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1. Introduction

Thank you for purchasing a Woollam Spectroscopic Ellipsometer (SE) system. We hope the information contained in this manual will help you develop a better understanding and appreciation for data analysis using the CompleteEASE software.

1.1. Manual Overview

The manual is organized into the following chapters:

Chapter 1: Introduction
Chapter 1 will introduce you to ellipsometry. There is information about software installation instructions, a list of shortcut keys, a list of the “conventions” used throughout this manual, and a brief overview of ellipsometry fundamentals. In addition, sections on File Structure and User Setup have been updated. If you are upgrading to a new version of CompleteEASE, please contact the Woollam Company for a special Application Note regarding “New Features of Interest.”

Chapter 2: Data Collection
Chapter 2 provides a quick overview of data collection. As the details can vary depending on your instrument and configuration, this chapter is meant only for quick reference and formal information should be sought from the Hardware manual matching your instrument.

Chapter 3: Data Analysis 1 – Basic
Chapter 3 begins a series of chapters devoted to data analysis procedures. Each chapter steps through examples provided with this manual. Chapter 3 describes the basic SE data analysis algorithms used by the CompleteEASE software. All users can benefit from understanding how the most basic models are constructed. Examples in this chapter include:

- Oxides on Silicon
- Transparent Thin Films
- Glass substrate considerations
- Surface Roughness and Index Grading
- Logging and Reporting Results from a Series of Films
Chapter 4: Data Analysis 2 – Intermediate
Chapter 4 continues to build on principles of data analysis, focusing on absorbing materials. The examples are intended to teach a few of the advanced features within CompleteEASE. Examples in this chapter include:

- Metal Substrates
- Organic monolayers on Gold
- Absorbing Thin Films (B-SPLINE approach)
- Absorbing Thin Films (GENOSC approach)
- Combining SE and Transmission data

Chapter 5: Data Analysis 3 – Advanced
Chapter 5 continues the description of data analysis for advanced applications. Examples in this chapter include:

- Working with Uniformity Maps
- Multi-Sample Analysis
- Model Non-Idealities
- Creating Alloy or Temperature Files
- Anisotropic Thin Films

Chapter 6: Data Analysis 4 – In Situ Data
Chapter 6 details how to work with time-dependent data that is measured during deposition or etch of a film. This chapter is also useful when working with liquid or heat stages, where real-time data can be measured dependent on sample conditions.

- Working with In Situ Data
- Growth Rate and Optical Constants model

Chapters 7 through 13: Feature References
These chapters are targeted toward advanced users that need an explanation of specific topics. The chapters are organized as follows:

Chapter 7: Basic Screen Layout and Graph Panel
Chapter 8: Measurement Tab
Chapter 9: Analysis Tab
Chapter 10: Gen-Osc Functions
Chapter 11: In-Situ Tab
Chapter 12: Hardware Tab
Chapter 13: Options Tab
Appendices: Additional Information

The Appendices include additional information for your reference.

- **Appendix 14.1** Trouble-Shooting Guide.
- **Appendix 14.2** Maximum Memory Usage
- **Appendix 14.3** Importing/Exporting WVASE32 data files
- **Appendix 14.4** Comparing WVASE32 and CompleteEASE results.
- **Appendix 14.5** Contacting the J.A. Woollam Company
1.2. Installing or Upgrading the Software

Before Installation

Verify that the computer used to operate the ellipsometer system meets all the requirements listed in Table 1-1.

Table 1-1. Computer requirements for the CompleteEASE software.

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<th>MINIMUM COMPUTER SPECIFICATIONS</th>
<th>REQUIREMENTS</th>
</tr>
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<tr>
<td>Operating System</td>
<td>Microsoft Windows 98, ME, 2000, XP, Vista. 32- and 64-bit Windows 7</td>
</tr>
<tr>
<td>Memory (RAM)</td>
<td>256 MB</td>
</tr>
<tr>
<td>Hard Disk Space</td>
<td>120 MB</td>
</tr>
<tr>
<td>USB Ports</td>
<td>1 required, version 1.1 or higher</td>
</tr>
<tr>
<td>Processor</td>
<td>Intel Pentium III, AMD Athlon, or higher 1 GHz clock speed or faster **Many operations in CompleteEASE benefit from multi-core processing power.</td>
</tr>
<tr>
<td>Display</td>
<td>Color display at least 800x600 pixel resolution</td>
</tr>
</tbody>
</table>

Upgrading versions prior to 4.45 (alpha-SE)

The software driver used for the alpha-SE was changed starting with CompleteEASE version 4.47. This change was made to better support Windows sleep and suspend features and to make it easier to add support for 64-bit Windows in a future release. On computers that were already running an older version of CompleteEASE, an extra step is required to update from the old driver to the new driver. This step is also required when the alpha-SE is plugged into a different USB port if that port was used with the old driver. Complete documentation is available from the Woollam Company that describes how to perform the update.

Running the SETUP Program

Run the “CompleteEASE_Setup_5.xx.EXE” application from the provided installation CD. This program is located on the CD along with a folder named “CNF” that contains the configuration files to match one specific ellipsometer system. Setup will install the CompleteEASE program files, the Java Virtual Machine (VM) files which are required to run CompleteEASE, and other related files (material optical constants files, optical model files, example files, etc.).
NOTE: The CompleteEASE_Setup program will not automatically install the calibration and configuration files for your specific instrument from the “CNF” directory. This is done at the factory and if your files become corrupt, you will need to copy all files from this folder into the corresponding C:\CompleteEASE\CNF\ folder after installation. These files will return the instrument to the factory setting and replace any calibrations that have been recently performed. Please contact JAWCo for further advice if this is required in the field.

Follow the onscreen prompts provided by the CompleteEASE_Setup program. The default directory for installation is C:\CompleteEASE. To minimize compatibility problems, it is recommended to install the software in this directory. This is shown in Figure 1-2. The installation CD includes configuration files for one specific ellipsometer system. If you are installing the CompleteEASE software to run this ellipsometer system, choose “Full Installation” as shown in Figure 1-3. This will install the drivers required for hardware operation. It is still required to copy the CNF files after installation, as discussed above.

The CompleteEASE software can be installed on multiple computers for data analysis purposes (most systems include 5-site license). If you are installing software on a computer that will not operate an ellipsometer system, choose “Data Analysis Only”, as shown in Figure 1-3.

NOTE: The CompleteEASE software is only compatible with the encoded Experimental Data files measured using CompleteEASE. Thus, you can install CompleteEASE at multiple computers, but only for use with data collected by CompleteEASE.

Figure 1-1. License Agreement for CompleteEASE.
Figure 1-2. Default folder location for CompleteEASE installation.

Figure 1-3. CompleteEASE Setup choices. The software can be installed for use with an ellipsometer system or for data analysis only.

The next message you will receive asks which tasks you would like the installation software to perform. The choice ranges from creating icons to associating CompleteEASE file types. These possibilities are shown in Figure 1-4. If you choose, a shortcut icon (Figure 1-6) will be placed on the Windows Desktop to start the CompleteEASE program. However, do **NOT** start CompleteEASE until
following the system connection instructions. The software installation process should take less than two minutes on most computers.

**Figure 1-4.** Additional Tasks desired during installation of CompleteEASE.

**Figure 1-5.** Details just before installation.
Operating Hardware

If you will be operating hardware with CompleteEASE, it requires the configuration files for your instrument. These are provided with the installation CD that accompanies each instrument. If upgrading to a newer version of CompleteEASE, the configuration files should already be installed from the factory and do not require any additional efforts. If you are installing CompleteEASE on a new computer to operate the hardware, or repairing a corrupted version, you can copy the contents of the "CNF" folder from the CD provided with the instrument to the associated folder on your hard drive: C:/CompleteEASE/CNF/.

NOTE: You will not want to copy configuration files from the CD if your system is already operating with CompleteEASE, as this will overwrite your latest calibration and return it to factory settings. This may require recalibration of your instrument. Please contact the Woollam Company for further details.
1.3. Conventions Used in this Manual

The CompleteEASE software is written in JAVA with a Tab-interface to conveniently organize software features. Figure 1-7 shows the front screen of CompleteEASE.

Figure 1-7. CompleteEASE Software

To help the user navigate this manual, the following conventions will be followed to describe features in the CompleteEASE software.

Tabs

When the software is used with an alpha-SE system, there are typically four Tabs, as shown across the top of the software screen: Measurement, Analysis, Hardware, and Options. An additional In Situ tab is available for M-2000 and RC2 systems that may be applied to real-time measurements. Throughout this manual, Tabs will be written in bold and italics.

Screen Panels

Within each Tab, there are screen areas referred to as panels. From the Measurement tab shown in Figure 1-7, the panels include System Status, Measurement Controls, and Fit Results. Throughout this manual, panels will be written in bold, blue text.
Buttons

Figure 1-8 shows the **Hardware** tab for CompleteEASE. There are six panels: **Hardware Status, Controls, Alignment, Other, Routine Test Measurement** and **System Information**. Within some panels are various buttons. Buttons will be designated with single ‘quote’ marks, such as ‘Align Sample’ and ‘Display Signal’.

![Figure 1-8. Hardware Tab, showing Screen Panels and Buttons.](image)

Model Commands

Figure 1-9 shows the **Analysis** tab. Within the **Analysis** tab, there is a **Model** panel. The model includes many choices – some of which are shown in bold, underlined, red letters. Within this manual, they will also be shown in bold, underlined, red letters. For example, **Add**, **Delete**, and **Save**. Model details are further divided into expandable sections, shown in bold letters preceded by a red + or - symbol. The symbol preceding the text shows whether the section is collapsed (+) or expanded (-). These sections will be listed using bold text with a red + symbol: **+MODEL Options**, **+FIT Options**, and **+OTHER Options**.

![Figure 1-9. Analysis Tab showing the Model section.](image)
Mouse-Selected Menus

The CompleteEASE software also utilizes both left- and right- mouse-buttons to access additional menus and features. Throughout this manual, if you need to select the right or left mouse button, you may see the following symbols: (Right) or (Left), respectively. Figure 1-10 shows the menu when the right mouse button (Right) is pressed within the **Fit: results** box.

![Figure 1-10](image.png)

**Figure 1-10.** Right-clicking mouse (Right) in Fit results section (white area) brings up a special menu related to the Fit.

Pipelining Descriptions

The descriptions used in above sections can be “pipelined” to form a convenient description of software features or menus. In this case, the description will take on a similar form as:

```
Tab>Panel>‘Button’
```

For example, the ‘Open’ button within the **Model: Panel** under the **Analysis** tab is conveniently described as:

```
Analysis>Model>‘Open’
```

As another example, the command to right-click the mouse within the **Fit: Panel** from the **Analysis** tab and then choose the “Add to Fit Log” menu choice would read as follows:

```
Analysis>Fit>(Right)>‘Add to Fit Log’
```
1.4. File Structure and File Types

The CompleteEASE software is installed on your hard-drive in a single directory. However, it is helpful to understand the sub-directories used by CompleteEASE. Figure 1-11 shows the sub-directories within the CompleteEASE folder.

![Figure 1-11. Structure within CompleteEASE folder.](image)

The following table provides more details about each of these subdirectories and their function within CompleteEASE.

<table>
<thead>
<tr>
<th>DIRECTORY</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>cnf</td>
<td>Hardware configuration files</td>
</tr>
<tr>
<td>DAT</td>
<td>Common Data files, including a sub-folder of Examples that are shipped with the CompleteEASE software. Any data files saved in this directory can be viewed and opened by all users, but are read-only. This directory is not intended for individual user files.</td>
</tr>
<tr>
<td>HTML Clipboard</td>
<td>Files and Graphs used by CompleteEASE to construct reports within the HTML Clipboard.</td>
</tr>
<tr>
<td>jar</td>
<td>Java Application files</td>
</tr>
<tr>
<td>MAT</td>
<td>Common Material Files to describe optical constants of materials you may encounter. This folder is further divided into: Advanced, Basic, Dielectric, Metal, and Semiconductor folders. Files within this directory can be viewed and opened by all users, but are read-only.</td>
</tr>
<tr>
<td>MOD</td>
<td>Common Model Files to describe a sample and the associated analysis strategy. Also location for Common snapshots. Further divided into Advanced, Basic, and Calibration Wafers.</td>
</tr>
<tr>
<td>Recipe</td>
<td>Common Recipe Files describe a complete measurement and model strategy. This is pertinent to variable angle or variable measurement point systems and is not used with the alpha-SE system.</td>
</tr>
</tbody>
</table>
Although the default directories will always be used by CompleteEASE, the user can create their own directories to store Data files, Models, Materials, Snapshots, and Recipes. To maintain the default files, any sub-directories created within the primary Directories (as listed in Table 1-2) will be treated as “read-only”. For example, the three subdirectories within MOD (Advanced, Basic, and Calibration Wafer) are read-only.

**COMMON Location**

CompleteEASE uses a “COMMON” location to store files that may be used by multiple users, or as a location where the software can always locate certain files needed to perform operations. For example, a Recipe may instruct CompleteEASE to run a measurement based on Acquisition Parameters that are saved in a file and then fit the data based on a Model. CompleteEASE needs to be able to locate both of these files in order to complete the Recipe – so they either have to be stored in the exact same folder as the Recipe or in the COMMON folder location.

The actual location of the COMMON folder is different depending on the file type. Table 1-3 lists the COMMON folder locations for material files, models, data files and recipes. Snapshots share the same COMMON folder as Models.

Table 1-3. Location of COMMON folders for different files types.

<table>
<thead>
<tr>
<th>Using CompleteEASE</th>
<th>Using Windows Explorer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Common Materials</td>
<td>C:\CompleteEASE\MAT\</td>
</tr>
<tr>
<td>Common Models</td>
<td>C:\CompleteEASE\MOD\</td>
</tr>
<tr>
<td>Common Data</td>
<td>C:\CompleteEASE\DAT\</td>
</tr>
<tr>
<td>Common Recipes</td>
<td>C:\CompleteEASE\Recipe|</td>
</tr>
</tbody>
</table>

CompleteEASE always maintains a folder link to the COMMON folder location, as shown in Figure 1-12 for Material files. When you select to Open or Save a File, the “Projects” tab of the Open/Save dialog box will have the COMMON folder listed.

Figure 1-12. Structure within the MOD directory along with the associated Open Model dialog box.
File Types

There are many different file types used within CompleteEASE. Table 1-4 provides a list of different file types and their corresponding file extensions.

*Table 1-4. List of File Types used by CompleteEASE and their corresponding extensions.*

<table>
<thead>
<tr>
<th>Type</th>
<th>Extension</th>
</tr>
</thead>
<tbody>
<tr>
<td>Experimental $\Psi,\Delta$ data</td>
<td>*.SE</td>
</tr>
<tr>
<td>In-Situ Experimental $\Psi,\Delta$ data (multiple times)</td>
<td>*.ISE</td>
</tr>
<tr>
<td>Model</td>
<td>*.mod</td>
</tr>
<tr>
<td>Material file (optical constants)</td>
<td>*.mat</td>
</tr>
<tr>
<td>Snapshot (Exp. data, model and fit results)</td>
<td>*.SEsnap</td>
</tr>
<tr>
<td></td>
<td>*.$SS$ (in old version)</td>
</tr>
<tr>
<td>Data Acquisition Parameters</td>
<td>*.pars</td>
</tr>
<tr>
<td>Mapping Pattern</td>
<td>*.scan</td>
</tr>
<tr>
<td>Recipe points to i) Acquisition Parameters, ii) Mapping Pattern and iii) Model</td>
<td>*.recipe</td>
</tr>
</tbody>
</table>
1.5. File Dialog

CompleteEASE has a custom File Dialog to help users organize and locate their files and folders. The key features are listed below:

- Tab for Recent Folders and Recently Opened Files
- Tab for Projects to list the most pertinent folder locations. These folders can be located anywhere on your computer, but are brought together in a single location within CompleteEASE
- Tab for Library files provided by J.A. Woollam Co.
- COMMON folder location is always shown
- Drag-and-drop files between folders
- Clear the listed folders without deleting the actual folders from your computer – keeps your workspace clean and pertinent.
- Maintain different folder and file lists for each user (based on login).

Figure 1-13 shows the File Dialog for the Material files. A new Tab-structure on the left-side has been added to provide more options on how to maintain your files and folders.

![Figure 1-13. New File Dialog Box.](image)

The Tab you select from this dialog box (Recent, Projects, Library) will remain “sticky” for each file-type. For example, if the Library tab was last selected in the Model Open dialog, that tab will remain selected the next time the Model Open dialog is shown.

In addition, the folder link will also remain “sticky” for each file type. For example, if “JAW Data” folder is selected in the Model Open dialog, then it will be displayed
when this dialog box is shown next. This allows the user to return quickly back to the same location again and again when working on the same project.

**Folder “Links”**

The folders listed in the Open and Save File Dialog boxes of CompleteEASE are actually “links” to folders on your computer. The actual folders can be located anywhere on the computer. In this manner, CompleteEASE helps organize the list of pertinent folders from various locations. The actual folder location is listed in lighter text just after the Folder name, as shown in Figure 1-14.

---

**Figure 1-14.** Folders listed in CompleteEASE are actually “links” to folders on your computer. Their actual locations is shown in lighter text to the right of the Folder name.

**Recent**

Each of the file dialog boxes used for opening or saving files contains a “Recent” tab with both a “Recently Opened Files” list and a “Recent Folders” list. Both enable the user to quickly find files and folders where they have recently worked. Each list contains the last 10 files or folders that were visited (excluding the COMMON folders). Figure 1-15 shows an example Recent file location list in CompleteEASE. The three yellow folders mark the three most recent locations where files have been opened. Library folders are not added to this list, as they are always available from a separate tab. The list of the most recent files is shown in the right-hand section of this dialog box when the top line, “Recently Opened Files”, is selected.

---

**Figure 1-15.** Recent File Locations Tab
**Add to Projects List**

If you right-click over a folder in the Recent list, you have two options that appear, as shown in Figure 1-16. Press “Add to Projects List” to include this same folder link in the Projects tab. There are two primary reasons you may want to do this: i) the folder will be regularly accessed, or ii) you would like to create or view sub-folders from this location. The Recent folder list does not allow the folders to be expanded if there are sub-folders below. This is done to maintain a “clean” list.

![Figure 1-16. Right-click on a Folder in the “Recent” list to Add that folder to the Projects list or to Clear the Recent Folders list.](image)

**Clear Recent Folders List**

The second right-click menu, which will show up no matter where you right-click within the “Recent” tab, is to clear the recent folder list. This does not affect the Recently Opened Files list.

**Browse File System**

**NOTE:** Use this button to open a single file.

If you would like to open a new file, you can do this by pressing the ‘Browse File System’ button from within the Recent folder tab. Please note that this button is used to open a file, but not necessarily to add a folder location (although the folder where the file is located will be added to the Recent list). When you press this button, a dialog box, as shown in Figure 1-17, will appear and allow you to navigate around your computer to locate the file to open.
**Projects**

The Projects tab contains a list of folder-links that are regularly visited. The COMMON folder is always listed on the top. Unlike the Recent tab, the folders listed here will not change order and will not “fall-off” the bottom of the list when you reach ten folders. In addition, each of these folder-links is expandable to view subfolders up to 3 levels deep. This is demonstrated in Figure 1-18, where each of the folders that has an arrow (▼) to the left can be expanded to show sub-folders. One of these folders has been expanded to show two sub-folders and one of these sub-folders also contains an arrow. This means it could also be further expanded.

**Remove From List**

The Project folders can be removed from the list by right-clicking on the Folder Link and selecting “Remove From List”, as shown in Figure 1-19. This will not delete the
folder from your computer; it only removes the link between CompleteEASE and that folder.

![Image of CompleteEASE software interface showing the Projects tab]

Figure 1-19. Right-click on a Folder in the “Projects” list to Remove that folder from the list or to create sub-folders within that Project folder.

**Create New Sub Folder**

In addition, you can create new sub-folders within the Projects tab by right-clicking on a Folder Link and selecting “Create New Sub Folder”. This will create the new sub-folder on your computer – wherever the original folder is located. This right-click menu is also shown in Figure 1-19.

**Add Folder Link**

NOTE: Use this button to Link to a Folder on your Computer.

Because the Projects tab is a listing of Folders that can be accessed, the button at the bottom of this tab is called “Add Folder Link”. This is different than the button at the bottom of the Recent tab (which was used to open a specific file). Press the ‘Add Folder Link’ button to add a specific folder link to the Projects tab. The file of interest can be opened after the Folder has been linked. The “Add Folder Link” dialog box is shown in Figure 1-20.
Figure 1-20. “Add Folder Link” dialog box opens up this view to search for a Folder to add to this list.

Library

The Library tab contains the read-only folders located within the COMMON folder location, as described in the previous section. For example, the three folders in Figure 1-21 (Advanced, Basic, and Calibration Wafers) are all located within the C:\CompleteEASE\Mod directory and contain common models provided with the software. The Library folder is only available for Models, Materials, Data, and Recipes.

Figure 1-21. Library File Location Tab.
The Library Tab is disabled when saving files, as it is not intended for users to add additional files into these read-only locations. However, additional folders can be created in Windows Explorer and populated with special files that you may choose to archive and wish to prevent users from changing (as they will be read-only files and folders). For example, five additional folders were added into the C:\CompleteEASE\Mod location in Figure 1-22 and they show up in the Library tab. These folders are read-only and no files can be added into these folders from within CompleteEASE. Files could be added or deleted from Windows Explorer.

**Opening a File from Windows**

Double-click to open data and snapshot files from Windows Explorer. This will add the folder location for that file to the Recent tab. It will also add the file into the Recently Opened Files list. This can be a convenient way to work with files. However, it only provides access to that single folder – no access to subfolders. To search sub-folders, right-click to move this folder link to the Projects tab. From the Projects tab, subfolders up to 3 total levels are shown.

**Adding Folder “Links” (Drag-and-Drop)**

It is convenient to add new folder “links” to CompleteEASE using Windows drag-and-drop features. To demonstrate, a folder is located within Windows Explorer and we left-click (L) on the folder name and hold the mouse button down while moving the folder over to the CompleteEASE File Location are. When the mouse button is let go in the Project Tab, the folder will appear in CompleteEASE. This is not the actual folder, but rather a “link” to the folder location on your computer. You can drag folders into either the Recent or Projects list, but not the Library.
Figure 1-23. Copying folders into CompleteEASE using drag-and drop.
1.6. Setting Up Multiple Users

A spectroscopic ellipsometer is often used by multiple users. In this case, it is more convenient to allow each user to establish a personal set of folder “links”. However, this requires Login of each user. To configure CompleteEASE for multiple users, make the following changes.

Go to the Options tab and select the ‘Edit Configuration’ button. The Configuration Parameters are divided into different areas. Expand the “General” section, as shown in Figure 1-24, and make the following changes:

1. Change “User Log-in Mode” to Must Log In.
2. Change “Users Have Own Folders List” to True, by selecting the parameter ON check-box.
3. OPTIONAL: You can decide whether to have all CompleteEASE folders visible to all users by changing the “Always Add CE Folder” to True or False. These are all folders within the CompleteEASE directories on your computer and would show up for all users if this is True.

NOTE: After completing these steps, you must restart CompleteEASE before the changes take effect.

After configuring CompleteEASE for multiple users, you will want to setup each User with their own account. Press Options>Miscellaneous>‘Manage Users’ and the User Management box will appear, as in Figure 1-25. Press ‘Add New User’ and type in the name and password for each user you wish to add.
NOTE: If the password area is left blank, no password will be required to log-in.

Figure 1-25. Setting up Multiple User Accounts.

After entering each of the Users with their own account, you should close CompleteEASE and restart the program. This will allow all of the changes to take effect. When you restart CompleteEASE, it will open by asking to enter your User Name and Password. The user names can be found in the drop-down list, as shown in Figure 1-26.

Any folders that are added by an individual user at this point will be shown for their account only. However, if the folder is located within the CompleteEASE directory, it will show up for all users. For this reason, it is preferred to create a folder outside of the C:\CompleteEASE directory for files that individual users want to access. For example, a separate folder can be created such as C:\Measurements. Within this folder, each user can have their own sub-directory and CompleteEASE can be configured by each user to link to only the folders they need to view.
1.7. Shortcut Guide

There are two types of shortcuts within CompleteEASE. First, like most software, if a letter is shown that is underlined, the shortcut for that command is to hold ALT button and press that letter. For additional shortcuts that are more global, incorporate the CTRL, SHIFT, and ALT buttons. These additional shortcuts are listed in Table 1-5.

Table 1-5. Shortcut reference.

<table>
<thead>
<tr>
<th>SHORTCUT</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTRL+M</td>
<td>Go to Measurement Tab</td>
</tr>
<tr>
<td>CTRL+A</td>
<td>Go to Analysis Tab</td>
</tr>
<tr>
<td>CTRL+I</td>
<td>Go to In situ Tab</td>
</tr>
<tr>
<td>CTRL+H</td>
<td>Go to Hardware Tab</td>
</tr>
<tr>
<td>CTRL+O</td>
<td>Go to Options Tab</td>
</tr>
<tr>
<td>CTRL+P</td>
<td>Graph the Psi data curves</td>
</tr>
<tr>
<td>CTRL+D</td>
<td>Graph the Delta data curves</td>
</tr>
<tr>
<td>CTRL+N</td>
<td>Graph the “N” data curves (N,C,S format)</td>
</tr>
<tr>
<td>CTRL+C</td>
<td>Graph the “C” data curves (N,C,S format)</td>
</tr>
<tr>
<td>CTRL+S</td>
<td>Graph the “S” data curves (N,C,S format)</td>
</tr>
<tr>
<td>CTRL+1</td>
<td>Graph the &lt;e1&gt; data curves</td>
</tr>
<tr>
<td>CTRL+2</td>
<td>Graph the &lt;e2&gt; data curves</td>
</tr>
<tr>
<td>CTRL+T</td>
<td>Graph the Intensity data</td>
</tr>
<tr>
<td>CTRL+Z</td>
<td>Graph the Depolarization data</td>
</tr>
<tr>
<td>CTRL+SPACE BAR</td>
<td>Switch view from single-point to Map/Dynamic</td>
</tr>
<tr>
<td>CTRL+L</td>
<td>Add to Fit Log</td>
</tr>
<tr>
<td>CTRL+ALT+L</td>
<td>View Fit Log</td>
</tr>
<tr>
<td>CTRL+R</td>
<td>Copy Analysis Report to Clipboard</td>
</tr>
<tr>
<td>CTRL+Y</td>
<td>Toggle between single and double-Y axis.</td>
</tr>
<tr>
<td>CTRL+ALT+O</td>
<td>Toggle Default Optical Constant units between “e1 &amp; e2” and “n &amp; k”</td>
</tr>
<tr>
<td>CTRL+ALT+W</td>
<td>Toggle Default wavelength units between “nm” and “eV”</td>
</tr>
<tr>
<td>CTRL+ALT+S</td>
<td>Show graph statistics</td>
</tr>
<tr>
<td>CTRL+ALT+SHIFT+I</td>
<td>Add the In Situ Tab</td>
</tr>
<tr>
<td>SHIFT+Mouse Roller</td>
<td>When positioned over model parameter, it will increase or decrease this parameter.</td>
</tr>
<tr>
<td>CTRL+SHIFT+Mouse Roller</td>
<td>Same as SHIFT-Mouse roller, but with smaller parameter increments.</td>
</tr>
</tbody>
</table>
If you are working with Uniformity Mapping data, the following shortcuts can help manipulate the data and graphs.

Table 1-6. Shortcut reference for mapping data.

<table>
<thead>
<tr>
<th>SHORTCUT</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTRL+Click on Point</td>
<td>Select/De-Select Point</td>
</tr>
<tr>
<td>CTRL+ALT+Click on Point</td>
<td>Delete Point</td>
</tr>
<tr>
<td>CTRL+ALT+SHIFT Click on Point</td>
<td>Show camera image from point (when available)</td>
</tr>
<tr>
<td>CTRL+SPACE Bar</td>
<td>Switch view from single-point to Map</td>
</tr>
</tbody>
</table>

Finally, when working with in-situ data or multi-sample analysis, there are additional shortcuts to help quickly switch between different data sets and models.

Table 1-7. Shortcut reference for mapping data.

<table>
<thead>
<tr>
<th>SHORTCUT</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTRL+(F1, F2, F3...)</td>
<td>Switch between models 1, 2, 3...</td>
</tr>
<tr>
<td>ALT+(1,2,3...)</td>
<td>Switch between data sets 1, 2, 3...</td>
</tr>
</tbody>
</table>
1.8. What is Spectroscopic Ellipsometry?

Ellipsometry is a non-destructive optical technique in which the sample to be characterized is illuminated with a beam of polarized light, as shown in Figure 1-27. Ellipsometry measures the change in polarization state of the measurement beam induced by reflection from (or transmission through) the sample. The change in polarization state is commonly characterized by the ellipsometric Psi (Ψ) and Delta (Δ) parameters defined in Eqn. (1-1):

\[ \tan(\Psi) \cdot e^{i\Delta} = \rho = \frac{r_p}{r_s} \tag{1-1} \]

In this equation, \( \rho \) is defined as the ratio of the reflectivity for p-polarized light \( (r_p) \) divided by the reflectivity for s-polarized light \( (r_s) \). \( \rho \) is a complex number, and the ellipsometric parameters simply report this value in polar form: \( \tan(\Psi) \) is the magnitude of the reflectivity ratio, and \( \Delta \) is the phase. Ellipsometry offers three main advantages over simple intensity-based reflection or transmission measurements:

1. **Precision** As the change in polarization state is defined by a ratio, ellipsometry is not sensitive to changes in the absolute intensity of the measurement beam (i.e., the sample itself is the ‘reference’ for the measurement). Thus, it can remain accurate without knowing “absolute” intensity values.

2. **Sensitivity** The phase information contained in the ellipsometric Δ parameter provides enhanced sensitivity to ultra thin films, even down to the sub-nm thickness level.

3. **Information** Ellipsometry measures 2 values (Psi and Delta) at each wavelength, doubling the information content compared to an intensity reflection or transmission measurement.

The high accuracy, precision, and sensitivity of the ellipsometric measurement make it highly suited for demanding thin film metrology applications.

![Figure 1-27. Interaction of polarized light with a sample.](image)

In Spectroscopic Ellipsometry (SE), Psi and Delta values are acquired as a function of wavelength. This greatly increases the information content in the data set, enabling the simultaneous determination of multiple sample properties. However, to extract sample parameters such as film thickness and optical constants from the measured SE data set, an optical model must be built to fit the data. The CompleteEASE software provides a graphical user interface for building models and displaying measured data and model fits. The fundamentals and theory of data...
For more information on ellipsometry theory, see the following references:

2. Data Collection

The first step to any ellipsometry experiment is collection of data with the ellipsometer. Measurement operations are better discussed in the hardware manual for each instrument. CompleteEASE is used with alpha-SE, M-2000, RC2, and AccuMap-SE ellipsometers from the Woollam Company. The details for data collection depend on both the system type and individual ellipsometer configuration. This chapter provides a brief overview of CompleteEASE data collection. More details are provided in Chapter 8, which lists all features of the Measurement Tab, and in the corresponding Hardware manual for your system.

NOTE: Instructions in this chapter are representative of various systems. For more complete instructions, including details on system alignment and calibration, please consult your corresponding Hardware manual.

2.1. alpha-SE Systems

CompleteEASE uses a simplified screen for data collection with the alpha-SE, as there are limited angles of incidence and no mapping to consider. The Measurement tab for an alpha-SE will appear as in Figure 2-1.

![Figure 2-1. Measurement tab for alpha-SE systems.](image-url)
There are five choices to make before collecting data. The user must select the i) measurement “Mode”, ii) type of “Sample Alignment”, iii) measurement Angles, iv) “Model” to use for data analysis, and v) whether to “Save Data after Measurement”. Three of these options provide a drop-down menu, as shown in Figure 2-2.

![Figure 2-2. Options for measurement Mode, Sample Alignment, and Model.](image)

For the majority of measurements, the “Standard” Mode and “Standard” Sample Alignment are used. Details regarding other choices are provided in Chapter 8.

### Measuring a Sample

Once the Measurement choices are selected, you can press the ‘Measure’ button to begin, as seen in Figure 2-3. The software will prompt to change angle if needed. As the alpha-SE has automated sample alignment, the system will align the z-height to detect reflected light from the sample (using the selected Sample Alignment) and then collect measured data (using the selected Mode). Finally, if a Model was selected, the data will be analyzed to determine the resulting film properties.

![Figure 2-3. Pressing ‘Measure’ will start the data collection procedure. The System Status will be updated to show the current hardware operation.](image)

If the “Save Data after Measurement” check-box was selected, a dialog box will appear, as shown in Figure 2-4. Find the location you wish to save the file within the File Location: area and type in a File Name and Comment. If the folder “link” does not exist, you can create a new folder or add a new link. These instructions were detailed in Section 1.4.

**NOTE:** CompleteEASE data files are encrypted & saved with “.SE” or “.iSE” extensions. These files are only accessible by the CompleteEASE program.
If a model was selected prior to measurement, the model analysis will be performed after SE data is acquired and the fit parameters displayed in the **Fit Results** panel. The model fit to the measured data is also displayed in the graph: the black dashed “Model” curves should lie essentially on top of the colored Psi and Delta measured data curves if the model fits the data well. If this is not the case, the wrong model was selected to analyze the data. The final result for a thin oxide on silicon is shown in Figure 2-5. The model parameters will be reported in the **Measurement>Fit Results** section, while a graph of Psi and Delta versus wavelength are displayed at the bottom of CompleteEASE – along with corresponding Fit curves from the model.

If no model was selected during measurement, the **Fit Results** panel will display “No Results Available.” The data can be analyzed later from the **Analysis** tab.
2.2. M-2000, RC2, and AccuMap-SE Systems

For M-2000, RC2 and AccuMap-SE systems, there are more data collection choices. This chapter will discuss the basic options for typical systems. As there are many different configurations, it is suggested you consult your Hardware manual for the details of your system and configuration. Figure 2-6 shows a representative CompleteEASE Measurement tab for an M-2000 with mapping and camera. Due to the number of choices for any measurement (how to measure, where to measure, etc.), CompleteEASE uses a “Recipe” to describe all details. These recipe files can be created and saved for any specific measurement or sample type.

![Measurement tab for an M-2000 system with mapping and camera. Due to the number of choices for any measurement (how to measure, where to measure, etc.), CompleteEASE uses a “Recipe” to describe all details. These recipe files can be created and saved for any specific measurement or sample type.]

Measurement Recipe

The measurement controls for this type of system are integrated into a “Recipe”. Each Recipe describes the three basic components of data collection: 1) Data Acquisition, 2) Mapping Scan settings, and 3) Modeling.

The drop-down box from Measurement>Measurement Controls: Recipe allow the following options: 1) Prompt for Acquisition Parameters, 2) Prompt for Recipe Components, 3) list of all Common recipes, 4) Choose From File Dialog, and 5) Create/Edit Recipe. These options are shown in Figure 2-7.
To get started without a full recipe, you can by-pass the Scan Pattern and Model, by selecting <Prompt for Acquisition Parameters>. This assumes you only want to measure at a single point and save data for modeling at a later time. You will be allowed to specify the Data Acquisition Parameters, from a box similar to shown in Figure 2-8. The options listed in the Acquisition Parameters box are described in Chapter 8.

<Prompt for Recipe Components>

When you are ready to map your sample at multiple locations and/or you know the model you would like to use for data analysis, you can choose these options as part of the overall recipe. Selecting <Prompt for Recipe Components> allows the user to select the individual recipe components, which include Acquisition Parameters, Scan Pattern, and Model – as shown in Figure 2-9. If the recipe components are not established, they can also be created from the Recipe Components dialog box.
‘Choose From File Dialog’

This button allows you to choose from an existing recipe previously saved by navigating to the file.

‘Create/Edit Recipe’

This button allows you to create or edit a recipe. This selection opens the same box as shown in Figure 2-9. Recipe options will be described later in this chapter.

The three primary components of any recipe are the Acquisition Parameters, Scan Pattern, and Model; as shown in Figure 2-10. Each of these components is discussed in further detail in the following sections.
2.3. Acquisition Parameters

The Acquisition Parameters describe how the data will be collected at each point of a Measurement Recipe (whether single-point or scan). A common box for Acquisition Parameter choices is shown in Figure 2-11. The options available will depend on your system configuration. The individual details for these options are discussed in the corresponding hardware manual.

![Figure 2-11. Acquisition Parameters Setup for M-2000 and RC2 systems. Choices listed above can vary from system-to-system.](image)

Data Type

The Data Type specifies the type of data to collect during the measurement, as shown in Figure 2-12. For 95% of applications, the “Standard” option will be selected.

![Figure 2-12. Different Options for Data Type.](image)

Sample Alignment

Sample Alignment options depend on the type of system and configuration. In general, there are two basic steps to sample alignment: Tilt alignment and Height alignment. Common selections for each are described in this section.
**Tilt Alignment**

Tilt alignment refers to the tip and tilt of sample relative to the incoming light beam such that the reflected beam is aligned on the receiver unit at the correct angle of incidence. Tilt alignment options include Skip, Manual, or Automated; as shown in Figure 2-13.

![Figure 2-13. Options for Sample Tilt Alignment.](image)

The Tilt alignment is often skipped when using focusing probes, as the tip-tilt alignment becomes less sensitive. It can also be skipped for applications where common samples are measured that remain flat. For example, if a system is continuously used to measure 200mm silicon wafers, the tip-tilt alignment of each wafer will probably be very similar from sample-to-sample.

For Manual or Automatic Alignment, the system will study the reflection of a light beam off the sample that is incident on a 4-quadrant detector. This detector allows the beam to be centered onto the 4-detector elements by adjusting tip and tilt until all 4 detectors have the same intensity. This method allows very repeatable alignment of the beam from sample-to-sample. An example of the Tilt Alignment step for an Automatic System is shown in Figure 2-14.

![Figure 2-14. Sample Tilt Alignment Window.](image)
Sample Height Alignment

Sample Height Alignment refers to the adjustment of ellipsometer beam relative to the surface of a sample to ensure correct positioning and adjustment for samples of different thickness. This adjustment becomes increasingly important for systems with smaller spot size (like Focusing ellipsometers). It is also more important to test the Sample Height Alignment when many samples of different thickness will be tested on the same Ellipsometer. The common choices for Sample Height Alignment are shown in Figure 2-15. For Automated Height Alignment, the reflected beam is detected by the Receiver unit and the intensity is recorded versus different Z-Height positions. This is demonstrated in Figure 2-16. The position of highest intensity is assumed to be the correct Z-Height for alignment of the sample.

![Sample Height Alignment options](image)

**Figure 2-15. Options for Sample Height Alignment.**

![Sample Height Alignment window](image)

**Figure 2-16. Sample Height-Alignment window.**
2.4. **Scan Pattern**

Scan patterns are available for systems with automated sample translation. The Scan Pattern is used to describe where each measurement should be performed across a sample. To make a new Scan Pattern, press ‘Edit/Create’ button to the right of the Scan Pattern selection of the Recipe Components box, as shown in Figure 2-17.

![Choose Recipe Components dialog box](image)

**Figure 2-17. Choose ‘Edit/Create’ Scan Pattern from the Recipe Components dialog box.**

The Scan Pattern Editor, shown in Figure 2-18, allows you to describe both Circular and Rectangular samples. There are many options to automatically fill-in the points to measure, including Grid Fill, R-T Grid Fill, and Line Fill. Most of these selections are intuitive and best learned by testing each button a few times.

![Scan Pattern Editor](image)

**Figure 2-18. Scan Pattern Editor**
2.5. Model

A model is a description of how the data will be “fit” by CompleteEASE to determine film thickness and optical constants (or other material properties). Models will be described in significant detail in the upcoming chapters, with examples of how to work with many common models. You can select any model of interest to analyze the data during the recipe. For scans of multiple points, the model will be used to analyze data from each point.

Basic Models

For convenience, CompleteEASE has a list of basic models in the “Basic” folder link. These include transparent or absorbing films on silicon or glass substrates, as shown in Figure 2-19.

Figure 2-19. From the Analysis tab, press ‘Open Model’ and choose the “Basic” Folder Link on the left – this provides access to many starting models on either glass or silicon substrates.
2.6. Running a Recipe and Viewing Results

After choosing the components of a Recipe, press the ‘Measure’ button. CompleteEASE will collect data per the Acquisition Parameters at each of the locations described in the Scan (if applicable). After each measurement, the data will be analyzed using the selected Model. An example of this process is shown in Figure 2-20. The System Status will show the current hardware operation. The Fit Results will be updated with the model fit results from each point. The Scan Map will show the current status of a pattern map (if applicable) – green points are already measured, blue is currently being measured, and red are yet to be measured. Finally, the Graph will show the data being collected from the most recent point.

![Figure 2-20. CompleteEASE during a Recipe measurement with multiple points.](image)

For maps of multiple points, the Graph can be toggled to Graph the parameters for all points measured up to the current point by pressing the ‘Graph Parms’ button within the Fit Results section. The parameter to graph can be selected from a drop-down list above the Graph, as shown in Figure 2-21.
Figure 2-21. Viewing Parameter Results of Scan during Measurement.

When the Scan is completed, a table of results will be presented in the Fit Results panel, as shown in Figure 2-22. The results of any model that are run during recipe are saved with the measurement data and can be opened at a later time by pressing the ‘View Previous Results’ button from the Fit Results panel. This opens the window shown in Figure 2-23.

Figure 2-22. Final Results after Scan.
Figure 2-23. Viewing Results after measurements are completed.
3. Data Analysis 1 – Basic

We have found that the best way to learn software features is through a series of real-world examples, where the user can follow along. To this end, we have developed examples that spread over multiple chapters to teach the common CompleteEASE data analysis procedures. Following this tutorial is instructive for both beginning and expert users. Chapter 3 begins with basic data analysis examples. Here, we concentrate on thin films that are transparent. Chapter 4 continues with intermediate concepts, concentrating on absorbing materials and the extra difficulties involved with their characterization. Chapter 5 introduces a few advanced topics such as multi-sample analysis and anisotropic materials. Finally, Chapter 6 introduces in situ data analysis methods.

The section examples from this chapter are listed below, along with the primary CompleteEASE features that are discussed.

Section 3.1 General Data Analysis Theory

Section 3.2 Samples with Known N,K: SiO₂ on Si

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3.1. **General Data Analysis Theory**

Data analysis is a very important part of spectroscopic ellipsometry (SE): without data analysis, SE measures only the ellipsometric parameters Psi and Delta versus wavelength. To determine sample properties of interest, such as layer thicknesses and optical constants, a model-based analysis of the SE data must typically be performed. The only exception to this is in characterizing the optical constants of bulk samples, which is described on the next page. The model-based analysis approach is summarized by the flowchart shown in Figure 3-1. The basic steps of this approach are:

1. SE data is measured on the sample.
2. A layered optical model is built which represents the nominal structure of the sample. This model is used to “generate” SE data.
3. Model fit parameters are defined, and then automatically adjusted by the software to improve the agreement between the measured and model-generated SE data. This is known as “fitting” the data.
4. The results of the fit are evaluated. If the results are not acceptable, the optical model and/or defined fit parameters are modified and the data is fit again.

![SE Data Analysis flowchart](image)

*Figure 3-1. SE Data Analysis flowchart.*

The above steps will be described in more detail in subsequent sections. Keep in mind that while the basic SE data analysis approach is straightforward, “real-world” samples can often be difficult to analyze. The CompleteEASE software is designed to simplify common SE data analysis procedures which work for many types of samples, but there is no substitute for experience when dealing with complex samples. Please contact your J.A. Woollam Co. representative if you require assistance.
“Pseudo” Optical Constants

If SE data is acquired on a bulk sample which does not have any oxide, surface roughness, or any other films or overlayers, then it is possible to directly determine the optical constants of the sample from the ellipsometric data. Equation 4-1 can be used to transform the ellipsometric parameters Psi and Delta into the material optical constants “n” and “k” (or equivalently, the complex dielectric function values “ε1” and “ε2”).

\[
\varepsilon = \varepsilon_1 + i\varepsilon_2 = (n + i(k))^2 = \sin(\phi)^2 \cdot \left[ 1 + \tan(\phi) \cdot \left( \frac{1-\rho}{1+\rho} \right)^2 \right]
\]  (2-1)

where \( \phi \) is the angle of incidence and \( \rho \) is defined in Equation 1-1.

However, the direct transformation from ellipsometric data to intrinsic material optical properties given by (4-1) is quantitatively valid only if all the assumptions are met, that is, the sample does not have any oxide, surface roughness, or other films or overlayers. Since all these assumptions are rarely met (or at least it is very difficult to independently verify that they are met), the term “pseudo” optical constants is used to describe optical constants which are derived from this analytic transformation. The “<>” brackets are used to denote “pseudo” optical constants.

In CompleteEASE, pseudo optical constants <n> & <k> (or the pseudo dielectric function <ε1> & <ε2>) of the current SE data set can be displayed by selecting a <Pseudo> Transforms option from the “Graph Type” menu above the Graph display. Viewing the data in terms of pseudo optical constants can be useful for bulk samples.

Figure 3-2. Select from <Pseudo> Transforms option of the Graph Type menu to display pseudo optical constants or pseudo dielectric function.

Layered Optical Model

The SE data analysis process for most samples begins by building a layered optical model which corresponds to the nominal sample structure as shown in Figure 3-3. Each layer is parameterized by a thickness (t1, t2, etc.) and optical constants. Optical constants describe how light interacts with and propagates through the layer. Using the optical model and standard textbook thin film equations (Snell’s law, Fresnel
equations, etc.), the software can calculate “generated” or “simulated” SE data. If the model is a good representation of the sample, the model-generated SE data will be in good agreement with the SE data measured on the sample, as is illustrated in Figure 3-4.

![Figure 3-3. Schematic representation of a layered optical model with 2 films, parameterized by thicknesses $t_1$ and $t_2$, and optical constants $n$ & $k$.](image)

![Figure 3-4. Layered optical model in CompleteEASE, with model generated data: (a) the model generated data (black dashed lines) are not in agreement with the measured SE data (colored curves), (b) when the top layer thickness is adjusted appropriately, the model generated data lies directly on the measured data.](image)
Sometimes an ideal layered model does not adequately describe the optical behavior of the actual sample. The CompleteEASE software can also model two common “non-idealities”: surface roughness and index gradients. Surface roughness is shown in Figure 3-5: to model the actual sample which may have a non-abrupt “rough” surface, an “effective” roughness layer is added to the model. The optical constants of the “effective” roughness layer are derived by mixing the optical constants of the underlying material with the optical constants of “void” (which has optical constants of n=1, k=0). The Bruggeman Effective Medium Approximation (EMA) is used to calculate the optical constants of this “mixed” layer assuming 50% void content. While the effective roughness layer approach is certainly an approximation to the actual sample, this approach works extremely well for modeling SE data when the size of the surface roughness is much less than the wavelength of light used to measure the sample. For most SE systems, this implies that surface roughness features must be less than ~40nm.

![Figure 3-5. Optical model for surface roughness: (a) actual sample with non-abrupt “rough” surface, (b) optical model with “effective” roughness layer.](image)

Sometimes the optical “constants” of a layer are not constant throughout the layer. This may be caused by process variations during the film deposition. Figure 3-6 shows how CompleteEASE can model a “graded” film: the layer is divided into sub-layers with small thicknesses, and each sub-layer has slightly different optical properties. In CompleteEASE, a linear variation in the film index “n” is assumed for the graded layer. This simplified approach, which is automated in CompleteEASE, works well for modeling many types of samples.

![Figure 3-6. (a) Continuous variation in the optical properties of the “graded” film, (b) CompleteEASE approximation of the graded film with discrete layers.](image)

“Goodness” of Fit: definition of the MSE

Quantifying how well the data generated by the optical model “fits” or “agrees with” the measured data is an important part of the SE data analysis process. One can easily see “by eye” that model (b) in Figure 3-4 fits the data much better than model
(a). However, to automate the analysis process, the data fit or “agreement” is quantified by the “MSE” value defined in Eqn. (2-2):

\[
MSE_{NCS} = \sqrt{\frac{1}{3n-m} \sum_{i=1}^{n} \left( N_{E_i} - N_{G_i} \right)^2 + \left( C_{E_i} - C_{G_i} \right)^2 + \left( S_{E_i} - S_{G_i} \right)^2}
\]  

where “n” is the number of wavelengths, “m” is the number of fit parameters, and

\[
N = \cos(2\Psi), \quad C = \sin(2\Psi)\cos(\Delta), \quad S = \sin(2\Psi)\sin(\Delta)
\]

“MSE” is an acronym for Mean Squared Error. Actually, the definition given in (2-2) should be called the “Root Mean Squared Error” (as the error summation is divided by the number of measurement points minus the number of fit parameters, and the square root is taken). However, for historical reasons, we still use the term “MSE” to quantify the “goodness of fit”.

Basically, the MSE sums over all the measurement wavelengths the differences between the measured data (parameters subscripted with “E”) and model generated data (parameters subscripted with “G”). In CompleteEASE, the difference between the measured and model generated data is calculated in terms of the “N”, “C”, and “S” parameters, which are derived from the ellipsometric Psi (\(\Psi\)) and Delta (\(\Delta\)) parameters. The “N, C, & S” parameters have the following properties which make them better suited for the MSE definition (compared to using \(\Psi\) and \(\Delta\)):

1. N, C, & S are always bounded between -1.0 and 1.0.
2. The rotating compensator ellipsometer configuration measures data with approximately the same precision and accuracy in N, C, & S on any sample.

The N, C, & S parameters for the current data set in CompleteEASE can be plotted by selecting N, C, or S from the “Graph Type:” menu above the Graph display.

**Figure 3-7.** Select N, C, or S option from the Graph Type menu to display corresponding data.

The lower the MSE value, the better the fit or “agreement” between the measured and model generated SE data. As the typical precision and accuracy of the measured ellipsometric data in terms of N, C, & S is \(\sim 0.001\), a multiplicative factor of “1000” is included in the MSE definition of (4-2). This implies that an ideal model fit
should have an MSE of ~1. While this is achieved on certain types of samples (single thin films on Si substrates, for example), the best model fits for more complex samples (with thick and/or multiple layers) may exhibit much larger MSE’s (>10) and still be considered acceptable. It is also possible to get an MSE smaller than 1, but this implies the match to data is better than accounted for with random measurement error. In other words, the results likely include some of the “noise” present from data set.

“Fitting” the Data

To obtain the best fit between the model generated and experimentally measured data (or in other words, to achieve the lowest MSE), parameters of the optical model must be “adjusted”. The user defines which model parameters should be “adjusted” in the fit. These adjustable parameters are also called fit parameters, and they typically consist of layer thicknesses and parameters which define the optical properties of the layers. The CompleteEASE software then uses a standard, iterative, non-linear regression algorithm (the Levenberg-Marquardt method) to automatically minimize the MSE by adjusting the fit parameters. The Levenberg-Marquardt method does an excellent job of quickly converging to the best (lowest) MSE value, assuming that: 1) the optical model accurately represents the sample, and 2) the initial starting values for the fit parameters are reasonably close to their correct (best fit) values. To help ensure that the best fit model is found, the CompleteEASE software includes a couple of powerful features. These features will be used in the data analysis examples which are presented later in this chapter.

- The “Try Alternate Models” command automatically fits the data using the ideal layered optical model, and then with surface roughness and index grading non-idealities added to the model. The results of the various model fits are summarized in tabulated and graphical format, and the user can then decide which model is most appropriate for the sample.

- The “Thickness Pre-Fit” and “Global Fit” options perform a large number of trial fits to the data set, using a wide range of initial starting values for selected fit parameters. The best fit model is then reported.

Evaluating the Fit Results

Evaluating the fit results is arguably the most important (and unfortunately the most overlooked) aspect of SE data analysis. For a model fit to be acceptable, the following requirements must be simultaneously met:

1. **The model generated data must fit the measured data.**

   Unfortunately, this is difficult to quantify exactly, but as a rough guideline the MSE value should be in the 0.5 – 2 range for thin film samples which exhibit relatively little structure in the SE data vs. wavelength curves. For thick film samples with much structure/oscillations in the data, MSE values as high as 10 – 20 may be considered acceptable. Another definition of an “acceptable” fit is that the model generated data lies essentially on top of the measured data set, reproducing all the structures and features (except noise). However, it is not always possible to achieve a perfect fit and still satisfy requirements #2 and #3 below.
2. **The model should be unique.**  
The general rule of thumb is to choose the simplest model that adequately fits the data set. It is always possible to reduce the MSE by adding more layers and/or fit parameters to the model, but unless the MSE is significantly reduced, the additional model complexity is not justified. It is not generally possible to guarantee that a given model is unique (i.e., it is the only optical model that can fit the data to the same MSE value). For more complex samples, one should try alternate models and different starting values for the fit parameters to partially verify model uniqueness.

3. **The model and fit parameters must be physical.**  
Evaluating this requirement depends on a fundamental understanding of the constituent materials and structure of the sample. For example, if the optical constants determined from a model fit are outside acceptable bounds for the material (e.g., k<0 or n>10), the results are unphysical and should be considered invalid. Often, a knowledge base of measurements on similar samples is helpful in evaluating the “believability” of the fit results. For example, from experience we know that index gradients would be acceptable (or even expected) when analyzing data acquired on indium tin oxide (ITO) films, but a model which reported an index gradient when analyzing a thermal oxide on silicon sample would be questionable. Complementary techniques such as stylus profilometry, AFM, XTEM, SIMS or Auger depth profiling can also be used to corroborate the physicality of the model fit results.

If the model fit is deemed unacceptable, try modifying the optical model and/or defined fit parameters, and repeat the fit procedure until an acceptable fit is achieved. Finding an acceptable balance between the quality of the fit and the model complexity requires patience and experience.
“Physical” Optical Constants

One important criterion for any model is that the resulting optical constants do not have an “unphysical” shape. This can be difficult to evaluate without experience. To help “jump-start” your understanding of optical constants, we show a few examples of material optical constants and explain what makes their shape “physical”. We will also show corresponding results that would be “unphysical”.

A. **Index (n) must increase toward shorter wavelengths when the material is transparent (k=0).**

![Figure 3-8](image1.png)  
*Figure 3-8. Index must increase toward shorter wavelengths if k=0. These optical constants are physical as they obey this rule.*

![Figure 3-9](image2.png)  
*Figure 3-9. Index must increase toward shorter wavelengths if k=0. These optical constants are not physical because n turns-over (decreasing toward shorter wavelengths) below 600nm without any absorption (k=0).*
B. **Index (n) will “turn-over” and decrease toward shorter wavelengths if material becomes absorbing (k increasing).**

![Graph showing the behavior of index (n) as the material becomes absorbing]

*Figure 3-10. Index will turn-over and decrease toward shorter wavelengths when material becomes absorbing (k increasing toward shorter wavelengths). These optical constants are physical as they obey this rule.*

C. **Index (n) will return to the normal behavior of increasing toward shorter wavelengths as the amount of absorption begins to decrease (k decreasing).**

![Graph showing the behavior of index (n) as the absorption decreases]

*Figure 3-11. Graphed optical constants appear “physical”, as the index is increasing for wavelengths above 400nm (where k = 0), index turns-over when absorption becomes larger (near 350nm) and then returns to normal shape with n increasing toward short wavelengths when the absorption decreases (below 320nm).*
D. Extinction coefficient (k) cannot be negative.

Figure 3-12. The graphed optical constants are “unphysical” because the extinction coefficient, k, goes negative.
3.2. Samples with Known N,K: SiO\(_2\) on Si

**FEATURES IN THIS EXAMPLE**

- Opening model and data files
- Fitting data
- Parameter error bars
- Hiding or renaming fit parameters
- Reporting “derived” parameters
- Thickness “Pre-Fitting”
- Defining fit parameters (i.e. turning on and off)
- Visualizing model changes

The simplest types of samples to analyze are those in which the optical constants of each material are known. For this case, only layer thickness needs to be determined in the optical model fit. Unfortunately, most samples do not fall into this category, as the “optical constants” for most materials are rarely constant. Instead, they often depend on the material deposition and processing conditions. The analysis examples in the following sections demonstrate how to determine both layer thickness and optical constants from SE data. However, in this section we show the “classic” ellipsometry example which assumes known optical constants for the materials: thermally-grown silicon dioxide films on silicon wafer substrates (SiO\(_2\) on Si).

For high quality thermally grown oxide films on semiconductor-grade Si wafers, the optical constants of both the Si wafer substrate and the SiO\(_2\) film are known to be constant. The CompleteEASE software has a standard model for analyzing this type of sample. To open this model:

1. Start the CompleteEASE software (if the program is not open).
2. Select **Analysis > Model:** ‘Open’.
3. Highlight (by clicking) the “Basic” folder in the Library tab of the **File Location:** panel on the left side of the screen.
4. Select “Si with Thermal Oxide.mod” in the “Files:” section and click ‘Open’ button.

Next, open an example data file that we previously acquired on an ellipsometer system:

1. Select **Analysis > Data:** ‘Open’.
2. Highlight the “Examples” folder in the Library tab of the **File Location:** panel.
3. Select the “25nm Oxide on Si.SE” file.
4. Click the ‘Open’ button to open the selected data file.

After opening the specified model and data files, the CompleteEASE screen should appear as shown in Figure 3-13. Note the experimentally measured SE data (the ellipsometric Psi and Delta parameters versus wavelength) appear in the graph panel as red and green curves, and the optical model contains a substrate (SI_JAW) and two layers (INTR_JAW and SIO2_JAW).
Figure 3-13. CompleteEASE screen after opening the “Oxide on Si” model and the “25nm Oxide on Si” example data.

Fit Results and Parameter Error Bars

To analyze the data, click on Analysis>Fit>‘Fit’. The analysis will proceed very quickly (less than one second on most computers) and the fit results will be displayed in the Fit panel (Figure 3-14). Of course, the main parameter of interest is the total film thickness (27.42nm). The value after the “±” is the estimated error bar on each fit parameter.

The error bar values reported by CompleteEASE come from the 90% confidence intervals that are calculated by the fit algorithm. Since the confidence interval calculation makes a number of statistical assumptions that are not in general rigorously satisfied, the error bar values reported by CompleteEASE should not be literally interpreted. In practice, the error bars provide an estimation of the measurement reproducibility. Large error bars (relative to the magnitude of the fit parameter) are a useful indicator that the model is not sensitive to the parameter (and the parameter should likely be removed from the fit). The “Total Thickness” parameter is a derived parameter (more on that later). Its error bar is calculated from the sum of error bars for each varied thickness in the fit. Also note that after the fit, the black dashed “Model” generated data curves are essentially on top of the colored experimental data curves.
As an exercise, use the “Si with Thermal Oxide” model to determine the total film thickness for each oxide on Si example file: “60nm Oxide on Si”, “120nm Oxide on Si”, “300nm Oxide on Si”, and “1600nm Oxide on Si”. Open each data file (Analysis> Data> ‘Open’) and click ‘Fit’; the results should agree with the values shown in Table 3-1. There are a few important trends to note in this table:

1. The MSE increases with thicker films (from ~1 to >10), though all data fits are very good to the eye (i.e. black dashed Model curves lie on colored experimental curves).
2. The reported “Total Thickness” values do not agree exactly with thicknesses given in the file names (this is acceptable, as the file names are only estimated film thickness values).

Table 3-1. Fit results from “Oxide on Si” example files.

<table>
<thead>
<tr>
<th>Example File</th>
<th>MSE</th>
<th>Angle Offset</th>
<th>Total Thickness (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>25nm Oxide on Si</td>
<td>0.943</td>
<td>0.001 ± 0.0023</td>
<td>27.42 ± 0.007 nm</td>
</tr>
<tr>
<td>60nm Oxide on Si</td>
<td>0.687</td>
<td>0.037 ± 0.0014</td>
<td>58.54 ± 0.006 nm</td>
</tr>
<tr>
<td>120nm Oxide on Si</td>
<td>2.510</td>
<td>0.071 ± 0.0044</td>
<td>124.61 ± 0.007 nm</td>
</tr>
<tr>
<td>300nm Oxide on Si</td>
<td>3.673</td>
<td>-0.051 ± 0.0070</td>
<td>273.02 ± 0.014 nm</td>
</tr>
<tr>
<td>1600nm Oxide on Si</td>
<td>13.885</td>
<td>-0.094 ± 0.0261</td>
<td>1640.58 ± 0.195 nm</td>
</tr>
</tbody>
</table>
JAW Oxide on Si Model, with Interfacial Layer

The J.A. Woollam Co. “Si with Thermal Oxide” optical model supplied with the CompleteEASE software has a number of unique features. First, the oxide film is modeled as 2 layers: INTR_JAW, which is a higher-index interface layer, and SIO2_JAW, which contains the optical constants for the SiO2 film (which are different from the optical constants of a SiO2 “fused silica” bulk sample). The optical constants for these layers and for the SI_JAW substrate were published by Herzinger et al.1 The interface layer adds model complexity, but it significantly improves the data fit when the model is applied over a wide oxide thickness range.

To simplify the model and provide more unique fit results, the interface layer (“Thickness #1”) is fixed at 1nm. This value is typical for thinner oxide films (thickness < 100nm); while for thicker oxides, a slight fit improvement can be obtained by also fitting the interface layer thickness. For example, open the “1600nm Oxide on Si” file and fit the data with the “Si with Thermal Oxide” model. Now add the interface layer thickness as a fit parameter by right-clicking the mouse on the “Thickness #1” value (that is, (R^Θ) on “1 nm”). The value is now shown in a bold font followed by the word “(fit)”, indicating that this value is now defined as a fit parameter. (To turn “off” a fit parameter, simply right-click (R^Θ) again on the parameter value.) Click on the ‘Fit’ button, and the results box shown in Figure 3-15 should appear. Note the fit value for Thickness #1 is 3.48 (an increase from the nominal value of 1nm) and the MSE is reduced slightly from 13.885 to 10.316. Unless the explicit goal of the sample analysis is to characterize the interface thickness, it is best to fix Thickness #1 at 1nm nominal value.

![Figure 3-15. Results from “Si with Thermal Oxide” model fit to “1600nm Oxide on Si” data with interface layer (“Thickness #1”) added as a fit parameter.]

Derived Parameters

Even though the interface thickness increased in the preceding fit, the Total Thickness remained essentially the same (1640.58 vs. 1640.52 nm) for both fits. The “Total Thickness” is not an actual fit parameter, as the value is “derived” by summing all layer thicknesses in the model. Derived parameters can be added to the results by turning “ON” the “Include Derived Parameters” option within Model: >FIT Options (Figure 3-16). In addition to “Total Thickness”, additional derived parameters include “n” and “k” values for a specified layer and wavelength, optical thickness, total optical thickness and many other possibilities. These will be addressed further in a later example.

---

If you wish to delete a Derived Parameters, simply right-click on the number (shown in blue) next to the parameter you wish to delete.

**Hiding, Renaming, and Bounding Fit Parameters**

In the “Si with Thermal Oxide” model, “Angle Offset” was also defined as a fit parameter. The “Angle Offset” is added to the nominal angle of incidence in the optical model calculation. For most analysis (and especially when fitting for optical constants) it is best to leave the Angle Offset fixed at 0. However, since the Si and oxide optical constants are well known and there is only one additional fit parameter in the model (Thickness #2), fitting the “Angle Offset” can slightly improve the SiO₂ thickness analysis results by compensating for small sample alignment errors. Best sensitivity to angle of incidence occurs for thin layers (<100nm).

“Thickness #2” is a fit parameter in the analysis, but this parameter doesn’t show up in the fit results. “Thickness #2” was hidden to prevent confusion with the “Total Thickness” that is reported. To hide a fit parameter so that it is not displayed in the Results, left-click (L) on the fit parameter value. For example, if you (L) on “1637.05 nm ”, the dialog box shown in Figure 3-17(a) will appear. “Minimum” and “Maximum” values for the parameter can be specified to constrain the parameter space for the fit. If the “Show Adv. Parameters” box is checked, the Parameter Name, Error Bars, and Specification sections are visible. If the “Hide” box is checked, the fit parameter value will not be displayed as a fit result.
Figure 3-17. Fit parameter dialogs, showing how to (a) “Hide” or (b) “Rename” fit parameters. High and Low specs for the fit parameter are also set in (b).

Edit “Thickness #2” as shown in the Figure 3-17(b) dialog box:

- Check the “Rename” box, type in a different “Name:” for the parameter (e.g., “Top SiO2”)
- Uncheck the “Hide” box
- Type in values for “Low Spec.” (1200) and “High Spec.” (1400).

Click “Ok” to close the dialog box, and fit the data again (with the “1600nm Oxide on Si” data file loaded). Fit results will appear as shown in Figure 3-18. Since the “Top SiO2” value (1639.58) is outside the Low and High parameter Spec (1200 – 1400), an (Out Of Spec.) message appears after the result.

Figure 3-18. Fit results with parameter renamed and “Top SiO2” parameter “Out-of-Spec”.

Thickness Pre-fitting

One further concept will be demonstrated using “Si with Thermal Oxide” model. For this example, reload both the “Si with Thermal Oxide” model and the “1600nm Oxide on Si” example data file. From the Model->FIT Options section, turn the “Perform Thickness Pre-Fit” option to “OFF” by clicking on the underlined blue value. Then press Fit->’Generate’. The screen should appear as shown in Figure 3-19. Note that the data generated by the model (black dashed curves) using the default thickness (100nm) does not match the experimental data. Now click the ‘Fit’ button: the data fit is still very poor, and the reported MSE is very high (677). This illustrates what happens if you start the fit with parameter values that are not reasonably close to their best fit values.
Figure 3-19. Data generated from “Si with Thermal Oxide” model, using default parameter values. The “Perform Thickness Pre-Fit” option is turned “OFF”.

Now set “Thickness #2” to “1650” and “Angle Offset” to “0” (by clicking on the respective parameter values), and click “Generate”. The screen should appear as in Figure 3-20. While the model and experimental data curves do not line up exactly (slight shift), at least the basic structure and features in the data sets appear similar. Click the ‘Fit’ button, and observe that a good quality fit is obtained.

Figure 3-20. “1600nm Oxide on Si” data set, with model data generated using a closer (1600nm) starting thickness value.
A good starting value for “Thickness #2” is critical to achieve the best data fit. Without the Thickness PreFit, the software tries to adjust the fit parameter (thickness) to lower the MSE. However, the MSE profile is not a single minimum at the correct thickness. It also contains many other minima, which are called local minima. These local minima can trap the normal fit process. Each of these local minima are at a much larger MSE value, as shown in Figure 3-21., so it is easy to distinguish the correct answer from a thickness result that does NOT match the Experimental curves. However, it is tedious to guess starting film thickness values for all samples.

![Parameter Uniqueness Fit](image)

**Figure 3-21.** MSE profile for 1600nm SiO2 on Silicon data using the Thermal Oxide on Silicon model. The only answer occurs when the SiO2 thickness is near 1640nm. However, a normal fit will only find this answer if the starting point is near the final answer. To overcome this problem, our patented “Thickness PreFit” automatically searches for a good starting thickness near the final answer.

The CompleteEASE “Thickness Pre-Fit” option automatically provides a good starting value for the thickest layer in the model, using a special patent-pending algorithm. To demonstrate this feature, turn the “Perform Thickness Pre-Fit” option back to “ON”. Input a “bad” starting value for the “Thickness #2” parameter, for example “50”. Click ‘Generate’ to verify that the model starting parameters are not close to the best fit. Now when you click the ‘Fit’ button, a good data fit appears on the screen, even though the initial thickness value (50nm) was not close to the final best fit value (1639.58nm). For most analysis, it is useful to have the “Perform Thickness Pre-Fit” option turned “ON”, especially if accurate starting thickness values are not available.

To better understand the Thickness Prefit, consider that the data will oscillate versus wavelength as the film becomes thicker due to interference between reflected light from the surface and bottom of a thin film. As thickness increases, more oscillations will occur versus wavelength. Of course this is not independent of the index of refraction, but the underlying principle is that a larger “optical thickness” will have more oscillations. The Thickness Prefit works by quickly interpreting the data oscillations to “estimate” the actual thickness, as illustrated in Figure 3-22. Often, this fast calculation is adequate to get a good starting point for the normal fit procedures.
**Visualizing Model Changes**

It is useful to understand how changes in the model affect the generated data. For example, an increase in thickness will shift the interference oscillations toward longer wavelengths. CompleteEASE allows convenient simulation as a model parameter changes.

- Position the mouse over a model parameter you wish to adjust.
- Hold down the “Shift” key as you roll the mouse wheel up and down. The model parameter is increased or decreased by a set increment and data are re-generated with the new model values.
- For smaller parameter increments, simultaneously hold “Ctrl-Shift” as you roll the mouse wheel.

*Figure 3-22. Thickness Prefit uses a proprietary algorithm to very quickly estimate the thickness of a film based on oscillations in the data.*
3.3. Transparent Thin Films - 1

FEATURES IN THIS EXAMPLE

- Cauchy dispersion equation
- Global Fits
- Renaming Layers
- Displaying optical constants
- Expanding Layers

The next set of examples will demonstrate how to determine the film thickness and index of refraction “n” for transparent films deposited on substrates with known optical constants. While there are certainly cases in which it is not known a priori if the film is transparent (films which are absorbing are covered in the following chapter), there are also many applications where the film can be assumed transparent, at least in the visible spectral range: optical coating materials (SiO₂, TiO₂, Ta₂O₅, MgF₂ etc.), some nitrides (Si₃N₄, AlN), many organic films (photoresists, PMMA, spin-on polymers, etc.), and so on. In addition, the presence of periodic data oscillations suggests interference between top and bottom of a semi-transparent film (as shown in Figure 3-23).

![Figure 3-23: Interference within a transparent thin film.](image)

The index of refraction for transparent films in the visible spectral range is often specified using the Cauchy dispersion relation given by Eqn. 4-3. The “A” parameter relates to the approximate amplitude for the material index, while “B” and “C” parameters provide the shape or curvature of the index versus wavelength. Figure 3-24 shows index dispersion curves for three common transparent materials. For most transparent materials in the visible spectral range, the index curves slightly upward at shorter wavelengths, a behavior which is well described by the Cauchy dispersion equation.

\[
n(\lambda) = A + \frac{B}{\lambda^2} + \frac{C}{\lambda^4}
\]  

(4-3)
Si with Transparent Film

This example uses the “Flow-on Glass on Si” example data file. First, fit this data set using the “Si with Thermal Oxide” model as described in the preceding section. The results should look like Figure 3-25; the data fit is reasonable, but discrepancies are observed between the model and experimental curves in certain spectral regions, and the MSE is fairly high (~28). In addition, there is a significant Angle Offset (~0.698°). This is unreasonable for the hardware measurement, which had an angle accuracy of 0.01°, which helps prove the model is not acceptable for this data set. The “Flow-on Glass” material coated on this sample is similar to SiO₂, but with slightly different refractive index. Thus, it requires a different model that can account for a different refractive index.

Figure 3-25. Fit using “Si with Thermal Oxide” model.
To determine the index and thickness of the Flow-on Glass film, open the “Si with Transparent Film” model (which is also found under the “Basic” folder in the Open Model dialog box), and click the ‘Fit’ button. A nice data fit is obtained (though the fit takes longer than the “Si with Thermal Oxide” data fit) with the fit results shown in Figure 3-26. The reported fit parameters are Thickness #1 (which is the film thickness) and the film’s Cauchy parameters: \( A, B, \) and \( C \) (which calculates the films refractive index). The film index of refraction at a specified wavelength (632.8nm) is also reported as a derived parameter.

![Graph showing film fit results.](image)

**Figure 3-26.** Fit results for “Flow-on Glass on Si” example file using the “Si with Transparent Film” model.

**Right-click the mouse () over a Layer name to show its optical constants \((n,k)\).**

To display the film’s index of refraction versus wavelength, right-click () on the material name “Cauchy Film” and choose the option “Graph Layer Optical Constants” (Figure 3-27). The optical constants will be displayed in the graph window as shown in Figure 3-28. These represent the film’s optical constants.

To display and edit the film Cauchy parameters, click (L) the red “+” symbol to the left of “Layer #1”. The layer expands (and the “+” turns to a “-”, which can be clicked to collapse the layer), and the layer fit parameters are now accessible, as shown in Figure 3-28. The “k Amplitude”, “Exponent”, and “Band Edge” parameters describe an exponential “Urbach” absorption tail - not required for this example.

**NOTE:** The Cauchy layer is described in further detail in Section 9.6
Figure 3-27. To view a layer optical constants, ( ) over the layer name and choose to “Graph Layer Optical Constants”.

Figure 3-28. Optical constants displayed versus wavelength. Also, the Cauchy layer is expanded, displaying its available fit parameters.

**Global Fits**

Like the “Si with Thermal Oxide” model, the “Si with Transparent Film” model uses the “Thickness Pre-Fit” option to provide a good starting value for the layer thickness. To provide starting values for the film index, a “Global” fit is performed. In a Global fit, a wide range of parameter starting values are attempted and the values which result in the lowest MSE are then used as starting values for the “final” fit. The “Use Global Fit” option is found within **Model:** > **FIT Options**. With this option “ON”, expand the section (click red “plus”) to display additional settings which are used to customize the Global fit procedure (Figure 3-29).
In the “Si with Transparent Film” model, the Cauchy “A” parameter for the film (which is basically the approximate index of the film) is defined as a Global fitting parameter. Up to 3 fit parameters can be used in the Global fit. The “Min.” and “Max.” values for “A” are 1.3 and 3.0, and the “# of Guesses” is 30. When the ‘Fit’ button is pressed, the Global fit procedure performs 30 “trial fits” with different starting values for “A”, distributed in equal increments between 1.3 and 3.0. At each trial fit (i.e. for each value of “A” - which corresponds to the film index), a thickness pre-fit is also performed such that the entire array of film thicknesses and indices are attempted. A final fit is performed using starting parameter values from the trial fit which had the lowest MSE.

Since the Global fit process is computation intensive, a status bar with a ‘Stop’ button is displayed during the Global fit (Figure 3-30). To speed up the Global fit, only a subset of the experimental data points are used during trial fits based on the “Number of Data Points to Use” option. The number of iterations used in the trial fits can also be limited. The default values for these options (20 and 5) are adequate for most samples, but they can be increased if necessary: larger values may improve the robustness of the global fit, but with the downside of increased analysis time. Likewise, the Min., Max., and # Guesses for the parameter(s) may also need to be “tuned” to handle specific types of samples.
Figure 3-30. Status box and ‘Stop’ button displayed during a Global Fit.

Rename Layer & Fit Parameters

If you notice from this example, the Fit Results shown in the **Fit**: panel list the Cauchy results, as shown in Figure 3-31.

![Fit Results](image)

**Figure 3-31. Fit Results for Flow-on Glass.**

When preparing this model for a new material, it can be helpful to rename the fit parameters to describe the type of film being measured. Right-click on the Cauchy layer and choose “Rename Layer & Fit Parameters”, as shown in Figure 3-32.

![Layer Commands](image)

**Figure 3-32. Right-click on a layer to “Rename Layer and Fit Parameters”.**
This brings up the following dialog box, as shown in Figure 3-33. Choose what you would like to rename – the layer as shown in the model, the layer parameters (for Cauchy, this would be A, B, and C parameters), and the thickness parameter as reported in the Fit Results. I have chose the first and third, which gives a final fit (press Fit again) as shown in Figure 3-34.

![Figure 3-33. “Rename Layer Options” dialog box.](image)

![Figure 3-34. Final Fit Results with the Layer Renamed (in model) and the Thickness Renamed (as shown in Fit: Results panel).](image)
3.4. Transparent Substrates

In the previous example, we introduced the Cauchy dispersion relation to model the index of refraction for transparent films. In addition, the Cauchy can also be applied to transparent substrates. This is the most common approach for substrates of glass or plastic and the “Glass Substrate” model will use a Cauchy to determine the index for glass substrates. This example will demonstrate the modeling of transparent substrates.

With transparent substrates, we must consider a possible complication – the light may reflect from the back surface of substrate and reach the detector. The backside reflections will join the surface reflection, but these two collected beams will be incoherent. This is an optical description for light beams that have lost “phase” information relative to each other. In this case, the backside reflection will only adjust the overall amplitude of the signal and not produce “oscillations” like we see for thin films. This amplitude “offset” from backside reflections can lead to an error in the reported substrate index if not handled correctly. Fortunately, the backside reflections can be corrected in the software (described below).

Backside Substrate Reflections

The backside reflections that occur with transparent substrates are unwanted. They will affect the measurement and can produce errors in the reported refractive index for either the substrate or the thin films coated on the surface of the substrate. There are multiple ways to deal with transparent substrates and this topic is reviewed by Synowicki². There are two general approaches to deal with transparent substrates. The first is to avoid collection of the backside reflection. Methods to avoid collecting backside reflections include:

- Spatial separation of the backside beam via 1) thick substrate, 2) small measurement beam size, or 3) wedged substrate.
- Scattering and/or absorption of the backside reflection via 1) roughening the backside to scatter light, or 2) applying index-matching material to the backside (cloudy tape has been shown to be successful to this end).

If the backside reflections can not be avoided, then they need to be included in the software calculation. This will be demonstrated during the example. Fortunately, most Woollam SE systems collect depolarization data, which provides evidence of whether backside reflections are present in the collected data. If the depolarization is essentially zero, then no backside reflections reached the detector. This is the preferred condition. If backside reflections reach the detector, the CompleteEASE software can incorporate incoherent interference modeling to correct for the situation. Let’s consider data from a glass substrate to better understand this situation.

Graphing Multiple Data Sets

Open the data file for “glass substrate_rough” from the Examples location. This measurement was taken on a glass slide which was mechanically roughened on the backside to eliminate any reflected light from the back surface. Next, (⌘R) on Data > ‘Open’ to access the “Append Data” command, as seen in Figure 3-35. Choose the “glass substrate_taped” data file from the Examples location. Repeat these steps to also append the “glass substrate_smooth” data file.

NOTE: While you can only open one file at a time, the append command allows multiple files to be selected simultaneously.

When multiple data sets are appended, CompleteEASE allows all three to be accessed in the Graph window (either individually or the complete set). Select the “Graph Type” command at the top-left of the graph panel and choose to compare Psi. From this same location, also turn off the Double Y Axis. Finally, choose the “Graph All Data Sets” at the top of the Graph panel. This will allow a comparison of the Psi curves collected from all three data sets as shown in Figure 3-36.
Figure 3-36. When multiple data sets are appended, there are new choices at the top of the Graph Panel to go to any specific data set. Selecting the “Graph All Data Sets” check-box, allows comparisons with all data sets graphed.

Using the Graph Scratchpad

While the “Graph All Data Sets” provides a quick comparison of the three data sets, it is not customizable. For example, we can’t change the colors or select a subset of the total curves. For this purpose, we turn to the Graph Scratch Pad. Right-click (⌘R) within the graph area and choose to “Copy Data to Graph Scratchpad”, as shown in Figure 3-37. This will copy all of the curves currently shown on the graph and place them in an internal “pad” for further manipulation. Next, choose to “View Graph Scratchpad” from the same (⌘R) menu. This opens the CompleteEASE Graph Scratchpad, as shown in Figure 3-38. Play with the Scratchpad by selecting which curves to compare, change their colors, and their styles. Finally, it is beneficial to change their names to correspond to the data set that they came from. Figure 3-39 shows a comparison of the taped and rough glass data – which are nearly identical. This shows that taping the backside can be as effective as roughening the backside of a transparent substrate for removing backside reflections. The data comparison also shows a large difference between the rough or taped data and the data with smooth backside. This large offset is caused by the light reflecting from the backside of the substrate. Next, we will model each data set to show how this backside reflection can produce error in refractive index measurement and finally, how to correct for this backside reflection.
Figure 3-37. Right-click (\text{CTR}L) within the graph area to access the “Graph Scratchpad”.

Figure 3-38. CompleteEASE Graph Scratchpad allows customization of the graph.
Figure 3.39. The Graph Scratchpad is used to compare data from glass substrate where the backside is roughened or taped to suppress backside reflections.

NOTE: Before proceeding, clear the contents in the Graph Scratchpad by viewing the Scratchpad and pressing ‘Delete All’.

Figure 3.40. Fit to "Glass Substrate" data.
In a later example, we will learn how to fit multiple data sets simultaneously. However, for this example, we will focus on fitting each individual data set. Start by opening the “glass substrate_rough” data file. Next, open the “Glass Substrate” model from the Basic folder. Press ‘Fit’ and you should get the result as shown in Figure 3-40. The data appear noisy, but this is primarily due to the graph scale.

![Opt. Const. of Cauchy Substrate vs. nm](image)

*Figure 3-40. Index of refraction from Cauchy model fit to bare glass substrate.*

The final index of refraction can be viewed by right-clicking on the Cauchy Substrate and choosing “Graph Layer Optical Constants”. Right-click (⌘R) inside the Graph and choose “Copy Graph to Clipboard” to transfer the graph shown in Figure 3-41 to Word.

**NOTE:** Please copy the optical constants graph to the “Graph Scratchpad” before proceeding as we will compare the results to future fits in this example.

### Graphing Depolarization

Ellipsometry measurements use polarized light. If the sample is isotropic and ideal, this measurement can be described by two values: \( \Psi \) and \( \Delta \). However, if a sample is partially depolarizing, two parameters are insufficient to fully describe the change in polarization. The patented Rotating Compensator technology used in the alpha-SE, M-2000, and RC2 systems permits measurements of three reported quantities: \( \Psi \), \( \Delta \), and %depolarization. The “%depolarization” should remain zero for ideal sample conditions.

**HINT:** Some ellipsometers require DC measurement to acquire accurate depolarization data. Please consult your hardware manual (or contact JAWCo) to best measure the depolarization.

**Shortcut:**

- **CTRL-Z** to graph Depolarization

Before proceeding, press on “Graph Type” at the top-left of the Graph. Change from “Psi & Delta” to “Depolarization”. Notice, the Depolarization graph in Figure 3-42 is near zero. This indicates that little light is reflected from the backside into the detector.
Next, open the “glass substrate_smooth” data file. The graph of Depolarization is now centered at almost 2%. This indicates the backside was smooth and incoherent reflections from the smooth backsurface reached the detector unit. Change the Graph Data back to “Psi”. Press Fit:’Generate’. The model no longer matches the experimental measurement even though this data is from the same glass substrate (only with backside reflections). The backside reflections do more than introduce depolarization – they also shift the ellipsometry values. Fit the data with the same model and you should get results as shown in Figure 3-43.

Before proceeding, view the optical constants and add these new results to the Graph Scratchpad - allowing comparison of the two results in the Scratchpad as shown in

**Shortcut:**

CTRL-P to graph Psi.

**Figure 3-42.** Depolarization from uncoated glass slide (no backside reflections).

**Figure 3-43.** Fit results and optical constants for glass substrate_smooth data modeled without backside reflections.
Figure 3-44. The first fit result should be correct, as the glass did not include backside reflections. The second fit, with a much lower index of refraction is incorrect because the backside reflections were present in the data, but were not considered in the model. Both fits had a good MSE, which shows that the wrong model can often lead to misinformation. In this case, there are a couple ways to ensure the correct model is found. For any bare transparent substrate, it is preferred to measure multiple angles of incidence. This data would have shown us that the smooth backside data was not correct, as more than 1 angle will not fit at the same time in the presence of backside reflections. Unfortunately, this data was collected from an instrument with only one fixed angle. Fortunately, there is another method that can account for backside reflections in measured data and help get correct results. This is considered in the following section as it is a common occurrence with transparent substrates.

![Graph Scratchpad](image)

**Figure 3-44. Comparison of fit results for refractive index from within the CompleteEASE Graph Scratchpad.**

### Including Backside Reflections

Backside reflections are common with transparent substrates. Substrates are generally thicker than the coherence length for the white light source of the ellipsometer. Thus, the returning light does not interfere in a coherent sense, as its phase information is lost compared to the reflections from the top-surface coatings.

Back-side reflections are undesirable when characterizing surface films. This is especially true if the substrate is anisotropic. To prevent back-side reflections the back-surface can be roughened to scatter light, as in Figure 3-45. For thicker substrates, the backside reflections may be spatially separated at the detector and thus not a concern.
If the backside reflections can’t be avoided, the “incoherent” summation of light beams can be handled mathematically. To include backside effects in the model, expand the **+MODEL Options** section of the Model. Turn “Include Substrate Backside Corrections” to ON, as shown in Figure 3-46. The correction depends on the amount of light collected from the backside, which can differ for different substrates. In the Model Options section, select the “# Back Reflections” as a fit parameter.

**Fitting Depolarization**

There is correlation between the substrate index and the amount of backside light collected when modeling ellipsometry data at one angle of incidence. However, depolarization data is only influenced by the amount of backside light collected. The default setting is to ignore the depolarization data and only fit Psi and Delta. In fact, the depolarization fitting selection is a hidden feature of the model. To show this feature, go to the bottom of the **Model: panel and press Configure Options**. This will bring up the dialog box shown in Figure 3-47. Check the box next to “Include Depolarization Data” in the Fit Options section and then ‘Ok’. With this option shown, choose **Model:>+Fit Options** and turn the “Include Depolarization Data” to ON, as shown in Figure 3-48.
Figure 3-47. Press **Configure Options** to show additional Model options. In this example, we will add “Include Depolarization Data” as shown.

+ **MODEL** Options
- **FIT Options**
  - Perform Thickness Pre-Fit = **OFF**
  - Use Global Fit = **OFF**
  - Fit Weight = **N.C.S**
  - Include Depolarization Data = **ON** % Weight = **100.00**
  - Limit Wvl. for Fit = **OFF**
  - Limit Angles for Fit = **OFF**
  - Max. Acceptable MSE = **100.000**
  - + Include Derived Parameters = **ON**

+ **OTHER Options**
  - **Configure Options**
  - **Turn Off All Fit Parameters**

Figure 3-48. The “Include Depolarization Data” section within the +Fit Options section of the model.

Press ‘Fit’ again for the “Glass Substrate_smooth” data including the backside reflections (with fit to # of Back Reflections collected) and using the Depolarization data in fit.

**NOTE:** You will probably have to change the starting # of Backside Reflections to 1 (from default of 5) to get the depolarization data to fit.

When done correctly, the result should appear as in Figure 3-49. The measurement of depolarization provides sensitivity to the amount of backside reflected light that is reaching the detector. In this case, the best fit is with 1.5 backside reflections, as shown in Figure 3-50 (where the number of backside reflections is varied to see whether the number of backside reflections is a unique result). This means the substrate is thin enough that the first reflection from backside is completely collected in the detector, while the second reflection is spatially shifted enough to only collect about half of this second reflection. Of course, as the number of reflections increases, the amount of additional light reaching the detector is much lower and the sensitivity goes down. Thus, there is little difference in the calculation between 3, 4, and 5 total reflections from the backside.
Before proceeding, let’s view the optical constants from this result and copy the graph to the Graph Scratchpad for further comparison with our previous results. The final comparison of optical constants is shown in Figure 3-51. The index for backside roughened glass is similar (but not exactly equal) to the results for smooth backside glass when the fit is corrected including depolarization. If the backside effects are not considered, the fit results significantly affect the index of refraction. There are two possibilities as to the difference in final results when corrected:

- Depolarization data not accurately determined.
- Some backside reflections for roughened glass still remain.

Figure 3-49. Fit results including backside reflections and depolarization data.
Figure 3-50. Uniqueness Test for the # of Backside Reflections, showing that the best fit results from about 1.5 backside reflections. The depolarization measurement helps to quantify this result.

Figure 3-51. Final comparison of glass index results. While the backside correction improves the substrate index results (to match more closely with roughened backside), they are not perfectly matched. When not corrected, the smooth backside measurement is very inaccurate.
3.5. Transparent Films - 2

FEATURES IN THIS EXAMPLE

- Surface Roughness
- Grade Layer
- Try Alternate Models

Glass with Transparent Film

For this example, load the “Glass with Transparent Film” model and the “ZrO2 on Glass” data file. Click the ‘Fit’ button: a Global fit will be performed and the fit results are displayed as shown in Figure 3-52. The refractive index for the ZrO\textsubscript{2} film can be displayed by right-clicking (\[\text{R}\]) on “Cauchy Film” and choosing “Graph Layer Optical Constants.” The results are shown in Figure 3-53.

![Graph showing fit results](image)

*Figure 3-52. Fit results from analysis of “ZrO2 on Glass” example file, using the “Glass with Transparent Film” model.*
The main difference between the “Glass with Transparent Film” and the “Si with Transparent Film” model which was used in previous examples is the material file used for the substrate: “7059_Cauchy” instead of “SI_JAW”. As the optical constants for 7059 glass are representative of many glass substrates, this model can be used as a starting point for many transparent films on glass. More accurate results could be achieved by replacing the 7059_Cauchy material with optical constants measured on your bare glass substrate (see section 3.4). In addition, because the substrate is transparent, backside reflections may be an issue. To test whether backside reflections are present in this data, press CTRL-Z to view the Depolarization data as shown in Figure 3-54. As the depolarization is centered at zero, it is likely that the backside reflections were suppressed to avoid collection at the detector. These issues are discussed in more detail in Section 3.4.

Figure 3-53. ZrO2 optical constants from fit using Cauchy dispersion layer.

Figure 3-54. Depolarization data is near zero percent, which indicates that backside reflections were not collected even though substrate was transparent.
Surface Roughness

An interesting feature to the current model is the inclusion of surface roughness (the “Include Surface Roughness” option is set to “ON”). Actually, the best fit surface roughness value for this sample is rather large: 11.17 nm. To test the importance of roughness in this model, try fitting the data without surface roughness: toggle the “Include Surface Roughness = ” value to “OFF” by clicking on the blue underlined value. Since we already know the nominal thickness and index, it is also wise to turn OFF both the “Use Global Fit” and “Thickness Prefit”. Go ahead and click the ‘Fit’ button, which will fit the data without roughness – resulting in a poor fit and higher MSE, as displayed in Figure 3-55.

NOTE: Surface Roughness automatically skips any layer that has zero thickness. This can be important when working with multi-sample analysis of multilayered structures.

Graded Index

Surface roughness, which is one of the model “non-idealities” discussed in section 3.1, is very important to achieve a good model fit on this sample. Another non-ideality described in section 3.1 is index grading. With the surface roughness turned off, try grading the film as described below:
• (CTRL) on the name “Cauchy Film” of Layer #1. You should see a drop-down menu, as shown in Figure 3-56. Choose “Grade Layer”.

• The “% Inhomogeneity” and Thickness are defined as fit parameter.

• Press ‘Fit’ and the results will appear as in Figure 3-57.

NOTE: Surface roughness and index grading often have the same effect on modeled data, as they can both reduce the optical density of a material toward the surface. If both give similar MSE values, it is up to the user to judge which is best for their sample.

While the fit appears quite good to the eye, the MSE is slightly higher than for the model which included surface roughness (2.1 vs. 1.175). It is also possible to combine surface roughness and index Grading in the model: turn the surface roughness back “ON” and press ‘Fit’ (leaving the “% Inhomogeneity” defined as a fit parameter). The results shown in Figure 3-58 indicate this model yields the lowest MSE yet (0.78).
Figure 3-57. Fit to “ZrO2 on Glass” data with index Grading ON.

Figure 3-58. Fit results for “ZrO2 on Glass” data fit with model that includes both surface roughness and grading.
Try Alternate Models

NOTE: This command will not work if the film is already graded. To remove grading from our current model, (CTRL on Graded Layer and select “Remove Grading”, as shown in Figure 3-59.

The process of trying optical models with and without surface roughness and index grading is so common in SE data analysis that CompleteEASE has a command to automate the procedure. Once a good starting model is established, click Model>OTHER Options> Try Alternate Models, as shown in Figure 3-60. Fits will automatically perform with the ideal optical model, with surface roughness, with index grading, and with roughness & grading. Fit results are summarized in tabulated and graphical formats, as shown Figure 3-61. The highlighted “green” column is a suggestion of the “best” optical model, based on 25% improvement in MSE for each additional fit parameter.
Figure 3-61. Results of the **Try Alternate Models** command.
3.6. Log and Report a Series of Films

**FEATURES IN THIS EXAMPLE**

- Copying to clipboard
  (Formatted or Table)
- Fit log
- Open/Save Snapshot
- Comparing optical constants in fit log
- Copy Analysis Report to Clipboard
- Re-analyzing data from log

One of the most common applications of any ellipsometer involves measurement of a series of thin films, which are to be compared and reported. The CompleteEASE software enables simple logging, comparing, and reporting of results. This example will step through a series of SiNx thin films on Silicon, using the analysis steps taught in Section 3.3. However, the emphasis will be on using CompleteEASE to log all fit results, compare optical constants for the various films, and finally create a record (or report) of the results.

To begin this exercise, open “SiNx on Si-1” data file and the “Si with Transparent Film” model in the Analysis tab of CompleteEASE. Go ahead and press ‘Fit’ and the results shown in Figure 3-62 appear.

![Figure 3-62. Fit to “SiNx on Si-1” data file using “Si with Transparent Film” model.](image)

**Figure 3-62. Fit to “SiNx on Si-1” data file using “Si with Transparent Film” model.**
Copy to Clipboard

Each pane (white area) in the Analysis tab of CompleteEASE offers right-click (Ctrl-R) menu choices. Examples of these right-click menus are shown in Figure 3-63. In addition, the Graph also features a right-click menu, as shown in Figure 3-64. To assist in creating reports, the model, fit results, and various graphs can be copied to the clipboard and then pasted in various programs such as Word, Excel, or PowerPoint. The Fit results can be copied to the clipboard using Formatted or Table choices, where the former is best for pasting into a word processor and the latter is preferred for pasting into a spreadsheet. The Model can be copied in its entirety or as the layers only.

![Figure 3-63. Right-click (Ctrl-R) menu choices from the Fit: and Model: panels.](image1)

![Figure 3-64. Right-click (Ctrl-R) menu choices from the Graph.](image2)
Fit Log

Now, work through all five SiNx on Si samples in the Examples folder. To record the final fit results for each film, right-click (⌘R) on the Fit: panel and “Add to Fit Log” after the fit is finished. This will save the current fit results. Type in an appropriate name for each fit result.

NOTE: The shortcut key to add results to the Fit Log is CTRL-L.

After all five data files have been fit and added to the Fit Log; choose to “View Fit Log” from the right-click (⌘R) menu of the Fit: panel. This should bring up a Fit Log with each of the five entries, as in Figure 3-65. You can press on any of the individual log entries to see the results from that sample.

NOTE: The shortcut key to View the Fit Log is CTRL-ALT-L.

Comparing Results

Now, select all five Log Entries by selecting the first entry with your mouse and then holding the shift-key down while you press the last Log entry. With multiple Log entries selected, press the ‘Compare’ button. This will create a Table of results for each sample, as in Figure 3-66. The columns/rows can be reversed by a simple button click. Also, the statistics can be added with a click. This table can be conveniently copied to the clipboard.

![Figure 3-65. Fit Log showing results from all five SiNx on Si data files.](image-url)
Compare Optical Constants

From the Comparison Table, press the ‘Compare Optical Constants’ button and dialog box appears as in Figure 3-67. Select each of the layers you would like to compare and choose the type of graph and you can compare the index for each of the SiNx layers, as shown in Figure 3-68. If you would like to select all of the SiNx layers, you can press ‘Select Layers by Number’ and choose 1. Otherwise, you can individually select each layer by clicking the box in the representative model graph. The final graph can be conveniently copied to the clipboard to be added to a report.

Figure 3-66. Comparison Table showing results from all five SiNx films.

Figure 3-67. Compare Optical Constants Dialog Box.
Re-analyzing Multiple Data Sets

The process we just finished for the series of SiNx samples is quick, but somewhat repetitive. There is a much faster way to analyze multiple data sets and build a master report. First, clear the Fit Log from our previous efforts. To clear the Fit Log, press CTRL-ALT-L to view the Fit Log and press the ‘Clear Log’ button. Now, you are ready to repeat this example using a faster method.

To start, press CTRL-ALT-L to view the Fit Log, if you are not already there. There should be no current entries, as we just finished clearing the log. Press the ‘ReAnalyze’ button, which will open the dialog window as shown in Figure 3-69. The basic concept is that we are going to add a “Row” describing each batch-analysis we would like to run. The “Row” will contain three items: a File Group (collection of data files), a Model, and an optional Log-Name.

NOTE: In older versions, CompleteEASE could only use a single model on a group of data sets. Now, multiple “Rows” can be added to provide different types of analysis (different data sets and/or modeling).

Start by pressing ‘Add Row’ to create the group of data sets to reanalyze. This collection of data sets can be from entries already in the Fit Log (in which case they need to be selected) or you can browse and select multiple data sets at this point by pressing ‘Add’ from the Choose Files dialog box, as shown in Figure 3-70. Go ahead and add all five SiNx data files that we just recently analyzed into a single File Group. You will be asked to enter a Group Name - I chose SiNx. If you liked to analyze other data files at the same time, you could add further Groups at this point.

Next, press “Choose a Model” and select the “Silicon with Transparent Film” model that we used for these nitride films previously – as shown in Figure 3-71. You can also add a “Log Name” that will be added to the end of each Log Entry after reanalysis. Go ahead and press ‘Reanalyze’. This will run through the analysis of all five data files and save the results of each to the Fit Log, as shown in Figure 3-72. This one step has reproduced the entire example we worked earlier in this section.
NOTE: The “Re-save Data” check-box allows the new fit results to be saved along with each data file. In CompleteEASE, the data file can also contain the results – to be reviewed at a later time.

Figure 3-69. ReAnalyze Data window opened from within the Fit Log.

Figure 3-70. Creating Groups of Data Files

Figure 3-71. Add Model and Log Name
Figure 3-72. After reanalysis, each data file shows as a Log Entry. Notice the Log Name “1” is added to the very end of each Log Entry.

‘Generate Report’

Now, use another approach to create a report for all the results. View the Fit Log (CTRL-ALT-L) and select all five entries. Next, press the ‘Generate Reports’ button, as shown in Figure 3-73. This saves the results of each log entry into an RTF file that can be opened by Microsoft WORD. This RTF file will be formatted as shown in Figure 3-74.

Figure 3-73. All log entries are selected and the ‘Generate Reports’ button is pressed to copy all results into the HTML Clipboard.
Figure 3-74. HTML Clipboard after a report has been generated for all entries in the Fit Log.


If you would also like to add a graph of the optical constants for the SiNx layer, this can be setup from the Model:>>OTHER Options> “Add Optical Constants to Report”, as shown in Figure 3-75. If you have a multi-layer model, you may need to change the Layer number to match the layer in your model that has the varying optical constants. You would need to resave this model (with feature turned on) and reanalyze all of the data sets again. We will skip this step, but demonstrate the capability on a single data file in the next section.

Figure 3-75. Use “Add Opt. Const. to Report” to include optical constant graphs when reporting CompleteEASE fit results. Note that you need to add the Layer number corresponding to the material of interest.
Copy Analysis Report to Clipboard

Next, we will demonstrate how the report can be created for a single result. After data analysis, right-click in the Fit: panel and choose "Copy Analysis Report to Clipboard", as shown in Figure 3-76. This will copy the same graphs that were created as part of a report, but into the Clipboard. For this example, the Optical Constants were copied to the clipboard as shown in Figure 3-77, since this feature was turned on in the Model: > OTHER Options section.

![Figure 3-76. Right-click in the Fit: panel to "Copy Analysis Report to Clipboard"](image)

![Figure 3-77. Example of Analysis Report, including Optical Constants graph.](image)
Save Log File

When finished with a project, you may want to save the results or send them to a colleague. The Fit Log is a nice way to store all of the modeling results you have completed. You can save a Fit Log, by selecting “View Fit Log” from the Fit: panel right-click menu. Then, choose the ‘Save Log’ button, as shown in Figure 3-65. The log will be saved with a “fit” extension. The log will include the list of all entries, with the filename for each data file needed and the model layers. To open the log, the data files must be in the same directory as the “fit” log file. Thus, if you intend to send your log to a colleague, they will also need all of the Data files to open the log file.

Snapshot

A better way to send single fits to a colleague, or save them for future use, is to save the Snapshot of your fit result. A snapshot will compress your data file, model, and fit results into a single file with “SESnap” extension. Thus, a single snapshot file can be stored or sent and then opened to get back to the same result that you saved. To save a Snapshot after performing a Fit on a SiNx data example, simply click the Model: > ‘Save Snapshot’ button, as shown in Figure 3-78. To open the Snapshot, simply click the ‘Open Snapshot’ button.
4. Data Analysis 2 – Intermediate

This chapter continues the series of examples with intermediate concepts, concentrating on absorbing materials. The section examples from this chapter are listed below, along with the primary CompleteEASE features that are discussed.

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### Section 4.6 Multiple Data Types (SE + T)

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4.1. Metal Substrates

Substrate optical constants are an important consideration when modeling thin films. Although certain materials, such as crystalline silicon are well understood and remain constant, others substrates such as glass and metals are more variable. In these cases, it is best to measure a bare substrate to determine optical constants. Glass substrates were considered in Section 3.4. We will consider a metal substrate for this example. Although the substrate is called a metal, it is actually a glass or silicon substrate that is coated with an optically opaque metal film. The measurement beam will not penetrate through the metal film, so we can consider it a substrate. For most metals, it only requires 100nm thickness to absorb the light and prevent measurement through the metal to layers or substrate below.

<Pseudo> Transforms

Open the “Au_Substrate” data file. Substrates offer a special case, as the measurement involves only a single-interface (reflection from surface). The experimental measurement of Psi and Delta can be directly inverted to determine n,k using the pseudo-substrate approximation. The <Pseudo> Transforms can be graphed by selecting the “Graph Type” and expanding the <Pseudo> Transforms section, as shown in Figure 4-1. This allows you to graph the <n>, <k>, <e1>, and <e2> values. It is important to note that these are a direct inversion of the measured Psi-Delta data to “Pseudo” optical constants under the assumption that a single reflection occurred at the surface. These values are not equivalent to the true optical constants in the presence of any thin films (even a few nanometers surface layer).

NOTE: <Pseudo> optical constants are represented with “< >” brackets to show that they are not the “true” optical constants, but simply an inversion of the raw data only valid for uncoated substrates.

Figure 4-1. Click on the “Graph Type” and Select <Pseudo> Transforms to expand to the “Pseudo” optical constant choices.
The <Pseudo> Transforms are shown for the Au data in Figure 4-2. Because the Au layer is opaque, only a single reflection occurs from the surface, so the <Pseudo> transformation is acceptable. We still need to model the data to create a layer that holds the optical constants as shown below. We will demonstrate this concept using two different layer types: the WvlbyWvl layer and the B-Spline layer.

![Spectroscopic Ellipsometric (SE) Data](image)

*Figure 4-2. <Pseudo> optical constants for the Au substrate data.*

**WvlbyWvl Layer**

Open the “Blank” Model from the Basic Folder. Press the “Substrate = none” to add a material file for the substrate. Choose the “WvlByWvl.mat” file from the Advanced Folder, as shown in Figure 4-3. Expand this layer, by pressing the + sign to the left of the layer and your model should look the same as seen in Figure 4-4.

![Advanced Folder Link](image)

*Figure 4-3. Choose the Advanced Folder Link and then select the WvlByWvl.mat file.*
The WvlByWvl layer starts with initial values of n=1.5 and k=0 (at all wavelengths). However, metals are always absorbing (k>0) and therefore these values are not representative of a metal. It can be important to have good starting values, as the Fit will proceed by moving the optical constants at each wavelength independently to find the best match to the Experimental data at that same wavelength. In this layer, there is no consideration of neighboring wavelengths, so it is possible for the optical constants to get “lost” and find a local minimum instead of the correct answer. To help the WvlByWvl layer find the correct result, it is better to “seed” the starting optical constants closer to the final results. There are two ways to accomplish this: entering “Initial Values” or entering a “Starting Material”. For this example, we can use the <Pseudo> Transform graph as a guide for the “Initial Values” – the <n> ranges from 0.2 to 1.6 while the <k> ranges from 1.7 to 5.8. A good choice of “Initial Values” would be near the center of these ranges. Because the Au values cover a very wide range, a better approach is to use the “Starting Mat”. Press to enter a new “Starting Mat” from within the WvlByWvl layer and choose the “Au” material file from the Metal Folder Link. This will enter the gold values from Palik’s Handbook of Optical Constants as the starting values at each wavelength within the WvlByWvl layer. You may be asking why we don’t stay with the Handbook values. Unfortunately, metal optical constants can vary significantly from source-to-source and are even discontinuous within Palik from his use of multiple references. Right-click on Au to Graph the Optical Constants and you should see the values as shown in Figure 4-5.
Figure 4.5. Optical Constants of Au from Palik’s Handbook of Optical Constants.

If you press ‘Generate’, you will see how closely the Palik Au values match our experimental data, as shown in Figure 4.6. As you can see, the discontinuity in Palik’s Au optical constants also produces a discontinuity in the generated data. Also, the match to our experimental data is not perfect because the optical constants from Palik are not from the same Au process.

Figure 4.6. Model generated curves from Palik’s Au values compared to our measured Au experimental data.

Now, go ahead and press ‘Fit’. The WvlByWvl layer allows the n,k values to vary independently at each wavelength to best match the experimental data. This produces a nearly perfect fit to single-angle data, as shown in Figure 4.7. Now, right-click on the WvlByWvl layer to graph the optical constants – as shown in Figure 4.8. These optical constants can now be saved as a reference material file for “our” Au process. Instructions for saving optical constants are found in the following sections. Before proceeding, please copy the Optical Constants graph to the Graph Scratchpad.
While the WvlByWvl layer works well for this simple substrate example, it does not use neighboring wavelengths to help determine the optical constants. This can produce small amounts of noise in the optical constants, due to any noise in data points at a specific wavelength. We see small evidence of this around 800nm in Figure 4-8. To keep the optical functions smooth, a B-Spline layer can be used for the model, instead of the WvlByWvl. We will save the intimate details of the B-Spline layer for a later example. In this exercise, we will build a model to fit any metal substrate and then save this model for future use.

**B-Spline**

While the WvlByWvl layer works well for this simple substrate example, it does not use neighboring wavelengths to help determine the optical constants. This can produce small amounts of noise in the optical constants, due to any noise in data points at a specific wavelength. We see small evidence of this around 800nm in Figure 4-8. To keep the optical functions smooth, a B-Spline layer can be used for the model, instead of the WvlByWvl. We will save the intimate details of the B-Spline layer for a later example. In this exercise, we will build a model to fit any metal substrate and then save this model for future use.
Let’s start fresh by re-opening the “Blank” model from the Basic folder. Left-click (⌘L) on the Substrate = “none”, as shown in Figure 4-9. This will bring up the “Open Material” dialog box. Choose the B-Spline.mat file from the Basic folder and press ‘Open’. Expand the B-Spline layer such that it can be edited.

- Set the resolution to 0.1eV.
- Set the Starting Mat = Au (from the Metals location)

The B-Spline layer specifies the optical constants versus wavelength using a series of control points (“nodes”) which are equally spaced in photon energy (eV). The approximate spacing of the control points is controlled by the “Resolution (eV) =” field. When a starting material is specified, the nodes are matched to the starting material file to provide an estimate before fitting. It is not necessary for the starting material to be the same as the material you have measured, but we improve our chances of finding the “correct” solution by choosing a starting point close to the correct values before fitting.

![Figure 4-9. Starting the example for Au_substrate with the “Blank” Model.](image-url)
Generate data and then attempt the fit. It should quickly converge to the answer as shown in Figure 4-11.

To view the optical constants as shown in Figure 4-12, right-click on “B-Spline” and choose “Graph Layer Optical Constants”. To view the B-Spline nodes as in Figure 4-13, press on ‘Draw Node Graph’ within the B-Spline layer. The red and green curves are the smooth optical constants (e1 and e2) that are derived from the B-Spline fit, with the open circles showing the individual spline nodes.
Before proceeding, view B-Spline layer optical constants again and copy the optical constants graph to the Graph Scratchpad. Open the Graph Scratchpad to compare to the results from the WvlByWvl fit. Notice we get more smooth optical constants, as shown in Figure 4-14.
As an exercise, change the Spline resolution to larger and smaller values and repeat the fit. Does the fit quality change? You can use the Graph Scratchpad for comparison.

**Save**

When happy with results, press the Layer Command: **Save** right above the model to save the gold optical constants. This opens the Save Layer Optical Constants dialog box as shown in Figure 4-15. First, choose the “Tabulated” choice from the right. Next, click on the substrate layer itself. After entering the file name and comment, the tabulated $n,k$ at each measured wavelength will be saved for future use.

**NOTE:** The Layer Command: **Save** is different than the ‘Save’ button in the Model: panel. The Layer Command: **Save** saves the layer optical constants, while the ‘Save’ button in the Model: panel saves the layered model structure and any +MODEL, +FIT, or +OTHER Options.

![Figure 4-14. Index results compared from fits using WvlByWvl and B-Spline Layers.](image1)

![Figure 4-15. Save Layer Optical Constants dialog box.](image2)
Save Layer Optical Constants

Alternatively, the layer optical constants can also be saved directly from a right-click menu over the Layer name (in this case “B-Spline”). This menu is shown in Figure 4-16. After pressing this selection, the user is asked whether to save the Parameterized optical constants (values used by equation to reconstruct the optical constants) or to save a tabulated list of optical constants at each measured wavelength.

![Figure 4-16. Right-click over a layer name to choose “Save Layer Optical Constants” from a drop-down menu.](image)

![Figure 4-17. Choose whether to save the Parameterized optical constants (values used by equation) or a tabulated list of optical constants at each wavelength.](image)

Au Substrate Model

After optimizing the analysis for a new sample, the model can be saved for future use. Let’s use this Au substrate as an example. If additional Au substrates will be measured in the future, a customized model would help reduce analysis time. For this example, let’s modify two additional settings:

- Replace the “Au” reference in the B-Spline layer with the metal optical constants saved in this example (previous step).
- Expand FIT Options and turn Include Derived Parameters ON. Add derived parameters for both ‘n’ and ‘k’ of the substrate (change layer # to
“0”). This will report \(n\) and \(k\) at the specified wavelength (default = 632.8 nm) in the Fit Results.

Press the ‘Save’ button at the top of Model: panel to save the final model as “metal substrate example” as in Figure 4-18. We will use this model in the next example.

NOTE: The Layer Command: Save is different than the ‘Save’ button in the Model: panel. The Layer Command: Save saves the layer optical constants, while the ‘Save’ button in the Model: panel saves the layered model structure and any +MODEL, +FIT, or +OTHER Options.

Layer Commands: Add Delete Save
Include Surface Roughness = OFF

- Substrate = B-Spline
  Init. values: \(n = 1.500\) \(k = 0.000\) Starting Mat = au_test
  Resolution (eV) = 0.100 19 Pts. (1.386-3.250 eV) Draw Node Graph
  Fit Opt. Const. = ON
  Use KK Mode = OFF
  Show Advanced Options = OFF

Angle Offset = 0.000
+ MODEL Options
- FIT Options
  Perform Thickness Pre-Fit = OFF
  Use Global Fit = OFF
  Fit Weight = N.C.S
  Limit Wvl. for Fit = OFF
  Include Derived Parameters = ON
  Add Derived Parameter
  1. Type = \(n\) Layer # = 0 Wavelength = 632.8 nm Name = \(n\) of B-Spline @ 632.8 nm
     Low Spec. = 0.000 High Spec. = 0.000
  2. Type = \(k\) Layer # = 0 Wavelength = 632.8 nm Name = \(k\) of B-Spline @ 632.8 nm
     Low Spec. = 0.000 High Spec. = 0.000

+ OTHER Options
  Configure Options
  Turn Off All Fit Parameters

Figure 4-18. Au Substrate model saved for future use.
4.2. **Self Assembled Monolayer on Au**

**FEATURES IN THIS EXAMPLE**

- Parameter Uniqueness
- Fixing B-Spline Optical Constants

This example uses the Au substrate result from Section 4.1. Thus, you will need to complete 4.1 before proceeding to this example. To ensure that the Au substrate model works before proceeding, open the “Au_substrate” data file again and press fit. If the fit does not work (error message appears, Fit Result returns “MSE out of spec”, model curves do not match experimental curves, etc.), then repeat Section 4.1 before proceeding.

**Self-Assembled Monolayers**

Self-Assembled Monolayers (SAMs) are very thin organic layers that consist of a single monolayer. Thus, they are generally less than 10 nm thick and often closer to 1-2 nm. Spectroscopic Ellipsometry is very sensitive to layers of this thickness, due to the phase information available from the measured Delta parameter. However, the index of refraction from such thin layers is not typically attainable. Instead, the nominal index of refraction must be assumed in order to fit the thickness. As layers become thicker than 10-20 nm, spectroscopic ellipsometry gains sensitivity to index of refraction in addition to thickness. A common method for thinner layers is to measure a similar material as a thicker film. From the thicker film, the index can be measured and then fixed when measuring thinner layers. Because this approximation may not always be true for such thin layers, the measurement of SAMs is often qualitative and best used for comparison between multiple samples.

Open the “Organic on Au” data file and press ‘Generate’. The graph should appear as in Figure 4-19. The Model dashed curves will represent a bare Au surface, as they come from our model with only Au. Notice a shift in Delta toward lower values (at all wavelengths). This indicates the presence of a thin surface coating. A larger shift indicates a thicker film on the Au surface.

![Figure 4-19](image-url)

*Figure 4-19. Data from “Organic on Au” compared to Generated Data from the bare Au surface. Notice the shift in Delta toward lower values for the coated sample, which indicates a surface coating.*
Add a new layer to the Model by clicking on the Layer Command: **Add**. Position the blue bar above the substrate (Au represented by B-Spline in this model) and click to add the layer. Choose Cauchy.mat from the Basic folder and press ‘Open’. The default values for Cauchy layer will be close to the values needed for generic organic films. Thus, we will not vary the optical constants for the Cauchy layer and only turn on the thickness as a fit parameter.

**NOTE:** Before fitting, it is important to set the Au optical constants so they do not vary during the fit. Without this step, the Au optical constants would also move and confuse our overall result.

To prevent the Au substrate optical constants from varying during the fit, make sure to do the following:

- Turn OFF the “Fit Opt. Const.” command inside the B-Spline layer, as shown in Figure 4-20.
- Ensure the Global Fit (under +FIT Options) is turned off.

![Figure 4-20](image)

Figure 4-20. Turn OFF the “Fit Opt. Const.” to ensure the Au optical constants stay fixed at the values fit from the bare substrate data.

Now we are ready to determine thickness of the SAM on Au. Turn on the Thickness of the Cauchy layer as a fit parameter (by clicking it with the right mouse button). Press ‘Fit’ and the resulting thickness of 7.48 nm should be reported. If you get a different thickness, make sure the Au optical constants did not adjust from the previous fit.

**NOTE:** Thickness units can be changed from the Options tab. Display Units: panel.

### Fit Parameter Uniqueness

We will use this result to demonstrate a feature in CompleteEASE that helps to visualize the model uniqueness. Go to the bottom of the Model: Panel and expand the +OTHER Options. In this section, press the Fit Parameter Uniqueness command. You will be asked which of the fit parameters you want to select for testing. In this case, we have only fit the thickness, so choose Thickness #1, as shown in Figure 4-21.
Figure 4-21. Choose fit parameter to test for Uniqueness.

Next, you need to enter the “Minimum Value”, “Maximum Value”, and “# of fit points”. These will define the test conditions. The parameter space between the minimum and maximum will be divided into equal spacing to match the total number of points entered. Then, the fit will proceed with the test parameter (Thickness #1 in this case) fixed at each individual value. The MSE is captured from each fit for graphing.

For this example, choose a Minimum Value of 2.5nm, Maximum Value of 15nm, and 100 total points, as shown in Figure 4-22. You will get a warning that no other fit parameters are selected, as shown in Figure 4-23. This is okay, as the test will still proceed to show the MSE profile as the Cauchy thickness is varied, as shown in Figure 4-24.

Figure 4-22. Settings for Parameter Uniqueness Test.

Figure 4-23. Warning message shown when the only fit parameter turned on is the test parameter. It is okay to proceed past this warning to see the MSE profile as that single fit parameter is varied.
Right-click on the graph area and copy the MSE profile data to the Graph Scratchpad. This will allow us to compare this MSE curve with our next test.

**NOTE:** You may want to clear the Graph Scratchpad (if results from previous examples are still present) before copying the Uniqueness test result to the Scratchpad. To do this, open the Scratchpad and click ‘Delete All’.

Next, allow the A, B, and C parameters in the Cauchy layer to fit by expanding the Cauchy layer (+) and right-clicking the A, B, and C parameters. Repeat the Parameter Uniqueness test for Thickness #1 with the same test conditions as before. This time, the result should appear as seen in Figure 4-26. Add this result to the Graph Scratchpad and then view the Scratchpad to allow comparison of the two tests as shown in Figure 4-27.

Figure 4-25. Model for SAM over Au with Cauchy parameters A, B, and C added as fit parameters.
Figure 4-26. MSE profile from Uniqueness test of the thickness when the Cauchy parameters for thin layer are allowed to vary at each point.

Figure 4-27. Comparison of MSE profiles from both tests, as seen in the Graph Scratchpad.

The shallow curvature of uniqueness test result for thickness and Cauchy parameters indicates poor sensitivity to both thickness and refractive index (determined by Cauchy equation). This is typical for thin films less than ~10nm. When working with ultra-thin organic films (<10nm), it is recommended to fit for thickness only, while setting refractive index at reasonable values (i.e. default Cauchy parameters).
4.3. Absorbing Thin Films (B-Spline)

**FEATURES IN THIS EXAMPLE**

- B-Spline Layer
- Starting Mat =
- Saving a Model
- Wvl. Range Expansion Fit
- Selecting a Graph/Fit Range
- Zoom All

SE measurements on absorbing films are more difficult to analyze compared to transparent films. This is because the general structure of the film optical constants \((n,k)\) versus wavelength is not always known before the measurement. To handle absorbing film analysis problems, CompleteEASE implements a couple special layer types: a “B-Spline” layer which allows arbitrary flexibility in \(n,k\) versus wavelength, and the “Gen-Osc” layer which uses a summation of standard oscillator lineshapes. The B-Spline is described in this example, while the Gen-Osc will be described in the next example.

The B-Spline layer is perfect for materials that are partially transparent and partially absorbing – such that the ellipsometer witnesses the onset of absorption. A few example materials of this nature include:

- a-Si, a-Ge, a-SiGe
- ZnS, ZnSe
- SiC
- TiO\(_2\)
- Most organics, such as polyimide, PEDOT, P2VP, Alq3, P3HT, photoresists, and many more.
- Diamond-like carbon (DLC)

**Identifying an Absorbing Film**

Open the “SiC on Si” data file from the EXAMPLES folder. Notice that Psi and Delta oscillate (Figure 4-28) as we have seen for transparent film examples. These oscillations are caused by interference between light that reflects from the film surface and light that travels through the film and returns from the bottom interface. The thicker the film is, the more oscillations that will appear.
Figure 4-28. Data from SiC film on Silicon. The interference oscillations in data suggest the film is transparent at longer wavelengths, while the decrease in amplitude of these oscillations at short wavelengths indicates UV absorption.

Unlike the transparent films we have witnessed, the interference oscillations decrease in amplitude toward shorter wavelengths. This is a strong indication of absorption in the ultraviolet spectrum. The oscillations are dampened, as less light is able to penetrate the film and return to the surface to interfere with the top-reflected light. This concept is illustrated in Figure 4-29. To model the SiC film on Silicon will require a material file that can describe both transparent and absorbing regions.

Figure 4-29. Interference oscillations occur when the light can travel through a thin film and return from the bottom interface (red). If the light is absorbed, the light only comes from the top-surface (blue) and will not produce interference.

### Si with Absorbing Film

Now, open the “Si with Absorbing Film” model from the BASIC folder and click ‘Fit’. The results should appear as shown in Figure 4-30. To view the optical constants of the film, right-click (\textbullet\textcircled{R}) on the B-Spline layer in the model and select “Graph Layer Optical Constants”. The resulting optical constants (which are the optical constants for the SiC film) are shown in Figure 4-31. The film is essentially transparent ($k\approx0$) at longer wavelengths ($\lambda>600\text{nm}$) with a gradual increase in absorption at shorter wavelengths. The film index (\textit{n}-red curve) also increases toward shorter wavelengths.
Figure 4-30. Data fit to SiC on Si using the Si with Absorbing Film model.

Figure 4-31. Optical constants for SiC film using the “B-Spline” layer.
The “B-Spline” Layer

The “Si with Absorbing Film” model settings are shown in Figure 4-32; the “+” next to “Layer #1” was clicked to expand the B-Spline layer. The B-Spline layer specifies the optical constants versus wavelength using a series of control points which are equally spaced in photon energy (eV). The approximate spacing of the control points is controlled by the “Resolution (eV) =” field and the resulting number of control points within the current spectral range is listed after this value. Basis (“B.”) spline curves (which are by definition smooth and continuous in the 0th, 1st, and 2nd derivatives) are used to interpolate the optical constants between the control points.

![Figure 4-32. Settings for the “Si with Absorbing Film” optical model.](image)

Clicking on **Draw Node Graph** displays the graph shown in Figure 4-33. This graph shows the location of the spline control points. The B-Spline layer always works in terms of the complex dielectric function \( \varepsilon = \varepsilon_1 + i\varepsilon_2 = (n + ik)^2 \), which is often written as “\( e_1 & e_2 \)”. One set of control points (and associated spline curve) is used to specify the dispersion in “\( e_1 \)” and a second set of control points is used to describe “\( e_2 \)”. Using a smaller “Resolution” value adds more control points, resulting in more flexibility to accommodate sharper features in the dielectric constant spectra. However, setting the Resolution too low can result in unwanted noise or unphysical artifacts in the dielectric function spectra.
There are two ways to initialize the B-Spline layer control point values:

1. Click on the “n=” and/or “k=” fields which sets all control points to this new value over the entire spectral range.
2. Click on “Starting Mat=” which chooses control points to match the optical constant values from the specified material file.

The “Si with Absorbing Film” model uses the first method combined with the “Global Fit” option to search a range of \( n \) and \( k \) values for the film. The “Use Global Fit” section in Figure 4-32, specifies twenty \( n \) values ranging from 1.3 – 5 and three \( k \) values ranging from 0 – 0.5.

**NOTE:** The Global Fit can be a slow procedure and should be turned off if you have a good starting guess for the optical constants. With good starting values, you can save significant time – especially when dealing with multi-angle data sets.

The “Fit Opt. Const.” button allows the user to turn on and off the fitting of spline points from within the B-Spline. Thus, an absorbing material could be fit using the B-Spline and then the “Fit Opt. Const.” button can be toggled to OFF to fix the optical constants of this layer in place for further analysis.

**NOTE:** To fix/set optical constants, the Global Fit also needs to be turned off

In the B-Spline layer, the “Use KK mode” is an advanced calculation that maintains Kramers-Kronig consistency between the resulting \( e_1 \) and \( e_2 \) spline curves. When this feature is ON, the optical constants will be constrained to a physical shape – thus it should provide further assurance that the resulting optical constants are correct. This feature will also reduce the total number of fit parameters, as only the \( e_2 \) spectra is described with a series of spline points while the \( e_1 \) spectra is calculated via the Kramers-Kronig transformation. Additional “Advanced Options” in the B-Spline layer are further documented in Section 10.
Wavelength Range Expansion (WVL-EXPAND) Fit

The “Si with Absorbing Film” model uses a Wavelength Range Expansion (WVL-EXPAND) fit within the Global Fit. This is explained in the subsequent section. First, let’s demonstrate how the WVL-EXPANd fit routine works. The principle behind the WVL-EXPAND fit is that a model may work better over a narrower selection of wavelengths as compared to the entire measured range. Organic films are an excellent example as they are typically transparent at longer wavelengths with absorption only in the UV. Thus, a Cauchy is often adequate for longer wavelengths but is not applicable to shorter wavelengths when the film becomes absorbing. The B-Spline layer can work in both transparent and absorbing regions, but may find the correct starting solution more readily in the transparent region. The transparent region minimizes the fit complexity as only the index needs to be determined.

Range-Selecting Wavelengths

To manually perform a WVL-EXPAND fit, we first need to learn how to select a subset of data over a specific wavelength range. This is simple in the CompleteEASE program. First, choose a wavelength on the graph that will serve as an end-point for the new data subset. Left-click (L) the mouse on this wavelength and hold the mouse button down while dragging to the opposite end-point wavelength. If the mouse drags off the side of a graph, all wavelengths in that direction will be included (even expanding the graph to include wavelengths not shown).

As an exercise, open any data file and try the following:

1. Range-select 600-700nm using the mouse (Figure 4-34).

![Figure 4-34. Range-selecting wavelengths between 600 and 700nm.](image)

As you drag the mouse across the graph with the left-button held down, it should create a “gray area” on the graph to show the range of wavelengths selected.

2. Range select wavelengths of 600nm and above. For this, start the mouse-button just inside the current graph (near 600nm) and then drag beyond the right graph boundary to incorporate all longer wavelengths (Figure 4-35).
3. Finally, select the entire wavelength range. This is accomplished by dragging the mouse over both graph boundaries or pressing ‘Zoom All’ at the top-right of the graph panel (Figure 4-36).

![Figure 4-35. Expanding the current wavelength range to longer region.](image1)

![Figure 4-36. Press ‘Zoom All’ to select the entire wavelength range.](image2)

Now, let’s return our attention to the WVL EXPAND fit. This fit works by taking a limited wavelength range (as selected by the user), applying the current fit, and then gradually increasing the range to both shorter and longer wavelengths until the entire data set is included in the fit. The fit is performed at each new increment of wavelengths.

To perform a WVL EXPAND fit, you choose the desired starting wavelength region, expand +OTHER Options, and click Wvl. Range Expansion Fit at the bottom of the Model panel.
WVL-EXPAND Fit with B-Spline Layer

The B-Spline layer works very well with the WVL-EXPAND fit. As the wavelength range is expanded, the number of spline control-points automatically increases to match the new wavelength range. The additional points are initialized by extrapolation of the existing model, but then are fit to match the experimental measurement. The true power of the WVL-EXPAND fit can be seen by extending this fit to include Global Fitting (as described in next section).

Global Fits with WVL-EXPAND Fitting

The WVL-EXPAND fit can also be used in conjunction with the Global Fit. When this option is “ON”, a WVL-EXPAND fit is performed at every trial of the global fit. Thus, each fit will begin over a restricted range of wavelengths, followed by an expansion of wavelengths to the entire range. The MSE of the final expanded fit is compared between all trial fits to determine which result gives the best agreement. The best fit from all the Global Guesses is reported.

- FIT Options
  + Perform Thickness Pre-Fit = ON
  - Use Global Fit = ON
    # of Data Points = 20 # of Iterations = 5
    Parm #1 = Spline Layer n
    Min. = 1.300 Max. = 5.000 # Guesses = 20
    Parm #2 = Spline Layer k
    Min. = 0.000 Max. = 0.500 # Guesses = 3
    Parm #3 = (none)
    Include Wvl. Range Expansion Fits = ON
    Limit Wvl. for Fit = ON Range = 600.0 nm - 5000.0 nm
    Fit Weight = N.C.S
    Limit Wvl. for Fit = OFF
    Include Derived Parameters = ON

Figure 4-37. Global Fit and Wavelength Range Expansion Fit regions of the model for absorbing film using B-Spline layer.

Figure 4-37 shows the additional options available when a WVL-EXPAND fit is turned “ON”:

Limit Wvl. for Fit

This first option chooses whether to start with the user-selected wavelength range (OFF) or the wavelength range chose within the model (ON). Typically, this option would be turned ON for automatic models, as it does not require the user to decide where to start. However, it can be turned OFF when first testing a model to determine how to setup the initial wavelength range.
Range

When the “Limit Wvl. for Fit” option is turned “ON”, this selection dictates the starting wavelength range for the WVL-EXPAND fit. This option is helpful when automating a model.

The Global fit with WVL-EXPAND fit is a very powerful data analysis algorithm which works for a wide range of samples and can be used to automate the fitting process. Here are some hints for “tuning” the algorithm to analyze data sets when the default values fail:

- Adjust the “Spline Layer n” and “Spline Layer k” ranges to values which closely bracket the nominal optical constants.
- For thicker films (i.e., data sets with many oscillations) it may be necessary to increase the “# Guesses”; thinner films can use fewer “# Guesses”, which will reduce the data analysis time.
- For thin films (<100 nm) the WVL-EXPAND Fit may not be required.
- For very thick films (>1 – 2 microns), it may be necessary to specify a more limited spectral range (it is preferable that the film is transparent in the limited spectral range).
- Unless the nominal layer thickness is accurately known (and set in model), “Perform Thickness Pre-Fit” option should be ON.
- Before beginning the global fit, make sure all other model parameter values are set appropriately (for example, if Surface Roughness is included in the fit, its starting value should probably be zero).
4.4. Absorbing Thin Films (Gen-Osc)

FEATURES IN THIS EXAMPLE

- Gen-Osc Layer
- Tauc-Lorentz
- Parameterize Layer
- Cody-Lorentz

In the previous example, we introduced the analysis of absorbing thin films via the B-Spline layer. Another method for absorbing films that can be used independently or after initial B-Spline modeling is the Gen-Osc layer. The Gen-Osc models the absorbing features by describing the shape of the absorption. This example demonstrates the common application of the Gen-Osc layer.

Parameterize Layer

The resulting B-Spline optical constants from the “Si with Absorbing Film” model fit may be completely adequate for many applications. While the B-Spline layer guarantees the optical constants versus wavelength will be “smooth”, it does not always guarantee they are physical (that is, Kramers-Kronig consistent). Furthermore, the B-Spline layer requires many fit parameters (two times the number of control points), which may lead to correlation in the data fit. Once the general shape of the optical constants are determined using the B-Spline layer, it is often desirable to parameterize the optical constants with a simple analytic expression. The “Parameterize Layer” command can conveniently perform this procedure.

NOTE: The goal of parameterizing a layer is to match the shape of n,k from the B-Spline fit using a dispersion model.

This example uses the SiC on Si result from Section 4.3. Thus, you will need to complete 4.3 before proceeding to this example.

To parameterize the B-Spline optical constants for the SiC film in the previous example, right-click on the B-Spline layer and choose “Parameterize Layer” from the drop-down menu, as shown in Figure 4-38. This opens “Parameterize Layer” dialog box as shown in Figure 4-39, with the B-Spline layer optical constants plotted in the graph. The default model (Gen-Osc) will automatically open in this window with a starting oscillator (in this case, Cody-Lorentz).
Figure 4-38. Right-click on a layer to drop-down menu, then press “Parameterize Layer”.

Figure 4-39. Parameterize Layer dialog box, showing starting optical constants of the previous B-Spline layer (blue, black curves) along with the default Cody-Lorentz oscillator (red, green curves).
The “Gen-Osc” material file (which stands for General Oscillator) allows a combination of various oscillator lineshapes. We want to match the shape of SiC optical constants with an oscillator. There are many choices which are detailed in Chapter 10. For this example, left-click on Cody-Lorentz and choose “Tauc-Lorentz” from the Input box, as shown in Figure 4-40.

![Input box with choices of different oscillator types.](image)

Figure 4-40. Input box with choices of different oscillator types.

The basic steps for matching optical constants with a new oscillator model are as follows:

1. Match e2 by adjusting and then fitting the oscillator parameters.
2. Match e1 by adjusting and then fitting the e1offset or Pole values.
3. Final adjustment to all parameter to fit both e1 and e2.

Each of these three steps will be demonstrated for the SiC example, where we match the SiC B-Spline results with the Tauc-Lorentz oscillator parameters.
Step 1: Match Imaginary Part Only.

Most oscillators are Kramers-Kronig consistent. This provides an internal coupling between the real ($e_1, n$) and imaginary ($e_2, k$) optical properties. As we adjust the oscillator parameters to match the shape of $e_2$ from our B-Spline result, we will in-turn be automatically adjusting the $e_1$ shape via the KK transformation. Step 2 will allow us to adjust additional parameters that will only affect $e_1$, which is why it is important to start with the $e_2$ matching.

To control the oscillator values, LEFT-click your mouse on the number next to the Tauc-Lorentz oscillator, as shown in Figure 4-41. Note that the selected oscillator will now have a shaded box surrounding the number.

**NOTE:** It is important to LEFT-click the mouse on oscillators you wish to adjust. RIGHT-clicking will delete that oscillator from the list (with warning before permanent deletion). If you do accidentally delete, simply press Add Oscillator to replace.

When an oscillator is selected, control boxes will appear in the graph to allow manipulation of the center energy, amplitude, broadening and possibly the bandgap. Not all of the control boxes will appear if the oscillator values are outside the graphed range. To view all of the control boxes, as shown in Figure 4-42, press the ‘Expand X-Axis’ button at the bottom of the dialog box.

![Figure 4-41](image)

*Figure 4-41. Left-clicking on the number next to an oscillator selects that oscillator for manipulation.*
Before we adjust any of the control points, it is easier to visualize the oscillator if we change the OC View/Fit Type to “Imaginary Part Only”. This will show the green curve of e2 (or k) from our B-Spline fit along with the dashed black curve from the oscillator we are adjusting. The goal is to match these two curves. If you position your mouse over any of the gray control boxes, your pointer should change to an XY arrow to show that you can now adjust that control box. This is shown in Figure 4-43, where the top control box is selected. This top control box can move both the amplitude and the center energy. Practice moving the different control boxes to better match the dashed curve to the green B-Spline data.
Figure 4-43. Cursor positioned above the top control point turns to an XY Arrow to show that this control point can be adjusted to change the shape of the selected oscillator.

After the oscillator parameters have been adjusted to better match the reference $e^2$ shape, right-click (\textcircled{R}) on the “Amp.”, “Br”, “Eo”, and “Eg” parameters. This will turn them ON as fit parameters. Click the ‘Fit’ button. In this case, we are not fitting the experimental data, but rather matching the imaginary oscillator shape (via the parameters) to the reference (B-Spline) optical constants.

The fit varies the oscillator parameters to match the imaginary curve shape as entered from the B-Spline. Figure 4-44 displays the fit results; the black dashed curve in the graph is calculated using the Tauc-Lorentz dispersion formula.
Step 2: Match Real Part Only.

When finished matching the imaginary part with an oscillator, change the OC View/ Fit Type to “Real Part Only”. The oscillator list will be hidden from view, as the only parameters allowed to vary during “Real Part Only” fitting are the Einf, and two Poles, as shown in Figure 4-45. Poles are unbroaded oscillators that are placed outside the measured spectral range to affect the real part of the optical constants in the same manner that a strong absorption in that spectrum would. Thus, the UV pole will pull the index up toward the short wavelengths while the NIR pole will pull the index down toward longer wavelengths. The Einf value adds a constant offset to all index values, which approximates the effects of absorptions that are significantly distant from the measured wavelengths.
The only parameters required when fitting “Real Part Only” are the $E_{inf}$ and two Poles.

For this example, it is best to adjust the parameter values before starting the fit. When values are changed in the **Layer Parameters** panel, the optical constant graph is automatically updated. Click on the UV Pole Amp. parameter, enter 100, and the curve raises upward. The parameters can also be “dialed in” by positioning the mouse above a parameter, holding down the “Shift” key, and rolling the mouse wheel. Figure 4-46 shows the curves after the UV Pole Amplitude has been raised to get closer to the reference curve. However, the “tilt” of the curve does not match. This is because the UV Pole Position may also need to be adjusted. In general, the closer the UV Pole energy to the measurement range, the more “tilting” that will occur.
Figure 4-46. Increasing the UV Pole Amplitude also raises the Gen-Osc “e1” value, but the blue curve does not match the “tilt” of the red reference curve from our B-Spline. This will require adjustment of the UV Pole Energy.

After adjusting the UV Pole Position and Energy to better match the reference B-Spline curve, the two values are “fit” to give the best match. Afterward, the Einf value was also added to the fit to get a good match, as shown in Figure 4-47.
Step 3: Match All.

If Step #1 and #2 were successful, the Gen-Osc will not match both real and imaginary values of the reference. However, there can be some correlation between values used to match the e2 curve and those used to match the e1 curve. Thus, a final fit is often recommended with the OC View/Fit Type set to “All”. The most common correlation occurs when one of the oscillators is centered outside the measurement spectral range and thus can contribute in different ways to the e1 value (via KK transformation), as the amplitude and center energy can adjust and still provide a similar “tail” of absorption in the measured wavelength range.

Replace Layer

After you are happy the Gen-Osc matches both real and imaginary components of the reference B-Spline, click ‘Replace Layer’ to apply the new oscillator model in place of the B-Spline layer, as seen in Figure 4-48. (If the parameterization is not successful, press ‘Cancel’ to revert to the existing layer optical constants.)

The new oscillator can now be used to fit data. Press ‘Fit’ to see how well the Tauc-Lorentz oscillator can describe the experimental data. If further samples with similar optical constants are measured, this model could be used directly – skipping the B-Spline layer step.
NOTE: Fitting within the Parameterize Layer dialog box is different than fitting your measured data. In the Parameterize Layer dialog box, you are fitting reference optical constants to obtain good starting values for the oscillator parameters. Fitting your measured data occurs from the **Fit** panel of the **Analysis** tab – here the oscillator parameters adjust to find the best match to the experimental SE data.

Figure 4-48. Gen-Osc has replaced B-Spline layer in the model.
Glass with Absorbing Film

In this example, we will repeat the demonstrated procedures with an absorbing film deposited on glass substrate. In general, the data analysis is similar, with different substrate optical constants. Another consideration for transparent substrates is the possible presence of backside reflections. In this example, the backside of the glass substrate was mechanically roughened to avoid this extra complication. However, section 3.4 described methods to handle transparent substrates when backside reflections are present.

Open the “a-Si on Glass” data file and the “Glass with Absorbing Film” model, and click the ‘Fit’ button. The results should appear as shown in Figure 4-49. The “Absorbing Film on Glass” model uses the same procedures as previously described for the “Si with Absorbing Film” model. As before, the optical constants are described via a spline with equal spacing of 0.3 eV between nodes. This spacing can be changed for materials that exhibit stronger optical dispersion. The initial values are determined via a global fit of n and k with WVL-EXPAND fitting. To view the optical constants of the film, right-click the mouse on the “B-Spline” layer in the model and choose “Graph Layer Optical Constants”. The resulting “B-Spline” optical constants from the fit are shown in Figure 4-50.

Figure 4-49. Results using “Glass with Absorbing Film” model to fit the “a-Si on Glass” data file.
The two models for absorbing films in this example are very flexible and can be used for many different materials. This is primarily due to the Global Fit of $n,k$. However, the data analysis takes longer than average, as the global fit attempts all possible combinations of $n,k$ starting values. To minimize analysis time, the global fit range could be reduced. However, this would require knowledge of the appropriate range for the material. If the material is known, it is also possible to use a similar material file as the “Starting Mat” inside the B-Spline layer.

For best comparison of analysis time, re-analyze the “a-Si on Glass” data set using the same model by simply pressing “Fit” again. Note the total time required to complete the fit. The analysis takes between 20 and 25 seconds on a Pentium D computer. Re-open the “Glass with Absorbing Film” model and make the following modifications:

- Expand the B-Spline layer and left-click “Starting Mat = none”. Open “a-si.mat” from the Semiconductor Folder.
- Turn OFF the global fit with Model: >>FIT Options.
- Press Fit: >>’Fit’.

The data analysis should return the same result as before, but in less than one second. Thus, this new model would be an alternative for a-Si when the optical constants for the film are similar to the reference material file.
Parameterizing Optical Constants with Cody-Lorentz

The a-Si optical constants can be parameterized, just as with the SiC values from the previous example. There are two common oscillators used for amorphous silicon: Tauc-Lorentz and Cody-Lorentz. To parameterize the B-Spline optical constants for the a-Si film in this example, right-click the B-Spline layer and then choose “Parameterize Layer” from the drop-down menu.

The “Parameterize Layer” dialog box appears with the B-Spline layer optical constants plotted in the graph. For this example, we will use a single “Cody-Lorentz” oscillator.

As before, go through the three steps with the Gen-Osc parameterization, namely:

1. Match \( e^2 \) using oscillator parameters (set View/Fit to Imaginary Part Only and press “Fit”). Amplitude (Amp), broadening (Br), center energy (Eo), and bandgap (Eg) are the most significant parameters in this oscillator. Start with these four as fit parameters; others may be added later in the example.

   For this step, there are two ways to manipulate the oscillator parameters before fitting. Try both ways to gain familiarity with the oscillator and process:

   A. Dial each oscillator parameter by holding the SHIFT key down while the mouse-wheel is rolled over top of the oscillator parameter. Holding CTRL-SHIFT will make reduce the step-size of each adjustment.

   B. Left-Click (L) on the number next to the Oscillator to highlight that oscillator. This shows “control” points on the graph that can be grabbed and adjusted with the mouse. Often, these control points are outside the measured spectral range, as in Figure 4-51. To view the control points of the oscillator, press the ‘Expand X-Axis’ button. This will widen the photon energy range to view the entire oscillator, as seen in Figure 4-52. Try adjusting the oscillator broadening, amplitude, and center energy by moving the gray control boxes on the graph. This will change the shape of the oscillator.
Figure 4-51. Click the number next to an oscillator to view the oscillator and its control points (in gray). As is often the case, the oscillator is outside the measured energy range, so it is difficult to locate.

Figure 4-52. Press ‘Expand X-Axis’ to better view the oscillator and control points (in grey). The control points can be adjusted with your mouse to change the amplitude, center energy, broadening and bandgap for the selected oscillator.
2. Match $e_1$ with poles and $E_{\text{inf}}$ (set View/Fit to Real Part Only and press "Fit"). In this example, we will only use the UV pole.
3. Match both (set View/Fit to All and press “Fit”).

Figure 4-55. Step 3 result (before clicking “Replace Layer”).

After the parameterization is complete, click “Replace Layer” and click Fit from the main window to fit the example data.

Figure 4-56. Fit result after replacing the previous B-Spline layer with new Gen-Osc layer.
Figure 4-56 displays the fit results; the black dashed curves in the graph are calculated using the specified oscillator. MSE is near 1.2.

Next, expand the oscillator layer in the model and add the following fit parameter to the fit: “Ep”. This allows added flexibility to the optical dispersion. The MSE should reduce to 0.8 as shown in Figure 4-57.

![Figure 4-57. Fit result after adding Ep parameter.](image)

Add the additional fit parameters (Einf, Et, and Eu) to the model to check for further improvement. The question becomes whether slight MSE improvement warrants the increased number of “variables” in the modeling process. As a general “rule-of-thumb”, each additional fit parameter should decrease the MSE by 20% to make it a clear choice.

NOTE: Due to correlation between oscillator parameters, it is likely that your results will not exactly match the above results. In this case, it is common practice to try and remove parameters until a minimum number of parameters is selected that still provides a good fit.
Saving a Model

After the appropriate Cody-Lorentz fit parameters are determined for the a-Si film, the model can be saved for future use. This process is quite simple: press **Analysis > Model:** ‘Save’. Choose the Folder where you would like to locate this model and give the model a name. The “Common” folder is a central location for easy access by all users. Figure 4-58 demonstrates this procedure. You will need this model for the next example.

![Figure 4-58. Press ‘Save’ from the Model: panel to bring up the “Save Model” dialog box.](image)

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4.5. Multiple Layer Film Stacks

NEW FEATURES IN THIS EXAMPLE

• Modifying a Model
• Global Fit/Pre-Fit with multiple layers

Open the data file “a-Si Multilayer”. This sample is similar to those discussed in Section 4.4. The substrate is silicon with a SiO$_2$ film underneath the a-Si. No model exists for this type of sample, so it will need to be constructed. It is preferred to start by modifying a similar model. Let’s start with the model saved in Section 4.4 for a-Si on glass using a Cody-Lorentz oscillator. Open this model from the “Common” location. If you did not save the model, then repeat Section 4.4 before proceeding and make sure to save the final model as suggested.

NOTE: Make sure to replace the substrate with silicon.

Left-click on the “7059_Cauchy” layer and open “Si_Jaw” from the Semiconductor location of the Open Material dialog box.

Select the Add command from the top of Model panel. This will bring up the dialog box shown in Figure 4-59. Move the mouse to locate the Blue bar between the two current layers (as shown). This will be the location of the newly added layer. Choose “SiO2_Jaw” from the Semiconductor location of the Open Material dialog box.

![Add Layer To Model](image)

*Figure 4-59. The Add command opens the dialog box shown. The blue bar can be moved up or down to designate the location of the added layer.*

Left-click the “Thickness #1= 0.00nm” designation for the SiO$_2$ layer to change its thickness to a nominal value of 100nm. Also select this thickness as a fit parameter. Generate data and the graph should appear as shown in Figure 4-60. The generated data does not match the experimental data. If a fit is attempted from this point, it will not be successful. Turn off the Global Fit in the Model and then press ‘Fit’. Go ahead and reset the fit using the ‘Reset’ button. If you forgot to turn-off the Global Fit, it may revert back to the spectral range-limited fit used by the Global Fit. In this case, right-click on the ‘Reset’ button to choose the “Show Reset List”. You can then choose to reset to the point right before you pressed ‘Fit’.
NOTE: Press ALT-R to reset the model to the previous values after verifying the poor fit quality.

The poor fit is due to the unknown thickness values. There are a couple approaches to determine better starting thickness values for each layer. Both approaches work better if the Cody-Lorentz oscillator parameters are turned OFF before proceeding:

- Expand General Osc layer and right-click on each of the fit parameters that was previously turned on, toggling them off.

Three layer thicknesses remain as fit parameters (SiO2, a-Si, roughness).

Figure 4-60. Generate data with a nominal thickness of 100nm for SiO2 layer.
**Approach #1:** Use a Global fit for both Thickness #1 and Thickness #2. Make sure to turn off the Thickness Pre-fit. This method can take a very long time if the search includes a wide range of possible thicknesses.

**Approach #2:** Use the Global fit for the SiO\(_2\) layer thickness with the Thickness Pre-fit turned on. This reduces the global fit dimension and works much more quickly.

**NOTE:** Approach #2 will not work if the a-Si layer (Thickness #2) is used within the Global Fit rather than the SiO\(_2\) layer (Thickness #1). The Thickness Pre-fit always operates on the thickest layer in the model.

Figure 4-61 shows the successful fit of the sample. The current model does not include the oscillator parameters, so the a-Si film must have similar optical constants to the previous sample.

![Figure 4-61. Fit result for Layer Thicknesses only.](image-url)
- Include Derived Parameters = ON

Add Derived Parameter

1. Type = n  Layer # = 2  Wavelength = 632.8 nm  Name = n of Gen-Osc @ 632.8 nm
   Low Spec. = 0.000  High Spec. = 0.000
2. Type = k  Layer # = 2  Wavelength = 632.8 nm  Name = k of Gen-Osc @ 632.8 nm
   Low Spec. = 0.000  High Spec. = 0.000

Figure 4-62. The Fit Options section of Model: panel allow Derived Parameters to report the optical constants from any layer in the model. With multilayer films, the correct layer # must be entered.

The Cody-Lorentz parameters can now be turned on to improve the fit quality by allowing the a-Si optical constants to vary. Figure 4-61 also points out that the Derived Parameters are pointing to the wrong layer number. Within the Fit Options section of the Model panel, change the Layer number from #1 to #2 for both the Derived n and Derived k as shown in Figure 4-62.

Add the Cody-Lorentz parameters Amp, Br, Eo, and Eg back as fit parameters (right-click on their values). Turn off the Global Fit and Thickness Prefit (as the thicknesses are already well known) and press the ‘Fit’ button. The results should appear as shown in Figure 4-63. Save the new model for future use on similar samples.

![Image](image-url)

**Figure 4-63.** Final fit results for a-Si multilayer.
Oxide or Roughness?

There is often a question whether the surface is oxidized or rough. In the model above, surface roughness was implemented with a final value of 2.29 nm. Turn the roughness off and AddLayer of “SiO2_Jaw” on top of Layer #2. Fit the data. The final result should have SiO2 surface layer of 2.70nm with an MSE of 1.25. Thus, there is very little difference between roughness and oxide for this sample. This is true in general when the film thickness is below about 5 nm. In the case of silicon, it is more likely the surface is oxidized than rough, but both models work.

4.6. Multiple Data Types (SE + T)

NEW FEATURES IN THIS EXAMPLE

- Appending Data
- Fitting SE and Transmission data simultaneously
- Parameter Uniqueness
- Graph Scratchpad

A common approach to increase the information content when measuring absorbing thin films on transparent substrates is to supplement the SE data with Intensity Transmission data. This extra information can help ensure a unique result for thickness and optical constants of the thin absorbing layer.

As a demonstration, open the “Cr on glass_SE” data file from the EXAMPLES folder. This data was collected from a thin Cr film on glass, with the backside taped to suppress backside reflections. Therefore, we can open the “Glass with Absorbing Film” model. This model was described in an earlier example and is designed for films that may be nearly transparent at longer wavelengths, but exhibit UV absorption. While this model will likely work without modification for the metal on glass, there is a better approach to start. Perform the following before pressing fit:

- Turn off the Surface Roughness.
- Turn off the Global Fit under the +Fit Options
- Choose the Cr.mat file from the metals directory as the “starting mat=” inside the B-Spline layer.

Parameter Uniqueness

Now, you are ready to press ‘Fit’. Your initial result should look similar to that shown in Figure 4-64. Because there is large correlation, pressing fit again-and-again will result in different results with nearly the same low MSE. To visualize the MSE profile, use the Parameter Uniqueness option under the +OTHER Options section of the model. Figure 4-65 shows the results for a Parameter Uniqueness test of the thickness (Thickness #1) from 10nm to 50nm. Before proceeding, copy this curve to the Graph Scratchpad for later comparison. Although there is a minimum value for the MSE around 45nm, the MSE value is below 0.4 for most of the thickness range.
Append Data

To help break this correlation apparent when only fitting the SE data, let’s add the Intensity Transmission to the fit to provide additional information. We want both the SE data and the Transmission Intensity data present in the fit. To this end, you need to right-click (⌘R) on the ‘Open’ data command and select “Append Data” as in Figure 4-66. Choose to append the “Cr on glass_T” data file from the EXAMPLES directory. This data was collected from the same sample. With both data files present, you can switch between graphs of SE and T data by choosing the data set right above the graph, as shown in Figure 4-67.
Figure 4-66. Right-click on the ‘Open’ command to choose “Append Data” when adding Transmission data to the already opened SE data.

Figure 4-67. Choose the data set to graph by selecting entry just above the graph. Here, you can see that both SE and T data for the Cr on Glass sample are open simultaneously. Thus, they will both be included in the fit.

Transmission Data % Weight

Any time Transmission Intensity data is fit, you should check the “Transmission Data % Weight”. This is the value that tells CompleteEASE how to weight the Transmission compared to Spectroscopic Ellipsometry data. Most models will have this value set to 100%, so the Transmission Intensity has equal weighting to the Spectroscopic Ellipsometry data. However, certain models can be saved with values other than 100%. To view the “Transmission Data % Weight”, you will need to press the Configure Options button at the bottom of the Model. In the Fit Options section, check the “Transmission Data Weighting” option, as shown in Figure 4-68.
After turning on the Transmission Data Weighting, you will need to set this value to 100% in the Fit Options section of the Model, as shown in Figure 4-69.

Go ahead and press ‘Fit’ again. If the model thickness starts close to the correct answer, the fit should find a thickness near 14nm. This is the final result. However, if the Cr thickness is started too far away from 14nm, the fit can get stuck in a “local” MSE minimum. Repeat the Parameter Uniqueness test to check the MSE profile, which should produce a curve as shown in Figure 4-70. It is obvious that there is only a single thickness that will provide the best MSE, although we can also easily see how it is possible to get stuck in the “local” minimum near 45nm.

Again, copy this curve to the Graph Scratchpad. Now, we can compare the MSE profiles from the previous two Parameter Uniqueness tests by viewing the Graph Scratchpad, as shown in Figure 4-71.
Figure 4-70. MSE profile when fitting both SE and T data simultaneously for a thin Cr layer on Glass.

Figure 4-71. Graph Scratchpad comparing the MSE profile when fitting both SE and T data or only the SE data from a thin Cr layer on glass.

This chapter includes further examples to demonstrate both advanced and specialized topics. Advanced topics include multi-sample analysis, creation of composition and temperature dependent material files and working with anisotropic or non-ideal models. A few of the examples use data from mapped samples, which help demonstrate how to work with data collected at multiple points from a sample. However, not all ellipsometers have this capability, so modeling of mapping data is only necessary for specific systems.

The section examples from this chapter are listed below, along with the primary CompleteEASE features that are discussed.

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### Section 5.3 Creating OC Library for Composition or Temperature-dependent Materials

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### Section 5.4 Anisotropic Films

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5.1. Map Data and Model Non-Idealities

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For this example, we will open a data file that contains multiple points scanned across a silicon wafer with thick SiO\textsubscript{2} coating. Our goal is to effectively determine the SiO\textsubscript{2} thickness and refractive index, along with the uniformity of the thin film properties across the wafer.

**Show Map Data**

Open the data file “SiO\textsubscript{2}_Map” from the EXAMPLES folder. You should see a color image of the Psi data across the center of a circular sample. To better understand what is shown, right-click on the graph panel and ensure “Show Symbols” is checked. This will mark the individual measurement locations, as in Figure 5-1. This uniformity map was created from a Radial scan of the wafer. We can visualize the uniformity of the ellipsometry parameters, but need to first analyze the data before we can visualize the thin film properties of interest.

![Figure 5-1. Map Data for “SiO2_Map” data file. With symbols shown, we can visualize each point on the wafer that was measured.](image)
Selecting a Point from the Map

If you press your left mouse button on any of the points of the map, it will load the spectroscopic ellipsometry data from that single location into the graph. This is a convenient way to choose a point to help construct a model to analyze the map. Go ahead and select a point with your mouse. The graph will change to Psi/Delta, as in Figure 5-2. Notice that an additional option is available at the top of the Graph Panel: “Show Map Data”. This allows you to go back to the colorful map of Psi and Delta. Make sure you select a single point before proceeding.

![Figure 5-2](Image)

*Figure 5-2. Spectroscopic Data from a single-point of the map. This view was gained by left-clicking the mouse on a map point.*

Silicon with Transparent Film

While this film is SiO₂, it is not necessarily a thermal oxide. Therefore, choose the “Silicon with Transparent Film” model that was first described in Section 3.3 for an unknown transparent layer. Go ahead and open this model and press ‘Fit’ which runs the model to fit the SINGLE point that is selected.

Before fitting all points on the map, we need to turn off the “Global Fit”, or the fit will take a very long time. The fit result should appear similar to Figure 5-3. The MSE is rather large which suggests there is a problem with this model. Thick films have a large number of interference oscillations, especially at short wavelengths. On this scale, it is difficult to see whether our model is adequate over the full spectrum.

Wavelength Units – eV

To better view short wavelengths, go ahead and convert the graph “Wavelength Units” to photon energy (eV) instead of nm. This will expand the UV region to give near-equal spacing between the interference oscillations. Wavelength and photon energy are inversely related. Thus, large photon energies are actually the shortest wavelengths. To convert to photon energy, either use the shortcut CTRL-ALT-W or go to the Options tab and change the Wavelength Units to eV within the Display Units panel. Figure 5-4 shows the data and model versus photon energy. For simplicity, I have turned off the Double-Y axis for the graph. It is now very clear that the short wavelengths (high eV) are not matched by the model. This is a problem with the final Cauchy parameters that were found.
Figure 5-3. Fit results for a single-point of the SiO2 map using the “Silicon with Transparent Film” model.

Figure 5-4. Data fit for Psi shown versus photon energy (in eV) to better visualize the discrepancies at short wavelengths (large eV).
Fixing Cauchy Dispersion

The Cauchy parameters work well to describe the index of refraction in the transparent region, but the Cauchy does have some limitations. It was not intended to cover very wide spectral ranges and it does not necessarily maintain a “physical” shape. The Cauchy parameters appear to fit the data well until about 4.5 eV. Above this range, there is not enough dispersion to match the correct interference oscillations. To fix this problem, we can either redo the fit with the B and C terms of the Cauchy included in the Global Fit (which will take a very long time) or range-select the region that is fit and slowly add more wavelength range. The second approach is demonstrated. First, range-select the data up to about 4.5 eV. Turn off the Global Fit and Thickness Pre-fit. They are no longer needed, as we are close to the final model. Press ‘Fit’. The Cauchy parameters will adjust to better match this range. Your result should look similar to those shown in Figure 5-5.

NOTE: For this example, do not worry about matching the manual results exactly, as you may have selected a different point from the uniformity map.

![Figure 5-5. Cauchy fit to 4.5eV.](image_url)

‘Set Ranges’

Next, extend the data range to 5.0eV. To add this extra range, let’s try using the Data: > ‘Set Ranges’ command. This will allow you to type in the new MAX photon energy. The ‘Set Ranges’ dialog box is shown in Figure 5-6. Enter Max: = 5eV. Click OK. Press ‘Fit’. Keep adding extra photon energy in 0.5eV increments and fitting the data until you have all data graphed. The final fit should appear as in Figure 5-7. The MSE is significantly improved – near 40 – compared to the previous MSE of over 200.
Figure 5-6. ‘Set Ranges’ command allows you to set the wavelength range and choose which data point you have selected.

Figure 5-7. Final fit to all wavelengths using Cauchy dispersion.

Sellmeier

Although we are able to get the Cauchy to better match the data with a little work, it is not designed to handle such a wide spectral range. A better model for transparent films that cover a wide spectral range is the Sellmeier relationship. It has the added benefit of remaining Kramers-Kronig consistent – thus ensuring “physical” shape for the resulting index of refraction.
Replace the Cauchy layer in your model by pressing the Cauchy name with left-
mouse click and go to the Dielectrics directory. Choose “SiO2 (Sellmeier)” material
file in place of the Cauchy. Press ‘Generate’ and the starting point should be close to
the experimental data, as in Figure 5-8. Go ahead and fit the SiO2 thickness and all
four Sellmeier parameters. Your MSE should be near 30 (as compared to values as
high as 200 and as low as 40 with the Cauchy). Thus, this model better matches the
true shape of SiO2 index dispersion over a wide spectral range.

![Spectroscopic Data At X=-0.26047, Y=1.47721](image)

*Figure 5-8. Generated data with the Sellmeier model for SiO2 (before fitting).*

**Fit Scan Data**

Now that we have “tuned” the model for a single point from the map, we are ready to
fit data over all points. Make sure to simplify the model by turning OFF:

- the Thickness Pre-Fit and
- the Global Fit

Both options are found in the **FIT Options** section of the Model. After both are
turned off, press the ‘Fit Scan Data’ button from the **Fit** panel.

**NOTE:** If you leave the thickness prefit and global fit ON during analysis of a full
scan, it will take excessively long. However, if there are large variations across the
sample, these may need to be turned back on – using caution and narrowing the
search range of a Global fit.

After the data from all points are analyzed, their results will be shown in the **Fit**
panel, as in Figure 5-9. Now, if you left-click on any of the “blue” parameters in the
**Fit** panel, the graph will update with uniformity map of that parameter. As an
example, the Thickness #1 parameter is graphed, as shown in Figure 5-10.
Figure 5-9. Scan Data results are shown in the Fit: panel. Press any of the “blue” parameters in this list to view the map of that parameter.

Figure 5-10. SiO$_2$ thickness uniformity map.

Model Calculation

Now, let’s return our attention to the fit at a single-point. Go ahead and left-click your mouse on a point toward the center of the graph. Again, the spectroscopic data from that point is graphed and automatically fit with our model. Psi and Delta are matched reasonably well. However, there may be more information about this sample to consider. Take a look at the depolarization curve, which was NOT currently included in the MSE calculation. Press CTRL-Z to see a depolarization graph as shown in Figure 5-11. The increasing depolarization toward short wavelengths (large eV) is very common for thick layers. The cause of this depolarization is due to a non-ideal consideration. The most common non-idealities for thick layers are thickness non-uniformity or instrument bandwidth. Both can be included in our model.
Model Calculation – Thickness Non-Uniformity

Go to the +MODEL Options section of the Model and change the “Model Calculation” from “Ideal” to “Include Thickness Non-uniformity”, as shown in Figure 5-12. This will calculate the effect on the data points if there was a thickness variation within the measurement spot. Unless the system used a focused beam, this is a valid possibility. Hold the SHIFT key down and roll your mouse wheel up/down over the Thickness Non-Uniformity value. When you get to ~2.4%, the model curve will better match the experimental depolarization, as shown in Figure 5-13. This value can be fit – right click on the thickness non-uniformity and fit the data. The MSE should reduce down below 20 (as compared to near 30 without thickness non-uniformity). This is due to improvement in the Psi/Delta fits, as we are not including the Depolarization data as part of the MSE calculation yet.

Figure 5-11. Depolarization curve for the thick SiO2 film.

Figure 5-12. Model Calculation choices.
Model Calculation – Bandwidth (nm)

Next, try changing the “Model Calculation” from “Include Thickness Non-uniformity” to “Include Bandwidth (nm)”. SHIFT-ROLL your mouse until the starting value matches the depolarization data. Then, fit for the Bandwidth. The result should have an even better MSE than with Thickness Non-Uniformity. The depolarization data are matched even better, as shown in Figure 5-14. This is due to the spectral differences between bandwidth and thickness non-uniformity. Thus, this model calculation appears to be the best. This is further verified by the final result that shows Bandwidth near 4nm, which is physically true for the M-2000 used for these measurements.
Include Depolarization Data

Thus far, we included Model Non-idealities to better match the depolarization data, but the depolarization data were not included in the MSE calculation. Thus, we have shown that the non-idealities will also better match the shapes of Psi and Delta (as the MSE was reduced). Now, we include the depolarization data in the MSE calculation. To do this, press the **Configure Options** menu at the bottom of the Model. Under Fit Options section, check the “Include Depolarization Data” choice, as shown in Figure 5-15. Press ‘Ok’. Now, we must turn this feature on in the +FIT Options section. After turning on the “Include Depolarization Data”, press ‘Fit’ again. The MSE should be further reduced, while the bandwidth stays close to where it was without including the depolarization data. As a final step, re-fit all points of the map. Press ‘Fit Scan Data’ and you should get resulting maps similar to those in Figure 5-17, Figure 5-18, and Figure 5-19.

**NOTE:** Analysis time is much longer when Model Non-Idealities are included. You will notice the analysis of all mapped points takes considerable time now that we include the Bandwidth.

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**Figure 5-15.** Press **Configure Options** to select hidden menu “Include Depolarization Data”.

**Figure 5-16.** After adding “Include Depolarization Data” to +FIT Options, it must be turned ON.
Figure 5-17. Final Thickness Map using Sellmeier model and including Bandwidth.

Figure 5-18. Final MSE Map using Sellmeier model and including Bandwidth.
Figure 5-19. Final Index at 633nm (1.960eV) using Sellmeier model including Bandwidth.
5.2. Multi-Sample Analysis

<table>
<thead>
<tr>
<th>FEATURES IN THIS EXAMPLE</th>
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<tbody>
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<td>• Selecting Multiple Points from Map</td>
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<tr>
<td>• CTRL+SPACE BAR to switch views.</td>
</tr>
<tr>
<td>• Set Symbol Size</td>
</tr>
<tr>
<td>• Graph All Data Sets</td>
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<tr>
<td>• Multi-sample Analysis</td>
</tr>
</tbody>
</table>

Once again, we will be working with “mapped” data from multiple locations across a sample. This sample is more complex than the previous example. In addition to a SiO$_2$ coating across the wafer, there are 5 patterned locations on the substrate with an extra layer of Cr over the SiO$_2$ film. The Cr layers were deposited to nominally 10, 15, 20, 25, and 40nm thick. These layers are thin enough to allow light to penetrate through to the underlying SiO$_2$ film. They were part of a study of different ellipsometric methods to measure thin absorbing layers.\(^3\) Figure 5-20 shows the location of each of the Cr regions, along with their nominal thicknesses in nanometers.

![Figure 5-20](image)

Figure 5-20. Overview of sample. The blue region represents a 100nm SiO$_2$ layer. At the five orange points, there is an additional Cr layer on the surface of the SiO$_2$ film, with the nominal thicknesses shown for each (in nm).

Shortcuts for Mapping Data

Before proceeding with this example, it is helpful to review the shortcuts when dealing with mapping data. The table below shows the important features we will use for this example.

Table 5-1. Shortcut reference for mapping data.

<table>
<thead>
<tr>
<th>SHORTCUT</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTRL+Click on Point</td>
<td>Select/De-Select Point</td>
</tr>
<tr>
<td>CTRL+ALT+Click on Point</td>
<td>Delete Point</td>
</tr>
<tr>
<td>CTRL+ALT+SHIFT Click on Point</td>
<td>Show camera image from point (when available)</td>
</tr>
<tr>
<td>CTRL+SPACE Bar</td>
<td>Switch view from single-point to Map</td>
</tr>
</tbody>
</table>

Blank Model

Open the “Cr on SiO2 MAP” data file. You will see the data from translation scan of the sample previously described – as shown in Figure 5-21. You can see “low” (red/yellow) points in Psi at the 5 locations that have been coated with Cr. To fit this data, we need to first build a model. Because we need multiple layers, let’s try building the model from the very beginning. Open the “BLANK” model from the Basic directory. This will appear as shown in Figure 5-22. Left-click on the Substrate = none and choose “Si_Jaw” material file from the Semiconductor directory. Next, press the Add command and locate the next layer above the silicon substrate. For our first layer, choose a Cauchy – which we will use to model the SiO$_2$ layer. Turn on the “thickness”, “A”, “B”, and “C” Cauchy terms on as fit parameters. Next, go to the +FIT Options section and turn on the Thickness Prefit. This model should be adequate to fit any of the areas that do not have a Cr layer.

![Translation Data: Psi](image)

*Figure 5-21. Translation scan graph for “Cr on SiO2 MAP”.*
Before continuing, let’s increase the symbol size to better visualize the locations of the measured points. Right-click on the graph panel. If “Show Symbols” is not checked, do so at this time. From the right-click menu, choose “Set Symbol Size” as shown in Figure 5-23. Increase the symbol size from default value of 1 to 2 or 3. The graph shown in Figure 5-24 has symbol size set to “3”.

![Figure 5-23. From right-click menu on graph, select the “Set Symbol Size” option.](image)
Zooming on Map

While this graph is not too complex, it is possible to collect hundreds or even thousands of points on the same sample. In these cases, it can be helpful to zoom into a certain location to better isolate a specific point. For demonstration purposes, drag your mouse across a region of the sample, as shown in Figure 5-25. The zoomed area is shown in Figure 5-26. To zoom back out to all points, simply right-click in the graph area and choose “Zoom All”.

![Figure 5-25. Drag your mouse over a graph area to zoom in to that region. A rectangle will show you the location of your “zoom” region.](image)

![Figure 5-24. Figure with Symbol Size set to “3”.](image)
Selecting Multiple Points on a Map

With our model ready to fit the SiO$_2$ locations, go ahead and press your mouse on any of the points away from the Cr patterns (blue, purple, and dark green locations). The data from those locations will be opened and fit automatically, as shown in Figure 5-27. Next, go back to the map by checking “Show Map Data”. We are going to select data from multiple points – all of which only contain the SiO$_2$ film – to be graphed and fit at the same time. To select multiple points, hold the CTRL button down when you press your mouse on the different areas. A larger grey circle will be shown around each data point that is selected, as shown in Figure 5-28.

NOTE: The spectroscopic data for this example may use eV for the wavelength – a carry-over from the previous example. I have converted back to wavelength units of nm.
Figure 5-27. Cauchy fit to a single-point with only SiO₂ film.

Figure 5-28. Graph with multiple points selected – as shown by the larger grey circle surrounding those points (4 pts in upper-left section).
Changing the Graph

To toggle the graph from the “map” view of the graph to the individual spectroscopic data from a single point, you can either un-check the “Show Map Data” menu, or simply hold down the CTRL button and press the space bar. This also holds for returning to the “map” view. Try this a few times.

From the Spectroscopic View, we now have multiple points selected, so there is a new section to describe the location of each selected point and a check-box of whether all data should be graphed together or one-at-a-time. This is shown in Figure 5-29. Before, we had an MSE near 5 when fitting a single point. Press ‘Fit’ again and the MSE is near 20. That is because all 4 points are being fit to the same model and it can not match all points with a single answer (i.e. thickness needs to be different for each point).

![Graph All Data Sets](image)

**Figure 5-29.** When multiple data points are selected, you can choose which point to view, or select the check-box to “Graph All Data Sets” simultaneously.

Graph All Data Sets

There is slight variation between the four selected points. This is easier to visualize by graphing all data sets at the same time. Before proceeding, simplify the graph by turning off the double-Y axis and selecting only the 65 degree data (from ‘Set Ranges’ menu, as shown in Figure 5-30. Now, it is okay to check the “Graph All Data Sets” option (above the Graph) and you see Psi curves from all four locations – see Figure 5-31.
Selecting and Unselecting Data Sets

There is a right-click menu over the words “Data Set” above the graph that allow for selecting, unselecting, and clearing data sets. This menu is shown in Figure 5-32. A good test of these menus is to select each individual data set and fit the thickness for each point. To do this, right-click on “Data Set”, and choose “Select Current Data Set ONLY.” This will exclude the other data points from the fit. Press “Fit” and if you only have a single data point selected, your MSE should be near six. Also, you can determine which data set is selected by dropping down the list of data sets. Notice in Figure 5-33 that all points that are not selected are marked as such.

Figure 5-32. Right-click over the “Data Set” text above graph to get extra choices for selecting, unselecting, deleting, and clearing data sets.
Multi-sample Analysis

Thus far, we have fit each of the four selected points individually and as a combined group. However, when we fit them all together, we had a single set of fit parameters for the entire group. This may be okay for the Cauchy parameters – as the film index is the same everywhere – but the thickness is NOT the same at each point. There is an advanced feature in CompleteEASE that allows multiple points to be fit simultaneously with some of the fit parameters common to all points and other fit parameters allowed to vary between points. Although the four points that are coated with SiO$_2$ do not benefit from this powerful technique, the demonstration for these points will help understand the basic concepts before applying to the Cr films – which can benefit from multi-sample analysis.

To use this method, first go to the bottom of the Model and choose Configure Options. Under the “Model Options” section, check “Multi-Sample Analysis” as an Available option. This is shown in Figure 5-34.

Next, go to the +Model Options section and expand the newly available +Multi-Sample Analysis section. Press the Add Fit Parameter text and choose Thickness #1. The thickness for each “selected” data point will show up in the list below (as a fit parameter). Now, right-click on the “Data Set” text and choose to “Select ALL Data Sets”. This will allow us to vary and fit the thicknesses for all 4 locations in the Multi-sample analysis section. The Cauchy parameters will still be fit – the same values will be used to fit all data sets simultaneously. Go ahead and press ‘Fit’. Figure 5-35 shows the final results after the multi-sample analysis. The graph shows the model matching each of the 4 curves – as four different thicknesses are fit along with a common set of Cauchy parameters.
Figure 5-35. Fit results for multi-sample analysis of four locations of SiO$_2$ on Si. A common set of Cauchy values is used, while the thickness of each SiO$_2$ layer is allowed to vary in the +Multi-Sample Analysis section of the Model.

Clear Multi-Data Set Mode

After the multi-sample analysis of four locations with SiO$_2$, the Cauchy values from the fit should be a good final result. We are now ready to turn our attention to the Cr coated areas. To remove the 4 locations with SiO$_2$ only that are currently selected, press the “Data Sets” text above the graph with right-mouse-button and choose “Clear Multi-Data Set Mode”. This returns to a single data set. Press CTRL-SPACE BAR to go back to the map view.

Before we proceed, we want to turn off the fit parameters for the SiO$_2$ film – including the thickness and Cauchy values. We are going to select the Cr coated areas, but we don’t want the model to “automatically” try to fit this data – as we haven’t added a layer for the Cr film. To turn off the “auto” fitting mode, right-click your mouse in the Fit panel and un-select the “Auto Fit” option at top, as shown in Figure 5-36. Now, we can select a point without automatically fitting that point. Go ahead and select the point in direct center of graph (as shown in Figure 5-37).
Figure 5-36. Right-click in the Fit panel and turn off the "Auto Fit" check-box.

Figure 5-37. Select the center point of graph (shown surrounded by grey).

I show this data in Figure 5-38 with double-Y axis turned back on to show both Psi and Delta. I have also generated data to show how different the center-point data are compared to a model for the single-layer SiO$_2$ film.

NOTE: Before fitting, we need to select ALL angles again.
To fit this data, we need to add another layer to our model. Press Add and choose location above SiO₂ film (represented by Cauchy layer) to add B-Spline layer from the Basic folder. Expand the B-Spline layer and press the Starting Mat = none. Choose Cr.mat from the Metal directory. This will calculate the B-Spline points to match Chromium optical constants as published in Palik’s Handbook of Optical Constants. Hold the SHIFT key and roll the mouse wheel over the thickness until it matches the data. Unfortunately, this film does not have the same optical constants as “book” values. This is common for metals, which will differ optically with different deposition methods and processing conditions. Roll the thickness to 40nm, which was the nominal thickness for the Cr layer at this location. The model with 40nm is compared to experimental data in Figure 5-39.

![Spectroscopic Data At Multiple Positions](image)

**Figure 5-38.** Data for center-point compared to model with single-layer SiO₂.

![Spectroscopic Data At Multiple Positions](image)

**Figure 5-39.** Model with 40nm of Cr (starting mat in B-Spline is Palik’s Cr) compared to experimental data at center point.

Because the results are not very close, we need to either global fit the B-Spline parameters or fix the thickness to start. We will choose the latter, as we were given the nominal Cr thickness. With thickness FIXED at 40nm, go ahead and press ‘Fit’. The data fit reasonably well (see Figure 5-40), with an MSE around 4. However, there is a small bump in the model at 3.5eV that don’t match. This is an “interference enhancement” from underlying SiO₂ film. We can only match this region with the correct Cr thickness. Now, turn on the Cr thickness and fit it as well. The final results appear as shown in Figure 5-41 with a good match across the spectrum and MSE near 2.
Figure 5-40. Fit to center point with 40nm fixed B-Spline thickness. The miss-match near 3.5eV is due to interference enhancement from the underlying SiO2 film. The correct Cr thickness will match this region as well.

Figure 5-41. Final fit to center point.
Multi-Sample Analysis (revisited)

Now that we have developed a model to match the center point – with Cr film on SiO$_2$ on Si – go ahead and try this model for all points that are coated with Cr. From the map view, press CTRL while choosing each point with the left mouse button. The points will appear with grey circles surrounding them, as shown in Figure 5-42.

Next, switch to the Spectroscopic View (CTRL+SPACE BAR) and turn off the double-Y axis and Set Ranges to only 65 degrees. This will allow a convenient comparison of the curves, as shown in Figure 5-43. From this data alone, we can estimate the points with thickest-to-thinnest Cr layers based on the shifts in data. The thickest point was at center (in purple). As the data shift, the Cr film gets thinner.

NOTE: The order of points will differ depending on the order they were selected. Thus, your colors may not match the same order in our Figure.

![Figure 5-42. Map view with all 5 locations selected that have Cr coating over SiO2/Si.](image)
Now, let’s prepare the Multi-Sample Analysis. In the +MODEL Options section, you will see that the Thickness #1 parameter is still selected (Figure 5-44(a)). Press the Delete All Parms option to remove Thickness #1, as we will start with a common SiO₂ thickness and only allow the Cr thickness to vary. Next, press the Add Fit Parameter selection and choose Thickness #2 (Figure 5-44(b)).

Before we begin fitting, we need to “tune” the different thickness values closer to their final result. For this purpose, un-check the “Graph All Data Sets”. Now, go down the list of points and SHIFT-Mouse Roll the different thicknesses shown under the Multi-Sample Analysis section until the Generated curve better matches the corresponding Experimental data curves. For more “fine” adjustment of the thickness, hold both the SHIFT-CTRL buttons and roll the mouse button. This decreases the thickness increment. An example is shown in Figure 5-45, where only Data Point #5 is selected and the 5th thickness is being rolled to better match the curves.
After all data points have been matched individually, press the “Graph All Data Sets” menu again. Your graph should look similar to that in Figure 5-46. You hopefully noticed that the curves for thinner layers were not perfectly matched when we rolled the thickness. This is okay, as the thickest Cr layer does not match the optical constants for the thinner layers. In fact, we will later remove this point from the multi-sample analysis. For now, go ahead and press ‘Fit’. All five data points will be fit simultaneously with a single B-Spline layer to describe the common Cr optical constants and individual thicknesses for the Cr layers. The fit result should appear as shown in Figure 5-47. The MSE should be just under 20.
Figure 5-46. Preliminary data match by rolling individual thicknesses to pre-tune our guess to all 5 data sets before fitting multiple sample data.

Now, let’s remove the thickest layer from the multi-sample analysis. To do this, choose the Point location for the center point from the drop-down list above the graph. Next, right-click the mouse over “Data Set:” and choose to “Un-Select Data Set”. The curve for the center point should disappear – leaving only four curves and their corresponding models. Press ‘Fit’ and the MSE should drop from above 19 to less than 14. The center (thickest Cr) point does not share the same optical constants as the thinner layers and thus is best left out of a multi-sample analysis. However, the problem is not confined to just the thickest layer. It turns out that the Cr layer optical constants appear to vary with thickness even for the thinner layers.

Figure 5-47. Fit result for all 5 locations with multi-sample analysis.
Some of this variation is probably due to microstructure, possible roughness at the surface and oxidation. To test this theory, turn on surface roughness in the model and add it as another fit parameter in the Multi-Sample Analysis list. This will allow each point to have different amounts of roughness. Press ‘Fit’ and the MSE drops to near 9. The thickness of the rough surface increases with the thickness of the film (as shown in Figure 5-48). With this model, we could now add the thickest Cr point back into the model to see if this trend (increasing roughness) will better match the center point and be consistent (we expect larger roughness). Sure enough, the MSE for all five points drops below 12 with the results shown in Figure 5-49. At this point, research would continue by testing the samples with other techniques — such as Atomic Force Microscopy — to see if the roughness result is true for this sample.

![Figure 5-48. Fit to four thin Cr films with roughness added to the multi-sample analysis. Notice the resulting roughness increases as the Cr film thickness increases.](image)
- Multi Sample Analysis

<table>
<thead>
<tr>
<th>Data Set</th>
<th>Thickness #2</th>
<th>Roughness</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td>35.59 nm</td>
<td>7.37 nm</td>
</tr>
<tr>
<td>#2</td>
<td>26.01 nm</td>
<td>3.02 nm</td>
</tr>
<tr>
<td>#3</td>
<td>20.30 nm</td>
<td>2.27 nm</td>
</tr>
<tr>
<td>#4</td>
<td>9.84 nm</td>
<td>0.86 nm</td>
</tr>
<tr>
<td>#5</td>
<td>16.16 nm</td>
<td>1.44 nm</td>
</tr>
</tbody>
</table>

Figure 5-49. Fit results for all five points, showing increasing roughness with increasing Cr layer thickness.

As a final exercise press Data: > ‘Set Ranges’ and select all the angles back. Notice when multiple data sets are selected (with more than 1 angle), each data set is graphed with the same color. For example, all curves for Data #1 are in magenta and all curves for Data #2 are in blue. This helps to group them and reduce the total number of different colors graphed. Before fitting again, turn on the SiO₂ thickness as a fit parameter and add Thickness #1 to the list of Multi-Sample parameters. With all the angles selected and three different thicknesses included in the Multi-Sample Analysis, press ‘Fit’ and the results should appear as shown below. You will notice that the center SiO₂ thickness is larger than the remaining 4. However, this was verified on a separate wafer that was not coated with Cr. Thus, the larger SiO₂ thickness at center is to be expected. The final result is shown in Figure 5-50.

Figure 5-50. Final fit to all angles for 5 locations with Cr, SiO₂, and Roughness thicknesses included in the multi-sample fit.
5.3. **Creating OC Library for Composition or Temperature-dependent materials**

<table>
<thead>
<tr>
<th>FEATURES IN THIS EXAMPLE</th>
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<td>• Opt. Const. Compare Model (OC Compare)</td>
</tr>
<tr>
<td>• Draw Graph</td>
</tr>
<tr>
<td>• OC Library Mode</td>
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<tr>
<td>• Build Library</td>
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</tbody>
</table>

In this example, we will demonstrate how to build a composition- or temperature-dependent optical constant library. This procedure can be used to create special material files for any series of samples that show a systematic shift in optical properties related to composition, temperature (or other material property or process condition). During the process, B-Splines are used to smooth the constituent optical constant spectra for each known composition or temperature. The library is then built using an interpolation algorithm based on the critical point shifting algorithm of Snyder et al.\(^4\) In the critical point shifting algorithm, the dielectric function at an arbitrary composition or temperature is evaluated from a weighted sum of the nearest reference dielectric function spectra, after the reference spectra are wavelength-shifted to line up the critical point features. CompleteEASE simplifies this process by automatically determining the wavelength shifting polynomials (that describe the path of critical points versus composition/temperature) via a non-linear regression analysis.

The full procedure for creating a composition or temperature dependent material file is to obtain optical constants from a series of samples with varying, known composition or temperature. This example will not demonstrate this initial step, as the procedures for each individual film will be similar to those taught in earlier examples. After successfully modeling the experimental data to determine accurate optical constants for a specific composition/temperature, save the tabulated optical constants for each material file. In this example, we work with a series of Si\(_{1-x}\)Ge\(_x\) material files that were obtained by fitting a series of layers with different germanium compositions (x). The actual composition was measured by other means – in this case X-ray Diffraction. Thus, we now have the optical constants (n,k) for a series of known Si\(_{1-x}\)Ge\(_x\) compositions.

**NOTE:** Before starting this example, change the Wavelength Units to “eV”, and the Optical Constant Units to “e1 & e2” from the Options Tab.

**Opt. Const. Compare Model**

To start this example, open the “Opt. Const. Compare” model from the Advanced Folder location. The model should appear as shown in Figure 5-51. Press the ADD MATERIAL line seven times to add seven layers to the model. Each of these layers

---

will be used to hold the optical constants for a specific composition of $\text{Si}_1\text{Ge}_x$.

Click on each layer where it says none to add a material file for that layer. There are 10 $\text{Si}_1\text{Ge}_x$ material files in the Examples Folder that were measured from a series of thin films with known composition (as determined from XRD). In addition, we can use crystalline Silicon and Germanium as the end-points for this material file. The compositions related to each material file are provided in Table 5-2.

![Figure 5-51. Opt. Const. Compare model.](image)

### Table 5-2. Compositions for each $\text{Si}_1\text{Ge}_x$ material file used in this example.

<table>
<thead>
<tr>
<th>MATERIAL FILE NAME</th>
<th>GERMANIUM COMPOSITION % (X)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Si}_\text{jaw}.\text{mat}$</td>
<td>0% (x=0.000)</td>
</tr>
<tr>
<td>$\text{SiGe-1}.\text{mat}$</td>
<td>5.2% (x=0.052)</td>
</tr>
<tr>
<td>$\text{SiGe-2}.\text{mat}$</td>
<td>9.8% (x=0.088)</td>
</tr>
<tr>
<td>$\text{SiGe-3}.\text{mat}$</td>
<td>15.5% (x=0.155)</td>
</tr>
<tr>
<td>$\text{SiGe-4}.\text{mat}$</td>
<td>20.4% (x=0.204)</td>
</tr>
<tr>
<td>$\text{SiGe-5}.\text{mat}$</td>
<td>26.3% (x=0.263)</td>
</tr>
<tr>
<td>$\text{SiGe-6}.\text{mat}$</td>
<td>31.9% (x=0.319)</td>
</tr>
<tr>
<td>$\text{SiGe-7}.\text{mat}$</td>
<td>37.1% (x=0.371)</td>
</tr>
<tr>
<td>$\text{SiGe-8}.\text{mat}$</td>
<td>42.3% (x=0.423)</td>
</tr>
<tr>
<td>$\text{SiGe-9}.\text{mat}$</td>
<td>49.3% (x=0.493)</td>
</tr>
<tr>
<td>$\text{SiGe-10}.\text{mat}$</td>
<td>53.9% (x=0.539)</td>
</tr>
<tr>
<td>$\text{Ge}.\text{mat}$</td>
<td>100% (x=1.000)</td>
</tr>
</tbody>
</table>

For this example, populate the seven layers with Silicon, $\text{SiGe-2}$, $\text{SiGe-4}$, $\text{SiGe-6}$, $\text{SiGe-8}$, $\text{SiGe-10}$, and Germanium (all shown in green in table). $\text{Si}_\text{jaw}.\text{mat}$ and $\text{Ge}.\text{mat}$ are found in the Semiconductor folder. The $\text{SiGe}$ files are found in the Examples folder. Before you will be able to set the compositions for each layer, you must turn ON the “OC Library Mode”. Type the corresponding compositions for each layer as shown in Figure 5-52. The temperatures can remain at 25° Celsius for
all files. If we were making a temperature dependent file, we would adjust the corresponding temperatures to match each material file.

ADD MATERIAL  CLEAR
Wvl. Range = 1.632 eV - 4.593 eV
# of Wvls = 300
Graph Type = e1 and e2
Show Difference = OFF
Fit Type = e1 and e2
OC Library Mode = ON  Show Birefr. = OFF

<table>
<thead>
<tr>
<th>Material</th>
<th>x</th>
<th>t</th>
<th>DELETE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ge</td>
<td>1.000</td>
<td>25.00°C</td>
<td></td>
</tr>
<tr>
<td>sige-10</td>
<td>0.539</td>
<td>25.00°C</td>
<td>DELETE</td>
</tr>
<tr>
<td>sige-8</td>
<td>0.423</td>
<td>25.00°C</td>
<td>DELETE</td>
</tr>
<tr>
<td>sige-6</td>
<td>0.319</td>
<td>25.00°C</td>
<td>DELETE</td>
</tr>
<tr>
<td>sige-4</td>
<td>0.204</td>
<td>25.00°C</td>
<td>DELETE</td>
</tr>
<tr>
<td>sige-2</td>
<td>0.088</td>
<td>25.00°C</td>
<td>DELETE</td>
</tr>
<tr>
<td>Si JAW</td>
<td>0.000</td>
<td>25.00°C</td>
<td>DELETE</td>
</tr>
</tbody>
</table>
Ref. Material = none

Build Library  Resolution (eV) = 0.050
Spectra Shifting Parameters

Figure 5-52. OC Compare model setup to build a Library for SiGe films of different composition.

Before proceeding to build the OC Library from these reference files, set the wavelength range to match the material file. For the SiGe material files, the optical constants were measured from 248nm to 1240nm (1eV to 5eV). Also, the “# of Wvls” may need to change to allow the B-Spline to more accurately match the reference files over this wide spectral range. The original material file contains about 540 individual wavelengths. Thus, I have set the “# of Wvls” to 500 – although this will significantly increase the amount of time needed for the calculation. When ready, press Build Library, and a graph will appear as shown in Figure 5-53. In this graph, the colored curves are the reference optical constants at the specified compositions, and the black dashed curves are interpolated spectra at reference and midpoint compositions. Thus, each colored graph should be matched with a corresponding dashed black curve. The additional dashed black curves should demonstrate whether the “shifting” algorithm has worked.
Draw Graph

This graph can be very busy, but you can change the view by selecting a different setting at the bottom of the model under **Draw Graph**. Pressing the **e1 & e2** selection will open the dialog box shown in Figure 5-54. Figure 5-55 and Figure 5-56 show the **e1 only** and **e2 only** spectra for this example, respectively. In addition, the graph can show the Shifted Spectra and Wvl Shift Function to help assess whether the composition library is successful. For advanced users, the Library Resolution and Spectra Shifting Parameters can be modified, as shown in Figure 5-57.

![Figure 5-54. Graphing selections after building an OC Library.](image-url)
Figure 5-55. Graph of $e_1$ only after building the SiGe composition library.

Figure 5-56. Graph of $e_2$ only after building the SiGe composition library.

Figure 5-57. Advanced features to customize the OC Library algorithm, including spectral shifting parameters and resolution in eV.

Build Library  Resolution (eV) = 0.050
Spectra Shifting Parameters
  Wvl Order = 3
  Max. Comp. Order = 3
  Max. Temp. Order = 3
Draw Graph  Graph Type: e2 only
Save OC Library

To further assess whether the library algorithm was successful, you can zoom in on each of the graphs. Figure 5-58 shows a zoomed view of the $E_1$ peak for the SiGe films. This shows that the peaks are shifted to higher energy with increasing silicon composition.
**Figure 5-58.** Zoomed view of the E1 peak for the SiGe compositions, showing a nice shift in critical point toward higher energy for increasing silicon composition.

**Save OC Library**

After successful calculation of a composition or temperature dependent optical constant file, press **Save OC Library** at bottom of the Model panel to save your library as a material file. This material file can now be used to fit future data files. To demonstrate what this file looks like and how it operates, click the “Reference Material = none”, which is the bottom layer in our OC Compare model. Add your newly created material file in this entry. Roll your mouse over the composition in this Reference Material. You will be able to visualize the optical constants shifting compared to your reference constants as the composition is varied. An example graph is shown in Figure 5-59, where the composition has been moved to x=0.75 (black curve) to see how well the shifting works between the two most widely spaced reference values.
Figure 5-59. The newly created Composition dependent file has been added as the Reference material to compare to the original reference material files as the composition is varied.
5.4. Anisotropic Films

In this example, we will allow a layer to have anisotropic optical constants, where the index normal to the sample surface is different than the index in the sample surface plane. This is common with many polymer films that orient along a specific direction.

To start, open the “Aniso-Organic on Si” data file from the Examples folder. The data should appear as shown in Figure 5-60. For simplicity, turn off the Double-Y Axis and view the Psi curves. The oscillations show that this film is transparent over the measured spectral range. Open the “Si with Transparent Film” model and press Fit. The results of this fit should appear as seen in Figure 5-61.

![Variable Angle Spectroscopic Ellipsometric (VASE) Data](image)

Figure 5-60. SE measurement from anisotropic organic film on silicon substrate.
Upon closer inspection, the data at 75 degrees shows a common “signature” of anisotropy in the film. The interference oscillations from anisotropic films will often “tilt” such that the curves are higher on one side of the peak and lower on the opposite side. This is demonstrated for anisotropic films on silicon at 75 degrees in Figure 5-62. The red curve represents an isotropic film. The green curves are higher on the long wavelength side of a peak and lower on the short wavelength side. These represent anisotropic films with \( N_z > N_{xy} \). The blue curves are lower on the long wavelength side of a peak and higher on the short wavelength side. These represent anisotropic films with \( N_{xy} > N_z \).
Convert to Anisotropic

Right-click over Cauchy Film to access the menu of options to modify the Cauchy layer. From this list, choose to “Convert to Anisotropic”. The Cauchy layer will now become embedded as part of a new layer named “Biaxial”. Biaxial stands for anisotropy where Nx, Ny, and Nz are all different. However, the layer name is used here to describe any anistropic film. Within the Biaxial layer, you can choose either Uniaxial or Biaxial anisotropy. For our case, we will use Uniaxial anisotropy, as it would require more advanced measurements to study the potential Biaxial anisotropy of a sample. This will be described in more detail in a later Example.

Biaxial Layer

The biaxial layer can be used in two different ways and both will work equally well for this example. The choice is whether to have Difference Mode = OFF, as shown in Figure 5-64 or Difference Mode = ON, as shown in Figure 5-65.
With the Difference Mode = OFF, each direction is described by an individual layer. In our case, each direction is seeded with the Cauchy layer, as that is the layer we started with. The layers could just as easily be B-Spline, Gen-Osc, or a mixture of these.

With the Difference Mode = ON, the x-direction is defined by a material layer, while all other layers are determined by summing the x-direction optical constants with the “difference” values calculated from the extended Cauchy type dispersion described as the dZ (z-difference). This method can be extremely useful when the fit is primarily sensitive to the index difference and not sensitive to the absolute value of the index.

For this example, turn the Difference Mode to ON and then hold shift key down while “rolling” the mouse wheel over the dZ_A parameter. If you roll this parameter to positive numbers, the simulated curve will look worse than the isotropic fit (tilt in wrong direction). However, if you roll the dZ_A parameter to negative values, it will start to match better. Go ahead and turn on dZ_A and dZ_B as fit parameters. Make sure you are still fitting the Cauchy parameters and the thickness, but it is wise to turn off the Global Fit and Thickness Prefit before fitting, as we are close to the final answer. Press Fit and the final result should appear as shown in Figure 5-66.

---

**Figure 5-64.** Biaxial layer with Uniaxial anisotropy and Difference Mode OFF.

**Figure 5-65.** Biaxial layer with Uniaxial anisotropy and Difference Mode ON.
Figure 5-66. Anisotropic fit using the Difference Mode = ON.

Now, right-click on the Biaxial layer and you will have two choices for viewing the optical constants, as shown in Figure 5-67. You can choose Graph Layer Optical Constants to view ordinary (Nx = Ny) and extra-ordinary (Nz) optical constants simultaneously in the Graph. Alternately, if you prefer to graph the index difference between the various directions, you can right-click on the biaxial layer and choose “Graph Layer Optical Constants (anisotropic differences)”, as shown in Figure 5-67. This option will show the difference between ordinary (Nx = Ny) and extra-ordinary (Nz). The graph will display “o-e” as shown in Figure 5-69.

If you prefer to see only the optical constants of a single direction, you can right-click on each individual material file describing that direction, as shown in Figure 5-68. In this example, you can view the ordinary optical constants (Nx = Ny, “Cauchy Film”). For access to all directions, you must fit with Difference Mode Off.
Figure 5-67. Right-click on Biaxial layer to graph the layer optical constants (as shown here) or to graph the “Anisotropic Differences”.

Figure 5-68. When Difference Mode is Off, you can right-click on the various material files describing the directions in the anisotropic “biaxial” layer to view them separately.
Figure 5-69. "Anisotropic difference" graph for this example.
6. Data Analysis 4 – *In Situ* Data

This chapter includes further examples pertaining to *in situ* data analysis. The most common data analysis for dynamic measurements of growth or etch use the “Multi-Time Slice” analysis discussed in the first example. However, we also demonstrate the use of a Growth Rate and Optical Constants model (GROC) that can be very easy to apply when the film matches the requirements (absorbing films with constant growth rate and optical constants).

The section examples from this chapter are listed below, along with the primary CompleteEASE features that are discussed.

**Section 6.1 In Situ Data Analysis**

<table>
<thead>
<tr>
<th>FEATURES IN THIS EXAMPLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Viewing individual time-slices</td>
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<tr>
<td>• Selecting multiple-time slices from dynamic scan</td>
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</table>

**Section 6.2 Growth Rate and Optical Constants**

<table>
<thead>
<tr>
<th>FEATURES IN THIS EXAMPLE</th>
</tr>
</thead>
<tbody>
<tr>
<td>• GROC Model</td>
</tr>
</tbody>
</table>
6.1. In Situ Data Analysis

**FEATURES IN THIS EXAMPLE**

- Viewing individual time-slices
- Selecting multiple-time slices from dynamic scan
- ‘Dynamic Fit’
- Multiple-Time Slice Analysis

In Situ data provides ellipsometry measurements that were taken versus time. This often involves data as the thin films are being processed. The model may require a description of changes in film thickness, optical constants, composition, or even temperature versus time. The navigation, manipulation, and analysis of in situ data have many parallels with the corresponding processes for uniformity maps and multi-sample analysis. Thus, it can be helpful to review these examples before proceeding with this example.

This example is split into two sections. First, we will discuss navigating and selecting data sets from in situ time-dependent data set. These topics can be considered by any user-level. Second, we will work to analyze the in situ data via a multi-sample analysis approach and finally a time-dependent data analysis procedure. As this data comes from an amorphous silicon film and will require analysis methods including the B-Spline and Genosc layers, those examples should be reviewed in detail before proceeding with this example.

**Shortcuts for Dynamic Data**

Before proceeding with this example, it is helpful to review the shortcuts when dealing with in situ dynamic data. The table below shows the important features we will use for this example.

*Table 6-1. Shortcut reference for dynamic data.*

<table>
<thead>
<tr>
<th>SHORTCUT</th>
<th>FUNCTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Click on Dynamic Graph</td>
<td>Opens Spectroscopic Data from that time point.</td>
</tr>
<tr>
<td>CTRL+Click Dynamic Graph</td>
<td>Selects Spectroscopic Data from that time point.</td>
</tr>
<tr>
<td></td>
<td>(Repeat for multiple time selections)</td>
</tr>
<tr>
<td>CTRL+SPACE Bar</td>
<td>Switch view from dynamic to spectroscopic data.</td>
</tr>
</tbody>
</table>
Navigating In Situ Data

Open “a-Si Grow Sim.iSE” data file from Examples folder. The time dependent data will be shown in the graph in Figure 6-1. Notice a selection of 5 wavelengths is graphed. These curves are not the only wavelengths measured, but represent the shape versus time for the spectral range selected.

![Figure 6-1. Graph shows in situ data versus time.](image)

Viewing Spectroscopic Data at a Specific Time

Left-click your mouse to select data for a specific time-slice. A line is drawn at that time slice, as show in Figure 6-2. Now, you can either press CTRL+SPACE BAR to toggle to the spectroscopic graph for that time-slice or “un-check” the box named Show Dynamic Data located just above the graph. Data for a time-slice selected near 3 minutes is shown in Figure 6-3.

![Figure 6-2. Left-click mouse on graph near 3 minutes selects this time and shows the selection as a vertical gray bar.](image)
Figure 6-3. To view the spectroscopic data at the selected time-slice, press “CTRL-SPACE BAR” or un-check the Show Dynamic Data button just above the graph pane.

**Turning off the “Auto-Fit” function**

Before we proceed, you will want to turn off the “Auto-Fit” function so the data are not automatically fit every-time we select a time slice or open a new model. To do this, right-click in the Fit pane region and un-check the Auto Fit option, as shown in Figure 6-4.

Figure 6-4. Right-click in the Fit pane region and turn off the “Auto-Fit” check-box.
Building a Model

Now, let’s build a model to fit this time-slice. You should be able to do this step with any time-slice between 2 and 3 minutes. We will start with the assumption that we measure a coating that has known optical constants. This will allow us to find the thickness at any point in time. For our model, open blank.mod. Start with “BK7 Glass” substrate from the Dielectrics subdirectory and then add “a-si parameterized” from the Semiconductor directory. Start by fitting the a-si parameterized layer thickness only. You will find the fit is not adequate, as shown in Figure 6-5.

![Figure 6-5](image1)

**Figure 6-5.** Fit result for data near 3 minutes when only fitting the a-Si layer thickness (no surface layer).

To improve the fit, we must consider that the surface of any silicon layer will be oxidized and possibly rough. To fit the surface, we can either add an SiO₂ layer or simply turn on the surface roughness thickness as a fit parameter. The improved fit result is shown in Figure 6-6.

![Figure 6-6](image2)

**Figure 6-6.** Fit result for time near 3 minutes when including the surface roughness.
‘Fit Dynamic’

Now, let’s fit these two parameters versus time to see how they change as the data vary across the measured time span. Simply click the Fit Dynamic button in the Fit pane and the software will progress from the earliest time slice and fit the data, continuing toward larger times, as shown in Figure 6-7.

You may notice problems starting a dynamic fit with model that is tuned for a thicker region of the data. If the fit fails, it may appear as shown in Figure 6-8. Here, the fit finally matches the data at around 3 minutes. To solve this problem, simply “tune” the fit for an earlier time before pressing Dynamic Fit. Go ahead and select a time slice near 0.3 minutes. Roll the thickness values for the roughness and a-Si layer until they are closer to correct for this new time. Press Fit for the single-point and you should get a result like shown in Figure 6-9. Now, press “Dynamic Fit” again and this time it should be successful. Press “Show Dynamic Data” to ensure the model curves match the Experimental curves over all times, as shown in Figure 6-10.

NOTE: It can be beneficial for many applications to start the fit from end-point rather than starting time, as the film thickness is often larger at the end of a data run. To fit the data in reverse-time order, right-click the ‘Fit Dynamic’ button and select “Fit Backwards in Time”.

Figure 6-7. Data during Dynamic Fit of all measurement times.
Figure 6-8. Fit fails until about 3 minutes because the starting thickness was not close for early time region.

Figure 6-9. Fit for a-Si and rough layer thickness at time slice near 0.3 minutes.
You will also notice in Figure 6-10 that the Fit Panel shows the final results (from last time slice). If you select the MSE, Roughness, or Thickness #1 with your mouse, the time-dependence of these values will be graphed, as demonstrated in Figure 6-11 and Figure 6-12.
Selecting Multiple Time-Slices

For many materials, the optical constants will not be known and in situ data can be very effective to determine thickness and optical constants. The main advantage of in situ data is that it provides information about the sample at different stages of growth. If multiple time-slices are fit simultaneously, we get the same advantages as with multi-sample analysis (see section). To setup a multi-time slice analysis, view the Dynamic Scan Data and then hold the CTRL key down when selecting the multiple positions with your left-mouse button. An example is shown in Figure 6-13. These can now be viewed simultaneously in a similar manner as shown for mapping data. Un-check the “Show Dynamic Data” box and then you can choose which time slice you would like to view from a drop-down box. You can also view all data sets simultaneously by checking the “Graph All Data Sets” box, as shown in Figure 6-14 (here I have also turned off the double-Y axis on graph for simplicity).

NOTE: To quickly select multiple consecutive time-slices, hold the CTRL key down and drag your mouse over the graph area you wish to select.
Manipulating Multi-Data Sets

Just like when manipulating multiple data sets from a “map”, there are additional options by right-clicking over the words “Data Set” just above the graph, as shown in Figure 6-15. For example, this is where we would go to “Clear Multi-Data Set Mode”.

Figure 6-14. Graphing spectroscopic data from multiple time-slices.

Figure 6-15. Right-click on the words “Data Set” above a graph to access additional options related to the multiple time-slice data sets.
Multi-Time Slice Analysis

Now, let’s setup the multiple time-slice analysis. From the Model pane, press the Configure Options menu at the bottom and then select “Multi Sample Analysis” from the choices of Available Model Options, as shown in Figure 6-16. This is a hidden feature as it is reserved for advanced users. Now, when we expand the Model Options section, there is a Multiple Sample Analysis section that can be further expanded. Click to “Add Fit Parameter” within this section and you will be able to define any of the fit parameters as variable for different time-slices. Let’s do this for Thickness #1, as shown in Figure 6-17. It will seed the current thickness for all time-slices. This will not be correct and it is important to individually adjust the thickness “starting-points” for each time slice. This can be done by selecting each time slice individually in the graph and then rolling the thickness of the corresponding time-slice from within the Multi-Sample Analysis section, as I have done in Figure 6-18.

Figure 6-16. Choose “Multi Sample Analysis” from the Available Model Options within the Configure Options selection.

Figure 6-17. Model Options section is expanded to view the newly added Multi Sample Analysis section. Press “Add Fit Parameter” to designate a fit parameter to vary with each time-slice.
Figure 6-18. Starting points for all time-slices have been adjusted by rolling the mouse over each individual thickness from within the multi-sample analysis section of the Model.

You can now press ‘Fit’ and all thicknesses (for each time slice) along with a single roughness value that is used to fit all data sets is varied. This result is shown in Figure 6-19. If the roughness was also changing with time, its thickness could be added as a time-dependent variable within the multi-sample analysis section.

Figure 6-19. Fit results when a-Si thicknesses were allowed to vary at each of the five time-slices, along with a single roughness for all time slices.
Fitting Optical Constants

The real benefit of multi-time slice analysis is that it allows better results when fitting optical constants for a layer. Let’s pretend we don’t know the optical constants for this amorphous silicon layer. Replace the coating in your model with a B-Spline layer. To get the B-Spline optical constants in the general vicinity of the final answer, we will click on the “Starting Mat =” and choose a-Si from the Semiconductor directory. This will seed the node values of the B-Spline to match the reference a-Si optical constants, as shown in Figure 6-20.

You will notice that the starting point is not the correct final optical constants for the time-dependent silicon simply by Generating data, as shown in Figure 6-21. However, we have five time slices to provide adequate information about this film at different thicknesses. Go ahead and press ‘Fit’. The node values will adjust in the B-Spline to best fit the amorphous silicon data from ALL time-slices simultaneously. This fit result is shown in Figure 6-22, with the B-Spline optical constants shown in Figure 6-23.
Figure 6-21. Generate Data and you see that the book-value a-Si optical constants used as a starting point in our B-Spline do not match the experimental data. However, they are close enough to give us a good starting point for fitting.

Figure 6-22. Fit results with B-Spline Optical constants allowed to vary.

Figure 6-23. Final optical constants from B-Spline fit.
6.2. Growth Rate and Optical Constants

In this example, we will consider a multi-layer of tantalum metal and amorphous silicon thin films, where the process conditions are varied for each layer. During deposition of each layer, the growth rate is constant and the optical constants should not vary. Our goal is to determine the process condition from these six different films.

Because of the ideal sample conditions, we are able to apply a simple model (called GROC) to the dynamic data analysis. This model can only be applied to cases where i) the growth rate is constant, ii) the optical constants do not vary during growth, and iii) a virtual interface approximation can be applied for the underlying structure. The latter requirement means the GROC is typically used for semiconductor and metal growth, and not likely to work with dielectric layers.

Virtual Interface

A virtual interface can simplify dynamic data analysis by allowing us to ignore the underlying sample structure and only consider the current thin film growth. The virtual interface is placed near the surface of the film that is being studied. For example, if we are determining the optical constants of Layer #3 in the stack of Figure 6-24, we would place the virtual interface inside Layer #3 and ignore the sample structure below this point.

Figure 6-24. A virtual interface allows the underlying sample structure to be approximated by a single interface. The optical constants at this interface are called the "pseudo-substrate".

There are different ways to calculate the virtual interface, but we will limit our discussions to the Common Pseudo-Substrate Approximation (CPA). This assumption replaces everything below the virtual interface by an effective, semi-infinite substrate that has optical constants equal to the pseudodielectric values for the stack below that level. This approximation works amazingly well for semiconductors and other high-index materials, but does not work for low index transparent materials, like glass.
Growth Rate and Optical Constants (GROC) Model

Go ahead and open the “Growth Rate and Optical Constants (GROC)” model from the Model panel, under the Advanced Folder. The GROC model is shown in Figure 6-25. As you can see, it does not contain a “layered structure” like a standard model we are accustomed to using.

<table>
<thead>
<tr>
<th>Rate = 3.00 Å/s (fit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fit Opt. Const. = ON</td>
</tr>
<tr>
<td>Layer = none</td>
</tr>
<tr>
<td>Angle Offset = 0.000</td>
</tr>
<tr>
<td>POI Offset = 0.000</td>
</tr>
<tr>
<td>Roughness = 0.00 nm</td>
</tr>
<tr>
<td>VI Mode = CPA (semiconductors)</td>
</tr>
<tr>
<td>Max. Times = 50</td>
</tr>
<tr>
<td>Spectral Resolution in eV = 0.050</td>
</tr>
<tr>
<td>Seed thickness = 10.00 nm</td>
</tr>
<tr>
<td>Surface Tracking Fit Mode = OFF</td>
</tr>
<tr>
<td>Limit Wvl. for Fit = OFF</td>
</tr>
<tr>
<td>Start Time for Data = 0.000 min.</td>
</tr>
</tbody>
</table>

Figure 6-25. Example of the GROC model.

The Growth Rate and Optical Constant (GROC) layer is designed to simplify the simultaneous extraction of growth rate and optical constants from dynamic in situ SE data. GROC utilizes a ‘virtual interface’, which means that the user does not have to specify an optical model for the underlying layers/materials. GROC also assumes a constant growth rate, constant surface roughness, and that the optical constants do not vary with film thickness over the specified time.

NOTE: The GROC assumptions are not valid during the nucleation phase of growth.

The Common Pseudo Substrate Approximation (CPA) used within the GROC is not generally valid for all materials, but does work quite well for growth of semiconductors on semiconductors, or the deposition of absorbing materials (such as metals, after about 5 nm of growth).

NOTE: The GROC is used for metals and semiconductors, such that the virtual interface assumption is valid.

**Rate**

Rate specifies the constant growth rate of the layer. For the fit algorithm to converge, the rate should be near actual value before starting the fit.
**Fit Opt. Const.**

When ‘ON’

- layer optical constants are fit using a piecewise-continuous polynomial representation for the optical constants. The spacing of the polynomial “control points” are specified by the “Spectral Resolution in eV” setting. ‘Draw Opt. Const.’ will graph the extracted optical constants, while the ‘Save Opt. Const.’ button will save the extracted optical constants in a tabulated material file.

- Starting values for the polynomial match the “Layer” entry, if present. This is helpful to “seed” reasonable starting optical constants.

When ‘OFF’

- the optical constants are specified by the ‘Layer’ entry. If the Layer entry is empty or no fit parameters are turned “on” within the layer, then the optical constants are fixed and only growth rate is varied. If parameters are turned on within the “Layer” entry, then the optical constants are fit using this layer and its dispersion equation fit results.

**NOTE:** Optical Constants can be “fit” whether Fit Opt. Const. is “on” or “off”.

**Layer**

The layer is used to “seed” optical constants when “Fit Opt. Const. = ON” and this material file specifies the optical constants of the layer when “Fit Opt. Const. = OFF”. Any fit parameters defined in the layer will be included in the GROC analysis of the data, when “Fit Opt. Const. = OFF”.

**Angle Offset**

**NOTE:** Feature only applied when ‘Fit Opt. Const. = OFF’ to avoid 100% correlation.

Specifies an offset to the nominal angle of incidence specified in the experimental data file. Fitting the angle offset is useful when “known” reference optical constants are specified in the “Layer” field.

**Roughness**

Specifies and/or fits the surface roughness of the growing film, which is assumed to be constant during the selected time range. It is not always possible to uniquely extract the optical constants, growth rate, and surface roughness: an adequate thickness of grown material (> > ¼ wave) must be encompassed by the selected time range, otherwise, the roughness should be fixed at 0.

**VI Mode**

Specifies how GROC implements the Virtual Interface. The “CPA” and “CPA – Fit VI” modes assume the common pseudo-substrate approximation, which is typically valid for the growth of semiconductors on semiconductors or absorbing materials such as metals.

- “CPA”: the ellipsometric \( \Psi \) and \( \Delta \) parameters at the first selected time are simply inverted into pseudo optical constants, which are then used to calculate the ellipsometric trajectory for the subsequent film growth.
• “CPA-Fit VI”: defines the VI pseudo optical constants as fit parameters, which minimizes the impact of random or systematic errors in the $\Psi/\Delta$ data used for the VI inversion.

**Max. Times**

To minimize analysis time, GROC limits the time slices during analysis to the number specified– evenly distributed across the selected time range.

**Seed Thickness**

To improve the robustness of the fit, GROC performs an initial fit with an initial subset of the selected time range before fitting all of the data. The ‘Seed Thickness’ specifies how much time should be included in this initial fit (the initial growth rate is used to calculate the time, given the ‘Seed Thickness’). For most fits, the default value is acceptable, but in some cases, smaller or larger (more likely) values could be tried if the fit does not converge properly.

**Surface Tracking Fit Mode**

When OFF (which is the default), the dynamic data in the selected time range is used in the GROC analysis. If ON, the dynamic data immediately preceding and including (the # of points is specified by the “Max. Times” setting) the selected timeslice is used in the analysis. Since the selected timeslice is automatically moved through the data during a “Fit | Dynamic Fit”, this mode enables a GROC analysis vs. time, which could be useful for extracting time dependent growth rate and/or surface roughness changes.

**Multilayer Example**

Open the “Dynamic Ta-Si Multilayer.iSE” data file from the Examples folder. This file is a dynamic, in situ measurement, so it will have “iSE” extension. There is a drop-down box in the “Open Data” dialog box, as shown in Figure 6-26.

![Figure 6-26](image)

Figure 6-26. The “Dynamic Ta-Si Multilayer” data file is an in-situ measurement, which has “iSE” extension. You can choose which files to view in the Data: > ‘Open’ dialog box.
The data graph should appear as shown in Figure 6-27, for Psi versus time. This data was collected during the growth of six consecutive films, as described in Table 6-1. Before the first minute, the substrate was monitored without film growth. This was followed by six films alternating between Ta metal and amorphous silicon. For each layer, the sputter gun current was varied to change the growth rate. Our goal is to determine the growth rate and optical constants for all six layers, to characterize the process conditions for this sputter chamber. To this end, we will be using the GROC layer.

![Dynamic Data](image)

*Figure 6-27. Dynamic Ta-Si Multilayer dynamic data.*

<table>
<thead>
<tr>
<th>Layer</th>
<th>Time Start</th>
<th>Time End</th>
<th>Film</th>
<th>Gun Current</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>6</td>
<td>Ta</td>
<td>0.2</td>
</tr>
<tr>
<td>2</td>
<td>6</td>
<td>11</td>
<td>a-Si</td>
<td>0.3</td>
</tr>
<tr>
<td>3</td>
<td>11</td>
<td>14</td>
<td>Ta</td>
<td>0.4</td>
</tr>
<tr>
<td>4</td>
<td>14</td>
<td>17</td>
<td>a-Si</td>
<td>0.5</td>
</tr>
<tr>
<td>5</td>
<td>17</td>
<td>24</td>
<td>Ta</td>
<td>0.1</td>
</tr>
<tr>
<td>6</td>
<td>24</td>
<td>31</td>
<td>a-Si</td>
<td>0.15</td>
</tr>
</tbody>
</table>

Open the “Growth Rate and Opt. Const.” model from the Advanced folder link. Next, we need to select the time range to be included in the analysis. Press the **Data: ‘Set Ranges’** button to choose the data from the first tantalum film, from 1 to 6 minutes, as shown in Figure 6-28. Before we press fit, we can improve our success by “seeding” optical constants into the polynomial by adding a “Layer”. Select the “Layer” and add “Ta.mat” from the metals directory. Now, press ‘Fit’ and you should get the results shown in Figure 6-29.
Figure 6-28. Press the Data: ‘Set Ranges’ button to choose the time range for the first tantalum film (1 to 6 minutes).

Figure 6-29. Results of GROC analysis over time range from 1-6 minutes for Ta film.
The fit is not perfect, but remember that the GROC analysis requires the optical constants and growth rate to be constant over the full time-range. The layer deposition may not have started at exactly 1 minute and the first few data points of growth may show nucleation effects where the metal optical constants are not consistent with the thicker layer. To improve the fit, range-select the data from 2-6 minutes, to ignore the early stages of film growth. Press ‘Fit’ again and the MSE should drop to near 1.5, as shown in Figure 6-30. Record the growth rate from this fit and copy the optical constants to the Graph Scratchpad.

Figure 6-30. GROC analysis for tantalum film growth from 2-6 minutes, ignoring the early stages of film growth where the film optical constants may vary during nucleation.

If the graph disappears, as shown in Figure 6-31, press the ‘Show Data’ button in the top-right of the Graph area.

Figure 6-31. If the dynamic data graph disappears, press the ‘Show Data’ button at the top-right of the Graph panel.
Now, let’s repeat this procedure for the other two regions of Tantalum growth (11-14 minutes, 17-24 minutes). Remember to avoid the nucleation region at the beginning of growth. When you start a new region, the GROC will calculate the new CPA, so the data at the first time will match. However, our growth rate may be too large or small, as it remains from the previous fit. For example, Figure 6-32 shows the initial generated curve for the third tantalum film, but the growth rate in the GROC is from the fastest film (#2), so the generated data do not match. Roll the growth rate to a better value to “seed” the GROC layer closer to the correct answer before pressing ‘Fit’.

Results for the second tantalum film (Figure 6-33) and third tantalum film (Figure 6-34) are shown, along with the final optical constants as copied to the Graph Scratchpad (Figure 6-35 and Figure 6-36).

Figure 6-32. Generate data for each new range. This will allow you to estimate whether the previous Growth Rate is too fast or slow for the new film.

Figure 6-33. GROC analysis for tantalum layer growth between 12 and 14 minutes.
Figure 6-34. GROC fit for the tantalum film growth between 18 and 24 minutes.

Figure 6-35. Graph Scratchpad comparing index of all three tantalum films.
Figure 6-36. Comparison of the extinction coefficient for all three tantalum films.

Feel free to continue analysis for a-Si films, and the final growth rates for both tantalum and silicon films should appear as shown in Figure 6-37.

Figure 6-37. Growth Rate for Ta and a-Si films versus sputter gun current.
7. Reference – Graph

In this chapter, the features and capabilities of CompleteEASE will be described, using a “reference” format. That is, the features will be listed in the order they appear on screen, but not necessarily in the order one would use to perform typical procedures. The previous chapters on Basic System Operation and Data Analysis should be studied for “procedural” documentation.

7.1. Basic Screen Layout

The basic layout of the CompleteEASE software consists of Tabbed Panes across the top of the window, with a Graph always visible in the bottom section (Figure 7-1). The window can be resized, maximized, or minimized using standard Windows keystrokes or mouse operations. The divider bar between the Tab Panes and Graph can be moved to adjust the relative size of the graph. The CompleteEASE screen layout is saved when the program exits, and restored upon restart.

![CompleteEASE software](image)

Figure 7-1. CompleteEASE software with the Measurement tab selected.
7.2. Graph Panel

The Graph panel is an important part of the CompleteEASE user interface. Spectroscopic data, model fits, and optical constants are all plotted in the Graph panel. The features described in this section can be used regardless of the type of information displayed in the graph window.

If the mouse is stopped in the plot area for a few seconds, a “tool tip” box will appear which reports the X and Y values corresponding to the mouse position. The mouse can be moved to other positions and the X-Y values will be updated.

The legend can be repositioned by clicking (\text{\textbullet}) in the legend area and dragging the legend box to the desired location.

‘Show Data’

If optical constants (or something else besides data) are being graphed, you can press this button to go back to the data graph.

Range-Select

To “zoom in” on a specific spectral range, hold down (\text{\textbullet}) and drag over the desired spectral range (Figure 7-2). The graph will then redraw, plotting only the selected spectral range.

\textit{Figure 7-2. To select a specific spectral range (hold down the left mouse button and drag the mouse over the desired range).}

The same is true for uniformity maps, as shown in Figure 7-3. Dragging the mouse over an area (left) will select this region to graph (right).

\textit{Figure 7-3. Drag mouse over area of map (left) to select that area of the map (right).}
**Zoom All**

To plot the entire spectral range (or entire uniformity map area), simply press the ‘Zoom All’ button at top-right of the Graph pane (Figure 7-4). This same option can be found at the top of the Right-Click menu from within the graph.

NOTE: ‘Zoom All’ does not appear if you change the graph to “Use Clipboard Aspect Ratio”.

![Figure 7-4. Right-Mouse menu from within the Graph.](image)

**7.3. Right-Click Menu (Graph)**

Press (^R) in the graph area to access the menu shown in Figure 7-5.

![Figure 7-5. Right-click (^R) menu from Graph panel for a spectroscopic scan and a map.](image)

**Zoom All**

Press to graph all data, whether spectroscopic scan or uniformity map.
Statistics

The graphed data undergo a variety of statistical calculations. For example, a straight-through measurement of air can be used to verify the measurement accuracy. The statistics of this measurement can then be calculated, as in Figure 7-6.

![Graph Statistics](image)

*Figure 7-6. Graph Statistics, taken from a straight-through measurement.*

Copy Graph to Clipboard

Graph is copied to the clipboard for use in other programs.

Copy Data to Clipboard

Graphed data curves are copied to the clipboard for use in a spreadsheet. Information will include a title line that matches the graph title; followed by columns of information with a heading for each column. The first column is the x-axis information – typically wavelength, followed by a column of information for each curve on the graph.

Copy Data to Graph Scratchpad

Places the data curves from graph into the CompleteEASE Scratchpad where they can be viewed and compared to other curves, along with customization of the curves being viewed.

Profile Along X and/or Y Axis

For uniformity maps, the 2D curve is not copied to the Graph Scratchpad. Instead, you are given a choice to copy the data points from along the X axis, Y axis, or both, as shown in Figure 7-7. An example graph from x-axis of a Uniformity map is shown in Figure 7-9.

![Copy to Scratchpad](image)

*Figure 7-7. When working with Uniformity Maps, you get the above options when you select “Copy Data to Graph Scratchpad.”*
View Graph Scratchpad

The Graph Scratchpad is a convenient place internal to CompleteEASE to compare curves. The curves can be any type of data that is graphed, including raw data, model-generated curves, optical constants, or even the MSE profile from a Parameter Uniqueness test. An example of the Graph Scratchpad is shown in Figure 7-8.

![Graph Scratchpad Example](image)

**Figure 7-8.** The Graph Scratchpad can be used to compare all types of graph curves, and customize the view for easier comparison.

![Graph Scratchpad Results](image)

**Figure 7-9.** Graph Scratchpad results from the X-axis points of a uniformity map.

Show Legend

Turn On/Off the graph legend.

Show Symbols

Turn On/Off graph symbols for the Experimental Data. The model curves are still represented as solid lines. Figure 7-10 shows an example data graph with symbols on.
Set Symbol Size

This menu option is available when viewing map data. Change the size of symbols representing map data points. The default size is 1, but this can be increased or decreased depending on the view required.

Auto X,Y,Z Scale

X,Y,Z-Axis is automatically selected to show all data.

Manual X,Y,Z-Scale

User can select lower and upper X,Y,Z-axis limits. All data are still included when fitting, but only points within this range are graphed.

Scale Z-Range by n-Sigma

User can quickly select the lower and upper Z-axis limits for mapped data based on a multiple of the standard deviation. This can be useful when there are a few “bad” points on an otherwise uniform sample. When selected, a dialog box appears requesting the number of standard deviations to use when calculating the range, as shown in Figure 7-11. All data are still included when fitting, but only points within this range are graphed.

Graph Points as a Line

Allows user to show Map data unwrapped in a line graph. For example, a map of SiO<sub>2</sub> thickness over a wafer is graphed as a Line in Figure 7-12.
Use Clipboard Aspect Ratio

The standard graph will scale with the size of the graph pane. However, when the graph is copied to the clipboard, it always uses the configured aspect ratio. Choose this button to view the graph as it will appear when copied to the clipboard. The clipboard aspect-ratio can be modified under the “Graph Clipboard Parms” section of Options > Configuration Controls: > ‘Edit Configuration’.

Cancel Menu

Close the Graph menu.

7.4. Graph Type

The "Graph Type" drop-down menu at the top-left of the Graph pane lists the different graph types, as shown in Figure 7-13.

Figure 7-12. Map data graphed as a Line.

Figure 7-13. Graph Type Menu choices for standard ellipsometry data.
There are additional options for this menu if the data type is either a Generalized Ellipsometry measurement or Mueller-Matrix measurement. In either case, the "Graph Type" menu will appear as in Figure 7-14.

![Figure 7-14. Graph Type drop-down menu for Anisotropic and Mueller-Matrix data.](image)

**Psi, Delta**

When Psi or Delta are selected, the graphs plot the standard ellipsometry parameters for amplitude ratio, Psi ($\Psi$) and phase difference, Delta ($\Delta$). These are traditional representations of an ellipsometry measurement, based on equation 6-1.

$$\rho = \tan(\Psi)e^{i\Delta} = \frac{R_p}{R_s}$$  \hspace{1cm} (6-1)

**Re(rho), Im(rho)**

Real and Imaginary rho graphs refer to an alternate view of the ellipsometry measurement, based on rho ($\rho$), as given in equation 6-1.

**N, C, S**

Another alternate form of viewing ellipsometry measurements is based on the Mueller-matrix formalism. For a standard, isotropic sample with no depolarization, the Mueller-matrix response would appear as shown in Equation 6-2.
Thus, NCS are the non-zero elements of the isotropic Mueller-Matrix and are related to the Psi-Delta curves traditionally graphed for ellipsometry. A few advantages to the NCS notation include i) they are always defined and bounded between -1 and 1, and ii) they are linearly related to the intensity harmonics measured by any ellipsometer system. Thus, they are closer to the direct measureables of the ellipsometer and there is approximately equal precision in N, C, and S.

**<Pseudo> Transforms (<e1>, <e2>, <n>, <k>)**

The <Pseudo> Transforms menu expands to include <e1>, <e2>, <n>, and <k>, as shown in Figure 7-15. These are referred to as the "Pseudo" optical functions to differentiate from the actual material optical functions. They are a direct calculation based on the measured ellipsometry data with the assumption of a single reflection from the surface. This calculation is given in Equation 6-4. The transformation can be performed for any data set, but the "Pseudo" curves lose meaning when the assumption of single reflection is not valid. In other words, they only transform the data correctly for a substrate. The "Pseudo" optical constants are NOT equal to the material optical constants if there are any thin films or surface layers present during measurement of the ellipsometry data.

\[
\langle \varepsilon \rangle = \langle \varepsilon_i \rangle + i\langle \varepsilon_2 \rangle = \langle \bar{n} \rangle^2 = (\langle \bar{n} \rangle + i\langle \bar{k} \rangle)^2 = \sin^2(\phi) \left[ 1 + \tan^2(\phi) \left( \frac{1-\rho}{1+\rho} \right)^2 \right]
\]  

**NOTE:** "Pseudo" optical constants do not represent a layer's optical constants. To view the layer optical constants, right-click on the layer and choose "Graph Layer Optical Constants".

Figure 7-15. <Pseudo> Transforms
**Mueller-Matrix Data**

When Mueller-Matrix data is present, an extra selection in the "Graph Type" menu will be available, as shown in Figure 7-16. These individual selections are described below, but are related to the Mueller-matrix description of a sample as shown in Equation 6-5.

![Figure 7-16. Mueller-Matrix Graphing Options](image)

\[
\begin{bmatrix}
m_{11} & m_{12} & m_{13} & m_{14} \\
m_{21} & m_{22} & m_{23} & m_{24} \\
m_{31} & m_{32} & m_{33} & m_{34} \\
m_{41} & m_{42} & m_{43} & m_{44} \\
\end{bmatrix}
\] (6-5)

**Mueller Matrix Diag, Off Diag**

The diagonal and off-diagonal choices select specific elements from the Mueller-matrix of Equation 6-5 to graph. The diagonal elements will include any of the measured values from \(m_{11}, m_{22}, m_{33}, \) and \(m_{44}\). The off-diagonal elements will include all remaining elements that have been measured.

**Mueller Matrix NCS**

When choosing Mueller-Matrix NCS, the standard Mueller-Matrix elements that are non-zero for an isotropic sample are graphed - which include all of the following that were measured: \(m_{31}, m_{12}, m_{33}, m_{34}, m_{43},\) and \(m_{44}\). These elements relate to NCS of Equation 6-2, via the following.

\[
\begin{bmatrix}
1 & -N & 0 & 0 \\
-N & 1 & 0 & 0 \\
0 & 0 & C & S \\
0 & 0 & -S & C \\
\end{bmatrix} = \begin{bmatrix}
1 & m_{12} & 0 & 0 \\
m_{21} & 1 & 0 & 0 \\
0 & 0 & m_{33} & m_{34} \\
0 & 0 & m_{43} & m_{44} \\
\end{bmatrix}
\] (6-6)
**Select Mueller Matrix Elements**

Opens a dialog box to allow you to select any number of the various measured Mueller-matrix elements for graphing, as shown in Figure 7-17. This dialog box then allows you to quickly select or clear all elements, select the on-diagonal blocks or select the off-diagonal blocks.

Figure 7-17. Select the Mueller-matrix elements you wish to graph by checking the appropriate boxes.

The Mueller Matrix elements available for graphing will coincide with the measured values. The last row/column of the Mueller matrix requires probing and detection of the handedness of the polarization. This requires a compensator before or after the sample. The RC2 has rotating compensators before and after the sample, so it can detect all 16 elements of the Mueller-matrix, though generally represented as ratios to the M_{11} element. Other ellipsometer types may have compensators before or after the sample, which allows them to collect the first three rows or columns, respectively. Rotating analyzer and rotating polarizer ellipsometers (without compensator before/after sample) can’t detect the fourth row or column. These various configurations are shown in Equations (6-7a to 6-7d).

\[
M_{RCE, RPE} = \begin{bmatrix}
\bullet & \bullet & \bullet & \times \\
\bullet & \bullet & \bullet & \times \\
\times & \times & \times & \times
\end{bmatrix}
\]  

(6-7a)

\[
M_{RCE(Compensator–before-Sample)} = \begin{bmatrix}
\bullet & \bullet & \bullet \\
\bullet & \bullet & \bullet \\
\times & \times & \times
\end{bmatrix}
\]  

(6-7b)
This graph helps determine whether the Mueller-matrix can be converted into a corresponding Jones matrix. The Mueller-matrix can handle both isotropic and anisotropic samples with and without depolarization. The Generalized Jones matrix can also handle both isotropic and anisotropic samples, but does not handle depolarization. Thus, the MM-Jones Quality factor will only be close to zero when the conversion is successful, signifying that the depolarization is not a significant issue.

The basic definition of the MM-Jones Quality factor is simply the RMS error between the measured MM parameters and the "closest" (in a least squared sense) Jones matrix. There are exact formulas for converting a Jones matrix (4 complex values) into a Muller Matrix (16 real values). However, there is no exact conversion from MM to Jones (can't convert 16 to 8 values in general), so we perform a non-linear regression fit to find the closest Jones matrix to the MM data, and report the RMS error. In CompleteEASE, the RMS error is then multiplied by 1000, such that a MM-Jones Quality of "1" means that the RMS difference between the closest Jones matrix and the MM data is 0.001, which is the target value for MM data accuracy.

**MM Derived Parameter**

Multiply the specified Stokes vector by the measured Mueller-Matrix to determine the "derived" outcome, as shown in Figure 7-18. The user gets a choice of source polarization state and polarization state detector, as listed in Figure 7-19. In addition, there are a large variety of derived parameters, as listed in Figure 7-20.
Figure 7-18. Mueller-Matrix Derived Parameter

Figure 7-19. Choice of Source Polarization States.

Figure 7-20. Choice of Derived Parameters.
Show Mueller Matrix Utility

This handy utility allows the user to view the Mueller Matrix elements graphically, as shown in Figure 7-21 for a spectroscopic MM scan (X-Y Type) and in Figure 7-22 for a rotational MM scan (Contour Type).

Figure 7-21. Mueller Matrix Graphing Utility.

Figure 7-22. Contour plot of the Mueller Matrix.
In addition to both X-Y and Contour graphing, the user can select the Data Type from choices of i) Experimental Only, ii) Exp and Gen, iii) Diff Exp and Gen, or iv) Generated Only. The standard X-Y Axis will be Wvl vs. Angle for spectroscopic single-position measurements. However, in the case of automated rotation stages that record the orientation of the sample for each measurement, there are also choices of Wavelength (Wvl) vs. Rotation Angle or Angle of Incidence (AOI) vs. Rotation Angle. In these two cases, the remaining parameter (AOI or Wvl) can be slid to different positions. Press the ‘Graph Settings’ button to access additional features, as shown in Figure 7-23. Finally, you can select to maintain this Graphing Utility “Always on Top” of other windows and you can copy both the graph and the data to the clipboard.

Figure 7-23. Graph Settings within the Mueller Matrix Graphing Utility

**Generalized Ellipsometry Ratios**

Anisotropic samples are often characterized using Generalized Ellipsometry measurements, which consider the entire Jones matrix, as shown in Equation 6-7.

\[ J_{Anisotropic} = \begin{bmatrix} \bar{r}_{pp} & \bar{r}_{sp} \\ \bar{r}_{ps} & \bar{r}_{ss} \end{bmatrix} \]  

(6-7)

As ellipsometry measurements are a ratio of Jones matrix elements, there are three measured values possible to access the entire Jones matrix. These values are referred to as AnE, Aps, and Asp. Each is defined in the equations of 6-8.

\[ AnE = \tan(\Psi)e^{i\Delta} = \frac{\bar{r}_{pp}}{\bar{r}_{ss}} \]  

(6-8a)

\[ Aps = \tan(\Psi_{ps})e^{i\Delta_{ps}} = \frac{\bar{r}_{ps}}{\bar{r}_{pp}} \]  

(6-8b)

\[ Asp = \tan(\Psi_{sp})e^{i\Delta_{sp}} = \frac{\bar{r}_{sp}}{\bar{r}_{ss}} \]  

(6-8c)
Select Ratios to Graph

You can select which of the Anisotropic ellipsometry ratios to graph from the dialog box shown in Figure 7-24.

![Figure 7-24. Selecting Generalized Ellipsometry Graphing Options.](image)

Intensity

Reflected or Transmitted Intensity that is collected at same time as SE data using CompleteEASE. The Intensity represented here is an average of the p- and s-polarization components. The accuracy of the Intensity data depends on the instrument calibration. Please consult your Hardware measurement to improve Intensity accuracy.

Because M-2000 and RC2 spectroscopic ellipsometers collect data from the AC signal, the Intensity value measured here is also from the AC signal. Similarly, the model calculations only consider the AC components. Depolarization is measured from the DC signal and will not be measured (or modeled) as part of the Intensity. See p-s (Total) Intensity in the More Options to collect the effects of depolarization on Intensity.

Depolarization

Percent depolarization is collected on ellipsometers that incorporate adjustable or rotating compensators. This information can be used to determine sample and measurement non-idealities such as thickness non-uniformity, spectrometer bandwidth, angular spread, and backside reflections.

In the presence of depolarization, the Mueller Matrix for an isotropic sample is written below, as shown in . In Ellipsometry, all Mueller Matrix elements are normalized to the first element, M11.

\[
M_{\text{sample}} = \begin{bmatrix}
1 & -P \cdot N & 0 & 0 \\
-P \cdot N & P & 0 & 0 \\
0 & 0 & P \cdot C & P \cdot S \\
0 & 0 & -P \cdot S & P \cdot C
\end{bmatrix}
\]

(6-9)

With the Polarization (P) defined as:

\[
P = \sqrt{N^2 + C^2 + S^2}
\]

(6-10)
This gives the Depolarization that is reported by Woollam Ellipsometers as:

\[
\%Depolarization = (1 - P^2) \cdot 100\%
\]  \hspace{1cm} (6-11)

**Graph vs. Angle**

Most SE data is graphed versus wavelength. When multiple angles of incidence have been collected, it can be useful to graph one wavelength versus angle to visualize where the pseudo-Brewster angle occurs (when \(\Psi\) goes through minimum and Delta goes through 90° or 270°). Figure 7-25 graphs the data for an indium tin oxide film (ITO) at 1500nm versus angle of incidence. At this high wavelength, the ITO has become conductive and appears like a metal.

![Variable Angle Spectroscopic Ellipsometric (VASE) Data](image)

*Figure 7-25. Graph of ITO data versus angle of incidence.*

**Double-Y Axis**

View two different data curves simultaneously, with each having its own Y-Axis.

**Difference Mode**

Difference Mode plots the difference between Experimental data and Generated data.

**More Options**

Additional graphing choices are listed in an expanded region, called “More Options”, as shown in Figure 7-26. These include various intensities and error bars.

![More Options from the Graph Type menu](image)

*Figure 7-26. More Options from the Graph Type menu.*
**p- s- Intensity**

The p- and s- Intensity values are measured from the AC signal. Thus, these values represent Rp and Rs for reflection and Tp and Ts for transmission. Because they are calculated strictly from the AC signal of the M-2000 or RC2 ellipsometers, these values will not contain any effect from depolarized light. When CompleteEASE models this data, it also calculates values corresponding to AC measurements.

**p- s- Intensity (Total)**

The p- and s- Intensity (Total) values are measured from both the AC and DC signal in an attempt to measure the “total” reflection or transmission in the p- and s-directions and include effects of depolarized light. This would be better comparison to most spectrophotometric measurements (or even measurements from WVASE32) as most do not distinguish between p-, s- and the unpolarized light along p- and s-.

**Error Bars**

The Error Bars of Psi, Delta, Depolarization, Intensity, and Mueller-Matrix are calculated during measurement to estimate the random error associated with each data point.
7.5. Special Graph Items

This section will detail special graph items that are shown across the top of the Graph panel when working with appended data, uniformity maps, rotation data, or dynamic data. Each of these data types shares some graphing features in common. There are extra items shown across the top of the Graph panel that allow selection and viewing choices specific to these data types. Most will have a drop-down list called “Data Set” that allows selection of each of the different data sets to be graphed, as shown in Figure 7-27.

![Figure 7-27. Drop-down list allows selection of various data sets.](image)

If you right-click just to the left of the drop-down list (over the words “Data Set”), you will be presented with data selection choices, as shown in Figure 7-28. These choices include:

“Select Data Set” adds currently listed data set to the group for viewing and fitting.

“Un-Select Data Set” removes currently listed data set from the group for viewing and fitting, but it remains in the list in case you would like to reselect it at a later time.

“Select Current Data Set ONLY” will un-select all other data sets and select only the current data sets. Again, none of the other data sets is removed from the list, so they can be re-selected at a later time.

“Select All Data Sets” will select all data sets from the list to be active for viewing and fitting.

“Delete Data Set” removes the selected data set from the Multiple-Data Sets List.

“Clear Multi-Data Set Mode” will remove all data sets from the Multiple Data Sets List.
Right-click over the word “Data Set” to get a drop-down list of choices for selecting, deselecting, and deleting data sets from the Multiple Data Set List.

**Multi-Sample (Appended Data)**

If multiple data files are opened at the same time (using the Append Data button), then they will all be included simultaneously in the fit. It is possible to graph each individual data set, by selecting it from the drop-down list, as shown in Figure 7-29. It is also possible to graph all data sets simultaneously, by checking the “Graph All Data Sets” box.

**Figure 7-28.** Right-click over the word “Data Set” to get a drop-down list of choices for selecting, deselecting, and deleting data sets from the Multiple Data Set List.

**Figure 7-29.** Graph window with multiple data sets selected. Only one graph is currently viewed, but all are included in the fit.

**Figure 7-30.** “Graph All Data Sets”.

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Rotation Data

Anisotropic samples are often characterized by measuring multiple sample orientations, which is facilitated by an automatic rotation stage. Each measured data set corresponds to a rotation azimuth and when this data is opened, each orientation is maintained separately, as shown in Figure 7-31. All data sets can be graphed at the same time, or individually, but all “selected” data sets will be included in the fit. As this is Rotation Data, an extra button appears called “Show Rotation Data”. When this check-box is selected, the data from a single wavelength are shown versus rotation angle, as demonstrated in Figure 7-32. To change the wavelength graphed for Rotation Data, choose the ‘Set Ranges’ button from the Data: panel.

NOTE: Hold the CTRL key down and drag the mouse over a range of rotation points to quickly select them all.
Map Data

When uniformity maps are collected – involving multiple points across a wafer, the graph will show this map data as Psi-Delta from a single wavelength versus wafer position. This graph appears when the “Show Map Data” check-box is selected, as shown in Figure 7-33. To change the wavelength graphed for Rotation Data, choose the ‘Set Ranges’ button from the Data: panel. If the “Show Map Data” check-box is de-selected, the spectroscopic data from a single point will be graphed. In addition, a slider-bar will be shown, as in Figure 7-34, that allows you to move “position” back and forth to view all locations across the sample.

Figure 7-33. “Show Map Data” selected will view the Psi-Delta data for a single wavelength across the sample.

Figure 7-34. With “Show Map Data” un-checked, the spectroscopic measurements from the uniformity map can be viewed by sliding the “Position” bar to change locations across the wafer.

With Uniformity maps, you can select multiple points by holding the CTRL key down and selecting the various desired points.

NOTE: Hold the CTRL key down and drag the mouse over a range of map points to quickly select them all.
Dynamic Data

When “Show Dynamic Data” is selected, five wavelengths across the measured spectral range are graphed versus time, as shown in Figure 7-35. If “Show Dynamic Data” is un-selected, a slider-bar appears to allow you to view different time-points of spectroscopic data, as shown in Figure 7-36.

Figure 7-35. Dynamic Data shows five wavelengths versus time.

Figure 7-36. With “Show Dynamic Data” un-selected, the spectroscopic measurement from a single time-point is graphed. A slider bar appears that allows you to move through different time-slices.
7.6. **Graph Scratchpad**

The Graph Scratchpad is used in CompleteEASE to add more control of how graphs appear. While not a complete graphing package, it does allow you to change line colors and offers a few different styles. Additionally, you can change the Legend Names for each curve and choose which curves to graph and whether to place them on a 2nd Y-Axis. An example of the Graph Scratchpad is shown in Figure 7-37.

![Graph Scratchpad Example](image)

*Figure 7-37. Example of the Graph Scratchpad, comparing multiple data curves.*

Select any curve and press the ‘Edit’ button and you will get an expanded dialog box, as shown in Figure 7-38. However, all of these items can be modified directly from the curve by pressing in the associated area. You can also delete any curve by selecting it and pressing the ‘Delete’ button. To delete all curves, there is a ‘Delete All’ button.

![Edit Selected Curve(s)](image)

*Figure 7-38. ‘Edit’ Button provides dialog choices similar to those listed in line.*
By default, each curve will maintain the same color as in the copied graphs. When multiple curves with the same color are compared, this can be confusing and require changes to many of the curve colors. To quickly facilitate this, CompleteEASE allows you to select multiple curves in the Graph Scratchpad, as shown in Figure 7-39. Then, pressing the ‘Edit’ button adds a new check-box called “Automatically Set Colors”, as shown in Figure 7-40. The changes affect all selected curves, as shown in Figure 7-41.

![Graph Scratchpad]

Figure 7-39. Multiple curves are selected in the Graph Scratchpad.

![Edit Selected Curve]

Figure 7-40. With multiple curves selected, check the “Automatically Set Colors” box to change the colors of all consecutive curves.
In addition to the curves that were added from the Graph, you can also create curves by pressing the ‘Create Curve’ button. This will allow you to enter an equation and manipulate the existing curve data using standard mathematical operations, as shown in Figure 7-42. The existing curves are all listed as parameters, as seen by pressing ‘Add Parameter’ button, shown in Figure 7-43.
Press ‘Add Trend’ to fit a Linear, Quadratic or Quartic curve to a selection of the data, as shown in Figure 7-44. You can extend this curve beyond the selected range by a selectable number of “x units” and you can add legend information regarding either the equation or the Y=0 intercept.

NOTE: Trends will be indicated in the Graph Scratchpad by T1, T2, etc. Curves will be indicated by C1, C2, etc. Trends and Curves offer different menu options.

You can select two linear trends and right-click to “Calculate Intersection” point, as shown in Figure 7-45. An example of the intersection is shown in Figure 7-46, where the linear extension of two trends has been added and then their intersection has been determined.
You can also Add Offset to any curve, Graph Difference (at least one curve must be selected), or Graph Ratio (at least two curves must be selected).

Press the ‘Graph Settings’ button to change titles, change font size and show or hide grid lines, as shown in Figure 7-47.

![Graph Settings](image)

*Figure 7-47. 'Graph Settings' button allows access to titles, font sizes, and grid lines.*

Right-click in the Graph Scratchpad graph display to access options similar to the Graph panel in the main CompleteEASE window (Copy Graph to Clipboard, etc.). Graphs created in the Scratchpad can be opened and saved with “Open” and “Save” buttons, respectively.
8. Reference – Measurement Tab

The Measurement tab is shown in Figure 8-1. This tab consists of three panels: System Status, Measurement Controls, and Fit Results. The buttons and choices within each panel can change depending on the hardware configuration. A brief description of all commands within this tab is provided.

![Figure 8-1. Measurement Tab for alpha-SE systems.](image)

8.1. System Status Panel

This panel describes the current hardware status. Before the ellipsometer is ready to measure, the message will state “Not Initialized”. If the ellipsometer is ready for operation, “Waiting to Acquire Data” will be displayed.

8.2. Measurement Controls: alpha-SE Systems

Mode:

Specifies the type of measurement to collect on alpha-SE systems. Further details on each mode are given in Table 8-1.

Sample Alignment:

Specifies how the Z-stage should be used to adjust the sample height before the measurement. Table 8-1 provides details for each choice.
Table 8-1. Measurement Settings for an alpha-SE.

<table>
<thead>
<tr>
<th>Category</th>
<th>Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mode</td>
<td>Standard</td>
<td>Use for most measurements (~10 seconds)</td>
</tr>
<tr>
<td></td>
<td>Fast</td>
<td>Quick measurements (~3 seconds)</td>
</tr>
<tr>
<td></td>
<td>Long</td>
<td>Low reflectivity samples or high precision measurements (~30 seconds)</td>
</tr>
<tr>
<td>Transmission</td>
<td>M.M.</td>
<td>Collects Intensity Transmission measurements.</td>
</tr>
<tr>
<td></td>
<td>M.M. Long</td>
<td>Mueller-matrix measurement – for anisotropic and depolarizing samples</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High Accuracy Mueller-matrix measurement</td>
</tr>
<tr>
<td>Sample Alignment</td>
<td>None</td>
<td>Stage location is below beam for straight-through (90°) measurements.</td>
</tr>
<tr>
<td></td>
<td>Standard</td>
<td>Use for standard reflected measurements. This setting does a quicker search for the light beam near the previous alignment position.</td>
</tr>
<tr>
<td></td>
<td>Robust</td>
<td>A thorough alignment of the light beam that searches over the full translation range. Use for glass substrates with multiple reflected beams.</td>
</tr>
<tr>
<td></td>
<td>Fixed Height</td>
<td>Moves sample stage to height described in hardware configuration file.</td>
</tr>
<tr>
<td></td>
<td>Prompt Height</td>
<td>Allows user to enter the sample stage height for measurement. It is often helpful to first perform a z-stage scan from the Hardware tab.</td>
</tr>
<tr>
<td>Model</td>
<td>Press “Choose from File Dialog.”</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>All default models are saved within various folders including: Common, Advanced, Basic, and Calibration Wafers</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Only models in the COMMON folder will appear in the drop-down list.</td>
</tr>
</tbody>
</table>

For the “Standard” and “Long” modes, the motor home sensors are checked before and after measurement. Also, they both implement Zone-averaging, which collects data with input polarizer set to +45° and -45° and then reports the average. This improves data accuracy.

The M.M. data modes allow measurement of 12 Mueller matrix elements. The absolute intensity of m11 is measured, while all other elements are normalized to m11.

**NOTE:** To improve accuracy of Transmission Intensity and Mueller matrix measurements, perform an “S-T Baseline” or “Off-Sample Baseline” prior to the sample measurement (from the Hardware tab).

**Model**

Specifies which model will analyze the data. If it is set to “None”, data will be acquired, but no data analysis will be performed.

**Save Data after Measurement**

Check this selection to ensure the data are saved after each measurement. The default setting for this option is ON, but it is “sticky” and will retain its new value until the program is turned off.
‘Measure’

Begins a measurement using the settings specified in the “Mode:” and “Sample Alignment:” fields. After data is acquired, the “Model:” is used to analyze the data. Results are placed in the box in the upper right corner of the screen.

Common Models

For convenience, you can move Models that are commonly encountered into the “Common” folder. To move into the Common folder location, go to the Analysis tab and press ‘Open’ from the Model: section. Search for the model of interest and then drag the model name (right side of Figure 8-2) with mouse to the Common folder (left of Figure 8-2, located under Projects tab). You can do this with multiple models.

Figure 8-2. From the Analysis tab, press ‘Open Model’ and then you can drag-and-drop models from any folder (listed on right) into the Common folder (shown in the Projects tab on left).

All models that you move to the Common folder will now appear in the drop-down menu from the Measurement>Measurement Controls>Model drop-down menu. This is shown in Figure 8-3, where the Common folder (left) has five models and these same models appear in the Model drop-down menu (right).

Figure 8-3. The models moved to the Common folder (shown on left) will be quickly accessible from the Measurement tab pull-down Model menu (seen on right).
8.3. M-2000, RC2, and AccuMap-SE Systems

For M-2000, RC2 and AccuMap-SE systems, there are more data collection choices. This chapter will discuss the basic options for typical systems. As there are many different configurations, it is suggested you consult your Hardware manual for the details of your system and configuration. Figure 8-4 shows a representative CompleteEASE Measurement tab for an M-2000 with mapping and camera. Due to the number of choices for any measurement (how to measure, where to measure, etc.), CompleteEASE uses a Recipe to describe all details. These recipe files can be created and saved for any specific measurement or sample type.

Figure 8-4. Measurement tab for an M-2000 system with automated angle of incidence, automated sample alignment, automated sample translation, focusing optics, and an integrated camera.

Measurement Recipe

The measurement controls for this type of system are integrated into a Recipe. Each Recipe describes the three basic components of data collection: 1) Data Acquisition, 2) Mapping Scan settings, and 3) Modeling.

The drop-down box from Measurement>Measurement Controls>Recipe: allow the following options: 1) Prompt for Acquisition Parameters, 2) Prompt for Recipe Components, 3) list of all Common recipes, 4) Choose From File Dialog, and 5) Create/Edit Recipe. These options are shown in Figure 8-5.
Figure 8-5. Recipe Options.

<Select a Recipe>
This listing is not an option, but rather a note to the user that they need to Select a Recipe or change this setting before they can proceed.

<Prompt for Acquisition Parameters>
This option will by-pass the Scan Pattern and Model, assuming you only want to measure at a single point and save data for modeling at a later time. You will be allowed to specify the Data Acquisition Parameters, from a box similar to the one shown in Figure 8-6. The options listed in the Acquisition Parameters box are described in Section 2.3.

Figure 8-6. Acquisition Parameters Setup dialog box.

<Prompt for Recipe Components>
This option allows the user to select the individual recipe components, which include Acquisition Parameters, Scan Pattern, and Model – as shown in Figure 8-7. If the recipe components are not established, they can also be created from the Recipe Components dialog box.
Common Recipes

All recipes that are saved in the Common Folder (C:\CompleteEASE\recipes) will appear in the drop-down list as shown in Figure 8-5. In this example, the Recipe called “55, 65, and 75° 3sec, 3mm thick quick” was created and moved to the Common Folder. To move a recipe to the Common Folder, select the ‘Choose From File Dialog’ option that is described next. This will open the Recipe Dialog box. Navigate through the projects tab on left to find the recipe you would like to add to Common folder. Then, drag-and-drop the recipe listed under Files: to the Common folder listed under File Locations: Projects tab.

‘Choose From File Dialog’

This button opens a window to view the File Location on your computer that may contain recipes, as shown in Figure 8-8. Navigate to find the desired recipe and select it to open.

‘Create/Edit Recipe’

The final button allows you to create or edit a recipe. Each of the options for a recipe will be described later in this chapter.
The three primary components of any recipe are the Acquisition Parameters, Scan Pattern, and Model; as shown in Figure 8-9. Each of these components is discussed in further detail.

![Figure 8-9. Components of a Recipe.](image)

### Save Data after Measurement

Check this selection to ensure the data are saved after each measurement. The default setting for this option is ON, but it is “sticky” and will retain its new value until the program is turned off.

### ‘Measure’

The ‘Measure’ button begins a measurement or recipe. There are also two right-click options for the ‘Measure’ button, as shown in Figure 8-10.

![Figure 8-10. Right-click on the 'Measure' button to choose "Acquire Multiple Measurements" or "Show Recipe Scheduler".](image)

### Acquire Multiple Measurements

Right-click on the ‘Measure’ button to “Acquire Multiple Measurements”.

### Show Recipe Scheduler

The Scheduler is a utility for running multiple recipes without user input. It is primarily designed for cases where multiple samples are loaded onto the translation stage at one time. The scheduler allows each sample to be measured with its own Recipe. To activate the Scheduler utility, right click on the Measure button on the
Measurement tab and choose the “Show Recipe Scheduler” menu item. Currently the scheduler supports the following steps:

- Run Recipe – run a predefined recipe
- Acquire Dynamic – acquire dynamic data for a specified amount of time
- Delay – pause for the specified amount of time
- Set Temperature – set the temperature and wait a specified amount of time
- Move Translator – move the translator to a specified position
- Move Angle – move angle to specified position
- Run Temperature Profile – run saved temperature profile
- Move Z Stage – move Z stage to specified position

A dialog box, as shown in Figure 8-11, will appear and allow the commands to be established in a list form.

![Scheduler dialog box](image)

Figure 8-11. Scheduler dialog box.

When a step is added, it can be formatted to be of any “type”, as listed in the dropdown list of Figure 8-12. Each of the step types are described below.

**Run Recipe**

The run recipe step type allows the user to specify a recipe to be run, as shown in Figure 8-13. A translator offset can be specified for the step that offsets the recipe’s translator positions by a specified amount, as shown in Figure 8-14. After this single entry is added, the Scheduler will list the “step” in a row, as shown in Figure 8-15.

**NOTE:** For the offset to apply, the recipe must be set up such that the Do Not Reposition Translator box is **not** checked.
The Acquire Dynamic step allows the user to specify a model number (as listed in the buttons of the In Situ tab) and an amount of time to acquire data, as shown in Figure 8-16.
**Delay**

The Delay step inserts a delay into the schedule. The delay time is specified in minutes, as shown in Figure 8-17.

![Figure 8-17. Adding a “Delay”.](image)

**Set Temperature**

The Set Temperature step is used to change the temperature of a supported temperature controller and then wait a specified amount of time for the temperature to stabilize, as shown in Figure 8-18.

![Figure 8-18. Setting Temperature in the Scheduler.](image)

**Move Translator**

The Move Translator step moves the translator to the specified position, as shown in Figure 8-19.

![Figure 8-19. Adding a Move Translator Step.](image)

**Move Angle**

Moves angle to specified position.
Run Temperature Profile

Choose saved temperature profile from file dialog to add to scheduler.

Move Z Stage

Moves Z Stage to specified position.

Setting up Filenames

The ‘Set up Filenames’ button allows the data files to be named when running a schedule. Figure 8-23 shows the Save Name Setup dialog. The default option is “Automatically assign file name - <custom name><step #><time>”. This option prompts the user for a text string when “Run Schedule” is pressed and prepends this text to the beginning of each automatically generated file name. The automatically generated file name consists of the step number and the current time.

The second option is similar to the default except that the user is not prompted for a text string. This results in a filename consisting of the recipe name, step number, and time.

The third option is to manually define the file names. When this option is chosen, the user must go to each step in the scheduler that requires a file name and double click on the table cell and choose a file name from the standard Save Data dialog.
8.4. Acquisition Parameters

The Acquisition Parameters describe how the data will be collected at each point of a Measurement Recipe (whether single-point or scan). A common box for Acquisition Parameter choices is shown in Figure 8-24. The individual details for these options are discussed in the corresponding hardware manual.

Figure 8-24. Acquisition Parameters Setup for M-2000 and RC2 systems. Choices listed above can vary from system-to-system.
Data Type

The Data Type specifies the type of data to collect during the measurement, as shown in Figure 8-25.

![Data Acquisition Parameters](image)

**Figure 8-25. Different Options for Data Type.**

**Standard**

Most applications require a “Standard” measurement. This is simply a standard Spectroscopic Ellipsometry measurement. The data collected will include Psi and Delta versus wavelength and angle of incidence. In addition, the depolarization and reflected intensity will be measured. However, the measurement accuracy of depolarization and reflected intensity depends on the accuracy of recent DC calibrations and intensity baselines, respectively. For more details, please consult your Hardware manual. Most models will ignore the depolarization and Intensity measurements, unless instructed to consider this extra data.

**Generalized Ellipsometry**

Generalized ellipsometry measurements include the complete Jones matrix of a sample, including the off-diagonal elements (which are assumed equal to zero for Standard SE measurements). This type of measurement is useful for anisotropic samples where the optical axis is not aligned parallel or perpendicular to the plane of incidence. Generalized Ellipsometry measurements are not compatible with measurements that exhibit depolarization, as it is based on a Jones matrix description of the interaction between measurement beam and sample.

Generalized Ellipsometry measurements consist of three ratios, so you will end up with three Psi curves and three Delta curves for each measured angle of incidence.

**Mueller Matrix**

Measurement of the polarization change using Stokes vectors to describe the light beam. The Mueller-matrix is a 4 X 4 matrix, but not all elements can be measured with each ellipsometer. In fact, the only ellipsometer to measure all 16 elements of the Mueller-matrix is the RC2 system. For most samples, there is significant redundancy in the Mueller-matrix and no need to measure all 16 elements. For isotropic samples (with or without depolarization), the complete interaction can be described by measuring 4 elements (M11, M21 or M12, M33 or M44, and M34 or M43). If the sample also contains anisotropy, the number of necessary elements climbs to 7 or 8. Only for the most advanced samples are more elements required for complete characterization. Thus, less than 1% of all Ellipsometry Applications require Mueller-matrix measurements.
**Transmission Intensity**

Measure Intensity of transmitted light through the sample. For M-2000 systems, the transmitted intensity is calculated from the AC components of the detected signal and will not include any measurement of unpolarized light.

**Reflection Intensity**

Measure Intensity of reflected light. For M-2000 systems, the intensity is calculated from the AC components of the detected signal and will not include any measurement of unpolarized light.

**Sample Alignment**

Sample Alignment options depend on the type of system and configuration. In general, there are two basic steps to sample alignment: Tilt alignment and Height alignment. Common selections for each are described in this section.

**Tilt Alignment**

Tilt alignment refers to the tip and tilt of sample relative to the incoming light beam such that the reflected beam is aligned on the receiver unit at the correct angle of incidence. Tilt alignment options include Skip, Manual, or Automated; as shown in Figure 8-26.

<table>
<thead>
<tr>
<th>Sample Tilt Alignment:</th>
<th>Automatic</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Skip</td>
</tr>
<tr>
<td></td>
<td>Manual</td>
</tr>
<tr>
<td></td>
<td>Automatic</td>
</tr>
</tbody>
</table>

*Figure 8-26. Options for Sample Tilt Alignment.*

The Tilt alignment is often skipped when using focusing probes, as the tip-tilt alignment becomes less sensitive. It can also be skipped for applications where common samples are measured that remain flat. For example, if a system is continuously used to measure 200mm silicon wafers, the tip-tilt alignment of each wafer will probably be very similar from sample-to-sample.

For Manual or Automatic Alignment, the system will study the reflection of a light beam off the sample that is incident on a 4-quadrant detector. This detector allows the beam to be centered onto the 4-detector elements by adjusting tip and tilt until all 4 detectors have the same intensity. This method allows very repeatable alignment of the beam from sample-to-sample. An example of the Tilt Alignment step for an Automatic System is shown in Figure 8-27.
Sample Height Alignment

Sample Height Alignment refers to the adjustment of ellipsometer beam relative to the surface of a sample to ensure correct positioning and adjustment for samples of different thickness. This adjustment becomes increasingly important for systems with smaller spot size (like Focusing ellipsometers). It is also more important to test the Sample Height Alignment when many samples of different thickness will be tested on the same Ellipsometer. The common choices for Sample Height Alignment are shown in Figure 8-28. For Automated Height Alignment, the reflected beam is detected by the Receiver unit and the intensity is recorded versus different Z-Height positions. This is demonstrated in Figure 8-29. The position of highest intensity is assumed to be the correct Z-Height for alignment of the sample.
Common Acquisition Parameters

When finished setting the Acquisition Parameters, you will be asked to save these details in a file. If you will use the same Acquisition Parameters for future measurements, the files can be saved or moved to the Common Folder. Then, they will appear in the drop-down list when selecting your Recipe components. To move Acquisition Parameters into the Common Folder, select the ‘Edit/Create’ Acquisition Parameters button from the Recipe Components box. Next, press the ‘Load’ button from the Acquisition Parameters box. This will open the File Dialog box, as shown in Figure 8-30. Find the Acquisition Parameter Files from the appropriate Folder and then drag them with the mouse from the list of Files: in the right to the Common folder under File Location: Projects tab in the left.

Figure 8-30. When opening Acquisition Parameters, you can find files that you would like to see immediately from the Recipe drop-down menu and drag them into the COMMON folder shown on the left of this box.
8.5. Scan Pattern

The Scan Pattern is used to describe where each measurement should be performed across a sample. This option is only needed for ellipsometers with automated sample translation. To make a new Scan Pattern, press ‘Edit/Create’ button to the right of the Scan Pattern selection of the Recipe Components box, as shown in Figure 8-31.

![Figure 8-31. Choose ‘Edit/Create’ Scan Pattern from the Recipe Components dialog box.](image)

The Scan Pattern Editor, shown in Figure 8-32, allows you to describe both Circular and Rectangular samples. There are many options to automatically fill-in the points to measure, including Grid Fill, R-T Grid Fill, and Line Fill. Most of these selections are intuitive and best learned by testing each button a few times.

![Figure 8-32. Scan Pattern Editor](image)
As an example, if you press the ‘Grid Fill’ button, you will get the options shown in Figure 8-33. The Scan Pattern also describes the overall Alignment methodology. For example, you can choose to Align at a single point only. You can also choose to do a full alignment at the first point and then z-alignment at all other points.

**Figure 8-33. Some Grid Fill Options.**

### Scan Region of Interest

Scan Region of Interest allows you to specify a grid over a portion of the substrate instead of the entire substrate. The Top, Bottom, Left, and Right values are entered relative to the center of the substrate.

For example, if the substrate is a 10x10 cm square, you can setup a grid over the bottom right portion of the substrate by entering:

Top=0, Left=0, Right=5, Bottom=-5

### Adding Common Scan Patterns

After you have developed Scan Patterns that meet your measurement requirements, you can save these files for future use. If they will be routine patterns, you can add the Scan Patterns to the Common Folder. To accomplish this, press ‘Load’ from the Scan Pattern Editor. This will open the “Open Scan Pattern” dialog box, as shown in Figure 8-34. Find the files you will routinely use and then simply drag them with your mouse from the listing on right to the COMMON folder (in the Projects tab of File Location: panel) shown on the left.

**Figure 8-34. Open Scan Pattern Editor. To move routine Scan Pattern files from any location, simply drag them with the mouse from the list on right to the COMMON folder listed to the left of this screen.**
8.6. **Fit Results Panel**

Data analysis results are placed in this text box, located in the upper right corner of the screen, as demonstrated in Figure 8-35.

![Figure 8-35. Fit Results panel in the Measurement Tab.](image)

8.7. **Right-click menu (Measurement>Fitt Results):**

**Copy to Clipboard – Formatted**

Copy results to the clipboard in a format ready to paste into a word processor, as shown below.

\[
\text{MSE} = 2.837 \\
\text{Thickness} \# 1 = 96.71 \pm 0.018 \text{ nm} \\
n \text{of Cauchy Film @ 632.8 nm} = 1.865
\]

**Copy to Clipboard - Table**

Copy results to the clipboard in a format ready to paste into a table or spreadsheet, as shown below.

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>VALUE</th>
<th>ERROR BAR</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSE</td>
<td>2.837</td>
<td></td>
</tr>
<tr>
<td>Thickness # 1 (nm)</td>
<td>96.71</td>
<td>0.018</td>
</tr>
<tr>
<td>n of Cauchy Film @ 632.8 nm</td>
<td>1.865</td>
<td></td>
</tr>
</tbody>
</table>

8.8. *‘View Previous Fit Results’*

Review results from past samples. This button will open a new dialog box, as shown in Figure 8-36. Choose the **Location** and **Files** you wish to review and a table of...
results is shown in the **File Information** panel. This table can be copied to the clipboard with the ‘Copy Text to Clipboard’ and ‘Copy Table to Clipboard’ buttons.

**Figure 8-36. View Previous Fit Results Dialog box.**

Select a single data file from the **Files** panel and you can ‘Open File’. This will open the data file, but not the model that was associated with the previous fit.

Use the mouse to select multiple results in the **Files** panel and you will be allowed to ‘Calc. Multi-Sample Stats’. An example of this is shown in Figure 8-37.

**Figure 8-37. Viewing statistics from multiple samples.**

If the selected data file is a map of multiple points across the sample, all points are shown in the list, along with the statistics across the top: average, min, max, standard deviation, %range, and %uniformity; as shown in Figure 8-40.

Alignment information will be shown if the “Include Alignment Parameters with Data” parameter is turned ON in the Hardware Configuration prior to measurement.
Figure 8-38. Viewing statistics from multiple samples – with alignment parameters included.

Figure 8-39. Include Alignment Parameters With Data.

%RANGE

The percent range is defined as:

\[
%\text{RANGE} = 100\% \left( \frac{\text{MAX} - \text{MIN}}{\text{AVERAGE}} \right)
\]

%UNIFORMITY

The uniformity is defined as:

\[
%\text{UNIFORMITY} = 100\% \left( \frac{\text{STDEV}}{\text{AVERAGE}} \right)
\]
Map results will show the average, min, max, standard deviation, %Range, and %Uniformity of each parameter across the top of list of all points.

For map data, the ‘Graph Data’ button can be selected to open a new window that allows the user to select the fit result they wish to view, as shown in Figure 8-41.
Figure 8-41. Uniformity map of total thickness.
9. Reference – Analysis Tab

The Analysis tab is shown in Figure 9-1. This tab consists of three panels: Data, Model, and Fit.

9.1. Data: Panel

‘Open’

Opens a data file. Right-click menu is shown in figure below.
Figure 9-2. Right-click menu for ‘Open’ from Data: panel, gives access to “Append Data”, “Append Intensity Data from Text File”, and the Recent List of Data Files to open.

(⌘R): “Append Data”

Appends data to currently open data. The data files must share the same wavelengths.

(⌘R): “Append Intensity Data From Text File”

Starting in CompleteEASE version 4.98 the software supports appending unpolarized transmission and reflection data from sources other than a J. A. Woollam ellipsometer. After loading or acquiring ellipsometry data, the transmission/reflection data can be added by right clicking on the data “Open” button in the Analysis tab and selecting “Append Intensity Data From Text File” from the menu. A file dialog then appears allowing the user to select the desired file. After appending, this data is treated just like transmission and reflection data acquired by the ellipsometer. The format for the tab delimited text file is as follows:

```
<Comment_Line>
<Data_Type>TAB<Optional_AOI#1> TAB <Optional_AOI#2> TAB<...>
<Wavelength_Units>
<Wavelength>TAB<Data_AOI#1>TAB<Optional_Data_AOI#2>TAB<...>
...
```

The first three lines in the file describe the content of the data. The first line is for a comment. The second line describes the type of data in the file. The options are “uT” for unpolarized transmission data and “uR” for unpolarized reflection data. By default the angle of incidence is assumed to be 0° but additional items on the second line are used to specify the angle(s). The third line specifies the wavelength units for the file. The choices are: “Å”, “nm”, and “eV” for angstroms, nanometers, and electron volts. The rest of the lines specify the data in ascending wavelength order. Example data files are shown below:
Opens data files are stored in a “Recent List”, accessed by right-click menu of ‘Open’ button. See Figure 9-2 for example.

‘Save’

Save a data file. CompleteEASE data files are saved as encrypted files that can only be opened with CompleteEASE. Their extension will be “.SE” for standard measurements or “iSE” for dynamic measurements.

Choose to “Save Data Subset” or “Export to Text File”.

“Save Data Subset”

Right-click on ‘Save’ to choose this option. The data subset will consist of the reduced data that are currently selected (wavelengths, angles, time, multiple points from a map, etc.).
“Export to Text File”

Right-click on ‘Save’ to choose this option. Data will be exported to a text file.

‘Info’

Shows information about the current data set opened in CompleteEASE. An example of the information box is shown in Figure 9-3.

![Information box showing details of data measurement](image)

Figure 9-3. Information box showing details of data measurement.

‘Set Ranges’

Choose the wavelength range and selected angles for use with current data set during fit. Also allows selection of translation point from uniformity maps, as shown in Figure 9-4. When mapped data are graphed, this dialog box also allows choice of the graph angle and wavelength.
9.2. **Fit: Panel**

**‘Generate’**

__(⌘L):__ Generate data based on the current model and data set. Shortcut keystroke to Generate data is Alt-G.

__(⌘R):__ “Clear Generated Data” allows removal of the generated data curves from the graph.

**‘Fit’**

Fit data based on the current model and data set. Shortcut keystroke to Fit data is Alt-F.

**‘Fit Scan Data’**

When uniformity maps are present, the third button will show ‘Fit Scan Data’ to allow the model to be fit to each point from the uniformity map in sequence. It is important to clear multi-sample mode before pressing this button.

**‘Fit Dynamic’**

__(⌘L):__ Fit dynamic data that has been taken versus time.

__(⌘R):__ Choose to “Append Fit Results”, “Fit Backwards in Time”, or “Fit Backwards and Append”.

---

Figure 9-4. Press ‘Set Ranges’ to select the wavelengths, angles, translation points and graphing angles/wavelengths.
“Append Fit Results”
When fitting Dynamic Data, it can be helpful to fit different regions of time separately. This menu option would be used after the first region has already been fit. The new range can be fit and appended to the previous fit results.

“Fit Backwards in Time”
Use this option for dynamic data where the “end” result is better understood than at the beginning of the experiment. For example, it can be used for Etching studies, where the thickness at the last time is easier to determine.

“Fit Backwards and Append”
This feature combines the previous two. It will perform the fit starting at the longest selected time and work backward – but then the results will be appended to previous dynamic data fit results.

‘Reset’

(⌘L): Restore the previous model parameters, from before the fit.

(⌘R): Choose to “Show Reset List”.

“Show Reset List”
Provides a list of reset points that have been saved automatically by CompleteEASE before and after each “fit” since the most recent data/model has been in use. An example is shown in Figure 9-5.

Figure 9-5. Reset List allows user to reset to any recent positions, which are automatically saved before and after each fit.
9.3. **Right-click menu (Analysis>Fit:)**

Right-click the mouse within the **Fit**: Panel to access the menu options as shown in Figure 9-6. As seen, the selections can vary depending on whether fitting a single point measurement, a uniformity map, or dynamic data.

![Figure 9-6. Right-click options from the Fit: panel after a fit to spectroscopic data and uniformity mapped data.](image)

**“Auto Fit”**

When this check-box is selected, the data analysis will automatically proceed when new data are loaded or a point is selected. When un-checked, the data analysis only begins after user presses the ‘Fit’ button.

**“View Fit Stats”**

After each fit, there are statistical values available that include the 90% confidence limits on each fit parameter and a correlation matrix which shows the correlation between each fit parameter, as shown in Figure 9-7. Correlation values close to +1 or -1 show two parameters are achieving the same purpose and are not unique.

![Figure 9-7. “View Fit Stats” shows statistics such as the 90% confidence limits and correlation matrix values after a fit.](image)

**“View Parameter Stats”**

After fitting a uniformity map, this selection allows you to view the statistics from all points in a map, along with the values from each location, as shown in Figure 9-8.
Figure 9-8. “View Parameter Stats” for all points of a Uniformity map.

“Copy to Clipboard – Formatted”
After fitting single point data, you can copy the results to the clipboard in a formatted layout to be used by word processors or presentations.

“Copy to Clipboard – Table”
After fitting single point data, you can copy the results to the clipboard in a tabulated layout to be used by spreadsheets.

“Copy Parmeters to Clipboard”
After fitting a Uniformity Map, you can copy the results from all points to the clipboard.

“Copy Analysis Report to Clipboard”
After fitting, you can easily place the results into a report. The format of the report will depend on the type of data analysis. For example, a single-point analysis will appear as shown in Figure 9-9. The inclusion of the Optical Constants graph for this report is optional and controlled within the Model: OTHER Options section.
After analysis of a uniformity map, the graphs from the map are copied as part of the report, rather than the spectral fit quality, as shown in Figure 9-10. Finally, the graphs from each spectral fit will all be copied if a multi-sample analysis has been performed, as shown in Figure 9-11.
Figure 9-9. Analysis Report from single point fit, with optical constants for layer added (from Model: +OTHER Options).

Figure 9-10. Analysis Report from uniformity map.
“Add to Fit Log”
Adds the current Fit results to the Fit Log for later viewing. CTRL-L is the shortcut to add to fit log.

“View Fit Log”
View the Fit Log, which lists any fit results recorded there since CompleteEASE was started, as shown in Figure 9-12. The Fit log is cleared when CompleteEASE is closed, but can be saved and opened at a later time.

‘Rename’
Select a Log Entry and then rename.
‘Delete’
Select a Log Entry and delete from the Fit Log.

‘Compare’
Select multiple Log Entries and compare statistics, as shown in Figure 9-13. From this Comparison dialog box, you can “Add Statistics”, “Reverse Columns/Rows” and even ‘Compare Optical Constants’, which brings up a new dialog box as shown in Figure 9-14.

Figure 9-13. Fit Log Comparison shows results from each fit log entry.

Figure 9-14. ‘Compare Optical Constants’ brings up a box that shows each model and allows you to choose the different layers to graph from each model.
‘Reanalyze’

The Reanalyze Data dialog box is shown in Figure 9-15. It uses a table for specifying analysis steps, consisting of a File Group, a Model, and a Log Name. A File Group is a list of files to be reanalyzed. It can consist of one or more files in the same directory. The model is a model file that has been saved to disk. The Log Name is a text string that will be appended to the data file name when adding the results of the reanalysis to the Fit Log.

![Reanalyze Data dialog box](image)

To add an Analysis Step to the table, press ‘Add Row’. This brings up the Choose Files dialog, as shown in Figure 9-16. By default the only file group available is the “Selected Log Entries” group which contains the files from any fit log entries that were selected before pressing the Reanalyze button. To add a group of files to the available file groups press the Add button. This button brings up the standard file dialog except that it allows multiple files to be selected. After selecting the desired files and closing the dialog, you are given the chance to name the file group, as shown in Figure 9-17. Note that if only one file is selected for a group then the name is automatically set to the name of the file. After giving the group a name, it now shows up in the Choose Files dialog, as in Figure 9-18. The file groups that are created will be available until the software is restarted.

![Choose files to add to groups for reanalysis](image)

![Enter File Group Name](image)
Figure 9-18. *New Group Name shows up with the associated files.*

To finish adding the row, choose a file group from the Choose Files dialog and press Ok. A row is now displayed in the reanalysis table with the file group specified, as shown in Figure 9-19. A different group can be specified by double-clicking on the cell you wish to change. This also applies to the Model and Log Name items. To finish setting up the analysis, specify a model and log name by double clicking the appropriate cell and entering these parameters, as demonstrated in Figure 9-20.

Figure 9-19. *File Groups show up as a row in the Reanalysis dialog box.*

Figure 9-20. *Click to add Models and optional Log Names for each row.*

Once the analysis is set up, the ‘Reanalyze’ button can be pressed to start the analysis. The software will spin through all of the files and add the results to the current Fit Log. The fit log for the example below is shown in Figure 9-21.
‘Generate Reports’
Copies results of selected Log entries into a report in RTF format, which is saved to a file.

‘Load Log Entry’
You can select a Log Entry from the list and open this result by pressing ‘Load Log Entry’. The same is accomplished by double-clicking on the Log Entry.

‘Clear Log’
Clears log of all entries.

‘Open Log’ / ‘Save Log’
Opens a saved log. This is a convenient way to save results on your computer, but requires that the data files are located in the same directory, as they are not saved as part of the Log.

‘Close’
Closes the Fit Log.

“Advanced Graph Options”
After fitting a uniformity map, there are “Advanced Graph Options”, which allow various combinations of X-Y parameters, as shown in Figure 9-22. This allows graphical interpretation of possible correlation between parameters.

More importantly, after a dynamic fit, you can graph various fit parameters versus time, as shown in Figure 9-23.
Figure 9-22. After fitting a uniformity map, there are “Advanced Graph Options”, which allow various combinations of X-Y parameters.

Figure 9-23. Using Advanced Graph Option, the Thickness is plotted versus time after a dynamic data analysis.

“Graph OCs vs. Time”

This choice allows the spectroscopic optical constants versus time to be graphed. When selected, you will be asked for the type of optical constant graph, as shown in Figure 9-24, the layer number, and the total number of curves to draw, as shown in Figure 9-25. These curves will be equally spaced versus time across the selected range of the data analysis. An example is shown for index of refraction of amorphous silicon growth in Figure 9-26.

Figure 9-24. Choose from n, k, e1, or e2.
9.4. **Model: Panel**

The box in the right half of the *Analysis* tab displays all the options and specifications for the current data analysis model. An example model is shown in Figure 9-27. Details of each option within the model are given below.

**Figure 9-27. Basic Model: Panel in the Analysis Tab.**
‘Open’

(⌘L): Open a new model.

(⌘R): Merge models or View Recent List of Models to open.

![Model: Si with Transparent Film]

Figure 9-28. Right-click on Model: ‘Open’ button to either “Merge Models” or get a list of recently opened models.

“Merge Models”

Right-click on the ‘Open’ button to merge models together.

‘Save’

(⌘L): Save current model and all model settings.

‘Clear’

Remove the current model.

‘Open Snapshot’

(⌘L): Open a snapshot.

(⌘R): View list of recently opened snapshots to open.

A snapshot is a single file that compresses all information from the CompleteEASE screen, including the data measurement, model, and fit results. This is a convenient way to save and store information. Because it includes all data, the snapshot can be overwhelming size if created from large uniformity map data.

‘Save Snapshot’

Saves a snapshot file, which is a single file that compresses all information from the CompleteEASE screen, including the data measurement, model, and fit results. This is a convenient way to save and store information. The file will be given “.SESnap” file extension.

Add

Press this button to add a layer. The Add Layer to Model dialog box appears. Select the position of the new layer by moving the mouse to adjust the blue-bar and press the mouse button (Figure 9-29). If Layer Type is changed to “Intermix” before pressing the blue-bar in place, the layer will be an EMA mixture between the layer below and above.
NOTE: Holding the CTRL key down when pressing the Add button will automatically add the new layer on top of the current model layers, rather than asking for the position.

NOTE: With Backside Correction turned on, it is possible to add layers below the substrate.

**Delete**

Press this button to delete a layer from the model. Choose the layer with the mouse (making it blue) and click to delete (Figure 9-30).

**Save**

Press this button to save optical constants for a layer. Before clicking on the layer you wish to save (Figure 9-31), choose the Save Type (either Dispersion Parameters or Tabulated).
Include Surface Roughness

Turn on to fit a thin layer on the surface of existing model. The surface is modeled with a Bruggemann Effective Medium Approximation mixing the top film with 50% void. The only fit parameter available is the thickness of the rough surface. If the thickness moves to negative values, the roughness layer is equivalent to -50% void mixing, which increases the surface region optical properties (rather than lowering them). The surface roughness layer is a “stealing” layer, in which it takes half of its thickness from the layer below. Thus, if a 100nm Cauchy layer has 20nm surface roughness on top, the model is actually calculated as 90nm Cauchy with 20nm surface roughness.

Model Pictorial

The model is shown as a layer-by-layer picture with details regarding thickness and other parameters to describe the individual layers. Any layer with a “+” can be expanded to show more details about that layer. This is demonstrated in Figure 9-32.

![Figure 9-32. (a) Model with layers collapsed and (b) with Cauchy Film layer expanded.](image)

Editing Layers

Each layer has a name, shown in blue. Place your mouse over the layer name to do the following:

(⌘L): Open a new material file in place of the current layer.

(⌘R): Choose from a menu to “Graph Layer Optical Constants”, “Append Text to Layer’s Fit Parameters”, “Save Layer Optical Constants”, “Parameterize Layer”, “View Layer Comment”, “Convert To EMA”, “Convert to Anisotropic”, or “Grade Layer”. This menu is shown in Figure 9-33.
“Graph Layer Optical Constants”
Right-click on a layer to select this option – the optical constants will be graphed.

“Rename Layer and Fit Parameters”
Right-click on a layer to select this option – will bring up a dialog box, as shown in Figure 9-34. There are three options for renaming the layer in the model, the layer parameters in both the model and the fit results, and the thickness in the fit results.

“Save Layer Optical Constants”
Right-click on a layer to save optical constants for that layer. You will be asked whether to save the Dispersion model parameters or a Tabulate list and then proceed to the File Location.

“Parameterize Layer”
Right-click a layer to parameterize the optical constants of that layer using a dispersion model, such as the General Oscillator layer. This will open the layer optical constants as reference values in the Parameterize Layer dialog box, as shown in Figure 9-35.
“View Layer Comment”

Right-click and choose this option to view the layer comments, which often tell the source of the material file.

“Convert to EMA”

Right-click to replace the layer with an EMA, in which the original layer is now embedded as the host material (Material 1). The second material is default to “Void” with EMA% default to 0, but selected as a fit parameter. An example is shown in Figure 9-36.

```
Layer #1 = EMA  Thickness #1 = 124.13 nm (fit)
# of Constituents = 2
+ Material 1 = Cauchy Film
  Material 2 = VOID
  EMA % (Mat 2) james = 0.0  (fit)
  depolarization_james = 0.333 Analysis Mode = Bruggeman
Substrate = Si_JAW
```

Figure 9-36. Example of the EMA layer where the Cauchy Film was automatically converted to the host (Material 1) with void as the second material.
“Unwrap from EMA”

After converting a layer to EMA, you can right-click on the EMA layer to unwrap the host film back to a single layer without the EMA.

“Convert to Anisotropic”

Right-click to replace the layer with an anisotropic Biaxial Layer, in which the original layer is now embedded as the starting material for both the Ex and Ez directions. The default configuration is to have no orientation (determined by Euler angles) and anisotropy set to Uniaxial (Nxy ≠ Nz). An example is shown in Figure 9-37.

<table>
<thead>
<tr>
<th>Layer # 1 = Biaxial</th>
<th>Thickness # 1 = 124.13 nm (fit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type = Uniaxial</td>
<td></td>
</tr>
<tr>
<td>Optical Constants:</td>
<td>Difference Mode = OFF</td>
</tr>
<tr>
<td>+ Ex = Cauchy Film</td>
<td></td>
</tr>
<tr>
<td>+ Ez = Cauchy Film</td>
<td></td>
</tr>
<tr>
<td>Euler Angles:</td>
<td>Phi = 0.000  Theta = 0.000</td>
</tr>
<tr>
<td>Substrate = SI_JAW</td>
<td></td>
</tr>
</tbody>
</table>

Figure 9-37. Example of Anisotropic layer.

“Convert to Isotropic”

After converting a layer to Anisotropic, you can right-click on the Biaxial layer to unwrap the host film back to an isotropic layer.

“Grade Layer”

Right-click to replace the layer with Graded Layer, where the original film is embedded as the average refractive index. The % inhomogeneity is default to 0%, but can be varied to allow the index to increase or decrease from the bottom to top of the layer. An example of this is shown in Figure.

<table>
<thead>
<tr>
<th>Graded Layer</th>
<th>Thickness # 1 = 124.13 nm (fit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grade Type</td>
<td>Simple</td>
</tr>
<tr>
<td>% Inhomogeneity</td>
<td>0.00  (fit)</td>
</tr>
<tr>
<td>Material</td>
<td>Cauchy Film</td>
</tr>
<tr>
<td>A = 1.460 (fit)</td>
<td>B = 0.00091 (fit)</td>
</tr>
<tr>
<td>C = 0.00034 (fit)</td>
<td></td>
</tr>
<tr>
<td>k Amplitude</td>
<td>0.000000 Exponent = 1.500</td>
</tr>
<tr>
<td>Band Edge</td>
<td>400.0 nm</td>
</tr>
<tr>
<td>Substrate</td>
<td>SI_JAW</td>
</tr>
</tbody>
</table>

Figure 9-38. Example of Graded Layer.

“Remove Grading”

After grading a layer, you can right-click on the Graded layer to unwrap the host film back to a layer without grading.

“Start Superlattice”

Choose this layer as the start of a repeated structure.
“End Superlattice”

This layer is designated as the top of the repeated structure. When selected, an additional section will be added to this layer to control the number of repeats (called SL Cts for superlattice counts), as shown in Figure 9-39.

<table>
<thead>
<tr>
<th>Layer #</th>
<th>Material</th>
<th>Thickness (nm)</th>
<th>SL Cnt</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>BaTiO3</td>
<td>20.00</td>
<td>5</td>
</tr>
<tr>
<td>1</td>
<td>Cauchy Film</td>
<td>40.00 (fit)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Substrate</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 9-39. Model with superlattice start, end, and number of counts (repeats).

“Clear Superlattice”

Removes layer from repeated structure. Select for both layers (start and end).

Angle Offset

Apply an offset to the angle when calculating the model. This parameter can be fit.

9.5. Right-click menu (Analysis>Model:)

Right-click the mouse within the Model: Panel to access the menu options as shown in Figure 9-40.

Draw Model Optical Constants Profile
Copy Model To Clipboard
Copy Model (Layers Only) To Clipboard

Figure 9-40. Right-click menu from within the Model: Panel.

Draw Model Optical Constants Profile

Shows the depth profile of the optical constants from the substrate (left) to the surface (right), as shown for a graded layer with a surface film, as shown in Figure 9-41.

Figure 9-41. Depth profile for a 2 layer stack, where the bottom layer is graded.
Draw All Layer Optical Constants

Plot optical constants from all layers. Choose $n$ or $k$ (or $e_1$ or $e_2$, depending on display units specified in the Options tab).

Figure 9-42. All Layer Optical Constants – $n$.

Draw All Layer Optical Constants (No Substrate)

Plot optical constants from all layers, excluding substrate. Choose $n$ or $k$ (or $e_1$ or $e_2$, depending on display units specified in the Options tab).

Figure 9-43. All Layer Optical Constants (No Substrate) – $n$.

Copy Model To Clipboard

This option copies the entire Model to the Clipboard, including the Model Options, Fit Options and Other Options that are saved along with the Layer description. To view the details from a section, such as the Model Options, it needs to be expanded by pressing the “+” next to that section. An example of this “full” model is shown in Figure 9-44.
Figure 9-44. “Full” Model as copied to Clipboard.

Copy Model (Layers Only) To Clipboard

This option copies the Layered-description of a Model to the Clipboard. If you prefer to see a layer collapsed or expanded, make sure to pre-select your view before copying. An example of this model is shown in Figure 9-45.

Roughness = $0.00 \text{ nm}$ (fit)

\[
\begin{array}{l}
+ \text{ Layer } \# \ 1 = \text{ B-Spline} \quad \text{Thickness } \# \ 1 = 100.00 \text{ nm} \ (\text{fit}) \\
\text{ Substrate } = \text{ SI_JAW} \\
\end{array}
\]

Figure 9-45. “Layer-Only” Model as copied to the Clipboard.
9.6. **Layer types**

CompleteEASE provides many different ways to represent the optical constants of a material versus wavelength. These representations range from simple tabulated lists for the optical constants to complex dispersion models. All of these different representations are classified as layers in the CompleteEASE program. The layers currently available are listed in Table 9-1. The rest of this chapter describes the operation and typical uses of the different layers in detail.

**Table 9-1. Layer types in CompleteEASE.**

<table>
<thead>
<tr>
<th>Layer Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard (Tabulated N,K)</td>
<td>Contains tabulated values for the optical constants of a material.</td>
</tr>
<tr>
<td>Void</td>
<td>Contains optical constants for air (vacuum): $n=1$, $k=0$.</td>
</tr>
<tr>
<td>EMA</td>
<td>A layer for mixing the optical constants of two materials together to create a composite material.</td>
</tr>
<tr>
<td>EMA-Coupled</td>
<td>A layer for mixing the optical constants of two materials together to create a composite material. In this case, the Material #1 is a “Coupled” layer which gets optical constants from a separate layer – referenced by the layer number.</td>
</tr>
<tr>
<td>Cauchy</td>
<td>A simple dispersion layer using three terms to describe the index and an exponentially decaying function to describe the extinction coefficient.</td>
</tr>
<tr>
<td>Cauchy_Wvl</td>
<td>A simple dispersion layer using three terms to describe the index. In this special layer, the index can be defined at a specific “design” wavelength.</td>
</tr>
<tr>
<td>Cauchy_Extended</td>
<td>Similar to Cauchy (described above) with additional terms.</td>
</tr>
<tr>
<td>Sellmeier</td>
<td>A simple dispersion layer useful for modeling dielectric materials.</td>
</tr>
<tr>
<td>Coupled</td>
<td>Obtains optical constants from a separate layer in model.</td>
</tr>
<tr>
<td>WvlByWvl</td>
<td>Allows optical constants to be individually determined on a wavelength-by-wavelength basis. This layer does not enforce Kramers-Kronig consistency and inherently contains a maximum number of fit parameters. Thus, it should be used with caution.</td>
</tr>
<tr>
<td>Biaxial</td>
<td>Layer to describe the anisotropic optical constants of both uniaxial and biaxial materials.</td>
</tr>
<tr>
<td>Graded</td>
<td>All other layer types can be graded to model the optical variation with depth through the film.</td>
</tr>
<tr>
<td>General Oscillator</td>
<td>A layer that allows multiple oscillators of various types to be used to represent the optical constants of a material.</td>
</tr>
<tr>
<td>B-Spline Layer</td>
<td>The B-Spline layer calculates a basis spline through evenly spaced points in eV for both $\varepsilon_1$ and $\varepsilon_2$.</td>
</tr>
<tr>
<td>Internix</td>
<td>Mixes the optical constants from the material below and material above in a 50-50% ratio.</td>
</tr>
<tr>
<td>Temperature/Composition</td>
<td>This layer contains optical constants of a material as a function of temperature and/or composition.</td>
</tr>
</tbody>
</table>
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<table>
<thead>
<tr>
<th>Layer Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uniaxial</td>
<td>Enter any material file into the Ordinary OC (in-plane optical constants) and Extra-Ordinary OC (out-of-plane optical constants) sections.</td>
</tr>
<tr>
<td>Uniaxial-Diff</td>
<td>The Uniaxial-Diff layer is similar, but you only enter a material file for the Ordinary optical constants. Extra-Ordinary optical constants are calculated as a “difference” from the Ordinary optical constants.</td>
</tr>
<tr>
<td>Additive</td>
<td>Sums optical constants together.</td>
</tr>
<tr>
<td>User Defined</td>
<td>Allows custom equation for optical constants of a material.</td>
</tr>
<tr>
<td>Virtual Substrate</td>
<td>Allows the underlying sample structure to be approximated by a single interface. The optical constants at this interface are called the “pseudo substrate.”</td>
</tr>
</tbody>
</table>

Tabulated n,k

Most of the layers in the CompleteEASE material library are standard tabulated optical constant lists. The optical constants for this type are represented as a table of values versus wavelength. The table can contain the real and imaginary parts of the materials' dielectric function ($\varepsilon_1$ and $\varepsilon_2$) or the materials' complex index of refraction ($n$ and $k$). If the user finds optical constant tables in the literature and wishes to use them in CompleteEASE, simply type the table into a file in the correct format and they can be opened.

Figure 9-46 shows how typical standard layers are represented in the Model. Unlike the rest of the layer types described in this chapter, these have no user adjustable parameters to define the optical properties making it the simplest layer type to use.

```
Layer # 2 = SiO2_JAW  Thickness # 2 = 25.00 nm
Layer # 1 = INTR_JAW  Thickness # 1 = 1.00 nm
Substrate = Si_JAW
```

**Figure 9-46.** A model constructed of standard layers – each set of optical constants is described by a table of values versus wavelength.

Void

The Void layer is a special case of Tabulated N,K layer where the optical constants are fixed for all wavelengths at the nominal values for air (vacuum): $n=1$, $k=0$.

EMA

The EMA layer uses the Effective Medium Approximation to calculate the optical constants of a mixed material. Calculation is based on mixing optical constants of two or three constituent materials, the percentage of each material, a depolarization “screening” factor, and the calculation type. An EMA layer is shown in Figure 9-47.

The EMA layer in CompleteEASE provides three different EMA mixing methods for the user to choose. These include a simple Linear combination, Bruggemann EMA, and Maxwell-Garnet EMA. The Linear mixing simply interpolates between the constituents dielectric functions to get the optical constants of the composite material. The Maxwell-Garnett and Bruggemann EMAs are more common. The Maxwell-Garnett assumes that spherical inclusions of materials #2 and #3 exist in a host matrix of material #1. The Bruggemann EMA makes the self-consistent choice...
of the host material. The Maxwell-Garnett and Bruggemann EMAs are very useful for modeling surface and interfacial roughness as well as modeling poly-crystalline materials by mixing together amorphous and crystalline optical constants of the material.

<table>
<thead>
<tr>
<th>Layer #1 = EMA</th>
<th>Thickness #1 = 50.00 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td># of Constituents = 2</td>
<td></td>
</tr>
<tr>
<td>Material 1 = a-si</td>
<td></td>
</tr>
<tr>
<td>Material 2 = Void</td>
<td></td>
</tr>
<tr>
<td>EMA % (Mat 2) = 23.0</td>
<td></td>
</tr>
<tr>
<td>depolarization = 0.333 Analysis Mode = Bruggeman</td>
<td></td>
</tr>
</tbody>
</table>

Figure 9-47. EMA model for mixed-constituent layers.

**EMA-Coupled**

This is a special case of EMA layer, just as described above, except with the first material using the Coupled description. This allows the Material #1 to obtain optical constants from a separate layer in the model, designated by the layer number. An example of the EMA-Coupled layer is shown in Figure 9-48.

<table>
<thead>
<tr>
<th>Layer #3 = EMA-coupled</th>
<th>Thickness #3 = 0.00 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td># of Constituents = 2</td>
<td></td>
</tr>
<tr>
<td>- Material 1 = Coupled</td>
<td></td>
</tr>
<tr>
<td>Coupled to Layer #2 = B-Spline</td>
<td></td>
</tr>
<tr>
<td>Material 2 = Void</td>
<td></td>
</tr>
<tr>
<td>EMA % (Mat 2) = 50.0</td>
<td></td>
</tr>
<tr>
<td>depolarization = 0.333  Analysis Mode = Bruggeman</td>
<td></td>
</tr>
<tr>
<td>Layer #2 = B-Spline</td>
<td>Thickness #2 = 0.00 nm</td>
</tr>
<tr>
<td>Layer #1 = Cauchy</td>
<td>Thickness #1 = 0.00 nm</td>
</tr>
<tr>
<td>Substrate = Si_JAW</td>
<td></td>
</tr>
</tbody>
</table>

Figure 9-48. Example of the EMA-Coupled layer, where Material #1 receives its optical constants from a separate layer in the model, referenced by the layer number.

**Cauchy**

The Cauchy layer is a commonly used layer for determining the optical constants of a transparent or partially transparent film (Dielectrics and Semiconductors below the fundamental bandgap). Over part of the spectral range, the optical constants of these materials can be represented by an index that varies slowly as a function of wavelength and an exponential absorption tail. The index of refraction of the Cauchy layer is represented by an inverse power series containing only even terms and the extinction coefficient is represented by a simple exponential tail. These representations are as follows:

The Cauchy is a dispersion layer that describes the index of refraction using a three-term equation, as in Equation 8-1.

**NOTE:** The wavelength units in the Cauchy layer, as expressed in Equation 8-1 as \( \lambda \) are microns.
In addition to the three terms for index of refraction \( (n) \), there are three additional terms to describe an Urbach absorption tail, as in Equation 8-2. The Cauchy layer and graph of the optical constants is shown in Figure 9-49.

\[
n(\lambda) = A + \frac{B}{\lambda} + \frac{C}{\lambda^2}
\]  

(8-1)

\[
k = k_{\text{amp}} \cdot e^{\exp(-E_{\text{band edge}})}
\]  

(8-2)

Figure 9-49 shows the Cauchy layer as displayed in the CompleteEASE model. The A, B, C parameters are variable fit parameters that determine the index dispersion. The \( k \text{ Amplitude} \) and \( k \text{ Exponent} \) are fit parameters for determining the shape of the extinction coefficient dispersion. The \( \text{Band Edge} \) parameter can be set manually but is not a fit parameter since it is directly correlated to the \( k \text{ Amplitude} \) parameter (the extinction coefficient is equal to \( k \text{ Amplitude} \) at the \( \text{Band Edge} \)).

\[
\left( \frac{A - B}{\lambda B} \right)^2 + \left( \frac{C}{\lambda^2} \right)^2 = 1
\]

\[
k = k_{\text{amp}} \cdot e^{\exp(-E_{\text{band edge}})}
\]

Cauchy_WVL

This special representation of the Cauchy layer is found in the ADVANCED materials directory. It provides the same shape as the Cauchy, but is described with the terms shown in Figure 9-50. This allows the user to specify a known index of refraction at a specific wavelength, without concern for the dispersion terms.

Figure 9-49. Cauchy dispersion layer in CompleteEASE.
- Substrate = \text{Cauchy\_Wvl}
  \begin{align*}
  n &= 1.450 \quad B = 0.01000 \quad C = 0.00000 \\
  \text{Design wvl} &= 632.8 \text{ nm}
  \end{align*}

Figure 9-50. Cauchy\_Wvl layer is same dispersion shape as Cauchy, but allows entry of the index at a design wavelength in place of the $\Lambda$ parameter of the standard Cauchy.

\textbf{Cauchy\_Extended}

This special representation of the Cauchy layer is found in the ADVANCED materials directory. It provides a similar shape to the Cauchy, but with ability to decrease index quickly toward NIR spectral region by adding an IR term. An example layer is shown in Figure 9-51 with the resultant optical constants shown in Figure 9-52.

- Layer \# 3 = \text{Cauchy\_Extended}  \quad \text{Thickness} \# 3 = 0.00 \text{ nm}
  \begin{align*}
  A &= 1.450 \quad B = 0.00500 \quad C = 0.00000 \\
  D &= 0.00000 \quad \text{IR} = 0.12000 \\
  \text{Urbach Amplitude} &= 0.30000 \quad \text{Exponent} = 2.000 \\
  \text{Cody Amplitude} &= 0.200 \\
  \text{Band Edge (eV)} &= 2.400
  \end{align*}

Figure 9-51. Example of the Cauchy\_Extended layer.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{fig9-52.png}
\caption{Optical constants from a Cauchy\_Extended layer.}
\end{figure}

\textbf{Sellmeier}

The Sellmeier layer is another dispersion layer that can be used to represent dielectrics. It is used to represent the optical constants of transparent dielectric materials. CompleteEASE uses a 2-term Sellmeier model to account for UV and IR absorption. The representation for the index of refraction is:
where \( \varepsilon(\infty) \) is an index offset, \( A \) is the amplitude, \( B \) is the center energy, and \( E \) is the position of a pole in the infrared. These are represented in the layer as ‘Einf’, ‘Amp.’, ‘Center En.’, and ‘IR Pole Amp.’ respectively. Figure 9-53 shows the layer as it appears in a model.

\[
 n = \left( \varepsilon(\infty) + \frac{A\lambda^2}{\lambda^2 - B^2} - E\lambda^2 \right)^{\frac{1}{2}} 
\]  

(8-3)

**Coupled**

Layer that has same optical constants as another layer in the model. An example is shown in Figure 9-54

**WvlByWvl**

Layer, as shown in Figure 9-55, which allows the optical constants to vary independently at each wavelength. This layer has the utmost flexibility, but also the largest number of fit parameters. As it does not maintain Kramers-Kronig consistency or even smooth continuous curves for the optical constant dispersion, it should be used with extreme caution.

**Biaxial**

The biaxial layer allows the user to model anisotropic materials. This layer is used for both uniaxial and biaxial anisotropy and has 3 Euler-angle terms to rotate the orientation of optical axes relative to the ellipsometer measurement orientation. There are two modes which can be used within the Biaxial layer. With Difference
Mode = OFF, the optical constants in all directions (2 for uniaxial, 3 for biaxial) are independently described by material files. This is shown in Figure 9-56. With Difference Mode = ON, the x-directed optical constants are described by a material file while an extended Cauchy “difference” is established for the index difference between the x-direction and the other orientations. This is shown in Figure 9-57.

**Figure 9-56. Biaxial anisotropic layer with Difference Mode = OFF**

![Figure 9-56](image1)

**Figure 9-57. Biaxial anisotropic layer with Difference Mode = ON**

![Figure 9-57](image2)

### Graded Layer

All other layer types can be graded to model the optical variation with depth through the film. This is achieved by right-clicking a layer and choosing to “Grade Layer”. An example Graded Layer is shown in Figure 9-58. There are four types of grading that can be achieved in this layer: Simple, Linear, Non-Linear, and Parametric. Grading is accomplished by dividing the thin film into a series of ‘slices’ that will have varying optical constants to approximate the index gradient profile. The number of slices can be designated. The primary fit parameter is the “% Inhomogeneity” which will be positive numbers when the index is larger toward the surface than the substrate and negative numbers if the index is smaller toward the surface. If you right-click on the title “Graded Layer”, the optical constants will be graphed, showing n and k at both top and bottom, as shown in Figure 9-59.

**Figure 9-58. Example of a Graded Layer.**

![Figure 9-58](image3)
Grade Type

There are four different types of grading available, as shown in Figure 9-60. Details for each are described below.

Simple Grade

Allows linear variation of the index by fitting or adjusting the “% Inhomogeneity”. The index can either increase or decrease from the bottom of film to the film surface, depending on whether “% Inhomogeneity” is positive or negative, respectively. An example of the Simple Grade is shown in Figure 9-61, where a Cauchy layer exhibits 4% inhomogeneity. The Cauchy layer remains embedded in the Graded Layer, so the “average” index of the layer can still be adjusted by the Cauchy parameters. The grading of index will occur in both directions of the Cauchy – with a total range of index between the top and bottom of the layer equaling the “% Inhomogeneity”.

NOTE: Simple grade does not affect the extinction coefficient.
Linear Grade

The Linear Grade uses an EMA to mix the current material with void. The amount of void through the film is set by the “Slope” and “Offset” of the line. The “Offset” defines the position where void goes to zero (or graded film exhibits same properties as original film). An example of the “Linear” grade is shown in Figure 9-62, with a slope of 0.1 and an offset of -0.2. The EMA void percentages are calculated for bottom and top of layer, based on these slope and offset.

The EMA percentage is not necessarily a physical mixture of the material with void. Rather, it is a mathematical calculation to adjust the optical constants up or down. In fact, the EMA% can go negative to specify the index and extinction coefficient go “up” (further away from void) rather than “down”. The main difference between Linear and Simple Grade is that the Linear Grade affects both index and extinction coefficient, where Simple grade only affects index.

Non-Linear Grade

The “Non-Linear” grade is very similar to the Linear grade, in that it uses an EMA mixing to vary the optical constants between bottom and top of the layer. The main difference is that it allows the linear variation to be modified to exponential variation with either symmetric profile about the center or a profile that proceeds from bottom to top with same exponential shape.

Figure 9-63 shows a “Non-Linear” graded layer with embedded Cauchy. The bottom and top EMA percentages can be set (or fixed) along with the exponent to describe the grading shape. Press the “Draw Profile” text to see how the void percentage is being distributed through the various slices.
A wide variety of grading profiles are possible, depending on whether the profile is symmetric or not and the value of exponent is below 1 or above 1, as shown in Figure 9-64 through Figure 9-67.

**Figure 9-63. Example of Non-Linear Grade.**

**Figure 9-64. Grade Profile for a 21-Slice “Non-Linear” graded layer with Exponent = 0.2 and Symmetry turned ON.**

**Figure 9-65. Grade Profile for a 21-Slice “Non-Linear” graded layer with Exponent = 2.5 and Symmetry turned ON.**
Figure 9-66. Grade Profile for a 21-Slice ‘Non-Linear’ graded layer with Exponent = 0.2 and Symmetry turned OFF.

Figure 9-67. Grade Profile for a 21-Slice ‘Non-Linear graded layer with Exponent = 2.5 and Symmetry turned OFF.

Slice Value

The profile is described by an equation between the top and bottom of the film. Then, the number of slices is evenly distributed through the film and the value of each slice is taken from the value at the center of each slice.

NOTE: The bottom and top “slice” do not have to be equal to the values of the equation at the bottom and top of film, as they get their value from the “slice” center.

Parametric Grade

The parametric grade allows select dispersion model parameters to vary with depth through the film, while the remaining parameters are constant. Thus, this layer only works with layers such as the Gen-Osc and Cauchy. An excellent example of parametric grading is used with indium tin oxide and other transparent conductive oxides. In these films, there can be significant grading due to variation in the “free-carrier” concentration with depth. The free-carriers lead to infrared absorption, which is modeled with a Drude oscillator. Thus, the variation in the film properties is not well represented by EMA mixing with void. Instead, the amount of IR absorption will change from bottom to top, requiring a variation in the Drude oscillator amplitude. The UV oscillator absorptions will generally stay constant through the film – thus we want to grade only the Drude. Figure 9-68 is an example of ITO layer with Drude grading by linear variation of the Resistivity of the Drude(RT) oscillator.
Figure 9-68. Example of “Parametric” grade with a linear variation of the Drude resistivity term.

After the Grade Type is converted to Parametric, there are four choices for Grade Equation, as shown in Figure 9-69. The Linear and Exponential grade equations allow the parameters to be varied through the film in a linear or exponential fashion. However, these grade types can be difficult for parameters that are close to zero.

Figure 9-69. Choice of Grade Equation for Parametric Grade types.

**Custom Grade Profile**

A custom grade equation has been added to the list of equations available under the Parametric Grade Type. The built-in variables for the Grade Profile are the position in the layer \([\text{pos}]\) which varies from 0 to 1 and value \([\text{value}]\) of the parameter being graded. Variable names must be enclosed in brackets. The figures below show how the custom grade equation can be set up to model a linear grade of the amplitude of a Cody Lorentz oscillator. The grade equation is evaluated at the center of each slice.
to determine the parameter value for that slice. If a grade equation does not include the [value] variable then the parameter being graded should be turned off as a fit parameter as it will be singular. If the user forgets to do this the fit will proceed correctly but the graded parameter will have an error bar of 9999. Also, when adding custom grade equations for more than one parameter that any fit parameter names that are the same between equations will be treated as the same fit parameter. This eliminates the need to manually couple parameters after entering the equations.

Figure 9-70. Custom equation for Parametric grade. Click <Enter an Equation> to open the Edit Equation window.

Figure 9-71. Edit Equation window.

Figure 9-72. Custom equation has been applied. % Grade is now available as fit parameter.

Two Segment

The two segment grade equation allows variation of the parameters between the bottom, top, and a middle node – with linear variation between bottom-middle and middle-top nodes. In addition, the position of the middle node can be varied. An example of “two-segment” grade equation is shown in Figure 9-73, where the bottom and middle resistivity values are set to 0.1 while the top value is 0.2. The middle
“Position” is set to 60% through the film. This gives the grade profile shown in Figure 9.74.

NOTE: Even though the grade profile has 2 segments, the 21 slices are distributed equally through the entire layer.

Figure 9-73. Example of a “Parametric” grade with “Two-Segment” equation. The bottom and middle resistivity values are set to 0.1 while the top value is 0.2. The middle “Position” is set to 60% through the film.

Figure 9-74. Grade profile for resistivity associated with the two-segment equation in the previous figure.

**Gen-Osc**

The Gen-Osc layer is a General Oscillator material that allows the summation of different oscillator line shapes. Figure 9-75 shows a Gen-Osc layer with two oscillators of different types, along with associated graph of the resulting optical constants.

Within the Gen-Osc layer you can add a new oscillator by pressing “Add Oscillator”. To change the oscillator type, click on the “Type=” selection and you will get a dialog box with your choices, as in Figure 9-76. The various oscillator functions are described in detail in Chapter 10.
Figure 9-75. Gen-Osc dispersion layer with both a Cody-Lorentz and a Gaussian oscillator.

Figure 9-76. Choices for Oscillator Types within the Gen-Osc layer.

Press “Show Dialog” to bring up the Editor to visualize and manipulate the oscillator parameters. This dialog editor is shown in Figure 9-77.
Figure 9-77. Dialog Editor to view and manipulate the Gen-Osc parameters.

NOTE: The oscillators can be manipulated by pressing the number to their left. Use care when pressing the number, as each mouse button does a different function:

- (R) Delete the oscillator from list.
- (L) Selects the oscillator and shows gray “drag-boxes” to allow manipulation of the oscillator parameters on graph.

The Dialog Editor allows the oscillator parameters to be matched to reference optical constants. This procedure is demonstrated in the Analysis Example on use of the Gen-Osc layer.

B-Spline

The B-Spline layer specifies the optical constants versus wavelength using a series of control points which are equally spaced in photon energy (eV). The approximate spacing of the control points is controlled by the “Resolution (eV) =” field and the resulting number of control points within the current spectral range is listed after this value. Basis spline curves are used to interpolate between the control points.

B-Spline curves are defined by a series nodes, which have a position and amplitude. Basis polynomial functions at each node are defined by the recursive B-Spline formula (Cheney and Kincaid, “Numerical Mathematics and Computing,” Third Edition, Brooks/Cole Publishing Company, 1994) in terms of the positions of the adjacent nodes. Summing the basis functions (weighted by the amplitude at each

---

node) produces the resulting curve. B-Spline curves have many desirable properties for modeling dielectric functions:

- B-Spline curves and their derivatives (up to the spline order minus one) are continuous (the B-Spline layer uses 3rd order B-Splines)
- the B-Spline node amplitudes affect only the “local” shape of the curve, e.g., changing a node amplitude in the UV does not affect the curve in the VIS
- B-Spline curves exhibit a “convex hull” property, e.g., if all the node amplitudes are non-negative, then the B-Spline curve is also non-negative, thus avoiding non-physical negative ε2 values
- since the basis functions depend only on the node positions, the node amplitudes which define the resulting curve are linearly independent, which greatly increases computation efficiency
- the Kramers-Kronig (K-K) integral can be applied to the B-Spline recursion formula to generate K-K consistent basis functions (i.e., the ε1 curve is defined by the K-K transform of ε2)

Figure 9-78 shows the default settings for the CompleteEASE B-Spline layer. The node amplitudes can be initialized to constant n and k values (vs. eV) by clicking on the “n=” or “k=” fields, or they can be initialized by specifying a “Starting Mat”. The “Resolution (eV)” field defines the nominal spacing of the nodes, which are equally distributed in eV over the selected spectral range of the data set. Clicking the “Draw Node Graph” field plots the positions and amplitudes of the B-Spline nodes, along with the calculated dielectric function of the layer (Figure 9-79). The “Fit Opt. Const.” setting turns ON and OFF all the optical constant fitting parameters for the layer (e.g., node amplitudes and positions, pole amplitudes, etc.). If “Use KK Mode = OFF”, separate B-Spline curves are calculated and fit for ε1 and ε2. When “Use KK Mode = ON”, a B-Spline curve is used to define ε2, and ε1 is calculated from via the K-K transform of ε2 (plus contributions from absorption outside the measured spectral range). Figure 9-81 shows the Node positions when KK Mode is turned on. Notice, there are nodes to describe ε2, along with a few nodes outside the measured spectral range – to help with the Kramers-Kronig integration.

NOTE: The B-Spline layer will automatically extrapolate/interpolate/seed node amplitudes and positions if the selected spectral range and/or resolution is changed, or if KK Mode is toggled.

Substrate = B-Spline
Init. values: n = 1.500 k = 0.000 Starting Mat = none
Resolution (eV) = 0.300 6 Pts. (1.379-3.177 eV) Draw Node Graph
Fit Opt. Const. = ON
Use KK Mode = OFF
Show Advanced Options = OFF

Figure 9-78. Default settings for the B-Spline layer.
Figure 9-79. B-Spline layer Node Graph with KK Mode = OFF

Figure 9-80. B-Spline layer with all advanced options displayed (KK mode off)
**Figure 9-81.** B-Spline layer Node Graph with KK Mode = ON.

**Figure 9-82.** B-Spline with all options shown, with KK Mode = On.
All the available options for the B-Spline layer are shown in Figure 9-80 for a B-Spline with KK mode off and Figure 9-82 for a B-Spline with KK mode on.

A brief description of each option follows:

**Init. Values**

Entering values for n or k in the Initial Values section will reset all node values across the entire spectrum to these values.

**Starting Mat**

Entering a material file in the “Starting Mat” section will reset the node values to match the reference material files across the spectrum. This can be an effective way to get reasonable “starting” node values in the B-Spline layer.

**Resolution (eV)**

Spacing of nodes across the spectrum.

**Draw Node Graph**

Press to show the node values across the spectrum. This is useful to view whether the node resolution is adequate to resolve the spectral features of a material.

**E Inf**

With KK mode = ON, this represents the $\varepsilon_1$ value at infinity. When in KK mode, this accounts for the absorption far outside the measured spectral range by adding a constant offset to the $\varepsilon_1$ curve.

**IR Amp**

With KK mode = ON, this value is the amplitude of a zero energy oscillator which accounts for absorption at low energies, typically due to free carrier absorption (i.e., a Drude term).

**IR Br**

With KK mode = ON, this is the broadening of the IR oscillator. Often this can be fixed at zero, but allowing to vary can improve the fit to some metallic dielectric functions.

**Fit Opt. Const.**

Turn this On when you want to allow the node values to vary to match the experimental data. After fitting is completed, this can be turned Off to hold the final optical constant values fixed.

**Use KK Mode**

When turned ON, the optical constants are represented by a single spline curve for $e_2$ with the KK transformation used to determine the $e_1$ curve. Adds Einf, IR Amp, IR Br, and extra nodes outside the measured range (tie-offs) to accommodate the KK transformation and determine the correct $e_1$ curve.
Query remote system for Opt. Const.

Used with multiple ellipsometers to “feed” the optical constants for a layer from a remote system (first ellipsometers) when fitting layers that need this updated information.

Show Advanced Options

Many additional options are hidden to avoid confusion. Turning this On will allow you to view the following Advanced options.

Show Nodes

Shows each node in the B-Spline, so you can turn on/off the fit for any of the individual values.

Spline_e2(photon energy) / Spline_e1(photon energy)

Each node that is shown is given as either the e1 or e2 value at a specific photon energy. These nodes are positioned equally across the spectrum. When KK mode is Off, there is a complete set of e1 and e2 nodes. When KK mode is On, only the e2 nodes are listed, as the e1 curve is generated from a Kramers-Kronig transformation.

Show Parms in Fit

Enables display of the all node positions and amplitude fit parameters in the Fit Results; by default they are hidden to reduce the number of displayed parameters.

Force E2 Positive

Setting this to ON prevents $\varepsilon_2$ nodes within the measured spectral range from going negative. The tie-off nodes outside the spectral range are still allowed to go negative, so it is possible that $\varepsilon_2$ may still go negative at the edge of the spectrum. While forcing $\varepsilon_2$ positive maintains “physical” dielectric functions, sometimes allowing slightly negative $\varepsilon_2$ values improves the fit convergence algorithms.

Prefit when Changing Wavelengths

It is difficult to maintain the optical constants of a B-Spline when changing wavelengths because of the effect of tie-off positions outside the selected spectral range on the optical constants. When the spectral range is modified, the tie-off positions and “values” will adjust the overall optical constant shape. When this feature is turned On, the B-Spline will save off optical constants before adjusting to the new spectral range that can be used to “prefit” the new nodes to match. This does not ensure the optical constants will not slightly adjust when changing wavelengths, but helps minimize the adjustment by prefitting to the previous values.

Node Spacing Spectral Ranges

A single node spacing across the entire spectrum may not be optimal for all materials. For this reason, CompleteEASE allows special spectral ranges to be “added” with smaller or larger node spacing. GaP is a good example, as shown in Figure 9-83, where the optical constants have no spectral features below the bandgap (above 500nm) but quickly change at shorter wavelengths. It may be desirable to reduce the total number of nodes by having a wide node spacing at longer wavelengths and narrow node spacing at shorter wavelengths. Press to “Add Node Spectral Spacing Range” and you will be presented with three dialog boxes, as shown in Figure 9-84. The first will ask for the minimum wavelength range, the
second for the maximum wavelength range, and the third for the custom node spacing to be used within this range. For GaP, I have selected a node spacing of 0.5eV everywhere, with custom node spectral range from 190-500nm of 0.1eV, as shown in the model in Figure 9-85. The final node graph is shown in Figure 9-86.

![Opt. Const. of GaP vs. nm](image)

**Figure 9-83.** Optical Constants for GaP from ultraviolet to near infrared.

![CompleteEASE Input](image)

**Figure 9-84.** Input the minimum/maximum wavelengths and the node spacing over this range.

![B-Spline for GaP](image)

**Figure 9-85.** B-Spline for GaP with custom Node Spacing between 190-500nm (0.1eV resolution).
Assume Transparent Region

A bandgap or spectral region can be specified for which $\varepsilon_2$ is fixed at zero.

Use Default TieOff Behavior

“TieOffs” are the nodes which account for absorption outside the spectral range when in KK mode. The “default” behavior (which specifies default TieOff node positions, with amplitudes being fit for each node) is adequate for most applications. Turning the Default TieOff Behavior = OFF allows for complete flexibility in the TieOff positions and amplitudes. This is typically only needed if there is a strong absorption feature right outside the measured spectral region that is dramatically influencing the optical constants.

View Tie Off Positions

The positions and amplitudes of the Tie Offs can be set and defined as fit parameters. The positions are defined relative to the ends of the spectral range, and relative to each other to guarantee monotonically increasing node positions. Five node positions are defined above and below the spectral range; the amplitudes of the first and last two nodes are fixed at zero, while the other six node amplitudes can be fit.

Intermix

An Intermix is added between two materials in the model using the Layer Command Add. Select Intermix as Layer Type, and then click on the desired position in the model. Intermix optical properties are modeled as a 50:50 mix of material above and material below. Intermix thickness can be entered or selected as a fit parameter.

An example of this is shown in Figure 9-88. Notice the Intermix is not a numbered layer. Intermix thickness is an independent thickness quantity, with half of its thickness extending into Layer 1, and half of its thickness extending into Layer 2. The thickness of the numbered Layers remains the same. Defining the Intermix “layer” in this manner maintains the total optical thickness of the film stack, independent of the intermix thickness. This helps to orthogonize the layer thickness and intermix thickness fit parameters, as the layer thickness essentially specifies the optical thickness of the film (which, for example, determines the period of any thin film interference oscillations), and the intermix thickness specifies how diffuse or
non-abrupt the interface is (which, for example, may affect the magnitude or dampin
do of any thin film interference oscillations).

Figure 9-87. Choose “Intermix” before adding a layer between two layers to create an
Intermix between materials.

Figure 9-88. Example of Intermix layer in a model.

**Composition Library**

Layers that are marked as “Comp Library” or “Comp & Temp Library” were created
to show variation in optical constants as the composition or temperature are varied.

Several of the material files provided with the CompleteEASE program contain
information on the optical constants of a material as a function of temperature and/or
alloy fraction. One example of this type of material is Al$_x$Ga$_{1-x}$As. The file
AlGaAs_T.MAT supplied with the CompleteEASE software contains information on
the optical constants of Al$_x$Ga$_{1-x}$As over $x$ values from 0 to 1 and temperature values
from 577°C to 650°C. Using this type of layer is very similar to using the standard
layer type for a material; the only difference being that there are temperature and/or
alloy parameters that can be edited and fit for. Figure 9-89 shows how such a layer
is represented in the model.

Figure 9-89. A Temperature/Alloy layer as displayed in the model. The composition and
temperature values are editable fit parameters.

**Uniaxial and Uniaxial-Diff**

Layers used to describe anisotropic materials. The Uniaxial layer is shown in Figure
9-90. Enter any material file into the Ordinary OC (in-plane optical constants) and
Extra-Ordinary OC (out-of-plane optical constants) sections. In this example,
Cauchy dispersions were used. The Uniaxial-Diff layer is similar, but you only enter
a material file for the Ordinary optical constants. The remaining terms are used to
express the index difference in a Cauchy-like summation to determine the Extra-
ordinary optical constants. An example of the Uniaxial-Diff layer is shown in Figure
9-91.

- **Substrate = Uniaxial**
  - **Ordinary OC = Cauchy**
    
    | A | B       | C       |
    |---|---------|---------|
    | 1.450 (fit) | 0.01000 (fit) | 0.00000 |
  
  
  k Amplitude = 0.00000 Exponent = 1.500
  Band Edge = 400.0 nm

- **Extra-Ordinary OC = Cauchy**
  
<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.550 (fit)</td>
<td>0.01000 (fit)</td>
<td>0.00000</td>
</tr>
</tbody>
</table>
  
  k Amplitude = 0.00000 Exponent = 1.500
  Band Edge = 400.0 nm

**Figure 9-90.** Example of the Uniaxial layer. Enter any material file for both the ordinary (in-
plane) and extraordinary (out-of-plane) optical constants.

- **Substrate = Uniaxial-Diff**
  - **Ordinary OC = Cauchy Wvl**
    
    | n | B       | C       |
    |---|---------|---------|
    | 1.450 | 0.01000 | 0.00000 |

  Design wvl = 632.8 nm
  
  A = 0.10000
  B = 0.00000
  C = 0.00000
  D = 0.00000

**Figure 9-91.** Example of the Uniaxial-Diff layer. Enter any material file for the ordinary (in-
plane) optical constants and then use the 4-parameter dispersion (Cauchy-like) to describe the
index difference.

**Additive**

Sums optical constants together in terms of e1 and e2. Click “**Add Material**” to
specify layers for summing. A simple example is shown in in

- **Layer # 1 = Additive** Thickness # 1 = 100.00 nm (fit)
  
  **Add Material**
  
  **Substrate = Si_JAW**

**Figure 9-92.** Additive layer.
Figure 9-93. Additive example: constant offset of $e_1=1$ is added to SiO2.

**User Defined**

The user defined layer can be found in the Advanced material folder. Click **Edit Equation** to open the Edit Equation dialog window. This dialog allows an equation to be entered in terms of $n$ and $k$ or $e_1$ and $e_2$. Variable names must be enclosed in brackets. The units of the wavelength variable [wvl] can be set to Å, nm, um, eV, or 1/cm. An example of how a Cauchy dispersion layer can be set up using the User Defined layer is shown in Figure 9-95. After clicking “Ok”, the variables “a”, “b”, and “c” will be available as fit parameters (Figure 9-96).

Figure 9-94. User Defined layer.
Virtual Substrate

Sample structure below the layer of interest may not be known or may be difficult to model. The Virtual Substrate models the underlying structure as a single “pseudo substrate” and simplifies the optical model of a multi-layer sample by allowing us to ignore the underlying sample structure and only consider the current thin film growth.
The “pseudo substrate” optical constants are obtained from a direct inversion of psi and delta using bulk substrate assumption. This approximation is nearly exact for materials with a high index of refraction and low index contrast (e.g. growth of semiconductors on semiconductors).

To set the time-slice for “pseudo substrate” optical constants calculation, ALT+Click on a time slice in dynamic data (use caution – the model will allow any time slice, even when VI approximation is not valid). Or set a range of times for averaging using the Start Time and End Time parameters. By default, CompleteEASE will ignore any time slices with out-of-spec Hardware parameters (i.e. insufficient intensity). Turn ON “Use All Times for VI” to override this function.

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**9.7. **+MODEL Options

The full list of +MODEL Options are shown in Figure 9-99. To view many of these, you first need to select Configure Options.

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![Diagram of Actual Sample and Simplified Optical Model with 'Virtual Interface' and 'pseudo substrate']
Include Substrate Backside Correction

Calculates the incoherent contribution of light reflecting from backside of transparent substrates in the model. When turned ON, the section will expand, as shown in Figure 9-100. Here, you can specify if the SE data were collected in transmission through the sample or through the reverse side of the substrate (opposite the coated side). There is also a selection for the number of backside reflections and the percentage of first reflection that is collected (from top side).

Model Calculation

Choose different model calculations to handle non-ideal sample situations. For most samples, the default setting of “Ideal” is preferred. However, there are many choices related to spectrometer bandwidth, film thickness nonuniformity, and beam angular spread, as shown in Figure 9-101, when these non-ideal situations are suspected. These options are best used in conjunction with Depolarization data fitting, as an ideal sample/measurement will not produce any depolarization.

When a “non-ideal” Model Calculation is selected, it will expand to allow fit parameters describing the amount of non-ideality, as shown in Figure 9-102. With advanced options selected, there is even the ability to describe two different bandwidths, with a selectable switching wavelength. This is useful for systems that have two separate spectrometers. Each non-ideality is modeled by convolving multiple data sets with different values. For example, model with thickness non-uniformity would be calculated by convolving the data from multiple different generated data sets with the model thickness varied over a range of values. The number of convolved data sets to approximate the “non-ideality” can be controlled, as seen in Figure 9-102, where the shape is set to Gaussian and “# of pts” is equal to 9.
Parameter Smearing

An advanced feature also allows other parameters to be “smeared.” This can include smearing of any of the fit parameters. When turned on, this section expands as shown in Figure 9-103, where the A parameter of Cauchy layer is smeared.

![Figure 9-103. Parameter Smearing expands to allow smearing of any fit parameter.]

Patterning

Patterning is another advanced feature that expands when turned on to calculate the model with only partial coverage of any designated layer. If all layers on the substrate are patterned, you can choose Layer # = 0 to pattern all of them, as shown in Figure 9-104. The %Patterned value lists the amount of film that is missing from the measurement area. Thus, 0% would be a fully intact film and 100% would be a non-existent film.

![Figure 9-104. Patterning expands to allow calculation with partial coverage of any single layer (or all layers).]

NOTE: Patterning may be usurped if “Multi-Model Patterning” is also turned on.

Ambient Index > 1

Use when working with liquid-cell where ambient is water or some other solution. When feature is turned On, the section expands as shown in Figure 9-105. Material can be selected to describe index of ambient material. If this material file is dispersion equation (such as Cauchy), the index of ambient can be fit and optical constants saved in this section.
**Ambient Index > 1 =** ON  
**Material: H2O Palm3**  
**Save Ambient Optical Constants**

Figure 9-105. Ambient Index > 1 expands to allow material file to describe ambient index.

**Wvl Shift (nm)**

Use to simulate a wavelength shift if the ellipsometer wavelength calibration is questioned.

**Return Path Ellipsometer**

Special configuration of M-2000 where source and detector are mounted side-by-side and the measurement beam reflects from the sample, then from a mirror and back off the sample surface on the way back to ellipsometer.

**Delta Offset**

Used to correct for windows. CompleteEASE accounts for Delta offsets from prisms (Cauchy) or windows (1/\(\lambda\)).

**Psi Offset**

The Psi offset can be used to model prisms in the beam path before and after the samples. J.A. Woollam company instruments that use prisms for beam steering already have a defined calibration method that accounts for offsets so this feature of the model is rarely used.

**Source and Receive Rot.**

Enter for rotation in degrees of either the source or receiver from the standard plane of incidence. This feature is typically used in special hardware configurations for in situ monitoring on chambers, where there is not room for the standard configuration.

**Multi-Model Patterning**

When Multi-Model Patterning is turned on, it overrides the simplified patterning feature discussed earlier in the Model Options section. Multi-Model patterning allows you to mix calculations from multiple model structures. All layers need to be added to the Model in the order they will appear from the different regions of the sample. For example, I have constructed a “layered structure”, shown in Figure 9-106, required for the hypothetical patterned sample shown in Figure 9-107.

<table>
<thead>
<tr>
<th>Layer # 4 = NTVE_JAW</th>
<th>Thickness # 4 = 2.00 nm (fit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Layer # 3 = a-Si parameterized</td>
<td>Thickness # 3 = 250.00 nm (fit)</td>
</tr>
<tr>
<td>Layer # 2 = Cr</td>
<td>Thickness # 2 = 10.00 nm (fit)</td>
</tr>
<tr>
<td>Layer # 1 = SiO2_JAW</td>
<td>Thickness # 1 = 1000.00 nm (fit)</td>
</tr>
<tr>
<td>Substrate = Si_JAW</td>
<td></td>
</tr>
</tbody>
</table>

Figure 9-106. Layered structure to be used in a Patterned model calculation.
To setup the multi-model pattern for this sample, we will need three different models – one for each of the areas that are different (within the measurement spot). The first area is just Cr film on Silicon (I assume 25% of the measurement area). The second area is Cr on SiO$_2$ on silicon (another 25% of area). Finally, we have the complete stack of surface oxide on a-Si on Cr on SiO$_2$ on silicon over the remaining 50%.

To build this using the Multi-Sample patterning, you first choose “# of Models” to be 3. Then, you can select the various areal percentages. Next, add a fit parameter for each of the layers that will vary from area to area – in our case, 3 of the 4 layers have less than 100% coverage. The Cr film is coating the entire surface and if we make the assumption that it is the same thickness in all areas, then we do not need to add this parameter to the list. However, it may be easier to add this parameter to the list, for visualization sake. The individual thicknesses in the different areas can be set, fixed, or even fit. The final model for this sample is shown in Figure 9-108.

Figure 9-107. Example of patterned sample, to be represented in CompleteEASE by a Multi-Model Patterning.

Figure 9-108. Model using Multi-Model patterning to represent a sample with 3 distinct areas.
Multi-Sample Analysis (MSA)

It is possible to fit multiple data sets simultaneously using the Multi-Sample Analysis (MSA). These multiple data sets can come from i) appended data files, ii) multiple selected points from uniformity map, iii) multiple rotations from automatic rotation scans, or iv) multiple time slice selections from dynamic data.

The key to each of these scenarios is that we wish to have some parameters consistent over all data sets and other parameters varied from data set to data set.

An example of multi-sample analysis is shown in Figure 9-109, where three oxides on silicon are fit with a common Cauchy layer to describe same index, but the thicknesses are allowed to differ for each data set.

Figure 9-109. Example of Multi-Sample Analysis, where all three data sets are fit simultaneously with common Cauchy optical constants and different film thickness.

It is possible to set different fit weightings for each data set active in the MSA. Right-click on the Data Set number and select Set Fit Weight. This will open an input box. By default all data sets are weighted equally (100%). You may wish to increase or decrease the fit weighting of a particular set or sets if for example, one set is significantly noisier than others. The Fit Weight can also be used to balance a fit result when a good fit can be achieve to one set but not others (be sure to consider sample structure and model first).
Parameter Coupling

Parameter coupling allows users to enforce relationships between different fit parameters. As an example in Figure 9-112 the Thickness of first data set in a multi-sample analysis is double the second Thickness in the same fit. This ratio will be enforced, and the thicknesses will adjust together during the fit.

The Coupling Mode can be a Ratio, Offset, or Custom Equation. The coupled parameters include all “fit” parameters and the final equation is shown in the dialog box so that you can verify it is correct.

Figure 9-112. Example of Parameter Coupling, where Thickness in a MSA is coupled by factor of 2X larger than another Thickness.
Use Previous Results

When CompleteEASE saves a data file, it also includes the results of any analysis that was performed (and saved). These results can be used for consecutive samples that may require information from the underlying structure.

For example, you may wish to map the uniformity of a layer on a substrate before it undergoes some processing. The fit results from this first map can be used during the analysis of the next measurement after processing. When working with multi-layers, you may wish to map the thickness of the first layer prior to depositing the second layer – and use these results when analyzing the second layer.

This feature requires that the same map locations are used for each run.

If using a B-Spline layer to fit for optical constants of first measurement, make sure that “ShowParms” parameter in the B-Spline layer is turned on. This will result in all of the B-Spline parameters being saved into the data file.

If a model is not used during data acquisition (i.e. as part of a recipe), then it will be necessary to use the fit log’s reanalysis feature to get the model results saved into the data file before this feature can be used.

Use Previous Results = ON
Results File: C:\CompleteEASE\DAT\205_backside_rough_SE

Figure 9-113. “Use Previous Results” expands to allow you to import results from a previous fit (inside the data file).

9.8. +FIT Options

The full list of +FIT Options are shown in Figure 9-114. To view many of these, you first need to select Configure Options.
- FIT Options
  - Include Initial Fit Stage = OFF
  + Perform Thickness Pre-Fit = ON
  - Use Global Fit = ON
    - # of Data Points = 20
      - # of Iterations = 5
      - Parm #1 = \( b \)
        - Min. = 1.300
        - Max. = 3.000
        - # Guesses = 30
      - Parm #2 = \( \text{(none)} \)
      - Parm #3 = \( \text{(none)} \)
      - Include Wvl. Range Expansion Fits = OFF
  - Selected Data = Standard Ellipsometric
    - Fit Weight = N.C.S
    - Include Depolarization Data = OFF
    - Include Intensity Data = OFF
    - Transmission Data % Weight = 0.50
    - Define MSE using \( L^2 \text{a}^2 \text{b}^2 \) = OFF
    - Limit Wvl. for Fit = OFF
    - Limit Angles for Fit = OFF
    - Max. Acceptable MSE = 100.000
    - Skip Data Points in Fit = 0
    - Max. Fit Iterations = 50
    - Auto Fit Parameter Reset = OFF
    - Include Derived Parameters = ON
    - Add Derived Parameter:
      - 1. Type = \( n \)
      - Layer # = 1
      - Wavelength = 632.8 nm
      - Name = \( n \) @ 632.8 nm
      - Hide = OFF
      - # of Decimal Places = 5
      - Low Spec. = 0.00
      - High Spec. = 0.00

Figure 9-114. List of all \textit{FIT Options}.

Include Initial Fit Stage

This feature allows a model to define a pre-fit where only a subset of the fit parameters is turned on before allowing all of the defined fit parameters to fit.

Perform Thickness Pre-Fit

When turned on, it uses Woollam’s patented algorithm to quickly estimate the thickness for the thickest layer in the model. This algorithm is based on calculation of zero-crossings when the data are oscillating due to the thickness.

Use Global Fit

Allows fit parameters to be varied over a range of values to find the best match to the data. The Global Fit is described in detail in the Example of Section 3.3.

Number of Data Points to Use

Used by Global Fit to improve speed. This refers to the number of wavelengths used during the Global search.

Number of Iterations

This is the number of fits performed at each global fit iteration in an effort to minimize the MSE from the various starting points.

Parm. #1, #2, #3

Up to three fit parameters can be included in the Global Fit.
**Min. / Max. and # Guesses**

Select the “search” range for each Fit Parameter that is included in the Global Search, along with the number of guesses between the minimum and maximum values.

**Include Wvl. Expansion Fits**

Decide whether the Global Fit should also use the Wvl Expansion Fit.

**Limit Wvl for Fit**

Select a limited spectral range during the fit to increase speed of the Global Fit. This feature only appears in conjunction with the wavelength expansion fit.

**Selected Data**

Choose which type of data to include in the Fit. Choices include Standard Ellipsometry (SE), generalized-SE, Mueller-matrix, and Intensity data only, as shown in Figure 9-115.

![Figure 9-115. Choose the selected data type to use during fit.](image)

**Fit Weight**

Select the fit weighting, with choices as shown in Figure 9-116.

![Figure 9-116. Select the Fit Weighting.](image)
Include Depolarization Data

Turn this setting on when you want the depolarization data to be included as part of the fit weighting. You can then select the percentage of weighting compared to other data curves – set to 100% for equal weighting, lower for less-importance, and higher for stronger importance.

Include Intensity Data

Choose whether intensity data should be included in the fit weighting, turn on this setting. You can then select the percentage of weighting compared to other data curves – set to 100% for equal weighting, lower for less-importance, and higher for stronger importance.

Transmission Data % Weight

When Transmission Intensity curves are appended to the data set, they are fit irregardless of the “Include Intensity Data” option above. This setting allows more or less importance to be placed on the Transmission Intensity data by increasing or reducing its Weight percentage, respectively.

Define MSE using L*a*b*

This feature defines the fit MSE by the difference between the model calculated L*a*b* color coordinates and target values set by the user.

Limit Wvl. for Fit

Reduce the wavelength used during fitting.

Limit Angles for Fit

Reduce the number of angles used during fitting. You can use comma-separated list to designate specific angles to include.

Max Acceptable MSE

Select this value to give a warning when fit exceeds this MSE.

Skip Data Points in the Fit

Reduce number of wavelength used during fitting by skipping this number of points.

Max Fit Iterations

Set the maximum number of fit iterations.

Auto Fit Parameter Reset

Will reset all fit parameters to their initial values before fitting new data. This can be very useful with uniformity maps to avoid the fit parameters getting “lost” and not able to recover after hitting a single “bad” data point.
Include Derived Parameters

To add a derived parameter to a model, open the Fit Options section and then turn the “Include Derived Parameters” flag to “ON”. This allows you to add multiple derived parameters. Figure 9-117 below shows how a derived parameter for displaying the index of refraction at 632nm would look.

![Figure 9-117. Derived Parameters in +FIT Options](image)

Clicking the mouse on the Type parameter allows you to change the derived parameter calculation. The available types are shown in Figure 9-118. Some of the types allow the specification of a wavelength and layer number for the parameter.

![Figure 9-118. Derived Parameter Types.](image)

After fitting data using a model with derived parameters, the fit results section shows the model fit parameters and the derived parameters. The derived parameters are shown in gray instead of black.
A new derived parameter type called “User Equation” has been added to CompleteEASE (new in version 3.38) so simple calculations can be performed after a fit has finished. If there is an error in the equation then the derived parameter will not be displayed in the **Fit** panel after a fit is completed. This derived parameter has 4 items that can be edited – a name, equation, and low and high specifications. Figure 9-120 shows how the derived parameter displays in a model.

To edit the equation press the **Edit Equation** item. This brings up the Edit Equation dialog as shown in Figure 9-121. In addition to specifying the equation this dialog allows the user to enter the number of decimal places that should be displayed.
The equation can contain numbers, operators, variables, and parentheses for grouping. The available operators are: ‘+’ (add), ‘-’ (subtract), ‘*’ (multiply), ‘/’ (divide), and ‘^’ (power). In addition there are a number of constants and functions that can be used. These are shown below.

**Functions**
- sin
- cos
- tan
- log
- abs
- sqrt

**Constants**
- PI
- e

Variables in an equation are fit parameter names in enclosing brackets. An example would be “[SiO2 Thickness]”. Any fit parameter name can be specified as well as any derived parameter names that appears in the list above the equation.

**Color Coordinates**
This derived parameter calculates the color coordinates based on the specular reflectivity of the final model. This was demonstrated by Johs et al. for specular coatings and rolled metal samples\(^6\). An example of color coordinates is shown in Figure 9-122, where it is listed as both a Derived Parameter in the **Model:** panel and the results are shown in the **Fit:** panel. When “Color Coordinates” are selected as a derived parameter, you have the additional choices of i) type of color coordinates, as shown in Figure 9-123, ii) angle of incidence for calculation of reflectivities, iii) choice of observer angle, as shown in Figure 9-124, and iv) choice of illumination light, as shown in Figure 9-125.

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Figure 9-122. Color Coordinates shown as a Derived Parameter, along with the fit results using $L^*a^*b^*$ type.

Figure 9-123. Choose type of Color Coordinates.

Figure 9-124. Choose type of observer.

Figure 9-125. Choose desired illuminant.
Sheet Resistance

This derived parameter, shown in Figure 9-126, calculates the sheet resistance of layers between the starting and ending layer numbers. It sums the resistivities as if they are parallel resistors, as shown in Figure 9-127. There are “Offset” and “Multiplicative Corrections” available for matching ellipsometry measured values to alternative techniques if needed (four point probe for example). The sheet resistance calculation rules are as follows:

- The calculation assumes all layers in the selected range “touch” the contacts. The user sets the selected range (start and end layer numbers) appropriately.
- By default all layers have an infinite resistivity.
- GenOsc layers with a Drude (RT) oscillator will use the Resistivity parameter for the calculation.
- GenOsc layers with a Drude (NMu) oscillator will calculate the resistivity from N and mu.
- Graded GenOsc layers using the parametric grade type will have the resistivity of the slices summed up as if they are individual layers.
- EMA layers and graded layers not using the parametric grade type are considered to have infinite resistivity even if they include a GenOsc layer with a Drude term.
- Internmix and Surface Roughness layers are ignored.

![Image: Derived Parameter showing “Sheet Resistance”](image)

![Image: Calculation of sheet resistance from multiple layers in a graded model](image)
9.9. **+OTHER Options**

There are other options available in the Model: panel, including all those shown in Figure 9-128.

**- OTHER Options**

- **Wvl. Range Expansion Fit**  Increment (eV) = 0.50
- **Try Alternate Models**
- **Fit Parameter Uniqueness**
- **Fit Parameter Error Estimation**
- **Simulate Data**
  - Add Opt. Const. to HTML Report = OFF
  - # of Processor Cores to use (available:8) = 0
  - Show Color Calculation Dialog = OFF
- **Configure Options**
- **Turn Off All Fit Parameters**

*Figure 9-128. List of all +OTHER Options.*

**Wvl. Range Expansion Fit**

Press this command to execute a Wavelength-Range Expansion Fit. The model will be fit over the “selected” wavelength range first. Then, the data range is increased toward shorter and/or longer wavelengths in increments. As each new range of data is added, the data are fit again. This method works well in combination with the B-Spline layer and is demonstrated in Example of Section 4.3

**Try Alternate Models**

The “Try Alternate Models” command automatically fits the data using the ideal layered optical model, and then with surface roughness and index grading non-idealities added to the model. The results of the various model fits are summarized in tabulated and graphical format, and the user can then decide which model is most appropriate for the sample. Example of report generated by this command is shown in Figure 9-129.
Fit Parameter Uniqueness

This command allows user to test the uniqueness of a fit parameter. A single fit parameter is selected to vary over a range of values. At each value, the fit parameter is fixed while all other fit parameters are varied to find the best fit (lowest MSE). The MSE is recorded to visualize the shape of MSE profile versus the varied fit parameter. An example MSE profile is shown in Figure 9-130.
Fit Parameter Error Estimation

This command tests the current model and data set for both random errors and systematic errors. When pressed, a dialog box will appear as shown in Figure 9-131.

![Fit Parameter Error Estimation dialog box.](image)

**Random Errors (Noise)**

The “true” fit parameter precision can be experimentally determined by calculating the standard deviation of the fit parameters obtained by multiple measurements on the same sample. However, multiple sample measurements can be very time consuming and impractical; it is desirable to estimate the parameter precision based on a single measurement. The Random Errors (Noise) section will test the current data set and model in three possible ways, as described below, to estimate the fit parameter precision. An example of these tests is shown in Figure 9-132 for measurement of a native oxide on silicon.

The Actual Standard Deviations are calculated by measuring the same sample 50 times and checking the statistics of the thickness “precision”. The Fit Stats are the 90% confidence error-bars that are calculated after fitting a single data set. The Error Bars MC and Bootstrap MC stand for the “Experimental Error Bars” and “Bootstrap” Monte-Carlo simulations, as described in the next sections. For this sample, these estimates all have similar magnitude, lending credibility to the value in estimating model precision based on a single measurement.

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Figure 9-132. Random Error tests for a native oxide on silicon, comparing results for all angles of incidence with results from various single-angle measurements.

Experimental Error Bars

The experimental data are fit multiple times, but are manipulated before each fit such that individual data from each wavelength are modified randomly based on their measured error bars. The percentage listed here is associated with the amount the experimental data are adjusted based on a percentage of the individual error bars.

Bootstrap Method

The Bootstrap method\textsuperscript{8,9} also fits the experimental data multiple times to check for “precision”. The sub-set of data points used by each trial-fit are selected at random from the total population of experimental data points. As the name suggests, each time an individual data point is selected, it is replaced back into the general population of experimental data points. This means that, during any single trial-fit, the trial-sample can contain one, two, or more copies of any given data point – or no copies at all, if it is never selected. The user determines the size of the trial-sample relative to the total available data points by entering a Refill%.

Parameter Perturbation

The experimental data are fit multiple times. However, before each trial-fit, the fit parameters in the model are perturbed by a percentage of their current value to check if the fit is robust enough to return to the same solution.

Systematic Errors

Random errors (precision) do not consider the effects of systematic errors on the model fit results. In many cases, the systematic errors can be more significant. Thus, CompleteEASE allows the accuracy to be estimated in the presence of different possible systematic errors. These errors may include measurement errors; such as angle offset, wavelength shift, or data offsets; or they may include modeling errors such as substrate optical constant inaccuracies. A new method is also proposed that adds the fit “errors” to the data to estimate the inaccuracy of the modeling.

The Systematic errors are either added to the experimental data sets (constant offsets in the ellipsometric data, and fit error magnitudes) or to the optical model (angle offset, wavelength shift, and offsets in substrate optical constants). The changes in the fit parameters in the presence of the assumed systematic errors are reported.

**Angle Offset**

Fit results are compared with and without an angle offset.

**Wavelength Shift**

Fit results are compared with and without a wavelength shift.

**Psi Offset**

Fit results from the standard data set are compared to a data set that is manipulated by applying a constant offset to the Psi values.

**Delta Offset**

Fit results from the standard data set are compared to a data set that is manipulated by applying a constant offset to the Delta values.

**Transmission Offset**

Fit results from the standard data set are compared to a data set that is manipulated by applying a constant offset to the Transmission Intensity values.

**Substrate ‘n’**

Standard fit is compared to a fit where the model substrate index is manipulated by applying offset.

**Substrate ‘k’**

Standard fit is compared to a fit where the model substrate extinction coefficient is manipulated by applying offset.

**Fit Errors Magnitude (+/-)**

After a fit, there will be differences between the Experimental Data curves and the Model-Generated Curves. These can be viewed as a “Difference Graph”. These represent systematic errors of an unknown kind (and combination) where the data are not perfectly matched by the model. Without knowing the type of systematic errors that produced these differences, CompleteEASE tries to estimate the effect of this error on the fit parameter accuracy by adding or subtracting the same “magnitude” of error at every wavelength to the existing data set and refitting the manipulated data.

For example, Figure 9-133 graphs the difference between Experimental Data and the Model-Generated curves for a thin organic film over a thick oxide coated silicon wafer. The Fit Error Magnitude takes the absolute value of this curve and then adds or subtracts this from the existing Experimental Data set (giving the +/- curves), as shown in Figure 9-134.

This test was applied to the P3HT film, which is anisotropic and absorbing in the visible spectral region. This makes it difficult to gain sensitivity to the extraordinary direction, which is out-of-plane (normal to sample surface). This is seen in the results of Figure 9-135 for the ordinary direction and Figure 9-136 for the extraordinary direction.
Figure 9-133. Difference Graph between Experimental Data and the Model Generated Curves for a P3HT film on SiO2-Si substrate.

Figure 9-134. Fit Error Magnitudes (+/-) calculated from the Difference Graph.
Figure 9-135. Optical constants for the P3HT layer ordinary direction (in-sample-plane) with Fit Error Magnitudes show the accuracy of the ordinary direction to be very good.

Figure 9-136. Optical constants and Fit Error Magnitudes for the extraordinary direction (out-of-plane, normal to surface) of the P3HT film show less sensitivity to the values in this direction.
**Simulate Data**

Experimental data can be simulated in CompleteEASE from the model using the Simulate Data feature. This feature generates ellipsometric data from the current model and uses it as though it is experimental data. To turn this feature on, click on **Configure Options** to open the Show Model Options dialog. Check “Simulate Data” (under Other Options) and click “Ok” to close the dialog.

![Show Model Options dialog](image)

*Figure 9-137. Show Model Options dialog.*

The model now has a Simulate Data item in the Other Options section.

- **OTHER Options**
  - *Wvl. Range Expansion Fit* Increment (eV) = 0.50
  - *Try Alternate Models*
  - *Fit Parameter Uniqueness*
  - *Fit Parameter Error Estimation*
  - *Simulate Data*
  - Add Opt. Const. to Report = **OFF**
  - **Configure Options**
  - **Turn Off All Fit Parameters**

![Simulate Data added under OTHER Options](image)

*Figure 9-138. Simulate Data added under OTHER Options.*

To use this feature, click on **Simulate Data** to open the Simulation Settings dialog. The settings dialog allows the user to specify the wavelength range and angle range for the simulated data as well as choose the data type to simulated. Click “Ok” to create the simulated data set.

![Simulation Settings](image)

*NOTE: To simulate transmission results, the Backside Correction needs to be turned to ON and “Transmission SE Data” also turned ON.*
Add Opt. Const. to Report
Choose this option to automatically add optical constants graphs to the report.

# of Processor Cores to Use
CompleteEASE can increase calculation speed by using multiple processor cores. By default, when set to “0” all core will be used.

Show Color Calculation Dialog
Press this button to show a dialog box representing the color coordinates of the current model, as shown in Figure 9-139.

![Color Calculation dialog box](image)

*Figure 9-139. Color Calculation dialog box shows the color coordinates and estimated graphical “color” based on current model, specified Angle of Incidence (AOI), observer angle and illuminant light.*

Configure Options
Choose this command to show “hidden” features in the +MODEL Options, +FIT Options, and +OTHER Options sections of the model. The choices are shown in Figure 9-140.
Figure 9-140. Configure which “hidden” options to show within the Model.

**Turn Off All Fit Parameters**

Turns off all active fit parameters in the model.
10. Gen-Osc Functions

This chapter describes the various oscillator functions available within the Gen-Osc layer. Figure 10-1 shows the Gen-Osc dialog box. The dialog box can be opened from any Gen-Osc layer by expanding the layer and clicking Show Dialog. The “type” of oscillator can be selected from a dialog box, as shown to the right.

![Figure 10-1. Left: Gen-Osc layer dialog box. Right: oscillator type input dialog box.](image)

10.1. Real-Part (e1)

There are four parameters at the top of the Gen-Osc Layer Parameters panel that are used to describe the real-part of the dielectric function. These include the Einf, UV Pole Amplitude, UV Pole Energy, and IR Pole Amplitude.

Einf

Einf, or “$\varepsilon_\infty$”, is a purely real constant added to $\varepsilon_1$. $\varepsilon_\infty$ is often referred to in the scientific literature. “$\text{Einf} =$ ” is located at the top of the Layer Parameters panel.
Pole

Poles are equivalent to Lorentz oscillators with zero-broadening. Their position should remain outside the “fit” spectral range. As such, they only affect the real part of the dielectric function ($\varepsilon_1$). They can describe dispersion created by absorptions that occur outside the measured spectral range. They are defined in the equation:

$$\varepsilon_{\text{pole}_n} = \frac{\text{Amp}_n}{\text{En}_n^2 - E^2}$$

Fit Parameters (units):
- $\text{En}_n$ (eV) & $\text{Amp}_n$ (eV$^2$)

10.2. Lorentz

This is a version of the classic Lorentz oscillator model, described as:

$$\varepsilon_{\text{Lorentz}} = \frac{\text{Amp}_n \cdot \text{Br}_n \cdot \text{En}_n}{\text{En}_n^2 - E^2 - i \cdot \text{EBR}_n}$$

Fit Parameters (units):
- $\text{En}_n$ (eV) & $\text{Br}_n$ (eV)
- $\text{Amp}_n$ (no units)

In this style of the Lorentz oscillator, $\text{Amp}_n$ approximately equals $\varepsilon_2$ at its peak value, and $\text{Br}_n$ is approximately the full width at half-maximum (FWHM).

10.3. Lorentz Im(Amp)

Adding an imaginary amplitude term to the Lorentz function allows $\varepsilon_2$ to be asymmetric around the center energy, $\text{En}$. Note however, that $\varepsilon_2$ will become negative when the imaginary term $i \cdot \text{Amp}$ is greater than zero. This means that another oscillator must be added to force $\varepsilon_2 \geq 0$ so that the total gen-osc function is Kramers-Kronig consistent.

$$\varepsilon_{\text{Lorentz,Im(Amp)}} = \frac{(\text{Amp}_n^2 + i \cdot \text{Amp}_n \cdot \text{Im(Amp)_n}) \cdot \text{Br}_n \cdot \text{En}_n}{\text{En}_n^2 - E^2 - i \cdot \text{EBR}_n}$$

Fit Parameters (units):
- $\text{En}_n$ (eV) & $\text{Br}_n$ (eV)
- $\text{Amp}_n$ & $i \cdot \text{Amp}_n$ (no units)

$\text{Amp}_n$ approximately equals the $\varepsilon_2$ at its peak value, and $\text{Br}_n$ is approximately the full width at half-maximum (FWHM).

10.4. Harmonic

The Harmonic oscillator function is similar to that derived from perturbation theory in quantum mechanical treatments of single-electron transitions, and is typically written in the literature as a two-term sum$^{10}$. The equation is:

$^{10}$ For example, see S. Adachi, *Optical properties of crystalline and amorphous semiconductors: materials and fundamental principles,* (Kluwer Academic Publishers, Norwell, MA, USA, 1999), p. 64.
\[ \varepsilon_{\text{Harmonic}} = \frac{\text{Amp}_n \cdot \text{Br}_n}{2} \left( \frac{1}{\text{En}_n - E - i \frac{\text{Br}_n}{2}} + \frac{1}{\text{En}_n + E - i \frac{\text{Br}_n}{2}} \right) \]

Amp\(n\) approximately equals \(\varepsilon_2\) at its peak value, and \(\text{Br}\) is approximately the full width at half maximum (FWHM). When the broadening becomes much less than the center energy (\(\text{Br}_n \ll \text{En}_n\)), the Harmonic oscillator is equivalent to the Lorentz function above.

### 10.5. Harmonic Im(Amp)

Adding an imaginary amplitude term to the Harmonic function above allows \(\varepsilon_2\) to be asymmetric around the center energy \(\text{En}\). Note however, that \(\varepsilon_2\) becomes negative when the imaginary term \(i\text{Amp}\) is greater than zero. This means that another oscillator must be added to force \(\varepsilon_2 > 0\) so that the total gen-osc function is Kronig consistent.

\[ \varepsilon_{\text{Harmonic,Im}} = \frac{[\text{Amp}_n + i(i\text{Amp}_n)] \cdot \text{Br}_n}{2} \left( \frac{1}{\text{En}_n - E - i \frac{\text{Br}_n}{2}} + \frac{1}{\text{En}_n + E - i \frac{\text{Br}_n}{2}} \right) \]

Amp\(n\) is approximately equals the \(\varepsilon_2\) (Im\{\(\varepsilon\}) peak value, and \(\text{Br}\) is approximately the full width at half maximum (FWHM).

### 10.6. Gaussian

This Oscillator type produces a Gaussian line shape in \(\varepsilon_2\) with a Kronig-Kronig consistent line shape for \(\varepsilon_1\).

\[ \varepsilon_{\text{Gaussian}} = \text{Amp}_n \left[ i \left( \Gamma \left( \frac{E - \text{En}_n}{\sigma_e} \right) + \Gamma \left( \frac{E + \text{En}_n}{\sigma_e} \right) \right) + \right] \]

\[ \sigma_e = \frac{\text{Br}_n}{2\sqrt{\ln(2)}} \]

where \(\text{Amp}\) is the amplitude, \(\text{En}\) is the center energy, and \(\text{Br}\) is the broadening. The function \(\Gamma\) is a convergence series that produces a Kronig-Kronig consistent line shape for \(\varepsilon_1\) (see footnotes \(^{11} \& ^{12}\) for details).

Amp\(n\) equals \(\varepsilon_2\) at its peak value (\(\sim \text{En}_n\)). By defining \(\sigma_n = \frac{\text{Br}_n}{2\sqrt{\ln(2)}}\), \(\text{Br}_n\) approximately equals the full width at half maximum value (FWHM).

### 10.7. Drude(RT) & Drude(NMu)

The classic Drude model describes free carrier effects on the dielectric response. Its form is equivalent to a Lorentz oscillator with center energy equal to zero. There are

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two versions of the Drude oscillator in the gen-osc layer: Drude(RT) and Drude(NMu).

Drude (RT) is described by the following equation\(^\text{13}\), and has resistivity, \(\rho\), and mean scattering time, \(\tau\), as fit parameters.

\[
\varepsilon_{\text{Drude,RT}}(E) = \frac{-\hbar^2}{\varepsilon_0 \rho E_n^2 + \hbar E} \quad \text{for} \quad n = \text{oscillator #}
\]

where \(\rho = \frac{m^* m_e}{Nq^2 \tau} = \frac{1}{q \mu N}\)

The physical constants are \(h\) (Planck’s constant)/\(\pi\), \(\varepsilon_o\) (the vacuum dielectric constant), \(m_e\) (the electron rest mass), and \(q\) (the electron charge).

The fit parameters for Drude(NMu) are \(N\) (the carrier concentration in \(\text{cm}^{-3}\)), \(\mu\) ("\(\mu\)" (the carrier mobility in \(\text{cm}^2\text{V}^{-1}\text{s}^{-1}\)) and \(m^*\) (\(m\star\), the carrier effective mass).

Note that \(\mu\) and \(m\star\) are completely correlated, so the user will have to choose which one to fit, and fix the other one.

\[
\varepsilon_{\text{Drude,RT}}(E) = \frac{-\hbar^2 q^2 N \mu m\star}{\varepsilon_0 (\mu \mstar m_e E_n^2 + i \hbar E)} \quad \text{for} \quad n = \text{oscillator #}
\]

\[
\text{Fit parameters (units):} \quad N \quad (\text{\text{cm}}^{-3}), \quad \mu \quad (\text{\text{cm}}^2\text{V}^{-1}\text{s}^{-1}), \quad m\star \quad (\text{no units})
\]

10.8. Tauc-Lorentz & Cody-Lorentz

The Tauc-Lorentz (Jellison and Modine\(^\text{14}\)) and Cody-Lorentz (Ferlauto, et al\(^\text{15}\)) dispersion functions both model the main absorption of amorphous materials using a broad Lorentzian line shape with zero absorption below a defined bandgap energy. The primary difference between the two functions is how they model absorption at energies slightly larger than the gap energy. In this region the Tauc-Lorentz model follows the Tauc law formula while the Cody-Lorentz follows the Cody formula:

\[
\text{Tauc Absorption Formula:} \quad \varepsilon_2(E) \propto (E - E_g)^2/E^2
\]

\[
\text{Cody Absorption Formula:} \quad \varepsilon_4(E) \propto (E - E_g)^2
\]

The Cody-Lorentz oscillator also includes an Urbach absorption term to model absorption at energies below \(E_g\).

\text{Tauc-Lorentz}

The equation for the Tauc-Lorentz oscillator function is given below. The bandgap energies of any Tauc-Lorentz oscillator can be tied to \(E_g\) of the first Tauc-Lorentz in the gen-osc layer by selecting \text{Common } E_g = ON.

---


\[ e_{r,n}(E) = e_{o,n} + i e_{s,n} \]

where

\[ e_{s,n} = \begin{cases} \frac{A\cdot E_0\cdot E}{E_0^2 + E^2} \left( 1 - \frac{1}{E^2} \right) & \text{if } E > E_{g,n} \\ \frac{A\cdot E_0\cdot E_0}{E_0^2 + E^2} & \text{if } E \leq E_{g,n} \end{cases} \]

\[ e_{s,n} = 0 \]

The Kramers-Kronig integral form is presented here for the sake of brevity. In CompleteEASE (as in Ferlauto\(^*\)), this Kramers-Kronig integration is solved analytically.

**Cody-Lorentz**

The Cody-Lorentz oscillator is similar to the Tauc-Lorentz in that it defines a bandgap energy \( E_g \), and a Lorentzian absorption peak (parameters \( Amp_n \), \( E_0n \), and \( Br_n \)). However, for the Cody-Lorentz \( e_{s,n}(E) \propto (E - E_g)^2 \) in the region just above \( E_g \). The Cody-Lorentz also defines two transition energies: \( Ep \) and \( Et \). \( Ep \) is where absorption transitions from Lorentzian to Cody behavior, and \( Et \) is where the absorption transitions from Cody to Urbach behavior.

\[ e_{s,c-l} = e_{o,c} + i e_{s,c} \]

where

\[ e_{s,c}(E) = \begin{cases} \frac{E}{E} \exp\left( \frac{E - E_{s,c} - E_{o,c}}{E_{o,c}} \right) & \text{if } 0 < E \leq \left( E_{s,c} + E_{o,c} \right) \\ \frac{A\cdot E_0\cdot E_0}{E_0^2 + E^2} \left( E - E_{s,c} \right)^2 + E_{o,c}^2 & \text{if } \left( E_{s,c} + E_{o,c} \right) < E \leq \left( E_{s,c} + E_{o,c} \right) \end{cases} \]

\[ e_{s,c}(E) = \frac{A\cdot E_0\cdot E_0}{E_0^2 + E^2} \left( E - E_{s,c} \right)^2 + E_{o,c}^2 \left( E - E_{s,c} \right)^2 + E_{o,c}^2 \]

\[ E_{s,c} = \left( E_{s,c} + E_{o,c} \right) \times G(E_{s,c} + E_{o,c}) \times L(E_{s,c} + E_{o,c}) \]

G(E)=near-bandgap function, L(E)=Lorentz oscillator function

\[ e_{s,c} = 2 \frac{A}{\pi} \int_0^{1/n} \frac{\tilde{G}(E)}{\tilde{E}^2 - \tilde{E}^2} d\tilde{E} \]

\( * \) The Kramers-Kronig integral form is presented here for the sake of brevity. In CompleteEASE (as in Ferlauto\(^*\)), this Kramers-Kronig integration is solved analytically.
Figure 10-2. Parameter definitions for the Cody-Lorentz oscillator in Gen-Osc.

In the equation, \( G(E) \) defines the Cody absorption behavior, while \( L(E) \) is a Lorentz oscillator. \( E_g + E_p \) is the energy where the function transitions from a Cody absorption \( \varepsilon_2(A) \propto (E - E_g)^2 \) to a Lorentzian absorption. Note that \( G(E) \to (E - E_g)^2/Ep^2 \) when \( E \to E_g^+ \), and \( G(E) \to 1 \) when \( E \gg E_p \).

\( (E_g + E_t) \) marks transition between Urbach absorption and Cody absorption. It is important to note that there are differences between the Ferlauto et al \(^{15}\) and CompletEASE definitions. Ferlauto et al defines \( E_t \) as an absolute energy, while CompletEASE defines it as a difference:

\[
E_t^{\text{Ferlauto}} = E_g^{\text{CompletEASE}} + E_t^{\text{CompletEASE}}
\]

The Urbach absorption is fully Kramer-Kronig consistent; that is, the exponential Urbach absorption in \( \varepsilon_2(E) \) has a Kramers-Kronig transformed counterpart in \( \varepsilon_1(E) \) function. The internal parameter \( E_1 \) guarantees that the \( \varepsilon_2 \) function transitions smoothly at the \( (E_g + E_t) \) transition point. \( E_u \) defines the exponential rate; specifically \( (E_g + E_t) \) is the energy at which the Urbach absorption equals \( e^{-1} \) of its maximum value. The ratio \( E_1/E_g \) guarantees that the Urbach exponential function exactly matches \( G(E)L(E) \) at \( (E_g + E_t) \).

10.9. Tanguy

Tanguy\(^{16,17}\) developed this expression as an analytical expression of Wannier excitons, including bound and unbound states. The function makes no distinction between bound and unbound states. As shown below, the function depends on five parameters: amplitude \( \text{Amp}_n \), fundamental bound state energy \( E_{g_n} \), unscreened binding energy \( R_n \), broadening \( B_{n} \), and \( \ln g \) \( (= \ln(g)) \). The parameter \( g \) is a screening factor that can vary between 0 to \( \infty \). \( g \to \infty \) is an unscreened Coulomb potential, and \( g \to 0 \) is total screening.

10.10. Psemi-M0 & Psemi-Tri

The Psemi-M0 and Psemi-Tri functions are a subset of the more general Herzinger-Johns Parameterized Semiconductor Oscillator function originally introduced in the WVASE32® psemi.mat layer\(^{18}\). The Psemi oscillator combines a highly flexible functional shape with Kramers-Kronig consistent properties.

The \( \varepsilon_2 \) part of the Psemi function consists of four polynomials splines connected end-to-end. Each spline connects smoothly with the adjacent spline, forming a single, continuous function, as shown in Figure 10-3.

Figure 10-3. Two views of Psemi-Tri oscillator. Top: \( \varepsilon_2 \) showing the Amp, Ec, endpoints and control points. Bottom: Psemi-Tri for 3 different Br values.

and Figure 10-4. The function’s shape is controlled by adjusting the center point, Ec, the end points WL and WR, and the mid-width control points, AL and AR. Thus

\[
\varepsilon_{\text{Psemi},n} = \frac{\text{Amp}_n \sqrt{R_n}}{(E + iBr_n)} \left\{ \tilde{g}(\xi(E + iBr_n)) + \tilde{g}(\xi(-E - iBr_n)) - 2\tilde{g}(\xi(0)) \right\} \quad n = \text{oscillator #}
\]

\[
\xi(z) = \frac{2}{\sqrt{E_n z_c}} \left( z - z_c \right)
\]

\[
\tilde{g}(\xi) = -2\nu\left( \frac{e^{\frac{\xi}{\xi_c}}}{\xi_c} \right) - \frac{\xi}{\xi_c} - 2\nu(1 - \xi) - 1/z
\]

\[
\nu(z) = \frac{d[\ln\Gamma(z)]}{dz} \quad \text{(digamma function)}
\]

Fit parameters (units):
- \( \text{Amp}_n \) (no units),
- \( E_n \) (eV),
- \( Br_n \) (eV),
- \( R_n \) (eV)
- \( \ln g_n = \ln(g_n) \) (dimensionless)

The \( \text{psemi.mat} \) layer is described in detail in the WVASE32® software manual.
the overall shape is controlled by the mid-points and location of the endpoints with respect to Ec.

The function is forced to equal zero near the endpoints, and a gaussian broadening parameter Br, rounds of the endpoints and peak (n = oscillator #)

**Fit parameters (units):**
- Ampn (no units)
- Brn (eV)
- Ecn (eV)
- WLn (eV)
- WRn (eV)
- ALn (no units)
- ARn (no units)

![Graph](image1)

![Graph](image2)

*Figure 10-3. Two views of Psemi-Tri oscillator. Top: $\varepsilon_2$ showing the Amp, Ec, endpoints and control-points. Bottom: Psemi-Tri for 3 different Br values.*

and Figure 10-4). The flexibility of these functions can be seen in the figure, as combinations of left and right control point amplitudes can produce a wide variety of asymmetrical $\varepsilon_2$ absorptions.
The Psemi-Tri oscillator shown in

\[ n \equiv \text{oscillator #} \]

Fit parameters (units):
- \( \text{Amp}_n \) (no units),
- \( Br_n \) (eV),
- \( Ec_n \) (eV),
- \( WL_n \) (eV),
- \( WR_n \) (eV),
- \( AL_n \) (no units),
- \( AR_n \) (no units)

Figure 10-3. Two views of Psemi-Tri oscillator. Top: \( \varepsilon_2 \) showing the Amp, Ec, endpoints and control-points. Bottom: Psemi-Tri for 3 different \( Br \) values.

is a general-use Psemi function. The fit parameters are: \( \text{Amp}, Ec, A, B, WL, WR, AL, \) and \( AR \).

\[ n \equiv \text{oscillator #} \]

Fit parameters (units):
- \( \text{Amp}_n \) (no units),
- \( Br_n \) (eV),
- \( Ec_n \) (eV),
- \( WL_n \) (eV),
- \( WR_n \) (eV),
- \( AL_n \) (no units),
- \( AR_n \) (no units)

Figure 10-3. Two views of Psemi-Tri oscillator. Top: \( \varepsilon_2 \) showing the Amp, Ec, endpoints and control-points. Bottom: Psemi-Tri for 3 different \( Br \) values.
The Psemi-M0 oscillator models the shape of an $M_0$ critical point seen in direct gap semiconductors such as GaAs and InP, in the bandgap energy region. Variable fit parameters are: $\text{Amp}_n$, $\text{Br}_n$, $\text{En}_n$, $\text{WR}_n$, $\text{PR}_n$, $\text{AR}_n$, and $\text{O2R}_n$. Figure 10-4 shows the effects on the $\varepsilon_2$ line-shape for three different combinations of $\text{AR}$ and $\text{PR}$ values.

![Figure 10-4. $\varepsilon_2$ plot showing the effects on the line-shape of the Psemi-M0 oscillator for different AR and PR values.](image)

10.11. CPPB (Critical Point Parabolic Band)

The CPPB (Critical Point Parabolic Band oscillator) was developed by Aspnes\textsuperscript{19} to model the shape of the dielectric functions at semiconductor critical points (CP’s). The shape of each CPPB oscillator depends on five parameters: the amplitude $\text{Amp}_n$, phase projection factor $\text{Phase}_n$, threshold energy $\text{En}_n$, broadening parameter $\text{Br}_n$, and exponent $\text{Mu}_n$. $\text{Mu}_n$ can have any one of three values:

- $\mu_n = +\frac{1}{2}$ for 1-D CP’s,
- $\mu_n = "0"$ (becomes $\ln(2\text{En}_n - 2E - i\Gamma)$) for 2-D CP’s, and
- $\mu_n = -\frac{1}{2}$ for 3-D CP’s.

\[
\varepsilon_{n,\text{CPPB}} = \text{Amp}_n e^{i\text{Phase}_n} \left( \frac{\text{Br}_n}{2\text{En}_n - 2E - i\text{Br}_n} \right)^{\text{Mu}_n}, \quad \text{Mu}_n = \pm 1/2
\]

\[
\varepsilon_{n,\text{CPPB}} = \text{Amp}_n e^{i\text{Phase}_n} \ln(2\text{En}_n - 2E - i\text{Br}_n), \quad \text{Mu}_n = 0
\]

Fit parameters:
- $\text{Amp}_n$ (no units),
- $\text{Br}_n$ (eV),
- $\text{En}_n$ (eV),
- $\text{Phase}_n$ (no units),
- $\text{Mu}_n = (\pm 0.5 \text{ or } 0)$ (no units)

### 10.12. Custom

The Custom oscillator provides allows the user to define a layer’s complex dielectric or refractive index function.

**Edit Equation dialog box**

Selecting **Edit Equation** (located under Type = Custom in the oscillator list) opens the Edit Equation dialog box. The box is shown in Figure 10-5. The user will need to select the following:

- **Optical Constant type**: $e_1 & e_2$ or $n & k$.
- **Wavelength [wvl]** in: Å, nm, μm, eV, or 1/cm.

The user then types the Real and Imaginary parts of the desired function into the appropriate boxes. Variables must be enclosed in brackets.

As an example we will recreate the Lorentz style “.1” oscillator from the WVASE32® genosc layer,

\[
\varepsilon_{\text{Lor}_\text{.1}} = \frac{\text{Amp} \cdot E_0}{E_0^2 - E^2 - iE \cdot B_r} = \frac{\text{Amp} \cdot E_0}{(E_0^2 - E^2)^2 + E^2 \cdot B_r^2} + i\cdot\frac{\text{Amp} \cdot E_0 \cdot E \cdot B_r}{(E_0^2 - E^2)^2 + E^2 \cdot B_r^2}.
\]

In this version of the Lorentz oscillator, $\text{Amp}$ is approximately proportional to the area under the $\varepsilon_2$ curve. That is, when $E_0 >> B_r$,

\[
\text{Amp} \approx \frac{2}{\pi} \int_{0}^{\infty} (\varepsilon_{\text{Lor}_{\text{.1}}}) dE
\]

Figure 10-5 shows the Re{$\varepsilon$} and Im{$\varepsilon$} parts for $\varepsilon_{\text{Lor}_{\text{.1}}}$ entered into the Custom Oscillator Edit equation dialog box.

After clicking “Ok”, new variables will be accessible as fit parameters in the model.
Operator List

A number of built-in operators and functions are available for use in the Custom Oscillator equation. They are listed in Table 1-2.

Table 10-1. List of built-in operators and functions available in the Custom Oscillator.

<table>
<thead>
<tr>
<th>OPERATOR/FUNCTION NAME</th>
<th>DESCRIPTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>+</td>
<td>Addition</td>
</tr>
<tr>
<td>-</td>
<td>Subtraction</td>
</tr>
<tr>
<td>*</td>
<td>Multiplication</td>
</tr>
<tr>
<td>/</td>
<td>Division</td>
</tr>
<tr>
<td>^</td>
<td>Power</td>
</tr>
<tr>
<td>sqrt(arg)</td>
<td>Square root(argument)</td>
</tr>
<tr>
<td>exp(arg)</td>
<td>Exponent(argument)</td>
</tr>
<tr>
<td>sin(arg)</td>
<td>Sine function(argument)</td>
</tr>
<tr>
<td>cos(arg)</td>
<td>Cosine function(argument)</td>
</tr>
<tr>
<td>tan(arg)</td>
<td>Tangent function(argument)</td>
</tr>
<tr>
<td>ln(arg)</td>
<td>Natural logarithm(argument)</td>
</tr>
<tr>
<td>log(arg)</td>
<td>Base 10 logarithm(argument)</td>
</tr>
<tr>
<td>real(arg)</td>
<td>Real part(argument)</td>
</tr>
<tr>
<td>imag(arg)</td>
<td>Imaginary part(argument)</td>
</tr>
<tr>
<td>parm(arg)</td>
<td>Argument can be any hardware parameter that is recorded during measurement. Two examples: parm(Temperature), parm(AlignX)</td>
</tr>
<tr>
<td>isPositive(test_val,eq_pos,eq_neg)</td>
<td>isPositive(test_val,eq_positive,eq_negative)</td>
</tr>
<tr>
<td>PI</td>
<td>π (uppercase “PI” required)</td>
</tr>
<tr>
<td>e</td>
<td>Natural number (lowercase “e” required)</td>
</tr>
<tr>
<td>h</td>
<td>Plank’s constant 6.626069E-34 Joules·sec (lowercase “h” required)</td>
</tr>
<tr>
<td>hbar</td>
<td>( h = h/2 \pi ) reduced Plank’s constant. 1.0545717E-34 Joules·sec (lowercase “hbar” required)</td>
</tr>
</tbody>
</table>
The next example is a modification of the Lorentz oscillator used in the previous example (see Figure 10-5). For this equation, the Amp parameter is replaced with $(2 \cdot \text{Area}/\pi)$. Now $\text{Area} \approx \text{area}(\varepsilon_2)$ when $E_0 >> B_r$.

Figure 10-6 shows the Re{$\varepsilon$} and Im{$\varepsilon$} parts in the Edit Equation dialog box. Note that the “PI” function is used in the equations.

![Figure 10-6. Custom Oscillator Edit Equation dialog box for $\varepsilon_{\text{Lor, area}}$ equation. Note the use of the “PI” function.](image)

**Edit Marker Parameters**

Selecting Edit Marker Parameters (located under Type = Custom in the oscillator list) opens the Edit Marker Parameters dialog box (Figure 10-7). For each custom oscillator function, the user can select three markers to represent user-defined variables.

![Figure 10-7. Custom Oscillator Edit Marker Parameters dialog box for $\varepsilon_{\text{Lor, area}}$ equation.](image)

Once selected, the user-defined variables can be varied by simply dragging the marker boxes around the Graph (Figure 10-8) using a mouse.

![Figure 10-8. Graph panel with Markers identified.](image)
11. Reference – In situ Tab

The In situ tab is used for real-time dynamic ellipsometry measurements where the sample properties might change with time. It provides a number of useful options for real-time dynamic data acquisition and display. The in situ tab may be required for M-2000 and RC2 systems mounted onto a chamber, or for accessories (i.e. liquid cells, heat stages, etc.) mounted onto M-2000, RC2, and alpha-SE systems.

11.1. Displaying the In situ tab (if not visible)

The In situ tab should be visible among the other tabs. If it is not visible, follow this procedure:

a. Select the Options>‘Edit Configuration’ button (Figure 11-1 Left). The CompleteEASE Configuration dialog box will open (Figure 11-1 Right).

b. Select General>‘Always Show In situ Tab = true’ option (Figure 11-1 Right).

c. Select ‘Yes’ when the program asks, “Do you want to apply the changes?”

Figure 11-1. Left: Options tab with ‘Edit Configuration’ button indicated. Right: CompleteEASE Configuration window with ‘Always Show In situ Tab = true’ option selected.

NOTE: To display the In situ tab during a single session of CompleteEASE, press CTRL-ALT-I.
11.2. *In situ* Tab features

The CompleteEASE *In situ* tab is shown in figure Figure 11-2. This tab consists of a Status window, Data Acquisition & Alignment section, the Models panel, the View panel and Graph.

![Figure 11-2](image.png)

*Figure 11-2. Left: The CompleteEASE In situ tab. Right: Data Acquisition & Alignment section prior to start of data acquisition.*

**Status window**

The Status window (upper left corner of Figure 11-2) provides information about the current status of the ellipsometer, such as “Initializing Hardware”, “Waiting to Acquire Data” etc.

**Data Acquisition & Alignment Section**

This section is shown in Figure 11-3, immediately below Status window. The user can start, stop, pause and trigger data acquisition, perform a hardware alignment and System Check, and display the raw signal from the spectrometer. The user can also choose the Fast Acquisition Mode.

![Figure 11-3](image.png)

*Figure 11-3. Data Acquisition & Alignment section prior to the start of data acquisition.*
**Start (End) Acquisition**

Starts the data acquisition process (Figure 11-3 and Figure 11-4). After the acquisition is started, the ‘Start Acquisition’ button changes to ‘End Acquisition’ which ends the data acquisition and saves the data to disk.

**NOTE:** The settings for Standard and Fast acquisition modes can be accessed through the Hardware Configuration dialog box, within the **Hardware** tab.

**Pause (Resume) Acquisition**

This pauses or resumes the data acquisition when Standard acquisition mode is selected (Figure 11-4 Left).

**Trigger Acquisition**

When the acquisition mode is set to Trigger mode (via Hardware Configuration dialog box), a ‘Trigger’ button will also appear (Figure 11-4 Right). Selecting the button will trigger a single data acquisition event.

**Align Hardware**

Selecting ‘Align Hardware’ button allows the user to align the ellipsometer Source and Detector units or sample stage.

**System Check**

Starts the System Check Operation.

**Display Signal**

This will display the raw intensity signal output by the ellipsometer’s CCD array spectrometer.

**Fast Acquisition Mode**

Checking this box changes the acquisition mode from Standard to Fast. The Standard mode uses a longer acquisition time with High Accuracy set to ON.

**NOTE:** With High Accuracy On, the measurement zone-averages the polarizer, which takes over twice as much time per measurement, but significantly improves accuracy as first order misalignments can be corrected.

Fast mode uses a shorter acquisition time with High Accuracy set to OFF. The exact settings for the Standard and Fast acquisition modes can be accessed through the Hardware Configuration dialog box.
**Models Panel**

The **Models** Panel, shown in Figure 11-5, contains the models that can be used for real-time analysis. Up to forty models can be added to the list (10 models per page times 4 pages). Pressing one of the model buttons before or during data acquisition causes that model to be loaded, and any data subsequently acquired will be fit to that model.

**NOTE:** In order for a model to be accessible in the model panel list, the model must be saved in the Common folder.

![Figure 11-5. The Models panel in the In situ tab.](image)

**Adding/Editing/Clearing a Models button**

To add, edit or delete a **Models** button, right-click (\&R) on the button. The Models Setup dialog box will appear, as shown in Figure 11-6.

![Figure 11-6. Left: Models Setup dialog box. Right: Model File list (only model files that have been saved in the Common folder will appear in this list).](image)

- **Selecting a Model file**
  Select the desired model from the Model file list. *Only model files that have been saved in the Common folder will appear in this list.*

- **Clearing a Model button**
  Select “(none)” if you wish to clear a button of any model.

- **Parameter Corrections**
  With a model selected, its fit parameters will appear in the parameter list. Highlighting any fit parameter in the list allows the user to include Offset, Multiply and AlignX Corrections to that fit parameter.
• **Do Thickness control**
The Thickness control check box activates CompleteEASE to send a signal to another program or device to stop growth or etch when a target thickness is reached.

• **Write Report**
Selecting the Write Report check box to activate a report writing function.

• **Merge**
The ‘Merge’ check box allows the user to merge the selected model with the current model in the Analysis tab. For example, suppose the current model consists of a Thermal oxide on silicon (Figure 11-7). Note that the Layer #2 thickness equals 568.15 nm, and is an active fit parameter in the model.

```plaintext
| Layer # 2 = SiO2_JAW  | Thickness # 2 = 568.15 nm (fit) |
| Layer # 1 = INTR_JAW  | Thickness # 1 = 1.00 nm          |
| Substrate = SI_JAW    |
```

*Figure 11-7. Initial Thermal oxide on silicon. Note that the thickness of layer #2 is 568.15nm and is an active fit parameter.*

We then add a ‘Metal on thermal oxide on silicon’ model as a button in the Models panel button list. This model has the same layers as the first model, but with an additional layer #3 B-Spline layer to represent the metal.

```plaintext
+ Layer # 3 = B-Spline  | Thickness # 3 = 0.00 nm (fit) |
Layer # 2 = SiO2_JAW  | Thickness # 2 = 568.15 nm      |
Layer # 1 = INTR_JAW  | Thickness # 1 = 1.00 nm        |
Substrate = SI_JAW    |
```

*Figure 11-8. Metal on thermal oxide on silicon model, which has been merged with the Thermal oxide on silicon model in Figure 11-7 above. Note that layer #2 is still 568.15 nm thick, but is no longer a fit parameter.*

**View Panel**
The View panel, as shown in Figure 11-9, is where the information that appears in the bottom graph is selected. Selecting ‘Parameters’ displays the parameters graph, selecting ‘Log’ displays the run information log, and selecting ‘Fit’ displays a spectroscopic view of the most recent time slice including the data fit.

*Figure 11-9. The View panel.*

**Parameters**
The “Parameters” radial button graphs the value of the fit parameters and hardware alignment information during data acquisition. Figure 11-10 shows the parameter graph during an in situ practice measurement of the 250Å SiO₂ Si calibration wafer. It is displaying the Thickness, MSE and Intensity versus time. The parameters that the graph shows are selected from the four drop-down list boxes above the graph. Parameters listed in the two left boxes are shown on the left y-axis and parameters listed in the two right boxes are displayed on the right y-axis. Besides the Fit parameters and MSE, the hardware parameters SigInt, Intensity, AlignX, AlignY, Hardware OK and Fit OK can also be displayed.
Log

The “log” radial button records information during data acquisition. Recorded information includes: the time when models are loaded or unloaded, and when the data acquisition is stopped (Figure 11-11).

The user can also record information in the log. To add a comment, type it into the textbox at the top (green arrow, Figure 11-11) and press the ‘Add’ button (red arrow, Figure 11-11). The comment is stamped with the time that the add button is pressed. Comments can only be added to the log during data acquisition. When data is saved, the parameters graph and log data are saved with it, so all can be retrieved by opening the experimental data file.

Fit

“Fit” radial button displays ellipsometric data vs. wavelength for the last recorded time slice, as well as data generated from the fit model (if present). This is shown in Figure 11-12 for a selected time-slice near 0.58 minutes.

Dynamic Data

The “Dynamic Data” displays one or more measured data values vs. time (Psi, Delta, N, C, S, Intensity, etc.). The Graph Type menu allows user to select which measured values are displayed. This is shown in Figure 11-13 for a section of dynamic data.
Figure 11-12. The Fit data screen.

Figure 11-13. The Dynamic Data screen.
12. Reference – Hardware Tab

The controls available in the Hardware tab are different for the different ellipsometer configurations. As a result, this chapter is divided into two major sections: one section describing the alpha-SE Hardware tab and another section describing the M-2000 and RC2 Hardware tab.

NOTE: Full details of the Hardware tab can be found in the Hardware manual for each individual ellipsometer. This Chapter is meant only as a general reference.

12.1. Configuration Files

The CompleteEASE software determines the hardware configuration based on the “cnf” files that are located in the C:\CompleteEASE\cnf folder. These files may include the following:

**hardware.cnf**
Describe the hardware configuration and options available with the current system.

**CompleteEASE.cnf**
Configuration file for user settable software parameters. These parameters should be accessed by choosing ‘Edit Configuration’ button from the Options tab.

**CompleteEASEhard.cnf**
Configuration file for user settable hardware parameters. These parameters should be accessed by choosing ‘Edit Hardware Config.’ from the Hardware tab.

**bview.cnf**
Used for in situ systems only

**Privileges.cnf**
User privileges when setup for password-protected User Log-in.
12.2. Hardware Tab – alpha-SE

The Hardware tab for an alpha-SE is shown in Figure 12-1. This tab consists of four panels: Hardware, Signal, Calibration, and Show Logs. A brief description of all commands within this tab is described.

![Hardware tab](image)

**Figure 12-1. CompleteEASE software with the Hardware tab selected.**

### Hardware

**Re-initialize**

This button re-initializes the ellipsometer hardware, which initiates communication over the USB link, scans the spectrometer, moves the motors to home position, and tests the motors. Details of any errors that occur during hardware initialization can be found in the “Hardware” and/or “Error” logs.

**Park Z-Stage**

Moves the Z-stage to the “park” position such that the Z-stage shipping lock can be engaged.

### Signal

**Display**

Displays signal intensity of the light collected by the spectrometer. Average, maximum, UV, and IR intensity values are also reported in the status box, along with the current spectrometer “Dark Count” and electronics “Temperature”. An example is shown in Figure 12-2.

**Max. Sig. Display**

Same as Display, except that the polarizer and compensator optics are adjusted to find the maximum signal intensity. Window will appear as shown in Figure 12-2, after the maximum signal is found.
Z-Stage Scan

Scans the Z-stage to generate intensity vs. sample height profile. A sample should be mounted and the ellipsometer unit set to the “Off-Sample” configuration before clicking this button. The resulting signal intensity profile should appear symmetrical, similar to the graph shown in Figure 12-3. The black dashed curve is a polynomial fit to the measured profile, and the reported “MSE=” value quantifies the symmetry of the profile. If the MSE value is greater than 0.02, contact your J.A. Woollam Co. representative.

Figure 12-2. CompleteEASE “Signal Display” screen.

Figure 12-3. Results of a typical Z-Stage scan.
Calibration

**S-T Baseline**

Acquires baseline spectra for the “Straight-Through” configuration such that accurate transmission intensity and Mueller matrix data can be acquired. For best data accuracy, always perform a S-T Baseline after moving the ellipsometer optics, or if it has been a long time (> 1 hour) since the previous baseline.

**Off-Sample Baseline**

Acquires an “Off-Sample” baseline such that qualitative reflection intensity data can be acquired. This option also calibrates and stores the angle of incidence which is used for subsequent measurements. For best angle of incidence accuracy, perform an Off-Sample Baseline after moving the source or receiver units, or after mounting the sample chuck.

**Angle Offsets**

Calculates the angle offsets. This is usually performed as part of a calibration procedure.

**Show Logs**

**Hardware and Error Logs**

These buttons display the Hardware and Error logs, which contain information useful for diagnosing and debugging problems with the instrument. If the instrument is not working properly, your J.A. Woollam Co. representative may require this information to diagnose the problem: click the “Save Debug File” button that appears in the log file, and email C:\CompleteEASE\CompleteEASE_Debug.zip file. An example Hardware Log is shown in Figure 12-4 and Error Log is shown in Figure 12-5.

![Figure 12-4. Hardware Log.](image-url)
NOTE: The old ErrorLog.txt files are archived in the "\CompleteEASE\cnf\Archive" folder. They are zipped to save space.

NOTE: When CompleteEASE starts, it checks the size of the ErrorLog.txt file. If greater than 5MB, then it is saved to the Archive folder and a new file is created.

12.3. Hardware Tab – M-2000 and RC2

Figure 12-6 shows the Hardware Tab for an M-2000 System. This tab consists of three panels: Hardware Status, System Information, and Controls. This section describes the panels and the controls found on them.

NOTE: Full details of the Hardware tab can be found in the Hardware manual for each individual ellipsometer. This Chapter is meant only as a general reference.

Figure 12-6. M-2000 and RC2 Hardware Tab.

Hardware Status

The Hardware Status panel displays the current state of the hardware. This is the same information displayed in the Hardware Status panel of the Measurement tab.

Controls

The Controls panel provides access to hardware functions other than those needed for acquiring data. The hardware functions are separated into groups displayed as tabs. The tabs are General, Calibration, System, and Misc.
General

The General tab shown in Figure 12-7 is nearly the same for all of the different ellipsometer configurations. The available controls are as follows:

![General Tab](image)

**Align Sample**

Align the sample without acquiring data. For systems with automated alignment capabilities a dialog is displayed allowing the user to choose between manual alignment and automated alignment (Figure 12-8). Choosing No in the dialog displays the manual alignment dialog. An Alignment screen with 4-quadrant detector position and z-intensity is shown in Figure 12-9.

![Sample Alignment](image)

**Figure 12-7. General Tab.**

**Figure 12-8.** The Sample Alignment dialog gives the user a choice of automated alignment or manual alignment.
Figure 12-9. Example of Alignment screen, showing a cross-hair to represent the position of beam reaching a 4-quadrant alignment detector and blue “signal” across the bottom tracking the signal that reaches the spectrometer.

**Display Signal**

This function displays the raw signal intensity. This screen, as shown in Figure 12-10, is useful for debugging light intensity problems. The Average AC Intensity includes all channels of the CCD spectrometer. The DUV AC Intensity is an average of the first 3 channels, while the UV AC Intensity is an average of the first 10% of CCD channels. The IR AC Intensity is an average of the last 10% of CCD channels.

Figure 12-10. Display Signal Screen.
System Check

A System Check is the most common procedure for calibrating the ellipsometer. See the hardware manual for details on the System Check procedure.

Move To Load Pos.

This function configures the system to make loading a sample easier. This applies only to systems with automated z-stages and translators. Generally, pressing this button will result in the z-stage moving to its maximum value and the translator moving towards the user.

Routine Test Measurement

This feature adds the ability to periodically measure a standard sample and then plot the fit parameters and hardware parameters versus date. Using this feature requires that a recipe be created and placed in the “\CompleteEASE\Recipe\Routine Test Measurement” directory and the recipe name needs to be listed in the Routine Test Measurements section of the Hardware Configuration, as shown in Figure 12-11. To make this feature useful, users will need to be trained to always use the same sample and to put the sample in the same position on the stage each time they perform the measurement.

![Figure 12-11. Routine Test Measurements section of the Hardware Configurations.](image)

Measure

Press to perform the “Routine Test”.

Show Results

Shows all past results of the “Routine Test”. This is a convenient way to track the performance of a system, with results on the same sample. The values that can be graphed include any fit parameters for the designated test sample (thickness, refractive index, etc.) and many hardware parameters, such as Intensity and Alignment values.
Calibration

The Calibration tab appears different depending on the type of ellipsometer. Figure 12-12 shows the Calibration tab for an M-2000 and Figure 12-13 shows the tab for an RC2. It is not common for a user to need the controls in this tab as a System Check will perform many of these functions.

![Calibration tab for M-2000 ellipsometers.](image1)

**Initialize Hardware**

This button re-initializes the ellipsometer hardware, which initiates communication over the USB/TCPIP link, scans the spectrometer and moves the motors to home position. Details of any errors that occur during hardware initialization can be found in the “Error” log.

**Coarse Calibrate and Fine Calibrate (M-2000 Only)**

These calibrations can be performed in place of a System Check when it is not possible to use the standard calibration wafer that is supplied with the instrument. This situation occurs during in situ measurements on a deposition chamber that requires a specific sample size. These calibrations will not determine the in-plane window effects. Always perform a System Check instead of a Coarse or Fine calibration when possible.

**DC Offset**

Measures the ambient light and electronic noise contribution to the measured raw intensity. This button is most often used before acquiring depolarization data to improve measurement accuracy, as the depolarization data is determined from the DC component of the measured signal. A System Check automatically performs the DC Offset as well.
**Phase Sensor Info (M-2000 Only)**

This diagnostic function displays information used to determine if the phase sensor is working and the compensator motor is spinning. When troubleshooting a hardware problem a J.A. Woollam Company representative may ask for this information.

**Standard Calibration (RC2 Only)**

A Standard Calibration is equivalent to a Coarse Calibration on the M-2000. The System Check is the preferred method for calibrating the instrument when the standard calibration wafer can be used.

**System Calibration**

See the hardware manual for information on the System Calibration.

**Light Intensity (RC2 Only)**

The Light Intensity box allows the user to change the current light intensity. Normally this is left in automatic mode.

**System Status (RC2 Only)**

The System Status button displays a diagnostics dialog with information on the phase sensors and USB communication. When troubleshooting a hardware problem a J.A. Woollam Company representative may ask for this information.

**System**

The controls in the **System** tab are dependent on the type of base the ellipsometer is attached to. The main types of bases are the ESM (horizontal sample mount, automated z-stage and angle), ES130 (horizontal sample mount, manual z-stage and angle), Fixed Angle Focused (horizontal sample mount, automated z-Stage), Vertical (vertical sample mount, automated angle), and Accumap. Some of the bases have multiple attachments which can also change the controls available on the System tab. Figure 12-14 through Figure 12-16 show the System tab for some of these bases. Brief descriptions of the different buttons on this panel can be found below. See the hardware manual for a detailed description of the use of these functions.

![Figure 12-14. System tab for ESM, ES130, and Fixed Angle bases.](image-url)
Angle, Z-Stage, Translator, Sample Tilt

These buttons allow the user to change the positions of the motorized base components. For example, to change the position of the Z-Stage, press the Z-Stage button and enter a position. The stage will then move to the specified position.

Set Angle (Vertical base)

The vertical sample-mount base does not have a home sensor for the angle so the Set Angle function allows the user to set the current angle.

Install New Sample Stage (ESM, ES130 and Fixed Angle)

This function steps the user through the process of installing a sample stage attachment. These attachments included items such as a Translator, Heat Cell, Liquid Cell, etc. On bases with automatic detection capabilities the software will automatically detect which stage is being installed. The other bases will display a dialog with the different stage options and the user will select the one being installed.
**Install/Remove Focus Probes (ESM, ES130 and Fixed Angle Focused)**

Steps the user through the focus probe installation/removal procedure.

**Configure System Components (Vertical base)**

This button displays a dialog that allows the user to specify the attachments that are currently installed. These attachments can include a translator, focus probes, camera, rotator, etc.

![Installed Components dialog](image)

*Figure 12-18. The Installed Components dialog for vertical bases.*

**Change Lamp (AccuMap)**

Steps the user through the lamp change procedure. See the hardware manual for a description of this procedure.

**Get Lamp Hours (AccuMap)**

Displays the number of hours the lamp has been on since it was installed.

**Misc.**

The controls in the Misc. tab cover functions that did not fit into the General, Calibration and System tabs and are dependent on the type of base the ellipsometer is attached to and which attachments are installed. Some systems will not have a Misc. tab.

![Misc. tab](image)

*Figure 12-19. Misc. tab for ESM bases.*
Z-Stage Scan (Systems with motorized z-stages)

This function scans the z-stage over a specified range and graphs the signal intensity versus z-stage position. This is especially useful when working with transparent substrates to determine if the backside beam is being collected.

Display Sample Image (Systems with USB camera installed)

Displays the camera image of the sample in a dialog. The dialog can stay open while other operations are performed.

Additional Camera Functions

Additional camera functions can include ‘Mark Measurement Beam Location’ and ‘Measure Camera Field of View’, as shown in Figure 12-21.

Mark Meas. Spot Location (Systems with USB camera installed)

This function is used to mark the position of the ellipsometer probe beam on the camera image when the sample is aligned. After performing this function the “Show Beam” check box in the camera dialogs will provide an accurate representation of where the probe beam is intersecting the sample.

Measure Camera Field of View (Systems with USB camera installed)

This function is used to determine the field of view of the digital camera and calibrate the automated movement of the translators with mouse-click on an image.

Disable MMC (AccuMap)

Pressing this button causes a user initiated fault on the translator. See the hardware manual for details on when this function may be used.
**Reset MMC (AccuMap)**

Pressing this button attempts to reset all translator faults. After pressing this button the translator is active and will move when commands are sent provided the door and all safety catches are engaged.

**Show Routine Meas. (AccuMap)**

Displays a dialog showing the results of routine measurements acquired on the Accumap. Routine measurements must be configured in the hardware configuration dialog.

Additional Hardware Functions

May include other Hardware Functions, specialized for the individual system. For example, an automated angle base may have an ‘Angle Correction’ button as shown in Figure 12-22.

**Angle Offset Correction**

Measure calibration sample over a range of angles on automated angle of incidence systems and then apply a correction to the mechanical positions to produce more accurate angle of incidence for future measurements.

**System Information**

The System Information panel displays details about the ellipsometer configuration and provides access to the hardware configuration dialog and hardware log. The information provided includes the type of ellipsometer, wavelength range, and current motor positions. Please have this information available when contacting the J.A. Woollam Company for assistance.

**‘Edit Hardware Config’**

Pressing this button will bring up the Hardware Configuration dialog, as shown in Figure 12-23. This dialog contains many settings for configuring the behaviour of the hardware. The dialog displays a description of each parameter’s function so only the “categories” will be discussed here.
‘View Hardware Log’

This button displays the Hardware log which contains information on calibrations and initializations.

**Data Acquisition**

Default parameters for data acquisition, including options as shown in Figure 12-24.
Normal Calibration

Configuration parameters describing the Normal Calibration, as shown in Figure 12-25.

Coarse Calibration

Configuration parameters describing the Coarse Calibration, as shown in Figure 12-25.
System Calibration

Configuration parameters relating to the System Calibration, as shown in Figure 12-26.

![System Calibration](image)

Routine Test Measurements

Configuration parameters relating to the Routine Test Measurements, as shown in Figure 12-26.

Window Effects

Specifies settings for Window effects, as shown in Figure 12-27.

![Window Effects](image)

Out-of-Plane Effects

The out-of-plane effects are measured when performing a calibration such as a System Check if Window Effects are enabled.

WinRet1 and WinRet2 are the out-of-plane window retardance for the entrance and exit windows respectively.
WinDisp1 and WinDisp2 are the dispersion terms for the out-of-plane window retardance of the entrance window and WinDisp2-1 and WinDisp2-2 are the dispersion terms for the exit window.

RotDisp1 and RotDisp2 are terms for the rotary power of the entrance window and RotDisp1-2 and RotDisp2-2 are the terms for the rotary power of the exit window.

**NOTE:** The effects of the out-of-plane window retardance on ellipsometric data are canceled out (at least to the 1st order) when performing a zone averaged (high accuracy) measurement.

### In-Plane-Effects

The in-plane effects are measured during a System Check or a Delta Offset Calibration if Window Effects are enabled.

Delta Offset Mode sets whether the effects are describing prism dispersion or window (1/λ) dispersion.

Delta Offset Err. Tol specifies the maximum error allowed on a delta offset parameter. If the error bar is above this tolerance then the calibration is considered to have failed.

Min Delta Offset1 Limit specifies the smallest Delta Offset 1 parameter before the higher order terms will be set to 0 (turned off).

# of Higher Order Terms specifies the number of Delta Offset terms to fit for beyond Delta Offset1 during a System Check or Delta Offset calibration.

The Delta Offset 1 – 4 parameters are the terms for describing the in-plane window retardance. The in-plane window retardance as a function of wavelength for a prism (Delta Offset Mode=1) is:

\[
\text{retardance}(\lambda \text{ in } \angstrom) = \Delta_{\text{Offset}1} + \Delta_{\text{Offset}2}w^2 + \Delta_{\text{Offset}3}w^4 + \Delta_{\text{Offset}4}/w^2
\]

where \( w = 5000/\lambda \)

The in-plane window retardance as a function of wavelength for a window (Delta Offset Mode=2) is:

\[
\text{retardance}(\lambda \text{ in } \angstrom) = w(\Delta_{\text{Offset}1} + \Delta_{\text{Offset}2}w^2 + \Delta_{\text{Offset}3}w^4 + \Delta_{\text{Offset}4}/w^2)
\]

where \( w = 5000/\lambda \)

### Psi Offsets

The Psi Offset parameters can be used to correct the Psi data when using prisms to bend the beam. It is only necessary to use this feature when the ellipsometer beam enters the prism(s) at non-normal angles. It is up to the user to determine the correct Psi offset values – they are not determined during calibration.

\[
\text{Psi\_correction}(\lambda \text{ in } \angstrom) = \psi_{\text{offset}0} + \psi_{\text{offset}1}w^2 + \psi_{\text{offset}2}w^4 + \psi_{\text{offset}3}w^6 + \psi_{\text{offset}4}w^8
\]

where \( w = (10000/\lambda) \times 0.2 \)

### System Check Settings

Specifies settings for System Check parameters.
**System Check Settings**
- Model = 25nm Oxide on Si
- Acq. Time = 4
- Angle = 70
- Measure Delta Offsets = True

*Figure 12-28. System Check settings.*

**Parm. Return Values**
Parameters for setting up how hardware and analysis parameter values are returned through the “get()” functions, as shown in Figure 12-29

*Figure 12-29. Parameter Return Values.*

**Channel Subsets**
Specifies subsets of the channels for data acquisition, as shown in Figure 12-30.
Miscellaneous

Contains miscellaneous hardware parameters that do not fit in other categories, as shown in Figure 12-30.

Alignment Parameters

Parameters to describe sample alignment, example shown in Figure 12-30.

Include Alignment Parameters With Data

CompleteEASE can be configured to save alignment information along with the measured ellipsometry data. This can be especially useful in conjunction with mapping measurements. To turn this option on, open the Hardware Configuration dialog, check “Show Advanced Config. Options”, and open the Alignment Parameters section. Select the parameter “Include Alignment Parameters with Data” and turn this parameter ON. Close the dialog and choose Yes when asked if you want to apply the changes. The alignment information will now be added to the data files.
To view the alignment parameters:

1. Select **Measurement** > **Fit Results** > ‘View Prev. Results’ button, which opens the View Data dialog box.
2. If necessary, ‘Choose’ the correct Location (upper left) where the file(s) of interest are located.
3. Select one or more files in the Files panel.

A table Acquisition Parameters will appear in the File Information panel. See figure below. If a model is present, the table will also list active model parameters. Select ‘Graph Data’ button (lower left in dialog box) in order to graph various parameters. You can also copy the table to Clipboard as text or HTML.

**NOTE:** If you choose more than one file you will also need to select ‘Calc Multi-Sample Stats’ to see parameters from more than one data file in the table and graphs.

**Pattern Rec. Parameters**

Describes settings for the Pattern Recognition features, as shown in Figure 12-33.
Camera Settings

Describes settings for a camera when included with system, as shown in Figure 12-33.

Simple Ex Situ Measu. Modes

Details for ex situ measurement modes, as shown in Figure 12-33.

Liquid Cell Attachment

Describes hardware and windows related to a Liquid Cell, as shown in Figure 12-33.

Heat Cell Attachment

Describes hardware and windows related to a Heat Cell, as shown in Figure 12-34.
Figure 12-34. Heat Cell and Transmission Attachment sections.

Transmission Attachment

Describes transmission attachment, as shown in Figure 12-34.

Focus Option

Describes hardware and windows related to a Focusing Probes, as shown in Figure 12-35.

Figure 12-35. Focusing Probe Options.
13. Reference – Options Tab

The Options tab is shown in Figure 13-1. This tab consists of four panels: Display Units, Configuration Controls, Miscellaneous, and About CompleteEASE. A brief description of all commands within this tab is described.

Figure 13-1. CompleteEASE software with the Options tab selected.

13.1. Display Units

Sets the “Wavelength Units,” “Thickness Units”, and “Optical Constants” settings display units.

Wavelength Units

Choose from Å, nm, μm, eV, and 1/cm for the wavelength units.

Wavelength is the distance between adjacent peaks of an electromagnetic wave, commonly given in Angstroms (Å), nanometers (nm), or microns (μm). Wavenumber refers to the number of waves in a length of one centimeter, with units of reciprocal centimeters (1/cm).

If light is treated as a quantum of radiant energy (photon), its energy is given in electron volts (eV) which is proportional to frequency. Choice of units depends largely on the spectral range and application. The following relationship shows the conversion between wavelength and photon energy.

$$\lambda_{nm} = \frac{1240}{E_{eV}}$$ (13-1)


**Thickness Units**

Choose from Å, nm, μm, and mm for the thickness units.

**Optical Constants**

Display and report optical constants as either complex refractive index \((n \& k)\) or complex dielectric function \((e_1 \& e_2\) , which stand for \(\varepsilon_1\) and \(\varepsilon_2\)). The two descriptions for optical constants are related by the following:

\[
\tilde{\varepsilon} = \tilde{n}^2
\]  

\((13-2)\)

**NOTE:** All settings are saved when the program is exited, and restored when the program is restarted.

**‘Use Defaults’**

Return all settings to their default values, which include wavelength in nm, thickness in nm, and optical constants as “n & k”.

### 13.2. **Miscellaneous**

The **Miscellaneous** panel contains four buttons, as shown in Figure 13-2.

![Miscellaneous panel from Options tab](image)

**Figure 13-2.** **Miscellaneous** panel from **Options** tab.

**Reset Win. Size**

Return the CompleteEASE window to the default size.

**Manage Users**

Press ‘Manage Users’ to setup the user list, along with access level and password, as shown in Figure 13-3. The User privileges can be defined for Engineer level, by further pressing ‘Define User Privileges’, which opens the window shown in Figure 13-4. The User customization is further defined in the CompleteEASE Configuration (next section).
Figure 13-3. Manage Users window, where multiple users, passwords and privileges can be established.

Figure 13-4. Define User Privileges for Engineer level.

Show Manual (F1)

Press to show a PDF of the CompleteEASE manual, in an integrated PDF viewer. Example is shown in Figure 13-5. This manual can also be viewed by pressing F1 key.

NOTE: The PDF manual file is located in C:/CompleteEASE folder on your computer. It can be opened with Adobe PDF viewer for more accessible viewing.
Figure 13-5. Press ‘Show Manual’ to view a PDF copy of the manual.

**View CompleteEASE Shortcuts**

Press this button to view a list of CompleteEASE shortcut keys, as shown in Figure 13-6.
13.3. Configuration Controls

‘Edit Configuration’

The ‘Edit Configuration’ button allows customization of the CompleteEASE software configuration. Press this button to open the window shown in Figure 13-7. Each item to the left can be expanded by pressing the circle to the left of the name. Also, there are many hidden options, which will not appear until you check the box at bottom of the screen: “Show Advanced Config. Options”. The list with all Advanced options grows to that shown in Figure 13-8.

![List of CompleteEASE shortcuts](image)
Most of the settings within the CompleteEASE Configuration are self-explanatory and describe the default settings for the software. If you press an item, it will give a small description to the right of the window. Each item is listed in this chapter, with a few items of particular interest discussed in detail.

**General Configuration Parameters**

The general configuration parameters are shown in Figure 13-9. A description of each option appears when you press the individual line. User Log-In Mode and Users Have Own Folders are two options that are used when managing multiple users.
Figure 13-9. General Configuration Parameters.

**Font Name**

Name of CompleteEASE font family. Changes are made from drop-down box selection, as shown in Figure 13-10.

![Figure 13-10. Choose Font Name from drop-down box.](image)

**Font Size**

Point size of CompleteEASE Font.

**Graph Font Size**

Point size of font used for text in graphs.

**Look & Feel**

Specifies the look & feel of the user interface elements, with choices from a drop-down list, as shown in Figure 13-11.
System Time Mode

 Specifies the method used by CompleteEASE to determine time, with the following choices:
  0. Java time call
  1. Windows GetTickCount
  2. Windows timeGetTime
  3. Windows Performance Counter
  4. CPU Counter

User Log-In Mode

 Specifies how users will log into and access software functions. This feature is important when setting up multiple users, as described in section 0.

Auto Fit Enabled

 Specifies if “Auto-Fit” is enabled when CompleteEASE is started. This is related to the “Auto Fit” check-box from the Analysis>Fit: panel.

Always Add CE Folders

 Indicates that folders under the base folder (usually C:\CompleteEASE) should always be added to the users location list when CompleteEASE is started. This feature is important when setting up multiple users, as described in section 0.

User Have Own Folders List

 When User Log-in is required CompleteEASE will keep each users location list separately. This feature is important when setting up multiple users, as described in section 0.

Clear Data on Logout

 When User Log-In is required CompleteEASE will clear the data, model, and selected recipe when a user logs out. This is important to prevent proprietary results from remaining within CompleteEASE after a user has logged out.

Always Show In situ Tab

 When this option is checked the insitu tab will always be displayed. When this item is not checked the insitu tab will only be displayed when the installed sample stage supports insitu measurements (Heat Cell, Liquid Cell,…).
NOTE: If the In situ Tab is not shown all the time and user desires it on case-by-case basis, just press CTRL-ALT-I to access the tab.

**Use Unicode**

Specifies the character encoding for saving configuration files and models. Must be set on ‘ON’ to use extended (non-Latin) character sets; setting to ‘Off’ improves backwards compatibility with earlier CompleteEASE versions.

**Force Read of Tabulated Mat Files**

Indicates that all tabulated material files should be read from the disk instead of from the model file. This should *almost* always be set to FALSE.

**# of Cores for Data Analysis**

Specifies the # of processor cores for data analysis: ‘0’ means all the available cores are used. This value can be over-ridden by the model options.

**Memory Management Configuration Parameters**

The memory is determined by the maximum number of wavelengths and maximum time slices allowed with in-situ data. See Figure 13-12.

![CompleteEASE Configuration](image)

*Figure 13-12. Memory Management and Display Units section of the Configuration Parameters window.*

**Max Wavelengths**

Controls the amount of memory required by CompleteEASE by setting a maximum number of wavelengths.

**Max Time Slices**

Controls the amount of memory required by CompleteEASE by setting a maximum number of time slices for dynamic data.

**Display Units Configuration Parameters**

The Display Units section sets the default units for CompleteEASE. See Figure 13-12.
**Light Units**
Units to display light wavelength or photon energy; with choices of Å, nm, μm, 1/cm, or eV.

**Time Units**
Units for time; with choices of seconds, minutes, and hours

**Rate Units**
Units for rate of dynamic growth/etch; with choices of Å/s, Å/min, Å/hr, nm/min, nm/hour, μm/min, and μm/hour.

**Thickness Units**
Units for thickness; with choices of Å, nm, μm, and mm.

**Opt. Const. Units**
Units for Optical Constants; with choices of “n&k” or “e1&e2”.

**Temperature Units**
Units for temperature; with choices of °C or K.

**Translator Units**
Units for translator coordinates; with choices of cm or mm.

**% Range Calculation**
This allows the choice of calculation for % Range in the map statistics tables.

**File Save Prompt**
Automatically prompt to save data after each measurement.

**Remote Communications Configuration Parameters**
The Remote Communications section helps establish communication for CompleteEASE. See Figure 13-13 for the list of items in this section.

![Figure 13-13. Remote Communications section of the Configuration Parameters window.](image)
**Communication Type**

Method of communication between a remote computer and the CompleteEASE program. Choosing None indicates no communication will take place. Other options include TCP/IP and RS232.

**TCP/IP Port #**

Port number that the CompleteEASE server listens on.

**RS232 Baud Rate**

Baud rate for serial communications.

**Serial Port**

Serial port to use with serial communications.

**Auto Report**

Turning Auto Report on will result in a string containing the values of current model fit parameters to be sent out on the Asynchronous TCP/IP port after data is acquired and fit. This is only done if the communication type is set to TCP/IP.

**Auto Report Format**

Specifies the format of data reported using Auto Report. Status Text will return the parameters as listed in the Simple Mode status box. All Parms will return all of the parameters including those that are hidden.

**Allow Parameter Query**

When “true” an additional checkbox appears in a fit parameters dialog. When this parameter is checked the model will try to ask via TCP/IP (remote computer must be connected to port “TCP/IP Port #” plus 2) for the value of the parameter before the fit.

**Save Results to Database**

When true the fit results from a recipe measurement will be added to a database. The URL for the database should be set in a file named DB-URL.cnf in the cnf folder.

**RunRecipe – Out of Spec**

Specifies that when a RunRecipe command is received, the return string should include :OOS for each parameter that is out of spec or :EEL for each parameter whose error bar exceeds the limit as defined in the model.

**RunRecipe – Time Out (seconds)**

This is the maximum amount of time to wait for a recipe to complete before giving up and returning an error message to the remote computer. The purpose of this feature is to allow CompleteEASE to inform the remote computer that a problem has occurred instead of having the remote computer wait indefinitely. The default is 180 seconds so if the user’s longest recipe longer than 3 minutes then this value should be changed to make sure that a false error message is not sent. For example, if the users longest recipe takes 3 minutes to complete then the timeout can be set to 240 seconds.
Window Setup Configuration Parameters

The Window Setup section sets the default window shape, size and location. See Figure 13-14.

![Figure 13-14. Window Setup Configuration Parameters.]

Title Text

The text that appears in the title bar of the CompleteEASE program.

Title Bar Color

The color of the title bar when the program is active; if specified, must be 'blue', 'green', 'red', 'orange', 'yellow', 'purple', or 3 integer RGB values separated by spaces (e.g., '160 250 240' is cyan).

Start Up Tab

The tab that is selected by default when CompleteEASE starts. If users are required to log in then this mode is the default mode when technician level is selected.

Width

Window width in pixels.

Height

Window height in pixels.

Top Left Corner X

X position of top left corner of window in pixels.

Top Left Corner Y

Y position of top left corner of window in pixels.

Divider Position

Position of graph divider in pixels.

Graph Clipboard Configuration Parameters

The Graph ClipboardParms section sets the default graph dimensions and resolution when copying graphs to the clipboard. See Figure 13-15.
**Width in Pixels**

Width of Graph on the Clipboard, in pixels.

**AspectRatio**

Ratio of Graph Width/Height when copied to the Clipboard.

**Resolution Factor**

Oversampling factor for the Graph. This value is multiplied by the graph width and aspect ratio to determine the total number of pixels stored in the image.

**Auto Legend Position**

Automatically place the legend so that it minimizes data coverage when copying a graph to the clipboard.

**RTF Left/Right Margin**

The left and right page margins to use when creating RTF content for the clipboard.

**RTF Top/Bottom Margin**

The top and bottom page margins to use when creating RTF content for the clipboard.

### Hardware Simulation Mode Configuration Parameters

The Hardware Simulation Mode section sets whether hardware can be simulated when not attached. See Figure 13-16.

![Hardware Simulation Mode Configuration Parameters](image)

*Figure 13-16. Hardware Simulation Mode and In Situ Tab Configuration Parameters.*
**Simulation Enabled**
When checked, CompleteEASE uses simulation mode instead of the physical ellipsometer hardware to acquire data.

**Simulation Data File**
Specifies the CompleteEASE data file to automatically load if Simulation is enabled.

**In Situ Tab Configuration Parameters**
The In Situ Tab Parameters section sets details regarding the In Situ Tab, which is generally hidden on systems that are not designated for in situ. See Figure 13-16.

**Show Error Bars in Parameter Graph**
When this setting is turned on, the parameter graph will display 2 curves instead of 4, with the error bars for curves 1 and 2 displayed on the right Y axis.

**System Check Button on In Situ panel**
When set to true, a System Check button will appear on the In Situ tab. When set to false, a Check Substrate button will appear.

**# of Model pages – Insitu**
Number of model pages available in the In Situ tab. Each page holds 10 models. There is a minimum of 4 pages and a maximum of 8.

**Rep. Converter Name**
If the name of a program is listed here, the program will be run after a model report (generated by the simple insitu mode) has been written to disk. The program will be given the report file name as a command line parameter. The program needs to be located in the CompleteEASE directory.

**Show Received Cmds**
Displays the text of Received Commands in the In Situ tab status box.

**Allow Second Model**
When this setting is on, the user can assign a second model to a button that will be fit to the data just as the first, but no report will be generated. The 2nd model’s parameters will show up in the parameter graph provided the fit parameters are named differently than the first model fit parameters.

**Single Model Mode**
Specifies which model button will be activated when CompleteEASE is started. Setting to ‘0’ disables this mode.

**Show Fast Acq. Box**
Includes the “Fast Acquisition” check-box in the In Situ tab.
Data Export Parameters

The Data Export Parameters relate to custom applications that send results out to external programs upon completion of analysis within CompleteEASE. The various setting parameters are listed in Figure 13-17.

Custom Export Application

This entry specifies the name of a custom user-supplied application that will be run after each measurement to export the results to an external program or database. The “Export Meas. Results” option should be checked, and the application must be located in the CompleteEASE directory.

Append Timestamp to Data File

When checked, the current time will be appended to the name of data files acquired through a recipe measurement.

Export Recipe Data as Text

Performing a recipe measurement when this parameter is set to true will result in the software saving an additional file with the data recorded in text format.

Include Hardware in 1Pt Export Format

The new export format only applies when exporting results from single point recipe measurements (no maps). When this parameter is set to True, the hardware related parameters will be added to the exported text file. When this parameter is false (default), only the fit parameters are added.

Shutter Controls Configuration Parameters

The Shutter Controls Parameters section sets details regarding in situ control of shutters for real-time process experiments. See Figure 13-17.

Treat as Signal

Treat the shutters as signals instead of shutters. The difference is that in process control conditions, a signal remains off until the target thickness is reached whereas the shutter opens (turns on) when the growth starts and closes (turns off) when the target thickness is reached.

# of Shutters

The number of shutters to be configured in the software. The maximum number is 4.
Import and Save Configuration

To copy the configuration from one computer to the next, you can ‘Save Configuration’ and then ‘Import Configuration’. This will help to setup the CompleteEASE software in the same way on multiple computers.

13.4. About CompleteEASE

Displays the “About” box shown in Figure 13-18, which contains the software version number, memory availability, and copyright information.

Create Debug File

Click this button to create a debug file for sending in details to Woollam Company. The name is: C:\CompleteEASE\CompleteEASE_Debug.zip. Email this file to your J.A. Woollam Co. representative to aid in debugging problems with your ellipsometer system.

View Error Log

View the error log.

Figure 13-18. About Box.
14. Appendices

14.1. Troubleshooting Guide

Please contact your J.A. Woollam Co. representative with any questions relating to CompleteEASE or your ellipsometer system. To reduce the support time, please create and email the debug file described in the following section before contacting your representative.

**NOTE:** Your closest Woollam Representative can be found in Section 0.

**Reporting Problems to the J. A. Woollam Co.**

In the event that you have a problem with the EASE software or your hardware, we have made it very easy to gather information about your current setup to send to the Woollam Company. By following this procedure, your current configuration, log files, and recent calibration data will be zipped into a file called `CompleteEASE_Debug.zip` that you can email to us along with a description of your problem and contact information. By providing all of this information in one place, we should be able to help you quickly identify the problem.

**Creating a Debug file**

1. Click on the *Options* tab.
2. Click on the ‘Create Debug File’ button.
3. The filename for the debug file is: `C:\CompleteEASE\CompleteEASE_Debug.zip`.

**NOTE:** The Debug File packages all hardware and software configuration files, along with the current Error Log into a single ZIP file.

Email this file to your J.A. Woollam Co. representative to aid in debugging problems with your ellipsometer system.
14.2. Maximum Memory Usage

CompleteEASE will grab up to 1GB of memory when it starts, depending on how much is available. To increase memory beyond 1GB, create a shortcut to the CompleteEASE program and add “-MEMxxxx” after the executable name in the target line. See figure below. The xxxx should be replaced with the number of MB. In the example -MEM2000 tells CompleteEASE to grab 2GB of memory when it starts. If there is not enough memory to do this then CompleteEASE will not start. On 32 bit Windows, we do not recommend setting this higher than 1500. On 64 bit Windows, we recommend no more than ½ of the available memory.

Figure 14-1. Add “-MEMxxxx” to the Target line to increases maximum memory usage.
14.3. **CompleteEASE Data Transfer (to/from WVASE32)**

The data files in CompleteEASE and WVASE32 are not directly compatible, as they have different file formats. The WVASE32 files are written in ASCII text, while the CompleteEASE files are “encrypted”. The file encryption in CompleteEASE is a “copy-protection” mechanism, as this software does not require an “activator key” to run, like WVASE32 requires.

If you have a copy of the WVASE32 software and the software key (activator) that accompanies it, you can easily import and export data files to modify their formats to be compatible with either WVASE32 or CompleteEASE.

The Experimental Data window is used to execute the data transfer of files already saved. Currently opened data is not modified.

**NOTE:** Any data open in the Experimental Data window will be removed (deleted if not saved) by performing the following steps.

**Importing CompleteEASE Data to WVASE32:**

1. From WVASE32, select “CompleteEASE Data Transfer” from the FILE menu of the EXPERIMENTAL DATA window and then select “Import Data from CompleteEASE”.

   ![Figure 14-2](image)

   *Figure 14-2. Choose the CompleteEASE Data Transfer menu from FILE menu of the Experimental Data Window.*

2. Browse to select data files you wish to import from CompleteEASE. Multiple files can be selected.
3. Browse to select the directory where you wish the new files to be saved. Imported files will be saved in WVASE32 format in the directory chosen.

4. After the files are imported, a message will appear, as in Figure 14-5. Previous “.SE” and “.iSE” files are now saved in WVASE32 “.dat” format.
Exporting WVASE32 Data to CompleteEASE:

1. From the Experimental Data window in WVASE32, select the file menu and choose “CompleteEASE Data Transfer”. Then, select the “Export Data to CompleteEASE” option as shown in Figure 14-6.

   Figure 14-6. Selecting “Export Data to CompleteEASE”

2. Browse and select the WVASE32 data files for exporting to CompleteEASE, as in Figure 14-7. Multiple files can be selected.

   Figure 14-7. Choose WVASE32 data files you wish to export.
3. Browse to select the directory for data exported to CompleteEASE, as shown in Figure 14-8. Exported files will be saved in CompleteEASE format in the directory chosen.

![Browse for folder]

Figure 14-8. Browse for location to place the newly created CompleteEASE files.

4. After the files are exported, the following message will appear. Previous “.dat” and “.bdt” files are now saved in CompleteEASE “.SE” and “.iSE” formats.

![CompleteEASE Data Export]

Figure 14-9. Message when all data files are exported. If files were unable to export, the total number of successful files is listed.
14.4. Comparing Results between CompleteEASE and WVASE32

For users that are familiar with using WVASE32, they may find a longer learning curve as they discover the features of CompleteEASE. When comparing results between analysis within WVASE32 and CompleteEASE, please consider the following differences:

**Fit Weighting**

Note the Fit Weighting during data analysis in both software packages. The WVASE default is “Experimental Standard Deviations”, while CompleteEASE uses “NCS” for its default.

**Depolarization Data**

If data contain depolarization measurement, WVASE32 will include this in the MSE calculation and fit. CE will not use the depolarization data during MSE calculation and fit unless you turn on special "Fit Option".

**Intensity Data**

If data are collected with CompleteEASE, the Intensity is also recorded (but not used in the Fit unless turned on with special Fit Option). If this CompleteEASE data is imported to WVASE32, it will also import the Intensity data and WVASE32 will automatically fit this data - thus it needs to be removed from the Experimental Data (using Range Select) to match CompleteEASE standard fit.

**Surface Roughness**

Surface Roughness is calculated differently in CompleteEASE and WVASE32.

In CompleteEASE, surface roughness is an independent thickness quantity with half of its thickness extending into the top layer and half of its thickness extending above the layer. In the actual model calculation (which is created in the background and not visible to users), an EMA layer is added above the top layer with the top layer thickness reduced by half of the surface roughness thickness. Thus, if your layer shows 100nm thick with 2nm surface roughness, the actual model calculation will use 99nm thick layer with 2nm roughness.

In WVASE32, surface roughness is simply an EMA layer over the top layer thickness. If your layer shows 100nm thick with 2nm surface roughness, this is the actual model calculation.

CompleteEASE and WVASE32 use the same method to calculate Intermix thickness. Contact J.A. Woollam Co. for more details if interested.
14.5. Contacting the J.A. Woollam Co.

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