

# Everything I think you should know about Inverse Adding-Doubling

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## Abstract

This document describes a method for measuring the optical properties of scattering and absorbing materials. The general idea is that you have made careful measurements of the total reflection and transmission of the sample. These observations are then fed into a program (the inverse adding-doubling `iad` program) to extract the intrinsic optical parameters for the sample. The program does this by repeatedly guessing the optical properties and comparing the expected observables with those that you have made. Upon matching, voilà, those optical properties used to generate the expected observables are the ones that characterize your sample.

If only it were that simple. I have written a couple of papers on the details of the program [1–3], on optical properties in general [4], and on making those measurements with integrating spheres [5,6]. Unfortunately, not one of these brings all the pieces together. Partly, this is my fault, partly it is because none of the articles is really concerned with dotting all the *i*'s and crossing all the *t*'s, partly it is because an appropriate forum for this information just doesn't exist other than in this manual.

This document describes version 3.6.0 of the `iad` program and constitutes a complete revision of the previous document. The program is documented in the journal article [7].

Over the last year there has been substantial improvement in the `iad` because, for good or ill, we have been preparing optical phantoms for a number of uses. Before releasing these phantoms, it is necessary to measure their optical properties. Since the phantoms have a much wider range of optical properties than any I had considered in the past, the `iad` broke down in various ways. In fact, Ville Keränen somehow managed to break the damn program essentially every week for a full year. As a consequence, `iad` is much more robust than it has ever been before.

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Sphere Properties		
$A$	area	$\text{mm}^2$
$a$	area divided by the total sphere wall area	—
$X_d$	(subscript $d$ ) detector	—
$X_w$	(subscript $w$ ) sphere wall (ports excluded)	—
$X_e$	(subscript $e$ ) entrance	—
$X_s$	(subscript $s$ ) sample	—
$X_b$	(subscript $b$ ) beam	—
Sample Properties		
$\delta$	sample thickness	$\text{mm}$
$\mu_a$	absorption coefficient	$\text{mm}^{-1}$
$\mu_s$	scattering coefficient	$\text{mm}^{-1}$
$g$	anisotropy, (average cosine of phase function)	—
$\mu'_s$	reduced scattering coefficient = $\mu_s(1 - g)$	$\text{mm}^{-1}$
Idealized Values		
$r$	true reflectance	—
$t$	true transmittance	—
Observables		
$R$	unnormalized reflection	arbitrary
$T$	unnormalized transmission	arbitrary
$U$	unnormalized unscattered transmission	arbitrary
$M_R$	measured total reflectance (eq. 2)	—
$M_T$	measured total transmittance (eq. 3)	—
$M_U$	measured unscattered transmittance (eq. 6)	—
$X_0$	(subscript 0) empty or nothing	—
$X_{100}$	(subscript 100) all	—
$X_{std}$	(subscript $std$ ) reflectance standard	—
$X_{dark}$	(subscript $dark$ ) blocked beam	—
$X^{diffuse}$	(superscript $diffuse$ ) diffuse illumination (generally omitted)	—
$X^{direct}$	(superscript $direct$ ) direct or normal illumination	—

Table 1: Nomenclature used in this document. For example,  $A_s$  is the area of the sample port and  $a_d$  is  $A_d/(A_w + A_e + A_s + A_d)$ . The symbol  $r_s^{direct}$  describes the true reflectance from the sample for direct (normal) illumination and  $r_s$  is the sample reflectance for diffuse illumination (where the *diffuse* superscript has been omitted).

## 1 Introduction

Inverse adding-doubling is a method for generating the optical properties of scattering and absorbing materials. Reflection and transmission measurements (typically made with an integrating sphere) are converted to the optical properties of the sample (scattering, absorption, and anisotropy) using the computer program `iad`. The `iad` has been extensively tested and validated for accuracy and precision; it is widely used [6, 8–110].

### 1.1 Overview

To use the `iad` program, create a data file with the appropriate header for your experiment and then append your experimental measurements of reflectance and transmittance. If the file is called `data.rxt`<sup>1</sup> then the file can be processed by the `iad` program by typing

```
prompt> iad data
```

You'll get back a file `data.txt` that contains a set of optical properties—the scattering, absorption, and anisotropy coefficients that correspond to each measurement set.

The `iad` program finds the optical properties characterizing a sample by using the measured total reflectance  $M_R$ , the measured total transmittance  $M_T$  values, and potentially the measured unscattered transmittance values  $M_U$ . A set of optical properties is guessed and values for  $M_R$  and  $M_T$  are calculated. These values are compared with the measured values. If they match then the optical properties for the sample have been found. If they do not match then a new set of optical properties is guessed and the process is repeated. As van de Hulst pointed out, the approach is fairly brute force. Oh, well it works.

The `iad` program has a number of parameters that can be modified to account for different experimental situations. One, two, or three measurements can be used to determine one, two, or three optical properties. `iad` is intended for batch operation, but can be used from the command line. The processed output file contains a summary of the experimental conditions as well as all the calculated optical properties.<sup>2</sup>

### 1.2 Assumptions

The adding-doubling method is used to calculate reflectance and transmittance. Since the adding-doubling method is an accurate solution of the radiative transport equation for all albedos, all optical depths, and all phase functions this method can be applied to any medium for which the radiative transport equation is valid. However, a number of restrictions apply

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<sup>1</sup>The `.rxt` extension is arbitrary and is the default extension for data files to be processed by this program. Any other extension is permitted, but you will have to type it out completely (e.g., `iad data.dat`).

<sup>2</sup>The optical properties are readily graphed using `gnuplot` because all the comment lines at the top of the file are preceeded by a hash mark (`#`).

1. distribution of light is independent of time,
2. samples have homogeneous optical properties,
3. tissue geometry is an infinite plane-parallel slab,
4. boundaries are smooth,
5. internal reflectance is governed by Fresnel's law, and
6. polarization effects may be ignored.

Despite these constraints, inverse adding-doubling has a number of positive features

1. works for any combination of optical properties.
2. takes into account all the interactions of a sample sandwiched between glass slides
3. incorporates the effects of the integrating spheres on the measured values
4. accounts for lost light (light that is not collected by the spheres)
5. has a reasonable trade-off between accuracy and speed.

### 1.3 Background

#### The graduate student days

The `iad` program started its life as an inverse diffusion program.<sup>3</sup> Steve Jacques (then at the Wellman Laboratories of Photomedicine) and I (a graduate student at the University of Texas at Austin) just wanted a way to find the optical properties of tissues. We published a paper [1] and thought that was that. We subsequently discovered that the diffusion model can generate negative reflectances for samples with large anisotropies ( $g = 0.9$ )! Steve encouraged me to improve the inverse diffusion program. I subsequently improved that program in two important ways. First, I used the  $\delta$ -Eddington approximation to eliminate the problem with highly anisotropic scattering. Second, I replaced my simple-minded iteration method with the simplex method of Nelder and Weaver. In this manner the inverse  $\delta$ -Eddington model was created. I used this in/for my thesis that was finished in 1982. During this time I my advisor A. J. Welch and the cash came from some Navy contract.

<sup>3</sup>This is somewhat ironic because when I translated the program into ANSI C, I did not convert the diffusion code. Version 1.4 of the `iad` program does not allow one to make calculations using a diffusion model.

### The post-doc era

About the time I was giving my final defense I realized that I could not recommend my inverse program to anyone because I had no idea how accurate it was. I started working on a paper to estimate the errors but it seemed pretty pointless. I decided to take the adding-doubling program that I had just written (for a different section of my thesis) and substitute it for the  $\delta$ -Eddington program in the inverse algorithm. This led to the inverse adding-doubling (**iad**) program. This was done in 1983 in Amsterdam under the auspices of Martin van Gemert and the necessary guilders came from the Fundamenteel Onderzoek voor Materie. This program was much more accurate. In fact, you can make the adding-doubling method as accurate as desired by increasing the number of quadrature points. There were a couple of major snags in the implementation—most notably the problem with internal reflectance at boundaries. It nearly killed me, but once found, the solution was surprisingly simple as long as really understand how quadrature works.

Since then, most of the implementation difficulties with the **iad** program can be traced to how the first guess is made. I spent a week working in Houston in 1984 with Steve Jacques (the at M. D. Anderson Cancer Center) trying to solve this problem. This problem is basically the same as trying to solve the entire inverse problem quickly and accurately, but with possibly poor precision. In the process of solving this problem I incorporated a version of the  $\delta$ -Eddington program into the **iad**. After a bit more fussing (a lot actually) this became the core of the current **iad**. The last little bit of work was done during a couple of months spent at the University of Utah Laser Institute in Salt Lake City with generous support from Dr. Richard Straight. Here the starting routine was completely rewritten. I developed the heuristics for calculating starting values that were eventually published in the adding-doubling paper [2]. Large portions of the code were split up into smaller files and the first version of this documentation was written.

While I worked at Wellman Laboratories (1984-1985) in Boston as a post-doc under Dr. Rox Anderson, I fixed a number of minor bugs and changed the headers for the input data files to bring the sphere corrections into accordance with the integrating sphere paper I worked on with Pickering and others. I also implemented the use of a stored grid to extract values from. This can yield immensely faster results when the optical depth or anisotropy is known *a priori*. However, if neither of these is the case, then using the stored grid does not help much, because a new grid must be calculated for each data point that is analyzed.

### The transition to C

Up to this point, **iad** was written in Pascal. I know that this is a horrible thing to admit, but I actually liked Pascal. Unfortunately, by 1986 it was clear that Pascal was seriously out of style and it was becoming harder and harder to find good Pascal compilers. All the cool programmers were using C. At the same



time I was taken by the extraordinary typesetting system of  $\text{\TeX}$  (this document is written using  $\text{\TeX}$ ). Since the `iad` is a mathematically intense program, I wanted my comments to be typeset nicely. This meant that I used the `CWEB` system<sup>4</sup> to write `iad`. This meant that I could create beautiful documentation at the cost of some extra mark-up and a pre-processing step. Therefore, one of things that I did when I moved to the Oregon Medical Laser Center, was to converted the entire program to ANSI C. I improved many things so that the program was much faster and more reliable (e.g., the stored-grid approach). This work was supported by the Collins foundation and finished in the early 1990's.

The next stage in the life of this program has been due to my most excellent interaction with David Royston at the FDA [111]. He did many careful experiments and then kept asking what was going wrong. I eventually tracked these problems down. Often they were the result of bugs that either had crept into the C port of the code, or they were the result of coding ineptitude on my part from the very beginning. Dave also provided the incentive to allow light absorption by slides on each side of the sample in the calculation.<sup>5</sup>

In the intervening ten years or so, I would correspond with users and add features or fix bugs as needed. In 2005, I looked at the code base and realized that I had added stuff (multiple-layers, conical irradiation, and a *Mathematica* interface) and that compiling the code was a huge hassle. I reorganized the code base and cleaned up the build process. Then I discovered that this document was severely out of date. I ignored the urge to update it.

### Trouble!

In 2008, I needed to revisit the whole problem of measuring optical properties. I had a contract to build a bunch of optical phantoms with closely specified tolerances for optical properties. We made a bunch of measurements to show that the optical properties were the same as the promised ones. These measurements included two thicknesses of identical polyurethane material as well as on phantoms with the same scattering coefficient, but with different absorbing properties. Unfortunately not all the measurements were consistent!

I knew that the optical properties derived by `iad` required careful measurements using integrating spheres. We made careful, measurements but the results were still inconsistent. I guessed that the difficulties might stem from the two areas that had always been problematic in the measurement process: integrating sphere corrections and lost light. The first problem could be solved experimentally by (1) measuring the integrating sphere properties, (2) using smaller sample port diameters, or (3) using a reflectance standard whose reflectance was closer that of sample. The second problem could presumably be solved by using larger samples. Little did I know.

<sup>4</sup>Link to Knuth's `CWEB` page

<sup>5</sup>This functionality is still present in the `iad` but is not one of the parameters in the usual `iad` input files. Recovering this feature just requires writing a few lines of code to explicitly specify the light absorption by the glass slides.

### Integrating sphere corrections

Integrating spheres are wonderful devices, but suffer from having a non-linear response to the presence of a sample. For example, even after correcting for dark noise, the detector on a sphere containing a sample with 25% reflectance will not measure exactly one-half the value of a 50% sample. (The presence of the sample changes the number of times that light bounces in the sphere. More reflective samples have more bounces and a greater chance of being captured by the detector. Thus the 50% sample will record slightly more than twice the value from the 25% sample. See figure 1).

Now `iad` has had code to account for the integrating sphere effect since the early 1990's. These corrections tended to be small and were easy to ignore. Moreover, the nomenclature we used was horribly obtuse *and* the derivation in the published papers contained errors [6]. This made it more or less impossible to use the integrating sphere correction factors unless one read the source code very carefully. Consequently, no one (including myself or my students) used it.

Instead of trying to find the errors in the earlier papers, my graduate student Ted Moffitt and I re-derived all the necessary formulas. We used sensible nomenclature and I insisted that all the quantities have actual physical significance. I also eliminated unnecessary frills (e.g., baffles are now required). As a consequence we came up with formulas that only required the experimentalist to know the various port sizes on the integrating sphere and the reflectance of the sphere walls. Incorporating these new formulas to the `iad` code also gave me the chance to develop a new input data file format. After forcing my students to use about twenty different versions, I finally came up with a file format that I liked. This new format forces the experimentalist to include all the relevant measurement parameters.

Unfortunately, despite a new file format and new sphere correction formulas, the optical properties of the phantoms were only consistent for the largest sample sizes (two inches or about 50 mm). This was good enough to finish the contract, but not really good enough to make me happy. I was confident that the experiments were performed correctly and I was equally sure that the light interactions with the spheres were correct. This indicated that the lost light was probably the source of any remaining discrepancies.

More recently, we now know that it is quite important to know the sphere wall reflectivity. For example on one sample the reduced scattering  $\mu'_s$  increased from 1.66 to 1.91 mm<sup>-1</sup> when the sphere reflectivity was changed from 0.960 to 0.973; consequently, it is important to measure the wall reflectivity of your spheres.

### Solving the light loss problem

The light loss problem is simple. Light hits the front of a sample and the reflected light is collected by a sphere with a finite port size. Consequently, some light reflected by the sample will not be collected. This light is lost. Furthermore, light collected by the sphere will uniformly and diffusely re-illuminate the sample.

Some of this light will hit near the the edge of the sample port; when this light re-emerges from the front of the sample it will have an excellent chance of not being collected by the sphere.

The only way that I knew of to properly account for these two types of light loss was to use a Monte Carlo program. I refused to consider just swapping out the adding-doubling code and replacing it with Monte Carlo code. A few other groups did exactly this, but I was stubborn. Monte Carlo programs are easily  $10^4\times$  slower than the adding-doubling method. I finally conceived of a hybrid approach that uses the fast inverse adding-doubling to find the optical properties for a sample.

A Monte Carlo program is used to estimate the amount of light lost for this set of optical properties. This lost light is incorporated back into the next inverse adding-doubling estimate of the optical properties. This process is repeated until the optical properties stabilize. Most experimental situations only require a few Monte Carlo iterations before stabilizing. Furthermore, when analyzed with the new `iad` that accounted for lost light, all the earlier results for the contract phantoms that we made were now consistent!

## Documentation

The last phase — much longer than I expected — has been to revise this manual.

## 2 Installation

### 2.1 Obtaining the code

The source code is available at

<http://omlc.org/software/iad/>

Links to compiled versions of the code for MacOS X (PPC and Intel) as well as for Windows (XP) are there as well. You Linux types will need to download `iad-3-6-0.zip` and compile as described below.

There are two basic distributions `iad-3-6-0.zip` and `xp_iad3-6-0.zip`. These are built from the same original source files. The former is the full distribution and contains all the CWEB<sup>6</sup> files. It is intended for unix systems and uses `make` to produce executables. The `zip` distribution contains a binary compiled version for Windows XP. It also has the minimal set of `.c` and `.h` files needed to compile a binary using Microsoft's compiler.

The actual source files `foo.w` are written in CWEB, from which the ANSI C source code files `foo.c` and `foo.h` files are generated. The documentation `foo.tex` is also generated from the same base `foo.w` files. This keeps the code and the documentation in one place and furthermore allows me to use the type-setting capabilities of T<sub>E</sub>X in the program comments (see `doc/ad.src.pdf` and `doc/iad.src.pdf`).

(I cautiously recommend the CWEB system. It really does produce beautiful documentation, but I do not think it is worth having code that is not readily sharable with others. So that you will not need to install CWEB the raw, the uncommented C source files are contained in both distributions.)

### 2.2 Compiling on Windows XP

The `iad` code has been tested and compiles cleanly under two windows development environments.

The first one is the standard Microsoft compiler Visual C++ 2005 Express<sup>7</sup> which uses the `xp_iad.zip` archive. The `xp_iad.zip` archive contains the minimum set of files required to build an executable using Visual C++. Unzip the file `xp_iad.zip`. Navigate to the directory `xp_iad` and then into `src` inside the directory. Double-click on the file named `xp_iad.vcproj` to start Visual C++ and open the program project. Select `Build`→`Build Solution` from the menu to compile the program to create the executable `iad.exe` in the `xp_iad` directory. The executable `iad.exe` is a command-line program that should be run in the `Cmd.exe` shell.

The second environment is the Minimalist GNU for Windows `mingW`<sup>8</sup>. `mingW` provides a simple Unix-like environment that runs on top of Windows XP. This second environment needs the standard `iad.tar.gz` distribution that all other

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<sup>6</sup><http://www-cs-faculty.stanford.edu/~knuth/cweb.html>

<sup>7</sup><http://msdn.microsoft.com/vstudio/express/visualc/>

<sup>8</sup><http://www.mingw.org/>

platforms use. To compile an executable, unpack the tarball, navigate to the top level directory in the distribution and type. Navigate to the top-level directory in the tarball and uncomment the line in the `Makefile` so that

```
#BINARY_EXTENSION=.exe
```

becomes

```
BINARY_EXTENSION=.exe
```

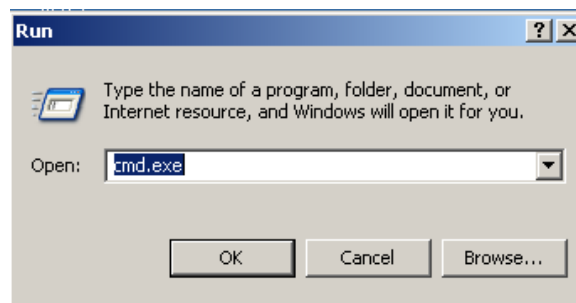
This causes the `Makefile` to produce executable files called `iad.exe` rather than just `iad`. To compile the program type

```
prompt> make
```

at the prompt. This should produce two executable files `ad.exe` and `iad.exe` in the same directory.

### 2.2.1 Testing

In Windows XP you can start `Cmd.exe` by going to the **Start Menu**, selecting **Run...**, and typing `cmd.exe`



and clicking 'OK'.

Next navigate to the directory containing `iad.exe`. (Alternatively you can add the directory containing `iad.exe` in your path variable.<sup>9</sup> and navigate to the directory containing your data files.

Test the executable by running

```
prompt> iad -v
```

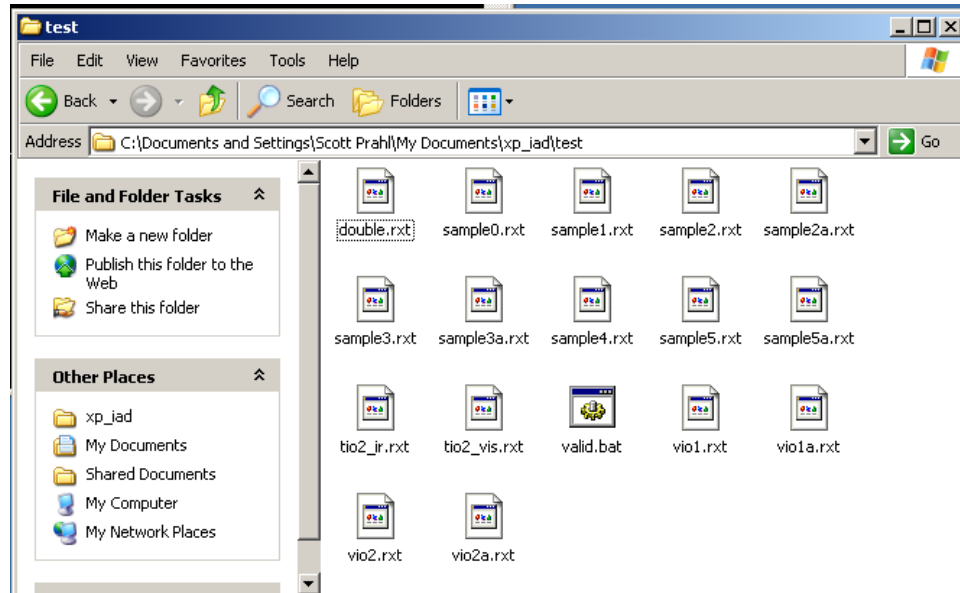
which should respond with something like

```
iad 3-11-1 (28 Mar 2019)
Copyright 2019 Scott Prahl, scott.prahl@oit.edu
(see Applied Optics, 32:559-568, 1993)
```

This is free software; see the source for copying conditions.  
There is no warranty; not even for MERCHANTABILITY or FITNESS.  
FOR A PARTICULAR PURPOSE.

<sup>9</sup>When started from a shortcut, `Cmd.exe` inherits the environment variables set in 'My Computer/Properties/Advanced/Environment'.

To run a quick validation, navigate to the `xp_iad/test` directory. Run the batch file `valid.bat` (located in the `test` directory) by double-clicking the `verify.bat` icon or by typing `valid.bat` at the `cmd.exe` prompt.



## 2.3 Installing on Unix systems

### 2.3.1 Compiling

This code has been compiled and tested on a variety of unix computers (MacOS X PPC, MacOS X Intel, Red Hat Linux x86, and SUSE Linux x86). If you just want to compile the code you just need a compiler and a `make` program. To compile an executable, unzip the compressed file, navigate to the top level directory in the distribution and type

```
prompt> make
```

at the prompt. This should produce two executable files `ad` and `iad` in the same directory.<sup>10</sup>

### 2.3.2 Testing

```
prompt> iad -v
```

should respond with something like

<sup>10</sup>To recreate all the files from the originals you'll need `perl`, `pdfLATEX`, `cweb` and `indent`.

iad 3-9-11 (10 Jul 2014)  
Copyright 2014 Scott Prahl, [scott.prahl@oit.edu](mailto:scott.prahl@oit.edu)  
(see *Applied Optics*, 32:559-568, 1993)

Once the code has been compiled, iad can be tested by at the prompt. Test the executable by running

```
prompt> make test
```

This will run a series of simple calculations. Check for validity by making sure that (column 1 = column 2) and (column 3 = column 4).

### 3 Experimental Decisions

OK, now you need to decide how you will actually make the measurements. The table below relates the basic premise that you can't get something for nothing; it gives the necessary measurements for a particular of optical properties.

Optical Property	Measured Values	Comments
$\mu_a$	$M_R$	The sample thickness is assumed to be too thick to get any light through. To estimate $\mu_a$ , the anisotropy is fixed (default $g = 0$ ), and $\mu_s = 1/\text{mm}$ .
$\mu_a, \mu'_s$	$M_R, M_T, \delta$	The anisotropy is fixed (default $g = 0$ ).
$\mu_a, \mu_s, g$	$M_R, M_T, M_C, \delta$	Three measurements are made.

#### 3.1 Decisions that you do not have to make

The `iad` code is quite flexible. In the initial releases, I thought that it was important to expose all this functionality. This had the unfortunate side effect of making easy things hard. This release is intended to make the easy things easy and while hard things remain possible (you can write your own code that calls the subroutines in `iad`).

*I have made a bunch of decisions for you.*

For example, the `iad` requires that the *total* reflection  $M_R$  and *total* transmission  $M_T$  be measured. You did not need to decide if the unscattered reflection or unscattered transmission should be included. Sometimes people like to bounce the specular reflection of the glass slide out of the entrance port (or into a beam block) so that their diffuse reflection measurement will be cleaner. Alternatively, some people like to allow the unscattered transmitted light to leave the sphere so that their diffuse transmittance measurement will be better. I decided that, in general, both of these are bad ideas.

My rationale was that experimentally separating the unscattered light from the slightly scattered light is difficult. Handling the unscattered light in the program is trivial, estimating the loss of scattered light accurately is hopeless. In the reflectance case, the specular beam reflected from the sample must travel a good distance before exiting the sphere. Ideally, this light would be a nice laser beam that would not diverge appreciably. Unfortunately, illumination is nearly always a white light source and an indefinite fraction of light will be lost out the port by which the unscattered light leaves. (Similar arguments hold for the transmitted light, with the additional caveat that the transmitted light that is only scattered a few times is strongly directional and therefore constitutes another unknown.)



Another decision I have made is how you must measure  $M_R$  and  $M_T$ . If you look at figure 3, then it seems like a pretty reasonable set of measurements. However, we tried a couple of other schemes, like making  $R(0,0)$  more like an  $R_{dark}$  experiment, i.e., just block the illumination beam. The problem with this method is that usually one does not have a small, tight illumination beam and therefore there is usually a non-negligible amount of light that misses the sample and hits the sphere walls directly,  $R(0,0) > 0$ . A similar situation exists for the transmission sphere, but because the light never enters the sphere, the background problem is much smaller.<sup>11</sup>

I have also selected the experimental parameters needed in the header of the data files. You are forced to enter these values for every experiment that you analyze. These are parameters that I, and hopefully you, will readily comprehend. For example, the header used to contain the value of  $a_s$ , but I could not look at the file header and know what the sphere size was. Similarly, instead of the easily understandable detector diameter, a weird  $b_2$  factor was used because it made things a little simpler for the code.

I have also decided that you must always have a baffle between the sample and the detector. As I said before, it is just too hard to estimate the amount of light that directly reaches the detector on the first bounce.

### 3.2 Which reflectance standard?

You will need some reflectance standards for the reflection measurement. Common NIST (no longer NBS) traceable standards are obtainable from LabSphere. I suggest buying from Avian Technologies<sup>12</sup> If you have little money, but lots of time, then you can make your own spheres and plates. These should be freshly coated with magnesium oxide (MgO) or you can use barium sulfate (BaSO<sub>4</sub>). Both coatings are somewhat fragile and the MgO coating has poor reflectance in the blue and UV region.

One thing that is not obvious is what value of reflectance for the standard is most desirable. It is important that the reflectance standard have higher reflectance than your sample.<sup>13</sup> The background reflection (nothing in the sample port) establishes the lowest amount of light reflected by a sample. The maximum amount of light should occur when 100% of the light is reflected by the sample. When such a standard is used, then all the sample measurements will fall between these two extremes (0% and 100%).

Unfortunately integrating spheres do not have a linear response to different sample reflectances (see Figure 1). This graph shows that the corrections needed for the sphere are least at the two ends, and highest in the center. The

<sup>11</sup>Using a two glass slides surrounding air or water sounds like a plausible 100% transmission (or 0% reflection) experiment. Unfortunately, interference within the slides leads to disturbingly large variations in this important reference experiment. Try it, you will be surprised how variable your results can be.

<sup>12</sup><http://www.aviantechnologies.com>

<sup>13</sup>It is axiomatic that you want to interpolate your measurements between two known points rather than extrapolate past them.

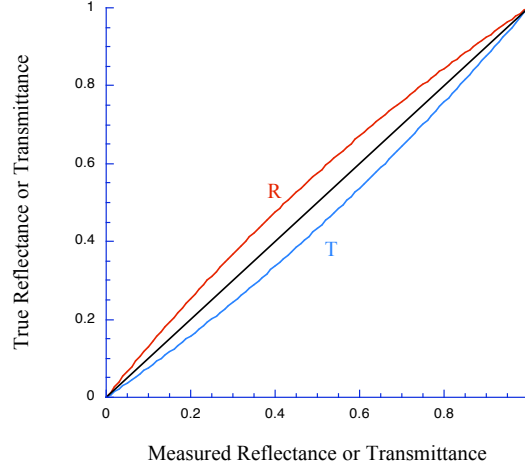


Figure 1: The relationship between measured values using an integrating sphere and the true values. In this graph, it is assumed that the dark measurement was completely dark and that the reflectance standard was 100%. These effects are typical for a relatively large sample port (50 mm port) in a moderate size integrating sphere (200 mm diameter). In general the measured values using an integrating sphere will tend to underestimate reflectance measurements ( $M_R < R$ ) and overestimate transmittance ( $M_T > T$ ).

corrections needed for the center depend on the geometric size of the sphere and ports as well as on the sphere wall reflectance. To minimize the need for these corrections, it is best to use a reflectance standard that is slightly higher than all the sample reflectances. This will reduce the magnitude of the sphere corrections and therefore the sensitivity of your measurement to integrating sphere parameters.

The magnitude of the sphere correction can be huge. For example, we were making measurements<sup>14</sup> on non-scattering, dark polyurethane samples using the substitution (single-beam) method. Now, the index of refraction of polyurethane is  $\sim 1.5$  and therefore the reflectance of the sample will be at least 4% due to specular reflectance. However, our measurements were consistently giving us 3%. The ultimate reason was that the sphere efficiency (gain) was so much greater for the standard reflectance measurement than it was for the sample reflectance measurement that when calculating  $M_R$  it came out 25% lower than expected. Obviously, this effect would have been mitigated if we had followed

<sup>14</sup>Hi Beatriz!

my own advice and used a dark reflectance standard, but I am sure that I had a good reason for ignoring myself.

### 3.3 Which integrating sphere?

There is a long history of the use of integrating spheres [5, 112–116]. Good 'ole Ulbricht proposed the first one a hundred years ago [113]. You can make your own easily. Just get a sphere, e.g., a child's ball and cut it in half. Paint the inside with  $\text{BaSO}_4$ , glue it back together attach detectors and voilà! Or you can plunk down a thousand bucks and buy one from LabSphere.

The integrating sphere is characterized by its diameter, its ports, its baffle, and its wall reflectance (typically  $\sim 98\%$ ). In all the discussion and code that follows, it is assumed that the baffle (inside the sphere) is located between the sample and the detector. This ensures that all the light leaving the sample hits the diffuse white coating on the sphere walls and is more-or-less uniformly distributed across the sphere walls. If the baffle is not present, then some (unknown) fraction of light directly propagates from the sample to the detector, while the rest of the light bounces around in the sphere. It is possible approximate the light captured by the detector and sample, but this depends on the extent of the emission profile from the sample, the geometry between the sample and the detector, as well as on the reflectance of the detector at that angle. It is much better just to require the presence of a baffle and avoid the whole mess.

The size of the sphere is usually determined by what sphere you happen to have in the lab. The sphere diameter pretty much determines the maximum sample port size. The theory of integrating spheres assumes that the sphere is spherical in shape. Obviously if you create a port in the sphere by slicing off a large chunk of the sphere and then place a flat sample on of the sample port you will no longer have a perfect sphere. A rule of thumb is that you do not want the sample area to be more than 10% of the sphere wall area.

The size of the sphere dictates the largest sample size, but the illumination beam determines the smallest sample size. Ideally the entire beam would hit the sample. In a transmittance experiment, light that misses the sample is a disaster because that light is difficult to properly measure experimentally.<sup>15</sup> Light that misses the sample in a reflectance measurement is less problematic. That light should be properly accounted for by the calibration measurements, but the light that misses the sample significantly adds to the background noise in the reflectance measurement.

The sample size is also connected to the sample thickness. Generally, one wants to have large aspect ratios for the sample (port diameter is ten times

<sup>15</sup>One might argue that the light does not enter the sphere at all and therefore will be properly accounted for by the calibration measurements. This is true, but (1) the calibration measurements will be exquisitely sensitive to shifts in the beam profile, (2) you are throwing away some signal immediately, and (3) by illuminating with a beam that is equal to the entrance aperture, you are maximizing losses out the edge of the sample—these are accounted for by `iad`, but will cause `iad` to run much more slowly because multiple attempts will be needed to properly calculate the light losses.

the sample thickness.) So for example, a beam 1 mm in diameter, a sample 1 mm thick, a port 10 mm in diameter, and a sphere 100 mm in diameter are all reasonable, practical values.

### 3.4 How many spheres?

This seems simple right? Why have a whole section devoted to counting the number of spheres? Well, mostly because the program treats the cases of 0, 1, and 2 spheres differently. The `iad` program always reads the same number of sphere parameters at the beginning of each data file (i.e., from the header). Therefore, even if you specify zero spheres, you will need to include values for two spheres in every data file.<sup>16</sup>

#### 3.4.1 Zero

When you specify 0 spheres, then the program assumes that you have made measurements of the sample in such a way that the values that you specify for  $M_R$  and  $M_T$  are devoid of integrating sphere errors and therefore `iad` *makes no corrections for the integrating spheres*. Good reasons for choosing this are

- You just want to get a quick estimate and can't be bothered to type in the five parameters needed to characterize the sphere (even though I made it really, really, easy for you.)
- You are making your measurements using the comparison method (dual beam) *and* you don't think that the uncollected light from the initial direct illumination is significant.
- You want to evaluate how much an effect the sphere corrections are having on your final derived optical properties.<sup>17</sup>
- You don't know the sphere wall reflectivity. If you want to have robust measurements then there is some merit in determining the reflectivity as outlined in this document. This measurement is consistent with the basic assumptions made in the program and helps to eliminate some sorts of systematic errors.

#### 3.4.2 One

When you specify 1 sphere, then a single sphere is assumed to have been used to measure the sample reflectance and transmittance. This does not have to be the same sphere, but it is assumed that a second sphere is not present in a way that

<sup>16</sup>Insane, I know, but the original `iad` did not require this, and automated processing became tricky. It was surprisingly hard to maintain the code and unnecessarily confusing. Now I figure that it is easier for me to make you copy-and-paste sphere parameters than it is for me to continually update `iad`.

<sup>17</sup>There is a simpler way to do this, just specify `-s` as a command line switch when processing your data.

it affects the results of the sphere making the measurements. *The program will use the values in the header to make corrections for a single sphere.* The values for the reflection sphere and the transmission sphere do not need to be identical — a single sphere means that one sphere is used at a time for all measurements.

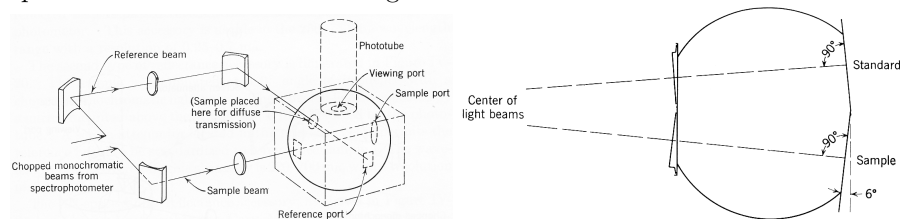
### 3.4.3 Two

Using two spheres is actually why I was originally interested in this problem.<sup>18</sup> There are a few good reasons to use two spheres. First, if the sample has dynamically changing optical properties, then it is critical that both  $M_R$  and  $M_T$  be measured at the same time. Another reason might be that you would like to measure the greatest number of samples in the shortest possible time.

Nevertheless, there are a couple of drawbacks to using two spheres at once. The first drawback is the prosaic problem of making good contact on both sides of the samples with the spheres.<sup>19</sup> One does not want to squish one side of the sample so that just barely adequate optical contact is reached on both sides. The second drawback is that it is a pain in the neck to make the calibration experiments.

## 3.5 What about dual-beam spectrometers?

Dual-beam spectrometers use two beams: the sample is placed in one beam and a reference is placed in the second beam. Two measurements are made (usually sequentially) and the reference beam is used to correct for system effects like lamp variation with wavelength or detector changes or anything that affects the light path. For example, two cuvettes might be used with one cuvette being a reference that allows most of the effects of specular reflection from surfaces to be corrected. We are most interested when two beams are used in an integrating sphere. The two common beam geometries are shown below



in these two (modified) images [117]. In both these examples the beam is incident perpendicular to the samples, but the angle between the beams might be ninety degrees (left) or shallow (twelve degrees on the right).

So why do I, or actually you, care? The geometry implies a different method of normalization. The good part is that “sphere corrections” are not needed any

<sup>18</sup>Back in the day, laser heating and coagulation of tissue was *the* application of interest. Since no one knew how the optical properties changed during high power irradiation, I tried to measure the changes in optical properties as they changed.

<sup>19</sup>One of our spheres was dropped and one port is not parallel to the port on the opposite face. Since the internal light baffles pretty much determine the necessary sphere orientation, this makes double integrating sphere measurements tricky.

longer. Because both the reference and the sample are present in the sphere during reference and sample measurements, the sphere efficiency remains constant and no “sphere corrections are needed.”

Light losses are another issue. The direct light loss is completely unaccounted for in the dual-beam sphere. Oddly, the diffuse light loss *is* properly accounted for because the diffuse sample reflectance only enters into the calculation when figuring out the integrating sphere efficiency. Since the sphere efficiency for the sample illumination and the standard illumination are equal, the diffuse light loss is not an issue.

Anyhow, if you are using a dual beam system, then if your system automatically gives you the ratio of the sample reflectance to the standard reflectance, you will want to calculate  $M_R$

$$M_R = r_{std}(R_s - R_0)$$

where  $R_0$  is a measurement obtained with nothing in the port and is present to make a first order correction for light that misses the sample and hits the sphere wall first. The value for  $M_T$  is similar

$$M_T = t_{std}(T_s - T_0)$$

although now  $T_0$  is made with a blocked sample port and represents the least amount of light that may be measured.

How does one analyze these in practice with IAD? Well, the simplest way is to just to use the -X option<sup>20</sup> and specify the sphere parameters that you used.

### 3.6 How should the sample be prepared?

There are a whole host of experimental problems which plague measurements of reflectance and transmittance. Inattention to experimental details will yield garbage. Just look at the variation in the optical properties gathered together in Cheong *et al.* [4]. These can be attributed to two sources of error—inattention to the details of the experimental measurement and to flaky optical models. Since presumably, the adding-doubling method is *not* flaky, you are saved from the second source of error. But I can do nothing about saving you from poor experimental technique except to warn you about various things.

#### 3.6.1 Freshness

If the sample is not fresh then optical properties will differ from those measured *in vivo*. Obviously, the tissue begins rotting as soon as its nutrient supply is removed. If the tissue is not in excellent condition then there will be problems. By the way, there has been essentially no work done on the changes in optical properties which take place when a tissue is removed. There has been minor work done on the changes in optical properties with heating, but nothing to that definitively says that the optical properties measured *ex vivo* are even close to those *in vitro*.

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<sup>20</sup>The ‘X’ is for crossed beams.

### 3.6.2 Boundaries

The need for glass or quartz slides at the boundaries is caused by the rough surface of most tissues. Since the boundaries are characterized by Fresnel reflection, it is relatively important for the experiment to match the quantities calculated by the code. Water, saline, or best phosphate buffered saline (PBS) or some other fluid should be used to ensure good index matching between the tissue and the slide. Care should be taken to ensure that no bubbles are formed between the slide and the tissue. Since the index of refraction of glass is 1.5 and that of tissue ranges from 1.33 to say 1.45, the boundaries are not perfectly matched. You just have to live with this or experiment with various immersion oils and various glasses and plastics achieve perfection.

### 3.6.3 Hydration

Optical properties of the sample definitely change with the amount of water in the tissue. It is important to store the tissue in air-tight containers sandwiched between moist (with PBS) towels at cool (above freezing) temperatures. If you don't believe me, then leave a piece of tissue exposed to air for 48 hours. It will appear quite different from fresh tissue.<sup>21</sup> Since your eye sees reflected and transmitted light, any measurements will also differ.

### 3.6.4 Sample Diameter

How big should the sample be? First, it must be larger than the sample port in the integrating sphere. If not, then make the port smaller. Second, you want the distance from the edge of the irradiating beam on the sample to the edge of the port ( $h$ ) to be as large as practicable. If,

$$h \gg \frac{1}{\mu_a + \mu'_s}$$

then relatively little light will be lost out the sides of the sample. “Relatively little” is a complex and depends on other things. In general, it is hard to predict without modelling carefully. Before the Monte Carlo correction was added to this code, all losses would be attributed to absorption. Consequently, any absorption coefficients generated for this sample will be too large. The ramifications now are less dire: lost light takes longer to analyze because the amount of lost light must be continually recalculated.

### 3.6.5 Variability

From spot to spot on one sample and from sample to sample. Make enough measurements that you have some idea of the average value and variance for each sample as well as the variability from sample to sample.

<sup>21</sup>Your co-workers will probably complain bitterly about the smell, but we are only concerned about the appearance of the tissue.

### 3.6.6 Blood

Should samples be measured with or without blood? Clearly neither really is a good example of the in vivo situation. If samples have no blood then the optical properties of the sample itself are being measured. The contribution to blood can be added later.

### 3.6.7 Freezing

To obtain very thin subsections it is necessary to freeze the sample. Rumor has it that the freezing process changes the optical properties. The evidence for this has been on muscle tissue. Other tissues like aorta and dermis are different and not so susceptible to freezing artifact.

### 3.6.8 Interference

Serious pain in the wahzoo. The interference usually comes from multiple internal reflections from the glass slides which sandwich the tissue sample. When a laser beam strikes the air-glass surface it is reflected. It is also reflected at the glass-tissue surface. This reflectance can interfere with the first reflectance. One would expect this to be a negligible effect, but in practice it can be nearly 10%. It is easily demonstrated by placing a microscope slide in the path of a laser beam and measuring the transmission. Then move the slide slightly and observe the transmission again. An even more dramatic effect is achieved when specular reflectance is measured.

What can be done? Sit down with a box of microscope slides and go through them one by one observing the reflectance profiles on a wall. Discard all those that show definite interference effects. These should be the least optically flat of the whole box.

Another possibility is to use very thick glass plates. By slightly rotating the plate the reflectance from the second surface will be displaced sufficiently that the two beams will no longer be aligned and consequently will not interfere. Thick plates are not ideal because they displace the integrating spheres and consequently all some diffuse light to be lost before it can enter the integrating sphere.

A final possibility is to use optically flat plates. Interference will be constant over the surface and can be measured and accounted for. This is the best option, but also the most expensive.



## 4 Measurements

### 4.1 Integrating sphere calibration

The sphere wall reflectance  $r_w$  is critical. Hopefully, you have a new, well-maintained sphere whose manufacturer-specified value for  $r_w$  is accurate. Measuring the wall reflectance  $r_w$  is challenging, but can be done. Two measurements are needed (figure 2),

$$\frac{1}{r_w} = a_w + a_d r_d (1 - a_e) + a_s r_{std} (1 - a_e) \frac{R_{std}^{diffuse}}{R_{std}^{diffuse} - R_0^{diffuse}}. \quad (1)$$

This equation illustrates the difficulty in making accurate measurements of the sphere wall reflectance. The two diffuse reflectances  $R_{std}^{diffuse}$  and  $R_0^{diffuse}$  will only differ by the amount of diffuse light leaking from the sphere when the port is empty. Consequently the difference will be small and any errors in the measurements will be magnified when the division is done.



Figure 2: The two experiments needed to determine the sphere wall reflectance. The reason that this is a hard experiment to do well is that the quantity of interest is the difference between the two measurements. This is a small difference and is roughly proportional to the relative area of the sample port  $a_s$ .

## 4.2 Single Sphere $M_R$ and $M_T$

Standard usage is that *reflection* is the light being reflected by the sample, while the *reflectance* is the light being reflected by the sample normalized by the incoming light. Reflectance has no units. A stable useful estimate for the reflectance of the sample is the *measured total reflectance*  $M_R$  which is defined in terms of easily measurable sample, standard, and background reflection measurements:

$$M_R \equiv r_{std} \cdot \frac{R(r_s^{direct}, r_s) - R(0,0)}{R(r_{std}, r_{std}) - R(0,0)}. \quad (2)$$

where  $R(r_s^{direct}, r_s)$ ,  $R(0,0)$ , and  $R(r_{std}, r_{std})$  are defined in Figure 3.<sup>22</sup>

The same idea applies to *transmission* and *transmittance*; the transmission has units while the transmittance is normalized to the incident power. The *measured total transmittance*  $M_T$  is defined as

$$M_T \equiv \frac{T(t_s^{direct}, r_s) - T_{dark}}{T(0,0) - T_{dark}} \quad (3)$$

where  $T(t_s^{direct}, r_s)$ ,  $T(0,0)$ , and  $T_{dark}$  are defined in Figure 3.

<sup>22</sup>The value of  $M_R$  differs from the true reflectance because the gain of the integrating sphere varies with the sample reflectance (see Figure 1).



Figure 3: The measurements needed to measure  $M_R$  and  $M_T$  in equations 2 and 3. Note that the baffle is always located between the sample and the detector. The back wall of the transmission sphere is never open (light that does not interact with the sample bounces around in the sphere). Finally, notice that  $T_{dark}$  and  $T(0,0)$  are very different.

### 4.3 Double Sphere $M_R$ and $M_T$

Double sphere values for  $M_R$  and  $M_T$  differ slightly from that for single spheres. The experimental arrangement for the spheres in double-sphere measurements is shown in Figure 4. The normalized reflectance is then

$$M_R = r_{std} \cdot \frac{R_2(r_s^{direct}, r_s, t_s^{direct}, t_s) - R_2(0, 0, 0, 0)}{R_2(r_{std}, r_{std}, 0, 0) - R_2(0, 0, 0, 0)} \quad (4)$$

and transmittance by

$$M_T = \frac{T_2(r_s^{direct}, r_s, t_s^{direct}, t_s) - T_2(0, 0, 0, 0)}{T_2(0, 0, 1, 1) - T_2(0, 0, 0, 0)}. \quad (5)$$



Figure 4: Measurements needed for  $M_R$  and  $M_T$  when two integrating spheres are used simultaneously.

#### 4.4 Measured Unscattered Transmittance

Finally, the *measured unscattered transmittance*  $M_U$  is the amount of light that gets through the sample without being scattered or absorbed. This has been called by a bunch of different terms: collimated transmittance, ballistic transmittance, on-axis transmittance, coherent transmittance.

$$M_U \equiv \frac{U_s - U_0}{U_{100} - U_0} \quad (6)$$

where  $U$  indicated an unscattered measurement. In this equation,  $U_s$  refers to the unscattered sample transmission,  $U_0$  is the background measurement for the unscattered transmission measurement (beam blocked) and  $U_{100}$  is the unscattered transmission measurement when no sample is present.



Figure 5: Measurements needed to determine  $M_U$ .

The measured unscattered transmittance is the most difficult measurement to make and it is frequently underestimated how hard it is to get correct. Be careful. Do yourself a favor and plot your measured values for unscattered transmittance versus the concentration or sample thickness. If this line is not exponentially decreasing then you have problems. Furthermore, you should check to make sure that your data passes through the correct value when concentration or sample thickness is zero. This is easily calculated using the indices of refraction. For example, for water ( $n = 1.33$ ) between glass slides ( $n = 1.50$ ) the transmittance should be roughly

$$M_U = (0.96)(0.996)(0.996)(0.96) = 0.914$$

where I have neglected to include multiple internal reflectances. The `iad` program does, but then it is good at that sort of thing.

Next, do a quick calculation to ensure that the amount of *scattered light* reaching the detector is significantly less than the estimated value for the unscattered transmittance. Since

$$M_U \propto e^{-(\mu_a + \mu_s)\delta}$$

the value for  $M_U$  gets small, very quickly! Do not underestimate the enormity of the smallness of  $M_U$ ! Remember, if your collimated transmittance data does not pass the above test, then the output from the `iad` program must be suspect. *Garbage in, garbage out.* It is undoubtedly better to assume a value for the anisotropy than to use a flawed value for the unscattered transmittance.

For scattering samples, measurement of unscattered transmittance is more complicated because any transmission measurement consists of contributions from both unscattered and scattered light.

$$U_{\text{measured}} = U_{\text{scattered}} + U_s$$

I can think of at least three methods for finessing this problem.<sup>23</sup> Reasonably accurate values for the measured unscattered transmittance can be obtained by make measurements at (1) various distances, (2) various angles, (3) with very thin samples in which the contribution from scattering is negligible. The last two methods are described in the paper by Jacques, Alter, and Prahl [118]. The other method has not been published (as far as I know—since I thought it up and have not had the time to go to all the trouble to write it up and review the literature.)<sup>24</sup>

If the light incident is perfectly collimated (no divergence) and the scattered light is isotropically distributed over all angles,<sup>25</sup> then the amount of light detected at a distance  $z$  from the sample is

$$\begin{array}{rcccl} \text{Measured} & & \text{Scattered} & & \text{Detector} \\ \text{Power} & = & \text{Intensity} & \times & \text{Solid Angle} & + & \text{Collimated} & & \text{Detector} \\ [\text{W}] & & [\text{W sr}^{-1}] & & [\text{sr}] & & \text{Irradiance} & \times & \text{Area} \\ & & & & & & [\text{W cm}^{-2}] & & [\text{cm}^2] \end{array}$$

where the solid angle subtended by the detector located at a distance  $z$  from the sample is

$$\text{detector solid angle} = \frac{\text{area of detector}}{\text{area of sphere with radius } z}$$

If the scattered light exiting from the back of the sample is isotropic then the intensity for this light will have a constant value  $I_{\text{scat}}$ . Therefore, if the

<sup>23</sup>Aside from using optical coherence tomography (OCT) or time-of-flight measurements

<sup>24</sup>I should probably mention that polarization can also be used to increase the collimated to scattered ratio. Unfortunately this only attenuates the diffuse contribution by a factor of two for thick samples (whose diffuse transmittance will be unpolarized.)

<sup>25</sup>This is a surprisingly good approximation when more than two scattering events take place in the sample.

irradiance is  $E_0$  then the unscattered transmission measurement will be

$$U_{measured} = \frac{I_{scattered}}{4\pi z^2} + U_s$$

Notice that the first term (the scattered light) decreases with the sample detector separation  $z$ . For a perfectly collimated beam  $U_s$  will be independent of the distance  $z$ , and therefore the first scattered term may be made as small as desired by moving the sample farther and farther from the sample.  $U_s$  is the limit of as the separation between sample and detector becomes infinite. Thus, if  $U_{measured}$  is known for several distances between sample and detector, linear regression might yield both  $U_s$  and the scattered intensity.<sup>26</sup>

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<sup>26</sup>A major assumption in this development is that the scattered light exiting the sample is isotropic. This is true for optically thick samples but definitely not for optically thin ones [118].

## 5 The Data File

The basic structure of the data file consists of a header section followed by your measurements of reflectance and transmittance. The header describes details of the experiment: the sample thickness, the beam diameter, the indices of refraction, the number and geometry of the spheres used.<sup>27</sup>

### 5.1 The header

The header was designed to be

1. easily annotated for humans
2. sufficiently flexible to accommodate all common experiments
3. readily parsed by machine
4. filled with entries that have a simple physical meaning
5. a complete description of experimental geometry
6. devoid of rarely-used parameters

These goals were achieved by carefully specifying a minimal set of parameters that describe a particular experiment and developing a simple comment style. At its simplest, the header consists of the identifying tag `IAD1` followed by eighteen numbers.

The first four characters of the file must be `IAD1`. This allows the program to do basic sanity checking of the file before beginning to parse the file. The final digit may one day be used as a version number. I hope not. Anyway, one example of a valid header is the sequence `IAD1` followed by 18 numbers,

```
IAD1 1.34 1.5 1 1 5 .96 1 203.2 25.4 12.7 1 .96 203.2 25.4 12.7 1 .96 2
```

This short example is functionally identical to the longer annotated example that is shown below. Of course, this version is not suitable for human consumption. Nevertheless, it demonstrates that (1) the comments help readability tremendously, (2) the quantity of the parameters is fixed, and (3) the ordering is fixed.

To make my life easier, I allow and nearly always use comments. A number sign (`#`), aka octothorpe, pounds sign, or hash symbol is used to start a comment; everything from the number sign to the end of the line will be ignored. White space and comments are ignored except when following a number. There is no difference between comments, tabs, spaces, linefeeds, or carriage returns; any one will end a number.

---

<sup>27</sup>Details about *how the measurements should be analyzed* are handled by command-line options (next section). This allows one (for example) to assume different default anisotropies, a different number of quadrature points, or different constraints on the Monte Carlo simulations. Of course, this is also a lie. I have extended the number of command-line parameters to allow one to specify all the parameters in the header of the data file. This allows one to do an IAD calculation without needing to create an input data file, which I find to be useful when scripting tasks — or asking *what if* questions from the command-line.

```

IAD1  # Must be first four characters

# Input Example with a single set of sphere coefficients
# The order of entries is important
# Anything after a '#' is ignored, blank lines are also ignored

1.34  # Index of refraction of the sample
1.50  # Index of refraction of the top and bottom slides
1.0   # [mm] Thickness of sample
1.0   # [mm] Thickness of slides
5.0   # [mm] Diameter of illumination beam
0.96  # Reflectance of the calibration standard

1     # Number of spheres used during each measurement

      # Properties of sphere used for reflectance measurements
203.2 # [mm] Sphere Diameter (8 in * 25.4 mm/in)
25.4  # [mm] Sample Port Diameter
12.7  # [mm] Entrance Port Diameter
1.00  # [mm] Detector Port Diameter
0.96  # Reflectance of the sphere wall

      # Properties of sphere used for transmittance measurements
203.2 # [mm] Sphere Diameter (8 in * 25.4 mm/in)
25.4  # [mm] Sample Port Diameter
0.00  # [mm] Entrance Port Diameter
1.00  # [mm] Detector Port Diameter
0.96  # Reflectance of the sphere wall

2     # Number of measurements, M_R, M_T

```

In this annotated header, the details of the geometry of the experiment are immediately apparent: reflectance and transmittance was measured using a single sphere in which the 12.7 mm diameter sample was illuminated by a 5 mm diameter beam. We also know that the sample was sandwiched between glass slides ( $n = 1.5$ ) that were 1 mm thick.

The last number in the header determines how many measurements are available. This number may vary from 1 to 6 depending on the experimental circumstances.

When using one reflectance sphere, the entrance port diameter defined as the size of the port through which light enters the sphere. The sample port is the size of the hole in the sphere that is adjacent to the sample.<sup>1</sup> Finally, the detector port diameter is the size of the port that houses the detector or optical fiber that goes to the detector. The reflectivity of all the ports is assumed to be zero — i.e., the ports are black.

If your detector is not round, but rectangular, then the detector diameter should be chosen to match the area of your detector. Thus if your detector is a rectangle  $10 \times 1 \text{ mm}^2$ , you would use a diameter of  $\sqrt{40/\pi} \approx 3.6 \text{ mm}$ . Furthermore, if you have multiple detectors in your sphere (e.g., for different wavelength regions), the detector diameter should correspond to area equal to the sum of all the detector areas.

When using one transmission sphere, the entrance port should be set to zero. We are assuming direct collimated illumination of the sample. The sample port

<sup>1</sup>The actual sample size is always assumed to be larger than the sample port. In fact, the sample is assumed to have infinite lateral extent. It is a challenge when working with small pieces of tissue to maintain a large port size and still have the sample port completely covered by the sample.



size should be set to the size of the port next to the sample. The detector size should be set as described above. The reflectance and transmittance spheres differ in the number of places that light can be lost. A reflectance sphere loses light out the entrance port, the detector port, and the sample port (through the sample). A transmittance sphere loses light only from through the detector ports and the sample port.<sup>1</sup>

## 5.2 The data section

After the header comes measurements of the reflectance  $M_R$  and transmittance  $M_T$ . This is sort of a wonky section because I wanted to have a reasonable amount of flexibility; this especially handy when combined with unix scripting to automatically process large numbers of integrating sphere measurement files. The organization is not *that* confusing, but you should read this section to understand why things are the way they are. My rationale will not make the layout any less kludgy, but it may provide a modicum of amusement where otherwise there would only be anger and frustration.

Ideally one would always have three (or more) measurements. Unfortunately, often the unscattered transmittance measurement is not available. For example the the amount of unscattered light that propagates through a millimeter of tissue is very small. For thicker samples the total transmitted light  $M_T$  is zero also. In this case then only one parameter can be measured. The following describes how you deal with multiple columns of data.

### 5.2.1 White space

The same rules for white space and comments that apply in the header section also apply to the data section. This means that you can liberally comment your data (or not) by using `#` to start a comment.

### 5.2.2 Optional wavelength entry

The `iad` program is mainly intended to analyze spectral measurements of reflectance and transmittance. For a number of reasons (e.g., graphing) it is convenient to precede the measured values by the appropriate wavelength,

<code>#lambda</code>	<code>M_R</code>	<code>M_T</code>
800	0.16830	0.24974
810	0.16271	0.26479
820	0.16289	0.25530

Since we don't want to require wavelength data, it would be nice if `iad` should also give identical results for input data like this

<sup>1</sup>Of course, in our current set up in the lab, we have mounted the reflectance fiber permanently in one sphere that is used for both transmittance and reflectance measurements. In this case, there are identical causes of light loss in both spheres and therefore the entrance port size in the transmittance sphere might be non-zero or this area could be lumped into the detector port size. In practice it is just easiest to make the parameters for both spheres be equal to one another.

#	M_R	M_T
0.16830	0.24974	
0.16271	0.26479	
0.16289	0.25530	

There are at least two possible solutions to solve this problem: add another parameter to the header or auto-recognize the presence of a wavelength. Since I wanted to keep the number of parameters in the header to a minimum, I went with the second option. This leads to the following rule.

*If the first number on a line is greater than one, that number is assumed to be a wavelength, otherwise it is interpreted as  $M_R$ .*

The value of the wavelength is not used anywhere except for being copied to the output file. The above rule means that you cannot measure your wavelength in microns if that forces the wavelength to fall between 0 and 1. For example 532.8 is fine as a wavelength but 0.5328 will be interpreted as a value for  $M_R$ . This is why all the example files in the test directory use nanometers for the wavelength range.

### 5.2.3 One measurement

#lambda	M_R
800	0.16830
810	0.16271
820	0.16289

When the only measurement specified, it is assumed to be the measured reflectance  $M_R$ . Since there is only a single measurement two assumptions are made:

- the optical thickness of the slab is assumed infinite and
- the anisotropy coefficient set to the default value ( $g = 0$  unless a specific command-line value has been specified).

The albedo is varied until the correct value for the measured reflectance  $M_R$  is obtained. Now `iad` generates both absorption and scattering coefficients. Because only the albedo is known, and `iad` reports both the scattering and absorption, some value for the scattering coefficient is assumed ( $\mu_s = 1$ ). There is really no good reason for this other than the general observation that scattering is relatively constant (as a function of wavelength) for most samples that are measured. One might process a reflectance-only data set to get a sense of how absorption varied across the range of input values.

More recently, I have expanded the command-line options for `iad`. This means that if you happen to know the absorption coefficient and have a reflectance measurement you might try

```
prompt> iad -A 0.01 -g 0.6 file.rxt
```

which will constrain the  $\mu_a = 0.01 \text{ mm}^{-1}$  and the  $g = 0.6$ . Therefore the scattering coefficient will be found.

### 5.2.4 Two measurements

#lambda	M_R	M_T	
800	0.16830	0.24974	
810	0.16271	0.00000	#treated as one measurement
820	0.16289	0.25530	

This is the most common case and

- the anisotropy coefficient set to the default value ( $g = 0$  unless a specific command-line value has been specified).

The first two values are the measured reflectance and the measured transmittance. As long as  $M_T \neq 0$  then the anisotropy coefficient set to the default value and the albedo and optical depth are varied until the correct values of  $M_R$  and  $M_T$  are obtained. The scattering and absorption coefficients are calculated from the final values for the albedo and optical depth and printed. When  $M_T = 0$  then the situation is identical to the one measurement case above and is treated in exactly the same way.

### 5.2.5 Three or more measurements

#lambda	M_R	M_T	M_U	
800	0.16830	0.24974	0.0012	
810	0.16271	0.00000	0.0000	#treated as one measurement
820	0.16289	0.25530	0.0000	#treated as two measurements

If both  $M_T \neq 0$  and  $M_U \neq 0$ , then  $M_U$  is used to calculate the optical thickness directly. The optical thickness is calculated by including all the multiple internal reflections in the slide and sample and using  $M_U$ . The albedo and anisotropy are varied until  $M_R$  and  $M_T$  are matched.

If  $M_U = 0$  but  $M_T \neq 0$ , then the situation is identical to the two measurement case above. Finally when both  $M_T = 0$  and  $M_U = 0$  then the single measurement case above applies.

### 5.2.6 Four measurements

#lambda	M_R	M_T	M_U	r_w	
800	0.16830	0.24974	0.0012	0.951	
810	0.16271	0.00000	0.0000	0.952	#treated as one measurement
820	0.16289	0.25530	0.0000	0.953	#treated as two measurements

This is how **iad** handles the situation in which the sphere wall reflectance varies with each measurement (e.g., as a function of wavelength). Both the reflectance sphere and the transmittance sphere wall reflectances are changed by this value. If you want to use this option when you only have one or two measurements, then indicate that you have four measurements and enter 0.00 for the unknown values (as in the 810 nm line above). This option was originally needed because we needed to use a sphere that was coated for use in the visible with wavelengths from 1,000–1,600 nm. In this region the wall reflectance changed significantly over the wavelength range that we were using.

The real advantage of including the sphere wall reflectance measurements  $r_w$  that were made according to section 4.1 are that it negates a variety of systematic errors that might be present in your measurement. Since the  $r_w$

measurement is made with the same light source, detector, spheres, and standards *and* is analyzed using the same set of integrating sphere assumptions as the  $M_R$  and  $M_T$  measurements, the two sets of measurements are consistent. To put it bluntly, if LabSphere says your sphere has a reflectance of 0.995 and you measure a value of 0.980, I would assert that your value is more appropriate for analyzing your integrating sphere measurements.

### 5.2.7 Five measurements

#lambda	M_R	M_T	M_U	r_w	t_w
800	0.16830	0.24974	0.0012	0.951	0.980
810	0.16271	0.00000	0.0000	0.952	0.981 #treated as one measurement
820	0.16289	0.25530	0.0000	0.953	0.982 #treated as two measurements

This is how `iad` handles the case of spheres with different reflectances as a function of wavelength. Typically if the wall re is changing, then so will the sphere wall reflectance. Of course, if the wall reflectance is not changing then just fill in the same value each time. Like the four measurement case, if  $M_U$  or  $M_T$  is unknown, then just enter zero.

### 5.2.8 Six measurements

The insanity continues and we allow the reflectance standard to have some spectral character.

# nm	M_R	M_T	M_U	r_w	t_w	rstd
550	0.0731	0.9307	0.0000	0.9788	0.9788	0.9800

Typically if the wall reflectance is changing, then so will the standard reflectance. Of course, if the wall reflectance is not changing then just fill in the same value each time. Like the five measurement case, if  $M_U$  or  $M_T$  is unknown, then just enter zeros.

### 5.2.9 Seven measurements!

The insanity finally ends. The default transmittance for 100% transmission gets specified also.

# nm	M_R	M_T	M_U	r_w	t_w	rstd	tstd
550	0.0731	0.9307	0.0000	0.9788	0.9788	0.9800	1.0000

### 5.2.10 Putting it all together

```

IAD1
#
# Tests using calculated values for M_R and M_T
# by Scott Prahl
#
1.4          # Index of refraction of sample
1.0          # Index of refraction of top slide
1.0          # [mm] Thickness of sample
1.0          # [mm] Thickness of slides
2.0          # [mm] Diameter of illumination beam
1.00         # Reflectance of calibration standard

0           # [mm] Number of spheres used during each measurement

           # Refection sphere properties (unused because n_spheres=0)
203.2        # [mm] Sphere Diameter
25.4         # [mm] Sample Port Diameter
12.7         # [mm] Entrance Port Diameter
12.7         # [mm] Detector Port Diameter
0.96         # Reflectivity of the sphere wall

           # Transmission sphere properties (unused because n_spheres=0)
203.2        # [mm] Sphere Diameter
25.4         # [mm] Sample Port Diameter
12.7         # [mm] Entrance Port Diameter
12.7         # [mm] Detector Port Diameter
0.96         # Reflectivity of the sphere wall

2           # [mm] Number of measurements

#           M_R           M_T           a     b     g
2.77865808457e-2  1.73065997660e-2 # 0.00 4.0000 0.00
4.20236699283e-2  1.91254597157e-2 # 0.19 4.0000 0.00
6.00960999727e-2  2.19067707658e-2 # 0.36 4.0000 0.00
8.34695920348e-2  2.63617299497e-2 # 0.51 4.0000 0.00
1.14361397922e-1  3.38563099504e-2 # 0.64 4.0000 0.00
1.56228601933e-1  4.70846891403e-2 # 0.75 4.0000 0.00
2.14577898383e-1  7.13708102703e-2 # 0.84 4.0000 0.00
2.97877311707e-1  1.16591498256e-1 # 0.91 4.0000 0.00
4.15349513292e-1  1.96420803666e-1 # 0.96 4.0000 0.00
5.54938077927e-1  3.06980103254e-1 # 0.99 4.0000 0.00
6.29535913467e-1  3.70464086533e-1 # 1.00 4.0000 0.00

```

Table 2: One of the test files `basic4.rxt`. This is a test file with accurately calculated values for  $M_R$  and  $M_T$ . As a consequence the number of spheres has been set to zero. Other test files in the `test` directory may be helpful starting templates for your data.

## 6 Running iad

The `iad` is run from the command-line. It is typically used with data files that contain values of  $M_R$  and  $M_T$  as a function of wavelength. `iad` can be used without a data file by using various command-line options or switches (see Table refswitches), but it is reasonably clumsy. In its simplest form one could analyze the data file `basic1.rxt` by typing

```
prompt> iad basic1
```

Note that the extension `.rxt` is automatically appended if necessary. This command will process the data file `basic1.rxt` and produce a file named `basic1.txt`.

## 6.1 Basic Program Information

```
-h  display help
-v  version information
```

These commands are pretty self-explanatory. The `-h` switch lists all the command line options. The `-v` switch just prints the version information of the program.

## 6.2 Controlling the Analysis

```
-e #  error tolerance (default 0.0001)
-q #  number of quadrature points (default=8)
-M #  number of Monte Carlo iterations
-p #  # of Monte Carlo photons (default 100000),
      (negative number is max time in milliseconds)
```

### 6.2.1 The Error Tolerance

The `-e #` option allows you to specify the tolerance for the relative error allowed before the program quits. The default value is 0.0001. It is unlikely that you will need to use this option since the default value seems to work pretty well in general. The relative error used by the program is<sup>28</sup>

$$error = \left| \frac{M_R^{measured} - M_R^{calculated}}{M_R^{measured} + 10^{-6}} \right| + \left| \frac{M_T^{measured} - M_T^{calculated}}{M_T^{measured} + 10^{-6}} \right|$$

As an example, the optical properties for  $M_R = 0.3$  and  $M_T = 0.1$  for a 1 mm thick sample are found using the command<sup>29</sup>

```
prompt> iad -m -r 0.3 -t 0.1
```

which produces

```
0.3000  0.3000  0.1000  0.1000  0.6730  3.0109  0.0000  0
```

The first two numbers are values for  $M_R$ ; the first one is measured and exactly matches the second one that is calculated using the given optical properties. The third and fourth numbers are values for  $M_T$  and again they match exactly. The next three are the calculated optical properties  $\mu_a = 0.67/\text{mm}$ ,  $\mu'_s = 3.01/\text{mm}$ , and  $g = 0.9$

If the error tolerance is changed to 0.01 then

```
prompt> iad -m -r 0.3 -t 0.1 -e 0.01
```

produces slightly different optical properties

<sup>28</sup>The value of  $10^{-6}$  is present to prevent division by zero.

<sup>29</sup>Here I have used the `-m` switch to eliminate the bulk of the output. The eliminated output tells you what sample thickness was assumed and other assumptions made by the program.

```
0.3000 0.3002 0.1000 0.0991 0.6750 3.0229 0.0000 0
```

where the calculated and measured values for  $M_R$  nearly match, but the : the measured value of 0.3000 and the calculated value (using the listed optical properties) of 0.3002.

### 6.2.2 The Number of Monte Carlo Iterations

The `-M #` option allows you to control the *maximum* number of Monte Carlo iterations. The default setting is to do as many Monte Carlo iterations as necessary (up to 20) to ensure that the calculated values for scattering and absorption change by less than  $0.01 \text{ mm}^{-1}$  between iterations. Now the number of Monte Carlo iterations is *not* the same as the number of photons used in the Monte Carlo calculation of lost light. Instead, the `-M #` option controls the maximum number of times that the lost light calculation is done.

This option is your friend.<sup>30</sup> You'll almost always want to turn off the Monte Carlo lost light iterations because the program will run *much, much* faster. Use<sup>31</sup>

```
prompt> iad -M 0 double
```

until your data file converts and you are reasonably certain that the data is worth spending more time on.

For example if the two sphere data is used `double.rxt` then

```
prompt> iad double
```

takes 41.9 seconds on my computer. This calculation does at least two Monte Carlo iterations for each of the 21 data points. Now if the Monte Carlo is completely omitted (0 iterations) then

```
prompt> iad -M 0 double
```

then the calculation only takes 0.3 seconds. Using the option `-M 0` is also handy because it allows you to quickly test the formatting of data files or whether you might have some crazy measurements.

### 6.2.3 The Number of Monte Carlo Photons

The `-p #` option allows you to specify the number of photons used in the Monte Carlo simulation *or* the length of time that one Monte Carlo iteration should take. The default is 100,000 photons. This option can be handy when you have samples with very low absorption and a single photon takes a long time to propagate. In this case the number of photons can be reduced explicitly. For example, the command

```
prompt> iad -p 10000 double
```

<sup>30</sup>`-M 0` is your bonus for reading this far and this deeply in the manual.

<sup>31</sup>`double.rxt` is found in the test directory that shipped with `iad`.



will take only 4.9 seconds. Alternatively, you might find it convenient to limit the calculation time to 100 milliseconds for each Monte Carlo simulation,

```
prompt> iad -p -100 double
```

#### 6.2.4 The Number Quadrature Points

The `-q #` option allows you to specify the number of quadrature points used in the adding-doubling calculation. There are a bunch of integrals that must be calculated numerically; quadrature is an efficient way to do these integrals accurately. `iad` uses a combination of Lobatto and Gaussian quadrature to achieve good results.

The default number of quadrature points or angles is 8. For a variety of reasons, the number of quadrature points should be a multiple of 4. Four quadrature points is more-or-less equivalent to the  $\delta - P_3$  approximation. The default value of 8 is accurate for highly anisotropic phase functions and runs quickly.

```
prompt> iad -q 4 double
```

takes only 0.1 seconds. While

```
prompt> iad -q 16 double
```

takes 1.3 seconds.

### 6.3 Constraining the Optical Properties

Command line constraints allow you to add information to the inversion process.

```
-g #  fixed anisotropy (default 0)
-a #  fixed albedo
-b #  fixed optical thickness
-A #  fixed absorption coefficient
-F #  fixed scattering coefficient
```

#### 6.3.1 Constraining the Scattering Anisotropy

The `-g #` option allows you to control the value of  $g$  that is used in the inverse calculation. The default value is 0. This option is particularly useful when you have only made two measurements, but know approximately what the value of the average cosine of the phase function should be. For example,<sup>32</sup>

```
prompt> iad -r 0.3 -t 0.1
```

produces

```
0.3000  0.3000  0.1000  0.1000  0.6730  3.0109  0.0000  0
```

<sup>32</sup>Here I have used the `-m` switch to eliminate the bulk of the output. The eliminated output tells you what sample thickness was assumed and other assumptions made by the program.

or  $\mu_a = 0.67/\text{mm}$ ,  $\mu'_s = 3.01/\text{mm}$ , and  $g = 0.9$ . Alternatively, if the value of  $g$  is specified then the resulting optical properties are

```
prompt> iad -r 0.3 -t 0.1 -g 0.9
```

produces

```
0.3000  0.3000  0.1000  0.1000  0.5522  3.1601  0.9000  0
```

or  $\mu_a = 0.55/\text{mm}$ ,  $\mu'_s = 3.16/\text{mm}$ , and  $g = 0.9$ . Interestingly in this example, the reduced scattering coefficient changes by only 5%, but the absorption changes nearly 40%!

### 6.3.2 Constraining the Albedo

Sometimes it is very handy to specify that the sample is totally absorbing or totally scattering. In this case, one can just type something like

```
prompt> iad -a 1 -r 0.3
prompt> iad -a 1 -t 0.3
prompt> iad -a 1 -g 0.9 -r 0.3
prompt> iad -a 1 -g 0.9 -t 0.3
prompt> iad -a 0 -t 0.3
```

### 6.3.3 Constraining the Optical Thickness

```
prompt> iad -b 1 -r 0.3
prompt> iad -b 1 -g 0.9 -r 0.3
prompt> iad -b 1 -t 0.5
prompt> iad -b 1 -g 0.9 -t 0.5
```

Constraining the optical thickness ( $b = (\mu_a + \mu_s)d$  where  $d$  is the thickness of your sample) is essentially the same as specifying the unscattered transmission through the sample, except that you need not worry about specular reflection at boundaries.

### 6.3.4 Constraining the Absorption Coefficient

```
prompt> iad -A 1 -r 0.3
prompt> iad -A 1 -g 0.9 -r 0.3
prompt> iad -A 1 -t 0.3
prompt> iad -A 1 -g 0.9 -t 0.3
prompt> iad -A 1 -r 0.1 -t 0.2
prompt> iad -A 1 -r 0.1 -t 0.2
prompt> iad -A 1 -g 0.9 -r 0.1 -t 0.2
```

Sometimes you know what the absorption coefficient is and it is handy to specify it directly. Oddly the constraint that I use most is the no absorption condition:

```

prompt> iad -A 0 -r 0.3
prompt> iad -A 0 -g 0.9 -r 0.3
prompt> iad -A 0 -t 0.3
prompt> iad -A 0 -g 0.9 -t 0.3

```

This is absurdly handy when dealing with white things like Intralipid and teflon and titanium dioxide samples.

### 6.3.5 Constraining the Scattering Coefficient

```

prompt> iad -F 1 -r 0.3
prompt> iad -F 1 -g 0.9 -r 0.3
prompt> iad -F 1 -t 0.3
prompt> iad -F 1 -g 0.9 -t 0.3

```

This is probably the least useful of all the constraints, but every once in a while it comes in handy.

## 6.4 Basic Experimental Description

-r #	total reflection measurement
-t #	total transmission measurement
-u #	unscattered transmission measurement
-n #	specify index of refraction of sample
-N #	specify index of refraction of slides
-d #	physical thickness of the sample
-D #	physical thickness of the slides
-S #	number of spheres used
-1 '# # # # #'	reflection sphere parameters
-2 '# # # # #'	transmission sphere parameters

Sometimes it is convenient to analyze data from the command line.

### 6.4.1 The Measured Reflectance $M_R$

The -r # option allows you to specify a reflectance to be used in the calculation. This number must be between 0 and 1.

### 6.4.2 The Measured Transmittance $M_T$

The -t # option allows you to specify a transmittance to be used in the calculation. This number must be between 0 and 1.

### 6.4.3 The Unscattered Transmittance $M_U$

The -u # option allows you to specify a unscattered transmittance to be used in the calculation. This number must be between 0 and 1.

#### 6.4.4 The Index of Refraction of the Slab

The `-n #` option allows you to specify the index of refraction of the sample. This number must be between 0.5 and 2.0. The default is 1.0.

#### 6.4.5 The Index of Refraction of the Glass Slides

The `-N #` option allows you to specify the index of refraction of the glass slides. This number must be between 0.5 and 2.0. The default is 1.0.

#### 6.4.6 The Physical Thickness the Sample

The `-d #` option allows you to specify the physical thickness of the sample. This number must be positive and in millimeters. The default is 1.0 mm.

#### 6.4.7 The Physical Thickness of the Glass Slides

The `-D #` option allows you to specify the thickness of the glass slides. This number should be positive. The default is 0.0 mm which means that there are no glass slides. This thickness is used only in the Monte Carlo simulations to figure out how much light is lost out the edges of the integrating sphere measurements. The units should be millimeters.

#### 6.4.8 The Number of Spheres

The `-S #` option allows you to specify the number of spheres used. This number must be 0, 1, or 2. The default is zero.

#### 6.4.9 The Reflection Sphere Properties

The `-1 '# # # # #'` option allows you to specify the properties of the reflection sphere. The five numbers (four diameters and the wall reflectance) must be surrounded by quotes. The five numbers must appear in this order (sphere diameter, sample port diameter, entrance port diameter, detector port diameter, and sphere wall reflectance). The units should be millimeters.

#### 6.4.10 The Transmission Sphere Properties

The `-1 '# # # # #'` option allows you to specify the properties of the transmission sphere. The five numbers (four diameters and the wall reflectance) must be surrounded by quotes. The five numbers must appear in this order (sphere diameter, sample port diameter, entrance port diameter, detector port diameter, and sphere wall reflectance). The units should be millimeters.

## 6.5 Controlling the Output

```
-o filename  explicitly specify filename for output
-V 0        verbosity low — no output to stderr
-V 1        verbosity moderate
-V 2        verbosity high
-x #        show intermediate calculations
```

### 6.5.1 The Output Filename

The `-o filename` option allows you to specify the name of the output file. The default operation is to take the name of the input file and append `.txt` to the name (or if the file ends with `.rxt` then the `.rxt` is replaced by `.txt`). For example,

```
prompt> iad -o foobar double.rxt
```

will put the optical properties for the data in `double.rxt` into the file `foobar`.

### 6.5.2 Controlling Verbosity

The `-V 0` option allows you to eliminate all the extra output from the `iad` program. This is not used by default. For example,

```
prompt> iad -r 0.3 -t 0.1
```

produces 45 lines of output. The abbreviated output from

```
prompt> iad -V0 -r 0.3 -t 0.1
```

has just one:

```
0.3000  0.3000  0.1000  0.1000  0.6730  3.0109  0.0000  0
```

The `-V 1` option is not so useful when a single line is being processed, but allows one to monitor the process of converting an `.rxt` file into optical properties. These result in the same output<sup>33</sup>

```
prompt> iad vio1
prompt> iad -V1 vio1
```

will produce the following on the screen

```
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 25 done ( 1.86 s/pt)
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 50 done ( 1.64 s/pt)
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 75 done ( 1.66 s/pt)
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 100 done ( 1.71 s/pt)
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 125 done ( 1.74 s/pt)
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 150 done ( 1.75 s/pt)
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 175 done ( 1.75 s/pt)
1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 1*1*1*1*1* 200 done ( 1.75 s/pt)
1*
```

<sup>33</sup>`vio1.rxt` is located in the test directory.

(The repeated sequence of `1*` above indicates that one Monte Carlo simulation is done and then the asterisk means that the inversion converged.) The primary reason for doing this is to keep the user informed that processing is taking place. When

```
prompt> iad -V0 vio1
```

is used, then nothing is displayed on the screen until the program finishes.

Use `-V 2` is the default. It is just like `-V 1` except it prints a message after processing the file.

### 6.5.3 Showing Intermediate Calculations

```
-x 1    show sphere and light loss effects
-x 2    show grid calculations
-x 4    show Monte Carlo iterations
-x 8    show lost light
-x 16   show sphere effects
-x 32   starting guess for optical properties
-x 64   show every calculation
-x 128  show information about search
-x 255  all of the above
```

### 6.5.4 Displaying light loss effects

The `-x 1` option allows you to collect an explicit record of how large the light loss effects are for a given calculation. To do this, each data point is analyzed four different ways (1) no sphere corrections and no Monte Carlo light loss corrections, (2) only sphere corrections, (3) only Monte Carlo light loss corrections, and (4) both sets of corrections. It takes a little more than twice as long to analyze the data.

For example the default output for

```
prompt> iad test/double
```

reports results like

#	Meas	Calc	Meas	Calc	Calc	Calc	Calc	Error
##wave	M_R	M_R	M_T	M_T	mu_a	mu_s'	g	Number
# [nm]	[---]	[---]	[---]	[---]	[1/mm]	[1/mm]	[---]	[---]
600.0	0.4378	0.4378	0.1808	0.1808	0.0190	0.8708	0.0000	0
601.0	0.4319	0.4319	0.1754	0.1754	0.0199	0.8702	0.0000	0
602.0	0.4265	0.4265	0.1706	0.1706	0.0208	0.8681	0.0000	0
...								

To show the effects of light loss, one uses the command

```
prompt> iad -x 1 test/double
```

reports results with the first eight columns as

#	Meas	Calc	Meas	Calc	Best	Best	Best
##wave	M_R	M_R	M_T	M_T	mu_a	mu_s'	g
# [nm]	[---]	[---]	[---]	[---]	[1/mm]	[1/mm]	[---]
600.0	0.4378	0.4378	0.1808	0.1808	0.0190	0.8708	0.0000
601.0	0.4319	0.4319	0.1754	0.1754	0.0199	0.8702	0.0000
602.0	0.4265	0.4265	0.1706	0.1706	0.0208	0.8679	0.0000
...							

Where the first column is the wavelength, the next is the measured value for  $M_R$  and the third is the value of  $M_R$  calculated using the optical properties shown in columns 6–8. The same holds for  $M_T$  in columns 4 and 5. The next four columns reveal the amount light lost

Lost	Lost	Lost	Lost
R_drct	R_diff	T_drct	T_diff
[---]	[---]	[---]	[1/mm]
0.0051	0.1055	0.0042	0.0364
0.0047	0.1018	0.0040	0.0358
0.0042	0.1020	0.0042	0.0331
...			

The first column is the light lost directly — collimated light that enters the sample, propagates in the sample and escapes out the front of the sample *but is not collected by the reflectance sphere*. The third column is the amount of this light that exits the back of the sample but is not collected by the transmission sphere. The second column is the amount of lost light arising from light that hits the sample as diffuse light. The two primary sources of this light are light from the source that misses the sample and hits the sphere wall first and light that exits the sample and is collected by the sphere. It is all very annoying.

The remaining columns show the effects that lost light has on the calculated properties of the sample.

None	SpOnly	MConly	None	SpOnly	MConly	MC	Error
mu_a	mu_a	mu_a	mu_s'	mu_s'	mu_s'	iter	Number
[1/mm]	[1/mm]	[1/mm]	[1/mm]	[1/mm]	[1/mm]	[---]	[---]
0.0209	0.0201	0.0220	0.6741	0.6739	0.8834	3	0
0.0220	0.0212	0.0228	0.6759	0.6752	0.8829	3	0
0.0230	0.0222	0.0237	0.6773	0.6752	0.8821	3	0
...							

Slight discrepancies between these results and the previous results are caused by variation in the Monte Carlo simulation. If one looks specifically at the reduced scattering numbers,

#	None	SpOnly	MConly	Best
##wave	mu_s'	mu_s'	mu_s'	mu_s'
# [nm]	[1/mm]	[1/mm]	[1/mm]	[1/mm]
600.0	0.6741	0.6739	0.8834	0.8708

```

601.0  0.6759  0.6752  0.8829  0.8702
602.0  0.6773  0.6752  0.8821  0.8679
...

```

One sees that the sphere-only corrections decrease the scattering coefficient. That is because the true reflectance is less than the measured value and vice versa for the transmittance. If the true reflectance and transmittance were used then the scattering coefficient would need to decrease which is what happens. Now if light losses are included then both transmittance and reflectance need to increase.

```

#          None      SpOnly  MOnly  Best
##wave    mu_a      mu_a    mu_a    mu_a
# [nm]    [1/mm]    [1/mm]  [1/mm]  [1/mm]
600.0  0.0209  0.0201  0.0220  0.0190
601.0  0.0220  0.0212  0.0228  0.0199
602.0  0.0230  0.0222  0.0237  0.0208
...

```



## 7 Validation

This program has been validated in a number of different ways. The first is using the test files `basic1.rxt`, `basic2.rxt` and `basic3.rxt`. These contain total reflectance and transmittance values for known optical properties.

For example `basic1.rxt` contains

```
#M_R      #mua  mus  g
0.2992621379268361  #0.1  1.0  0.0
0.1722475379306846  #0.3  1.0  0.0
0.1308339701159747  #0.5  1.0  0.0
0.1096241197070925  #0.7  1.0  0.0
0.0966223181570366  #0.9  1.0  0.0
```

and the output of `iad basic1` is

Meas	Calc	Meas	Calc	Calc	Calc	Calc	Error
M_R	M_R	M_T	M_T	mu_a	mu_s'	g	Number
[---]	[---]	[---]	[---]	[1/mm]	[1/mm]	[---]	[---]
0.2993	0.2993	0.0000	0.0000	0.1008	1.0000	0.0000	0
0.1722	0.1722	0.0000	0.0000	0.3022	1.0000	0.0000	0
0.1308	0.1308	0.0000	0.0000	0.5036	1.0000	0.0000	0
0.1096	0.1096	0.0000	0.0000	0.7049	1.0000	0.0000	0
0.0966	0.0966	0.0000	0.0000	0.9062	1.0000	0.0000	0

and the absorption coefficients are recovered with an error of about 0.5% when only the reflectance measurement is available.

Tables 4 and 5 show that these results remain valid when experiments with two or three measurements are analyzed. The effect of including spheres is detailed in a paper that will soon be submitted to JBO.

True $\mu_a$ [1/mm]	Calc $\mu_a$ [1/mm]	True $\mu'_s$ [1/mm]	Calc $\mu'_s$ [1/mm]
0.1	0.1006	5	4.9840
0.3	0.3016	5	4.9850
0.5	0.5025	5	4.9853
0.7	0.7034	5	4.9853
0.9	0.9042	5	4.9853
0.1	0.1005	10	9.9597
0.3	0.3015	10	9.9613
0.5	0.5024	10	9.9626
0.7	0.7033	10	9.9632
0.9	0.9041	10	9.9642
0.1	0.1005	15	14.9353
0.3	0.3015	15	14.9383
0.5	0.5023	15	14.9417
0.7	0.7033	15	14.9413
0.9	0.9042	15	14.9413
0.1	0.1005	20	19.9106
0.3	0.3015	20	19.9130
0.5	0.5024	20	19.9148
0.7	0.7033	20	19.9169
0.9	0.9041	20	19.9197
0.1	0.1005	25	24.8870
0.3	0.3014	25	24.8929
0.5	0.5024	25	24.8901
0.7	0.7033	25	24.8928
0.9	0.9042	25	24.8947

Table 3: Analysis of `basic2.rxt` that shows an inversion error of approximately 0.5% when two measurements  $M_R$  and  $M_T$  are available.

True $\mu_a$ [1/mm]	Calc $\mu_a$ [1/mm]	True $\mu'_s$ [1/mm]	Calc $\mu'_s$ [1/mm]	True $g$ [—]	Calc $g$ [—]
0.1	0.1008	0.5	0.5035	0.9	0.8993
0.3	0.3023	0.5	0.5017	0.9	0.8996
0.5	0.5045	0.5	0.4995	0.9	0.9000
0.7	0.7072	0.5	0.4969	0.9	0.9005
0.9	0.9105	0.5	0.4940	0.9	0.9010
0.1	0.1007	1.0	0.9994	0.9	0.9000
0.3	0.3023	1.0	0.9976	0.9	0.9002
0.5	0.5041	1.0	0.9955	0.9	0.9004
0.7	0.7061	1.0	0.9932	0.9	0.9006
0.9	0.9085	1.0	0.9909	0.9	0.9008
0.1	0.1007	1.5	1.4967	0.9	0.9002
0.3	0.3022	1.5	1.4949	0.9	0.9003
0.5	0.5037	1.5	1.4929	0.9	0.9004
0.7	0.7055	1.5	1.4909	0.9	0.9006
0.9	0.9073	1.5	1.4888	0.9	0.9007
0.1	0.1007	2.0	1.9938	0.9	0.9003
0.3	0.3020	2.0	1.9921	0.9	0.9004
0.5	0.5035	2.0	1.9903	0.9	0.9005
0.7	0.7050	2.0	1.9885	0.9	0.9006
0.9	0.9066	2.0	1.9865	0.9	0.9006
0.1	0.1007	2.5	2.4910	0.9	0.9004
0.3	0.3020	2.5	2.4896	0.9	0.9004
0.5	0.5033	2.5	2.4880	0.9	0.9005
0.7	0.7047	2.5	2.4862	0.9	0.9005

Table 4: Analysis of `basic3.rxt` that shows an inversion error of approximately 0.5% when three measurements  $M_R$ ,  $M_T$  and  $M_U$  are used.

## 8 Experimental Procedure

1. Prepare your sample. Sandwich it between glass microscope slides. Use normal saline to improve contact between tissue and glass slides. Do not squish your sample. You may need to use spacers. Refer to section 3.5 for more information about sample preparation.
2. Set up the transmission sphere with the proper port sizes. Generally, you want your sample to be larger than the entrance (sample) port. The beam size to be much smaller than the entrance port size. If you are using an optical fiber to collect light for your detector, then you may need to build or fabricate an insert so that the detector port is completely white except for where the optical fiber is. The diameter of your detector (as used by the program) should be the size of the non-white portion of the collection system.
3. Align your light source and sphere for transmission measurements (right side of Figure 3). Measure your beam size. You will need to make sure that you use the same beam size when making the reflection measurements later.
4. Do the  $T(0, 0)$  measurement first, followed by  $T_{dark}$ . Then place the sample over the entrance port and measure  $T(r_s^{direct}, r_s)$ . Make sure the baffle is located as shown in Figure 3. Make sure that the sample is flat against the sphere.
5. Change the sphere to make a set of reflectance measurements. You may need to insert your optical fiber in a metal tube (painted white on the outside) so that the fiber face can be placed near your sample. Make sure that your beam diameter is the same as for the transmission experiment.
6. Do the  $R(0, 0)$  measurement first and try to align so that this measurement is relatively small. Next do the  $R(r_{std}, r_{std})$  measurement, make sure that you do not saturate your detector. Finally, place your sample over the sample port and measure  $R(r_s^{direct}, r_s)$ . Make sure the baffle is located as shown in Figure 3. Make sure that the sample is flat against the sphere.
7. Collect your results into a data file. Prepare the header as outlined in section 5.1. Use equations 2 and 3 to calculate the values for  $M_R$  and  $M_T$ . Place these values after the header.
8. Convert your data using the `iad` program as outlined in Section 6.

## Appendix 1: Error Codes

Error	Description
0	No error
1	The algorithm did not converge after 5000 iterations.
2	Too little reflected light, $M_R < 0$ .
3	Too much reflected light, $M_R > 1$ .
4	Too little transmitted light, $M_T < 0$ .
5	Too much transmitted light, $M_T > 1$ .
6	Too little unscattered light, $M_U < 0$ .
7	Too much unscattered light, $M_U > 1$ .
8	Too much light, $M_R + M_T > 1$ .
9	Estimated non-absorbing sample but $M_T > M_U$
10	Estimated unscattered transmittance exceeds $M_T$
11	Estimated Fresnel reflectance exceeds $M_R$ .
12	Estimated total reflectance $M_R > 1$ .
13	Estimated total transmittance $M_T > 1$ .
14	Estimated total reflection and transmittance $M_R + M_T > 1$ .
15	Estimated total light $M_R + M_T + M_U > 1$ .
16	Diameter of sample port is invalid.
17	Diameter of entrance port is invalid.
18	Diameter of detector port is invalid.
19	Reflectance of sphere wall is invalid.
20	Reflectance of detector is invalid.
21	Reflectance of calibration standard is invalid.
22	Internal value $\gamma$ is not valid.
23	Internal value $f$ (fraction of light hitting the wall first) is invalid.
24	Internal value for phase function is invalid.
25	Value for number of quadrature points must be a multiple of 4.
26	Value for default anisotropy must be $-1 < g < 1$
27	Value for number of layers is too large.
28	Insufficient memory to allocate necessary data structures.
29	File error
30	Too much lost light.
31	Too little reflectance for the measured transmittance

## Appendix 2: Optical Properties

Light propagation through tissue is characterized by radiative transport theory with three different optical properties:

- the absorption coefficient  $\mu_a$  [ $\text{m}^{-1}$ ],
- the scattering coefficient  $\mu_s$  [ $\text{m}^{-1}$ ], and
- the phase function  $p(\hat{s}, \hat{s}')$  [ $\text{sr}^{-1}$ ].

The reciprocal of the absorption or of the scattering coefficient ( $1/\mu_a$  or  $1/\mu_s$ ) is the average distance that light will travel before being absorbed or scattered respectively. The phase function describes the fraction of light scattered from the direction  $\hat{s}$  into the direction  $\hat{s}'$ . The phase function is normalized so that its integral over all directions is one,

$$\int_{4\pi} p(\hat{s}, \hat{s}') d\omega' = 1.$$

Here  $d\omega'$  is a differential solid angle in the  $\hat{s}'$  direction. The functional form of the phase function is usually not known. In these cases the phase function is characterized by its average cosine

$$g = \int_{4\pi} p(\hat{s}, \hat{s}') (\hat{s} \cdot \hat{s}') d\omega'.$$

The average cosine of the phase function  $g$ , is usually called the anisotropy coefficient. The anisotropy coefficient varies between isotropic scattering ( $g = 0$ ) and complete forward scattering ( $g = 1$ ). Notice that the integrals in equations 2 and 3 are independent of the choice of  $\hat{s}$ . This means that the scattering profile is independent of the incoming light direction. Not surprisingly, a single anisotropy coefficient is inadequate to describe tissues that have special directions along which light scatters better or worse. For example, tissues (e.g., tendon) that have preferential scattering along oriented collagen fibers in a tissue.

The default phase function used by this program is the Henyey-Greenstein

$$p(\cos \theta) = \frac{1}{4\pi} \frac{1 - g^2}{[1 + g^2 - 2g\mu]^{3/2}}$$

Other phase functions can be used, but currently this is the only phase function implemented.

Often the the anisotropy coefficient is not known. In this case the reduced scattering coefficient is useful. It comes from similarity relations that reduce the number of necessary parameters from three down to just two.

$$\mu'_s = (1 - g)\mu_s$$

### Appendix 3: Dimensionless Optical Properties

Light propagation is characterized by three parameters: the albedo, the optical depth or thickness, and the phase function. The albedo  $a$  is a dimensionless parameter defined as the ratio of scattering coefficient to the sum of scattering and absorption coefficients

$$a = \frac{\mu_s}{\mu_s + \mu_a}$$

The optical depth  $b$  is the product of the tissue thickness and the sum of the scattering and absorption coefficients

$$b = \delta(\mu_s + \mu_a).$$

The parameter  $g$  is sometimes referred to as the anisotropy coefficient. The scattering coefficient  $\mu_s$  and the absorption coefficient  $\mu_a$  are given by

$$\mu_s = \frac{ab}{\delta} \quad \mu_a = \frac{(1-a)b}{\delta}.$$

The reduced optical albedo and optical depth are given by

$$a' = \frac{a(1-g)}{1-ag} \quad b' = (1-ag)b.$$

The inverse relations are

$$a = \frac{a'}{1-g+a'g} \quad b = b' + \frac{a'b'g}{1-g}.$$

and

$$\mu'_s = \frac{a'b'}{\delta} \quad \mu_a = \frac{(1-a')b'}{\delta} \quad \mu_s = \frac{a'b'}{(1-g)\delta}.$$

## Appendix 4: Beam and sample size

This appendix is included to show how significant the light loss problem can be. It is excerpted from Niek Van Wieringen's report *The limitations of the determination of optical properties of tissue with a double integrating sphere set-up* (1990). This describes an experiment of some relevance to the issue of selecting a beam size and an integrating sphere port size. It has a few problems, most notably in how the total light measurement was calculated, but the original data is gone so it can't be fixed.

### Introduction

One problem associated with making measurements of the optical properties of materials, is light lost out the sides of the sample holder. Lost light leads to erroneously high absorption coefficients because all lost light is attributed to absorption by the sample. The total amount of light collected is the sum of:

- the light specularly reflected out of the reflectance sphere,
- the light collected by the reflectance sphere,
- the light collected by the transmittance sphere, and
- the light transmitted unscattered out of the transmittance sphere.

As light is scattered by the sample some may leave the holder laterally (Figure 6). The amount of light lost will depend on the thickness  $\delta$  of the sample, the diameter of sphere  $d$ , the diameter of the sample port  $d_s$ , and the diameter of the beam  $d_b$ .

Reflectance and transmittance measurements with a double integrating sphere set-up were used to find the total collected light for non-absorbing samples with

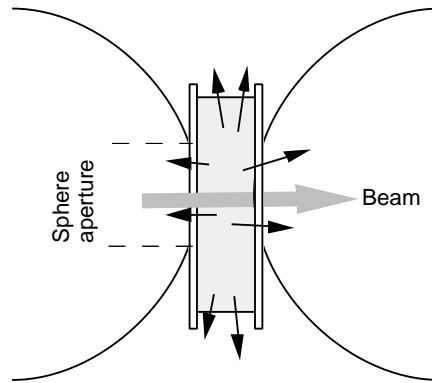


Figure 6: Light losses via the sides of the sample holder. The dashed lines indicate the diameter of the hole in the integrating sphere.



various physical thicknesses  $\delta$ , scattering coefficients  $\mu_s$ , and optical thicknesses  $\mu_s\delta$ . All experiments were done with a 1 mm HeNe laser and various concentrations of Intralipid-10% to create sample optical thicknesses that varied from zero to about 55.

## Theory

The straightforward way of figuring out the total light collected would be to add the normalized outputs from the reflectance, diffuse transmittance, and unscattered detectors. The problem is a simple normalization process ignores light losses caused by (1) the absorption by the sphere walls and (2) light exiting out ports is ignored. These effects should be taken into account when using the formulas for a double integrating sphere apparatus. This is what Niek did, but he used the formula for a single integrating sphere to correct his numbers. He also used a slightly incorrect form of this equation and therefore his results will be slightly off (I would guess no more than 5%). Rather than repeat an incorrect derivation, I'll just skip to the results.

The total light collected was measured for different combinations of the sample port size, different sample thicknesses, and different Intralipid concentrations. The results of these experiments are shown in Figure 7.



Figure 7: The total collected light versus the optical thickness of the sample. Here  $\delta$  is the physical thickness of the sample and  $d_s$  is the diameter of the sample of the integrating sphere. The beam diameter was 1 mm.

## Discussion

The total light collected has roughly the same dependence on the optical thickness for the three smaller port configurations (Figure 7).

1. When the optical thickness is zero, then there is no scattering and no light is lost. (The total amount of light does not equal 100% because the specular reflectance from the sample was not collected.)
2. As the optical thickness increases, more and more light is scattered. More and more light leave out the edges of the sample until a maximum loss is reached.
3. As the optical thickness increases, so does the optical distance in the radial direction. The radial optical thickness eventually increases to the point that the number of scattering events encountered by photons is so large that they travel only a short distance laterally and can no longer escape out the edges.
4. Eventually the radial optical distance becomes so large that no light escapes laterally and the total collected light increases asymptotically to a fixed value.

Consider the curves for a sphere aperture of  $d_s = 10$  mm. The maximum discrepancy occurs for an optical thickness of about four. When the sample thickness is 2 mm, the total collected light is always less than when the sample thickness is 0.5 mm. The relative radial path lengths are nearly a factor of four greater in the 2 mm case, and photons will encounter four times more scattering events before they might be lost. When the sphere aperture is doubled in size  $d_s = 19$  mm, then the total light collected is increased for all optical thicknesses.

Summarizing, it appears that:

- the overall collection efficiency is determined primarily by ratio of the radial optical distance to the sample optical thickness,
- the limiting value of for the total light collected as the optical thickness becomes large is primarily determined by the size of the hole in the sphere.

The second point bears closer examination. In short, the argument is because the radial optical distance is always larger than optical depth and therefore as the optical depth becomes large it may be expected that no light can escape via the sides of the sample holder.

However, *light does escape* and one mechanism<sup>34</sup> by which this can happen is via the the glass slides of the holder, as in Figure 8. The circle on the top represents light incident at all angles on a point on the liquid/glass boundary.

---

<sup>34</sup>A second mechanism might be caused by the diffuse light in the integrating sphere. Unlike the relatively small incident beam, all the light in the integrating sphere is diffuse and uniformly impinges on the entire exposed sample surface. The diffuse light that hits near the edge of the port has a roughly 50% chance of being lost. If this lossy rim area is assumed to



Figure 8: A glass slide with light incident from all possible angles from the water side. Since there is no critical angle for the water-air transition most light will not be reflected by this boundary. However, total internal reflectance will occur in the glass slide for all light incident at angles greater than  $40.5^\circ$ . This corresponds to angles in the water of greater than  $48.8^\circ$ . Thus any light incident from the water at angles larger than this stand a chance of “walking” down the inside of the glass slide.

The light that passes this boundary makes angles of at most  $59.7^\circ$  with the normal of the slide (represented by the cone inside the glass slide). At the glass-air boundary, light incident at angles larger than the critical angle  $\theta_c = 40.49^\circ$  will be reflected totally and otherwise partially. Also at the glass-liquid boundary light will be reflected. If light has been reflected inside the glass slide, a couple of times it will have travelled some distance in the radial direction. For small test apertures the possibility arises that light is conducted sideways and escapes.

This possibility is tested with the following experiment. A thick piece of white perspex, strongly scattering and weakly absorbing, is placed in front of the test aperture of a single sphere set up and percentage of light reflected is measured. This measurement is repeated, this time however with a glass slide against the perspex (‘glued’ together with a very thin layer of water in order to minimize specular reflectance). The experiment is done for two different sample

be roughly one mean free path wide, then the area is given by

$$\text{lossy rim area} = \frac{\pi d_s}{\mu'_s}$$

The loss will be relative to the total sample area  $\pi d_s^2/4$ , and consequently we would expect the light losses due to this effect to drop as

$$\frac{\pi d_s / \mu'_s}{\pi d_s^2 / 4} = \frac{4}{\mu_s d_s} = \frac{1}{\delta \mu_s} \frac{4\delta}{d_s}$$

This means that the losses should continue to decrease with increasing optical thicknesses. This was not observed.

	spot radius	$R = 5 \text{ mm}$	$R = 12.5 \text{ mm}$
without glass	$5 \pm 1 \text{ mm}$	94.3%	94.3%
with glass	$12 \pm 2 \text{ mm}$	83.5%	90.7%

Table 5: The fraction of light collected using a perspex plate with and without a glass slide in front. The spot radius is the size of the glow ball on the perspex. Both experiments used the same beam diameter of 1 mm .

port diameters. The percentage light that is reflected, is shown in table 5.

The values with and without glass slides cannot and should not be directly compared.<sup>35</sup> However, the results for different sample apertures are comparable and more light is lost when a glass slide is present. The glass slide ‘conducts’ light to the sides of the sample holder.

This is substantiated by another simple experiment that consists of measuring the diameter of the glow ball of scattered light in the perspex plate. When a glass slide is present the glow ball is more than twice the diameter measured with the perspex alone (table 5). A similar effect also appears when Intralipid is used instead of the perspex plate. Thus more light can be recovered by using a larger sample aperture. This effect is present in Figure 6, where the largest sphere aperture gathered the most light. However, for this combination the total percentage of collected light exceeds 100% . This of course is physically impossible, but it has to be taken into account that an approximation is used to determine the percentage of reflected and diffusely transmitted light. Still, this approximation is used in all four combinations and therefore a relation to one remains useful.

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<sup>35</sup>More light will be transmitted through the sample without glass slides because there are fewer boundaries that can lead to reflected light.

## Appendix 5: Beckman UV 5270 Spectrophotometer

### Introduction

This memo<sup>36</sup> is intended to describe how reflectance and transmittance experiments might be made with the Beckman UV 5270. Particular attention is paid to the idiosyncrasies and limitations of the Beckman.

First things first. The wavelength is not what it might seem and it should be understood that the number displayed by the Beckman is not the number recorded by the PC attached to the Beckman. For simplicity we<sup>37</sup> choose not to use the digital signal generated by the Beckman, but rather just took the analog signal out of the back and ran it through an A/D board. The output from the A/D board is recorded by the computer.

The measurement need to be divided into four wavelength regions. The UV (250–400 nm), the visible (400–800 nm), the near-infrared (800–1300 nm), and the less-near-infrared (1300–2500 nm). Each will be discussed in turn and skin will be assumed as the sample. However, before we do this some remarks which are applicable regardless of wavelength should be mentioned.

### General

- The Beckman should be turned on into the warm-up mode and left there for about 10–15 minutes before turning it on completely. The tungsten lamp is used exclusively except in the UV when the deuterium lamp is needed for extra photo ummph.
- For reflectance measurements, the skin sample is placed against a quartz or glass plate and water was used to match the refractive index of the skin with this plate. Oil was not used<sup>38</sup>. Before performing all measurements a 0% and 99% reflectance scan must be done. Also when doing this scan the 0% and 100% knobs must be adjusted so that the Beckman never reads below 0% or above 100% for either of these scans over the whole wavelength region. Note that the Spectralon reflectance standards should never be put in contact with oil as it renders them defective (they absorb oil, but are hydrophobic). When doing the reference scans no matching fluid or glass/quartz plate is used.
- When performing the transmittance measurements, the sample is placed between two glass or quartz or glass plates. This sample is placed in

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<sup>36</sup>Vasan Venugopalan at Wellman helped revise this subsection. So he shares the blame and the glory.

<sup>37</sup>Steve Jacques and I spent a week taking data with the Beckman back in 1986. Steve found the A/D converter and I hacked a BASIC program together that week just to get data. This is the pretty much the system that you have now.

<sup>38</sup>Why the worry about oil and water, well if you are interested in hydration effects then it could be an issue. Really, I only considered oil because it has been used on psoriatic lesions to reduce backscattering from the flakey outer layers

front of the sample beam which enters the sphere. In this case a  $\text{BaSO}_4$  plate is placed in both the sample and reference ports. The 0% and 100% transmittance measurements are performed by placing either an opaque or no sample in the path of the beam. When doing these scans the 0% and 100% knobs must be adjusted so that the Beckman never reads below 0% or above 100% for either of these scans over the whole wavelength region. For the tissue samples, water must also be included to keep the tissue sample hydrated. However when testing epidermis and dermis make sure that the amount of water added does not artificially increase the thickness of the sample. Of course for the transmittance measurements the thickness of the sample must be measured. This is done by measuring the thickness of the quartz/glass plates used with and without the sample inserted in between. You need to make several measurements in different places because most samples are not very uniform in thickness.

- These reference scans should be performed three additional times (e.g., once just before lunch, once after lunch and definitely once just before shutting the machine down for the day.) It is also advisable to scan the 10% or 20% standards at this time if they are to be used for the days measurements. The times when this is necessary will become clearer as you read the rest of this memo.

## The Ultraviolet

There are a few problems associated with this region.

- First, the Beckman illuminates with only a single wavelength and detects all light falling on the detector in the integrating sphere. This means that if light is absorbed and the generates fluorescence, much more light will be detected for that wavelength than should be. This was pointed out by Rox, in his (in)famous skin optics paper [119]. Rox used a solar-blind photomultiplier tube (Hamamatsu R456) to prevent detecting visible fluorescence. Thus the same photomultiplier should be used when making the measurements in the wavelength region 250–300 nm. (It should be in one of the drawers in the Beckman room.) Ideally these measurements should also be made with some sort of fluorimeter with the excitation and emission monochromators scanning at the identical speed and wavelength.
- The reflected and transmitted light can be very small (especially for darker skin types). The measurements are correspondingly noisy. We used a 20% reflectance plate as the reference target and adjust the 0% and 100% dials on the Beckman to effectively utilize the full range of the machine. As a result 0–100% will then correspond to a reflectance between 0 and 20%. This improves the generated signal significantly. It might be that the electronics in the Beckman just don't work so well for very small voltages.

- Because glass transmits poorly in this region, quartz plates must be used as the window against which the sample is placed for the reflectance measurement.
- Probably most troubling is that in the ultraviolet region the amount of light falling on the detector is necessarily small. The Beckman opens and closes the slits on the monochromator to ensure that the signal stays reasonably constant. This ensures good signal. Unfortunately, it also increases the bandpass. Watch the dial below the wavelength indicator to get some idea of how large the bandpass is during a typical scan. You might be surprised.
- Of course this is where you need to use the deuterium lamp.

### The Visible and Near Infrared (400–1300 nm)

- There is quite a lot of light in this region. The detector begins to fail at around 800 nm (or was it 700 nm?). Anyway the spectrophotometer must be adjusted for infra-red operation. This means that the machine must be turned off, and the setting on the sphere accessory (the lever on the back of the spectrophotometer behind the sphere) switched to IR mono sphere as well as some dinky little switch near the wavelength indicator.
- Once the machine is working in the infrared, you should be aware that the wavelengths now change four times as fast as in the visible. The bandpass indicator is likewise increased in magnitude.
- Also once we are in the infrared you must specify the higher wavelength as the starting wavelength  $\lambda_s$ . The ending wavelength  $\lambda_e$  must be

$$\lambda_e = \lambda_s - \frac{\lambda_s - \lambda_e}{4}$$

because the computer has no idea that the Beckman is operating in the infrared. For example if you want to scan from 800 nm to 2500 nm, You set the dial to 2500 nm and specify the starting wavelength as 2500 nm and the ending wavelength as 2075 nm.

- The scan speed is automatically increased by a factor of 4 once we switch into the IR. Thus scan speed should reduced by depressing the 1/4 nm/s switch from the 4/16 nm/s switch to keep the scan speed at 4 nm/s in the IR.

### The Near Infrared (1300–2500 nm)

- The reflectance of skin is very poor in this region. The strong absorption by water in skin is comparable to or exceeds the scattering coefficient. Anyway there is not a whole lot of signal.

- The solution once again is to replace the reference plate with a 10 or 20% Spectralon standard. The experiments proceed as for the ultraviolet. The absorption by glass is still pretty small and quartz need not be used.
- The 0% reflectance measurements are a problem. It turns out that the back of the sphere door reflects significant amounts of light in this wavelength range. I remove the door completely and turn off the room lights and sneak off during these calibration scans. All the scans are done in a dark room.



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