



# **SlideBook™ 5.0**

**Version 5.0.0.1 for Windows XP**

**User Manual**

**Olympus America Inc. Reseller Edition**

**Latest Version 4/1/09**

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This software interfaces with LibTiff version 3.6.1.1

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## Technical Support

If you have any questions or experience any problems with SlideBook™, please contact Olympus America ([slidebooksupport@olympus.com](mailto:slidebooksupport@olympus.com) or 1-800-446-5967 option 4).

## Manual Conventions

### 1.1 *Typographic Conventions*

Menu commands are written in bold and follow the order of menu navigation. For instance, choosing “Open” from the “File” menu is written as **File > Open**.

Dialog fields and other interface items are written in bold, as in **Initial Offset**.

References to other sections of the manual are underlined, as in Image Capture.

New terms are *italicized*.

**NOTES** will be written in bold and small capitals. Notes that you should pay particular attention to are prefaced **CAUTION**.

### 1.2 *Of Pixels and Voxels*

In other contexts, *image* typically connotes a two-dimensional entity. However, SlideBook™ was specifically designed to operate on three-dimensional data as easily and intuitively as two-dimensional data. In light of this, a two-dimensional data set can be seen as a special case of a three-dimensional set where data has been collected for only one value of the *z*-axis. An image corresponds to *all* data collected during a particular capture.

To simplify this manual, we always refer to a single data element as a *voxel* and a region as a *volume*, regardless of the depth of the image. These terms map directly to *pixel* and *area*, respectively, for a two-dimensional set.

## Using this Manual

The SlideBook™ Manual describes all of the steps necessary for performing digital microscopy, from installing the software to analyzing images. The chapters are outlined below. If you are new to SlideBook™, you may wish to start by taking a step-by-step Quick Tour as outlined in Chapter 2. This will familiarize you with basic elements and commands in SlideBook™.

### **1.3 Chapter 1 - Installing SlideBook™**

If your system was installed by 3i or a 3i reseller, then you will not need to install software on your system. However, if it was not installed by 3i or a 3i reseller, or you need to upgrade your software, you can learn how to do this in Chapter 1.

### **1.4 Chapter 2 – Quick Tour**

This chapter will take you on a step-by-step guided tour of the basic elements of SlideBook™. This is the perfect starting point for the beginning user.

### **1.5 Chapter 3 – SlideBook™ Organization**

This chapter describes how hardware control, image capture, data storage, and data analysis are organized into SlideBook™ menus, windows, and objects.

### **1.6 Chapter 4 - Configuring Your System**

If your system was installed by 3i or a 3i reseller, then you will not need to configure your system prior to use. However, if it was not installed by 3i or a 3i reseller, or you need to make changes to your system configuration, you may do so by selecting **Edit > Define Optics** or **Edit > Hardware Configuration**. Chapter 4 teaches you how to configure hardware and define optics.

### **1.7 Chapter 5 - Controlling the Camera and Microscope**

Next, you will learn how to set your microscope to the proper optics and bring your sample into view and focus. Any motorized aspect of the microscope can be controlled through SlideBook's **Focus Window**. You may bring up this window by selecting **Window > Focus Window** or clicking on the focus window button in the toolbar .

### **1.8 Chapter 6 - Image Capture and Import**

Parameters for image capture will be set in the **Capture Window**. Here you will set camera parameters, such as exposure and binning, by performing test exposures. You may bring up this window by selecting **Image > Capture New Image** or by clicking on the capture button in the toolbar . You may also import images that were captured using other programs by selecting **Image > Import**.

### **1.9 Chapter 7 – Advanced Capture**

In this chapter, you will learn to perform more advanced capture sequences, such as timelapse, 3D, 4D, and *montage* capture.

### **1.10 Chapter 8 - Image Display and Manipulation**

Once you have captured an image set, it may be displayed in various modes using the **View** menu. Interactive data views allow you to display any combination of different channels and masks (ROIs) as well as change lookup table parameters. You may also perform operations such as volume rendering. Note that operations from this menu do not alter the underlying intensity data contained in an image.

### **1.11 Chapter 9 - Preparing an Image for Analysis or Export**

Before performing analysis on an image, you may want to perform additional operations on an image. For example, you may wish to eliminate any constant artifacts from the image by performing *flat field correction*. It may also be useful to deblur an image by performing deconvolution. Other operations that may be necessary or useful prior to analysis are cropping, aligning channels, or applying filters. Operations such as these are found under the **Image** menu.

### **1.12 Chapter 10 - Image Analysis**

Mathematical analysis can be performed on regions of the image that you select. You may select these regions by creating a binary overlay called a mask. Masks are generated from the **Mask** menu. Once you have made a mask, you may obtain information such as average intensity and number of pixels in the region of interest by using the **Statistics** menu.

NOTE: Screenshots used in this manual were taken with SlideBook running in Windows Vista™. Appearance may vary slightly with different operating systems.

## 1 Installing SlideBook™

### 1.1 Computer System Requirements

Your computer must meet the following system requirements in order to run SlideBook™ 5.0.

#### 1.1.1 Operating System

Windows XP Professional for computers driving hardware, also known as *online* systems.  
Windows 32-bit or 64-bit Vista for computer running analysis only, also known as *offline* systems.

#### 1.1.2 Hardware

##### 1.1.2.1 Processors

Recommend at least dual quad-core Xeon 3.2GHz processors

##### 1.1.2.2 Memory Requirements

RAM: recommended 3 GB, minimum 2GB.

Hard drive: minimum 200 GB required for installation

##### 1.1.2.3 Graphics Card

(for Interactive Volume Rendering only)

VRAM: minimum 256 MB, recommended 512 MB or more

Minimum models: NVIDIA GeForce or NVIDIA Quadro FX

Recommended models: NVIDIA GeForce 9800, NVIDIA Quadro FX 3540 or higher

Please contact Olympus America ([slidebooksupport@olympus.com](mailto:slidebooksupport@olympus.com) or 1-800-446-5967 option 4) with questions regarding hardware compatibility.

### 1.2 Connecting the Hardware Key

SlideBook™ comes with a hardware key, or *dongle*, that permits the application to run on the machine that has the key attached. The key is a Sentinel series black (with purple accent) USB device manufactured by SafeNet. Before you begin the software installation; please connect the dongle to a USB port in the computer.



**NOTE:** Given the hardware key system, it is possible to install the SlideBook™ software on several different machines. However, the software will only run on the machine that has the key attached. Networked computers may run SlideBook™ if one of the computers on the network has a network key.

### 1.3 About SlideBook™ Installers

SlideBook installers are available as “compact” or “complete”.

- **Compact** – installs SlideBook program files, Sentinel dongle driver, user manual, and QuickTime.
- **Complete** – installs all SlideBook program files, user’s manual, QuickTime program files, and Sentinel dongle drivers. Also allows the option to copy camera driver files, Edgeport driver files, and QuickTour sample files to the Intelligent Imaging Innovations, Inc directory.

### 1.4 Copying the Software from CD-ROM

1. Insert the SlideBook™ CD in your CD-ROM drive.
2. Navigate to the CD-ROM and copy the SlideBook™ setup program, InstallSlideBook.exe to your desktop.
3. Double-click on the icon to begin installation and proceed to the section “Installing SlideBook 5.0” (below).

### 1.5 Copying the Software from your Download Site

Every SlideBook™ user may access a private download site in order to obtain current installers and updaters. Please contact Olympus America ([slidebooksupport@olympus.com](mailto:slidebooksupport@olympus.com) or 1-800-446-5967 option 4) to obtain your user name and password.

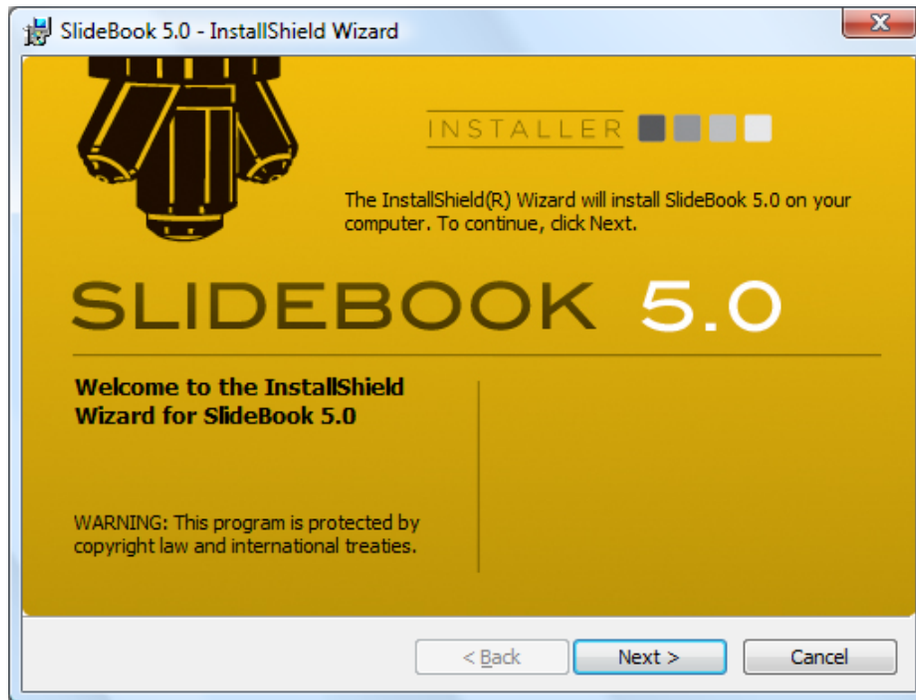
To access your download site:

1. Go to [www.intelligent-imaging.com](http://www.intelligent-imaging.com)
2. Go to Downloads, and then Customer Login, and click on link “Click Here to Login.”
3. Enter your username/password (case-sensitive).
4. Click on the desired link to download the software. When prompted, select **Save** and then navigate to the desired destination for the file.
5. Click **OK** to save the file.

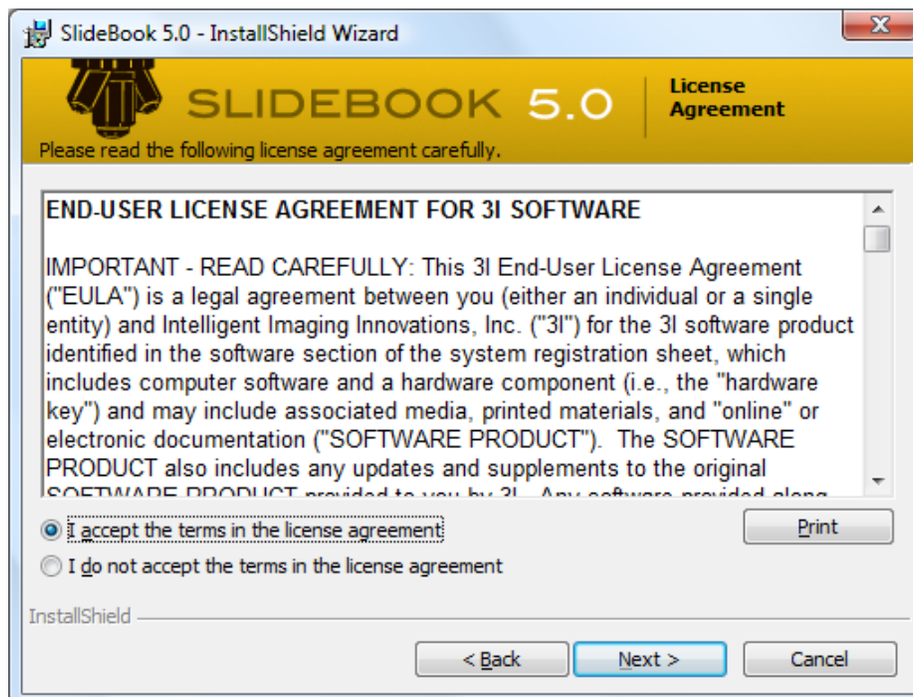
### 1.6 Installing SlideBook 5.0

The SlideBook installer allows you to install the program, drivers, adjunct programs, the manual, and tutorials. If you are installing SlideBook for the first time, continue with this section. If you already have SlideBook installed on your machine, please see [Installing Updates](#).

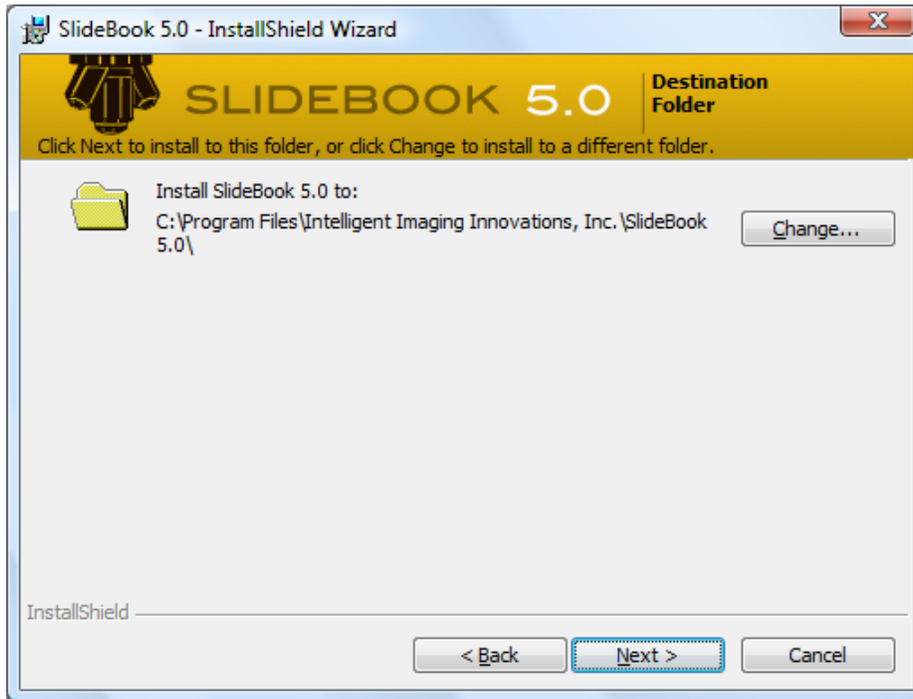
1. Double-click on the InstallSlideBook icon to start the setup utility. The following dialog box will be displayed.



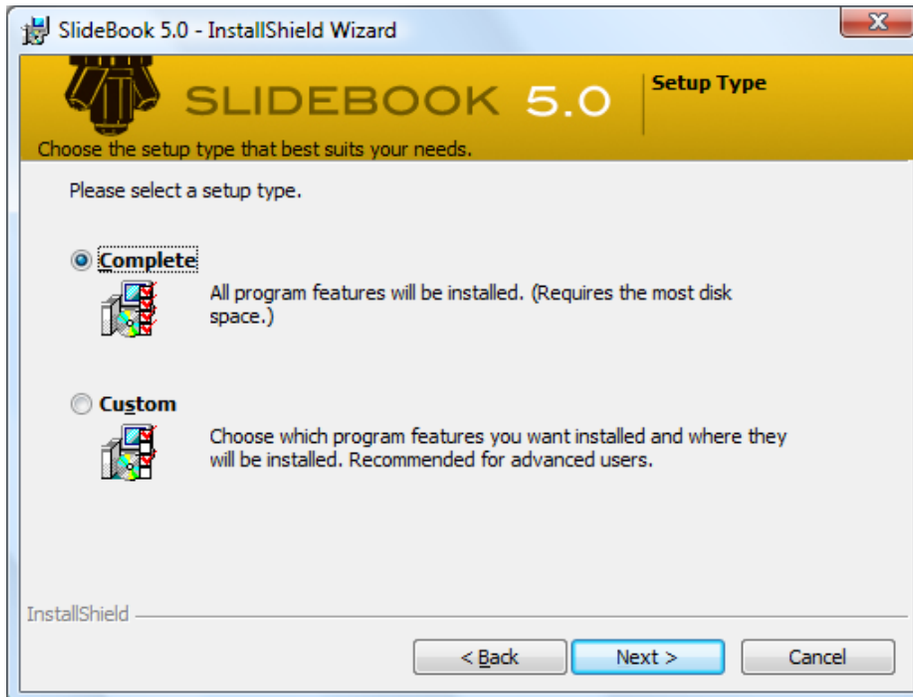
2. Select **Next** and you will be prompted to review and accept the terms of the license agreement.



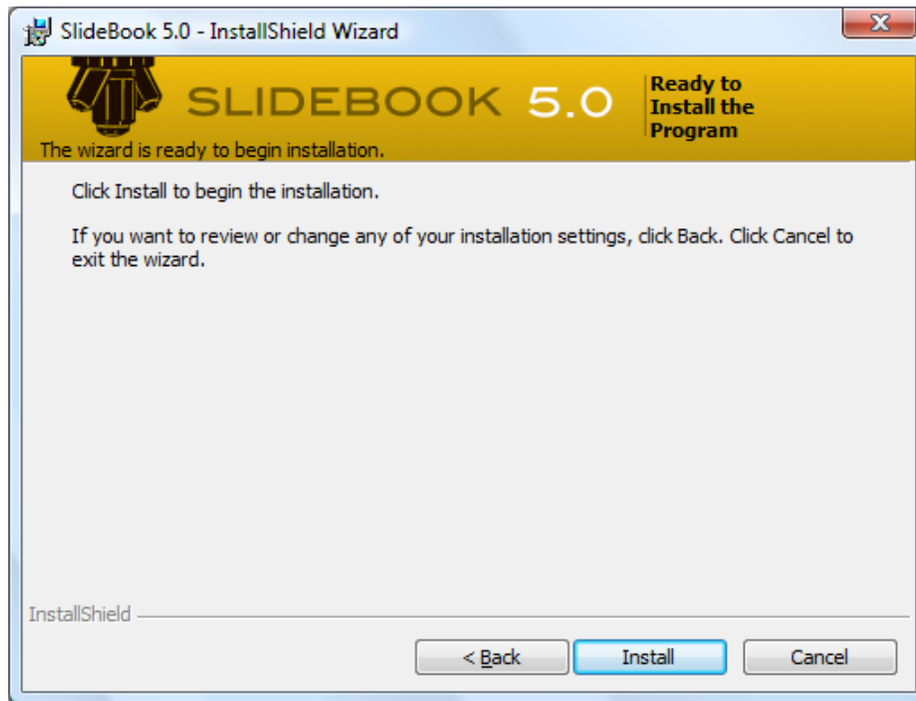
3. Select **Next** and the following window will appear:



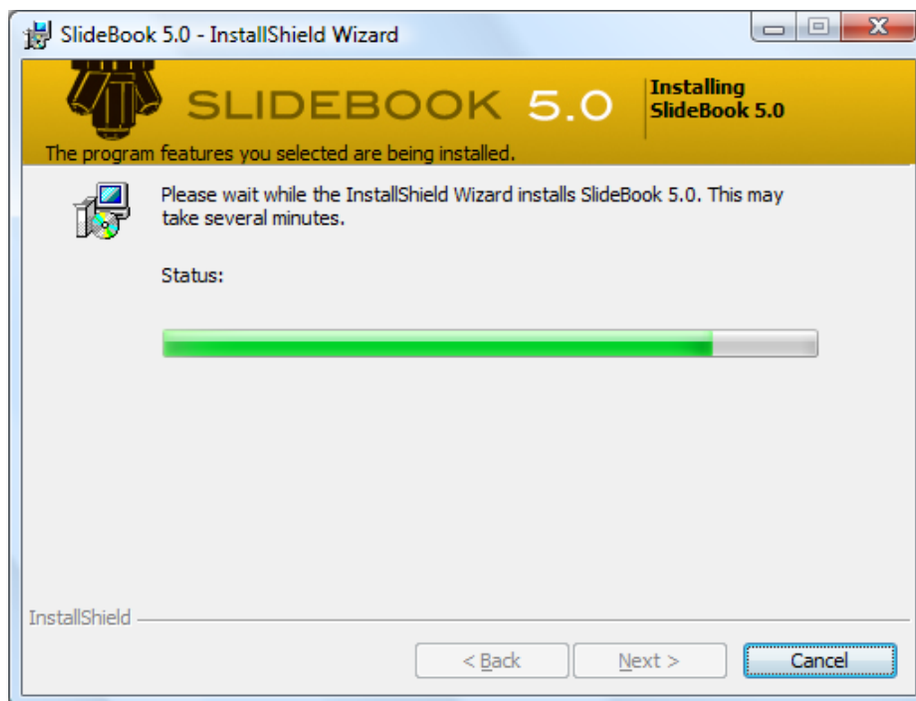
4. If the default destination directory is acceptable, select **Next**. If you would like to install to a different directory, click **Change** and navigate to the desired location and then select **Next** to continue installation. The following dialog box will appear:



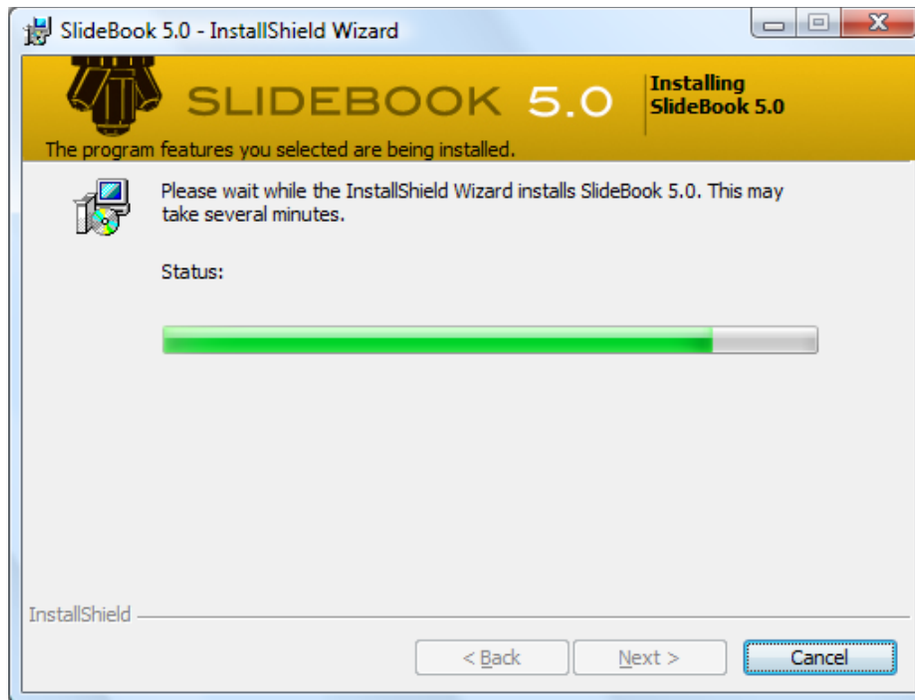
5. Select **Complete** then **Next**. The following dialog box will appear:



6. Select **Install** and the setup utility will display status of the installation:



7. When the installation is complete, the following message will confirm installation:



8. Click **Finish** to complete the setup.

You may now launch SlideBook™ from either the **Start Menu > Programs** or from the **Shortcut to SlideBook.exe**.

**NOTE:** When you start SlideBook™ for the first time, it will notice that a *preferences file*, which is stored along with the application, does not exist. SlideBook™ will create a default preferences file. As you configure your hardware and define optics (discussed in Chapter 4, [Configuring Your System](#)), SlideBook™ will update this preferences file with specific information about your setup.

An electronic copy of this manual, called SlideBook 5.0 Users Manual.pdf, is in Adobe Acrobat format, and can be opened by an Acrobat reader. Feel free to distribute the manual to all users of SlideBook™.

### 1.7 Installing Updates

You may download software updates at [www.intelligent-imaging.com](http://www.intelligent-imaging.com) as described above in [Copying Software from your Download Site](#). Before installing an update, be sure to back up your configuration files. To backup your files, go to **Edit > Setup Guides > Backup Preferences**. You will be prompted for a location. If you use the defaults settings, a folder called Global Preferences will be created in the SlideBook 5.0 directory. After downloading is complete, double-click on the file that you have just downloaded and follow the installation directions. Install the update in **Program Files\Intelligent Imaging Innovations, Inc\SlideBook**. If SlideBook™ is not located in the above path; refer to section 4.6 Working with SlideBook Preferences and Hardware Properties on page 74.

## 2 Quick Tour

This Quick Tour will introduce you to some basic elements of SlideBook. By the end of this tour, you will know how to do the following:

- Open *slides* and *images*
- Display a **Main View**, **Three View**, and **Tile View**
- Use basic tools in the **Info/Tool Bar** to Maneuver within a *view*
- Perform volume rendering
- Create a *mask* and obtain statistics

**NOTE:** Menu operations, such as choosing “Open Slide” from the “File” menu are written as **File > Open Slide**.

In order to run the Quick Tour, first make sure that the SlideBook program and Quick Tour slides have been installed. If not, please go to Chapter 1, Installing SlideBook™ and follow the procedures for installing the software and the Quick Tour slides.

### 2.1 Opening Slides and Images

To begin the tour, we will first open a slide file, SlideBook’s native file type (\*.sld). A slide file corresponds to a single file on the disk and can be thought of as SlideBook’s document type. It can contain many multi-plane, multi-channel images in any combination.

1. Launch SlideBook by pointing to **Programs > SlideBook** from the **Start** menu.

An empty slide will appear in the window under the SlideBook menus.

2. Close the new slide by choosing **File > Close Slide** or by clicking on the  in upper right corner.
3. Choose **File > Open Slide** and navigate to the Documentation folder in the SlideBook 5.0 directory. Double click on QuickTourWIN1.sld.

The following slide window appears that shows a thumbnail of one image. This slide contains a single 3D image of a dividing B cell.

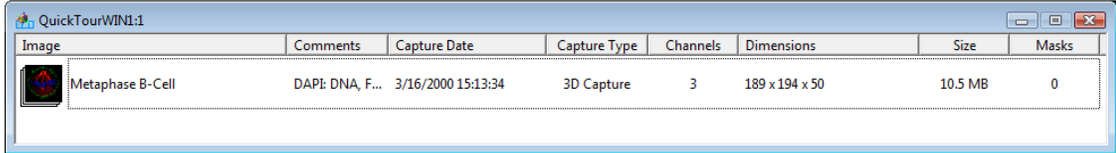


Image	Comments	Capture Date	Capture Type	Channels	Dimensions	Size	Masks
 Metaphase B-Cell	DAPI: DNA, F...	3/16/2000 15:13:34	3D Capture	3	189 x 194 x 50	10.5 MB	0

### 2.2 Displaying a Main View

SlideBook can display any image using a variety of *views*. Each data set can have as many views open at one time as you desire. SlideBook has seven kinds of views that display individual images:

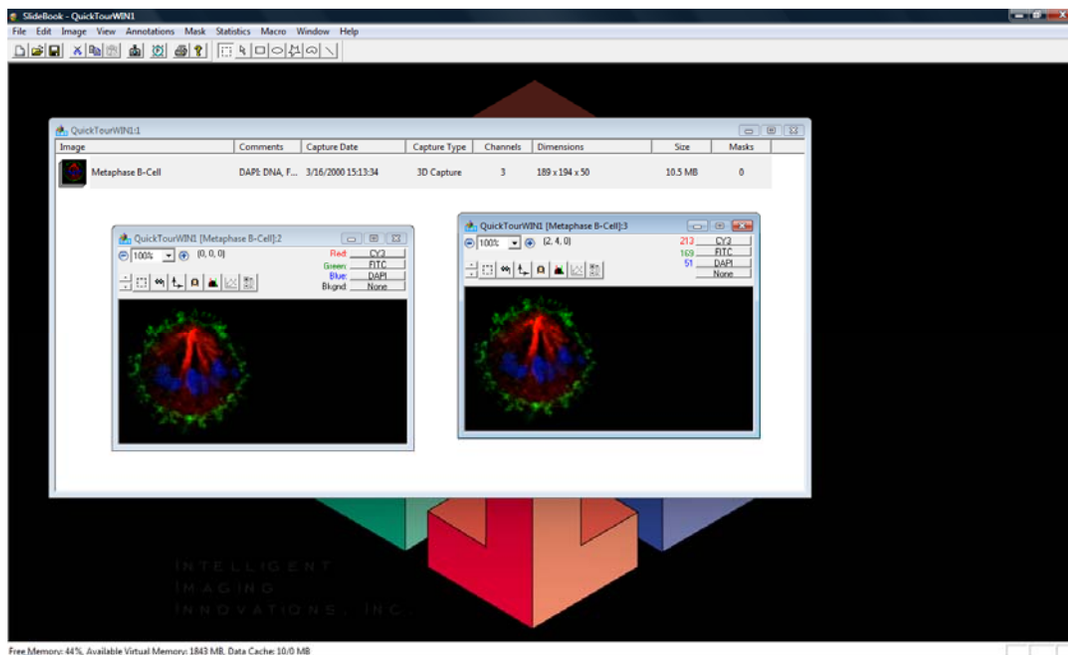
- **Main View**
- **Three View**
- **Tile View**
- **Channel View**
- **Multidimensional Channel View**
- **3D Volume View**
- **3D Surface View**

**NOTE:** Views are only used to display and alter the look of an image. The underlying data are not affected.

Now, we will open a *Main View*.

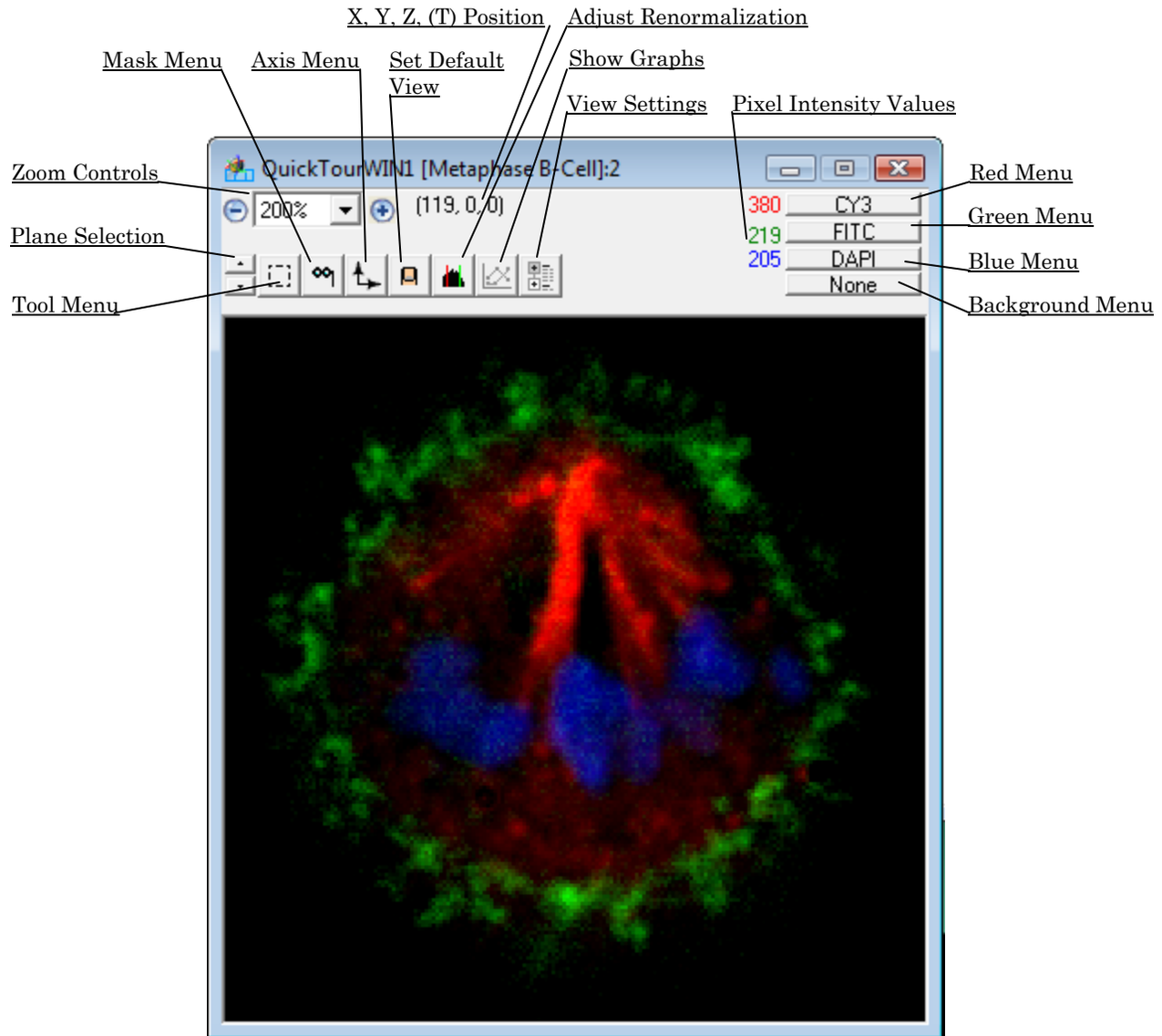
1. Double-click anywhere on the thumbnail or within the shaded region that surrounds the thumbnail. A new Main View window will open that displays the image.
2. Create another Main View of the Dividing B Cell image by first selecting the image with a single click, then selecting **View > New Main View**.

At this point you should have three open windows: the Slide Window and two Main View windows. Both Main Views show the dividing B cell in three colors, where each color corresponds to a different *channel*, or fluorophore. DNA appears in blue (AMCA channel), surface immunoglobulin appears in green (FITC channel), and tubulin appears in red (Cy3 channel).



### 2.3 Using the Info/Tool Bar

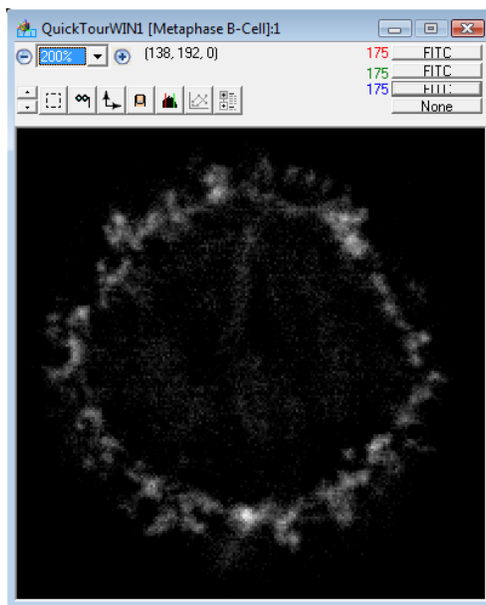
The **Main View** window contains an *Info/Tool Bar* placed above the image of the data. The Info/Tool Bar displays a variety of useful information and gives you quick access to tools you will use to explore and manipulate the view. The Info/Tool Bar is present in all data views, namely the **Main View**, **Three View**, **Tile View**, and **Channel View**.



#### 2.3.1 Channel Menus and Data Values

On the far right of the Info/Tool Bar are three pop-up menus with colored red, green, and blue text next to them. These menus allow you to select the fluorescence channel that is displayed for each color. The Main View in this data set has the *red menu* set to the CY3 channel, the *green menu* set to the FITC channel, and the *blue menu* set to the AMCA channel.

1. Click on one of the **Main View** images to select it as the active window.
2. Click on the **CY3** bar (red menu) and select **FITC** from the pop-up menu. Notice that the FITC label in the view now appears yellow from the combination of red and green pixels.
3. Click on the red menu and select **None**. Note that there are no data displayed in red.
4. Next, set all color menus to the same channel, **FITC**, and observe that the image is now displayed in black and white.



Try changing the settings for each menu, noting how the display changes.

5. Reset the red menu to **CY3**, the green menu to **FITC**, and the blue menu to **AMCA**.

To the left of the channel menus are colored numbers. These numbers correspond to the *intensity* of the voxel that the cursor is positioned over for each channel.

6. Move the cursor over the image displayed in the Main View. Note how the intensity values increase as you move over brighter regions and decrease as you move to darker regions.
7. Visualize just the DNA by setting the red, green, and blue menus to **DAPI**. As you move the cursor over the image, note that the red, green, and blue values are equal.

- Close this Main View window by clicking on the  in upper right corner. One Slide View and one Main View window should now be open.

### 2.3.2 Up/Down Arrows, Scrolling through a 3D image

The “Dividing B-Cell” image is actually a 3D data set containing 50 planes captured at 0.2 micron spacing. You may navigate through the stack in a variety of ways.

- Click on the remaining Main View. Scroll through the  $z$ -axis using the small arrow buttons   on the far left of the info/tool bar.

Note that the coordinates in the upper left corner give you the  $x$ ,  $y$ ,  $z$  position of the mouse pointer. As you scroll through the image stack, the  $z$  position will update to your current location.

- Select **View > Go To Plane**. Enter “2” in the data entry field and select **OK**. The value of the  $z$  coordinate is now 2, which corresponds to the third plane in the image, as the count starts at 0.



- Next, select **View > Previous Plane** and **View > Next Plane** commands, observing the value of the  $z$  coordinate in the upper left corner of the info/tool bar.
- If your mouse is equipped with a scroll wheel it can also be used to move through the planes of the image.

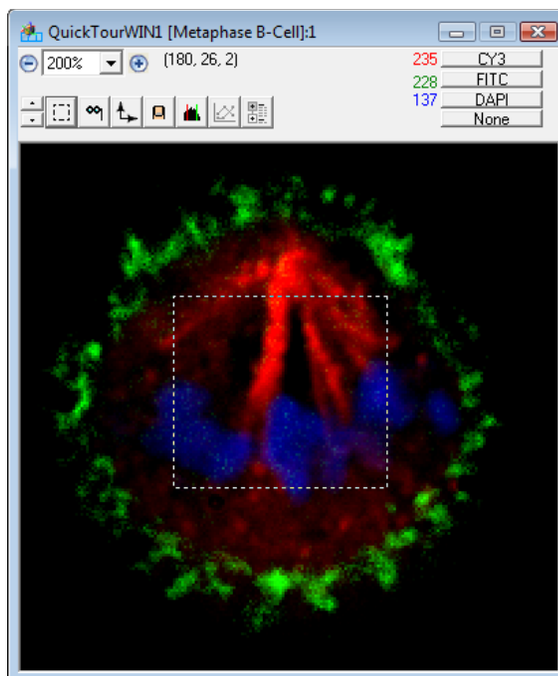
### 2.3.3 Tool Menu

The *tool menu* lets you choose the function the mouse cursor performs.

#### 2.3.3.1 The Marquee Tool

The *marquee tool* is the default tool of the tool menu and allows you to select a two- or three-dimensional region of the image.

- Click and drag the mouse over a region in the **Main View**. Note that the cursor appears as cross-hairs. Release the mouse button.



2. You have just selected a 2D region using the marquee tool. In the visible axes ( $x$  and  $y$ ), the marquee tool works as in any drawing program, but because SlideBook is designed around the 3D images, a marquee must be manipulated in the invisible plane as well. To illustrate this point move to different  $z$  positions in the image using one of the methods outlined above. Notice that the marquee selection only appears in the plane in which it was originally drawn.
3. With the marquee region still selected, go to **View > Select Rect in All Planes**. Again, scroll through the different  $z$  planes in the image. Notice that now the marquee selection exists in all planes of the image.
4. Click once on the image to remove the marquee from the view.

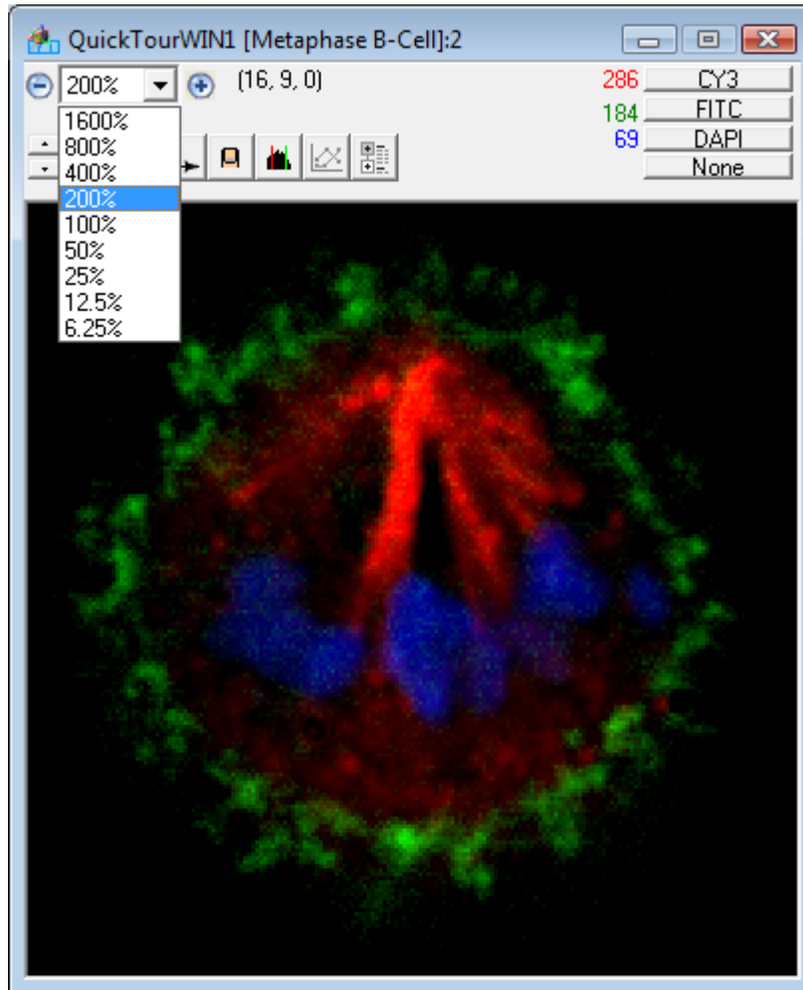
**NOTE:** There are other ways to select 3D regions that will be demonstrated later in the tour.

### 2.3.3.2 Zoom Tool

The *zoom tool* is used to expand or shrink the size of the window (and the corresponding zoom factor of the displayed image).

1. Click on the tool menu and select the zoom tool.
2. Click on the **Main View** image two times, zooming in on the image. Note that the percent magnification is displayed in the upper left hand corner.
3. To zoom out on the image simply hold the shift button down while the zoom tool is selected and click on the Main View image.

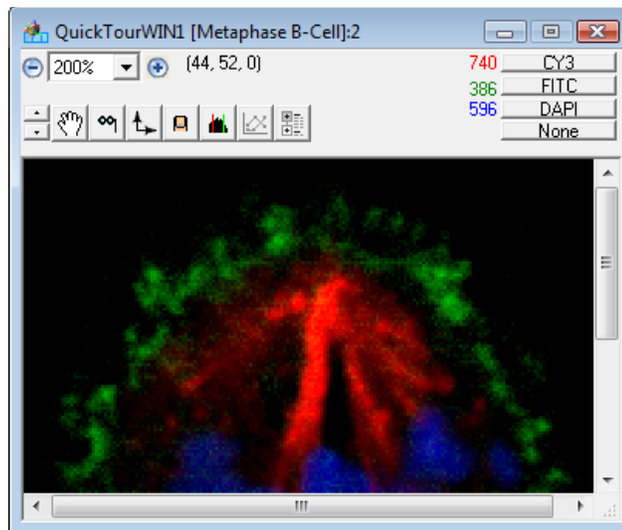
4. Alternately, you may click on the “-“ and “+” buttons or utilize the drop down menu in the upper left hand corner of the view window as illustrated below.



### 2.3.3.3 Hand Tool

The *hand tool* lets you move the displayed portion of an image that is in a window smaller than the extent of the displayed axes.

1. Zoom in on the **Main View** image until it is larger than the confines of the window as illustrated on the next page.



2. Select the hand tool button from the tool menu.
3. Click and drag the hand tool in the window, moving the portion of the image that is displayed.

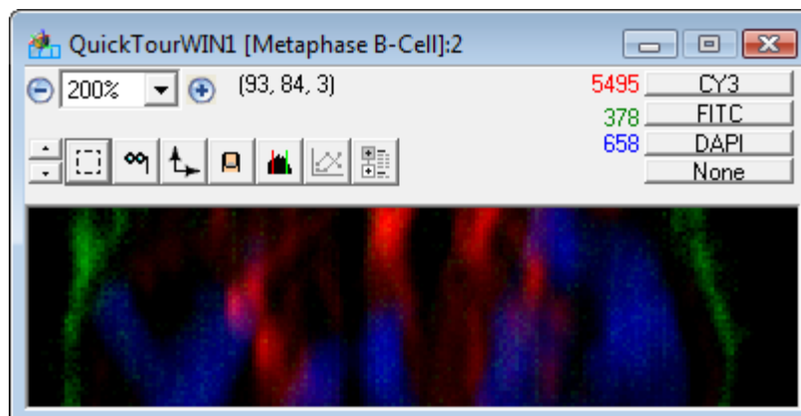
Notice that the hand tool performs the same function as the scroll bars on the right and bottom sides of the Main View.

4. Return to a 100% zoom factor by any of the methods discussed previously.

### 2.3.4 Axis Menu

SlideBook is capable of visualizing data along all three axes.

1. Click on the *axis menu* on the tool bar and select *y*. Enlarge the image by using the zoom tool.
2. Use the up and down arrows to move through the planes.



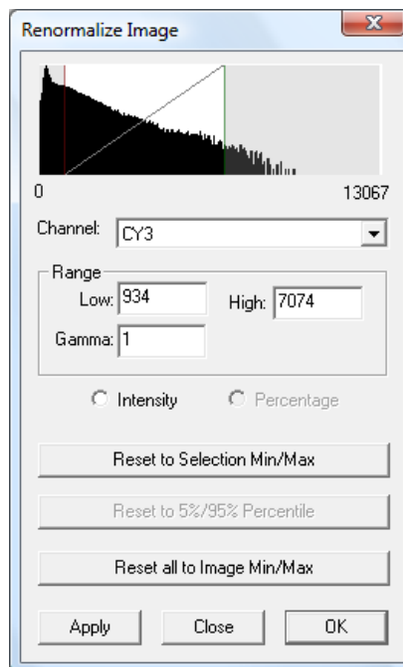
The  $y$ -axis is now the invisible axis. You are observing the  $xz$ -plane and scrolling through the  $y$ -axis. You may view all three Cartesian planes simultaneously using the *Three View*, which will be discussed later in the tour. The axis menu options are unavailable in the Three View because all three planes are already displayed.

3. Select the  $z$  axis from the axis menu to return the view to the  $xy$ -plane.

### 2.3.5 Renormalize Button

The *renormalize button* brings up the Renormalize Image dialog box. This tool allows you to change the range of intensities that are displayed for any channel that you choose.

1. Click on the renormalize button. The following dialog box will appear:

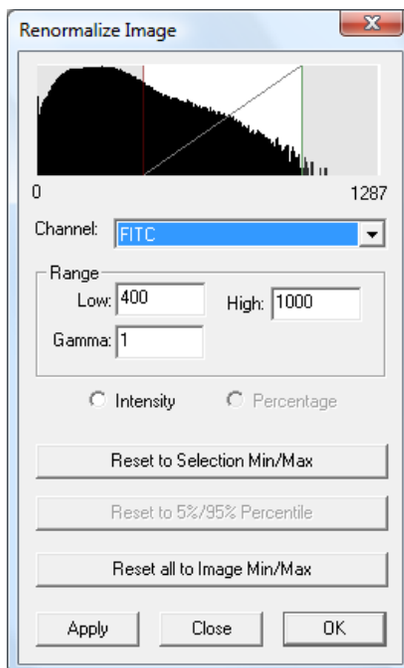


A histogram is displayed showing the relative number of pixels on the  $y$  axis and intensity values on the  $x$  axis for the specified channel. In this image, the CY3 channel intensities range from 0 to 14281.

2. Click on the drop-down channel menu and select **FITC**.
3. Click on the red and green bars in the histogram window and drag them to the left or right, observing the change in the display of the green data.

The red and green bars allow you to select the minimum and maximum intensities that correspond to the absence (black) or full saturation of color in the display, respectively. Again, the underlying values in the image are not affected.

4. Now enter the numbers 300 and 800 in the **Low** and **High** data entry fields. Note that the image does not automatically update as it does when moving the red and green bars.
5. Click **Apply** to register the change and update the image.
6. Enter 400 and 1000 in the **Low** and **High** data entry fields and click on **Apply** to register the change.



7. Choose **OK** to exit the Renormalize Image dialog box.

**NOTE:** You may also bring up the Renormalize Image dialog box by selecting **View > Renormalize**.

### 2.3.6 Thumbnail Button

The *thumbnail button* lets you set the default view settings for a particular image: color assignment of particular channels, renormalization information, and the *z* position of a newly opened image. If you close the Main View that is currently open without setting a default, the changes that you just made using the renormalize button will be lost.

1. Select **View > Go to plane...** and enter “10” in the edit box. (Note that plane number 10 is actually the eleventh plane, as plane numbering starts at zero.)
2. Select **OK**.
3. Click on the thumbnail button to select your current display as the default.

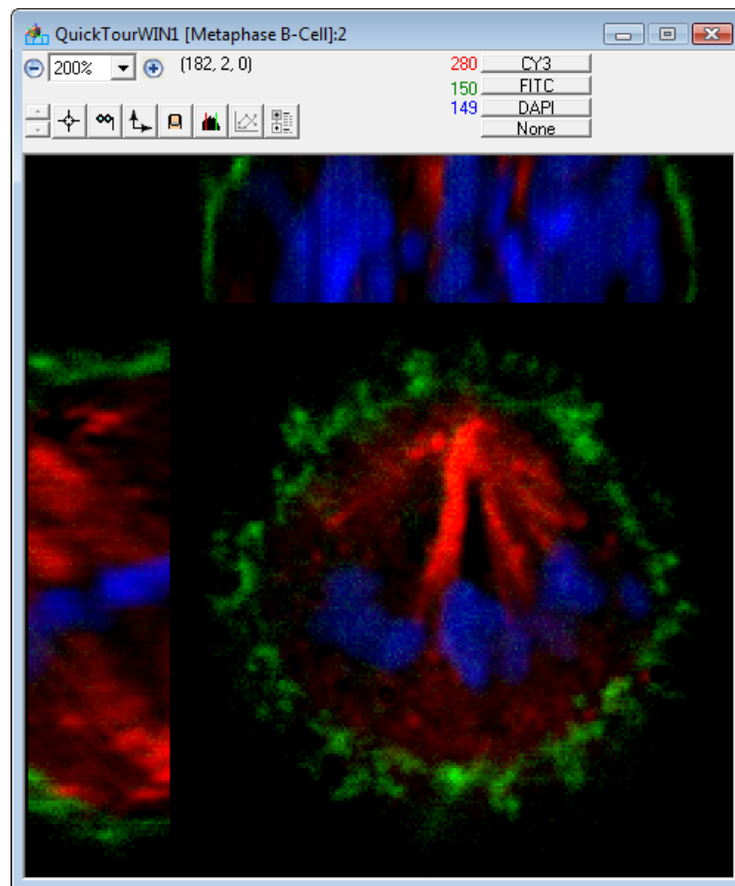
Now all subsequent views of the image will default to the same display. You have also changed the “thumbnail” image that is displayed in the slide window.

4. Open another **Main View** of the image by double-clicking on the thumbnail. Note that the new settings that you entered previously apply to this new **Main View**. Also, notice that the eleventh plane is displayed.
5. Close the **Main View** that you just created.

## 2.4 Generating a Three View

SlideBook allows you to view all three axes simultaneously using a **Three View**.

1. Select **View > Three View** to generate a new **Three View**.



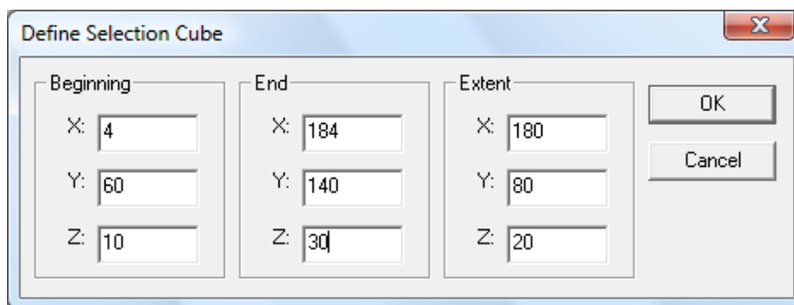
A new window will open with an  $xz$  pane on the top, a  $yz$  pane on the left and an  $xy$  pane in the middle. The most useful tool for exploring this view is the *point selection tool*.

2. Choose the point selection tool icon in the tool menu .
3. Click on the  $xy$  pane (center pane) of the **Three View** with this tool. Notice that you update the  $xz$  and  $yz$  panes.

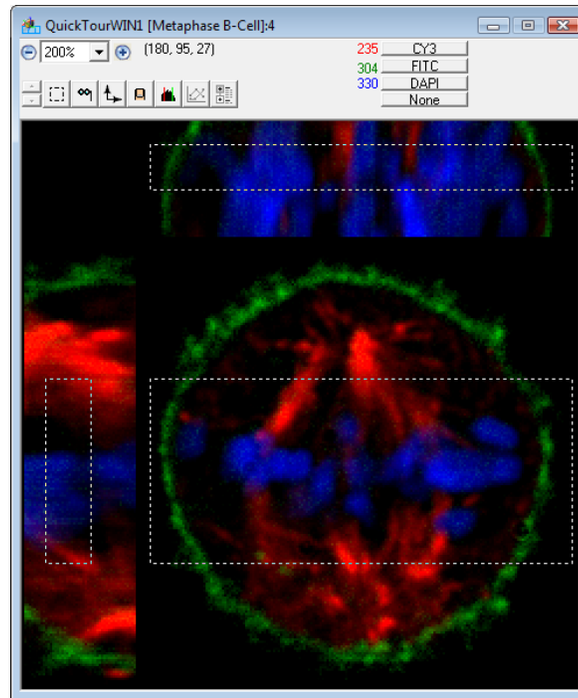
4. Hold the mouse down and drag it around the center pane to get a cutting plane effect.
5. Click and drag the mouse along the  $x$  axis of the center pane, and note that the  $yz$  plane is rapidly changing, while the  $xz$  plane is relatively still.
6. Next, click and drag the mouse on the  $yz$  pane to the left, and observe that the other two planes are being updated.

Now we will make 3D selections in the **Three View** using the marquee tool.

7. Select **View > Go To Plane**. Enter plane 20 and select **OK**.
8. Next, choose the marquee tool from the tool menu in the info/tool bar.
9. Select the chromosomes by clicking and dragging a rectangle around them.
10. Then go to **View > Define Selection Cube**. The following dialog box will appear:



11. The Define Selection Cube dialog box allows you to manually enter coordinates for the selection box. Change the **Beginning Z** value to 10 and the **End Z** value to 30. Notice that **Extent Z** has updated to indicate 20 planes are selected. Leave the  $x$  and  $y$  coordinates unchanged and press **OK**.



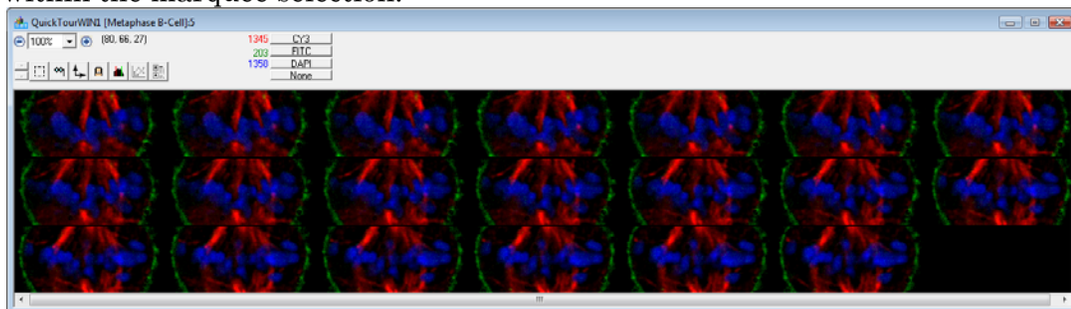
Notice that the  $z$  extent of the marquee selection only spans the 20 planes that were selected.

Next, we will spawn a Tile View that contains the data defined by the selection cube.

## 2.5 Spawning a Tile View

SlideBook can generate a plane-by-plane mosaic called a *Tile View* from any Main View or 3D selection. Now we will spawn a Tile View of the selection cube that we just created.

1. Select **View > Tile View**. A large mosaic of the chromosomes is displayed with each panel representing one of the 20 different  $z$  planes that was contained within the marquee selection.



2. Select the *hand tool*  from the tool menu in the info/tool bar.
3. Click and drag around a single pane of the mosaic. This adjusts the position of the display cube within the image, and allows all of the tiles to be readjusted simultaneously.

4. Close the Tile View.
5. Open a Tile View from the Main View by first clicking on the Main View to remove the marquee selection and then selecting **View > Tile View**. Notice that the entire image space is now displayed.

**NOTE:** In order to display a **Tile View** of a portion of the image from the **Main View**, you must first define a 3D selection.

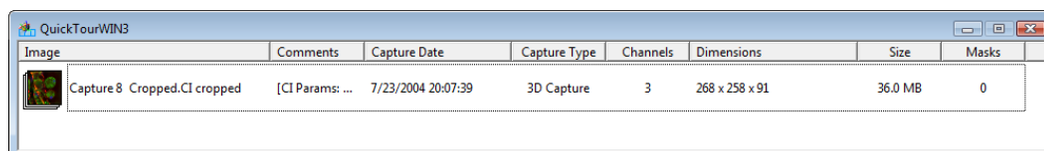
6. Close the Tile View.
7. You may close the slide by either clicking on the  in the upper right hand corner, or by selecting **File > Close Slide**.

### 2.6 Performing Volume Rendering

SlideBook can perform interactive volume renderings using the *3D Volume View*. The following rendering types are available. When you go to **View > 3D Volume View**, two options will be available, **High Speed** and **High Quality**. **High Speed** will render the image at the highest speed possible. **High Quality** will render the image with a more rigorous algorithm that may improve the quality of the image with a small decrease in the speed performance. These settings apply to all modalities of volume rendering but are most noticeable when using **Dynamic Lighting** mode.

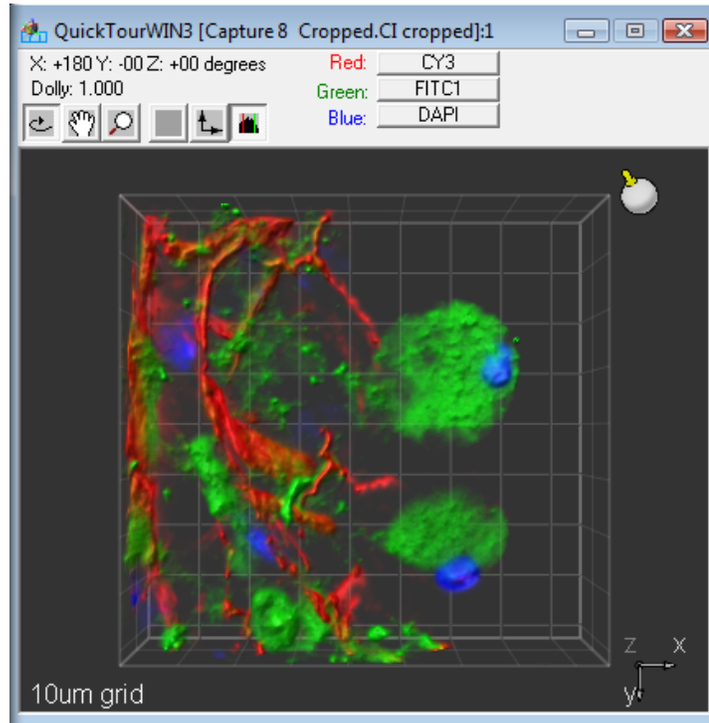
- **Dynamic Lighting** – allows you to illuminate your object as you wish by allowing control of the lighting angle. Useful for determining surface characteristics.
  - **Fixed Lighting** – uses a fixed lighting pattern for shading, approximates surface characteristics.
  - **MIP** – Displays the pixel of maximum intensity along the axis perpendicular to the viewing plane. This type of rendering does not require an advanced graphics card; however, performance of certain functions will be improved with one.
  - **X Ray** – Displays the summed intensity along the axis perpendicular to the viewing plane.
1. Choose **File > Open Slide** and navigate to the Documentation folder in the SlideBook directory. Double click on `QuickTourWIN3.sld`.

The following slide window appears that shows a thumbnail of one image. This slide contains one 3D image of a mouse mammary gland section.



2. Click anywhere in the area of the image in the list and select **View > 3D Volume View > High Speed** or **High Quality**, depending on the desired mode.

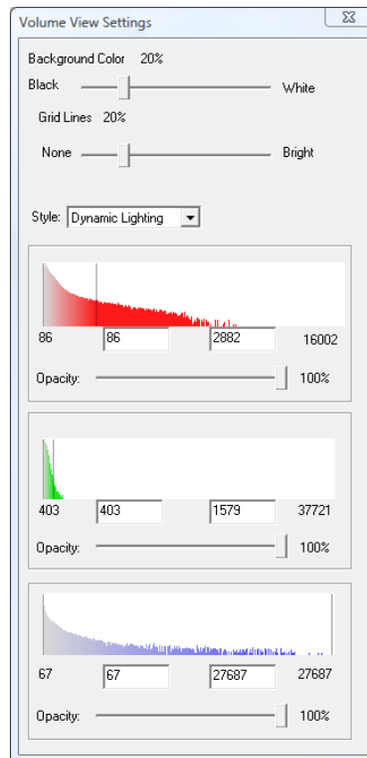
**NOTE:** If the image contains a marquee selection, SlideBook will only render the portion of the image that is selected.



1. To change the channels that are displayed, use the channel menus as you would for the Main View, Three View, or Tile View.
2. To alter the shading, grab the *lighting tool* by clicking on the icon in the upper right hand corner  and rotating it to achieve the desired lighting.
3. To zoom into the view, click on the zoom tool  and then click and drag up and down in your view to zoom in and out.
4. To rotate the image, click on the *rotate tool*  and then click and drag in the direction that you wish to move the image. You may click and drag either directly on the image or on the coordinate image in the lower right hand corner.
5. To move the image in the field of view, click on the hand tool icon  and then grab and move the image by clicking and dragging on it.

- To instantly reorient your image, you may choose to display a specific face of the volume by choosing the axis tool  and selecting **Top**, **Bottom**, **Left**, **Right**, **Front**, or **Back**.
- To select a different rendering mode, or to alter the colors displayed, select the renormalization icon . The Volume View Settings window will appear.

**NOTE:** The Volume View Settings window automatically opens when you open a 3D Volume View.



Several features are offered in addition to the ability to renormalize the image (change the lookup tables).

- Alter the background color using the **Background** slider and change the brightness of the gridlines using the **Gridlines** slider.
- You may alter the opacity of an individual channel using the **Opacity** slider control that corresponds to each color.
- Once you have explored this view, you may close the slide by selecting the  in the upper right hand corner, or by choosing **File > Close Slide**.

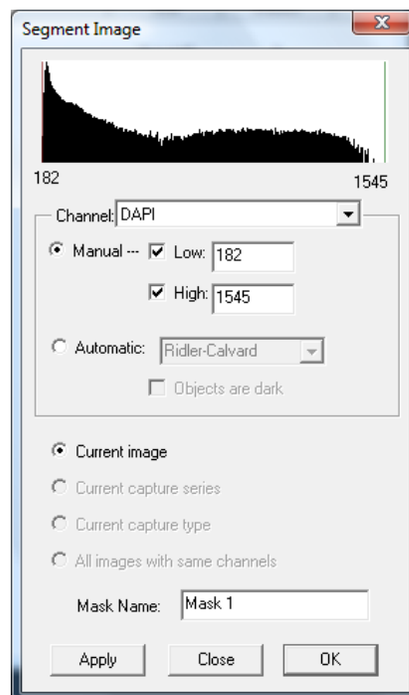
For further information about volume rendering, please see [Displaying a 3D or 4D Volume View \(Volume Rendering\)](#) on page 189.

## 2.7 Creating Masks and Generating Statistics

A *mask* is a two- or three-dimensional set of binary values that has the same extent as the image to which it is assigned. Masks are often termed regions of interest (ROIs) in other imaging programs. Masks are used for performing advanced selection and analysis of image data and can either be created and edited manually or generated automatically through threshold segmentation and other techniques. A single image can contain several different masks. We will now learn how to create masks using a variety of techniques.

### 2.7.1 Creating a Mask using Threshold Techniques

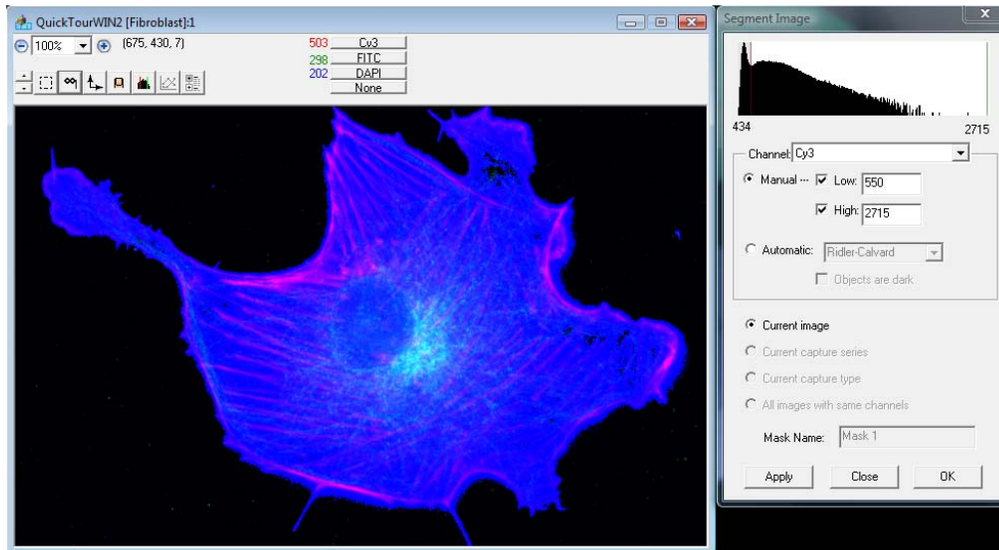
1. Select **File > Open Slide** and navigate to `QuickTourWIN2.sld` in the Documentation folder within the SlideBook directory. A slide window will open that contains two images.
2. Double click on the “Fibroblast” thumbnail to open a Main View of the image.
3. Create a threshold mask by clicking on the mask menu icon  and selecting **Create Segment Mask**. Alternately, you can reach this dialog by selecting **Mask > Segment**. The following dialog box will appear:



4. Select **Apply**. The Generating Mask dialog box will appear and report the segmentation progress by plane.

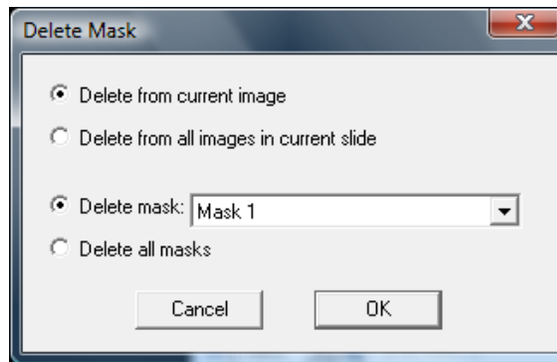
Once segmentation is complete, your entire image will have a blue overlay. SlideBook has selected, or masked, any pixel that has intensity values that lie between the red and green bars in each of the three channels.

5. Select **Cy3** from the drop-down menu.
6. Move the red bar to the right by clicking and dragging it. Note that when you release the mouse button, the image will be segmented again with the new parameters and the mask will be updated.
7. Set the low threshold edit box to “550” and click on **Apply**.



Now SlideBook is masking any pixel with a Cy3 channel intensity value between 550 and 2715.

8. Scroll through the image using the scroll wheel or the up and down arrows in the info/tool bar. Note that the mask persists throughout the entire image.
9. Close the Segment Image dialog box by pressing the **OK** button.
10. Change the red channel from Cy3 to **None**. Observe the mask that is displayed.
11. Click on the mask menu button  in the tool bar and select **None**. The mask has been removed from the display.
12. Select **Mask 1** from the mask menu. The mask is now displayed.
13. To delete the mask, select **Mask > Delete**. The following dialog box will appear.

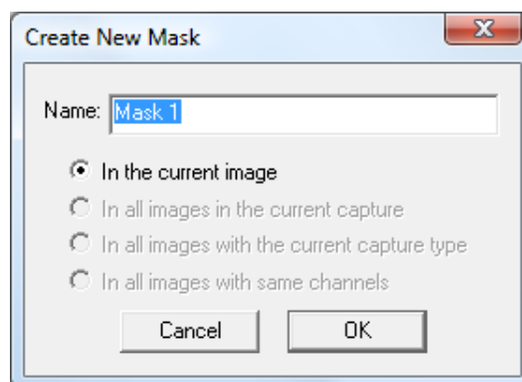


14. Click **OK** to delete the mask from the image.

### 2.7.2 Creating Masks Manually (Background Subtraction Example)

SlideBook also offers a variety of tools that can be used to manually select regions of interest. Note that this example illustrates multiple features of SlideBook.

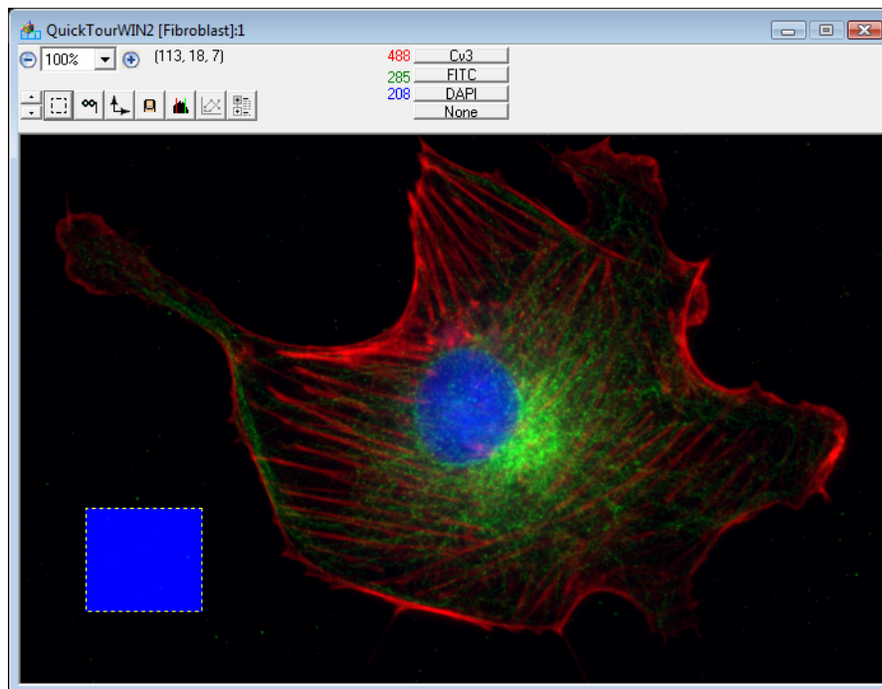
1. Click on the Main View of the “Fibroblast” image.
2. Go to the middle of the focus range by selecting **View > Go to Plane**, entering “7” in the edit box, and selecting **OK**.
3. Next, select **Create Empty Mask** from the mask menu or go to **Mask > Create**. The following dialog box will appear.



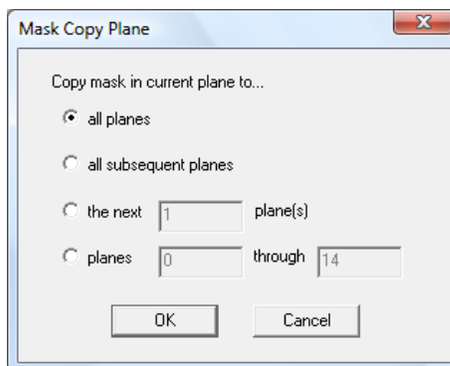
4. Click **OK** to create a new mask for the current image.

Often when you perform analysis, you would like to specify a region to be used for background subtraction. This can easily be done using the marquee tool.

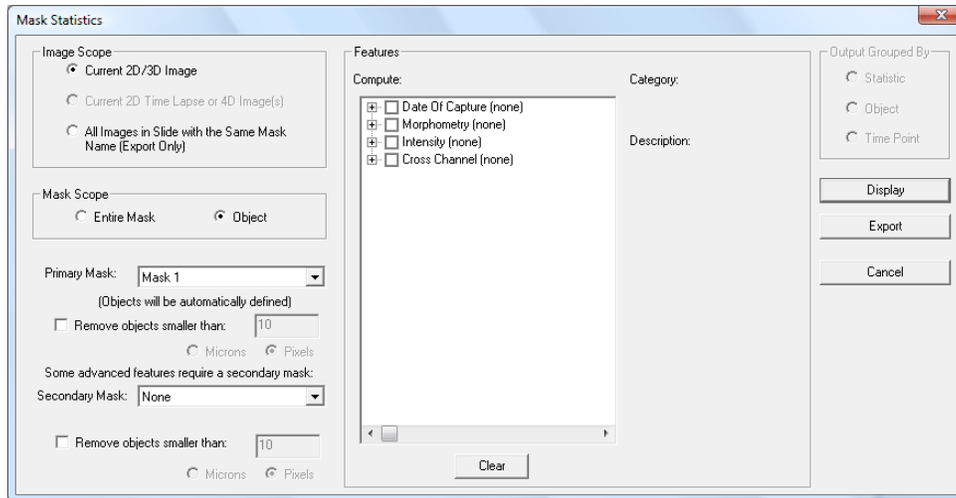
5. Click and drag over a region that would be suitable for calculating the background intensity.
6. Select **Mask > Mark Selection**, and notice that the region is highlighted in blue.



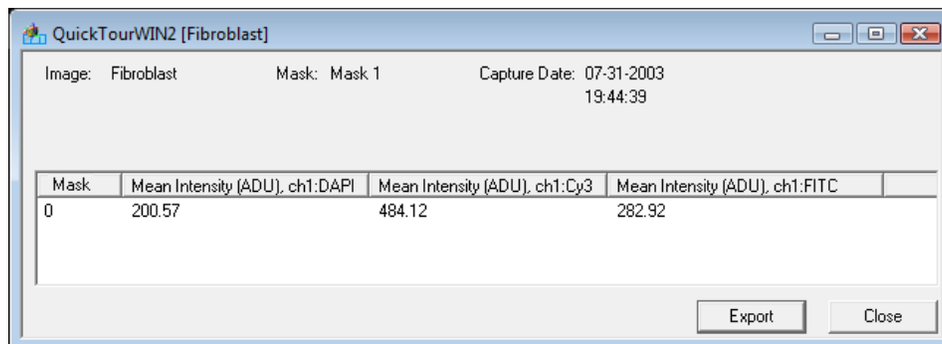
7. This mask only exists in the current plane. To copy this region to multiple planes go to **Mask > Copy This Plane**. The following dialog box will appear:



8. Leave the **all planes** radio button selected and click **OK**.
9. Scroll through the image and note that now the same masked region is selected in all planes.
10. You can use SlideBook to get statistics from masked regions. Go to **Statistics > Mask Statistics**. The following dialog box will appear:



11. Check the **Entire Mask** radio button in the Mask Scope section and then expand the **Intensity** menu in the Features section of the Mask Statistics dialog box by clicking on the **+** to the left of the checkbox. Then check **Mean Intensity**, and then click **Display**. A table will appear that resembles the following.

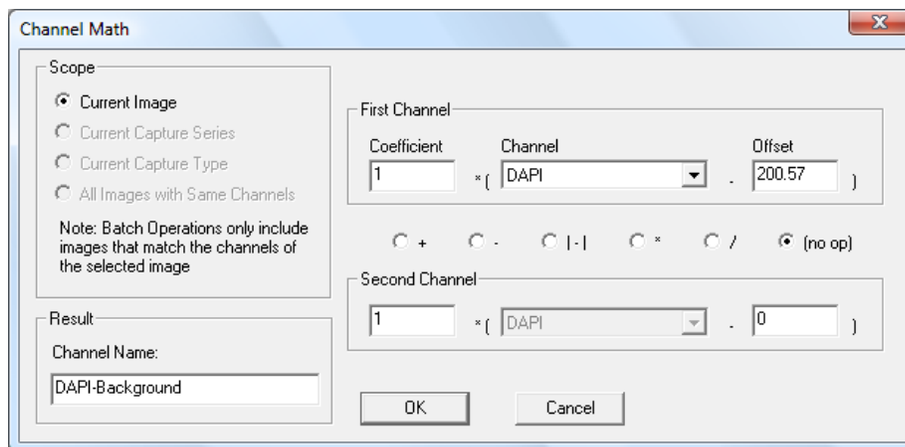


These statistics can be exported as a tab-delimited file by selecting **Export**.

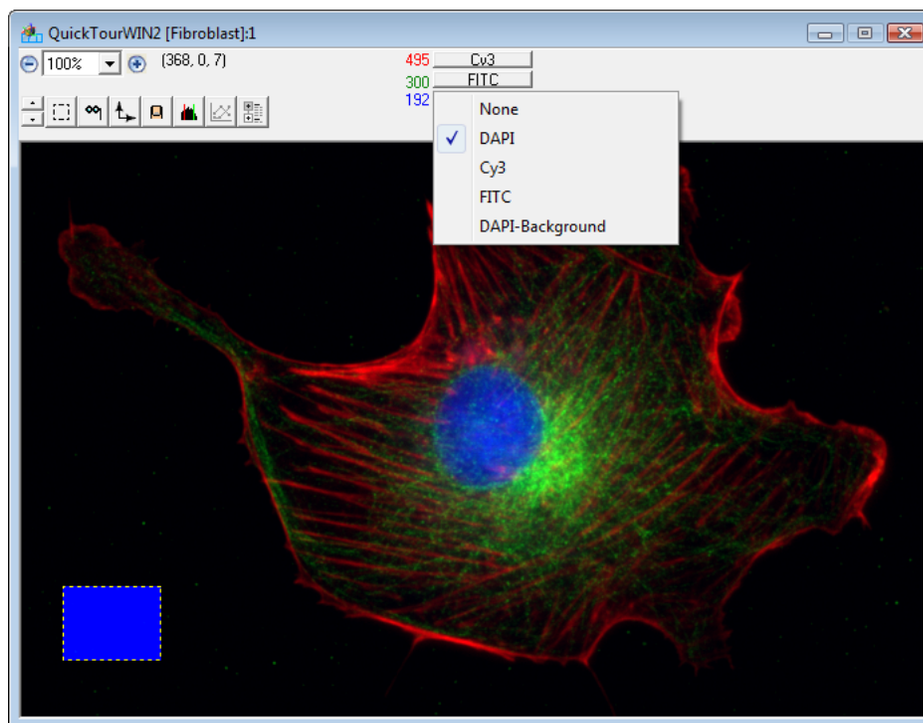
12. To perform background subtraction, choose **Image > Channel Operations > Channel Math**. The following dialog box will appear:



- Enter the mean background intensity value for the given channel (DAPI) to the offset data field for the first channel, then select **(no op)**. Enter a name, such as “DAPI-Background”, for the new channel that will be generated. The dialog box will look like this:



- Select **OK** to create the background subtracted channel. This new channel will now be present in the channel pull down menu within the Main View:

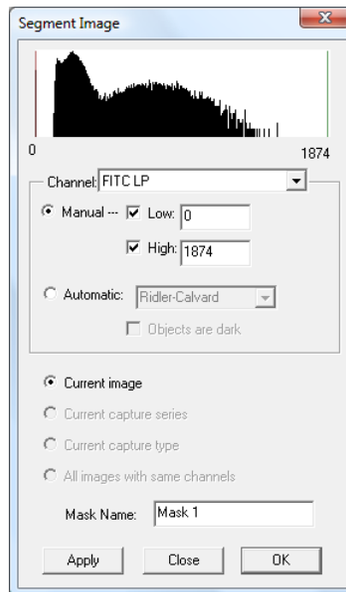


- You may repeat steps 12-14 for each channel by selecting the appropriate channel in the Channel Math dialog box and subtracting the relevant background intensity value. When you are finished, close the **Main View** of the Fibroblast image.

### 2.7.3 Defining Multiple Objects

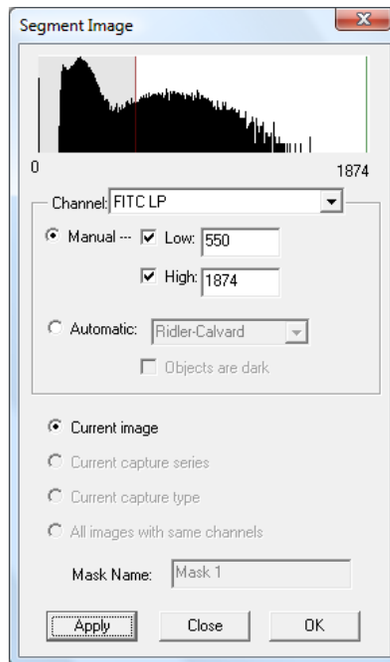
Often you may have several distinct objects that will have intensities that are very similar, and thus will segment together. A mask that contains several distinct objects may be separated into objects, as we will now demonstrate.

1. Open a **Main View** of the “HeLa” image within the QuickTourWIN2.sld by double clicking on the thumbnail.
2. Create a new threshold mask by selecting **Create Segment Mask** from the mask menu. The following dialog box will appear:

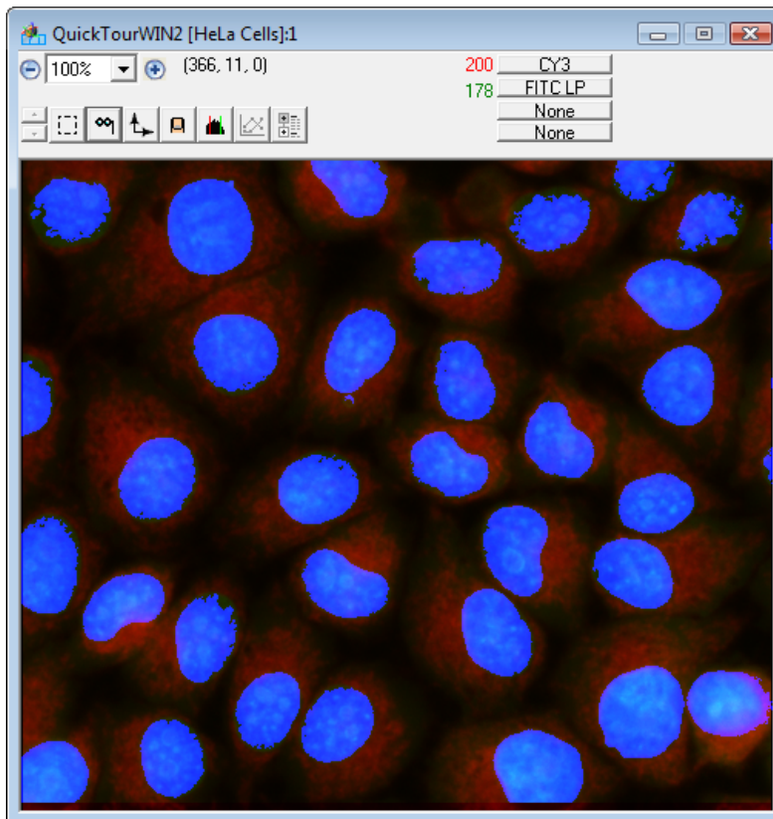


3. Drag the red bar to the right until the low threshold value is approximately 550. As you drag the bar, note the masked areas that appear on the image. Also, note that the data that is being selected is shown in white on the histogram. Pixels with intensity levels that lie in the gray area are not included in the mask.

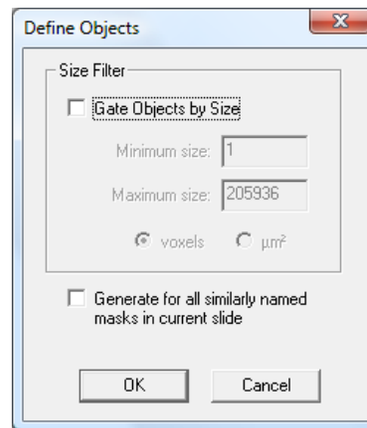
Alternatively, type “550” in the **Low** data entry field and select **Apply** to set the low threshold value. Your dialog box will appear as follows:



4. Select **OK** to create the mask.

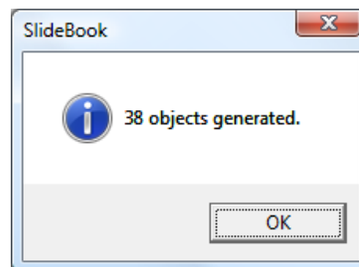


5. Next, select **Masks > Define Objects in Mask**. The following dialog box will appear:

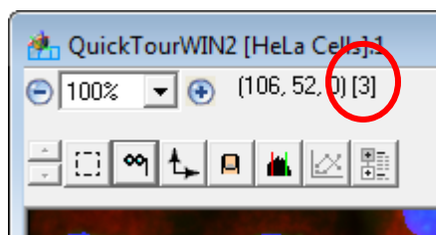


Here you may choose to exclude objects based on minimum and maximum size gates. Leave the box unchecked for now.

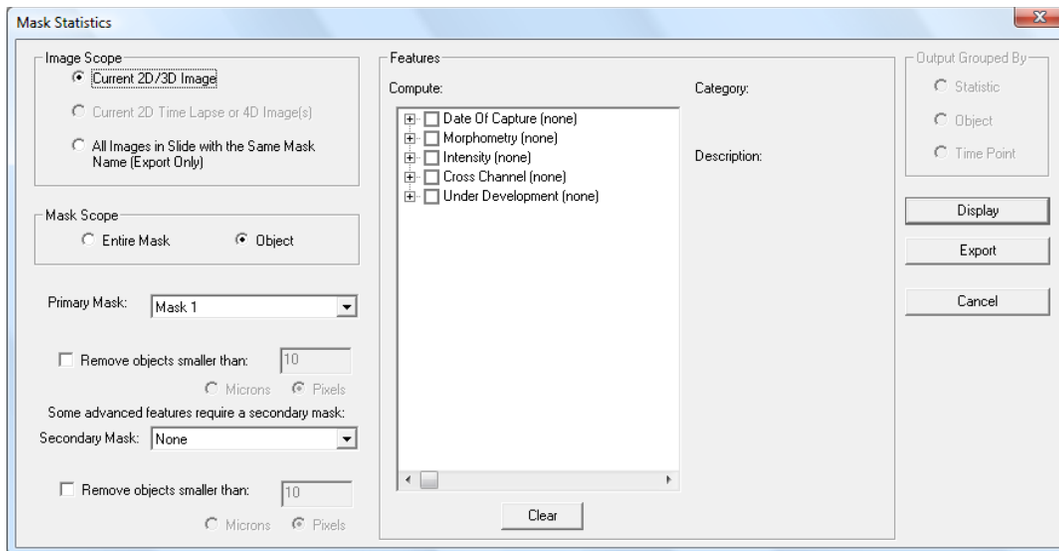
6. Select **OK**. A dialog box will appear that gives the status of the object generation. When object generation is complete, the following box will appear, noting the number of objects generated.



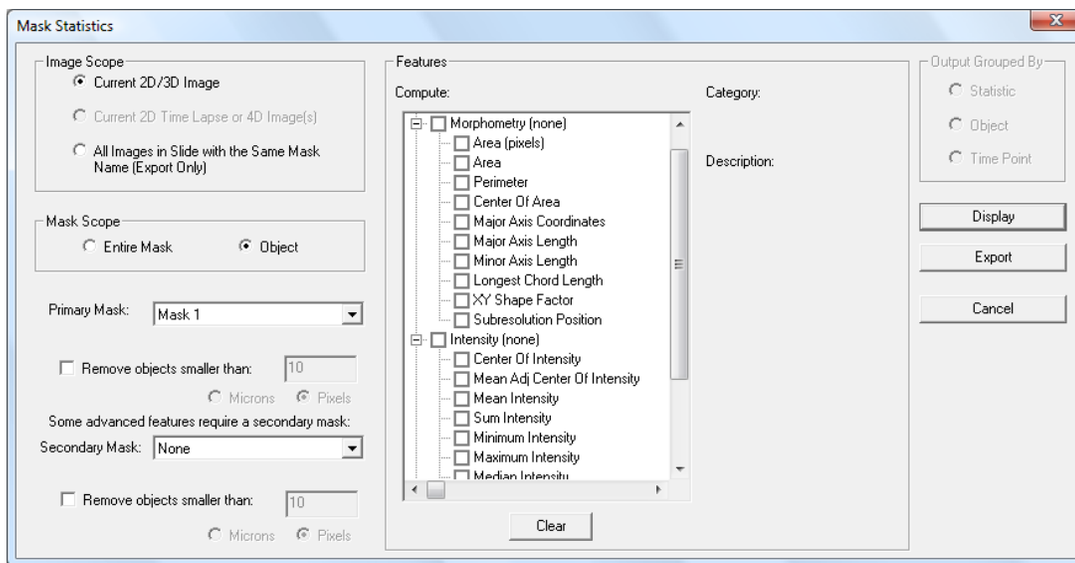
7. Click on **OK**, and now position the cursor over one of the objects in the **Main View**. Note that the object number appears to the right of the cursor location coordinates in the info/tool bar.



8. SlideBook can generate individual statistics for each object. Select **Statistics > Mask Statistics**. The following dialog box will appear:



- Expand the **Morphometry** and **Intensity** statistics selections by clicking on the + signs to the left of the checkboxes.



- Select **Area (pixels)** and **Mean Intensity**, then select **Display** to generate statistics. The following table will appear.

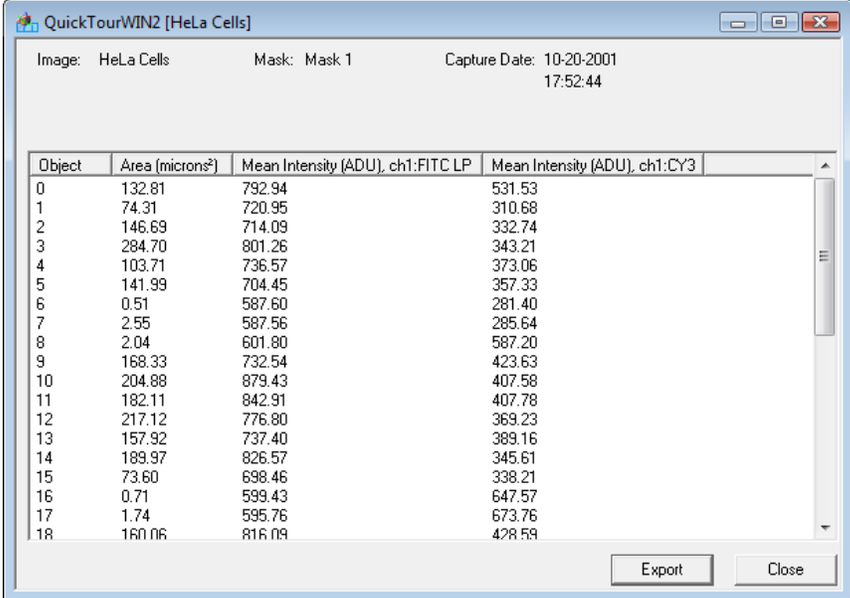
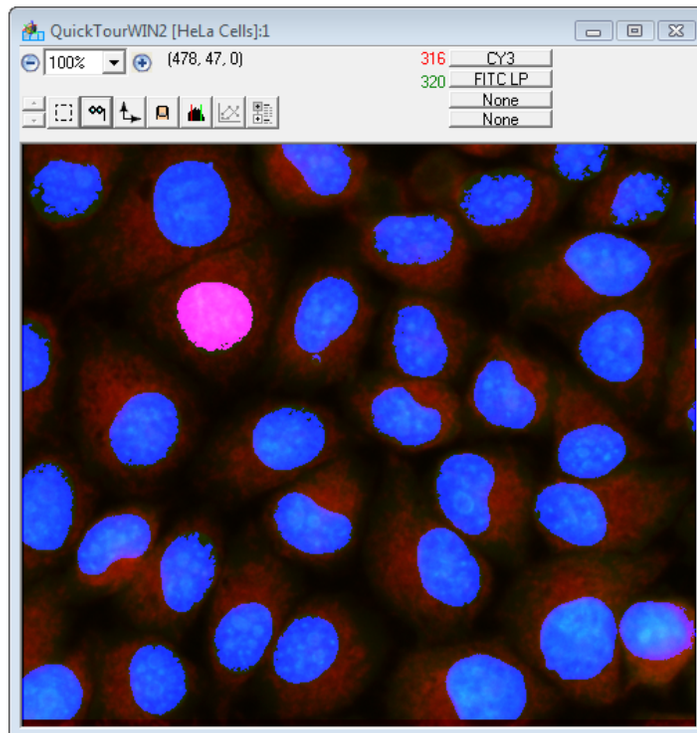


Image: HeLa Cells      Mask: Mask 1      Capture Date: 10-20-2001  
17:52:44

Object	Area (microns <sup>2</sup> )	Mean Intensity (ADU), ch1:FITC LP	Mean Intensity (ADU), ch1:CY3
0	132.81	792.94	531.53
1	74.31	720.95	310.68
2	146.69	714.09	332.74
3	284.70	801.26	343.21
4	103.71	736.57	373.06
5	141.99	704.45	357.33
6	0.51	587.60	281.40
7	2.55	587.56	285.64
8	2.04	601.80	587.20
9	168.33	732.54	423.63
10	204.88	879.43	407.58
11	182.11	842.91	407.78
12	217.12	776.80	369.23
13	157.92	737.40	389.16
14	189.97	826.57	345.61
15	73.60	698.46	338.21
16	0.71	599.43	647.57
17	1.74	595.76	673.76
18	160.06	816.09	428.59

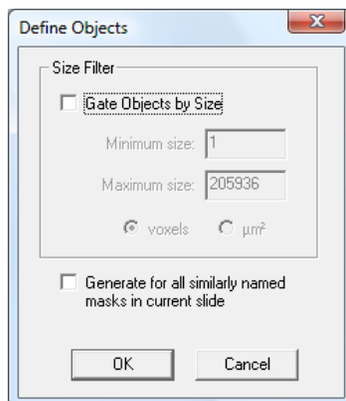
Export      Close

11. Expand the column headers by clicking and dragging the edge of the header to the right. Click on any object number in the table. Notice that the corresponding object in the image is highlighted in pink.



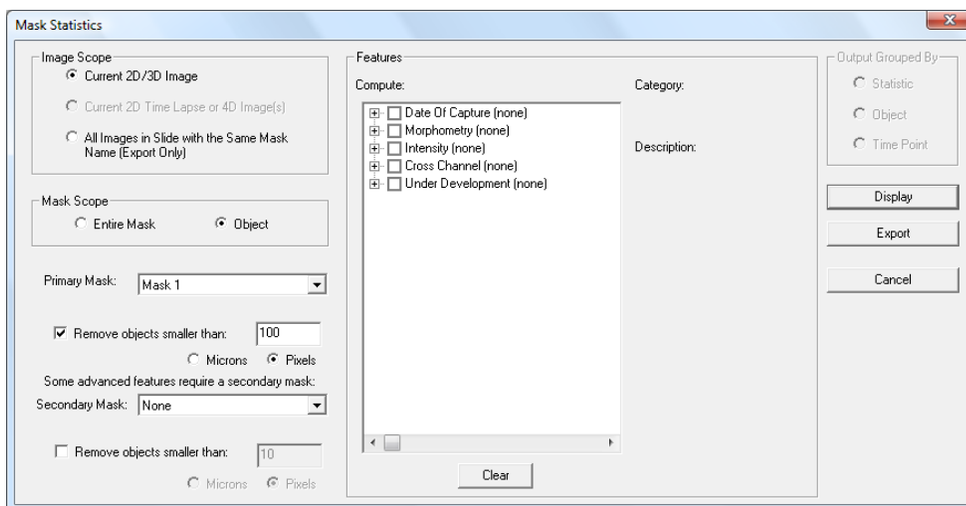
Notice that some of the objects are of very small size. If you would like to exclude those regions when making objects, you may gate the objects by size. This can be done either by going back to the Define Objects dialog box or through the Statistics View itself.

12. To gate objects by size using the Define Objects window, first close the statistics window. Go to **Mask > Update Object Definitions** (Note the menu item is now named “Update Object Definitions” as opposed to “Define Objects” since objects already exist for this image). The following dialog box will appear:



For the purposes of this demonstration, select the **Gate Objects by Size** checkbox and enter a minimum size gate of 100 voxels. This will exclude any objects from the mask that are less than 100 voxels.

13. Select **OK** to update the object definitions. Notice that SlideBook has generated fewer objects than when this operation was performed without a size gate.
14. Statistics can now be generated on these new objects as described above.
15. To gate objects by size using the Statistics window select **Statistics > Mask Statistics**. The following dialog box will appear:



16. Now click the **Remove objects smaller than:** check box located under the **Primary Mask** drop down menu and enter a minimum size gate of 100 in the active edit box.

17. Select the statistics you would like to have computed and click **Display**. SlideBook will automatically remove objects smaller than the minimum size gate during statistics computation.
18. The statistics table may be exported for use in other programs by clicking the **Export** button. A dialog box will appear that will allow you to choose the file destination. The file will be saved as a tab-delineated text file (\*.txt), which can be opened by various programs, such as Excel.
19. Close the slide by clicking on the  in the upper right corner of the slide view or by selecting **File > Close Slide**.

### 3 SlideBook™ Organization

SlideBook was designed to seamlessly integrate microscope control and image operations such as capture, display, and analysis. In this chapter you will learn about the organization of SlideBook elements. The following topics will be covered:

- Hardware Control and Image Capture
- Data Storage, Display, and Analysis

#### 3.1 Hardware Control and Image Capture (Chapters 5, 6 and 7)

In order to capture an image, two distinct processes are performed. First, a sample is brought into view and focus using the *Focus Controls*. Second, image capture parameters are set and capture is initiated in the *Capture* dialog box. Once the capture begins, it can be monitored in the *Capture Controls*.

##### 3.1.1 Focus Controls

In this window, the microscope parameters are selected and moved into position, if motorized. A semi-live stream of images are captured and displayed in the Focus Controls. Additionally, parameters for the XY stage, such as points of interest, and for the Z stage, such as the depth for a 3D capture, can be set in this window. Once the hardware is set as desired and the sample is in focus, the user will move to the Capture dialog box. See Chapter 5 for further explanation of the Focus Window.

##### 3.1.2 Capture Dialog Box

Camera and capture parameters are set in the Capture dialog box. The appropriate exposure time and capture sequence are determined in this window. For instance, the user will decide whether the capture will be two-dimensional, three-dimensional, two-dimensional over time or three-dimensional over time. Parameters set in the Focus Controls, such as multiple points in the  $xy$  plane and  $z$  distances, can be imported into the **Capture** dialog box and utilized during the capture sequence. See Chapters 6 and 7 for further explanation of the **Capture** dialog box.

##### 3.1.3 Capture Controls

The **Capture Controls** window shows the progress of a capture sequence. The features in this window are most useful when performing timelapse acquisition and are discussed in Chapter 7.

#### 3.2 Data Storage, Display, and Analysis (Chapters 8, 9, and 10)

SlideBook's image visualization and analysis tools are designed around five principal objects: *image*, *channel*, *view*, *mask*, *statistics*, and *slide*. This organization is somewhat different than other image processing programs. For instance, with this organization, operations that change data are kept separate from those that only affect its display or analysis.

### 3.2.1 Images and channels

An *image* is a two- or three-dimensional set of data collected by a digital microscope or imported from another application. It consists of one or more *channels*, where each channel corresponds to a different fluorophore. For instance, one channel may represent a FITC labeled protein while another represents a CY3 labeled protein.

An image (and its constituent channels) represents the underlying data. Operations that can alter the underlying data (such as deconvolution, flat field correction, and cropping) are found under the **Image** menu.

### 3.2.2 Views

Unlike images, *views* represent a visual interpretation of the image's data. SlideBook currently has two classes of views: *data views* and *display views*. It is within a view that data can be renormalized (altering the mapping from image intensity levels to red, green, or blue intensities on the display) and colors can be assigned to different channels. You can have multiple views open on a given image at one time. Commands used to create views are found under the **View** menu.

### 3.2.3 Masks and Statistics

A *mask* is a two-, three-, or four-dimensional set of binary values that has the same extent as the image to which it is assigned. Masks are often termed regions of interest (ROIs) in other imaging programs. Masks are used for performing advanced selection and analysis of image data and can either be created and edited manually or generated automatically through threshold segmentation and other techniques. A single image can contain several different masks. Commands used to create and manipulate masks are found under the **Mask** menu. Statistics can be viewed and exported for both masks and image tools from the **Statistics** menu.

### 3.2.4 Slides

Images in SlideBook are organized into *slides*. Typically, a slide corresponds to a number of images captured from a single sample, and it is the native file type saved by SlideBook. By collecting related images into a single slide, SlideBook makes managing files easier. Slides appear on your file system with a `.sld` extension.

## 4 Configuring Your System

In order for SlideBook to control any motorized aspects of your microscope system and to perform deconvolution and analysis of images, it must know the optical and system parameters of the microscope used to collect the image. By the end of this chapter, you will know how to:

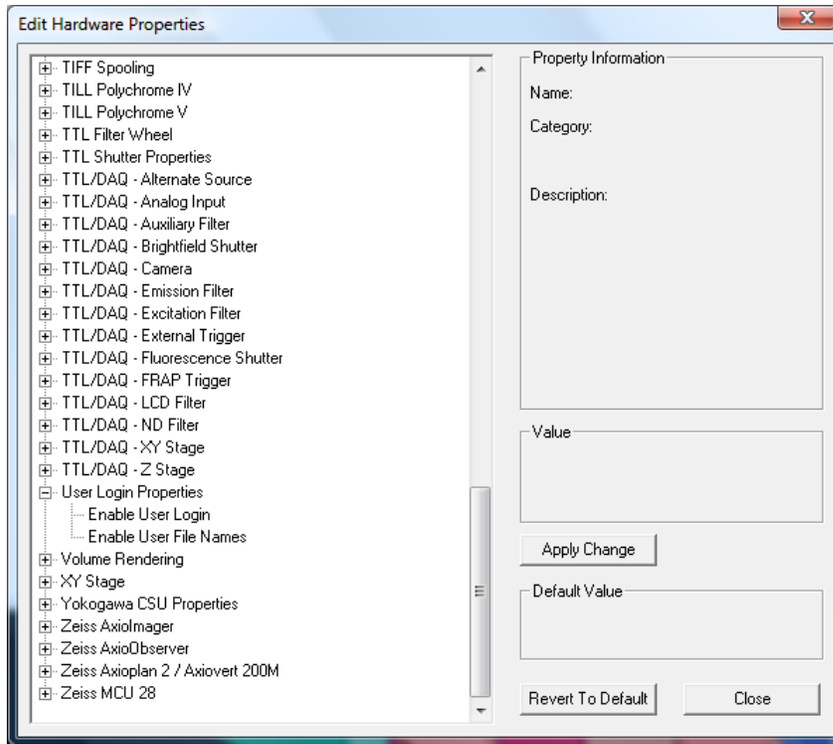
- Designate User Logins
- Configure Hardware
- Define Objectives
- Define Filters
- Define Magnification Changers
- Work with SlideBook Preferences and Hardware Properties
- Define System Parameters

Once the hardware is configured and the optical parameters are defined, they can be utilized in several different ways. If the optics are motorized, they may be selected and brought into the light train using the focus window. Once an image is captured using the selected motorized optics, the capture information is automatically stored with the image. If the optics are not motorized, they may be selected in the capture window and then automatically stored with the image information. Also, optics information can be added or edited post-capture, which is especially useful for imported images.

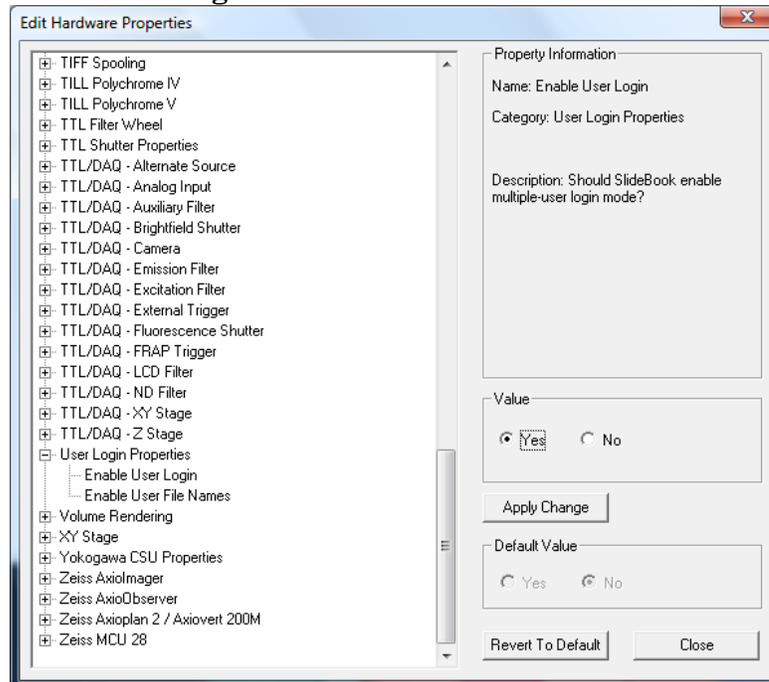
### 4.1 Designating User Logins

You may define multiple user logins to separate user preferences. The user login will allow individual users to access their capture preferences (see [Saving Capture Parameters](#) on page 154). Hardware, filter, and objective configurations are shared among all users. To configure user logins:

1. Go to **Edit > Hardware Properties** and scroll down until **User Login Properties** is visible.



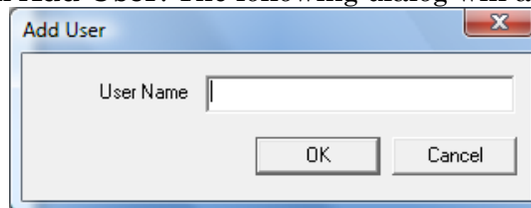
2. Click on the + next to **User Login Properties** to expand the section, and then click on **Enable User Login**.



3. Click **Yes** and then **Apply Change**.
4. Restart SlideBook. Upon startup, you will see the following screen:



5. To add a user, click **Add User**. The following dialog will appear:



6. Enter the desired user name in the **User Name** in the edit field and click **OK**.
7. Upon restart, choose your user name from the drop-down list and then click **Login**. Your capture preferences will be available in the Capture dialog.

## 4.2 Configuring Hardware

SlideBook has the ability to control a range of hardware, including the following:

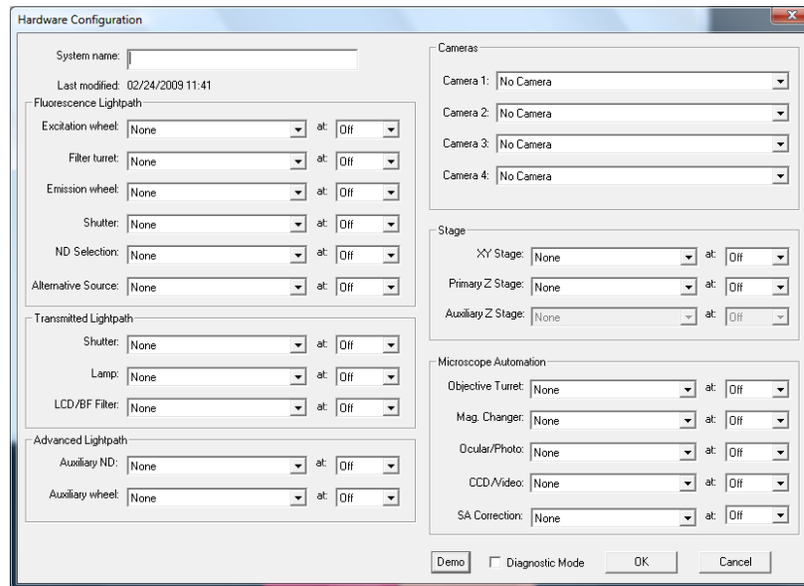
- CCD cameras
- Automated microscope components
  - Objective turret
  - Magnification changer turret
  - Ocular/Photo switch
- Fluorescence lightpath hardware
  - Filter wheels
  - Filter turret
  - Shutter
- Automated *xy* and *z* stages
- Image Splitters
- Transmitted (brightfield) lightpath hardware
  - Shutter
  - Lamp
  - LCD filter
- TTL controllable hardware

When you start SlideBook for the first time, it will notice that a *preferences file*, which is stored along with the application, does not exist. SlideBook will create a default preferences file. As you configure your hardware and define optics, SlideBook will update this preferences file with specific information about your setup.

### 4.2.1 General Hardware Configuration

To configure your hardware:

1. Start SlideBook by selecting the program from the **Start** menu.
2. Select **Edit > Hardware Configuration**. The following dialog box will appear:



3. Name the system in the **System name** edit field. This should be descriptive and include the laboratory name so that the preferences file can be identified by 3i technical support.
4. Select the components that are present from the drop-down menus (set to **None** in the dialog box above).
5. For each component that is controlled through a COM port (or USB port mapped to a COM port), select the COM port to which it is connected in the drop-down menus to the right of each component. If a component is not present, or you would not like it to communicate with SlideBook, be sure the COM port is set to **Off**.

**NOTE:** A typical PC has two serial COM ports, port 1 and port 2, located at the rear of the machine. Additional COM ports are often added (e.g., through use of an Edgeport adapter or PCI card). These additional ports are then typically termed port 3, port 4, etc. instead of following the labels on the adapter.

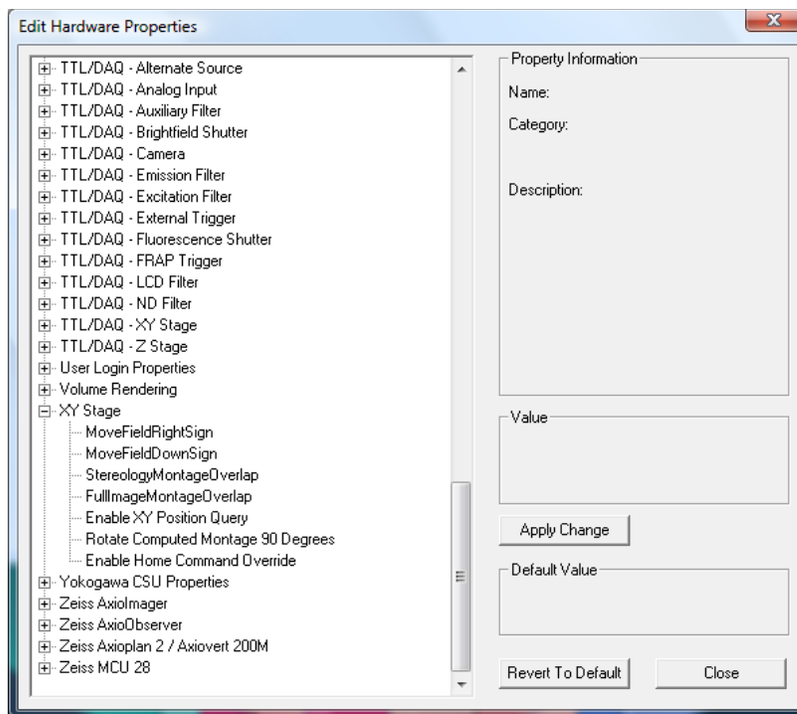
6. Select **OK**.
7. Restart SlideBook to register the changes and begin hardware communication.
8. If you plan to control your hardware via TTL control, please consult the SlideBook TTL Module Manual.

**NOTE:** There are instances when you will need to select your objective and filter in the Focus Window, even if these components are not motorized. For example, if you would like to perform auto focus you will need to select the objective and filter you are using. In these instances, you will define Manual Filter Turret or Manual Objective Turret (**COM** port set to **OFF**). Make sure that you specify the position of your objectives in the Objective Definition section (see page 62).

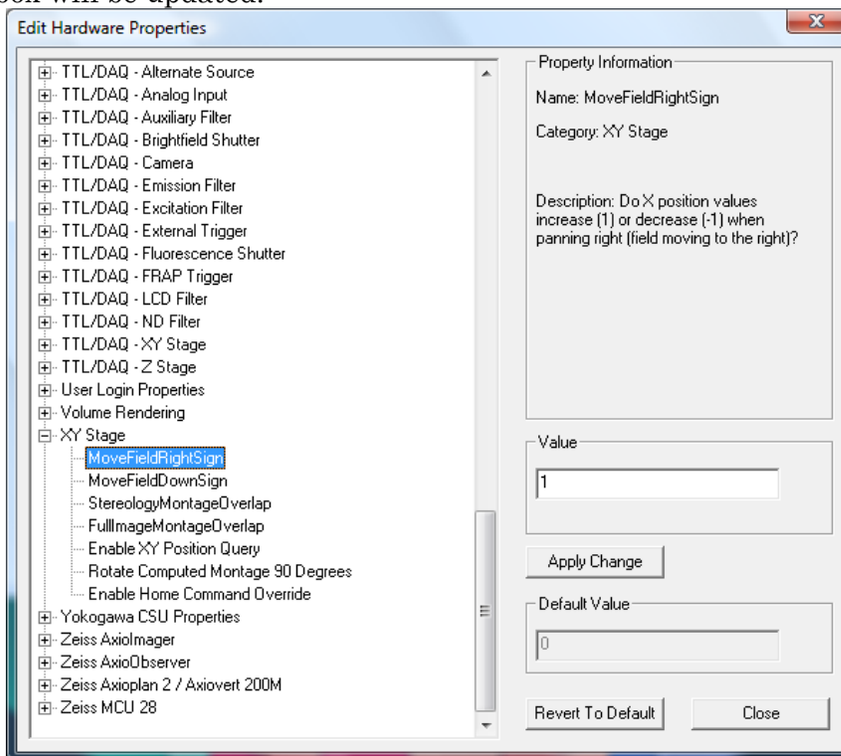
### 4.2.2 Configuring a Motorized XY Stage

In order for your motorized *xy* stage to work properly when collecting montage images, you will need to specify the relationship between the stage and camera.

1. To do so, you will first need to bring a sample into view and focus in the focus window (see Using the Focus Window on page 85).
2. Using the joystick, move the stage so that the image on the screen moves to the left (in other words, pan and move the image field to the right). When panning to the right, notice whether the *x* coordinate position in the **XY Stage** section of the Focus Controls window increases or decreases.
3. Repeat for the *y* axis by panning down so that the image appears to move up. Note the change in the *y* axis coordinate position (increasing or decreasing) when panning.
4. Select **Edit > Hardware Properties**. The following Edit Hardware Properties dialog box will appear.
5. Click on the + sign next to **XY Stage** to expand the list.



- Click on the item in the list titled **MoveFieldRightSign**. The left side of the dialog box will be updated.



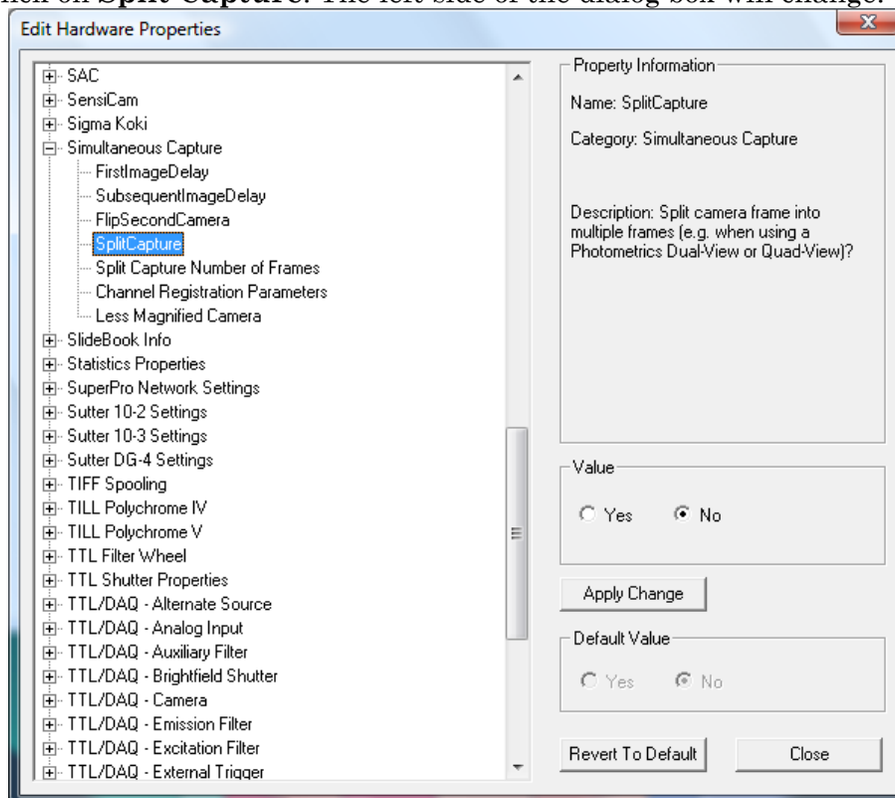
- Enter the appropriate value, 1 or -1 based on the change in x position values when panning right, in the edit field and select **Apply Change**.

- Repeat steps 6 and 7 for the *y* axis by clicking on the **MoveFieldDownSign**.
- Close the dialog box using the **Close** button.

### 4.2.3 Configuring an Image Splitter

SlideBook supports the use of image splitter devices. An image splitter allows you to capture two separate channels for the right and left halves of the camera. To configure an image splitter:

- Select **Edit > Hardware Properties** and click on the **+** sign to expand the **Simultaneous Capture** section.
- Click on **Split Capture**. The left side of the dialog box will change.



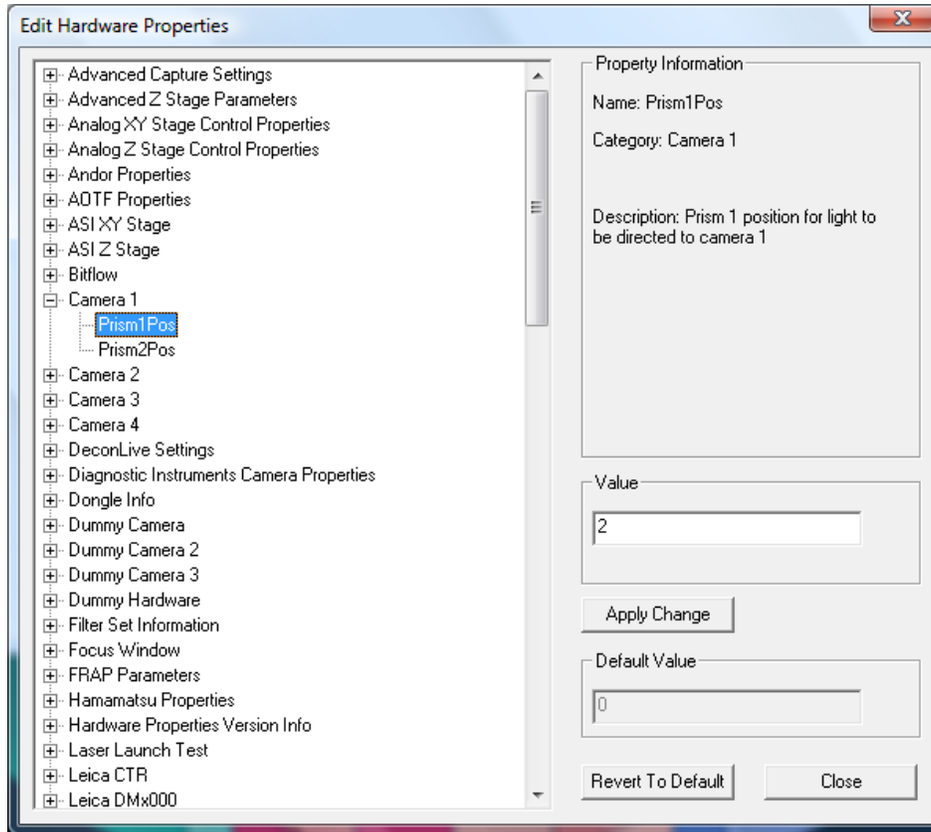
- Select **Yes**, **Apply Change**, and then **Close**.
- Restart SlideBook to register the changes.

To use the image splitter, you must define filters appropriately (see [Defining Filters when using an Image Splitter](#) on page 72). To align your image splitter, use the **Test Dual-View** function in the Focus Window as described on page 85. To capture images using your image splitter, use the **Simultaneous Capture** checkbox as described on page 133.

#### 4.2.4 Configuring Camera and Ocular Ports

If your system has motorized ocular photo and CCD/Video prisms as well as multiple cameras, you may define the positions of the prisms that correspond to each camera. To do so:

1. Select **Edit > Hardware Properties** and expand the section titled **Camera 1**.
2. Click on **Prism1Pos**. The right side of the dialog will be updated.



3. Enter the value of the ocular photo prism (Prism 1) position that corresponds to the position of Camera 1 in the edit field.
4. Select **Apply Change**.
5. Repeat steps 4 and 5 for **Prism2Pos** (CCD/Video prism).
6. Repeat steps 2 through 5 for any other cameras and the ocular lightpath.

Once you have defined the prism positions, you can click on any of your cameras in the **Camera** tab and the prisms will move to direct light to that camera. Additionally, if you have defined filters for specific cameras, the prism positions will move to the appropriate camera when you select a filter in the Focus Controls.

### 4.3 Defining Objectives

Once an objective is defined, it can be controlled (if the objective turret is motorized) from the focus window. If the objective turret is not motorized, it may be selected in the capture window and the objective definition will automatically be stored with the image information. If the objective was used to capture an image that has been imported, the objective information can be added to the image information.

#### 4.3.1 Adding a New Objective

In order to define a new objective for your system, do the following:

1. Select **Edit > Define Optics > Objectives**. The following dialog box will appear:

2. To add a new objective definition, click on the **Add** button.
3. Enter the objective name, magnification, numerical aperture, working distance, and  $\mu\text{m}/\text{pixel}$ . You must give the lens a name and fill out all parameter fields, except those in the **Parfocality and Parcentricity** and **Transmitted Light Configuration** groups.

NOTE: For optimal results, objective pixel size should be measured by traditional means using a stage micrometer. If this is not possible, a close estimate can be calculated by dividing the camera pixel size by the magnification of the objective. Please note that this will not take into account any optics that alter magnification such as the Spherical Aberration Correction device by 3i.

4. Select the location of objectives from the **Position** drop-down menu. If you have a motorized objective turret, **Position** should reflect the position of the objective

on the microscope; if you are using SlideBook offline or with a non-motorized turret, you should select **Unmounted**. If you are using a non-motorized turret, but have selected **Manual Turret** in your hardware configuration, select the appropriate turret position.

5. Select the **UV** checkbox for UV objectives and select the appropriate medium from the radio button menu.
6. Select the appropriate configurations for the desired transmitted light condenser turrets using the drop-down menus on the right. Once you have defined the condenser positions for a given objective, you may define filters that utilize the appropriate condenser positions. Filter definitions are discussed in the next section.
7. Repeat steps 1 through 6 for any additional objectives that you wish to add.
8. Select **OK** when you are finished.
9. Restart SlideBook to register the changes that you have made.

### 4.3.2 Modifying Information for an Existing Objective

You can modify existing settings as follows:

1. Select **Edit > Define Optics > Objectives** to bring up the Objective Parameters dialog box.
2. Select the appropriate lens using the **Objective Lens** dropdown list.
3. Change the appropriate fields as desired.
4. Select **OK** to save the changes.
5. Restart SlideBook to register the changes that you have made.

### 4.3.3 Removing an Objective

1. Select **Edit > Define Optics > Objectives** to bring up the Objective Parameters dialog box.
2. Select the appropriate lens using the **Objective Lens** dropdown list.
3. Choose **Remove** then **OK** to save the changes.
4. Restart SlideBook to register the changes that you have made.

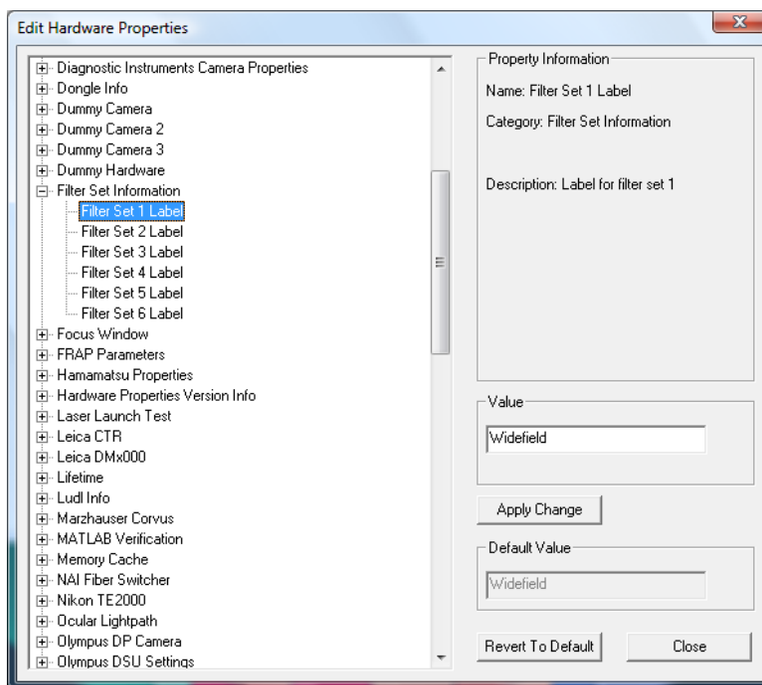
## 4.4 Defining Filter Configurations

You can add, modify, and remove filter configurations using the **Filter Configuration Parameters** dialog box. The components of this dialog box and instructions for adding, modifying and removing filter configurations are explained below.

### 4.4.1 Defining Filter Sets

SlideBook allows you to assign a particular filter to one of six groupings, or *filter sets*. These sets can be chosen in the focus window and capture window. The default sets are **Fixed**, **Live**, **User 1**, **User 2**, **User 3**, and **User 4**. Sets allow you to work with many different filter definitions in an orderly fashion. To change these default group names:

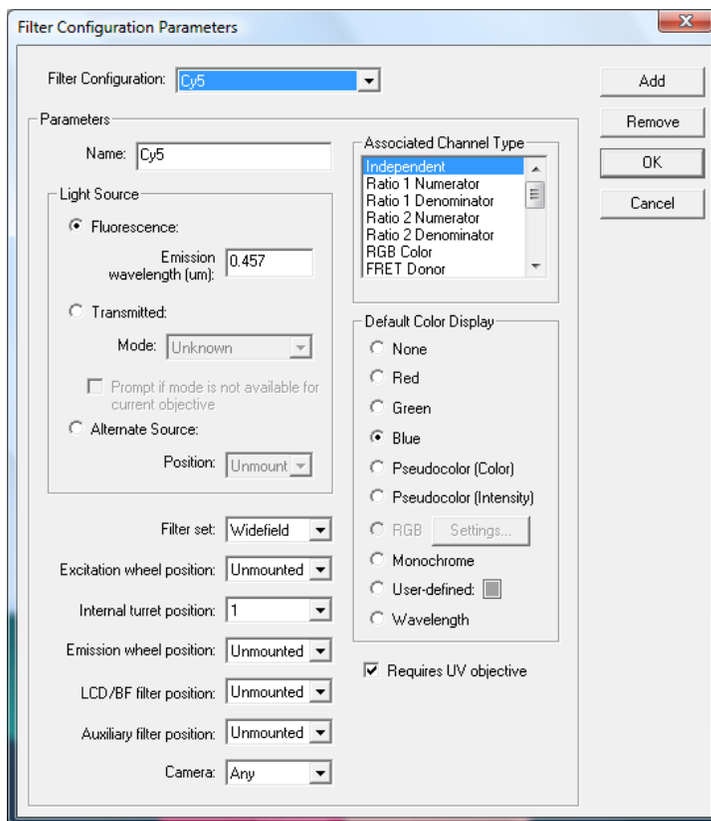
1. Select **Edit > Hardware Properties**. The Filter Configuration Parameters dialog box will appear.
2. Scroll down to find the property **Filter Set Information** and click on the + sign to expand as shown below:



3. Click on the **Filter Set Label** that you wish to change.
4. Enter the text for your filter set. Examples may be a specific user name or a type of experiment.
5. Click **Apply Change** and then **Close** to register your changes.

## 4.4.2 Filter Configuration Parameters

The Filter Configuration Parameters dialog box is found by selecting **Edit > Define Optics > Filter Configurations**.



### 4.4.2.1 Light Source

Here you may choose which light source will be used. If your microscope has an automated brightfield lamp, the lamp will be switched on and off (or shuttered) for each brightfield image that is acquired. Also, if you have a motorized condenser turret, you may select whether a transmitted light channel is brightfield, darkfield, DIC (Nomarski), or phase. If you check the **Prompt if mode is not available for current objective** checkbox, SlideBook will automatically warn you if objective and filter definitions are inconsistent. For fluorescent channels, input the emission wavelength in microns in the edit field. You should also indicate whether the fluor requires a UV objective. SlideBook will automatically check for consistency between objective type and fluorescent channel type and prompt you if there is a problem.

**NOTE:** You **MUST** have condenser turret positions defined for objectives if you would like the condenser turret position to correspond to a specific transmitted light channel (see [Adding a New Objective](#)).

### 4.4.2.2 Filter Set

SlideBook allows you to assign a particular filter to one of six groupings, or sets. The default sets are **Widefield, Confocal, Transmitted, TIRF, Life Time, and FRAP**. You may also edit these set names as shown on page 64. These sets can be chosen in the Focus

Controls and the Capture Controls. Sets allow you to work with many different filter definitions in an orderly fashion.

### 4.4.2.3 Filter Positions

SlideBook permits filters to be defined in up to five positions within the optic path. For systems equipped with filter wheels or a rapid filter switcher (such as the Sutter DG-4 or TILL Polychrome V), use the **Excitation Wheel Position** and **Emission Wheel Position** fields. If your system has a motorized filter cube turret, you must specify the **Internal Turret Position** for the filter configuration's cube. If your system does not have a motorized filter cube turret, you should still enter the position of the cube. Then, when you select the filter from the Focus Controls, SlideBook will prompt you to move the turret to the appropriate position. Some examples of how to use filter positions are described in the section [Adding a New Filter Configuration](#) on page 68.

**NOTE:** If you have a manual filter turret, SlideBook will prompt you to move the beam splitter when performing multi-channel capture. In cases where you do not want a prompt for a different beam splitter (or in the motorized case, you do not want the internal turret to move), you can simply set **Internal Turret Position** to **Unmounted**. If you believe you are seeing a prompt in error, you may wish to review your filter configuration to make sure that your positions are defined correctly and also check your hardware configuration to ensure that any motorized elements are defined properly.

Finally, if you have an LCD color filter as part of your system, the **LCD/BF Filter Position** field can be set to select positions for individual colors.

### 4.4.2.4 Camera

SlideBook supports use of multiple cameras. If you have more than one camera, you may select the camera that will be used for capturing the given filter configuration.

### 4.4.2.5 Channel Types

All fluorescent filter configurations that are not used for ratio imaging should be set to **Independent**. Configurations for ratio and FRET channels are discussed in the Ratio/FRET Module manual. Brightfield channels that use a color camera should be set to **RGB Color**. Brightfield channels that use an LCD color filter, or three separate color filter cubes, should be set to **Virtual RGB (red), (green), or (blue)**.

### 4.4.2.6 Default Color Display

You can always change the colors (or color display type) of data captured with different filter configurations. The **Default Color Display**, however, is a convenient way to choose how data from different filter configurations is displayed in the capture status dialog box as well as the initial view created for an image. Channels may be set to standard RGB, pseudocolor, monochrome, user-defined, or set to the emission wavelength.

#### 4.4.2.6.1 Pseudocolor

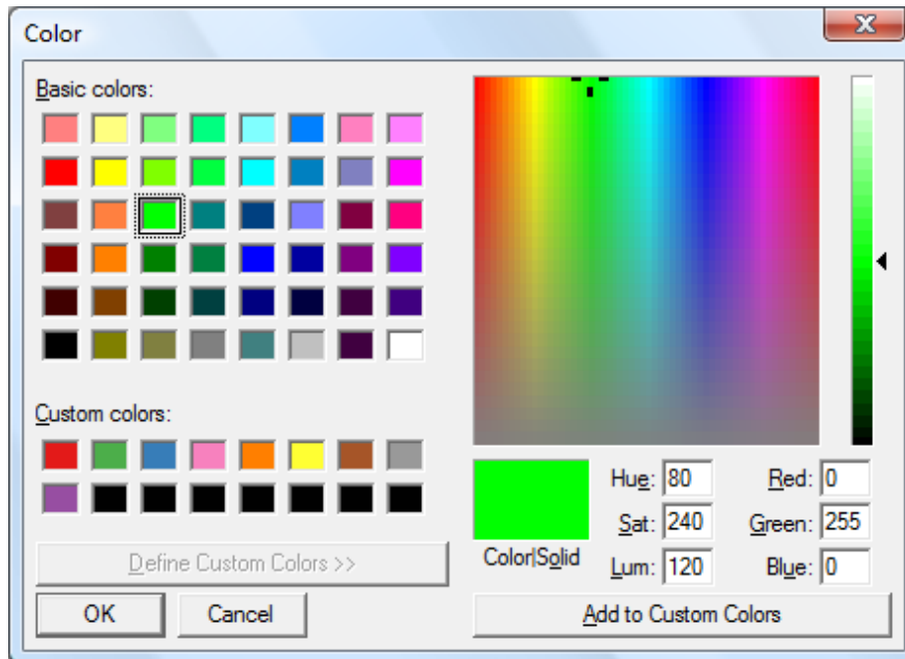
**Pseudocolor (Color)** and **Pseudocolor (Intensity)** together determine the look of a pseudocolored image. A filter configuration that has **Pseudocolor (Color)** as its default will set the hue value of the display according to the normalized (i.e., scaled from minimum

to maximum) value of each voxel. A filter configuration that has **Pseudocolor (Intensity)** as its default will gate the intensity of the pseudocolored image according to the normalized value of each voxel. If a captured image contains a channel with a filter configuration set to **Pseudocolor (Color)** but not **Pseudocolor (Intensity)**, the corresponding view will show a range of hues at maximum intensity (i.e., ranging from bright blue to bright red). If the image also contains a channel with a filter configuration set to **Pseudocolor (Intensity)**, the corresponding view will show a range from dark blue (black) to bright red. If the image only contains a channel with a filter configuration set to **Pseudocolor (Intensity)** and no **Pseudocolor (Color)** counterpart, the default setting will be treated as if it were set to **None**.

Pseudocolor color types cannot be displayed along with red, green, or blue channels. Monochrome settings may only be used for a background channel when displayed with RGB channels. In multiple channel images, the pseudocolor setting will override any red, green, or blue settings. Red, green, or blue settings will in turn override a monochrome setting. Again, this doesn't mean you can't easily get back to viewing any other channels, but simply determines what will be visible initially.

#### 4.4.2.6.2 User-Defined Color

You may elect to set the default color to one of your choosing. To do so, select the **User-Defined** radio button, and then click on the color block to the right. A dialog box similar to the following will appear.



Select one of the basic colors by clicking on the color block in the upper left hand color, or design a custom color using the color spectrum on the right and choose **Add to Custom Colors**.

### 4.4.3 Adding a New Filter Configuration

1. Select **Edit > Define Optics > Filter Configurations**. The Filter Configuration Parameters dialog box will appear.
2. To add a new filter configuration definition, click on the **Add** button.
3. Enter the filter name in the data entry field.
4. Select the light source, and enter the emission wavelength in  $\mu\text{m}$  (for fluorescent filters). Also, if the filter requires a UV objective, select the checkbox under the emission wavelength data field. The **Light Source** field permits sequential brightfield (or Nomarski) and fluorescence imaging on systems equipped with both a fluorescence shutter and either a brightfield shutter or electronic voltage control.

**NOTE:** You may configure multiple transmitted light channels if you have a motorized condenser turret. For instance, you may define a DIC channel by selecting **Transmitted** light source and **DIC** from the **Mode** dropdown menu. You could also define a phase channel by creating a new channel with **Phase** selected from the **Mode** drop-down menu. Then, if you performed capture with either of these channels, the condenser turret would move to the appropriate position automatically. You must define the condenser turret positions when defining objectives for this feature to work properly (see section [Adding a New Objective](#)).

5. Select the desired **Filter Set** from the dropdown menu.
6. If the exciter is located in an external excitation wheel, select the position of the filter from the **Excitation Wheel Position** dropdown menu. If it is mounted in a cube in the internal turret or if it is not currently mounted on the system, select **Unmounted**.
7. Select the internal turret position of the dichroic mirror from the **Internal Turret Position** dropdown menu. If the dichroic mirror is not currently mounted, you should select **Unmounted**.
8. If the emitter is located in an external emission wheel, select the position of the filter from the **Emission Wheel Position** dropdown menu. If it is mounted in a cube in the internal turret or if it is not currently mounted on the system, select **Unmounted**.
9. Repeat steps 1 through 8 for any additional filter configurations that you wish to add.
10. Select **OK** when you are finished.
11. Restart SlideBook to register the changes that you have made.

#### 4.4.3.1 Example Filter Configuration for Fluorescence Channels

*Example – Quad Pass and Dual Pass sets on a system with filter wheels and motorized turret*

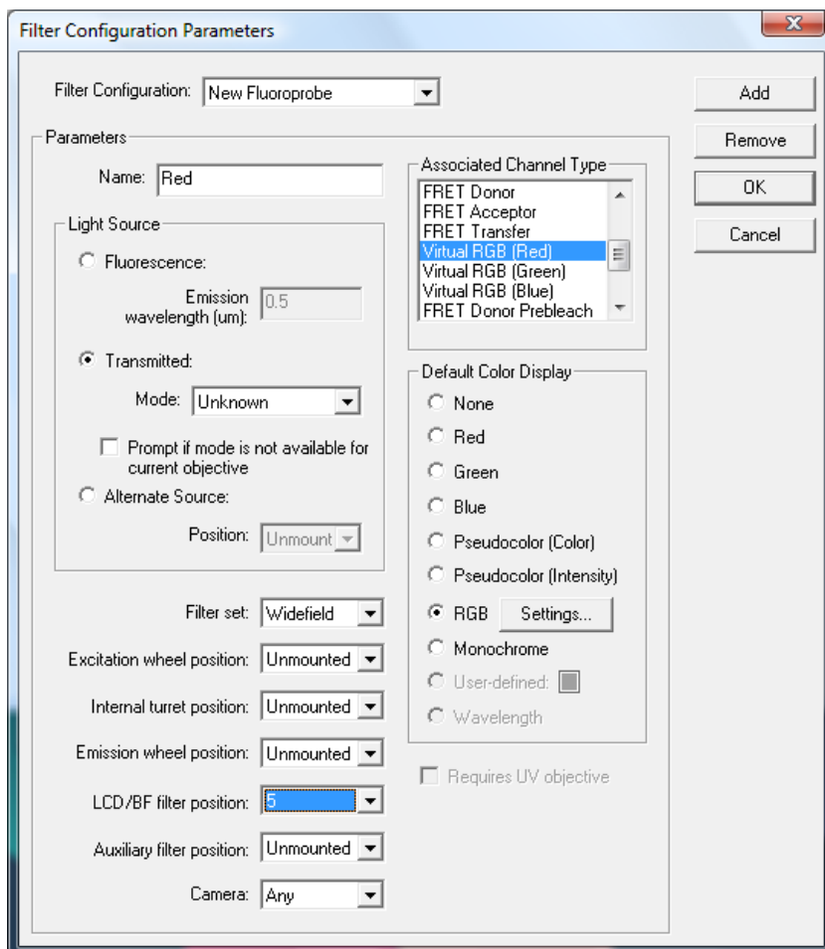
Some systems with filter wheels will need different beam splitters to be in position for different sets of fluorophores. For instance, a system may have excitation and emission filter wheels loaded with filters for DAPI, CFP, FITC, YFP, CY3, and CY5. The DAPI, FITC, CY3, and CY5 filters require a quad pass beam splitter designed for that set of fluorophores, while CFP and YFP require a dual pass beam splitter which is designed specifically for those two fluorophores.

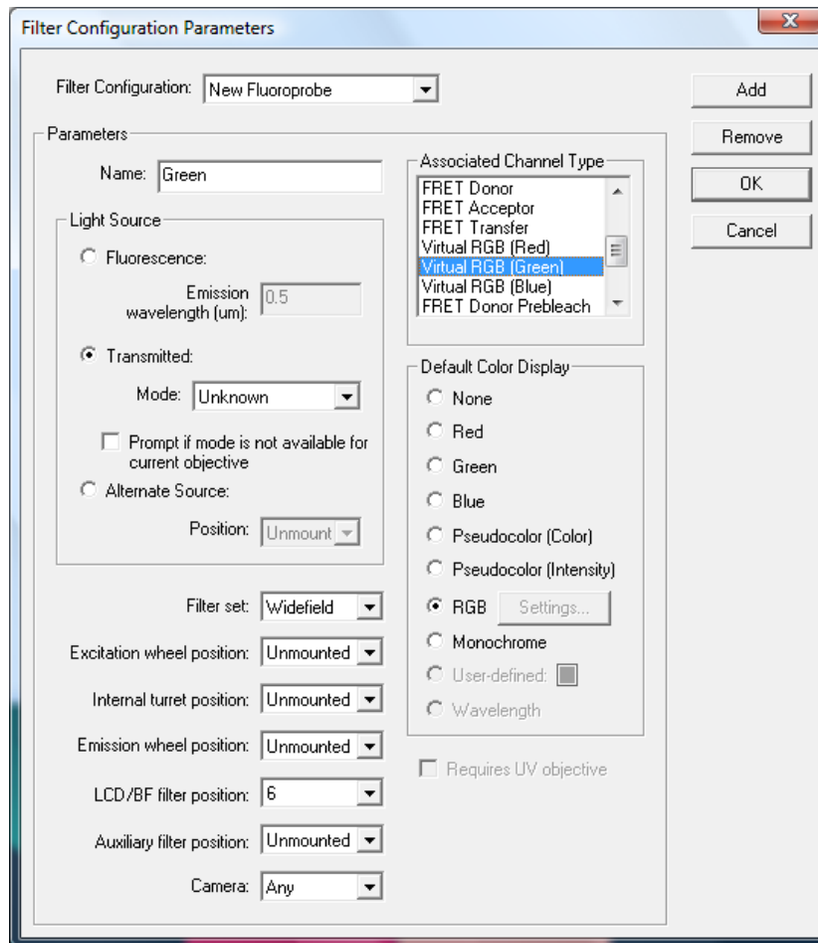
In a system with filter wheels and a motorized turret, **Internal Turret Position** should be set to the appropriate beam splitter (or dichroic) position for each filter configuration. In a system with filter wheels but no motorized turret, the **Internal Turret Position** field should be set differently for each set of fluors that require a different beam splitter if you want SlideBook to prompt you to move to the necessary beam splitter. Given the example above, even if the system does not have a motorized turret, the filter configurations for DAPI, FITC, CY3, and CY5 should have **Internal Filter Position** set to say, 1, while CFP and YFP have **Internal Filter Position** set to 2.

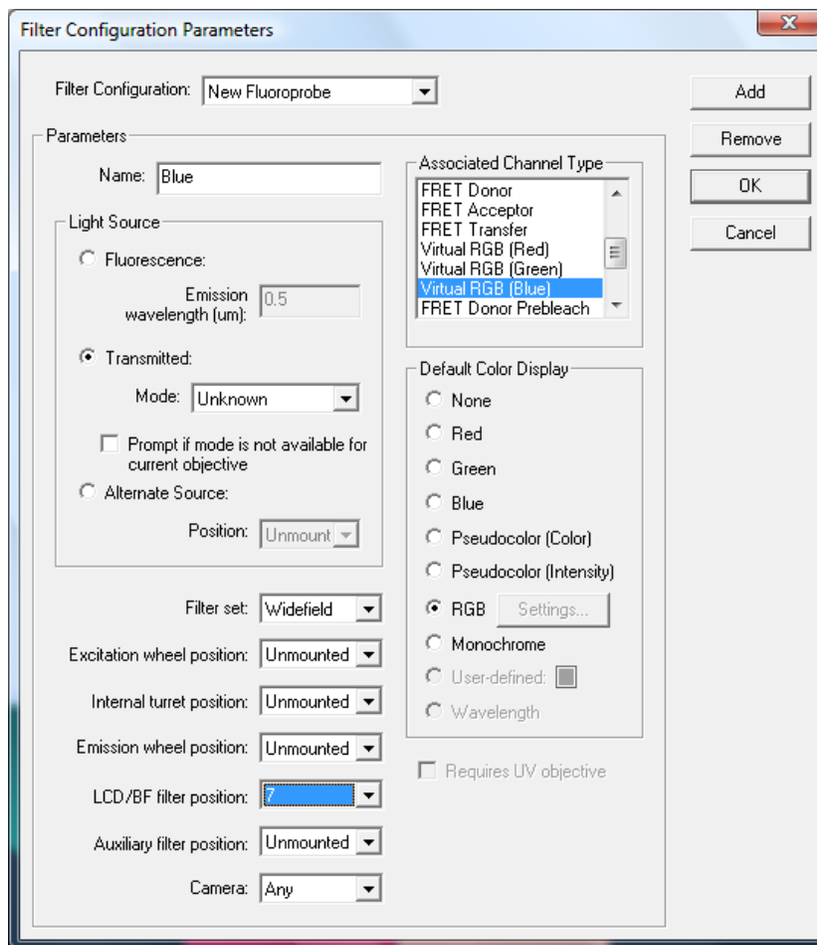
#### 4.4.3.2 Configuring Channels for Color Cameras or Color LCD sliders

If you have a color camera, you will need one filter definition that appears as follows (you must enter the appropriate location of an empty turret and emission wheel position if applicable):

If you have a LCD slider, such as that made by CRI, you will need to configure three separate channels; however, these three channels will appear as one channel in the capture dialog. You must define one channel each for red, green, and blue. For the color LCD slider, the filter definitions must appear as follows. Create the definitions in the following order: red, blue, green. This is because you will most often perform autofocusing with the green channel, and autofocusing will be performed with the last channel captured. For the red channel, you may also choose “Red” as the **Default Color Display**. RGB Settings allows you to manually enter white balance values. You may find that using the white balance function in the capture dialog is sufficient (please see [Performing Auto-White Balance for Color Capture](#) on page 95).







#### 4.4.3.3 Defining Filters when using an Image Splitter

If you have configured an image splitter, such as the Dual-View from Optical Insights, you should define two filters that have identical excitation wheel positions (if applicable) and identical internal turret positions. Then, depending on the position of the image splitter, select one channel **Camera** position as **Camera 1**, and the second channel as **Camera 2**. For instance, if you are performing CFP/YFP FRET, you may wish to define four filter configurations:

1. YFP excitation, YFP emission,
2. YFP excitation, CFP emission (this is a dummy filter that must be configured, but will not contain information),
3. CFP excitation, YFP emission,
4. CFP excitation, CFP emission.

In this example, filters 1 and 3 will use **Camera 1**, while filters 2 and 4 will use **Camera 2**. Also, filters 1 and 2 will use the same excitation position, as will filters 3 and 4. You must define the filters in order so that the channels within each pair that share an exciter are adjacent. In the example above, channels would be defined in order of 1, 2, 3, 4.

#### 4.4.4 Modifying Information for an Existing Filter Configuration

1. Select **Edit > Define Optics > Filter Configurations**. The **Filter Configurations Parameters** dialog box will appear.
2. Select the appropriate filter using the **Filter Configuration** dropdown list.
3. Modify the appropriate fields as desired.
4. Select **OK** to save the changes.
5. Restart SlideBook to register the changes that you have made.

#### 4.4.5 Removing a Filter

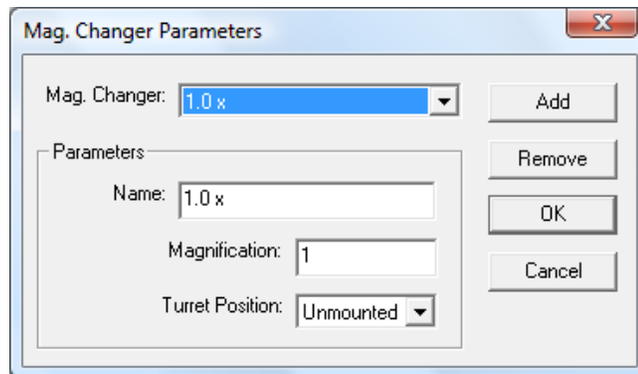
1. Select **Edit > Define Optics > Filter Configurations**. The **Filter Configurations Parameters** dialog box will appear.
2. Select the appropriate filter using the **Filter Configuration** dropdown list.
3. Choose **Remove**, then **OK** to save the changes.
4. Restart SlideBook to register the changes that you have made.

### 4.5 Defining Magnification Changers

A magnification changer, or optovar, is a component which your microscope may or may not have. It acts to increase the effective magnification of an objective. If you do not have a magnification changer, you should use the predefined “1.0 x” definition.

#### 4.5.1 Adding a New Magnification Changer

1. Selecting **Edit > Define Optics > Mag. Changers**. The **Mag. Changer Parameters** dialog box will appear:



2. Click on the **Add** button.

3. Enter a **Name** in the data entry field, and the **Magnification** value.
4. Click **OK** to save the definition.
5. Restart SlideBook to register the changes that you have made.

### 4.5.2 Removing or Modifying Magnification Changer Definitions

You may remove and alter magnification changer definitions as you would for objectives and filter configurations.

Note that if you switch between several camera adapters with different relay lens magnifications, you can also use magnification changer definitions to specify which relay lens is in place.

## 4.6 Working with SlideBook Preferences and Hardware Properties

SlideBook stores two different types of configuration files, *User Preferences* and *System Preferences*. User Preferences refers to settings that are user-specific such as saved capture settings or saved particle tracking protocols. System Preferences refers to system wide settings such as hardware configuration and hardware properties. There are several tools built into the software to help you manage and backup preference files as discussed below.

### 4.6.1 Backing Up SlideBook Configuration

1. To back up your SlideBook configuration, go to **Edit > Setup Guides > Backup Preferences**, the default behavior is to save the backup files in `C:\Program Files\Intelligent Imaging Innovations, Inc\SlideBook 5.0`.
2. Select where you would like the backup files to be located. If desired, create and name a new folder.
3. Click **OK**.
4. Three folders containing system configuration will be placed in the directory designated. The folders are as follows:
  - **Exceptions** – This folder contains backup data from interrupted sessions and will be discussed later in this section.
  - **Global Preferences** – This folder contains the system configuration files such as `SlideBookHardwareProperties.dat`, `SlideBookPrefs.dat`, `PSFs.dat`, and `FlatFields.dat`.
  - **Users** – This folder will contain files specific to individual users or the default user. This is where saved capture settings, point lists, and analysis protocols are saved.

### 4.6.2 Restoring SlideBook Configuration

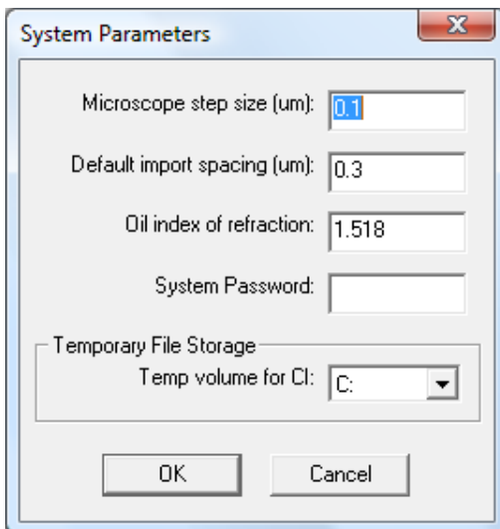
1. In order to restore your system preferences, you must first open the backup configuration folder. This file may be located at C:\Program Files\Intelligent Imaging Innovations, Inc\SlideBook 5.0 or a user-designated location as specified in Step 2 above.
2. Double-click on the **Global Preferences** folder and select all of the files and folders. Copy the data by right-clicking and selecting **Copy** from the menu.
3. Now, return to SlideBook and go to **Edit > Setup Guides > Open System Preferences Folder**.
4. Click inside of the Windows Explorer folder view that opens. Right-click and select **Paste** from the menu.
5. You will be informed that there is already a file in this location with the same name. Select the option to overwrite all files or folders and click the box to apply to all files and continue.
6. You have now replaced the active system configuration files with the backup copies. Now you will restore the user preferences.
7. Go back to the backup configuration from Step 1.
8. Double-click on the **Users** folder and select all of the files and folders. Copy the data by right-clicking and selecting **Copy** from the menu.
9. Now, return to SlideBook and go to **Edit > Setup Guides > Open User Preferences Folder**.
10. Click inside of the Windows Explorer folder view that opens. Right-click and select **Paste** from the menu.
11. You will be informed that there is already a file in this location with the same name. Select the option to overwrite all files or folders and click the box to apply to all files and continue.
12. Close all folder views and restart SlideBook.

### 4.6.3 Exceptions

**Exceptions** is a folder that is backed up with your system and user configurations. This folder may contain a number of folders that begin with “ExceptDump”. These files are created when a process is interrupted in SlideBook. If your SlideBook session is interrupted and your data was not saved, you can recover the data by looking in the **Exceptions** folder. It may be useful to monitor this directory and delete unnecessary files.

#### 4.7 Defining System Parameters

You can adjust the  $z$ -axis increment size, the default interplane spacing, the index of refraction of the oil you are using, and the disk drive on which to put temporary files using **Edit > Define Optics > System Parameters**:



For microscope automation to work correctly, you need to enter the size of a single  $z$ -axis increment of the stage (in microns) in the **Microscope Step Size** field. When you import files from other applications, SlideBook will assume a default interplane spacing (in microns) that is defined in the **Default Import Spacing** field.

## 5 Controlling the Camera and Microscope Hardware (*Focus Window*)

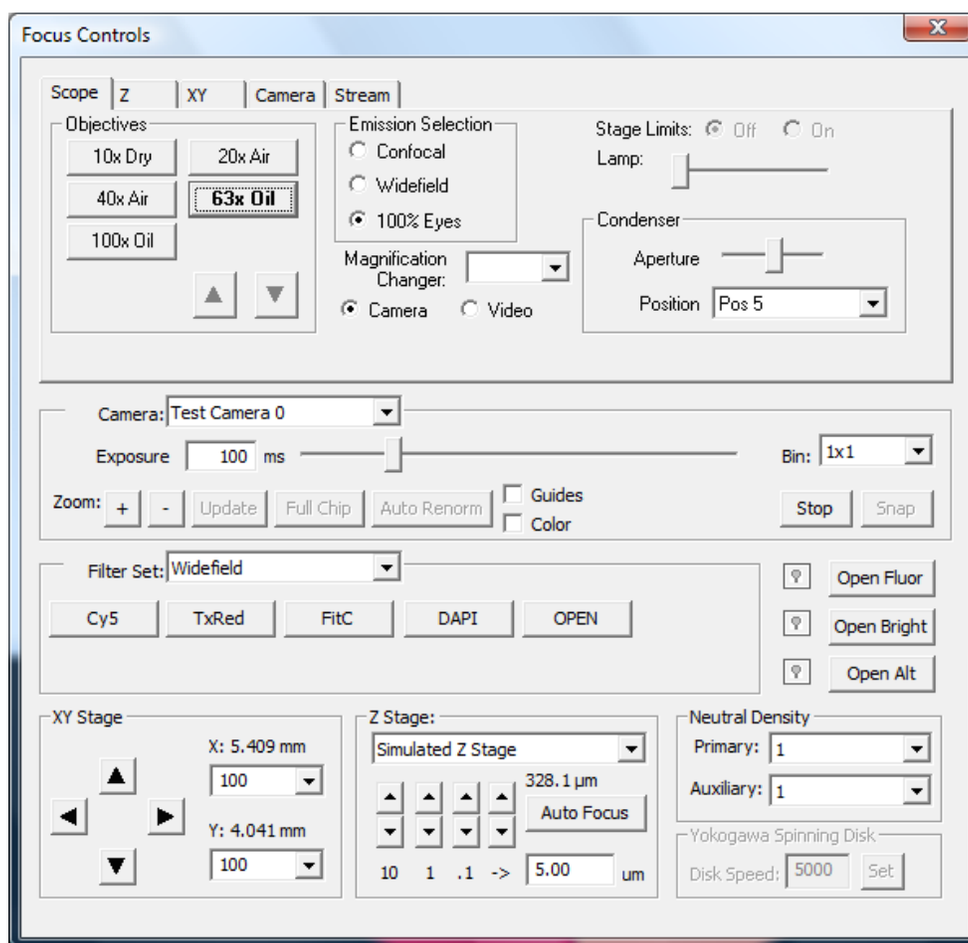
SlideBook provides a Focus Window for automated hardware control and semi-live camera readout. The Focus Window will allow you to choose the appropriate optics and bring your sample into focus. It will also allow you to set parameters for advanced capture, including top and bottom  $z$  limits for 3D imaging,  $xy$  coordinates for multipoint acquisition, and boundaries for a generating a montage. This chapter covers the following topics:

- Focus Window Features
- Using the Focus Window

The use of the Focus Window for advanced capture is discussed in detail in Chapter 7.

### 5.1 Focus Window Features

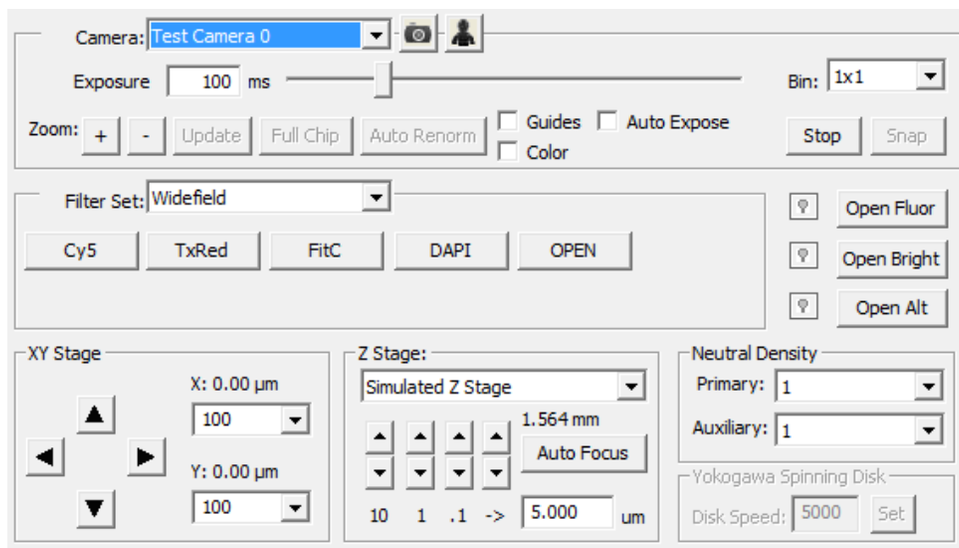
The Focus Window can be opened by either selecting **Window > Focus Window** or clicking on the focus window button in the toolbar . The Focus Window consists of a Focus Controls dialog box and the Live Window, which operates in a live mode when the Focus Window is selected. The Live Window is also used for capture as discussed in Chapters 6 and 7.



The Focus Controls dialog box always has at least two tabs: **Scope**, and **Camera** that control and report the status of automated microscope components and cameras. Typically systems also have either a **Z** tab or an **XY** tab or both. Additionally, there are a number of permanent controls in the lower half of the dialog box that are displayed no matter which tab is selected.

### 5.1.1 Permanent Controls

The lower half of the Focus Controls dialog is present regardless of the tab selected.



#### 5.1.1.1 Camera and Display Settings

The **Exposure** time slide bar allows you to select the duration of a single camera exposure (between 1 and 500ms). This exposure is repeated continuously to form the semi-live image display. You can also select between four bin factors, all of which are based on square pixel groupings (i.e., 1x1, 2x2, etc). Binning increases camera sensitivity and readout speed. This is useful in the focus window because it allow shorter exposure times and faster image refresh rates, ultimately making it easier to find a sample and bring it into focus.

Importantly, the focus window exposure time and bin factor adjustments are used only for optimizing image display in the focus window. Adjusting the camera settings for capturing an image is covered in Chapter 6 on page 89.

On systems with multiple cameras and motorized camera port selection, the **Camera** drop-down menu lets you direct the light to the desired camera and display the resulting image. On systems with two cameras running simultaneously from a single port, these buttons let you select which camera's output will be displayed in the Focus Window.

The **Start/Stop** button starts (or stops) continuous camera capture. This can be particularly useful if you have a mechanically shuttered camera since the focus window can be left open without taking unnecessary camera exposures. You may also press **Snap** to grab a single image that will be stored in an open slide.

The  and  buttons in the bottom left corner are used to zoom in and out on the image display. Zooming in on the display can be particularly useful for focusing on small objects.

Selecting the **Guides** check box will bring up yellow guides that mark a  $256 \times 256$ ,  $512 \times 512$ , and  $1024 \times 1024$  pixel region in the image. The **Color** check box is used to display a color brightfield image in the focus window. Use of this feature requires specific hardware for capturing color images, such as an LCD color filter changer or color camera.

### 5.1.1.2 Open Fluor, Open Bright, Open Alt

The **Open Fluor** button is used to toggle the fluorescence light source if there is a computer-controlled shutter or dark slider in the reflected light path.

The **Open Bright** button is used to toggle the brightfield light source if there is either a computer-controlled shutter in the transmitted light path or the microscope has computer control over the transmitted light source voltage.

The **Open Alt** button is used to toggle an alternate source if it is present. For example, if you have a NEOS AOTF laser system, or a shuttered TIRF laser, **Open Alt** will toggle the laser light on and off.

### 5.1.1.3 Filter Controls

You may select a filter set using the filter set drop-down menu. Each button corresponds to a filter configuration that you have defined for the system. (For more information on creating new filter configurations, please refer to the section [Defining Filter Configurations](#) on page 64). If filter configurations have been assigned to a particular filter set, they can be accessed using the drop-down menu. Depending on the configuration of your microscope system, clicking on a filter configuration tab will change one or more of the following:

- excitation filter wheel position
- emission filter wheel position
- internal filter turret (or slider) position
- camera that is used for display (when the system has multiple cameras)

The filter configuration that is currently selected will be highlighted and displayed in bold letters, while the rest of the configurations will have a normal button appearance.

### 5.1.1.4 XY Stage

On systems equipped with a motorized *xy* stage, you can use the directional buttons to move the stage in increments specified by the value in the adjacent edit field. This value is always given in microns (or fractions thereof). You can change this increment distance at any time. The defaults are 100, 10, 1 and FOV (Field of View is an integer value defined by the pixel array of the CCD) which can be selected from the drop-down menu.

**TIP:** If you double-click on an object in the image display, the *xy* stage will move so that the object is centered.

The current *x* and *y* positions read from the stage are also displayed in this section.

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### 5.1.1.5 Z Stage

On systems equipped with motorized *z* focusing, you can move the focus up and down using the directional buttons. There are three sets of buttons with predefined stage movement increments (10, 1, and 0.1 microns). In addition, an user-defined stage movement increment value can be entered in the edit field below the last set of buttons on the right. The default value of the edit field will be set to the theoretical *z* resolution for the selected objective.

With few exceptions, the up and down buttons correspond to the direction of mechanical movement. For instance, on an upright microscope with either an integrated or external stepper motor, the direction corresponds to stage movement, so “up” means focusing further past the cover slip and towards the slide since the stage is moving up. On an inverted microscope, “up” corresponds to moving the nosepiece up, and therefore also means focusing further past the cover slip.

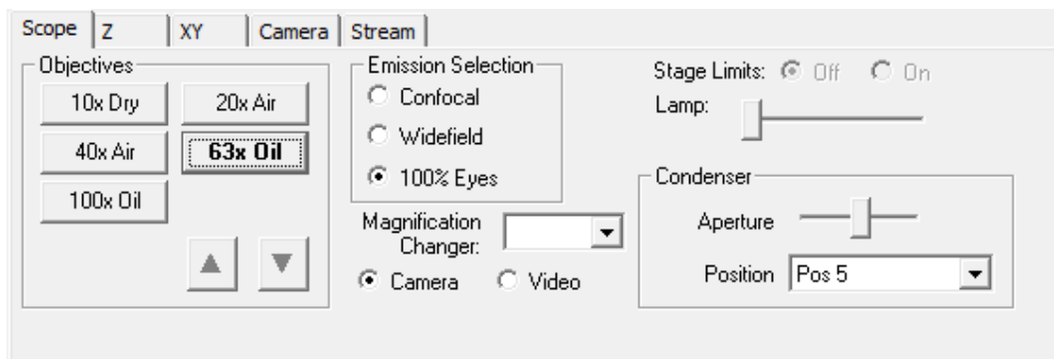
The **Auto focus** Button is used to define and test parameters for auto focusing during capture. Use of this feature is covered in Chapter 7.

### 5.1.1.6 Neutral Density

If you have defined neutral density positions in your hardware configuration, you may select the position from the **Neutral Density** drop-down menu.

## 5.1.2 Scope

The **Scope** tab lets you observe and control other encoded and automated features of your system.



### 5.1.2.1 Emission Selection

On systems with a motorized emission light path, these radio buttons let you choose between letting the light go 100% to the camera, 100% to the oculars (eyepieces), or to 50% each (if available).

### 5.1.2.2 Magnification Changer

On systems with an encoded magnification changer, this field reports its current position.

### 5.1.2.3 Objectives

On systems with a motorized objective turret, you can move to a different objective by selecting the corresponding button. (For information on adding objectives to those defined for your system, please refer to the section [Defining Objectives](#) on page 62).

If you try to move between objectives that use differing immersion media, such as an oil objective to an air objective, SlideBook will bring up a dialog box warning about the pending move.

**CAUTION:** On most systems with motorized objective turrets, you can safely move between objectives of similar immersion media (e.g., air, oil). However, there are situations where you will definitely not want to move between objectives automatically. For instance, if you are using a water immersion objective on an upright microscope, where the objective is dipped into the chamber, trying to move the objective could break the chamber or damage the motorized turret.

### 5.1.2.4 Stage Limits

On certain microscopes with motorized  $z$  focus drives, stage limits are used to control the maximum and minimum position of the stage. These limits prevent the microscope from being focused above or below certain  $z$  positions. The **Stage Limits** radio buttons are used to enable or disable this feature.

### 5.1.2.5 Lamp

On microscopes with a computer controlled lamp voltage adjustment, the intensity of the brightfield illumination can be controlled using this slider. Moving the slider to the far left corresponds to turning the lamp off, while moving the button to the far right corresponds to turning the lamp to maximum intensity.

### 5.1.2.6 Condenser

On microscopes with an automated condenser turret, you may select condenser positions from the drop-down menu.

## 5.1.3 Z

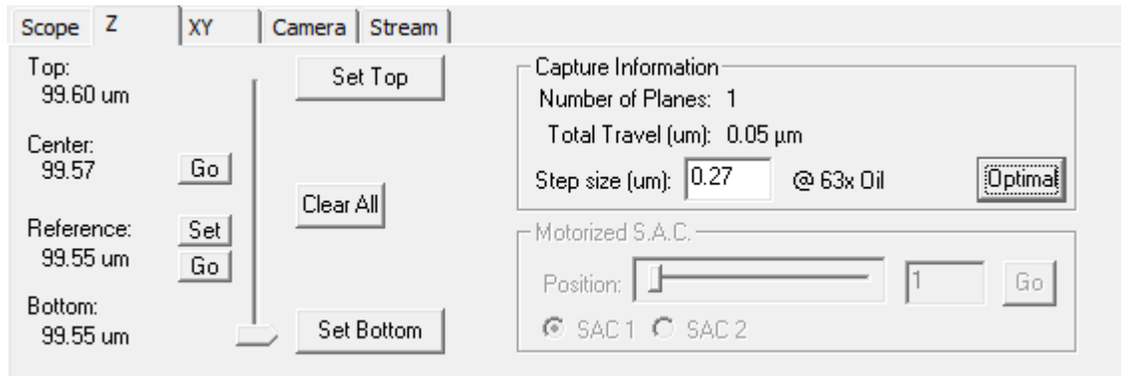
There are several possible ways to set up 3D capture using the Focus Window. These methods are explained in further detail in Chapter 7, [3D Capture](#) on page 105. You must first choose the basis for your capture. You may define your capture based on the current  $z$  position, a selected reference position, or by defining the top and bottom of capture.

Examples of using the Focus Window for 3D capture are as follows:

- Current  $z$  position - Use the Focus Window to find a plane of interest (for example the focal plane), and then choose to range your capture around that position in the Capture dialog.
- Define a reference position in the Focus Window, and then define a total range in microns or number of planes to capture in the Capture dialog.

- Define a top and bottom position in the Focus Window, and choose to use those positions in the Capture dialog. You may define the step size in either the Focus Window or the Capture dialog.

The **Z** tab contains controls that let you make an interactive selection of a *z* axis range that can be imported into the Capture dialog when performing 3D acquisition. It is available on all systems that have motorized *z* focusing.



**NOTE:** You can also select a *z* axis range directly in the Capture dialog box (see [3D Capture](#) on page 105). The **Z** tab allows you to interactively define the 3D capture range and is generally used under circumstances where the precise *z* extent of the sample is unknown.

### 5.1.3.1 Set Top, Set Bottom, Set Reference, Center

The **Set Top** and **Set Bottom** buttons let you specify the end points of a 3D capture range. You can select these in either order. For instance, suppose you have found a cell that you want to capture and are in the middle of the focus range. To set the boundaries of capture in the *z* direction, first move the stage (or nosepiece) up until the focal plane is at the top of the volume that you wish to capture. This can be done either by repeatedly clicking on one of the “up” buttons or by adjusting the fine focus of the microscope itself. After you are finished, click on the **Set Top** button. A number will appear above the **Set Top** button indicating that the top *z* position has been set. Next, focus in the opposite direction until the focal plane is at the bottom of the volume of interest. Click on the **Set Bottom** button. Again, a number will appear below the **Set Bottom** button indicating that the bottom *z* position has been set. Once you have set a top and bottom position, you may drag the *z* position slider to return to either of those positions. In addition, SlideBook will calculate the **Center** position based on your top and bottom positions. You may select **Go** to return to the center position.

You may also set a reference, or *home*, position. This reference position can be the point of best focus, or some other position that is of importance for your experiment. Once you set this reference position, it may be used for 3D capture. To set the reference position, press **Set**. This position will be saved until you reset the position or close SlideBook.

**NOTE:** If you do not wish to continuously expose your sample while performing *z* focusing, you may set these ranges using single images generated in the Capture dialog box. Please see [3D Capture](#) on page 105.

### 5.1.3.2 Slider

Once top and bottom  $z$  positions have been selected, the slider to the left of the buttons provides the ability to move the  $z$  focus of the microscope to any position along the specified range. This is useful for confirming that the specified region covers all of (and not significantly more than) the volume you wish to capture and provides a means of returning the  $z$  focus to the brightest part of the sample without altering the 3D capture range. When you later wish to find optimal exposure times that won't saturate the camera, it is best to be focused on the  $z$  position with the highest peak intensities.

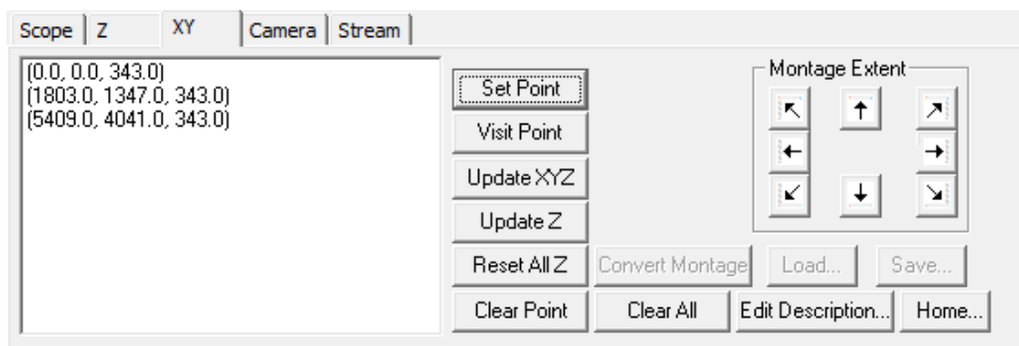
### 5.1.3.3 Capture Information

Choosing top and bottom positions in the Focus Window to set the extent of  $z$  travel is only one part of setting up a 3D capture. You also need to specify how many *planes* (discrete  $z$  focus positions) should be collected. Typically, the most important consideration is the *step size*, as this governs the ultimate  $z$  axis resolution of the acquired image. The step size (in microns) is specified in the corresponding edit box. SlideBook suggests a  $z$  step size based on the theoretical resolution of the lens. The theoretical resolution is a function of wavelength of the chosen filter and numerical aperture. You may adjust this step size by entering a value in the **Step Size** edit field. Once top and bottom positions are defined, SlideBook will use the step size to compute how many planes must be acquired to cover the specified range. At any point a different step size can be entered and the computed number of planes will be updated. Note that you can also specify the direction of travel of the stage (or nosepiece). The stage and objective should always travel toward each other. Thus, the direction of the arrow should normally point upwards.

You may also choose to set either number of planes or the step size in the Capture dialog, as shown in [3D Capture](#) on page 105.

### 5.1.4 XY

If your system is equipped with a motorized  $xy$  (and  $z$ ) stage, the **XY** tab lets you specify, store and visit multiple  $xy$  (or  $xyz$ ) locations. These points can be used for visiting multiple locations of interest in a sample and returning to them later as well as for specifying a sequence of locations to repeatedly visit during timelapse acquisition.



- **Set Point** - adds the current  $xyz$  location to the list
- **Clear Point** - removes the currently selected  $xyz$  location from the list

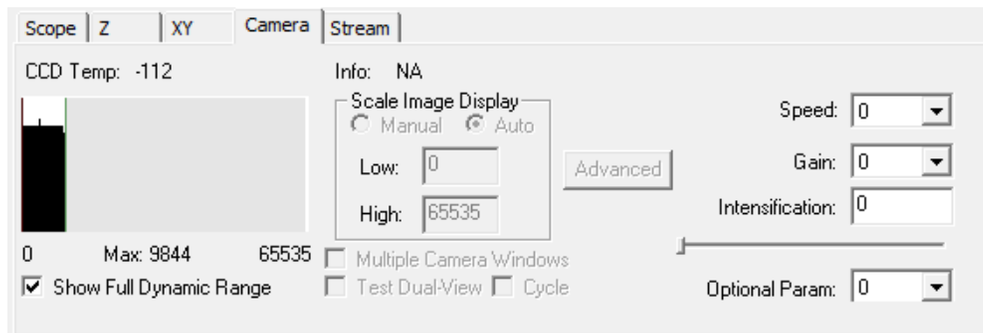
- **Visit Point** - moves the stage (and  $z$  focus) to the selected  $xyz$  location
- **Clear All** - removes all locations from the list
- **Reset All Z** – updates the  $z$  positions in all points to the current  $z$  position
- **New Point Z** - updates the  $z$  position of the selected point to the current  $z$  position
- **Montage Extent** - sets the boundaries that will be used to create a montage image.
- **Load** – allows you to load a saved  $x,y,z$  point list
- **Save** – allows you to save an  $x,y,z$  point list
- **Home** – allows you to calibrate your  $x,y$  stage before loading a point list.

Using these features in the context of preparing to capture an image is also discussed in [Chapter 7-Advanced Capture](#).

**NOTE:** If your system has a motorized  $xy$  stage, but not motorized  $z$  focus, then the **XY** tab works the same way except with  $xy$  rather than  $xyz$  locations.

### 5.1.5 Camera

The **Camera Tab** lets you control various settings on the CCD camera as well as view a histogram of the pixel intensities for the image in the focus window.



#### 5.1.5.1 Histogram

The histogram shows a plot of pixel intensities values for the focus window image. Pixel intensity values are plotted on the  $x$  axis, while numbers of pixels at a given intensity are plotted on the  $y$  axis. In addition, the minimum, maximum, and mean pixel intensity values are reported beneath the histogram. This data is continuously updated when the camera is actively acquiring images. The histogram can be operated in two modes as shown in the **Scale Image Display** region of the window, **Manual** and **Auto**. In manual mode, you may drag the red and green bars to set the min and max values that are displayed in the live image. You may also enter numbers in the **Low** and **High** edit fields. In auto mode, the live image will use the minimum and maximum intensity values of the live image to set

the contrast in the image. Thus, if you increase the exposure time, the live image will update and automatically reset the contrast of the image to match the range of intensities in the live image. You also have the option to display the full dynamic range of the camera using the **Show Full Dynamic Range** checkbox. When this box is checked, the histogram will not show the minimum and maximum intensities of the image, but rather the minimum and maximum *possible* intensities based on your camera's readout.

### 5.1.5.2 CCD Temp

On systems with a camera that has electronic temperature readout, this field reports the current sensor temperature in degrees Celsius.

### 5.1.5.3 Speed

On cameras that support this adjustment, the CCD read-out speed can be adjusted using this dropdown menu.

### 5.1.5.4 Gain

On cameras that support this adjustment, the gain can be set using this dropdown menu.

### 5.1.5.5 Intensification

On cameras that support this adjustment, the intensification factor can be adjusted using the slider and edit field located below the speed menu.

### 5.1.5.6 Parameters

This dropdown menu allows for support of additional features on certain cameras. For instance, on Hamamatsu cameras, the parameters menu allows you to select the camera offset.

### 5.1.5.7 Test Dual-View

The **Test Dual-View** check box is used with systems capable of performing dual camera simultaneous capture. It is also available when SlideBook is configured with an image splitter such as the Optical Insights Dual-View (see [Defining Filters when using an Image Splitter](#) on page 72 and [Configuring an Image Splitter](#) on page 60) This option is only available when multiple cameras or an image splitter have been configured in SlideBook. When this box is checked, the focus window display will rapidly alternate between images from camera 1 and camera 2.

## 5.2 Using the Focus Window

The following general procedure describes how to use the focus window to focus on a sample. If the microscope that you are using is not automated, you will need to move the optics manually. Procedures for using more advanced features of the focus window are detailed in [Chapter 7 –Advanced Capture](#).

1. Open the Focus Window by either selecting **Window > Focus Window** or clicking on the focus window button in the toolbar . The Focus/Capture Window will begin updating as soon as the Focus Window is opened.
2. Choose the desired objective and magnification changer from the **Scope** tab.

3. Choose the desired filter configuration from the **Focus Controls** below the **Scope** tab.
4. Place the sample on the stage.
5. Click on **Open Fluor** or **Open Bright** to open the desired lightpath.

Next, you will need to bring the sample into focus. This may be accomplished by either viewing the sample through the eyepieces or viewing the sample in the focus window. If your microscope is equipped with a motorized emission light path, the image can be directed to the eyepieces or the camera using the **Scope** tab. Moving the *xy* stage and changing the *z* focus can be accomplished by either using the manual controls on the microscope or the computer controls (see above).

6. Using the slider, set the exposure time so that the image is not overexposed.
7. Using the controls in the **Focus Controls** window (or joystick or manual controls), adjust the *x* and *y* position of the stage and *z* position of the stage or nosepiece to find and focus on an area of interest.

**TIP:** If you double-click on an object in the image display, the *xy* stage will move so that the object is centered.

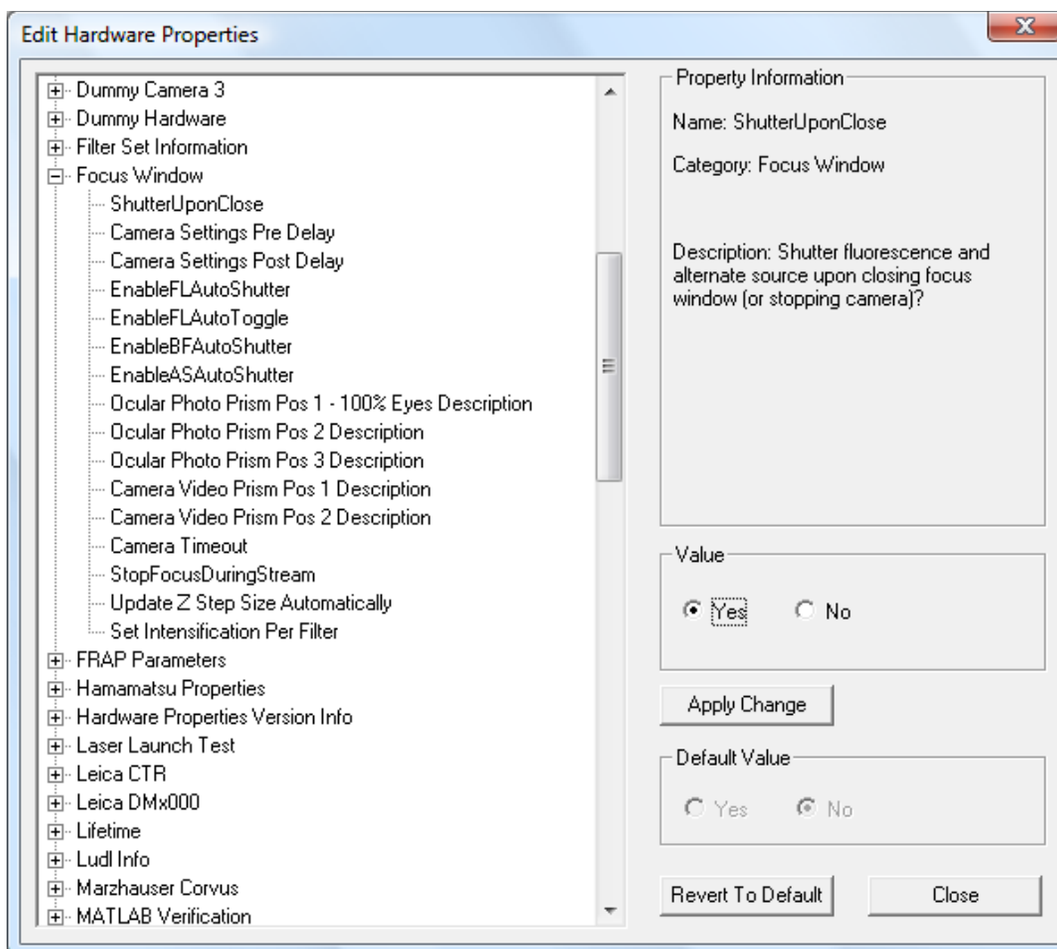
8. Once the sample focus is satisfactory, you may wish to close the lightpath for light sensitive samples, and then close the focus window by pressing the **X** button in upper right corner. If desired, the Focus Window and Capture dialog box may be open simultaneously.

### 5.2.1 Snapping an Image

You may find that you want to capture a single image as you are using the Focus Window. To do so, simply press **Snap** once you are happy with the image in the live view. The image will be stored into the open slide file.

### 5.2.2 Closing the Fluorescence Shutter Automatically when Closing the Focus Window

If you would like your fluorescence shutter to close automatically when you close the focus window, select **Edit > Hardware Properties**, then expand the **Focus Window** section. Click on **ShutterUponClose**.



Set the value to **Yes** and then click on **Apply Change**. Click **Close** and then restart SlideBook.

### 5.2.3 Labeling the Ocular Photo Prism Positions

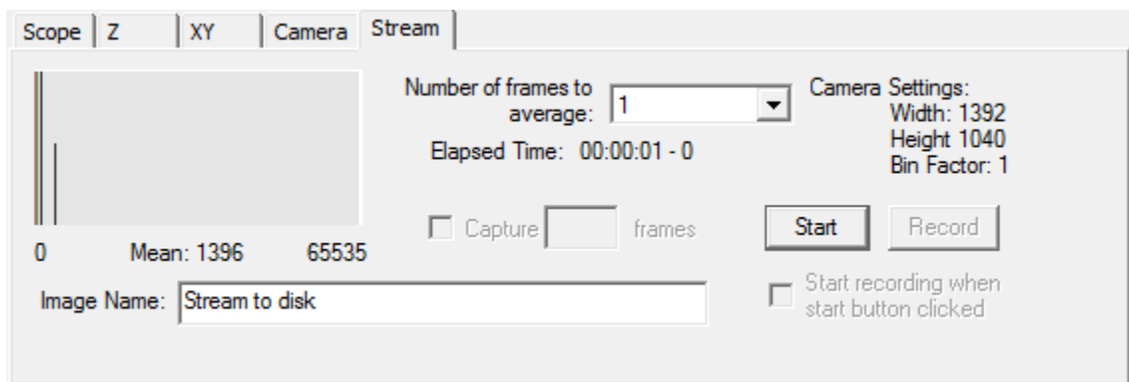
You may also provide custom labels for the emission selection positions (ocular photo prism positions) that appear in the **Scope** tab of the focus window. To do so, select **Edit > Hardware Properties**, then expand the **Focus Window** section. Click on the **Ocular Photo Prism Pos** you wish to edit and enter the label in the edit field.

Click **Apply Change**, **Close**, and restart SlideBook.

### 5.2.4 How to Perform Streaming Capture

The *streaming capture* function allows you to run your Hamamatsu or Photometrics camera at the fastest possible speed. There are limitations with Streaming Capture. When a camera is recording in this mode, you may only alter other hardware via the Focus Window. Streaming Capture does not work using the Capture dialog box, so you may not perform automated multi-channel, 3D, 4D, multipoint or montage capture. This capture mode is ideal for rapid 2D timelapse capture. To run your camera in streaming capture mode:

1. Open the **Focus Window** and select the **Stream** tab.



2. Select the **Start** button. A new image capture window will open with a counter for number of images captured.



3. Your camera will continuously generate images with the exposure time that you have selected in the Focus Window. If desired, select the **Number of frames to average** from the drop-down list.
4. Name the file in the **Image Name** field.
5. Press **Record** to save the captures to disk. Once you have pressed record, images will be saved to disk and the Record button will convert to **Pause**. To end the recording, press **Stop**. This will also stop the streaming update. To resume the streaming update, press **Start**. You will not be able to change camera settings while the camera is streaming. You must press **Stop** and make changes to camera settings, then press **Start** again.

**CAUTION:** Recording at this speed will rapidly use computer disk space.

Spool files will be opened automatically once you press **Stop**. They will also be placed in the open slide with the name designated in Step 4.

## 6 Image Capture and Import

Image capture parameters are set in SlideBook's Capture window. In order to capture an image, you must first set your optics and bring your sample into view and focus using the **Focus Window** (please see [Chapter 5 - Controlling the Camera and Microscope Hardware \(Focus Window\)](#) on page 77). Next, you must set camera parameters such as binning, exposure time, and image extent, as described in this chapter. Once all of these parameters have been set, you may initiate capture. You may also choose to import an image created using another program. This chapter covers the following topics:

- Selecting Channels and Setting Exposure Times
- Selecting the Area to be Imaged
- Entering Image Information and Optical Parameters before Capture
- Capturing a 2D Image
- Performing Auto-White Balance
- Importing an Image
- Getting and Editing Image Information after Capture

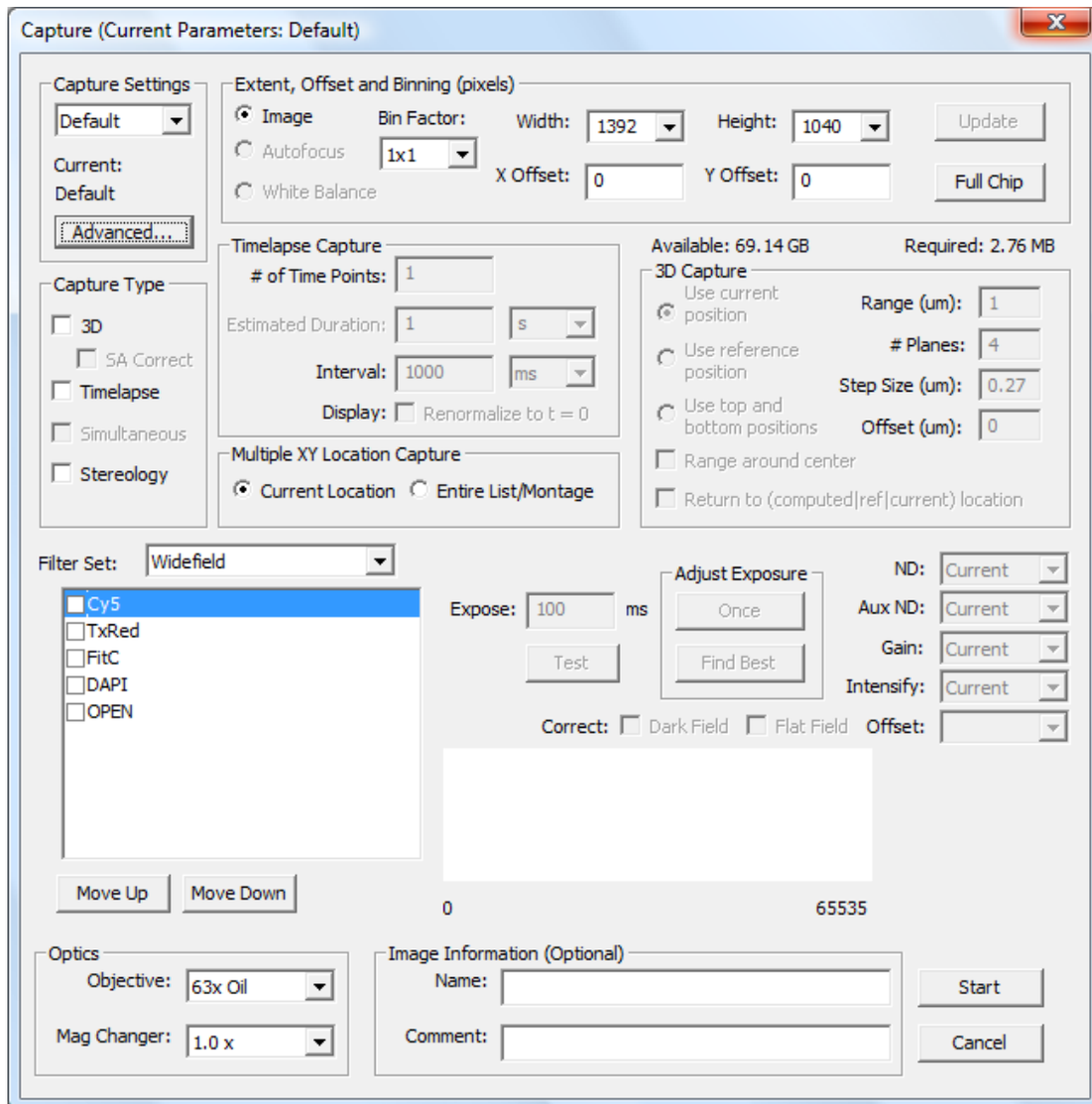
Advanced topics such as 3D, timelapse, 4D, montage, and multipoint imaging are covered in [Chapter 7 – Advanced Capture](#).

### 6.1 Selecting Channels and Setting Exposure Times

Exposure times are set by generating a series of test images for each channel that will be exposed. If you are performing a transmitted light capture with a color camera or color LCD filter, you may wish to first perform an auto-whitebalance (see below).

1. Once you have a sample in focus, open the Capture dialog by selecting **Image > Capture** or by selecting the Capture icon in the toolbar . Note that you must have a slide file open in order to open the Capture dialog.

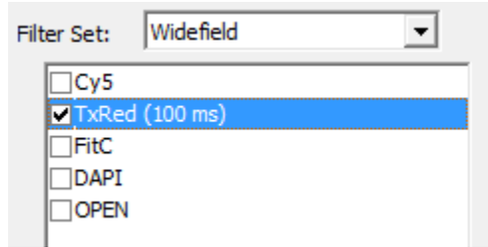
The dialog box that appears is divided into eight parts starting from the upper left corner: *capture settings*, *image extent and binning factor*, *multiplane capture*, *multipoint capture*, *exposure settings by channel*, *image histogram*, *optical parameters*, and *image info*. The Capture dialog uses the same Live View as the Focus Window.



2. First select the binning factor in the upper left drop-down menu.

*Binning* lets you group several pixels together to increase sensitivity (at the cost of resolution). For instance, selecting  $2 \times 2$  binning effectively gives you a camera with half the horizontal and vertical resolution, but four times the sensitivity. Note that as you increase binning, you will need to reduce the exposure time.

3. Select the desired **Filter Set** from the dropdown menu.
4. Click on the checkbox of the channel that you would like to expose. The channel that you select will be highlighted in blue.



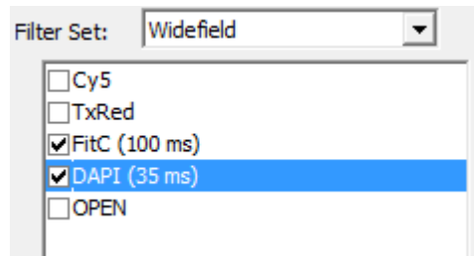
5. Enter an exposure time in the **Expose** field, if a different one is desired, and press **Test**.

Pressing the **Test** button for a particular filter configuration will give you a sample single-plane image at the specified exposure time for that channel. You may resize the image by dragging and dropping the corners or edges of the Live View. An *intensity histogram* and the minimum and maximum data values will be displayed in the Capture dialog in order to show you where in the dynamic range of the CCD your exposure lies.

Making a test image for each channel gives you the opportunity to choose the best exposure times (and of course make sure that the light is directed towards the CCD camera and not the eyepieces).

6. Press the **Once** button in the **Adjust Exposure** section of the dialog and notice that the exposure time has been adjusted to take advantage of the CCD camera's full dynamic range.
7. Press the **Test** button again to update the image in the test view.
8. Repeat the test and adjust exposure process until the histogram shows an adequate intensity distribution (see note below).

For multi-channel capture, simply repeat this process for each channel that you wish to expose. For instance, a two channel capture using both the DAPI and the FITC filter configurations will appear as follows:



**NOTE:** The maximum value for a pixel depends on the bit depth of the camera. For instance, on 12-bit cameras, the pixel values can range from 0 to 4095. In order to take advantage of the full dynamic range of the camera, it is important to try to get the maximum pixel value well into the four digit figures. Pressing the **Once** button will automatically adjust the exposure time to try to achieve a maximum pixel value

of about 3000. In extremely dim samples, this may not be practical, but it is a good target value. Conversely, it is important for deconvolution that no pixel values are at the camera's maximum value, as that means that the image is overexposed and past the linear range of the CCD detector. Pressing the **Once** button will reduce exposure times if necessary to try to reach a lower maximum value.

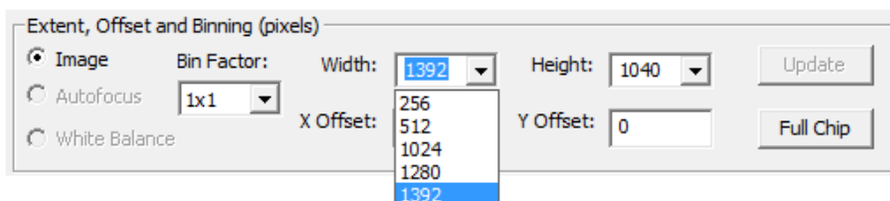
## 6.2 Selecting the Area to be Imaged

Typical cameras contain either a  $1392 \times 1040$  or a  $512 \times 512$  pixel chip, but often you will want to capture from a smaller portion of the chip in order to reduce the memory required for capture and/or increase the camera frame rate. You may select the area to be imaged in two ways.

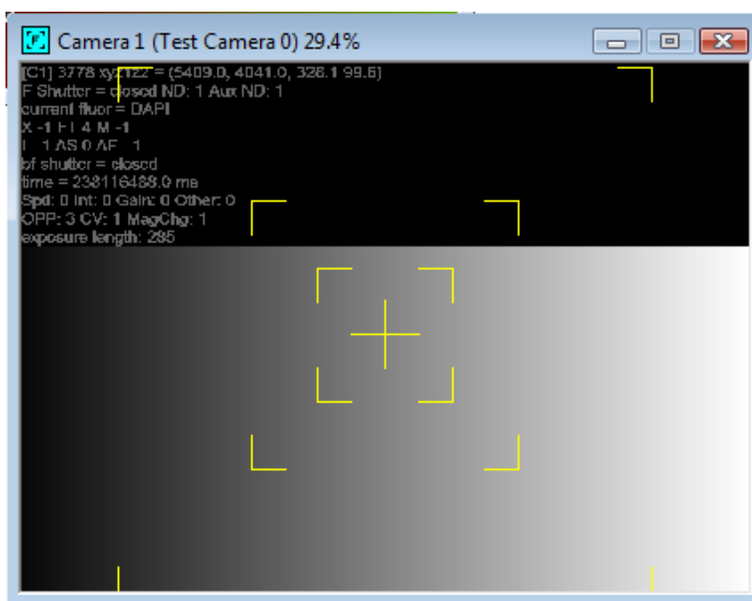
### 6.2.1 Using the Image Extent Menu

The image area can be selected using the features in the Image Extent and Binning section of the Capture window.

1. Select the x- and y-extent of the image from the **Width** and **Height** drop down menus. You may also manually enter the desired values in the edit field.



You may select to add yellow guides in the test view that correspond to  $256 \times 256$ ,  $512 \times 512$ , and  $1024 \times 1024$  pixels. By default, SlideBook reads from an area centered on the chip with the provided dimensions. To add the guides, you must select the **Guides** checkbox in the Focus Controls.



2. If you want to move the region that is read to some other location on the chip, enter offset values for both the  $x$ - and the  $y$ -axes in the **X Offset** and **Y Offset** fields.
3. Select **Test** to update the Live View so that it has the desired size.

### 6.2.2 Using the Test View

Alternatively, you may choose a region of the chip to be imaged by directly selecting it from the Image Window.

1. Be sure that the  button at the top of the window is depressed.
2. Click and drag the mouse over the desired portion of the test view.
3. Select **Update** to register the new image size.
4. Select **Test** to update the Live View so show the new image size.

Clicking on the **Full Chip** button will automatically select the maximum extent of the camera.

**NOTE:** After you have selected the area to be imaged, you may wish to adjust the exposure times using the **Test** and **Once** buttons to take advantage of the full dynamic range of the camera.

### 6.3 Entering Image Information and Optical Parameters before Capture

You may specify the following image information and optical parameters in the Capture dialog box.

- **Image Name** and **Comments** – can be entered in edit fields
- **Objective** and **Mag. Changer** - can be selected (or will automatically be selected if they are either encoded or automated) using the dropdown lists

The channels to be captured and their exposure times are automatically stored with the image. Although these settings don't immediately affect the microscope settings or capture, they are stored with the image and are important for correct deconvolution and measurement.

You can always go back and add or alter this information by selecting **Image > Get Info**, which is described in the section [Getting and Editing Image Information After Capture](#) on page 101.

### 6.4 Capturing a Single or Multi-Channel 2D Image

The default mode of the capture window is for a single 2D image. The 2D image may be either single or multi-channel depending on the number of channels that you have checked. If you wish to capture 3D, timelapse, 4D, multipoint, or montage images, please see [Chapter 7 –Advanced Capture](#).

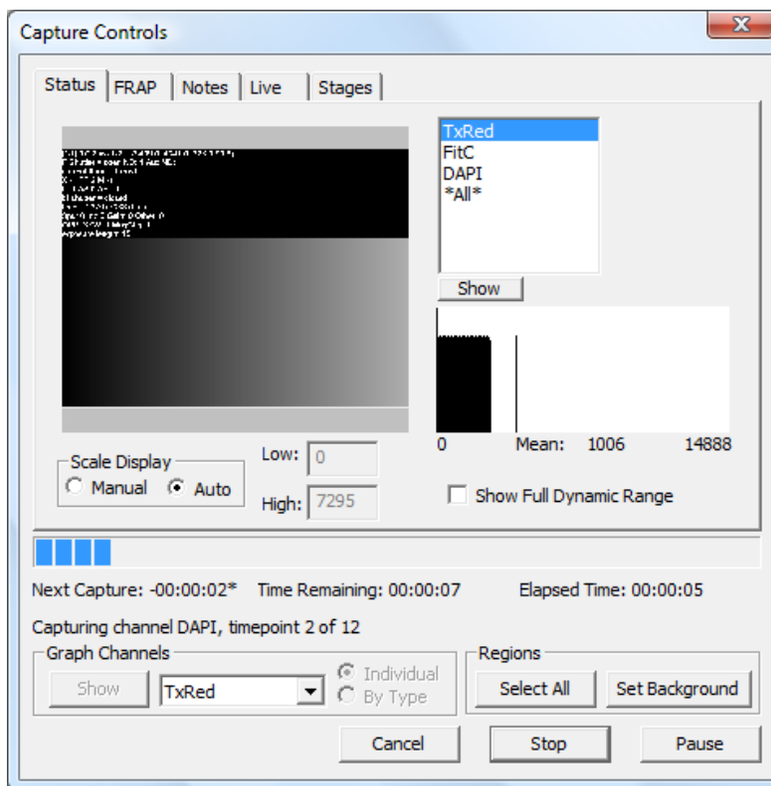
1. Once you have set all of the parameters (channels, exposure times, binning, etc.) in the Capture dialog box, select **OK** to begin capture.

If your system does not have automated filter control, you will be prompted to move the filters into the correct position before the image is captured.

**NOTE:** If you see these prompts in error, you need to make sure that you do not have positions defined for hardware components that are not motorized. For example, if you have a filter cube (reflector) turret that is not motorized, make sure that your filter definitions are designated **Unmounted** in the filter turret position.

2. Move the hardware into position and click **OK**.

The Live View will show the image in color as it is being acquired. If you have more than one channel selected for capture, it will show a combined image. Thus if you have designated your DAPI and FITC filters to display as blue and green, respectively, the Image Window will first display the DAPI image in blue and then overlay the FITC image in green. In addition to the Image Window updates, the Capture Controls will become active and report the progress of your capture.

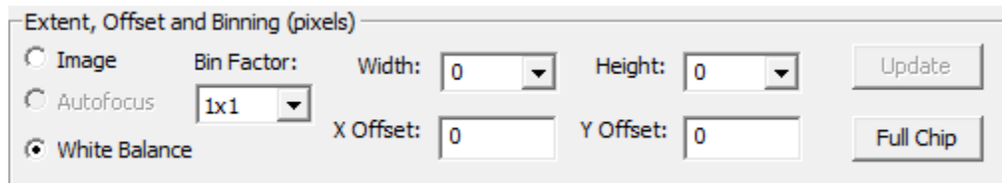


The default **Status** tab of the Capture Controls displays the current channel in monochrome. The features of the Capture Controls are best utilized when performing timelapse imaging and are discussed in [Timelapse Capture](#) on page 112.

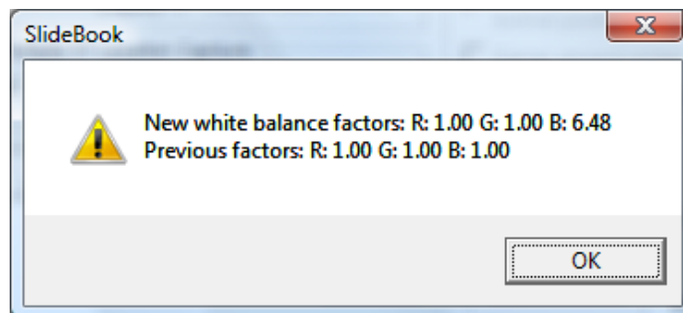
### 6.5 Performing Auto-White Balance for Color Capture

If you are performing a color capture with an LCD filter, you may first wish to perform a white balance procedure (you must first define your filters as discussed on page 69). This will yield the truest color rendering. To do so:

1. In the focus window, find a clear portion of your slide. Alternatively, use a piece of white lens paper.
2. Make sure that the halogen lamp is set to 3200K and that your microscope is set for Köhler illumination.
3. Select the green filter and check that your exposure time is at least 25ms (this yields the best white balance results). If you are saturating the camera at 25ms, you should use neutral density filters.
4. Once you have achieved an adequate exposure time, go to **Image > Capture** or select the image capture icon .
5. Select the filter configuration checkbox corresponding to the color channel and then **Test**.
6. Select the area to be imaged and any binning desired.
7. Adjust the exposure time as described above.
8. Select the area of the test view that you would like to use for performing white balance by clicking and dragging over the region.
9. Select **White Balance** from the radio buttons and then click **Update**.



10. A white balance operation is performed and a dialog box similar to the following will appear.

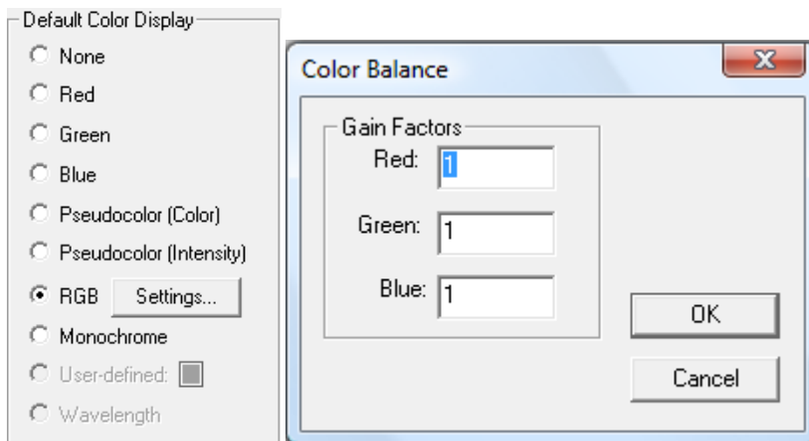


**NOTE:** If you are getting inconsistent color maximums when performing capture and the RGB factors are varying significantly each time you perform a white balance, then you should adjust your illumination. Bright, dim or inconsistent lighting may cause this phenomenon.

11. Select **OK** to close the dialog box.
12. You may now capture an image by selecting **OK** in the Capture dialog box.

The white balance settings will remain in the system memory until the slide file is closed.

13. If you would like to manually enter white balance settings, go to **Edit > Define Optics > Filter Configuration** and select the red filter configuration for an LCD filter, or to the color filter configuration for a color camera.
14. In the default **Color Display** section, select the **Settings** button. The following dialog box will be displayed:



15. Enter the white balance gain factors. In general, the green factor will be the largest.
16. Click **OK** to save the settings.

## 6.6 Importing an Image

You may also use SlideBook to display and analyze images captured using other programs. The following file types are supported by SlideBook:

- SlideBook Spool (\*.sp1)
- TIFF
- TIFF Sequence

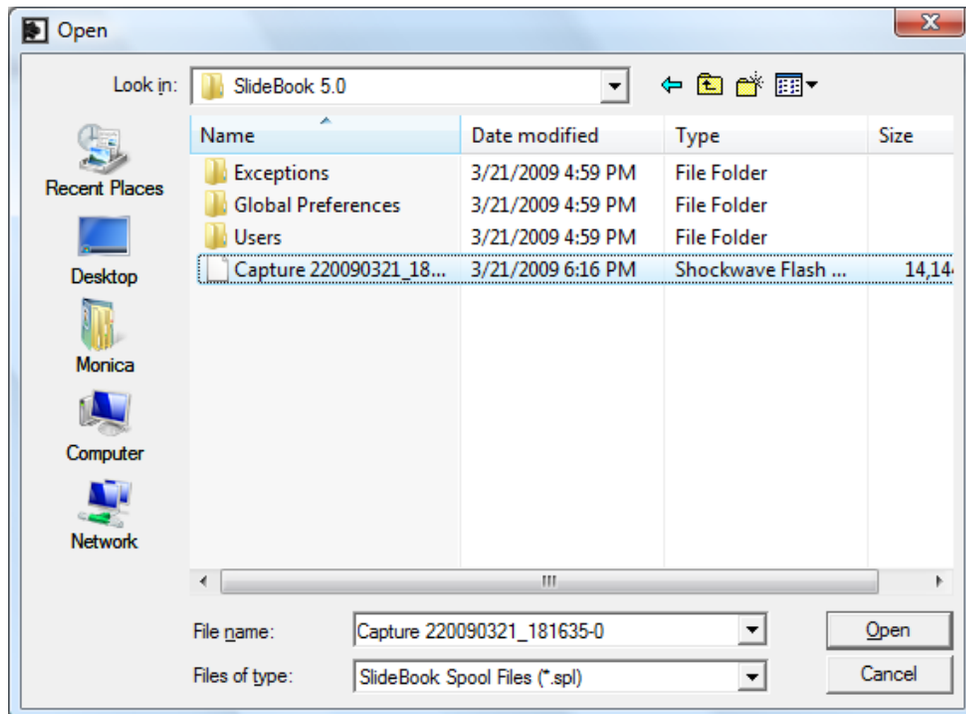
SlideBook Spool is the file type used when capturing images to disk. Please see [Saving Images to Disk \(Spooled Capture\)](#) on page 153.

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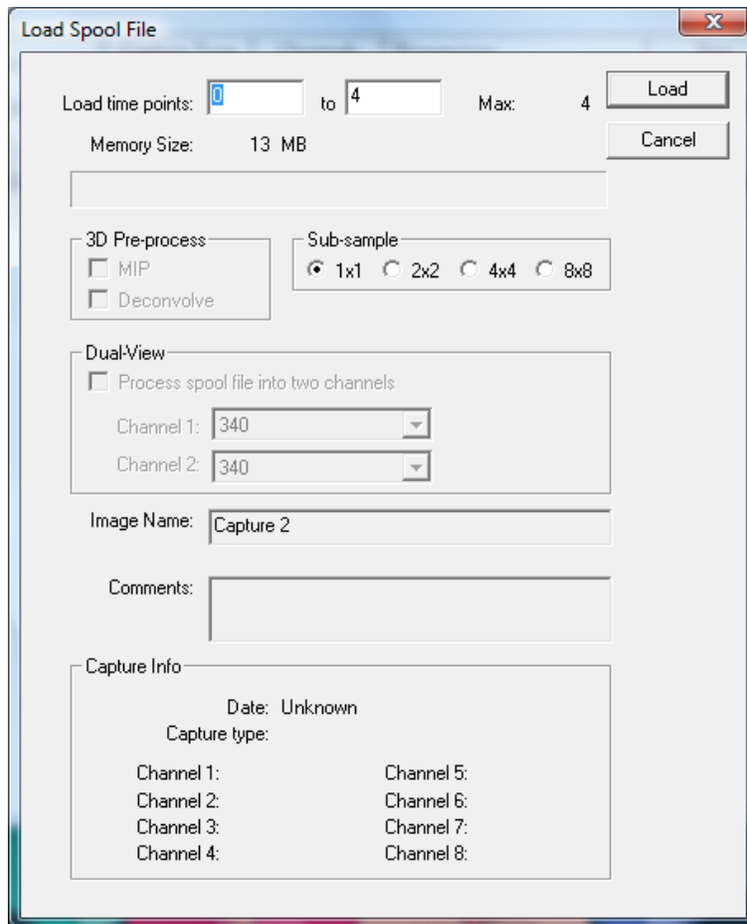
### 6.6.1 Importing SlideBook Spool Files

To import a SlideBook Spool file:

1. Select the destination for import by either opening a new slide or opening an existing slide file. You may do this by selecting **File > New Slide** or **File > Open Slide**. If the slide is already open, make it the active window by clicking on it.
2. To import an image, select **Image > Import > SlideBook Spool**. The following dialog will appear:



3. Navigate to the desired spool file and select **Open**. The **Load Spool File** dialog will appear.



4. Select the timepoints that you wish to load. For large data sets, it can sometimes be easier to load a subset of data.
5. Select if you would like to subsample the data. This may speed up processing for large data sets but will decrease the image resolution.
6. If applicable, select pre-processing and Dual-View parameters.
7. Name the image and add comments.
8. Click **Load**. The image will be added to the active slide.

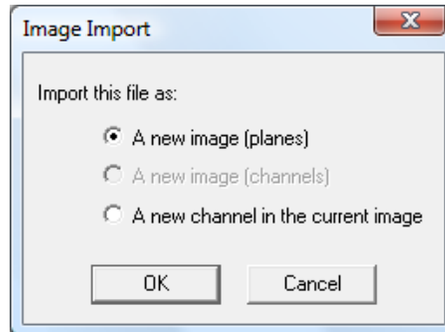
### 6.6.2 Importing TIFF Files

To import individual TIFF files or TIFF sequences:

1. Select the destination for import by either opening a new slide or opening an existing slide file. You may do this by selecting **File > New Slide** or **File > Open Slide**. If the slide is already open, make it the active window by clicking on it.

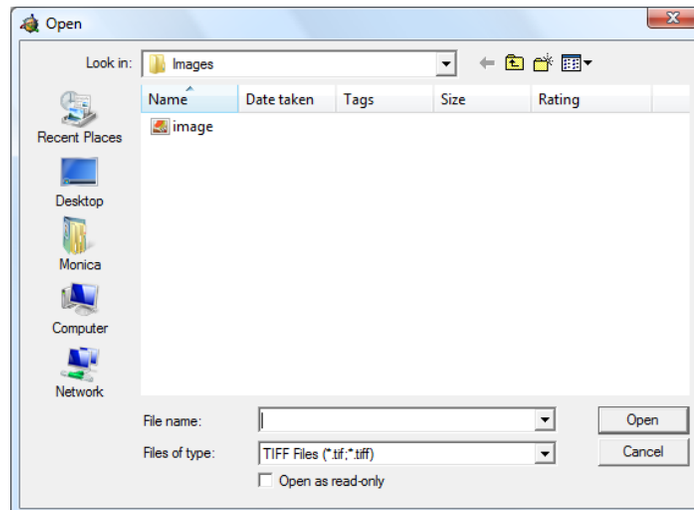
2. To import an image, select **Image > Import** and select the file type that you wish to import.

If there are already images in your slide file, and one is selected, the following dialog box will appear for TIFF files (but not TIFF Sequence):

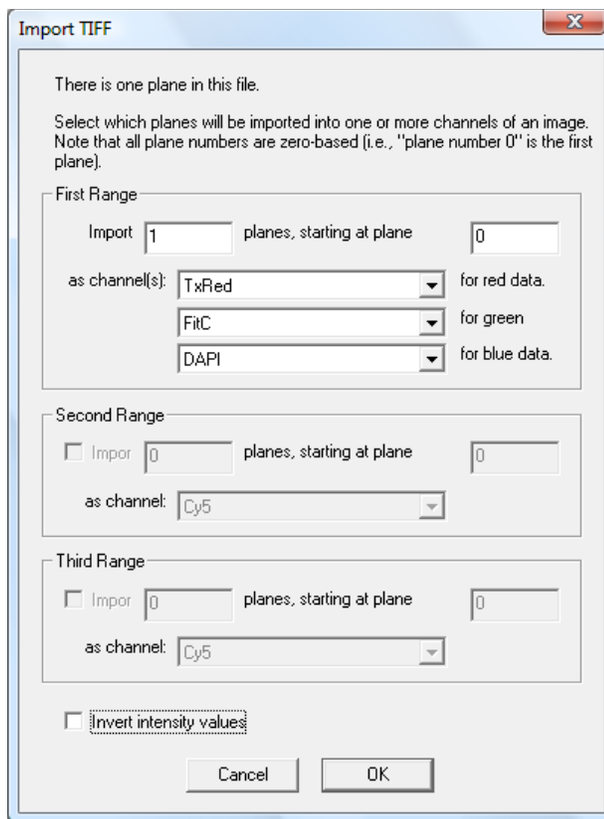


Select whether you would like to import the file as a new image or as a new channel in the current image, and click **OK**.

3. The following dialog box will appear:



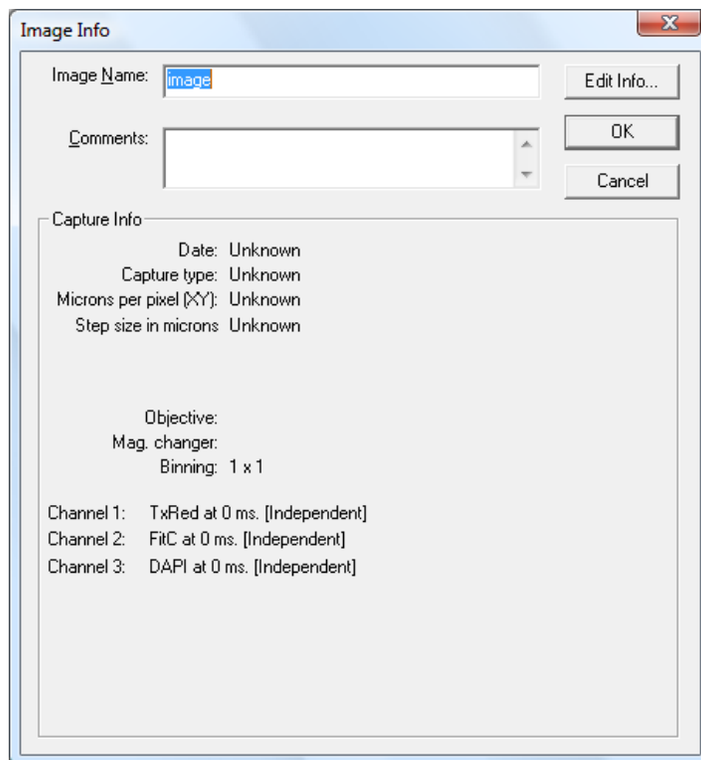
4. Navigate to the file and select **Open**. The following dialog box will appear if you have selected TIFF. (If you have selected TIFF sequence, proceed to step 6.)



5. Select the planes you would like to import as well as the channels to be used for each color in the image and select **OK**.

For example, if you have an image that displays FITC as green data, you will designate the green data as FITC in the dropdown menu.

6. The following dialog box will appear:



7. Enter image information as outlined in the following section.
8. Click **OK** after you have entered the information. The imported image will appear as a **Main View**. You may always go back and edit the image information by selecting **Image > Get Info**.

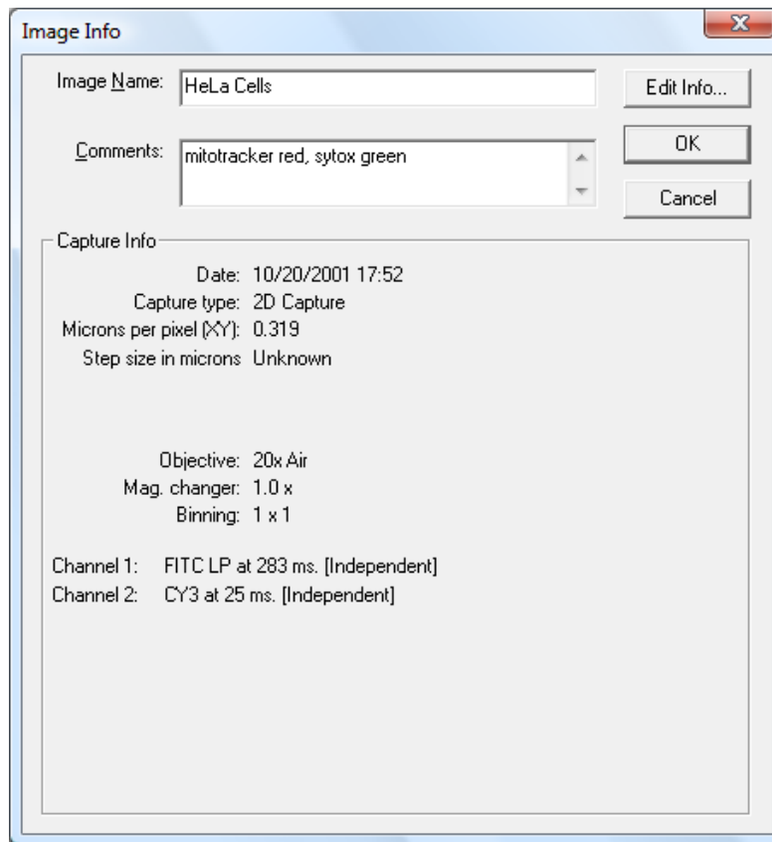
### 6.7 Getting and Editing Image Information After Capture

You can get information about an image after capture or import. The Image Info dialog box gives you the name of the image, comments associated with the image, and the time and date of capture if the image was collected using a digital microscopy workstation (as opposed to imported). It also gives you the optical settings as well as which filter configurations were used to collect each channel along with the exposure time. If you would like to add optical information to an imported image, you must first define the objective, magnification changer, and filter configurations as described in Chapter 4 - [Configuring Your System](#).

You may view and edit information as described below.

#### 6.7.1 Images that were captured in SlideBook

1. Open a Main View of the desired image by either double-clicking on the thumbnail in the slide file, or by clicking once on the thumbnail and then selecting **View > New Main View**.
2. Select **Image > Get Info**. The following dialog box will appear. View and edit the **Image Name** and **Comments** fields as desired.



3. Press the **Edit Info** button if you wish to change any of the optical configuration or collection information. This brings up the following dialog box:

**Edit Image Info**

Capture Type: 2D Capture Interval (ms): 0

Microscope Components

Objective: 20x Air

Mag. Chg.: 1.0 x

Optical Parameters

Microns/pixel: 0.649

Num aperature: 0.5

Working distance: 2000

Eta: 1

3D Collection

Interplane spacing (um): 0

Channels

	Filter Configuration	Lambda (um)	Time (ms)
Channel 1:	Other (FITC LP)	0.54	283
Channel 2:	Other (CY3)	0.59	25
Channel 3:		0	0
Channel 4:		0	0
Channel 5:		0	0
Channel 6:		0	0
Channel 7:		0	0
Channel 8:		0	0

OK

Cancel

**NOTE:** If you open a SlideBook file that was created on another computer or with a different SlideBook preferences version, the filter and objective configurations will be displayed as **Other**. To prevent this, make sure that the SlideBook preferences match those that were used for capture.

4. Select the **Objective** and **Mag. Changer** used during capture from the drop down menus.

**NOTE:** If you have imported an image from another program, you must first enter the objective and magnification changer definitions as discussed in Chapter 4. Then you may select the proper objective and magnification changer from the drop down menus.

5. Enter the **Interplane Spacing (um)** for 3D images.
6. Select the filter configurations from the dropdown menus, and enter the exposure times for each channel.
7. Select **OK** when you are finished. The data that you entered is saved with the image.

**NOTE:** Even if you are working with 2D images, it is a good idea to provide a value for **Interplane Spacing** as this will be used in the no neighbors deconvolution algorithm for the theoretical spacing value. That way, you will not have to change any values that appear

in the deconvolution dialog box. As a guideline, you should use the approximate  $z$  resolution for the objective used.

### 6.7.2 Imported Images

You may also add information to images that are imported into SlideBook from other programs.

1. Import the image as discussed above in [Importing an Image](#) on page 96.
2. Make sure that the objective definitions exist by selecting **Edit > Define Optics > Objectives** and looking at the available objectives in the dropdown list. If the objective is not available, add the objective definition as described on page 62.
3. If a magnification changer (optovar) or relay lens was used, make sure that its definition exists by selecting **Edit > Define Optics > Mag. Changers** and looking at the available magnification changers in the dropdown list. If the magnification changer is not available, add the definition as described on page 73.
4. Make sure that the filter definitions exist by selecting **Edit > Define Optics > Filter Configurations** and looking at the available filters in the dropdown list. If the filters are not available, add the filter definitions as described on page 68.
5. Open a Main View of the imported image by either double-clicking on the thumbnail in the slide file, or by clicking once on the thumbnail and then selecting **View > New Main View**.
6. Follow steps 2-7 above for getting and editing info for images that were captured in SlideBook. **You should also select the Capture Type for your imported image and make sure that the image information is accurate.**

## 7 Advanced Capture

SlideBook can automate many different types of capture. This chapter covers the following topics:

- 3D Capture
- Timelapse Capture
- 4D Capture
- Multipoint Capture
- Montage Capture
- Multiwell Capture
- Simultaneous Capture
- Configuring Your Hardware for Speed
- Auto Focus
- Varying Capture Rates for Timelapse Capture (Sequence Feature)
- Saving Images to Disk (Spooled Capture)
- Saving Capture Parameters

### 7.1 3D Capture

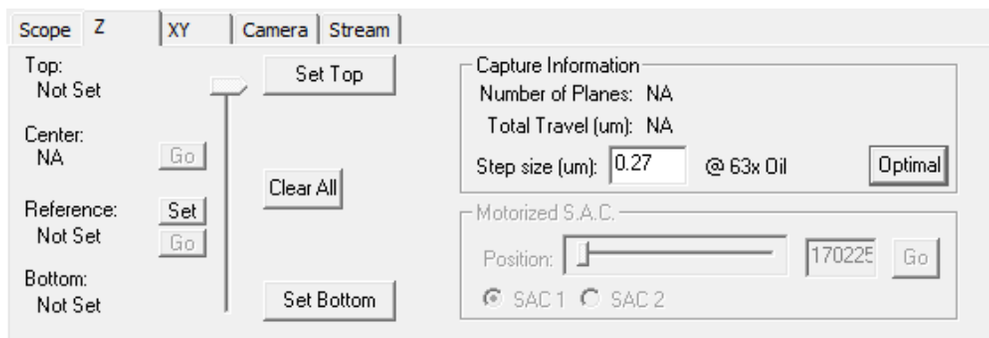
Similarly to 2D capture, 3D capture is accomplished by setting microscope parameters in the Focus Window and then setting capture parameters in the Capture dialog box. In order to perform 3D capture, your system must have motorized  $z$  control.

There are two procedures for performing 3D capture. In Method 1, you will use the focus window both to bring the sample into view and to interactively set the capture extent in the  $z$  dimension. These  $z$  parameters will then be imported into the capture dialog box. In Method 2, you will use the focus window to bring the sample into view and focus, but then set the  $z$  parameters in the capture dialog box. You may wish to use Method 2 if you already know the  $z$  range of capture. For example, if you are imaging an object that you know to be 30  $\mu\text{m}$  in height, you may bring the cell into focus and then capture 20  $\mu\text{m}$  above and below the plane of focus. However, if you are unsure of the extent of the object of interest in the  $z$  dimension, you should use Method 1 since it allows the range of capture to be interactively defined. Both methods are outlined below.

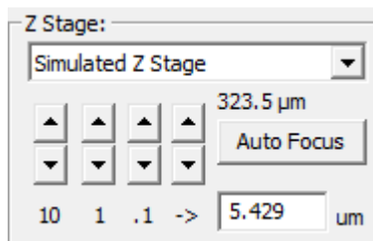
#### 7.1.1 Method 1

##### 7.1.1.1 Setting Focus Window Parameters

1. Open the Focus Window by either selecting **Window > Focus Window** or clicking on the focus window button in the toolbar . The current camera image will begin updating as soon as the Focus Window is opened.
2. Select your objective, magnification changer, and filter configuration, and bring your sample into view and focus as discussed on page 85 in [Chapter 5 - Controlling the Camera and Microscope Hardware \(Focus Window\)](#).
3. Click on the **Z** tab. Now, you will need to set the  $z$  limits for capture.



- Starting at the middle of the focus range, move the stage (or nosepiece) up until you are at the top of the volume that you wish to capture. This can be accomplished either by repeatedly clicking on one of the “up” arrow buttons or by manually adjusting the fine focus of the microscope.



**NOTE:** When moving the *z* position by computer control you may either use one of the fixed increment buttons (10, 1 or .1 micron steps) or manually enter a step size in the adjacent edit box.

- Click on the **Set Top** button. SlideBook will save the current *z* position and display it to the left of the **Set Top** button.
- Focus in the opposite direction until you are at the bottom of the volume of interest.
- Click on the **Set Bottom** button. SlideBook will save the current *z* position and display it to the left of the **Set Bottom** button.
- Return to the point of best focus (the brightest plane) using the slider, **Go** button, or microscope controls. Returning to the brightest point in the sample is necessary so that the correct exposure times can later be defined in the capture window.
- Close the lightpath for light sensitive samples by choosing **Close Fluor**.
- Enter the step size that you would like to use for capture in the **Step Size (µm)** edit field of the **Capture Information** group. The number of planes that will be

required to capture the entire  $z$  range will be calculated and displayed above the step size.

**Note:** You may wish to alter the capture step size so that the size of the captured image is not excessive. Alternately, you may decrease the total distance traveled by resetting the top and bottom parameters.

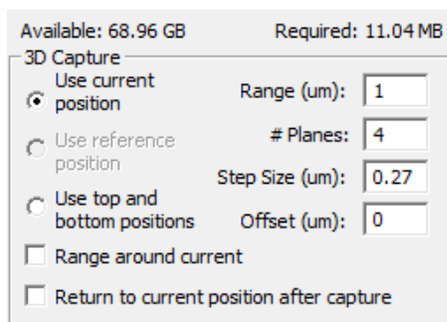
11. Close the focus window by pressing the **Close** button.

### 7.1.1.2 Setting Capture Window Parameters

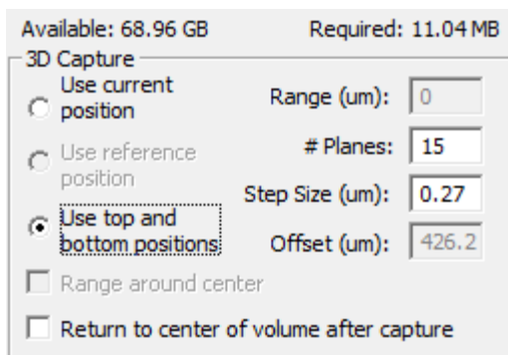
1. Once you have set the  $z$  parameters in the Focus Window, open the Capture dialog box by selecting **Image > Capture** or by selecting the capture icon in the tool bar .
2. Select channels and set exposure times as described in the section Selecting Channels and Setting Exposure Times in Chapter 6 (page 89).

**NOTE:** It is important that  $z$  parameter is set at the middle of the focus range (brightest) when setting exposure times so that the brightest planes are not overexposed.

3. Select the area to be imaged as described in the section Selecting the Area to be Imaged in Chapter 6 (page 92).
4. Enter image information and settings as described in the section Entering Image Information and Optical Parameters before Capture in Chapter 6 on page 93.
5. Click on the **3D** checkbox in the **Capture Type** section. The radio buttons and edit fields in the **3D Capture** section will become active. Note that default value of the **# Planes** field is one, signifying a single plane capture.



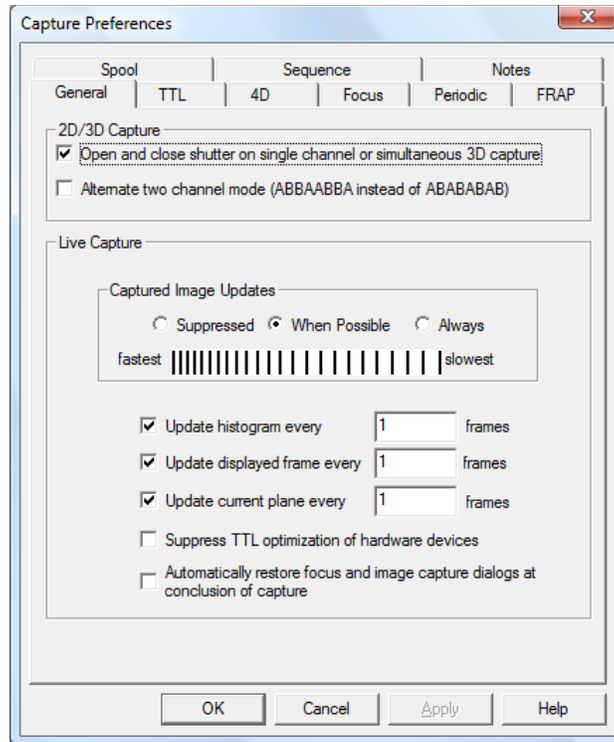
6. Import the  $z$  parameters that you set in the focus window by selecting the **Use top and bottom positions** radio button. The **# Planes** field will display the number of planes to be captured (see dialog box below). The **Spacing** field determines how many microns the stage needs to move between planes. The **Offset** field is not used with this method of setting 3D capture parameters.



7. If you wish for your  $z$  focus position to return to the center of the volume once capture is completed, select the **Return to center of volume after capture** checkbox.
8. Click **OK** to begin capture. The Capture Controls window will appear and display each plane as it is captured. You may choose to **Pause**, **Cancel**, or **Stop** your capture any time after it has started.
  - **Pause** – pauses the capture until you press **Continue**
  - **Cancel** – cancels capture and does not save any images
  - **Stop** – stops capture and saves a truncated sequence

#### 7.1.1.3 Setting Capture Preferences to Open and Close Shutter during Capture

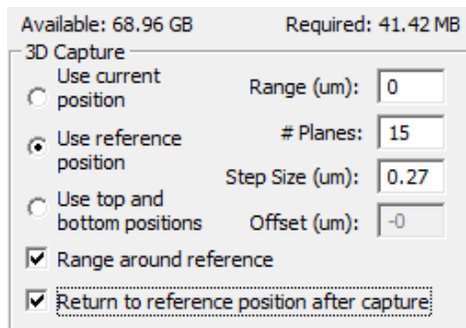
If you are performing single channel 3D capture, the default fluorescence shutter action in SlideBook is to leave the shutter open during capture. If you would like the shutter to open and close after every capture, select the **Advanced** button in the **Capture Settings** section of the Capture dialog box. Select the 2D/3D capture checkbox labeled **Open and close shutter on single channel or simultaneous 3D capture** in the **General** tab. The default action for multiple channel capture is to open and close the shutter after every capture.



### 7.1.2 Method 2

Use this method if you already know the distance above and below the plane of focus or other reference position that you would like to capture.

1. Bring your sample into view and focus by following steps 1-2 as described above in [Method 1: Setting Focus Window Parameters](#). If desired, you may set another position of interest by pressing the **Set** button next to **Reference**.
2. Set capture preferences as described above in [Method 1: Setting Capture Preferences](#).
3. Open the Capture dialog box by selecting **Image > Capture** or by selecting the capture icon in the SlideBook toolbar .
4. Follow steps 2 through 5 as described above in [Method 1: Setting Capture Window Parameters](#). If you have set a reference position, the dialog will appear as follows:

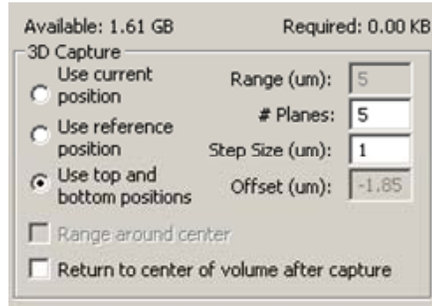
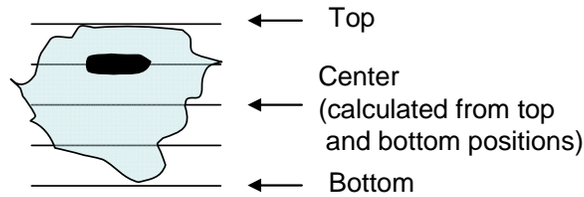


5. The edit fields that appear are described below.
- **Range (µm)** – Total distance traveled in microns.
  - **# (Number of) Planes** - Total number of z planes acquired during capture.
  - **Step Size (µm)** – Distance the stage (upright) or objective (inverted or fixed-stage) must move between z planes. A positive number typically corresponds to the stage or objective moving up during capture, while a negative number corresponds to the stage or objective moving down during capture.
  - **Offset (µm)** – The distance above or below the current stage location that the stage or objective will move before starting image collection. For example, on an inverted microscope, a positive number corresponds to the stage moving up while a negative number corresponds with the stage moving down. Thus, on an inverted microscope, you will need to use a positive spacing and negative initial offset.

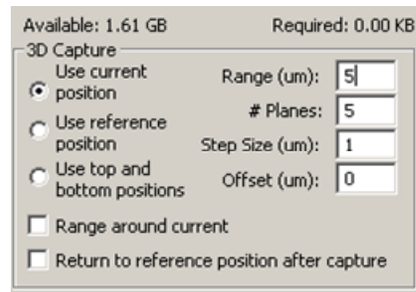
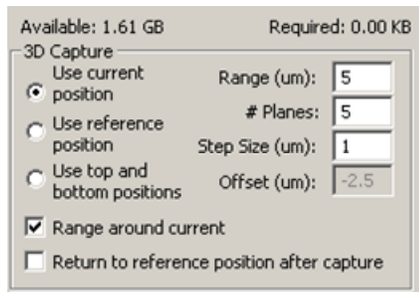
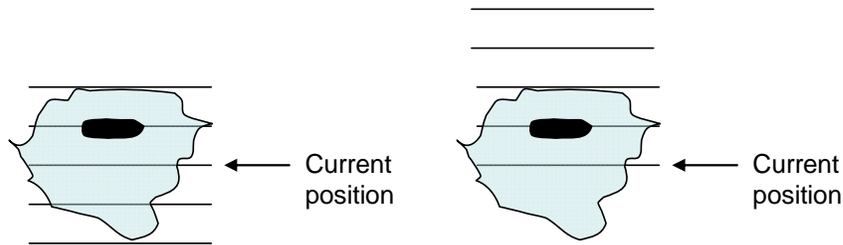
If your sample is already in focus, you may select the **Use current position** radio button and choose the **Range around current** checkbox and simply enter in the total range of travel (**Range**) and either the number of planes or the step size. The remaining capture parameters will be calculated for you. If you have set a reference position, you may choose to use the reference position as the basis for capture.

6. If you would like your z focus to return to the current position or reference position after capture check the **Return to current position** or **Return to reference position** checkbox as appropriate.
7. Click **OK** to begin capture. The Capture Controls window will appear and display each plane as it is captured. You may choose to pause, cancel, or stop your capture any time after it has started, as described above in Method 1.

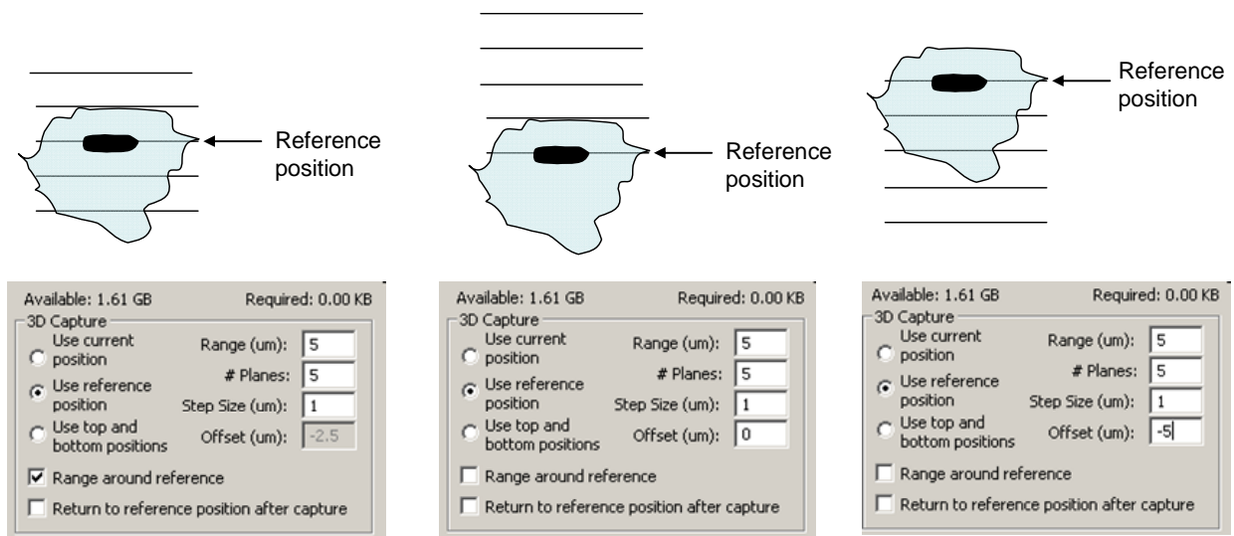
A comparison of these capture situations are shown in the figures below.



**Fig. 7.1** Method 1 is accomplished by setting top and bottom positions in the Focus Window and then setting the capture window parameters as shown.



**Fig. 7.2** Method 2 can be accomplished using either the current position or a chosen reference position. This figure shows how capture is performed when using the current position of the z stage with the given capture window parameters.



**Fig. 7.3** This figure shows how Method 2 is performed when a reference position is selected in the Focus Window. Note that the offset is different for the two situations on the right.

## 7.2 Timelapse Capture

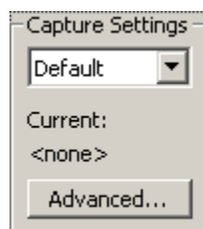
In order to perform timelapse capture, first bring a sample into view and focus as described in the [Using the Focus Window](#) section of Chapter 5 on page 85. Then, proceed as follows.

### 7.2.1 Setting Capture Preferences

Before beginning a timelapse capture, you may alter your capture preferences to do the following:

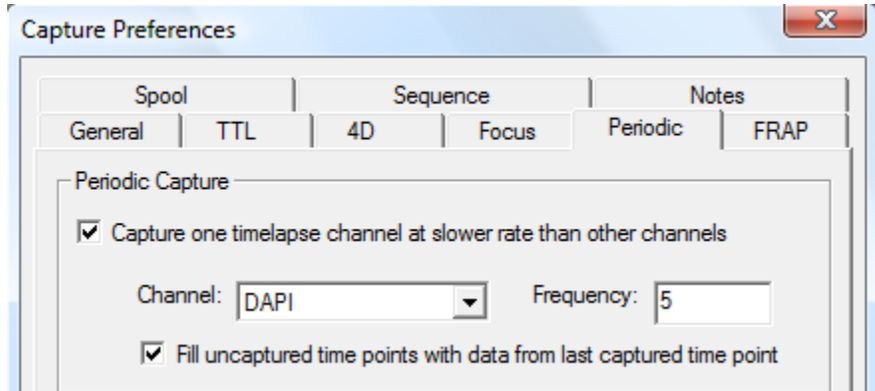
- Change capture frequency for a single channel in a multi-channel timelapse capture
- Open and close shutter between exposures for single channel capture
- Change the rate of live updates (to increase capture speed)
- Autofocus during capture
- Change capture frequency
- Save images to disk

All of these options can be accessed in the **Capture Preferences** dialog. This dialog box is displayed when you select the **Advanced** button in the **Capture Settings** portion of the **Capture** dialog.



### 7.2.1.1 Performing Periodic Capture

You may choose to capture a single channel at a slower rate than other channels when performing a multi-channel 2D timelapse capture. You may do so by bringing up the **Capture Preferences** dialog and selecting the **Periodic** tab. Select the check box, the channel that you would like to capture at a slower rate, and the frequency at which you would like to capture that channel. For example, a frequency of 5 indicates that you would like to capture a channel every 5th image. If you would like to copy the image of the selected channel to timepoints where you are not collecting the channel, select the check box to **Fill uncaptured timepoints with data from last captured timepoint**.

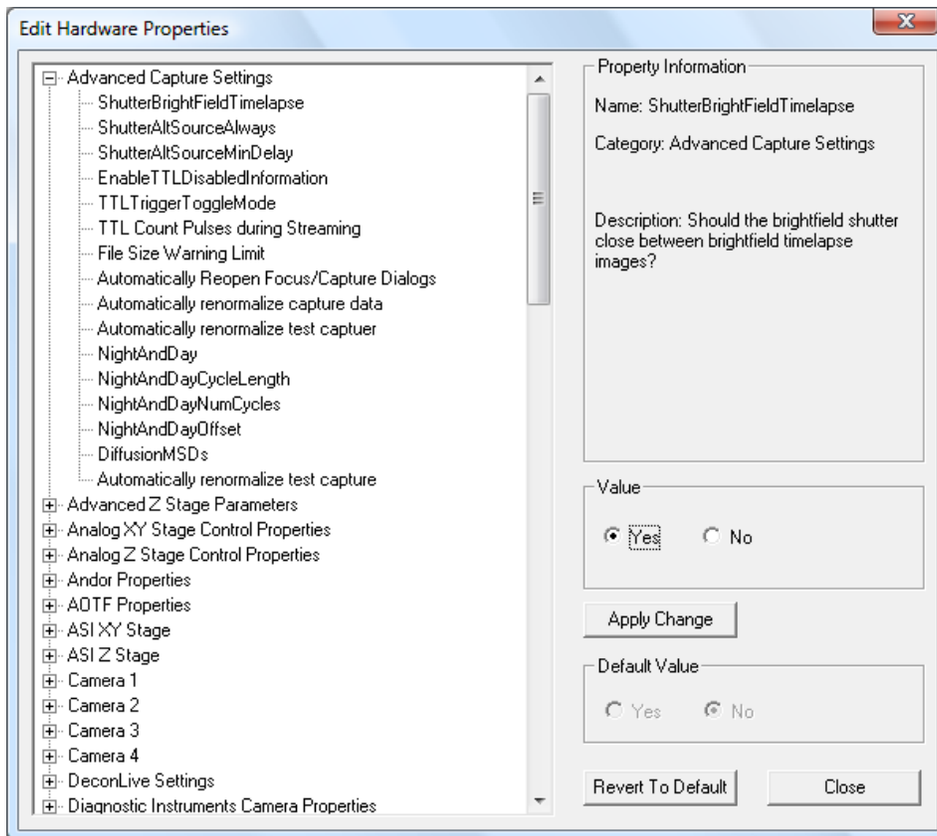


### 7.2.1.2 Opening and Closing Brightfield Shutter Between Exposures

For fluorescence shuttering, please see the above section [Setting Capture Preferences to Open and Close Shutter during Capture](#) on page 108. If you are performing 2D timelapse capture, the default fluorescence shutter action in SlideBook is to open and close the shutter if your timelapse interval is non-zero. If you have a 0ms timelapse interval, the shutter will remain open to allow you to capture as quickly as possible. This default behavior is designed to protect the sample at non-zero timelapse intervals, yet maximize acquisition speed at 0ms timelapse intervals. Thus, the **Open and close shutter on single channel or simultaneous 3D capture** checkbox is only applicable when you wish to shutter for 2D timelapse images with a timelapse interval of 0ms.

If you would like your brightfield shutter to close between transmitted light images:

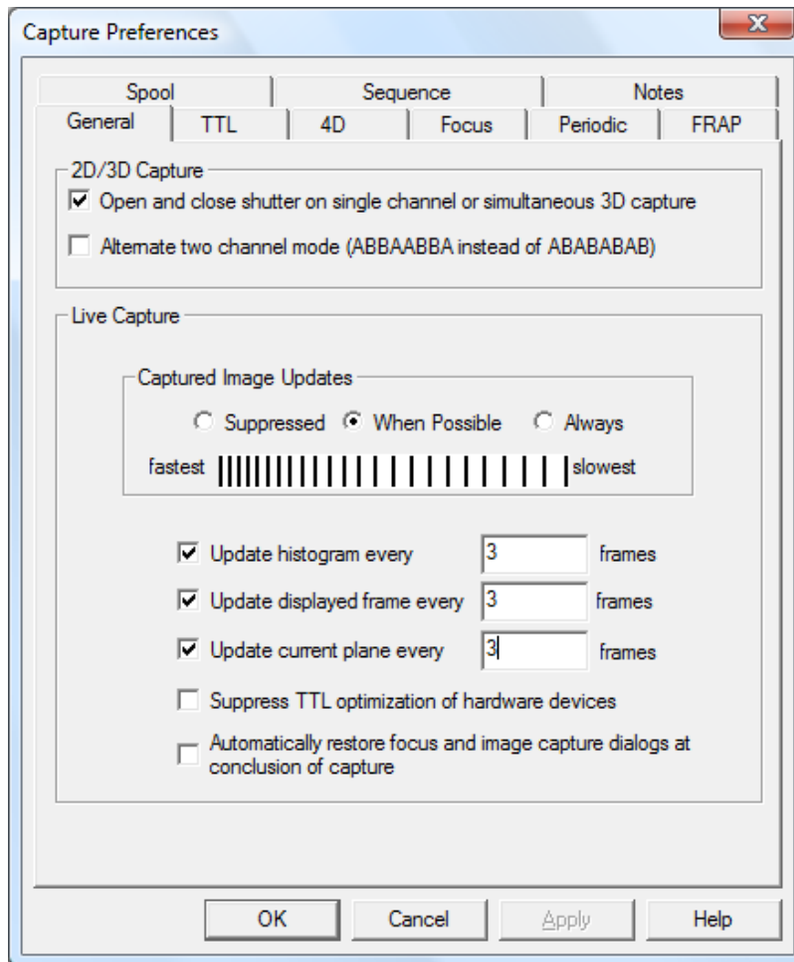
1. Select **Edit > Hardware Properties** and expand the **Advanced Capture Settings** list.
2. Click on **ShutterBrightFieldTimelapse**. The right side of the dialog box will update.



3. Select **Yes** and then **Apply Change**.
4. Select **Close** and then re-launch SlideBook to register the changes.

### 7.2.1.3 Changing the Status Update Frequency

You may change the frequency with which the elements in the **Capture Status** window are updated by choosing **Advanced** in the Capture dialog box to bring up the Capture Preferences dialog and then entering the desired update frequency in the edit boxes in the **General** tab. An update frequency of “3” would update the given element at the conclusion of every 3rd capture. Reducing the update frequency is primarily useful for increasing capture rate with single processor computers.



You may suppress all updates for optimal speed performance. To do so, select the **Suppressed** radio button. If **When Possible** is selected, updates will be provided when the computer processor allows. Some updates will be dropped in the interest of speed. If **Always** is selected, each frame requested will be displayed (at the expense of speed).

#### 7.2.1.4 Autofocus During Capture

Please see the section [Autofocus](#) on page 144.

#### 7.2.1.5 Changing Capture Frequency

Please see the section [Varying Capture Rates During Timelapse Capture](#) on page 149.

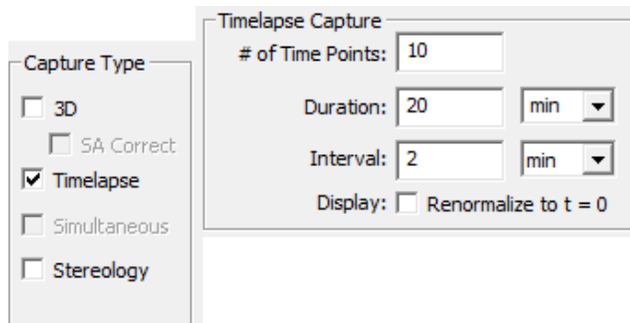
#### 7.2.1.6 Saving Images to Disk

Please see the [Saving Images to Disk \(Spooled Capture\)](#) section on page 153.

### 7.2.2 Setting Capture Window Parameters

1. Open the Capture dialog box by selecting **Image > Capture** or by selecting the capture icon in the SlideBook toolbar .

2. Follow steps 2-4 in the above section [3D Capture: Setting Capture Window Parameters](#).
3. Create any annotations that you would like to record during capture (see [Creating Notes](#) below).
4. Check the **Timelapse** checkbox in the **Capture Type** section of the Capture dialog box. You now have the option to define a variety of timelapse parameters. Enter the desired values in the following fields:



- **# of Timepoints** – Number of timepoints that will be captured.
- **Duration** – Total length of time for the experiment. Units of time, in milliseconds (ms), seconds (s), minutes (m) or hours (h), can be selected from the dropdown menu. Due to the inability to predict time required for hardware movement, this option is only available when the interval for capture is over 30 seconds.
- **Interval** – Delay between the beginning of one timepoint and the beginning of the next timepoint. The interval unit can be selected from the dropdown menu. If the capture sequence at one timepoint takes longer than the interval, SlideBook will capture the next timepoint immediately following the preceding timepoint.

**NOTE:** As you type in two of these values, the third field will be calculated automatically.

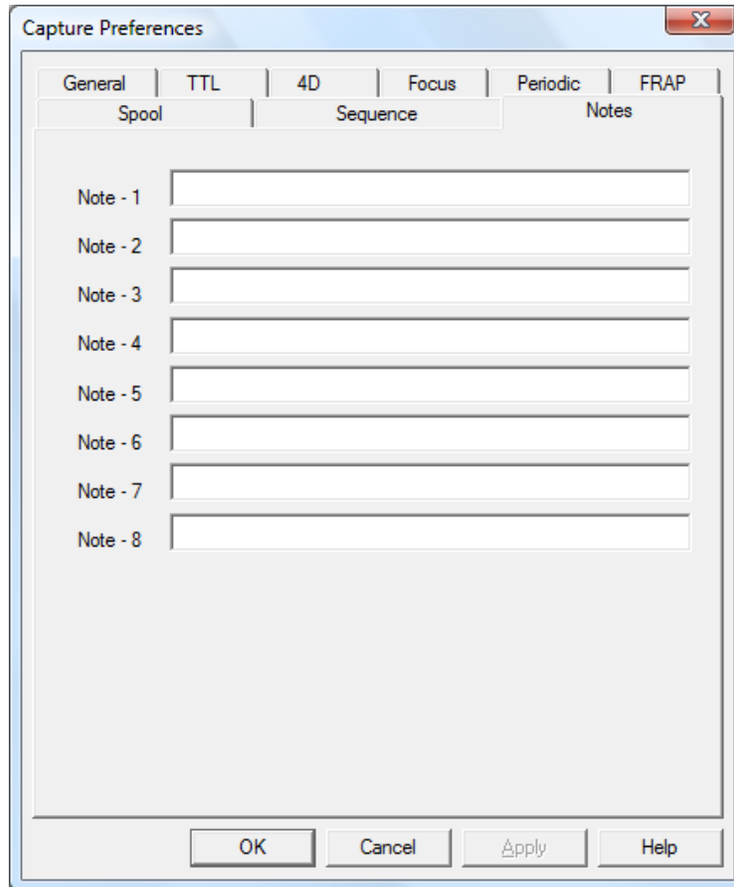
- **Renormalize to First Timepoint** – When checked, the minimum and maximum pixel intensity values for the first timepoint in a timelapse capture series will be used to determine the renormalization values of all subsequent timepoints. This option only affects the image display during capture and will not affect the captured image.

### 7.2.3 Creating Notes

SlideBook allows you to record the exact time of experimentally relevant actions. To do this, you may either create notes prior to capture or during capture. These notes will then appear as buttons in the Capture Status Window during capture. If you do not wish to create notes, proceed to the next section, [Initiating and Monitoring Timelapse Capture](#).

---

1. To create annotations, first bring up the Capture Preferences dialog from the Capture dialog by clicking on the **Advanced** button.
2. Select the **Notes** tab. The following dialog box will appear.

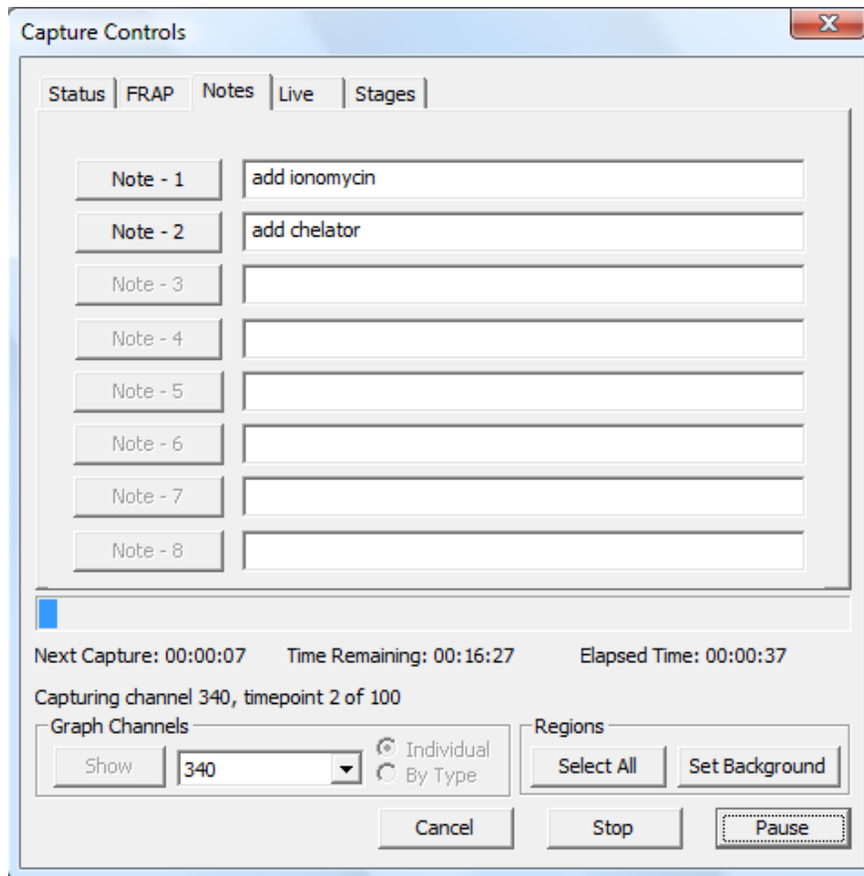


You may specify up to eight notes, which will appear in Capture Controls and the exported data.

3. Click **OK** once you have entered the desired annotations.

Once capture is initialized (by clicking **OK** in the **Capture** dialog box), the notes will appear in the Capture Status Window (see below).

4. To record an event, simply click on the desired annotation button during capture. You can access notes from the **Notes** tab of the Capture Controls. You may also add new notes at this point. You need not specify the notes before capture.



**NOTE:** You can make multiple clicks during capture (even of the same event), but currently only one event can be recorded with any single timepoint. After capture, elapsed time and timelapse annotations can be displayed as a text overlay in the image by selecting **Annotations > Timestamps**. In addition, these events are displayed next to the annotated timepoint when data is exported using **Statistics > Ratio/Timelapse Data**.

## 7.2.4 Creating ROIs and Graphs to Monitor Regions of Interest

Before beginning your capture, you may wish to create regions of interest in order to monitor intensities during. You may also do this during capture. To draw regions before capture, perform the following:

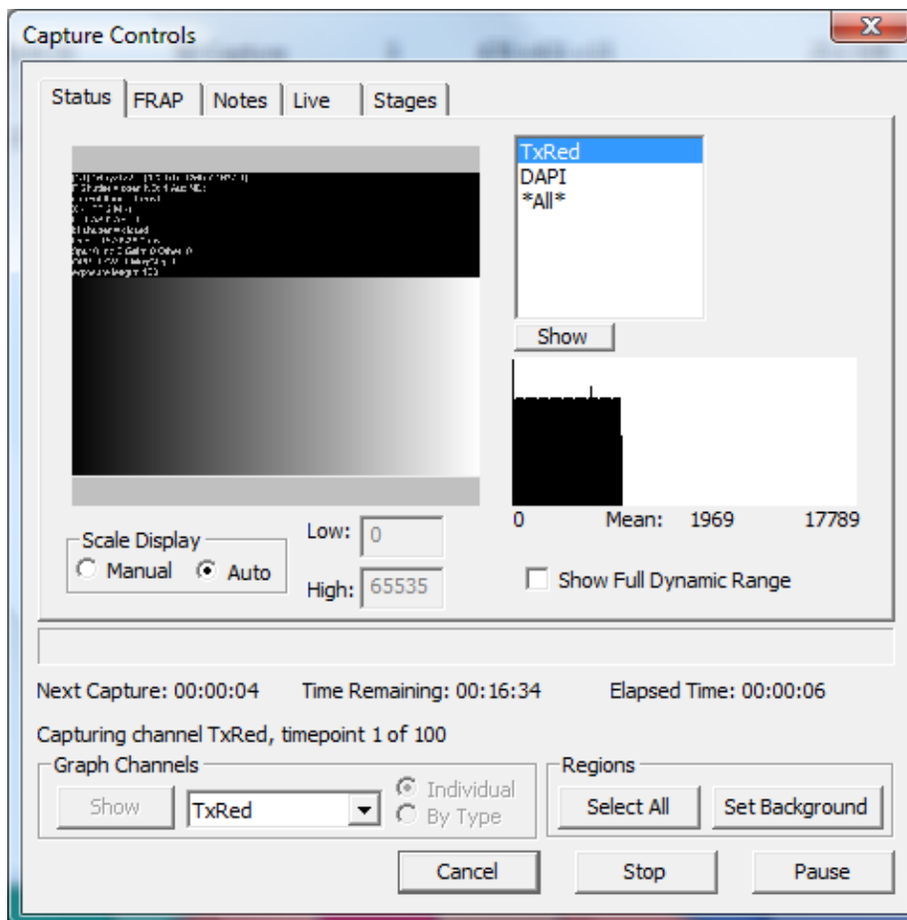
1. Make sure that the Live View is displaying the specimen. You can do this by pressing **Test** in the Capture dialog box.
2. Choose one of the ROI tools, which are located on the SlideBook toolbar at the top of the application window.
  -  - Rectangle Tool, click and drag to draw a rectangle
  -  - Ellipse Tool, click and drag to draw an ellipse

-  – Polygon Tool, click to draw a series of straight lines, double-click to close the shape
  -  – Free Hand Tool, click and drag to draw freehand, release mouse button to close the shape
  -  – ROI Selection Tool, use to select an ROI. Holding shift and clicking on multiple regions will allow you to select more than one region. Clicking and dragging on a selected region will move the selected ROIs. **If the ROI number is displayed in white, this indicates that the ROI is selected.**
3. Continue to draw regions as desired. The following operations can be performed by selecting the ROI with the ROI selection tool, and then right-clicking to bring up the ROI menu.
    - **Set as background** – uses the selected ROI for background subtraction during graphing
    - **Duplicate** – creates a copy of the selected ROI
    - **Delete** – removes the selected ROI (s)
  4. Once you have created the desired regions of interest and begun capture, you can display a graph by selecting the desired channel in the **Graph Channels** section at the bottom left of the Capture Controls and then clicking on the **Show** button.

The regions of interest that you have created can also be displayed after capture as discussed in [Displaying Annotations](#) on page 181. To recreate the graph post-capture, please see page 251.

### 7.2.5 Initiating and Monitoring Timelapse Capture

In the Capture dialog box, click **Start** to begin capture. The Capture Controls will appear and display the current single channel image in the **Status** tab. The Live View will show a larger, resizable version of the current capture which can combine multiple channels.



### 7.2.5.1 Creating Graphs to Monitor Regions of Interest

If you have created ROIs, you may choose to display intensity graphs. To do so:

1. Select the channel that you wish to graph from the drop-down in **Graph Channels** section and click **Show**. You may choose to display the graphs for multiple channels individually or by type.



2. Use the *zoom in* and *zoom out* tools to explore your data.
3. Once your capture is finished, your graph will remain open for viewing.
4. To export your graph, simply select the icon.

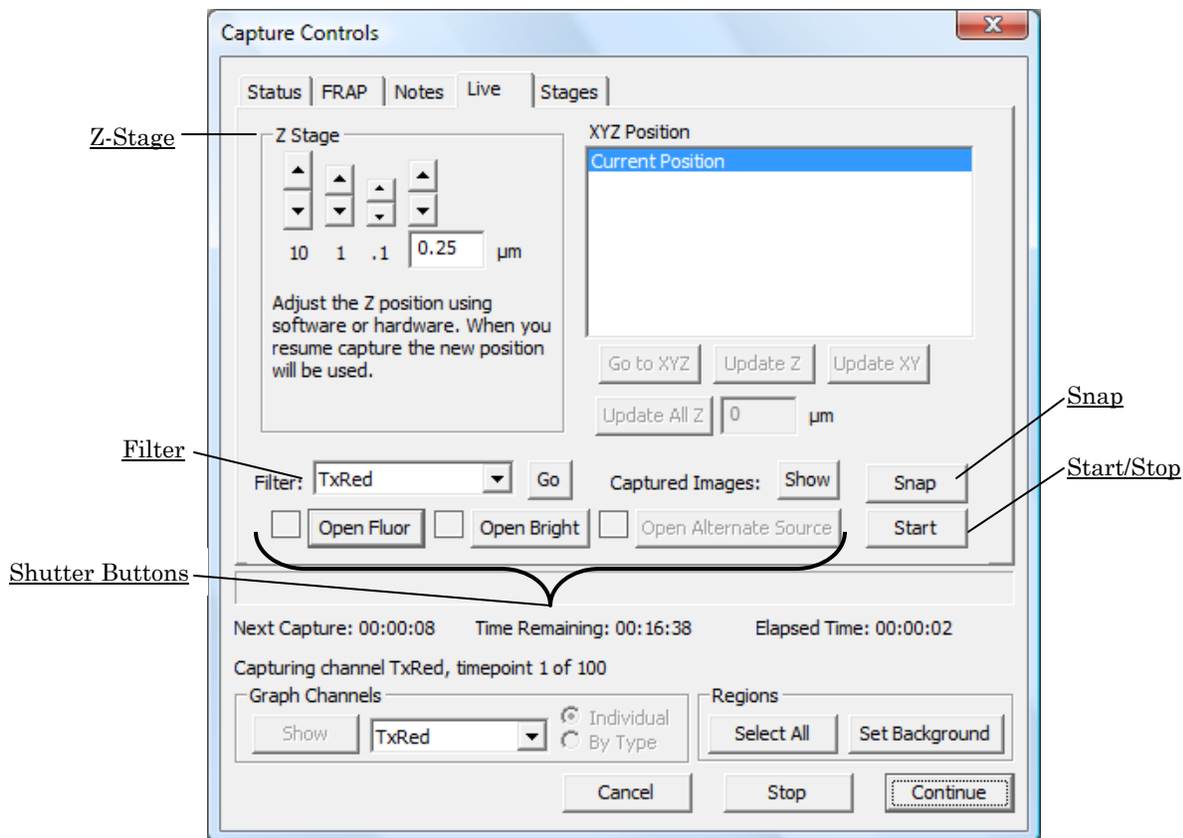
The regions of interest that you have created can be displayed as discussed in [Displaying Annotations](#) on page 181. To recreate the graph post-capture, please see page 251.

### 7.2.5.2 Focusing During Capture

You may adjust the focus of your image during capture. To do so, go to the **Live** tab. The **Live View** works similarly with the Capture Controls as it did with the Focus Controls.

**NOTE:** Before using the **Live** tab, you must first select **Pause**. Once you are finished focusing, you must select **Continue** to resume capture.

The window has the following features:



- **Start / Stop** – begins (or ends) the semi-live camera readout
- **Snap** – snaps a single image
- **Shutter Buttons** – toggle fluorescence, brightfield, or alternate source shutters open and closed, boxes to the left of the buttons indicate status
- **Filter** – moves motorized filters into position when you select **Go**
- **Z Stage** – allows you to refocus your sample using precise movements

To use this window, press **Start**, open the desired shutter, select the appropriate filter, and focus your sample. When you are finished, close the shutter, press **Stop**, and then **Continue**. Alternately, you can press **Snap** to check a single focal plane then adjust the z-stage, then **Snap** again to update the view. The later approach may minimize exposure to the sample.

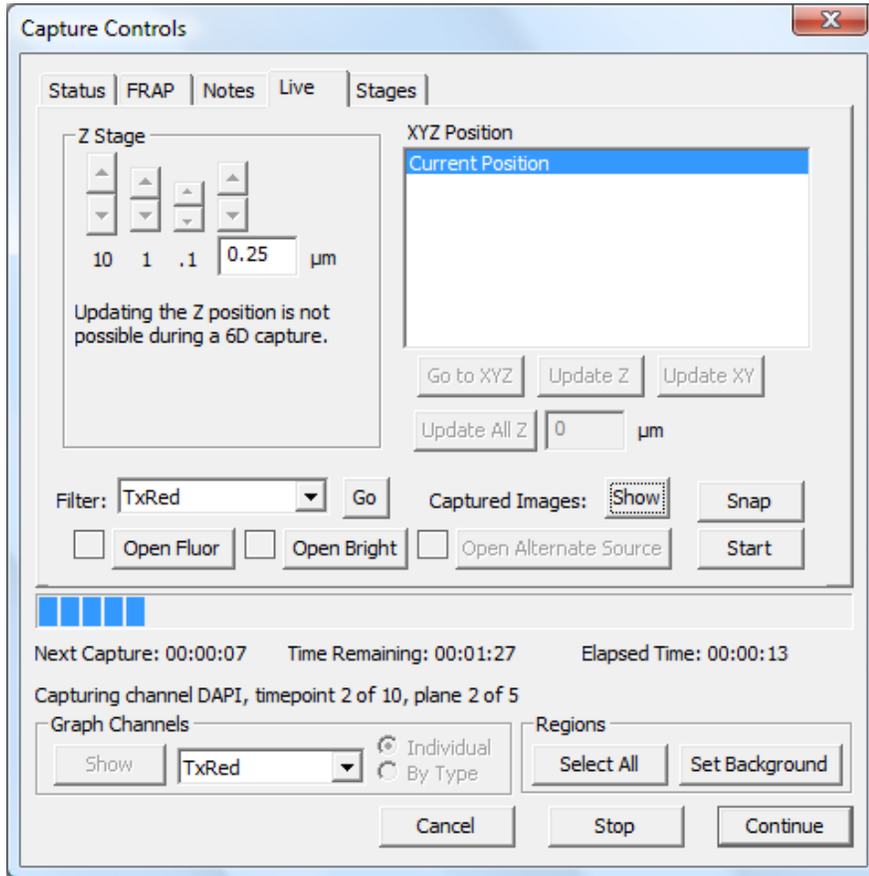
### 7.2.5.3 Viewing Previous Timepoints

You may choose to view previously captured images while your 2D timelapse or 4D experiment is in progress. To do so:

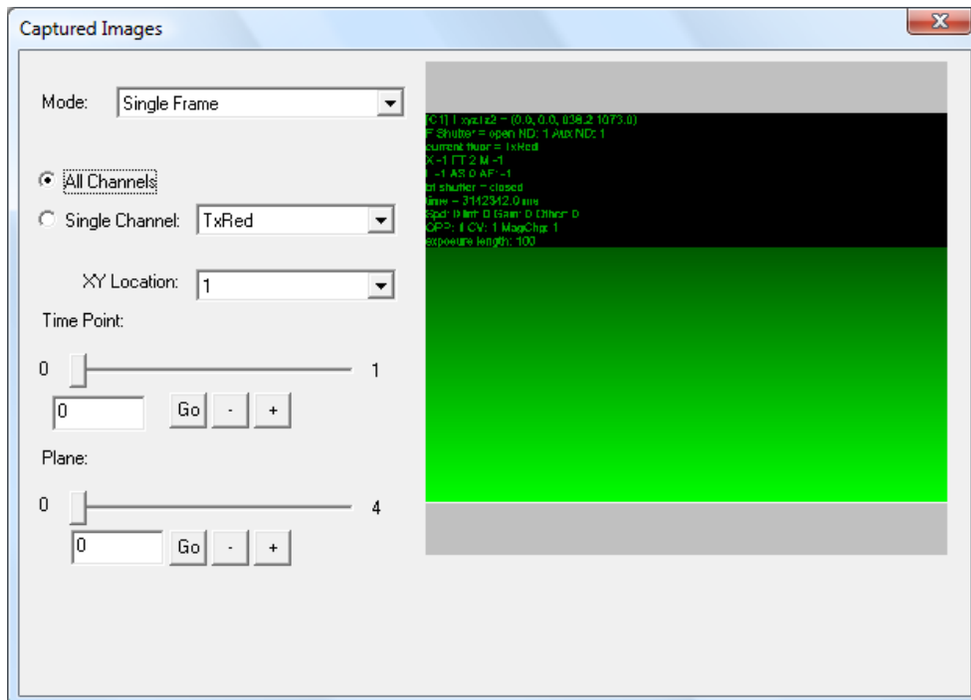
1. Set up your capture as you would normally in the Capture dialog box. Make sure that you are saving images to memory, or to memory and disk (see page 149).

You can do this by selecting **Advanced** to bring up the Capture Preferences dialog. Then, select **Spool** and be sure that one of the first two radio buttons is selected.

2. Once you have set up your capture, click **OK**. The Capture Controls will appear.
3. Select the **Live** tab.



4. Select the **Show** button next to **Captured Images** and the following window will appear:



5. Use the slider to scroll through previously captured images or enter a specific timepoint into the edit field and select **Go**. You may also use the **+** and **-** buttons to move one timepoint at a time.
6. If you would like to view a single channel, select the **Single Channel** radio button and the desired channel from the dropdown list.
7. The default mode is “Single Frame.” If you are capturing a 3D stack over time, you may also choose **MIP** to view a maximum intensity projection of each z stack at any timepoint.
8. Once you are finished viewing images, close the dialog box by clicking on the **X** in the upper right hand corner of the dialog box or selecting **Hide** from the Live tab.

### 7.3 4D Capture

SlideBook allows you to capture 3D images over time. To set up for 4D capture simply set up for 3D capture and timelapse capture as described in the above sections, then make sure that both the **3D** and **Timelapse** checkboxes are selected in the Capture dialog box and the appropriate parameters are entered in the edit fields before selecting **OK**. The resulting slide file will contain a single 4D image that includes all of the z sections and timepoints.

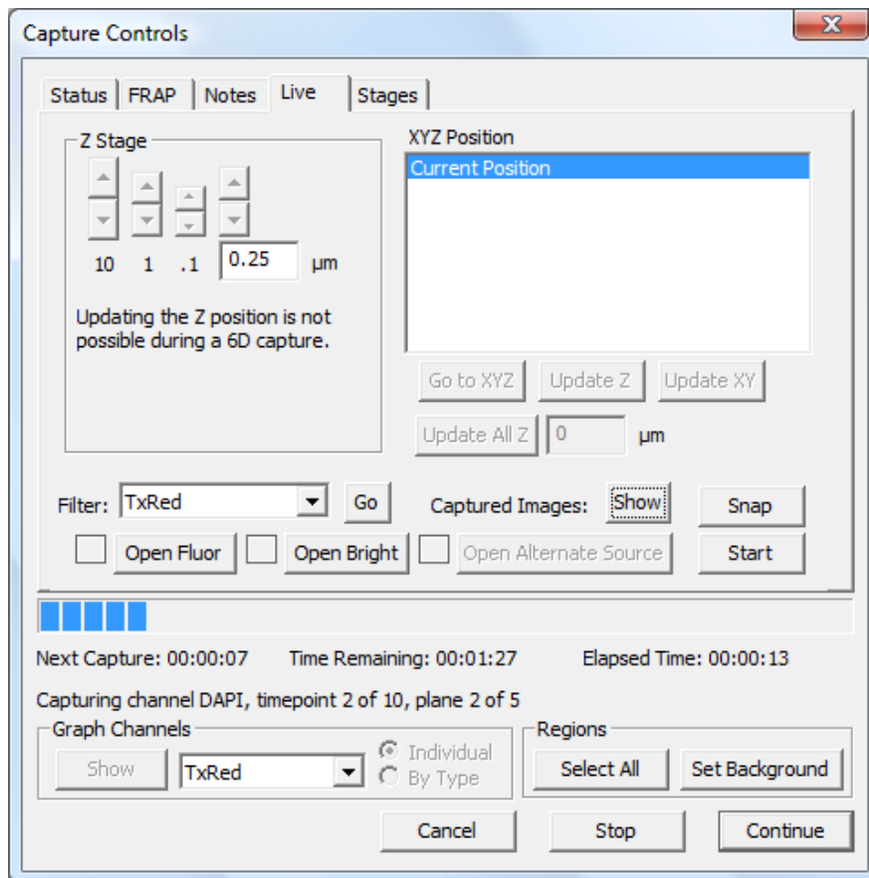
**NOTE:** If you have specified a timelapse interval that is shorter than the time required to capture a single 3D stack, SlideBook will default to capturing 3D stacks as quickly as possible.

In addition to all of the options available for 2D timelapse imaging, SlideBook also allows you to perform mid-volume capture and sequential multi-channel z series, as discussed below.

### 7.3.1 Shifting the Volume During 4D Imaging

You may find that your sample exhibits focus drift during image acquisition. You may correct for this by selecting the **Live** tab during capture. To move your volume:

1. Click on the **Live** tab and press Pause to pause the capture.
2. Click **Start** to go into “live” mode in the Live View. Refocus your sample either manually or by using the **Z Stage** controls and select **Recenter Z**. You may also shift the whole 3D volume by checking the **Shift 3D Volume** box.

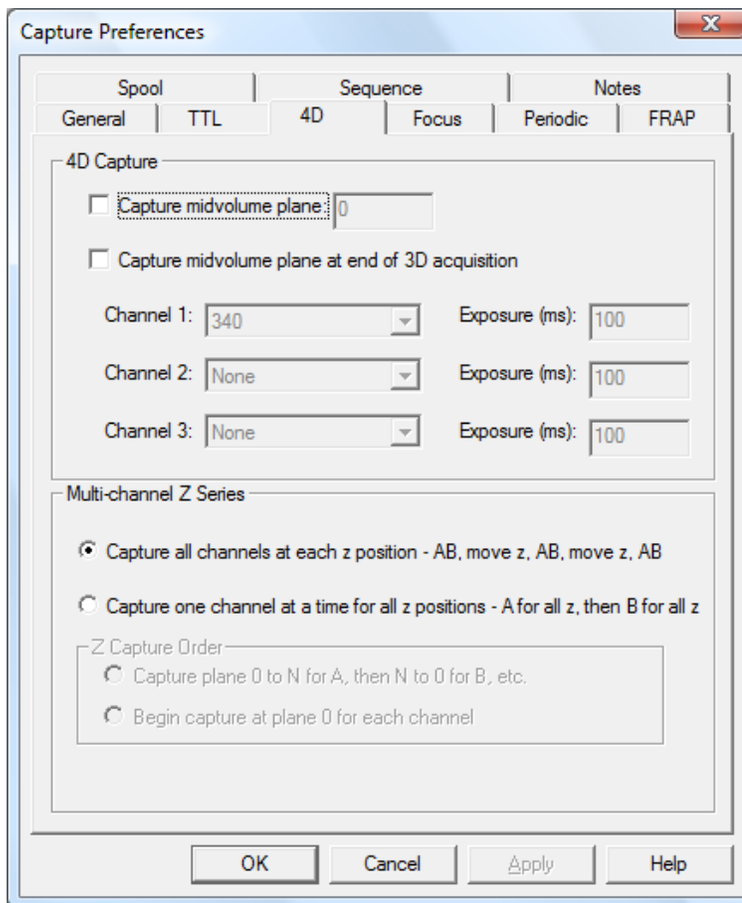


### 7.3.2 Mid-volume Capture During 4D Imaging

SlideBook offers the ability to capture up to three 2D mid-volume images during each timepoint of a 4D capture. This feature is useful for acquiring channels that often do not need to be imaged in three-dimensions (such as DIC or Fura-2) during a 4D capture. To setup a 4D mid-volume capture:

1. Go to Capture Preferences dialog box by pressing the **Advanced** button in the Capture dialog box.

2. Select the **4D** tab from the capture preferences dialog box. The following window will appear:



3. Select the **Capture midvolume plane** checkbox. This will activate the edit box to the right allowing you to specify the plane at which a midvolume image will be acquired. The midvolume plane must fall within the plane limits of the 4D capture as specified in the Capture dialog box.
4. SlideBook defaults to capturing the midvolume plane during each 3D acquisition. For instance, consider the case of a 4D capture with a 15 plane 3D image being acquired at every timepoint. If the midvolume capture plane is set to 10, SlideBook will acquire 10 planes of the 3D image, acquire the midplane channel and then acquire the remaining 5 planes of the 3D image. To have SlideBook capture the midvolume image at the end of the 3D capture select the **Capture midvolume plane at end of 3D acquisition** checkbox. When this option is selected, SlideBook will capture the entire 3D volume, return to the specified midvolume plane and then capture the selected channels.
5. Select the midvolume channels that will be captured at each timepoint during the 4D capture. In the edit box to the right of each channel specify the exposure time that will be used to capture the channel. Importantly, channels that will be

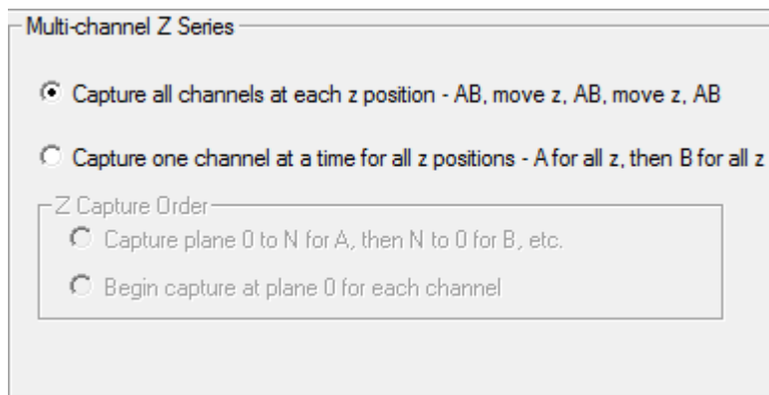
captured at midvolume planes should not be selected for exposure in the Capture dialog box when setting up the 4D capture parameters.

- Click **OK** and then proceed to the Capture dialog box to set up for 4D capture as described above.

### 7.3.3 Multi-Channel Z-Series

In order to speed up capture, you may wish to capture multi-channel z stacks one channel at a time.

- Go to **Capture Preferences** dialog box by pressing the **Advanced** button in the Capture dialog box.
- Select the **4D** tab from the capture preferences dialog box. The lower half of the dialog box appears as follows:



- Select the desired capture order. If you would like to reduce the number of filter switches that occur during capture, select **Capture one channel at a time for all z positions**.
- Click **OK** and then proceed to the Capture dialog box to set up for 4D capture as described above.

## 7.4 Multipoint Capture

If your stage is equipped with a motorized *xy* stage, SlideBook allows you to select multiple locations to visit throughout the course of capture. You may perform timelapse and/or 3D capture at each location. As with the other types of capture, you will set parameters in the focus window and in the capture window.

### 7.4.1 Setting Capture Preferences

Depending on the type of capture you plan to perform (2D timelapse, 3D or 4D), you may wish to select capture preferences such as periodic capture, midvolume capture, multi-channel *z* options, autofocus and spooled capture as described in the sections above and below (see pages 113, 115, 125, 127, and 146).

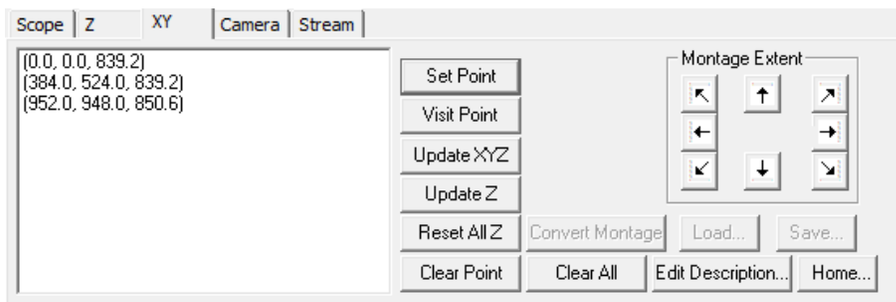
## 7.4.2 Setting Focus Window Parameters

You may choose to select new points, or you may load points that you have saved previously.

### 7.4.2.1 Setting Points

1. Open the Focus Window by either selecting **Window > Focus Window** or clicking on the focus window icon in the SlideBook toolbar . The Live View will open and begin updating.
2. Select your objective, magnification changer, and filter configuration, and bring your sample into view and focus as discussed in [Chapter 5 - Controlling the Camera and Microscope Hardware \(Focus Window\)](#) on page 77.
3. To select a field to be visited during capture, first click on the **XY** tab.
4. Choose **Set Point** to add a field of view to be visited during capture. The *xy* location coordinates will be added to the window of the **XY** tab.

**NOTE:** If your system also has encoded *z* focusing, you may specify *xyz* coordinates.



Once you have added a location to the list, you may remove, visit, or add locations. To perform these operations, first highlight the location by clicking on the coordinates in the list, then click on the appropriate button to perform the following actions:

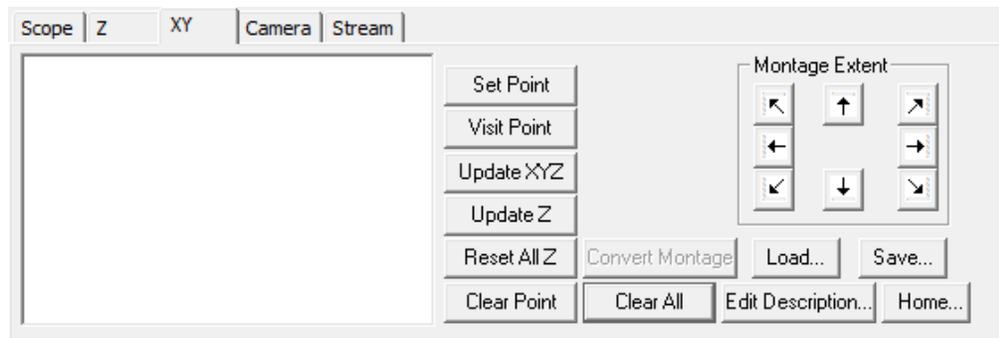
- **Visit Point** – moves the stage (and *z* focus) to the selected *xyz* location.
- **New Point Z** – allows you to alter the *z* position of a given *xyz* location. This allows you to update the focus position of a certain sample if it has drifted.
- **Reset All Z** – resets the *z* location on all points to the current *z* location.
- **Clear Point** – removes the selected location from the list.
- **Clear All** – clears all locations from the list.

5. Once you are satisfied with your list of locations, close the lightpath for light sensitive samples by choosing **Close Fluor**.
6. Return to the brightest field that you are planning to capture by selecting the location from the list in the **XY** tab and choosing **Visit Point**. It is important that the brightest field is in view when setting exposure times in the Capture dialog box.

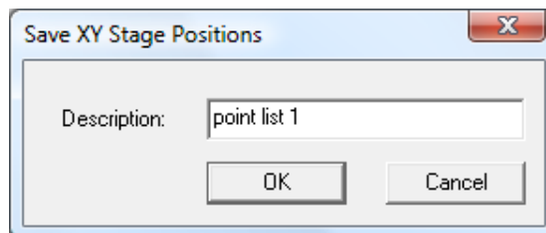
#### 7.4.2.2 Saving and Loading a Multipoint List

This feature allows you to save the *xyz* multipoint list so that you may visit the same points in future experiments. To do so:

1. Before placing your sample on the stage, open the Focus Controls and go to the **XY** tab.



2. Before setting points, you will need to home the stage. Some stages do not maintain an absolute coordinate system, but rather reset to zero at whatever position the stage is in when it is turned on. If it is your first time performing the home operation with the stage, make sure that your objectives are in the “load” position, as far from the stage as possible. Once you have all possible obstructions cleared, select **Home**. If **Home** is grayed out, you must manually home the stage by taking the stage to its limit at any of the four corners, then resetting your stage to zero (for instance, on an ASI stage, you would take the stage to the limit, then turn the stage off and on to zero the stage). You must perform this procedure each time you plan to save or load a multipoint list.
3. Once the stage is homed, place your sample on the stage and proceed with setting your point list. Once the list is complete, select **Save**. You will be prompted to enter a description for your point list.



4. Enter a description and then select **OK**.

Now that you have saved XY stage positions, you can load them for subsequent experiments. To load them, simply home the stage before loading your sample, then select **Load** to populate the point list.

### 7.4.3 Setting Capture Window Parameters

1. Once you have set the appropriate parameters in the Focus Window, open the Capture dialog box by selecting **Image > Capture** or by selecting the capture icon in the SlideBook toolbar .
2. Follow steps 2 through 4 in the above section [3D Capture: Setting Capture Window Parameters](#) in order to select channels to be captured and set parameters for image extent and binning.
3. Click on the **Entire List/Montage** radio button in the **Multiple XY Location Capture** section. Alternately, if you would only like to capture the current point, you may choose the **Current Location** radio button.

If you wish to take a single image at each location, proceed to step 8.

#### 7.4.3.1 3D Multiple Location Capture

4. If you would like to capture 3D stacks at each location, set up for 3D capture as discussed in [3D Capture](#) on page 105.
5. Click on the **3D** checkbox in the **Capture Type** section.

#### 7.4.3.2 Timelapse Multiple Location Capture

6. If you would like the points to be visited repeatedly, click on the Timelapse checkbox in the **Capture Type** section of the Capture dialog box.
7. Enter the number of timepoints to be collected and the interval between capture in the appropriate edit fields.

**NOTE:** In multiple location timelapse capture, the timelapse interval corresponds to the time that elapses between the completion of capture for the last point in the list and the beginning of the capture for the first point in the list.

8. Click **OK** to begin capture. The Capture Status Window will appear and display each plane as it is captured. You may choose to pause, cancel, or stop your capture any time after it has started.

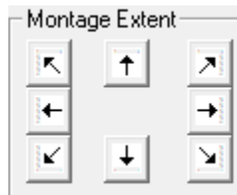
## 7.5 Montage Capture

Often, you may wish to capture an image that is larger than a single field of view. SlideBook can stitch multiple fields together into what is called a *montage* image. It does this by taking successive camera fields and moving the motorized *xy* stage a computed amount. The procedure for capturing a montage image is described below.

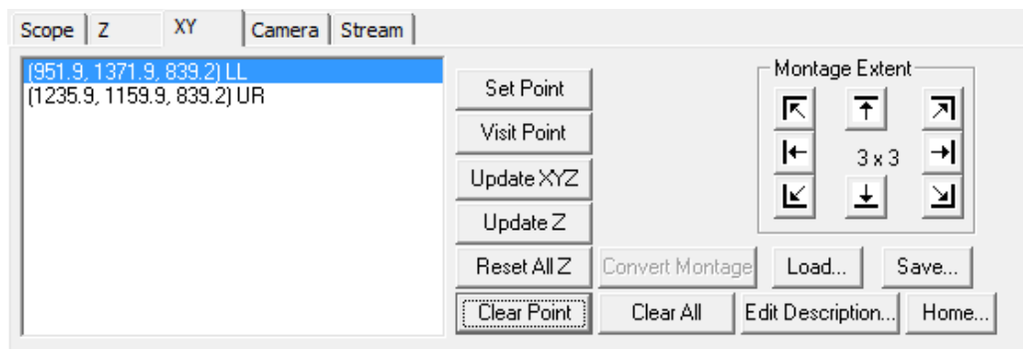
**NOTE:** Before attempting to perform montage capture, make sure your *xy* stage is properly configured as discussed in [Configuring a Motorized XY Stage](#) on page 58.

### 7.5.1 Setting Focus Window Parameters

1. Open the Focus Window by either selecting **Window > Focus Window** or clicking on the focus window icon in the SlideBook toolbar . The Live View will begin updating.
2. Select your objective, magnification changer, and filter configuration, and bring your sample into view and focus as discussed in [Chapter 5 - Controlling the Camera and Microscope Hardware \(Focus Window\)](#) on page 77.
3. The **XY** tab of the focus window has a panel of buttons that assist in selecting the range of the montage:



4. These buttons are used to define the boundaries of the entire region to be included in the montage image. SlideBook needs enough coordinate information to generate an imaginary “box” around the region to be imaged. Thus, the extent of the montage image can be defined by selecting one of the following sets of coordinates: upper left/lower right, upper right/lower left, top/bottom/left/right. In the following example the upper left and lower right buttons are used to define the montage extent.
5. Navigate to the upper right corner of the region that you want to image by either moving the stage controller’s joystick or using the up, down, left, and right arrows that appear in the Focus Controls.
6. Click on the  button. The *x*, *y*, and *z* coordinates of that position will appear in the **XY** tab’s list of points with a **UR** designation next to it.
7. Now, search for the lower left corner of the selection and click on the  button. The *x*, *y*, and *z* coordinates of that position will appear in the **XY** tab’s list of points with a **LL** designation next to it.



Selecting these two corners lets SlideBook automatically compute how many camera fields need to be captured in order to stitch together a montage image that will cover the entire region. When a montage extent has been successfully defined, the number of vertical and horizontal camera fields needed to cover the region will be displayed (in this case a  $2 \times 3$  grid of 6 images). If **X axis?** or **Y axis?** is displayed, SlideBook cannot compute a montage extent because the selected coordinates are not correctly orientated with respect to one another. For instance, **Y axis?** would be displayed if the selected lower right position is actually located above the selected upper left position. If you continue to see these errors, please check that your *xy* stage is configured properly (page 58).

If you want to change the montage extent, you can simply go to a different location and click on one of the buttons again, as appropriate. The previous coordinates will remain on the list of points, but the **LL** or **UR** designation will move to the new point.

Once you have added the locations to the list, you may visit or adjust them. To perform these operations, first highlight the location by clicking on it and then click on the appropriate button to perform the following actions:

- **Visit Point** – moves the stage (and *z* focus) to the selected location
  - **Clear All** – clears all locations from the list
  - **Reset All Z** – resets the *z* coordinate of all locations to the current *z* position
  - **New Point Z** – resets the *z* coordinate of the selected location to the current *z* position
8. Once you have selected two corner boundaries, close the lightpath for light sensitive samples by choosing **Close Fluor**.

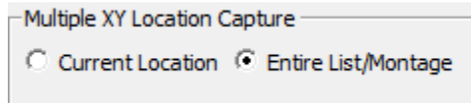
### 7.5.2 Setting Capture Window Parameters

1. Once you have set the boundaries of the montage image in the Focus Window, open the Capture dialog box by selecting **Image > Capture** or by selecting the capture icon in the SlideBook toolbar .

2. Select channels and set exposure times as described in the section [Selecting Channels and Setting Exposure Times](#) in Chapter 6 on page 89.
3. Enter image information and settings as described in the section [Entering Image Information and Optical Parameters before Capture](#) in Chapter 6 on page 93.

**NOTE:** It is critical to verify that the **Objective** and **Mag. Changer** fields under **Optical Parameters** are set correctly. SlideBook uses its knowledge of the size of the camera field at the current magnification to compute the number of images needed to span the montage extent. If your microscope has an automated objective nosepiece, SlideBook can read its position and the **Objective** field should already be set correctly. If it has an automated magnification changer turret (or only a single magnification changer setting) then the **Mag. Changer** field should also be set correctly. However, if your microscope has manual selection of objectives or more than one manually selected magnification changer, you must make sure the **Objective** and **Mag. Changer** fields are correct before proceeding.

4. Click on the **Entire List/Montage** radio button in the **Multiple XY Location Capture** section of the Capture dialog. The image extent is adjusted to reflect the montage extent in pixels.



**NOTE:** If the **Entire List/Montage** button is not enabled, a montage extent was not properly defined in the Focus Window.

5. Click on the **OK** button to start the capture.

SlideBook will return the *xy* stage to the upper left corner and begin its acquisition sequence. The Capture Status Window will appear and display each plane as it is captured. You may choose to pause, cancel, or stop your capture any time after it has started. Once the capture is complete, the current slide will contain images of all of the fields that contributed to the montage. To generate a montage image please see page 169, [Displaying a Montage](#).

## 7.6 MultiWell Capture

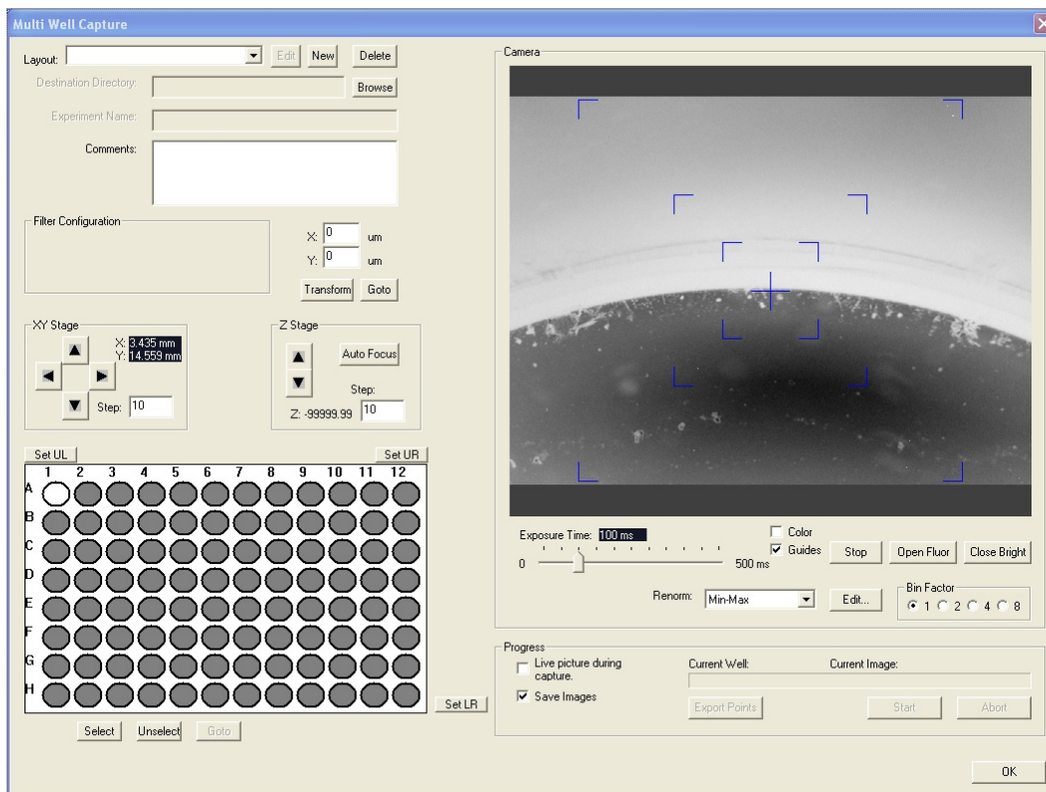
Multi-well capture uses a different interface than other forms of capture. In this section, you will learn how to:

- Create a Layout
- Manipulate Hardware and Focus on the Sample
- Calibrate the Stage Position
- Select Wells for Capture
- Initiate and Monitor Capture

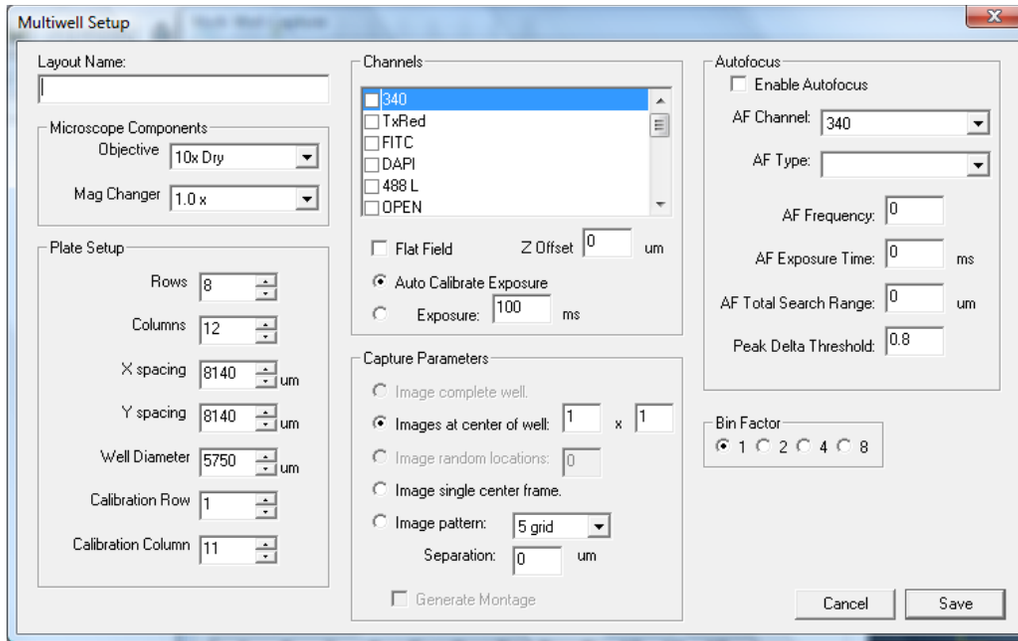
### 7.6.1 Creating a Layout

In order to perform multi-well capture, you must first create a *layout*, or template, that defines capture parameters. These layouts may be saved and applied for future captures. To create a layout:

1. Go to **Image > Advanced Operations > Capture Multi-well Plate**. The **Multi Well Capture** dialog will appear:



2. Go to the upper left corner and select **New** to generate a new layout. The following dialog will appear:



3. Enter the **Layout Name** in the edit field. This layout will be saved so you may use it again for future experiments.
4. Select the **Microscope Components** from the **Objective** and **Mag Changer** dropdown menus.
5. Enter **Plate Setup** parameters by either entering numbers or using the up and down arrows.
  - **X spacing, Y spacing** – center-to-center distance in *x* and *y* directions in microns (can be determined by contacting the plate manufacturer; values show in above dialog box are typical for a standard 96-well plate)
  - **Well Diameter** – can also be determined by contacting the plate manufacturer
  - **Calibration Row** – row used for setting top of upper left and upper right wells (see below). You can generally specify this to be 1 (e.g. row A) unless you cannot move your stage to focus on the top edge of the top row and must use row 2.
  - **Calibration Column** – column used for setting right side of lower right well (see below). You can generally specify this to be the rightmost column of the plate. For a 96 well plate often you cannot reach the right edge of the wells in the 12th column and must set this value to the 11th column.
6. Select the channels that you wish to capture by clicking in the appropriate checkboxes. You may choose to perform an autoexposure or you may set the exact exposure time.

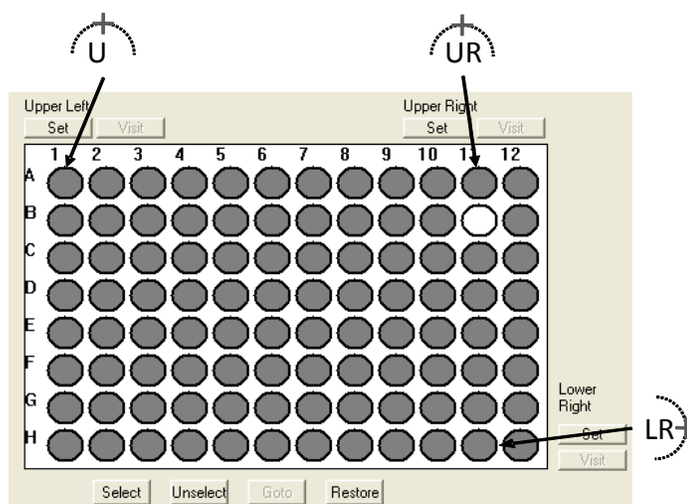
7. Select the type of capture that you wish to perform for each well as shown in the **Capture Parameters** section of the dialog. You have the following choices:
  - **Images at the center of the well** - you may choose the size of the grid, and you may generate a montage of the grid
  - **Single center frame**
  - **Image pattern** - for the predefined grid pattern, you may enter a separation value, which will allow you to sample at various points in the well
8. Select autofocus parameters. You may wish to determine these parameters first using the Focus Window (see [Autofocus](#) on page 144.)
9. Click **Save** to exit the Multiwell Setup dialog box.

### 7.6.2 Manipulating Hardware, Focusing on the Sample

Once you have determined your layout, you can interface with hardware in the Multi Well Capture dialog. This works very similarly to the Focus Window. You can select filters, move  $x,y$ , and  $z$  positions, and open the fluorescence or transmitted light shutters. Before proceeding to the next section, make sure that your sample is in view and focus.

### 7.6.3 Calibrate the Stage Positions

1. Go to a column towards the left of the plate and find the top of a well in the calibration row (as defined by **Calibration Row** in your layout, generally row A).
2. Focus on the upper middle edge of the meniscus of the well and center it in the camera image (see picture below), and click **Set UL**.
3. Go to a column towards the right of the plate and find the top of a well in the calibration row (as defined in your layout, generally row A).
4. Focus on the upper middle edge of the meniscus of the well and center it in the camera image (see picture below), and click **Set UR**.
5. Go to a row towards the bottom of the plate in and find the right of a well in the calibration column (as defined by **Calibration Column** in your layout, column 11 by default).
6. Focus on the right middle edge of the meniscus of the well and center it in the camera image (see picture below), click Set next to Lower Right.
7. Test calibration by clicking on a well (it will turn white), and selecting **Goto**.



#### 7.6.4 Save and Restore Calibration Positions

If the currently selected layout has been calibrated before, then you can use the **Restore** button to load the previously set **Upper Left**, **Upper Right** and **Lower Right** positions. Caution: Your *xy* stage **MUST** be **Home'd** before you set the positions or they will not be saved (see [Setting Points](#) on page 128). After you restore calibration positions you should use the corresponding **Visit** buttons to view and then the **Reset** button to adjust for changes in the plate placement.

#### 7.6.5 Selecting Wells for Capture

1. Select the wells you wish to image by clicking on the well and then clicking **Select**. If you wish to select multiple wells, first select a single well (it will be gold). Starting from the selected well, click and drag over the wells you want to image. The selected wells will turn gold.
2. To unselect wells, click on the well that you don't want to image and press **Unselect**. To unselect multiple wells, first unselect a single well, then click and drag over the wells you wish to unselect. The unselected wells will turn gray.

#### 7.6.6 Initiating and Monitoring Capture

To begin imaging, you will first make sure that the first well is in focus.

1. Go to the first well that you will capture (by clicking on the first well and using the **Goto** command).
2. Open the desired light path by clicking **Open Fluor** or **Open Bright**.
3. Select the desired filter from the **Filter Configuration** section.

4. Obtain focus on your sample in the camera view.
5. If you would like to view images as they are captured, select the **Live Picture during Capture** checkbox.
6. Click **Start** to begin capture.

### 7.6.7 Exporting Point List

To capture 3D images or to take advantage of other advanced capture options you must use the **Export Points** button to transfer the list of positions for each selected well to the **XY** tab of the **Focus Controls** window. Once transferred, you can perform a multipoint capture using the exported points (see Multipoint Capture on page 127). The well name and position index within the well will be saved as the image name and spool file name for each image captured.

### 7.7 Simultaneous Capture

SlideBook allows for simultaneous dual camera or image splitter operation for systems so equipped (see [Configuring an Image Splitter](#) on page 60 and [Defining Filters when using an Image Splitter](#) on page 72). Simply set up for the type of capture that you would like to perform as described in the sections above. Before initiating capture, make sure that the **Simultaneous Capture** checkbox is selected in the **Multiple XY Location Capture** section of the Capture dialog box.

### 7.8 Configuring Your Hardware for Speed

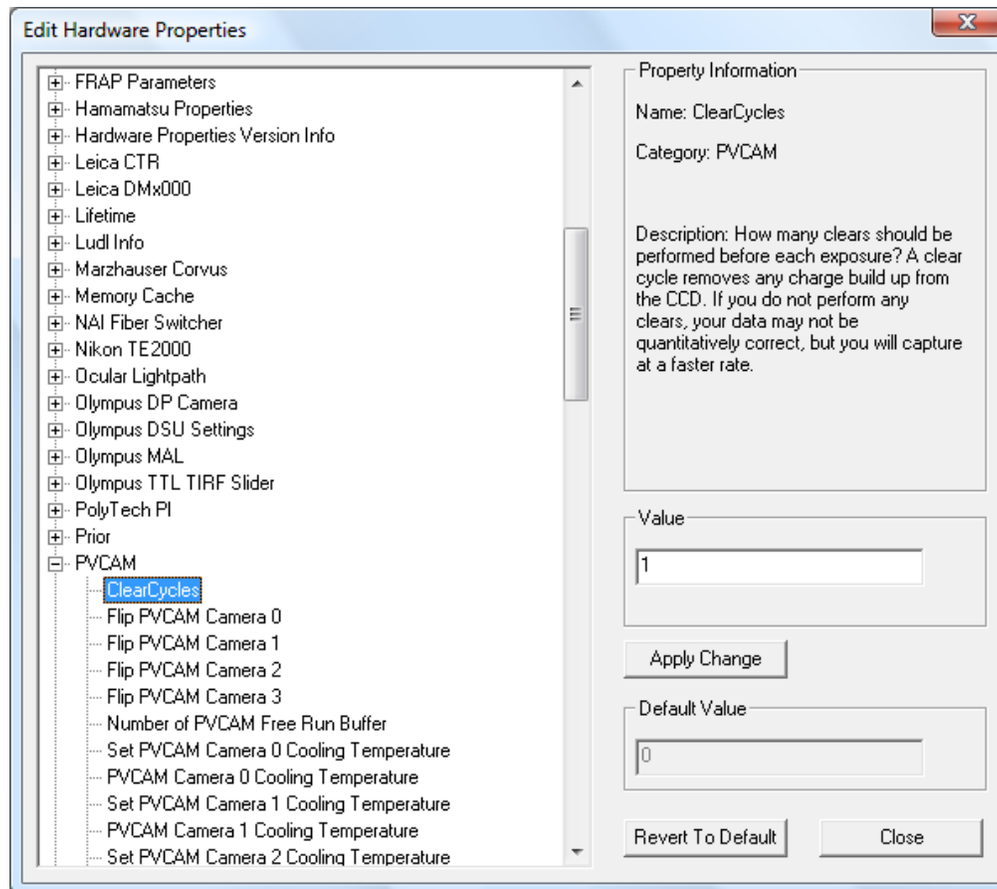
The following sections describe how to get the best speed performance when performing timelapse imaging. In addition, to get best speed performance, you may wish to change the frequency of image and histogram updates during capture (see page 114).

#### 7.8.1 Photometrics Cameras

If you feel that your camera is not running at appropriate speeds, go to the **Camera** tab in the **Focus Window** and make sure that the drop-down **Speed** menu is set to 0. Also, you may choose to set the clear cycle on the camera so that charge is not cleared from the chip before each image is captured. Charge accumulates between image captures, and the default action of SlideBook is to clear the accumulated charge before capturing an image. If you choose not to perform the clear action, the camera will run at a faster rate. However, the accuracy of the intensities measured will be reduced.

To change the clear cycle setting, do the following:

1. Select **Edit > Hardware Properties**.
2. Click on the + sign to expand the **PVCAM** section.
3. Click on **Clear Cycles** and the right hand side of the dialog box will update.



4. Enter “0” in the edit field and then **Apply Change**.
5. Click **Close** and then restart SlideBook to register the change.
6. Before performing an experiment, make sure to capture a test image immediately before beginning the experiment. This will help reduce the effect of charge build up in the first frame of the image capture.

### 7.8.2 Sutter DG-4

You may increase the switch time on your Sutter DG-4 by employing TTL control. You must purchase the SlideBook TTL Synchronization Module to enable TTL control (please contact Olympus America ([slidebooksupport@olympus.com](mailto:slidebooksupport@olympus.com) or 1-800-446-5967 option 4) if you wish to purchase this module). Note that in order to use TTL control, the exposure times of the various channels to be captured must be equal. Please see the TTL module documentation for details on configuring a system for TTL control. The TTL module for SlideBook includes software plus a National Instruments multifunction DAQ board and breakout box.

### 7.8.3 Z Stages

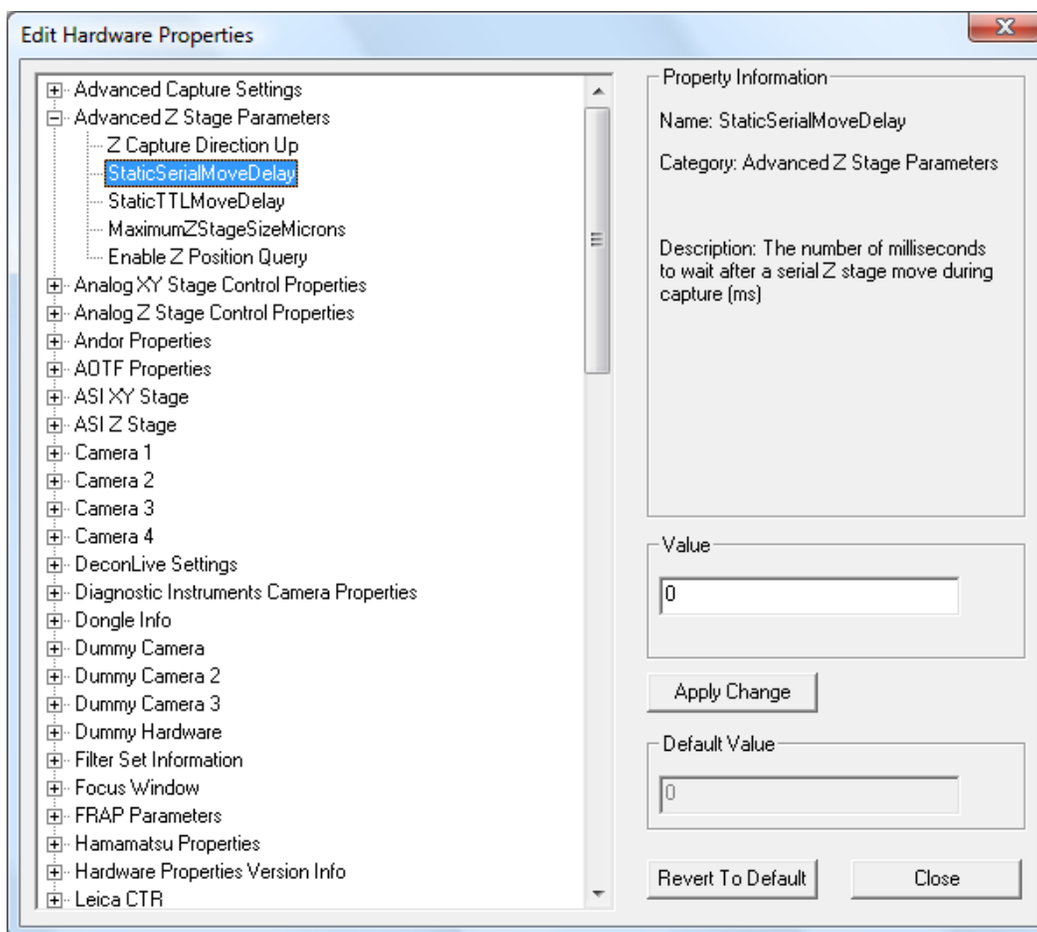
There are delays that may be reduced in order to increase the speed performance of the ASI/Mad City piezoelectric stage and the Physik Instrumente PIFOC piezoelectric focusing collar. The following adjustments allow the user to fine-tune the amount of time the

software waits for a Z movement to complete. The appropriate delay depends on system configuration and experimental conditions. The following parameters must be set to give you the minimum possible delay to achieve maximum system performance. Real-world conditions may make it necessary to increase the delays to achieve scientifically acceptable results. Should you experience issues with image quality we recommend you reset these parameters to the default value.

### 7.8.3.1 ASI/Mad City Piezoelectric Stage

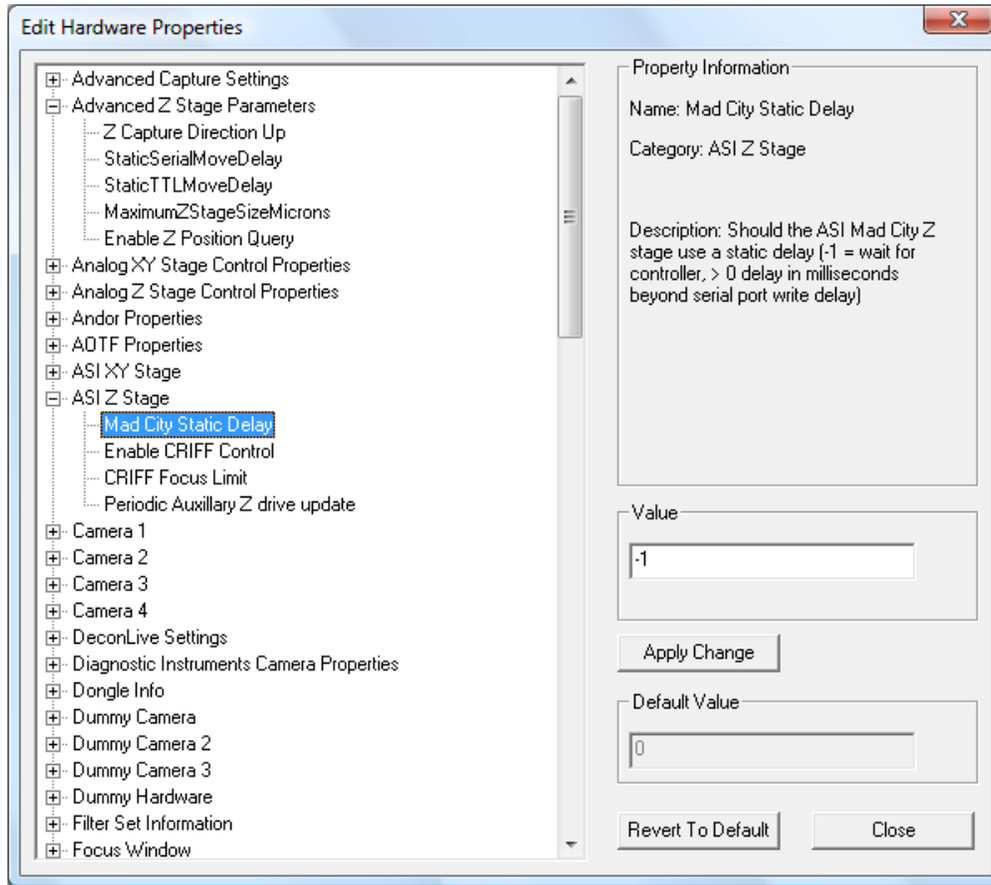
To increase the speed of this device, perform the following steps:

1. Select **Edit > Hardware Properties**.
2. Click on the + sign to expand the section next to **Advanced Z Stage Parameters**.



3. Click on **StaticSerialMoveDelay** and enter "0" in the edit field on the right side of the dialog box.
4. Click on the + next to **ASI Z Stage** to expand the section.

- Click on **Mad City Static Delay** and the right hand side of the dialog box will update.



- Enter the number “1” in the edit field and then select **Apply Change**.
- Select **Close** and then restart SlideBook to register the change.

You may find that you need to increase the delay above 0ms in order to achieve scientifically acceptable results.

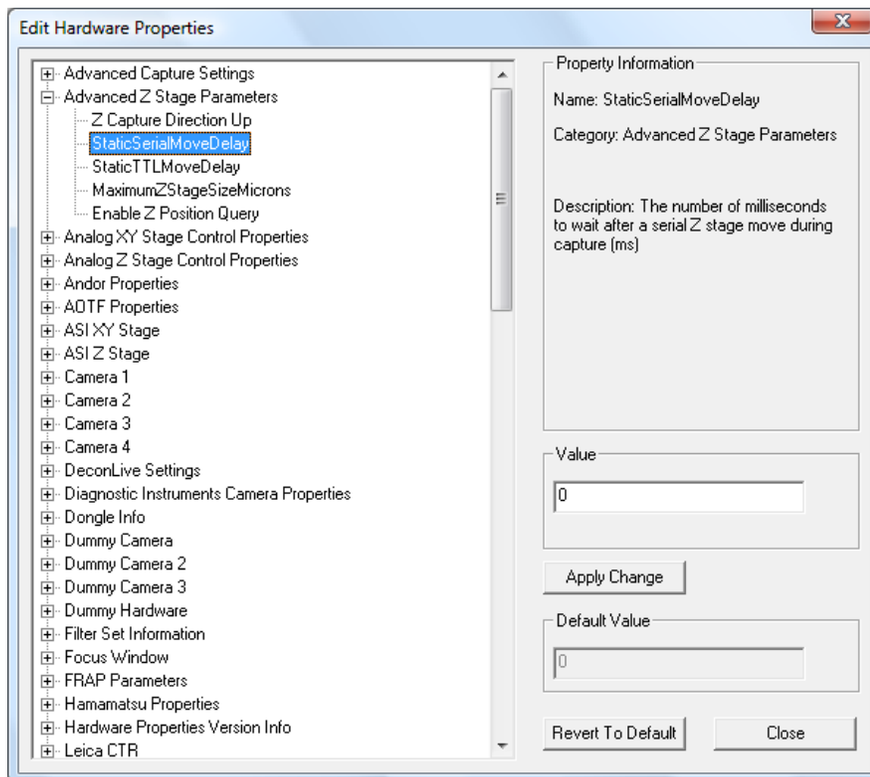
### 7.8.3.2 Physik Instrumente PIFOC Piezoelectric Focusing Collar

The PIFOC collar can be operated in TTL mode (see TTL module documentation) or in serial mode. In order to operate in TTL mode, you must purchase the TTL module for SlideBook.

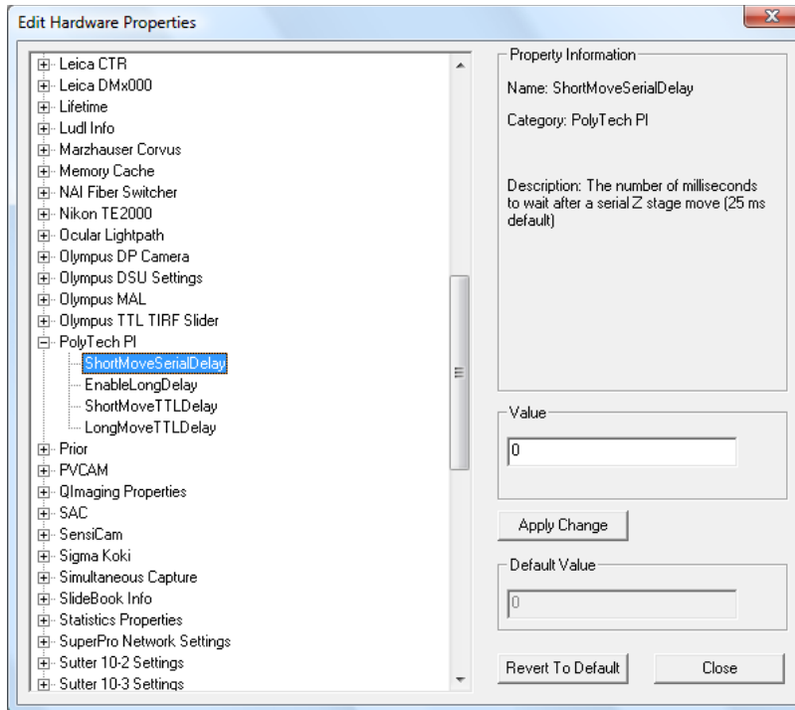
You should perform the following procedure to increase speed in serial mode whether you are configured for TLL or serial mode. There may be instances where the TTL mode will not be activated, so you will need to increase the performance in serial mode. To increase the speed of this device in serial mode, perform the following steps:

- Select **Edit > Hardware Properties**.

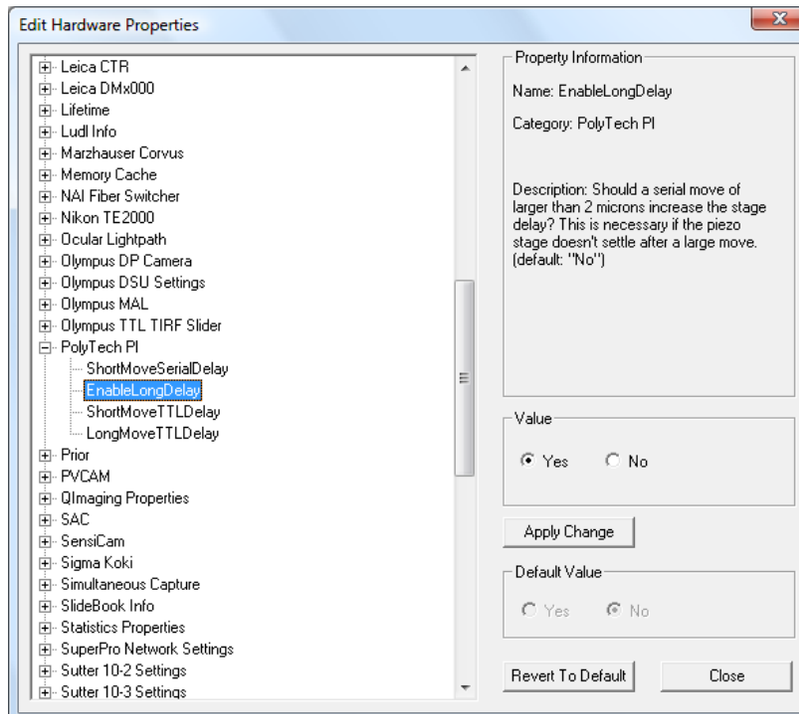
2. Click on the + sign to expand the section next to **Advanced Z Stage Parameters**.
3. Click on **StaticSerialMoveDelay** and enter “0” in the edit field on the right side of the dialog box.



4. Click on the + next to **Polytech PI** to expand the section.
5. Click on **ShortMoveSerialDelay**
6. Enter “0” in the edit field and then select **Apply Change**.



7. Click on **EnableLongDelay**, the right side of the dialog will update.



8. Select **Yes** and then **Apply Change**.
9. Select **Close** and then restart SlideBook to register the change.

You may find that you need to increase the delay above 0ms in order to achieve scientifically acceptable results

### 7.9 Autofocus

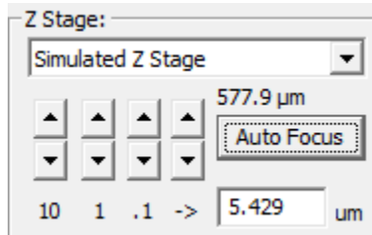
SlideBook offers the ability to perform an automatic focus routine during a 2D timelapse, a 2D multipoint timelapse or a montage capture. As with the other types of capture, you will set parameters in the focus window and in the capture window. The basic procedure for adding an auto focus routine to a multiplane capture is given below:

#### 7.9.1 Determining Auto Focus Parameters

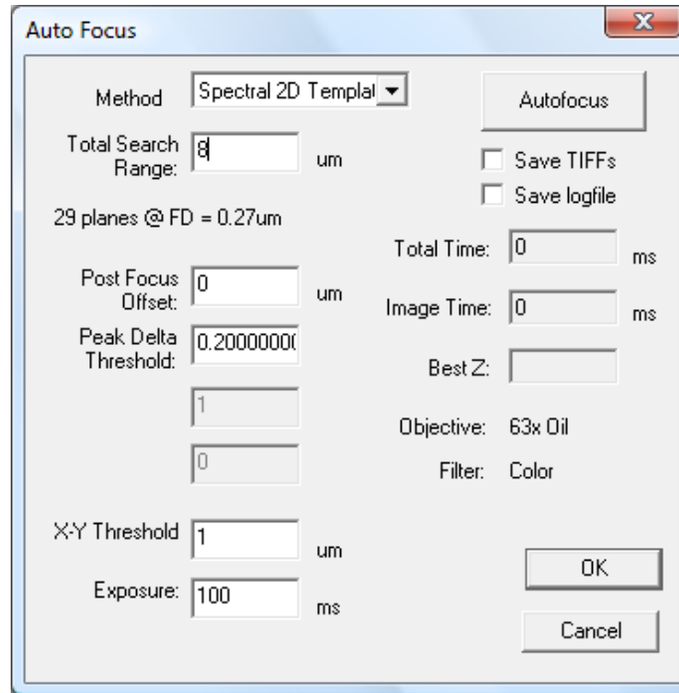
1. Open the Focus Window by either selecting **Window > Focus Window** or clicking on the focus window button in the SlideBook toolbar .
2. Select your objective and filter configuration then bring your sample into view as discussed in [Chapter 5 - Controlling the Camera and Microscope Hardware \(Focus Window\)](#) on page 77.

**NOTE:** It is critical that the **Objective** information is set correctly under **Edit > Optical Parameters > Objectives**. SlideBook uses its knowledge of the objective to compute the *z* spacing that it will use during auto focus. If your microscope has an automated objective nosepiece, SlideBook can read its position; however, if your microscope has a manual objective turret you must make sure the proper **Objective** is selected. To view objectives in the focus window on a manual microscope, go to **Edit > Hardware Configuration** and select **Manual Objective Turret** from the **Objective Turret** dropdown menu.

3. Select the **Auto Focus** button from the Focus Window.



4. The **Auto Focus** dialog box is used to determine the optimal auto focus parameters for your particular sample.



5. Enter parameters into the edit boxes according to the following descriptions:

- **Method** – The computational method used to evaluate and compare focal planes. Changing the method can affect the success of the auto focus routine. The default method of **Spectral 2D Template** will work best in almost all cases. The other methods are experimental. You should consult 3i if you are not having success with the default method.
- **Total Search Range** – The total distance in the  $z$  dimension that will be searched during the auto focus routine starting from half the specified distance below the current  $z$  position. SlideBook will automatically compute the number of planes to image based on the total search range and the calculated depth of field of the microscope objective. This information is displayed below the **Total Search Range** edit box. For instance, in the above example in order to span the 8 micron search distance with an objective that has a focal distance (FD) of 0.27 microns, a total of 29 planes will be imaged.
- **Post Focus Offset** – A positive or negative offset distance that the  $z$  stage will be moved from the optimal focus position. This feature is useful for always focusing a constant distance above or below the automatically determined focal plane.
- **Peak Delta Threshold** – Used to determine when no optimal plane of focus should be computed from a particular image position. When performing a montage capture, for example, some fields might not contain tissue and should not be used to change the current  $z$  position. This value can range between 0.0 and 1.0, but should not be decreased below 0.1 in

most cases. For captures where you know that each image in the sequence will have sufficient detail to focus, this threshold can be raised. If the value is set to 1.0 then the threshold will be ignored.

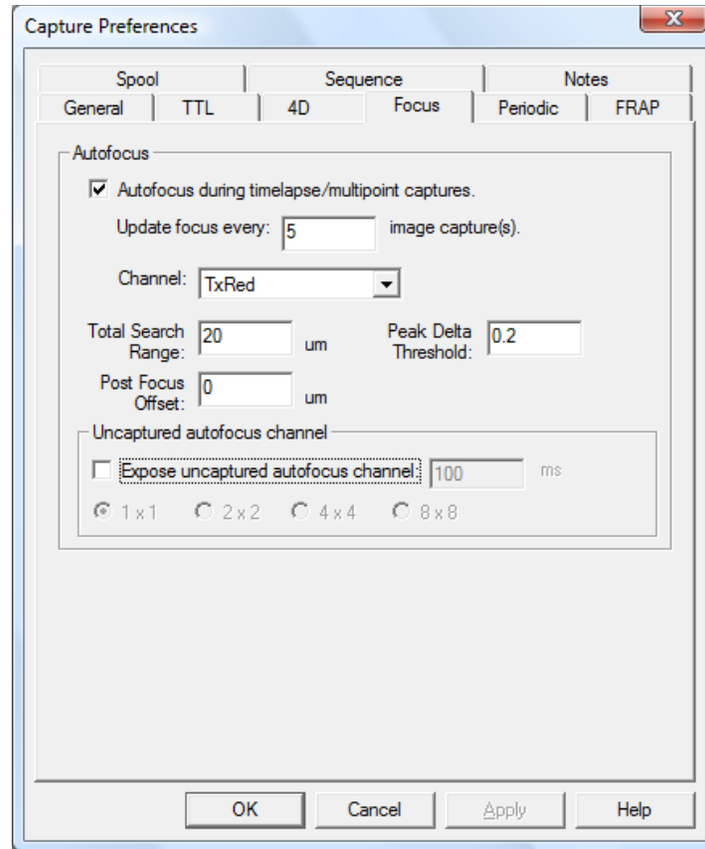
- **X-Y Threshold** – Not currently used.
  - **Exposure** – The exposure time used for image capture during the auto focus routine. This edit box will have the same value as the exposure time used in the focus window.
  - **Save TIFFs** – An advanced debugging feature that will generate a series of TIFF files in a numbered directory created in SlideBook program directory.
  - **Save Logfile** – An advanced debugging feature that creates an autofocus log file in the SlideBook program directory.
6. Press the **Autofocus** button to initiate a single auto focus scan. When finished, SlideBook will report the optimal focal  $z$  position in the **Best Z** information box and automatically move the  $z$  stage to this location. In addition, timing statistics for the autofocus routine are given in the **Total Time** and **Image Time** information boxes.

**NOTE:** If SlideBook is unable to determine the optimal focal plane in the sample it will return one of two messages in the **Best Z** information box. The **No Gradient** message indicates that SlideBook was unable to find a focal gradient within the specified search range. In this case, the peak delta threshold used for autofocus computation may need to be increased or the search range may need to be increased. The **At Edge** message indicates that the optimal focal plane appears to lie above or below the specified search range. Since SlideBook will automatically move the  $z$  stage to the best focal plane within the specified search range, simply running the auto focus routine again (starting from this new  $z$  position) will often find the optimal focal plane.

7. Press the **OK** button in the Auto Focus dialog box.
8. Use the focus window to evaluate the success of the autofocus routine. Using the zoom button **+** to enlarge the image often aids in determining the best plane of focus (see page 78). Once an acceptable set of auto focus parameters has been determined, an auto focus routine can be added to the capture sequence.

## 7.9.2 Adding Auto Focus to an Image Capture Sequence

1. To include an auto focus routine in an image capture sequence go to **Capture Preferences** dialog box (via the **Advanced** button in the Capture dialog box) and select the **Focus** tab. The following window will appear:

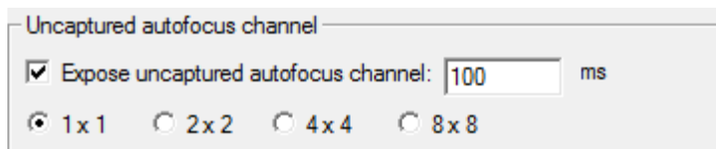


2. Select the **Auto focus during timelapse/multipoint captures** checkbox. This will enable specification of several auto focus parameters.
3. Enter the autofocus frequency for the image capture series in the **Update focus every \_\_ image capture(s)** edit box. The autofocus routine will always be initiated before the first image of a capture series. Specifying the autofocus frequency will determine at which subsequent timepoints the autofocus will occur. For instance, setting a frequency of 2 would cause an auto focus to occur every other timepoint.

**NOTE:** SlideBook keeps track of the optimal auto focus plane during the image capture series. Thus, after an auto focus routine has determined an optimal focal plane, all subsequent images will be captured at this  $z$  position until another auto focus routine is initiated. The  $z$  position of this new auto focus routine will be determined by the results of the previous auto focus.

4. Select the channel that will be used for auto focus. This should be the same channel that was used to test the auto focus parameters in the focus window.
5. The **Total Search Range**, **Post Focus Offset**, and **Peak Delta Threshold** values will be automatically entered based on the auto focus parameters determined in the focus window. If needed, these values can be further modified.

- If the autofocus channel will be a captured channel during the image series, click the **OK** button and proceed with setting up the timelapse image capture parameters as described on page 112. Make sure to expose the autofocus channel in the image capture window and set an appropriate exposure time. SlideBook will use these capture parameters (image extent, bin factor and exposure time) when acquiring the image stack for auto focus computation.
- If you do not want to capture the autofocus channel, or you want to use a different image extent, bin factor or exposure time for auto focusing, select the **Expose uncaptured auto focus channel** checkbox.



Enter the exposure time and bin factor for the uncaptured auto focus channel. These parameters will be used to determine image capture for the purposes of autofocusing. Click the **OK** button and proceed with setting up the image capture. Now the channel that is being used for autofocusing does not need to be exposed during the capture series.

- Selecting **Expose uncaptured auto focus channel** also provides the ability to specify a region of interest that will be used for auto focusing. This is useful for auto focusing on samples that have features with heterogeneous optimal focal planes. In this case, a full chip image extent can be captured at each timepoint while using only a small subregion to determine optimal focus. Specifying an auto focus region of interest is done in the Capture dialog box:



- When the **Expose uncaptured AF channel** checkbox is selected in the Capture Preferences dialog box, the **Autofocus** radio button is active in the **Extent, Offset, and Binning (pixels)** section of the Capture dialog box. This menu is used to alternate between designating the image extent for captured channels and the image extent for auto focus. To define a subregion for autofocus, simply select the **Autofocus** radio button and specify a **Width, Height, X Offset, and Y Offset**. Alternatively, draw a selection in the Live View and press the **Update** button. For more information on defining image subregions refer to the [Selecting the Area to be Imaged](#) section of Chapter 6 on page 92.

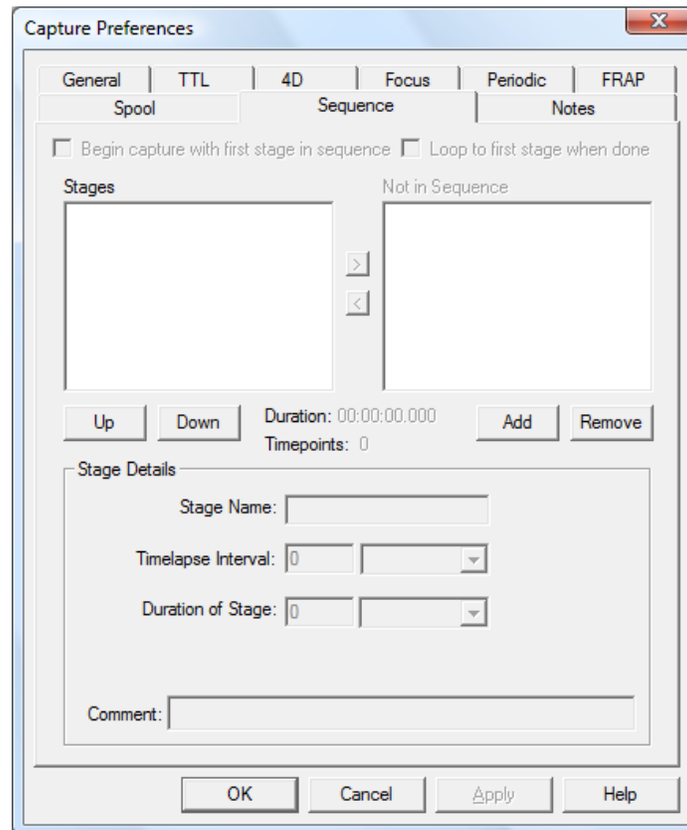
## 7.10 Varying Capture Rates During Timelapse Capture (Sequences)

You may choose to capture at various capture rates throughout your experiment when performing a multi-channel 2D or 3D timelapse capture. The capture rates may be changed either automatically in a preset protocol or manually during capture.

### 7.10.1 Setting Capture Preferences

To use this feature, you must first set up capture preferences. To do so:

1. Open the **Capture** dialog (**Image > Capture**) and press the **Advanced** button.
2. In the Capture Preferences dialog, select the **Sequence** tab.

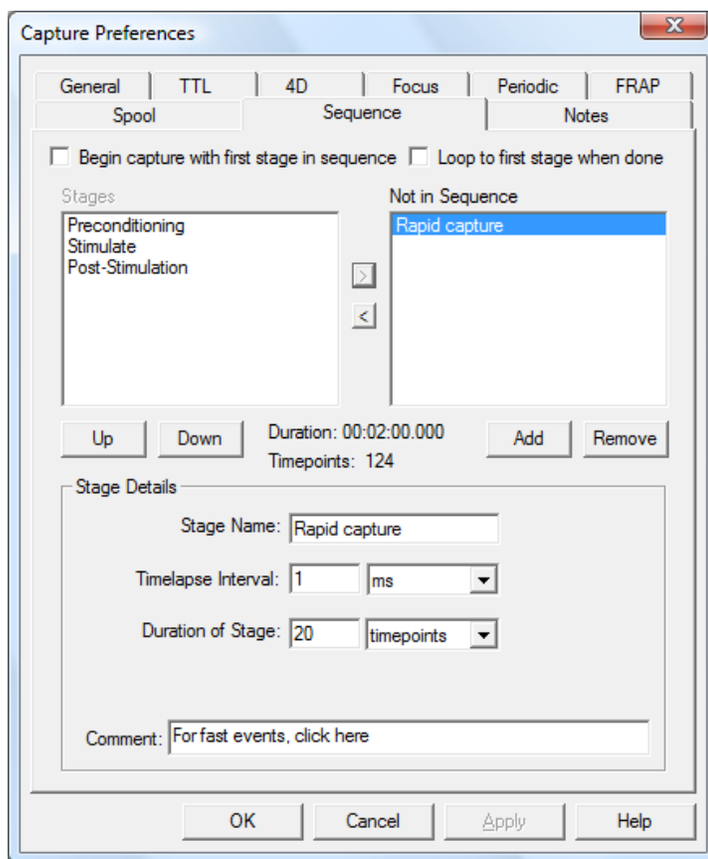


The elements of the dialog box are described below. A sequence is made up of a series of stages. Each stage may have a specified duration and timelapse interval (time between timepoints).

- **Begin capture with first event in sequence** – if checked, capture will start with the first event in the sequence
- **Loop to first stage when done** – if checked, capture will return to first stage
- **Stages** – each stage will be automatically performed in the order they are displayed, from top to bottom

- **Not in Sequence** – these stages must be manually selected in the Capture Status dialog; they will not be performed automatically
- **Stage Name** – defined by the user
- **Timelapse Interval** - the delay between the beginning of one timepoint and the beginning of the next timepoint. Available units are milliseconds, seconds, minutes, and hours.
- **Duration of Stage** - how long capture will occur at the defined rate. You may define the length as timepoints (e.g., 5 timepoints), a time period (e.g., for 10 min), or **continuous** (ending when manually selecting the next stage)

An example of a completed dialog box is shown below.



3. To add a stage, select **Add**. Enter the stage information described above. The stage will appear in the **Stages** list.
4. To add a stage that will be started manually in the Capture Controls, click on the **Not in Sequence** list to activate it.

5. Add a stage as described in Step 3. The stage will appear in the **Not In Sequence** frame. You may use the > or < buttons to move events from the **Stages** frame to the **Not In Sequence** frame and vice versa.

To change the sequence of stages, use the **Up** and **Down** buttons to move the stage in the list up or down one place.

6. To remove a stage, select the stage from the list by clicking on the stage name and then clicking **Remove**.

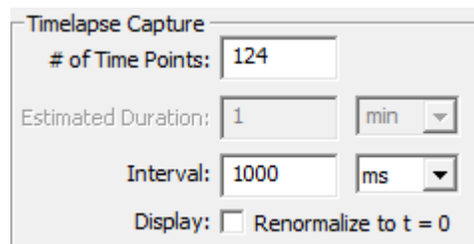
**NOTE:** The overall time (duration) and the number of timepoints of all sequential stages is shown.

7. After you have set up your sequence, press **OK**. If you have selected to perform an automated sequence, the appropriate parameters will be loaded into the Capture dialog box.

### 7.10.2 Setting Capture Dialog Box Parameters

1. Open the Capture dialog box by selecting **Image > Capture** or by selecting the Capture icon in the tool bar .
2. If it is not already checked, check the **Timelapse** checkbox in the **Capture Type** section of the Capture dialog box. You now have the option to define a variety of timelapse parameters. Enter the desired values in the following fields:
  - **# of Timepoints** – The number of timepoints that will be captured.
  - **Duration** – The total length of time for the experiment. Units of time, in milliseconds (ms), seconds (s), minutes (m) or hours (h), can be selected from the dropdown menu.
  - **Interval** – The delay between the beginning of one timepoint and the beginning of the next timepoint. The interval unit can be selected from the dropdown menu. If the capture sequence at one timepoint takes longer than the interval, SlideBook will capture the next timepoint immediately following the preceding timepoint.

**NOTE:** As you type in two of these values, the third field will be calculated automatically.



Timelapse Capture

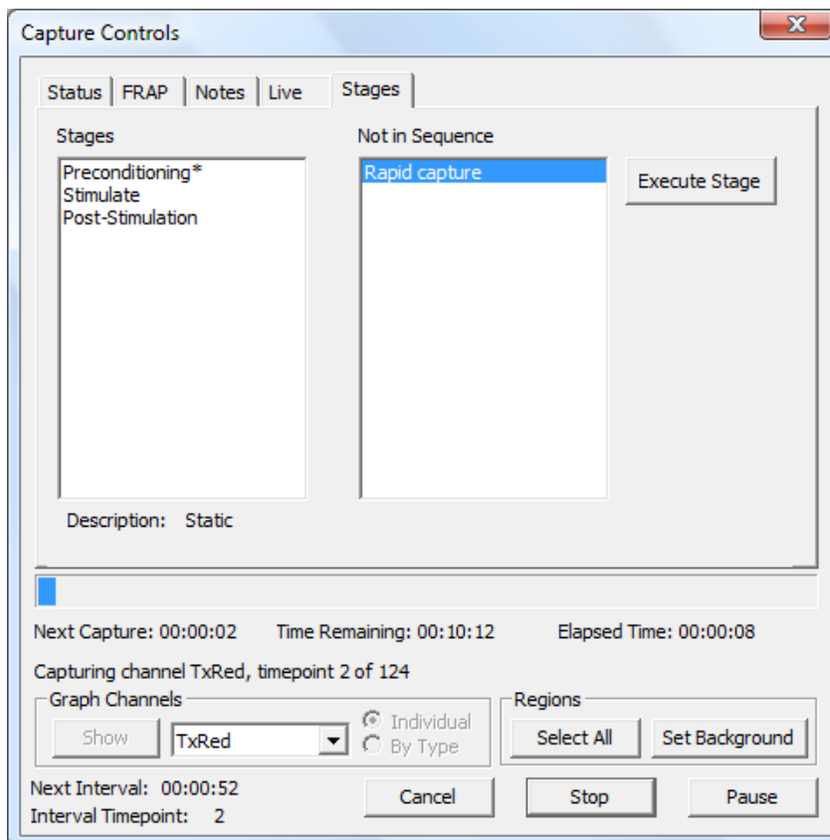
# of Time Points: 124

Estimated Duration: 1 min

Interval: 1000 ms

Display:  Renormalize to t = 0

3. To begin a capture click **OK**. The Capture Status Window will appear.
4. Click on the **Stages** tab.



This tab has two lists, **Stages** and **Not in Sequence**.

5. If you have opted to start a sequence at the beginning of capture, the first stage in the top frame will be marked with a “\*”. To manually execute a stage, select the stage by clicking on it in the list and then click on the **Execute Stage** button. The number of timepoints in the stage and the timelapse interval will be displayed at the bottom of the window.

### 7.10.3 Examples for Variable Capture

#### 7.10.3.1 Example 1

Suppose you have set up a stage with 22 timepoints and a 200ms interval, and have checked the **Begin capture with first stage in sequence** checkbox in the Capture Preferences dialog. In the Capture window, you select a timelapse capture with 100 timepoints, and a 1000ms interval. When you click **Start**, capture will start by executing the sequential events in the displayed order in the **Stages** list. After the total sequence has been captured (22 timepoints) the remaining 78 timepoints ( $100 - 22 = 78$ ) will be captured

with the interval set in the **Timelapse Capture** section of the Capture dialog box (1000ms).

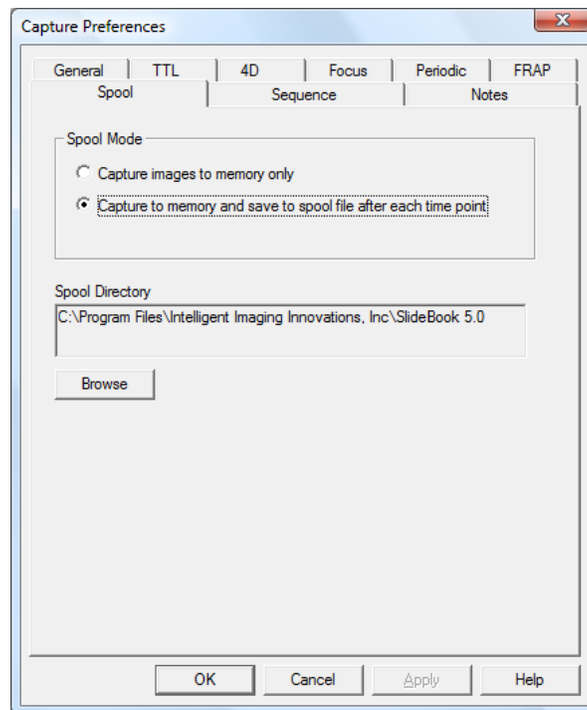
### 7.10.3.2 Example 2

Suppose you have set up capture as in Example 1, but did not check the **Begin capture with first stage in sequence** checkbox. Capture will begin with 1000ms interval between timepoints. To execute the stage, click on the stage to be executed and click **Execute Stage**. Twenty-two timepoints will be captured. The remaining timepoints will be capture at 1000ms interval until the total of 100 timepoints have been performed.

## 7.11 Saving Images to Disk (Spooled Capture)

SlideBook allows you to save your images directly to disk. This is especially important when performing long timelapse captures that will exceed your computer's memory limit. To save images to disk:

1. From the Capture dialog box, select **Advanced** to bring up the Capture Preferences dialog box.
2. Select the **Spool** tab. The following window will be displayed:



2. Select the desired spool capture mode:
  - **Capture images to memory only** – Images are saved in memory and only become permanent after capture if **Save Slide** is selected.
  - **Capture to memory and save to spool file after each timepoint** – Images are both kept in memory and saved to disk. This feature allows

you to permanently save images as they are captured, and immediately view the images post-capture (no need to import a spool file). This is only useful for captures that are not larger than the memory available.

3. Select the location for storage of spool files by selecting **Browse** and then navigating to the desired location.
4. Click **OK** to save these preferences, and then set up for your desired capture sequence. Once your capture is finished, you may import it for viewing as discussed in the section [Importing SlideBook Spool Files](#) on page 97.

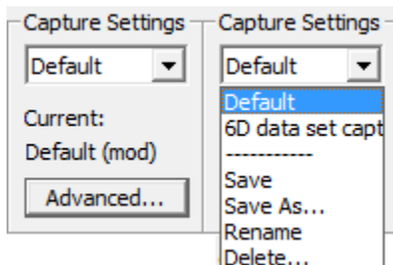
### 7.12 Saving Capture Parameters

You can store and recall frequently used capture parameters. The following parameters can be saved:

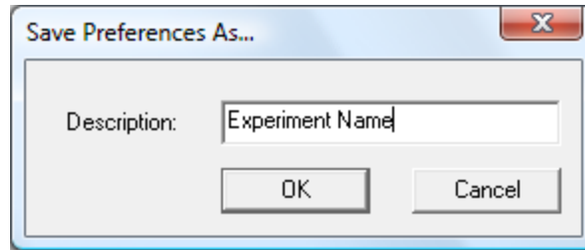
- Image extent
- Binning
- Capture type
- Channels and exposure times
- Camera intensification and gain settings (if applicable)
- Capture preferences (e.g., Periodic, 4D, Spool features)

To do so:

1. Begin a new capture by selecting **Image > Capture**. The Capture dialog will appear.
2. Select your capture parameters. To access advanced parameters such as periodic capture select the **Advanced** button. The Capture Preferences dialog will appear.
3. Configure the desired preferences and select **OK**.
4. To save these preferences, select **Save As** from the **Capture Settings** drop-down menu in the Capture dialog. If you choose **Save**, the default parameters will be overwritten.



The following dialog will appear:



5. Enter a name for these preferences and select **OK**.
6. If you would like to load a different set of parameters, select one from the dropdown list. If you modify your capture parameters, the name of the currently loaded parameters will say "(mod)" next to the name. You may choose to save them under the same name by selecting **Save** or under a different name by selecting **Save As** as discussed above.

## 8 Image Display and Manipulation (Views)

SlideBook stores images in *slides* and displays images using *views*. There are two kinds of views: *data views* and *display views*. Data views are interactive—the user can display any combination of different channels and masks as well as change lookup table parameters (renormalization) and edit the image’s masks. A new data view can be spawned either from the slide, or in one case, from another view. Display views, on the other hand, are always spawned from a data view and are set to the channel, mask, and renormalization selections of that view. Data views focus on interactive analysis, while display views focus on visualization.

This chapter covers the following topics:

- Introduction to Slide and Image Display in SlideBook
- Using Data Views to Display Images
- Manipulating Data Views Using the Info/Tool Bar
- Using the Tool Menu
- Using Display Views (Creating Renderings and Movies)
- Exporting Views

### 8.1 Introduction to Slide and Image Display in SlideBook

When you start SlideBook a new slide called **slide1** is created, and unless you open another slide, it is into this slide that all new and imported images are inserted. Typically a slide corresponds to all of the images taken from a particular sample.

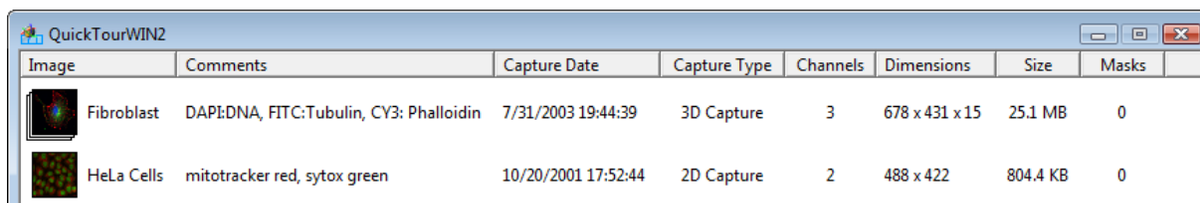


Image	Comments	Capture Date	Capture Type	Channels	Dimensions	Size	Masks
	Fibroblast DAPI:DNA, FITC:Tubulin, CY3: Phalloidin	7/31/2003 19:44:39	3D Capture	3	678 x 431 x 15	25.1 MB	0
	HeLa Cells mitotracker red, sytox green	10/20/2001 17:52:44	2D Capture	2	488 x 422	804.4 KB	0

Each image in a slide is represented in a slide window as a thumbnail and some associated data. The image may be either a single image or a set of images depending on the type of capture that was performed (e.g., 2D, 3D, or timelapse). The thumbnail is either a shrunken rendering of the first plane of an image (the default) or of some portion of an image (if you have specified a particular thumbnail using the thumbnail button—see [Changing the Default Display](#) on page 180). The information includes the following:

- **Image name** - automatically generated in the case of captured images or corresponds to a file name in the case of imported images.
- **Comments** – entered by the user either before or after capture
- **Capture date and time**
- **Capture type** – 2D, 3D, 2D timelapse, or 4D

- **Number of channels** – number of fluorophores
- **Dimensions** – image size in pixels and planes (can be X\*Y\*Z/T or X\*Y\*Z\*T)
- **Image size** – image size in MB
- **Masks** – number of masks stored with the image

### 8.1.1 Working with Slides

The Slide (.sld) file is SlideBook's native document type. Opening, closing, and saving slide files are performed in a manner similar to other software programs.

#### 8.1.1.1 Saving Slides

To save a newly created or updated slide:

1. Click on the slide that you wish to save to make it the active slide.
2. Select **File > Save Slide**. If you have not previously saved the slide, you will be prompted for a file name.
3. Enter a filename and select **Save**.

To save a copy of a slide:

1. Click on the slide that you wish to save to make it the active slide.
2. Select **File > Save Slide As**.
3. Navigate to the location where you would like to save the file, enter a filename, and select **Save**.

#### 8.1.1.2 Opening Slides

To open an existing slide:

1. Select **File > Open Slide**.
2. Navigate to the desired slide and select **Open**.

#### 8.1.1.3 Closing Slides

To close a slide:

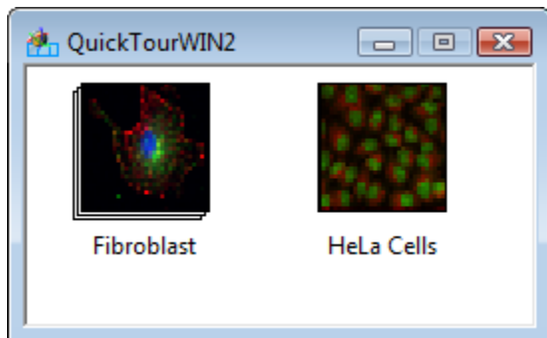
1. Click on the slide that you wish to close to make it the active slide.
2. Select **File > Close Slide** or click on the  in upper right corner.

#### 8.1.1.4 Deleting Slides

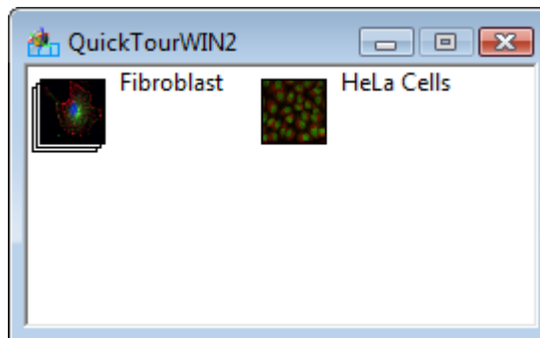
Slides can be deleted using standard Windows operations for deleting files. You may not delete a slide from within SlideBook; however, you may delete the images that are present in the slide.

### 8.1.1.5 Changing the Slide View Display

SlideBook offers several ways to view your Slide. To change your Slide View display, go to **View > Slide View Display** and select one of the following options:



**Large Icons**



**List**

The default display is the **Details** display (shown at the beginning of the chapter).

## 8.1.2 Working with Images

As discussed above, each slide may contain several images. Using the same clipboard metaphor as other applications, SlideBook lets you remove images from a slide, duplicate images, and insert images. You can cut or copy an image from one slide and insert it into another.

**NOTE:** There is currently no mechanism for multiple selection; you can only manipulate one image at a time.

### 8.1.2.1 Selecting an Image

Before performing any operations on a given image, it must first be selected, or made the active image. To make an image the active image, click on an image thumbnail in an active slide window (or anywhere in the box surrounding an image's information).

The icon (and attributes) of the selected image will turn blue. It is on the selected image that all data manipulation commands (such as deconvolution), mask creation commands, and image information commands will operate.

### 8.1.2.2 Removing an Image from One Slide and Placing it in Another

To move an image from one slide to another, do the following:

1. Click on the slide that contains the image you would like to move; this makes it the active slide.
2. Click on the image you would like to move; this makes it the selected image.
3. Select **Edit > Cut Image**, or click on the Cut Image icon . This will remove the image from the slide.

4. Click on the slide that will hold the pasted image.
5. Select **Edit > Paste Image**, or click on the paste image icon .

**NOTE:** Since the image is stored internally in a proprietary format, these operations do not use the Windows clipboard. In other words, if you quit SlideBook without pasting a cut image, you will not be able to get to that image again.

### 8.1.2.3 Copying an Image

You may duplicate an image and place it in another slide, or in the same slide. It is often useful to copy an image before performing manipulations that affect the underlying data.

1. Make the slide that contains the image that you would like to copy the active slide by clicking on it.
2. Make the image that you would like to move the selected image by clicking on it as discussed above.
3. Select **Edit > Copy Image**, or click on the copy image icon .
4. Click on the slide that will hold the copy of the image. This may be the current slide, or another slide.
5. Select **Edit > Paste Image**, or click on the paste image icon .

### 8.1.2.4 Deleting an Image

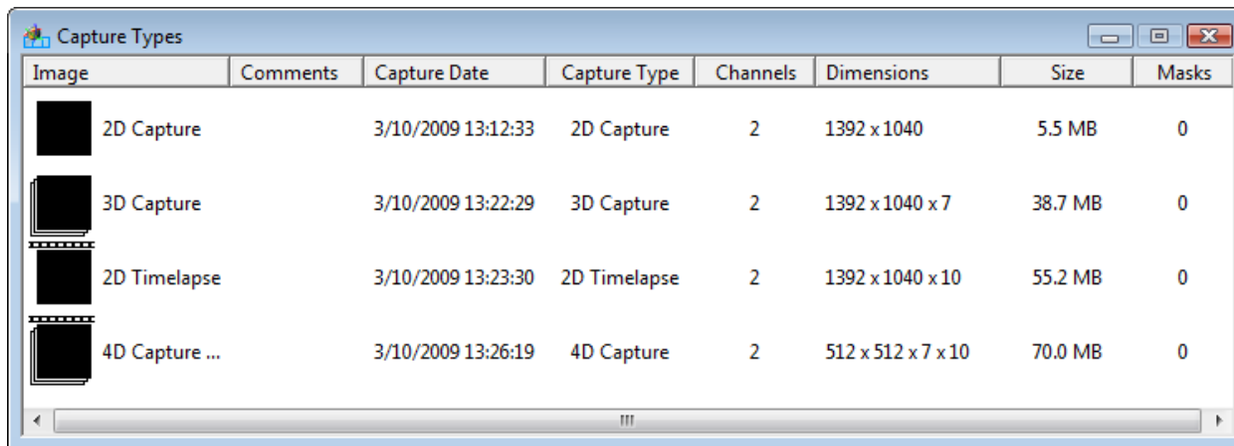
To permanently remove an image:

1. Make the slide that contains the image that you would like to delete the active slide by clicking on it.
2. Make the image that you would like to delete the selected image by clicking on it as discussed above.
3. Select **Edit > Delete Image**.

**CAUTION:** This is a permanent removal that cannot be undone.

## 8.1.3 Image Display

SlideBook contains four different kinds of data views: the *Main View*, the *Three View*, the *Tile View*, and the *Channel View*. Data views may be used not only to look at data, but to generate statistics. Display views are strictly used for viewing data. The following display views are available: *Volume View*, *Surface View*, and *Timelapse Intensity Plot*. The displays that are most appropriate for different modes of imaging or, capture types, are shown in the table below. Capture type is designated by the icon for any image as well as identified in the default .sld view as shown below.



		2D Capture	3D Capture	2D Timelapse	4D Capture
Data Views	Main View	✓	✓	✓	✓
	Three View		✓	✓	✓
	3D Tile View		✓		
	Timelapse Tile View			✓	✓
	Multidimensional Channel View	✓	✓	✓	✓
Display Views	3D Volume View		✓		✓
	4D Volume View				✓
	3D Surface View		✓		✓
	2D Timelapse Intensity Plot			✓	

Next, we will discuss how to generate data views and manipulate those views using the info/tool bar. Finally, we will discuss display views.

## 8.2 Using Data Views to Display Images

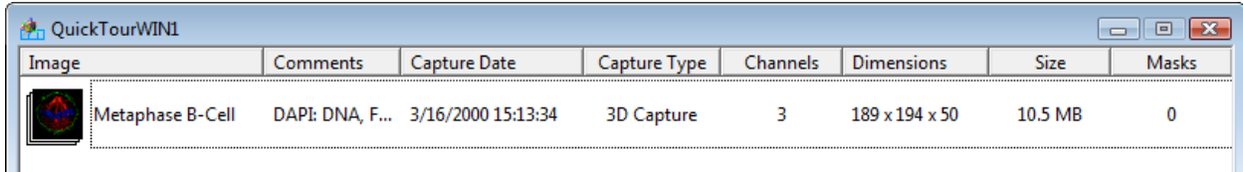
This section describes how to display data views. SlideBook allows multiple data views of a single image to be open simultaneously.

### 8.2.1 Displaying a Main View

The *Main View* can be used to display any type of data. The Main View displays a single plane at a time where the *z* or *t* axis can be scrolled using arrows or a slider.

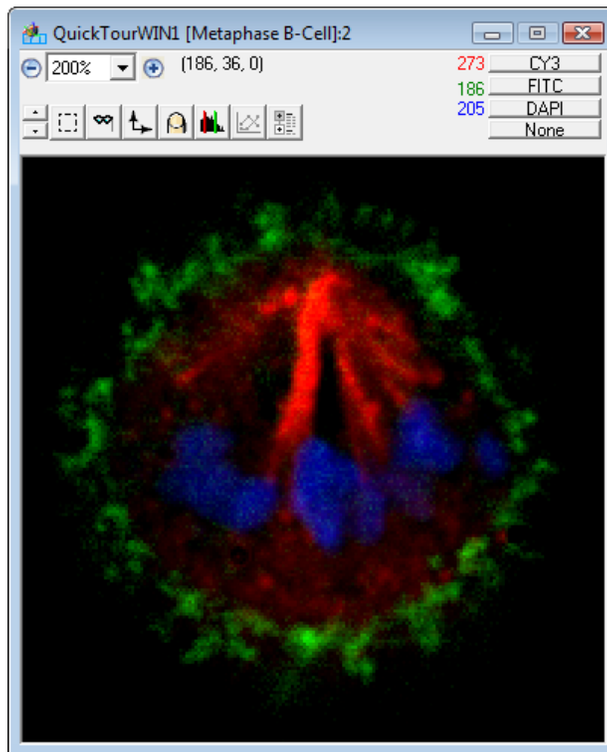
To generate a Main View:

1. Choose **File > Open Slide** and navigate to the desired slide file. Double click on the slide file to open it. A slide window will appear.



2. Double-click anywhere on the thumbnail or within the shaded region that surrounds the thumbnail.

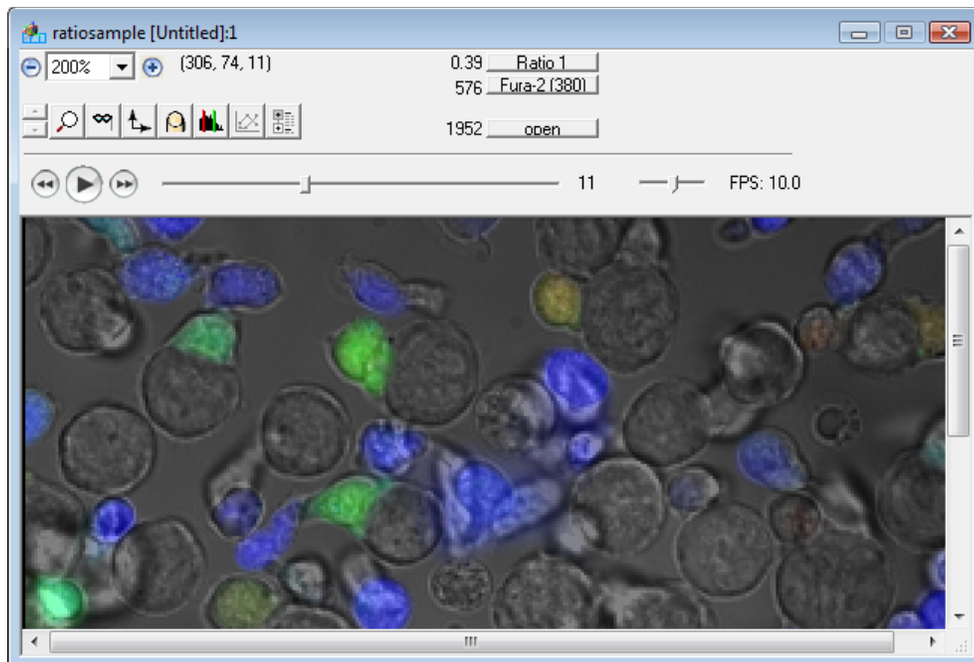
A new window will pop up that contains the image. This window is a Main View of the image.



Alternatively, you may open a Main View using the **View** menu. To do this:

1. Click once anywhere on the thumbnail or within the shaded region that surrounds the thumbnail to highlight the image.
2. Choose **View > New Main View** to display a new Main View.

When working with 2D Timelapse or 4D Capture types, the Main View window will have controls for time as shown below:



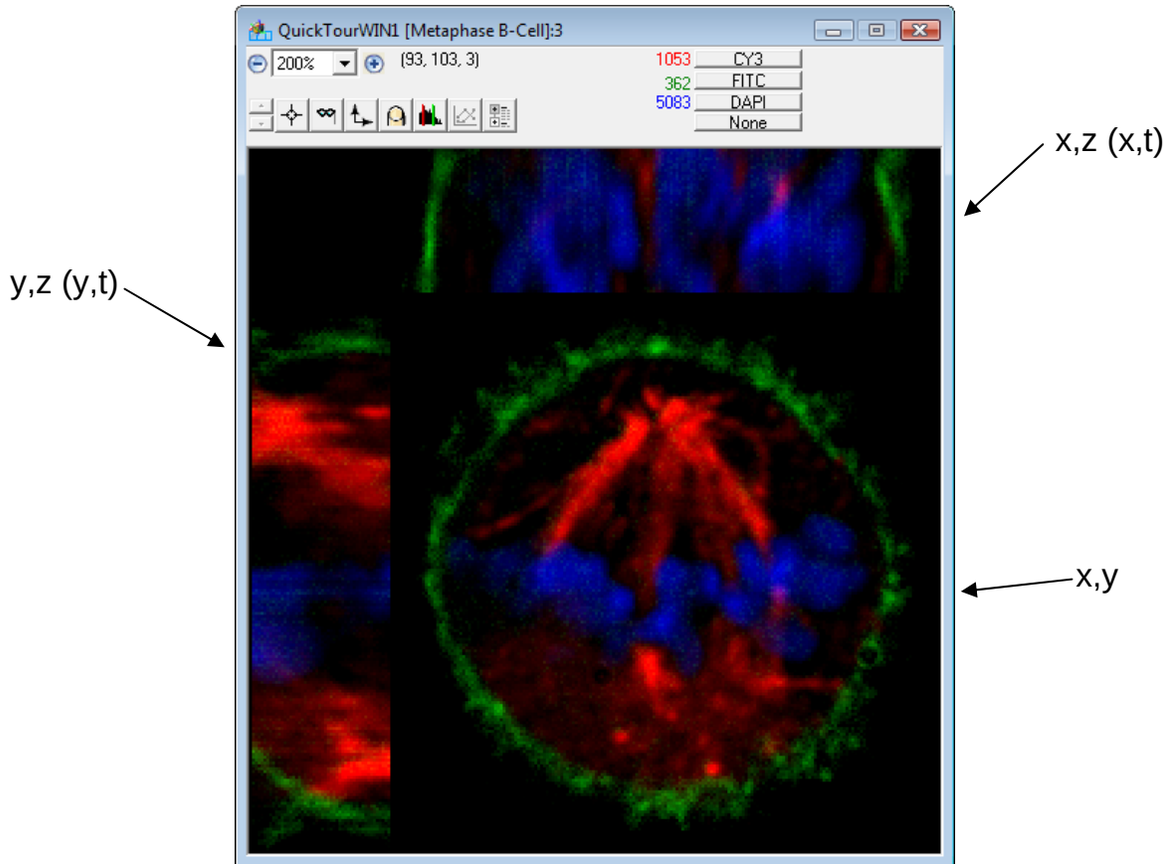
For 4D capture types, the Main View operates similarly to Main Views for 3D and 2D data. Additionally, there is a slider that you may click and drag to view individual timepoints of the 4D capture. You may scroll through the *z* axis by using the up and down arrow bars in the tool bar (see [Scrolling through the Invisible Axis in a 3D or Timelapse Image](#) on page 180).

### 8.2.2 Displaying a Three View

The *Three View* is equivalent to three Main Views put together, one for each axis. It is useful for exploring 3D, 4D, and 2D Timelapse data.

To open a Three View from an open slide:

1. Click once anywhere on the thumbnail or within the shaded region that surrounds the thumbnail to highlight the image.
2. Choose **View > Three View** to display a new Three View.



**NOTE:** You may notice that your object may appear disproportionately small, or “squashed” in the  $z$  direction. When performing 3D imaging, each plane that is captured is represented by a set of pixels in the image set. Each pixel is a perfect cube. You will notice that the  $z$  dimension in the 3D image set appears to be different than what was actually captured. This is because the  $z$  step size that was selected during capture is not actually the same as the length of the  $z$  dimension of the pixel. In the image set, planes are simply stacked on top of one another with no regard for the actual  $z$  distance traveled (this is done in order to preserve data fidelity). You may interpolate between planes and expand your  $z$  dimension to represent the actual  $z$  distance traveled by the microscope by generating an isotropic image (see page 213).

### 8.2.3 Spawning a Tile View

The *Tile View* can display many planes simultaneously. Both the 3D Tile View and the Timelapse Tile View are designed to enhance interaction with 3D and timelapse data.

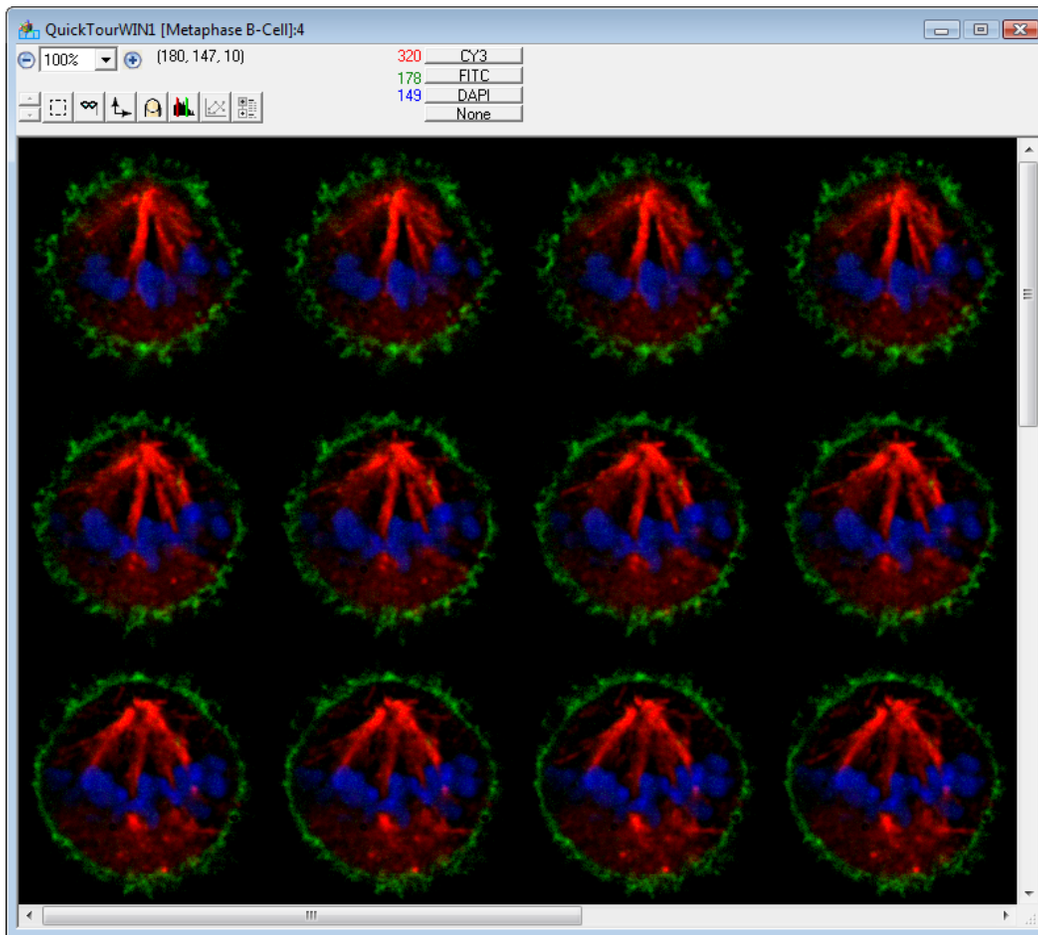
#### 8.2.3.1 Displaying a 3D Tile View

The 3D Tile View is designed for use with 3D Capture data. This view will display each  $z$ -plane in a montage.

From an open slide:

1. Click once anywhere on the thumbnail or within the shaded region that surrounds the thumbnail to highlight the image.

2. Choose **View > 3D Tile View** to display a new 3D Tile View.

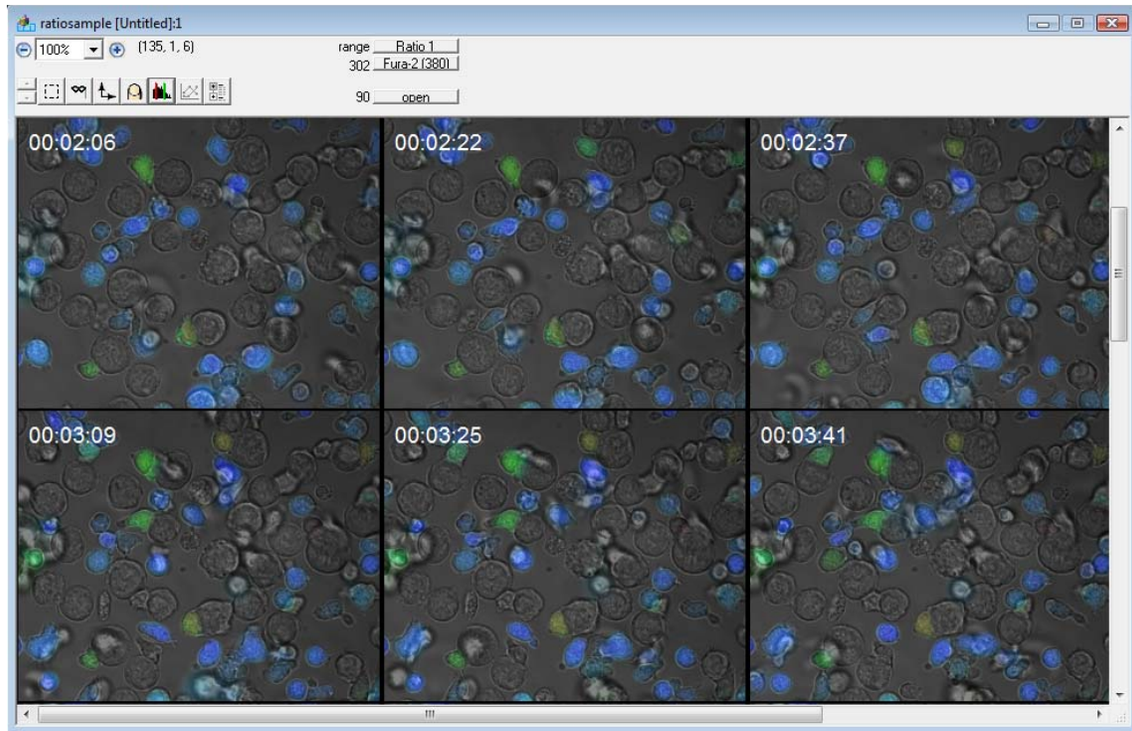


### 8.2.3.2 Displaying a Timelapse Tile

The 3D Tile View is designed for use with 2D Timelapse or 4D Capture data. This view will display each time-point in a montage. The arrows for z-movement will be available for 4D Capture data

From an open slide:

3. Click once anywhere on the thumbnail or within the shaded region that surrounds the thumbnail to highlight the image.
4. Choose **View > Timelapse Tile View** to display a new Timelapse Tile View.



### 8.2.3.3 Displaying a 3D or Timelapse Tile View of a Portion of the Image

The Tile View is slightly different than the Main and Three Views in that it is tailored to display a portion of the image rather than the entire image. Since the Tile View creates a pane for every plane that it displays, a Tile View of the entire image will often be far larger than the size of the monitor, which defeats the Tile View's purpose of allowing simultaneous viewing of multiple planes along the same axis. So, although a Tile View can be created directly from the slide window, you will more likely be making a 3D selection in either a Main View or a Three View (or even another Tile View) and spawning a Tile View from there.

In order to spawn a Tile View from another view:

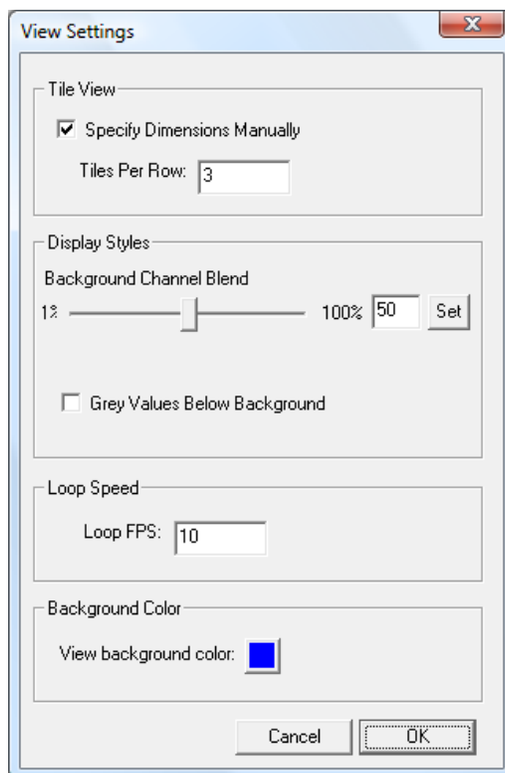
1. Make the desired view the active window by clicking on it.
2. Define the selection. There are several ways to do this, and they are discussed [Making a 2D or 3D \(x,y,z or x,y,t\) Selection](#) on page 186.
3. Choose the desired Tile View from the **View** menu to display a new 3D or Timelapse Tile View of the selected portion of the image.
4. To change the portion of the image that is displayed, select the hand tool from the tool menu and click and drag in one of the panes (see table on page 184 for a description of this function).

You can create a Tile View that shows slices along any of the three axes simply by making your selection in a view (or pane) that is currently set to the desired invisible axis (see description of the axis menu in the table below). In other words, if you make a selection in a Main View that has an invisible y-axis, and the selection spans ten units in the y-axis, the resulting Tile View will have ten panes that display the x axis horizontally and the z axis vertically.

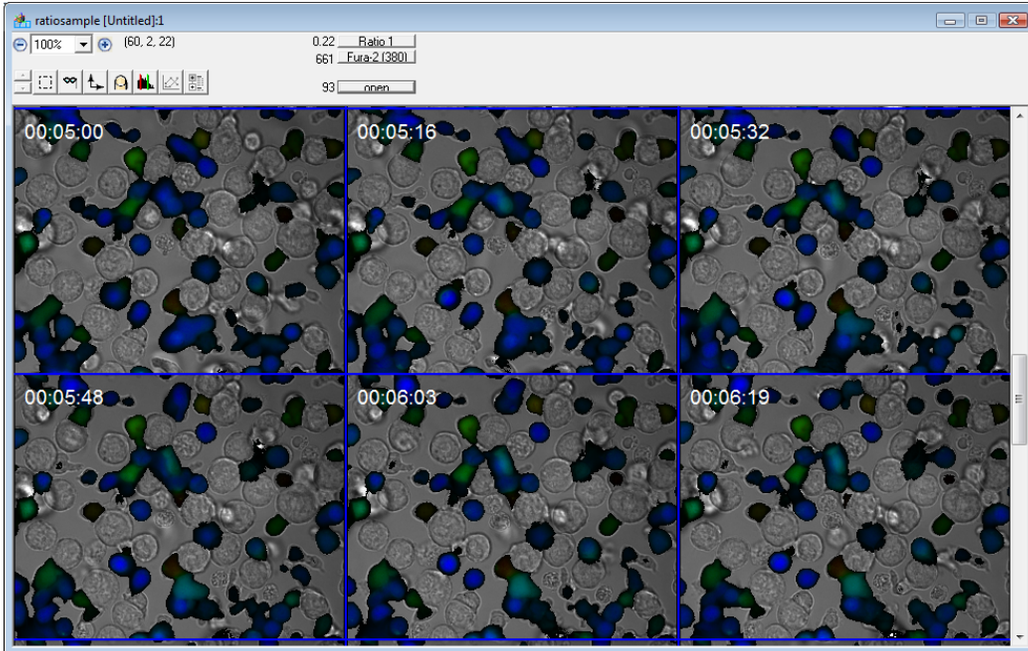
### 8.2.3.4 Arranging your Tile View

You may wish to change the way your **Tile View** is displayed. For instance, the default tile display may choose a 4 × 6 arrangement, while you desire a 6 × 4 arrangement. You may also wish to change the background color between the tiles. To alter these settings:

1. With a Tile View displayed, select **View > Settings** or click on the  Setting icon in the View window. The following dialog box will appear:



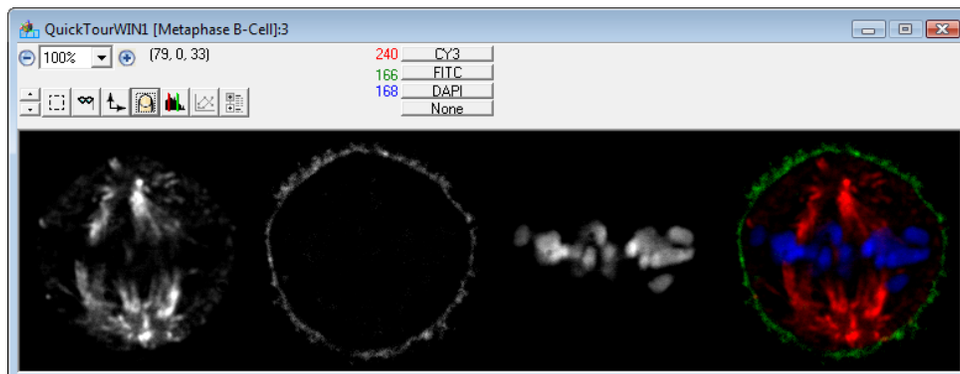
2. Select the **Specify Dimensions Manually** checkbox, then enter the **Tiles Per Row** that you desire in the edit field.
3. In the Background Color section, click on the colored square next to **View background color**.
4. Select the desired background color from the **Color** dialog and click **OK**.
5. Click **OK** to update your **Tile View**.



### 8.2.4 Displaying a Multidimensional Channel View

The *Multidimensional Channel View* is most useful for displaying multi-channel images. The Multidimensional Channel View displays each channel independently plus a composite RGB image. Users can also choose to display individual channels with their default coloring or to display multiple timepoints or z-sections of the image. To display a Multidimensional Channel View from an open slide:

1. Click once anywhere on the thumbnail or within the shaded region that surrounds the thumbnail to highlight the image.
2. Choose View > Multidimensional Channel View to display a new Multidimensional Channel View.

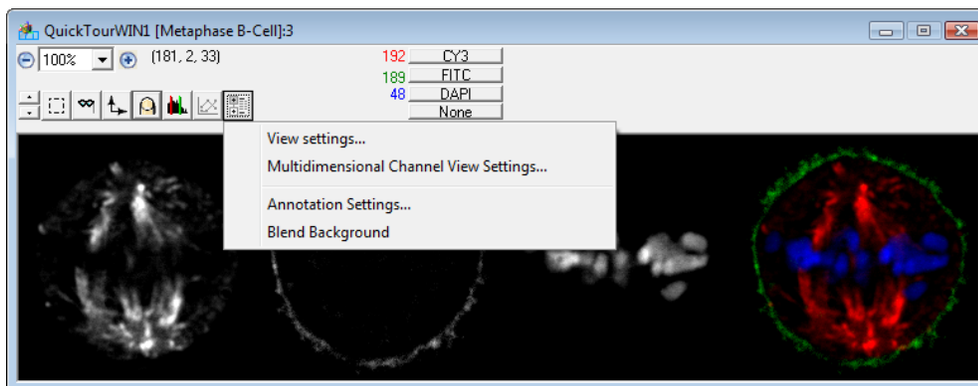


#### 8.2.4.1 Multidimensional Channel View Settings

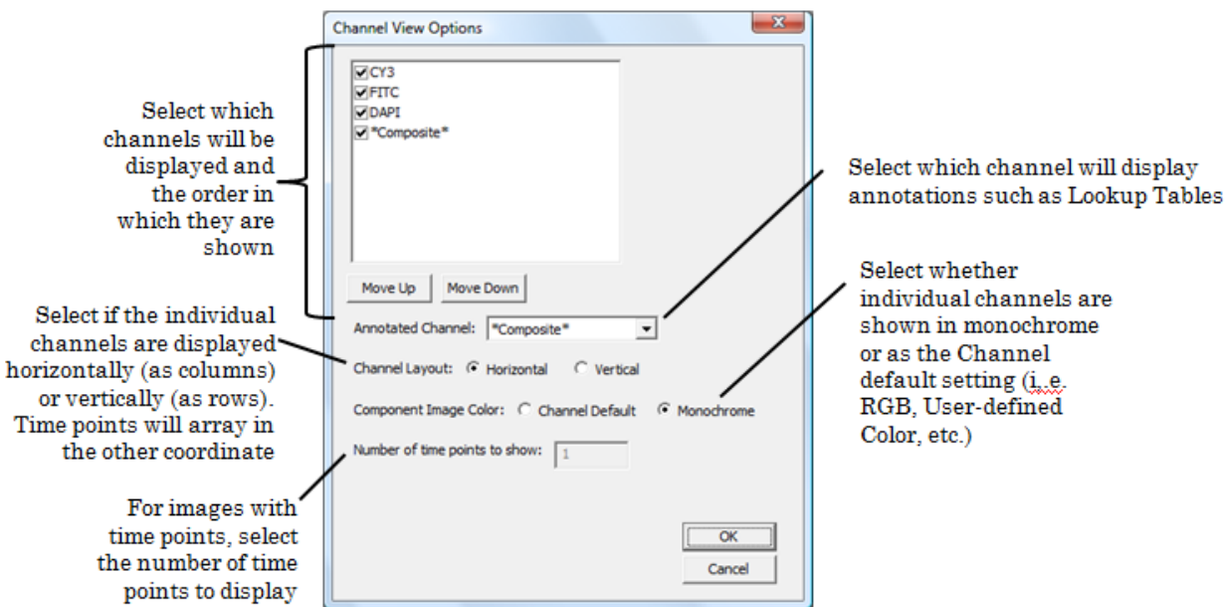
The Multidimensional Channel View allows users the flexibility to create images for export from 2D, 3D, 2D timelapse, or 4D data types. This section discusses adjusting the Multidimensional Channel View Setting in order to maximize the information shown in the

exported image. You may wish to choose a region before generating the Multidimensional Channel View in order to improve viewing ability (see [Making a 2D or 3D \(x,y,z or x,y,t\) Selection](#) on page 186). To create a Multidimensional Channel View:

1. Select an image in a slide by clicking on it.
2. Select **View > Multidimensional Channel View**. A Multidimensional Channel View will appear.
3. You may then alter the display of channels, composite images, and timepoints/z-planes by going to the **View Settings** icon in the View window (shown below) or by going to **View > Multidimensional Channel View Settings**.



4. The Multidimensional Channel View Settings dialog will open. Adjust the following parameters to achieve your desired display. Display Parameters are shown below.

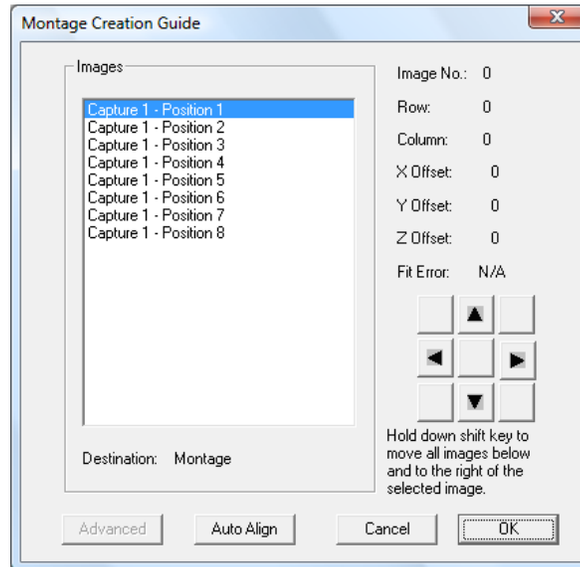


5. Once the display parameters are selected as desired, click **OK**.

### 8.2.5 Displaying a Montage

SlideBook allows you to generate montages from images captured in SlideBook using the montage feature (see page 131). To generate a montage:

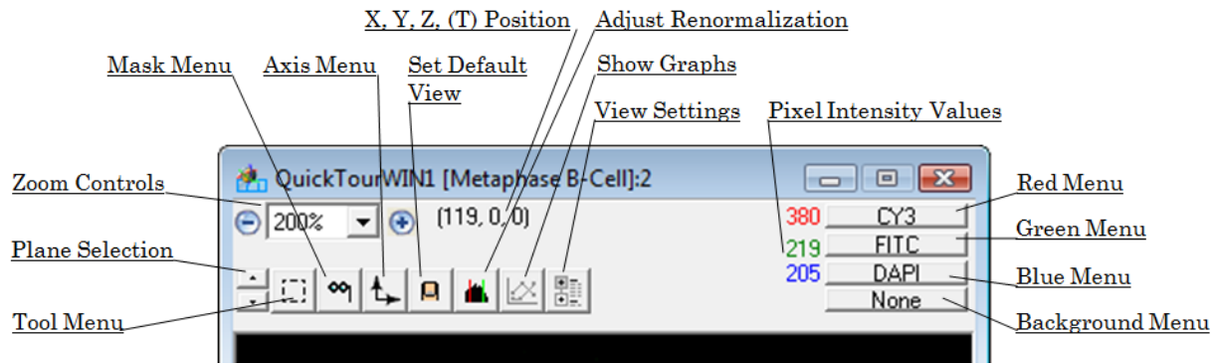
1. Open the slide that contains montage data.
2. Select **Image > Generate Montage**. SlideBook will attempt to create a montage image using only stage coordinates and pixel size information. A dialog box similar to the following will appear.



3. You may choose to manually align the images by choosing an image from list and then pressing any of the arrow buttons. Only the selected image will move. You may also hold down the shift key to move all of the images to the right and below the selected image. If you click on an image in the montage display, the appropriate image will be highlighted in the image list. Alternatively, you may choose to perform an auto-alignment by pressing the **Auto Align** key.
4. Once you are satisfied with the alignment, select **OK** to close the dialog box and save your alignment settings.

### 8.3 Manipulating Data Views Using the Info/Tool Bar

All Data Views have, at the top, a set of buttons, menus, and readouts collectively called the *info/tool bar*.



The various features are described in the table below.

Feature	Function
X,Y, Z, (T) Position	Shows and continuously updates $x,y,z$ (3D images) or $x,y,t$ (timelapse images) position of the mouse pointer. The location follows $x,y,z$ (or $t$ ) order regardless of the invisible axis that is displayed.
Zoom Controls	Displays and manipulates the magnification, or zoom factor of the image
Channel Menus	Allow the user to select the channels that correspond to R,G, and B or monochrome displays; default display corresponds with filter configuration definitions ( <a href="#">Defining Filter Configurations</a> on page 64)
Pixel Intensity Values	Shows and continuously updates pixel intensity values corresponding to the mouse pointer location for a given channel
Up and down buttons 	<p><i>Main View:</i> used to navigate through the invisible axis</p> <p><i>Three and Channel View:</i> used to navigate the <math>z</math>- or <math>t</math>-axis</p> <p><i>4D Tile View:</i> used to navigate <math>z</math>-axis</p> <p>Clicking once will move the displayed plane by one; holding continuously will scroll through the planes. Not used for 2D images.</p>
Tool Menu 	Allows the user to choose the function that the mouse and its cursor perform, as described in the section <a href="#">Using the Tool Menu</a> . This menu has different functions when a mask is displayed (see Chapter 10).
Mask Menu 	Allows the user to create and/or display a mask, or region of interest, as a blue overlay on the image. It is usually best to display image data in only red and green while displaying or editing masks.

Axis Menu 	Allows the user to choose the invisible axis in <b>Main Views</b> of 3D and timelapse data, and in <b>Tile Views</b> of 4D data. The up and down buttons are tied to the invisible axis.
Thumbnail Button 	Allows the user to alter the default view of a particular image. Once the thumbnail button is selected, subsequent views and the slide thumbnail image will retain channel settings, renormalization information, and invisible axis plane.
Renormalize Button 	Allows the user to alter the look-up table, or renormalization parameters, for each channel of the view

Now, we will explore several data view manipulations that require features of the info/tool bar.

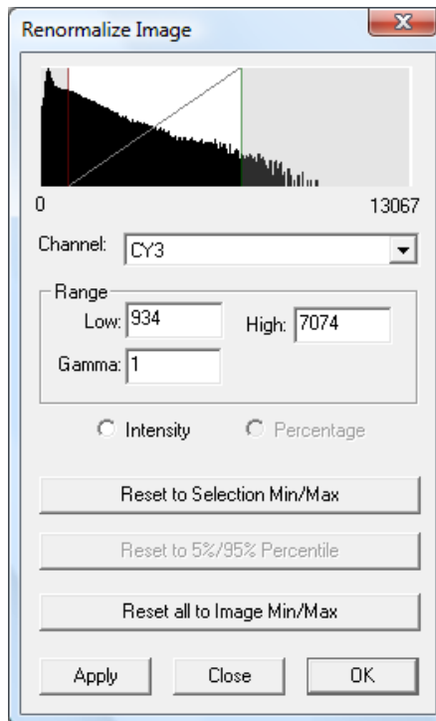
### 8.3.1 Altering the Renormalization Parameters (Lookup Table)

You will often want to highlight a particular range of data values in an image. Perhaps a particular feature is dim relative to other features, so you will want to saturate the mapping between the data and the display so that this feature will become more apparent. Or, perhaps in one channel you have a histological marker that might otherwise overpower the data and want to dim its contribution to the color values. To alter the range of data that is displayed, you will choose the minimum and maximum data values that correspond to the absence or full saturation of the color associated with each channel. You also have the ability to apply nonlinear lookup tables to your view.

#### 8.3.1.1 Changing Renormalization Parameters for a Single View

To alter renormalization parameters:

1. Create a Main View, Three View, Tile View, or Channel View of an image. If you have several views open, make sure that the desired view is the active window by clicking on it.
2. Click on the renormalize button  or select **View > Renormalize**. The Renormalize Image dialog box will appear.



A histogram is displayed showing the relative number of pixels on the y axis and intensity values on the x axis for the specified channel. The x axis value shown on the left is the lowest data value across the entire image. The x axis value shown on the right is the highest data value. In the dialog box above, the CY3 channel intensities range from 0 to 13067.

3. Select the channel that you wish to renormalize from the **Channel** dropdown menu.

The red and green bars allow you to select the minimum and maximum intensities that correspond to the absence (black) or full saturation of color in the display. Thus, any data whose value is lower than the low bar will not appear in the view. Any data whose value is above the high bar will show as fully saturated color. The display's dynamic range, then, is a linear interpolation of the values in between.

4. You may move the red and green bars and alter the display two ways.
  - a. Click and drag the red and green bars in the histogram window to the left or right. The change will automatically register once you release the mouse button.

OR

- b. Enter the desired minimum and maximum intensities in the **Low** and **High** edit fields and click on **Apply** to register the change and update the image. This gives you the opportunity to see the effect of the values and make

changes without closing and reopening the dialog. Choosing **Close** after entering intensities manually will close the dialog box without registering the change.

Again, the underlying intensity values are not affected by the renormalization process.

5. Follow steps 3 and 4 for any other channels that you wish to renormalize.
6. Choose **OK** to exit the Renormalize Image dialog box. If you would like subsequent views of the image to contain the same renormalization parameters, click on the thumbnail button .

**NOTE:** You can reset the low and high values for a single channel to correspond to the minimum and maximum intensities for that channel by pressing the **Reset to Selection Min/Max** button. You can also set all channels to the global min/max (the lowest and highest values of any channel present in the image) by pressing the **Reset all to Image Min/Max**. This is particularly useful when working with transmitted light color images.

Remember that renormalization is a property of a particular view, not the entire image. It is possible to have multiple views on the same image each with their own renormalization values. If, however, you decide that a particular selection of values is best for most viewings of an image, you can press the thumbnail button in the corresponding view and make those values the default for that image (see [Changing the Default Display](#) on page 180).

### 8.3.1.2 Changing Renormalization Parameters for a Group of Images in a Slide

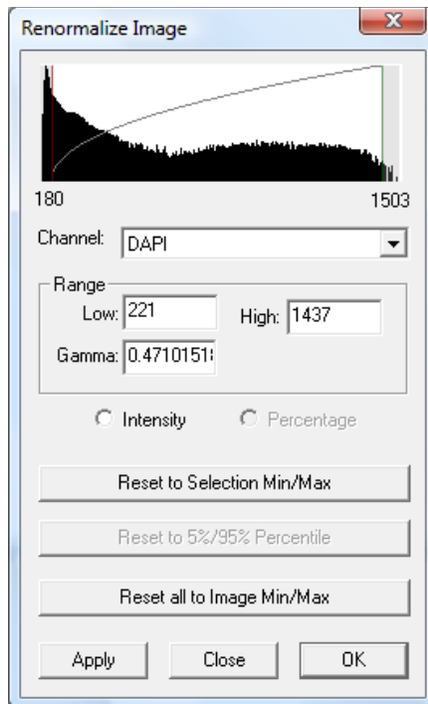
You may apply the default renormalization parameters of an open view to all images in a slide by selecting **Image > 4D Operations > Renormalize All**.

**NOTE:** You **MUST** set the display as default using the thumbnail button before you apply the settings across the images in a slide.

You may also set the renormalization to the global minimum and global maximum for each channel for all images in a slide. To do so, select **Image > 4D Operations > Renormalize All to Global Min/Max**. This function finds the lowest minimum and highest maximum pixel value for the group of images in the slide on a per channel basis, and then uses those values to set the renormalization of all images in the slide.

### 8.3.1.3 Applying Nonlinear Lookup Tables

SlideBook allows you to apply nonlinear lookup tables using a gamma correction. This feature is particularly useful for emphasizing dim features in an image. To apply a nonlinear look-up table, simply click and drag the gray line that crosses the histogram diagonally. This adjusts the value of gamma. Alternatively, you may enter a value for gamma manually and select **Apply**. If you wish to return the gamma to 1 (no gamma correction), simply click-and-drag the line off of the histogram and it will return to 1.



### 8.3.2 Changing the Display Colors

Images may be displayed using a variety of color schemes, including RGB, monochrome, inverted monochrome, pseudocolor, and user-defined color. SlideBook also supports a mixed display that blends a monochrome background image (e.g., DIC) with an RGB image. The default display is determined by the filter configuration definitions (see [Defining Filter Configurations](#) on page 64), though you may easily change the colors that are displayed for a given view. Generally, the filter configurations will be set to R, G, or B. However, monochrome display may be useful for visualizing a single channel. Pseudocolor display may be useful for visualizing one or two channels.

#### 8.3.2.1 Changing the Channels Displayed on an RGB Image

1. Make the desired view the active window by clicking on it.
2. Select the fluorophore that you would like to display from the corresponding drop-down channel menu in the info/tool bar.

**NOTE:** If your image is displayed as something other than RGB, you may change the display to RGB by choosing **View > RGB Color**.

#### 8.3.2.2 Changing an Image Display to Monochrome

1. Make the desired view the active window (by clicking on it).
2. Select **View > Monochrome** to change the display to monochrome.

3. Click on the channel menu, and select the fluorophore that you would like to display from the dropdown list

### 8.3.2.3 Changing an Image Display to Pseudocolor

1. Make the desired view the active window by clicking on it.
2. Select **View > Pseudocolor** to change the display to pseudocolor.

Two channel menus will be available. The first channel will be displayed as a pseudocolored image ranging from saturated red to saturated blue, known as **Pseudocolor (Color)**.

3. Click on the first channel menu, and select the fluorophore that you would like to determine the pseudocolor hue from the dropdown list.

The second channel menu will act to gate the pseudocolor image based on intensity. This second channel is called **Pseudocolor (intensity)**. Thus, an area with a high pixel intensity for the first channel and a low pixel intensity for the second channel will appear as dim red. An area with a low pixel intensity for the first channel and a high pixel intensity for the second channel will appear as bright blue.

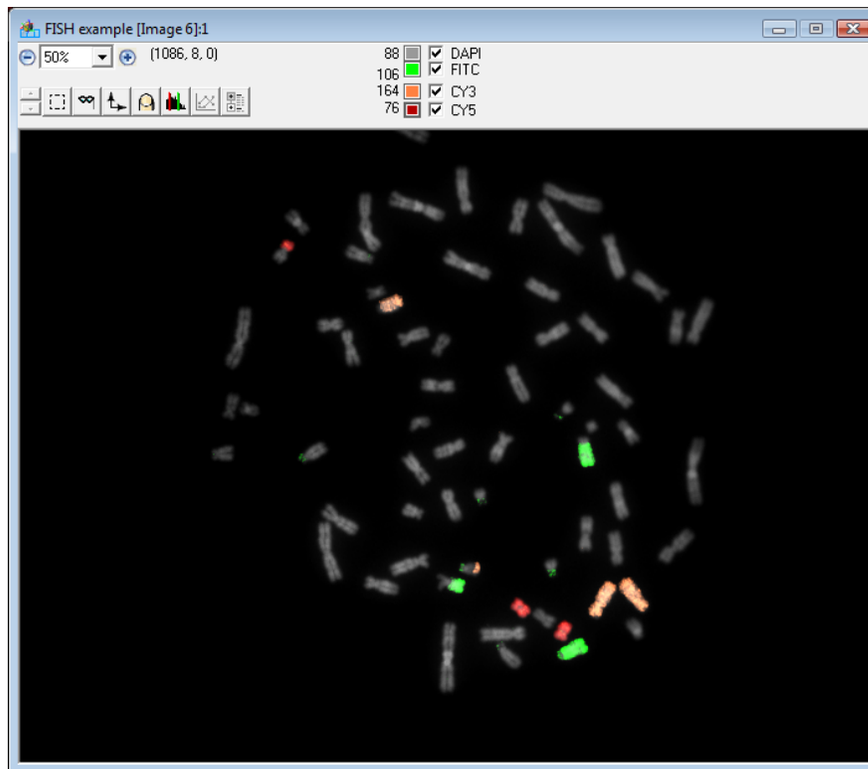
4. If you would like to gate the first channel by the intensity of a second channel, select a second channel from the dropdown menu.

### 8.3.2.4 Changing the Display to a User-Defined Color Palette

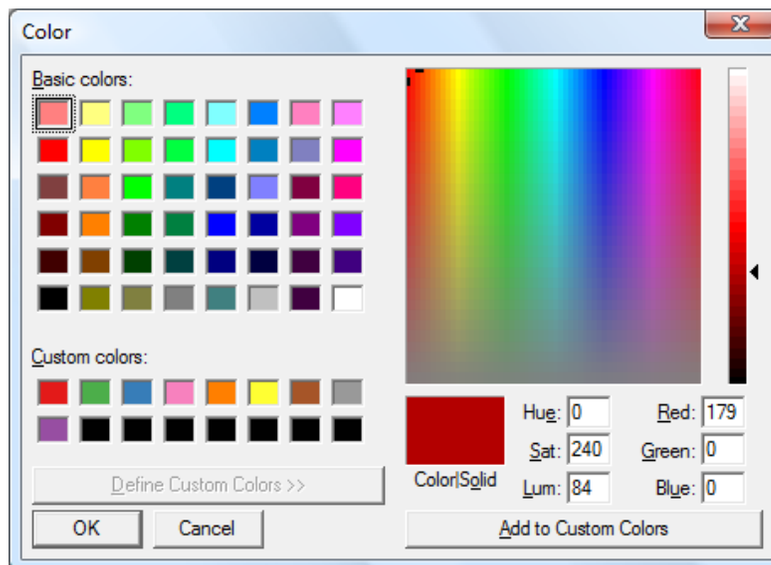
You can now display up to eight channels in colors of your choosing. This is useful for applications such as fluorescence *in-situ* hybridization (FISH).

To use this feature:

1. Open a Main View, Three View, or Tile View.
2. Select **View > User-Defined Color**. Your image will appear in the following format. All channels that were captured will be listed in the channel list.



3. To change the color of one of the channels, click on the colored box next to the channel whose color you would like to change. The following dialog box will appear.



4. To change the color, either click on one of the **Basic colors** in the upper left section of the dialog, or click on one of the **Custom colors** in the lower left, or create your own color on the right side. If you would like to add a color to the **Custom colors**, use the color picker on the right to select a color, then select **Add to Custom Colors**.

5. To add a channel to display, simply select the checkbox next to the channel.

You may choose to always display a channel using a color of your choosing by specifying it in the **Filter Definition** (see page 66).

### 8.3.2.5 Displaying a DIC image as Background of an RGB Image

In an RGB image, the first three popup menus control which channel is visible in red, green, and blue respectively. The last popup menu, labeled **Bkgnd**, selects a background display channel. In multi-channel images, one channel can be chosen to be displayed in monochrome wherever the other channels would not otherwise be visible. This can be particularly useful for showing DIC information alongside fluorescence. To display DIC alongside a fluorescent RGB image:

1. Create a Main View, Three View, Tile View or Channel View of an image that contains a DIC channel and at least one fluorescent channel.
2. Make sure that the fluorescent channels are displayed as RGB. If they are not, select **View > RGB Color** to change the display to RGB, and select the fluorophores to be displayed from the channel menus.
3. Set the **Bkgnd** channel menu to DIC.

The DIC image will show through in any pixel that has RGB intensities that are below the minimum intensity threshold of the fluorescent channels. If you would like to increase the amount of DIC that shows through, you must raise the minimum intensity threshold on all of the RGB channels (See [Altering the Renormalization Parameters \(Lookup Table\)](#) on page 171).

4. To change the display so that the DIC and fluorescent channels are blended at 50% opacity, select **Blend Background** from the **View Settings** button  or select **View > Blend Background**. You will not need to alter the minimum intensity threshold on the RGB channels to see DIC.

### 8.3.3 Cropping an Image

You may wish to crop an image in order to reduce its size or center on a specific portion of the image.

**CAUTION:** Cropping is an irreversible process. You may wish to make a copy of the image before cropping (see [Copying an Image](#) on page 159) or select **Crop to New Image**.

To crop an image:

1. Open a **Main** or **Three View** of the desired image.
2. Define a 2D or 3D selection as described in [Making a 2D or 3D \(x,y,z or x,y,t\) Selection](#) on page 186.

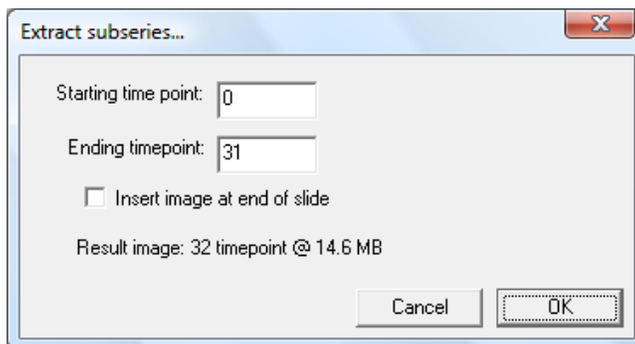
3. Select **Image > Crop** to replace the existing image, or **Image > Crop to New Image** to generate a new, cropped image.

### 8.3.3.1 Extracting Timepoints from a Timelapse Image

You may wish to extract only a subset of timepoints from a 2D Timelapse or 4D data series.

To extract timepoints:

1. Select an image in an open slide or open a Main View of the desired image.
2. Select **Image> Extract Subseries to New Image**. The following dialog will appear:



3. Enter the starting and ending timepoint of the subseries you would like to extract.
4. Check the box for **Insert image at end of slide** if you would like to save the subseries in the same slide as the original data. If you do not check this box, the image will save below the active image in the Slide View.

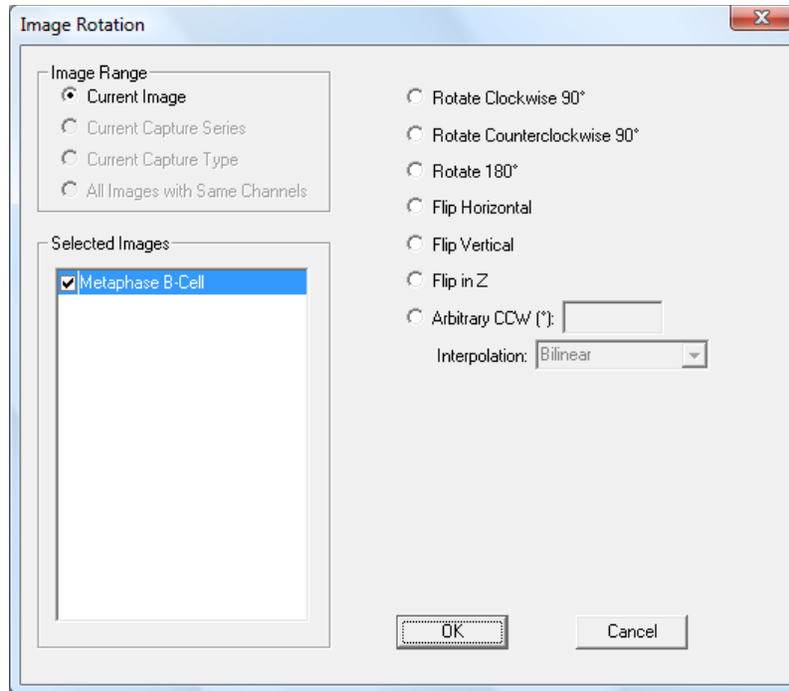
### 8.3.4 Rotating an Image

You may wish to rotate a view of the image for publication purposes.

**CAUTION:** Rotation is an irreversible process. You may wish to make a copy of the image before rotating (see [Copying an Image](#) on page 159).

To rotate an image:

5. Open a Main View of the desired image.
6. Select **Image> Rotate**. The following dialog box will appear:



7. Select the **Image Range** as described below. You may also select and deselect images using the checkboxes in the **Selected Images** list.

- **Current Image** – Operation will be performed on the selected image only.
- **Current Capture Type** – Operation will be performed on all images in the slide with the same capture type (2D, 3D, 4D, etc.) as the selected image.
- **All Images with Same Channels** – Operation will be performed on all images in the slide with the same channels as the selected image.

8. Select the desired rotation angle and select **OK**. The image will be rotated as specified.

### 8.3.5 Changing the Invisible Axis

SlideBook allows for 3D and timelapse data to be viewed in all possible dimensions. By default, a Main View will display the  $x,y$  plane (invisible  $z$  axis for 3D and  $t$  axis for timelapse). However, you can change the invisible axis to the  $x$  or  $y$  axis if you wish. To view your image with a different invisible axis:

1. Open a Main View of the desired image as discussed above.
2. Click on the axis menu  in the info/tool bar and select the desired invisible axis from the dropdown menu. The up and down arrows in the info/tool bar will be tied to the new invisible axis.

**NOTE:** The axis menu is not available in the Tile View or Channel View. If you would like to view a different invisible axis in the Tile View you should first open a Main View, change the invisible axis, make a 3D selection and then spawn a Tile View as described above ([Displaying a Timelapse Tile](#) on page 164).

### 8.3.6 Scrolling through the Invisible Axis in a 3D or Timelapse Image

There are several ways to scroll through the invisible axis in Main Views, Channel Views, and 4D Tile Views.

- Using the up and down arrows  in the info/tool bar: one click on either arrow will move one plane up or down; holding continuously will scroll.
- Using menu commands: **View > Prev Plane**, **View > Next Plane**, **View > Go to Plane** and **View > Loop** will move the invisible axis location one plane down, one plane up, to a specified plane, or in a continuous loop, respectively. See page 27 for instructions on how to use the **Go to Plane** feature.

**NOTE:** To scroll around Three View images, see the section titled [Scrolling through a Three View](#) on page 187.

### 8.3.7 Changing the Default Display

Once you have an image that has the color scheme and renormalization parameters that you desire, you may save those parameters so that all subsequent views retain those parameters. To do so:

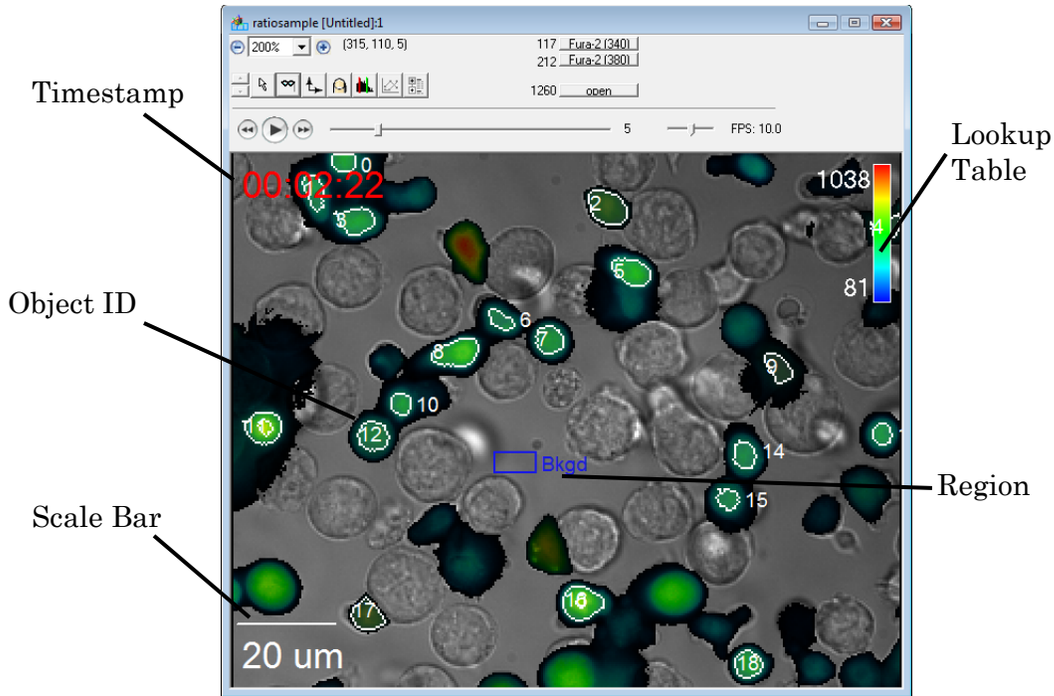
1. Using a Main View, Three View, or Tile View, adjust the color display and renormalization parameters as described in the sections above.
2. Click on the thumbnail button  in the info/tool bar to set new default display parameters. The thumbnail image in the slide view and all subsequent views will display the new default parameters.

**NOTE:** When using a Main View, you may wish to scroll to a desired plane in the invisible axis before clicking on the thumbnail. The selected invisible axis plane will be displayed in subsequent Main Views.

### 8.3.8 Displaying Annotations

The following annotation features are available in SlideBook 4.2:

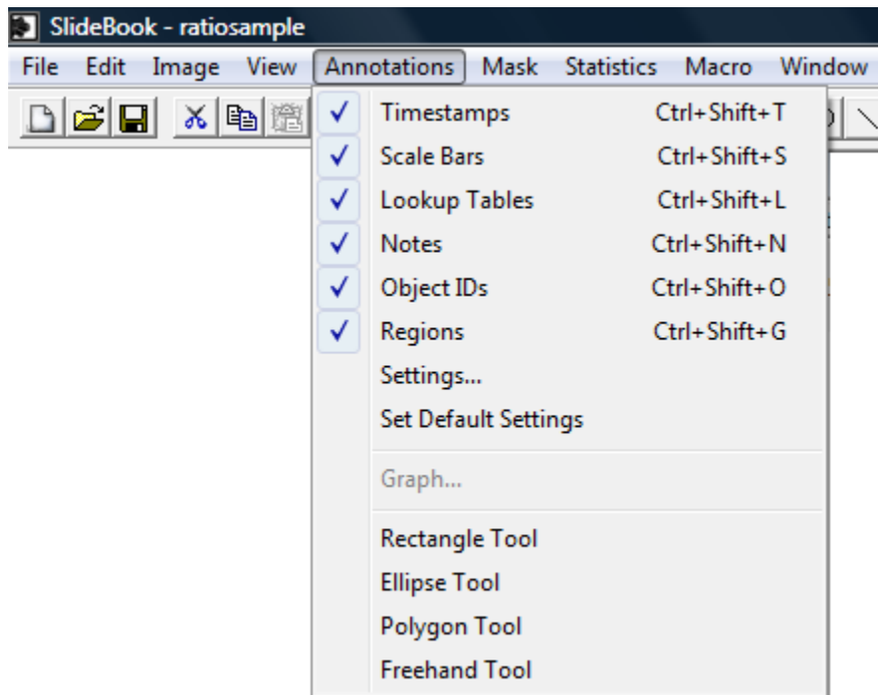
- **Timestamps** – available for timelapse captures
- **Scale Bars** – bar showing relative scale/size of image
- **Lookup Tables** – range of colors that correspond to intensities for a given channel
- **Notes** – formerly called *Annotations* in previous SlideBook versions, can be used to record experimental actions (see [Creating Notes](#) on page 116)
- **Object IDs** – corresponding to objects in a displayed mask
- **Regions** – selections made using ROI tools



### 8.3.8.1 Displaying Annotations

To display any of these annotations on an open view:

1. Open a Main View, Three View, or Channel View of your data.
2. Select the item you wish to display from the **Annotations** menu.

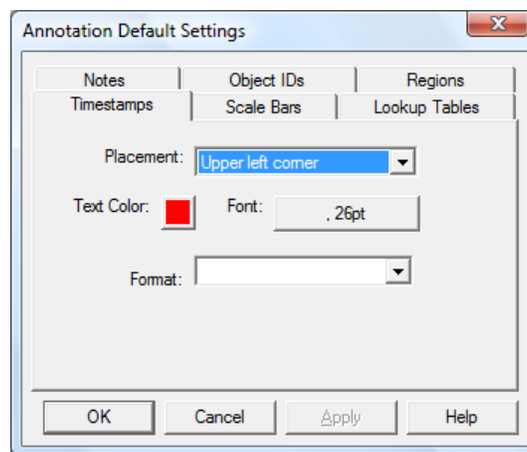


3. When an item is selected for display, a checkmark will appear next to the item on the menu.

### 8.3.8.2 Formatting Annotations

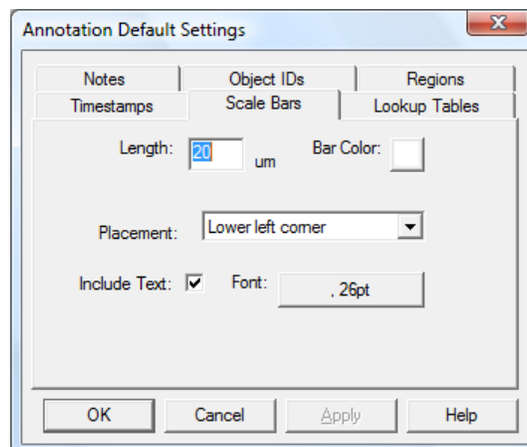
Several formatting options for annotations are available. Once you have displayed your annotations, you may format by doing the following:

1. Open the Annotation Default Settings dialog by selecting **Annotation Settings** from the **View** icon  or by going to **Annotations > Settings**. The following dialog will appear:

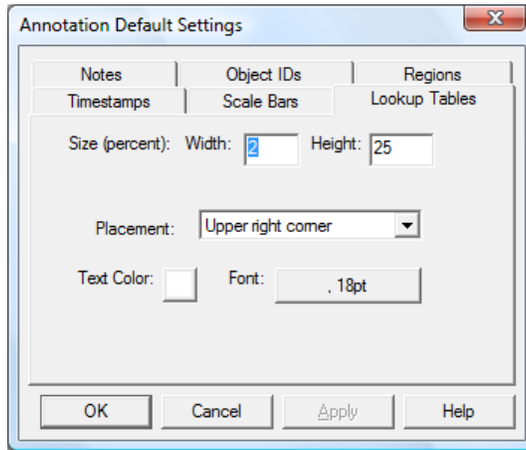


2. Select the annotation that you would like to format. The dialog boxes for annotations are shown below.

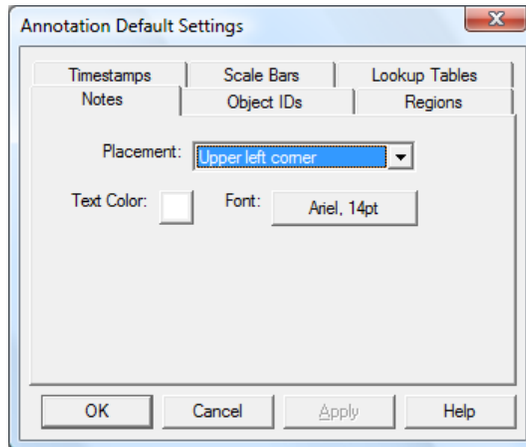
**Scale Bars** – You may define the length of the scale bar; choose color, placement and font; and include text or not.



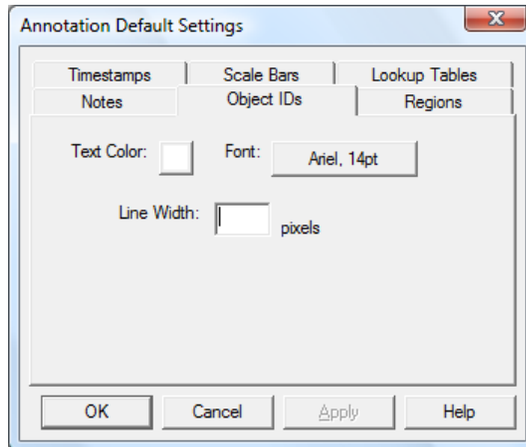
**Lookup Tables** – You may choose the size of the lookup table (as a percentage of the image width and height) as well as placement, text color and font.



**Notes** – You may choose placement, text color, or font.



**Object IDs** – You may select text color and font. Note that object IDs only appear if you have a mask displayed that has been divided into objects.

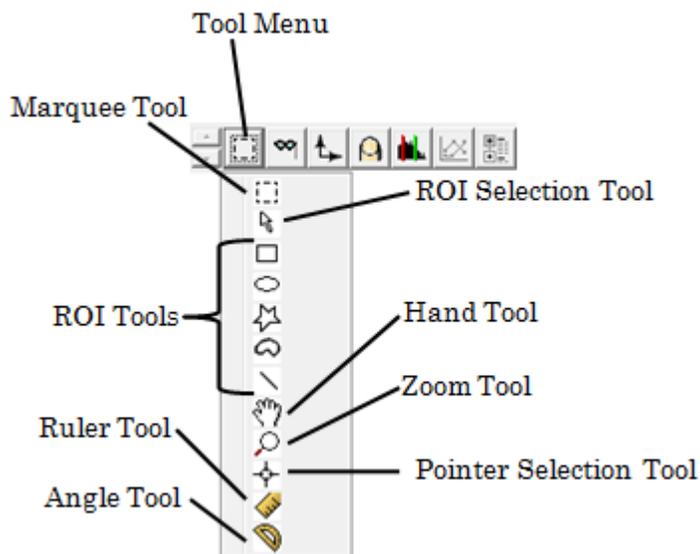


### 8.3.8.3 Setting Default Annotation Settings

Once you have adjusted the Annotations to the desired display, you can save the settings as defaults by going to **Annotations > Set Default Settings**.

## 8.4 Using the Tool Menu

The tool menu of the info/tool bar allows the user to choose the function that the mouse and its cursor perform.



The tool menu may also be accessed by selecting **View > Tool**. The various features are described in the table below:

Feature	Function
Marquee Tool 	Lets you select a 2D or 3D region of the image and is used when spawning a Tile View or a volume rendering (Making a 2D or 3D (x,y,z or x,y,t) Selection on page 186). This tool can also be used to make masks (see Chapter 9).
ROI Selection Tool 	Lets you select an ROI in order to perform further operations. Holding shift allows you to select multiple ROIs. See <a href="#">Creating ROIs and Graphs to Monitor Regions of Interest</a> on page 118 for more information on working with ROIs. This tool is also available from the SlideBook toolbar.
Rectangle ROI Tool 	Allows you to draw rectangular ROIs on an open view by clicking and dragging. This and all other ROI drawing tools are available from the SlideBook toolbar at the top of the application window.

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<p>Ellipse ROI Tool </p>	<p>Allows you to draw elliptical ROIs on an open view by clicking and dragging.</p>
<p>Polygon ROI Tool </p>	<p>Allows you to draw closed polygons by clicking to draw a series of straight connected lines. Double-clicking closes the shape.</p>
<p>Freehand ROI Tool </p>	<p>Allows you to draw freehand by clicking and holding. Shapes close automatically when mouse button is released.</p>
<p>Hand Tool </p>	<p><i>Main and Three Views:</i> lets you move the displayed portion of an image in a window smaller than the dimensions of the displayed axes (performs the same function as the horizontal and vertical scroll bars).</p> <p><i>Tile View that displays a portion of an image:</i> adjusts the position of the display cube within the image, therefore scrolling the portion of the image that is visible in all planes.</p>
<p>Zoom Tool </p>	<p><i>Main and Three Views:</i> grows or shrinks the size of the window and image by a factor of two.</p> <p><i>Tile View:</i> the overall size of the window and its panes remains constant, but the zoom factor of the displayed image is adjusted by a factor of two.</p> <p>Clicking on a view will zoom in (double the size) and clicking while holding down the shift key will zoom out (halve the size).</p>
<p>Point selection Tool </p>	<p><i>Three View:</i> updates the displayed position in the other two panes by clicking or dragging in any of the three panes. Dragging over a pane will continuously update the other two panes.</p> <p><i>Main View:</i> Ctrl-click on a region in a 2D timelapse image to display a graph of intensity vs. time.</p> <p><i>No function in Tile View or Channel View.</i></p>
<p>Ruler Tool </p>	<p>Measures a desired distance in 2D and 3D images, and velocity in 2D timelapse images. See <a href="#">Making Measurements with the Ruler Tool</a> below.</p>
<p>Angle Tool </p>	<p>Measures a desired angle in 2D and 3D images. Click and drag to draw a single line, then make a second click at the angle you wish to measure. The angle measurement appears next to the zoom factor.</p>

Now, we will explore several data view manipulations that require features of the tool menu.

### 8.4.1 Making a 2D or 3D ( $x,y,z$ or $x,y,t$ ) Selection

As discussed above, the ability to make a 3D selection is useful when spawning a Tile View. You may also want to make a 3D (or 2D) selection in order to perform volume rendering (see page 189), crop an image (see page 177), or generate statistics for analysis (see page 227). 3D selections will be visible in all three panes of the Three View, and when scrolling through the invisible axis in the Main and Channel Views.

#### 8.4.1.1 Making a 2D Selection in any Data View

1. Select the Marquee tool  from the tool menu. This is the default tool in Main, Tile, and Channel Views.
2. Click and drag to designate a 2D selection.

There are three ways to make 3D selections. 3D selections may be for 3D volumes ( $x,y,z$ ) or 2D timelapse ( $x,y,t$ ) images. One of the methods does not apply in Tile Views.

#### 8.4.1.2 Making a 3D Selection in a Main, Three, and Channel View

1. Select the Marquee tool  from the tool menu.
2. Click and drag to designate a 2D selection, and continue to hold the mouse button down.
3. Hold the shift key down while continuing to hold the mouse button and move the mouse up or down to move the display along the invisible axis.
4. Release the shift key, and then the mouse button.

#### 8.4.1.3 Making a 3D Selection in any Data View

1. Select the Marquee tool  from the tool menu.
2. Click and drag to designate a 2D selection and release the mouse button
3. Extend the 2D selection across the third dimension by
  - a. Selecting **View > Select Rect in all Planes**. This will extend the selection to all planes of the third dimension.

OR

- b. Selecting **View > Define Selection Cube** and entering the location range of the  $z$  or  $t$  planes that you would like to include in your selection. You must click **OK** to complete the selection.

**NOTE:** You may also use the **View > Define Selection Cube** menu selection to enter  $x$ ,  $y$ , and  $z$  (or  $t$ ) dimensions. The  $x$  and  $y$  dimensions are pixel locations, while the  $z$  and  $t$  dimensions are plane numbers.

#### 8.4.1.4 Selecting an Entire Image

If you would like to return your selection to the entire image, simply select **View > Select All**. This will negate any previous selections.

### 8.4.2 Scrolling through a Three View

The point selection tool  allows you to easily maneuver around the Three View, and it is the default tool for the Three View. Click the point selection tool somewhere in the pane where  $z$  is the invisible axis (the  $x,y$  pane), and the  $x$  axis ( $y,z$  pane) and  $y$  axis ( $x,z$  pane) will both scroll to the same coordinates that you selected in the  $z$  axis pane. Clicking and dragging around any of the panes will continuously update the other two panes.

### 8.4.3 Making Distance or Velocity Measurements

This tool can be used in any of the views to measure a desired distance (2D, 3D images) or velocity (2D timelapse images). In all cases, it is critical that both the optical parameters of the objective (those that determine  $\mu\text{m}/\text{pixel}$ ) are accurately defined (see [Defining Objectives](#) on page 62) and the  $z$  or  $t$  interval is specified in the image info (see [Getting and Editing Image Information After Capture](#) on page 101). The ruler tool makes one measurement at a time, and each measurement replaces the previous measurement. In other words, the ruler tool is not used for permanent measurements, but it can be used to make scale bars or to make quick and easy measurements.

#### 8.4.3.1 Making a Measurement on a 2D Image

This measurement can be made in any of the views.

1. Select the ruler tool  from the view's tool menu.
2. Click and hold the mouse button on the image at the point where you would like to begin measurement.
3. Drag the mouse to the desired end point and release the mouse button.

Alternatively, you may click and release once at the start point and click and release at the end point.

The length of the line that you have just drawn will be displayed in microns in the info/tool bar. Note that these distances cannot be saved to disk from within SlideBook. In order to perform measurements that are to be saved, see Chapter 10.

4. To remove the line from the view, click once anywhere on the view.

#### 8.4.3.2 Making a Measurement on a 3D or 2D Timelapse Image

1. Open a Three View of the desired image by clicking once on the thumbnail image, and then selecting **View > Three View**.
2. Click the mouse button on the image at the point where you would like to begin measurement. This point may be in any of the three panes.
3. Click the mouse button on the image at the point where you would like to end measurement. Again, this point may be in any of the three panes.

The length or velocity between the two points will be displayed in microns in the info/tool bar area. Note that these distances can not be saved to disk from within SlideBook. In order to perform measurements that are to be saved, see Chapter 10.

4. To remove the line from the view, click once anywhere on the view.

#### 8.4.3.3 Making a Measurement on an Imported Image

1. Import the image as discussed in Importing an Image on page 96.
2. Make sure that the objective definitions exist by selecting **Edit > Define Optics > Objectives** and looking at the available objectives in the dropdown list. If the objective is not available, add the objective definition as described on page 62. It is extremely important that the **microns/pixel** data field is reported accurately.
3. If a magnification changer or relay lens was used, make sure that its definition exists by selecting **Edit > Define Optics > Mag. Changers** and looking at the available magnification changers in the dropdown list. If the magnification changer is not available, add a new magnification changer definition as described on page 73.
4. Add the objective and magnification changer information to the image by selecting **Image > Get Info**, pressing the **Edit Info** button, and then selecting the objective and magnification changer information from the dropdown menus. You will also want to select the capture type from the dropdown menu.
5. Select **OK** to exit the **Edit Image Info** dialog box, then **OK** in the **Image Info** box to save the image information.
6. Use the ruler tool as discussed above to perform distance and velocity measurements.

#### 8.4.3.4 Using the Ruler Tool to make a Scale Bar

1. Open a Main View or Three View of the image.
-

2. Select the ruler tool as discussed above to draw a line of the desired length, observing the distance reported in the info/tool bar.
3. To preserve the line, export the view of the image as discussed below in [Exporting Views](#) on page 196. The scale bar will be present in the exported image. This image can then be further modified using a drawing program.

### 8.5 Using Display Views (Creating Renderings and Movies)

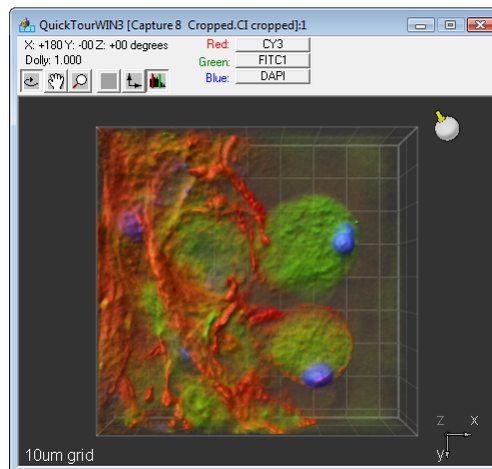
Besides data views, SlideBook can generate two types of movie displays, volume renderings and series movies. Both must be created from an open data view (Main, Three, Tile, or Channel View) and will use the same color assignments and renormalization values as that data view. The volume rendering will be generated and displayed from within SlideBook, while the series movie must first be saved and then later viewed in QuickTime. The volume rendering can also be exported and viewed using QuickTime.

#### 8.5.1 Displaying a 3D or 4D Volume View (Volume Rendering)

SlideBook can perform interactive volume renderings using the *3D Volume View*. The following rendering types are available. When you go to **View > 3D Volume View**, two options will be available, **High Speed** and **High Quality**. **High Speed** will render the image at the highest speed possible. **High Quality** will render the image with a more rigorous algorithm that may improve the quality of the image with a small decrease in the speed performance. These settings apply to all modalities of volume rendering but are most noticeable when using **Dynamic Lighting** mode.

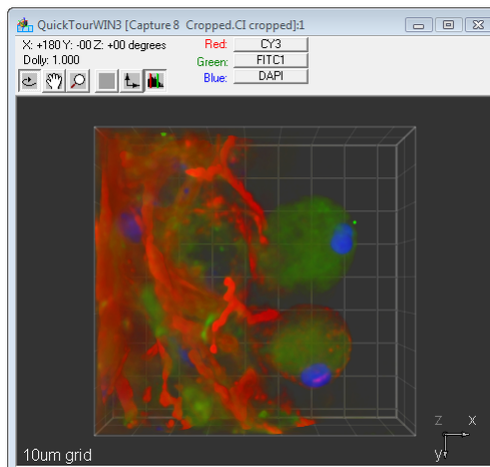
Examples of the various modes of volume rendering are shown below:

- **Dynamic Lighting** – allows you to illuminate your object as you wish by allowing control of the lighting angle. Useful for determining surface characteristics.



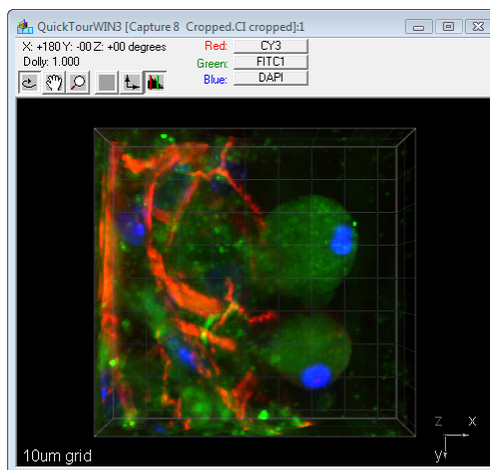
Dynamic Lighting

- **Fixed Lighting** – uses a fixed lighting pattern for shading, approximates surface characteristics.



**Fixed Lighting**

- **Maximum Intensity Projection (MIP)** – The brightest pixels through the axis perpendicular to the screen are displayed. This algorithm requires the least processing power and is particularly useful for rendering large data sets (and does not require advanced graphics card).



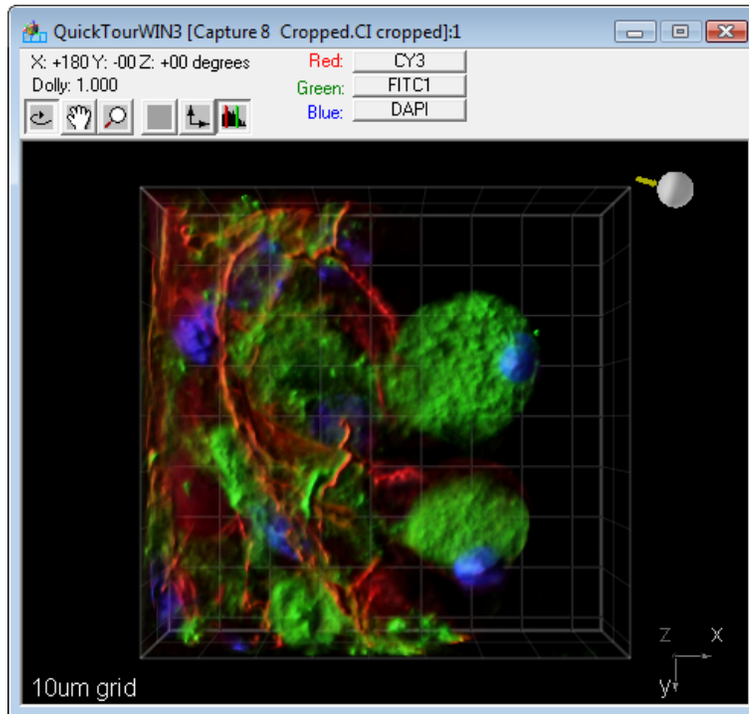
**MIP**

- **X Ray** – Similar to MIP; however, the sums of pixels through the axis perpendicular to the screen are displayed. This mode is useful when trying to visualize density or concentration differences.

### 8.5.1.1 Working with 3D Volume Views

To open a new 3D Volume View:

1. In a Slide View, click once anywhere on the thumbnail or within the shaded region that surrounds the thumbnail to highlight the image.
2. Select **View > 3D Volume View** or **View > 4D Volume View** and select either **High Speed** or **High Quality**. A new window will appear:

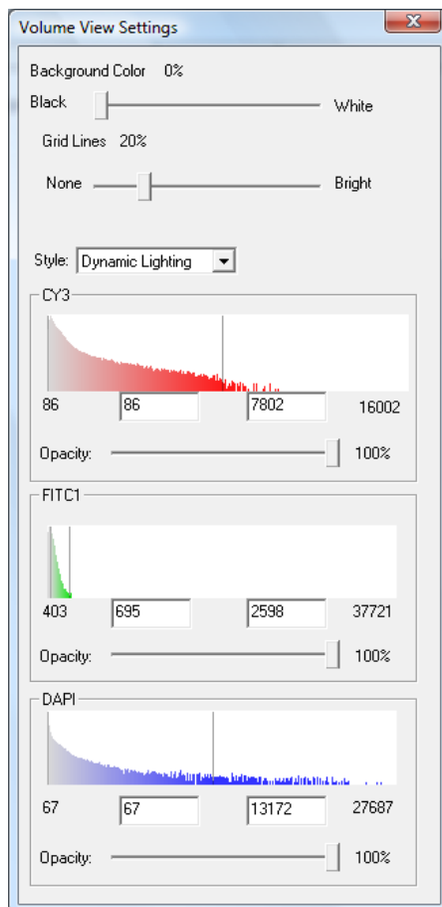


### 3-D Dynamic Lighting Volume View

3. To change the channels that are displayed, use the channel menus as you would for the Main View, Three View, or Tile View.
4. To alter the shading, grab the Lighting Tool in the upper right hand corner  and rotate it to achieve the desired lighting.
5. To zoom into the view, click on the zoom tool  and then click and drag up and down in your view to zoom in and out.
6. To rotate the image, click on the rotate tool  and then click and drag in the direction that you wish to move the image. You may either click and drag directly on the image or you on the coordinate image in the lower right hand corner.
7. To move the image in the field of view, click on the Hand Tool icon  and then grab and move the image by clicking and dragging on it.

- To instantly reorient your image, you may choose to display a specific face of the volume by choosing the Axis Tool  and selecting **Top**, **Bottom**, **Left**, **Right**, **Front**, or **Back**.
- To select a different rendering mode, or to alter the colors displayed, select the Renormalization icon . The **Volume View Settings** window will appear.

**NOTE:** The **Volume View Settings** window also automatically opens when you open a 3D Volume View.



Several features are offered in addition to the ability to renormalize the image (change the lookup tables).

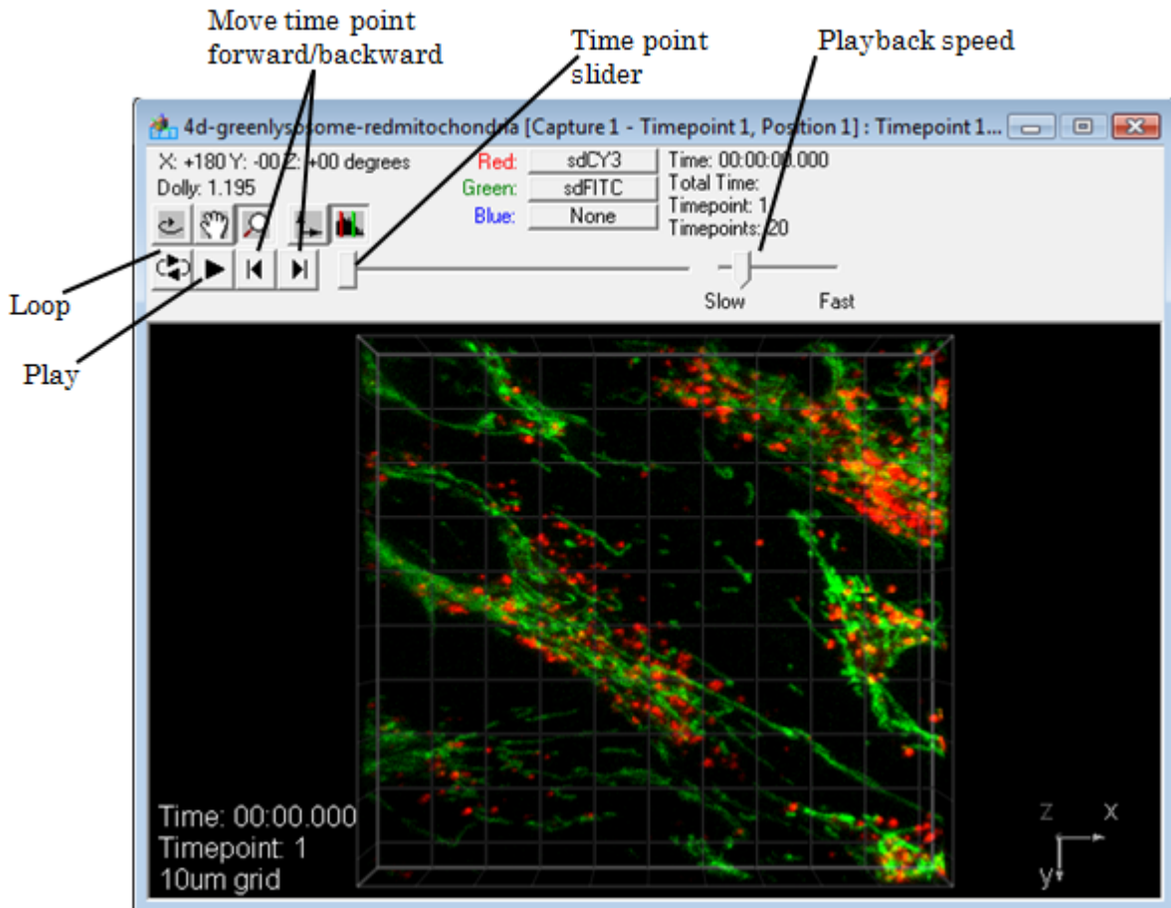
- Alter the background color using the **Background Color** slider and change the brightness of the gridlines using the **Gridlines** slider.
- You may alter the opacity of an individual channel using the corresponding **Opacity** slider control.
- Select the rendering mode from the **Style** drop-down menu.

13. You may also change the channels that are displayed just as you would in other standard views.

### 8.5.1.2 Working with 4D Volume Views

The 4D Volume View has the same functions as the 3D Volume View plus a timepoint navigation feature.

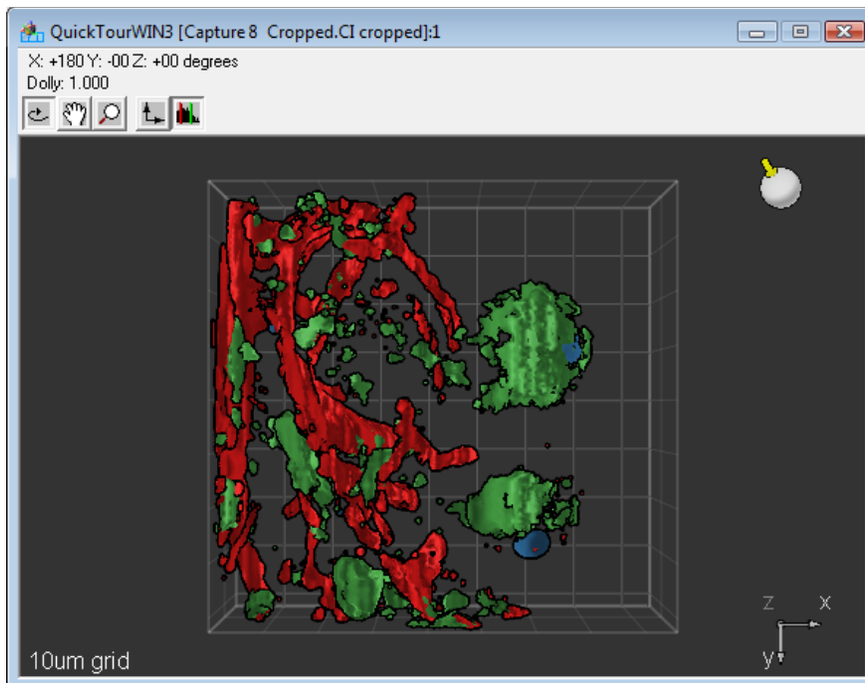
To display a 4D Volume View, click on a thumbnail in your slide and select **View > 4D Volume View**.



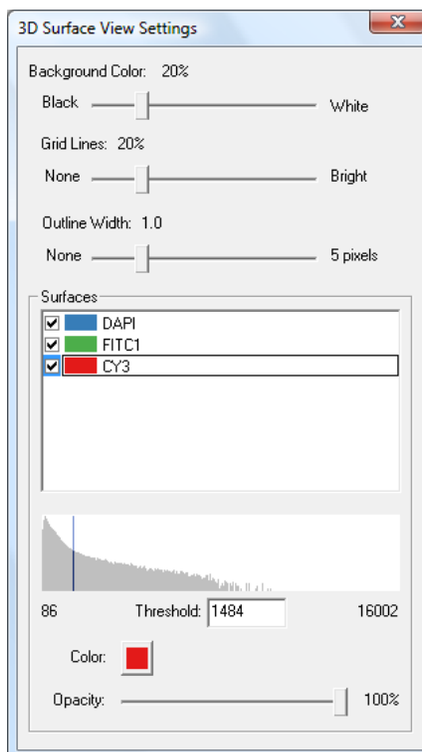
You may use any of the timepoint controls to move between timepoints or play the images like a movie. Other controls such as rotate, zoom, hand, axis, and histogram function exactly as in the **3D Volume View**.

### 8.5.2 Displaying a 3D Surface View

You may also display your image as a surface; to do so, click on a thumbnail in your slide and select **View > 3D Surface View**. A new window and corresponding renormalization dialog will appear.



This view has functions similar to the 3D Volume View. The rotate tool, hand tool, zoom tool, axis tool, and lighting tool all function as described above in the section on 3D Volume View. The **3D Surface View Settings** function similarly to the Volume View Settings. To change the intensity at which the surface is drawn, adjust the bar on the histogram of pixel intensities.



To display or remove surfaces, simply check or uncheck the appropriate channels in the **Surfaces** list. You may also change the color of the surface by selecting the color box for the corresponding channel. You may also change the width of the outline using the **Outline Width** slider.

### 8.5.3 Generating a Physically Proportional Rendering of 3D Data

You may notice that your object may appear disproportionately small, or “squashed” in the  $z$  direction. When performing 3D imaging, each plane that is captured is represented by a set of pixels in the image set. Each pixel is a perfect cube. You will notice that the  $z$  dimension in the 3D rendering appears to be different than what was actually captured. This is because the  $z$  step size that was selected during capture is not actually the same as the length of the  $z$  dimension of the pixel. In the image set, planes are simply stacked on top of one another with no regard for the actual  $z$  distance traveled (this is done in order to preserve data fidelity).

To generate a proportionate rendering:

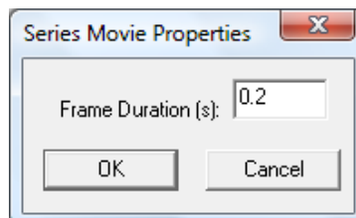
1. Open a data view (Main, Three, Tile, or Channel View) of the image as discussed above.
2. Alter the renormalization parameters and color display as desired (see above).
3. Interpolate between planes and expand your  $z$  dimension to represent the actual  $z$  distance traveled by the microscope by generating an isotropic image (see page 213).
4. Create a Volume or Surface View as discussed in the sections on pages 189 and 193.

### 8.5.4 Creating Series Movies

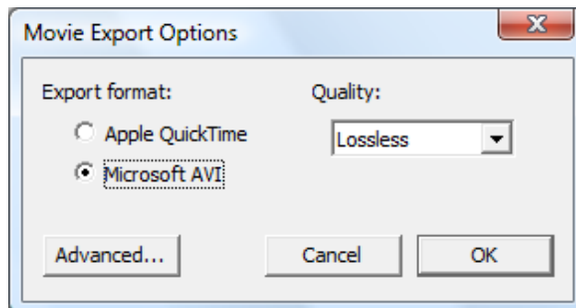
#### 8.5.4.1 2D Timelapse Images

Scans through time lapse images can be converted into QuickTime movies. To do so:

1. Open a Main View of the image as discussed above.
2. Alter the renormalization parameters and color display as desired (see pages 171 and 174). You may also wish to crop the image (see page 177).
3. Select **View > Create Series Movie**. The Series Movie Properties dialog box will appear:



4. Enter the speed at which you would like the movie to play in the **Frame Duration (s)** edit field. A value of 0.2 seconds generally yields acceptable results.
5. Click **OK**. The following dialog box will appear:



6. Select the export format and quality that you prefer and then click **OK**. Lossless has the highest quality and the largest file size. A standard “Save As...” dialog box will appear.
7. Type in a filename and click **Save** and you are finished.

#### 8.5.4.2 4D Images

There are two ways to generate movies of 4D images.

1. Open a slide that contains a 4D Capture.
2. Generate a projection image as discussed in [Creating a Projection Image](#) on page 213. SlideBook will automatically generate a 2D timelapse image from the series of projection images.
3. You may now generate a series movie as discussed above in steps 3-7.

OR

1. Select an individual  $z$  plane for the movie by scrolling to the desired  $z$  plane and selecting the thumbnail button.
2. You may now generate a series movie as discussed above in steps 3-7.

### 8.6 Exporting Views

Views may be exported as a variety of different file types. These images will be 8-bit for each color and should not be used for intensity analysis. However, an exported view may be loaded into Adobe Photoshop (or any other illustration program) or inserted directly into a word processor document.

**NOTE:** If you would like to use the image in another analysis program for purposes of deriving intensity information, you will need to export the image as described in [Exporting Images](#) on page 221.

**NOTE:** If you would like to show annotations such as scale bars, time stamps, etc, you should export the image from the **View** menu.

### 8.6.1 Exporting a Main View, Three View, Channel View, or Tile View as a TIFF

1. Open a Main View, Three View, Tile View, or Channel View of the image and adjust the look of the data as desired. This may include altering the color display and renormalization parameters or cropping the image.
2. Select **View > Export > TIFF** to export the view as a single TIFF (.tif) file.
3. Enter the file name and select **Save**. The TIFF image can now be used in other programs.

### 8.6.2 Exporting a 3D or 2D Timelapse View as a TIFF or TIFF Series

Three-dimensional Main Views may be exported as a single TIFF file containing multiple-planes or as a series of TIFF images.

1. Open a **Main View** of the image and adjust the look of the data as desired.
2. Select **View > Export > TIFF** to export the view as a single TIFF (.tif) file or **View > Export > TIFF Series** to export the view as a series of TIFF images (one image for each  $z$  or  $t$  plane).
3. Enter the file name and select **Save**. The TIFF image(s) can now be used in other programs.

### 8.6.3 Exporting Default Views of All Images in a Slide

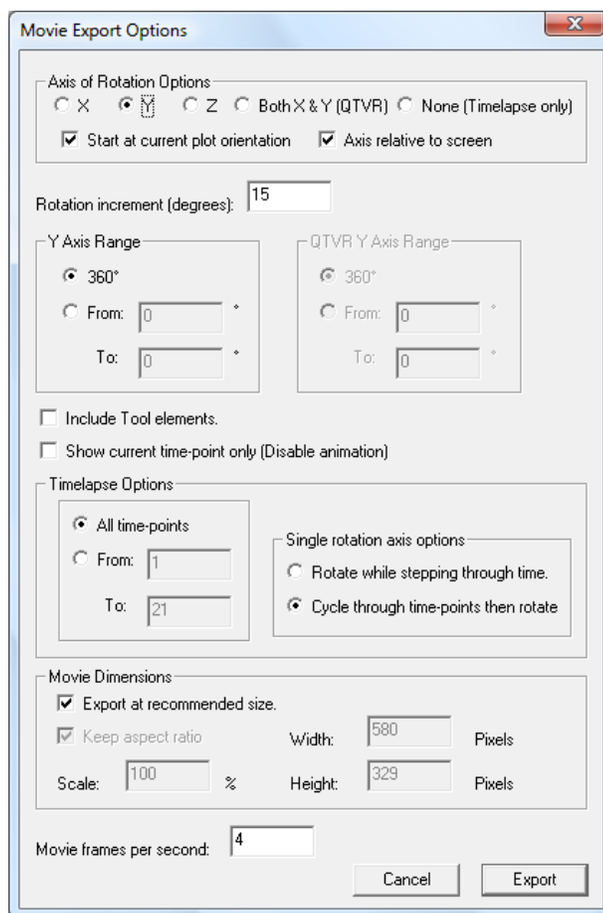
SlideBook allows you to use a single command to export a series of tiff files, one for each image in the slide. Each TIFF file will be a single plane that has the same look as the thumbnail image (default view). To use this feature:

1. Alter each image to the desired look as described in this chapter. This may include altering the color display and renormalization parameters or cropping the image.
2. Select **View > Export > Default Views of All Images as TIFFs**.
3. Choose a directory for the files and select **OK**. The TIFF images will be saved under the individual image names in the chosen directory and can be used in other programs.

## 8.6.4 Exporting Volume View and Surface View Movies

You may wish to export your volume or surface views so that they may be displayed in a presentation or online publication. You may export either as a QuickTime movie, or as a QTVR (QuickTime Volume Rendering) file. QTVR files can be rotated at arbitrary angles in QuickTime (available at [www.apple.com](http://www.apple.com)). To export your Volume or Surface View:

1. In the Slide View, click on an open Volume or Surface View to make it the active view.
2. Select **View > Export > Movie**. The following dialog box will appear:



The dialog has the following sections:

- **Axis of Rotation Options** – you may generate either QuickTime movies where your volume or surface rotates around a single axis (*x*, *y*, or *z*) or QTVR movies where you may rotate your volume as desired using QuickTime.
  - **Start at current plot orientation** – check this box if you want to start your movie with the view that is currently displayed. If left unchecked, the movie will begin with a view with surfaces at right angles to the selected axis.

- **Axis relative to screen** – check this box if you'd like the image to rotate around an axis relative to the screen. If left unchecked, the image will rotate around the image axis.
  - **Rotation Increment** - enter the desired rotation increment in the data field. The rotation range that you selected is computed in the increments you provide. Choosing a smaller increment increases the smoothness of the rotation and adds to the depth cue provided by moving the image. Projection is an expensive computation, so choosing a larger increment will make the projection process faster. To generate a single head-on projection of the entire image, choose 360.
  - **Rotation Range** – select 360 to rotate the data completely. You may select a smaller range if desired.
  - **Show current timepoint only (disable animation)** – available for 4D images only. Allows you to render a single timepoint to conserve time and memory.
  - **Timelapse Options** – select the timepoints you would like to render and how you would like your timelapse movie to proceed (either step through time as the object is rotating, or complete a rotation before proceeding to the next timepoint).
  - **Movie Dimensions** – allows you to change the dimensions of the movie
    - **Keep Aspect Ratio** – keeps the current height/width ratio
    - **Scale** – change the size of the movie with respect to the original image dimensions
    - **Width/Height** – manually enter the dimension you wish to export
  - **Movie frames per second** – adjust the speed of your movie
3. Once you have selected the appropriate parameters, select **Export**. A standard “Save As...” dialog box will appear.
  8. Type in a filename and then click **Save** and you are finished.

## 9 Preparing an Image for Analysis or Export

Before performing statistical analysis on your images, you may wish to perform a variety of pre-processing operations. In this chapter, you will learn how to:

- Manipulate Individual Channels
- Crop an Image
- Align an Image
- Manipulate Timelapse Data
- Perform Flat Field Correction
- Perform Photobleach Correction
- Perform Background Subtraction
- Create a Projection Image
- Create an Interpolated Image
- Apply Filters
- Perform Deconvolution

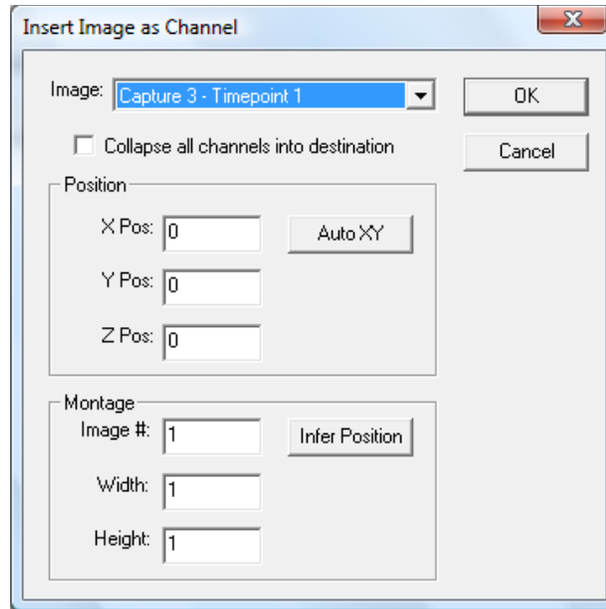
### 9.1 Manipulating Individual Channels

A *channel* denotes the image data collected for a specific fluorophore. SlideBook allows you to manipulate channels prior to performing quantitative analysis on your images. You may insert a channel from one image into another image, remove a channel, or create new channels using channel math. For timelapse images, you may also create a timelapse composite image. The timelapse composite channel adds the previous timepoints to each timepoint. Thus, timepoint 3 in a timelapse composite channel is the sum of timepoints 1, 2, and 3.

#### 9.1.1 Inserting an Image as a Channel

You may wish to insert an image containing a single channel, such as a background image, into another image that you have collected. To do this:

1. Make sure that the image you would like to insert is in the same slide as the destination image (see [Copying an Image](#) on page 159).
2. Select the destination image (the image that will contain the newly inserted image) by clicking on it. You may also start from an open view of the image.
3. Select **Image > Channel Operations > Insert Channel**. The following dialog box will appear:

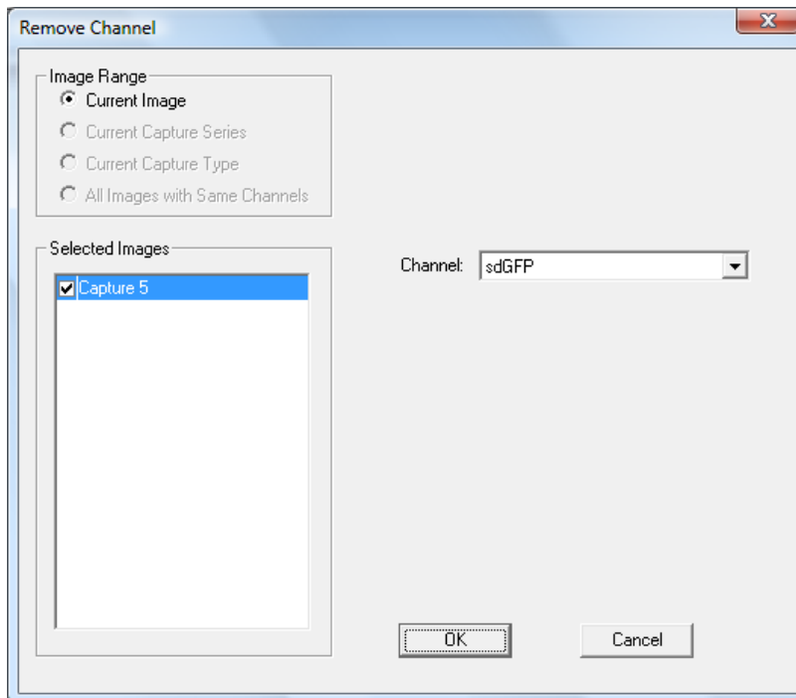


4. Select the image that you would like to insert as a channel from the dropdown menu. If the image that you select contains more than one channel, the channel that is listed as channel 1 (see [Getting and Editing Image Information After Capture](#) on page 101) will be the channel that is inserted.
5. Select the  $x$ ,  $y$  and  $z$  coordinate positions for insertion in the **X Pos**, **Y Pos**, and **Z Pos** edit fields. If left at 0, 0, 0, the single channel image will be inserted in the upper right hand corner of the image in the first plane (if the destination image has multiple planes). Images that are the same size will overlay exactly.
6. Click **OK** to insert the channel. The new channel can now be selected from the channel menus in the info/tool bar and displayed in any view. The channel may also be selected when performing channel math (see [Using Channel Math](#) on page 202).

### 9.1.2 Removing a Channel

You may also wish to remove a channel that was either added improperly or does not contain useful data. To do so:

1. Click on the image that contains the channel you would like to remove. You may also start from an open view of the image.
2. Select **Image > Channel Operations > Remove Channel**. The following dialog box will appear:



The dialog box has the following options for **Image Range**:

- **Current Image** – Operation will be performed on the selected image only.
  - **Current Capture Type** – Operation will be performed on all images in the slide with the same capture type (2D, 3D, 4D, etc.) as the selected image.
  - **All Images with Same Channels** – Operation will be performed on all images in the slide with the same channels as the selected image.
3. Select the channel that you would like to remove from the dropdown menu.
  4. Select the image extent for which you would like to remove channels. You may select and deselect images using the checkboxes in the **Selected Images** list.
  5. Click **OK** to remove the channel(s).

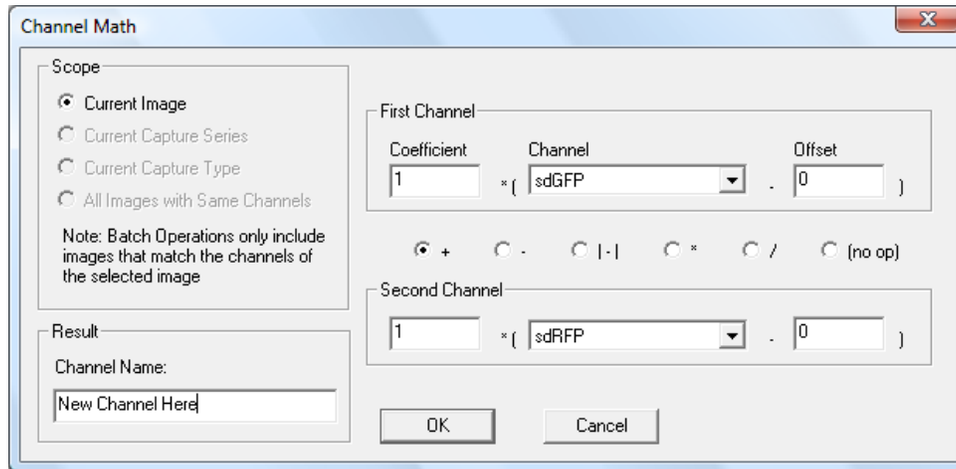
**NOTE:** Removing a channel is an irreversible process. You may wish to keep a copy of the original image (or entire slide).

### 9.1.3 Using Channel Math

SlideBook allows basic mathematical operations to be performed on specified channels in an image. This is appropriate when doing background subtraction or when calculating bleedthrough.

To perform channel math, do the following:

1. Make sure that the channels that are to be used in the mathematical operation are present in a single image. You may wish to insert a channel as discussed above.
2. Select the image by clicking on it. You may also start from an open view.
3. Select **Image > Channel Operations > Channel Math**. The following dialog box will appear:

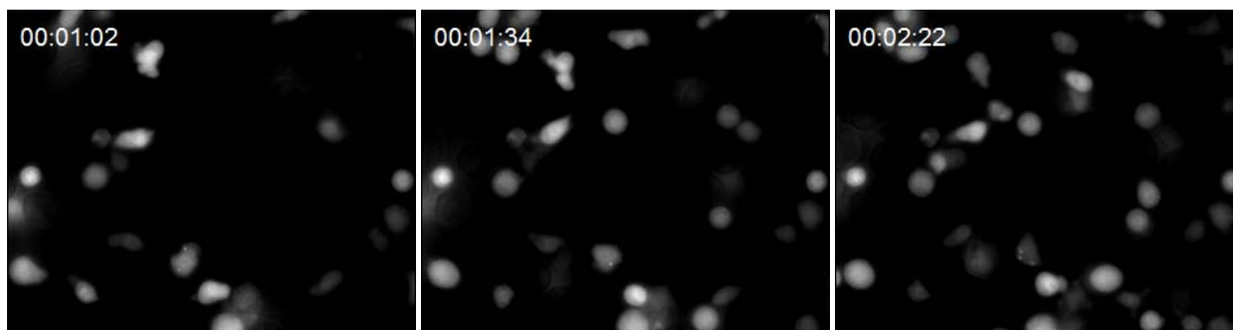


4. Setting the scope section of the Channel Math window allows channel math operations to be applied to multiple images in the slide. Select one of the following options:
  - **Current Image** – Operation will be performed on the selected image only.
  - **Current Capture Type** – Operation will be performed on all images in the slide with the same capture type (2D, 3D, 4D, etc.) as the selected image.
  - **All Images with Same Channels** – Operation will be performed on all images in the slide with the same channels as the selected image.
5. Select the channels that will be used for the mathematical operations from the dropdown list. For instance, if you would like to subtract the FITC channel from the CY3 channel, you would select the FITC channel from the second channel menu, and the CY3 from the first channel menu.
6. Select the mathematical operation that you would like to perform using the radio buttons. From left to right, the operations are: addition, subtraction, absolute difference, division, and no operation. If you select (**no op**), the second channel selection will become unavailable.
7. Enter any desired **Coefficients** or **Offsets** in the edit fields. The coefficient will be multiplied by the selected channel, and the offset will be subtracted.

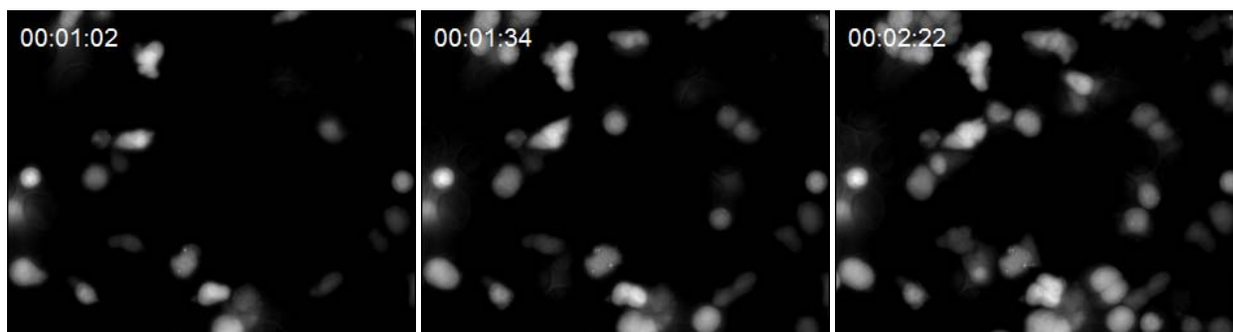
8. Enter a **Channel Name** in the edit field and select **OK** to generate the new channel. This new channel will now appear in the channel selection drop-down box in the image views.

### 9.1.4 Creating a Timelapse Composite Channel

To create a timelapse composite channel, select **Image > Channel Operations > Create Timelapse Composite Channel**. Select the desired channel from the dropdown channel and then **OK**. An example of a timelapse composite channel is shown below. The timelapse composite channel shows the image data as accumulation over time. This is useful if you wish to see the “path” of travel for migrating cells or other objects of interest.



Original Data



Timelapse Composite Channel

## 9.2 Cropping an Image

Before analyzing an image, you may wish to crop an image. This is especially important to increase the speed of computationally intensive processes such as deconvolution. Please see [Cropping an Image](#) on page 177.

## 9.3 Aligning an Image

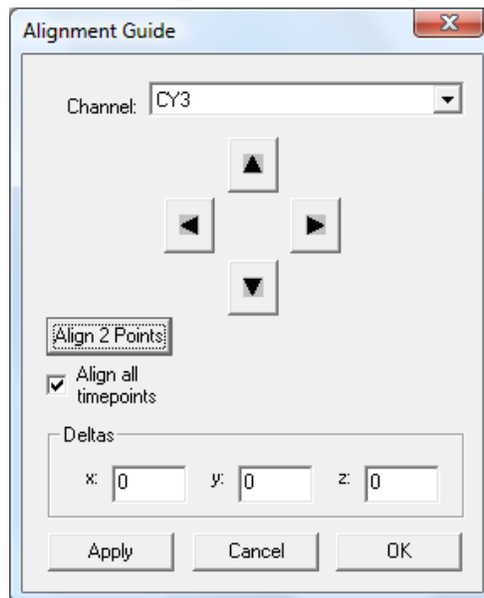
Sometimes, due to misalignments of filters or lack of mechanical repeatability, data for one channel of an image may be slightly offset from data for another channel. To solve this problem, SlideBook offers an alignment guide that lets you adjust relative positions of any channel in the  $x$ ,  $y$ , and  $z$  axes. The alignment guide works alongside a Main View, a Three

View, or a Tile View (see [Using Data Views to Display Images](#) on page 160) in order to shift how each channel lies in the image with respect to the other channels.

Changing channel alignment for an image makes a permanent change to the image itself and will be saved with the image. In other words, every view that you open from that point on will reflect the new alignment. Of course you can re-adjust the alignment later if you wish.

To align an image:

1. Select the image in the Slide View by clicking on it.
2. Open a Main View, Three View, or a Tile View of an image (see [Using Data Views to Display Images](#) on page 160)
3. Select **Image > Align Channels**. The following dialog box will appear.



4. Select the channel that you wish to align from the dropdown menu.
5. Use the left and right arrows to move the selected channel in the  $x$  direction, and the up and down arrows to move in the  $y$  direction, or enter an offset for any axis directly into the corresponding edit controls.

**NOTE:** The view will update automatically when you use the arrow buttons, but you must press the **Apply** button if you manually enter the offsets. Clicking **Cancel** will ignore any changes to the edit field that were not previously applied.

6. Click **OK** when you are satisfied with the image alignment.

**NOTE:** If you find it necessary to align certain filters consistently, your system may need repair. Please contact Olympus America ([slidebooksupport@olympus.com](mailto:slidebooksupport@olympus.com) or 1-800-446-5967 option 4).

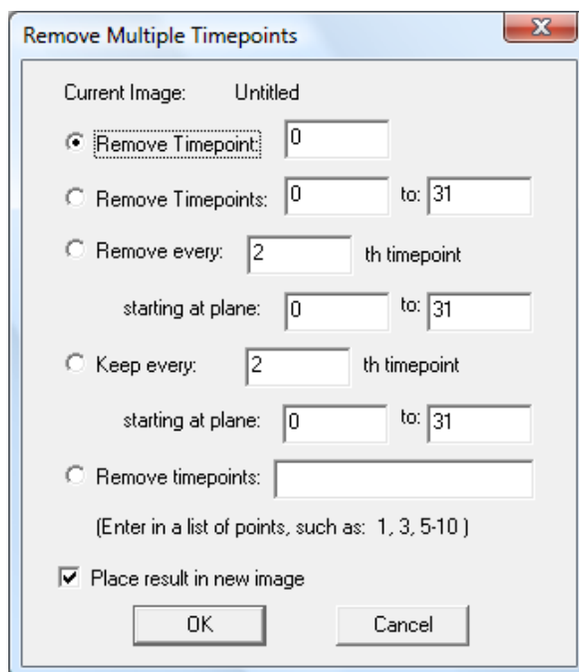
## 9.4 Manipulating Timelapse Series

SlideBook allows you to remove timepoints from a timelapse series or merge two timelapse series into one series.

### 9.4.1 Removing Timepoints from a Timelapse Series

To remove timepoints from an image:

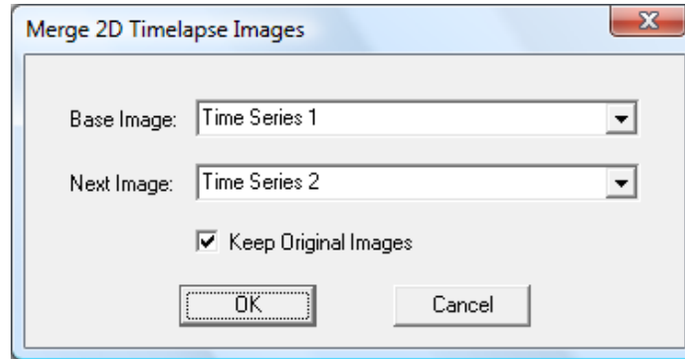
1. Click on an image's thumbnail in the Slide View, or have an open view and select **Image > Advanced Operations > Remove Multiple Timelapse Planes**. The following dialog box will appear:



2. Enter the timepoint (or group of timepoints) that you wish to remove. You may also choose to subsample the image by selecting **Remove every *n*th timepoint** or **Keep every *n*th timepoint**. Alternately, you may enter a list of points to be removed. If you display timestamps, you will see that the timepoints have been removed.
3. Select the checkbox **Place result in new image** if you choose to do so, then select **OK**. If you do not choose to do so, the timepoints will be irreversibly removed from the current image.

### 9.4.2 Merging Timelapse Series

If you have an experiment that was captured into multiple timelapse images, you may merge these images by selecting **Image > Advanced Operations > Merge Timelapse Images**. The following dialog box will appear:



In the example dialog above, “Time Series 2” will be appended onto “Time Series 1” and placed into a new image. The original images will remain unchanged.

## 9.5 Performing Flat Field Correction

As with any optical system, specks of dust or pieces of lint will find their way into the microscope’s optical path. There may also be some variation in the level of illumination across the field of capture. SlideBook offers an operation called flat field correction, which eliminates many of these constant artifacts from an image. Flat field correction is particularly useful in brightfield and DIC (Nomarski) imaging, where such artifacts are usually more evident than in fluorescence imaging. Flat field correction (or shading correction) is not the same as background correction (background subtraction is strictly subtraction). When flat field correction is applied, the desired image is multiplied by a matrix of coefficients that is determined from a series of flat field images (see [Collecting Flat Fields](#) below). To generate this matrix, flat field images are collected, and the average of each pixel for the set of images is calculated to generate a single averaged image (flat field image). The average intensity of all the pixels in the resulting averaged image is then calculated. Finally, a coefficient matrix (pixel by pixel map) is calculated so that when the intensity at any pixel location is multiplied by its coefficient, the generated pixel intensity matches the average pixel intensity of the flat field image. If you are planning to perform both background subtraction and flat field correction, perform flat field correction first.

### 9.5.1 Collecting Flat Fields

In order to perform flat field correction, SlideBook needs to know what an image with no sample in focus looks like. This image should contain all the constant artifacts, but little that will change from sample to sample. From this image, SlideBook can compute a coefficient matrix that can be applied to an image to eliminate most of its constant artifacts. SlideBook, in fact, can average several “empty images” to better ensure that the flat field contains only constant artifacts of the optical system and not some peculiarity of a single “empty image.”

Since constant artifacts might exist anywhere along the optical path, a flat field matrix is necessary for each combination of objective, filter, and magnification changer setting. To

keep things simple, SlideBook stores all collected flat fields in a single database and keeps track of the optical parameters for each one.

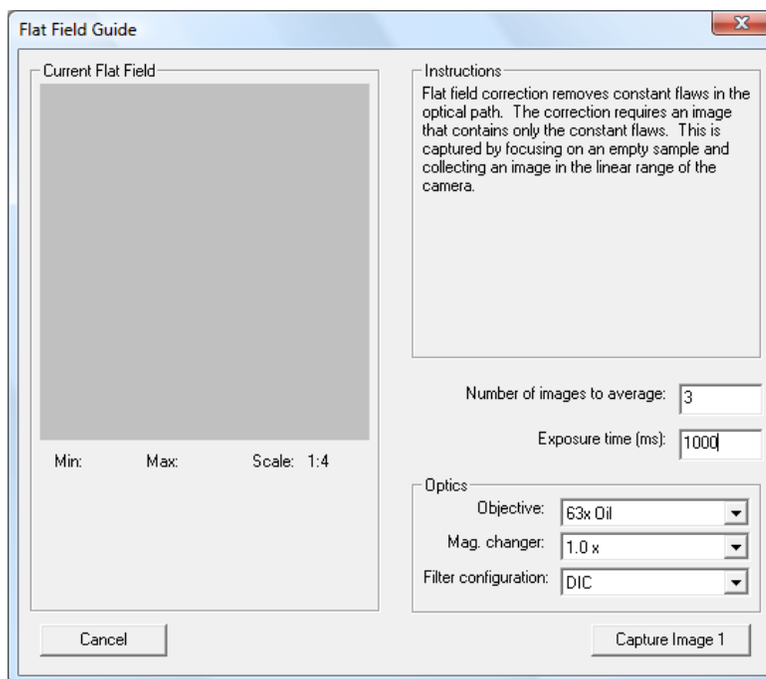
### 9.5.1.1 Adding or Replacing a Flat Field

If you want to add a flat field matrix to the database (or replace an existing entry), you will need to capture a series of images on an empty sample using the desired set of optics, including the filter, objective, and magnification changer. To add or replace a flat field, do the following:

1. Place a blank slide containing no sample (or a uniformly fluorescent slide if collecting a fluorescent flat field) on the stage.
2. Bring the desired optics into place as described in Chapter 5 - [Using the Focus Window](#).

In order to bring the  $z$  stage of the microscope to the correct position, focus on any irregularities (e.g., dust or scratches) on the blank slide. Then move the  $xy$  position of the slide to a location where there are few irregularities and adjust the microscope so it is slightly out of focus. This will minimize the contribution of irregularities with the slide to the flat field images.

3. Select **Edit > Setup Guides > Flat Field Guide** to bring up the flat field guide. The following dialog will appear:



4. Specify an exposure time and number of images to take.

Later, you will be able to adjust the exposure time so that the linear range of the camera is being used. The number of images required depends on the user: the

more images, the more accurately the flat field matrix will reflect only constant artifacts, though the collection process will take more time. Typically three will be sufficient.

5. Select **Capture Image 1**. SlideBook will capture an image and display it in the upper left corner of the guide.

Adjust the exposure time such that the maximum intensity shown in the histogram is 3000-3500 for 12 bit CCDs (on a 0-4095 scale) or 48000-56000 for 16 bit CCDs (on a 0-65535 scale)

6. If necessary, make an appropriate adjustment to the exposure time and click **Recapture Image 1**. Otherwise, move the *xy* position of the slide slightly and capture the next image by clicking **Capture Image 2**.

After the first image, it's unlikely you'll need to recapture any of the subsequent images, but you will still have that option.

7. Acquire the remaining images making sure to move the slide slightly between capture.
8. Once you've captured all the images, click on the **Finish** button to complete the process.
9. Repeat the process for other optical combinations.

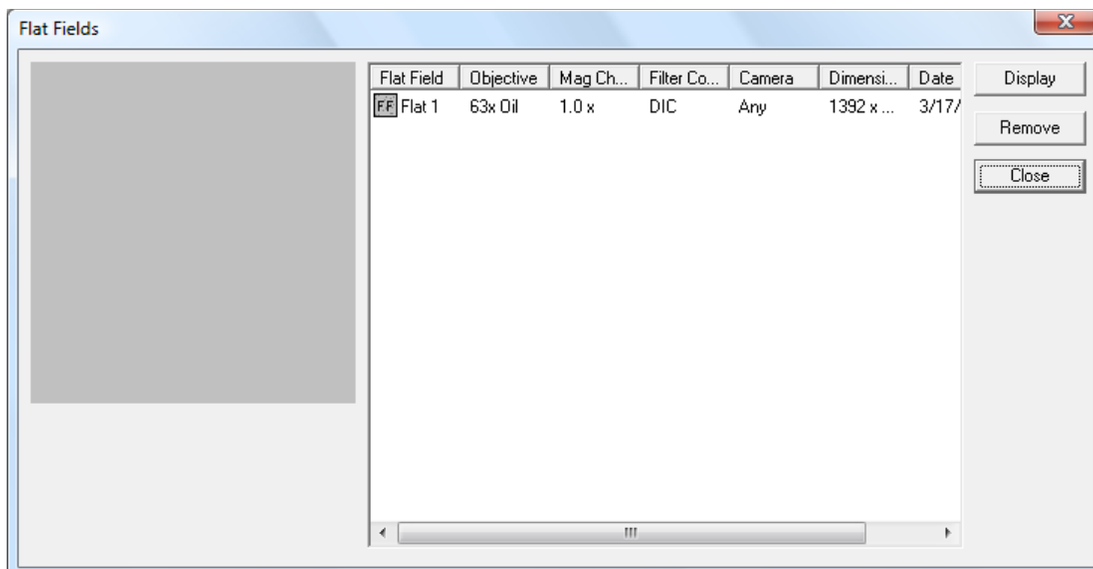
### 9.5.1.2 When to Update the Flat Field Database

If you try to add a flat field matrix for optics where one is already defined, SlideBook will alert you and ask if you want to replace the existing one. SlideBook does not keep multiple flat field matrices for identical optics.

As time passes, it is possible for a flat field matrix to become outdated. Specks may have changed position, or other constant artifacts have either appeared or disappeared. A telltale sign that it is time to update a flat field is when performing the flat field operation no longer does its job—marked by white specks appearing after applying the flat field.

### 9.5.1.3 Displaying Existing Flat Fields

You can get a list of the existing flat fields (and their corresponding optical combinations) by selecting **Edit > Setup Guides > Display Flat Fields**. To display the flat field, click on the desired flat field icon in the table and press the **Display** button.

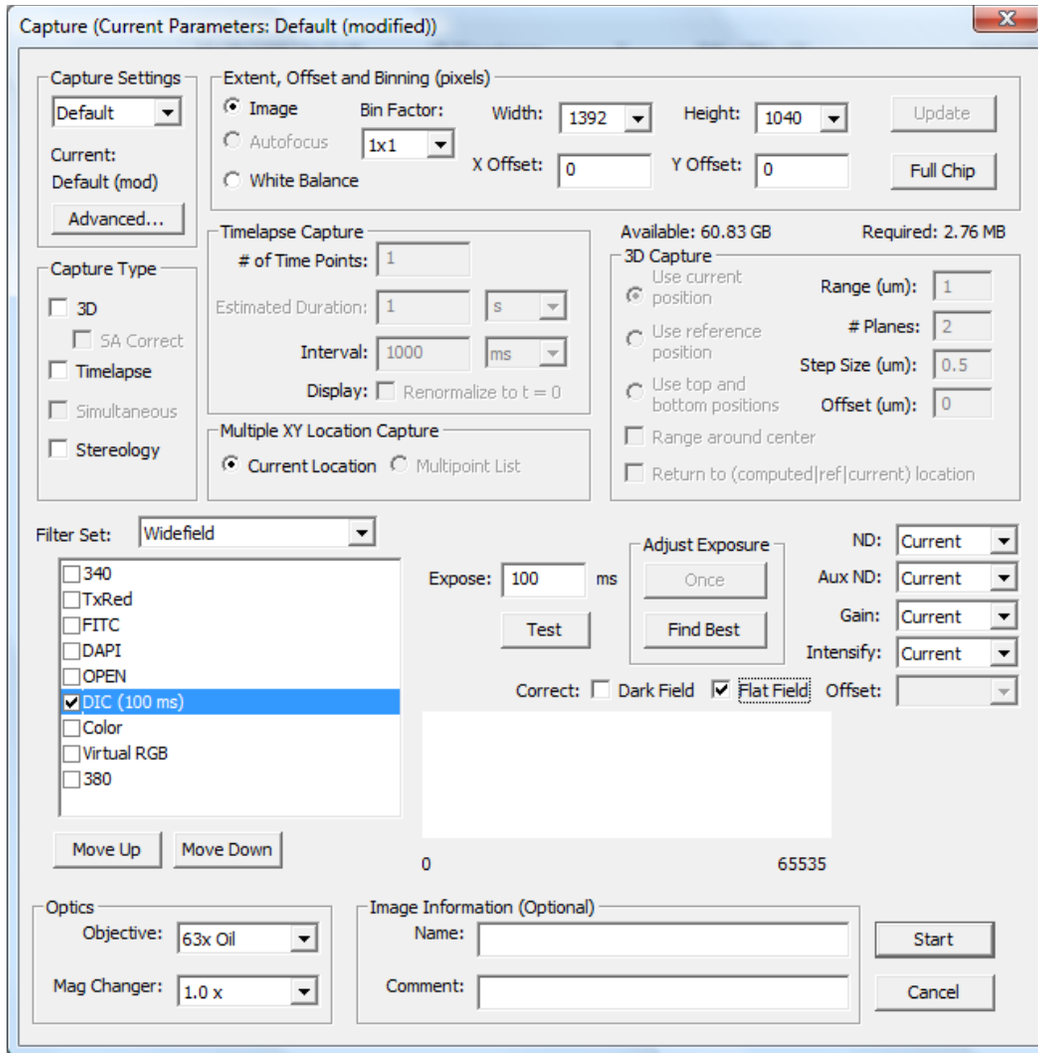


## 9.5.2 Applying Flat Fields

If you wish to flat field correct an image that was captured using an optical combination that has a corresponding entry in the flat field database, you may choose to apply the flat field correction during or after image capture.

### 9.5.2.1 Applying Flat Field Correction During Image Capture

1. Bring a sample into view and focus as discussed in [Using the Focus Window](#) on page 85.
2. Open the Capture dialog box by selecting **Image > Capture** or by selecting the Capture icon in the SlideBook toolbar .
3. Set all other capture parameters as discussed in Chapters 6 and 7.
4. Select the Flat Field Correction checkbox, which is located below the exposure information.



5. Click **OK** to begin capture. The resulting images will be flat field corrected.

### 9.5.2.2 Applying Flat Field Correction after Image Capture

1. Select the image that you would like to correct by clicking on it. You may wish to copy the image before applying the flat field, as it is an irreversible operation (see [Copying an Image](#) on page 159). You may also start from an open view of the image.
2. Select **Image > Apply Flat Field**. The flat field correction will be applied to the image.

**NOTE:** If you do not have an entry in the flat field database that match the optics used to capture an image, SlideBook will alert you that it is unable to perform a flat field correction.

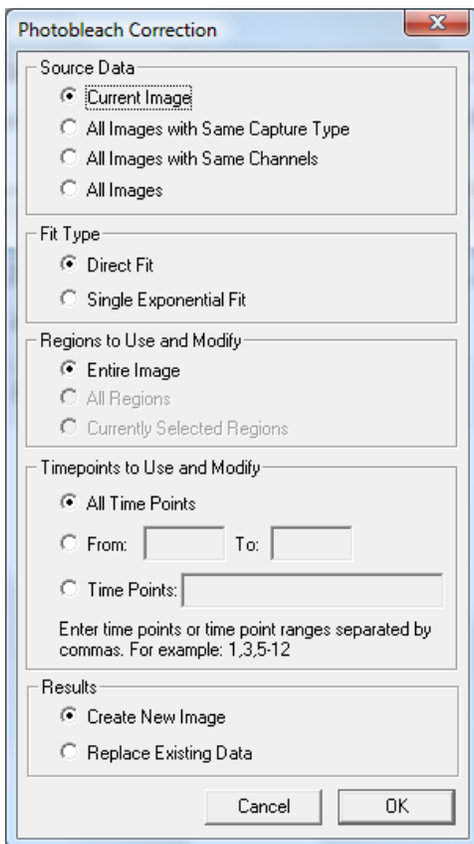
## 9.6 Performing Photobleach Correction

Photobleach correction can be used to correct for photobleaching in 2D timelapse images. The photobleach correction operation takes the mean intensity at each timepoint and fits a line or curve through the mean intensity versus time data. Then, it determines a multiplication factor for each timepoint that is used to adjust the intensities at each pixel, resulting in constant intensities over time.

Before performing photobleach correction, you should examine your data to determine what type of fit is most appropriate. To do so, you need to view the intensity-time profile of your data set. You can do this by drawing a box over the entire field for a single plane, then selecting **Image > Statistics > Ratio/Timelapse Data**. If the intensities decrease linearly over time, you may perform a direct fit. If they seem to decrease exponentially, choose exponential fit.

To perform photobleach correction:

1. Select **Image > Photobleach Correction**. The following dialog will appear:



2. Select the fit, and the scope of operation, as described below:

- **Current Image** – Operation will be performed on the selected image only.

- **All Images with Same Capture Type** – Operation will be performed on all images in the slide with the same capture type (2D, 3D, 4D, etc.) as the selected image.
  - **All Images with Same Channels** – Operation will be performed on all images in the slide with the same channels as the selected image.
  - **All Images** – Operation will be performed on all images in the slide.
3. Select the timepoints to modify and whether or not you would like to overwrite your data.
  4. Select **OK** to perform the operation.

**WARNING:** Photobleach correction is an irreversible operation. Please be sure to save a copy of your original data.

### 9.7 Performing Background Subtraction

You may also select a region of interest of arbitrary size and subtract the background mean intensities from your image using channel math. Please see the example in Chapter 2, [Creating Masks Manually \(Background Subtraction Example\)](#) on page 41. You may also permanently subtract a background ROI:

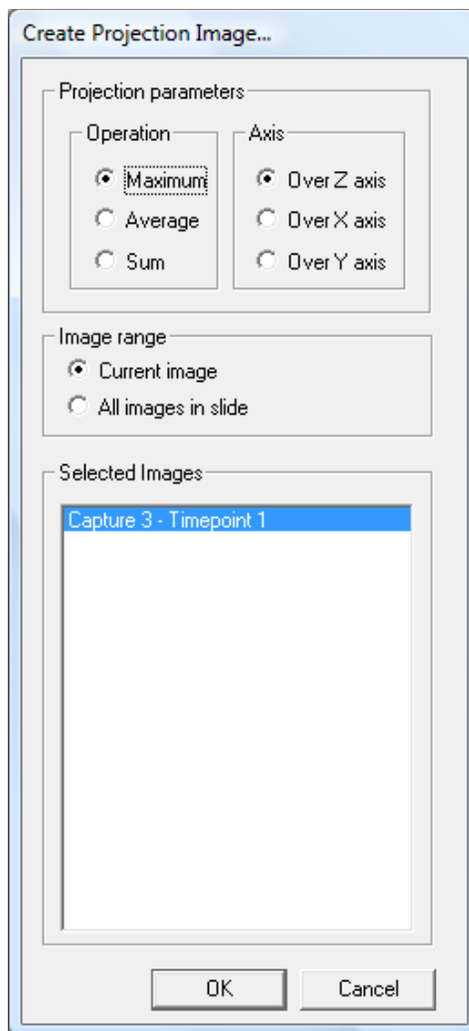
1. Make a copy of the data that you are working with.
2. Use the marquee tool to select a background region in your image.
3. Select **Image > Select Background**.
4. Select **Image > Advanced Operations > Subtract Region Background**. The average values from your background ROI will be subtracted on a channel by channel basis.

**WARNING:** The background subtraction operation described above is an irreversible operation. Please be sure to save a copy of your original data.

### 9.8 Creating a Projection Image

You may wish to view 3D data as a single 2D projection image. To do so:

1. Make the desired image the active image by clicking on it in the slide view. You may also start from an open view of the image.
2. Select **Image > Create Projection Image**. The following dialog box will appear:



3. Select the **Operation** as described below:
  - **Maximum** – The brightest pixel throughout the chosen axis will be displayed.
  - **Average** – An average of all the pixels through the chosen axis will be displayed.
  - **Sum** – The sum intensity of all the pixels through the chosen axis will be displayed.
4. Select the **Axis** you would like to compress to create the projection image.
5. Select the **Image Range** as described below. You may also select and deselect images using the checkboxes in the **Selected Images** list.
  - **Current Image** – Operation will be performed on the selected image only.

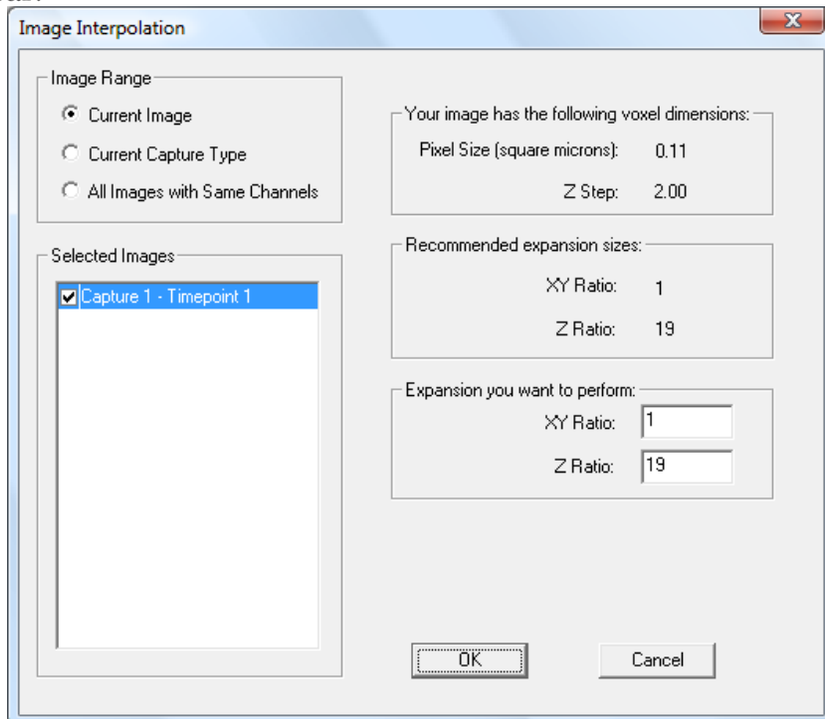
- **All Images in Slide** – Operation will be performed on all images in the slide.
6. Click **OK**. The projection images will be added to the bottom of the slide. The type of projection will be displayed in the **Comments** section of the slide view.

### 9.9 Creating an Interpolated (Isotropic) Image

You may notice that your object may appear disproportionately small, or “squashed” in the *z* direction (especially when using the **Three View**). When performing 3D imaging, each plane that is captured is represented by a set of pixels in the image set. Each pixel is a perfect cube. You will notice that the *z* dimension in the 3D image set appears to be different than what was actually captured. This is because the *z* step size that was selected during capture is not actually the same as the length of the *z* dimension of the pixel. In the image set, planes are simply stacked on top of one another with no regard for the actual *z* distance traveled (this is done in order to preserve data fidelity). Generating an interpolated image will expand your *z* dimension to represent the actual *z* distance traveled by the microscope. Creating interpolated images can also help smooth the *z*-axis when less than optimal *z*-step size is used.

To create an interpolated image:

1. Select the image that you would like to expand by clicking on it. You may also start from an open view of the image.
2. Select **Image > Create Interpolated Image**. The following dialog box will appear:



The dialog box recommends expansion parameters based on the calibrated pixel size and z step size used for capture. These recommended parameters will generate an image with the approximate dimensions of the actual sample. You can use the recommended expansion parameters or enter custom values in the edit boxes.

3. Select the **Image Range** as described below. You may also select and deselect images using the checkboxes in the **Selected Images** list.
  - **Current Image** – Operation will be performed on the selected image only.
  - **Current Capture Type** – Operation will be performed on all images in the slide with the same capture type (2D, 3D, 4D, etc.) as the selected image.
  - **All Images with Same Channels** – Operation will be performed on all images in the slide with the same channels as the selected image.
4. Select **OK** and a new image(s) will be generated.

**NOTE:** The  $\mu\text{m}/\text{pixel}$  value for the objective that was used must be accurate for this feature to work properly. Before generating the isotropic image, check the image information to make sure the  $\mu\text{m}/\text{pixel}$  value is set properly (see [Getting and Editing Image Information After Capture](#) on page 101).

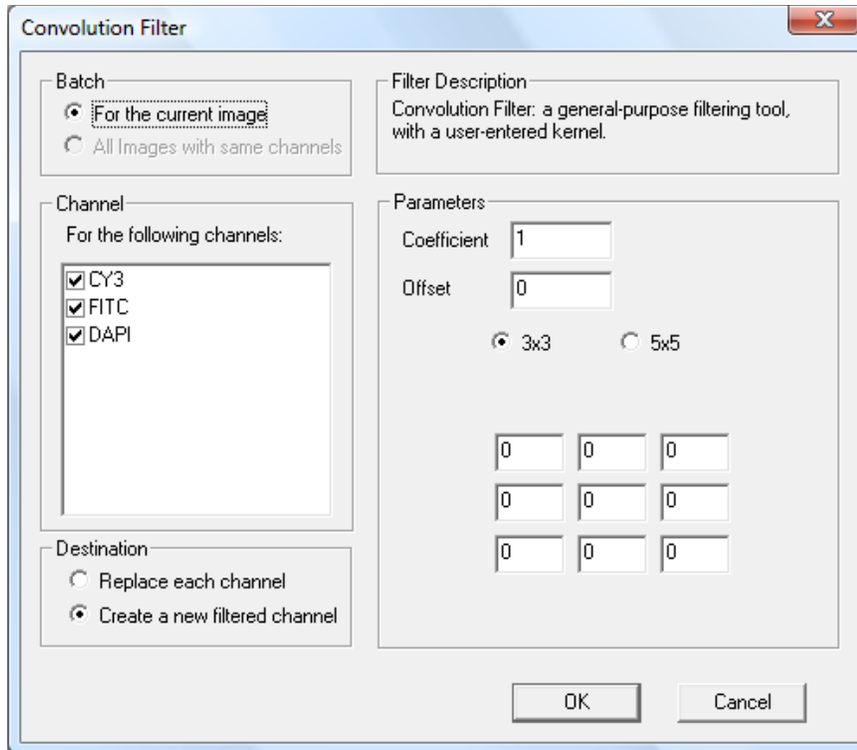
### 9.10 Applying Filters

SlideBook offers a variety of filtering options, including a user-defined kernel and semi-defined Gaussian, Gaussian derivative, hot pixel, median, and mean filters. These filters perform local, rather than global, calculations on a pixel by pixel basis. In other words, each pixel is altered based on the intensities of the neighboring data.

#### 9.10.1 User-Defined Convolution Kernels

This filtering option allows you to define a custom kernel. To apply a custom kernel:

1. Select the image that you would like to correct by clicking on it. You may wish to copy the image before applying the filter, as it is an irreversible operation (see [Copying an Image](#) on page 159 of Chapter 5.). You may also start from an open view of the image.
2. Select **Image > Filter > Convolve**. The following dialog box will appear:



3. Select the **Batch** as to determine if you will apply the filter to the current image or all images with the same channels.
4. Select which channels will be filtered using the convolution kernel by checking the desired channels in the **Channel** list.
5. Select the destination of the filtered channel:
  - **Replace each channel** – The selected channels will be replaced with the filtered data.
  - **Create a new filtered channel** – New channels containing the filtered data will be created within the image.
6. Select either a **3x3** or a **5x5** kernel size.
7. Enter the desired **Coefficient** and **Offset** in the edit field. The output value of the kernel will be multiplied by the coefficient and added to the offset. These parameters can be used to scale the convolved data to a desired range and to account for negative kernel output values.
8. Enter the desired matrix elements in array of edit fields.
9. Select **OK** to perform the convolution.

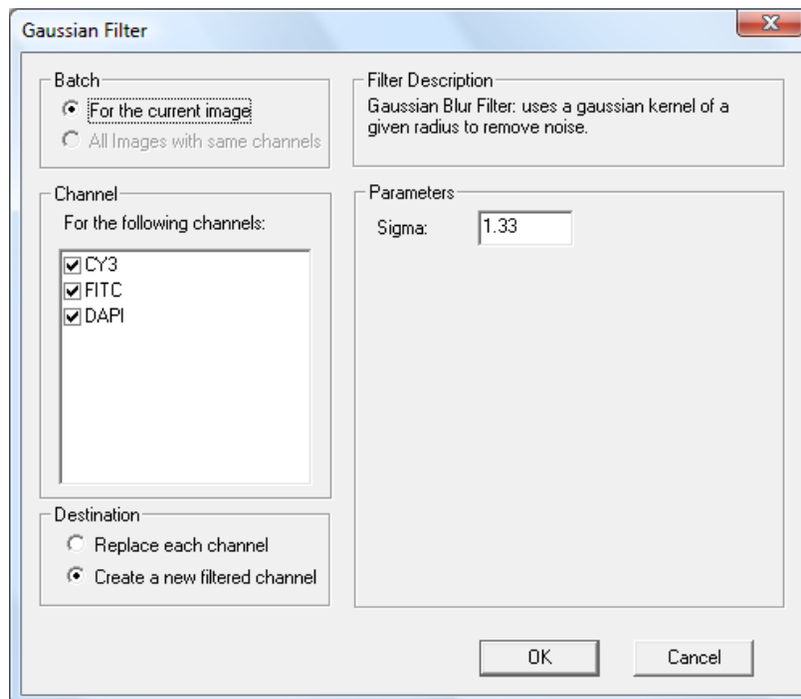
### 9.10.2 Gaussian, Hot Pixel, Mean, and Median Filters

SlideBook offers a set of standard filters for digital image processing.

- **Gaussian** – A blurring filter that utilizes a Gaussian kernel of a given radius. Useful for removing noise from an image.
- **Gaussian Derivative**- An edge detection filter that uses a first derivative Gaussian kernel of a given radius.
- **Mean** or **Median** – Blurring filters that replace the intensity value of a pixel with the mean or median intensity value of pixels within a given radius.
- **Hot Pixel** – A nonlinear filter used to remove “hot pixels” in an image. If a pixels intensity value is some factor greater than the mean of its neighbors it is replaced with this mean value. A hot pixel in an image is generally caused by a bad pixel on a CCD chip.

To apply the Gaussian, Gaussian Derivative, Mean, Median and Hot Pixel Filters:

1. Select the image that you would like to filter by clicking on it in the Slide View. You may wish to copy the image before applying the filter, as this is an irreversible operation (see [Copying an Image](#) on page 159). You may also start from an open view of the image.
2. Select the appropriate filter from the **Image > Filter** hierarchical menu. A dialog box will appear that is similar to that shown below:



3. Select the batch, channel and destination options for the filter as described in the above section [User-Defined Convolution Kernels](#) on page [216](#).
4. Each filter has different user-definable parameters. Enter the parameters for the filter based on the following descriptions.
  - **Gaussian filter** – A Gaussian kernel of a designated **Radius** will be used to determine a pixels intensity value.
  - **Gaussian Derivative** – A first derivative Gaussian kernel of a designated **Radius** will be used to determine a pixels intensity value.
  - **Mean or Median** – A pixel will be replaced with the mean or median intensity value of pixels within a designated **Radius**.
  - **Hot Pixel** – A pixel whose intensity value is higher than the mean of its neighboring pixels by more than the designated **Factor** will be set to the mean of its neighboring pixels.
5. Select **OK** to filter the image.

### 9.11 Performing No Neighbors Deconvolution

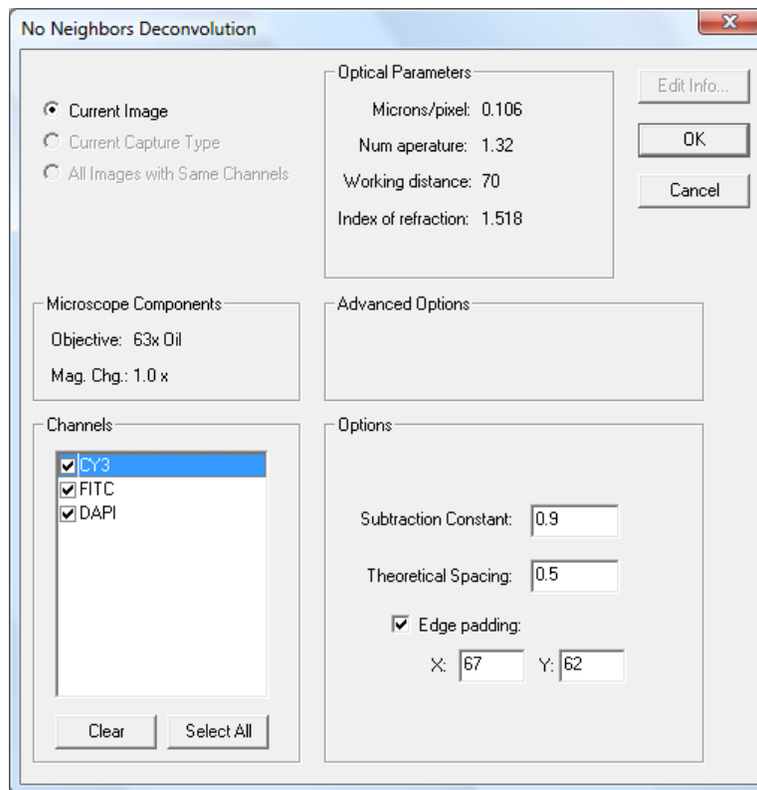
*Deconvolution* is the process by which SlideBook mathematically removes out-of-focus information from a fluorescence image set. SlideBook can implement three deconvolution approaches: *constrained iterative*, *nearest neighbors*, and *no neighbors*. No Neighbors deconvolution is incorporated into the base SlideBook package. Nearest neighbors and constrained iterative deconvolution are available as an additional SlideBook module. For more information about nearest neighbors and constrained iterative deconvolution refer to the supplemental [SlideBook 3D Deconvolution Module](#) manual or contact Olympus America ([slidebooksupport@olympus.com](mailto:slidebooksupport@olympus.com) or 1-800-446-5967 option 4).

No neighbors is the least computationally intensive method of deconvolution offered in SlideBook and the only method that can be used on 2D images. While no neighbors deconvolution can be applied to three dimensional data sets, it does not use information across multiple planes to calculate and remove out of focus information. Rather, it functions by creating a theoretical blurred image from the original two-dimensional image. This blurred image is then subtracted from the original image, leaving objects that are more likely to be in focus.

#### 9.11.1 Running No Neighbors Deconvolution

1. Select the image that you would like to correct by clicking on it. You may also start from an open view.
2. Make sure that the image information is correct as described in [Getting and Editing Image Information After Capture](#) on page 101.

3. Select **Image > Deconvolve > No Neighbors**. The following dialog box will appear:



4. Select the scope of the deconvolution operation.
  - **Current Image** – Operation will be performed on the selected image only.
  - **Current Capture Type** – Operation will be performed on all images in the slide with the same capture type (2D, 3D, 4D, etc.) as the selected image.
  - **All Images with Same Channels** – Operation will be performed on all images in the slide with the same channels as the selected image.
5. Select the Channels that will be deconvolved by toggling the checkbox next to the channel name.
6. Select the deconvolution **Options** according to the following descriptions:
  - **Subtraction Constant** – Adjusts the amount of intensity subtraction used in the deconvolution algorithm. Increasing the subtraction constant will increase the amount of deblurring but can also result in a loss of image data.

- **Theoretical Spacing** – Adjusts the distance between theoretical image planes used for de-blurring. The closer the theoretical planes, the greater the amount of de-blurring. Make sure to enter a value greater than zero for this parameter.
- **Edge Padding** – creates a space around the border of the image in order to resolve fluorescent point sources that exist outside of the image.

7. Once the parameters are set, select **OK** to begin the deconvolution.

In addition to the user-definable parameters, the No Neighbors Deconvolution dialog box also displays the following information:

- **Microscope Components** – Displays the optical components that were used to take the image.
- **Optical Parameters** - Displays the optical parameters associated with the microscope components used for image capture. To change these parameters refer to Chapter 4 - Configuring Your System on page 54.

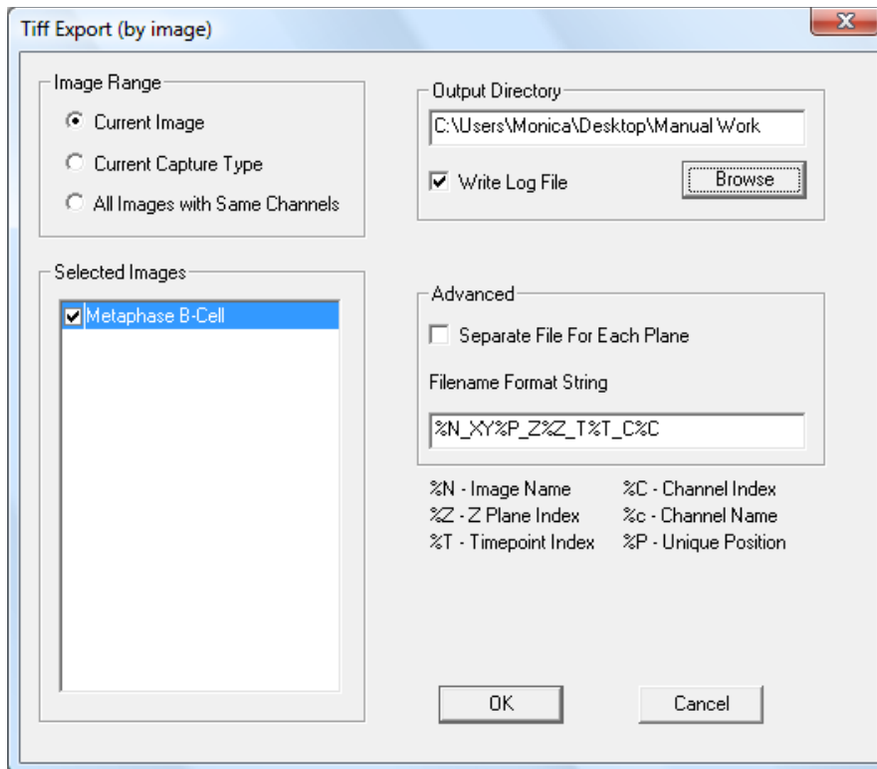
### 9.11.2 Memory Requirements for Deconvolution

Deconvolution is a memory-intensive process and is limited by the size of the image that is being deconvolved. On the PC, any given program can use up to 2 GB of memory. This results in an 800MB size restriction for no neighbors deconvolution.

### 9.12 Exporting Images

If you would like to perform intensity analysis or image processing in a program other than SlideBook, you may want to export the image from the **Image** menu. This will export a 16-bit file that retains all of the intensity data from the original SlideBook image. In contrast, exporting an image from the **View** menu will generate an 8-bit file. This will result in a loss of data assuming the images were captured on a 12-bit or higher digital camera. To export an image from the **Image** menu perform the following steps.

1. Select the image that you would like to export by clicking on it. You may also start from an open view of the image.
2. Select the appropriate file type from the **Image > Export** hierarchical menu. A dialog box similar to the following will appear.



3. Select the **Image Range** as described below. You may also select and deselect images using the checkboxes in the **Selected Images** list.
  - **Current Image** – Operation will be performed on the selected image only.
  - **Current Capture Type** – Operation will be performed on all images in the slide with the same capture type (2D, 3D, 4D, etc.) as the selected image.
  - **All Images with Same Channels** – Operation will be performed on all images in the slide with the same channels as the selected image.
3. Press **OK** and to export a single multi-plane image for each channel in the selected image(s).

## 10 Using Masks for Image Analysis

Masks are the principal means to perform analysis in SlideBook. A mask is a binary overlay on image that can be used to select an arbitrary region (or set of regions) of the image at the same resolution as the image itself. Masks are often referred to as Regions of Interest (ROIs) in other imaging programs. Once you have selected a region of interest(s) using masks, you may generate a variety of morphometric measurements and intensity statistics on those regions, including:

- Morphometrics
  - Area and Volume
  - Center of Mass
  - Estimated perimeter or surface area
  - Length along major axis
  - Center of area
  
- Intensity Statistics
  - Mean
  - Minimum and maximum
  - Integrated intensity
  - Center of intensity
  - Correlation (Colocalization)

*Plus many others....*

In this chapter, you will learn how to:

- Display an Image Histogram and Mean Intensity
- Create Masks and Objects
- Display or Delete Masks
- Perform Boolean Mask Operations
- Use Masks to Perform Smooth Curve Analysis
- Generate and Export Mask and Object Statistics
- Display Graphs

### 10.1 Displaying Image Histogram and Mean Intensity

You may quickly get a statistical snapshot of your image using the **Statistics > Intensity Statistics** function. A dialog box will display a histogram of the intensities of the entire image by channel along with the mean intensity and standard deviation. You must first open a view of your image before selecting the **Intensity Statistics** menu selection.

### 10.2 Creating Masks and Objects

A *mask* is a 2D or 3D set of binary values that has the same extent as the image to which it is assigned. Once created, a mask is permanently stored with its corresponding image. Masks are used for performing advanced selection and analysis of image data. They can either be created and edited manually or generated automatically through threshold segmentation and other techniques. Selecting **Mask > Create** creates a mask with the first

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method; selecting **Mask > Segment** creates a mask with the second. These menu items can also be found under the mask icon  in any **View** window. A single image can contain multiple masks.

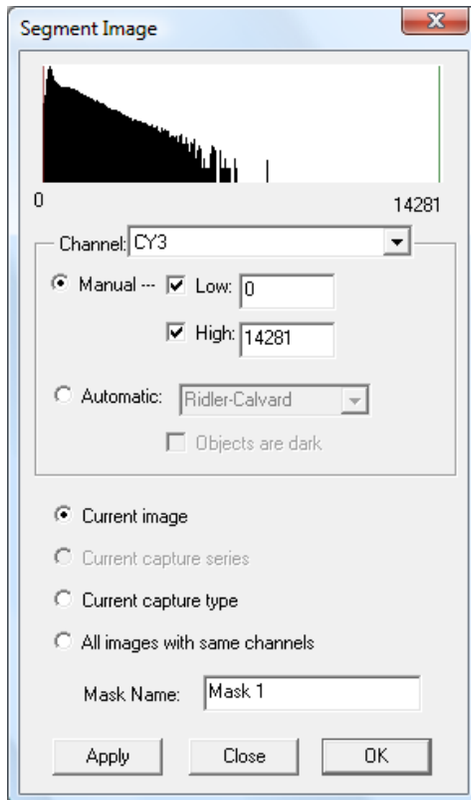
Often you may have several distinct objects that will have intensities that are very similar, and thus will segment together. A mask that contains several distinct entities may be divided into *objects*. For instance, if an image contains a field of many cells, performing threshold segmentation of a nuclear stain such as Hoechst will result in a mask that contains distinct “islands.” Defining objects permits functions such as counting of objects and generating statistics for each subregion.

We will now learn how to create masks using a variety of techniques.

### 10.2.1 Creating a Mask using Threshold Techniques

Threshold techniques allow the user to perform a global selection based on image intensities. SlideBook’s segmentation process generates regions of interest that persist throughout the entire image in all dimensions.

1. Click on an open view to make it the active view.
2. Create a mask by selecting the mask icon  > **Create Segment Mask** or by going to **Mask > Segment**. A dialog box similar to the following will appear. Note that it resembles the **Renormalization** dialog box.



3. Select the channel that you would like to segment from the dropdown menu.
4. Enter a name for your mask.
5. Select the scope of the operation using the radio buttons. For any operation that segments more than one image, you must select **OK** to segment all of the desired images. If you select **Apply**, only the current image will be segmented.
  - **Current Image** – Segments the current image only
  - **Current Capture Type (on OK)** – Segments the current capture type only. For instance, if you have a 2D timelapse image open in a slide with at least one other timelapse image, as well as other capture types, only the 2D timelapse images would be segmented.
  - **All Images with Same Channels (on OK)** – Segments images that have the exact same number and type of channels. For instance, if you wish to segment a FITC channel, and one image contains a FITC channel while another contains both DAPI and FITC, both images will NOT be segmented. Only the active image will be segmented.

**NOTE:** All images in the slide must be of the same size for this operation.

6. Set the thresholds for segmentation. You may do this in one of two ways:
  - a. Move the red and green bars to the right and left by clicking and dragging them. Note that when you release the mouse button, the image will be segmented and the mask will be displayed.

OR

- b. Enter intensity values in the **Low** and **High** edit fields, then click **Apply** to register the change and move the red and green bars.

SlideBook has selected, or *masked*, any pixel that has intensity values that lie between the red and green bars in *all* of the three channels. In other words, segmentation is performing an AND operation across the bounds defined in all channels.

7. If working with a 3D image, scroll through the image as discussed in [Scrolling through the Invisible Axis in a 3D or Timelapse Image](#) on page 180 and [Scrolling through a \*\*Three View\*\*](#) on page 187. Note that the mask persists throughout the entire 3D extent of image.
8. If you are satisfied with your mask, click on **OK**. If you are not repeat step 6.

**NOTE:** Once you have created a mask using threshold techniques, you may use editing tools to add or remove pixels in the mask (see [Editing the Mask](#) on page 227).

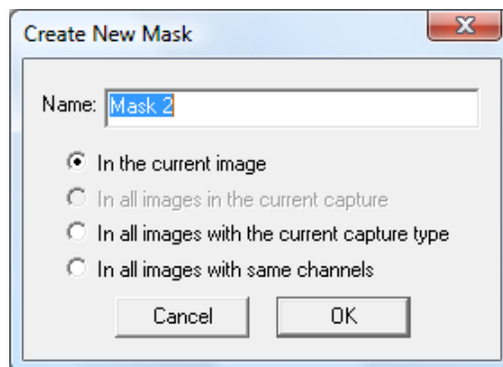
## 10.2.2 Creating Masks Manually

SlideBook also offers a variety of tools that can be used to manually select regions of interest. An example demonstrating the creation and use of a manual mask is described in the Quick Tour section [Creating Masks Manually \(Background Subtraction Example\)](#) on page 41.

### 10.2.2.1 Creating an Empty Mask

To make a mask manually, you will first create an empty mask, then you will select your regions of interest using editing tools.

1. Click on an open view to make it the active view.
2. Create a mask by selecting the mask icon  > **Create Empty Mask** or by going to **Mask > Create**. The following dialog box will appear:



3. Enter a name for the mask in the edit field.
4. Choose to create the mask for the current image or for multiple images using the radio buttons, which are defined as follows:
  - **In the current image** – Creates mask for the current image only.
  - **In all images with the current capture type** – Creates mask for the current capture type only. For instance, if you have a 2D timelapse image open in a slide with at least one other timelapse image, as well as other capture types, only the 2D timelapse images would be segmented.
  - **In all images with same channels** - Creates mask for images that have the exact same number and type of channels. For instance, if you wish to segment a FITC channel, and one image contains a FITC channel while another contains both DAPI and FITC, both images will NOT have a newly created mask. A mask will only be created for the active image.

5. Click **OK** to create a new mask for the current image.

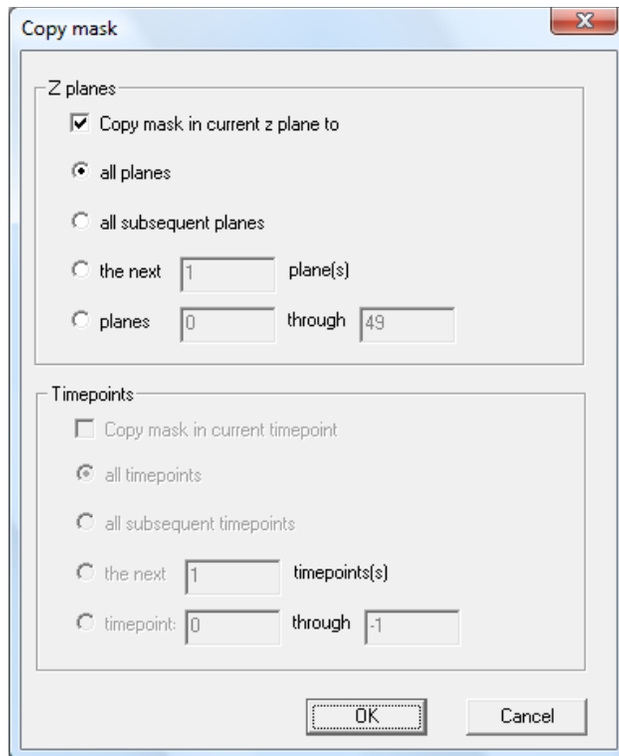
### 10.2.2.2 Editing the Mask Manually

Once you have created an empty mask, you can use a variety of tools to construct a useful mask. There are two means of editing the mask: using drawing tools from the tool menu, and performing fine adjustments using tools from the mask menu on the info/tool bar.

1. Make sure that your mask is the currently selected mask. To do this, click on the mask menu  and select the desired mask from the list.
2. Select a tool from the drop-down tool menu  in the info/tool bar (see [Using the Tool Menu](#) on page 184).
  -  **Marquee tool** – click and drag to draw a box outline, then select **Mask > Mark Selection** to fill in your selection. To remove the box, choose **Mask > Unmark Selection**.
  -  **Single pixel pencil** – click and hold the mouse button to generate a single pixel-width freehand line (**NOTE:** the resulting line is guaranteed to be continuous—no gaps—as long as you have the mouse button down)
  -  **Small pencil** – click and hold the mouse button to generate a thin freehand line
  -  **Large pencil** – click and hold the mouse button to generate a thick freehand line
  -  **Small eraser** - click and hold the mouse button to remove small amounts
  -  **Large eraser** - click and hold the mouse button to remove large amounts
  -  **Paint bucket** – fill a shape by single clicking within any closed shape drawn with the small or large pencil

Additionally, you may generate a 2D or 3D rectangular volume by first defining a volume as discussed in [Making a 2D or 3D \(x,y,z or x,y,t\) Selection](#) on page 186, then choosing **Mask > Mark Selection**.

6. Use the tool to add or remove pixels. The selected pixels will be shown in blue.
7. If you are working with a 3D or timelapse image, and would like to copy your selections to other planes in the image, select **Mask > Copy This Plane**. The following dialog box will appear:



8. Use the radio buttons to select the desired action depending on your capture type, and click **OK**.
9. To further refine your mask, you may select a variety of operations by selecting **Mask > Mask Filter**. The menu items are described below. You may use these operations in any order or as many times as desired.
  - **Erode** - Applies a kernel to the entire mask that removes a user-defined number of voxels from the perimeter of the mask. This function can be applied to in only the  $x, y$  dimension (2D) or in the  $z$  dimension (3D) as well.
  - **Dilate** - Applies a dilation kernel to the entire mask and thickens features. The user defines a radius of the number of pixels the current mask will be expanded by and whether the mask should be dilated in 2D or 3D.
  - **Skeletonize** – Reduces the mask to a set of connected junctions. This function can only be applied to single plane (2D) images.

**NOTE:** These operations do not create new masks, but simply edit the current mask.

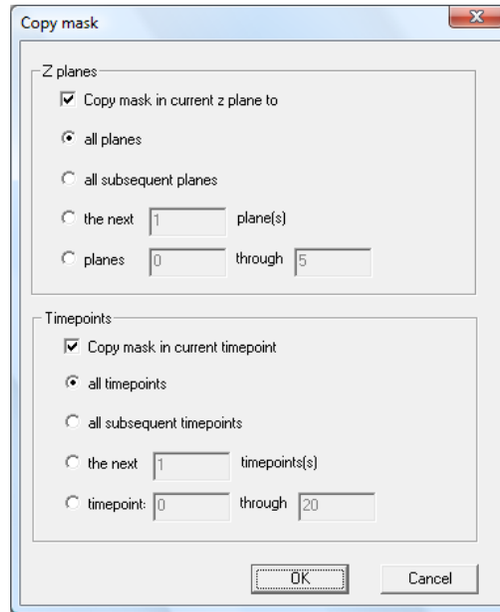
### 10.2.2.3 Moving the Mask

You can shift the mask in any direction by selecting **Ctrl**-arrow keys.

### 10.2.2.4 Copying a Manually Created Mask to Other Planes or Other Images

Once you have created a mask, you may choose to copy the mask to other planes in the image (if you have a 2D timelapse or 3D image). To do this:

1. Select **Mask > Copy This Plane**. The following dialog box will appear:



2. Select the desired data you would like to copy the mask to and select **OK**.

You may also copy the mask to all of the images of the same size in the same slide. To do so, from an open view of an image:

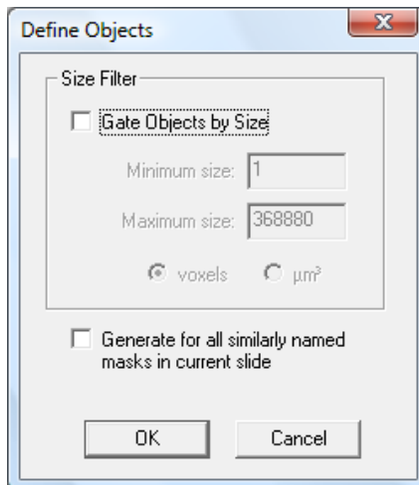
1. Display the mask that you wish to copy by selecting it from the mask menu  in the info/tool bar.
2. Select **Mask > Copy to all images in slide**.

The mask will be copied to all of the images in the slide provided they are the same size.

### 10.2.3 Defining Objects

Objects are defined when you wish to get data for individual objects in a mask. The mask statistics dialog box allows you to automatically define objects when generating statistics. However, in order to view timelapse graphs of individual objects or to obtain a quick object count, you will need to define objects. To do so:

1. Generate a mask that contains distinct regions, either using manual editing or segmentation techniques.
2. Select **Mask > Define Objects in Mask**. The following dialog box will appear:

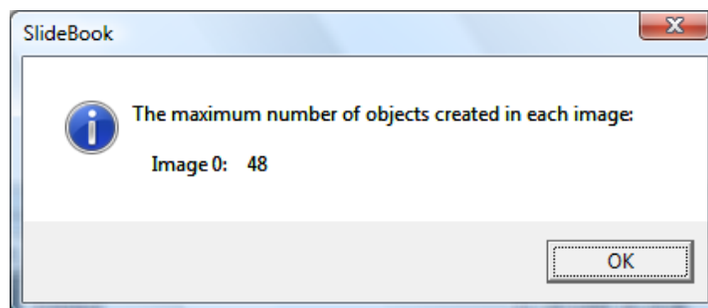


The dialog box allows you to set upper and lower bounds on the size of the regions that should be made into objects.

3. If you would like to set limits on the size of the objects, select **Gate Objects by Size** and choose a minimum and maximum size. The size bounds can be specified in either voxels or cubic microns (square microns for a 2D image).
4. Select the **Generate for all similarly named masks in current slide** checkbox to perform the object definition operation on the entire slide.

**NOTE:** This process does not create multiple masks, but rather assigns a number to each isolated, contiguous subregion of the existing mask.

5. Select **OK**. A dialog box will appear that gives the status of the object generation. When object generation is complete, a dialog box will appear, noting the number of objects generated. When objects are defined for a 2D timelapse image, SlideBook will display the maximum number of objects generated at any given timepoint.



6. Click on **OK**. If you now position the cursor over one of the objects in the Main View, the object number appears to the right of the cursor location coordinates in the info/tool bar.

7. If you would like to remove objects, select **Masks > Update Object Definitions**. The Define Objects dialog will appear as it did when selecting **Masks > Define Objects in Mask**.
8. Select **Gate Objects by Size** and increase the lower threshold, or decrease the upper threshold, then select **OK**. The number of objects generated will decrease. This is particularly useful when working with noisy data that may contain a very large number of objects.

**NOTE:** You may not decrease the lower threshold to increase the number of objects. If you have removed too many objects, then you will need to remake your mask.

### 10.2.4 Splitting Objects

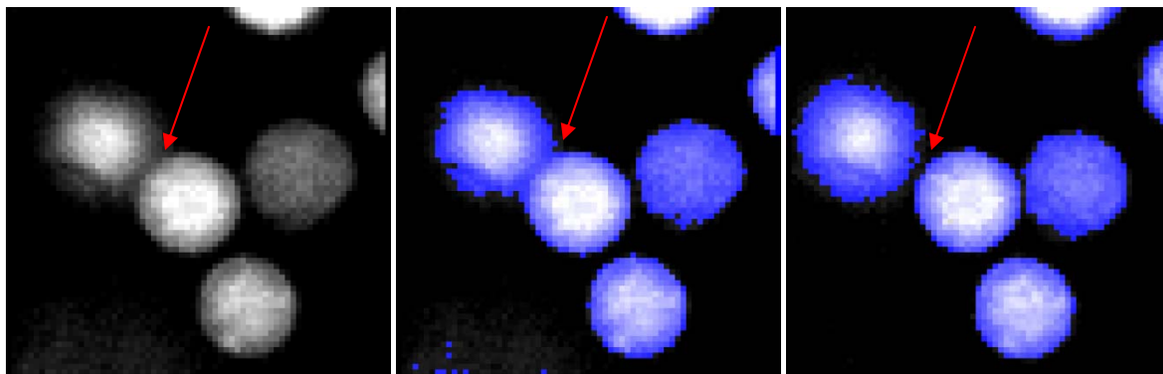
If you have two or more objects that are touching but exist under one contiguous region of the mask (a mask object), you may choose to split the mask into multiple objects. This can be achieved either manually by drawing lines along which to split the object or in an automated fashion.

#### 10.2.4.1 Manually Splitting Objects

To split objects manually, use the line ROI tool to draw lines where you would like to separate the objects. Then select **Mask > Advanced Operations > Split Objects Along Lines**. To see the split objects, turn the ROI display off by going to **Annotations > Regions**. You may undo the split by selecting **Mask > Advanced Operations > Undo Split Objects Along Lines**.

#### 10.2.4.2 Automatically Splitting Objects

To split objects automatically, SlideBook employs a “watershed” algorithm. This is useful to separate objects that may be joined by a thin connector as shown below:



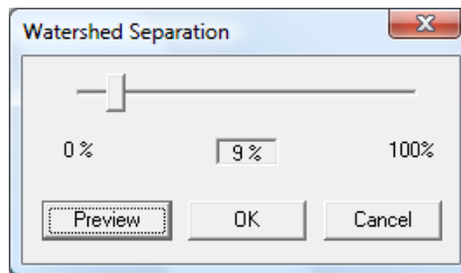
Raw Data

Initial Mask

After Split Objects

To perform a watershed operation:

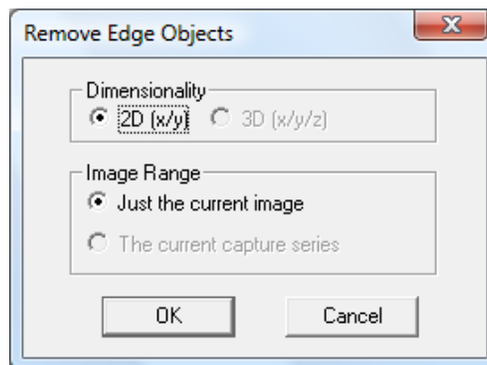
1. Generate a mask that contains distinct regions, either using manual editing or segmentation techniques.
2. Go to **Mask > Advanced Operations > Split Object Automatically**. The following dialog will appear:



3. The slider bar controls how aggressive the splitting algorithm will be. A value of 100% refers to no splitting while 0% refers to the most aggressive splitting algorithm. Adjust this slider to a position you feel will give you clear object separation.
4. Click **Preview**. You will notice that the mask is updated on the image.
5. If the splitting is adequate, click **OK**. If the splitting is inadequate, repeat steps 3 and 4 until you are satisfied with the result, then click **OK**.

### 10.2.5 Removing Objects from Edge of Image

In some instances, you may wish to eliminate objects that touch the edges of your image. To do so, select **Mask > Advanced Operations > Remove Edge Objects**. The following dialog box will appear. Select the desired Image Range and select **OK**.



### 10.2.6 Tracking Objects (Particle Tracking)

Once you have defined objects, you may wish to track them over time. SlideBook offers two methods for particle tracking, automated and manual.

#### 10.2.6.1 Automated Particle Tracking (Tutorial)

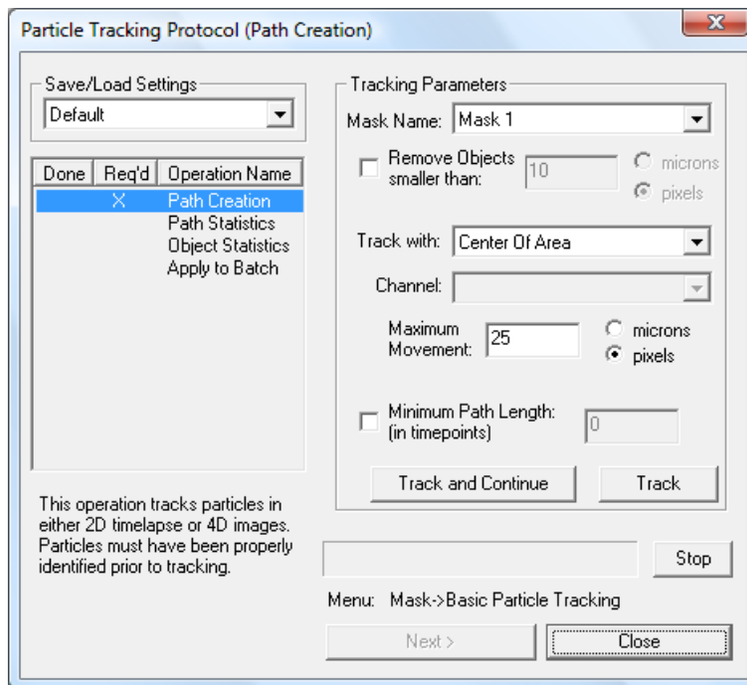
To perform particle tracking, you may either use the Particle Tracking Protocol or individual dialog boxes. The Particle Tracking Protocol has the advantage of being able to save and reuse your particle tracking parameters, and apply them in a batch fashion. Below we describe how to use the Particle Tracking Protocol. Each step in the protocol can be accessed via a menu item, which is also noted. You may wish to practice particle tracking

using the sample file “ParticleTrackingDemo.sld”. You may download this file at the SlideBook download site.

1. Select the objects you wish to track by generating a mask using thresholding and/or manual marking (see [Creating a Mask using Threshold Techniques](#) on page 224 or [Creating Masks Manually](#) on page 226). You may need to separate objects that are touching using mask filters (**Mask > Mask Filter > Erode**) or an object splitting algorithm (see [Splitting Objects](#) on page 231).

For ParticleTrackingDemo.sld, create a mask using a threshold of 217 for the i-485 channel.

2. Begin by selecting **Mask > Particle Tracking > Particle Tracking Protocol**. The following dialog will appear:



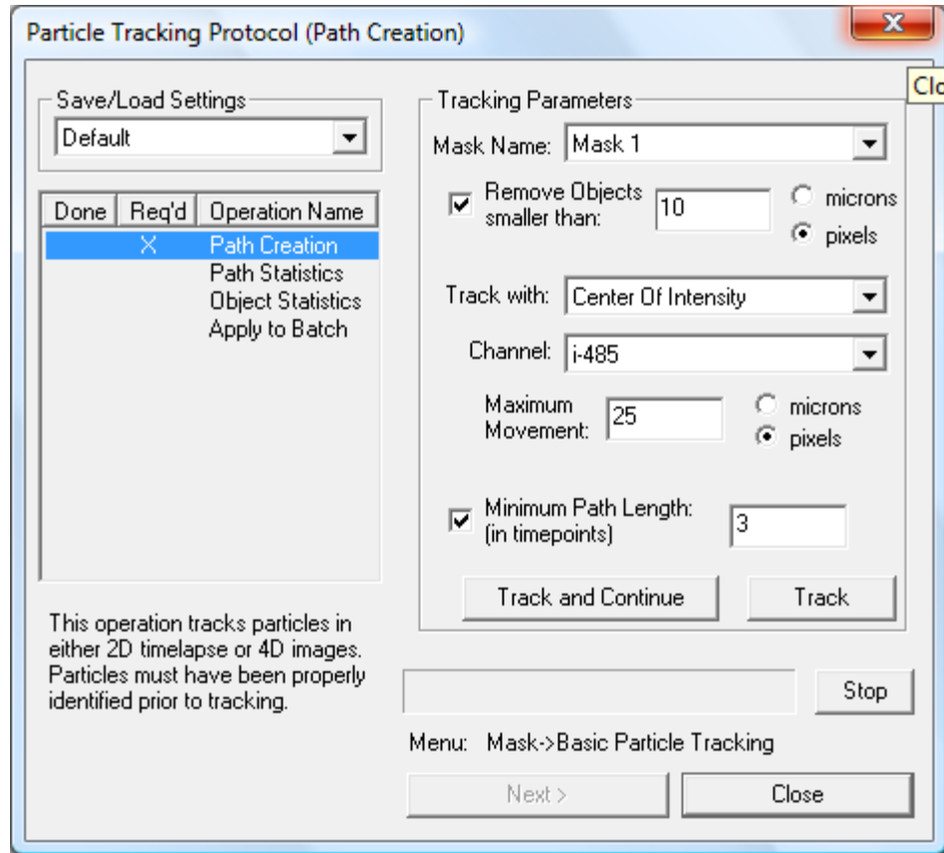
The dialog box contains the list of protocol steps on the left along with a dropdown menu for saving and loading settings. The list of steps is underneath. Required steps must be completed before moving to the next step in the protocol. Steps that are not required may be skipped. Once a step is complete, an **X** will appear in the **Done** column. The right side of the dialog will update as you progress through the protocol.

3. Select **Tracking Parameters** as described below.
  - **Mask Name** – Select the Mask that contains the objects you would like to track. The menu will default to the mask that is currently displayed on your open view.

- **Remove Objects smaller than** – Check this box to selectively remove objects smaller than a given size. Enter the size in the edit field and choose either microns or pixels as the unit.
- **Track with** – Select the parameter that you wish to track from the dropdown list:
  - **Center of Area** – the coordinates of the center of the object, unweighted by intensity values.
  - **Center of Intensity** – the coordinates of the center of the object, weighted by intensity values. If you select this feature, you will also need to select the channel that is used for weighting.
  - **Mean Adjusted Center of Intensity** – the coordinates of the center of the object, weighted by intensity values that are above the mean intensity. This is a useful feature to track if you have delineated your objects by hand. By using values that fall above the mean, any imprecision in your hand delineation does not impact the center of intensity calculation. If you select this feature, you will also need to select the channel that is used for weighting.
- **Maximum Movement** – enter the maximum movement that you expect from any given object from one timepoint to the next. This will help the particle tracking algorithm distinguish between objects that may have similar centroid values.
- **Minimum Path Length** – enter the minimum number of timepoints that must be tracked in order to determine a path. For instance, if you enter “3,” any object that cannot be followed for at least three timepoints will be removed for the mask.

**NOTE:** This dialog box may also be found via the **Mask > Particle Tracking > Basic Particle Tracking** menu.

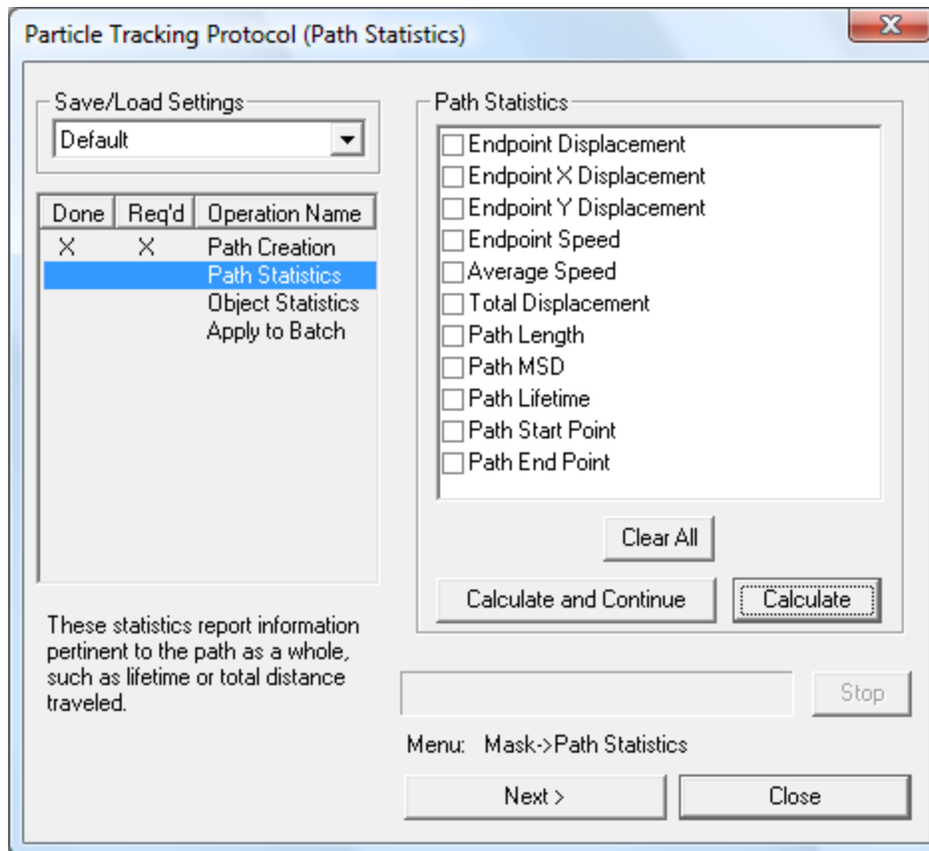
4. For the tutorial, select the parameters as shown in the dialog below:



5. Select **Track** and the tracking algorithm will be executed. When completed the paths will be displayed on the image. You may click on the image and scroll through the timepoints to see the paths. The first point in the path appears blue, and the last appears red, with a gradation of color between. At this point, you may wish to view the object IDs by selecting **View > Annotations > Object IDs**. Also, you may wish to perform the tracking algorithm using new parameters. You may enter those parameters and then press **Track** again. If you already know the parameters in advance you may select **Track and Continue**. The tracking algorithm will be executed and the protocol will automatically advance to the next step.

**NOTE:** You may turn path display on and off by selecting **Mask > Particle Tracking > Display Paths**.

6. Once tracking is complete, select **Next**. The protocol will advance to the next step.



Here, you may choose to calculate the following statistics for each path.

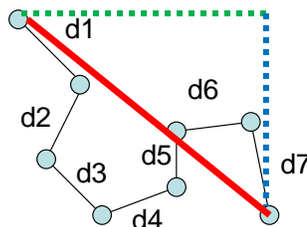


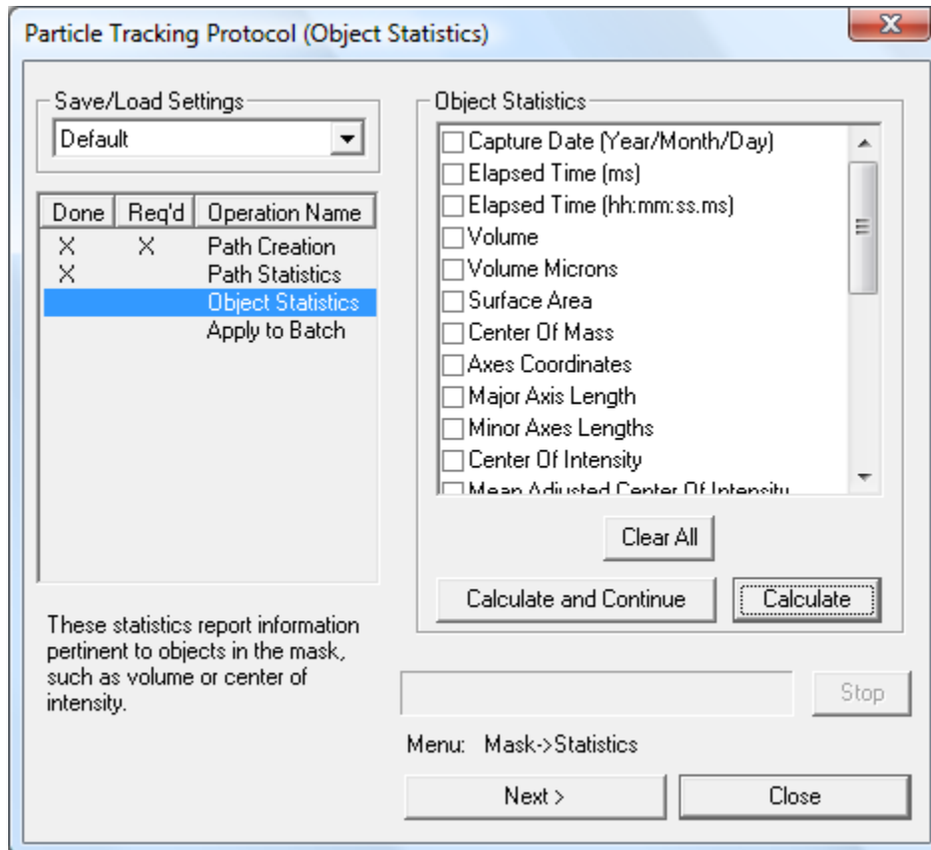
Fig. 10.1 – Schematic of tracked object. Blue circles represent centroids

- **Endpoint Displacement** – the distance traveled between the first and last timepoints of the path, as represented by the length of the red line in the figure above.
- **Endpoint X Displacement** – the distance traveled in the x-axis between the first and last timepoints of the path, as represented by the length of the dotted green line in the figure above.
- **Endpoint Y Displacement** – the distance traveled in the x-axis between the first and last timepoints of the path, as represented by the length of the dotted blue line in the figure above.
- **Endpoint Speed** – the distance represented by the length of the red line divided by the time required to travel that distance.

- **Average Speed** – the total distance traveled (the sum of distances d1 to d7 as shown in the figure above) divided by the time required to travel that distance.
- **Total Displacement** – the total distance traveled by the object (the sum of distances d1 to d7 as shown in the figure above).
- **Path Length** – number of timepoints in the path.
- **Path MSD** -
- **Path Lifetime** – the total elapsed time for a given path.
- **Path Start Point** – centroid of the first point in the path.
- **Path End Point** – centroid of the last point in the path.

**NOTE:** The Path Statistics dialog box can also be found by selecting **Mask > Particle Tracking > Path Statistics**.

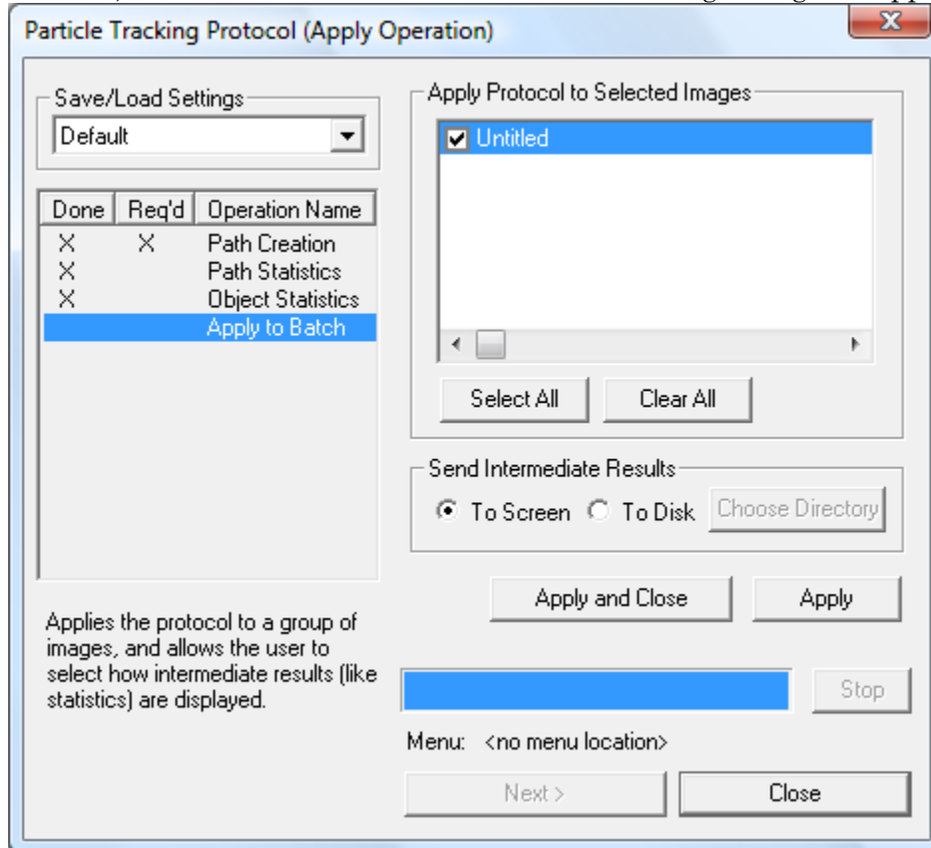
7. Select the statistics you wish to calculate using the checkbox. Select **Calculate and Continue** if you wish to progress to the next step, or simply select **Calculate** if you wish to view your statistics without going to the next step, then select **Next** when you are ready to go to the next step. The following dialog will appear:



**NOTE:** Object statistics can also be calculated via the Mask Statistics dialog (**Mask > Statistics**).

Statistic definitions are given in the section [Generating and Exporting Mask and Object Statistics](#) on page 243.

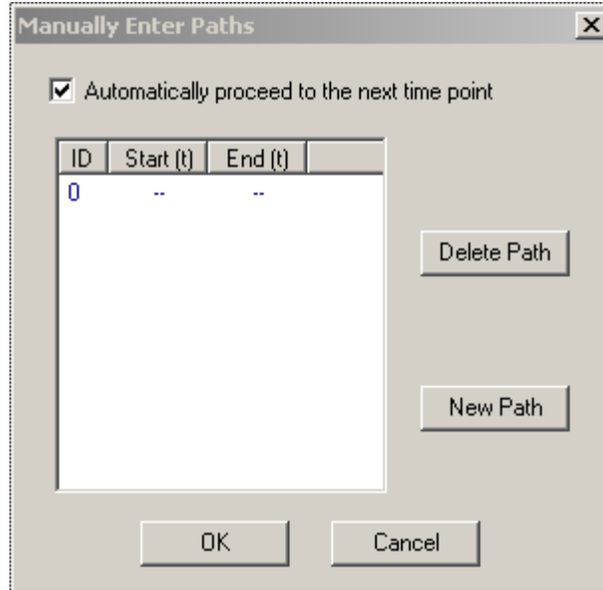
8. Select the statistics that you wish to calculate and either select **Calculate and Continue**, or **Calculate** and then **Next**. The following dialog will appear:



9. If you have more than one image in the slide, you may choose to apply the protocol to other images in the slide. Simply check the checkboxes of the images you wish to analyze, and either select **Apply and Close** to finish the protocol, or select **Apply** to perform the batch analysis. You may select whether you would like to send results to the screen or straight to disk.
10. If you would like to save the particle tracking parameters for processing another slide, select **Save As** from the **Save/Load Settings** dropdown menu. Enter a name for the settings and select **OK**. You may now select those settings the next time you open the Particle Tracking Protocol dialog. You may also delete settings or modify the settings and then choose **Save** to save the settings with the same name.

### 10.2.6.2 Performing Manual Particle Tracking

1. Open a main view of the image containing objects you wish to track.
2. Go to **Mask > Particle Tracking > Manual Tracking**. The following dialog box will appear:



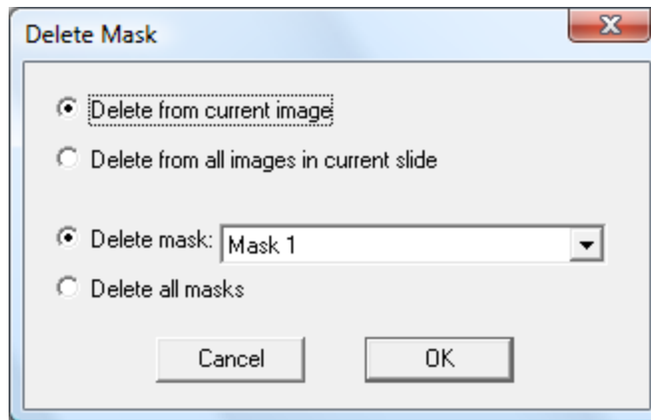
3. Select either the medium or large pencil tool from the tool menu. The view will go to the first timepoint automatically.
4. Click near the center of the first object you would like to track. The view will automatically move to the next timepoint.
5. Continue clicking on the object you would like to track until the path is complete. When complete, either choose **New Path** to go back to the first timepoint and start over, or **OK** to close the dialog. You may now generate object statistics as described in [Generating Statistics for Masks and Objects](#) on page 243.

### 10.3 Displaying or Deleting a Mask

If you have just issued either the **Mask > Create** or **Mask > Segment** commands, the new mask is already selected and displayed. If not, you must select the mask you want to edit using the mask pop-up menu  in an open view of the image. Note that while you are editing from within a particular view, the mask is owned by the image, not just the particular view. Any changes you make to the mask can be seen in any other view in which the same mask is displayed.

To delete a mask:

1. Select **Mask > Delete**. The following dialog box will appear:



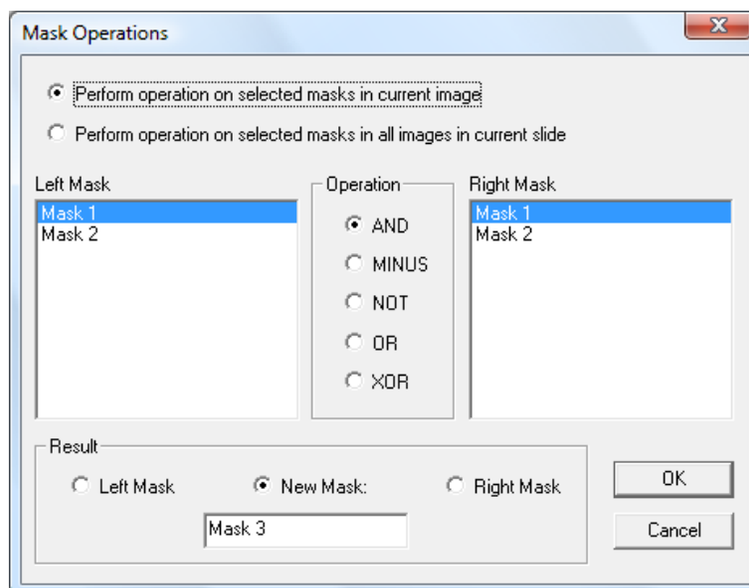
2. Choose the mask(s) you wish to remove using the radio buttons and dropdown menu. You may also choose to delete the mask from the current image or from all images in the slide.
3. Choose **OK** to remove the mask.

#### 10.4 Boolean Mask Operations

Masks can be combined or altered using Boolean operations. For instance, the intersection of two masks can be generated by an AND operation. The Mask Operations dialog lets you either generate a new mask that contains the result of a Boolean operation or replace an existing mask with the result of an operation.

To perform Boolean mask operations:

1. Open a view that contains the masks you would like to manipulate.
2. Select **Mask > Mask Operations**. The following dialog will appear:

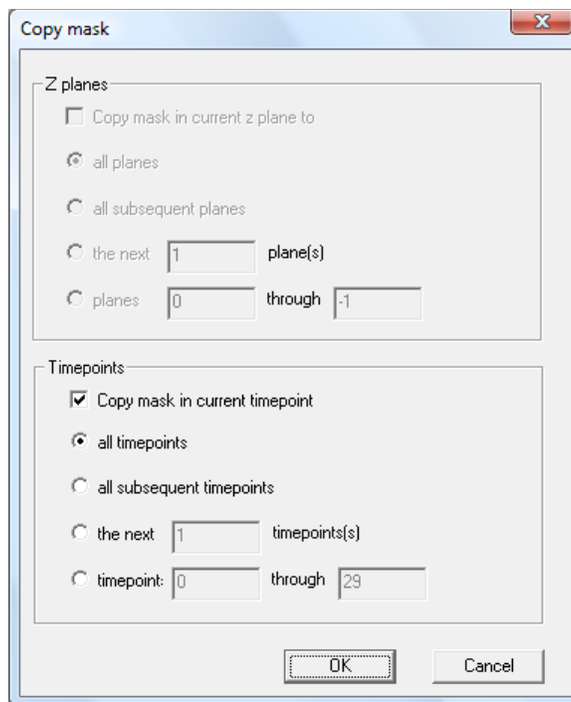


- Using the radio buttons, select whether the operation will take place for the current image or the entire slide.
- Select the masks to be used from the **Left Mask** and **Right Mask** lists.
- Select the operation to be performed using the radio buttons.
  - AND** – selects pixels that are in both masks
  - MINUS** – removes any pixels that are present in the right mask from the left mask
  - NOT** – selects all pixels that are not marked in the left mask
  - OR** – selects pixels that are in either of the two selected masks
  - XOR** – adds both masks and removes any common pixels
- Select the destination for the mask operation using the radio buttons. If you want the result to replace the contents of either the left or right mask, select the corresponding radio button for that mask. If you want to generate a new mask, select the **New Mask** radio button and give the new mask a name in the edit field.
- Click **OK** to perform the Boolean operation. You may display the new or edited mask as described above.

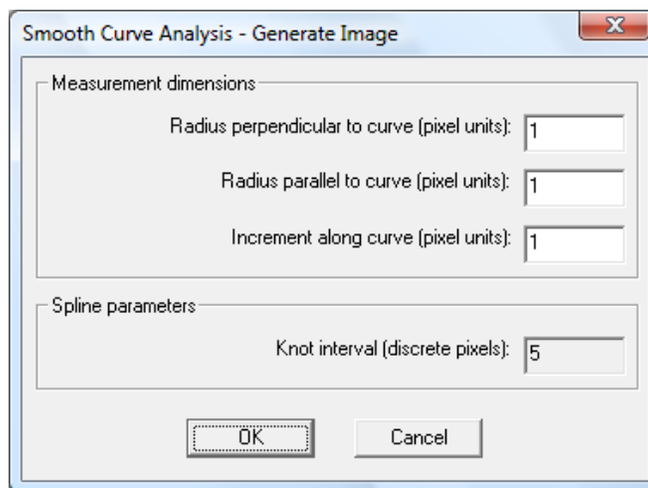
### 10.5 Using Masks for Smooth Curve Analysis (Kymograph)

You may use masks to identify regions to be used for smooth curve analysis, also known as kymograph analysis. Smooth curve analysis allows you to follow time-dependent lateral intensity changes along a static object. For example, an action potential moving along a neural spine could be displayed as an  $x$  vs.  $t$  image using smooth curve analysis. You could then extract velocity and displacement information from this image. To perform smooth curve analysis:

- Open a Main View or Three View of your timelapse data.
- Create a new mask as discussed above in [Creating an Empty Mask](#) on page 226.
- Select the single pixel pencil tool from the tool menu  and draw a line along the desired object.
- Copy the mask to all planes in the image by selecting **Mask > Copy This Plane**. The following dialog box will appear:



5. Use the radio buttons to select the desired action, and click **OK**.
6. Select **Mask > Advanced Operations > Smooth Curve Analysis**. The following dialog box will appear:



7. Adjust the dimensions as necessary (please see Appendix A for a detailed description of Smooth Curve parameters) and select **OK**. A new image will appear in your slide.
8. Open a main view of the Smooth Curve Image and select the y axis from the axis menu  on the info/tool bar. You are now able to display a 2D view of your 3D

selection. You may perform velocity measurements of timelapse data with the ruler tool (see [Making Distance or Velocity Measurements](#) on page 187).

### 10.6 Generating and Exporting Mask and Object Statistics

You can answer many quantitative questions about a mask through mask statistics. SlideBook lets you compute three types of mask statistics: statistics for the entire mask, statistics for each timepoint in a time lapse image, and statistics for objects (defined above on page 223). SlideBook allows for a wide range of statistics to be generated, from simple to sophisticated.

#### 10.6.1 Counting Objects in a Mask

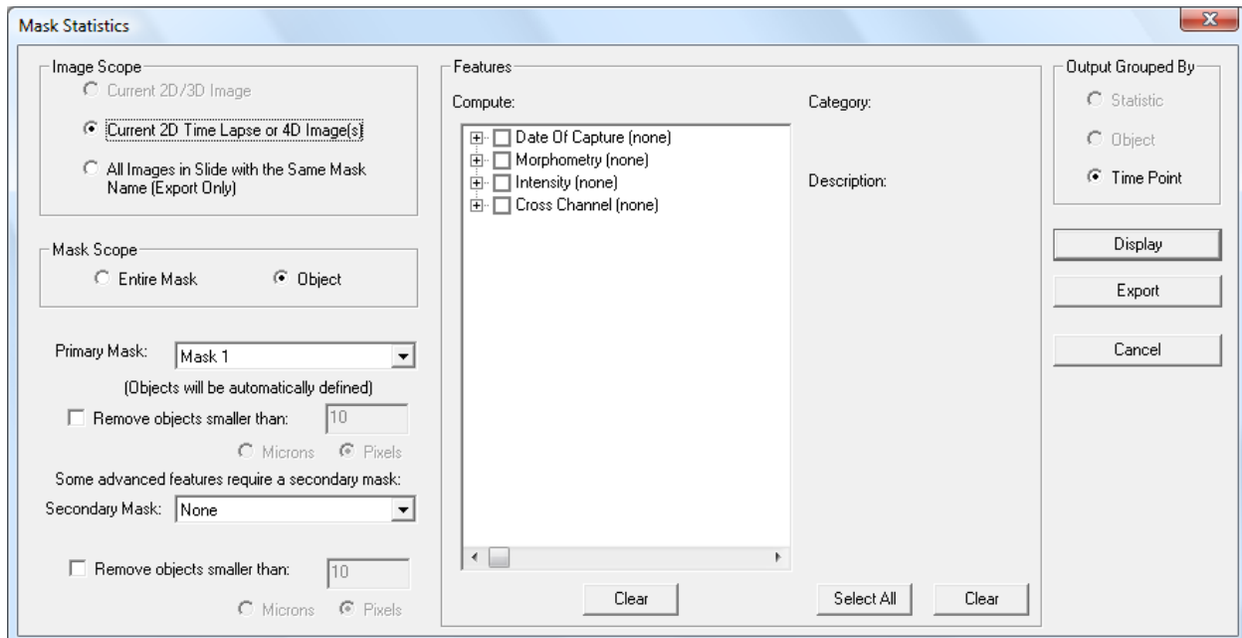
You may wish to count the individual objects, in your mask. If you do not wish to obtain other statistics, you may count objects by generating objects (see page 229). You may also get an object count by generating the **Object Count** statistic (see next section). This is especially useful if you wish to count objects for multiple images in a slide.

#### 10.6.2 Generating Statistics for Masks and Objects

SlideBook allows you to generate a wide variety of statistics for regions of interest in 2D and 3D data.

To generate statistics from an open view of an image:

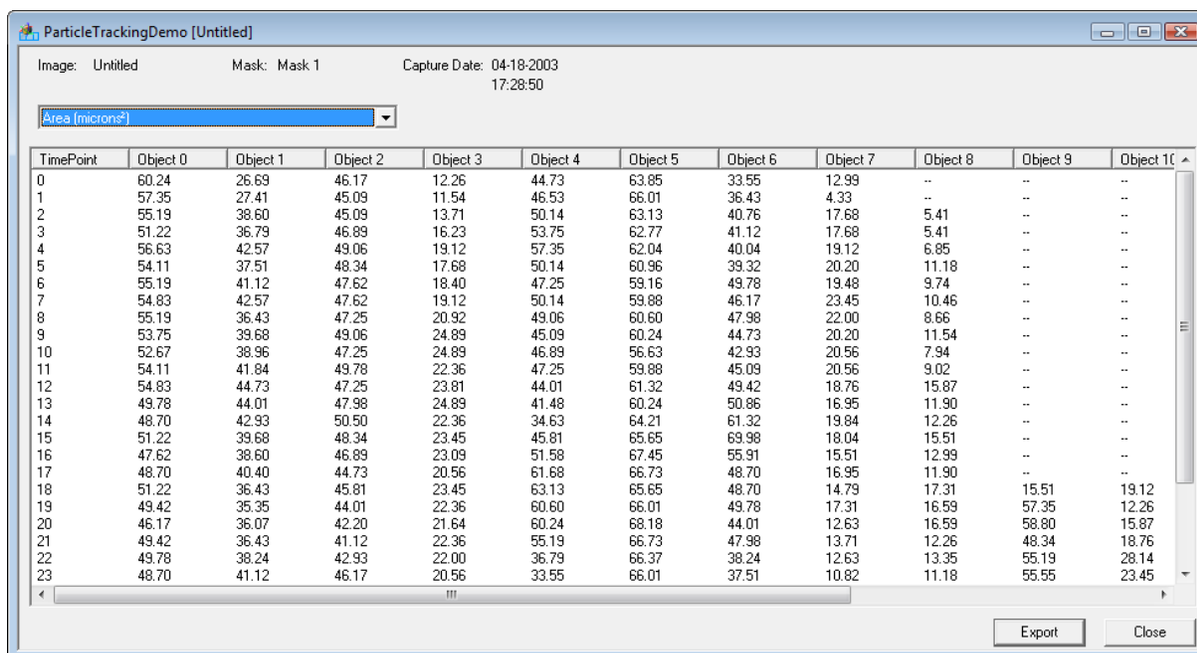
1. Display the mask of interest by selecting it from the mask menu .
2. Select **Statistics > Mask Statistics**. The following dialog box will appear:



The dialog box has the following sections:

- **Image Scope** – allows for generation of statistics in multiple images. If you are working with a 2D or 3D image, the first radio button will be selected by default. If you are working with a 2D timelapse or 4D data set, the second radio button will be selected by default. You may select the third radio button if you have multiple images with the same mask name.
- **Mask Scope** – allows you to select whether you want statistics for the entire mask or on a per object basis. If you have chosen to obtain statistics for multiple images, the **Object** option will be unavailable.
- **Primary Mask** – defaults to the mask that is currently displayed but can be altered via the dropdown menu. The primary mask will automatically be made into objects. You may also remove objects under a specified pixel size (gating by size, see page 229). To get statistics for all pixels in the mask, leave this box unchecked.
- **Secondary Mask** – can be selected to obtain additional statistics (see below).
- **Features** – lists all available statistics that may be calculated. Note that the list of features will depend on the dimensions of the mask. For instance, area will be displayed as a statistic for 2D masks, while volume will be displayed for 3D masks.
- **Output Grouped By** – these choices are only activated if you are performing object statistics on 2D timelapse or 4D images. Sample output is shown below:

#### Output Grouped By Statistic:



The screenshot shows a software window titled "ParticleTrackingDemo [Untitled]". At the top, it displays "Image: Untitled", "Mask: Mask 1", and "Capture Date: 04-18-2003 17:28:50". Below this is a dropdown menu currently set to "Area [microns²]". The main part of the window is a table with 12 columns: "TimePoint", "Object 0", "Object 1", "Object 2", "Object 3", "Object 4", "Object 5", "Object 6", "Object 7", "Object 8", "Object 9", and "Object 10". The table contains 24 rows of data, with the first row (TimePoint 0) starting with values 60.24, 26.69, 46.17, 12.26, 44.73, 63.85, 33.55, 12.99, and so on. The last row (TimePoint 23) ends with values 48.70, 41.12, 46.17, 20.56, 33.55, 66.01, 37.51, 10.82, and so on. At the bottom right of the window are "Export" and "Close" buttons.

TimePoint	Object 0	Object 1	Object 2	Object 3	Object 4	Object 5	Object 6	Object 7	Object 8	Object 9	Object 10
0	60.24	26.69	46.17	12.26	44.73	63.85	33.55	12.99	--	--	--
1	57.35	27.41	45.09	11.54	46.53	66.01	36.43	4.33	--	--	--
2	55.19	38.60	45.09	13.71	50.14	63.13	40.76	17.68	5.41	--	--
3	51.22	36.79	46.89	16.23	53.75	62.77	41.12	17.68	5.41	--	--
4	56.63	42.57	49.06	19.12	57.35	62.04	40.04	19.12	6.85	--	--
5	54.11	37.51	48.34	17.68	50.14	60.96	39.32	20.20	11.18	--	--
6	55.19	41.12	47.62	18.40	47.25	59.16	49.78	19.48	9.74	--	--
7	54.83	42.57	47.62	19.12	50.14	59.88	46.17	23.45	10.46	--	--
8	55.19	36.43	47.25	20.92	49.06	60.60	47.98	22.00	8.66	--	--
9	53.75	39.68	49.06	24.89	45.09	60.24	44.73	20.20	11.54	--	--
10	52.67	38.96	47.25	24.89	46.89	56.63	42.93	20.56	7.94	--	--
11	54.11	41.84	49.78	22.36	47.25	59.88	45.09	20.56	9.02	--	--
12	54.83	44.73	47.25	23.81	44.01	61.32	49.42	18.76	15.87	--	--
13	49.78	44.01	47.98	24.89	41.48	60.24	50.86	16.95	11.90	--	--
14	48.70	42.93	50.50	22.36	34.63	64.21	61.32	19.84	12.26	--	--
15	51.22	39.68	48.34	23.45	45.81	65.65	69.98	18.04	15.51	--	--
16	47.62	38.60	46.89	23.09	51.58	67.45	55.91	15.51	12.99	--	--
17	48.70	40.40	44.73	20.56	61.88	66.73	48.70	16.95	11.90	--	--
18	51.22	36.43	45.81	23.45	63.13	65.65	48.70	14.79	17.31	15.51	19.12
19	49.42	35.35	44.01	22.36	60.60	66.01	49.78	17.31	16.59	57.35	12.26
20	46.17	36.07	42.20	21.64	60.24	68.18	44.01	12.63	16.59	58.80	15.87
21	49.42	36.43	41.12	22.36	55.19	66.73	47.98	13.71	12.26	48.34	18.76
22	49.78	38.24	42.93	22.00	36.79	66.37	38.24	12.63	13.35	55.19	28.14
23	48.70	41.12	46.17	20.56	33.55	66.01	37.51	10.82	11.18	55.55	23.45

**Output Grouped By Object:**

Time Point	Area (microns <sup>2</sup> )	Mean Intensity (ADU), ch1:i-405	Mean Intensity (ADU), ch1:i-485	Ratio[i-405][i-485]
0	60.24	61.80	278.14	0.2222
1	57.35	62.79	277.25	0.2265
2	55.19	54.79	269.29	0.2035
3	51.22	53.07	278.85	0.1903
4	56.63	50.58	279.26	0.1811
5	54.11	52.45	279.08	0.1880
6	55.19	52.25	278.51	0.1876
7	54.83	54.30	278.27	0.1951
8	55.19	53.99	282.18	0.1913
9	53.75	54.68	283.89	0.1926
10	52.67	55.18	281.82	0.1958
11	54.11	56.33	288.63	0.1951
12	54.83	56.72	297.45	0.1907
13	49.78	54.42	295.08	0.1844
14	48.70	53.72	289.09	0.1858
15	51.22	53.15	288.26	0.1844
16	47.62	53.64	291.02	0.1843
17	49.70	53.95	292.64	0.1896

**Output Grouped by Timepoint:**

Object	Area (microns <sup>2</sup> )	Mean Intensity (ADU), ch1:i-405	Mean Intensity (ADU), ch1:i-485	Ratio[i-405][i-485]
0	60.24	61.80	278.14	0.2222
1	26.69	48.81	243.81	0.2002
2	46.17	60.00	266.77	0.2249
3	12.26	46.29	229.97	0.2013
4	44.73	48.90	281.26	0.1739
5	63.85	48.92	302.68	0.1616
6	33.55	45.09	267.49	0.1685
7	12.99	52.83	228.06	0.2317

3. Check all the statistics you want to compute. SlideBook offers the following options for statistics for a single mask:

**Morphometry**

- **Object Count** – number of objects. This statistic is only available when you select **Entire Mask** in the Mask Scope section of the statistics dialog.
- **Volume (voxels) / Area (pixels)** - total number of voxels. Note that in a 2D or time lapse image, you will always get an area measurement.

- **Volume ( $\mu\text{m}^3$ ) / Area ( $\mu\text{m}^2$ )** - volume in cubic microns (or square microns in 2D or timelapse images). Note that the image must have the correct objective and optovar selected for these values to be accurate (see [Defining Objectives](#) on page 62 and [Defining Magnification Changers](#) on page 73).
- **Surface area ( $\mu\text{m}^2$ ) / Perimeter ( $\mu\text{m}$ )** – the area of exposed voxel faces for 3D objects (or exposed pixel faces in 2D or time lapse images). Note that these values should only be used as rough estimates, as area or perimeter measurements are highly affected by the resolution of the image.
- **Center of Area/Volume** – the coordinates of the center of the object, unweighted by intensity values.
- **Major Axis Coordinates** – the coordinates of the two edge voxels that lie along the major axis.
- **Major Axis Length** – the length of the longest line that runs through the center of volume, also called the ‘axis of minimal inertia’.
- **Minor Axis Length** – the length of the minor axis (in 2D) or both minor axes (in 3D).
- **Longest Chord Length** – the distance between the two furthest voxels in the object.
- **Major/Minor Shape Factor** – The ratio of major axis to first minor axis (i.e. width vs. height).

### Intensity

- **Center Of Intensity** – the coordinates of the center of the marked object weighted by intensity, calculated for each channel.
- **Mean Adj Center Of Intensity** – the coordinates of the center of the object, weighted by intensity values (minus the mean intensity of the object).
- **Mean Intensity** – average marked voxel intensity value for each channel.
- **Sum intensity** – sum of all marked voxel intensity values for each channel. Note that this value is equivalent to the product of the mean intensity and the volume or area.
- **Minimum Intensity** – the minimum voxel value in the object.
- **Maximum Intensity** – the maximum voxel value in the object.
- **Median Intensity** – the median voxel value in the object.

- **Variance** – the variance of intensities of the marked voxels.
- **Standard Deviation** – the standard deviation of intensities of the marked voxels.
- **Background Intensity** – the average background intensity in the channel.

**Cross-Channel**

- **Pearson’s Correlation** – the correlation of intensity in one channel with another; a 0 value signifies no correlation, while a value of 1.0 signifies perfect correlation. A negative value signifies anti-correlation.

This statistic can be used for measuring colocalization. The equation used is shown below.  $R_i$  is the intensity in channel 1 for pixel  $I$ , and  $G_i$  is the intensity in channel 2 for the same pixel.  $R_{av}$  and  $G_{av}$  are the average (mean) intensity values over all pixels.

$$r = \frac{\sum_i (R_i - R_{av}) \cdot (G_i - G_{av})}{\{\sum_i (R_i - R_{av})^2 \cdot \sum_i (G_i - G_{av})^2\}^{\frac{1}{2}}}$$

- **Mander’s Coefficients** – these coefficients indicate the percent of protein 1 colocalized with protein 2 ( $M_1$ ) and the percent of protein 2 colocalized with protein 1 ( $M_2$ ) as given by the equation below. The value  $T$  (threshold) is considered to be zero intensity. Thus, adequate background subtraction must be performed.

$$M_1 \cong \frac{\sum_{I_1 > T} I_1}{\sum_{All I_1} I_1}, \quad \text{and} \quad M_2 \cong \frac{\sum_{I_2 > aT+b} I_2}{\sum_{All I_2} I_2}$$

SlideBook offers the following statistics to describe relationships between objects that are present in two different masks. You must first set the **Mask Scope to Object**, then select a secondary mask to view these statistics.

**Cross-Mask**

- **Mask Overlap (objects)** - for each object in the primary mask, this returns a count of the number of overlapping objects in the secondary mask.
- **Mask Overlaps (voxels)** – for each object in the primary mask, this returns the number of overlapping pixels/voxels in the secondary mask.
- **Mask Overlaps (largest object volume)** – for each object in the primary mask, this returns the area or volume of the largest overlapping object in the secondary mask.

- **Mask Contains Other Masks** – returns the IDs of the objects in the secondary mask that overlap with the primary mask.
4. Select **Display** to generate a new window that shows the selected statistics. Alternatively, you may directly export the data by pressing **Export** (see below).  
  
If you have generated object data, you may click on any of the object numbers in the table, and the corresponding object will be highlighted in pink in the data view.
  5. To save the data, select the **Export** button. A dialog box will appear that will allow you to choose the file destination.
  6. Select a destination and a file name, then select **Save**. The file will be saved as a tab-delimited text file (.txt) file, which can be easily importable into a spreadsheet program, such as Excel.

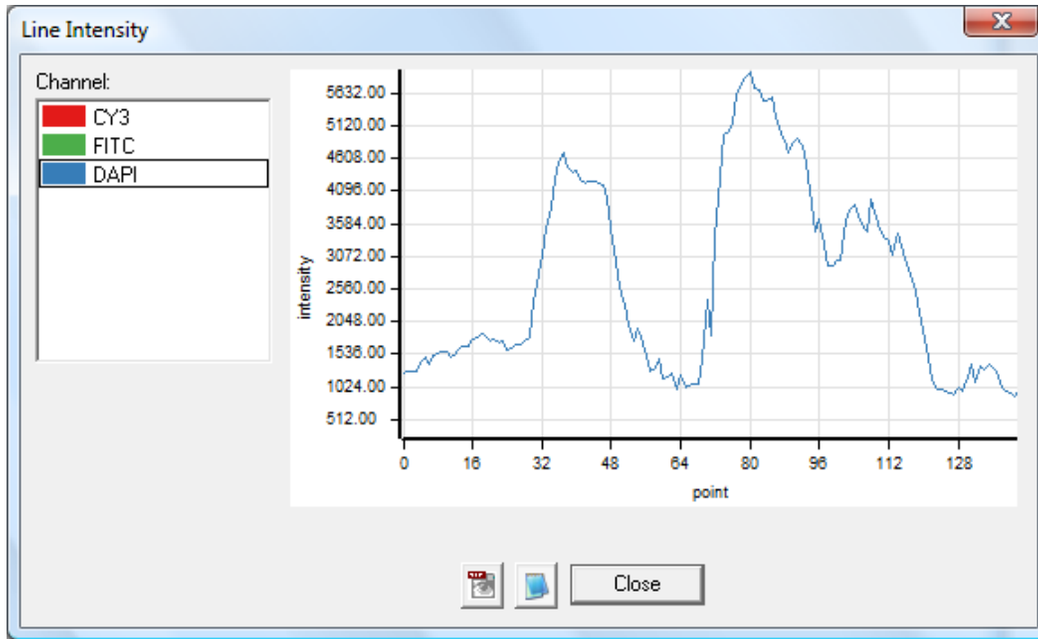
## 10.7 Displaying Graphs

SlideBook allows you to view intensity profiles of selected data. You may view line intensity or timelapse intensity profiles.

### 10.7.1 Viewing a Line Intensity Profile

SlideBook allows you to generate a line intensity profile using the **Statistics** menu. To view the intensity profile of a line, do the following:

1. From an open view, use the ruler tool to draw a line (see [Making Distance or Velocity Measurements](#) on page 187). You may draw this line in two or three dimensions.
2. Select **Statistics > Line Intensity**. A dialog box similar to the following will appear:



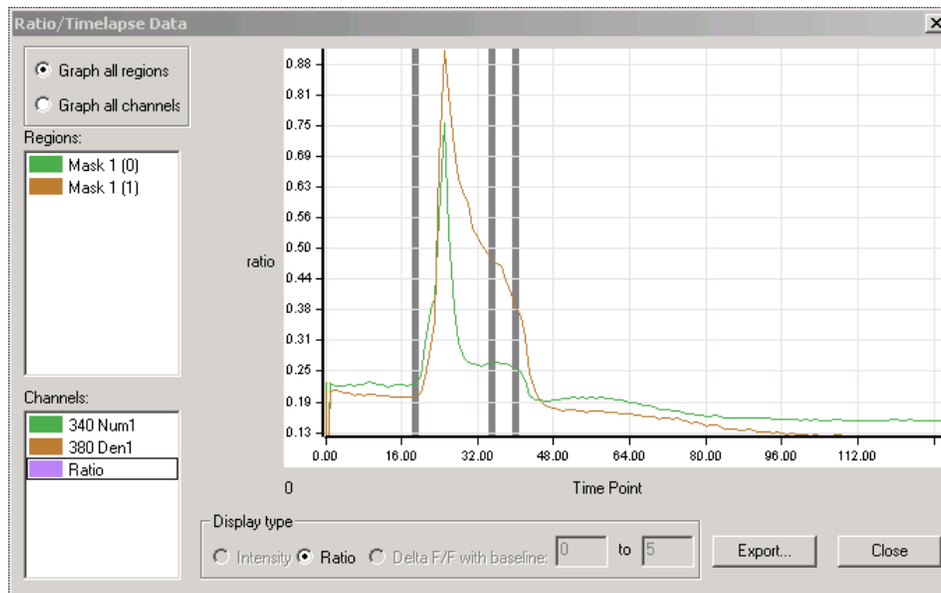
3. Select the channel that you would like to view from the **Channel** menu. The line intensity profile displays pixel distance on the *x* axis and intensity on the *y* axis.
4. To export the graph as a tiff, click on the tiff icon . You will be prompted to enter a filename. To export the data as a text file, click on the notepad icon . You will be prompted to enter a filename.
5. Click **Close** to close the dialog box.

### 10.7.2 Viewing Timelapse Intensity Profiles

SlideBook also allows you to view timelapse intensity profiles. There are four methods for viewing profiles.

#### 10.7.2.1 Method 1 – Using Masks

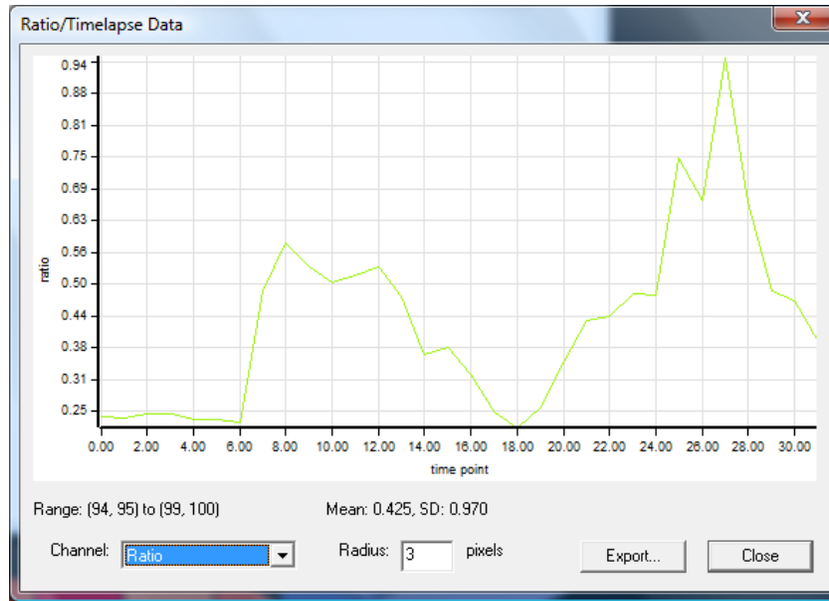
1. Select the regions that you would like to profile using masks (see above).
2. If there are multiple objects in the mask for which you would like to plot intensity profiles, generate objects by selecting **Mask > Define Objects in Mask** (see above for a discussion of objects).
3. Select **Statistics > Ratio/Timelapse Data** to generate a timelapse intensity graph:



4. Use the radio buttons in the upper left hand corner to either **Graph all regions** or **Graph all channels**.
  - **Graph all regions** - displays all regions for a single channel. If you have multiple floors, you may select the channel to display from the **Channels** list.
  - **Graph all Channels** – displays all channels for a single region. If you have multiple regions defined, you may select the region to display from the **Regions** list.
5. To export the data, click **Export**. You will be prompted to enter a filename.
6. Navigate to the desired file location, enter a filename, and choose **Save**. The intensity profile data will be saved as a text file that can be opened in programs such as Excel.
7. Click **Close** to close the dialog box.

### 10.7.2.2 Method 2 – Point and Click

From a main view, select the point selection tool from the tool menu . Press **Ctrl** and left click on any object in the image. An intensity-time profile will appear as shown below:

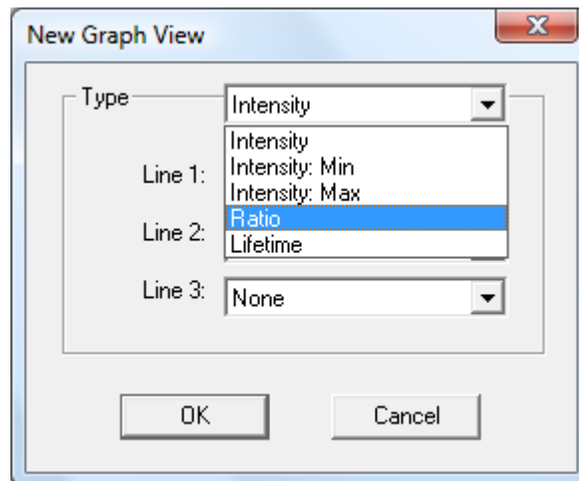


The intensity displayed is the average intensity of an area around the point where you clicked of the specified radius. You can alter the radius edit field, and choose to export the statistics as a tab-delimited text file (.txt).

### 10.7.2.3 Method 3 – Using ROIs

If you have drawn regions of interest to monitor timelapse capture, you can create graphs of these regions post-capture.

1. Open a view of the desired image.
2. Select any of the region tools or go to **Annotations > Regions** to display the ROIs.
3. Select **View > Graph**. The following dialog will appear:



4. Select the type of graph you wish to display and the channels that you wish to display.

5. Select which lines you would like to display (only applies when graphing intensities, not ratios).
6. Click **OK**.

The graphs can be explored as described in [Creating Graphs to Monitor Regions of Interest](#) on page 120.

**NOTE:** You can add ROIs by using any of the ROI tools to draw regions. Please see [Creating ROIs and Graphs to Monitor Regions of Interest](#) on page 118 for more information on creating and deleting ROIs.

## Appendix A: Smooth Curve Analysis (Kymograph)

Smooth curve analysis allows you to follow time-dependent lateral intensity changes along a static object. For example, an action potential moving along a neural spine could be displayed as an  $x$  vs.  $t$  image using smooth curve analysis. You could then extract velocity and displacement information from this image.

SlideBook's Smooth Curve Analysis is a mathematically rigorous and flexible operation with several adjustable parameters. These parameters are defined using the following figure.

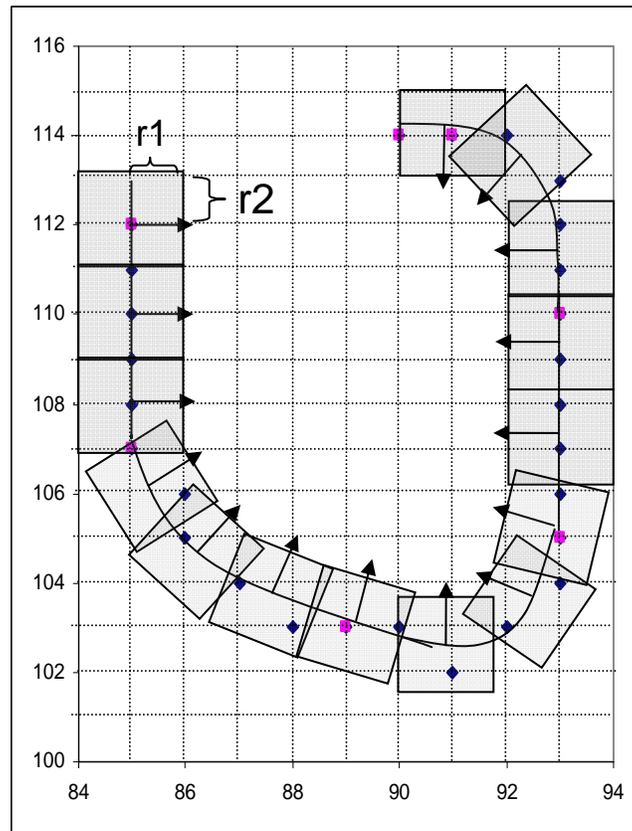


Figure A1: Representation of Smooth Curve Analysis (Kymograph)

The above figure is a schematic of a pixel map of a 2D image. When performing kymograph analysis, you first define a curve along a static object that will serve as a “backbone” for your analysis. Once this curve is defined, you may perform a smooth curve analysis. Smooth curve analysis uses the following parameters:

- **Spline parameters** – Knot Interval (discrete pixels), the default is set to five. A spline is fit between two “knots.” Multiple splines are used to fit a given curve. The knots are shown in pink in the figure above.
- **Measurement dimensions**

- **Radius Perpendicular to Curve (pixel units)** – shown as dimension **r2** in the figure above
- **Radius Parallel to Curve (pixel units)** – shown as dimension **r1** in the figure above
- **Increment Along Curve (pixel units)** – center to center distance between rectangular areas used to calculate smooth curve. In the above figure, the boxes are drawn every two pixels units along the curve. The resultant smooth curve analysis will have half the resolution of the original image.

Once these parameters are defined, the smooth curve algorithm averages the areas at each timepoint and creates a new image in two dimensions (distance in  $x$  vs. time in  $y$ ).

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