

Imaging System User's Manual (Revision B)



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Revision B of the DeltaVision Imaging System User's Manual. Part number 04-720160-000 Rev B.

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### **Other Reference Documents**

User instructions for *DeltaVision* include the following publications:

Document	Purpose	Available for
Online Help	Provides reference information for <i>softWoRx</i> and procedures that show how to use <i>softWoRx</i> tools	All <i>softWoRx</i> workstations
Product Notes	Provide examples and tips for using <i>softWoRx</i>	All <i>softWorRx</i> users (online at www.appliedprecision.com)
softWoRx Imaging Workstation User's Manual	Shows how to use the <i>softWoRx</i> software to process, visualize, and analyze data.	All <i>softWoRx</i> workstations

In addition to the above Applied Precision documentation, you will be provided with manufacturers' manuals for the microscope components. You may also be provided with manufacturers' manuals for cameras and other devices, depending on your system configuration. (Some manufacturers do not provide manuals for these devices.)

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# Preface

This manual shows how to use the *DeltaVision* system to acquire images. It also shows how to maintain the system.

- *About This Manual* describes the information in each chapter.
- Document Conventions explains the typography, notes, and other conventions used in this manual.
- *Contacting Applied Precision, Inc.* provides information about how to contact customer support.

# **About This Manual**

This manual provides instructions for scientists who are using *DeltaVision* to acquire data. It also includes instructions and references for maintaining the system.

The *Introduction* provides a brief summary of the *DeltaVision* system features. Safety warnings and guidelines are provided in Chapter 2.

Chapters 3 - 6 show how to use the system to acquire data.

- Chapter 3, *Getting Started*, describes how to turn the system on, acquire an image, and run an experiment macro.
- Chapter 4, Setting Up and Running Experiments, shows how to set up experiment macros for 3D sectioning, Time lapse, Multiple wavelengths, Paneling (for stitching), and Point visiting.
- Chapter 5, Acquiring Data From Live Specimens, provides information on how to use the DeltaVision system to collect images from live specimens.
- Chapter 6, *Data Collection Techniques*, describes how to determine the proper exposure time and provides guidelines for finding the areas of interest on a sample.

The remaining chapters provide information on how to maintain and configure the system.

- Chapter 7, Facility Requirements and Components, lists requirements and describes the DeltaVision system components.
- Chapter 8, *Changing Cameras and Filters*, describes how to replace cameras and how to install or replace filters.
- Chapter 9, *Maintenance*, shows how to change and align the xenon lamp, align the light path, clean the system, and change fuses.

The appendices include reference information and procedures for configuring the system.

# **Document Conventions**

To make the information provided in this manual as easy as possible for you to locate and use, the following conventions are observed.

### Lists

- Round bullets indicate options in procedures.
- 1. Numbered items are sequential steps for completing a procedure.
- Square bullets indicate items in a list.
- Arrows indicate single step procedures.

### Notes, Warnings, and Cautions

**Note** Indicates information about the previous paragraph or step in a procedure.

Important Indicates important or critical information about the previous paragraph or step in a procedure.

Ý Tip Indicates helpful advice.



**WARNING:** Indicates important information regarding potential injury.



WARNING: Indicates risk of explosion.



WARNING: Indicates risk of shock.



**CAUTION:** Indicates important information regarding potential damage to equipment or software.

### **User Interface Description Conventions**

Boldface indicates the names of buttons, menus, window options, and fields.

Initial Capitals indicate the names of windows and tabs.

ALL CAPITALS SAN SERIF indicates the name of a key on your keyboard or keypad, such as ENTER, DELETE, or STEP INCREASE.

Uniform width font indicates text to enter on a command line or in the GUI.

# **Contacting Applied Precision, Inc**

If you have questions about *DeltaVision*, first refer to this manual or consult the online Help system. If you don't find the information you need, contact us at one of the following addresses.

# **Customer Service Hotline**

Phone: 800-862-5166

email: servicehotline@api.com

Hours: 8:00 AM - 5:00 PM, Pacific Time, Monday - Friday

# Corporate Office

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Internet Address: www.api.com

# 1

# Introduction

This chapter provides an introduction to DeltaVision.

- What is DeltaVision? introduces the DeltaVision system and provides a history of its development.
- What Can You Use DeltaVision for? lists the supported imaging modes and summarizes the data acquisition options that are supported by DeltaVision.
- What Should You Know to Use DeltaVision? summarizes the background and experience required to run the system.

# What is DeltaVision?

The *DeltaVision* Imaging System can be used to collect and analyze threedimensional microscope images, acquired over long periods of time and on multiple samples. With the sophisticated *softWoRx* image analysis and modelbuilding software, the system is a comprehensive package for biological image data collection, interpretation, and display.



# History

The original restoration microscopes were designed and developed in the laboratories of Dr. John W. Sedat and Dr. David A. Agard at the University of California, San Francisco. Their first working system actively collected images as early as 1983. At that time, a *small* deconvolution (128x128x64) required overnight processing on a million dollar mainframe computer.

During the evolution of the UCSF microscope, it became clear that micropositioning was a critical part of the optical sectioning process. In particular, controlled movement of the focal plane relative to the specimen (the Z axis) was identified as a key to reliable deconvolution. To accomplish adequate Z scans, Dr. Sedat built a microscope stage using Applied Precision's Nanomotion<sup>™</sup> micropositioning technology.

Before long, Dr. Sedat contacted Applied Precision cofounder Ron Seubert for detailed information about Nanomover performance. The relationship between UCSF and Applied Precision grew steadily. Later, in 1993, Applied Precision licensed the image restoration technology from UCSF and began development of *DeltaVision*. Collaboration between Applied Precision and UCSF still continues, for the benefit of both parties. In October of 1993, Applied Precision shipped the first Applied Precision/UCSF hybrid to Michael Paddy at the University of Florida. In 1994, Applied Precision designed, built, and delivered a *DeltaVision* prototype to Paul Goodwin at the Fred Hutchinson Cancer Research Center, Seattle. In August 1994, the first commercial *DeltaVision* microscope was shipped to Bethe Scalettar at Lewis & Clark College, Portland, Oregon. All three of these systems are still active. The *DeltaVision* software has grown continuously since 1983, with contributions from scientific programmers, faculty, and graduate students at UCSF. Applied Precision's contribution to the software started in earnest in 1994.

The advances in computer and camera technology in the early 1990s resulted in the emergence of optical sectioning technology. For example, in 1993 Applied Precision's benchmark deconvolution (512 × 512 × 64) required 3 hours of processing time on a \$35,000 workstation. (The benchmark deconvolution is 16 times larger than the original deconvolutions performed in 1983.) Although only a few laboratories were able to afford \$35,000 for a computer, the lower cost warranted commercialization of a deconvolution microscope. In 1996, the same deconvolution required about a quarter of an hour with a \$14,000 workstation, and half an hour with an \$8000 machine. This thirty-fold increase in performance/price allowed a more widespread acceptance of deconvolution microscopy.

Advances in computing power and data storage have continued to benefit *DeltaVision* users. In 2003, a \$2500 workstation could perform the standard benchmark in less than four minutes, representing another twenty-fold increase in performance/price since 1996. The current workstations can now perform this benchmark in less than 30 seconds (under 20 seconds for *DeltaVision* and under 30 seconds for *personalDV*). Vast amounts of data generated by these experiments can be stored on local hard drives, by burning DVDs, or by transferring the data to other locations using high-speed network connections.

# What Can You Use Delta Vision for?

*DeltaVision* imaging systems use research grade microscopes to collect optical images in one of the following imaging modes:

- Fluorescence
- Brightfield
- Phase contrast\*

<sup>\*</sup> Phase contrast requires additional equipment that is available from Olympus distributors.

### Differential Interference Contrast (DIC)

Capability	Description
Digital Microscopy	Fluorescence Imaging.
	Also capable of Brightfield, Phase Contrast, and DIC imaging.
Automated Optical Sectioning, Time-lapse, Point visiting	Optical sectioning, filter changes, and shutters are coordinated by the controller.
Quantitative Processing	Image processing and 3-D reconstructions of multi-dimensional data files.
Image Display and Analysis	3-D reconstructions can be visualized, rotated, and enhanced.

The following table summarizes the capabilities of the *DeltaVision* system.

*DeltaVision* supports a wide array of imaging applications, including: Cytoskeletal Studies, RNAi experiments, Live Cell Imaging, Cell Cycling Studies, Protein Translocation, and Protein Pathway Analysis.

# **Standard Data Collection Options**

*DeltaVision* supports the following types of data acquisition:

### **3D Imaging**

To acquire "3D data," you can set up *DeltaVision* to acquire a series of images along the Z axis. The *softWoRx* workstation provides a sleek interface that allows you to control the optical sectioning through a specimen. Behind the scenes, macro language provides automated computer control of sample position, optical filters, and shutters. After image data acquisition, a series of image processing algorithms improve image resolution. Three-dimensional information can be reconstructed and then visualized in a variety of ways that allow quantitative measurement and analysis.

### **Multiple Wavelengths**

*DeltaVision* can sequentially acquire images through multiple wavelengths and combine them into a single image file.

### **Panel Collection**

Panel collection acquires a series of images with adjacent fields of view. You can stitch these images together to create images that are much larger than a single field of view. This is especially useful when you want to collect data at a high magnification over a large area.

### Time-Lapse

You can run macros to acquire time-lapse images and use the data to create timelapse movies. This is especially useful for studies of live samples or for experiments that use lasers.

### **Point Visiting**

Point Visiting allows you to acquire data from several areas of interest during a single experiment. You can select which points to monitor on your sample and save them in a list that contains the exact stage coordinates of each point. When you run the experiment, *DeltaVision* reads the coordinates for the points in the list, moves the stage to each point, and captures an image. This process is repeated at specified time-lapse intervals. For live specimens, this significantly improves the lab efficiency of experiments by allowing you to monitor multiple points of interest in a single session.

### Autofocus

You can use Autofocus to automatically focus *DeltaVision* when you are viewing a sample or when you are running an experiment.

After you have found focus and marked a point of interest, the <u>optional</u> UltimateFocus<sup>™</sup> feature adds the ability to maintain focus during an experiment. The UltimateFocus Module uses an infrared laser that follows the illumination path and bounces off the cover slip/sample interface. The reflected beam is evaluated and the software returns an offset to the Z motor for automatic stage adjustment to maintain focus.

### Köhler and Critical Illumination

You can easily switch between Critical and Köhler Illumination. Köhler Illumination provides very even specimen illumination across the field of view. You will typically use Critical Illumination for most of your data collection. Critical Illumination only illuminates the area detected by the CCD. It therefore helps to preserve the biology and associated dyes outside the CCD's field of view. Köhler Illumination spreads the intensity of the light over the entire field of view of the oculars. The higher intensity light of Critical Illumination is also useful for low abundant probes that require more light.

### Live Cell Imaging

You can set up a controlled environment to acquire data from live specimens. *DeltaVision* supports an optional environmental chamber that you can use to control temperature and inject carbon dioxide.

*DeltaVision* also includes software that is specially designed to acquire data from live specimens:

- Cell tracking is used for point visiting time-lapse experiments. It automatically changes the coordinates of a point to follow a cell as it moves.
- Real-Time Z Sweep Acquisition (or Optical Axis Integration) allows you to quickly acquire 2D projections of specimens.
- Real-Time Deconvolution provides previews of deconvolved images as they are acquired.
- Reference Imaging is useful for acquiring reference images that can be used for Differential Interference Contrast (DIC) and other techniques.

# Laser Photo-bleaching and Photo-activation

If your system is equipped with the optional X4 Laser Module, you can use *DeltaVision* to run and analyze laser photo-bleaching and photo-activation experiments. The X4 Laser Module supports up to four lasers and provides software to control and to analyze the data that is obtained from these experiments.

# **TIRF (Total Internal Reflection Fluorescence)**

If your system has the optional X4 Laser Module and API FI, you can use *DeltaVision* to run and analyze multiline *TIRF* experiments. The TIRF technique is used in a number of research applications, such as cellular protein and vesicle trafficking, focal cellular adhesions, single biomolecule dynamics, studies in neuroscience, and cell-to-cell communications. The ability to excite molecules on the surface of a specimen while eliminating the fluorescence from the depth of the sample makes TIRF a valuable tool for examining cell surface structure and protein dynamics.

For more information on the TIRF technique, see the section "*X4 Module and TIRF*" on Page 191.

# What Should You Know to Use Delta Vision?

This document assumes that you are familiar with the basics of microscopy. Correct operation of the microscope is fundamental to obtaining quality images with *DeltaVision*. In addition, an understanding of image processing basics will help you use the system to its full potential. To manage the computer systems, some familiarity with Linux workstations and Windows-based personal computers is helpful.

We have taken care to ensure that *DeltaVision* is straightforward to use, reliable, and complete. Please report errors and problems with *DeltaVision* to Applied Precision using the e-mail address servicehotline@api.com, or alternatively, use the problem report form in *Appendix B: Troubleshooting*.

# 2



This chapter covers safety and damage prevention issues to be aware of when using a *DeltaVision* imaging system. The precautions detailed in this chapter must be carefully observed to prevent possible personal danger:

Note All laser-related safety issues, including proper labeling, are discussed in *Appendix F: Lasers and Safety Issues.* 

- *UV Exposure* discusses potential for UV exposure from the xenon arc lamp.
- Bright Light Exposure warns about bright light exposure from the transmitted light source installed in the microscope.
- Laser Hazards explains that a *DeltaVision* system with lasers installed is considered a Class 3B device and warns that visible and invisible radiation from the installed lasers can exit the device at the same time and are powerful enough to potentially cause damage to the human eye.
- *Burn* provides guidelines for avoiding burns from the xenon arc lamp. (The arc lamp reaches very high temperatures.)

- Shock includes warnings about potential shock hazards. (Hazardous voltages are present even when the system is disconnected from the AC main power outlet.)
- Damage Prevention describes actions that can damage the system.
- *Warning Labels* describes the system labels.

Additional safety guidelines for maintenance and alignment are detailed in *Chapter 9: Maintenance*.

For laser safety considerations and laser safety labeling information, see *Appendix F: Lasers and Safety Issues*.

# **UV** Exposure

Since the excitation illumination module (xenon arc lamp or InsightSSI) emits ultraviolet (UV) light, there is a danger of exposing your eyes and skin. Loss of eyesight could occur if unfiltered light from the xenon arc lamp reaches your eyes.

### To prevent UV exposure:

- Open the shutter only when an excitation filter is engaged.
- Do not open the xenon arc lamp housing during operation. See *Chapter 9, Maintenance* for detailed instructions on changing the xenon arc lamp bulb.

# **Bright Light Exposure**

While the transmitted light source installed in the microscope does not present possible UV exposure, it could cause discomfort under certain conditions.

- You must be aware of the eyepiece filter wheel when viewing a specimen through the microscope oculars. Make sure that the proper filters are in place so that your eyes are not suddenly exposed to a bright flash of light when the transmitted light shutter is opened.
- Do not look through the eyepiece while switching filters. Your eyes can be exposed to unfiltered light during the filter transition.

# Laser Hazards

The optional laser modules available for *DeltaVision* are considered Class 3B devices. The power level is high enough to cause damage to the human eye instantaneously and possible damage to the skin.

The optional *UltimateFocus*<sup>™</sup> Module complies with CFR 1040.10 and 1040.11 except for deviations pursuant to Laser Notice No. 50, dated June 24, 2007 IEC 60825-1, 2007-03.

OSHA regulations require (via ANSI Z136.1) and IEC 60825-1 recommends that a Laser Safety Officer be identified who will be responsible for the safe use of Class 3B lasers. This includes training users, installing all necessary warnings and controls in the laser area, and other duties.



### WARNING:

Given the inherent exposure possible with an inverted frame microscope stand, users of the system must be trained in laser safety before using this instrument. Contact your lab administrator for information about Laser Safety training at your institution. Training information is also available online at: www.kenteklaserstore.com The International Electrotechnical Commission (IEC) and the FDA recommend

that Class 3B and Class 4 lasers be used only in restricted areas.

Please see Appendix F, Lasers and Safety Issues on Page 327 for detailed laser safety information.

### Burn

The xenon arc lamp reaches a very high temperature when lit. Never touch the housing during operation. Never remove the housing during operation or before allowing it to cool completely. Carefully follow the directions found in *Chapter 9*, *Maintenance,* for changing the lamp.

# Shock

Hazardous voltages are present even when the system is disconnected from the AC main power outlet.

To replace the xenon arc lamp, follow the instructions found in *Chapter 9*: Maintenance. Refer also to the following section, "Xenon Lamp Safety," for additional related safety issues. No other system components contain userserviceable parts and do not warrant disassembly.

# **Xenon Lamp Safety**

Xenon lamps are under high pressure and emit high levels of radiation. Proper handling procedures and safety precautions should be observed to assure the safety of the users of this product. Only operate this lamp within the recommended operating specifications as detailed in this manual. Refer to *"Replacing the Xenon Lamp"* in Chapter 9 of this manual for details on the proper procedures for lamp replacement.

# **Explosion**

These xenon lamps are under high pressure. Use of face shields or safety glasses during handling is recommended. Avoid applying excessive shock or stress to the lamp during handling.

# **High Voltage**

The ignition voltage of the xenon lamp presents a very high voltage hazard. Do not touch the lamp during operation. To avoid the risk of electrical shock, the input power should be disconnected prior to servicing the lamp.

# UV, Visible, and IR Radiation

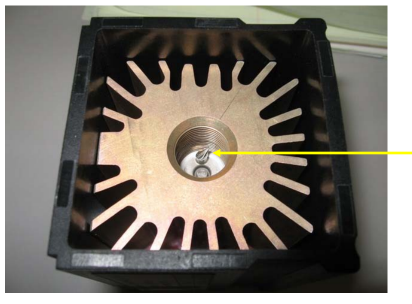
Xenon lamps emit high levels of radiation that can cause severe skin burns and permanent eye damage. Avoid direct exposure to the emitted or reflected beams.

# Thermal Hazards

Xenon lamps can get very hot during and after operation (up to several hundred degrees C). To avoid potential for serious burns, do not touch the lamp during operation. Do not touch the lamp after operation until the lamp has adequately cooled.

# Disposal

It is recommended that the xenon lamp's internal pressure be relieved prior to disposal. This is accomplished by squeezing off the tip (see illustration below) with needle-nosed pliers until the pressurized gas escapes.



Pinch off the tip with needle-nosed pliers to relieve pressurized gas before disposal.

Relieve the xenon lamp's internal pressure before disposal! Pressurized xenon lamps should not be incinerated, but disposed of in a landfill.

# **Damage Prevention**

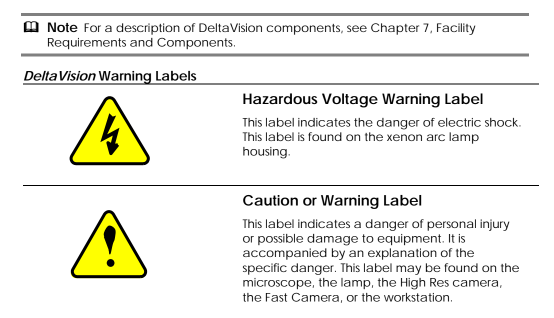
The following actions could damage the system:

- Moving the stage to the home position with the objective up could break or scratch the objective. The stage could be driven into the objective and may potentially scratch the top lens or compress the lens housing, causing a leak, crack, or lens misalignment.
- Disconnecting cables before the system is completely shut down may damage one or more of the electronic circuit boards.
- Disconnecting cables to the camera when the power is on may damage the camera.
- Touching optical filters or the polychroic beam splitter contaminates them with oil and can lead to premature failure or poor image quality. For cleaning information, see *Chapter 9: Maintenance*.
- Bending the fiber optic cable into a coil with a diameter less than 24" will damage the cable.
- Using improperly rated replacement fuses can create a fire hazard and may result in damage to components. Use only the fuse types listed on the component or in the manual.

 Leaving a camera out of the tray when it is not in use presents an opportunity for the camera to fall to the floor and break. Always place the camera in the camera tray when it is not in use.

# Warning Labels

Warning labels have been applied to the components of the system that pose a potential hazard to the user. The labels have been duplicated here and carefully explained. Please read this section carefully.



See *Appendix F, Lasers and Safety Issues* on Page 327 for laser-specific safety warning labels.

# 3

# **Getting Started**

This chapter shows how to get started with *DeltaVision*.

- *Before you start* includes a checklist of things that you need to have before you can acquire images for your sample.
- *Getting Familiar with DeltaVision* describes the key controls that you will use to direct the light path, focus, and select filters. It also introduces the keypad and joystick.
- *Turning DeltaVision On* shows how to turn on the system.
- *Acquiring an Image* shows how to place the slide and the objective, find the sample, and acquire an image.
- Saving Image Data describes how to set up a personal data folder and how to save an image file.
- *Turning DeltaVision Off* shows how to turn off the system.

# Before you start

Before you start, make sure that you:

- Select the proper oil for your objective and specimen. The immersion oil kit includes 18 oils (6 for *personalDV*) with refractive indexes that range from 1.500 to 1.534, in increments of 0.002. (For personalDV, the kit includes 6 oils that range from 1.512 to 1.522.) To calculate the best refractive index for your application, follow the instructions in *Appendix A: The Immersion Oil Kit* on Page 247. If you are working at standard temperature and pressure, the oil with a 1.516 refractive index is generally a good place to start. For work at 37 degrees C, use the oil with a 1.520 refractive index.
- Know your login ID and password for the workstation.
- Prepare your sample using the recommended practices that are documented in the following references. (If you are untrained in sample preparation, consider attending a course like the one shown below.)

### **Books on Cell Biology Methods**

<u>Current Protocols in Cell Biology</u>. Bonifacino, Dasso, Lippincott-Schwartz, Harford, Yamada ed. Wiley Press, http://www.wiley.com/legacy/cp/cpcb/

<u>Cells, A Laboratory Manual</u>. Spector, Goldman, Leinwand ed. Cold Spring Harbor Press, 1998.

<u>Video Microscopy: The Fundamentals (2nd Edition)</u>. Inoue and Spring, Plenum Press, 1997.

<u>Digital Microscopy (3rd Edition</u>), Methods in Cell Biology Vol. 81. Sluder and Wolf, Academic Press, 2007.

### Papers on Sample Preparation

Rines DR, He X, Sorger PK.Quantitative microscopy of green fluorescent proteinlabeled yeast. Methods Enzymol. 2002;351:16-34.

Hutchins JR, Moore WJ, Hood FE, Wilson JS, Andrews PD, Swedlow JR, Clarke PR. Phosphorylation regulates the dynamic interaction of RCC1 with chromosomes during mitosis. Curr Biol. 2004 Jun 22;14(12):1099-104.

### Courses in Microscopy and Cellular Imaging

In Situ Hybridization, Immunocytochemistry, and Live-cell Imaging. Dernberg, Hu, and Murray Course Directors, Cold Spring Harbor, October (annual).

Analytical and Quantitative Light Microscopy, Sluder and Wolf Course Directors, Marine Biological Laboratory, Woods Hole, MA, May (annual).

# Getting Familiar with DeltaVision

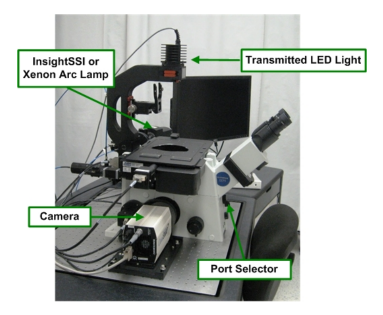
Before you acquire an image, become familiar with the key *DeltaVision* controls for:

- Controlling the Light Path
- Focusing
- Choosing Filters
- Using the Keypad and Joystick

Most of the manual controls for controlling the light path, focusing, and choosing filters are similar to those that you will find on any microscope. Additional controls for moving the stage and acquiring images are provided by the keypad and joystick.

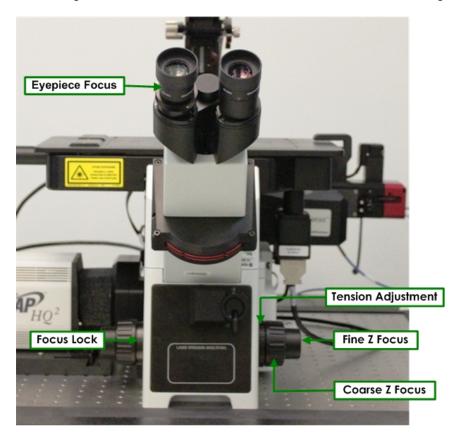
# Controlling the Light path

*DeltaVision* provides a transmitted LED light and either a xenon arc lamp or a solid state illumination source. The transmitted light works the same as the light source for a traditional microscope, with the light path directed on the specimen from above. Both the xenon arc lamp and the InsightSSI (solid state illumination source) provide excitation light directed through the back of the microscope and focused on the specimen from below. You can use the Port Selector to direct the light path either to the **Eyepiece** or to the **Camera**.



# Focusing

There are three manual focus controls and a Focus Lock on the *DeltaVision* microscope. These controls are similar to those on other microscope systems.



**Eyepiece Focus** – Use the Eyepiece Focus (on the left ocular) to focus the eyepiece.

**Focus Lock** – Use the Focus Lock to set a maximum height for Z focus. This can keep users from hitting the sample with the objective.

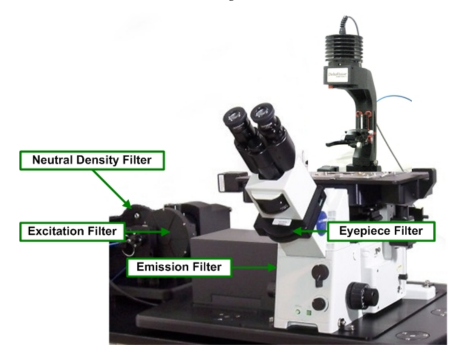
**Tension Adjustment** – Use the tension adjustment on the focus knob to help keep the knob from moving inadvertently.

**Fine Z Focus** – Use the Fine Z Focus knob to move the objective in very small increments. It is used to focus on the focal plane.

**Coarse Z Focus** – Use the Coarse Z Focus knob to move the objective in large increments. It is typically used to lower the objective when the system is initialized or to move the objective up to the slide until the oil is touching the slide.

# **Choosing Filters**

Choosing and controlling filters is a key for any fluorescent probe experiment. When a fluorescent probe is excited by a specific wavelength, it emits light at another wavelength. Choosing the correct filters for the dyes in your sample allows you to obtain a complete set of data specifically from your probe without interference from other wavelengths.



*DeltaVision* provides several different types of filters for controlling the fluorescent light path:

- **Neutral Density** filters (often denoted as "%T") reduce the amount of light that illuminates your sample when you are using fluorescence. *DeltaVision* provides six neutral density filters that block from 0 to 99.9% of all light.
- **Excitation** filters block all but a single band of wavelengths, providing a specified range of light to excite the fluorescent probes in the sample.

Note With the InsightSSI solid state illumination module, the excitation filters are included as part of the illumination module. Also, no neutral density filters are necessary when using InsightSSI as your broadband light source.

- **Polychromatic Beam Splitter** reflects the excitation wavelengths to the sample and transmits the emission wavelengths from the sample. *DeltaVision* ships with a standard polychromatic beam splitter for DAPI, FITC, TRITC, and Cy5. Other beam splitters ship with optional live-cell sets.
- **Emission** filters allow only a narrow band of light from the excited probe to reach the camera.

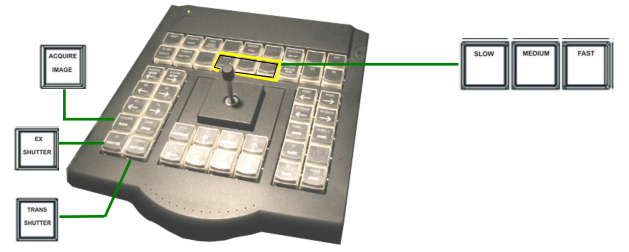
• **Eyepiece** filters are emission filters that allow only a narrow band of light from the excited probe to reach the eyepiece and your eyes.

These filters are arranged in sets that are associated with specific dyes. (For example, a dye such as DAPI is typically used with a DAPI Excitation filter, a DAPI Emission Filter, and a DAPI Eyepiece Filter.)

You can choose filter sets manually by rotating the eyepiece filter wheel. The filter sets are synchronized so that when you change an eyepiece filter, the neutral density filter, excitation filter, and emission filter automatically change.

# Using the Keypad and Joystick

The keypad and joystick are used to move the stage, open shutters, acquire images, and control other acquisition options. Key controls are shown below.



Key controls on the Keypad

### Acquire Image

With the microscope set to Camera Mode, this button acquires an image and displays it on the monitor. Use this key when you are scanning through your sample and want to get a quick look at the specimen on the monitor. (Note that Resolve3D must be running at the time.)

### **Ex Shutter**

Opens or closes the Excitation (i.e., Fluorescence) shutter. You will use this control frequently to open and close the shutter. Because the shutter is designed to protect your eyes from exposure to ultraviolet light, it automatically closes each time that the eyepiece filter wheel is moved. It must be reopened with the EX SHUTTER button.

### **Transmitted Light Source**

Toggles the transmitted light LED between off and on. (Subsequent to changing the transmitted light source from halogen to LED, an actual shutter is no longer necessary.)

### Slow, Medium, and Fast

Control the speed that the stage is moved using the joystick. It's usually best to start with Medium.

### The Joystick

Controls stage movement. Use the joystick to move the stage in the direction that you point with the joystick (for example, moving the joystick up moves the stage away from you, moving joystick left moves the stage to the left, and so on).

# Turning DeltaVision On

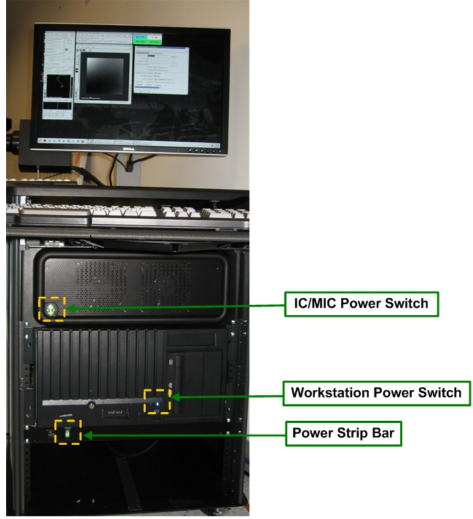
Use the following instructions to turn the system on for day-to-day use. For instructions that show how to turn on *DeltaVision* after a system shutdown, see *Shutting Down and Starting the System* on Page 237.

### To turn on DeltaVision:

1. Turn on the power strip bar.

Note For *personalDV*, there is no switch on the power strip bar. Begin the powerup process with Step 2.

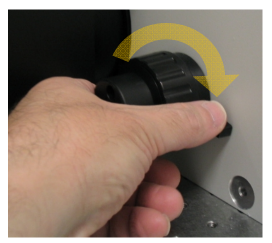
- 2. Turn on any additional equipment such as cameras or the heater for the Environmental Chamber.
- 3. Turn on the IC/MIC.



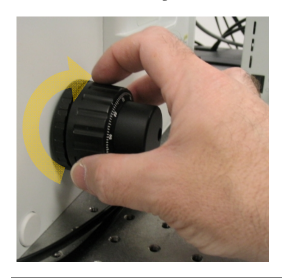
Main Power Switches (DeltaVision Cabinet)

- 4. If the monitor is off, turn it on.
- 5. If the Workstation is off, turn it on and wait for it to boot up.
- 6. Log on to the Workstation.
- 7. Remove any slides from the stage.
- 8. On the desktop, double-click the **Start** *softWoRx* icon to open *softWoRx*.
- 9. On the *softWoRx* menu, choose **File** | **Acquire** (**Resolve3D**).

10. Release the Focus Lock by turning it clockwise (when facing the lock) until it is loose.



11. Lower the objective by turning the Coarse Z Focus knob away from you (clockwise) when facing the knob.

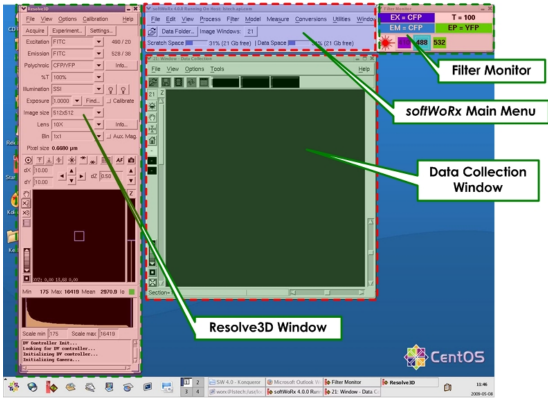




**CAUTION:** Always lower the objective before you initialize the system to prevent damage to the objective lens.

12. A prompt is displayed reminding you to lower the objective before continuing. After you lower the objective, select **Initialize** to initialize the system.

The Resolve3D window, the Data Collection Window, and the Filter Monitor window open on the desktop.



#### softWoRx Desktop Display

The Resolve3D window includes acquisition parameters and controls for moving the stage, the Data Collection window displays images as they are acquired, and the Filter Monitor displays the filters currently selected.

■ Note You will use the Resolve3D window throughout the data collection process. For more about this window, see *The Resolve 3D Window* on Page 278.

## Acquiring and Saving Data

To acquire an image of your sample:

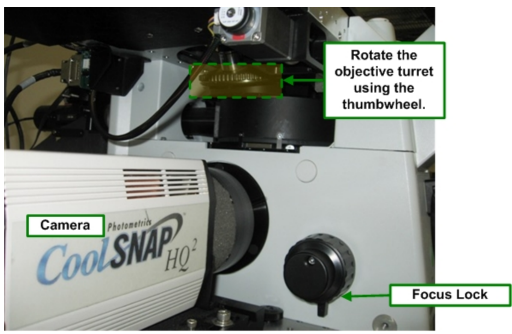
- Set up *DeltaVision* by placing the sample slide and selecting the appropriate filters.
- Find the sample so that it is visible in the eyepieces.
- Set the *DeltaVision* microscope to Camera mode.
- Acquire an image with the camera.

## Setting Up the Sample for Image Acquisition

Setting up *DeltaVision* for imaging includes placing the sample on the stage and selecting the appropriate filter set for the fluorescent probe used to label your sample.

## To prepare for image acquisition:

- Note This procedure assumes you are using an oil immersion lens. If you're using a water, air, or silicone lens, you may need to modify this process accordingly.
- 1. If your system includes a xenon lamp for its primary light source, turn it on now.
- 60
- 2. Rotate the Coarse Focus knob so that the top of the knob rotates away from you (while you are sitting in front of the microscope) to move the objective all the way down.
- 3. Rotate the objective turret (using the thumbwheel located at the leftunderside of the stage) to select an objective.



Adjust the objective turret beneath the left side of the stage to select an objective.

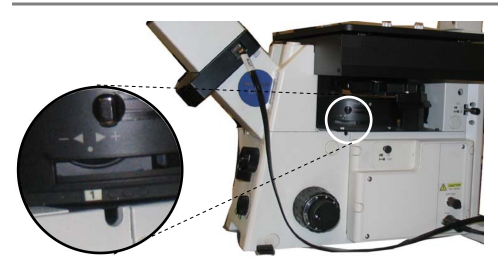
4. In the Resolve3D Lens list, select the matching objective.

Acquire	Experiment Set	tings	
Excitation	FITC	490 / 20	Bulb icons appea
Emission	FITC	▼ 528 / 38	only if you have xenon arc lamp(s
Polychroic	CFP/YFP	▼ Info	installed. An icon i
%T	100%	-	displayed for each
Illumination	LASER	88	
Exposure	1.0000 <b>T</b> Find.	🔲 Calibrate	Soloottho
Image size	512x512	-	Select the corresponding
Lens	10X	I Info	- objective in the
Bin	1x1 ·	Aux. Mag.	Resolve3D <u>Lens</u> field.

**CAUTION:** Make sure that the correct objective is selected in the Resolve3D Lens list.

5. Rotate the polychroic filter wheel (located on the right-underside of the stage) to select the appropriate dichroic mirror.

**Note** If your *DeltaVision* system is equipped with a motorized polychroic turret, you will control the turret's position using the *softWoRx* software.



/!\

6. In the Resolve3D Window, click the **Info** button next to the **Lens** field to open the Lens Information window.

Lens Information				- X	
<u>File L</u> ens				Help	
Lens ID 10105		Name	Olympus 10X/0.40, D Plan Apo UV, phase, fluor fr	-	
Manufacturer [Olymp	us				
Manufacturer P/N [1-LP3	31	APLLC P/N	Unknown		
Magnification	[10.0		NA (0.40		Refractive
Working Distance (µm)	3100.0	F	ocal Length (μm) [15690.0		Index
Standard Refractive Index	[1.000 R	ecommended	Refractive Index [1.000		
Optical Conditions					1
Distance From Coverslip	o Specimen (	um) [0.0	Coverslip Thickness (µm) [170.0		Optical
	Temperature	(C) 25.0			Conditions
Specimer	Refractive Ind	dex [1.470	1.470 - Pure Glycerol	-	
Resolution Calculation	IS				
Resolution Ratio (Z/>	Y) 9.2		Depth of Field (µm) 3.16		
Maximum XY Pixel Size (µ	m) 0.48	Recommende	ed Z Step Size (µm) 4.35		

#### Lens Information Window

- 7. In the **Optical Conditions** fields, enter the conditions for the sample.
- 8. If you're using an oil emersion lens, note the displayed value in the **Recommended Refractive Index** field and apply an oil with that refractive index. See Page 30 for details.



**CAUTION:** Before you start each data collection session, calculate the oil. Your oil selection should be the same if the sample and conditions are the same.



CAUTION: Do not touch the glass dropper to the objective.

- 9. Mount the slide on the stage with the coverslip down.
- 10. If installed, use the Adjustment knob on the Repeatable Slide Holder and the joystick on the Keypad to center the coverslip over the objective.

Tip If you have the Repeatable Slide Holder, you can record the position of your slide. This is useful when you are performing a Point Visiting experiment and you need to remove the slide before you are finished with the experiment (see Page 176).

- 11. Rotate the Coarse Focus knob toward you to move the objective up until the coverslip is just in contact with the oil. From this point on, use only the fine focus knob to raise and lower the objective.
  - Note If you DeltaVision system includes the UltimateFocus Module, you can use alternate methods for finding the focal plane.
- 12. Rotate the eyepiece filter wheel (below the oculars on the scope) to select the filter for the probe that you used to stain your sample. If your sample has more than one probe, select the one with the brightest fluorescence. The selected filter is displayed on the Filter Monitor window.



#### Filter Monitor Window

The filter names are displayed in the Filter Monitor window on the right side of the workstation screen. As you rotate the eyepiece filter wheel, the filter name next to **EP** (eyepiece) changes and the **EM** and **EX** (emission and excitation) filters change automatically to match.

The displayed colors match the wavelengths of the installed filters. If an X4 Laser Module is installed, the Filter Monitor also displays the wavelengths of the lasers that are available on your system.

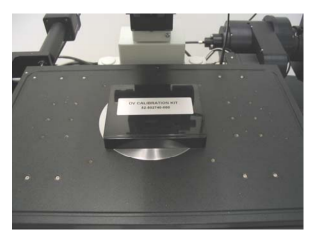
To determine the probes to use with each filter and the EX and EM wavelengths for each, refer to *Standard Fluorescence Filters* in Appendix D: Reference Information (Page 266).

## Finding the Sample

To set the focal plane for imaging, you will need to find the sample in the eyepieces and then focus on the sample.

## To find the sample:

- 1. With the sample on the stage and the filters selected (shown in the previous procedure), set the Port Selector to **Eyepiece**.
- 2. Turn <u>off</u> the lights in the lab. If you cannot turn off the lights, place a box over the sample to reduce the amount of ambient light.



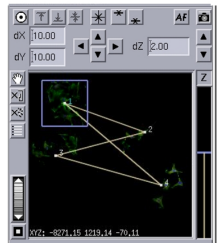
3. Open the Excitation shutter by pressing the EX SHUTTER button in the lower left corner of the keypad.

You should see light through the objective. The light on the stage should be the same color as the Excitation filter that you selected. (For example, if you selected DAPI, the light should be very deep violet. It may be hard to see.) Be sure the eyepiece filter position matches the desired excitation filter position.

- **Note** The Excitation shutter is designed to protect your eyes. Each time the eyepiece filter wheel position is changed, the shutter automatically closes and must be reopened with the EX SHUTTER button.
- 4. Focus to find the focal plane. Turn the Fine Focus knob toward you to slowly raise the objective until you see a cloud of emission color in the eyepiece. Continue to slowly raise the objective until the sample image is sharp and clear.
- 5. Use the joystick to move the stage around. Change the speed of the movement with the SLOW, MEDIUM, and FAST keys on the Keypad. When you find a region of interest, place it in the middle of the field of view.

On the Stage View, note the stage trails that show where you have moved the stage in XY.

EX SHUTTER



Stage Trails in the Stage View Window

## Acquiring an Image

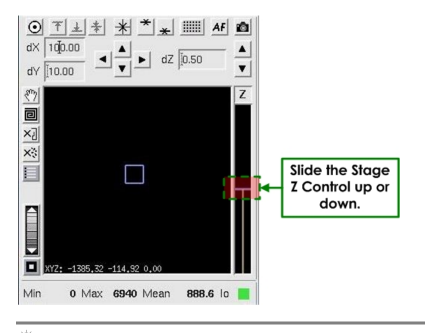
To acquire a *DeltaVision* image, you'll need to set the port selector to **Camera** and work with the images that are displayed in the Data Collection window until you are satisfied. Then you can create and run experiments, as described in *Creating and Running an Experiment Macro* on Page 52.

## To acquire an image:

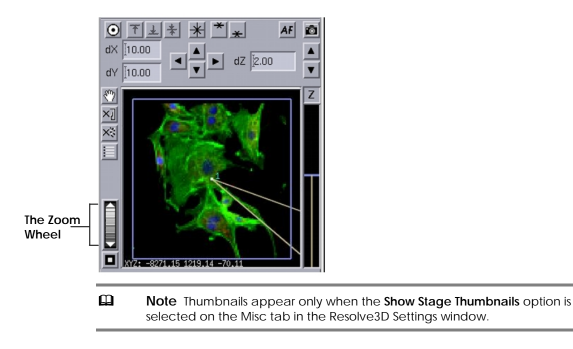
- 1. On the keypad, press **EX Shutter** to close the shutter.
- 2. Switch the Port Selector to **Camera**.
- 3. In the Resolve3D window **Exposure** field, enter an exposure time (in seconds). A good starting exposure time is 0.1 second for fixed cell and 0.01 seconds for live cell applications (see *Finding Exposure Time* on Page 104 for details).

Tip You can also click the Find button on the Resolve3D window to find a good exposure time. Use Find carefully. Overuse of this option can photobleach the specimen. Find is not recommended for use with live cell imaging.

- 4. Click 🏙 to acquire an image.
  - Tip You can also acquire an image by pressing the ACQUIRE IMAGE button on the keypad, right-clicking on the stage view, and selecting Acquire; or by choosing File | Acquire Image on the Resolve3D menu.
- 5. To focus, use the mouse to slide the Stage Z Control bar up or down.



- You can also use software AutoFocus to focus the sample as follows: On the Resolve3D window, click **AF** to auto focus the sample. If your image is far out of focus, you may need to click **AF** more than once. Software AutoFocus is not recommended for live cell applications.
- 6. To center the image, click (above the Stage View window) and then click on the object that you want to center in the Image window.
- 7. To enlarge the thumbnail image displayed in the Stage View, drag the mouse down over the zoom wheel.



8. To clear the thumbnail image, click **Clear Thumbnails** button next to the Stage View window.

## Saving Image Data

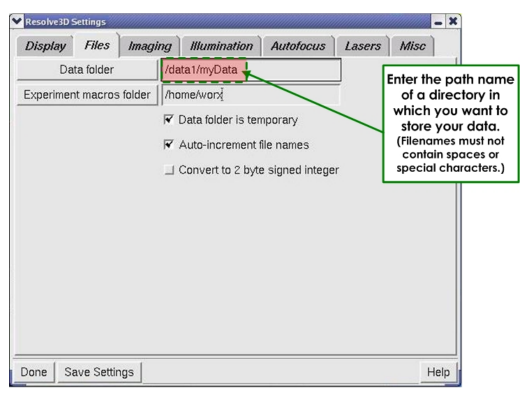
After you acquire an image, use these instructions to set up a personal data folder and use the Snapshot utility to acquire a single multi-channel image.

## Setting up a Personal Data Folder

You can set up a personal data folder in which to save your images. After you create and save this folder, *DeltaVision* automatically uses it as the data folder when you log on to the system.

## To set up a personal data folder:

1. At the top of the Resolve3D window, click the **Settings** button to open the **Resolve3D Settings** window. Then click the **Files** tab.



2. In the **Data folder** field, enter the directory (e.g., /data1/myData) in which you want to save your images. To create a new folder, type the name of the folder after /data1 (e.g., /data1/myNewData) and select **Save Settings**.

- Important Store all files under the /data1 directory unless you are instructed to do otherwise by your system administrator. If you store files outside the /data1 directory on the workstation, you can render the system unusable.
- Tip If you are using an existing folder for your data, you can drag and drop the folder from the Linux File Manager to the Data folder field.
- 3. To reset the default Data folder when you log out, select the **Data folder is temporary** option.

## Saving a Single-Section, Multi-Channel Image

Sometimes it is desirable to acquire a single-section, multi-channel image. The Snapshot utility helps you quickly create a 2-D, multi-wavelength image without having to run a full experiment.

#### To acquire a Single-section, Multi-channel image:

1. From the Resolve3D window, choose **File** | **Snapshot** to open the Resolve3D Snapshot window.

Y Resolve	e3D Snap	shot	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	<u> </u>
Image F	ile Nam	e hotCell001		
🔽 Deco	involve	(2D) Images		
🔽 Do O	Al Sca	n Thicknes	s (5.00	
Select C	hannel	5		
	<b>R</b> 1	CFP		
	<b>R</b> 2	YFP		
	🛛 З	EGFP	•	
	□ 4	mCherry	*	
	⊒ 5	CFPLaser	*	
Done	Do It	1		Help

#### Resolve3D Snapshot Window

- Tip Alternatively, you can open the Resolve3D Snapshot window by rightclicking on the Resolve3D window and choosing Snapshot from the shortcut menu.
- 2. In the **Image File Name** field, enter a file name. Then select which channels to save. (Snapshot uses the exposure conditions that were last used in the Resolve3D window for those channels.)

- Note *softWoRx* adds the \_R3D.dv extension to the file name. If you enter a file name without a directory path, the file will be located in the current data folder. You can specify to place the file in another directory by including the path in the file name (e.g., /data1/myfolder/myfile).
- 3. Click Do It.

An image window opens and displays the new image. The image is saved as a *DeltaVision* file that you can open in *softWoRx*.

## Turning Delta Vision Off

Use the following instructions to turn off *DeltaVision* on a daily basis. For instructions that show how to shut down the system, see *Shutting Down and Starting the System* on Page 237.

#### To turn off DeltaVision:

If your light source is the xenon arc lamp, turn off the lamp using the bulb
 icon on the Resolve3D window. Clicking the icon will switch it to the

off 🔋 state.

Note If you are using InsightSSI for your main light source, skip this step and continue to Step 2.

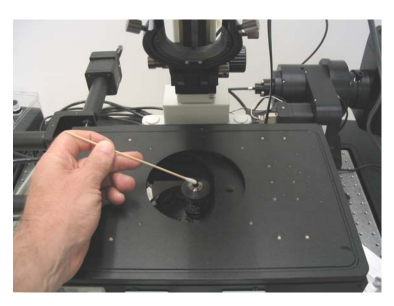
- 2. Save all data on the workstation.
- 3. On the *softWoRx* menu bar, select **File** | **Exit**. Then exit all other workstation applications.
- 4. Log out of the workstation account.
- 5. Press the IC/MIC power switch to turn off the IC/MIC. Wait 30 60 seconds for the IC/MIC to power down.

Note If the IC/MIC fails to turn off within 30 – 60 seconds after pressing the power switch, contact your system administrator.

- 6. Turn off any additional equipment such as cameras or the heater for the environmental chamber.
- 7. Dab the objective with a clean, unused cotton swab or lens paper to remove all of the oil on the objective. Then apply chloroform to a cotton swab, gently roll it over the objective lens once, and discard it.



**CAUTION:** After you finish imaging, always clean the objective with a clean, unused swab or lens paper. Never reuse swabs or lens paper.



8. Lower the objective by turning the Coarse Z Focus knob away from you (clockwise) when facing the knob.



# 4

## Setting Up and Running Experiments

A *DeltaVision* experiment is a macro that runs a set of instructions for collecting image data. You will use experiments to acquire almost all of your data. This chapter shows how to set up and run experiments that use several *DeltaVision* data collection options. It includes:

- Creating and Running an Experiment Macro
- Sectioning Specimens for 3D Images
- Selecting Filters
- Setting Up Time-lapse Experiments
- Point Visiting
- Collecting Panel Images Over Large Areas
- Using the Multiplexed Wavelength Option

## **Creating and Running an Experiment Macro**

Resolve3D Experiment macros are files that include settings and commands for acquiring data with *DeltaVision*. Macros allow you to automate data collection. After you use the Resolve3D interface to create a macro, you can run the macro to collect data. Macros can be very simple or they can be very complex and include numerous options. After you create a macro, you can use it as a template to modify and create other macros.

## To create and run an experiment macro:

1. At the top of the Resolve3D window, click the **Experiment** button to open the Design/Run Experiment window.

Design Experiment	Design PK Experiment	Run Experiment
Experiment name	Resolve3D	
Estimated file size	32.01 Mb	(21.43 Gb Available)
Use Fast Acquisition	Fast Acquisition Option	ns 🔄 🗆 Lamps Off when finished
Sectioning * Char	nnels Time-lapse Poi	nts Panels Plate Actions
Z Sectioning	Z Scan Options	
ocus point when scan	starts Middle of Sample	· · · · · · · · · · · · · · · · · · ·
Optical section sp	pacing 0.20	
Number of optical se	ctions (64	
Sample thic	kness (12.80 Get	hickness
	☐ Enable OAI Sca	

2. Click the **Design Experiment** tab.

- 3. In the **Experiment name** field, enter a name for the macro only if you need to have it saved as a unique macro. By default, *softWoRx* uses Resolve3D as the experiment name. If you do not intend to reuse an experiment, using Resolve3D for the experiment name is the recommended method.
- 4. From the top of the Design/Run Experiment window, click the **Save** icon. The macro name appears in the Design/Run Experiment window title bar.
- 5. Click the **Run Experiment** tab.

	nt Design PK Experiment	Run Experiment	
Image file na	ume 2_01	Settings	DMS Setup
Image	title I		
Add note to	log [	Do It	]
Change next time la	ose (0.00	Do It	1
Disk space	required: 0.00 Mb	_	
Current o	command:		
	ommand:	imated Finish	

6. In the **Image file name** field, enter the file name.

#### Notes

**#1** Files names can include alpha-numeric, dash, and underscore characters.

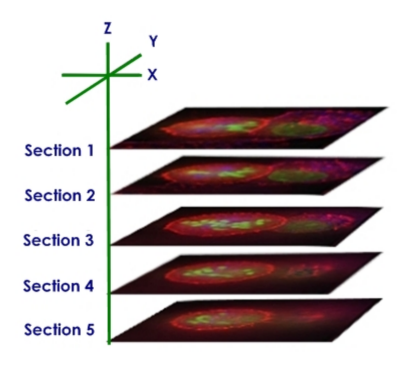
**#2** The resulting image file name has an \_R3D.dv extension. This is the unprocessed (i.e., raw) Resolve3D image. A file with a .log extension is also created. The .log file is a text file containing information about the image file, including the macro and objective that were used to collect the image.

7. In the toolbar at the top of the screen, click the green arrow button 💆 to start the scan.

The macro collects the image and saves it in your data folder (see *Setting Up a Personal Data Folder* on Page 46). To process the image, use the *softWoRx* program, as described in the *softWoRx Imaging Workstation User's Manual*.

## Sectioning Specimens for 3D Images

Using the *DeltaVision* system's precision XYZ stages, you can acquire images of multiple sections. This type of data can be used to create 3D projections, view cross sections, and create volumetric and line models.



If you are new to 3D microscopy, you may have never had to deal with the specimen's absolute three-dimensional size (except in relation to the rest of the specimen). Manually operated microscopes provide very little information about absolute distances in *Z*, or even X and Y. Once you start viewing your specimens in three-dimensions, however, you will become accustomed to the relationship between the microscope stage, sample, and optics, and will find that designing a 3D experiment is usually straightforward.

When you set up a Z sectioning (3D) experiment on *DeltaVision*, you normally begin by defining a Z range (or scanning thickness) to use. You start by finding a central plane of interest within the sample. The sectioning scan range is then defined by finding and marking the top and bottom of the sample. The target specimen data is captured by acquiring images at different Z planes within the defined scan range.

Use the following procedure to create and run a 3D experiment. To determine the Focus point when scan starts, Optical section spacing, Number of optical sections, and Sample thickness, refer to *Guidelines for Designing/Running 3D Experiments* on Page 57.

**Note** The *softWoRx* software reports all distances in microns.

## To design and run a 3D experiment:

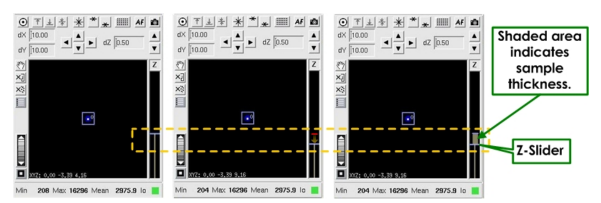
1. You'll need to set the proper exposure for each channel you plan to image. In the Resolve3D window, use the blue Z Slider on the right side of the Resolve3D window to find and focus the object of interest. Select your

primary channel of interest and find its focal plane. This plane tends to provide an image with the highest Max Counts value.

- 2. Adjust the %T and exposure time until the Max Counts is ~50-60% of the camera saturation value for fixed samples and 2-3 times the minimum value for live cells.
  - Tip Generally shorter EX times with higher %T is preferred over long EX times with lower %T.

Repeat the process of finding the exposure time and %T for each active channel.

- 3. Select the filters that correspond to the fluorophore that occupies the greatest volume of your sample.
- 4. Drag the Z Slider in the Resolve3D window up to find the top of your sample. (When the mouse button is pressed, the line turns red. When you release the mouse button, an image is acquired and displayed in the Data Collection window.) Drag the slider or use the Z arrows and acquire images until you are satisfied with the position. Then press the Mark Top to mark that location. Next, use the slider to find the bottom of the sample. When you are satisfied, press the Mark Bottom to mark that location. The region between the top and bottom marks becomes shaded and the represented thickness is used when running the experiment.
- $\square$  Note To reduce the risk of running the objective lens into the sample, the Z slider is limited to 5  $\mu$ m steps at a time.



Use the Z slider to find the top and bottom of a sample. The sample thickness is indicated by the shaded area on the Z slider (shown in the image on the right).

Note The Z stage view displays the relative position of the objective lens to the sample for an inverted microscope. Dragging the Z slider down focuses the microscope closer to the cover glass (negative direction). Dragging the Z slider up moves the focus from the cover glass toward the microscope slide (positive direction).

5. Click **Experiment** to open the Design/Run Experiment window and click the Design Experiment tab. Then click the Sectioning tab.

	Design/Run Experiment: Aug19_test.exp     File	- 3 Help
Click the <u>Design</u> Experiment	Design Experiment Design PK Experiment Run Experiment	
tab.	Experiment name Aug19_tes[ Estimated file size 20.01 Mb (20.59 Gb Available)	
Click the Sectioning tab.	Sectioning *     Channels *     Time-lapse *     Points     Panels     Plate     Actions       V Z Sectioning     Z Scan Options	
	Focus point when scan starts Middle of Sample Optical section spacing (µm) 0.20 Number of optical sections 1 Sample thickness (µm) 0.20 Get thickness	
	Sample thickness (µm) 0.20Get thickness Enable OAI Scan	

6. Specify the focus point of the microscope at the start of the experiment and move the stage to that focus point as follows:

Choose one of the following focus points in the Focus point when scan starts list.	Then click the corresponding button in the Resolve3D window to move the stage to that focus point
Top of the sample	T
Middle of the sample	*
Bottom of the sample	L

**CAUTION:** Be sure to set the focus consistently with this setting before you start the experiment. (For example, if you specify to use the **Middle of Sample** option, be sure to focus midway between the top and bottom of the sample.) *DeltaVision* generates the sectioning commands based on this relative starting point. If this is not done correctly, you may not capture the desired data.

7. Specify the separation (in microns) between each optical section in the Optical section spacing field. If you are using a 60X objective, start by using the default value (.2 µm). If you are using another objective, click the Info button (next to the Lens field) on the Resolve3D window to open the Lens Information window. Use the value in the Recommended Z Step Size field

(at the bottom of the window). As a general rule of thumb, the Z step size is approximately 1/3 of the Z resolution for each objective.

- 8. Specify the number of sections and the sample thickness as follows:
  - To specify the number of sections, enter the number of sections in the **Number of optical sections** field.
  - To get the thickness of the specimen, as defined in Step 2, click **Get thickness**. *softWoRx* adjusts the number of sections to span the specimen thickness.
- 9. Save and run the experiment.

## Guidelines for Designing/Running 3D Experiments

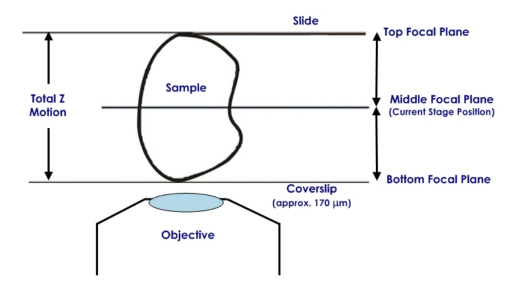
Use the following guidelines to determine values for the fields on the Sectioning tab under the Design/Run Experiment window > Design Experiment tab.

#### Focus point when scan starts

The **Focus point when scan starts** field allows you to specify the focal plane of the microscope stage at the start of an experiment.

Choose one of the following options:

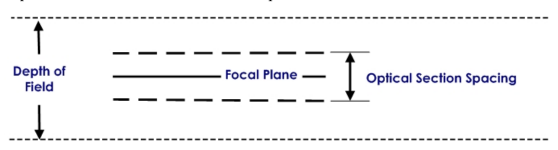
- Middle of Sample (Middle Focal Plane) The focal plane is midway between the top and bottom of the sample. This is the default and is usually the "plane of best focus" for the sample. This is also the best point to set for Point Visiting.
- Bottom of Sample (Top Focal Plane) This represents a plane where the sample is farthest away from the objective.
- Top of Sample (Bottom Focal Plane) This represents a plane where the sample is closest to the objective.



## **Optical section spacing**

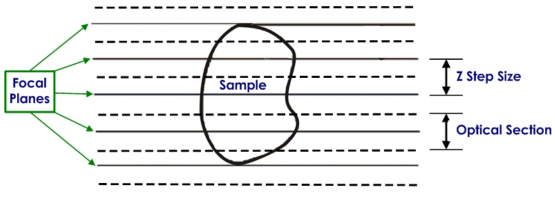
The **Optical section spacing** field indirectly defines the size of the incremental Z stage movement (the distance between focal planes).

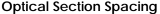
When *DeltaVision* acquires an image, the objective collects information within the depth of field above and below the focal plane.



The Optical section spacing specifies the upper and lower boundaries for which information is collected. It is centered on the focal plane and includes the information above and below that plane.

When you select an Optical section spacing, *DeltaVision* uses a Z step size that is the same distance as the Optical section spacing. (The Z step size is the vertical distance that the stage moves between focal planes.)





For example, the standard step size for a 1.42 NA lens is 0.2  $\mu$ m, although other step sizes are usually fine. Due to the inherent Z resolution of a 1.42 NA lens, there is little need to take steps finer than 0.1  $\mu$ m.

When scanning a PSF (see *Appendix C* on Page 255), use 0.1 µm steps. For low power lenses, the step size should be appropriately larger. The Lens Info window for each lens contains the suggested Z Step size for each lens (derived using Nyquist sampling). This value is approximately 1/3 of the theoretical Z resolution for each objective.

#### Number of optical sections

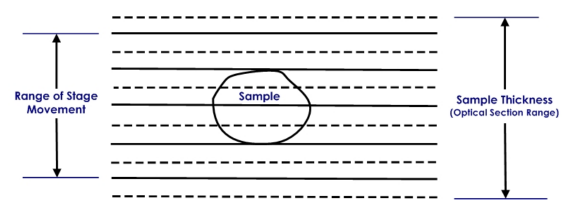
The **Number of optical sections** field defines the number of optical sections in the Z series of the experiment. Enter the number of sections necessary to span from the lower to the upper boundaries of the sample.

The required number of Z sections depends upon the thickness of your specimen. A typical scan consists of between 16 and 64 sections, but there is no set number. For best results, use at least three optical sections. Using one or two optical sections, however, will also provide acceptable results.

The maximum number of optical sections is limited by memory and time. Applied Precision's benchmark deconvolution consists of 64 optical sections that have XY dimensions of 512x512.

#### Sample thickness

The **Sample thickness** field displays the thickness of the optical section range. It is approximately one section thicker than the actual stage movement, as shown below.



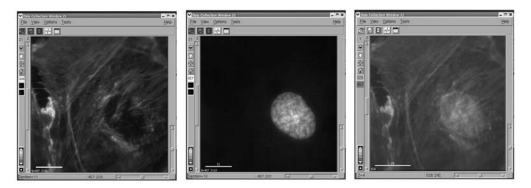
For optimal deconvolution, the Z sections should extend beyond the volume of interest by 1-3 sections.

Sample thickness = Optical section spacing x Number of optical sections

A number of methods have been used to determine the necessary Z scan settings. There is not really a single best way to approach the problem. The best method for your work depends largely on preference and experience.

## **Selecting Filters**

Channels define which wavelengths and filters are used. From the Channels tab on the Design Experiment tab in the Design/Run Experiment window, you can define which filter sets to use in the experiment for imaging. *DeltaVision* acquires images from each selected filter set (channel) and creates a single *DeltaVision* file that contains all of the channels that were collected. There can be up to five channels per experiment.



FITC (left) and DAPI (center) filtered images are combined in the final DeltaVision file (right).

## **Choosing Filters**

Your choice of filters will be determined by the types of fluorescent probes with which you have labeled your sample and the filters available on your system.

The excitation and emission spectrum of your filters should match the spectra for the absorption and emission of the fluorescent probes you want to use.

Some *DeltaVision* systems use interchangeable Standard and Live Cell filter modules. The Standard filter module filter sets are typically used for fixed specimens while the Live Cell filter module sets are commonly used for live specimens.

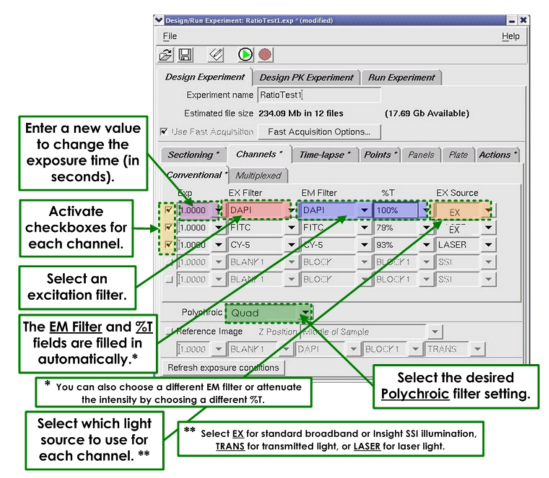
You can select other combinations of excitation and emission filters. You can also add your own custom filter sets. The spectra for the standard *DeltaVision* filter sets and the optional Live Cell sets are provided in *Appendix D* on Page 265.

Note The CY-5 filter set does not include an eyepiece filter due to the low sensitivity of the human eye at this wavelength.

For instructions on how to change filter wheel modules and calibrate filter wheels, see *Changing Filter Wheel Modules* and *Calibrating the Filter Wheels* in Chapter 8.

#### To specify which filters to use:

- 1. On the Design/Run Experiment window, click the **Design Experiment** tab. Then click the **Channels** tab.
- 2. Select a checkbox on the far left next to an **Exp** field. The **EX Filter** field controls which excitation wavelength will be used. The **%T** field attenuates the intensity of the excitation light. The **EM Filter** field controls which filter will be placed in the emission pathway between the microscope and the camera. The **EX Source** field specifies which light source to use for that channel.
- 3. On the **EX Filter** list for the same line as the checkbox you selected, use the drop-down list to select the excitation filter. Entries for the **EM Filter** field and the **%T** field are automatically specified. (*DeltaVision* specifies the most recent filters that were selected for that filter set in the Resolve3D window. If no filters have been selected, *DeltaVision* specifies default filters for that filter set.)



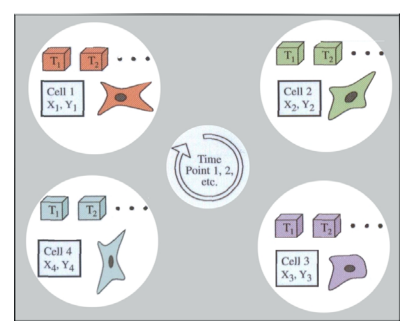
- 4. To change the exposure time, enter a value in the **Exp** field. Values in the **Exp** field are in seconds. For example, 0.100 is 1/10<sup>th</sup> of one second. If you do not enter a value from Resolve3D in this field, the most recently selected exposure value is automatically set for each filter.
- 5. To select a different emission filter, choose a filter from the EM Filter list.
- 6. To select a different neutral density setting to attenuate the intensity, choose a value from the **%T** list.
- 7. You can choose which light source to use for each channel by selecting a light source from the **EX Source** field. Use **EX** for either the xenon or InsightSSI fluorescence light source, **TRANS** is the transmitted light source, and **LASER** is an optional laser source (using the *DeltaVision* X4 Laser module).
- 8. Repeat Steps 2-7 for each set of filters that you want to use.
- 9. In the **Polychroic** field, use the drop-down list to select the appropriate polychroic filter sets. (This field is available only when your *DeltaVision* system is equipped with a motorized polychroic turret.)

- 10. If you are repeating an experiment and you have entered new exposure times or %T in the Resolve3D window, click the **Refresh exposure conditions** button at the bottom of the window to update the chosen settings.
- 11. Save and run the experiment.

## **Point Visiting**

You can use point visiting to monitor areas of the slide that are in different fields of view. Instead of recording one cell or field in a single experiment, multiple sites can be imaged in a single experiment, increasing data collection efficiency.

In practice, the number of sites is limited only by the minimum acceptable time interval between each time point at a single site. This makes time-lapse imaging much more efficient, and allows you to collect enough data to generate statistically significant results. In addition, variability between cells within an experiment can be assayed, eliminating uncertainty as to the behavior of cells in a single experiment.



Point Visiting monitors points that are in different fields of view. To set up an experiment that visits points:

- Mark the points to visit with the Keypad or with Resolve3D and save a point list if desired.
- Edit the point list if necessary.
- Load the point list and specify which points to visit.

#### Notes

#1 If you are placing your slide directly on the microscope stage (without the Repeatable Slide Holder), you can store the X, Y, and Z coordinates of the stage positions you mark; however, if the slide is removed from the stage or the Z focus knob is adjusted, these coordinates no longer apply.

#2 If you are using Applied Precision's Repeatable Slide Holder, you can remove the slide and then place it in the same position when you want to revisit the points. To do this, you must record the Repeatable Slide Holder position (A-G) of the slide before you remove it. Then put the slide in the same position when you return it to the slide holder and update the Z coordinates.

## **Marking Points**

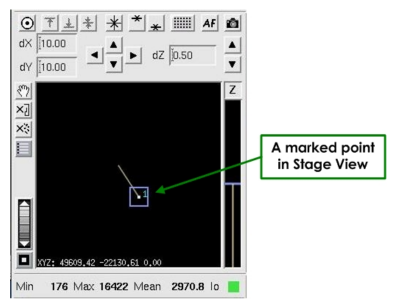
There are two ways to mark points within a sample: using the keypad or using Resolve3D. An efficient way to mark points is to use both methods together. First, mark the points using the keypad, which is quick and approximate, and save the point list. Then, in Resolve3D, specify a more exact location for each point.

The points marked using the keypad are automatically communicated to Resolve3D and displayed in the Stage view. You can view the list of points by opening the Point List window.

#### To mark points using the keypad and Resolve3D:

- 1. Use the arrows on the keypad or the joystick to move to the desired point.
- 2. Press the POINT MARK key on the keypad.

The number of the marked point is displayed in the Stage view.

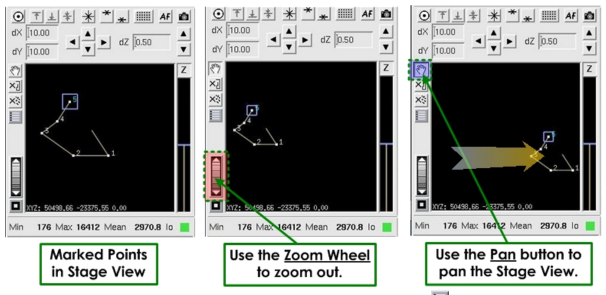


3. Repeat Steps 1 and 2 to mark multiple points.

 $\odot$ TŁ ¥ !!!!! AF Ô dX 10.00 dZ (0.50 . dY 10.00 ধ্য 7 XJ X **Point Numbers** 50416 58 0.00 -221Z0\_61 in Stage View 175 Max 16417 Mean 2970.9 lo Min

As you mark points, the stage trails and points (along with their point numbers) are displayed in the Resolve3D Stage view.

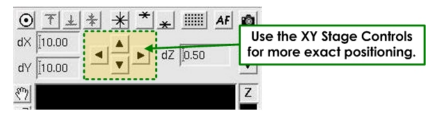
4. To view all of the points, use the Zoom Wheel to zoom out on the Stage View. To pan the Stage View, use the Pan ♥ Button. Activate the Pan tool by clicking the Pan ♥ Button and then dragging the image. To return to normal mode, press the Pan ♥ Button again.



5. In the Resolve3D window click the **Marked Point List** icon to open the Point List window.

Mark Point	Visit Point	1:	+49609.42	-22130		nt and then clic
Delete Point	Replace Point	3:	+51141.65	-22650 -22924	me visit roll	nt button to move to that point.
Clear List	Compact List	5:	+50498.66	-23375	- 5 A.A 5 A.A 5 A.A.	
Save List	Open List	7:	+49408.39	-24069.		
Calibrate All	Optimize List	9: 10:	+48441.17	-23274.	93 +0.00	
		11: 12:	+49183.86 +49615.65	-22860. -22860.		
Done	Help					IZ

- 6. From the Point List window, click on a point to select it and then click **Visit Point** to move the stage to that point.
- 7. Use the XY Stage Controls on the Stage View to move the stage to refine the XY position using the camera.



8. Use the Resolve3D Z controls (the Z buttons or the Z slider on the Resolve3D window) or the *AF* (autofocus) button to find the focal plane.



**CAUTION:** To record and maintain accurate stage coordinates, focus by moving the stage with the Resolve3D controls instead of using the focus knobs to move the objective. Resolve3D cannot track the movement of the objective.

- 9. On the Point List window, click **Replace Point** to replace the old coordinates with the new exact coordinates for the point.
- 10. Repeat Steps 6-9 to replace all of the points in the list with the exact X, Y, and Z coordinates.

## **Editing a Point List**

## To delete a point using Resolve3D:

- 1. In the Point List window, select the point that you want to delete.
- 2. Click Delete Point.

		the <u>Dele</u>	a point and then click <u>elete Point</u> button to the point from the list.			
Point List					,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	-
Mark Point	Visit Point	1:	+49609.42	-22130.61	+0.00	
		2:	+50416.58	-22130.61	+0.00	
Delete Point	Replace Point	3:	+51141.65	-22650.48	+0.00	
		4: 5:	+50785.96	-22924.09	+0.00	
Clear List	Compact List	5:	+49822.91	-23603.09	+0.00	
Coundint	OnenList	7	+49408.39	-24069.43	+0.00	
Save List	Open List	8:	+48924.78	-23637.64	+0.00	
Calibrate All	Optimize List	9:	+48441.17	-23274.93	+0.00	
Galibrate All		10:	+48596.61	-22739.50	+0.00	
		11:	+49183.86	-22860.40	+0.00	
		12:	+49615.65	-22860.40	+0.00	
Done	Help	1				12

#### To save the point list:

1. In the Point List window, click the **Save List** button. The Save Point List File window is displayed.

Preview:	Directory Mome/worx					
	Date ^	File Name				
Created (acquired):  Last Modified:  File Size: 	[07/02/10 14:19] [03/29/11 11:42] [04/12/11 16:45] [11/29/10 16:50]	Desktop/ softworx-logs/				
Image Info: 	File name kplist.	pts OK				
	File Types *.pts	Cancel				

#### Save Point List File Window

- 2. In the Prompt window, enter a name for the point list.
- 3. Press the Enter key or click **OK**. The point list file will have a .pts extension.

#### To visit a point using Resolve3D:

Use one of the following methods to visit a point:

• In the Stage view, double-click on the point.

• In the Point List window, click the point that you want to visit and click the **Visit Point** button.

## Loading a Point List and Specifying Points to Visit

If you are using a previously saved point list, you will need to load the point list in Resolve3D before you specify which points to visit.

Tip You can mark and then visit a point list without saving it when you are setting up an experiment.

## To load a point list into Resolve3D:

1. On the Resolve3D window, click 🗉 or choose **View** | **Points List** to open the Point List window.

Mark Point	Visit Point	1:	+49609.42	-22130.61	+0.00	4
Delete Point	Replace Point	2: 3: 4:	+50416.58 +51141.65 +50785.96	-22130.61 -22650.48 -22924.99		pen List buttor
Clear List	Compact List	5:	+50498 66		+0.00	aved point list.
Save List	Open List	7:	+49408.39	-24069.43	+0.00	
Calibrate All	Optimize List	8: 9: 10:	+48924.78 +48441.17 +48596.61	-23637.64 -23274.93 -22739.50	+0.00 +0.00 +0.00	
		11: 12:	+49183.86	-22860.40 -22860.40	+0.00	
Done	Help	<u>.</u>				

2. Click **Open List**. The Open Point List File window is displayed.

Preview:	Directory //hom	e/worx	<u></u>
	Date ^	File Name	3
Created (acquired):  Last Modified:  File Size: 	[07/02/10 14:19 [10/01/10 09:07 [11/08/10 10:20	7] share on LS Demo Data (Demodisk)/ 9] SW ScreenShots/ 7] Desktop/ 0] softworx-logs/	
Image Info:	<u>الم</u>		K1
	File name 🛽		OK

#### **Open Point List File Window**

3. In the **File name** field, select a file from the directory or enter a name for the list and click **OK**.

Note You may need to load a list when the list of points has been cleared from the Point List window and you want to reload the saved list.

#### To specify which points to visit:

1. Make sure that the point list is loaded in the Point List window.

Calibrate All	Optimize List Help	<u> </u>	for th	ing <u>Optim</u> lers the po ne most ef sitation po	oint list ficient		
Save List	Open List	7:	-2044.35	+329.73	+0.00		l
Clear List	Compact List	5: 3:	-571.54 +769.38	-1274.97 -1824.53	+0.00 +0.00		
Delete Point	Replace Point	4: 1:	+747.40 +0.00	-87.93 +0.00	+0.00 +0.00		
Mark Point	Visit Point	2: 6:	+373.70 +1033.17	+1209.03	+0.00		
Point List						-	

- Note Click the Optimize List button to run an algorithm that determines the fastest route through all of the selected points, beginning with the currently selected point. The sequence of points in the list is reordered accordingly. (In the example above, Point #2 was selected as the starting point.)
- 2. Open the Design/Run Experiment window and click the **Design Experiment** tab. Then click the **Points** tab and activate the **Visit Point List** checkbox.

	▶ Design/Run Experiment: * (modified)		- ×
	File		Help
Click the <u>Points</u> tab.	Design Experiment Design PK Experiment name Resolve3D	ment Run Experiment	
Activate the Visit Point List checkbox.	Estimated file size 32.01 Mb	(17.30 Gb Available) Options	
	Sectioning * Channels Time-laps	Points * Panels Plate Acti	ions
	Visit Point List	1-3,5-7,9	Enter the points to visit.
	☐ Image-based Autofocus before imaging	Autofocus Options	

The Point Visiting window is linked to the point list that is open.

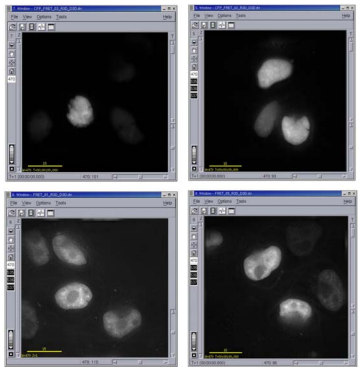
3. Enter the points that you would like to visit in the **Visit Point List** field (to the right of the **Visit Point List** checkbox). For example, entering 1-3 specifies

to visit the first 3 points in the point list. Entering 1-3, 5-7, 9 specifies to visit Points 1-9, but not 4 or 8.

Note It's important to note that the point names do not really reflect their position within the list. The values you enter into this field represent the point positions in the list, not necessarily the displayed names. (For example, if your list has point names of 1, 3, 7, 8 and you enter 2, 4 into the field, the experiment would visit point names 3 & 8.)

## Monitoring Point Visiting Experiments with Point Track

You can set up *DeltaVision* to display a separate Data Collection window for each point in your point visiting experiment.



With the Point Track Display Option enabled, each point in a point visiting experiment is displayed in a separate window.

## To set the Point Track display option:

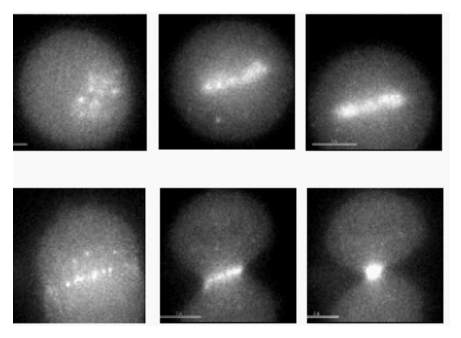
- 1. On the Resolve3D window, click the **Settings** button to open the Resolve3D Settings window.
- 2. Click the **Display** tab.

Display	Files	Imaging	Illumination	Autofocus	Lasers	Misc
m <mark>age</mark> disp	lay mode	Point Track	· ·			
	Window	21 💌	]			
	Wave	1				
7 Calculat	e statistic	s				
7 Calculat	e histogra	am				
7 Auto his	togram ra	ange				
7 Display	images					
] Deconv	olve prev	iew images				
7 Auto int	ensity sca	ale				
7 Acquire	after poir	nt visit				

- 3. In the Image Display mode list, select **Point Track**.
- 4. Click the **Save Settings** button, then click **Done** to close the Resolve3D Settings window.

## Setting up Time-lapse Experiments

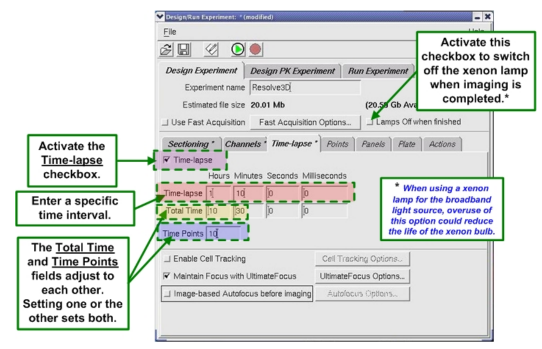
Time-lapse images are very useful for analyzing live specimens.



Time-lapse images showing cell mitosis

## To design a time-lapse experiment:

1. Select the Time-lapse tab under the Design Experiment tab on the Design/Run Experiment window.



- 2. Activate the Time-lapse checkbox.
- 3. If your *DeltaVision* system is using a xenon arc lamp for its broadband illumination source and you want the lamp to turn off when the experiment is completed, activate the **Lamps Off when finished** checkbox. (The **Lamps Off when Finished** checkbox is displayed only if your DeltaVision system is equipped with a xenon arc lamp as its broadband light source.)

Note When using the xenon lamp, use discretion when activating the Lamps Off when finished checkbox feature. Cycling the *DeltaVision* lamp on or off more than is necessary can reduce the xenon bulb life.

4. Enter the desired time interval in the **Time-lapse** field.

Note The minimum time interval is limited by acquisition time. If you specify a time that is less than the acquisition time, the experiment proceeds at the quickest possible rate.

5. Set up the remaining parameters for your experiment in one of the following ways:

- Enter the desired number of time points in the **Time Points** field. The **Total Time** field displays the total time that will elapse during the experiment.
- Enter the total time in the **Total Time** field. The **Time Points** field displays the number of time points for the experiment.

### Using the Actions Tab with Time-lapse Experiments

When designing time-lapse experiments, you can also use the Actions tab in the Design/Run Experiment window to further customize your experiments. The Actions tab allows you to set up additional tasks for your experiment macro to complete before, during, or after a time-lapse interval.

#### To use the Actions tab:

- 1. At the top of the Resolve3D window, click the **Experiment** button to open the Design/Run Experiment window.
- 2. Click the Design Experiment tab and then click the Actions tab.

♥ Design/Run Experiment: Tes	st1_withTTL.exp * (modified)		- ×	
<u>F</u> ile			Help	
Design Experiment	Ocsign PK Experiment R	un Experiment		Select the Design
			1	Experiment tab,
	2.34 Gb in 12 files	(18.55 Gb Available)		then select the
Use Fast Acquisition				Actions tab.
	Past Acquisition Options			
		ints * Panels Plate	Actions	
? Time Visited Points Points	Action	When	Options	
+ સૉ્રિ	Fatio Image 💷	After Imaging 💷		

3. Fast Acquisition is useful for highly dynamic experiments in which data must be captured quickly. In general, Fast Acquisition prioritizes the acquisition process so that the imaging is completed as quickly as the system can handle it. Other processes that the system would normally be checking during the acquisition are placed "on hold" until the system has completed the task. The advantage of this method is that the time sequences within the experiment can remain more predictable when the system is focused solely on the acquisition.

If you plan to use the Fast Acquisition method for this experiment, activate the **Use Fast Acquisition** checkbox.

- 4. If you activated the **Use Fast Acquisition** checkbox, click the **Fast Acquisition Options** button and continue with Step 5. Otherwise, skip to Step 8 of this procedure.
- 5. The Fast Acquisition Options window is displayed, as shown.

♥ Fast Acquisition Options		×
Image Scan Sequence	Wavelength then Z	-
Open Shutter For Each	Z Scan	•
Camera Readout Mode	Medium	•
	✓ Use Conventional Time-laps	se
Done	H	elp

The choices for this screen are as follows:

**Image Scan Sequence** – Two options are available:

- Wavelength then Z causes the system to switch filter positions before changing the Z position. Use this mode when channel-to-channel alignment is critical.
- Z then Wavelength causes the system to collect an entire Z stack before switching filters and rescanning the specimen. This method is faster because the filter wheels are moved only once for each wavelength during the Z scan. The tradeoff is that different wavelengths may not be as finely aligned as with the Wavelength then Z method.
- **Open Shutter for Each** Selects when the shutter should open and close during the acquisition. Three options are available:
  - **Exposure** should be used for any acquisitions that require more than one wavelength and for acquisitions using a full-frame CCD chip.
  - **Z** Scan should be used for single wavelength acquisitions.
  - Entire Scan should be used for time-lapse acquisitions with only one Z section and one wavelength. The shutter should be closed after each exposure unless the time-lapse duration is zero. The Entire Scan option should also be used for acquisitions utilizing an interline chip.

Camera Readout Mode – Selects the camera readout mode to best match the experiment requirements. The implementation of the readout mode depends on the camera model and manufacturer. Not all cameras have noticeable differences between the modes. Five options are available: Normal, Very Fast/Aggressive, Fast/Aggressive, Medium, Slow/Careful.

- **Normal** is the default camera configuration, which is usually the same as Slow/Careful.
- Fast/Aggressive and Very Fast/Aggressive readouts attempt to move electrons more quickly off of the CCD chip, which can cause electrons to be "spilled" into adjacent CCD wells. This is usually not a problem unless the CCD has collected enough light (electrons) to be near full-well capacity. In most cases, high-speed imaging does not allow sufficient time for collecting lots of photons, so the more aggressive modes usually do not cause problems in this regard.
- 6. Activate the **Use Conventional Time-lapse** checkbox. This mode allows the use of the tools on the Actions tab, including **UltimateFocus**, **Autofocus**, and **Cell Tracking**, even when **Fast Acquisition** is selected.

**Note** This may slow down acquisition.

- 7. When you are satisfied with your selections in the Fast Acquisition Options window, click the **Done** button to return to the **Actions** tab of the Design/Run Experiment window.
- 8. Add new items to the list of actions by clicking the "+" button in the far-left column. Remove unwanted items from the list of actions by clicking the "-" button in the far-left column.

Note Actions must be removed from the list in the reverse order in which they were added.

- 9. In the following two columns, **Time Points** and **Visited Points** may be indicated as follows:
  - By the keyword: **all**
  - As a range: 1–5
  - As a list of values: **1**, **3**, **5**, **8**
  - As an Nth expression: every 3, or 7 then every 4 (indicates 7, 11, 15, 19...)
  - As a compound specification: 1–3, 5, 7 then every 3

- Note The above information is also available on your screen whenever you click the "?" button just left of the Time Points heading, on the left.
- 10. Each row in the list of actions represents an individual action. In the **Time Points** column, select the time points at which you want the selected action to occur.
- 11. If your experiment includes point visiting, select the appropriate points in the **Visited Points** column. Note that this column defaults to **all** if left blank.
- 12. In the **Action** column, use the drop-down list to select the type of action to include in this experiment. Note that the details and conditions relating to each action are provided in the **Options** column at the far-right.

♥ Design/Run Experiment: T	est1_withTTL.exp * (modified)	- *	
<u>F</u> ile	He	elp	
28 🖉 🖸			
	Design PK Experiment     Run Experiment       e     Test1_withTTU       e     2.34 Gb in 12 files       n     Fast Acquisition Options		Click to view a drop-down list of the available actions.
Sectioning Cha	annels * Time-lapse * Points * Panels Plate Actions	••]	
?     Time Points     Visite Points       -     ali     ali       -     svery 10     0       -     3     0       -     every 5     0			Click to view options for each selected action.
Click the "+" button	Change Time-lapse Ratio Image Add Log Note Trigger TTL Custom		Click to view a drop-down list of when each action is to occur.
to include additional actions.	<u></u> :		

The actions available in the Action column are as follows:

- Laser Event places a laser event into the experiment macro at each time point indicated.
- **UltimateFocus** uses the hardware-based UltimateFocus tool at each time point indicated.
- **Autofocus** uses the software-based Autofocus tool at each time point indicated.
- **Cell Track** applies cell tracking for each time point indicated.

- **Pause** stops the acquisition process at each indicated time point until the user intervenes to continue the experiment.
- **Wait** stops the acquisition process at each indicated time point, and then waits for a specified amount of time before continuing the experiment.
- **Change Time-lapse** changes the subsequent time-lapse interval at each time point indicated.
- **Ratio Image** generates a ratio image at each time point indicated. A ratio image is a simple two-channel image displayed with a ratio graph showing the mean value vs. time.
- Add Log Note inserts a note in the Resolve3D acquisition file.
- **Trigger TTL** inserts a TTL command into the experiment macro at each time point indicated.
- **Custom** allows you to insert macro command directly into the experiment macro. This action should be used by advanced users only.
- 13. In the **When** column, use the drop-down list to select the time frame during which you want the action to occur. The choices for this column are **Before Imaging**, **After Imaging**, or **During Time-lapse**.
- 14. In the **Options** column, click the button to view an Options window specific to the action you have chosen. The following subsections describe each of the possible Action Options windows.

#### Laser Event

When you select **Laser Event** as the action and then select **Options**, the Laser Action Options window is displayed as shown.

Laser Action Options			
Laser focus point Mi	ddle of Sample 📃 💌		
- Laser Event Settin	gs	**	
Laser channel	488 💌 Duration (secs) [0.0	50 F	Refresh
Number of events	<u>[</u> 1		
Bleach event motion	Optimize for Position	•	
Event pattern	Use Current Center Point	•	
- Create Bleach Eve	nts		
	Active set	A - Red	
Pattern Spacing 20	🛛 🔽 Show Event Graphics		
Done			

On the Laser Action Options window:

- Use the drop-down list in the Laser focus point field to select Middle of Sample, Bottom of Sample, or Top of Sample.
- Select the Laser channel to use and specify the Duration (in seconds), or use the **Refresh** button to update the active laser and duration to that currently used in the Laser Module Settings window.
- Select how many laser events you want for this action in the **Number of events** field.
- Use the drop-down list in the **Bleach event motion** field to select either **Optimize for Position** or **Optimize for Speed**.
- Use the drop-down list in the **Event pattern** field to select either **Use Current Center Point** or **Use Bleach Event Specification**.
- If you selected **Use Bleach Event Specification** in the previous step, use the now active tools in the Create Bleach Events section of the window to generate specific bleach events, otherwise, skip to the next step.
- Click the **Done** button.

#### **UltimateFocus**

When you select **UltimateFocus** as the action and then select **Options**, the UltimateFocus Action Options window is displayed as shown.

♥ UltimateFocus Action O	ptions 📈 🗙
Move Threshold ( $\mu m$ )	0.100
Maximum Iterations	Ĭz
Done	

On the UltimateFocus Action Options window:

- In the **Move Threshold** field, enter the desired maximum measured focus error (in µm) before a corrective action should be taken.
- In the **Maximum Iterations** field, enter the maximum measure/move sequences for this action to reach the calibrated focus point.

**Note** More iterations will take longer, but may provide better focus.

• Click the **Done** button.

#### Autofocus

When you select **AutoFocus** as the action and then select **Options**, the AutoFocus Action Options window is displayed as shown.

Autofocus Action Options	<u>×</u>
Auto-Determine Parameter	rs
Channel for Autofocus	DAPI
Contrast calculation method	Auto
Autofocus Z test step (µm)	j3.12
Maximum Z test range (µm)	<u></u> 50.00
Post-autofocus Z offset (µm)	[0.00
Done	

On the Autofocus Action Options window:

- To have the options in this window set automatically, activate the Auto-Determine Parameters checkbox. Then click the Done button.
   If you <u>do not</u> activate the Auto-Determine Parameters checkbox, set the remaining fields in the window as follows:
- Use the drop-down list in the **Channel for Autofocus** field to select the channel you want to use for Autofocus.

- Use the drop-down list in the **Contrast calculation method** field to select **Auto**, **Fluorescence (Light on Dark)**, or **Brightfield (Dark on Light)**.
- Select the Autofocus Z test step (in µm) for this experiment.
- Choose the Maximum Z test range (in µm) for this experiment.
- Indicate any **Post-autofocus Z offset** (in µm) for this experiment.
- Click the **Done** button.

#### **Cell Track**

When you select **Cell Track** as the action and then select **Options**, the Cell Tracking Action Options window is displayed as shown.

Cell Tracking Options	×
Tracking Method Center of Inte	nsity 🗾 💌
Reference Channel 1	
Move Threshold (µm) 1.000	
ROI Percent	50
Done	Help

On the Cell Tracking Options window:

- Use the drop-down list in the **Tracking Method** field to select either **Center of Intensity** or **Center of Shape**.
- Use the drop-down list in the **Reference Channel** field to choose the channel to use for reference.
- Enter the desired **Move Threshold** (in µm).
- Use the **ROI Percent** slider to indicate the appropriate ROI percentage.
- Click the **Done** button.

#### Pause

The **Pause** action does not have a Pause Action Option window. Use this action to pause the experiment either before or after imaging. The **Pause** action requires user intervention before the experiment can continue.

Note For additional information on these settings, see the *Settings Window Autofocus Options* section on Page 315 in *Appendix E*.

#### Wait

When you select **Wait** as the action and then select **Options**, the Wait Action Options window is displayed as shown.

💙 Wait Acti	on Options	×
Wait For	[1	Seconds
Done		

On the Wait Action Options window:

- Enter the amount of time (in seconds) for the experiment to wait. The **Wait** action does not require user intervention for the experiment to continue.
- Click the **Done** button.

#### **Change Time-lapse**

When you select **Change Time-lapse** as the action and then select **Options**, the Time-lapse Action Options window is displayed as shown.

♥ Time-lapse Action Option	IS	×
New Time-lapse Value	<u>)</u> 0.000	seconds
Done		

On the Time-lapse Action Options window:

- Enter the new time-lapse (in seconds) for the remainder of this experiment.
- Click the **Done** button.

#### Ratio Image

When you select **Ratio Image** as the action and then select **Options**, the Ratio Image Action Options window is displayed as shown.

♥ Ratio Image Action Options	×
🔽 Use Channel 1 As Numerato	r
Channel 1 Background 0.000	
Channel 2 Background 0.000	
Done	

A ratio image is basically a method of viewing experiment status. On the Ratio Image Action Options window:

- To use Channel 1 as the ratio's numerator, activate the **Use Channel 1 as Numerator** checkbox.
- In the **Channel 1** and **Channel 2 Background** fields, enter the intensity values to subtract from each channel.
- Click the **Done** button.

While a ratio imaging experiment is running, a two-channel time-lapse image is displayed along with a ratio graph showing the mean value vs. time. The mean value for a ratio graph is calculated using a specified portion of the image. See *Using Ratio Imaging* on Page 87 for details.

#### Add Log Note

When you select **Add Log Note** as the action and then select **Options**, the Log Note Action Options window is displayed as shown.

Log Note Action Options	×
Use text from Run Experiment dialog	
- or -	
Put this note in the log:	
Y	
Done	_

On the Log Note Action Options window:

- Activate the **Use text from Run Experiment dialog** checkbox to use the text from the **Add note to log** field on the Run Experiment tab.
- If you leave the checkbox inactive, enter the text for your log note in the field provided.
- Click the **Done** button.

#### Trigger TTL

You can insert TTL triggers into an experiment macro to initiate specific events from various components during your experiment. When you select **Trigger TTL** as the action and then select **Options**, the TTL Action Options window is displayed as shown.

✓ TTL Action Options	×
TTL Channel	<b>]</b> 1
Pulse Duration (msec)	
Done	

On the TTL Action Options window:

- Enter the appropriate channel in the TTL Channel field.
- Enter the desired pulse duration for this TTL trigger in the **Pulse Duration** field.
- Click the **Done** button.

#### Custom



**CAUTION:** The use of **Custom Actions** should be considered by advanced users only. In particular, adding any command to the macro that changes the number of image frames may cause Resolve3D to fail or behave unexpectedly.

When you select **Custom** as the action and then select **Options**, the Custom Action Options window is displayed as shown.

Help on Macro Commands	Available Macro Commands
Macro commands to execute	ACTIVATELASER ACTIVE ACTSHUT ACTSHUT ACTOFOCUS ATTORAMOE ATTOSCALE ATOSCALE ATOSCALE ATOSCALE ATOSCALE ATOSCALE ATOSCALE ATOSCALE ATOSCALE ATOSCALE COD COD CLP CLP CLP CLP CLP CLP CLP CLP CLP CLP

• Use the Custom Action Options window to specify macro commands that will be executed at the specified time points.



**CAUTION:** Specifying any macro that changes the size (number of X, Y, Z, Time, or Channels) can result in image files that cannot be used by *softWoRx*.

• When you are satisfied with the Custom Actions Options window, click the **Done** button.

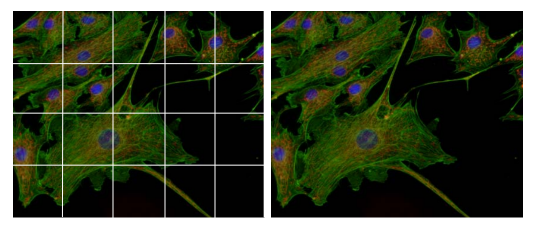
# Sending TTL Triggers to External Devices

For sending TTL signals to devices external to the *DeltaVision* system, you'll need a cable to run from the back of the IC/MIC to the external device. You can order this cable directly from Applied Precision (ask for Part Number 54-852384-000G), or you can ask your Applied Precision representative for a pin-out diagram of the cable so you can build your own.

Note All outputs from a *DeltaVision* system's TTL interface are Active High by default.

# **Collecting Panel Images over Large Areas**

Panel collection macros are useful when you want to scan a large area with a relatively high magnification lens. You can use the panels as a means of reviewing a large area of a slide, or as data that you want to stitch together to form a single, large image.



A set of panel images (left) can be stitched together to create a single image (right). Image files that are collected with Panel Collection enabled may be stitched to create a single . dv file. For details, see Stitching in the *softWoRx Imaging Workstation User's Manual*.



**CAUTION:** Do not try to reuse panel collection experiment macros. These macros are sensitive to microscope settings such as image size and magnification because the number of panels required depends upon many factors.

# **Determining Border Rolloff Voxels**

Before you collect panel images, you must first determine the number of *voxels* in the Border Rolloff that is used in Deconvolution.

#### To determine Border Rolloff Voxels:

1. Determine the size of the panels that you plan to collect.

i Tip Smaller panels work best because they have the flattest intensity distribution.

- 2. From the *softWoRx* main menu, choose **Process** | **Deconvolve** to open the Deconvolve window. In the **Input** field, enter the file name of an image that is similar to the panel size. (You can acquire and save a blank image of that size for this purpose.)
- 3. Click the **More Options** button to open the More Deconvolution Options window and record the value that is displayed in the **Border Rolloff (voxels)** field.

## **Collecting Panel Images**

#### To collect 3D panel images:

- 1. From the Resolve3D window, choose **View** | **Point List**. In the Point List window, choose **Mark Point** and mark points in two opposite corners to define the area of your final composite image.
- 2. In the Resolve3D window, click **Experiment** to open the Design/Run Experiment window.
- 3. Click the **Design Experiment** tab. Then set up the Sectioning and Channels information in the same way that you would for standard data collection.
- 4. In the Point List window, select one of the points that you marked in Step 1 and click **Visit Point**.
- 5. Click the Panels tab under the Design Experiment tab. Then select the **Collect Panels** option.
- 6. Click **Get Start** under the Panels tab. Then visit the other point that you marked in Step 1 and click **Get End**. (Alternatively, you can enter the coordinates that are displayed in the Point List window.)
- 7. In the **Overlap (pixels)** field, enter at least twice the number of border rolloff voxels as you recorded when you determined border rolloff voxels (see Page 84).
- 8. Save and run the experiment.

For instructions on how to use *softWoRx* to stitch panels together, see the *softWoRx Imaging Workstation User's Manual*.

**Note** You cannot easily use Panel Collection with time-lapse or point-visiting experiments. It is possible only by editing the header.

# 5

# **Acquiring Data from Live Specimens**

This chapter shows how to use the following features for imaging live cells with *DeltaVision*.

- Set up *Experiment Macros* to automatically focus before acquiring each image. This is useful for cells that move during the experiment.
- Use *Cell Tracking* to follow cells as they move laterally and move the stage to keep them in the field of view.
- Use Optical Axis Integration (or OAI; also referred to as Z Sweep Acquisition) to acquire a 2D projection. This method collects and integrates one continuous image through an extended Z movement. Z Sweep Acquisition is much faster than the traditional technique of collecting an individual image at each focal plane. It also reduces the risk of specimen damage and has less total camera read noise.
- Acquire Reference Images that can be used for Differential Interference Contrast (DIC) and other types of analysis.

# **Using Autofocus in Experiment Macros**

You can set up your experiment macro to automatically focus before each image is acquired. This is especially useful for long time-lapse experiments that are susceptible to changes in environmental conditions. Autofocus allows you to acquire focused images without closely monitoring the experiment. The contrastbased autofocus algorithm is a software method that uses both image contrast and peak image intensity to determine when the specimen is in focus.

To optimize the focus position for *DeltaVision*, you must find the Z position where the contrast is greatest. However, this position may not always be at the optimal in-focus plane of the sample. Autofocus allows you to adjust several parameters within the standard algorithm so that the final Z position is most favorable for the specific sample. Limiting the step size and maximum Z range of travel can limit the effects of objects that are not of interest. In cases where the contrast outside of the actual middle of the sample is highest, you can create an offset so that the middle of the sample is located during the Autofocus process.

The Autofocus option works the same way as the **AF** button on the Resolve3D window. Use the following instructions to include this option in the experiment macro.

# To set up an experiment that automatically focuses before acquiring points:

- 1. At the top of the Resolve3D window, click the **Experiment** button to open the Design/Run Experiment window.
- 2. Click the **Design Experiment** tab and then click the **Time-lapse** tab.

Design/Run Experim	ient: * (mod	ified)				– X Help	
Design Experiment Experiment Estimated fil ▼ Use Fast Acqu	name R	esian PK Exper esolve3D <u></u> 2.01 Mb Fast Acquisitio	-	un Experiment (19.07 Gb Av	anabi	<u>Design E</u> and <u>Tir</u>	rate the Experiment me-lapse abs.
Time-lapse		s Seconds Mil	-	Panels Plate	e Act	ions	
<ul> <li>✓ Enable Cell Trans</li> <li>✓ Maintain Focu</li> <li>✓ Image-based a</li> </ul>	s with Ulti		UltimateF	king Options			

- 3. Select the **Autofocus before imaging** option. When this option is selected, the software automatically determines the parameters for the Autofocus process.
- 4. To change the various Autofocus parameters, click the **Autofocus Options** button. The Resolve3D Settings window is displayed with the **Autofocus** tab selected.

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🗆 Auto	omatically determine	parameters Test			
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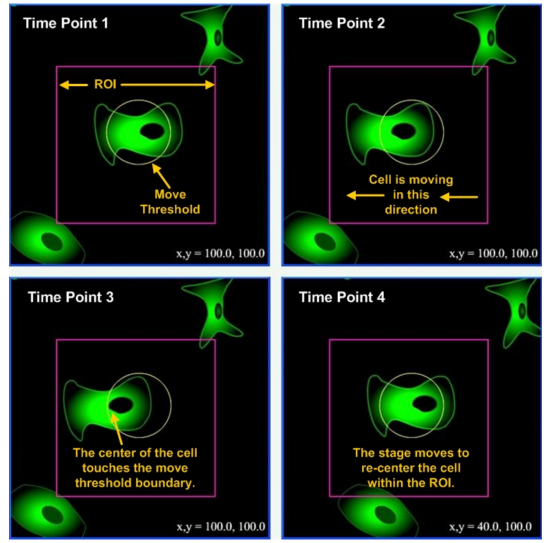
5. Deselect the **Automatically determine parameters** option. The remaining parameters in the window become active.

The Autofocus parameters in this window are described in *Settings Window Autofocus Options* on Page 315.

# **Tracking Cells**

Cell Tracking moves the stage laterally to follow cells as they move during a time-lapse experiment. With the Enable Cell Tracking option selected, *DeltaVision* automatically keeps cells in the field of view as they move during a time-lapse experiment.

After you specify a region of interest (or ROI) around a specific portion of the sample, the software determines the center of the cell and establishes a recognizable pattern within the ROI. In subsequent images, the software recognizes this pattern and recalculates the center of the cell on-the-fly. The position of the new center is compared with the position of the previous center and, if the cell has moved beyond a specified threshold, the system automatically moves the stage and re-centers the cell in the field of view.



At Time Points 1 and 2, the cell is moving through the ROI (the inner square), but the center is still within the Move Threshold (the circle). At Time Point 3, once the center of the cell touches the threshold boundary, the stage moves to re-center the cell within the ROI. (Note the new stage position displayed in the lower-right corner of Time Point 4.)

#### To use Cell Tracking:

- 1. Set up a time-lapse experiment as shown in *Setting Up Time-lapse Experiments* on Page 71.
- 2. On the **Time-lapse** tab in the Design/Run Experiment window, activate the **Enable Cell Tracking** checkbox.

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3. Click the **Cell Tracking Options** button. The Cell Tracking Options window is displayed.

Cell Tracking Options	ilinnin manan in an
Tracking Method	Center of Intensity
Reference Channel	1
Move Threshold (microns)	1.000
ROI Percent	50
Done	Help

#### **Cell Tracking Options Window**

4. Manually acquire an image for each channel and determine which single channel best identifies the features of your specimen. In the **Reference Channel** list, specify that channel. (The channels are numbered in the order in which they are listed on the Design Experiment Channels tab.)

**Note** *Delta Vision* uses the reference channel for image recognition.

- 5. In the **Move Threshold** field, enter the distance in microns that the cell must move to trigger stage movement.
- 6. Specify the area around the cell (the region of interest or ROI) that you want *DeltaVision* to use for image recognition. For more about this option, see *ROI Percent* on Page 95.
- 7. If you are performing an experiment on a single cell, use the stage controls to center the cell (laterally) in the field of view. If you are performing a point

visiting experiment, make sure that the points that you are monitoring are centered before you start your experiment.

8. Click the Run Experiment tab and then click the **Green Arrow** button to start the scan.

## **Guidelines for specifying Cell Tracking Options**

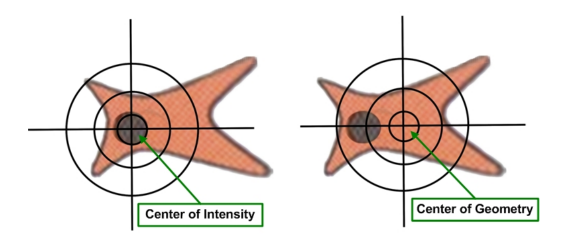
You can specify several parameters that *DeltaVision* uses to track cells:

- Tracking Method
- Reference Channel
- Move Threshold
- ROI Percent

#### **Tracking Method**

The tracking method is the process *DeltaVision* uses to determine the center of the feature. This center is recalculated after each image is acquired. You can choose between two tracking methods:

- Center of Intensity calculates the center of the feature based on intensity values.
- Center of Geometry calculates the center of the feature based on its geometry.



#### **Reference Channel**

*DeltaVision* uses the reference channel for pattern recognition. If you are acquiring more than one channel, choose the channel that has the most contiguous features.

Note The term, *Reference Channel* should not be confused with *Reference Image*, which is something very different. Reference images are described later in this chapter in *Acquiring a Reference Image*.

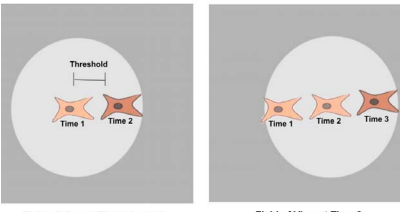
Channels are numbered as they appear in the Experiment Designer window. In the following example, CFP is Channel 1 and YFP is Channel 2.

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28								
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Refresh exp	osure con	ditions						

#### Move Threshold

The move threshold is the distance the center of the cell (as defined by the tracking method) must travel before the system resets the stage. When the cell moves beyond this threshold, the stage moves so that the center of the cell retains its original position in relationship to the center of the field of view.

Live cells, by their very nature, are constantly on the move. When choosing a move threshold, it is important to select one that is small enough to keep the cell in the field of view, yet large enough to buffer out needless stage movement that could result from numerous small cell movements.



Field of View at Times 1 and 2

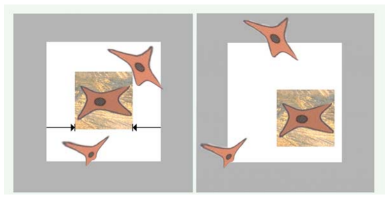
Field of View at Time 3

Cell tracking lags cell movement. Cell movement exceeds the threshold between Time Points 1 and 2 (left). In response, *DeltaVision* moves the stage so that it is centered on the cell position at Time Point 2. The stage is at this position when Time Point 3 is acquired (right).

#### **ROI** Percent

With rare exceptions, few live cells are isolated on the substrate. Most are genetically "programmed" to seek attachment and communication with other cells. When performing live cell microscopy, not only is it important to filter out unnecessary stage movement, it is also important to define an appropriate region of interest within the field of view. This region of interest must be large enough to include the entire cell or structure of interest, yet small enough to exclude other cells or structures that may tend to wander in and out of the field of view throughout the experiment.

The ROI Percent parameter defines how much of the field of view the software will use for image recognition. In the software, ROI units are specified as a percentage of the width of the field of view. For example, a 50% ROI has a width that is 50% of the width of the field of view.

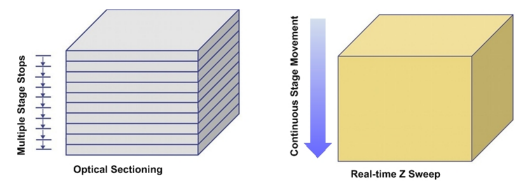


In the first image, the width of the ROI is defined as 50% of the width of the field of view. In the second image, the stage has moved to keep the cell of interest within the ROI. Perimeter cells have also moved, almost entirely out of the field.

Tip During a time-lapse experiment, it's possible to lose the cell as it moves out of the field-of-view. When this occurs, you can update the point coordinates by manually finding the cell and centering it in the field-of-view. The point list is automatically updated.

# Acquiring 2D Z Projections with OAI

Optical Axis Integration (or OAI), also referred to as Z Sweep Acquisition, is useful for acquiring 2D Z projections of live specimens. Instead of collecting an individual image at each focal plane, OAI collects and integrates one continuous image through an extended Z movement.



Instead of acquiring multiple images, Real-Time Z Sweep acquires one image during a continuous stage movement and instantly creates a 2D Z Projection.

Z Sweep Acquisition has significant advantages for applications such as Leading Edge Motion Analysis, Fast Organelle Dynamics, Microtubule Dynamics, and Fluorescence in situ Hybridization (FISH). It is especially useful for studies of objects that are moving in 3D space (e.g., kinetichores in a cell nucleus or other rapidly moving structures).

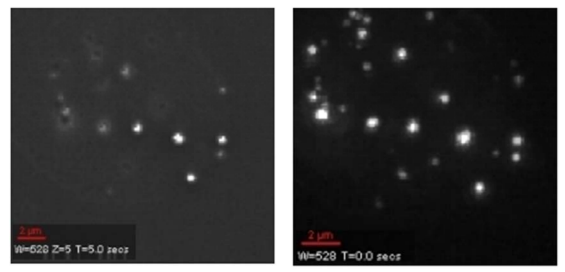
## Z Sweep Acquisition versus Traditional Projections

If you are acquiring data objects that you plan to use for 2D projections, Z Sweep Acquisition provides several advantages over creating 2D projections from multiple optical sections:

- Complete Z data acquisition collects all data in the interval of interest (with 2D imaging or Z section sampling, some data is lost.)
- Fast data acquisition provides accurate image registration of rapidly moving objects.
- Low total exposure time reduces the risk of damage to the specimen.

- Low total read noise (the camera is only read once) improves signal-to-noise ratio.
- Note For samples that contain a large amount of fluorescence throughout 3D space (for example, a tumor spheroid that has a lot of fluorescence), optical sectioning may provide better results than OAI.

The following 2D image of endosomes in a HeLa cell (left) and a 2D Z projection (right) of the same area were acquired under similar conditions. The additional data in the 2D Z projection include objects that moved out of the depth-of-field of the 2D image during the data acquisition process.

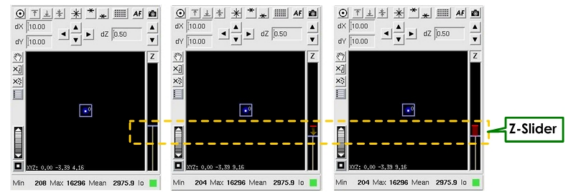


endosomes in a HeLa cell (left)and an instant 2D Z projection (right) of the same area

# Using Z Sweep Acquisition

#### To set up a Z Sweep Acquisition Experiment:

- 1. Set up your experiment as shown in *Creating and Running an Experiment Macro* on Page 52.
- 2. In the Resolve3D window, use the Z Slider on the right side of the Resolve3D window to find the best focal plane. Drag the slider up to find the top of your sample. (When you release the mouse button, an image is acquired and displayed in the Data Collection window.) Drag the slider and acquire images until you are satisfied. Then press the the button to mark that location. Next, use the slider to find the bottom of the sample. When you are satisfied, press the to mark that location.



Use the Z slider to find the top and bottom of a sample. The sample thickness is indicated by the wide line on the Z slider (shown on the right image).

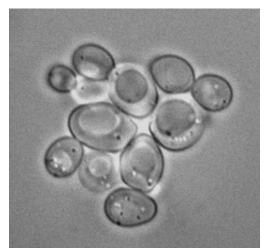
- 3. Click **Experiment** to open the Design/Run Experiment window. Then click the Sectioning tab.
- 4. Select the **Z** Sectioning and Enable OAI Scan options.

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File	Help
Design Experiment         Design PK Experiment         Run Experiment           Experiment name         Resolve3Di	
Estimated file size 32.01 Mb (21.43 Gb Available)  Very Use Fast Acquisition Fast Acquisition Options Lamps Off when finished  Sectioning * Channels Time tapse Points Panels Plate Actions	Activate the <u>Z Sectioning</u> and <u>Enable OAI Scan</u> checkboxes.
Z Sectioning       Z Scan Options         Focus point when scan starts       Middle of Sample         Optical section spacing       [0.20         Number of optical sections       [64         Sample thickness       [12.80         Get thickness       [12.80         Enable OAI Scan	

- 5. Click the **Run Experiment** tab and click the **Green Arrow** button in the toolbar at the top of the screen to start the experiment.
- Note When deconvolving OAI images, select the More Options button on the Deconvolve window, and then select the Deconvolve Projections option in the More Options window.

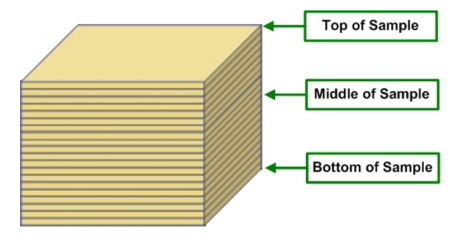
# Acquiring a Reference Image

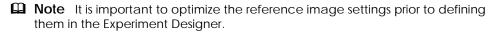
You can use either an alternate filter with the excitation light source or the transmitted light to acquire a reference image that can be combined with other images. This option is useful for Differential Interference Contrast (DIC) analysis. It can also be useful for other types of reference images.



A reference image of yeast cells

A reference image is a single image taken from a Z stack. You can specify to acquire the reference image at the top, middle, or bottom of the sample.





#### To create a reference image:

1. Set up a 3D Sectioning experiment (see *Sectioning* Specimens for 3D Images on Page 53).

- 2. In the Design/Run Experiment window, click the **Channels** tab and select the channels for your experiment (see *Selecting Filters* on Page 60).
- 3 Activate the **Reference Image** checkbox on the bottom-left side of the window.
- 3. In the Z position list, choose whether to acquire the reference image at the top, middle, or bottom of the sample.
- 4. Under the **Reference Image** checkbox, select which filters to use.
- 5. In the last field (bottom-right), select the light source to use for the image. **EX** is for a broadband light source (the xenon lamp or InsightSSI), **TRANS** is for transmitted light, and **LASER** is for a laser light source.

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زا <b>ک</b>		ted	nt name RatioT file size 234.11 julsition Fast	•		Gb Available)		Activate the <u>Reference Image</u> checkbox.
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I	Exp		EX Filter	EM Filter	%Т	EX Source		top, middle, or bottom of
<b>r</b>	[1.0000	•	DAPI	▼ DAPI	100%	▼ EX		the sample.
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	]1.0000	*	BLANK 1	▼ BLOCK	BLOCK	▼ SSI	-	for the reference image.
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]	Referenc	e In	GFP/mCherry nage Z Posit DAPI ure conditions	ion Middle of Sam	ple 100%			Select the type of light to use for the reference image.

This experiment specifies using the DAPI and Cy-5 filters to acquire images of each section and using the transmitted light to acquire a reference image using the DAPI filter in the middle of the sample.

6. Save and Run the Experiment.

After *DeltaVision* acquires the data, it creates two image files. One file contains the reference image data (with the extension \_R3D\_REF.dv) and the other file contains all of the other data.

# 6

# **Data Collection Techniques**

This chapter provides guidance and suggestions for:

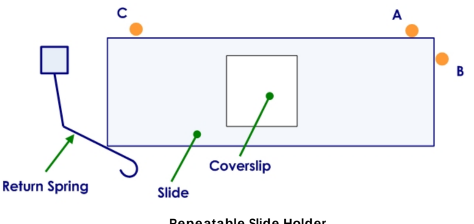
- Finding a Specimen and Recording its Position
- Finding Exposure Times
- Using Köhler and Critical Illumination
- Monitoring Data Acquisition
- Editing Experiment Macros

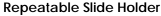
# Finding a Specimen and Recording its Slide Holder Position

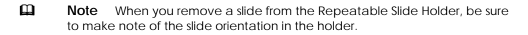
Recording the position of your slide is useful when you are conducting a point visiting experiment and you need to remove the slide before you are finished. Use the following instructions to find a sample and record its position on the slide with the Repeatable Slide Holder. When you resume your experiment, you can place the slide in the position that you recorded.

#### To find and center the specimen:

- 1. Choose the objective by rotating the objective turret. Be sure to select the same objective in the Resolve3D window.
  - Note If you are using an oil immersion lens, you must apply immersion oil to the sample. See Appendix A: The Oil Immersion Kit, for more information about the oil calculator.
- 2. Secure the slide. Since *DeltaVision* is an inverted microscope, the sample must be placed with the cover slip facing down toward the objective. If you are using a standard microscope slide (1" x 3" or 25 mm x 75 mm), you can use the Repeatable Slide Holder to hold the slide. To do this, pull the return spring to the left and place the slide cover slip down onto the Repeatable Slide Holder so that the upper right corner is pressed laterally against the brass locators (shown as **A** and **B** in the figure below) and the upper left edge is pressed laterally against the brass locator **C**. Then gently release the return spring to secure the slide. Lightly push down on the slide to make sure that it is fully seated in the holder.







- 3. Use the Coarse Focus knob to raise the objective.
- 4. Select the desired eyepiece (EP) filter.
- 5. Select the desired excitation (EX), neutral density (%T), and emission (EM) filters.
- 6. Rotate the Port Selector at the base of the microscope to direct the light collected by the objective to the Eyepiece ( $\checkmark$ ).

7. To use the transmitted light to locate the specimen, open the transmitted light shutter by pressing TRANS SHUTTER on the keypad. This allows white light to transilluminate the sample. (The intensity for the transmitted light is controlled using the %T field in the Resolve3D window.)

Alternatively, open the EX shutter and find the desired focal plane optically using either the focus knobs on the microscope or the Resolve3D Z stage controls.

- Note When you are sitting in front of the microscope, rotate the top of the microscope's focus knob toward you to move the objective up. Rotate the top of the microscope's focus knob away from you to move the objective down.
- 8. Use one of the following methods to maneuver the slide and find an area of interest:

#### Keypad

The buttons on the keypad are used for movement in the X, Y, and Z directions. Using the arrow keys on the keypad causes the stage to move by steps. The size of each step is doubled each time the STEP INCREASE button is pressed and halved each time the STEP DECREASE button is pressed. By adjusting the step size to the frame size, you can create a condition where each press of a step arrow will move the sample one frame. This is a convenient way to scan a large area for rare events (e.g., mitosis). Since the step movements are rapid, using the arrow keys in this way can be much less fatiguing than using the joystick.

#### Joystick

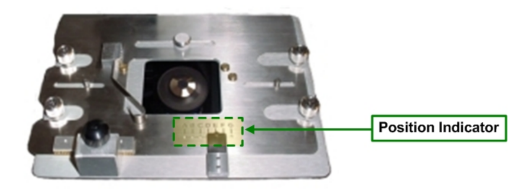
The joystick controls stage movement in the X and Y directions. Set the stage speed using the keypad SLOW, MEDIUM, or FAST buttons. MEDIUM is typically the best speed to start with. These buttons control the stage speed when you are using the joystick to move the stage.

#### Workstation

The Resolve3D module allows you to finely control stage motion. The arrow buttons move the stage in discrete increments that are indicated by the values in the **dX**, **dY**, and **dZ** text boxes. Resolve3D provides the only mechanism from which to obtain discrete step sizes.

**CAUTION:** To record and maintain accurate stage coordinates, focus by moving the stage with the Resolve3D controls instead of using the focus knobs to move the objective. Resolve3D cannot track the movement of the objective.

- 9. Tighten the Focus Lock on the left side of the focusing knob (see Page 32).
- 10. Use one of the methods from Step 8 to find the approximate center of the specimen and place it in the center of the field of view.
- 11. Direct light to the camera port using the Port Selector knob on the microscope base.
- 12. If you want to repeat the experiment using the same slide, record the position of the slide on the Position Indicator (the letter scale at the bottom of the Repeatable Slide Holder).



# **Finding Exposure Times**

The exposure time correlates the signal intensity level for the image acquisition. Finding the appropriate exposure time for each channel in an experiment is crucial to acquiring the best image data. Many factors must be considered. Although you want to see the maximum intensity in each wavelength, you must not saturate the camera or photobleach your sample.

High-Speed CCD Camera saturation occurs when the intensity values reach 4095 counts. The 12-bit High-Speed camera has an intensity range of 0 to 4095 counts. (The EMCCD 16-bit High-Speed camera has an intensity range of 0 to 65535 counts.) For successful deconvolution, a minimum intensity of 50 counts above background is recommended.

Note For live cell imaging, cells tolerate shorter exposures better than longer exposures, even at increased illumination intensity.

The response of increasing the exposure time is roughly linear. Therefore, if an exposure time of 0.2 sec results in a maximum intensity level of 1000 counts above background, then an exposure time of 0.4 sec will result in a maximum intensity level of approximately 2000 counts.

Use the following instructions to find the best exposure time for each excitation wavelength. This procedure should be performed for each excitation wavelength used in the experiment.

#### To find exposure time:

- 1. From the Resolve3D window, set the neutral density filter (in the %**T** field) to 100%. (For live cells, set this field to 10%.)
- 2. Set the exposure time to a low value. (0.5 seconds is recommended for live cell experiments.)
- 3. The **dZ** field represents the step size in microns.
  - Click in the **dZ** field.
  - Enter the desired value for the Z step.
- 4. Move through the area of interest by pressing **A** and **Y** in the Stage window portion of the Resolve3D window to move the stage in the Z direction.
- 5. Click the Acquire button.
- 6. Continue performing Steps 4 and 5 to obtain a sampling of Z sections. Find the maximum intensity in these sample images by observing the value of the **Max** field in the Resolve3D window. Typically, the focal plane with the highest maximum intensity value is the plane of optimal focus.
- 7. Move to the plane of optimal focus. This is the Z section with the highest intensity value.
- 8. From the Resolve3D window, choose View | Point List.
- 9. Click Mark Point to record its position.
- 10. Increase exposure time, but consider the amount of exposure the specimen can tolerate. Intensity values increase in direct proportion to exposure time.
  - Note There could be a higher intensity value in the image data that you did not sample. Therefore, do not set the exposure time to the limits of the sampled data because a higher intensity, un-sampled plane may saturate the camera.
     For fixed cell experiments, Max should be 50 60% of the saturation value. For live cell experiments, Max should be 2 3 times the Min (background),
- 11. Repeat Step 10 for all desired wavelengths.

# **Using Köhler and Critical Illumination**

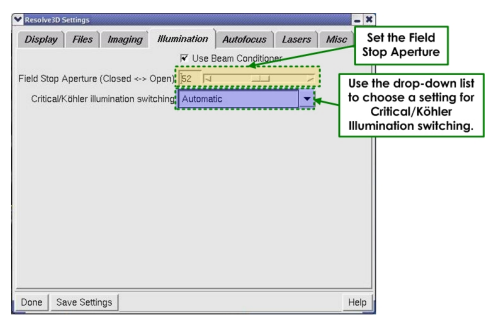
DeltaVision allows you to easily switch between two types of illumination:

- Köhler Illumination is the most commonly used form of illumination. It provides very even specimen illumination across the field-of-view. The light uniformly drops off as the distance from the focal plane increases.
- Critical Illumination directs the entire light source to the size of the detected area, and not the entire sample. With Critical Illumination, more light is directed to the focal plane and the out-of-focus light drops off more rapidly than in Köhler Illumination, based on the size of the field-of-view. Critical illumination also provides better axial (Z) and lateral (X,Y) contrast.

# Köhler/Critical Switching with API FI

If your *DeltaVision* system is using the Applied Precision Fluorescence Illuminator (API FI), use the following procedure to switch between Köhler and Critical illumination:

 On the Resolve3D main menu, click Settings and then select the Illumination tab.



#### Field Stop Aperture (Closed <-> Open)

This field allows you to set the field stop aperture to **Closed**, **Open**, or any percentage in between. You can either enter a percentage in the field or use the slider to set the percentage.

■ Note When the setting is set to Closed, the field stop aperture is about 20µm in diameter.

#### Critical/Köhler Illumination Switching

On an Applied Precision Fluorescence Illumination Module, there are three choices for the motorized Critical/Köhler illumination position:

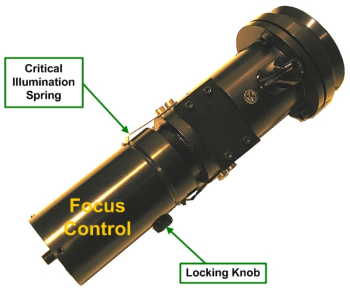
- Automatic DeltaVision automatically adjusts the position of the critical/Köhler illumination when the EX shutter is opened using the keypad (typically when the eyepieces are being used). The position is automatically returned to critical illumination when an image is requested, either via Resolve3D or by running an experiment.
- Always Critical DeltaVision moves the critical/Köhler motor to critical illumination and leaves it in its position regardless of the current shutter states.
- Always Köhler DeltaVision moves the critical/Köhler motor to Köhler illumination and leaves it in its position regardless of any new image requests or experiments running.

#### Köhler/Critical Switching with Olympus FI

If your *DeltaVision* system is using an Olympus Fluorescence Illuminator, you'll use the *DeltaVision* system's Fiber Optic Module (FOM) to align the light path from the fiber optic cable to the fluorescence illuminator. This module allows you to adjust the tilt, horizontal, and vertical orientation of the light path.

#### To switch to Critical Illumination:

Loosen the Locking Knob on the FOM and pull the Focus Control back (away from the stand) until it snaps into place in the Critical Illumination position. Then tighten the Locking Knob.



The FOM in Critical Illumination position, the Focus Control is extended so that the Critical Spring is in the groove.



**CAUTION:** When you are adjusting the Fiber Optic Module, be careful not to disconnect the Fiber Optic cable or to bend it in a diameter that is less than 24".

#### To switch to Köhler Illumination:

Loosen the Locking Knob on the FOM and push the Focus Control toward the stand until it snaps into place in the Köhler Illumination position. Then tighten the Locking Knob.



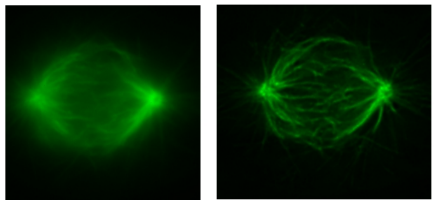
The FOM in the Köhler Illumination position, the Fiber Optic Module is pushed in so that the Köhler Illumination Spring is in the groove.

### **Monitoring Data Acquisition**

You can set options that control how images are displayed as they are acquired.

### **Viewing Deconvolved Image Previews**

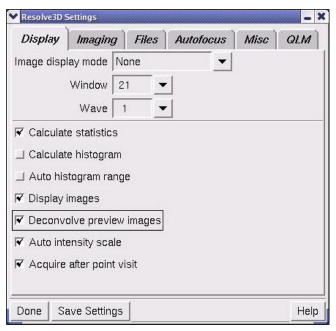
You can use 2D deconvolution to get a more accurate representation of what data will look like after deconvolution.



These images show how images are displayed during data acquisition, with and without Real-Time 2D Deconvolution.

#### To view previews of deconvolved images:

1. From the Resolve3D window, click **Settings** to open the Resolve3D Settings window. Then click the **Display** tab.



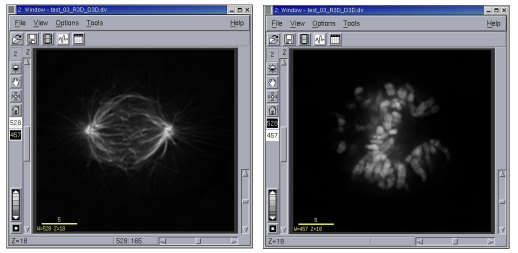
Resolve3D Settings Window | Display Tab

2. Activate the **Deconvolve preview images** checkbox.

Note This option does not provide a full iterative deconvolution, but it allows you to preview images as you collect them.

### **Selecting Viewing Modes**

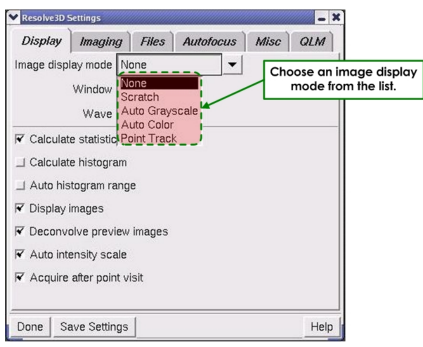
You can select from several modes for displaying images in the Data Collection window (or windows) as they are collected. You can choose other modes to display images in color, display each point in a point visiting experiment in a separate window, or to display each channel in a separate window, as shown below.



Auto Grayscale mode displays a separate window for each channel.

### To select the display mode:

- 1. From the Resolve3D window, click **Settings** to open the Resolve3D Settings window. Then click the **Display** tab.
- 2. On the **Image display mode** list, select a display mode.



**Resolve3D Settings Window** 

#### Image display modes

Mode	Description					
None	Displays images in the current window.					
Scratch	Displays all images in the default Data Collection window (Window 21).					
Auto Grayscale	Displays images in a separate window for each emission filter.					
Auto Color	Displays images in color as they are collected. When using Point Visiting, images are automatically displayed in separate windows for each point. This option should be used only when you are running an experiment.					
Point Track	Opens a separate window for each visited point in a point-visiting experiment.					

### **Displaying Statistics and the Histogram**

You can choose to display statistics and a histogram of the intensity values of each image as it is collected. These values are displayed below the Stage View in the Resolve3D window. The **Min**, **Max**, and **Mean** values are the minimum, maximum, and mean intensity values of the most recently acquired image. The histogram displays the intensity distribution.

### To display statistics and the histogram:

- 1. From the Resolve3D window, click **Settings** to open the Resolve3D Settings window.
- 2. Click the Display tab. Then select from the following options.

То	Select
Calculate image intensity statistics	Calculate Statistics.
	Note: You can improve readout speed by disabling this option.
Calculate and display an image intensity histogram	Calculate histogram
Automatically scale the histogram width for each	Auto Histogram Range
image that is analyzed.	Note: This option changes only the display of the histogram.
	It does not change the image data.

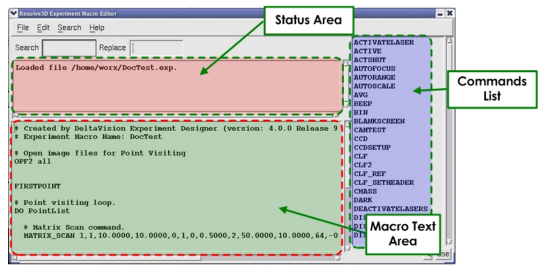
### **Editing Experiment Macros**

The Experiment Macro Editor is used to create or edit Resolve3D experiment macros (command scripts) that control the *DeltaVision* microscope. Most experiment macros are automatically generated using the Design/Run Experiment window, but you may need to create custom macros for certain types of experiments.

The best way to get started is to modify an existing macro with the Experiment Macro Editor. One approach is to use the Design/Run Experiment window to generate a macro and then edit it. Another approach is to use a reference macro.

### To open the Experiment Macro Editor:

- 1. On the Resolve3D window, click the Experiment button.
- 2. In the Design/Run Experiment window, choose **File** | **Edit** or the **Edit** button to open the Experiment Macro Editor.



**Experiment Macro Editor** 

### Search & Replace

These text fields are used to specify a search and replace pattern. (You will need to enter strings in these fields before you use the **Search** | **Replace Text** menu command.)

#### **Status Area**

The text area below the **Search** and **Replace** fields is used by the Macro Editor to provide status and command information.

### Macro Text Area

The Macro Text Area below the Status Area is the main working area of the Macro Editor.

#### **Commands List**

This is the list of available commands. When you highlight a command in the list by clicking it, you are provided with information about the command in the Status Area. If you double-click or select the **Use Item** button below this list, the command is inserted in the Macro Text Area at the current cursor location.

#### To edit a reference macro:

1. On the Resolve3D Experiment Macro Editor window, choose **File** | **Open Reference Macro**. The Open Experiment Macro File window is displayed.

/home/worx/*į́	
Directories 	Files Exp01.exp Resolve3D.exp Resolve3DPK.exp
.gconf .gconf .gconfd .gnome .gnome2	Resolve3DSettings_Misc_01.jpg Resolve3D_FW_LiveCell.exp Resolve3D_FW_Standard.exp Resolve3D_Startup.exp Resolve3D_startup.exp
Gelection	
/home/worx/Resolve3DPK.ex	4

#### **Open Experiment Macro File Window**

2. In the Open Experiment Macro File window, select a macro file and click **OK** to open it. The file is opened in the Reference Macro window.

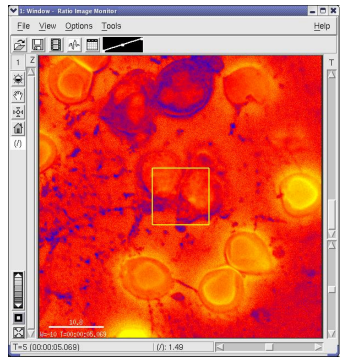
🗙 Reference Macro	_ ×
<u>File</u> <u>E</u> dit	
# Created by DeltaVision PhotoKinetics Designer (Resolve3DPK)	2
# Open the image file OPF	
# PhotoKinetics Scan request SCANPK 0.100,17,0,3,0,0,1,0,2,-1.000,0.050,0,3,1,3,32,1.000,120.000,4,1,0.500,1,0.000	
# Close the Image file CLF BEEP	
# #	
SI	

#### **Reference Macro Window**

- 3. Use the mouse to select the part of the reference macro that you want to copy.
- 4. On the Reference Macro window, choose Edit | Copy.
- 5. On the Resolve3D Experiment Macro Editor, choose **Edit** | **Paste** to copy the macro into the macro editor.
- 6. Edit and save the file as a new experiment macro.

### **Using Ratio Imaging**

*softWoRx* provides a ratio imaging acquisition function that allows you to view a graphic representation of the ratio of two channels as the images are being collected. In addition, a ratio graph displays the mean value (of an area in the middle of the image) vs. time. Both the ratio image and the ratio graph are for monitoring purposes only. The ratio imaging experiment results in a two-channel time-lapse image.



Sample image for ratio imaging experiment

#### To set up a ratio imaging experiment:

- 1. From the *softWoRx* main menu, select **File** | **Acquire (Resolve3D)** to open the Resolve3D window.
- 2. On the Resolve3D window, click the **Experiment** button to open the Design/Run Experiment window. Then, select the **Design Experiment** tab.

Acquire	Experiment	ttings	
Excitation	FITC	490 / 20	Click the
Emission	FITC	✓ 528 / 38	Experiment button.
Polychroic	Quad	▼ Info	
%Т	100%	•	
Illumination	SSI	•	
Exposure	[1.0000 <b>•</b> Find.	Focus	
Image size	512x512	🖌 🗌 Calibrate	
Lens	10X	▼ Info	
Bin	1x1 ·	- Aux. Ma	a

3. Select the **Sectioning** tab and unselect the **Z Sectioning** checkbox.

Design/Run Experin File	nent: *(modi	fied)	Select <u>Desia</u> <u>Experir</u> tab	gn nent			- Help
<i>Design Experin</i> Experimen Estimated f ⊔ Use Fast Acq	t name Re ile size 0.5	sign PK Experi solve3Dí 50 Mb Fast Acquisition			Gb Avail	<b>able)</b> en finished	
Sectioning	Channels	Time-lapse	Points	Panels	Plate	Actions	
Z Sectioning	5	Z Scan Opt	ions				
Focus point whe Optical section Number of op Sample th	spacing (µ	n) (b.co ns (b.d	Giet thick	ness			
	[	Deactivate <u>Z Sectionii</u> checkbo	the ng				

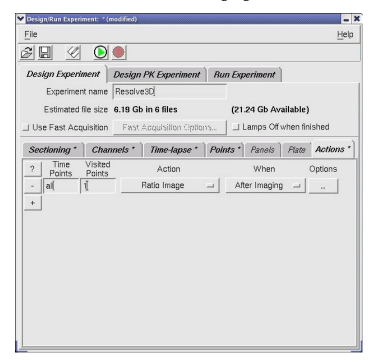
4. Select the **Channels** tab and specify the two channels you want to use for this experiment.

		Ø	2							
De	sign Exp	peril	ment Des	ign P	K Experiment	1	Bun Exp	eriment	1	
	Experi	imer	nt name Ratio	Test	1]					
	Estima	ted	file size 156.	07 MI	b in 12 files		(17.6	9 Gb A	vailable)	
n,	lse Fast	Á.CQ	ulsitlon Fa	ist Ac	quisition Optio	ns				
Se	ectioning	<i>,.</i>	Channels	• ]	Time-lapse *	P	oints *	Panels	Plate	Actions
ca	onventio	nal	Multiplexe	d ]						
	Ехр		EX Filter		EM Filter		%Т		EX Sourc	e
~	0.200	•	CY-5	-	CY-5	•	50%	•	EX	•
7	0.200	•	FITC	•	FITC	•	32%	•	EX	-
1	[1.0000	*	CY-S	+	CY-S	*	100%	*	\$\$1	-
_	[1.0000	٠	BLANK 1	-	DAPI	*	0.1%	*	SSI	*
1	[1.0000	*	BLANK 1	*	DAPI	*	0.1%	Ψ.	SSI	-
	Polychr	oic	Quad		•					

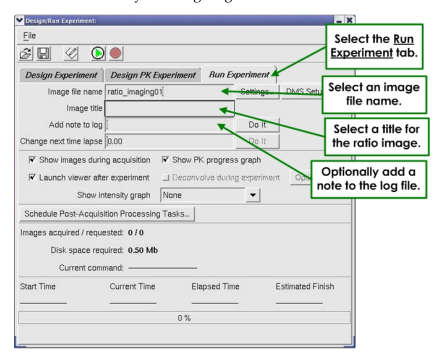
5. Select the **Time-lapse** tab and specify the time-lapse and total time for this experiment.

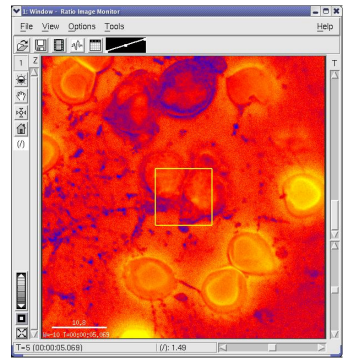
Design/Run Experiment: * (modified)	- X
<u>F</u> ile	<u>H</u> elp
Design Experiment Design PK Experi	iment Run Experiment
Experiment name Resolve3D	
Estimated file size 64.52 Mb	(20.59 Gb Available)
Use Fast Acquisition Fast Acquisition	n Options 📃 Lamps Off when finished
Sectioning * Channels * Time-lapse	Points Panels Plate Actions
✓ Time-lapse	
Hours Minutes Seconds Mill	iseconds
Time-lapse 10 10 10	
Total Time 10 10 10	
Time Points 1	
,	
Enable Cell Tracking	Cell Tracking Options
Maintain Focus with UltimateFocus	UltimateFocus Options
$\overline{\boldsymbol{v}}$ Image-based Autofocus before imaging	Autofocus Options

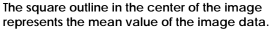
6. Select the **Actions** tab and select **Ratio Image** from the drop-down list of actions for this experiment. The **Time Points** field will default to **all** and the **When** control will be **After Imaging**.



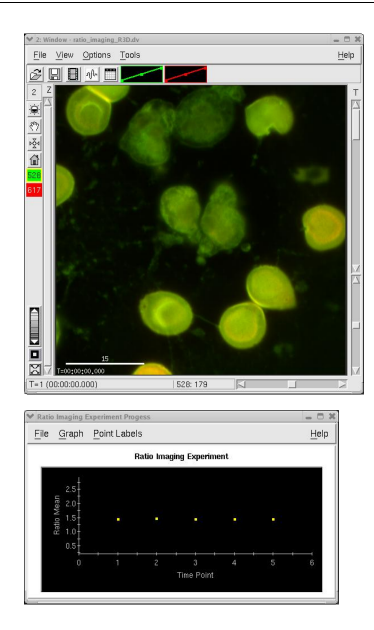
7. Select the **Run Experiment** tab and enter the image file name and a title for the ratio image. You can also enter text into the **Add note to log** field to include the text in your image log file.







When the ratio imaging experiment is complete, a two-channel time-lapse image is displayed along with a ratio graph showing the mean value (the outlined area in the middle of the image) vs. time. The square outline represents the portion of the image used to calculate the mean for a ratio graph.



## Using the Microtiter Stage Option

The addition of a Microtiter stage to a *DeltaVision* system enhances the imaging system's capability by providing the ability to scan microtiter plates. With a Microtiter stage on a *DeltaVision* system, you can design a plate scan by generating a point list that corresponds to the nominal center point of each well of interest, and then defining how the wells are to be sampled.

The model used for sampling plates is a matrix of N rows by M columns of *panels* for each well. Each panel can be a Z stack, a single Z, or an *OAI* (optical axis integration) scan.

Note These panels are never stitched together like the panels defined in the Panels tab of the Experiment window.

The panels can be separated by spacing defined through the software. You can specify autofocus for each individual well, or for every panel within the matrix. You can also set up parameters to specify the autofocus sampling range that represents the well-to-well and panel-to-panel variability of the focal plane.

Note Although it's possible to use Z stacks with panels, it is not recommended. The data sets can get extremely large very quickly.



Microtiter Stage for Delta Vision Systems

### Before Operating the Microtiter Stage

The Microtiter stage requires 3 hours of warm-up time before the stage has stabilized. This warm-up time is necessary even if you are using an environmental chamber.

- The motors need ample warm-up time due to the long lead screws.
- Significant X, Y, and Z drift has been observed prior to 3 hours of warm-up. To avoid this drift, a complete system warm-up is essential.
- You can help the system warm up faster by running a continuous time-lapse experiment with 6 points covering the span of a 96-well plate.
- Aurora and MatriCal Microtiter plates are recommended for use with the Microtiter stage.
- Cell culture plates are not of optical quality and therefore will not work with the Microtiter stage.

- If you are using the plate-scanning feature, an air objective is recommended. It is difficult to be successful scanning large areas of a plate using an oil objective.
- It is imperative for proper scanning autofocus that all of the objectives have the correct lens information attached to them in the Resolve3D.SYS file. If you click on the **Info** button next to the **Lens** dropdown list, all of the fields for the objective should be filled.

### Loading a Microtiter Plate

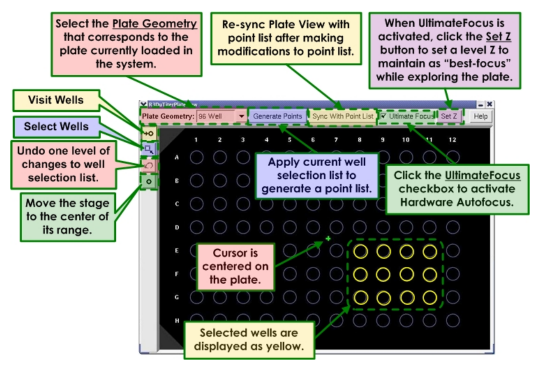
### To load a microtiter plate:

1. From the Resolve3D main menu, select the **Microtiter Plate** icon.

LIIISSION	p no		30	
Polychroic	Quad	✓ Info	J	
%T	100%	<b>_</b>		
Illumination	SSI	<b>_</b>		
Exposure	[1.0000 ▼ Fir	nd Focu	IS	
Image size	512x512	💽 🗆 Calib	orate	
Lens	10X	✓ Info	o	Select the
Bin	1x1	💌 🗆 Aux.	Mag.	Microtiter
Pixel size	0.6680 µm			Plate icon.
<ul> <li></li></ul>		HZ (0.50		
			Z	

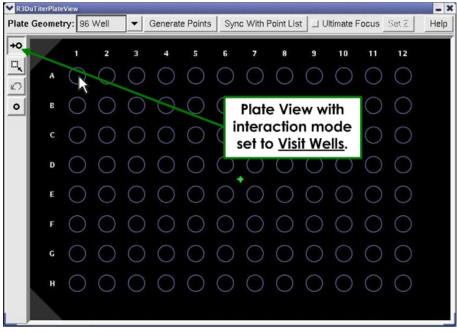
Resolve3D | Microtiter Plate Icon

A representation of the microtiter plate is displayed in the Plate View window. The following illustration provides an overview of the functions available from the Plate View window.



#### Plate View Overview

The Plate View window initially opens with the interaction mode set to **Visit Wells** as shown.

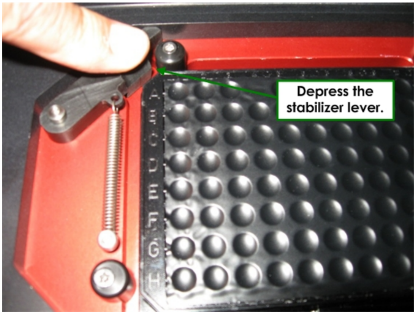


### Plate View Window – Visit Wells

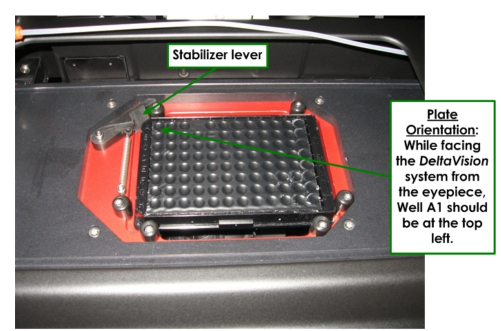
- 2. Load a plate into the plate carrier as shown:
  - 1. Insert the right edge of the selected plate onto the right edge of the plate carrier.



2. Depress the spring-loaded stabilizer in the top-left corner of the plate carrier.



3. Release the stabilizer so that it rests against the corner of the plate.



3. When loaded properly, the microtiter plate should be level in the plate carrier.

### **Exploring a Microtiter Plate**

### To explore a microtiter plate:

 From the Resolve3D main menu, select the Microtiter Plate icon. A representation of the microtiter plate is displayed in the Plate View window.

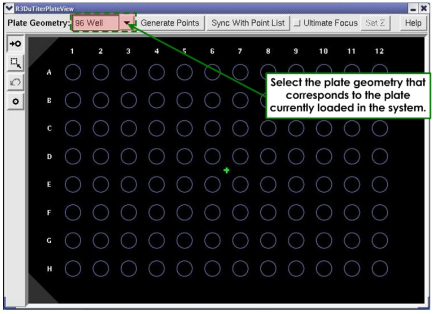


Plate Geometry Overlay

- 2. From the **Plate Geometry Overlay** field, select the plate geometry that corresponds to the plate you have loaded into the system for this experiment.
- 3. Click the **Visit** icon to put the Plate View window in Visit mode.
- 4. Click on a well to visit the nominal X/Y center of the well.
- 5. Through the oculars, perform a rough focus on the first well of the plate using the microscope's focus knobs.
- Note After obtaining a rough focus on the first well, you should not need to use the focus knobs.
- 6. Click the **Autofocus** AF icon from the Resolve3D window to activate Autofocus.

If the UltimateFocus Module is installed on your system, you can also set up the UltimateFocus feature to help maintain the plate focus. See *Using UltimateFocus* on Page 143 for more information.

- Note If you're still having trouble focusing, you may need to adjust the setting in the Maximum Z test range (μm) field (Settings window | Autofocus tab).
- 7. Explore the plate to set up filters, exposure times, binning, and the rest of the variables you would normally configure for an acquisition.

### **Designing a Plate Scan Experiment**

### To design an acquisition for a microtiter plate:

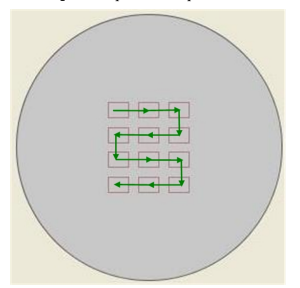
- Note Before you begin designing a plate acquisition, be sure to wait until the system has completely stabilized.
- 1. From the Resolve3D window, click **Experiment** to display the Design/Run Experiment window.
- 2. Click the **Design Experiment** tab.

File		
28 🖉 🔍 🌒	Select the Des	ign Experiment tab.
Design Experiment	Experiment	
Experiment name Reso	lve3Dį́	-
Estimated file size 2.01 I	db	(124.21 Gb Available)
Vuse Fast Acquisition Fast	st Acquisition Options	Lamp Off when finished
Sectioning * Channels	Time-lapse Points	Panels Plate Actions
Z Sectioning	Z Scan Options	
Focus point when scan starts	Middle of Sample	<b>•</b>
Optical section spacing	0.20	
Number of optical sections	<u></u> [64	
Sample thickness	[12.80 Get thickr	ness
	Enable OAI Scan	

- 3. Set up Z sectioning and channels as usual. The experiment design elements of **Time Lapse**, **Points**, **Panels**, and **Actions** are incompatible with Plate Scanning. These features are not used in this process.
- 4. Select the **Plate** tab and activate the **Do Microtiter Plate Scanning** checkbox.

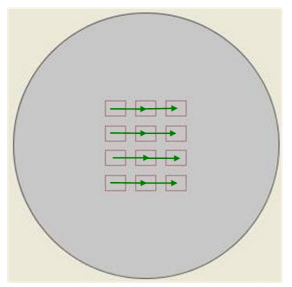
💙 Design/Run Exp	eriment: * (r	nodified)			,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	_ ×			
File							Help			
28 4										
Design Expe	eriment	Design PK Experin	nent Rui	n Experiment	ľ.					
Experim	nent name	Resolve3Dį				_				
		32.01 Mb	0-1	(20.58 Gb Av			-	ect the <u>e</u> Tab.		
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Sectioning ✓ Do Microtit		nels   Time-lapse	Points Plate View	Panels Plate				he <u>Do M</u> ing che		
	f panels pe							of rows o each w		
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Pa	nel Scan P	attern Raster 🔺		5) Select	scan	patte	rn for f	he pane	əls.	
🔽 Do Auto	ofocus							Do Aut		
Au	utofocus M	ethod Image Contra	ast (Software	e) 🗸 🗸	· · · ·			o activa us optio		
Autof	ocus Freq	uency Once Per Wi	ell	- More	e Setting	Met C	thod fie ontras iutofoc	e Autofo eld, use <u>t</u> for soft cus, <u>Ultin</u>	<u>Image</u> ware <u>nate</u>	2
[		utofocus Freque Per Well only c						or hardv cus, or <u>B</u>		

- 5. In the **Rows of panels per well** and **Columns of panels per well** fields, specify the number of rows and columns of panels to be used to sample each well's contents. Keep in mind that using large numbers of panels can generate huge amounts of data.
- **Note** Panels are saved as time points within one image.
- 6. In the **Panel Scan Pattern** field, select **Serpentine** or **Raster**. The **Serpentine** panel scan pattern is bi-directional.



Serpentine Panel Scan Pattern

7. The Raster panel scan pattern is unidirectional.

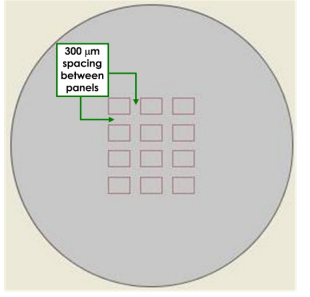


### **Raster Panel Scan Pattern**

8. The length of time it takes to perform a Serpentine scan is less than the length of time for performing a Raster scan. This is because, with a raster scan, the microtiter stage must return to the beginning of each row of panels to start

the next row. However, due to the unavoidable hysteresis involved with the bi-directional Serpentine scan, a Raster scan will always be the more accurate.

- Note *Hysteresis* refers to the position differential of the stage when approaching a specific location from different directions due to mechanical conditions. In this case, the change in the direction of the scan results in a very slight, but nevertheless measurable, position difference between similar locations visited from the opposite direction.
- 9. On the positive side, using a Serpentine scan pattern can actually increase the reliability of panel-to-panel autofocus.
- Note It is recommended that the Serpentine scan pattern is used whenever the experiment is relying on autofocus to scan.
- In the Panel Separation field, specify the distance (in μm) for panel separation. The single value is used for both horizontal and vertical separation of panels. The following graphic uses 300 μm as an example.



#### **Panel Separation**

- 11. Select the **Do Autofocus** checkbox to activate the autofocus fields.
- 12. In the **Autofocus Method** field, choose an appropriate Autofocus model to use for this acquisition. The options for this field are as follows:
  - **UltimateFocus (Hardware)** This method uses a laser (hardware) to perform autofocus on the sample.
  - **Image Contrast (Software)** This method uses the software autofocus feature to perform autofocus on the sample.
  - **Both** The UltimateFocus <u>and then</u> Image Contrast methods are used. If you would like to modify the Image Contrast search parameters, click **Settings** on the Resolve3D control panel and see the **Autofocus** tab for Plate Scan parameters. The "long" range is used when doing the **Once Per Well**

model or on the first panel of an **Each Panel** Autofocus style. The "short" range is used when focusing on each panel within a well. These values are retained, but are not part of the experiment macro.

- 13. In the **Autofocus Frequency** field, select how often you want the autofocus process to occur for this acquisition. The options for this field are as follows:
  - Once Per Well the stage moves to the center of the well, Autofocus finds the best plane of focus, and then the stage moves back to the first panel of the well.

• **Each Panel** – Autofocus is used on every panel in every selected well. For the default panel separation of 10  $\mu$ m, you may not need to use autofocus on every panel. When autofocus is used on every panel, you run a greater risk of photo-bleaching the sample. Check to be sure you need to autofocus every panel by scanning only the first well in your experiment.

14. To adjust the Autofocus parameters, click the **More Settings** button next to the **Autofocus Frequency** field.

Design/Run Experiment: * (modified)	- *
File	Help
Design Experiment Design PK Experiment Run Experim	nent
Experiment name Resolve3Dj	
Estimated file size 32.01 Mb (20.58 0	Sb Available)
Use Fast Acquisition Fast Acquisition Options	os Off when finished
Sectioning * Channels Time-lapse Points Panels	Plate * Actions
To Microtiter Plate Scanning Microtiter Plate Viewer	
Rows of panels per well $\left[1 \frac{\lambda}{2}\right]$	
Columns of panels per well	
Panel separation (microns)	
Panel Scan Pattern Raster	
I Do Autofocus	
Autofocus Method Image Contrast (Software)	Click the <u>More</u>
Autofocus Frequency Once Per Well	More Settings
	adjust autofocus
	parameters.

15. The Plate Autofocus Settings window is displayed.

Plate Autofocus Settings	
- Software Autofocus Options	3
Channel for Autofocus	DAPI 🗸
Autofocus Exposure	Į́0.500
Plate Scan Focus Binning	2
Plate Scan Long Focus Range	<u></u> 40.00
Plate Scan Short Focus Range	40.00

The fields in the Plate Autofocus Settings window are used as follows:

- Channel for Autofocus: Select the desired channel for autofocus. You are selecting the EX Filter. The parameters for EM and ND are taken from the last image acquired with that EX filter in the Resolve3D window.
- Autofocus Exposure: Set the autofocus exposure time. You will want to use the same or a shorter exposure time than what will be used during the acquisition time in your experiment. In order to make sure that your Autofocus Exposure time is not longer than necessary, test the exposure time using the same binning value in the Resolve3D Window.
- Note You may have to adjust the Resolve3D Autofocus settings in order to find the best focal plane using the **Autofocus** icon. Remember that the Plate Autofocus Settings are not the same as the Resolve3D Autofocus settings.
- Plate Scan Focus Binning: The default is set to a bin of 2x2. It is recommended to use binning for the autofocus for most samples in order to reduce the amount of light exposure to the cells. Please enter a single digit for this value. A Plate Scan Focus Binning of 2 is similar to a 2x2 in the Resolve3D window.

Note The software is set up so that you can autofocus with a different bin than acquisition. For example, you can autofocus with a binning of 2x2 and acquire data with a binning of 1x1.

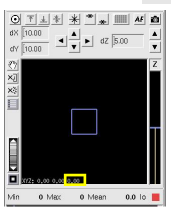
- Plate Scan Long Focus Range: This value is the Maximum Z Test Range for the first image in every well. If autofocusing Once Per Well, this is the only Focus Range that is used for the experiment. If autofocusing Each Panel, this Focus Range is used only on the first panel in each well.
  - The **Plate Scan Long Focus Range** is combined with the Lens' Depth of Focus in order to find the best focal plane during the scan. It is imperative

that the correct lens has been selected in the Resolve3D window and that the Lens has a valid Lens ID.

- You cannot change the Z Step Size for the **Plate Scan Long Focus Range**. The system will take the same sized steps no matter the value of the **Plate Scan Long Focus Range**. Therefore, a scan of 100 microns will take twice the amount of time when compared to a scan of 50 microns.
- When you are imaging multiple wells, the Plate Scan Autofocus starts from the best plane of focus from the previously scanned well. It then searches for the best signal-to-noise ratio in the distance ½ of the value of the **Plate Scan Long Focus Range** above that focal plane and ½ of the value of the **Plate Scan Long Focus Range** below that focal plane.

### To determine an estimate for the Long Focus Range:

In order to determine a good estimate for the **Plate Scan Long Focus Range** value, you will need to manually determine the distance between focal planes in the four corners of your plate (or the 4 corners of the wells filled within your plate).



•

Wells A1 and A2:

• Use the Resolve3D Window to find the best focal plane in A1.

Note: The microscope's focus knobs should <u>not</u> be used at this point.

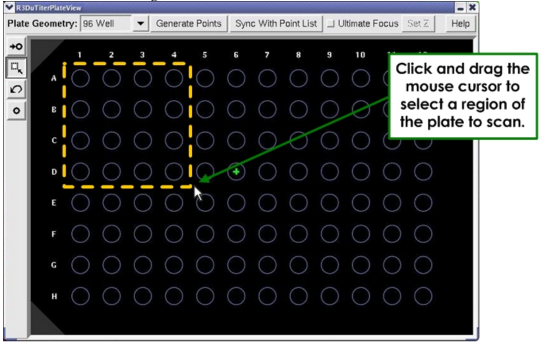
- Write down the Z coordinate from the Resolve3D
   Window (location highlighted in the yellow box to the left).
- Move to A2. Use the Resolve3D Window to find the best focal plane.
- o Subtract: Z(focal plane A2) Z(focal plane A1)
- Record the difference.
- Repeat for Wells A11 and A12.
  - Don't be concerned about the focal plane difference from A2 to A11. The focal plane will adjust as you move along Row A.
- Repeat for Wells H11 and H12.
- Repeat for Wells H1 and H2.
- Find the largest distance between wells from the tests above. Double this value and enter it as the estimate for the **Plate Scan Long Focus Range**. Make sure that this value is within the working distance of

the objective.

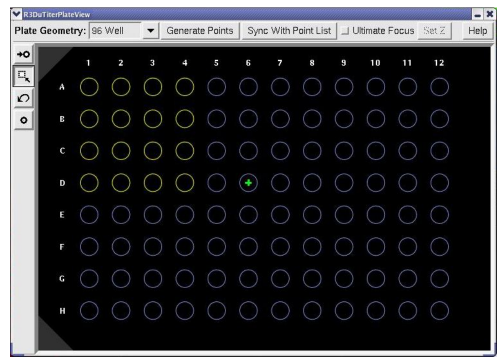
Plate Scan Short Focus Range: This is the Maximum Z Test Range between panels in a plate scan. If imaging more than one panel in each well, this value will be used for each panel after the first one.

- The Plate Scan Short Focus Range is combined with the Lens' Depth of Focus in order to find the best focal plane within panels during the scan. It is imperative that the correct lens has been selected in the Resolve3D window and that the lens has a valid Lens ID.
- You cannot change the Z Step Test Size for the Plate Scan Short Focus Range, which is the same as with the Plate Scan Long Focus Range.
- To determine if autofocusing within wells is necessary or to determine the value for the Plate Scan Short Focus Range, measure the change in Z throughout one well. Find the best focal planes in different fields of view within close proximity in a well and use the Z coordinates to find out the distance between focal planes. Double the largest difference in Z to get the Plate Scan Short Focus Range.
- Note If the distance between Z planes is not significant, you may want to consider autofocusing Once Per Well only. Running the autofocus once per well instead of per panel will save time for your experiment and will save your sample from the effects of photobleaching and/or phototoxicity.
- 16. Use the Plate View window to make a selection of wells to scan. From the Plate View window, click the icon to change to **Select** mode.
- 17. With the mouse, click and drag an outlined region to select a group of wells on the plate. Click and drag while holding the Shift key down to extend the

selection or to de-select specific wells. You can use the **Undo** icon to undo one level of changes to the well selection list.



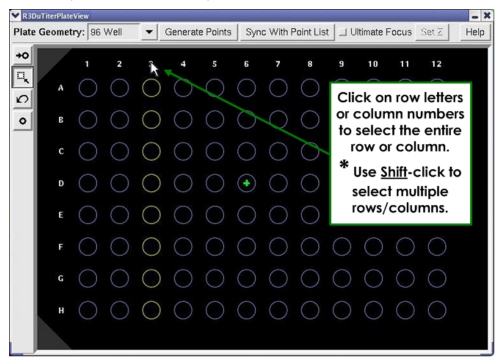
### **Plate Region Selection** 18. The selected wells are highlighted in the Plate View window.



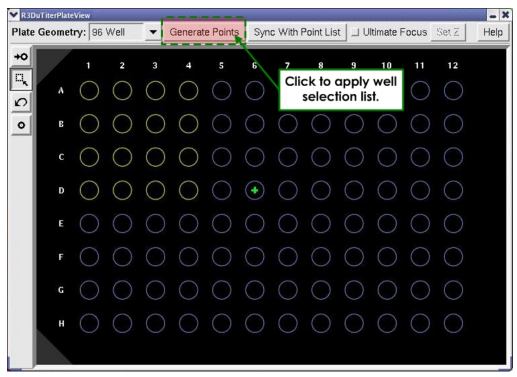
Selected wells are highlighted.

Alternatively, you can click on the row letters or column numbers to select the entire row or column.

Use this method in combination with the Shift key to select multiple rows or columns (or rows *and* columns).



- 19. When you are satisfied with the well selection, click the **Visit Wells** icon and click on the top left well in your selected list.
- 20. Bring this well into focus to set the best Z estimate for the start of the scan.
- 21. In the Plate View window, click **Generate Plate Point List** to apply your well selection list.



22. You can check your point list by selecting the **Marked Points List** icon in the Resolve3D window. The Point List window is displayed.

Mark Point	Visit Point	*	1:	+49500.00	-31500.00	+0.00	A01	1
Tyrearty Forne	wisiter onite	*	2:	+40500.00	-31500.00	+0.00	A02	
Delete Point	Replace Point	*	3:	+31500.00	-31500.00	+0.00	A03	
Deleterreinte	Tiopicoo Forne	*	4:	+31500.00	-22500.00	+0.00	B03	
Clear List	Compact List	*	5:	+40500.00	-22500.00	+0.00	в02	
Glear List	- vonprocurer	*	6:	+49500.00	-22500.00	+0.00	B01	
Save List	Open List	*	7 :	+49500.00	-13500.00	+0.00	C01	
ouve List	opencise	*	8:	+40500.00	-13500.00	+0.00	C02	
Calibrate All	Optimize List	*	9:	+31500.00	-13500.00	+0.00	CO3	
Galibrate All	opumze cise	*	10:	+31500.00	-4500.00	+0.00	D03	
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		*	<b>12</b> :	+49500.00	-4500.00	+0.00	D01	
Done	Help	1						

#### **Point List Window**

- 23. Notice in the Point List window that the points are labeled with the Well Row/Column specification.
- 24. Return to the Design/Run Experiment tool and save the experiment.

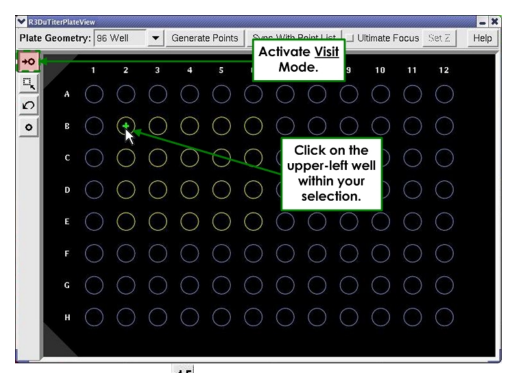
esign Experiment	Design PK Experiment Run Experiment
Experiment name	Resolve3Dj
Estimated file size	32.01 Mb (20.58 Gb Available)
Use Fast Acquisition	Fast Acquisition Options
Sectioning * Channe	els Time-lapse Points Panels Plate * Actions
Do Microtiter Plate Sca Rows of panels per	
	well $\overline{[1]} \xrightarrow{\lambda}$ well $\overline{[1]} \xrightarrow{\lambda}$ rons) $\overline{[10.00]}$
Rows of panels per Columns of panels per Panel separation (micr Panel Scan Pa	well $\boxed{1}$ $\frac{1}{7}$ well $\boxed{1}$ $\frac{1}{7}$ rons) $\boxed{10.00}$ ttern Raster

■ Note The saved Experiment Macro tells the system how you want the well scanning to be done. The Plate Point List tells the system what you want to scan. Since the Experiment Macro and the Plate Point List are relatively independent from one another, you can re-use the Experiment macro with a different list of selected wells, or you could re-use the list of selected wells and perform multiple styles of imaging on the same list.

### **Running a Microtiter Plate Scan**

### To run a scan:

1. For best Autofocus results, visit the first well (the upper-left corner) of your selection using the Plate Viewer. On the Marked Point list, this should be the top entry.



- 2. Use the **Autofocus** icon **AF** in the Resolve3D window to find the best focus. When you are satisfied with the focus, choose **Replace Point** from the Point List window to restate the Z position of that point.
- 3. In the Design/Run Experiment tool, provide a name for the experiment.

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File	Help
Design Experiment Desig	ign PK Experiment Run Experiment the experime
Experiment name Reso	
Estimated file sze 32.01	1 Mb (20.58 Gb Available)
🔟 Use Past Acquisit <mark>on 🔤 F</mark> a	Click the <u>Run</u> icon to start the scan. d
Sectioning * Channels	Time-lapse Points Panels Plate Actions
▼ Do Microtiter Plate Scannin	ng Microtiter Plate Viewer
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Columns of panels per well Panel separation (microns)	
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Panel separation (microns)	[10.00
Panel separation (microns) Panel Scan Pattern	[10.00 Raster

Resolve3D automatically creates a subdirectory of your current data folder with the name you provide here. All of the scan data for this experiment are placed in that folder.

4. Click the **Run** () icon to start the scan.

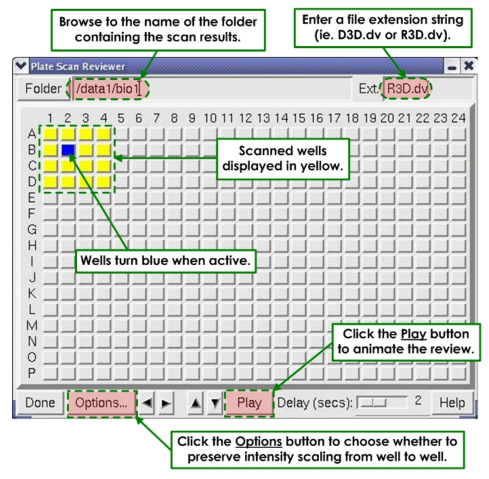
You can stop the scan at any time using the **Stop** sicon. When you abort the scan in this manner, scan data is lost for all unfinished wells.

The most effective way of monitoring the scan is using the Plate Viewer. It updates its current location indicator whenever it moves in X and Y.

### **Reviewing a Finished Plate Scan**

### To review a finished scan:

1. From the *softWoRx* main menu bar, select **Utilities** | **Review Plate Scan**. The Plate Scan Reviewer window is displayed.

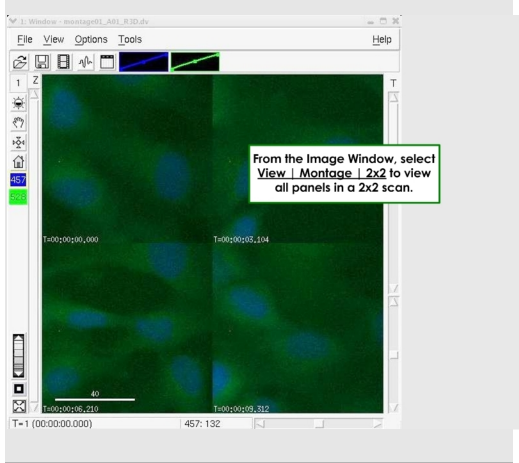


- 2. Click the Folder button to select the folder where you saved the scan.
- 3. If the data is already processed, you may need to specify the file extension string (like D3D.dv or R3D.dv) to view the files of interest. If files are

found that adhere to the plate scan file naming convention, they are displayed as yellow buttons on the Plate Scan Reviewer tool. Click on a yellow button to open the Image Viewer and display an image from that well. The button turns blue when active.

4. Select the **Play** button to animate the review. The Play mode steps through all of the images in order.

**WHint:** Use the Image Window's Montage View to view all of the panels within a well. If, for instance, you scanned 2x2 panels, choose **View | Montage | 2x2** to see all the regions. This is a "Montage by Time" situation since the panels are considered the Time dimension of the *DeltaVision* file.



### **Re-scanning Selected Wells**

If you determine that you want to re-scan some of the wells (due to Autofocus miscalculations, etc.), you can re-scan an individual well or group of wells using the same objective and experiment macro. The difference is that you'll be modifying the Plate Point List to select only the well(s) to be re-scanned.

Note For instructions on re-scanning selected wells using a *different* objective, refer to the next section (below).

### To re-scan selected wells:

- 1. From the Plate View window, select the well or wells to be re-scanned and click **Generate Plate Point List** to replace the current Marked Point list with this modified one. Remember, that the points are generated with the current stage Z value, so make sure the *DeltaVision* system is in focus.
- 2. In the Design/Run Experiment window, use the same experiment name as you used the first time. You may be prompted to re-save the experiment since the point list has changed.
- 3. From the Design/Run Experiment window, click the **Run** window icon to restart the scan. You are prompted to confirm that you want to overwrite the file(s) you scanned before.

# Re-scanning Selected Panels with a Different Objective

If you determine that you want to re-scan some panels with a different objective (i.e. you scanned the plate with an air objective, but you'd like to reimage with a 60x oil objective) you will need to edit the coordinates of a points list in order to return to the panel location.

Since the field of view is different for each objective magnification, you cannot simply rescan the panels. The method below describes the process of going back to the center coordinates of your image. If the object of interest isn't in the center of the field of view, you may have to make small movements to search for it. The region of interest will be within the field of view of the oculars, so it may also be helpful to switch to the oculars to locate it.

### To re-scan selected panels with a different objective:

Find the coordinates for your panel of interest as follows:

- 1. In the Resolve3D log file (\*R3D.dv.log), find the image of interest. It will be displayed as a time point. Write down the time point number.
- 2. To find the image number, multiply the time point number by the number of wavelengths acquired in the experiment (for example, if the time point is 25 and you have DAPI and FITC in your channel, the image number is 50).

	bio1_A03_R3D.dv.log - KEdit	- • ×
<u>File</u> <u>E</u> dit <u>G</u>	o Tools Settings Help	
3 🗁	🖬 🗁 🔗 🤌 🗣 🗇 👘 🕵	
Image 49.		ultiply the time point number
Image 50.	Time: Tue Sep 8 10:16:16 2009 Time Point: 0.000 secs Intensity: Min = 204, Max = 8268, Mean = 1593.22107 Photosensor reading: 626203.5	to find the Image Number.
	Exposure time: 1.000 secs EX filter: CFP EM filter: CFP ND filter: 100% [Stage coordinates: (+31500.000,-31500.000,-3.400)]	Coordinates for yo panel of interest a displayed.
Image 51.	Time: Tue Sep 8 10:16:17 2009 Time Point: 0.000 secs Intensity: Min = 204, Max = 8264, Mean = 1593.22485 Photosensor reading: 626203.5 Exposure time: 1.000 secs EX filter: CFP EM filter: CFP ND filter: 100% Stage coordinates: (+31500.000,-31500.000,-3.600)	
Image 52.	Time: Tue Sep 8 10:16:17 2009 Time Point: 0.000 secs	
	T . '. W' DOA W DOGD W 1500 04000	

- 3. Every experiment acquired creates a log file (image\_name.dv.log). Open the log file for your image.
- 4. Find the image number calculated in Step 1 and write down the X, Y, and Z coordinates for all of the points that you would like to revisit.

Note It's critically important that you include both positive (+) and negative (-) indicators when noting coordinates.

Edit the Point List as follows:

- 5. From the Resolve3D window, click on the **Marked Points List** icon. The Point List window is displayed.
- 6. Press the **Clear List** button.
- 7. Use the Joystick to navigate to a location in X, Y to revisit with the new objective.
- 8. Find the best plane of focus.
- Note Upon returning to a previously visited point, you may need to search around to find the exact location. Also you will always need to adjust the Z focus. This is especially true when you change objectives.
- 9. Press the Mark Point button.
- 10. Repeat Steps 3 through 6 for as many points as you want to revisit.

Click the <u>Clear</u> List button.		k the <u>Ma</u> nt Button		Click the <u>List</u> Butt			
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Delete Point	Replace Peint	* 4:	+31500.00	-22500.00	+0.00	B03	
Clear List	Compact List	* 5:	+40500.00	-22500.00	+0.00	в02	
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							17
Done	Help	100			_		- "
		<b>I</b> A					1

11. Click on the **Save List** button. The Save Point List File window is displayed.

Preview:	Directory //home/	/worx	è 🕂 🔳 ?
	Date ^	File Name	
Created (acquire  Last Modified:	[07/02/10 14:19] [10/01/10 09:07] [11/08/10 10:20]	softworx-logs/	a file name
 File Size:  Image Info:	A		Point List.
	File name		OK

12. Give the Point List a Name and click OK.

### Adding Overlays to the Microtiter Overlay Window

Your system was set up with the overlay options of the 96-well and 384-well plates. The instructions in this section describe the method for adding 2, 4, 6, 12, 24, and 48-well plates to the Microtiter Overlay window.

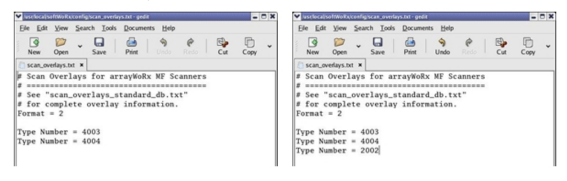
### To add plate overlays to the Overlay window:

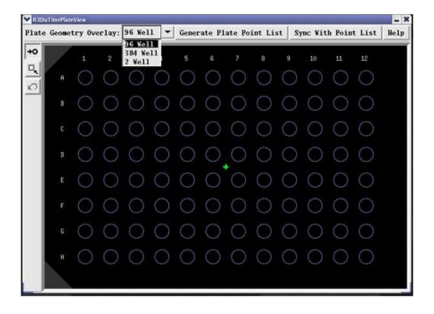
- 13. Open the scan\_overlay\_standard.db.txt file from the /usr/local/softWoRx/config folder.
- 14. Locate the desired plate configuration in the **Name** field.
- 15. Record the **Type Number** corresponding to the plate configuration. For example, if you were to add the 2-well plate, you would record a type number of 2002.

- 16. Save a backup copy of the scan\_overlays.txt file in the /usr/local/softWoRx/config folder.
- 17. Open the scan\_overlays.txt file.
- 18. Add "Type Number = xxxx" for the overlay(s) that you would like to add, where xxxx is the **Type Number** recorded in Step 3.

**Note** Do <u>not</u> alter the line in the scan\_overlays.txt file that reads **Format = 2**.

19. Add more overlays in the same manner.





### Using UltimateFocus<sup>®</sup>

If your *DeltaVision* system is equipped with the UltimateFocus Module, you can use this feature to help maintain focus within a specific move threshold. The UltimateFocus Module is always on and is enabled through a checkbox toggle available at key points from within the *softWoRx* software. You can use UltimateFocus either while exploring a sample, or when collecting experimental data.

After the initial plane of focus is determined, you can then scan the sample area with the UltimateFocus correction mechanism applied as follows:

1. From the Resolve3D main menu, select **File** | **Continuous Acquire**. The Continuous Acquire window is displayed.

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	ocus [1 [0.100	

2. Activate the **Use UltimateFocus** checkbox.

Continuous Acquire		re	Activate the <u>Use</u> <u>UltimateFocus</u> checkbox.		
Mainta Move TI	ain every	[1 [0.10] [1	frames 0	-	Other UltimateFocus options are activated.
Delay Ti	me (sec)	0.0			
Done	Start Ac	quirin	g Images		

### Using the Microscope's Oculars:

The default parameters for the Continuous Acquire window are shown in the example above. With these settings, you can scan the sample looking through the oculars, with the joystick speed set to SLOW.

### **During Camera Acquisition:**

When you press the **Start Acquiring Images** button on the Continuous Acquire window, the camera collects images using the excitation and exposure settings currently selected in the Resolve3D main menu.

From the Continuous Acquire window, you can adjust the following focus parameters:

- The Maintain every \_\_\_\_\_ frames field defines how often the stage corrections for UltimateFocus are made. Entering 1 tells *softWoRx* to perform the Move Threshold check for every image, entering 2 indicates performing the process for every other image acquired, and so on. Entering a higher number in this field will increase the speed of your acquisitions.
- The Move Threshold field defines the minimum Z position change from the point of origin. If the change in the Z position is less than the value indicated in the Move Threshold field, the position of the stage *will not* change. If the change in position is greater than the Move Threshold, the stage *will* move to correct the Z position. The default value is set to 0.100 microns.
- Note For experiments in which fast acquisition speed is the most important factor, limiting the number of corrections will increase the speed of the image acquisition.
- The Max Iterations field defines the number of stage corrections the UltimateFocus module will make using the Move Threshold. This field sets the upper limit for corrective moves allowed to the stage position at one time. Limiting the number of iterations increases the speed of your image acquisitions at the possible cost of focus accuracy. For fast imaging, it is recommended that the setting in the Max Iterations field be set to no greater than 1.

## **Monitoring Stage Corrections**

In the following Resolve3D main menu example, a sample was scanned using the UltimateFocus module. The Message Pane at the bottom of the menu lists each correction to the focal plane made by the module.

Resolve	3D	_ ×		
<u>File</u> <u>View</u>	v Options Calibration	n <u>H</u> elp		
Acquire	Experiment Setting	S		
Excitation	561Laser 💌	561 / 4.00		
Emission	mCherry 💌	63Z / 60		
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Exposure	0.1000 🔻 Find	Calibrate		
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## **Tips for Focus Maintenance**

#### Focus Maintenance with Long Time-lapse Experiments

- For time-lapse intervals much greater than the imaging time requirement.

A common problem with long time-lapse experiments has been that objects drift out of Z focus due to physical changes in the microscope stand and stage as a result of environmental conditions. To address this, softWoRx has the ability to allow periodic adjustments to be done *during* the time-lapse wait interval. To accomplish this, you can specify an UltimateFocus Action to be triggered for all time points, and further to use the **When** control to tell the system to perform the focus maintenance periodically *During* the time-lapse wait. It will be necessary to toggle the **Maintain Focus with UltimateFocus** button **Off** in the **Time-lapse** tab as the Action will be sufficient to maintain the focus. This should provide maximum flexibility to minimize Z drift focus artifacts in the experiment.

#### Focus Maintenance with Medium Time-lapse Experiments

- For short time-lapse intervals that are safely greater than the imaging time requirement.

In this scenario, you can specify **Maintain Focus with UltimateFocus** in the **Time-lapse** tab of the Experiment Designer. When you encounter this situation while running the experiment, the system will determine whether it has enough time to do the UltimateFocus maintenance and if so, do it *just before* the time lapse expires and imaging starts. The idea is to get focus adjusted as close to the start of imaging as possible.

#### Focus Maintenance with Very Short Time-lapse Experiments

- For time-lapse intervals nearly the same or less than the imaging time requirement.

When imaging as fast as possible, it seems very unlikely that significant enough environmental drift would occur from one frame to the next to make a discernible difference in focus. Therefore, to maximize frame rate, you can easily design an experiment that triggers an UltimateFocus Action with a certain periodicity. For example, you can choose to do UltimateFocus maintenance **every 20** frames. You would need to toggle the **Maintain Focus with UltimateFocus** button **Off** in the **Time-lapse** tab of the Experiment Designer and add an UltimateFocus Action with **every 20** as the specification for which time points to apply the Action. This allows fairly fast imaging (as fast as conventional time lapse can do), but with an occasional UltimateFocus maintenance event to compensate for drift.

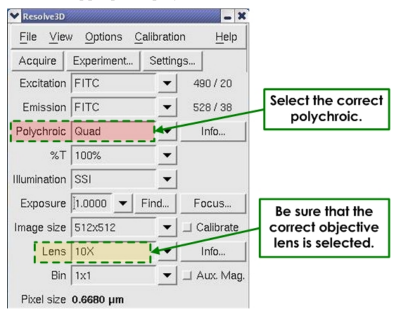
## Multi-well Plate Scanning with UltimateFocus

If your DeltaVision system is equipped with the optional Microtiter Stage and the UltimateFocus Module, the UltimateFocus can be used with the Microtiter stage to maintain focus when running a plate scanning experiment. This laserbased focus method monitors the change in the index of refraction as it goes through the coverslip and sample media. UltimateFocus will only perform well with an aqueous-based sample media.

UltimateFocus for plate scanning requires a calibration profile of the long-range focus for the plate. This profile is unique to each objective lens, polychroic, and plate type used on the DeltaVision. You'll need to perform a calibration for each objective / polychroic / plate-type combination. The profile is automatically saved for use in future experiments after you have collected the calibration. For best results, however, the calibration should be done each time the objective lens, polychroic, or plate type is changed. You'll use the sample plate to be scanned in the experiment to perform the calibration step. This will provide the best results.

#### To calibrate a multi-well plate:

- 1. Ensure that the correct objective lens is selected in the **Lens** field of the Resolve3D main menu.
- 2. Select the appropriate polychroic in the **Polychroic** field.



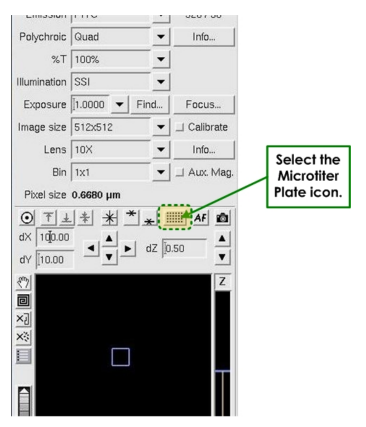
3. Click the **Settings** button in the Resolve3D main menu to open the Settings menu. Then activate the **Autofocus** tab.

Resolve3D Settings	Click the Autofocus tab to
Display Files Imaging Illumination Autofocus	ers Mis activate the tab.
Image Contrast Software Autofocus	
Channel for Autofocus BLANK1	
Contrast calculation method Auto	
Autofocus Z test step (µm) [3.12	
Maximum Z test range (µm) [50.00	
Post-autofocus Z offset (µm) 0.00	Click the Calibrate
UltimateFocus Hardware	<ul> <li><u>UltimateFocus for plate</u></li> </ul>
Calibrate UltimateFocus for plate scanning	scanning button.
Done Save Settings	Help

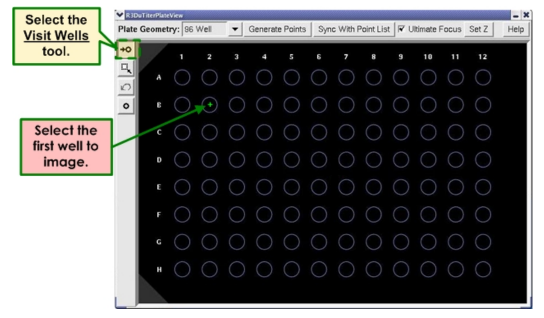
4. In the **Autofocus** tab, click the **Calibrate UltimateFocus for plate scanning** button. The UltimateFocus Plate Calibration window is displayed.

UltimateFocus Plate Calibration	×
UltimateFocus Plate Calibration Protocol	
1) Mount a microtiter plate containing sample in aqueous media	•
2) Find a well with material that you will be able to see, preferably on the well bottom.	У
3) Manually lower the microscope objective to its minimum posit	tio
<ol> <li>"Zero" the microtiter stage Z coordinate using the Home Stag button below.</li> </ol>	le
5) Using only the eyepieces and the coarse and fine focus knobs raise the objective to focus on the sample on the well bottom	
6) Verify that the Polychroic and Lens choices are correct in the main Resolve3D control panel.	
7) Choose the Calibrate Focus button below to perform the calibration that a few minutes to complete.	bra
Z Sweep Min, Max, Step Size 500.00 500.00 0.500	
	Click the Hom
Done Home Stage Z Calibrate Focus He	In Stage Z button

- 5. Follow the instructions in the UltimateFocus Plate Calibration window to complete the calibration process as in the steps that follow.
- 6. Place a multi-well plate on the microtiter stage. (See *Loading a Microtiter Plate* on Page 122 for details.)
- 7. From the Resolve3D main menu, select the **Microtiter Plate** icon.



A representation of the microtiter plate is displayed in the Plate View window.



- 8. Select the **Visit Wells** tool and then select a well with visible sample material, preferably at the bottom of the well.
- 9. Manually lower the objective turret to its minimum position.
- 10. Click the **Home Stage Z** button at the bottom of the Ultimate Focus Plate Calibration Protocol window to "zero" the microtiter stage Z coordinate.

- 11. Using the eyepieces and the microscope's manual focus controls, raise the objective to focus on the sample material at the bottom of the well.
- 12. Click the **Calibrate Focus** button at the bottom of the Ultimate Focus Plate Calibration Protocol window to start the calibration process.

**Note** This procedure may take a few minutes to complete.

VUltimateFocus Plate Calibration × UltimateFocus Plate Calibration Protocol 1) Mount a microtiter plate containing sample in aqueous media. 2) Find a well with material that you will be able to see, preferably on the well bottom. 3) Manually lower the microscope objective to its minimum positio 4) "Zero" the microtiter stage Z coordinate using the Home Stage button below. 5) Using only the eyepieces and the coarse and fine focus knobs, raise the objective to focus on the sample on the well bottom. 6) Verify that the Polychroic and Lens choices are correct in the main Resolve3D control panel. 7) Choose the Calibrate Focus button below to perform the calibra This procedure may take a few minutes to complete. Click the Z Sweep Min, Max, Step Size -500.00 0.500 500.00 Calibrate Focus button. Home Stage Z Calibrate Focus Done Help

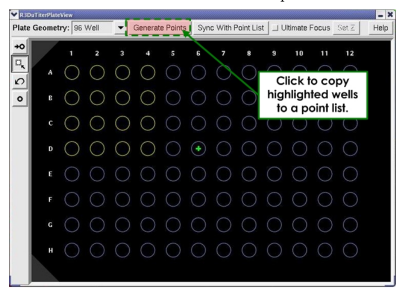
#### To set up and run an UltimateFocus plate experiment:

- 1. Ensure that the port selector is set to the **Camera** is setting.
- 2. Use the controls in Resolve3D to focus on the sample and refine exposure conditions for each channel in your experiment. If your experiment will be using Z sectioning, define the desired Z thickness.
- 3. In the Microtiter Plate View window, use the **Select Wells** mode to highlight the wells you want to image.



Click and drag while holding the Shift key down to extend the selection or to deselect specific wells. You can use the **Undo** icon to undo one level of changes to the well selection list.

- Note Alternatively, you can click on the row letters or column numbers to select the entire row or column. Use this method in combination with the Shift key to select multiple rows or columns (or rows *and* columns).
- 4. Click the Generate Points button at the top of the Plate View window.



- 5. To use UltimateFocus to explore various wells on the plate, activate the **UltimateFocus** checkbox at the top of the Plate View window.
- 6. Click the **Set Z** button to calibrate the current focus point of the current well as the desired Z position for the experiment. **Set Z** helps the system by setting a Z

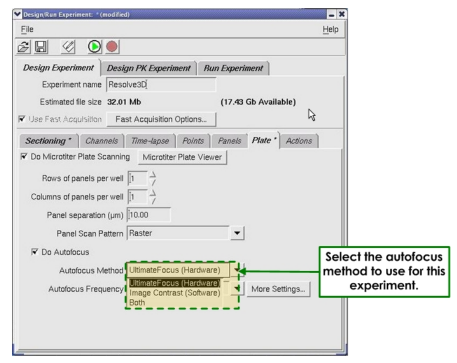
Activate the UltimateFocus checkbox. \_ X R3DuTiterPlateVie Plate Geometry: 96 Well 

Generate Points Sync With Point List 

UtimateFocus Set Z Help Click Set Z to +0 4 10 calibrate the current 5 8 9 12 6 Q, focus point as the  $\bigcirc$  $\bigcirc \bigcirc$  $\circ \circ$  $\bigcirc$ 0 desired Z position. 0 B () (+)  $\bigcirc \bigcirc$ () () $\bigcirc \bigcirc$  $\bigcirc$ 0000000000000

offset from the interface boundary to the plane of focus which you are interested in maintaining.

- 7. Select the first well in the region of the plate you want to scan and click the *AF* button on the Resolve3D main menu.
- 8. In the Resolve3D main menu, click **Experiment** to open the Design/Run Experiment window.
- 9. Set up the **Sectioning**, **Channels**, and **Plate** tabs. In the Plate tab, select the desired autofocus method as shown.



10. Click the **Run** licon to start the scan.

You can stop the scan at any time using the **Stop** icon. When you abort the scan in this manner, scan data is lost for all unfinished wells.

# acility Requirements & Components

## Facility Requirements & Components

This chapter describes the main components of the *DeltaVision* System. It includes the following sections:

- *Electrical and Environmental Requirements* describes the *DeltaVision* operating and facility requirements.
- *Overview of Components* shows the location of the key system components.
- *Optical Components* describes the system light sources, cameras, and filters.
- Desktop Components provides information about the monitor, the keypad and joystick, the vibration isolation table that supports the microscope, and other components.
- Cabinet Components describes the combined Instrument Controller and Microscope Interface Chassis (IC/MIC) and the Workstation.
- Other Standard Components describes the Repeatable Slide Holder, the Fiber Optic Module, the tool kit, and standard software.

- Optional Components describes DeltaVision options such as the X4 Laser Module, the EMCCD Camera, the Environmental Chamber, the Microtiter Stage, the InsightSSI solid state light source, the UltimateFocus Module, Differential Interference Contrast Components, and the *softWoRx Explorer* application.
- *Consumable Parts* lists the fuses and other components that you may need to replace to maintain the system.

## **Electrical and Environmental Requirements**

An important aspect of collecting high quality images is meeting the proper electrical and environmental requirements for the system.

## **Electrical Requirements**

#### Line Requirements

**Operating Frequency: 50/60 Hz** 

**Operating Power:** Up to three separate 100-120 VAC 15A circuits. (One for the Instrument Controller, a second one for the Environmental Chamber (if included), and a third one for the rest of the *DeltaVision* system.

Note If users want to have the *DeltaVision* workstation powered on with the rest of the system off, the workstation can be plugged into a power connection separate from the rest of the imaging system.

Up to three separate 200-240 VAC 6A circuits. (One for the Instrument Controller, a second one for the Environmental Chamber (if included), and a third one for the rest of the *DeltaVision* system.

**Transients:** Transient over-voltages in accordance with Installation Category II in IEC 664

Maximum Power: 1200 VA

#### **Power Cord Set Requirements**

The power cord set received with *DeltaVision* meets the requirements for use in the country where the equipment will be used.

#### **General Requirements**

The requirements listed below are applicable to all countries:

The length of the power cord set can be a maximum of 9.75 feet (3.0 m).

- All power cord sets must be approved by an acceptable accredited agency responsible for evaluation in the country where the power cord set will be used.
- The power cord set must have a minimum current capacity of 10A for 230 -240 VAC systems or 15A for 100-120 VAC systems, as required by each country's power system.
- The appliance coupler must meet the mechanical configuration of an EN60320/IEC 320 Standard Sheet C13 connector, for mating with the appliance inlet on the Isolation Transformer Assembly.

#### **Country-specific Requirements**

The following table shows the accredited agency and power cord set requirements for each country.

Country	Accredited Agency	Power Cord Set Requirements	
Australia	EANSW	The flexible cord must be <har> Type HO5VV-F, 3-</har>	
Austria	OVE	conductor, 1.0 mm <sup>2</sup> conductor size. Power cord setfittings (appliance coupler and wall plug) must bear	
Belgium	CEBC	the certification mark of the agency responsible for – evaluation in the country where it will be used.	
Denmark	DEMKO	evaluation in the country where it will be used.	
Finland	SETI	_	
France	UTE		
Germany	VDE	_	
Italy	IMQ		
Norway	NEMKO SEMKO	_	
Sweden			
Switzerland	SEV		
United Kingdom	BSI		
United States	UL	The flexible cord must be Type SJT or equivalent, No.	
Canada	CSA	14 AWG, 3-conductor. The wall plug must be a tw pole grounding type with a NEMA 5-15P (15A, 125 configuration.	
Japan	JIS	The appliance coupler, flexible cord, and wall plug must bear a "T" mark and registration number in accordance with the Japanese Dentori Law. The flexible cord must be Type VCTF, 3-conductor, 2.00 mm <sup>2</sup> conductor size.	

## **Environmental Requirements**

An important aspect of collecting high quality images is having the proper environment for the system. The following environmental requirements are outlined in this section.

#### **General Environmental Requirements**

#### Floor space:

*DeltaVision* - 3 ft × 6 ft (90 cm × 180 cm). Include 18 in (45 cm) space behind instrument rack.

*personalDV* - 22in x 54 to 62in (depending on whether or not the keyboard is kept in a slide-out tray). Include 18 in (45 cm) space behind instrument rack.

Maximum System Weight: 940 lbs (425 kg).

Service: Indoor use only.

**Temperature:** 65 - 77 °F (18 - 25 °C), daily variation of no more than 3 °F (1.8 °C). The actual room temperature should be stable to within 1 degree (Fahrenheit or Celsius) per hour. Fluctuations in temperature will affect microscope optics, which can cause the specimen to drift approximately 1  $\mu$ m per 0.1 degree Celsius.

**Humidity:** Stable humidity levels under 50%, with daily variations of less than 10%. High humidity can result in condensation on the CCD camera window that obscures image formation. Excessive humidity may also reduce filter life and may result in chromatic aberrations in the images.

Altitude: up to 6550 ft (2000 m).

Pollution: POLLUTION DEGREE 21 in accordance with IEC 664.

Ingress Protection Level: IP20

#### **Communication Recommendations**

Connect the workstation to a local area network for data storage and access to Applied Precision's web site.

**CAUTION:** Applied Precision, Inc. is not responsible for damage or harm to the workstation or imaging system due to network security breaches.

Having a telephone in the same room as the system will facilitate communication with Technical Support if needed.

<sup>&</sup>lt;sup>1</sup> IEC 61010-1: 2<sup>nd</sup> ed. International Electrical Commission defines POLLUTION DEGREE 2 as follows: "Normally only non-conductive POLLUTION occurs. Occasionally, however, a temporary conductivity caused by condensation must be expected."

#### Air Movement

Air movement around the microscope can cause specimen drift on the scale of several microns. Two common sources of air movement are window air conditioners and open windows. Central air conditioning is recommended. However, the system should not be placed in the direct path of the incoming air.

#### **Vibration Isolation**

The vibration absorbing design of the system minimizes motion artifacts from internal vibration due to shutters, filter wheels, and stage movement. The system is also designed to damp out external vibration as well. Avoiding locations near refrigerators, elevators, ventilation equipment, and other sources of vibration will improve image resolution.

#### **Ambient Illumination**

For best results, minimize ambient illumination during data collection. A lighttight room is recommended. Ensure that there are no light sources pointed downward into the lens.

A small desk lamp located near the workstation is recommended for preparing and monitoring experiments. The workstation keyboard is backlit for working in low light situations.

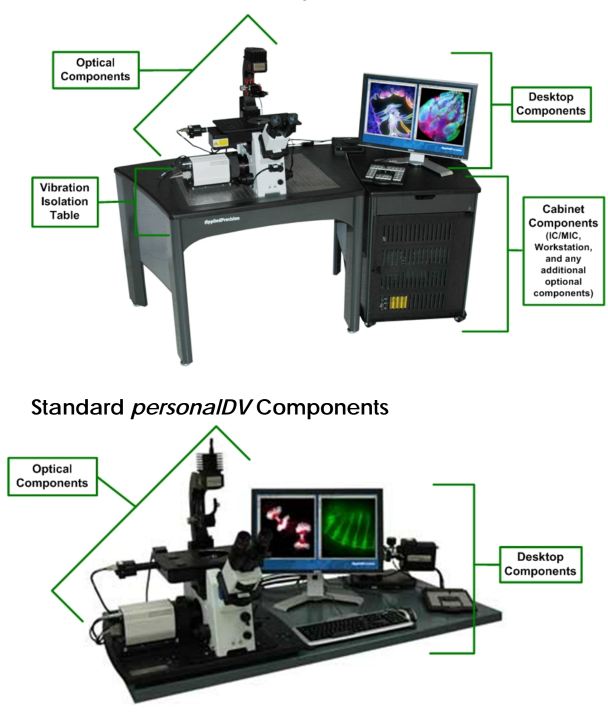
You can press the BLANK SCREEN key on the keypad to darken the monitor for improved image quality. Pressing any key on the keyboard restores monitor function.

#### Dust

It is important to minimize dust on the microscope components because dust on components can cause spots on microscope images. Minimize contamination by maintaining a clean room and covering the microscope when it is not in use. Store all extra filter sets, cameras, objectives, plates, etc. away from dust.

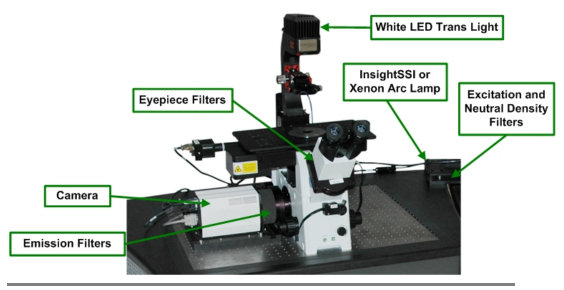
## **Overview of Components**

The standard *DeltaVision* components are shown below. Detailed descriptions of these components and descriptions of optional components are included in the following sections.



## Standard DeltaVision Components

## **Optical Components**



Note Optical components are shown for a typical installation, which includes the Olympus IX71 Microscope. Other microscope configurations vary slightly. The Olympus IX71 is currently the only microscope available.

## Fluorescence Microscope

The microscope is an advanced research grade epifluorescence inverted microscope. Each objective is qualified by Applied Precision to guarantee the highest possible quality. The point spread function (PSF) is measured in order to uphold the image quality. The microscope supports a Differential Interference Contrast (DIC) module (this is an optional component). A transmitted light is included with your system to enable automated DIC and Brightfield image acquisition.

## **Optical Filters**

The system uses a polychroic filter and filter wheels rather than a simple dichroic filter cube. The excitation, neutral density, and emission settings are selected in one of three ways:

- Selecting options in the Resolve3D window of the *softWoRx* acquisition software.
- Selecting a mode from the keypad.
- Rotating the eyepiece filter wheel (if Resolve3D is running).

Note The eyepiece filter wheel is operated manually but reports its position to the instrument controller, which in turn adjusts the excitation and emission filters automatically.

Depending upon the environment, filters last between one and three years. To ensure optimal performance, filters should be periodically inspected and replaced if necessary.

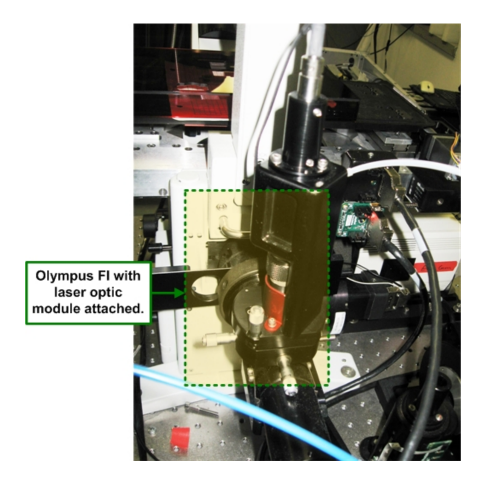
The Standard and Live Cell *DeltaVision* Filter sets are listed in *Appendix D*, *Reference Information* on Page 266.

## **Fluorescence Illuminators**

A *DeltaVision* system can be configured with either an Olympus Fluorescence Illuminator or an Applied Precision Fluorescence Illuminator. The key differences between the two fluorescence illuminators are outlined in the following sections.

#### **Olympus Fluorescence Illumination Module**

The Olympus Fluorescence Illumination Module (Olympus FI) is attached to the rear of the *DeltaVision* imaging system. The Olympus FI is a less expensive choice for fluorescence illumination if you are *not* planning to use multi-line TIRF or the UltimateFocus Module. You are still able to use broadband light sources, such as Solid State Illumination (InsightSSI) or a xenon lamp. You will still be able to use the single-line TIRF technique, but you must have an optional laser module to do so. In general, the configuration settings for the Olympus FI (field stop aperture, etc.) are set manually.



#### **API Fluorescence Illumination Module**

The Applied Precision Fluorescence Illumination (API FI) Module is attached to the rear of the *DeltaVision* system and extends to the right side. The API FI Module is actually a hardware interface that supports a combination of three smaller modules, each adding its own special utility to enhance the capabilities of the *DeltaVision* system. The Broadband Module (described below) is <u>always</u> included with the API FI Module. The TIRF/PK Module and the UltimateFocus Module are optional components. The configuration settings for the API FI Module are set up through software, which can be especially convenient if an Environmental Chamber is being used with the *DeltaVision* system. See *The Environmental Chamber* on Page 181 for details.

The three components that may be included with the API FI Module are as follows:

The Broadband Module – is an input module for broadband light sources, such as Solid State Illumination (InsightSSI) or a xenon lamp. The Broadband Module provides a user-configurable illumination function for selecting either Critical or Kohler Illumination. The module also houses the beam conditioner, a field stop aperture, and a photo-sensor interface. The

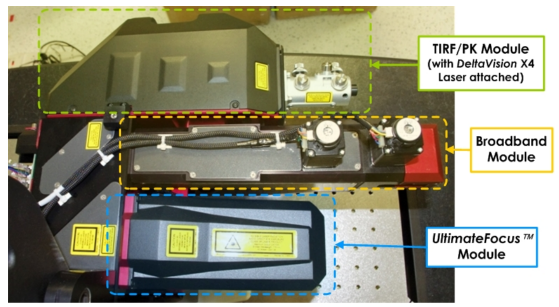
Broadband Module is standard on a *DeltaVision* system with the API FI Module installed.

The TIRF/PK Module – provides the ability to introduce a laser beam into the optical path of the *DeltaVision* microscope. With lasers attached (like the *DeltaVision* X4 Laser Module), users can generate photokinetic events and TIRF (Total Internal Reflective Fluorescence) experiments. TIRF is an imaging method that illuminates a thin section of the sample. The excitation light emits from the objective at a shallow angle, enters into the cover slip, and bounces inside the cover slip creating an *evanescent* field of fluorescence. The TIRF/PK Module is available on a *DeltaVision* system with the API FI Module installed. For more information on the X4 Laser Module and the TIRF technique, see X4 Laser Module and TIRF on Page 191.

**Note** Your TIRF/PK Module may include TIRF only, PK only, or both TIRF and PK.

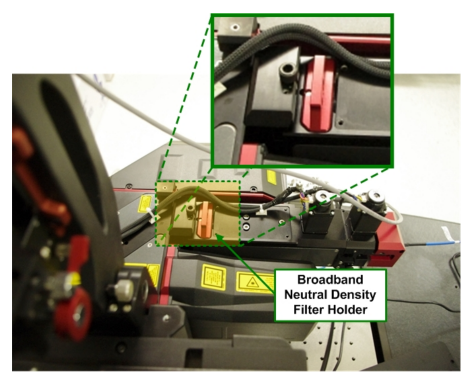
➤ The UltimateFocus<sup>™</sup> Module – maintains focus during an experiment, after the operator has found focus and marked a point of interest. The UltimateFocus Module uses an infrared laser that follows the illumination path and bounces off the cover slip/sample interface. The reflected beam is evaluated and the software returns an offset to the Z motor for automatic stage adjustment to maintain focus. The UltimateFocus Module is available on a DeltaVision system with the API FI Module installed.

Each of these components is mounted to the rear of the *DeltaVision* imaging system (while facing the microscope) as shown below.



DeltaVision Fluorescence Illumination Module

Note Particularly when using the InsightSSI Solid State Illumination Module, you may find that you need to decrease the imaging system's illumination output by placing an additional neutral density (ND) filter directly into the Broadband Module. Pull up on the red ND Filter Holder (shown below) and insert an ND filter (p/n 52-852283-000-9).



**Broadband Neutral Density Filter Holder** 

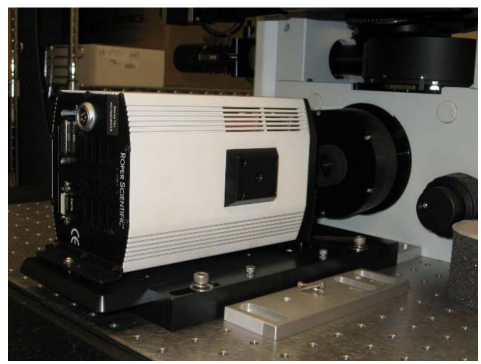
#### Cameras

*DeltaVision* comes standard with the CoolSNAP *HQ*<sup>2</sup> CCD (charge coupled device) camera, referred to as the High-speed Camera, and also supports an upgrade option of the Evolve EMCCD camera. For more about the Evolve camera, see *Evolve Camera* on Page 167.

personalDV comes standard with the CoolSNAP ES<sup>2</sup> Camera.

#### **High-speed** Camera

The High-speed CCD camera (CoolSNAP  $HQ^2$ ) collects digital image data from the microscope. This camera is designed to collect image data at a high frame rate. At the fastest speed, the camera can collect 30 frames per second of a 64 × 64 pixel image. Increased acquisition speed is useful when collecting images of live cells that deteriorate over short periods of time.



High-speed CoolSNAP HQ<sup>2</sup> CCD Camera

The High-speed Camera is air-cooled. The cooling apparatus is incorporated into the camera assembly; therefore, when the High-speed Camera is turned on, the cooler is also operating. When using cooled cameras in humid conditions, it is possible for condensation to form on the CCD camera window. If this happens, a mottled pattern is superimposed on the images. You will need to lower the ambient humidity level to avoid condensation.

Note If no images are acquired over a 4-hour period, the High-speed Camera cooling automatically shuts down.

**CAUTION:** Do not disconnect cables from the High-speed Camera when the power is on. Be sure to leave a 1 in (2.5 cm) minimum space around the cooling fan. Use only the PCI cards, cable, and power supplies that are designated for this system.

#### High-speed Camera Power Supply

The High-speed Camera power supply provides electric power to the camera. It is housed on the bottom shelf of the cabinet.

#### ES<sup>2</sup> Camera

The CoolSNAP  $ES^2$  CCD Camera is basically the same as the CoolSNAP  $HQ^2$  Camera, except the  $ES^2$  is not as deeply cooled. This makes it more affordable,



but allows for slightly higher noise. However, the noise difference is virtually unnoticeable unless you are using long exposure times (~8s).

CoolSNAP ES<sup>2</sup> CCD Camera

#### **Evolve Camera**

The Evolve camera is an EMCCD-type camera with a 512 x 512 imaging array and 16 x 16  $\mu m$  pixels.



Photometrics<sup>®</sup> Evolve Camera

The Evolve Camera was designed by Photometrics to perform very well in lowlight applications and is therefore an excellent camera for live-cell imaging. This camera also reads out pixel data in electrons, resulting in more accurate and reproducible data. Some of the key features of the Evolve Camera are:

- EM gain
- Back-illuminated EMCCD
- Optimized field of view and very high sensitivity
- PAR feedback system
- ACE (Advanced Clocking Enhancement) technology
- 10-MHz readout
- 5- and 1- readout
- Dual amplifiers
- 16-bit digitization

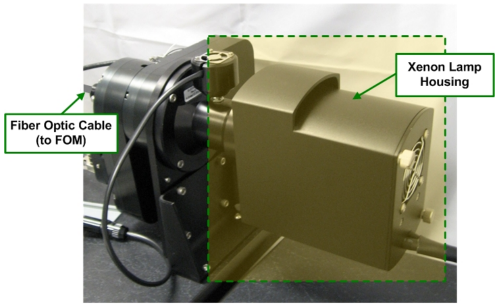
## **Light Sources**

A *DeltaVision* imaging system provides at least two, and sometimes three light sources: either a solid state illumination (InsightSSI) module or a xenon arc lamp for the broadband light source, a white LED for transmitted light, and an optional laser module for laser-based light.

Note A *DeltaVision* system can optionally be configured with both broadband light sources (InsightSSI and a xenon arc lamp). Note that only one broadband light source can be used at any given time.

#### **Xenon Lamp**

One of the choices offered for a *DeltaVision* broadband light source is the xenon arc lamp. Illumination for the microscope is delivered from a xenon lamp to the specimen through a fiber optic cable.



#### **Xenon Lamp Housing**

For instructions on how to replace the xenon bulb, see *Replacing the Xenon Bulb* on Page 240.



**WARNING:** Do not disconnect the xenon arc lamp power cable when the power is on.



**WARNING:** The xenon arc lamp presents potentially harmful risks to the user, including the possibility of UV exposure to skin and eyes. Before operating the microscope, consult *Chapter 2: Safety* for important information regarding arc lamp operation.

Note The illumination path alignment is critical to acquiring the highest possible image resolution. If you have reason to suspect that the illumination path alignment is inaccurate for your *DeltaVision* system, contact the Applied Precision Customer Hotline.

#### Photo sensor

The light passes through the fiber optic module (FOM), where approximately 1% of the light is diverted to the Photo sensor. The remaining 99% of the light is delivered to the microscope. The Photo sensor measures illumination intensity by sampling a small percentage of the light from the broadband light source during an exposure. The photo sensor signal is recorded by the instrument controller and then used to correct for variations in the xenon lamp intensity during an experiment.

The Corrections tool in *softWoRx* normalizes each image based on its photo sensor value. These corrections are automatically applied during deconvolution. This enables quantifiable intensity comparisons between images, even if the brightness of the excitation light varies between the images.

#### InsightSSI™ (Solid State Illumination)

The InsightSSI Module, another choice offered for a *DeltaVision* broadband light source, adds a solid state light source to your imaging system. Because the light produced by the InsightSSI Module is solid state and can be switched on and off very quickly, no shutters are necessary for imaging. The InsightSSI Module increases power across all wavelengths. This feature can greatly reduce exposure times. Not having to wait for a xenon bulb to stabilize is another advantage of using a solid state light source. The InsightSSI Module turns on and off instantly.



Four-color InsightSSI Module

Depending on the type of imaging you plan on doing, the InsightSSI Module is available with a choice of three different excitation filter sets:

- Four-color InsightSSI Module for fixed cell applications
- Four-color InsightSSI Module for live cell applications
- Seven-color InsightSSI Module for both fixed and live cell applications



#### Seven-color InsightSSI Module

The three available filter sets are detailed in the following tables:

#### Four-color Standard Filter Set

Wavelengths (nm)	Suggested Fluorophores	
• 381-399	DAPI, Hoechst, Alexa Fluor <sup>®</sup> 350, Dylight™ 350, CF <sup>®</sup> 350, Cascade Blue, Lucifer Yellow, AMCA, CellTracker <sup>®</sup> Blue, all Qdot <sup>®</sup> nanocrystals	
• 461-489	FITC, Alexa Fluor <sup>®</sup> 488, Dylight™ 488, CF <sup>®</sup> 488, Oregon Green 488, Calcium Green, Flou-3, Flou-4, CellTracker <sup>®</sup> Green, Rhodamine 123, MitoTracker <sup>®</sup> Green, LysoTracker <sup>®</sup> Green	
• 529-556	TRITC, Cy3 <sup>®</sup> , Alexa Fluor <sup>®</sup> 546, Alexa Fluor <sup>®</sup> 555, Alexa Fluor <sup>®</sup> 568, Dylight™ 549, CF <sup>®</sup> 555, MitoTracker <sup>®</sup> Orange	
621-643	Cy5 <sup>®</sup> , Cy5.5, APC, Alexa Fluor <sup>®</sup> 647, Alexa Fluor <sup>®</sup> 635, Alexa Fluor <sup>®</sup> 660, Dylight™ 633, Dylight™ 649, CF <sup>®</sup> 633, CF <sup>®</sup> 647, MitoTracker <sup>®</sup> Deep Red	

#### Four-color Fluorescent Protein Filter Set

Wavelengths (nm)	Fluorescent Proteins	Suggested Fluorophores	
• 426-450	CFP	Pacific Blue™	
• 461-489	GFP, EGFP	FITC, Alexa Fluor <sup>®</sup> 488, Dylight™ 488, CF <sup>®</sup> 488, Oregon Green 488, Calcium Green, Flou-3, Flou-4, CellTracker <sup>®</sup> Green, Rhodamine 123, MitoTracker <sup>®</sup> Green, LysoTracker <sup>®</sup> Green	

<b>•</b> 505-515	YFP	Alexa Fluor <sup>®</sup> 532
<b>563-588</b>	mCherry	Alexa Fluor <sup>®</sup> 568, Alexa Fluor <sup>®</sup> 594, Dylight™ 594, MitoTracker <sup>®</sup> Orange, MitoTracker <sup>®</sup> Red, LysoTracker <sup>®</sup> Red, CellTracker <sup>®</sup> Red

#### Seven-color Combined Filter Set

Wavelengths (nm)	Fluorescent Proteins	Suggested Fluorophores
• 381-399	BFP	DAPI, Hoechst, Alexa Fluor <sup>®</sup> 350, Dylight™ 350, CF <sup>®</sup> 350, Cascade Blue, Lucifer Yellow, AMCA, CellTracker <sup>®</sup> Blue, all Qdot <sup>®</sup> nanocrystals
• 426-450	CFP	Pacific Blue™
• 461-489	GFP, EGFP	FITC, Alexa Fluor <sup>®</sup> 488, Dylight™ 488, CF <sup>®</sup> 488, Oregon Green 488, Calcium Green, Flou-3, Flou-4, CellTracker <sup>®</sup> Green, Rhodamine 123, MitoTracker <sup>®</sup> Green, LysoTracker <sup>®</sup> Green
<b>•</b> 505-515	YFP	Alexa Fluor <sup>®</sup> 532
• 529-556	OFP, RFP, DsRed	TRITC, Cy3 <sup>®</sup> , Alexa Fluor <sup>®</sup> 546, Alexa Fluor <sup>®</sup> 555, Alexa Fluor <sup>®</sup> 568, Dylight™ 549, CF <sup>®</sup> 555, MitoTracker <sup>®</sup> Orange
<b>-</b> 563-588	mCherry	Alexa Fluor <sup>®</sup> 568, Alexa Fluor <sup>®</sup> 594, Dylight™ 594, MitoTracker <sup>®</sup> Orange, MitoTracker <sup>®</sup> Red, LysoTracker <sup>®</sup> Red, CellTracker <sup>®</sup> Red
621-643		Cy5 <sup>®</sup> , Cy5.5, APC, Alexa Fluor <sup>®</sup> 647, Alexa Fluor <sup>®</sup> 635, Alexa Fluor <sup>®</sup> 660, Dylight™ 633, Dylight™ 649, CF <sup>®</sup> 633, CF <sup>®</sup> 647, MitoTracker <sup>®</sup> Deep Red

A critical advantage of using the InsightSSI Module for your *DeltaVision* system's broadband light source instead of the xenon lamp is speed. When mechanical shutters are eliminated from the system, shorter exposure times can be achieved.

Note Additional high-speed polychroic options are available for the four-color fluorescent protein InsightSSI and the seven-color InsightSSI.

#### **LED Transmitted Light**

An LED transmitted light source is also provided.



LED Transmitted Light Source

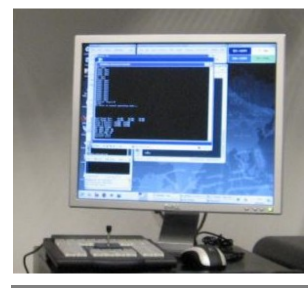
Note Most of the *Delta Vision* light sources (the xenon lamp, solid state illumination, or the LED transmitted light) can be used with the eyepieces or the cameras. Lasers cannot be used with the eyepieces.

## **Desktop Components**

Desktop Components include the flat panel display monitor, the keyboard, the mouse, the keypad/joystick. *DeltaVision* includes a vibration isolation table. *personalDV* includes a bench-top microscopy isolation platform.

## Flat-Panel Display Monitor

All *DeltaVision* systems are equipped with Flat-Panel LCD monitors. These monitors offer a very high level of performance in several areas pertaining to the quality of displayed images, the most critical of which is contrast ratio.



Note For instructions that show how to adjust the monitor, see the Flat-Panel Display manual.

**CAUTION:** *DeltaVision* is configured to work with the monitor that is included with the system. Other monitors are not necessarily supported.

## The Keypad and Joystick

Many of the functions accessible through Resolve3D are also available on the keypad/joystick (see *Keypad/Joystick Operation* on Page 322).



## Vibration Isolation Table

Within the vibration isolation table is a breadboard surface that is supported by mechanical vibration isolators. These components provide optimal performance

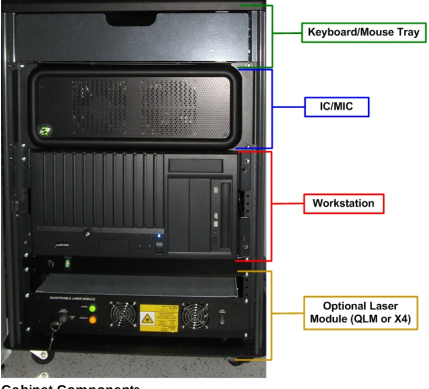
without an external air source. The isolators are sized for the system weight as delivered. If significant additional weight is added to the breadboard, higher capacity isolation may be required. Contact Applied Precision for more information.

Note The vibration isolation table is not included with *personalDV*. Instead, *personalDV* includes a bench-top isolation platform to provide similar stability for the microscope. The *DeltaVision* vibration isolation table is available for personalDV as an optional upgrade.

## **Cabinet Components**

Note The component cabinet is included with *DeltaVision* only. The cabinet is not available for *personalDV*.

The cabinet contains all of the electronic control equipment and provides surfaces for the keyboard and the Flat Panel Display. The front two wheels should remain locked and the cabinet left in place. There are many cables connecting the microscope that can be damaged if they are pulled. The standard cabinet components are shown below.



#### Cabinet Components

Solution of DeltaVision cabinet components may vary slightly.

## Instrument Controller / Microscope Interface Chassis (IC/MIC)

The Instrument Controller (IC) is the portion of the computer that interfaces with all of the microscope hardware (including the microscope stage motors, filter wheel motors, and cameras). It coordinates all activities related to positioning the stage and collecting images. Data from the camera feeds through the controller to the workstation. The controller also receives instructions from the workstation and issues commands to the motors through the Microscope Interface Chassis.

The Microscope Interface Chassis (MIC) side of the computer provides power and control for the filter motors, stage drive, and shutters. It also contains the Photo sensor, which is connected to the microscope through a fiber optic cable.

#### Notes

**#1** The Instrument Controller / Microscope Interface Chassis has no userserviceable parts inside.

**#2** Occasionally, you may need to replace fuses on the back panel of the chassis. Consult Chapter 9 for a description of this procedure. Unless there is an obvious reason why the fuse blew, you should contact Applied Precision Customer Service when you need to replace fuses.

#### Workstation

The workstation hosts the *softWoRx* application that is the primary interface used to control the system. The workstation can also provide a server for data storage when used with the optional Data Management Software (DMS).

Note The workstation has no user-serviceable parts inside.

## Other Standard Components

Other standard components include the Repeatable Slide Holder, the Slide Holder Adapter, the Fiber Optic Module, and the keypad/joystick.

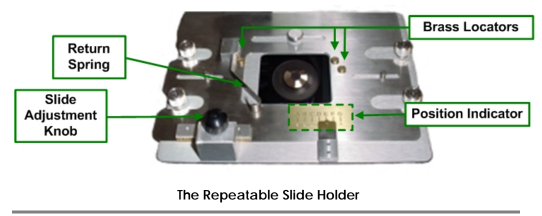
#### The Repeatable Slide Holder

The Repeatable Slide Holder holds the slide on the stage. It also allows you to move the slide across the stage and to mark the slide position when you remove the slide.

The ability to move the slide across the stage allows you to view the full slide on the  $1" \times 1"$  stage area. With the slide held against the three brass locators by the

Return Spring, you can use the Slide Adjustment Knob to move the slide laterally.

Marking the position of your slide is useful when you are conducting a point visiting experiment and you need to remove the slide before you are finished. You can use the Position Indicator (the letter scale at the bottom of the slide holder) to record the position of your slide. When you resume your experiment, you can place the slide in the position that you recorded.



Note If you are using a Petri dish or any other format that is not similar to a 1" x 3" slide, you will need to remove the Repeatable Slide Holder or use the Slide Holder Adapter.

For more about using the Repeatable Slide Holder, see *Finding a Specimen and Recording its Slide* Holder Position on Page 101.

## **Slide Holder Adapter**

The Slide Holder Adapter holds a chambered coverglass. It is mounted on the Repeatable Slide Holder. NUNC Lab-Tek<sup>™</sup> II chambered coverglasses with one, two, four, or eight wells are available. The Slide Holder Adapter also supports Petri dishes from 25 – 40 mm in diameter.



Slide Holder Adapter

## **Calibration Kit**

The Calibration Kit is provided for calibration operations on the DeltaVision system. The kit includes the following six slides:

- Three plastic fluorescent calibration slides for flat-field calibration:
  - Blue plastic (EX 408nm, EM 440 nm) Good for DAPI, Hoechst, etc.
  - Orange plastic ((EX 488nm, EM 519nm) Good for FITC, GFP, TRITC, CY-3<sup>®</sup>, etc.
  - Red plastic (EX 590nm, EM 650nm) Good for CY-5<sup>®</sup>, etc.
- Silicon mirror slide for light path alignment and troubleshooting.
- Silicon grid slide for pixel size measurement.
- 100nm Rhodamine<sup>®</sup> bead slide for PSF measurement.

## The Fiber Optic Module

Use the Fiber Optic Module to align the light path from the fiber optic cable to the Fluorescent Illuminator. This module allows you to adjust the tilt, horizontal, and vertical orientation of the light path.



Fiber Optic Module

## The Tool Kit

A tool kit is provided with all *DeltaVision* systems that includes tools for maintaining the system. This kit includes:

- Ties and belts for suspending cables, screws, and other fasteners
- Cleaning material (Q-tips and lens paper)
- A hex wrench set (5/64" 3/16")
- A set of metric L keys (1mm –5mm)
- A #1 Phillips screwdriver
- A 5mm T Handle hex key
- An acrylic bulls eye level
- A micrometer nut wrench
- An Applied Precision Hotline sticker

## Software

All *DeltaVision* systems include *softWoRx* for Linux (1 copy).

Note DeltaVision systems can be ordered with a number of different software configurations. Talk with your Applied Precision representative to determine the best configuration for your applications.

#### softWoRx

*softWoRx* is the Linux software application that runs the acquisition workstation. The software allows you to perform the following tasks:

- Acquire image data
- Set up and run experiments
- Deconvolve data
- Measure point spread functions
- Calculate optical transform functions
- Process 2-D and 3-D images
- Perform quantitative analysis

- Archive data
- Configure task chains
- Manage user accounts

## **Optional Components**

You can purchase several optional *DeltaVision* components. Some options are available for all *DeltaVision* systems, but many are available only for specific *DeltaVision* models. Check with your Applied Precision representative for details.

- The Environmental Chamber provides a controlled temperature environment for live cell imaging. The chamber also supports CO<sub>2</sub> injection, which allows you to control humidity and pH by maintaining a CO<sub>2</sub> flow over the sample. For additional information about the Environmental Chamber, see Page 181.
- Microtiter Stage for *DeltaVision* provides the system with the ability to scan microtiter plates. For more details on the Microtiter Stage, see Page 120.
- Additional Filter Modules can be loaded with custom filters or with the Live Cell filter set to provide greater flexibility for your lab. For more information about these Filter Modules, see Page 190.
- The X4 Laser Module adds multi-line TIRF (Total Internal Reflection Fluorescence) capabilities to your *DeltaVision* system. This is an optical technique in which laser light is introduced to the microscope in a way that limits fluorescence imaging to a thin area at the surface of the substrate. Typically, the TIRF sample is only illuminated 100-200nm into the specimen, resulting in an enhanced signal-to-noise ratio and increased imaging contrast. With the module's lasers, you can also perform Fluorescence Recovery After Photo-bleaching (FRAP) analysis and other photokinetic experiments. For more information on the X4 Laser Module and the TIRF technique, see X4 *Laser Module and TIRF* on Page 191. For details on FRAP analysis, see "Analyzing <u>Fluorescence Recovery After Photo-bleaching</u>" in the <u>softWoRx Imaging</u> Workstation User's Manual.
- The InsightSSI (Solid State Illumination) Module is a solid state light source available for *DeltaVision* systems. The InsightSSI Module is available with either a set of four filters for fixed cell imaging <u>or</u> a set of four filters for live cell imaging, or seven filters for both fixed <u>and</u> live cell imaging. For more information about the SSI Module, see *InsightSSI™(Solid State Illumination)* on Page 170.

- The Evolve EMCCD-type camera is an optional camera that provides high signal-to-noise ratios for low-light fluorescence, TIRF, or single molecule fluorescence. For more information on the Evolve Camera, see Page 167.
- The Multiplexed Wavelength module for *DeltaVision* includes a second shuttered xenon light source and provides nearly simultaneous two-channel imaging. For more information about the Multiplexed Wavelength option, see Page 200.
- The UltimateFocus<sup>TM</sup> Module is a tool for maintaining focus during an experiment. After the operator has found the proper focus and marked a point of interest, the UltimateFocus Module maintains focus during the experiment. The UltimateFocus Module uses an infrared laser that follows the illumination path and bounces off the cover slip/sample interface. The reflected beam is evaluated and the software returns an offset to the Z motor for automatic stage adjustment to maintain focus. The UltimateFocus Module is optional for most *DeltaVision* systems. For more information on the UltimateFocus Module, see *Using UltimateFocus* on Page 143.
- Analysis Workstations include the *softWoRx* Linux workstation and the *softWoRx* Suite Windows workstation. For further information on these workstations, see Page 199.
- Available software includes two optional modules for *DeltaVision* systems: the *softWoRx* Explorer option and the *softWoRx* Suite advanced option. More information on software options is provided on Page 201.
- A selection of Differential Interference Contrast (DIC) components are supported by *DeltaVision* systems. For alignment and setup details, see *Differential Interference Contrast Module* on Page 202.
- The following optional objectives are tested by Applied Precision to ensure that they meet our rigid quality standards.

Optional Objectives
U-APO 40X Oil, 0.65-1.35NA, 0.10mm WD
U-PLAN S-APO 100X OII, 1.4NA, 0.12 WD
U-PLAN APO 60X W/PSF Water, 1.20 NA, 0.25mm WD

The following subsections describe each of the optional DeltaVision components.

# The Environmental Chamber

Environmental control is an essential factor in live-cell microscopy or any other type of experiment requiring system stability. Focus instability can be difficult to

resolve for critical experiments. Small changes in ambient temperature can lead to thermal expansion or contraction in the microscope stand, the stage, or the objective. These subtle changes can completely shift the optimal plane of focus. The Environmental Chamber is designed to precisely control temperature and CO<sub>2</sub> levels making it ideal for long term time-lapse applications and high-resolution imaging.



**Environmental Chamber - Opaque** 



**Environmental Chamber - Transparent** 

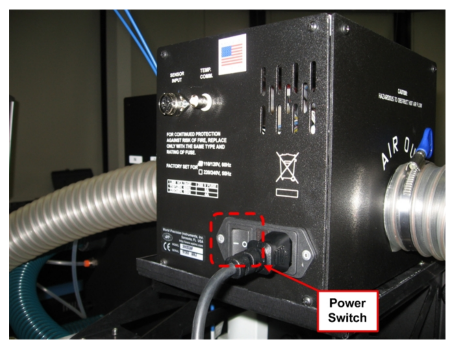
The Environmental Chamber includes a temperature controller, a CO<sub>2</sub> humidifier, and a CO<sub>2</sub> chamber. This Applied Precision-designed product is generally installed by a trained Applied Precision service representative.

## **Basic Environmental Chamber Components**

Use the functional descriptions in this section for a basic understanding of the utility provided by the Environmental Chamber option installed on a *DeltaVision* system.

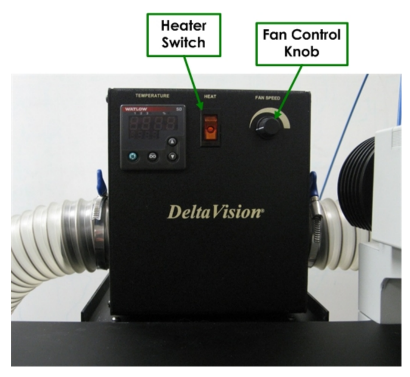
## **Temperature Controller**

The power switch for the temperature controller is on the back panel next to the power cord connection (see below). This switch controls the main power for the fan, the heater, and the microprocessor controller.



**Temperature Controller Power Switch** 

The Heater Switch on the front panel of the temperature controller controls <u>only</u> <u>the heater</u>. When the Heater Switch is off and the Power Switch is on, the fan and microprocessor controller are still powered.



#### Heater Switch and Fan Control Knob

The temperature controller is preset for use with *DeltaVision* Environmental Chamber by default. The out-of-the-box settings for the PID controller should <u>not</u> be changed. All testing and calibration have been performed using the preset configuration.

**CAUTION:** Changing the settings on the PID controller from the default configuration may cause problems when using the device for the Environmental Chamber on a *DeltaVision* system.

## Fan Control Knob

The Fan Control Knob on the front panel adjusts the airflow rate from 20 to 50 cubic feet per minute. A lower flow rate can provide quieter operation and may be suitable for some applications. In order to keep the temperature more even throughout the Environmental Chamber, however, a higher flow rate setting is generally recommended. A higher fan speed setting will generate more air flow within the environmental chamber so that the temperature in the chamber remains more uniform.

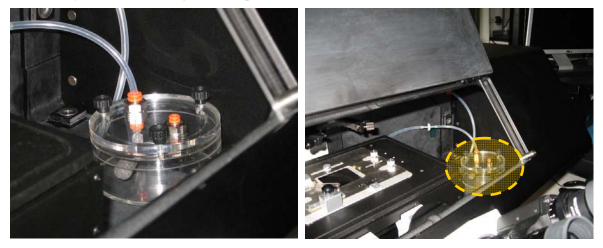
Important It is recommended that, for normal use, the Fan Control Knob is set to the middle of the full range of the knob's rotation and that you wait approximately 3 hours for the Environmental Chamber to stabilize before beginning any imaging tasks.



**WARNING:** Avoid blocking the airflow. Although a built-in thermal cutoff switch in the temperature controller is designed to reduce the hazard, a fan failure or obstruction in the airflow has the potential for overheating the system and damaging the *DeltaVision* system or even causing a fire.

## CO<sub>2</sub> Humidifier

When filled with pre-warmed water, the CO<sub>2</sub> Humidifier contains enough water to last for seven days of continuous use. The CO<sub>2</sub> Humidifier can be cleaned with any of the following: lab soap (such as Alconox), 70% Ethanol, or 10% bleach.



#### Flowmeter

The Flowmeter for the Environmental Chamber controls the flow of air in and out of the chamber. Users will need to provide premixed air cylinders for the Environmental Chamber. The Flowmeter provides a flow range from 1.3 ml/min to 5.8 ml/min. The Flowmeter should be set so that bubbles are slowly emerging from the stone, similar to the bubbles in carbonated soda.



### Flowmeter

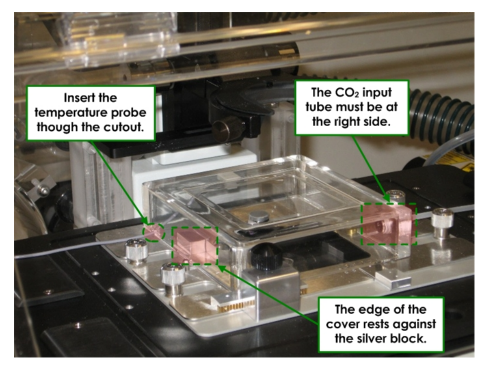


**CAUTION:** The maximum inlet pressure for the Flowmeter is 200 psi, however, the fittings have a maximum pressure of only 145 psi.

## CO<sub>2</sub> Chamber

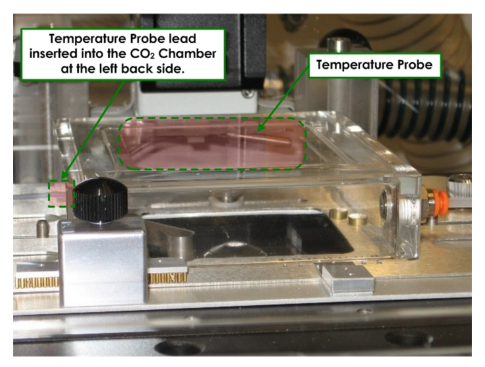
The CO<sub>2</sub> Chamber fits directly over the sample area and provides CO<sub>2</sub> to the sample during imaging. Place the chamber over the sample before imaging.

The CO<sub>2</sub> Chamber is not attached in any way to the *DeltaVision* stage however, <u>proper orientation of the CO<sub>2</sub> Chamber is essential</u> for successful imaging. The CO<sub>2</sub> input tube should be on the right side (as shown below) and the chamber should be flush with the silver block on the left side.



#### CO<sub>2</sub> Chamber - Positioning

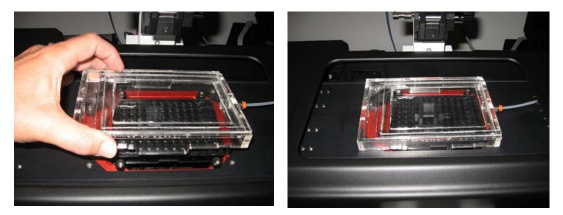
When you use the CO<sub>2</sub> Chamber, the temperature probe must be relocated from its stand near the transmitted light tower and inserted through the cutout on the left side of the chamber as shown.



CO<sub>2</sub> Chamber - Temperature Probe

Note Using the CO<sub>2</sub> chamber limits the use of the Repeatable Slide Holder. With the CO<sub>2</sub> chamber installed, some of the detent positions are not available.

The CO<sub>2</sub> chamber for the Microtiter Stage (see next section) is larger to accommodate a microtiter plate, but otherwise is set up with the same method used for the standard CO<sub>2</sub> chamber.



Microtiter Stage CO<sub>2</sub> Chamber

## Air Filter

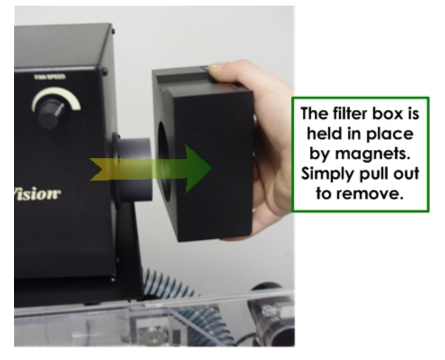
The Environmental Control System is equipped with an intake air filter. This filter is removable and cleanable. You are advised to clean this filter every six months.



#### **Heater Filter Housing**

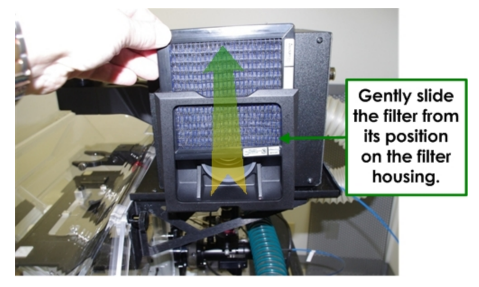
To remove and clean the air filter:

- 1. Turn off the heater.
- 2. The filter can be removed from the filter box without removing the filter box from the heater. If desired, remove the filter box by pulling it away from the heater. Magnets hold the filter box in place.



Removing the Filter Housing

3. Slide the filter out of the filter box as shown below.



#### **Removing the Filter**

- 4. Using a vacuum cleaner, thoroughly clean the filter mesh.
- 5. Reinstall the filter and box in reverse order of removal.
- 6. Continue with normal operation of the system.

# The Microtiter Stage Option

The addition of a Microtiter stage to a *DeltaVision* system enhances the imaging system's capability by providing the ability to scan microtiter plates. With a Microtiter stage on a *DeltaVision* system, you can design a plate scan by generating a point list that corresponds to the nominal center point of each well of interest, and then defining how the wells are to be sampled.

The model used for sampling plates is a matrix of N rows by M columns of *panels*. Each panel can be a Z stack, a single Z, or an *OAI* (optical axis integration) scan. The panels can be separated by spacing defined through the software. You can specify autofocus for each individual well, or for every panel/stack within the matrix. You can also set up parameters to specify the autofocus sampling range that represents the well-to-well and panel-to-panel variability of the focal plane.



Microtiter Stage for DeltaVision Systems

For instructions on basic operation of the Microtiter Stage option, refer to "Using the Microtiter Stage Option" on Page 120.

# **Additional Filter Modules**

You can purchase additional *DeltaVision* filter modules and use them to quickly change the filter sets to meet different imaging requirements.

You can purchase a Live Cell Filter Module that is preloaded with four filter sets that are commonly used for live cell imaging. You can also purchase empty filter modules and insert customized filter sets into those modules.

# **TIRF/PK Module**

The TIRF/PK Module provides the ability to add a laser beam into the optical path of the *DeltaVision* microscope. A laser beam is introduced into the back aperture of the microscope objective to provide a focused illumination spot in the center of the optical field. With lasers attached (like the *DeltaVision* X4 Laser Module), users can design photokinetic experiments and TIRF (Total Internal Reflective Fluorescence) experiments. TIRF is an imaging method that offsets the laser excitation light slightly off-axis using a special TIRF-based objective. The excitation light emits from the objective at a shallow angle, enters into the cover slip, and bounces inside the cover slip creating an evanescent field of fluorescence. The TIRF/PK Module is optional for the *DeltaVision* system and is offered in three varieties: TIRF/PK, TIRF only, and PK only.

# X4 Laser Module

The X4 Laser Module houses up to four lasers. The lasers are enclosed in a rackmounted module that fits in the cabinet. All lasers in the X4 Laser Module are active at the same time and controlled with electronic shuttering. Light is directed from the lasers to the light path through the Laser Optic module that mounts on the back of the Fiber Optic module.

If your system has the X4 Laser module, you can use *softWoRx* to run Photokinetic laser-based experiments and/or TIRF (Total Internal Reflection Fluorescence) experiments.

## Laser Safety using the X4 Laser Module

The laser interlock is attached to the port selector knob on the front of the microscope. When the port selector is set to the eyepieces, the laser shutters are prohibited from opening. Only when the port selector knob is set to the Camera port will the laser shutters open.

For complete information on laser safety and proper labeling for the X4 Module, refer to *Appendix F*, *Lasers and Safety Issues*.



The port selector knob and the UltimateFocus and X4 Laser Module indicator lights are shown on the front of the *DeltaVision* microscope.

For more information on the X4 Laser Module and the TIRF technique, see X4 *Laser Module and TIRF*.

# Using the X4 Laser Module

The X4 Laser Module can contain up to four lasers. All of these lasers can be active at the same time to allow for extremely fast sequential activity. When using any or all of these lasers, your first task should be to check their status.

You can view and adjust the status of all available lasers from the Resolve3D Settings window, in the **Lasers** tab. The lasers are displayed with one of four levels of status:

- Off The laser is not in use and is in rest mode. The laser may enter rest mode if it has not been used for a specified period of time. In this mode, laser lifetime is preserved.
- **Stabilizing** The laser is coming up to full power. The length of time for each laser to stabilize at full power will vary between 5 and 25 seconds.
- **Ready** The laser is now available for use.

 Locked – The laser interlocks are open, prohibiting all laser function (for example, the eyepiece port is selected).

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	o test 488	☐ Show spot a	lignment target		
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Laser t Image Do Te	o test 488  The Spot Con	tinuous (Sp	litter: 50% PK <-	-> 50% TIRF)	

#### Resolve3D Settings Window | Lasers tab showing status for all four lasers

#### Activating a Laser:

• Click the checkbox next to the specific laser to enable/disable emission.

#### Setting Laser Power:

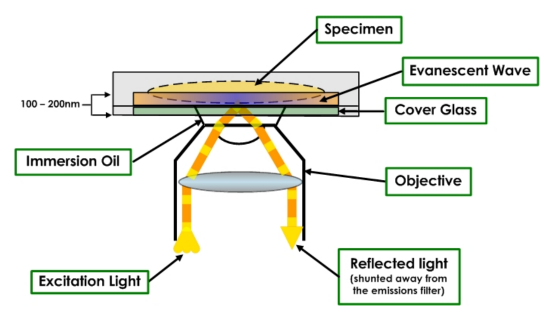
Laser power is displayed as a percentage of the full power of each laser.
 Adjust the laser's power by entering a numerical value.

### X4 Laser Module and TIRF

With the X4 Laser module, you can use *softWoRx* to run Photokinetic laser-based experiments or TIRF experiments. The module houses up to four lasers and all lasers can be active at the same time and controlled by electronic shuttering.

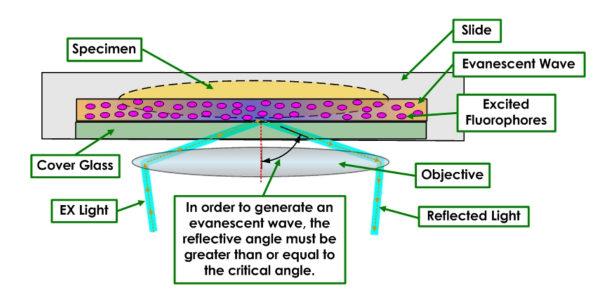
TIRF is an optical sectioning technique that limits fluorescence imaging to a thin area at the surface of a specimen, typically restricted to a depth of only 100 – 200nm, resulting in an enhanced signal-to-noise ratio and increased imaging contrast.

This technique uses a simple law of physics to improve biological imaging. When light passes from a medium of high refractive index to a medium of low refractive index and the angle of incidence is greater than or equal to the critical angle, the light will reflect off of the interface and not actually enter the second medium. Under these conditions, an electromagnetic wave traveling perpendicular to the interface is created. This electromagnetic wave has the same wavelength as the light that created it and decays very rapidly in the direction of the optical axis such that most of the energy is lost within a couple of hundred nanometers of the interface. This electromagnetic wave is known as an *evanescent* wave.



If there are fluorescent molecules in close proximity to the interface, and these molecules are capable of absorbing the wavelength of the evanescent wave, then these molecules (known as *fluorophores*) will become excited and fluoresce. Since the evanescent wave decays rapidly, molecules that are more than about 200nm from the surface of the interface will not be sufficiently excited to fluoresce. These molecules contribute to neither the signal nor the background (noise) fluorescence.

For TIRF slides, the sample must be grown on a glass cover slip and the sample is bathed in a water-based buffer with a refractive index of approximately 1.33. The glass cover slip typically has a refractive index of about 1.518. When light is introduced from the cover slip to the buffer at an angle that is greater than about 61°, the light reflects off the glass-water interface and establishes an evanescent wave that will travel into the buffer. Fluorophores close enough to the interface and that can be excited by the evanescent wave will generate fluorescence that is then detected through the objective lens.



## **TIRF-specific Laser Safety Considerations**

Due to the TIRF illumination optics, the light being emitted from the *DeltaVision* objective is collimated and has high power density. The TIRF system also has the ability to direct this light to sharp off-axis angles relative to the objective axis.

# **Imaging Using TIRF**

## To generate a TIRF image:

- 3. Locate the sample through the eyepieces using widefield fluorescence.
- 4. From the Resolve3D Settings window under the **Lasers** tab, click the **Multiline TIRF Settings** button near the bottom of the window.

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Module Status	On	Connect			
Laser	₹ 405	<b>₹</b> 488	<b>□</b> 514	<b>F</b> 561	
Status	Ready	Ready	Off	Stabilizing	
Power (%)	50	<u></u> [50	Įso	Iso	
	The Spot Con				
Do Te	% st Bleach	Bleach Bac	kground		Click the <u>Multi-line</u> <u>TIRF Settings</u>
TIRF Multi-line TIR	F Settings				button.
Done Save S	Settings			Help	

<ul> <li>Multi-line TIRF Settings</li> <li>■ X</li> <li>TIRF Depth</li> <li>↓ 405 [1090.0]</li> </ul>	Activate the radio button next to the desired laser.
514       990.0         561       990.0         Adjust channels independently         Block       Epifluorescence         Restore TIRF Depth	Activate the <u>Adjust</u> <u>Channels Independently</u> checkbox to make separate angle adjustments to individual wavelengths.
Back Aperture Focus For Current Lens	Deactivate the Locked checkbox to unlock the Back Aperture Focus.

The Multi-line TIRF Settings window is displayed.

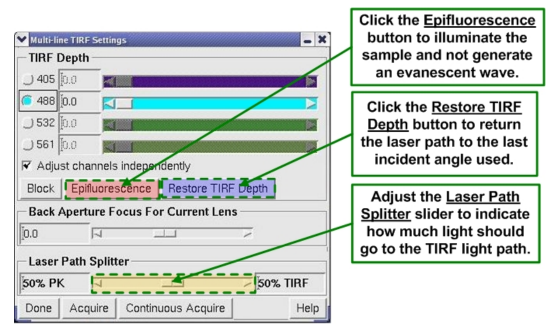
- 5. Activate the radio button next to the laser you want to adjust.
- 6. (*Advanced only not required*): Activate the **Adjust Channels Independently** checkbox to make unique angle adjustments for each wavelength (channel).
- 7. (*Advanced only not required*): In the **Back Aperture Focus For Current Lens** field, deactivate the **Locked** checkbox to release the slider tool for use. By default, this checkbox is activated (slider is locked).
- 8. From the Resolve3D main menu, use the **Acquire** button with the Z control to adjust the focal plane.

When you control the incident angle of illumination, an evanescent field is generated. Adjusting the angle changes the depth of light penetration into the sample.

You can adjust all laser angles together, or you can adjust them separately when you activate the **Adjust channels independently** checkbox.

- 9. Adjust the TIRF angle as follows:
  - **Coarse adjustments** are made to the TIRF angle by dragging and dropping the box in the slider bar using the left mouse button. An image is acquired when the mouse button is released.
  - **Fine adjustments** are made to the TIRF angle by clicking the arrows on either end of the slider bar or by using the left or right arrows on the workstation keyboard.
- 10. To illuminate the sample and not generate an evanescent wave, click the **Epifluorescence** button.
- 11. To return the laser to that last incident angle used, click the **Restore TIRF Depth** button.

12. Adjust the slider for the **Laser Path Splitter** to determine the amount of light to send to the TIRF light path.

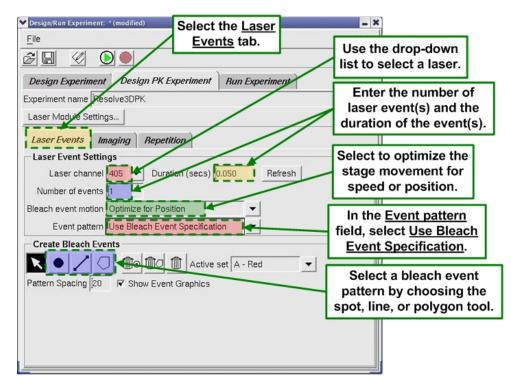


## Photokinetic (PK) Experiments

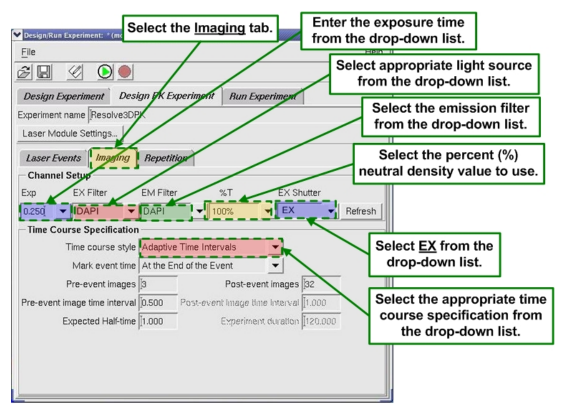
A template within softWoRx, in the Resolve3D Design/Run Experiment window contains basic photokinetic experiments such as FRAP (fluorescence recovery after photobleaching).

### To run a basic PK experiment:

- 1. Click the **Experiment** button in the Resolve3D main menu to open the Design/Run Experiment window.
- 2. Select the **Design PK Experiment** tab.
- 3. On the Laser Events tab, select the desired laser for the drop-down menu.



- 4. To generate multiple bleach events, the stage moves the sample to the site of laser activity. You can optimize this motion for either speed or position. When **Optimize for Position** is selected, the stage goes through an LMC (Lost Motion Compensation) move to achieve the most accurate position. This move takes more time than when **Optimize for Speed** is selected.
- 5. Bleach events occur at either the center point of the field of view or in a specified position. Select **Use Bleach Event Specification** in the **Event pattern** field.
- 6. In the Create Bleach Events section of the window, select an event pattern by choosing the spot, line, or polygon tool.
- 7. Click on the image in the Data Collection window to generate the selected bleach pattern.
- 8. Click on the **Imaging** tab to select the **Channel Setup** (exposure time, wavelength, and illumination source settings) and **Time Course Specification** conditions.



- 9. When you select the Time course style, the drop-down list presents three choices:
  - Adaptive Time Intervals allows you to select the total number of images to be collected. Images are collected faster after the bleach event and then slower over time.
  - Uniform Time Intervals allows you to select the total number of images collected, as well as the time interval between images. The value in the Experiment Duration field is calculated based on these factors. Also, if you change the value in the Experiment Duration field, the Post-event images field automatically updates.
  - As Fast As Possible collects the images in rapid succession with no time-lapse interval. For this style, you enter only the total number of Postevent images.

# The UltimateFocus® Module

After the operator has found the proper focus and marked a point of interest, the UltimateFocus Module maintains focus for that point during the entire experiment. The UltimateFocus Module uses an infrared laser that follows the illumination path and bounces off the cover slip/sample interface. The reflected beam is evaluated and the software returns an offset to the Z motor for automatic stage adjustment to maintain focus. The UltimateFocus Module is optional for a *DeltaVision* system. For more information on the UltimateFocus Module, see

*Using UltimateFocus*<sup>®</sup> on Page 143. For complete information on laser safety and proper labeling for the UltimateFocus Module, refer to *Appendix F*, *Lasers and Safety Issues*.

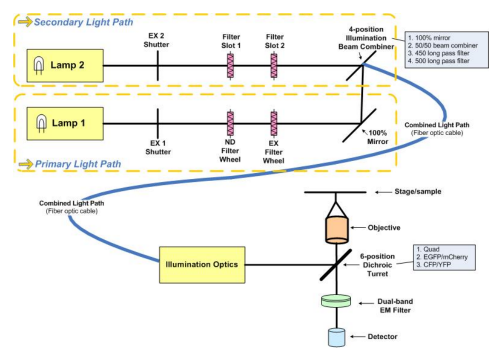
# **Analysis Workstations**

You can purchase additional *softWoRx* Linux analysis workstations.

The *softWoRx* Linux workstation includes all of the *softWoRx* Analysis modules, including 3D Visualization tools, Colocalization, Fluorescence Resonance Energy Transfer (FRET) Analysis, Fluorescence Recovery After Photo-bleaching (FRAP) Analysis, Intensity and Distance Measurement, and Modeling.

# Multiplexed Wavelength Module

The optional Multiplexed Wavelength module for the *DeltaVision* systems allows you to perform nearly simultaneous two-channel imaging without the drawbacks associated with true simultaneous two-channel imaging. This option uses two shuttered illumination sources and a dual-band emission filter to eliminate filter wheel movement between channels, and therefore greatly reduces the time required for the *DeltaVision* system to acquire a set of two-channel images. The combined light path ensures no registration artifacts are introduced and independent excitation of probes helps to ensure minimal crosstalk.



Conceptual View of Multiplexed Wavelength Functionality

Before you use the Multiplexed Wavelength option, you must first have it installed and configured correctly. Your Applied Precision representative will assist you in setting up this option and help you ensure that all hardware and software to support Multiplexed Wavelength functionality is installed properly.

After the option has been installed and configured, the menus, tools, and other infrastructure necessary to use the feature will be available on your workstation.

# Software

## softWoRx Explorer

*softWoRx*<sup>°</sup> Explorer is a cross-platform image viewer that is available for many commonly used operating systems.

*softWoRx* Explorer allows you to view and explore *DeltaVision* images and images from other sources that contain spatial, temporal, and spectral ranges. In addition to displaying data in the X and Y plane, you can scroll through Z sections and time-lapse data. Individual spectra (i.e., channels or fluorescent wavelengths) can be hidden or displayed in a variety of colors.

## softWoRx Suite for Windows Option

*softWoRx* Suite is a Windows-based ensemble of software developed by Applied Precision in collaboration with Bitplane AG. It provides sophisticated multidimensional data visualization, analysis, image restoration, image correction, and image viewing management, all within an easy-to-use streamlined browser interface.

## softWoRx Suite Advanced Option

The Advanced Option for *softWoRx* Suite includes two sophisticated analysis features: *4-D Particle Tracking* and *ImarisColoc. softWoRx* Suite Advanced Option is offered by Applied Precision in collaboration with Bitplane, AG.

4-D Particle Tracking allows users to observe temporal changes of objects. This tracking module offers a choice of methods for both detection and tracking and allows analysis and measurement of various object properties.

The Colocalization feature enables users to easily isolate, visualize, and quantify regional overlap in 3D and 4D images. Results can be presented in two ways—as a new 3D or 4D channel or as a statistical report.

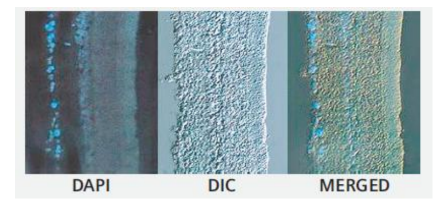
## softWoRx DMS

*softWoRx* DMS (Data Management Solution) provides a functional infrastructure for the storage of biological images and their associated metadata in a centralized database. Like *softWoRx* Suite, *softWoRx* DMS includes the Browser and the Explorer programs, but unlike *softWoRx* Suite does not have the ability to perform any type of image processing. The DMS product actually has two distinctive parts. One part is a *DMS Server* and the other part is represented by the *DMS clients*.

A DMS Server is made up of a specific configuration of the centralized database. The DMS clients are the different programs and methodologies of accessing the image data within the centralized database (ie. *softWoRx*, *softWoRx* Suite, and *softWoRx* DMS).

# Differential Interference Contrast (DIC) Module

Differential Interference Contrast (DIC) microscopy is a method for imaging unstained live or fixed specimens that may appear nearly invisible with conventional brightfield microscopy. The resulting image looks as if the specimen is casting a shadow to one side.



The shadow primarily represents a difference in the refractive index and thickness of the specimen rather than its topology. With this method of imaging, a DIC prism (known as a Wollaston Prism) splits an incident beam of plane-polarized light so that one part of the beam passes through one region of the specimen and the other passes through a closely adjacent region. A second prism then reassembles the two beams. Minute differences in the thickness or in the refractive indices between adjacent parts of a sample are converted into bright regions (if the two beams are in phase when they recombine) and dark regions (if they are out of phase).



**DIC Image Samples** 

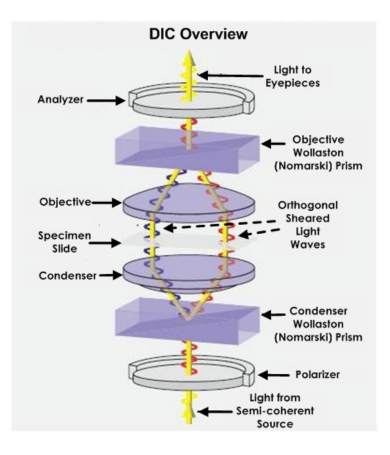
DIC microscopy has proven to be a very important technique for observing even very small objects, such as single microtubules (after digital enhancement) or acquiring useful reference images in combination with fluorescence images from the same focal plane.

# **DIC Optics**

The arrangement of the four essential DIC optics in the order of their location in the optical pathway from the transmitted light source to the image plane include:

- A polarizer in front of the condenser to produce a plane polarized light source.
- A condenser DIC prism mounted close to the front aperture of the condenser to act as a beam splitter.
- An objective DIC prism mounted close to the back aperture of the objective lens to recombine the two beams in the objective back aperture.
- An analyzer to "analyze" rays of plane and elliptically polarized light coming from the objective and to transmit plane polarized light that is able to interfere and generate an image in the image plane.

The following illustration shows the basic layout for DIC.

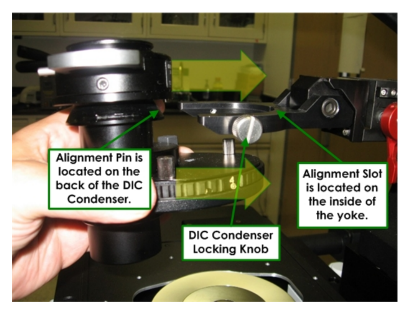


## Installing the DIC Module

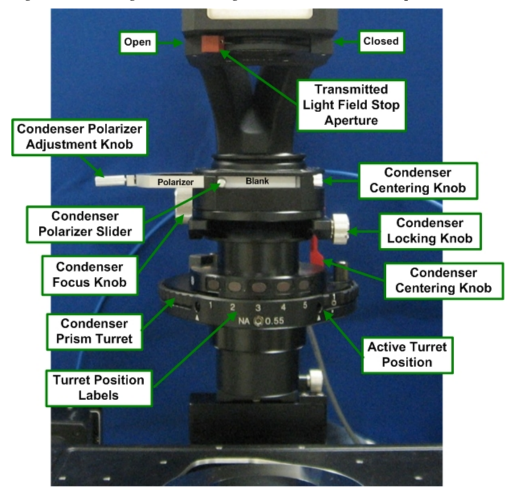
The DIC module is installed into the yoke on the *DeltaVision* system's transmitted light pillar.

### To install a DIC module on a DeltaVision system:

1. Align the pin on the back of the DIC condenser with the slot on the inside of the yoke.



2. Tighten the locking knob on the right to fix the condenser in place.



**Overview of DIC Controls** 

# Aligning the DIC Module

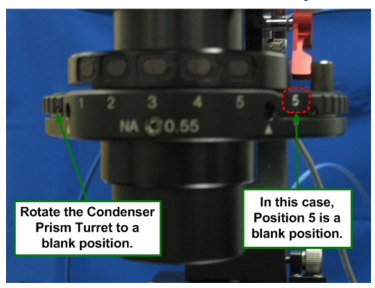
To acquire successful DIC images, it is necessary to properly align all of the DIC components. The alignment process includes setting up Köhler Illumination for the transmitted light source as well as crossing (aligning to 90 degrees) the Condenser Polarizer with both the Emission Polarizer and Eyepiece Polarizer. The Condenser Polarizer is left stationary and the other two polarizers are oriented according to the Condenser Polarizer's position. To accomplish this, you will first remove the Eyepiece Polarizer and use it to find the proper position for the Emission Polarizer. When satisfied with the position of the Emission Polarizer in relation to the Condenser Polarizer, you then replace and cross the Eyepiece Polarizer with the Condenser Polarizer.

## To set up Köhler Illumination:



1. Move the Condenser Polarizer slider to the Blank position, as shown.

2. Rotate the Condenser Prism Turret to a blank position.

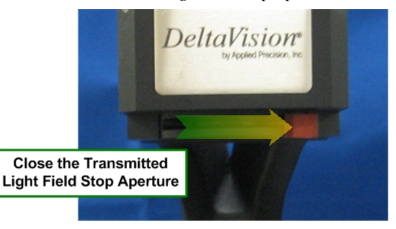


3. From beneath the right underside of the microscope stage, remove the Principal Prism Slider from the light path, as shown.



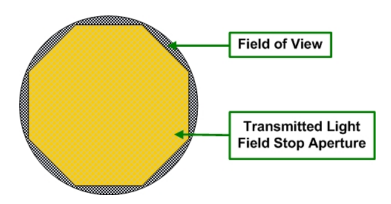
Removing the Principal Prism Slider from the Light Path

- Note It is not necessary to completely remove the Principal Prism Slider from the system. The slider clicks into place in both the In and Out positions.
- 4. Using the smallest magnification objective available, mount a specimen slide and focus on the specimen using either brightfield or fluorescence.
- Note With higher magnification objectives, 60x or greater, it may be difficult to see the edges of the Transmitted Light Field Stop Aperture.
- 5. Move the Transmitted Light Field Stop Aperture to the Closed position.



- 6. Use the *DeltaVision* keypad to open the Transmitted Light Shutter.
- 7. Observe the specimen through the oculars while turning the Condenser Focus Knob. You will see the Transmitted Light Field Stop Aperture encroaching on the image as you get close to the focal point. Keep focusing the Condenser until the edges of the Transmitted Light Field Stop Aperture (the blades of the iris) are in sharp focus.

Note If you lose light as you approach the focal point, you'll probably need to re-center the condenser until the entire field becomes bright, and then continue focusing.



If you cannot see the entire iris while moving the Condenser Focus Knob, it may be necessary to alter the centering before you are able focus on the edges of the iris.

- Center the Transmitted Light Field Stop Aperture within the field of view using the two Condenser Centering Knobs.
- Repeat this step as necessary to achieve optimum focus and alignment.
- 8. After the DIC Condenser has been focused and centered, move the Transmitted Light Field Stop Aperture toward the Open position until the aperture is just out of view.

The DIC Module will remain in Kohler alignment when different objectives are used, however, you'll need to adjust the Transmitted Light Field Stop Aperture (opened or closed to keep the aperture just outside of the field of view) for each different magnification. Also, focus can change when moving the DIC condenser so it's important to check Köhler illumination frequently.

## Before you proceed to crossing polarizers:

If you are able to verify *maximum extinction* as follows, you may not need to perform the procedure for crossing the Emission Polarizer with the Condenser Polarizer:

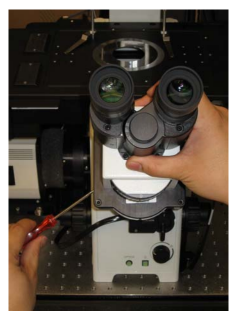
- Verify that both the Condenser Polarizer and the Emission Polarizer are in the light path.
- Acquire an image. Adjust Exposure Time and %T to obtain maximum intensity between 500 and 1500 counts.
- Acquire images continuously (File | Continuous Acquire) as you slowly move the Condenser Polarizer Adjustment knob until the lowest maximum intensity is reached. This point is maximum extinction. This point should be close to the center of the knob's travel, with the maximum intensity increasing on either side.

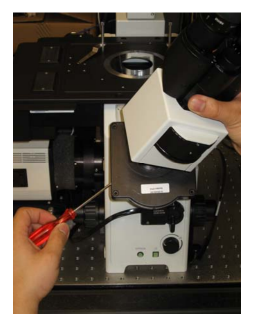
Note If a minimum intensity (maximum extinction) cannot be found, the orientation of the Emission Filter Wheel must be adjusted as described in the following procedures.

• If you are able to attain maximum extinction, tighten the Condenser Polarizer Adjustment knob to lock the polarizer in place at that position. If this is the case, you will not need to cross the Emission and Condenser Polarizers. Skip the following section and continue with the procedure for crossing the Eyepiece Polarizer with the Condenser Polarizer.

### To cross the Emission Polarizer with the Condenser Polarizer:

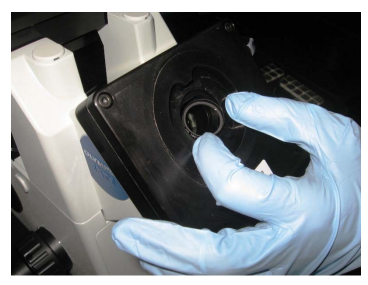
- 1. Remove any specimens from the stage and clean the objective.
- 2. Set the Eyepiece and Emission filter wheels to the Polarizer position.
- 3. Remove the Eyepiece Polarizer from the Eyepiece filter wheel. (See *Changing Filter Wheel Modules* on Page 225 for illustrated details on removing the Eyepiece assembly.)
  - Support the white Eyepiece assembly and use a 3mm hex key to remove the Eyepiece assembly from the Eyepiece Filter Wheel. Set the Eyepiece assembly aside. Leave the Eyepiece Filter Wheel attached to the microscope base.





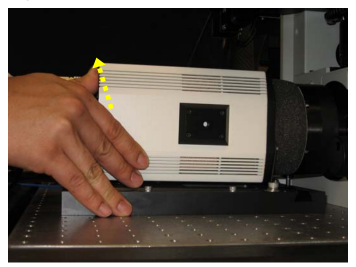
**Removing the Eyepiece** 

• Carefully remove the Polarizer from the Eyepiece Filter Wheel and set it aside. Be careful not to touch the surface of the filter.



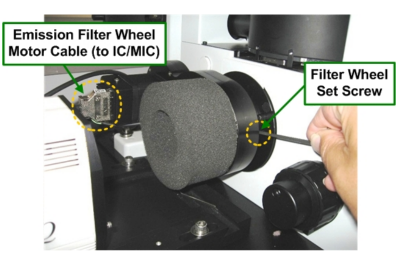
Removing the Polarizer from the Eyepiece Filter Wheel

- Leave the white Eyepiece assembly off for now.
- 4. To gain access to the Emission Filter Wheel, remove the camera by pushing in and pulling up on the camera end as shown. Refer to *Changing Cameras* on Page 221 for removal details.



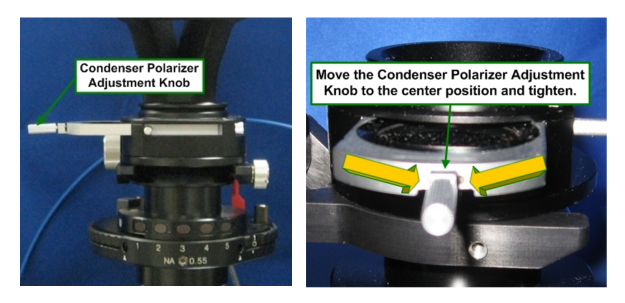
### Removing the Camera

- 5. Using a 3mm hex key, loosen the set screw to remove the Emission Filter Wheel assembly.
- Note Leave the Emission Filter Wheel Motor Cable connected, but use care not to strain this connection.



Removing the Emission Filter Wheel

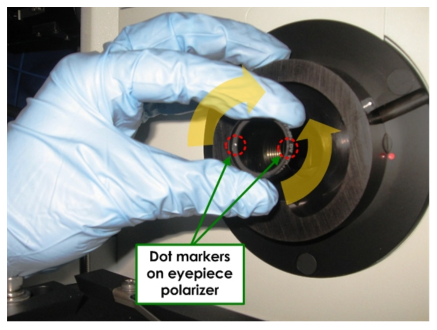
6. Using the Condenser Polarizer Slider, move the Condenser Polarizer into the light path. Move the Condenser Polarizer Adjustment Knob into the center of its travel and tighten the knob to secure it.



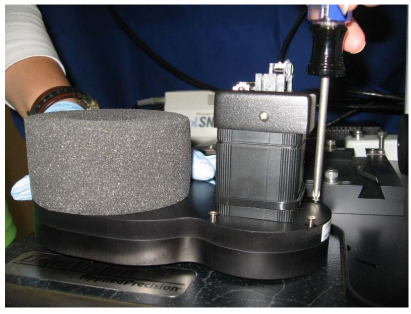
- 7. Verify that the Condenser Prism Turret is in a blank position and that the Principal Prism Slider is not in the light path.
- 8. Switch the Port Selector on the microscope to the **Camera** icon.



- 9. Adjust the T% to 50% and turn on the transmitted light.
- 10. Hold the Eyepiece Polarizer you removed in Step 3 in front of the camera side port and rotate the polarizer until maximum light extinction occurs. Note the orientation of the dots on the rim of the polarizer. This is the correct orientation for the Emission Filter Wheel Polarizer.

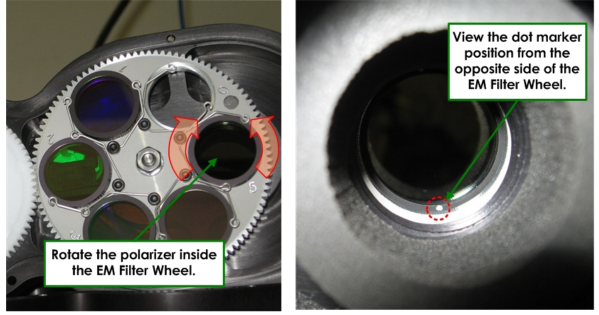


11. Use a Phillips screwdriver to remove the top plate from the Emission Filter Wheel assembly.

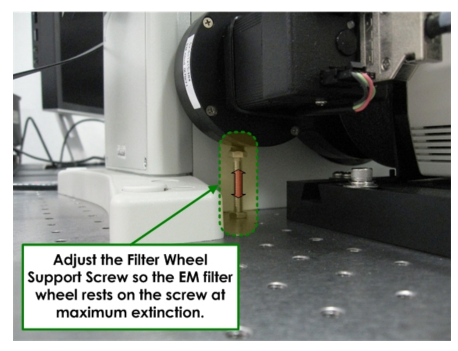


Removing the Emission Filter Wheel Top Plate

- Note Eight Phillips screws hold the EM filter cover in place. Some of the screws are located beneath the foam collar.
- 12. Holding the filter wheel in the same position in which it will be installed on the system, rotate the Polarizer within the Emission Filter Wheel until the orientation of the dots matches those determined for the Eyepiece Polarizer.



13. Reassemble the Emission Filter Wheel and reinstall the unit back onto the microscope. Adjust the Filter Wheel Support Screw so that the Emission Filter Wheel rests on the screw at maximum extinction as observed through the EM filter.



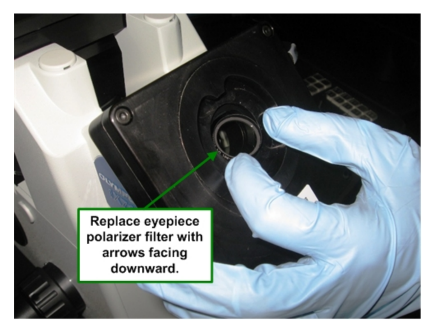
- 14. Repeat Steps 10 13 to align any additional filter wheels.
- Note When aligning <u>additional</u> filter wheels, make sure the filter wheel rests on the Filter Wheel Support Screw, but **Do NOT** adjust the Filter Wheel Support Screw to achieve maximum extinction. You <u>must</u> rotate the Polarizer within each filter wheel assembly.
- 15. Move the camera back into its operating position on the *DeltaVision* system and verify extinction as follows:
  - Verify that both the Condenser Polarizer and the Emission Polarizer are in the light path.
  - Acquire an image. Adjust Exposure Time and %T to obtain maximum intensity between 500 and 1500 counts.
  - Acquire images continuously (File | Continuous Acquire) as you slowly move the Condenser Polarizer Adjustment knob until the lowest maximum intensity is reached. This point is maximum extinction. This point should be close to the center of the knob's travel, with the maximum intensity increasing on either side.

**Note** If a minimum intensity cannot be found, the orientation of the Emission Filter Wheel must be adjusted.

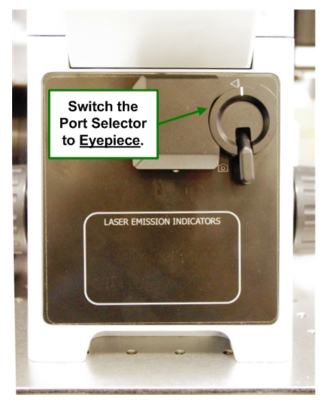
• Tighten the Condenser Polarizer Adjustment knob at maximum extinction.

### To cross the Eyepiece Polarizer with the Condenser Polarizer:

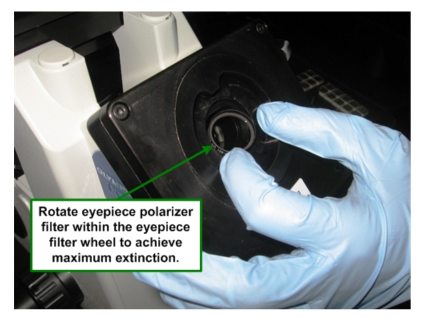
1. Return the Eyepiece Polarizer to the Eyepiece Filter Wheel with the arrows facing away from you.



2. Switch the Port Selector on the front of the microscope to the **Eyepiece** <



3. As it rests in its holder, rotate the Eyepiece Polarizer until maximum extinction is achieved.



4. Return the white Eyepiece assembly to the Eyepiece Filter Wheel on the microscope base and tighten with a 3mm hex key.



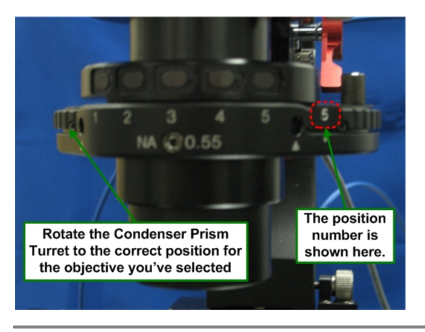
**Reconnecting the Eyepiece** 

# Setting Up DIC

After you have completed all of the DIC alignment tasks described previously, you are ready to perform the DIC setup procedure.

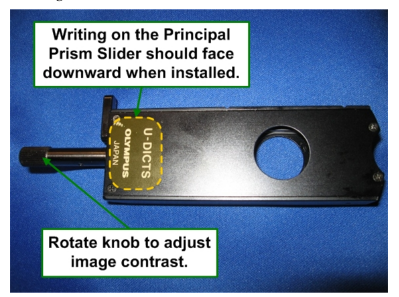
## To set up DIC:

- 1. Verify that all three of the polarizers (Condenser, Emission, and Eyepiece) are in the light path.
- 2. Move the Condenser Prism Turret to the correct position for the selected objective.



**Note** The above image does not show the objective labels that would normally be visible above the numbers.

3. Move the Principal Prism Slider into the light path under the objective (it will click into place). For proper orientation, be sure that the writing on the slider is facing down.





The Principal Prism Slider has two positions: **Normal** and **BFP1** (as shown below). The selected position will depend on the type of objective being used. If you are unsure, check the objective. It will say **BFP1** if that is the correct position.



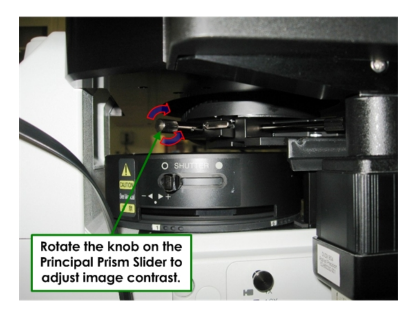


4. Switch the Port Selector to the **Eyepiece** < icon.

- 5. Return the sample to the stage to begin imaging.
- 6. Switch the Port Selector to the **Camera** icon.



7. Acquire images continuously (**File** | **Continuous Acquire**) as you rotate the knob on the Principal Prism Slider to adjust the contrast displayed in the images. For example, move the shadows from bottom to top and move the black line into or out of the image.



# **Consumable Parts**

#### Common to 100-120 V and 220-240 V Systems

Component	Fuse for Component	API Part Number
IC/MIC	6.3A 250V UL High Break Capacity	19-170045-000
Microscope	Т5АН	Used in some older DeltaVision systems

#### Bulbs

Component	API Part Number
250W Xenon Arc Lamp Bulb	34-100390-000
LED Transmitted Light (must replace entire assembly)	52-851243-000

#### **British Power Cord**

Component	Fuse for Component	API Part Number
Power cord with Bussman TDC180-10A built into plug. (Must meet British Standard BS1362.)	10A	19-210046-000

In addition to these specific parts, objectives, filters, laser heads, bulbs, slides, and batteries for *DeltaVision* are also considered consumable parts. Check with your Applied Precision representative or contact the Applied Precision Customer Hotline for additional information on consumable parts.

# 8

# **Changing Cameras and Filters**

This chapter provides the following instructions for cameras and filters:

- Changing Cameras
- Using Live Cell or Custom Filter Wheel Modules
- Changing Filter Wheel Modules
- Calibrating Filter Wheels

# **Changing Cameras**

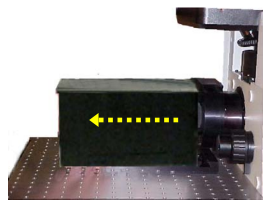
Your DeltaVision system may include up to two connected cameras. If you have the optional EMCCD camera, you can change cameras to better meet your imaging needs.

These instructions show how to change the camera and how to select the new camera in Resolve3D.

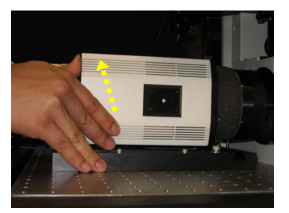
**CAUTION:** Contamination from fingerprints or dust on the camera window or inside the Emission filter wheel will degrade image quality.

#### To change the camera:

1. Remove the camera cover by lifting it up and sliding it away from the microscope.



2. Remove the camera by pushing in and then pulling up on the camera end. You do not need to remove cables from the camera.



3. Remove the other camera from the spare camera tray and set it aside. Place the camera that you just removed on the spare camera tray.

**CAUTION:** Always place the camera in the camera tray when it is not secured to the system. Dropping the camera will cause severe damage.

- 4. Install the new camera by sliding it into place and then pushing in and pressing down on the end of the camera.
- 5. Replace the camera cover.

#### To select the camera:

- 1. In the Resolve3D window, click **Settings** to open the Resolve3D Settings window.
- 2. On the **Imaging** tab, click the Camera list and select the camera that is currently installed.

Display	Imaging	Files	Autofocus	Misc	QLM
	Camera	COOLS	NAP_ES2/IC	X285 -	·
Fram	es to average*	Ĭ1			
	Gain	4.00	•		
	Transfer speed	5000	•		
Tarç	jet temperature	0.00	L		
Curre	ent temperature	0.00	Refresh		
'EX 1" b	ulb age (hours)	557.0	Refresh	Reset	[
'EX 2" b	ulb age (hours)	488.0	Refresh	Reset	[
		🔽 Use j	hotosensor		
Setting	s change base	d on Excit	ation Filter)		
	Save Settings	1			He

#### Notes

**#1** The EMCCD camera is listed twice in the Camera list:

CASCADE2\_512 Conv./... sets the camera in Conventional mode. Cascade2\_512 EMCCD/... sets the camera in electronmultiplication

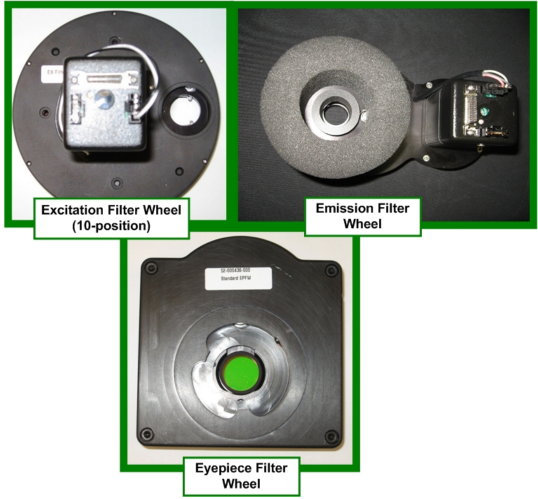
mode.

**#2** When the camera is changed, the binning number is reset to 1.

**#3** If the camera cables loosen or are accidentally disconnected, you may need to restart the IC software. To reseat the cables, power the camera down and consult the camera documentation.

# Using Live Cell or Custom Filter Wheel Modules

A filter wheel module includes an Excitation filter wheel for xenon light sources, an Emission filter wheel, and an Eyepiece filter wheel.



# DeltaVision Filter Wheel Module

A filter wheel module includes the Excitation, Emission, and Eyepiece filter wheels and can support up to six filter pairs.

Note The appearance of the filter wheels will depend on the options included on your system. For example, if your system is equipped with the InsightSSI broadband light source, it may not include the excitation filter wheel described in this section.

If your system has an alternate filter wheel module, you can swap modules to meet your imaging needs.

You can purchase a module of Live Cell filter wheels that includes factoryinstalled filters. You can also purchase a module of empty filter wheels and customize it with your own filters.

### **Changing Filter Wheel Modules**

Changing the filter wheel modules for your system includes changing the Eyepiece and Emission filter wheels and, depending on how many filter pairs are supported, may also include changing the Excitation filter wheel.

If you purchased a filter wheel module as a separate component after you purchased your system, you must configure the instrument controller for the new module the first time that you use it (see the instructions included with your filter wheel module).

#### **Before You Begin**

Before you begin the process of changing filter wheel modules, it's helpful to understand the entire procedure in general.

Basically, the procedure for changing filter wheels is as follows:

- a. Turn the IC/MIC off.
- b. Change the Excitation filter wheel if necessary (not available on SSI).
- c. Change the Emission filter wheel.
- d. Change the Eyepiece filter wheel.
- e. Restart and initialize the IC/MIC.
- f. Select the new filter modules in the R3D **Misc** tab.

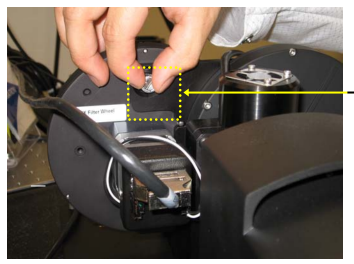
The following subsections will walk you through the procedures for changing each of the filter wheels.

**WARNING:** To avoid eye injury, before you unplug any motor cables, make sure that the IC/MIC and any xenon lamps are turned OFF.

#### Changing the EX Filter Wheel

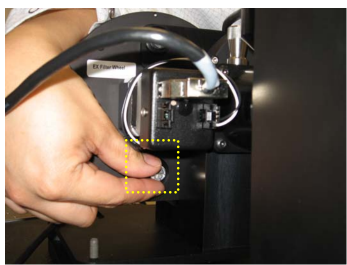
#### To change the Excitation Filter Wheel (if necessary):

- 1. Ensure that the IC/MIC is turned off, and make sure that any attached xenon arc lamps have been off and allowed to cool for several minutes.
- 2. Loosen the two silver thumb screws at the top and bottom of the filter wheel housing.



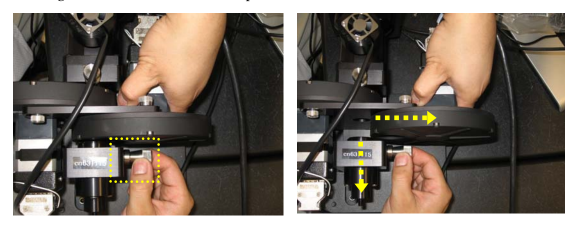
Thumb screws should loosen without the use of tools, but can also be removed with a flat-blade screwdriver if necessary.

Loosening top thumb screw from filter wheel housing



Loosening bottom thumb screw from filter wheel housing

3. Loosen the Light Seal thumb screw to disengage it from the beveled support on the Excitation filter wheel housing and gently slide the filter wheel housing outward from its mounted position.



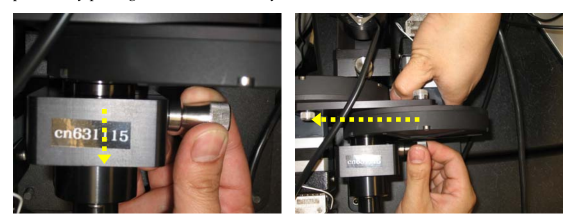
Removing the Excitation Filter Wheel housing from its mounted position

4. Disconnect the cable to the IC/MIC from the Excitation filter wheel and connect it to the alternate Excitation filter wheel.



Disconnecting the IC/MIC cable from the Excitation Filter Wheel

5. Slide the alternate Excitation filter wheel housing back into its mounting position by pulling back on the assembly as shown.



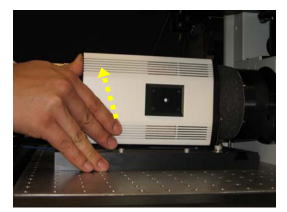
Pulling back the housing and re-engaging the filter wheel

- 6. Re-engage the housing into the beveled support on the new Excitation filter wheel housing and align and tighten the two thumb screws at the top and bottom of the Excitation filter wheel housing.
- 7. Tighten the thumb screw on the Focusing Lens housing.

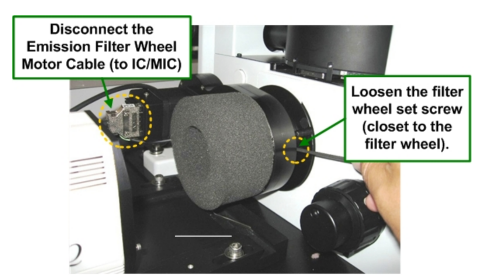
#### Changing the EM Filter Wheel

#### To change the Emission filter wheel:

- 1. Turn off the IC/MIC (if it's not already off).
- 2. Remove the camera cover. Then remove the camera by pushing in and pulling up on the camera end as shown below.



3. On the Emission filter wheel motor, disconnect the cable that connects the motor to the IC/MIC. Use a 3 mm hex key to loosen the set screw that holds the Emission filter wheel to the microscope.



4. Remove the Emission filter wheel.



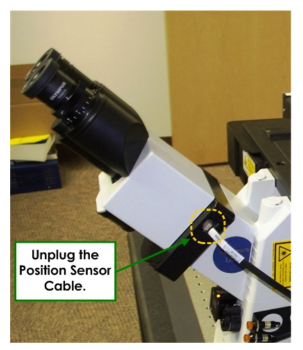
5. Install the alternate Emission filter wheel. Tighten the Emission filter wheel set screw and connect the cable from the IC/MIC to the new Emission filter wheel.

6. Reinstall the camera by sliding it into place and then pushing in and pressing down on the camera.

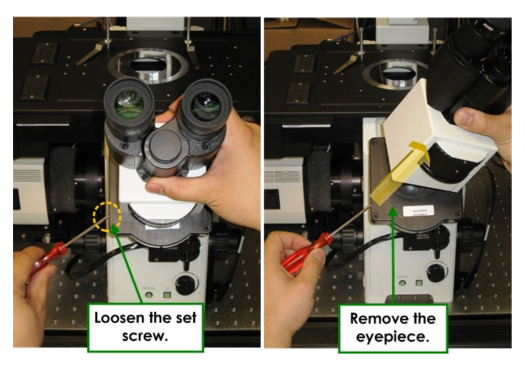
### Changing the EP Filter Wheel

#### To change the Eyepiece filter wheel:

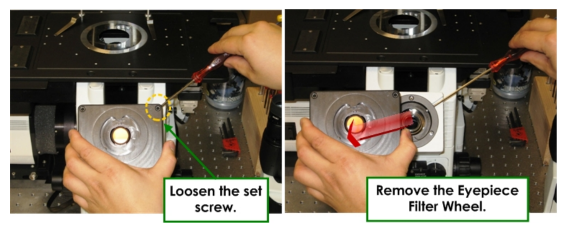
- 1. Turn off the IC/MIC (if it's not already off).
- 2. Disconnect the Position Sensor cable that connects the filter wheel to the IC/MIC. *DeltaVision* displays a message that indicates the cable is removed.



3. Holding the eyepiece in one hand, use a 3 mm hex key (Olympus provides one with the microscope) to loosen the set screw that holds the eyepiece to the Eyepiece filter wheel and set the eyepiece on the table.



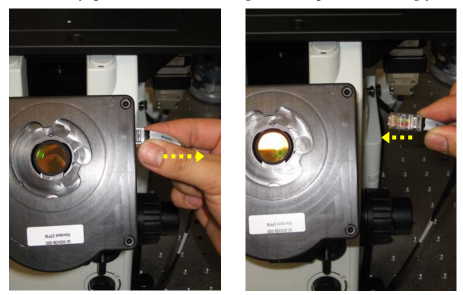
4. Loosen the set screw that holds the Eyepiece filter wheel to the beveled mount on the stand and remove the Eyepiece filter wheel.



- 5. Place the new Eyepiece filter wheel on the beveled mount on the stand and tighten the set screw that holds it in place.
- 6. Connect the Position Sensor cable to the new Eyepiece filter wheel. The IC/MIC will automatically start to re-initialize the EM and EX filter wheels even though the current EX and EM filter wheels have not yet been replaced. You can ignore this re-initialization.
- 7. Place the oculars on the microscope and tighten the set screw that holds them in place.

# **Completing the Filter Wheel Change**

- 1. When you are finished changing all filter wheels, restart the IC/MIC.
- 2. Open Resolve3D and wait for the system to initialize.
- 3. If Resolve3D does not automatically display the correct filter module, there are two ways to expressly let the system know you have installed a new filter set:
  - a. From the Resolve3D window, select **Settings**, change to the **Misc** tab, and select the current filter set.
  - b. Unplug and then re-plug in the Eyepiece filter wheel position sensor cable (see below). The system will automatically recognize which filter set the Eyepiece filter wheel belongs to and update accordingly.



Note that it is not necessary to remove the oculars before plugging or unplugging the Eyepiece filter wheel.

4. To select the current filter set from the **Misc** tab, click **Settings** in the Resolve3D main menu to open the Resolve3D Settings window. Then click the **Misc** tab.

Resolve3D S	Settings				_ ×
Display	Imaging	Files	Autofocus	Misc	QLM
Stage View ▼ Show st ▼ Show st ▼ Show pc	ost Motion Co v Options – age trails age thumbna pint numbers		tion (LMC)		
Filter Whe Excitation fi Emission fi Eyepiece fi Activate F	Iter wheel L	iveCell iveCell iveCell	• • •		
Done Sa	ave Settings				Help

- 5. Select the newly installed Excitation, Emission, and Eyepiece filter wheels in their respective fields in the **Misc** tab.
- 6. Click the Activate Filter Sets button.

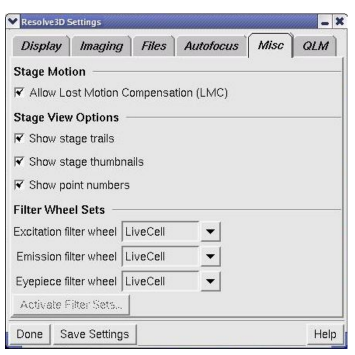
# **Calibrating the Filter Wheels**

Calibration initializes the filter positions and ensures that the filters are centered in the filter wheel openings.

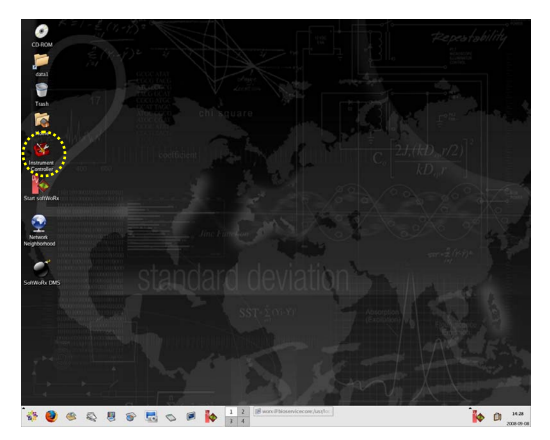
If you notice poor light transmittance or poor image quality, one possible cause is a misaligned filter wheel. The filter wheel calibration re-establishes the zero position of the filter wheel for the Instrument Controller.

To calibrate the Neutral Density, Excitation, and/or Emission filter wheels:

1. On the Resolve3D window, click **Settings** to open the Resolve3D Settings window. Then click the **Misc** tab.



- 2. Verify that the **Excitation**, **Emission**, and **Eyepiece** filter wheel fields all display the expected and identical filter wheel module name (if any of the names are different, you may need to select or replace one or more of the filter wheels; see *Changing Filter Wheel Modules* on Page 225).
- 3. Press Save Settings and close Resolve3D.
- 4. Click on the Instrument Controller icon on the workstation desktop as shown.

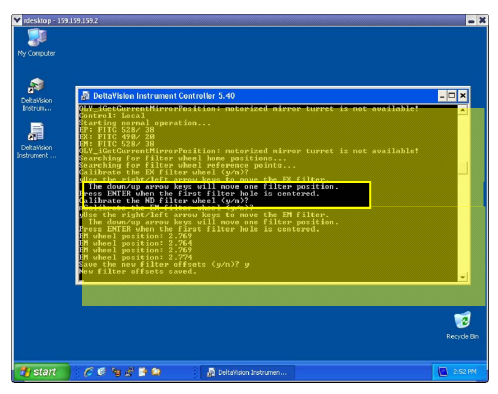


- 5. Log on to the Instrument Controller with the user name worx and password 4delta if prompted.
- 6. Press Alt + H. The filter wheels will rotate as they are being initialized.

You are then prompted with, "Calibrate the EX filter wheel (y/n)."

- 7. Remove the filter wheel to be calibrated (Excitation, Emission, or Neutral Density) from the system so the opening is visible.
- 8. If you are calibrating the Excitation Filter Wheel, press **y**. Otherwise press **n** to move to the next filter wheel (**ND** or **EM**).

When you press **y**, the following message is displayed:



Pressing the up or down arrow keys advances the filter wheel one complete position. The left and right arrow keys move the filter wheel in small increments.

- 9. Find the Home filter position on the filter wheel. The Home filter position is marked on each of the filter wheels as follows:
  - EX = **1**
  - ND = **1**
  - EM = 0 (for 6-position filter wheels) or 1 (for 10-position filter wheels)

The position is stamped on the surface of the filter wheel and may be viewed through the filter wheel hole.



Filter position not centered

Filter position centered

- 10. Once the filter wheel is centered, press Enter. After all necessary filter wheels have been centered you are prompted with, "Save the new filter offsets (y/n)."
- 11. If you made a mistake during calibration you can press **n** and start the process over; otherwise press **y** to save your offsets.
- 12. Re-install the filter wheels in the system.
- 13. To shut down the IC/MIC, go to the Start tab and select **Windows Security**. Then select the **Shutdown** option.

Yrdesktop - 159.159.159.2			- ×
rdesktop - 159.159.159.2 WOTX Internet Internet Explorer E-mail Outlook Express ic540_dv.ini072808.txt ic540_dv.ini072808.txt ic540_dv.ini072808.txt OelkaVision Instrument Controller Notepad	My Documents My Recent Documents My Pictures My Pictures My Music My Computer My Computer Control Panel Set Program Access and Def Printers and Faxes	sitions use points )? n)? n)? nove the EM filter. nove filter position. hole is centered.	X
All Programs	<ul> <li>Help and Support</li> <li>Search</li> <li>Run</li> <li>Windows Security</li> <li>Log Off O Disconnect</li> </ul>	y	r Recycle Bin
🦺 start 👘 🖉 🙆	🛃 🃴 😂 🔡 DeltaV	ision Instrumen	🚇 2:56 PM

The IC/MIC will power down.

14. Power the IC/MIC back on and boot the *DeltaVision* as normal.

# 9

# Maintenance

This chapter provides the following instructions for the basic maintenance of the system:

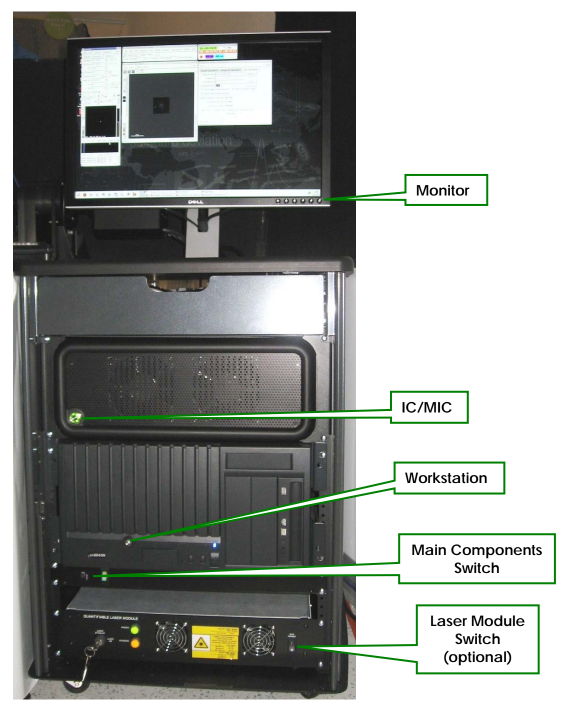
- Shutting Down and Starting the System
- Replacing the Xenon Bulb
- Replacing the Transmitted Light
- Replacing IC/MIC Fuses
- Cleaning
- Moving the System

# Shutting Down and Starting the System

Use the following instructions for shutting the system down during occasions that require total shutdowns.

# Delta Vision Power Switches

The main *DeltaVision* power switches are shown below. (The Master Switch on the isolation transformer that plugs into the wall is not shown.)



# **Guidelines for Using Switches**

#### Main Components

Use this switch to turn power on and off for the *DeltaVision* Workstation, the Camera Power Supplies, the *DeltaVision* Microscope, and some optional DeltaVision equipment.

#### Workstation, IC/MIC, and Monitor

Leave these switches on except on rare occasions (such as power outages) when you need to shut down the entire system.

## Shutting Down the System

In some situations, such as power outages, you will need to shut down the entire *DeltaVision* system.

#### To shut down the DeltaVision system:

- 1. Save all data on the workstation.
- 2. On the *softWoRx* menu bar, choose **File** | **Exit**. Then exit all other workstation applications.
- 3. From the main menu button, choose **Logout** and then **Shut Down**. Wait until the monitor displays Power Down.
- 4. Press the power button on the IC/MIC once to shutdown.
- 5. Turn off the monitor.
- 6. Turn off the main component switch.

**Note** For *personalDV*, there is no power strip switch. For *personalDV*, skip this step.

- 7. Clean the objective if necessary.
- 8. Lower the objective.

### Starting the System

Use the following instructions to start the system after a total shutdown.

#### To start the DeltaVision System:

1. Turn on the power strip bar.

- Note For *personalDV*, there is no power strip bar. Begin the power-up process with Step 2.
- 2. Turn on the IC/MIC.
- 3. Turn on the workstation.
- 4. Turn on the monitor.
- 5. Follow the instructions for turning on *DeltaVision* on Page 35.

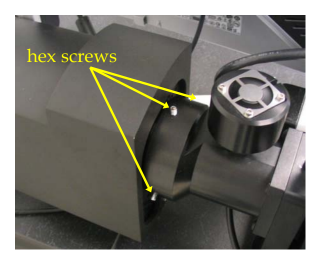
## **Replacing the Xenon Bulb**



**WARNING:** Ensure the xenon lamp is off and has had plenty of time to cool before starting this procedure. Refer to the "Xenon Lamp Safety" section in Chapter 2 of this manual for details on safety issues and proper disposal of the lamp.

Follow these steps to replace the xenon bulb on *DeltaVision*:

- 1. If the system is on, exit Resolve3D and ensure that the IC/MIC is off prior to proceeding. The fan on the lamp housing must be off before you begin this procedure.
- 2. Loosen the three hex screws in the flange of the xenon lamp housing.



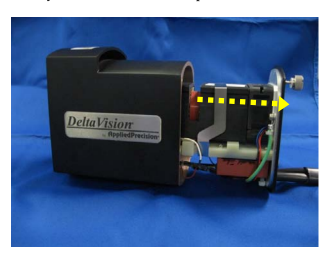
3. Gently slide the lamp housing away from the flange to remove it from the *DeltaVision* excitation module.

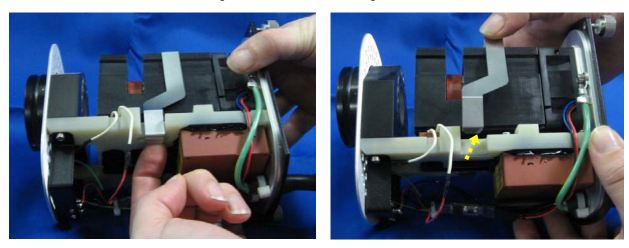


4. Loosen the two thumb screws on the opposite end of the xenon lamp housing.

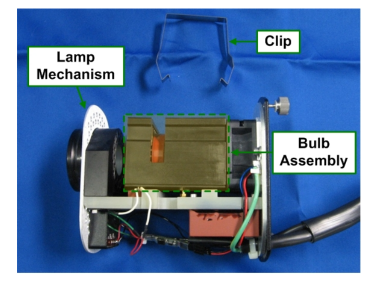


5. Gently slide the internal lamp mechanism from the lamp housing.

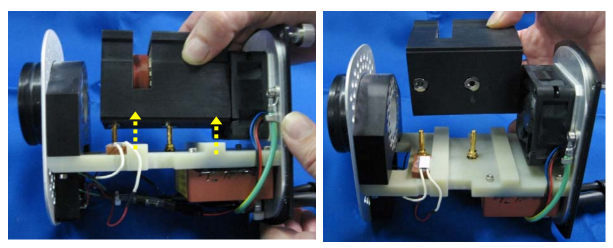




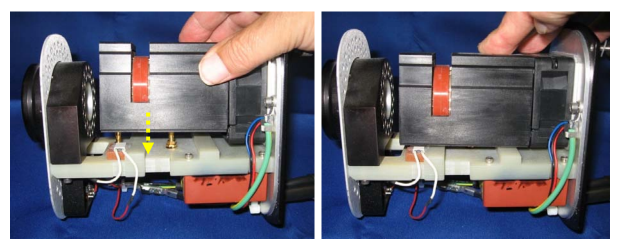
6. Remove the center clip from the internal lamp mechanism as shown.



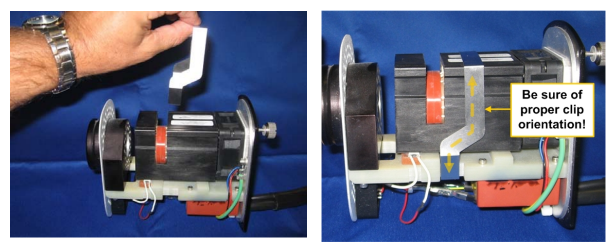
7. Lift the bulb assembly (small black box) from the two supporting pins in the lamp mechanism.



8. Replace the xenon bulb assembly with a new one (Part #34-100390-002). Insert the new bulb assembly onto the two supporting pins and press down firmly.



9. Replace the clip around the internal lamp assembly, making sure the clip is properly oriented.



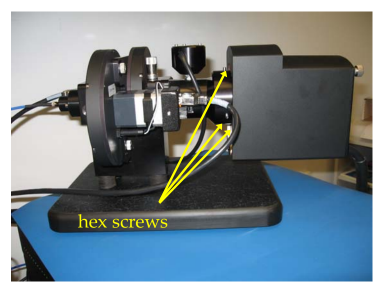
10. Gently slide the internal lamp mechanism into place within the lamp housing.



11. Tighten the two thumb screws on the end of the lamp housing.



12. Place the open end of the lamp housing over the flange on the *DeltaVision* and tighten the three hex screws as shown.



- 13. Turn the *DeltaVision* on as usual and start Resolve3D.
- 14. Before resetting the bulb age, write down the age of the bulb you just replaced. This will help you to keep track of when you may need to replace the next one.
- 15. Open the **Imaging** tab in the Resolve3D Settings window and click the **Reset** button to reset the bulb age.

Display	Imaging	Files	Autofocus	Misc	QLM	
	Camera	COOL	SNAP_ES2/IC	X285 🗨	·	
Frame	s to average*	Ĭ1				
	Gain	4.00	•			
т	ransfer speed	5000	•			
Targe	et temperature	0.00	L	В	ulb Age	
Currer	nt temperature	0.00	Refresh		-	
'EX 1" bul	b age (hours)	557.0	Refresh	Reset	$\mathbb{H}$	<u>Reset</u> B Age
'EX 2" bul	b age (hours)	488.0	Refresh	Reset		
		😿 Use p	photosensor			
Settings	change base	d on Excit	ation Filter)			
Done   S	ave Settings	0			Hel	

Note For *DeltaVision* systems with the Multiplexed Wavelength option installed, the procedure for changing the xenon bulb in the secondary lamp housing is identical to the procedure described above.

# **Replacing IC/MIC Fuses**

Follow these instructions to replace a fuse in the IC/MIC. To replace fuses for other components, follow the instructions in the manuals that are provided for those components.



**CAUTION:** Installation of improperly rated fuses can cause damage to the system.

#### To replace a fuse:

- 1. Shut down the system.
- 2. Unplug the power cord on the back of the IC/MIC.
- 3. Remove the fuse holder.
- 4. Test the fuses with a continuity meter.
- 5. Replace any bad fuses with 5X20mm 6.3A 250V UL high break capacity fuses (API P/N 19-170045-000).
- 6. Install the fuse holder.
- 7. Plug in the power cord.

# Cleaning

Most system surfaces are best cleaned with a lint-free cloth or lint-free swabs and spectroscopy-grade isopropyl alcohol or chloroform. Avoid contaminating the cleaning solution by never reusing the cleaning cloth or swabs. Operators should be trained in the handling of flammable liquids such as alcohol. Material Safety Data Sheets (MSDS) should be maintained for the cleaning solutions, as with any hazardous material.

The exceptions to this cleaning practice are the polychroic mirror and the optical filters. These components should be cleaned with low-pressure air. For example, use a bulb designed for cleaning camera lenses, which blows air across the surface. Do not use high pressure. Do not use canned air, as this often leaves a fluorescent residue.

To clean the microscope, follow the instructions in the manufacturers' manuals that are provided for these components.



**CAUTION:** Improper cleaning of the polychroic mirror and the optical filters will result in damage.

# Moving the System

If you need to move your *DeltaVision* system, call Applied Precision for instructions.



# **Appendix A: The Immersion Oil Kit**

The immersion oil kit is a collection of oils with refractive indexes that range from 1.500 to 1.534. Use of the correct immersion oil decreases the spherical aberration in the image data.

• For *DeltaVision*, the immersion oil kit includes eighteen oils that range from 1.500 to 1.534, in increments of 0.002.

Note For *personalDV*, the immersion oil kit includes six oils that range from 1.512 to 1.522, in increments of 0.002.

Many factors influence the optimum refractive index of the immersion oil, including specimen preparation, temperature, humidity, and atmospheric pressure.

# The Oil Calculator

In order to calculate the desired refractive index, *softWoRx* is equipped with the Lens Information function. This function is located in the Utilities menu in *softWoRx*. It can also be accessed from Resolve 3D by clicking the **Info** button. The following parameters are explained here to help you enter the appropriate information and use the resulting calculations.

#### Distance from Coverslip to Specimen (microns)

Establishes the distance from the surface of the coverslip to the desired focal plane.

#### Temperature

Defines the temperature of the specimen and the immersion medium.

#### Specimen Refractive Index

Defines the refractive index of the specimen, which is usually that of the mounting medium. In some cases, the specimen itself contributes significant refraction.

#### **Recommended Refractive Index**

Displays the resulting optimal refractive index of the immersion oil. Actually experimenting with oils with refractive indexes very close to this value is the best way to select the optimal oil.

#### **Resolution Ratio**

Displays the ratio between the Z resolution and the XY ratio. This serves as a reference to the degree of Z elongation.

#### Maximum XY Pixel Size

Displays the maximum recommended XY pixel size for deconvolution.

#### **Recommended Z Step**

Displays the smallest possible Z step for this objective. Choosing a smaller Z step will add to the size of the image file, but will not improve image quality.

# B

# **Appendix B: Troubleshooting**

This appendix was designed to help you diagnose and correct the most common problems encountered on the *DeltaVision* system. Two types of troubleshooting tasks are covered:

- Diagnosing System Problems
- Analyzing Reasons for Poor Image Quality

If you are unable to correct a problem, fill out the *DeltaVision* Problem Report Form at the end of this appendix and either e-mail it to hotline@api.com or fax it to 425-557-1055, attn: Bio Service Hotline.

# **Diagnosing System Problems**

# Troubleshooting the Controller

The following table shows the most common Instrument Controller problems and their resolutions.

Indication	Cause	Correction
Encoder Error when	Poor cable	Power down system, including IC/MIC.
initializing stage.	connection.	Reseat X, Y, and Z motor cables.
		Reseat motor cables on excitation module.
		Power up system.

Table B-1: Controller Troubleshooting Chart
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## Troubleshooting the Workstation

Other system troubles are indicated by messages or readings in the software. The Resolve 3D message window displays Resolve 3D activity. Observe the messages in this window when troubleshooting. This table shows possible problems and corrective actions.

Indication	Cause	Correction
"File system full"	There is no more	Delete unwanted files.
message when trying to save	storage space for image data.	Save image files to CD or DVD or LAN.
images.	5	See softWoRx Imaging Workstation User's Guide for more information.
"Camera not found" message.	Power up sequence was incorrect.	Shut down the system and then restart it using the steps described in <i>Chapter 9: Maintenance</i> .
Resolve 3D settings are not updating changes made using the keypad or joystick (for example, the filter selection, or Z position).	Lack of communication between the Instrument Controller and Workstation.	Shut down the system and then restart it using the steps described in <i>Chapter</i> <i>9: Maintenance</i> .
At login, user name not recognized.	User has not been added.	Add user. See the softWoRx Imaging Workstation User's Guide.

Table B-2: Workstation Troubleshooting Chart

# Analyzing Reasons for Poor Image Quality

The following table documents the most common acquisition difficulties and abnormalities in image data.

Indication	Cause	Correction
Dim images or long	Poor illumination.	Fully open field aperture.
exposure times.		Ensure that filter cube turret is locked in position on rail mount.
		Ensure shutter on polychroic filter wheel is open.
		Ensure slider behind microscope is seated in an open position.
		Ensure proper filter cube is in position and seated in detent.
Dim illumination. When fiber optic cable and focusing lens are removed, the projected light	Filter wheel(s) out of alignment.	Shut down and start up as described on Pages 35 and 48 or unplug and plug in the Eyepiece filter wheel. This will reset the home position of the filter wheels and align filter wheel position.
does not form a circle.		Calibrate filter wheels following instructions on Page 232.
Dark, out of focus spots on image.	Dust interference.	Clean polychroic filter, emission filter, and camera window using low-pressure air. Do not use canned air. See Page 246 for further recommendations regarding cleaning system components.
Image is distorted around edges or throughout. Occlusion seems to	Condensation on camera window, possibly due to improper camera	Clean camera window using low- pressure air. Do not use canned air. See Page 246 for further recommendations regarding cleaning system components
creep in toward center.	temperature.	Check camera temperature in Resolve3D. Consult camera documentation for proper setting.
Brightness of Z section images	A broken Photo sensor cable	Disconnect the Photo sensor cable and the EX module cable.
varies greatly		Connect the Photo sensor cable to the EX module.
		Set the Excitation filter to FITC or some other visible light.
		Open the EX shutter.
		Bend the cable and examine it for light leaks. If you observe a light leak, replace the cable.
		Direct the light to a wall. If you observe inconsistencies in the light output as you bend the cable, replace the Photosensor cable.

Table B-3: Image Quality Troubleshooting Chart

Cause	Correction
Air bubble in immersion oil.	Clean front and back surfaces of objective and coverslip.
	Reapply immersion oil and restart experiment.
Possibly dirt, dust, oil, or air bubble.	Clean front and back surfaces of objective and coverslip.
Camera saturation.	Use lower exposure time and/or higher neutral density filter.
Poorly aligned illumination.	Align xenon lamp and fiber optic cable.
	See Chapter 9, Maintenance.
Knob at base of microscope is directing light to the eyepiece.	Move knob to direct light to camera.
Stage was not centered within sample at start of experiment.	Position stage in center of the sample and run experiment again.
	Air bubble in immersion oil. Possibly dirt, dust, oil, or air bubble. Camera saturation. Camera saturation. Poorly aligned illumination. Knob at base of microscope is directing light to the eyepiece. Stage was not centered within sample at start of

# DeltaVision Problem Report Form

Research Facility:	System Serial Number:
Contact Person:	softWoRx version: (use Help → Software Versions
Phone:	to display)
E-mail:	Date:
Problem Encountered: Please write a detailed descrip	tion, answering as many of the following questions
as possible.	
Questions:	
1 When did this first occur?	
2 Was there any recent change or update to the system	prior to the problem occurring?
<ul> <li>What sequence of operations produces the problem?</li> <li>What other programs were running when you encount</li> </ul>	torod the problem?
<ul><li>4 What other programs were running when you encount</li><li>5 What error messages, if any, were shown?</li></ul>	
6 Is the failure the same each time or does it show differ	rent symptoms?
7 Does it occur consistently or is it random?	
8 Does it go away after the workstation is re-booted?	
9 Does it go away after the instrument controller is re-bo	
10 How often do you re-boot the workstation and instrum Additional Comments:	ent controller?
(See Page 2 for additional clarification issues)	
Please supply the following log files:	
Workstation: (/home/userName/softworx-logs)	/softworxlog.txt)
Instrument Controller: (c:\ic540_dv\log\IC540_dv.	log)
	1031

Please e-mail this form to <u>hotline@api.com</u> or Fax to: 425-557-1055, attn: Bio Service Hotline

# С

# **Appendix C: Acquiring a PSF**

This appendix shows how to acquire a Point Spread Function (PSF) and convert it to an Optical Transfer Function (OTF).

- *Acquiring a PSF* shows how to measure a Point Spread Function.
- Converting PSF to OTF shows how to convert the Point Spread Function to the Optical Transfer Function that is required to process images.

## **Before You Start**

Before you attempt to measure a PSF, check the OTF library included with *softWoRx* to find out if the library provides an OTF for your objective.

# Acquiring a PSF

To measure the point spread function (PSF), you need to optically section a fluorescent bead. Since the properties of the objective lens are the most important elements of determining a PSF, it is necessary to have the proper PSF whenever new lenses are added to your microscope. The deconvolution software adapts to the PSF (actually the OTF) wavelength, so it is unnecessary to measure the PSF at more than one wavelength.

■ Note If you do not have the tools necessary to acquire a PSF, *softWoRx* includes a utility that allows you to calculate a theoretical OTF based on the numerical aperture of the camera lens, index of refraction, and emission wavelength. If the aperture of the lens is lower than 0.75 N.A., the calculated OTF may work as well, or even better, than a measured OTF. However, if the aperture is greater than 0.75 N.A., a measured OTF will generally give you better results. For information about calculating an OTF, see the *softWoRx* online Help.

A well-measured PSF is a key to successful deconvolution. For this reason, make sure that you:

- Thoroughly check all imaging conditions.
- Take the time you need to get a good signal-to-noise ratio in the image.
- Find a bead that is completely isolated from others in XYZ. (Use the field stop aperture to block fluorescence if needed.)
- Completely scan the bead.

# Tools

This procedure requires the following tools:

- A clean and aligned DeltaVision system
- A bead slide with 0.1µm, or smaller, fluorescent beads
- A grid slide or other microscopic ruler
- An immersion oil set, if appropriate (see *Selecting the Correct Oil* on Page 259.)

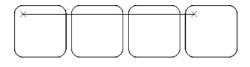
The following steps describe how to calculate pixel size, measure the PSF, and obtain the corresponding OTF.

#### To calculate pixel size:

- 1. Place the new objective in the Objective Turret and set the maximum image size, best camera speed, and auxiliary magnification slider at 1X.
- 2. Use the Eyepiece Filter Wheel to select the FITC eyepiece filter.
- 3. In the Resolve3D window, select the following values:

In this Field	Select
Excitation	TRITC
Emission	FITC
%T	2%

- 4. Place the silicon target grid on the stage and focus it (9.995 μm/square). Align the grid image to the vertical and horizontal axis and maximize the image.
- 5. Switch the Port Selector to **Camera** and click **Acquire**. Leave Data Collection Window 21 open.
- 6. In the Image window, choose **Tools** | **Measure Distances**. Then set the **Units** to **Pixels** in the Measure Distance window.
- 7. Draw a line across the grid slide image from a point on the top left square to a point in the same relative position on the top right square.



- Note Your measurement will be more accurate if you choose points on the vertices of the squares.
- 8. If the vertical delta is more than four pixels, re-align the slide. Repeat this process at the middle and bottom. Then count and record the number of grid elements (3 in the above image) and record the distance in pixels.
- 9. Repeat Steps 7 and 8 in the vertical direction.
- 10. Calculate the pixel size for each of the six measurements (top, middle, bottom, left, center, and right) as follows:

Pixel Size( $\mu$ m) =

9.995 µm/box

Measured distance (in pixels)/(Number of grids per measurement)

Your calculation should be accurate to four decimal places.

- 11. Average the six pixel sizes to obtain the correct pixel size.
- 12. Repeat these steps for each camera attached to your *DeltaVision* system.

#### To add an objective to the microscope configuration:

If you are adding a new objective to the microscope, follow these steps:

- 1. To obtain the objective lens ID number, select **Conversions** | **Convert PSF to OTF** from the *softWoRx* main menu. The PSF to OTF Conversion window is displayed. The current lens identification number is shown in the **Lens ID** field.
- 2. From the *softWoRx* main menu, choose **Utilities** | **Revise Microscope Configuration**. Then enter the root password to open the RESOLVE3D.SYS file.
- 3. In the RESOLVE3D.SYS file, under the Microscope Specifications section:

- a. Increase the number of lenses next to MS\_Number\_Lenses: by 1.
- b. Add the name of the objective to MS\_Lens\_Names: (e.g., 100Xoil, 60Xwater).
- c. Add the lens ID to MS\_Lens\_ID\_Numbers:
- d. Enter the pixel size for the new lens.

```
For example:
If the desired lens is 40X/1.35 with ID=10403 (the third lens in the list), then the pixel size is 0.1656.
```

```
MS_Lens_Names: 10X 20X 40X 60X 100X
MS_Pixel_Size_1: 0.6680 0.3313 0.1656 0.1103 0.06631 (for Coolsnap HQ2)
MS_Pixel_Size_2: 0.01 0.01 0.01 0.01 0.01 (for Evolve EMCCD - conventional mode)
MS_Pixel_Size_3: 0.01 0.01 0.01 0.01 0.01 (for Evolve EMCCD - EM mode)
MS_Lens ID_Numbers: 10105 10205 10403 10602 10002
```

4. To apply the new information, save and close Resolve3D.sys, then close and restart *softWoRx*.

#### To acquire a PSF:

It is easiest to find beads in a very dark room. Bead slides from Applied Precision include  $1\mu m$  beads and  $0.1\mu m$  beads. Both fluoresce brightly at 617 nm. Coarsely focus on the slide by positioning the lens near the slide. Scan the slide while looking for fluorescent haze from the  $1\mu m$  beads. When you focus on the fluorescence haze from the  $1\mu m$  beads you should also find the  $0.1\mu m$  beads.

Note Although a replacement bead slide is included in the Slide kit, bead slides have a limited shelf life. To purchase bead slides from Applied Precision, contact us at the appropriate number or address listed in *Chapter 1: Getting Started*.

- 1. Mount a bead slide on the microscope and focus on the beads to obtain the maximum intensity. Find a bead that is located by itself.
- 2. Use the **Center Object** tool to center a single bead in the X and Y directions. (It is helpful to collect large images, such as 1024×1024.)
- 3. Now use a 256×256 image.
- 4. Adjust the CCD exposure time so that the maximum intensity at the plane of best focus is at least 2000 counts. Make sure that the camera does not saturate at the plane of best focus.

- 5. Ensure there is only one bead in the field of view and that, as you go out of focus, no rings from other nearby beads enter the image. If necessary, use the field stop aperture to block out undesired fluorescence.
- 6. Verify that your microscope and software are accurately configured for lens and auxiliary magnification.
- Execute the Standard PSF Measurement Macro described in the online Help to measure the standard point spread function (or run a Z series through the bead consisting of 128 sections acquired in 0.1 µm Z increments).
- 8. Run the *softWoRx* PSF to OTF program that converts the optical sections into an OTF. (Refer to *Converting PSF to OTF* later in this appendix.)

# Selecting the Correct Immersion Oil

Accurate PSF measurements depend on the selection of the correct immersion oil. Our experience has shown that the oils recommended by microscope manufacturers are often not ideal for 3-D microscopy. We recommend that PSFs are measured with a minimal amount of spherical aberration. Inappropriate immersion oils yield asymmetric PSF measurements as a result of spherical aberration. In the case of Olympus microscopes, an index of refraction equal to 1.518 is ideal for measuring beads that are mounted in glycerol using #1.5 coverslips. There are many variables that can affect the selection of the correct immersion oil. The *softWoRx* Lens Information program can help you select the proper oil.

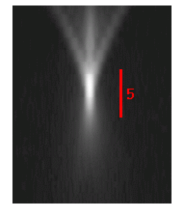
Note An oil kit is included with your *Delta Vision* system. To purchase replacement oil, please contact Applied Precision.

#### To confirm that you are selecting the correct immersion oil:

- 1. Collect the image data. Then, in the Image window, select **Tools** | **Orthogonal Viewer**.
- 2. Move the blue crosshair to the center of the bead in the viewer. To better see the shape of the PSF, it is helpful to do an exponential scaling—an exponent of 0.3 usually works well.
- 3. Look for symmetric flare in the resulting image. Symmetry indicates that the oil is correct, and in virtually all situations, the most symmetric PSF along the Z axis is also the smallest and has the highest intensity. In other words, symmetry corresponds with the highest resolution.
- 4. Repeat the process with different oils until you determine the optimal immersion oil.

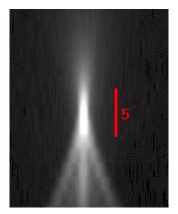
different immersion oils.

Correct Immersion Oil



The following figure demonstrates how image flare can be affected by the use of

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Immersion Oil Index Too Low

Immersion Oil Index Too High

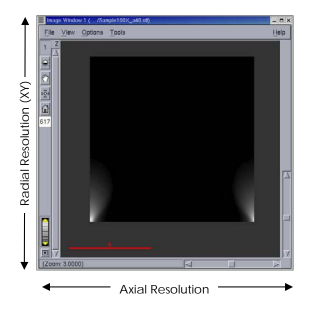
Flare from Immersion Oils (Orthogonal Views)

# **Converting PSF to OTF**

The PSF to OTF program converts a measured point spread function (PSF) to an optical transfer function (OTF). Essentially, the OTF is the Fourier transform of the PSF. The pixel size of the resulting image is given in cycles/ $\mu$ m. To reduce problems associated with measurement noise, the PSF is radially averaged during the conversion and, as a result, the 3D PSF image becomes a 2D OTF image.

The horizontal axis of the OTF represents axial (Z) frequency and the vertical axis represents radial (XY) frequency. The brightness of the OTF image elements, on a scale of 0 to 1, represents the frequency response of the microscope system at the corresponding radial and axial frequencies.

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#### Sample OTF Image

✓ PSF to OTF Conversion						/ - ×
PSF File						
OTF File						
X Range:	Start: [1	ļ.	End:	[1	-	
Y Range:	Start: 1	l.	End:	[1	Ī	
Z Range:	Start: 1	l.	End:	[1	Ī	
T Range:	Start: [1		End:	<b>[</b> 1	Increment 1	
Wavelengths:			Ĭ			
Sub-Image:						
Center: 0 0 0	Si	ze: [0	0 0			
Lens ID: 0						
Additional Parameters	;					
Done Do It Opti	ons					Help

#### **PSF to OTF Conversion**

Each option in PSF to OTF Conversion is described briefly below. For additional information regarding these options, refer to the online Help.

#### **PSF File**

Defines the name of the PSF image file to be converted to an OTF.

#### **OTF File**

Displays the name of the resulting axially symmetric OTF. (For your convenience, the OTF filename is created by appending "\_otf" to the PSF filename.)

#### X Range

Defines the start and end pixel numbers in X.

#### Y Range

Defines the start and end pixel numbers in Y.

#### Z Range

Defines the start and end pixel numbers in Z.

#### T Range

This field is not used for PSF to OTF conversion.

#### Wavelengths

Determined by PSF wavelength.

**Lens ID** Specifies the lens identification number (e.g., 12004).

#### Sub-Image: Center

Specifies the central XYZ coordinates of the point spread.

#### Sub-Image: Size

Specifies the XYZ image dimensions about the central coordinates. (The standard *softWoRx* point spread measurement is 256×256×128.)

#### Additional Parameters: Border Rolloff (voxels)

Specifies the number of voxels to roll off at the edge of the image. This reduces edge effects resulting from the Fourier Transform used in the PSF to OTF conversion.

The procedure for converting a PSF to an OTF is very simple. After the PSF file has been identified in PSF to OTF Conversion, *softWoRx* assigns default settings to the rest of the options in the window. In almost every instance, these settings will be appropriate to use for the conversion.

#### To convert a PSF to an OTF:

- 1. Click **Conversions** on the main menu bar of *softWoRx*.
- 2. Click **Convert PSF to OTF** in the **Conversions** menu. PSF to OTF Conversions will appear.
- 3. Use one of the following options to enter the PSF file to convert.
  - Drag the appropriate PSF file from the File Manager into the **PSF File** text box.
  - Click **PSF File** to display a small version of the File Manager and then choose the PSF file that you wish to convert.

- Type the desired path and filename into the **PSF File** text box.
- 4. Click Do It.

#### To place OTF into OTF Library

If the objective used is in addition to those already present, you'll need to modify *softWoRx* to use the new objective by adding the file to /usr/local/softWoRx/config/system.swrc as follows:

- 1. Log in to Linux as root.
- 2. Navigate to /usr/local/softWoRx/config/system.swrc
- 3. Find the section labeled, "Lens-to-OTF matching" and follow the instructions provided for the OTF file.

The following is an example of this section of the file:

# Lens-to-OTF matching. These are of the form LENS\_<lensIDNumber>\_OTF
# and are defined to be the file name in the OTF directory of the OTF that is
# to be used for this lens ID.

LENS\_12\_OTF LENS\_10602\_OTF LENS\_10003\_OTF LENS\_10403\_OTF LENS\_10603\_OTF LENS\_10205\_OTF

60X140\_sample.otf 60X140.otf 100X135.otf 40X135\_sample.otf 60Xw\_120.otf 20X0.75c.otf

# D

# **Appendix D: Reference Information**

The appendix includes the following topics:

- Standard Filename Extensions lists the filename conventions used by *DeltaVision*.
- Standard Fluorescence Filters shows the excitation and emission peaks of the standard filters for both fixed and live cell experiments included with DeltaVision.
- *Reference List* includes references for microscopy, Linux, image processing, optics, microscopy, and sample preparation.

# **Standard Filename Extensions**

The following is a list of filename conventions used by DeltaVision.

Type of File
Standard <i>DeltaVision</i> image
Optical Transfer Function
Resolve3D image
Deconvolved image
Volume Rendered image

# Standard Fluorescence Filters

Depending on which broadband light source the system uses, the *DeltaVision* imaging system provides a variety of different filter sets for both fixed and live cell imaging, a polarizer, and multiple sets of optional filters. These filters are designed to be used with many common fluorescent probes. If you are using fluorescent probes that are not well matched with the standard *DeltaVision* filters, contact Applied Precision for assistance.

# InsightSSI Module

The 4-color Fixed InsightSSI includes four of the most common sets of filters used for fixed cell imaging.

Filter Name	Excitation	Emission	Appropriate Probes
DAPI	UV, 381-401nm	Blue, 409-456nm	DAPI, Hoechst, Coumarin, Alexa 350 <sup>®</sup> , Alexa 405 <sup>®</sup>
FITC	Blue Green, 464- 492nm	Green, 500-523nm	Fluorescein, GFP, Cy3, Alexa 488 <sup>®</sup>
TRITC	Yellow Green, 531- 556nm	Yellow, 564-611nm	Rhodamine, Texas Red, Phycoerythrin, Alexa 568 <sup>®</sup> , Alexa 594 <sup>®</sup>
Су5	Red, 619-644nm	Infrared, 652-700nm	Cy5, Alexa 647 <sup>®</sup>

#### InsightSSI 4-color Fixed

The 4-color Live Cell InsightSSI includes four of the most common sets of filters used for live cell imaging.

#### InsightSSI 4-color Live Cell

Filter Name	Excitation	Emission	Appropriate Probes
CFP	Blue, 400-454nm	Blue Green, 463- 487nm	Cyan FP (CFP)
GFP	Blue Green, 425- 495nm	Green, 500-550nm	E-GFP
YFP	Green, 496-528nm	Yellow Green, 537- 559nm	Yellow FP (YFP)
mCherry	Yellow, 555-590nm	Orange, 600-675nm	mCherry, tdTomato, mRFP, DsRed

The 7-color Live Cell InsightSSI includes seven of the most common sets of filters used for fixed *and* live cell imaging.

InsightSSI 7-color	(for Fixed and Live	Cell Imaging)
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Filter Name	Excitation	Emission	Appropriate Probes
DAPI	UV, 381-401nm	Blue, 409-456nm	DAPI
CFP	P Blue, 400-454nm Blue Gre 487nm		CFP
FITC-GFP	Blue Green, 425- 495nm	Green, 500- 550nm	Fluorescein, E-GFP, Alexa 488 <sup>®</sup> , Cy3
YFP	Green, 496-528nm	Yellow Green, 537-559nm	Yellow FP (YFP)
TRITC	Yellow Green 531- 556nm	Yellow 564- 611nm	Texas Red, Rhodamine
mCherry	Yellow 555-590nm	Orange 600- 675nm	mCherry, tdTomato, mRFP, DsRed
Су5	Red 619-644nm	Infrared 652-700	Cy5, Alexa 647 <sup>®</sup>

# Xenon Arc Lamp

The excitation and emission peaks of the *DeltaVision* filters are provided in the following table.

#### Xenon Standard Fixed Cell Filter Set

Filter Name	Excitation	Emission	Appropriate Probes
DAPI	UV, 325-375nm	Blue, 438-478nm	DAPI, Hoechst, Coumarin, Alexa 350 <sup>®</sup> , Alexa 405 <sup>®</sup>
FITC	Blue Green, 481- 502nm	Green, 506- 543nm	Fluorescein, GFP, Cy3, Alexa 488 <sup>®</sup>
TRITC	Green, 547-563nm	Yellow/Orange, 576-630nm	Rhodamine, Texas Red, Phycoerythrin, Alexa 568 <sup>®</sup> , Alexa 594 <sup>®</sup>
Cy5®	Red, 636-656nm	Infrared, 667- 719nm	Cy5, Alexa 647®

The optional Xenon Live Cell filter wheel module includes four of the most common sets of filters used for live cell imaging.

Filter Appropriate Name Excitation Emission Probes CFP Blue, 415-445nm Blue Green, 455-Cyan FP (CFP) 485nm YFP Green, 490-510nm Yellow Green, 520-Yellow FP (YFP) 550nm mCherry Yellow, 555-590nm Orange, 600-675nm DsRed, tdTomato, mCherry, mRFP GFP Blue Green, 425-Green 500-530nm E-GFP 495nm

Xenon Live Cell Filter Set

# Additional Reading Material

Material for further reading is available on the following pages. Contact Applied Precision for the most recent list. If you notice omissions from the list, please inform Applied Precision.

### Microscopy

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# E

# Appendix E: Resolve3D and Keypad Options

This chapter describes the following:

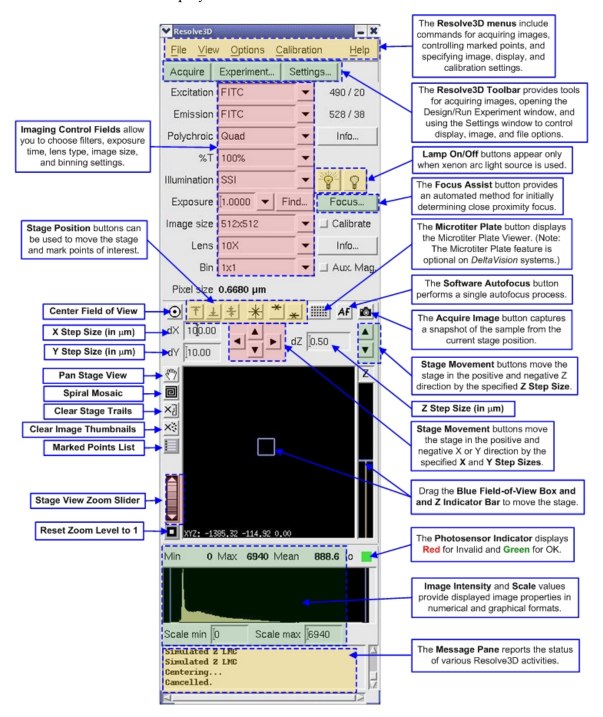
- *The Resolve3D Window* is the main window for data acquisition.
- The Design/Run Experiment Window provides tools to select, design, edit, and execute experiment macros.
- *The Settings Window* used to control how images are displayed, select camera settings, and specify file output.
- *Keypad/Joystick Operation* is a reference for the buttons on the keypad. Many of the Resolve3D functions are also available on the keypad and joystick.

# The Resolve3D Window

The Resolve3D window is the main data acquisition window. In addition to providing many of the acquisition options and controls, it provides access to the other windows that are used for data acquisition.

#### To open Resolve3D:

► From the *softWoRx* menu, choose **File** | **Acquire** (**Resolve3D**). The Resolve3D window is displayed.



# The Resolve3D Menu

The Resolve3D menu has the following menu items:

- The File menu includes commands to acquire images and to open the key windows for setting up and running experiments.
- The View menu includes commands to manage marked points and to create a blank screen.
- The Options menu includes commands to open the Settings window (where you can set display and image options) and to save settings.
- The Calibration menu opens the Calibration tool.
- **The Help menu** provides options to show or hide ToolTips and to get Help.

#### The File Menu

Use the following Resolve3D File menu commands to acquire images, create scratch files, and open the Design/Run Experiment window.

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	Acquire Image		F1		
	Continuous Ac	quire			
	S <u>n</u> apshot				
	Scratch File				
	Experiment		Ctrl+E		
	<u>Q</u> uit		Ctrl+Q		
L					

The Resolve3D File Menu

#### Acquire Image

Collects and displays an image from the microscope. This image is only displayed in the Data Collection window. It is not saved to a disk file for later use.

#### **Continuous Acquire**

Opens the Continuous Acquire window that you can use to collect and display images continuously. These images are only displayed; they are not saved to a disk file.

#### Snapshot

Launches a tool to let you collect a single 2-D, multi-wavelength "snapshot" image. If you need to collect a Z series, time-lapse image, or other complex scheme, you

will need to design an experiment with the Resolve3D Experiment Designer. You can also do an OAI as part of your snapshot.

#### Scratch File

Creates a "scratch file" to which you can save individual image frames for later use. After a file is opened, clicking **Save Current Image** saves the most recently collected image frame. Clicking **Close Scratch File** or **Done** closes the file. Note that the image size cannot be changed while a scratch file is open.

#### Experiment

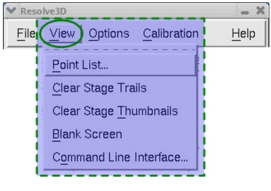
Opens the Design/Run Experiment window. You can use Design/Run Experiment to select a previously created experiment, to design a new experiment, or to open the Experiment Macro Editor to create or edit an experiment macro.

#### Quit

Closes the Resolve3D window.

#### The Resolve3D View Menu

Use the View menu to manage points that you have marked, to clear the history of the path that you took while exploring your sample, or to create a black screen for light-sensitive conditions.



The Resolve3D View Menu

#### Point List

Opens the Points List window. This window helps you manage a list of points (with X, Y, and Z coordinates) that you want the system to remember. These points can be interactively "visited" at any time and they can be used in experiments.

#### **Clear Stage Trails**

Clears the Stage Trails history. (The system maintains a history of the paths of motion that you take while exploring your sample. These paths are displayed as "Stage Trails" on the Stage View.)

#### **Clear Stage Thumbnails**

Clears all of the thumbnail images currently displayed on the Stage View.

#### **Blank Screen**

Turns the computer screen blank. This is useful when you are imaging under very light-sensitive conditions. Clicking anywhere on the screen restores it.

#### Command Line Interface

Opens the Command Line window that provides advanced users with the ability to issue individual Resolve3D commands to the system.

**CAUTION:** The Command Line Interface should be used carefully because it can put the system in an unstable state.

#### The Resolve3D Options Menu

Use the Options menu to open the Settings window (where you can set display and image options) and to save configuration settings and state information.

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	Zero the Stage Z	
	Center the Stage XY	
	<u>S</u> ettings	
	S <u>a</u> ve Settings	

The Resolve3D Options Menu

#### Zero the Stage Z

Selecting **Zero the Stage Z** from the Options menu provides a protocol for moving the stage in the Z direction to the middle of its working range.

#### Center the Stage XY

Selecting the Center the Stage XY from the Options menu provides a protocol for moving the stage in the XY direction to the middle of its working range.

#### Settings

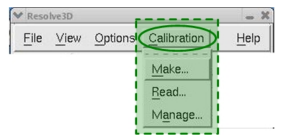
Opens the Settings window which allows you to control display, imaging, and file output options.

#### Save Settings

Saves the configuration settings and information (current filters, image size, etc.) to be used the next time Resolve3D is opened.

#### The Resolve3D Calibration Menu

Use the Calibration menu to make calibration tables, read calibration tables, and designate which calibration tables are active.



The Resolve3D Calibration Menu

#### Make

Opens the Calibration tool used to create flat-field calibration tables and optional tables of "bad" pixels. These tables may be applied to images, either when they are acquired or at a later time. (To apply the tables to images after they are acquired, use the Calibrate utility available from the Process item on the main *softWoRx* menu.)

#### Read

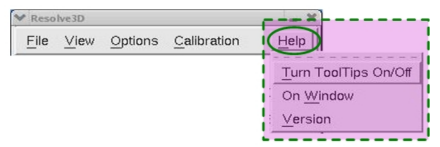
Opens the Read Calibration Files window that you can use to read calibration tables.

#### Manage

Opens the Manage tool that you can use to designate which calibration tables are active and to remove tables from the system's session memory.

### The Resolve3D Help Menu

Use this menu to turn ToolTips on or off, get help on the Resolve3D window, or find out which version of *softWoRx* you are using.



#### The Resolve3D Help Menu

#### Turn ToolTips On/Off

Turns tool tips on or off. Tool tips display "pop-up" information about buttons on the interface. They open when the mouse pointer is held over a button for a few seconds.

#### On Window

Opens Help for the Resolve3D window.

#### Version

Displays version numbers for the *softWoRx* components.

# The Resolve3D Toolbar

Use the buttons on the Resolve3D toolbar to acquire images, open the Experiment Designer/Run window that allows you to set up and run experiments, and open the Settings window that allows you to control display, imaging, and file output options.

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Acqu	uire	E	xperiment.		Settings			

The Resolve3D Toolbar

#### Acquire

Collects and displays an image from the microscope with the current settings. This image is only displayed in the Data Collection window. It is not saved in a file.

#### Experiment

Opens the Design/Run Experiment window that you can use to select a previously created experiment, to design a new experiment, or to open the Experiment Macro Editor to create or edit an experiment macro.

#### Settings

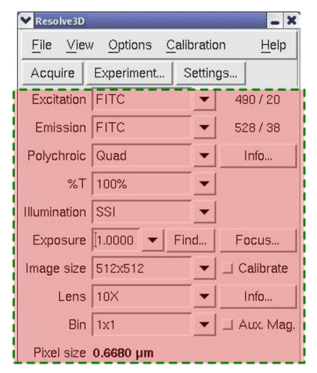
Opens the Settings window that allows you to control display, imaging, and file output options.

### Image Control Fields

Use the Resolve3D Image Control fields to:

- Select the Excitation, Emission, and Neutral Density filters.
- Select exposure time.
- Determine whether to calibrate the image.
- Select the desired shutter.
- Select the image size.
- Select the lens and get lens information, including the oil calculation.
- Select whether auxiliary magnification is used.
- Determine binning parameters.
- Get pixel size.

Select the polychroic.



The Resolve3D Image Control Fields

#### Excitation

Specifies an excitation channel. When this field is changed, the following parameters are set to the last values that were used for that excitation channel.

- Emission filter
- Exposure time
- Neutral Density value
- Active illumination shutters
- Number of frames to average
- Target intensity
- Camera gain

You can choose **Save Settings** from the Options menu to store the configuration. The wavelength/bandwidth of the selected filter is indicated to the right of the filter choice box.

#### Emission

Specifies an emission filter. The wavelength/bandwidth of the selected filter is indicated to the right of the filter choice box.

#### Polychroic

Specifies a polychroic to place into the active position. The selections available are displayed in the drop-down list in this field. Your choices will completely depend upon which polychroic filters are installed on your system.

#### Illumination

Indicates which light source will be used. Your choices are as follows:

- For broadband illumination, select SSI.
- For transmitted light, select **TRANS**.
- For laser light, select LASER.

#### %T

Specifies a Neutral Density value. The relative illumination intensity is indicated in the menu.

#### Exposure

Specifies camera exposure time (in seconds). The minimum and maximum exposure times allowed for this field depend upon the camera type.

#### Find

Opens the Target Intensity window which allows you to find the exposure time necessary to reach a Max intensity near the Target Intensity Value. For dim samples, 300-900 counts are acceptable. For bright samples, 200-2500 counts provide a high dynamic range without saturation. This tool is not recommended for use with live cells.

#### Calibrate

Calibrates images when they are collected from the camera. If you use this option, you must load and activate a calibration table that fits the current imaging parameters (wavelength, image size, objective, bin choice) before you acquire images.

#### **Image Size**

Specifies image size (pixels or CCD detector elements) for acquired images. The pull-down list contains predefined sizes for convenience. You can enter special sizes by choosing **Other...** from the pull-down list. (Image size must be a multiple of four.)

#### Lens

Specifies the objective lens name. The pull-down list contains the lenses that are known to be part of the microscope system.

#### Info

Opens the Lens Information window which displays information about the current objective lens, including the oil calculator.

#### Bin

Specifies the number of CCD detector elements to add together to form one image element. Binning is applied in both the X and Y directions. It increases intensity, but it decreases resolution. Pixel size is a function of binning.

#### Aux. Mag.

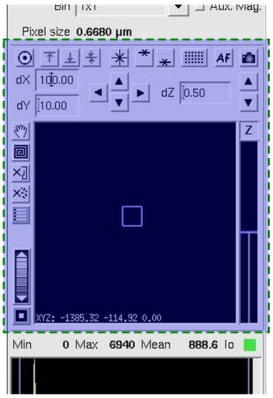
Specifies whether the microscope's 1.6x manual auxiliary magnification is in use. (On IX70 stands, the manual auxiliary magnification is 1.5x. For EMCCD camera systems, the auxiliary magnification is 2.0x.)

#### **Pixel size**

Displays pixel dimensions in microns/pixel. The value is calculated from the Lens, Bin, and Auxiliary magnification settings.

# **Stage Position Control Fields and Buttons**

The Resolve3D Stage Position Control area allows you to control the microscope's X, Y, and Z stage positions. The Center, Mark, and Visit tools and the Stage Motion Controls are shown below.



The Resolve3D Stage Position Controls

## Stage Control and Display Tools



#### Center Object

Centers the stage on an object selected in an Image Window. The pixel size must be correct in order for object centering to work properly, which means that the correct lens and auxiliary magnification setting must be selected. An image will be acquired after the object is centered.



#### Mark Point

Marks the current X, Y, and Z stage coordinates as points to be visited later.



#### Mark Top of Sample

Marks the current stage Z position as the "top" of your sample. Because the scanning process always moves the stage toward the objective, this position also represents the point where the stage is closest to the objective (the most negative Z value). This also represents the focal plane that is the closest to the slide side of a sample. Use this along with the **Mark Bottom of Sample** button to establish the thickness of the sample. You can use these marked positions to aid with the Z sectioning setup.

■ Note All scans that are set up using the Experiment Designer scan in Z, using relative coordinates. The Mark Top of Sample and Mark Bottom of Sample buttons are most helpful in determining the thickness of the sample to be scanned. When an experiment is started, the scan region is determined by the current focus point and the thickness of the sample. So, for example, if you have marked three points to visit in an experiment and they all have different middle-Z positions, the experiment will calculate the scan based on these different Z positions and the fixed thickness.

#### Visit Top

Moves the stage to the position marked as the top of the sample.

×

#### Mark Bottom of Sample (end of scan)

Marks the current stage Z position as the "bottom" of your sample for a potential scan. This corresponds to the positive stage Z coordinate value or the coverslip side of the sample.



#### Visit Bottom

Moves the stage to the position marked as the bottom of the sample.



#### Visit Middle

Visits the point in the middle of the defined scan region.



#### **Plate Viewer**

Displays the Microtiter Plate Viewer window. You must have the microtiter stage option installed on your *DeltaVision* system in order to scan microtiter plates.



#### Autofocus

Automatically focuses using a contrast-based software method.



#### Acquire Image

Collects and displays an image from the microscope. The current Resolve3D image control settings are used to collect the image. This image is only displayed in the Data Collection window. It is not saved in a file for later use.

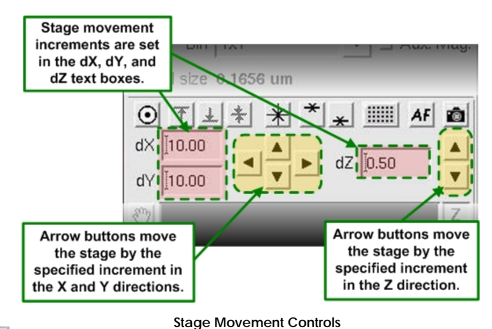


#### XY Stage Controls

Moves the stage in the X and Y axis. The left and right arrows move the stage in the X axis in the increment set in the dX field and the up and down arrows move it in the Y direction in the increment set in the dY field.

#### Z stage Motion Controls

Moves the stage in the Z axis in the increment set in the **dZ** field.





#### Pan

Places the stage view in Pan mode. Click and drag the current view to pan it. Note that this tool is "sticky." To disable Pan mode, click the **Pan** tool again.

#### **Spiral Mosaic**

Starts a preview collection pattern centered on the current stage location and using the active Resolve3D wavelengths. The pattern begins with the current stage location and then continues acquiring thumbnail images directly adjacent to it, spiraling outward in a counter-clockwise rotation so that the entire area centered around the initial stage position is previewed in the Resolve3D stage view. Thumbnail collection continues until either the preset spiral mosaic size (set in **Resolve3D** | **Settings** | **Misc** tab) is reached or you click the **Spiral Mosaic** button again. The stage is always returned to the initial position regardless of how the collection ends.

# ×J

#### **Clear Stage Trails**

Clears all of the stage trail lines from the stage view.



#### **Clear Thumbnails**

Clears all of the thumbnail images currently displayed on the stage view.

## **Marked Points list**

Opens the Point List window to manage the list of marked points.

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7

# Zoom Tool

Controls zoom of the stage view. Drag the thumbwheel down to zoom in; up to zoom out. The button under the thumbwheel resets the zoom to 1:1.

# Z Slider

Moves the stage up or down. Click and drag the blue horizontal bar to move a maximum of  $\pm$  5  $\mu m.$ 

# dX

Specifies the X step size, in microns.

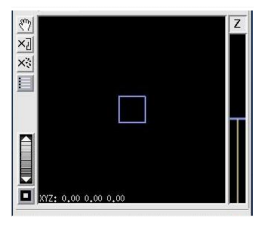
# dY

Specifies the Y step size, in microns.

# dZ

Specifies the Z step size, in microns.

# **Stage Trails Window**



The blue box represents the current stage location. Drag the box to move the stage in X and Y.

# **Image Intensity and Scale Values**

The Resolve3D image intensity and scale show the intensity values of the data collection window in numerical and graphical formats. The scale of the image is also shown.

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The Resolve3D Image and Intensity and Scale View

## Min, Max, Mean

Displays the minimum, maximum, and mean intensity values of the most recently acquired image. For a 12 bit CCD camera, these values range between 0 and 4095. A value of 4095 indicates camera saturation, unless image calibration is in effect. (If you are using 0.5X Gain, the saturation is less than 4095 counts.)

## Histogram

Shows the intensity distribution for the most recently acquired image. The vertical blue bars indicate the **Scale min** and **Scale max** and can be dragged interactively with the mouse to change how the data collection window display is scaled. The X-axis represents intensity and the Y-axis represents the number of pixels.

## Scale min/Scale max

Specifies the settings for the minimum and maximum display values. These numbers can be changed manually or by moving the histogram threshold bars.

### lo

Indicates valid (or invalid) photo sensor values or saturation. The indicator is green if the photo sensor value is valid. It is red before the first image is acquired. After the first scan, a red color indicates that the most recently acquired image has an invalid photo sensor value. A red indicator can also signify unreasonable saturation or an improperly functioning photo sensor device.

# The Message Pane

The Resolve3D Message pane reports the status of various Resolve3D activities. Use the scroll-bar to view messages that have scrolled off the top of the pane.

# **Resolve3D Shortcuts**

You can right-click anywhere in the Resolve3D window to open a shortcut menu that allows you to acquire images, mark points, and create a blank screen (for imaging under very light-sensitive conditions).

Acquire
Snapshot
Mark Point
Blank Screen

## **Resolve3D Shortcut Menu**

## Acquire

Collects and displays an image from the microscope. The current settings are used to collect the image. This image is only displayed in the Data Collection window. It is not saved in a file for later use.

# Snapshot

Launches a tool to let you collect a single 2-D, multi-wavelength "Snapshot" image. (The current settings are used to collect the image.)

# Mark point

Marks the current X, Y, and Z stage coordinates as a point to be visited later.

# **Blank Screen**

Turns the computer screen black for imaging under very light-sensitive conditions. Clicking anywhere on the screen restores it.

# The Design/Run Experiment Window

The Design/Run Experiment window provides tools to select, design, edit, and execute experiment macros. (Experiment macros are "scripts" of commands that guide the *DeltaVision* system to collect images.)

# To open the Design/Run Experiment window:

From the Resolve3D window, click Experiment and click the Run Experiment tab.

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# The Design Experiment Tab

The **Design Experiment** tab is used to generate experiment command macros.

To open the Design Experiment tab:

► From the Resolve3D window choose **Experiment**. The Design/Run Experiment window opens with the **Design Experiment** tab selected.

Note The Design PK Experiment tab is available only for systems equipped with a laser module.

Design Experiment	Design PK Experiment	Run Experiment
Experiment name	Resolve3Dí	
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ectioning * Chai	nnels   Time-lapse   Poir	nts Panels Plate Actions
Z Sectioning	Z Scan Options	
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**Design/Run Experiment Window** 

### **Experiment name**

Specifies the name of the experiment macro.

## **Enable Fast Acquisition**

Enables fast acquisition experiments (see Page 294).

### Sectioning Tab

Specifies sectioning for 3D images (see Page 294).

### **Channels** Tab

Specifies channels and exposure time (see Page 296).

## Time-lapse Tab

Specifies criteria for time-lapse experiments (see Page 297).

## **Points Tab**

Specifies a list of marked points (see Page 300) and options for UltimateFocus and Autofocus.

#### **Panels Tab**

Allows you to set up panel collection that you can use to acquire a large area of a slide to stitch together to form a single image (see Page 302).

## Plate Tab

Provides tools for setting up and running microtiter plate scanning experiments when your *DeltaVision* system is equipped with the Microtiter Stage option.

## **Actions Tab**

Lets you designate new Laser, Autofocus, and UltimateFocus events, add Pause and Wait times, alter Time-lapse intervals and image status methods, and other specific actions to occur during the experiment.

# **Experiment Name and Enable Fast Acquisition**

## **Experiment name**

Specifies the name of the file that is generated by the Experiment Designer. A file extension of .exp will be added to the name.

## **Enable Fast Acquisition**

Enables fast image acquisition of 2D images. (This should not be used with cameras that require a shutter.)

This data acquisition mode should be used carefully. Fast Acquisition uses a single command to set up a data stream to the Instrument Controller.

There is an upper memory limit for data collection that is based on the size of RAM memory for the Instrument Controller (typically about 350-400 MB). If you run into this limitation, turning Fast Acquisition off allows you to collect larger data sets.

If the system is unstable after collecting a large data set with Fast Acquisition, restart the workstation or stop using Fast Acquisition unless it is strictly necessary. Sometimes Fast Acquisition isn't much faster than standard acquisition, and more is risked than gained by using it.

Note You can set Fast Image Acquisition options to control the scan sequence, the shutter open mode, the camera readout mode, and the starting Z location for the scan. (See the online Help for more information.)

# **Sectioning Setup**

Once your microscope is focused near the middle of the vertical zone of interest of your sample, you can use the parameters in the Design Experiment Sectioning tab to control the Optical Sectioning procedure.

**Note** The standard scan direction moves the objective lens towards the specimen.

	<i>Design PK Experiment</i> Resolve3Dį́	Run Experiment
, Estimated file size		
	32.01 Mb	(21.43 Gb Available)
Jse Fast Acquisition	Fast Acquisition Option	s   💷 Lamps Off when finished
ectioning * Chann	els   Time-lapse   Poin	ts Panels Plate Actions
Z Sectioning	Z Scan Options	
cus point when scan s	tarts Middle of Sample	•
Optical section spa	acing 0.20	
Number of optical sect	tions (64	
Sample thick	ness (12.80 Get th	nickness
	🗆 Enable OAI Scar	1

**Design Experiment Sectioning Setup Options** 

#### Focus point when scan starts

Specifies the Z location of the sample when the experiment starts. The recommended selection is **Middle of Sample**.

### **Optical section spacing**

Specifies the spacing (in microns) between each optical section. The focal point will be changed by this value after each image is collected.

### Number of optical sections

Specifies the number of sections to collect (for each wavelength) for the experiment. *softWoRx* automatically calculates and displays this value if the **Sample thickness** and **Optical section spacing** are entered. If you specify this value, the **Sample thickness** value is changed.

#### Sample thickness

Displays the **Number of optical sections** multiplied by the **Optical section spacing**.

## **Get Thickness**

Retrieves the Sample thickness, based on the locations defined with the  $\mathbf{T}$  and  $\mathbf{I}$  buttons in the Resolve3D window.

## OAI

Acquires a 2D Z projection of the interval defined by the marked top and bottom of the sample. The optical section spacing and the number of optical sections are determined automatically, based on the depth-of-field of the objective and the exposure time.

# **Channels Setup**

Use the Design Experiment Channels tab to select wavelengths (filters) and to specify an exposure time for each filter.

Design Ex	periment	Desig	ın PK Experin	nent	Run Exp	erii	ment	
Expe	riment name	Reso	lve3Dį́					
Estim	ated file size	1.50	db		(22	.28	Gb Availabl	e)
Use Fast	. Acquisition	Fas	st Acquisition	Optio	ns 🗍 🗆 l	am	ps Off when	finished
Sectionin	g * Chani	nels *	Time-lapse	Poi	nts Pane	Is	Actions	J.
Conventio	onal * Multi	plexed						
Exp	EX Filter		EM Filter		%Т		EX Shutter	
7 [1.000	DAPI	•	DAPI	•	100%	-	EX	-
7 1.000	FITC	•	FITC	-	100%	•	EX	•
1 [1.000	BLANK 1	*	BLOCK		BLOCK 10	*	EN	<b>_</b>
1.000	BLANK 1	*	BLOCK	*	BLOCK 10	*	EN	<b>.</b>
1.000 I	BLANK 1	*	BLOCK	*	BLOCK 10	*	EX.	<u>_</u>
7 Referen	ce Image	Z Pos	ition Middle o	f Sarr	ple		•	
Ĭ1.000	CY-5	1 10	CY-5		)0% –	ſГт	RANS -	1

# **Design Experiment Channels Options**

# **Refresh exposure conditions**

Updates the filter and exposure settings to those last used in the main Resolve3D window.

# Active Wavelength Toggle Buttons

Enable the exposure time, filters, and display settings for specific wavelengths. Select the buttons that activate the wavelengths that you want to collect. (You must select one button for each wavelength.) If no wavelengths are selected, the exposure and filters that are set in the Resolve3D window are used.

# Ехр

Specifies the exposure time (in seconds) to be used when acquiring an image for the selected wavelength. If left blank, the value specified in Resolve3D will be used when the experiment is run.

# **EX** Filter

Specifies the excitation filter to use for this experiment or image. When it is changed, the currently "paired" emission filter is automatically selected.

## **EM Filter**

Specifies the emission filter to use. This filter may be selected independently of the EX Filter setting.

# %T Filter

Specifies the Neutral Density value to use. The % value indicates light transmission. A value of 100 % indicates that no light is blocked.

## **Ex Source**

Specifies the excitation (illumination) source to use for each channel of the image.

## Polychroic

Specifies the polychroic filter to use for the experiment. If you have the optional motorized mirror turret installed, it will automatically move the selected polychroic into the light path. The polychroic is moved when the experiment is run.

**Note** This field is available only if the polychroic turret is motorized.

## **Reference Image**

Specifies to use an alternate filter or the transmitted light to acquire a reference image that can be combined with other images. This option is useful for Differential Interference Contrast (DIC) analysis. It can also be useful for other types of reference images. Only one reference image per Z stack is collected.

# **Time-lapse Setup**

Use the Design Experiment Time-lapse tab to specify the number of time points, the time periods, and the total time for a time-lapse experiment.

✓ Design/Run Experiment: * (modified)	- ×					
File	Help					
Design Experiment Design PK Experi	iment Run Experiment					
Experiment name Resolve3D						
Estimated file size 64.52 Mb	(20.59 Gb Available)					
Use Fast Acquisition Fast Acquisition	n Options					
Sectioning * Channels * Time-lapse	Points Panels Plate Actions					
✓ Time-lapse						
Hours Minutes Seconds Milliseconds						
Time-lapse 10 10 10						
Total Time 🚺 🚺 🚺						
Time Points 1						
F						
Enable Cell Tracking	Cell Tracking Options					
✓ Maintain Focus with UltimateFocus	UltimateFocus Options					
✓ Image-based Autofocus before imaging	Autofocus Options					

Design Experiment | Time-lapse Setup Options

# **Time-lapse option**

Specifies running a time-lapse experiment.

## **Time-lapse**

Specifies the hours, minutes, and seconds between each time period.

## **Total Time**

Displays the total time of the experiment, which can also be calculated as follows:

Total time = (The number of time points - 1) x time-lapse

Specifying a total time will update the **Time Points** field.

## **Time Points**

Specifies the number of time samples to collect in a time-lapse experiment. Changing this value will update the **Total Time** field.

## **Enable Cell Tracking**

Moves the stage laterally to follow cells as they move during a time-lapse experiment. With the **Enable Cell Tracking** option selected, *DeltaVision* automatically keeps cells in the field of view.

## **Cell Tracking Options**

Opens the Cell Tracking Options window that allows you to set the parameters for cell tracking. For more information on using the tools on this window, see Tracking Cells on Page 90.

Cell Tracking Options		/// ×
Tracking Method	Center of Intensity	•
Reference Channel	1	
Move Threshold (microns)	1.000	
ROI Percent	50	

## Maintain Focus with UltimateFocus

Specifies focus should be maintained during the time-lapse experiment. If this experiment does not involve point visiting, a calibration (characterizing the UltimateFocus response at the current location) is performed when the experiment is started.

Notes #1 The initial focus is essential to the success of focus maintenance.
 #2 The actual focus maintenance process happens just before each time-lapse event.

#### **UltimateFocus Options**

Opens the UltimateFocus Options window, in which you can indicate a specific **Move Threshold** and the **Maximum** (number of) **Iterations** to move within the threshold before re-running UltimateFocus.

♥ UltimateFocus Options	×
Move Threshold ( $\mu m$ )	Į0.100
Maximum Iterations	[]1
Done	Help

On the UltimateFocus Action Options window:

- In the **Move Threshold** field, enter the desired maximum measured focus error (in μm) before a corrective action should be taken.
- In the **Maximum Iterations** field, enter the maximum measure/move sequences for this action to reach the calibrated focus point.

**Note** More iterations will take longer, but may provide better focus.

• Click the **Done** button.

## Image-based Autofocus before imaging

Using the software-based Autofocus function, automatically focus the camera before each time point. For point visiting experiments, the camera is automatically focused every time that a point is visited.

## **Autofocus Options**

Opens the Resolve3D Settings window to the Autofocus tab, in which you can manipulate the settings for the software-based Autofocus function.

Display       Files       Imaging       Illumination       Autofocus       Lasers       Misc         Image Contrast Software Autofocus	Resolve3	D Settings					-
L Automatically determine parameters Test Channel for Autofocus DAPI Contrast calculation method Auto Autofocus Z test step (µm) [3.12 Maximum Z test range (µm) [50.00 Post-autofocus Z offset (µm) [0.00 UltimateFocus Hardware	Display	Files	Imaging	Illumination	Autofocus	Lasers	Misc
Contrast calculation method Auto Autofocus Z test step (µm) [3.12 Maximum Z test range (µm) [50.00 Post-autofocus Z offset (µm) [0.00 UltimateFocus Hardware	-				[	0.	
Autofocus Z test step (µm) [3.12 Maximum Z test range (µm) [50.00 Post-autofocus Z offset (µm) [0.00 UltimateFocus Hardware	Ch	annel for A	utofocus D.	API		•	
Maximum Z test range (µm) [50.00 Post-autofocus Z offset (µm) [0.00 - UltimateFocus Hardware	Contrast	calculation	n method A	uto		•	
Post-autofocus Z offset (µm) (0.00	Autofoc	us Z test s	step (µm) [3.	12			
- UltimateFocus Hardware	Maximur	m Z test ra	nge (µm) (50	).00			
	Post-auto	ofocus Z of	fset (µm) (0.	00			
Calibrate UltimateFocus for plate scanning	Ultimat	teFocus H	ardware				
	Calibrat	e UltimateF	Focus for pla	te scanning			
	Done	Save Settin	ngs				Hel

Resolve3D Settings Window | Autofocus Tab

# **Point Visiting Setup**

Use the Design Experiment Points tab to specify a list of marked points to visit during the experiment.

Tip Before you specify these points, make sure that there are active points in the point list.

Design/Run Experiment: *() File	nodified)			 Help
38 Ø 🕑				
Design Experiment	Design PK Experin	nent Rui	n Experiment	
Experiment name	Resolve3Dį			
Estimated file size	32.01 Mb		(20.59 Gb Available	)
⊔ Use Fast Acquisition	Fast Acquisition	Options	Lamps Off when fi	nished
Sectioning * Chan	nels   Time-lapse	Points	Panels Plate	Actions
✔ Visit Point List	yana			
✓ Maintain Focus with	JltimateFocus	UltimateFo	cus Options	
✓ Image-based Autofor	us before imaging	Autofocu	us Options	
	-			

Design Experiment | Point Visiting Setup Options

## Visit Point List

Specifies the list of points (described by number) to visit during the experiment. All sectioning and wavelength procedures are repeated at each of the listed points. A point list can be entered as a series of numbers separated by commas or dashes. Separating two numbers with a dash '-' (as shown in the following example) indicates that all point numbers in between should also be visited.

For example:

1,2,5,7-10

Note When you use Point Visiting with Z sectioning, the microscope uses the Z value of the current focus of each point as the initial reference for that point. Values specified for Z sectioning are incremented relative to that point.

### Maintain Focus with UltimateFocus

Specifies focus should be maintained during the point visiting experiment. A UltimateFocus calibration (characterizing the UltimateFocus response at the current location) is performed when the point is initially marked.

### **UltimateFocus Options**

Opens the UltimateFocus Options window, in which you can indicate a specific **Move Threshold** and the **Maximum** (number of) **Iterations** to move within the threshold before re-running UltimateFocus.

♥ UltimateFocus Options	×
Move Threshold (µm)	Į0.100
Maximum Iterations	<u>[</u> 1
Done	Help

On the UltimateFocus Action Options window:

- In the **Move Threshold** field, enter the desired maximum measured focus error (in μm) before a corrective action should be taken.
- In the **Maximum Iterations** field, enter the maximum measure/move sequences for this action to reach the calibrated focus point.

**Note** More iterations will take longer, but may provide better focus.

• Click the **Done** button.

## Image-based Autofocus before imaging

Using the software-based Autofocus function, automatically focus the camera at each visited point before acquiring an image.

## **Autofocus Options**

Opens the Resolve3D Settings window and the Autofocus tab, in which you can manipulate the settings for the software-based Autofocus function.

# **Panel Collection Setup**

Use the Design Experiment Panels tab to create panel collection macros. These macros are useful when you want to scan a large area with a relatively high magnification lens. You can use the panels as a means of reviewing a large area of a slide, or as data that you want to stitch together to form a single, large image.

Note Panel collection macros are sensitive to microscope settings such as image size and magnification. Because the number of panels required depends upon many factors, it is usually not a good idea to reuse panel collection macros.

File 월 🔛 🛷 (			<u>H</u> elp
Design Experiment Experiment na Estimated file s	me Resolve3E ize 64.52 Mb	Run Experimen (21.42 Gb	
Collect Panels	Overlap (pixels) 0.00 0.00		Plate Actions

Design Experiment | Panel Collection Setup Options

## **Collect Panels**

Specifies to use a panel collection macro.

## **Overlap** (pixels)

Specifies the amount of overlap (in pixels) between adjacent panels.

### Start Coordinates

Specifies the XYZ coordinates at which to start collecting panels. Use the **Get Start** button to obtain the current XYZ stage coordinates.

### **End Coordinates**

Specifies the XYZ coordinates at which to finish collecting panels. Use the **Get End** button to obtain the current XYZ stage coordinates.

Note These coordinates will be included at minimum. Based on CCD size, the final collection area may be larger than specified.

# **Design PK Experiment Tab**

The TIRF/PK Module provides the ability to add a laser beam into the optical path of the *DeltaVision* microscope. A laser beam is introduced into the back aperture of the microscope objective to provide a focused illumination spot in the center of the optical field. With lasers attached (like the *DeltaVision* X4 Laser Module), users can design photokinetic experiments and TIRF experiments. TIRF is an imaging method that offsets the laser excitation light slightly off-axis using a special TIRF- based objective. The excitation light emits from the objective at a shallow angle, enters into the cover slip, and bounces inside the cover slip creating an evanescent field of fluorescence. The TIRF/PK Module is optional for the *DeltaVision* system and is offered in three varieties: TIRF/PK, TIRF only, and PK only.

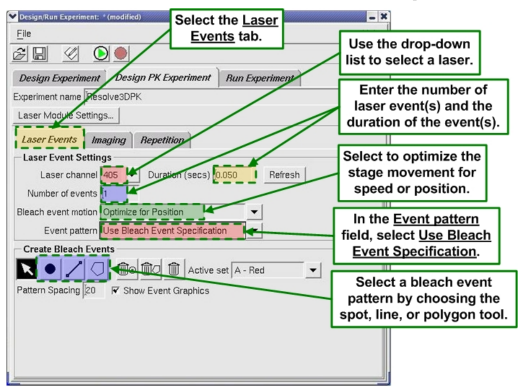
The **Design PK Experiment** tab provides options for designing the nature of a photokinetic event, setting up the event timing, setting image conditions to associate with the event, and defining the locations of photokinetic events.

# **Running Photokinetic (PK) Experiments**

The Resolve3D Design/Run Experiment window contains basic photokinetic experiments such as FRAP (fluorescence recovery after photobleaching).

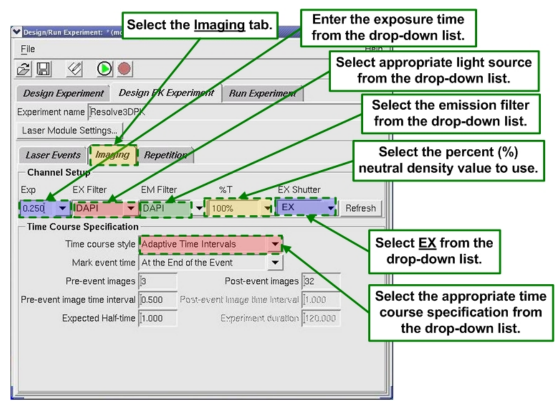
# To run a basic PK experiment:

- 8. Click the **Experiment** button in the Resolve3D main menu to open the Design/Run Experiment window.
- 9. Select the **Design PK Experiment** tab.
- 10. On the Laser Events tab, select the desired laser for the drop-down menu.



11. To generate multiple bleach events, the stage moves the sample to the site of laser activity. You can optimize this motion for either speed or position. When **Optimize for Position** is selected, the stage goes through an LMC (Lost Motion Compensation) move to achieve the most accurate position. This move takes more time than when **Optimize for Speed** is selected.

- 12. Bleach events occur at either the center point of the field of view or in a specified position. Select **Use Bleach Event Specification** in the **Event pattern** field.
- 13. In the Create Bleach Events section of the window, select an event pattern by choosing the spot, line, or polygon tool.
- 14. Click on the image in the Data Collection window to generate the selected bleach pattern.
- 15. Click on the **Imaging** tab to select the **Channel Setup** (exposure time, wavelength, and illumination source settings) and **Time Course Specification** conditions.



- 16. When you select the Time course style, the drop-down list presents three choices:
  - Adaptive Time Intervals allows you to select the total number of images to be collected. Images are collected faster after the bleach event and then slower over time.
  - Uniform Time Intervals allows you to select the total number of images collected, as well as the time interval between images. The value in the **Experiment Duration** field is calculated based on these factors. Also, if you change the value in the **Experiment Duration** field, the **Post-event images** field automatically updates.
  - As Fast As Possible collects the images in rapid succession with no timelapse interval. For this style, you enter only the total number of **Post-event** images.

# The Run Experiment Tab

Use the Run Experiments tab options to access tools that allow you to select and execute experiment macros. There are also controls to monitor and interact with a running experiment.

Design/Run Experime			Hel
28 0			
Design Experime	ent Design PK Experiment	Run Experimen	1
Image file n	name [	Settings	s DMS Setup
Image	e title	ĺ	
Add note to	o log [	Do It	
Change next time la	apse 0.00	Do It	
✓ Launch viewe		PK progress graph volve during experim	ient Options
✓ Launch viewe	er after experiment 🔄 Decon		ient Options
✓ Launch viewe Sh	er after experiment 🔄 Decon		ient Options
✓ Launch viewe Sh Schedule Post-Ac	er after experiment Decon ow intensity graph None equisition Processing Tasks		ient Options
✓ Launch viewe Sh Schedule Post-Ac Images acquired / I	er after experiment Decon ow intensity graph None equisition Processing Tasks		ient Options
✓ Launch viewe Sh Schedule Post-Ac Images acquired / I Disk space	er after experiment ow intensity graph None equisition Processing Tasks requested: 010		ient Options
✓ Launch viewe Sh Schedule Post-Ac Images acquired / I Disk space	er after experiment Decom ow intensity graph None equisition Processing Tasks requested: 0/0 e required: 32.01 Mb command:		ent Options

### The Run Experiment Tab

Note The Design PK Experiment tab is available only for systems equipped with a laser module.



# The Green Arrow ⊵ Button

Starts the selected macro to run an experiment.

### Image file name

Specifies the file name to use for the image in this text field. If you do not provide a file name, you will be asked to provide one when the experiment starts running. If you have the **Auto-increment file names** setting (in the Settings window) turned on, you will only need to provide a name once for each session. The names will have incrementing numbers appended to them automatically.

### Image title

Specifies text to save in the header of the image file created by the experiment. The title can be viewed later using the Header Labels button of the Image window's Image Information window.

## Add note to log

Inserts a note in the experiment's log file at any time before or while an experiment is running. (You will need to click **Do It** to insert the note.)

# Change next time lapse

Changes the time lapse value while an experiment is executing. (You will need to type the desired time, in seconds, in the text field and click the **Do It** button to change the value.) This value only affects experiment macros that contain a TLAPSE command. It does not permanently change the macro.

## Note This value does not change the current time lapse.

# Show images during acquisition

Displays images in a Data Collection window as they are collected. In cases where acquisition speed is important, deselecting this option may increase performance.

# Show PK Progress graph

Opens a new window to display and update a preliminary results graph of a photokinetic experiment.

## Launch viewer after experiment

Launches an Image window when the experiment is finished. Note that if more than two files are created in a point-visiting experiment, the Image windows will not launch even if this option is selected.

## Show Intensity Graph

Provides three choices for displaying an intensity graph: **None**, **Maximum Intensity**, and **Mean Intensity**.

**Note** This option is available only for time-lapse images.

**Schedule Post-Acquisition Processing Tasks** Displays the Post Processing Options window.

Post Proces	ssing Options		*
🛛 Enable F	ost-Acquisition Proces	sing	
☐ Apply Pr	ocessing Tasks to Refe	rence Images	
Processi	ng Options		
🔟 Automat	ically Run the Queue		
_ Move In	put Files Before Proces	sing	
Choose F	older		
Processi	ng Tasks		1
	Task	Options	
	Deconvolution		
Add			
Done			

Refer to the section *"Scheduling Post-Acquisition Processing Tasks,"* in the <u>softWoRx</u> <u>Imaging Workstation User's Manual</u> for complete information on the options available from this window.

## Images acquired/requested:

Displays the current number of images acquired, compared to the total number requested. (During fast acquisition, the reported number of acquired images may not be updated regularly.)

**Note** The request portion of this is updated as the experiment is designed.

## **Disk space required**

Displays the estimated size of the experiment (i.e., the size of the image file). Resolve3D checks the disk space to make sure that enough space is available on the selected volume before it runs the experiment.

**Note** The disk space required value is updated as the experiment is designed.

## Elapsed time:

Displays the elapsed time of the experiment, in seconds.

## **Estimated Finish**

Displays the estimated clock time in which a running time-lapse experiment will finish.

## Current command:

Reports each macro command as it is executed.

# The Cancel Scan 💻 Button

Terminates a running experiment. (The images that are collected before the experiment is cancelled are automatically saved.)

```
Help
```

Opens the online Help for the Design/Run Experiment window.

# The Settings Window

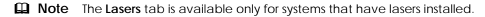
Use the Resolve3D Settings window to control how images are displayed, select camera settings, and specify file output.

## To open the Settings window:

From the Resolve3D Window, click the **Settings** button.

Resolve3D S	Settings					- 1
Display	Files	Imaging	Illumination	Autofocus	Lasers	Misc
Image disp	lay mode	None	•			
	Window	21 🔻	·]			
	Wave	1 🗸	-			
🔽 Calculat	e statistic	s				
🗸 Calculat	e histogra	am				
🗸 Auto his	togram ra	inge				
🗸 Display	images					
🗸 Deconv	olve previ	ew images				
🗸 Auto inte	ensity sca	ale				
🗸 Acquire	after poin	t visit				
🛛 Show ali	gnment ta	arget (cross	hairs)			
Done Sa	ave Settin	gs				Help

### The Settings Window | Display Options



# **Settings Window Display Options**

The Settings window Display tab options allow you to control how the images are displayed and what information about them is available.

## Image Display Mode

Specifies which window and wave are used for image display. Five modes are currently available:

Mode	Description
None	Displays images in the current window.
Scratch	Displays all images in the default Data Collection window (Window 21).
Auto Grayscale	Displays images in a separate window for each emission filter.
Auto Color	Displays images in color as they are collected. When using Point Visiting, images are automatically displayed in separate windows for each point. This option should be used only when you are running an experiment.
Point Track	Opens a separate window for each visited point in a point-visiting experiment.

## Window

Specifies the number of the data collection window, sometimes referred to as the "Scratch" window. Select a new window to change where the next image will be displayed. (Temporary windows are numbered 21 or higher.)

## Wave

Specifies the number of the wave (channel) that will receive the next image in the Data Collection window. Select a new wave to change which display window channel to use.

## **Calculate statistics**

Calculates image intensity statistics. The typical reason for disabling this feature is to improve readout speed.

## Calculate histogram

Calculates and displays the image intensity histogram in the Resolve3D status area during image collection. (This option does not set scaling.)

### Auto histogram range

When enabled, automatically scales the Resolve3D status area histogram width for each image that is acquired, so that the histogram display ranges from the minimum to the maximum intensity. When disabled, the status area histogram is scaled to the full range of a typical CCD.

### **Display images**

Displays images when the **Acquire** button is clicked or an experiment is running. Deselecting this option can increase performance.

### Deconvolve preview images

Displays instantly processed 2D image previews that closely resemble images processed with advanced 3D image restoration techniques.

## Auto intensity scale

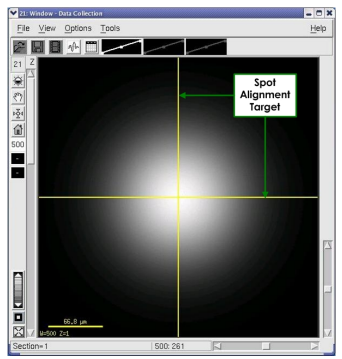
Provides automatic scaling of the image intensity between the minimum and maximum brightness. (This switch applies only to the appearance of the displayed image, not the actual data.)

## Acquire after point visit

Automatically acquires an image when the Visit Point option is selected.

## Show alignment target (crosshairs)

Displays a crosshair over the data collection window indicating the location of the image center.



Show Alignment Target

# **Settings Window Files Options**

Use the Settings window File options to set the image output directory, set the directory where experiment macros are stored, and select auto incrementing for file names.

Resolve	3D Settings					
Displa	y Files	Imaging	Illumination	Autofocus	Lasers	Misc
	Data folder	/di	ata 1į́			
Experin	nent macros	folder //h	ome/worxį́		-	
		<b>R</b>	Data folder is ter	mporary		
		₹	Auto-increment	file names		
			Convert to 2 byt	e signed intege	r	
Done	Save Settir	ngs				He

The Settings Window | Files Options

## **Data Folder**

Specifies the destination directory for Resolve3D output images. If you supply a directory that you do not have permissions to use, you will be warned. If you provide a directory name that does not exist, you will be presented with the option of creating the directory. (You can also change the destination directory by changing the global *softWoRx* Image Data Directory in the User Parameters tool.)

## **Experiment macros folder**

Specifies the directory where your experiment macros are stored.

## Data folder is temporary

Specifies that any directory entered in the **Data Folder** field only applies to the current Resolve3D session. In subsequent Resolve3D sessions, the Data folder value reverts back to the global *softWoRx* Data Directory as specified in User Parameters.

Use this option in environments where several users are running the microscope and using a single system login. Data folders can be created for each user under a common parent folder. As users start Resolve3D, they can choose to use their own folders.

In most cases, you should not use this option if each user has their own system login in your environment. With the option deselected, the Data folder assignment modifies the global *softWoRx* Data Directory definition and the system remembers the directory used in the last Resolve3D session.

## Auto-increment file names

Creates a new file name by appending a serialized number to the base file name each time you run an experiment.

# Convert to 2 byte signed integer

If enabled, Resolve3D saves all images as signed 16-bit (the softWoRx default). This setting is designed to increase compatibility with older software. Note that this setting cannot be enabled for EMCCD cameras.

# **Settings Window Imaging Options**

The Settings window Imaging options allow you to select camera settings.

	e3D Settings					•
Displa	ay Files	Imaging	Illumination	Autofocus	Lasers	Misc
	Camera		AP_ES2/ICX285	•		
rame	s to average*	· [1				
	Gain	4.00	•			
Т	ransfer speed	5000	•			
Targe	et temperature	0.00				
urrer	nt temperature	0.00	Refresh			
		₩ Use pł	notosensor			
Settir	ngs change bi	ased on Ex	citation Filter)			

# The Settings Window | Imaging Options

# Camera

Specifies which camera to use. For the EMCCD camera, this list also specifies whether to use Conventional or Electron Multiplication mode.

# Frames to average

Specifies the number of successive camera images to average into a result that is displayed in the window, increasing signal to noise. (This is similar to the AVG macro command.)

# Gain

This field allows you to specify a gain value for the selected camera.

# **Transfer speed**

Specifies a transfer rate for the selected camera (in kHz). Higher speeds may boost performance at the expense of image noise.

### Target temperature

This field displays the cooled CCD camera's target temperature. This can be changed only by modifying the appropriate configuration file on the Instrument Controller computer.

**Note** Not all cameras can be temperature controlled.

### Current temperature

This field displays the current temperature. Resolve3D monitors the temperature and updates this value occasionally. If you want to force a request for the current temperature, click the **Refresh** button.

## Use photosensor

Enables use of the photosensor.

# **Settings Window Illumination Options**

Use the options on the Resolve3D Illumination tab to set up illumination parameters.

Display	Files	Imaging	Illumination	Autofocu	s Lasers	Misc
	"EX 1	" bulb age (h	iours) <b>997.</b>	0 Refresh	Reset	
	"EX 2	" bulb age (h	iours) <b>997.</b> :	2 Refresh	Reset	
			🔽 Use	Beam Condit	ioner	
eld Stop /	Aperture (	(Closed <-> (	Dpen) 74 🗔		~	Ŧ
Critical/ł	<öhler illu	mination swit	ching Automa	atic	•	1

# "EX 1" and EX 2" bulb age (hours)

These fields provide figures for the current number of hours of use on each xenon bulb.

Note The EX 1 and EX 2 bulb age (hours) settings are displayed only if your *Delta Vision* system is using a xenon lamp for broadband illumination.

## Field Stop Aperture (Closed <-> Open)

This field allows you to set the field stop aperture to **Closed**, **Open**, or any percentage in between. You can either enter a percentage in the field or use the slider to set the percentage.

■ Note When the setting is set to Closed, the field stop aperture is about 20µm in diameter.

#### Critical/Köhler Illumination Switching

On an Applied Precision Fluorescence Illumination Module, there are three choices for the motorized Critical/Köhler illumination position:

- Automatic DeltaVision automatically adjusts the position of the critical/Köhler illumination when the EX shutter is opened using the keypad (typically when the eyepieces are being used). The position is automatically returned to critical illumination when an image is requested, either via Resolve3D or by running an experiment.
- Always Critical DeltaVision moves the critical/Köhler motor to critical illumination and leaves it in its position regardless of the current shutter states.
- Always Köhler DeltaVision moves the critical/Köhler motor to Köhler illumination and leaves it in its position regardless of any new image requests or experiments running.

# Settings Window Autofocus Options

Use the options on the Resolve3D Autofocus tab to set up Autofocus parameters.

Autofocus Z test step (µm) 3.12 Maximum Z test range (µm) 50.00	
Contrast calculation method Auto	
Autofocus Z test step (µm) (3.12 Maximum Z test range (µm) (50.00	
Maximum Z test range (µm) 50.00	
Post-autofocus Z offset (µm) [0.00	
UltimateFocus Hardware	
Calibrate UltimateFocus for plate scanning	

Settings Window | Autofocus Options

## Automatically determine parameters

Autofocus parameters are determined from the depth of field of the objective lens, which is calculated from the lens' numerical aperture. You must select the proper objective lens in order to get the appropriate Autofocus settings. When the **Automatically determine parameters** checkbox is activated, the remaining fields in this window are set automatically.

## **Channel for Autofocus**

This setting indicates the wavelength to use for Autofocus.

## Contrast calculation method

This setting determines the polarity of the contrast calculation. There are three choices for image contrast calculation methods:

- **Auto** The instrument controller usually can determine which contrast calculation method to use, but not always.
- Fluorescence for light objects on a dark background
- Brightfield for dark objects on a light background

## Autofocus Z test step (µm)

This option sets the step size used for Autofocus within the maximum Z range.

## Maximum Z test range (µm)

This setting indicates the maximum range that the Autofocus will search.

## Post-autofocus Z offset (µm)

This setting can provide a constant offset after the Autofocus position determines the best plane of focus. Often times Autofocus will find a plane that is consistently different from the desired plane.

## Calibrate UltimateFocus for Plate Scanning

This button displays the UltimateFocus Plate Calibration window.

UltimateFocus Plate Calibration	•
UltimateFocus Plate Calibration Protocol	
]1) Mount a microtiter plate containing sample in aqueous media	-
<ol><li>Find a well with material that you will be able to see, preferabl on the well bottom.</li></ol>	У
3) Manually lower the microscope objective to its minimum posi	tion.
<ol> <li>"Zero" the microtiter stage Z coordinate using the Home Stag button below.</li> </ol>	je Z
5) Using only the eyepieces and the coarse and fine focus knob raise the objective to focus on the sample on the well botton	
6) Choose the Calibrate Focus button below to perform the calibrate This procedure may take on the order of 20 minutes to complete the complete the calibrate complete complete the calibrate complete complete the calibrate comp	
Z Sweep Min, Max, Step Size 500.00 500.00 0.200	
Done Home Stage Z Calibrate Focus	Help

This window provides a procedure for plate calibration. The **Home Stage Z** and the **Calibrate Focus** buttons at the bottom of the window are used in the calibration procedure. You can also set the **Z Sweep Minimum**, the **Z Sweep Maximum**, and the **Z Step Size** from this window.

When working with microtiter plates, many factors affect the performance of UltimateFocus, such as stage angle, plate flatness, plate angle and more. These types of issues are mitigated by characterizing UltimateFocus response over a larger Z range in a single well. This characterization (or calibration) is then used during the course of an entire plate scan. The boxes at the bottom of the calibration tool are used to limit the calibration Z range (**Z Sweep Min/Max**) and the **Step Size** which is used between UltimateFocus calibration measurements. Typically, the entire Z range is used with a step size of 0.20  $\mu$ m. Using step sizes smaller than this can cause calibration to take longer to finish and may not increase performance.

# **Settings Window Lasers Options**

Use the options on the Resolve3D Lasers tab to set up parameters for the attached lasers.

Note If no lasers are installed on your *DeltaVision* imaging system, the Laser tab is not displayed.

Resolve3D S	ettings				
Display	Files	Imaging	Illumination	Autofocus	Lasers Misc
Laser Sta	itus —			U.C.	
√lodule Sta	tus 0	'n	Connect		
La	ser 🖡	7 405	<b>⊮</b> 488	<b>5</b> 14	<b>R</b> 561
Status Ready		Ready	Off	Stabilizing	
Power	(%) 5	D	 įso	Įso	Iso
Im	iage Th	ne Spot   Cor	ntinuous (Sp	olitter: 50% Pk	< <-> 50% TIRF)
		est   488 🔄	1	alignment targe blitter: 50% Pk	
		%	Bleach Ba	ckground	
Do	o Test (	Bleach	1		
TIRF					
Multi-line	TIRF S	Settings			
Done   Sa	ive Set	tings			Н

Settings Window | Lasers Options

## Laser

This field displays the peak wavelength for each laser currently installed on the *DeltaVision* system. Activate or deactivate the individual checkboxes to turn laser emission on or off.

# Status

Each field shows the current status of the associated laser. The fields will display **Off**, **Stabilizing**, or **Ready** for each installed laser.

# Power (%)

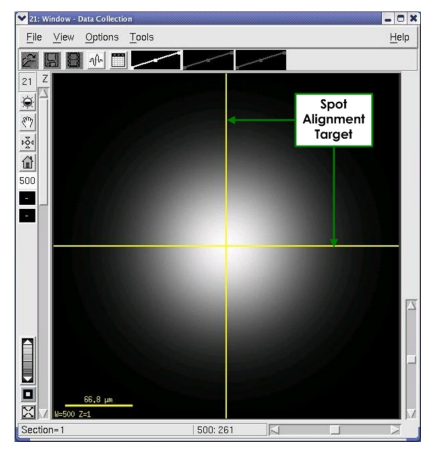
Each field displays the percentage of power currently applied to each laser.

## Laser to test

Use the drop-down list in this field to select a specific laser on which to run PK tests.

# Show spot alignment target

Activate this checkbox to include a target alignment reticle on the displayed test image as shown below.



PK Laser Tests | Image The Spot

## **Image The Spot**

Selecting this button opens the laser shutter and acquires a single image of the laser spot. The event duration is set using the **Exposure** field in Resolve3D.

## Continuous

Selecting this button opens the laser shutter and acquires continuous images of the laser spot. Select **Stop Imaging** to discontinue the imaging process.

# **Do Test Bleach**

Perform a bleach test using the current Power and PK Duration settings.

## % Bleach

The value entered in this field displays a before and after comparison of the intensities at the bleach spot: 100x (Signal After/Signal Before).

# Background

The value in this field provides a measurement of the signal (intensity value) after the bleach event.

# **Multi-line TIRF Settings**

Selecting this button opens the Multi-line TIRF Settings window.

Y Multi-li	ne TIRF Sett	ings			
TIRF I	Depth —				
<u>)</u> 405	[1090.0				
<b>6</b> 488	712.5	4			Z
0514	[390.0				
J 561	[390.0	4			
🔽 Adju	st channe	ls indeper	ndently		
Block	Block   Epifluorescence		Restore TIRF	Depth	
Back	Aperture	Focus Fo	r Current Len	s	
J-099.0				🐨 🔽 La	ocked
Laser	Path Spl	itter —			
50% PI	<u>د</u>			50%	TIRF
Done	Acquire	Contin	uous Acquire		Help

## Multi-line TIRF Settings Window

From the Multi-line TIRF Settings window, the following actions are available:

- **TIRF Depth** allows you to select a laser to adjust its evanescent wave depth.
- Adjust channels independently selects whether to independently adjust the TIRF Depth sliders. With the checkbox activated, you can adjust the depth setting for each laser separately.

■ Note Keep in mind that adjusting these separately means a significant time penalty for imaging.

- Block sets the TIRF depth adjuster to a position that prohibits illumination of the sample.
- **Epifluorescence** sets the TIRF depth adjuster for epifluorescence (the highest value of the TIRF depth slider).
- Restore TIRF Depth restores the TIRF depth adjuster to the last-used TIRF position.
- Back Aperture Focus for Current Lens focuses the laser on the back aperture of the objective resulting in a collimated illumination laser and an even TIRF depth across the field of view. This value should be set once for each installed objective. To make sure this position is set correctly for imaging, verify the proper objective is selected from the Resolve3D dropdown menu. The Locked checkbox (activated by default) keeps the slider from being inadvertently moved.

Laser Path Splitter – sets the percent of laser light to be directed to the TIRF illumination path versus the PK path.

# **Settings Window Misc Options**

Use the options on the Resolve3D Misc tab to select or clear Lost Motion Compensation or to select a filter wheel configuration (if your system is configured to use alternate filter wheels).

Display	Files	Imaging	Illumination	Autofocus	Lasers	Misc
- Stage Mo		n Compensa	ition (LMC)			
- Stage Vie Spiral Mose Show st Show st Show pt	aic 5x5 age trails age thum	• bnails				
– Filter Wh Excitation fi		Standard				
Emission fi	lter whee	Standard	<u> </u>			
Eyepiece fi Activate F						
Done Sa	ave Settin	as				Hel

The Settings Window | Misc Options

# Allow Lost Motion Compensation (LMC)

Enable Lost Motion Compensation to remove the effect of hysteresis in the stage. This option should be selected for most applications. When the option is turned off, the Z focal plane may shift depending on the direction of approach. Clear this option if you want to improve speed at the expense of position repeatability.

# **Spiral Mosaic**

Sets the size for the Spiral Mosaic preview collection pattern centered at the current stage location. The pattern begins at the center and then continues acquiring thumbnail images directly adjacent to it, spiraling outward in a counter-clockwise rotation so that the entire area centered around the initial stage position is previewed in the Resolve3D Stage View. Thumbnail collection continues until either the preset spiral mosaic size (set here) is reached or you click the **Spiral Mosaic** button again. The stage is always returned to the initial position regardless of how the collection ends.

## Show stage trails

Display the path of stage movement on the Resolve3D Stage View.

### Show stage thumbnails

Display a thumbnail image of each image on the Stage View as the image is acquired.

## Show point numbers Display the number of each point in a point list on the Stage View.

### **Filter Wheel Sets**

Use these fields to switch between active filter sets.

## **Activate Filter Sets**

This button reinitializes the filter wheels after changes have been made in any of the above fields. Finalizes any new selections made to the **Filter Wheel Sets** fields and may include reinitializing filter wheels.

# **Action Buttons**

Done

The **Done** button closes the Resolve3D Settings window.

### **Save Settings**

The **Save Settings** button preserves the current options for your next Resolve3D session. In addition to options in the Resolve3D Settings window, current state information such as currently selected filters and exposure time is saved.

### Help

The Help button opens the online Help for the Settings window.

# **Keypad/Joystick Operation**

Many of the functions accessible through Resolve3D are also available on the keypad/joystick. This section describes each key on the keypad/joystick.

**Note** Some buttons on your keypad may not be active.

BLANK SCREEN	CONTROL MODE CAMERA SHUTTER	LMC RESET ON/OFF	REMOVE TRAIL W MEDIUM	FAST	EYLIGHT ON/OFF DISABLE MOTION KEYS	YES	ΝΟ
ACQUIRE MODE ND EX EX	ACQUIRE MODE ND EX MODE		•			ETRACE	POINT
ACQUIRE IMAGE EX SHUTTER	SAVE IMAGE TRANS SHUTTER	STEP DECREASE	Y INCRE				POINT MARK

DeltaVision Keypad/Joystick

## RESET

Clears communication buffers, closes shutters, stops all motors, and clears encoder errors. Use the RESET button when you suspect that the workstation and controller are not synchronized.

# **CONTROL MODE**

Toggles the controller mode between local mode and remote mode. Local mode enables control by the keypad and joystick. The remote mode disables the joystick and the keypad buttons (except for CONTROL MODE and a few other buttons) and shifts control to other internal components.

## LMC RESET ON/OFF

Executes a Lost Motion Compensation (LMC) move.

Tip You can disable LMC on the Resolve3D Settings window. (For more information, see the online Help.)

## **REMOVE TRAIL**

Clears the stage movement history from memory.

## **KEYLIGHT ON/OFF**

Turns the keypad backlight on and off.

# **BLANK SCREEN**

Suspends (or activates) the monitor's light display (BLANK SCREEN is a toggle button). Use this feature when viewing dim samples or performing light sensitive experiments.

# SLOW/MEDIUM/FAST

Adjusts the joystick stage movement speed.

# **DISABLE MOTION KEYS**

Disables the eight keys below the joystick. This prevents accidental input from the keys when using the joystick.

# ACQUIRE MODE ARROWS

Changes the acquisition mode as set in Resolve3D. (This includes the Excitation filter, exposure time, shutter configuration, and many of the options defined in the Settings window.)

# JOYSTICK

Moves the stage in X and Y.

# POINT ARROWS

Scrolls through the list of marked points. Press ACQUIRE IMAGE to view the image for a selected marked point.

# ND and EX ARROWS

Changes the neutral density (ND) or excitation (EX) selection up or down the list.

# ACQUIRE IMAGE

Commands the system to collect image data. This key function is identical to clicking **Acquire** in Resolve3D. Use this key when you are scanning through your sample and using the eyepiece to find a region of interest, or when you want to get a quick look at the specimen on the monitor.

# SAVE IMAGE

Saves the last acquired image to the currently open file.

# STEP DECREASE/INCREASE

Changes the step size of the movement controlled by the arrow keys located beneath the joystick. Pressing either button many times will change the step size to the minimum/maximum. There are 8 possible step sizes: 50nm, 100nm, 200nm, 400nm, 800nm, 2250nm, 4500nm, and 9000nm.

# X AND Y ARROW KEYS

Move the stage in the direction shown.

# Z ARROW KEYS

Move the stage in the direction shown.

#### Z1 MARK/Z2 MARK

Mark the bottom (Z1) and the top (Z2) of the focal plane for your sample.

#### **EX SHUTTER**

Toggles the excitation shutter between open and closed.

#### TRANS SHUTTER

Toggles the transmitted light source between On and Off.

#### POINT MARK

Adds the stage position to the marked points list.

# F

## **Appendix F: Lasers and Safety Issues**

This appendix describes the necessary precautions to take when working with instruments containing lasers, including required safety labeling and label locations. The appendix is divided into two main sections.

- DeltaVision X4 Laser Module Safety provides laser safety information for the DeltaVision X4 Laser Module when used in conjunction with the Olympus Fluorescence Illumination Module.
- API Fluorescence Illumination Module Safety provides laser safety information for the DeltaVision X4 Laser Module when used in conjunction with the Applied Precision (API) Fluorescence Illumination Module.

Much of the information presented is redundant from one section to the next, but not all. Please read the sections carefully to determine the proper safety measures to take for your particular *DeltaVision* laser configuration.

## DeltaVision X4 Laser Module Safety

This section describes the hazards and precautions to take when using the *DeltaVision X4* Laser Module in conjunction with the Olympus Fluorescence Illumination Module. These hazards and precautions must be fully reviewed and understood, and proper safety protocols followed during any use of the *DeltaVision X4* Laser Module.

The DeltaVision X4 Laser Module is a Class 3B laser system.

#### **Important Safety Recommendations**

The X4 Laser Module contains up to three 50mW lasers and one 100mW laser, and is considered a Class 3B device. This means that radiation from all installed lasers can exit the device at the same time. The power level is high enough to cause damage to the human eye instantaneously.

OSHA regulations require (via ANSI Z136.1) and IEC 60825-1 recommends that a Laser Safety Officer be identified who will be responsible for the safe use of Class 3B lasers. This includes training users, installing all necessary warnings and controls in the laser area, and other duties.



#### WARNING:

Given the inherent exposure possible with an inverted frame microscope stand, users of the system *must be trained in laser safety before using this instrument*. Contact your lab administrator for information about Laser Safety training at your institution. Training is also available online at:

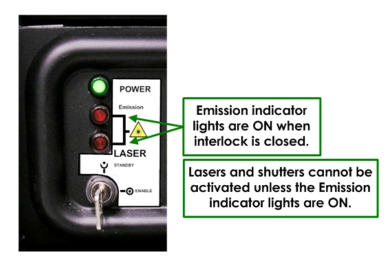
www.kentek-laser.com/edu/lasercrs.htm

The International Electrotechnical Commission (IEC) and the FDA recommend that Class 3B and Class 4 lasers be used only in restricted areas.

## **Safety Features**

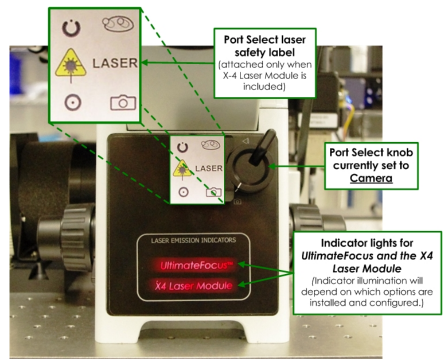
The Laser Module provides several features that control access to the device and improve its safety.

The Key Lock switch on the front panel of the Laser Source Chassis must be in the LASER ON position to enable use of the lasers. The key can be removed only when this switch is in the LASER STAND-BY position.



Laser Source Key Switch

A Port Selector Interlock prevents laser output unless the Port Selector knob on the microscope stand is set to the Camera (left side port) position. This device prevents exposure to the beam while looking through the eyepieces.



Port Selector Knob for Laser Module

Indicator lights on the system illuminate when the lasers are powered on and the interlocks are closed. The lasers can be fired at any time when the system is in this state, so the lights warn users to take any necessary precautions. A pair of indicator lights is visible on the front of the Laser Source Chassis. Another pair of indicator lights is visible on the Port Selector switch. All lights should be OFF or ON at the same time in proper operation.

- A User Interlock Connector on the back panel of the Laser Source Chassis allows you to install a door interlock to prevent laser output when the door is open. When the door interlock is active, the lasers turn ON but cannot emit light past the shutter.
- If you are not using a remote interlock, you must insert Interlock Jumpers into the User Interlock Port and the Fiber1 Interlock Port. These jumpers must be in place for the lasers to emit light.



#### Interlock Jumpers

For the lasers to operate, all of the following safety devices must be set as follows:

- The Key Lock switch must be ON.
- The Port Select knob must be set to the Camera position.
- The User Interlock Connector must be closed.
- At least one of the fiber interlocks must be closed.

If any of these devices are not set as described above, the shutters between the laser heads and the fibers are prevented from opening.

## **Avoiding Specific Hazards**

The hazard of being exposed to the laser beam through the objective turret when no objective is in place can be mitigated by turning off the key lock switch on the front of the Laser Source Chassis prior to any system reconfiguration or maintenance.

Specific hazards during alignment or system maintenance include:

- Exposure to the beam when disassembling the Fiber Optic Module from the microscope.
- Exposure to the beam when disassembling the Laser Optics Module from the Fiber Optic Module.

- Exposure to the beam when disassembling the optical fiber from the Laser Optics Module or the Laser Source Chassis.
- Exposure to the beam while the polychroic beam splitter turret is removed from the stand.

Radiation can be emitted as follows:

- Through the objective. The beam that comes through the objective is emitted during imaging experiments. It can be as powerful as 26mW and includes all wavelengths of lasers available in the Laser Source Chassis. The beam is highly divergent and depends on the objective used; using various API-approved objectives, the divergence angle can be as low as 45 degrees or as high as 140 degrees. With the maximum power on all lasers, using the lowest-NA objective, the Nominal Optical Hazard Distance (NOHD) is less than 10cm (4 inches).
- From the objective turret when no objective is in place. This beam is only visible during maintenance (for example, when aligning the Laser Optics Module) or when the lasers are triggered accidentally without an objective being in place. This beam is collimated and small (a few mm across). The beam power may be as high as 30mW.
- From the fluorescence illuminator when no polychroic is in place. This beam is only visible during maintenance, when adding or removing cubes from the polychroic turret. This beam is collimated and small (a few mm across). The beam power may be as high as 45mW.

Whenever you use the *X4* Laser Module, it is critical to keep your own safety and the safety of those around you a top priority. This is particularly important when maintaining the system, such as aligning the Optics Module or focusing the beam. *Some maintenance tasks involve potential exposure to dangerous levels of laser radiation.* According to ANSI Z135.1 (which is the standard the United States government uses for the safe use of lasers), the operator of a laser is responsible for the safety of everyone in the area, so you must be aware of the risks and keep others in the area safe. Some basic precautions will dramatically reduce the potential for injuries and damage.



WARNING: CAUTION - USE OF CONTROLS OR ADJUSTMENTS OR PERFORMANCE OF PROCEDURES OTHER THAN THOSE SPECIFIED HEREIN MAY RESULT IN HAZARDOUS RADIATION EXPOSURE.



**WARNING!** EVEN IF ALL OF THESE PRECAUTIONS ARE TAKEN, A RISK OF INJURY CAN STILL EXIST.

#### DURING USE:

• Make sure the system is ready for use before enabling the lasers. An objective should be in place and the polychroic turret should be installed. The indicator lights on the system (two small lights on or near the Port Selector Interlock, depending on whether or not the system includes the *API FI* Module) and two more on the Laser Source Chassis) should be either all ON or all OFF.

• **Do not lean close to the objective** to view the sample or make adjustments to parts of the system, etc. while the laser is on.

#### BEFORE BEGINNING MAINTENANCE:

• Whenever possible, work with the lasers disabled, for example, by turning the key to the "LASER STAND-BY" position.

• When lasers must be used, begin work by reducing the beam power as much as possible. Only increase the beam power if the work cannot be performed with lower laser power settings.

• Do not turn the laser on until the entire beam path is safe. Determine where the beam is going to go before turning on the laser and make sure the beam is blocked as soon as possible. Clear all reflective surfaces from the beam path—a reflected beam can be dangerous for several meters in many cases.

• **Be extremely vigilant about putting items in the beam path.** Tools, watches, rings and microscope samples can all make excellent reflective surfaces, and when inserted into the beam path, they can steer the beam in dangerous and unpredictable ways.

• If anyone is in the room, brief them on the procedures to be performed and what hazards will be present. Reiterate that the area may be dangerous and that they must comply with any instructions you give regarding safety.



WARNING: CAUTION - USE OF CONTROLS OR ADJUSTMENTS OR PERFORMANCE OF PROCEDURES OTHER THAN THOSE SPECIFIED HEREIN MAY RESULT IN HAZARDOUS RADIATION EXPOSURE.

## **TIRF-specific Laser Safety Considerations**

Due to the TIRF illumination optics provided by the TIRF Module and its laser component, the light being emitted from the *DeltaVision* objective is collimated and has high power density. The TIRF system also has the ability to direct this light to sharp, off-axis angles relative to the objective axis. Appropriate laser safety goggles selected for the specific wavelength being used are recommended.

## X4 Laser Module Safety Labeling

## **Standard Configuration**

**Avoid Exposure Caution Label** 



#### Avoid Exposure Label



#### Avoid Exposure Label

CAUTION

CLASS 3B VISIBLE AND INVISIBLE LASER RADIATION WHEN OPEN. AVOID EXPOSURE TO BEAM.

AVOID EXPOSURE.

VISIBLE AND INVISIBLE LASER RADIATION IS EMITTED FROM THIS APERTURE.



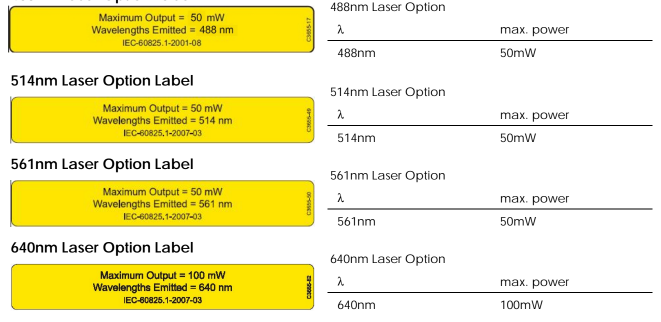
#### CAUTION

CLASS 3B VISIBLE AND INVISIBLE LASER RADIATION WHEN OPEN. AVOID EXPOSURE TO BEAM.

## **Optional Configurations**

#### 405nm Laser Option Label

Maximum Output = 100 mW	405nm Laser Option			
Wavelengths Emitted = 405 nm	λ	max. power		
	, 405nm	100mW		
445nm Laser Option Label				
Maximum Output: 40mW	445nm Laser Option			
Wavelengths Emitted: 445nm IEC-60825.1-2007-03	λ	max. power		
	445nm	40mW		



#### 488nm Laser Option Label

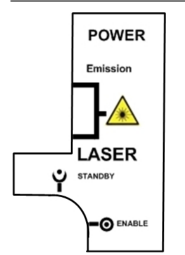
## X4 Safety Label Locations

Laser Safety labels and notifications should be installed on the X4 Laser Module product as illustrated below. In the event that a label is not installed or installed improperly, please notify Applied Precision by contacting hotline@api.com or calling 800.862.5166.

### X4 Laser Source Chassis

The following labels are placed on the front of the Laser Source Chassis.

Note: Only safety labels that correspond to the laser(s) installed in the Laser Source Chassis will be present.



Laser Module Key Switch Safety Label



405nm Laser Safety Label

Maximum Output = 50 mW Wavelengths Emitted = 488 nm IEC-60825.1-2001-08

488nm Laser Safety Label

Maximum Output = 50 mW Wavelengths Emitted = 514 nm IEC-60825.1-2007-03

514nm Laser Safety Label

Maximum Output = 50 mW Wavelengths Emitted = 561 nm IEC-60825.1-2007-03

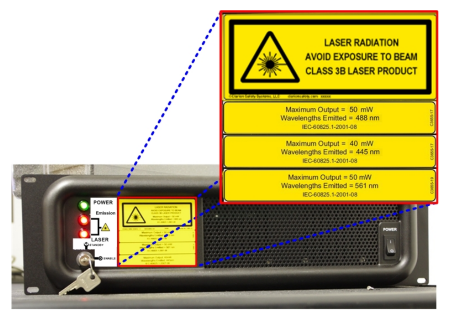
561nm Laser Safety Label

640nm Laser Safety Label

The following photos show the locations for the safety labels above to be placed on the front of the Laser Source chassis.



Laser Module Key Switch Label Location



Laser Source Chassis Laser Safety Label Locations

Note: Safety labels on the Laser Source Chassis will vary depending on which lasers are installed in your system.

## X4 Laser Optics Module

The X4 Laser Optic Module connects to the *DeltaVision* Fiber Optic Module on non-*API FI* systems. The Laser Optics Module has the following laser safety label attached in two places on the module as shown.



**Optics Module Laser Safety Label** 

One safety label is attached to the beam cover of the Optics Module as shown.



**Optics Module Safety Label (Location 1)** 

The second safety label is attached to the upper portion (opposite side) of the Optics Module as shown.



**Optics Module Safety Label (Location 2)** 

## **DeltaVision Stage**

The following safety label is placed on the edge of the *DeltaVision* stage as shown.



Laser Safety Label Location - DeltaVision Stage

## Polychroic Beam Splitter Removal Screw

The following label is placed on the side of the *DeltaVision*, next to the screw for removing the Polychroic Beam Splitter.



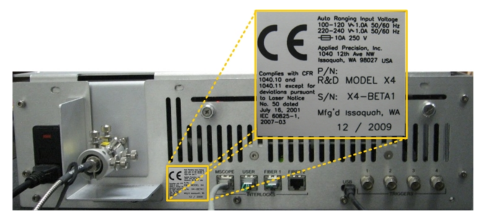
Polychroic Beam Splitter Removal Warning

This label warns users that, with the turret removed, laser radiation can come out through the fluorescence illuminator and be accessible. The label is attached to the *DeltaVision* as shown.



## CE Label

The CE Label for the *X4* Laser Module is attached to the back of the Laser Source Chassis as shown.



## **API Fluorescence Illumination Module Safety**

This section describes the hazards and precautions to take when using the *API FI* Module with lasers installed. These hazards and precautions must be fully reviewed and understood, and proper safety protocols must be followed during any use of the module, particularly those components which contain lasers.

#### **Important Safety Recommendations**

A *DeltaVision* system configured with the *API FI* Module can incorporate up to three 50mW lasers and one 100mW laser, and is considered a Class 3B device. This means that laser radiation from the installed lasers can exit the device at the same time. The power level is high enough to cause damage to the human eye instantaneously.

The *UltimateFocus*<sup>TM</sup> Module complies with CFR 1040.10 and 1040.11 except for deviations pursuant to Laser Notice No. 50, dated June 24, 2007 IEC 60825-1, 2007-03.

OSHA regulations require (via ANSI Z136.1) and IEC 60825-1 recommends that a Laser Safety Officer be identified who will be responsible for the safe use of Class 3B lasers at your site. This includes training users, installing all necessary warnings and controls in the laser area, and other duties.



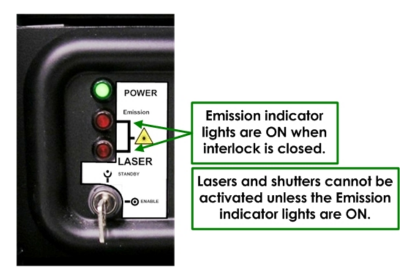
#### WARNING:

Given the inherent exposure possible with an inverted frame microscope stand, users of the system *must be trained in laser safety before using this instrument*. Contact your lab administrator for information about Laser Safety training at your institution. Training information is also available online at: **www.kenteklaserstore.com/** The International Electrotechnical Commission (IEC) and the FDA recommend that Class 3B and Class 4 lasers be used only in restricted areas.

## Safety Features

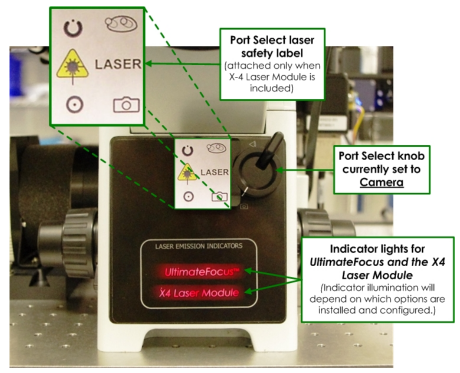
The Laser Module component of the API Fluorescence Illumination Module provides several features that control access to the device and improve its safety.

The Key Lock switch on the front panel of the Laser Source Chassis must be in the LASER ON position to enable use of the lasers. The key can be removed only when this switch is in the LASER STAND-BY position.



#### Laser Module Key Switch

A Port Selector Interlock prevents laser output unless the Port Selector knob on the microscope stand is set to the Camera (left side port) position. This device prevents exposure to the beam while looking through the eyepieces.



#### Port Selector Knob

Indicator lights on the system illuminate when the lasers are powered on and the interlocks are closed. The lasers can be fired at any time when the system is in this state, so the lights warn users to take any necessary precautions. A pair of indicator lights is visible on the front of the Laser Source Chassis. Another pair of indicator lights is visible on the Port Selector switch. All lights should be OFF or ON at the same time in proper operation.

- A User Interlock Connector on the back panel of the Laser Source Chassis allows you to install a door interlock to prevent laser output when the door is open. When the door interlock is active, the lasers turn ON but cannot emit light past the shutter.
- If you are not using a remote interlock, you must insert Interlock Jumpers into the User Port and the Fiber1 Interlock Port. These jumpers must be in place for the lasers to emit light.



For the lasers to operate, all of the following safety devices must be set as follows:

- The Key Lock switch must be ON.
- The Port Select knob must be set to the **Camera** position.
- The User Interlock Connector must be closed.
- At least one of the fiber interlocks must be closed.

If any of these devices are not set as described above, the shutters (between the laser heads and the fibers) are prevented from opening.

## **Avoiding Specific Hazards**

The hazard of being exposed to the laser beam through the objective turret when no objective is in place can be mitigated by turning off the key lock switch on the front of the Laser Source Chassis prior to any system reconfiguration or maintenance.

Specific hazards during alignment or system maintenance include:

- Exposure to the beam when servicing the Fiber Optic Module from the microscope.
- Exposure to the beam when servicing the Laser Optics Module from the Fiber Optic Module.

- Exposure to the beam when servicing the optical fiber from the Laser Optics Module or the Laser Source Chassis.
- Exposure to the beam while the polychroic beam splitter turret is removed from the stand.

Radiation can be emitted as follows:

• **Through the objective.** The beam that comes through the objective is emitted during imaging experiments. It can be as powerful as 26mW and include all wavelengths of lasers available in the Laser Source Chassis. The beam is highly divergent and depends on the objective used; using various API-approved objectives, the divergence angle can be as low as 45 degrees or as high as 140 degrees. With the maximum power on all lasers, using the lowest-NA objective, the Nominal Optical Hazard Distance (NOHD) is less than 10 cm (4 inches).

WARNING! When using low NA air objectives (anything below 0.45 NA), the *UltimateFocus*<sup>™</sup> laser beam does not diverge as much as it does with higher NA objectives. The beam, a Class 3B Invisible Laser light up to 2.2mW, is nearly collimated and is emitting straight up through the objective. Whenever the Laser Emission Indicators show that *UltimateFocus*<sup>™</sup> is on, users <u>must not</u> look down the objective turret.

- From the objective turret when no objective is in place. This beam is only visible during maintenance (for example, when aligning the Laser Optics Module) or when the lasers are triggered accidentally without an objective being in place. This beam is collimated and small (a few mm across). The beam power may be as high as 30mW.
- From the fluorescence illuminator when no polychroic is in place. This beam is only visible during maintenance, when adding or removing cubes from the polychroic turret. This beam is collimated and small (a few mm across). The beam power may be as high as 45mW.

Whenever you use any of the *DeltaVision* lasers, it is critical to keep your own safety and the safety of those around you a top priority. This is particularly important when maintaining the system, such as aligning the Optics Module or focusing the beam. *Some maintenance tasks involve potential exposure to dangerous levels of laser radiation.* According to ANSI Z135.1 (which is the standard the United States government uses for the safe use of lasers), the operator of a laser is responsible for the safety of everyone in the area, so you must be aware of the risks and keep others in the area safe. Some basic precautions will dramatically reduce the potential for injuries and damage.



WARNING: CAUTION - USE OF CONTROLS OR ADJUSTMENTS OR PERFORMANCE OF PROCEDURES OTHER THAN THOSE SPECIFIED HEREIN MAY RESULT IN HAZARDOUS RADIATION EXPOSURE.



**WARNING!** EVEN IF ALL OF THESE PRECAUTIONS ARE TAKEN, A RISK OF INJURY CAN STILL EXIST.

#### DURING USE:

• Make sure the system is ready for use before enabling the lasers. An objective should be in place and the polychroic turret should be installed. The indicator lights on the system (two small lights on the Port Selector Interlock and two more on the Laser Source Chassis) should be either all ON or all OFF.

• **Do not lean close to the objective** to view the sample or make adjustments to parts of the system, etc. while the laser is on.

#### **BEFORE BEGINNING MAINTENANCE:**

• Whenever possible, work with the lasers disabled, for example, by turning the key to the "LASER STAND-BY" position.

• When lasers must be used, begin work by reducing the beam power as much as possible. Only increase the beam power if the work cannot be performed with lower laser power settings.

• Do not turn the laser on until the entire beam path is safe. Determine where the beam is going to go before turning on the laser and make sure the beam is blocked as soon as possible. Clear all reflective surfaces from the beam path—a reflected beam can be dangerous for several meters in many cases.

• **Be extremely vigilant about not putting items in the beam path.** Tools, watches, rings and microscope slides can all create excellent reflective surfaces, and when inserted into the beam path, they can steer the beam in dangerous and unpredictable ways.

• If anyone is in the room, brief them on the procedures to be performed and what hazards will be present. Reiterate that the area may be dangerous and that they must comply with any instructions you give regarding safety.

• Always wear appropriate laser safety goggles.



WARNING: CAUTION - USE OF CONTROLS OR ADJUSTMENTS OR PERFORMANCE OF PROCEDURES OTHER THAN THOSE SPECIFIED HEREIN MAY RESULT IN HAZARDOUS RADIATION EXPOSURE.

## **TIRF-specific Laser Safety Considerations**

Due to the TIRF illumination optics provided by the TIRF/PK Module and its laser component, the light being emitted from the *DeltaVision* objective is collimated and has high power density. The TIRF system also has the ability to direct this light to sharp, off-axis angles relative to the objective axis. When servicing the TIRF system, use extreme caution that the emitted light is not directed into the user's eyes. Appropriate laser safety goggles selected for the specific wavelength being tested are mandatory.

## Safety Labeling

## **Standard Configuration**

AVOID EXPOSURE

VISIBLE AND INVISIBLE ASER RADIATION IS EMITTED FROM THIS APERTURE

No C30

#### **Avoid Exposure Caution Label**



#### CAUTION

CLASS 3B VISIBLE AND INVISIBLE LASER RADIATION WHEN OPEN. AVOID EXPOSURE TO BEAM.

AVOID EXPOSURE.

VISIBLE AND INVISIBLE LASER RADIATION IS EMITTED FROM THIS APERTURE.

#### Avoid Exposure Label

alety Systems LLC



clarionsafety com xxxx

#### CAUTION

CLASS 3B VISIBLE AND INVISIBLE LASER RADIATION WHEN OPEN.

AVOID EXPOSURE TO BEAM.

405nm Laser Option Label					
Maximum Output = 100 mW	405nm Laser Option	1			
Wavelengths Emitted = 405 nm IEC-60825.1-2001-08	δ.	max. power			
	405nm	100mW			
445nm Laser Option Label					
Maximum Output: 40mW	445nm Laser Optior	445nm Laser Option			
Wavelengths Emitted: 445nm IEC-60825.1-2007-03	λ	max. power			
	445nm	40mW			
488nm Laser Option Label					
Maximum Output = 50 mW	488nm Laser Optior	1			
Wavelengths Emitted = 488 nm IEC-60825.1-2001-08	<u>λ</u>	max. power			
IEC-60825.1-2001-08	488nm	50mW			
514nm Laser Option Label					
•	514nm Laser Optior	1			
Maximum Output = 50 mW Wavelengths Emitted = 514 nm	λ	max. power			
IEC-60825.1-2007-03	514nm	50mW			
561nm Laser Option Label					
-	561nm Laser Optior	1			
Maximum Output = 50 mW Wavelengths Emitted = 561 nm	- λ	max. power			
IEC-60825.1-2007-03	561nm	50mW			
640nm Laser Option Label					
	640nm Laser Optior	1			
Maximum Output = 100 mW Wavelengths Emitted = 640 nm	λ 	max. power			
IEC-80825.1-2007-03	640nm	100mW			

## **Optional Configurations**

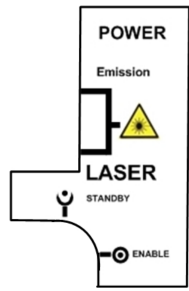
## **Safety Label Locations**

Laser Safety labels and notifications should be installed on the API Fluorescence Illumination Module product as illustrated below. In the event that a label is not installed or installed improperly, please notify Applied Precision by contacting hotline@api.com or calling 800.862.5166.

## Laser Source Chassis

The following labels are placed on the front of the Laser Source Chassis.

Note: Only safety labels that correspond to the laser(s) installed in the Laser Source Chassis will be present.



Laser Module Key Switch Safety Label



Primary Laser Safety Label

Maximum Output = 100 mW Wavelengths Emitted = 640 nm IEC-60825.1-2007-03

R

640nm Laser Safety Label

Maximum Output = 50 mW Wavelengths Emitted = 561 nm IEC-60825.1-2007-03

561nm Laser Safety Label

Maximum Output = 50 mW	q
Wavelengths Emitted = 514 nm	and a second
IEC-60825.1-2007-03	ĉ

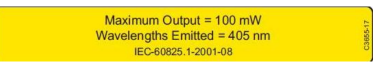
#### 514nm Laser Safety Label

Maximum Output = 50 mW Wavelengths Emitted = 488 nm IEC-60825.1-2001-08

488nm Laser Safety Label

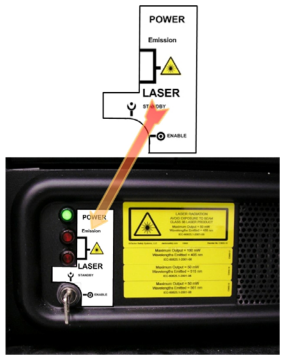


#### 445nm Laser Safety Label

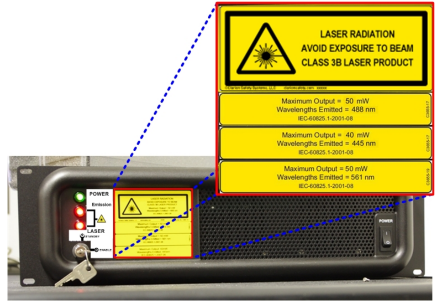


#### 405nm Laser Safety Label

The following photo shows the locations for the safety labels above to be placed on the front of the Laser Source chassis.



Laser Module Key Switch Label Location



Laser Source Chassis Laser Safety Label Locations

Note: Safety labels on the Laser Source Chassis will vary depending on which lasers are installed in your system.

## Laser Safety Labels with API FI

Depending on how the API Fluorescence Illumination Module is configured, it can result in adding several different visible and invisible laser wavelengths to the standard *DeltaVision* system. Since the different options available with this module are so many and varied, you should consult the following table to determine the location and type of Laser Safety Labels that should be attached to a particular system.

The following table shows the location and type of Laser Safety Labels, depending on the various options installed, that should be attached to a system equipped with the *API FI* Module. Use this table in conjunction with the illustrations immediately following, in which the label positions are called out by number.



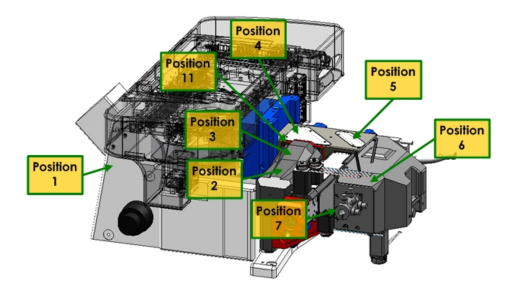
APIFI Module Laser Safety Label Types and Locations on the Delta Vision system

Visible and Invisible Laser Warning			9
Class 3B Invisible when Open Laser Warning		1, 4, 8, 10	1, 4, 8, 10
Class 3B Visible when Open Laser Warning	1, 4, 5 <sup>≜</sup> , 6, 7, 10, 11 <sup>≜</sup>		1, 4, 5, 6, 7, 10
Invisible Laser Radiation, Class 3B		2	2
Class 3B Visible and Invisible when Open		3	3

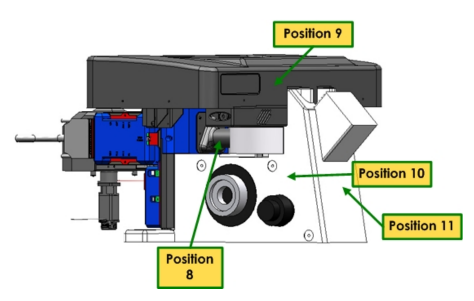
Port Select Laser Safety Label	11	11	
0			

 $\underline{A}$ Label is also present underneath this panel.

<sup>B</sup>Label is present only when the Ultimate Focus Module is not installed.



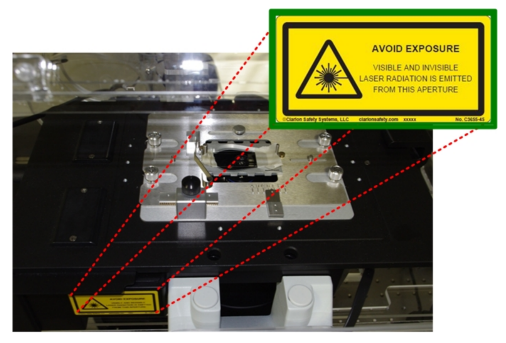
Laser Safety Label Locations (as viewed from the right rear of the system)



Laser Safety Label Locations (as viewed from the left front)

## **DeltaVision Stage**

The following label is placed on the edge of the *DeltaVision* stage as shown.



Laser Safety Label Location - Delta Vision Stage

### CO<sub>2</sub> Cell Box / TIRF Cover

The following label is placed on the CO<sub>2</sub> Cell Box / TIRF Cover as shown.

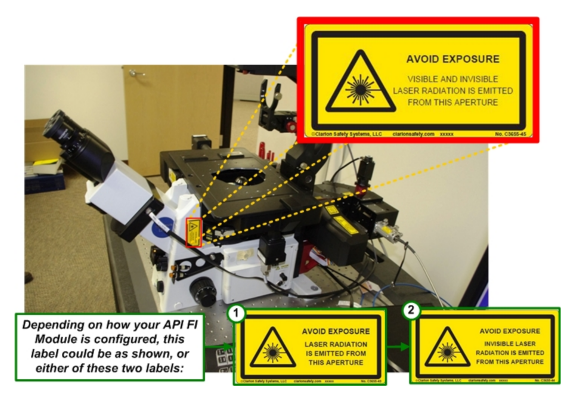


Laser Safety Label Location - CO2 Cell Box / TIRF Cover

## Polychroic Turret Removal

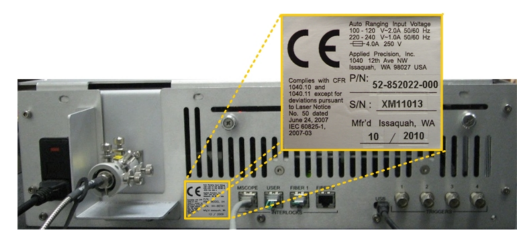
The following label is placed on the right side of the *DeltaVision*, next to the screw for removing the Polychroic Turret.

This label warns users that, with the turret removed, laser radiation can be accessible coming out through the fluorescence illuminator. The label is attached to the *DeltaVision* as shown.



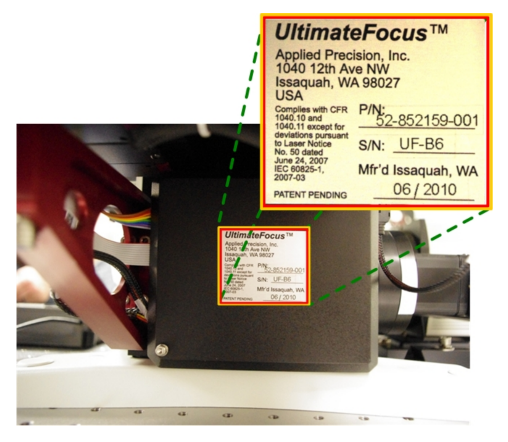
## **Compliance Labels**

The CE Label for the *X4* Laser Module is attached to the back of the Laser Source Chassis as shown.



#### X4 Laser Module - CE Label

The Compliance Label for the *UltimateFocus*<sup>™</sup> Module is attached to the back of the *DeltaVision* Microscope as shown.



*UltimateFocus™* Compliance Label

# G

# Appendix G: China\_RoHS

该标志表明本产品含有超过SJ/T11363-2006

电子信息产品中有毒有害物质的限量要求》中限量的有毒有害物质。标志中的数字为本产品的环保使用期,表明本产品在正常 使用的条件下,有毒有害物质不会发生外泄或突变,用户使用本产品不会对环境造成严重污染或对其人身、财产造成严重损害 的期限,单位为年。

为保证所申明的环保使用期限,应按产品手册中所规定的环境条件和方法进行正常使用,并严格遵守产品维修手册中规定的定 期维修和保养要求。

产品中的消耗件和某些零部件可能有其单独的环保使用期限标志,并且其环保使用期限有可能比整个产品本身的环保使用期限短。应到期按产品维修程序更换那些消耗件和零部件,以保证所申明的整个产品的环保使用期限。

本产品在使用寿命结束时不可作为普通生活垃圾处理,应被单独收集妥善处理。

This symbol indicates the product contains hazardous materials in excess of the limits established by SJ/T11364-2006 Marking for Control of Pollution caused by Electronic Information Products. The number in the symbol is the Environment-friendly Use Period (EFUP), which indicates the period during which the toxic or hazardous substances or elements contained in electronic information products will not leak or mutate under normal operating conditions so that the use of such electronic information products will not result in any severe environmental pollution, any bodily injury or damage to any assets, the unit of the period is "Year". In order to maintain the declared EFUP, the product shall be operated normally according to the instructions and environmental conditions as defined in the product manual, and periodic maintenance schedules specified in Product Maintenance Procedures shall be followed strictly.

Consumables or certain parts may have their own label with an EFUP value less than the product. Periodic replacement of those consumables or parts to maintain the declared EFUP shall be done in accordance with the Product Maintenance Procedures.



志表明本 品不含有超 中国标准SJ/T11363-2006 子信息 品中有毒有害物 的限量要求"中限量的有毒有害物,后可以 行回收 理,不能随意 弃

This symbol indicates that this electronic information product does not contain any toxic or hazardous substances or elements above the maximum concentration value established by the Chinese standard SJ/T11363-2006, and can be recycled after being discarded, and should not be casually discarded.

0: 表示该有毒有害物质在该部件所有均质材料中的含量均在SJ/T11363-2006 标准规定的限量要求以下

- X: 表示该有毒有害物质至少在该部件的某一均质材料中的含量超出SJ/T11363-2006 标准规定的限量要求
- 此表所列数据为发布时所能获得的最佳信息
   由于缺少经济上或技术上合理可行的替代物质重

由于缺少经济上或技术上合理可行的替代物质或方案,此医疗设备运用以上一些有毒有害物质来实现设备的预期临床功能,或给人员或环境提供更好的保护效果。

O: Indicates that this toxic or hazardous substance contained in all of the homogeneous materials for this part is below the limit requirement in SJ/T11363-2006. X: Indicates that this toxic or hazardous substance contained in at least one of the homogeneous materials used for this part is above the limit requirement in SJ/T11363-2006.

Data listed in the table represents best information available at the time of publication
 Applications of hazardous substances in this medical device are required to achieve its intended clinical uses, and/or to provide better protection to human beings and/or to environment, due to lack of reasonably (economically or technically) available substitutes.

部件名称	有毒有害物质或元素					
	Hazardous substances' name					
	铅	汞	镉	六价铬	多溴联苯	多溴二苯醚
Component Name	(Pb)	(Hg)	(Cd)	(Cr6+)	(PBB)	(PBDE)
Printed Circuit Board Assemblies	х	0	0	0	0	0
Arc Lamp and Filter Assembly	х	х	0	0	0	0
Power Supplies	х	0	0	0	0	0
Cables	Х	0	0	0	0	0
Enclosures	0	0	0	0	0	0
Microscope Assembly	0	0	0	0	0	0
Table Assembly	0	0	0	0	0	0

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