm1;
    int n;

n = local\_n\_size;
    if (x \ge 1.0) {
    \langle \text{Phi and phiprime at } x = 1 \ 280 \rangle
}

else if (x \le -1.0) {
    \langle \text{Phi and phiprime at } x = -1 \ 281 \rangle
}

else {
    Pn\_and\_Pnm1(n, x, \&Pnm1, \&Pn);
    *phi = (Pn + Pnm1)/(1 + x);
    *phiprime = ((n * x - 1 + x + n) * Pnm1 + (-n * x + x - n - 1) * Pn)/(1 + x)/(1 - x);
}
```

**280.** To find  $\phi(1)$  and  $\phi'(1)$  we need to recall a few facts about Legendre polynomials. First,

$$P_n(1) = 1$$

Therefore

$$\phi(1) = 1$$

The value of the first derivative is somewhat trickier. Recall that the Legendre polynomials are solutions to

$$(1 - x^2)P_n''(x) - 2xP_n'(x) + n(n+1)P_n(x) = 0$$

Now if x = 1 then the first term on the left hand side will be zero. Therefore

$$P_n'(1) = \frac{n(n+1)}{2}$$

Therefore

$$\phi'_{n-1}(1) = \frac{n^2 - 1}{2}$$

```
\langle Phi and phiprime at x=1 280 \rangle \equiv { *phi=1; \\ *phiprime=(n*n-1)/2; \\ }
```

This code is used in section 279.

**281.** To evaluate  $\phi(-1)$  we must return to the original definition, i.e. So

$$\phi_{n-1}(x) = P_{n-1}(x) + \frac{x-1}{n}P'_{n-1}(x)$$

To evaluate this we need to remember some stuff, namely that

$$P_n(-x) = (-1)^n P_n(x)$$
 so  $P_n(-1) = (-1)^n$ 

The value of the first derivative is again obtained from the differential equation and

$$P'_n(-1) = -\frac{n(n+1)}{2}P_n(-1) = (-1)^{n+1}\frac{n(n+1)}{2}$$

Now we just substitute to get

$$\phi_{n-1}(-1) = (-1)^{n-1} \cdot n$$

The first derivative is more difficult. Mathematica says that it is

$$\phi'_{n-1}(-1) = (-1)^n \frac{n(1-n^2)}{4}$$

**282.** For Radau quadrature, we want to find the n-1 roots of

$$\phi_{n-1}(x) = P_{n-1}(x) + \frac{x-1}{n}P'_{n-1}(x)$$

F. B. Hildebrand notes that by using a recurrence formula this becomes

$$\phi_{n-1}(x) = \frac{P_{n-1}(x) + P_n(x)}{1+x}$$

This is particularly convenient, because we must find  $P_{n-1}(x)$  before we can find  $P_n(x)$  and this is exactly what  $P_{n-1}(x)$  does.

It is noteworthy that this routine uses the recurrence formula

$$P_{n+1}(x) = \frac{(2n+1)xP_n(x) - nP_{n-1}(x)}{n+1}$$

to calculate the Legendre polynomial  $P_n(x)$ . This recurrence relation is given in H. H. Michels, "Abscissas and weight coefficients for Lobatto quadrature," *Math Comp*, **17**, 237-244 (1963).

```
\langle \text{ Prototype for } phi \text{ 282 } \rangle \equiv
static double phi(\text{double } x)
```

This code is used in sections 263 and 283.

```
283. \langle \text{Definition for } phi | 283 \rangle \equiv \langle \text{Prototype for } phi | 282 \rangle
{

double Pn, Pnm1;

if (x \leq -1.0) {

if (local\_n\_size \% 2 \neq 1) return (-local\_n\_size);

else return (local\_n\_size);
}

Pn\_and\_Pnm1 (local\_n\_size, x, \&Pnm1, \&Pn);
return ((Pn + Pnm1)/(1 + x));
}

This code is used in section 263.
```

# 284. Radau Tables.

Here is a selection of commonly used number of quadrature points.

```
285. \langle Values for n \equiv 2 285\rangle \equiv x[1] = 0.333333333333333333; <math>w[1] = 1.500000000000000000; break;
```

```
286. \langle \text{Values for } n \equiv 4 \text{ 286} \rangle \equiv x[3] = -0.5753189235216942; x[2] = 0.1810662711185306; x[1] = 0.8228240809745921; w[3] = 0.6576886399601182; w[2] = 0.7763869376863437; w[1] = 0.4409244223535367; break;
```

This code is used in section 267.

```
287. \langle \text{ Values for } n \equiv 8 \ 287 \rangle \equiv
  x[7] = -0.8874748789261557;
  x[6] = -0.6395186165262152;
  x[5] = -0.2947505657736607;
  x[4] = 0.0943072526611108;
  x[3] = 0.4684203544308211;
  x[2] = 0.7706418936781916;
  x[1] = 0.9550412271225750;
  w[7] = 0.1853581548029793;
  w[6] = 0.3041306206467856;
  w[5] = 0.3765175453891186;
  w[4] = 0.3915721674524935;
  w[3] = 0.3470147956345014;
  w[2] = 0.2496479013298649;
  w[1] = 0.1145088147442572;
  break;
```

```
288.
       \langle \text{ Values for } n \equiv 16 \text{ 288} \rangle \equiv
  x[15] = -0.9714610905263484;
  x[14] = -0.9054008198116666;
  x[13] = -0.8045734013587561;
  x[12] = -0.6728619212112202;
  x[11] = -0.5153294780626855;
  x[10] = -0.3380303900599197;
  x[9] = -0.1477783218133717;
  x[8] = 0.0481153830735303;
  x[7] = 0.2421226227060438;
  x[6] = 0.4267878274849459;
  x[5] = 0.5950144898997919;
  x[4] = 0.7403379488928179;
  x[3] = 0.8571740937696823;
  x[2] = 0.9410354027041150;
  x[1] = 0.9887186220549766;
  w[15] = 0.0477022269476863;
  w[14] = 0.0839852814449645;
  w[13] = 0.1170203531038591;
  w[12] = 0.1455555452202026;
  w[11] = 0.1684963978499219;
  w[10] = 0.1849617814886653;
  w[10] = 0.1849617814886653;
  w[9] = 0.1943190897115679;
  w[8] = 0.1962087882390318;
  w[7] = 0.1905582942553547;
  w[6] = 0.1775847927527395;
  w[5] = 0.1577869218042020;
  w[4] = 0.1319256999330681;
  w[3] = 0.1009956796217840;
  w[2] = 0.0661895086101364;
  w[1] = 0.0288971390168143;
  break;
```

**289. AD Phase Function.** This section contains all the routines associated with generating the necessary matrices for Henyey-Greenstein phase functions. This is the place to put code to implement other phase functions.

```
⟨ad_phase.c 289⟩ ≡
#include <stdlib.h>
#include <math.h>
#include "nr_util.h"
#include "ad_globl.h"
#include "ad_phase.h"
⟨Definition for Get_Phi 295⟩
290. ⟨ad_phase.h 290⟩ ≡
⟨Prototype for Get_Phi 294⟩;
```

**291.** Redistribution function. The single scattering phase function  $p(\nu)$  for a tissue determines the amount of light scattered at an angle  $\nu = \cos \theta$  from the direction of incidence. The subtended angle  $\nu$  is the dot product of the unit vectors  $\hat{\mathbf{s}}_i$  and  $\hat{\mathbf{s}}_j$ 

$$\nu = \hat{\mathbf{s}}_i \cdot \hat{\mathbf{s}}_j = \nu_i \nu_j + \sqrt{1 - \nu_i^2} \sqrt{1 - \nu_j^2} \cos \phi$$

where  $\hat{\mathbf{s}}_i$  is the incident and  $\hat{\mathbf{s}}_j$  is the scattered light directions

The redistribution function  $\mathbf{h}_{ij}$  determines the fraction of light scattered from an incidence cone with angle  $\nu_i$  into a cone with angle  $\nu_j$ . The redistribution function is calculated by averaging the phase function over all possible azimuthal angles for fixed angles  $\nu_i$  and  $\nu_j$ ,

$$h(\nu_i, \nu_j) = \frac{1}{2\pi} \int_0^{2\pi} p(\nu_i \nu_j + \sqrt{1 - \nu_i^2} \sqrt{1 - \nu_j^2} \cos \phi) \, d\phi$$

Note that the angles  $\nu_i$  and  $\nu_j$  may also be negative (light travelling in the opposite direction). The full redistribution matrix may be expressed in terms a  $2 \times 2$  matrix of  $n \times n$  matrices

$$\mathbf{h} = \begin{bmatrix} \mathbf{h}^{--} & \mathbf{h}^{-+} \\ \mathbf{h}^{+-} & \mathbf{h}^{++} \end{bmatrix}$$

The first plus or minus sign indicates the sign in front of the incident angle and the second is the sign of the direction of the scattered light.

When the cosine of the angle of incidence or exitance is unity ( $\nu_i = 1$  or  $\nu_j = 1$ ), then the redistribution function  $h(1, \nu_j)$  is equivalent to the phase function  $p(\nu_j)$ . In the case of isotropic scattering, the redistribution function is a constant

$$h(\nu_i, \nu_j) = p(\nu) = \frac{1}{4\pi}.$$

For Henyey-Greenstein scattering, the redistribution function can be expressed in terms of the complete elliptic integral of the second kind E(x)

$$h(\nu_i, \nu_j) = \frac{2}{\pi} \frac{1 - g^2}{(\alpha - \gamma)\sqrt{\alpha + \gamma}} E\left(\sqrt{\frac{2\gamma}{\alpha + \gamma}}\right)$$

where g is the average cosine of the Henyey-Greenstein phase function and

$$\alpha = 1 + g^2 - 2g\nu_i\nu_j$$
 and  $\gamma = 2g\sqrt{1 - \nu_i^2}\sqrt{1 - \nu_j^2}$ 

The function E(x) may be calculated using algorithms found in Press *et al.* This method of calculating the phase function is slower than the method that is used in this program.

Other phase functions require numerical integration of the phase function. If the phase function is highly anisotropic, then the integration over the azimuthal angle is particularly difficult and care must be taken to ensure that the integration is accurate. This is important because errors in the redistribution function enter directly into the reflection and transmission matrices for thin layers. Any errors will be doubled with each successive addition of layers and small errors will rapidly increase.

An alternate way to calculate the redistribution function is the  $\delta$ -M method of Wiscombe. This method works especially well for highly anisotropic phase functions. The number of quadrature points is specified by M. The  $\delta$ -M method approximates the true phase function by a phase function consisting of a Dirac delta function and M-1 Legendre polynomials

$$p^*(\nu) = 2g^M \delta(1 - \nu) + (1 - g^M) \sum_{k=0}^{M-1} (2k+1) \chi_k^* P_k(\nu)$$

where

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$$\chi_k^* = \frac{\chi_k - g^M}{1 - g^M}$$
 and  $\chi_k = \frac{1}{2} \int_0^1 p(\nu) P_k(\nu) d\nu$ 

When the  $\delta$ -M method substitutes  $p^*(\nu) \to p(\nu)$ , then both the albedo and optical thickness must also be changed,  $a^* \to a$  and  $\tau^* \to \tau$ . This approximation is analogous to the similarity transformation often used to improve the diffusion approximation by moving a part  $(q^M)$  of the scattered light into the unscattered component. The new optical thickness and albedo are

$$\tau^* = (1 - ag^M)\tau$$
 and  $a^* = a\frac{1 - g^M}{1 - ag^M}$ 

This is equivalent transforming the scattering coefficient as  $\mu_s^* = \mu_s(1-g^M)$ . The redistribution function can now be written as

$$h^*(\nu_i, \nu_j) = \sum_{k=0}^{M-1} (2k+1)\chi_k^* P_k(\nu_i) P_k(\nu_j)$$

For the special case of a Henvey-Greenstein phase function,

 $\langle \text{Prototype for } Get\_Phi \ 294 \rangle \equiv$ 

This code is used in section 289.

$$\chi_k^* = \frac{g^k - g^M}{1 - g^M}.$$

Calculate the renormalization matrix for a Henyey-Greenstein phase function using the delta-M method. This version has been optimized for isotropic and Henyey-Greenstein phase functions.

```
void Get_Phi(int n, int phase_function, double g, double **h)
This code is used in sections 290 and 295.
        \langle \text{ Definition for } Get\_Phi \ 295 \rangle \equiv
   \langle \text{ Prototype for } Get\_Phi \ 294 \rangle
     (Local variables for Get_Phi 296)
      Test for bad calling parameters 297
     (Initialize the phase function matrix 298)
     We're done if phase function is isotropic 299
      Calculate the quadrature coefficients 300 \
      Create Legendre Polynomial matrix 301
      Calculate the coefficients 305
      Add the symmetric part of the matrix 306
     Free p and chi 307
```

```
296.
       \langle \text{Local variables for } Get\_Phi \ 296 \rangle \equiv
  int i, j, k;
  double g2M, gk, x;
  double *chi;
  double **p;
This code is used in section 295.
297. \langle Test for bad calling parameters 297 \rangle \equiv
  if (g \neq 0 \land phase\_function \neq \texttt{HENYEY\_GREENSTEIN})
     AD_{-error}("Only_{\sqcup}the_{\sqcup}Henyey-Greenstein_{\sqcup}phase_{\sqcup}function_{\sqcup}has_{\sqcup}been_{\sqcup}implemented\\");
  if (fabs(g) \ge 1) AD_{error}("Get_Phi_was_called_with_a_bad_g_calc_value");
This code is used in section 295.
298. (Initialize the phase function matrix 298) \equiv
  for (i = -n; i \le n; i++)
     for (j = -n; j \le n; j ++) h[i][j] = 1; /* zero the zero column and zero row */
  for (i = -n; i \le n; i++) {
     h[i][0] = 0.0;
     h[0][i] = 0.0;
This code is used in section 295.
299. \langle We're done if phase function is isotropic 299 \rangle \equiv
  if (g \equiv 0) return;
This code is used in section 295.
       To avoid extra calculation let's define
                                                   chi[k] \equiv (2k+1)\chi_k^*
This will slighly simplify things later on
\langle Calculate the quadrature coefficients 300\rangle \equiv
  chi = dvector(1, n);
  g2M = pow(g, n);
  qk = 1.0;
  for (k = 1; k < n; k++) {
     gk *= g;
     chi[k] = (2 * k + 1) * (gk - g2M)/(1 - g2M);
```

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Allocate the matrix for the Legendre values this is much more efficient than calculating them as they are needed. Since the Legendre polynomial  $P_n(x)$  is generated using recurrence relations, all Legendre polynomials  $P_k(x)$ , where  $0 \le k \le n$  must also be calculated. Now the formula

$$h^*(\nu_i, \nu_j) = \sum_{k=0}^{n-1} (2k+1) \chi_k^* P_k(\nu_i) P_k(\nu_j)$$

requires all those to be found as well. There are 2n+1 values that must be calculated for  $-\mu_n \dots 0 \dots \mu_n$ different arguments. A simple way is just to put all of the necessary values in a two-dimensional array and define  $p[i][j] \equiv P_i(\mu_j)$ .

```
\langle Create Legendre Polynomial matrix 301\rangle \equiv
  ⟨ Allocate the polynomial matrix 302⟩
   \langle Fill in all the unique values 303\rangle
  (Fill in the symmetric values 304)
This code is used in section 295.
```

It is not at all clear that zeroing is needed.

```
\langle Allocate the polynomial matrix 302\rangle \equiv
  p = dmatrix(0, n, -n, n);
This code is used in section 301.
```

303. Here I use the recurrence relation

$$P_{k+1}(\mu_j) = \frac{(2k+1)xP_k(\mu_j) - kP_{k-1}(\mu_j)}{k+1}$$

(which should be stable) to find all the values for all the positive angles.

```
\langle Fill in all the unique values 303\rangle \equiv
  for (j = 1; j \le n; j++) {
    p[0][j] = 1;
     x = angle[j];
    p[1][j] = x;
     for (k = 1; k < n; k++) p[k+1][j] = ((2*k+1)*x*p[k][j] - k*p[k-1][j])/(k+1);
This code is used in section 301.
```

**304.** I make use of the fact that

$$P_k(-\nu_j) = (-1)^k P_k(\nu_j)$$

to fill in all the negative angles in the phase function matrix. This eliminates half the calculation. I do two at a time. This way there does not need to be a flag. Since I know that the dimension of the matrix will be even, this should not be a problem. If the matrix is not then you have problems.

```
\langle Fill in the symmetric values 304\rangle \equiv
  for (j = 1; j \le n; j ++)
     for (k = 1; k < n; k++) {
       p[k][-j] = -p[k][j];
       p[k][-j] = p[k][j];
```

**305.** Just a straightforward calculation of

$$h^*(\nu_i, \nu_j) = \sum_{k=0}^{n-1} (2k+1)\chi_k^* P_k(\nu_i) P_k(\nu_j)$$

and since  $\chi_0^* = 1$  and  $P_0(x) = 1$  this is

$$h^*(\nu_i, \nu_j) = 1 + \sum_{k=1}^{n-1} (2k+1)\chi_k^* P_k(\nu_i) P_k(\nu_j)$$

Since h has many symmetries, there are only about  $n^2/4$  unique entries. We only need to calculate those. Oh yeah, recall that chi[k] includes the factor 2k+1 for speed.

This code is used in section 295.

**306.** Several symmetries in the redistribution matrix are used. to fill in some entries that begin with a negative angle

$$h(-\nu_i, \nu_j) = h(\nu_j, -\nu_i)$$

and secondly

$$h(-\nu_i, -\nu_i) = h(\nu_i, \nu_i)$$

Next, some entries along the diagonal are filled in using

$$h(-\nu_i, -\nu_i) = h(\nu_i, \nu_i)$$

Finally, the lower triangle is filled in using the values from the upper half using

$$h(\nu_i, \nu_i) = h(\nu_i, \nu_i)$$

This could probably be more elegant, but it hurts my brain to think about it. This works and should take advantage of all the symmetries present.

 $\langle$  Add the symmetric part of the matrix 306 $\rangle \equiv$ 

```
\begin{array}{l} \textbf{for} \ (i=n; \ i \geq 2; \ i--) \\ \textbf{for} \ (j=1; \ j < i; \ j++) \ \{ \\ h[-i][j] = h[-j][i]; \\ h[-i][-j] = h[j][i]; \\ \} \\ \textbf{for} \ (i=1; \ i \leq n; \ i++) \ h[-i][-i] = h[i][i]; \\ \textbf{for} \ (i=-n; \ i \leq n; \ i++) \\ \textbf{for} \ (j=i+1; \ j \leq n; \ j++) \ h[j][i] = h[i][j]; \end{array}
```

This code is used in section 295.

```
307. \langle \text{Free } p \text{ and } chi \text{ 307} \rangle \equiv free\_dmatrix(p, 0, n, -n, n); free\_dvector(chi, 1, n);
```

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# 308. Main Program.

Here is a quick program that I put together on the 18th of July 1996 to calculate the change in reflection and transmission when a small change in the absorption coefficient is made. Specifically, the absorption coefficient will change from  $\mu_a$  to  $\mu_a + \mu_a \Delta$ .

The program reads and input file that contains the optical properties of the slab. The output file will have the same name, but appended by ".out" and contain the change in the reflection and transmission calculated for normal irradiance using 8 quadrature points.

Note that the streams get redirected so that I can use the standard streams for reading, writing, and error messages. This makes interactive stuff problematic, but this whole thing is a batch sort of problem.

All the output for this web file goes into ad\_main.c but to simplify the Makefile, I create an empty ad\_main.h.

```
\langle ad_{main.h} 308 \rangle \equiv
```

```
309.
       The program begins here
\langle ad_main.c 309 \rangle \equiv
#include <stdio.h>
#include <string.h>
#include <stdlib.h>
#include <math.h>
#include "ad_globl.h"
#include "ad_prime.h"
#include "ad_cone.h"
#include "mygetopt.h"
#include "version.h"
  extern char *optarg;
  extern int optind;
  ⟨ print version function 317⟩
  ⟨ print usage function 318⟩
  ⟨stringdup together function 319⟩
  ⟨validate slab function 320⟩
  int main(int argc, char **argv)
    \langle Declare variables for main 310 \rangle
    if (argc \equiv 1) {
       print_usage();
       exit(0);
    \langle Handle options 314\rangle
    if (argc \ge 1) {
       \langle Prepare file for reading 315\rangle
       ⟨ Prepare file for writing 316⟩
       while (feof(stdin) \equiv 0) {
         slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};
         \langle Read line from input file 312\rangle
         (Calculate and Print the Results 313)
    else {
       (Put optical properties into slab 311)
       ⟨Calculate and Print the Results 313⟩
    return 0;
```

```
310. \langle Declare variables for main\ 310 \rangle \equiv struct AD_slab_type slab; int nstreams = 24; double anisotropy = 0; double albedo = 0.5; double index\_of\_refraction = 1.0; double index\_of\_slide = 1.0; double optical\_thickness = 100; char *g\_out\_name = \Lambda; double g\_incident\_cosine = 1; int machine\_readable\_output = 0; double R1, T1, URU, UTU; int failed; This code is used in section 309.
```

**311.** I assume that the optical properties are in the following order — albedo, optical thickness, anisotropy, the index of refraction of the slab, the index of refraction of the top slide, the index of refraction of the bottom slide. The slides are assumed to have no absorption.

```
\langle \text{Put optical properties into } slab | 311 \rangle \equiv
  slab.phase\_function = \texttt{HENYEY\_GREENSTEIN};
  slab.a = albedo;
  slab.b = optical\_thickness;
  slab.q = anisotropy;
  slab.n\_slab = index\_of\_refraction;
  slab.n\_top\_slide = index\_of\_slide;
  slab.n\_bottom\_slide = index\_of\_slide;
  slab.b_{-}top_{-}slide = 0.0;
  slab.b_-bottom\_slide = 0.0;
  slab.cos\_angle = g\_incident\_cosine;
This code is used in section 309.
312.
\langle \text{ Read line from input file } 312 \rangle \equiv
     int fileflag;
     fileflag = scanf("%lf", &slab.a);
     slab.cos\_angle = g\_incident\_cosine;
     if (fileflag \neq EOF) fileflag = scanf("%lf", &slab.b);
     if (fileflag \neq EOF) fileflag = scanf("%lf", &slab.g);
     if (fileflag \neq EOF) fileflag = scanf("%lf", &slab.n_slab);
     if (fileflag \neq EOF) fileflag = scanf("%lf", &slab.n_top_slide);
     if (fileflag \neq EOF) fileflag = scanf("%lf", &slab.n_bottom_slide);
    if (fileflag \neq EOF) fileflag = scanf("%lf", &slab.b_top_slide);
     if (fileflag \neq EOF) fileflag = scanf("%lf", &slab.b_bottom_slide);
     if (fileflag \neq EOF) fileflag = scanf("%d", &nstreams);
This code is used in section 309.
```

```
313.
          \langle Calculate and Print the Results 313\rangle \equiv
   failed = validate\_slab(slab, nstreams, machine\_readable\_output);
  R1 = failed;
  T1 = failed;
  URU = failed;
  \mathtt{UTU} = failed;
   if (\neg failed) RT(nstreams, \&slab, \&R1, \&T1, \&URU, \&UTU);
   if (machine_readable_output) printf("%9.5fu\t%9.5fu\t%9.5fu\t%9.5fu\t%9.5f\n",R1,T1,URU,UTU);
   else if (\neg failed) {
      printf("UR1<sub>□</sub>=<sub>□</sub>Total<sub>□</sub>Reflection<sub>□□□</sub>for<sub>□</sub>Normal<sub>□□</sub>Illumination\n");
      printf("UT1<sub>□</sub>=<sub>□</sub>Total<sub>□</sub>Transmission<sub>□</sub>for<sub>□</sub>Normal<sub>□□</sub>Illumination\n");
      printf("URU<sub>□</sub>=<sub>□</sub>Total<sub>□</sub>Reflection<sub>□□□</sub>for<sub>□</sub>Diffuse<sub>□</sub>Illumination\n");
      printf("UTU_{\square} = \Box Total_{\square} Transmission_{\square} for_{\square} Diffuse_{\square} Illumination \n');
      printf("$\cup$\cup$\cup$UR1$\cup$\cup$\cup$\cup$\cup$VT1$\cup$\cup$\cup$\cup$URU$\cup$\cup$\cup$\cup$\cup$VTU\n");
      printf("\%9.5f_{\perp}\t\%9.5f_{\perp}\t\%9.5f_{\perp}\t\%9.5f_{n}",R1,T1,URU,UTU);
```

```
use the mygetop to process options. We only handle help at the moment
\langle Handle options 314\rangle \equiv
    char c;
    double x;
    while ((c = my\_getopt(argc, argv, "h?vma:b:g:i:n:o:q:s:")) \neq EOF) {
      switch (c) {
      case 'i': x = strtod(optarg, \Lambda);
         if (x < 0 \lor x > 90) fprintf(stderr, "Incident_angle_must_be_between_0_and_90_degrees_n");
         else g_{incident\_cosine} = cos(x * 3.1415926535/180.0);
         break;
      case 'o': g_-out_-name = strdup(optarg);
         break;
      case 'n': index\_of\_refraction = strtod(optarg, \Lambda);
         break;
      case 's': index\_of\_slide = strtod(optarg, \Lambda);
         break;
      case 'm': machine\_readable\_output = 1;
         break;
      case 'q': nstreams = (int) strtod(optarg, \Lambda);
      case 'a': albedo = strtod(optarg, \Lambda);
         break;
       case 'b': optical\_thickness = strtod(optarg, \Lambda);
       case 'g': anisotropy = strtod(optarg, \Lambda);
         break;
       case 'v': print_version();
         break;
      default: case 'h': case '?': print_usage();
         break;
      }
    argc -= optind;
    argv += optind;
```

```
315.
       Make sure that the file is not named '-' and warn about too many files
\langle Prepare file for reading 315\rangle \equiv
  if (argc > 1) {
    fprintf(stderr, "Only_a_single_file_can_be_processed_at_a_time\n");
    fprintf(stderr, "try" 'apply ad file1 file2... fileN'\n");
  if (argc \equiv 1 \land strcmp(argv[0], "-") \neq 0) {
                                                  /* filename exists and != "-" */
    if (freopen(argv[0], "r", stdin) \equiv \Lambda) {
      fprintf(stderr, "Could_not_open_file_', s'\n", argv[0]);
    if (g\_out\_name \equiv \Lambda) g\_out\_name = strdup\_together(argv[0], ".rt");
This code is used in section 309.
316. Take care of all the output files
\langle \text{ Prepare file for writing 316} \rangle \equiv
  if (g\_out\_name \neq \Lambda) {
    if (freopen(g\_out\_name, "w", stdout) \equiv \Lambda) {
      fprintf(stderr, "Could_not_open_file_<%s>_for_output", g_out_name);
       exit(1);
This code is used in section 309.
317. \langle \text{ print version function } 317 \rangle \equiv
  static void print_version(void)
    fprintf(stderr, "ad_{\sqcup}%s\n", Version);
    fprintf(stderr, "Copyright_2020_Scott_Prahl,_scott.prahl@oit.edu\n");
    fprintf(stderr, "This_is_ifree_isoftware;_isee_ithe_isource_ifor_icopying_iconditions.\n");
    fprintf(stderr, "There\_is\_no\_warranty; \_not\_even\_for\_MERCHANTABILITY\_or\_FITNESS. \n");
    fprintf(stderr, "FOR_ A_ PARTICULAR_ PURPOSE. n");
    exit(0);
This code is used in section 309.
```

```
319. returns a new string consisting of s+t  \langle \text{stringdup together function } 319 \rangle \equiv \\ \text{static char } *strdup\_together(\text{char } *s, \text{char } *t) \\  \{ \\ \text{char } *both; \\ \text{if } (s \equiv \Lambda) \{ \\ \text{if } (t \equiv \Lambda) \text{ return } \Lambda; \\ \text{return } strdup(t); \\ \} \\ \text{if } (t \equiv \Lambda) \text{ return } strdup(s); \\ both = malloc(strlen(s) + strlen(t) + 1); \\ \text{if } (both \equiv \Lambda) \text{ } fprintf(stderr, "Could\_not\_allocate\_memory\_for\_both\_strings.\n"); \\ strcpy(both, s); \\ strcat(both, t); \\ \text{return } both; \\ \}
```

This code is used in section 309.

```
108
```

```
320.
       Make sure that the input values are correct
\langle \text{ validate slab function } 320 \rangle \equiv
  static int validate_slab(struct AD_slab_type slab, int nstreams, int machine)
    if (slab.a < 0 \lor slab.a > 1) {
       if (\neg machine) printf("Bad_Albedo_a=%f\n", slab.a);
       return (-1);
     if (slab.b < 0) {
       if (\neg machine) printf ("Bad_\Optical_\Thickness_\b=\f\n", slab.b);
       return (-2);
     if (slab.g \le -1 \lor slab.g \ge 1) {
       if (\neg machine) printf("Bad_\Anisotropy_g=%f\n", slab.g);
       return (-3);
     if (slab.n\_slab < 0 \lor slab.n\_slab > 10) {
       if (\neg machine) printf("Bad_Slab_Index_n=%f\n", slab.n_slab);
       return (-4);
    if (slab.n\_top\_slide < 1 \lor slab.n\_top\_slide > 10) {
       if (\neg machine) printf("Bad_{\sqcup}Top_{\sqcup}Slide_{\sqcup}Index_{\sqcup}n=\%f\n", slab.n_top_slide);
       return (-5);
     if (slab.n\_bottom\_slide < 1 \lor slab.n\_bottom\_slide > 10) {
       if (\neg machine) printf("Bad_Top_Slide_Index_n=%f\n", slab.n_bottom_slide);
       return (-6);
     if (slab.b\_top\_slide < 0 \lor slab.b\_top\_slide > 10) {
       if (\neg machine) printf("Bad_{\square}Top_{\square}Slide_{\square}Optical_{\square}Thickness_{\square}b=%f\n", slab.b_top_slide);
       return (-7);
     if (slab.b\_bottom\_slide < 0 \lor slab.b\_bottom\_slide > 10) {
       if (¬machine) printf("Bad_Bottom_Slide_Optical_Thickness_b=%f\n", slab.b_bottom_slide);
       return (-8);
    if (nstreams < 4 \lor nstreams \% 4 \neq 0) {
       if (\neg machine) {
         printf("Bad_Number_of_Quadrature_Points_npts=%d\n", nstreams);
         printf("Should_be_a_multiple_of_four!\n");
       return (-9);
     return 0;
This code is used in section 309.
```

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