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The single-color miniature fluorescence microscopy systems are configured to image a single fluorophore within small animal subject brains. An imaging cannula is used to send excitation light to fluorophores and receive the resulting fluorescence. A miniature fluorescence microscope body (Section 2.1, 2.2) uses a CMOS camera to take fluorescence images transmitted by the cannula and sends them to a computer. Multiple configurations and modalities are available for a multitude of specialized applications.
1.0.1 Basic Fluorescence Microscopy Systems

The **Basic Fluorescence Microscopy System (BFMS)** (Fig. 1.1a) allows the imaging of a single fluorophore using a single wavelength light source. These systems are available in freely-moving and head-fixed configurations. The standard GCaMP6 system (Fig. 1.2) uses a Connectorized LED as a light source, while the standard RCaMP2 system (Fig. 1.3) uses a Ce:YAG optical head. Each system uses a different Microscope Body optimized for each fluorophore excitation/emission wavelengths.

![Figure 1.2: GCaMP6 Excitation Filter Set Spectra](image1)

![Figure 1.3: RCaMP6 Excitation Filter Set Spectra](image2)
1.0.2 Optogenetically Synchronized Fluorescence Microscopy Systems

![GCaMP6+NpHR3.0 Excitation Filter Set Spectra](image1)

The **Optogenetically Synchronized Fluorescence Microscopy (OSFM) system** (Fig. 1.1b) allows the imaging of a single fluorophore while performing optogenetic activation/deactivation. These systems are available in freely-moving and head-fixed configurations. When using a **GCaMP6 + NpHR3.0** microscope body a Ce:YAG + LED optical head is needed for illumination (Fig. 1.4). When using a **RCaMP2 + ChR2** microscope body a Ce:YAG + Laser Diode optical head is needed for illumination (Fig. 1.5).

![RCaMP6+ChR2 Excitation Filter Set Spectra](image2)

1.0.3 Freely-moving and Head-fixed Configurations

The **freely-moving** (Fig. 1.1a) configuration allows fluorescence microscopy to be performed while the subject is active thanks to a Pigtailed assisted fiber-optic & electric rotary joint. The **head-fixed** configuration is used in combination with stereotaxic instruments. It is also used for high-precision cannula implantation.
Device Overview

2.1 Basic Snap-in Fluorescence Microscope Bodies

The SFMB Bodies allow in vivo imaging of fluorescence within small animal subject brains. The microscopes are optimized to image calcium indicators like GCaMP6 and RCaMP2. The Basic Snap-in Fluorescence Microscope Body is offered in two models: Model L for deep brain imaging and Model S for brain surface imaging. The Model L has a 0.5 NA objective lens within its body and is used with an implantable imaging cannula that transfers the image to the microscope CMOS camera. The Model S has a plan-parallel plate instead and relies on the objective lens within the model S imaging cannula to create an image on the CMOS. The SFMB are composed of the following elements.

- **Fiber-optic patch cord** connects to the **M3 optical connector**, bringing light to the microscope. The patch cord length is chosen to correspond to the electrical cable length.
- **Electrical cable** sends and receives electrical signal for the microscope using an HDMI connector, with a standard length of 1 m. The cable is jacketed in either the light and flexible Ultralight fiberglass jacket or the robust but heavier Lightweight metal jacket.
- **Microscope clamps** secure the microscope on an imaging cannula. This allows the removal of the microscope body between each imaging session.
- **CMOS sensor** records fluorescence images. Each **CMOS sensor** has a serial number stored in memory that links each microscope to a specific set of mask correction filters recognizable to our software package.
- **Cannula Model L** and **Protrusion adjustment ring** are described in Section 2.3.

![Figure 2.1: Basic Snap-in Fluorescence Microscope Body](image)
2.2 Optogenetically Synchronized Fluorescence Microscope Bodies

The OSFM bodies combine fluorescence imaging and optogenetic stimulation/inhibition capabilities within the Single-color miniature fluorescence microscope. To avoid cross talk between optogenetic stimulation and fluorescence imaging, the OSFM hardware provides for at least two distinct spectral bands for optogenetic activation or fluorophore excitation (like blue and yellow). As the field of opsins and calcium indicators is very dynamic, spectral bands can be tailored to specifications. GCaMP6 + NpHR3.0 and RCaMP2 + ChR2 microscope versions are currently available. The appearance (Fig. 2.1) and general specifications (Table 5.1) of the OSFM Body are identical to the BSFM Body.

2.3 Snap-in Imaging Cannulas

Snap-in imaging cannulas transmit images of structures located inside of the brain to the surface of the skull. Cannulas have the following elements.

- The Cannula clamp groove is used to secure the microscope body in place using Cannula clamps.
- The Implantation relay lens is found on the Model L imaging cannula (Fig. 2.2 right). It is an image guiding gradient-index rod lens that brings the image from inside the brain to the skull surface.
- The Objective lens is found on the Model S imaging cannula (Fig. 2.2 left). For areas near the brain surface (less than 150 μm below the skull) it provides higher image quality and a larger FOV than the Model L imaging cannula.

A set of rod lenses with different lengths is available to reach different depth ranges of the brain tissue. Fine focusing of the Model L imaging cannula is done with the protrusion adjustment ring that comes with each cannula (Fig. 2.3). The Model S Imaging Cannula uses a single protrusion adjustment ring (height of 4.5 mm).

Figure 2.2: Snap-in imaging cannula type S (left) and type L (right)

Figure 2.3: Model L Imaging Cannula Protrusion Adjustment Rings. Height from left to right: 2.05 mm, 2.77 mm, 3.48 mm, 4.20 mm and 4.92 mm
2.4 Dummy Microscope

The **Dummy Microscope** is of similar weight and size as the microscope bodies. It is used to habituate animal subjects to the weight and feel of the microscope.

2.5 Pigtailed Assisted Fiber-optic & Electric Rotary Joint

To use the microscope in experiments using freely-moving animals, the **Pigtailed Assisted Fiber-optic & Electric Rotary Joint** (Fig. 2.5) is provided. This rotary joint allows effectively frictionless rotation of optical fibers and electrical cables connected to the microscope.

- The **Fastened Patch Cords** transmit light from the light source to the microscope. They are integrated into the rotary joint for maximal optical signal stability. The fixed and rotary patch cords use an FC connector.
- The **HDMI connectors** are used to provide electrical communication between the driver and the microscope.
- The **5 V mini-USB-B connector** connects to the power supply to allow assisted rotation.
2.6 Fluorescence Microscope Driver

This driver allows for computer control over the excitation LED light source as well as microscope image capturing and broadcast at video rate to a single computer via high speed Ethernet communication. It can be triggered by or synchronized with external recording devices and it can also trigger other devices using 5V TTL signals (Fig. 2.6). This Fluorescence Microscope Driver is used with the SFMB System, the OSFM System and the eSFMB System. The standard model has an integrated LED light source driver.

![Fluorescence Microscope Driver](image)

Figure 2.6: Fluorescence Microscope Driver

2.7 Fluorescence Microscope Snapping Tool

![Fluorescence Microscope Snapping Tool](image)

The Fluorescence Microscope Snapping Tool is used to remove the Microscope clamps from the Cannula clamp groove.
2.8 Fluorescence Microscope Holder

The Fluorescence Microscope Holder (FMH) and the Clamp for Fluorescence Microscope Holder are used to secure the microscope in a stereotaxic system.

- The Clamp for Fluorescence Microscope Holder (Fig. 2.8b) can be secured to a rod in a stereotaxic system using the Slot. The Clamp secures the Fluorescence Microscope Holder while still allowing easy removal of the holder.
- The FMH FC Connector allows the holder to be connected to an FC-connectorized patch cord.
- The FMH Ferrule is inserted into the microscope M3 Optical Connector. The ferrule is secured by screwing the FMH Extremity onto the M3 Optical Connector.

![Fluorescence Microscope Holder Elements](image1)

![Clamp for Fluorescence Microscope Holder](image2)

Figure 2.8: Fluorescence Microscope Holder
3.1 Connecting the Microscope Driver

- Connect the microscope driver to the power outlet using the 12 V power supply.
- Connect the microscope driver to the computer or to the router using the Ethernet cable.
- Connect the microscope HDMI cable to the CMOS port on the driver.
- Push the power button. After \( \sim 5 \) seconds the LED will turn ON and start to blink.

3.2 Installing the software

1. Run the Doric Neuroscience Studio Installer from the supplied USB key or download the latest version of the software from our website. See Table 5.8 for computer requirements.
2. Select the language to use during the installation.
3. In the license agreement window (Fig. 3.1), accept the agreement and click Next to continue the process.

![License Agreement Window]

**Figure 3.1: Doric Neuroscience Studio License Agreement**

4. Click Next in the Information window.
5. **Choose** where to install the software (Fig. 3.2) and click **Next**.

![Select Destination Location](image1)

Figure 3.2: *Select Destination Location*

6. **Choose** if desired to create a shortcut in the Start Menu folder and click **Next**.

7. **Choose** if desired to create a desktop icon and click **Next**.

8. When ready, click **Install** to begin the process. This should take a few moments. When the installation is done, the message in figure 3.3 will show up.

![Successful Installation](image2)

Figure 3.3: *Successful Installation of the Doric Neuroscience Studio*

9. Click **Next** and **Finish** to exit the setup.

10. Now the software is ready for use.

### 3.3 Setting Up The Communication

In order to communicate with the driver, the IP address of the computer must be static. If the driver is connected to a router, jump to section 3.3.3. If the driver is connected directly to the computer, continue to section 3.3.1.
3.3.1 Configuring Static IP Address

To change the computer’s IP address in Windows 7, type `network and sharing` into the Search box in the Start Menu and select Network and Sharing Center. If you are in Windows 8, it will be on the Start Screen itself (Fig. 3.4).

![Figure 3.4: Open Network and Sharing Center.](image)

On the side menu, select Change Adapter Settings (Fig. 3.5).

![Figure 3.5: Click on Change Adapter Settings.](image)

Right-click on the local adapter and select Properties (Fig. 3.6).
Select Internet Protocol Version 4 (TCP/IPv4) in the list, and click on Properties (Fig. 3.7).

Use the following IP address, and set the new IP address to 192.168.1.149, and the Subnet mask to 255.255.255.0. Leave the Default gateway and the DNS settings empty. Finally, click OK and close the Network Center (Fig. 3.8).
3.3.2 Activating The JUMBO Ethernet Frames

In order to reduce the load on the computer CPU, the Doric Neuroscience Studio is using JUMBO frames for the image transfer. In order to activate the JUMBO frames, open the Ethernet interface Properties menu as shown in Fig. 3.6 and click Configure (Fig. 3.9).

Figure 3.9: Interface configuration.

In the Configuration window, click on the Advanced tab, and on jumbo frames in the list. From the choices, select a value >4 KB MTU, the bigger the better (Fig. 3.10).

Figure 3.10: Jumbo frame configuration.
3.3.3 Configuring The Driver IP Address

The microscope must be connected to the driver in order to proceed.

When connected for the first time, the microscope driver will broadcast its MAC address. To set the IP address, you can use the built-in ethernet setup interface in the controller software.

- Turn ON the microscope driver.
- Launch the Doric Neuroscience Studio software.
- Allow ~10 seconds for the microscope initialization.
- Choose the proper network interface from the list in the Ethernet Settings box, and click on Pair and Remember to configure the microscope driver.

When the system is ready to use, the power switch LED will stop blinking, and the software status will display The microscope is ready.

3.4 Updating The Driver Firmware

To update the driver firmware version, contact us at sales@doriclenses.com. The updated firmware and an installation guide will be provided.

3.5 Updating Doric Neuroscience Studio

To update Doric Neuroscience Studio see the Neuroscience Studio Manual.
3.6 General Setup Guidelines

3.6.1 Optical fiber patch cord use

1. Clean the optical fiber connector before insertion. Use isopropanol and a lint-free wipe.

2. With an FC connector, the connector key must be oriented to enter within the receptacle slot to ensure proper connection (Fig. 3.11).

![Figure 3.11: FC connector, Fiber Installation](image)

⚠️ To reduce the risk of eye injury, it is sound practice to NOT CONNECT/DISCONNECT OPTICAL FIBERS when the light source is turned on.

3.6.2 Microscope Clamping Procedure

For details concerning the imaging cannula handling and implantation, as well as the microscope clamping procedure, it is important to read the Snap-in Miniature Fluorescence Microscope Surgical Procedure, as well as our Instructional Video.

3.7 Connecting the Fluorescence Microscope System

There are multiple different fluorescence microscope system configurations depending on the desired freedom of the animal and the fluorophores targeted. The animal must be held in one of the two following configuration:

- The **Head-fixed** configuration (Section 3.7.1, Fig. 3.13a) allows the observation of the animal while placed in a stereotaxic apparatus.

- The **Freely-Moving** configuration (Section 3.7.2, Fig. 3.13b) allows the observation of the animal while moving freely in a cage.

Depending on the **Microscope Body** type, several different lighting configurations are used. The most common are presented in this manual:

- The **SFMB GCaMP6** configuration (Section 3.7.3, Fig. 3.14a) allows the imagery of **GCaMP6** fluorescence.

- The **SFMB RCaMP2** configuration (Section 3.7.3, Fig. 3.14b) allows the imagery of **RCaMP2** fluorescence.

- The **OSFM GCaMP6 + NpHR3.0** configuration (Section 3.7.4, Fig. 3.15a) allows the imagery of **GCaMP6** fluorescence with **NpHR3.0** activation.

- The **OSFM RCaMP2 + ChR2** configuration (Section 3.7.4, Fig. 3.15b) allows the imagery of **RCaMP2** fluorescence with **ChR2** activation.
Figure 3.12: Full RCaMP2 Freely-Moving Configuration, combination of (Fig. 3.13b) and (Fig. 3.14b)

Figure 3.13 shows the head-fixed and freely-moving system configurations. Each microscope body and lighting configuration uses the same devices in each of these configurations.

### 3.7.1 Head-fixed configuration installation

1. Install the Fluorescence microscope holder into the Stereotaxic clamp (Fig. 3.13a). Ensure the holder is tightly secured inside the clamp.

2. To secure the microscope on the Fluorescence microscope holder:
   
   a) Remove the Connector caps from the microscope M3 optical connector and the Fluorescence microscope holder ferrule.
   
   b) Insert the ferrule into the M3 optical connector. Secure them in place by screwing the Fluorescence Microscope holder extremity.

3. Install the Fluorescence microscope holder in a stereotaxic apparatus.

4. Connect the Fluorescence microscope holder and the light source using a fiber-optic patch cord. The patch cord characteristics (MFP_200/230/LWMJ-0.48_1m_FC-FC) are set to achieve optimal coupling efficiency with the Fluorescence microscope holder.

5. Connect the Pigtailed microscope cable to the driver HDMI connector.

6. When ready for use, remove the cannula protective cap by unscrewing it. If using a type-L microscope, take great care to remove the cap in a slow, straight motion so as not to break the rod lens inside.
3.7.2 Freely-moving configuration installation

1. Link the driver and the Pigtailed assisted fiber-optic and electric rotary joint using the HDMI cable (Fig. 3.13b).

2. Connect the top pigtailed patch cord to the light source Beam aperture.

3. Connect the microscope Pigtailed HDMI Cable to the rotary joint.

4. Connect the lower pigtailed patch cord of the rotary joint to the FC mating adapter, then connect the mono fiber-optic patch cord (FC-CM3) to the microscope. The patch cord (MFP_200/230/LWMJ-0.48_0.8m_FC-CM3) linking the Pigtailed Assisted Fiber-optic & Electric Rotary Joint (AHRJ) and the microscope is chosen to ensure proper illumination characteristics.

5. Remove the Input Protective Cap from the cannula and insert the microscope body.
3.7.3 Connecting the BFMS

The BFMS body comes with two standard models; the first allows GCaMP6 fluorescence imagery while the second allows RCaMP2 fluorescence imagery.

![Diagram of BFMS Body Illumination System Connections](image)

Figure 3.14: BFMS Body Illumination System Connections
Microscope Driver and Light Source Installation

⚠️ If using the Ce:YAG Optical head, connect the Interlock connector plug to a Laser safety interlock circuit for safe use. See the application note Important Laser Safety Information for more information, or contact your institution’s laser safety officer.

1. Connect the 12 VDC power supply to the microscope driver 12 VDC power input.
2. Connect the driver and the light source.
   - If using an LED light source (such as for the GCaMP6 system (Fig. 3.14a)), connect the driver to the LED using the M8 connector.
   - If using a Ce:YAG Optical Head (such as for the RCaMP2 system (Fig. 3.14b)), see the Ce:YAG manual for details on driver and light source installation. Once complete, connect the Fluorescence microscope driver BNC Output to the Ce:YAG Driver BNC Input. The Ce:YAG Driver must be put in TTL mode to receive proper signals from the Microscope driver.
3. Connect the driver to the computer using the CAT5E Ethernet Cable.
4. Turn on the Fluorescence Microscope Driver.
3.7.4 Connecting the OSFM system

The indications in section 3.7, notably Fig. 3.13b and Fig. 3.13a are to be followed for the installation of the OSFM. As the OSFM uses two light sources, driver and light source installation is more elaborate. Both standard configurations use the Ce:YAG + LED/LD fiber Light Source to illuminate the sample.

Driver and light source installation

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1For details and specifications concerning the Ce:YAG + LED/LD Fiber Light Source, see their manual.
**GCaMP6 + NpHR3.0 configuration**

For the **GCaMP6 + NpHR3.0 configuration** (Fig. 3.15a), the Ce:YAG + LED optical head is used as the light source. The LED light source is used for excitation, while the Ce:YAG light source is used for opsin activation.

1. Connect the OTPG to the computer using a USB-A/USB-B cable.

2. In this configuration, the LED light source needs to be triggered by the Fluorescence microscope driver. Connect the driver BNC output to the Ce:YAG + LED driver LED BNC input.

3. Connect the Ce:YAG + LED driver Ce:YAG BNC input to an OTPG channel.

4. Connect the Fluorescence microscope driver Input BNC to an OTPG channel.

5. When installing in a head-fixed or freely-moving configuration, the rest of the system is connected as described in section 3.7.1 and 3.7.2.

**RCAmp2 + ChR2 configuration**

For the **RCAmp2 + ChR2 configuration** (Fig. 3.15b), the Ce:YAG + LD optical head is used as the light source. The LD light source is used for opsin activation, while the Ce:YAG light source is used for excitation.

1. Connect the OTPG to the computer using a USB-A/USB-B cable.

2. In this configuration, the Ce:YAG light source needs to be triggered by the Fluorescence microscope driver. Connect the driver BNC output to the Ce:YAG + LD driver Ce:YAG BNC input.

3. Connect the Ce:YAG + LD driver LD BNC input to an OTPG channel.

4. Connect the Fluorescence microscope driver Input BNC to an OTPG channel.

5. When installing in a head-fixed or freely-moving configuration, the rest of the system is connected as described in section 3.7.1 and 3.7.2.
4.1 Microscope

The Microscope module of the Doric Neuroscience Studio provides an interface to control our Fluorescence Microscope Driver. The module enables image acquisition and its export in 16 bit .tif or in .dor (hdf5-based) files. The TIF format can easily be read with any standard imaging software. Doric files can be read by the Doric Neuroscience Studio Image Analysis Module or using an HDF5 library. Despite the fact that the images are saved with a 16 bit pixel depth, the true image pixel depth is 10 bit, so pixel gray values are contained between 0 and 1020 counts.

Below is the user interface (Fig. 4.1) and a complete description of all the functions.

![Microscope Module Interface](image)

**Figure 4.1: Microscope Module Interface**

1. The **Image Box** (Fig. 4.1) displays images from the microscope and allows region of interest (ROI) drawing by clicking and dragging the mouse over the image.

   a) The **Sensor Tabs** (Fig. 4.1) display the sensors available to view. For multi-sensor microscopes, changing tabs allows you to see the image available to each.

   b) The microscope **Status** (Fig. 4.1) will indicate the current microscope state (Live/Stopped).

   c) The **Exposure** (in ms) (Fig. 4.1) indicates the exposure time of the microscope sensor.
d) The **FPS** (Frames Per Second) (Fig. 4.1) indicates the number of frames per second taken by the sensor.

e) The **Gain** (Fig. 4.1) indicates the electrical gain of the sensor.

f) The **Size** (Fig. 4.1) indicates the resolution of the sensor images (in Pixels x Pixels).

g) The **Bin** (Fig. 4.1) status indicates whether or not the sensor image is being binned (yes/no).

2. The **Capture** tab (Fig. 4.1) contains different image-capturing functions of the microscope.

![Figure 4.2: Capture Tab](image)

- a) The **Live** button (Fig. 4.2), when pressed, displays images from the microscope. These images are not saved.
- b) The **Snap** button (Fig. 4.2), when pressed, takes a snapshot of the current image and saves it in the requested directory with the desired name (**Saving Options**) as a single image.
- c) The **Album** button (Fig. 4.2), when pressed, acquires a snapshot and adds it to an album stack. The whole stack can be saved as one image stack.
- d) The **Record** button (Fig. 4.2), when pressed, acquires a continuous image stream, until **Stop** is pressed, and saves it in the requested directory with the desired name (**Saving Options**) as one image stack.

![Figure 4.3: Time Series Window](image)

- e) The **Time Series** button (Fig. 4.2), when pressed, opens the time series interface (Fig. 4.3).
  - i. The **Number of time points** (Fig. 4.3) defines the number of moments when a set of images will be recorded.
  - ii. The **Images per time point** (Fig. 4.3) defines the number of images taken in each set.
  - iii. The **Time interval between points** (Fig. 4.3), defined in ms, s and min, defines the duration between each image set. This duration always has a minimum value of **Exposure time/Images per time point**.
  - iv. The **Summary** box (Fig. 4.3) shows many values related to the time series, including the **Total images** recorded, the **Total memory** occupied by the full series, the **Time point duration** and the **Total duration** of the full series.
  - v. The **Progression bar** (Fig. 4.3) displays the progress (in %) of the time series.

![Figure 4.4: Saving Options Window](image)
f) The **Saving options** button opens the **Saving options window**.
   i. The **Filename** box (Fig. 4.4) is used to define the recorded file name.
   ii. The **...** button (Fig. 4.4) opens a window used to choose the save file location.
   iii. The **File type** drop-down menu (Fig. 4.4) is used to decide which file type is used to save images. For files larger than 4 GB, the .doric extension is recommended.
   iv. The **Index** box (Fig. 4.4) displays the current index that will be added to the filename.
   v. The **Target File** box (Fig. 4.4) shows the full location and name of the file being saved when an image sequence is recorded.

3. **Microscope settings** tab (Fig. 4.5) is used to set parameters related to the microscope recording images.

   ![Figure 4.5: Microscope Settings Tab](image)

   a) The **Exposure** box (Fig. 4.5) sets the exposure time of the sensor. The time can be set between 22 and 1000 ms.
   b) The **SENSOR** section (Fig. 4.5) defines characteristics for a single sensor and the associated excitation source. When a microscope used has multiple sensors, multiple **SENSOR** sections will be displayed, one for each sensor.
      i. The **Gain** box defines the sensor gain.
      ii. The **Illuminator power (%)** box defines the power emitted by the excitation light source. The light sources will be activated when the image acquisition is started. The maximum optical power (in mW) depends on the light source model.
   c) The **Working Distance Adjustment Slider** appears when an eFocus Miniature Fluorescence Microscope is connected to the driver. This slider will adjust the working distance from a central spot (0 μm) to the extremes (± 45 μm).
   d) The **External Trigger** button (Fig. 4.5) opens the **external trigger** window.

   ![Figure 4.6: External Trigger Settings Windows](image)

   i. The **Number of images per trig** box (Fig. 4.6a) defines the number of images acquired at each trigger pulse.
   ii. The **File name/location** (Fig. 4.6a) box displays the location where the images are saved as well as their file name.
   iii. The **Select...** (Fig. 4.6a) button allows the selection of the **File name/location**.
   iv. The **Progression bar** (Fig. 4.6a) displays the advancement of the triggered sequence (in %).
   v. The **Gated mode** checkbox (Fig. 4.6a) will change the external trigger to gated mode (Fig. 4.6b). In this mode, the microscope will only acquire images when a high TTL signal is received on the TRIG IN input.
e) The **Save configuration** button (Fig. 4.5) will save all **Microscope settings** and **Image settings** in a `.doric` format file.

f) The **Load configuration** button (Fig. 4.5) will load a selected configuration file.

g) The **Select mask file** button opens a window to select a mask file for the microscope used. This section only appears when a 2-color Fluorescence Microscope or an efocus Microscope is connected. The mask file currently loaded will be shown just above it. For more information on masks, see section ??.

Figure 4.7: Image Settings Tab

4. The **Image settings** tab (Fig. 4.1) is used to define certain settings related to the displayed and recorded images.

a) The **Crop Image** button (Fig. 4.7) allows a square to be drawn onto the image. When a new **Capture** sequence is activated, only the cropped region will be captured.

b) The **Reset crop** button (Fig. 4.7) resets the cropped image to its original state. The change will only appear when a new **Capture** sequence is activated.

c) The **Binning** drop-down list (Fig. 4.7) allows the binning of pixels. This reduces the number of pixels for smaller save file sizes.

5. The **View** tab (Fig. 4.1) is used to change viewing parameters of the sensor image. These changes will only appear on the sensor image when a new **Capture** sequence is started. Any adjustments made affect only the displayed image and not the recorded images.

Figure 4.8: View Tab

a) The **Zoom In/Zoom Out** buttons (Fig. 4.8) will increase/decrease the zoom of the sensor image.

b) The **Reset Zoom** button (Fig. 4.8) will reset the **Zoom factor** to 100%.

c) The **Zoom Factor** drop-down list (Fig. 4.8) allows the selection of a zoom factor from a pre-set list. The box will also display the current zoom if it was changed using different buttons.

d) The **Roi shape** drop-down list (Fig. 4.8) allows the selection of the shape used when drawing a **Region Of Interest** onto a sensor image. These shapes include **Freehand**, **Circle**, **Rectangle** and **Square**.

e) The **SENSOR** section (Fig. 4.8) is used to adjust contrast on a given sensor. When a microscope used has multiple sensors, multiple **SENSOR** sections will be displayed, one for each sensor.

i. The **Contrast** slider (Fig. 4.8) allows the adjustment of contrast from 0.1 to 5.

ii. The **Min/Max** sliders (Fig. 4.8) indicate the minimum/maximum number of counts displayed. Should the **Min** be above 0, all pixels with lower count will display a minimal value. Should the **Max** be below 1020, all pixels with a higher count will appear saturated.

iii. The **Auto contrast** slider button (Fig. 4.8) will active an automatic contrast adjustment algorithm.

iv. The **Reset** button resets contrast functions to their default settings.

f) The **Pseudocolor** drop-down list (Fig. 4.8) allow the sensor image color palette to be changed.

g) The **Show saturation** checkbox (Fig. 4.8) allows all saturation on the sensor image to be displayed in red. This function is only available if no pseudocolor is selected.
6. The **Ethernet** tab (Fig. 4.1) is used to define the ethernet connection used to connect the computer to the microscope driver.
   
   a) The **Refresh** button (Fig. 4.9) will identify any accessible IP addresses and add them to the drop-down list.
   
   b) The **Ethernet** drop-down list (Fig. 4.9) includes all IP addresses connected to an ethernet adapter. The proper one must be selected to properly connect the microscope.
   
   c) The **Pair** button (Fig. 4.9) connects the software to the driver.
   
   d) The **Remember** checkbox (Fig. 4.9) will keep the chosen IP address so that the chosen microscope driver will be connected automatically next time the software is opened.

7. The **ROI Manager** (Fig. 4.1) displays the live mean pixel intensity from a drawn ROI.
   
   a) The **Average Intensity in ROI** plot (Fig. 4.10) displays the average intensity over time inside a drawn ROI. CTRL + mouse wheel will adjust the x-axis zoom, while SHIFT + mouse wheel will adjust the y-axis zoom.
   
   b) The **Clear** button (Fig. 4.10) will clear any data displayed in the ROI manager and the ROI on the **Image Viewer**.
   
   c) The **Optimal Zoom** button (Fig. 4.10) sets the zoom factor on the plot to best display all data.
   
   d) The **Reset Zoom** button (Fig. 4.10) resets the zoom to its default setting.

To update the driver firmware version, connect the driver to the computer using a USB-A/USB-B male cable **only**. In the **Hardware** drop-down menu, select the **Update Microscope** to start the process. Turn the driver power ON.

### 4.2 Image Analyser

This module provides an easy way to extract relevant data from the images acquired by the Doric miniature fluorescence microscopes. The software loads images in .TIF format, implements image processing functions and an export tool...
to save the fluorescence data in .CSV or .doric format. This software does not replace standard analysis tools such as Matlab, ImageJ or Excel, but aims to offer useful processing algorithms developed for the microscope images. All the underlying algorithms are implemented from the OpenCV library. In this section, we will describe the different functions available, and how to use them.

![Image Analysis Module Interface](image1.png)

**Figure 4.11: Image Analysis Module Interface**

1. The **Image Viewer** displays the loaded images, allows navigation through the image stack and the drawing of regions of interest (ROIs) by clicking and dragging the mouse over the image. Multiple image sets can be opened, appearing as tabs in the upper left of the image box.

2. The **ROI Manager** displays the different ROI parameters and traces the mean signal intensity for each image.

3. The **Function Toolbar** contains all the buttons and functions accessible.

### 4.2.1 Function Toolbar

1. The **File tab** (Fig. 4.12) is used to save/load data.

![File Tab](image2.png)

**Figure 4.12: File Tab**

- The **Load Images** function loads a square, 16 bit TIF file or multipage file.
- The **Save Images** function saves the current image tab to a 16 bit TIF multipage file.
- The **Export Traces** function saves the average fluorescence intensity values for each ROI of the current tab to a .CSV or .doric file.
2. The **ROI tab** (Fig. 4.13) is used to save/load data relating to regions of interest drawn on an image.

![ROI Tab](image)

**Figure 4.13: ROI Tab**

- The **Load ROIs** function loads `.CSV` file containing informations about the saved ROIs.
- The **Save ROIs** function saves the current ROIs information to a `.CSV` file.
- The **Clear All** button clears all ROIs.
- The **ROI shape** function is a drop-down list that allows the selection of the ROI shape. These include **Freehand**, **Circle**, **Rectangle** and **Square**.

3. The **View tab** (Fig. 4.14) is used to manipulate the appearance of an image without changing base data.

![View Tab](image)

**Figure 4.14: View Tab**

- The **Contrast** function applies a different luminance response curve (gamma). See section 4.2.3 for details.
- The **Min** function applies a lower threshold with the cut-off value defined by the slider. See section 4.2.3 for details.
- The **Max** function applies an upper threshold with the cut-off value defined by the slider. See section 4.2.3 for details.
- The **Autocontrast** function directly applies the `equalizeHist` function of the OpenCV library.
- The **Reset** function returns the contrast and range values to their default.
- The **Image Info** button displays the image information window.
- The **Pseudocolor** function is a drop-down list for selecting alternate coloring schemes for the images presented.

4. The **Processing tab** (Fig. 4.15) is used to process the image data.

![Processing Tab](image)

**Figure 4.15: Processing Tab**
• The **Align Images** function aligns the image stack to the user-defined key frame. See section 4.2.3 for computational details. Selecting this button will open the **Align Images** window (Fig. 4.16). By selecting the **Save Alignment Values** checkbox, the image alignment values will be preserved when saving the processed images. There are 4 different methods available.

![Select Alignment Method](image)

**Figure 4.16: Align Images Window**

- The **First Frame Of Current Image Set** method uses the first image in the set to align the rest.
- The **Select Frame From Current Image Set** method allows the selection of a single image in the set to use for alignment of all other frames.
- The **Select Other Image Set And Frame** method aligns the current set using data from a different image set.
- The **Select From Alignment Value File** method uses a previously-defined alignment for another image set. This method is most valuable when trying to align images from the 2-color fluorescence microscope, to align one color channel using the data from the other.

• The **Remove Background** function removes the average value of a selected ROI from all images in the stack.
• The **ΔF/F₀** function calculates the normalized fluorescence variation of the images and displays the results in a new tab. When selected, see section 4.2.3 for details.
• The **Find Cells** function detects the cells and creates the ROI automatically. See section 4.2.3 for details.
• The **Stack Projection** function projects all movie frames to a single frame using the method selected in the Settings dialog. See section 4.2.3 for details.
• The **Batch Processing** function opens the **Batch Processing Window** (Fig. 4.17). This allows the processing of large datasets in sequential order, without needing to activate each individual function.
a) The **Available processes** box lists all processes available. Processes on the list will be greyed out if the work-flow order prevents them from being used. Each process has a number of parameters that are identical to those used outside of batch processing.

   - The **Align Images** process aligns the image stack to the user-defined key frame. See section 4.2.3 for computational details.
   - The $\Delta F/F_0$ process calculates the normalized fluorescence variation of the images and displays the results in a new tab. See section 4.2.3 for details.
   - The **Stack Projection** process projects all image frames to a single frame using the method selected in the Settings dialog. See section 4.2.3 for details.
   - The **Find Cells** process detects the cells and creates the ROI automatically. See section 4.2.3 for details.
   - The **Convert** process is used to convert an image stack to .doric or .tif format.

b) The **Workflow** box displays the order in which image processing actions will be taken.

c) The **Choose a Folder** button allows the selection of a folder to save batch processing results.

d) The **File Type** list is used to defined the file extension used when the images are saved.

e) The **Save Intermediate files** option will save intermediary files in the image processing process alongside the completed files.

---

**Figure 4.17: Batch Processing Window**
4.2.2 ROI Manager

The ROI manager extracts average intensity of a defined section of the image over an entire image stack. There is no limit to the number of ROI allowed per image stack.

1. The Image Viewer contains the image stack and the ROI, numbered according to the order they where set. The ROI can be saved independently from the image stack on the ROI toolbar. The ROI are drawn directly on the Image Viewer in a freehand manner.

2. The Intensity Plot panel shows the plot of average intensity as a function of the frame index. The Y-axis represent the average count of all the pixels of the ROI. It is separated in Overview and Detailed tabs.
   - The Overview tab displays all the traces on the same graph, on the same scale (see Fig. 4.19a).
   - The Detailed tab displays each trace on a separate graph, allowing for precise intensity measurements (see Fig. 4.19b).

3. The ROI Data list shows the parameters defining each ROI. Selected items will be displayed in orange on the Image Viewer and in the Overview graph.
   - The ID shows the order of the ROI (starting at 0).
   - The Area shows the area (in pixels) contained in the ROI.
4.2.3 Algorithms

Contrast

The contrast adjustment applies the following operation to each pixel of the image: \( V_{\text{out}} = AV_{\text{in}}^\gamma \), where \( V_{\text{out}} \) is the corrected pixel value, \( A = 1 \), \( V_{\text{in}} \) is the initial pixel value, and \( \gamma \) is the value as selected by the contrast slider.

Min and Max ranges

When the values of the display range are other than the default \( \text{min} = 0 \) and \( \text{max} = 1020 \), the following operation is applied to each pixel: \( V_{\text{out}} = 1020 \times (V_{\text{in}} - \text{min})/\text{(max} - \text{min}) \), where \( V_{\text{out}} \) is the corrected pixel value, \( V_{\text{in}} \) is the initial pixel value, \( \text{min} \) and \( \text{max} \) are respectively the minimum and maximum slider values.

Image Alignment

The algorithm is inspired from Manuel Guizar-Sicairos, Samuel T. Thurman, and James R. Fienup, *Efficient subpixel image registration algorithms*, Opt. Lett. 33, 156-158 (2008). The basic idea is to obtain an initial estimate of the crosscorrelation peak by a Fourier transform and then refine the shift estimation by upsampling the Fourier transform only in a small neighborhood of that estimate by means of a matrix-multiply Fourier transform. With this procedure, all the image points are used to compute the upsampled crosscorrelation. In order to increase the precision of the algorithm, we use the laplacian of the images as inputs, instead of using the raw images. Briefly, the algorithm applies the following steps:

1. Calculate gaussian blur of the reference image with window of size 39 to smooth high frequency noise.
2. Calculate the laplacian of the blurred reference image.
3. Use the absolute values as the final reference image.
4. Reproduce steps 1 to 4 for the following image.
5. Calculate the 2D Fourier transform of the reference and the target image.
6. Multiply both images.
7. Calculate the inverse Fourier transform of the product image.
8. Get the position of the maximum correlation peak.
9. Create an upsample array around the maximum correlation peak to refine the shift calculations.
10. Calculate the Fourier transform of the larger array.
11. Do the matrix multiplication.
12. Locate the maximum correlation and map it back to the original space.

\( \Delta F/F_0 \)

The algorithm calculates a standard \( \Delta F/F_0 \) with \( F_0 \) corresponding to the temporal average intensity, with an optional preprocessing step to remove the illumination variation artefacts. In order to properly calculate the \( \Delta F/F_0 \), the algorithm uses a dark frame to account for the sensor electronic offset. Calculating the \( \Delta F/F_0 \) without subtracting the offset will lead to artificially lower values. To record a dark frame, set the microscope driver to the desired exposure and gain, the LED power to zero and take a snapshot. Before calculating the \( F_0 \), the average temporal variations can be compensated to get a flat temporal average profile (Fig. 4.20). Keep in mind that removing the average temporal profile can also remove global activity patterns.
Briefly, the algorithm applies the following steps:

1. Calculate the average image intensity as a function of time ($C$).

2. If the global variation removal option is selected, apply the following correction to each image: 
   \[ I_{\text{out}} = (I_{\text{in}} - I_{\text{dark}}) \times \frac{\text{mean}(C - I_{\text{dark}})}{(C - I_{\text{dark}})} \]
   where $I_{\text{out}}$ is the LED illumination corrected image, $I_{\text{in}}$ the input image and $C$ is the average temporal trace.

3. Calculate $F_0$ as the average projection of the movie.

4. Calculate the relative change $R(t)$ of fluorescence signal 
   \[ R(t) = \frac{F(t) - F_0}{F_0} \].

**Find Cells**

The algorithm is inspired by Eran A. Mukamel, Axel Nimmerjahn and Mark J. Schnitzer, *Automated analysis of cellular signals from large-scale calcium imaging data*, Neuron 63(6), 747-760 (2009). The basic idea is to use a principal component analysis (PCA) as input of an independent component analysis (ICA) to separate the different temporal signals contained in the movie. This method is used as a starting point to determine the position of the different active cells. It is coupled with a segmentation routine optimized for reducing the false positives. The *Find Cells* algorithm uses user-defined boundaries shown in Fig. 4.21. The first parameter is an estimate of the number of cell present in the movie. By design, it must be lower than the number of frames minus five. The next parameters are the smallest and biggest object diameter in microns. These values are used to filter the object found by the PCA/ICA.

Briefly, the algorithm applies the following steps:

1. Calculate and remove the spatiotemporal average from the movie, as the PCA/ICA algorithm requires zero-mean data.

2. Run OpenCV PCA algorithm on the centered data.

3. Normalize data by standard variation.

4. Calculate ICA with PCA as input data.

5. Apply segmentation to each ICA found.

6. Filter contours found at the previous step using user-defined boundaries.
**Stack Projection**

This function can be used to help for ROI drawing. It calculates a temporal projection using the user-defined method (see Fig. 4.22).

![Stack Projection Settings](image.jpg)

Figure 4.22: Stack Projection Settings

- **Maximum**: the output is the maximum value found in all frames for each pixel.
- **Average**: the output is the mean value of all frames for each pixel.
- **Sum**: the output is the sum of all frames for each pixel.
- **Minimum**: the output is the minimum value found in all frames for each pixel.
Specifications

Table 5.1: Single-color Fluorescence Microscope Body Specifications

<table>
<thead>
<tr>
<th>SPECIFICATION</th>
<th>Microscope Body</th>
<th>Model S</th>
<th>Model L</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass without cables (g)</td>
<td></td>
<td>2.2</td>
<td></td>
</tr>
<tr>
<td>Dimensions without cables in mm (W x L x H)</td>
<td></td>
<td>8.8 x 13.9 x 16.6</td>
<td></td>
</tr>
<tr>
<td>Frame rate (fps)</td>
<td></td>
<td>49</td>
<td></td>
</tr>
<tr>
<td>Objective lens NA</td>
<td></td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>FOV at image plane (pixel)</td>
<td></td>
<td>630 x 630</td>
<td></td>
</tr>
<tr>
<td>FOV at object plane (µm)</td>
<td></td>
<td>700 x 700</td>
<td>350 x 350</td>
</tr>
<tr>
<td>Lens magnification</td>
<td></td>
<td>3.3x</td>
<td>6x</td>
</tr>
</tbody>
</table>

Table 5.2: Single-color Fluorescence Microscope Body Excitation and Detection Spectra

<table>
<thead>
<tr>
<th>Microscope Bodies</th>
<th>SPECTRUM (nm)</th>
<th>Excitation</th>
<th>Detection</th>
<th>Opsin activation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basic</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GCaMP6</td>
<td>445/62</td>
<td>525/40</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>RCaMP2</td>
<td>549/15</td>
<td>609/57</td>
<td>—</td>
<td></td>
</tr>
<tr>
<td>OSFM</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>GCaMP6 + NpHR3.0</td>
<td>445/62</td>
<td>525/40</td>
<td>616/76</td>
<td></td>
</tr>
<tr>
<td>RCaMP2 + ChR2</td>
<td>549/15</td>
<td>609/57</td>
<td>Compatible with 450, 473, 488</td>
<td></td>
</tr>
</tbody>
</table>

Table 5.3: Model S Cannula general specifications

<table>
<thead>
<tr>
<th>SPECIFICATIONS</th>
<th>VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnification</td>
<td>2 x</td>
</tr>
<tr>
<td>Working distance</td>
<td>2.4 mm</td>
</tr>
</tbody>
</table>

Table 5.4: Model L Cannula general specifications

<table>
<thead>
<tr>
<th>SPECIFICATIONS</th>
<th>VALUE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lens diameter</td>
<td>500 µm</td>
</tr>
<tr>
<td>Working distance</td>
<td>80 µm</td>
</tr>
</tbody>
</table>

1Center wavelength/bandwidth
Table 5.5: Fluorescence Microscope Driver general specifications

<table>
<thead>
<tr>
<th>SPECIFICATIONS</th>
<th>VALUE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power supply</td>
<td>110 - 240 VAC, 50 - 60 Hz</td>
<td></td>
</tr>
<tr>
<td>DC Power supply</td>
<td>12 VDC</td>
<td></td>
</tr>
<tr>
<td>Dimensions</td>
<td>186 x 90 x 77 mm</td>
<td>Including connectors</td>
</tr>
<tr>
<td>Data link</td>
<td>Gigabit ethernet</td>
<td></td>
</tr>
<tr>
<td>TTL input voltage</td>
<td>0 to +5 V</td>
<td></td>
</tr>
<tr>
<td>LED Analog input voltage</td>
<td>400 mV light source current</td>
<td>Standard 1 A LED Driver</td>
</tr>
<tr>
<td></td>
<td>40 mA/V light source current</td>
<td>Low power mode enabled</td>
</tr>
<tr>
<td>LED BNC output voltage</td>
<td>2.5 V/A</td>
<td></td>
</tr>
<tr>
<td>LED Maximum output current range</td>
<td>200, 2000 mA</td>
<td></td>
</tr>
<tr>
<td>LED Maximum forward voltage</td>
<td>7 V</td>
<td></td>
</tr>
<tr>
<td>LED Minimum output current</td>
<td>2.5 mA</td>
<td>Low power mode enabled</td>
</tr>
<tr>
<td>LED Rise/Fall time</td>
<td>&lt;10 μs</td>
<td>Typical</td>
</tr>
<tr>
<td>LED Connector</td>
<td>M8</td>
<td>See Figure 5.1</td>
</tr>
</tbody>
</table>

Figure 5.1: M8 Female Pinout (Microscope Driver)

Table 5.6: Fluorescence Microscope Driver Software specifications

<table>
<thead>
<tr>
<th>SPECIFICATIONS</th>
<th>VALUE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Current adjustment steps</td>
<td>1 mA</td>
<td>Internal complex mode: 0.000054 Hz</td>
</tr>
<tr>
<td>Modulation minimum frequency</td>
<td>0.01 Hz</td>
<td>Internal complex mode: 0.0054 Hz</td>
</tr>
<tr>
<td>Modulation maximum frequency</td>
<td>50 kHz</td>
<td>-3 dB attenuation</td>
</tr>
<tr>
<td>Minimum ON or OFF time</td>
<td>0.005 ms</td>
<td>Internal complex mode: 2 ms</td>
</tr>
<tr>
<td>Maximum ON or OFF time</td>
<td>100 s</td>
<td>Internal complex mode: 5 h</td>
</tr>
<tr>
<td>Maximum number of pulses per sequence</td>
<td>16.68 millions</td>
<td>Internal complex mode: 65 535</td>
</tr>
<tr>
<td>Maximum number of sequences</td>
<td>4.2 billions</td>
<td>Internal complex mode: 65 535</td>
</tr>
<tr>
<td>Minimum step increments</td>
<td>39 μsec</td>
<td>Internal complex mode only</td>
</tr>
<tr>
<td>Number of steps per period</td>
<td>128²</td>
<td>Internal complex mode only</td>
</tr>
<tr>
<td>Scope Acquisition speed</td>
<td>10 kS/s</td>
<td>Single channel</td>
</tr>
</tbody>
</table>

²For all operation modes, except the internal complex mode
Table 5.7: Connectorized LEDs general specifications

<table>
<thead>
<tr>
<th>SPECIFICATION</th>
<th>VALUE</th>
<th>NOTE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Input current</td>
<td>Min: 0 mA - Max: 1000 mA</td>
<td>Recommended 700 mA</td>
</tr>
<tr>
<td>Forward voltage</td>
<td>3.0 - 4.0 V Typical</td>
<td></td>
</tr>
<tr>
<td>Dimensions</td>
<td>63 x 59 x 21 mm³</td>
<td>Without cable, with baseplate</td>
</tr>
<tr>
<td>Mass</td>
<td>~100 g</td>
<td></td>
</tr>
<tr>
<td>CLED cable connector</td>
<td>M8-4pins - Male</td>
<td>See Figure 5.2</td>
</tr>
<tr>
<td>Output NA</td>
<td>0.55</td>
<td>NA of up to 0.63 NA will slightly increase power;</td>
</tr>
<tr>
<td>Output optical fiber core diameter</td>
<td>&lt;960 μm</td>
<td>Power scales up to this core diameter</td>
</tr>
</tbody>
</table>

![Figure 5.2: M8 Male Pinout (CLED)]

Table 5.8: Doric Neuroscience Studio Hardware Requirements

<table>
<thead>
<tr>
<th>SPECIFICATIONS</th>
<th>VALUE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Operating System</td>
<td>Windows 7, 8, 10</td>
<td>64-bit</td>
</tr>
<tr>
<td>Memory (Minimum/Recommended)</td>
<td>4 GB/16 GB</td>
<td></td>
</tr>
<tr>
<td>Processor Speed (Minimum/Recommended)</td>
<td>2 Ghz Quad-Core i5/ 3.46 Ghz Eight-core i7</td>
<td></td>
</tr>
<tr>
<td>Hard Drive</td>
<td>500 MB</td>
<td></td>
</tr>
</tbody>
</table>
Annex 1: Cleaning and Handling

6.1 Important Handling Information

*Warning*: Handle the microscope and the cannula with care.

Miniature fluorescence microscopes are composed of sensitive electronic components and should always be handled with care. When they are not used, the microscope body with its cannula should be stored in a closed, dust-free environment. Some microscope components must be handled with extra care:

- **Electrical cable**: *Do not twist or pull on the cable.* This cable is pigtailed to the CMOS sensor and cannot be easily replaced.

- **Relay lens**: The cannula lens is made of glass and is unprotected. *Abrasive materials can scratch the surface* and reduce the image quality.

The microscope bodies and the cannula lenses are made of glass, metal, plastic and the contact with organic tissues or liquids, like blood or salted water solution is not recommended. If the microscope comes in contact with these substances, clean the optics (section 6.2) to avoid the apparition of stains.

6.2 Cleaning Optics

The microscope objective lens should be cleaned before each use. The procedure explained here can also be used to clean the cannula relay lenses.

- Turn the driver OFF.

- **Wear gloves to manipulate the microscope**. Finger oil can stain the glass and is often hard to remove properly.
- Use isopropyl alcohol on a cotton swab to gently clean the lens.

- **Do not blow on the optics**. Saliva particles will often stain the surface. Larger dust particles can be removed using a dust-free blower before cleaning with a cotton swab.
Annex 2: Troubleshooting

7.0.1 Software

How to make the Doric Neuroscience Studio detect the Microscope Driver?

1. Ensure that the Microscope Driver is plugged into the computer using an Ethernet cable.
2. Ensure that the each Electrical Cable Connector is plugged into the appropriate device. The Microscope Driver must be linked to the Microscope.
3. Ensure that the IP address is static (see section 3.3.1)
4. Ensure the Jumbo Frames are activated (see section 3.3.1)
5. Windows Firewall can prevent communication. To ensure the communication is not being blocked, open the Windows Firewall configuration window, then click on Allow an app through the firewall. From there, select the Change Settings button, find the Doric Neuroscience Studio and check the Private and Public checkboxes.
6. In the Network & Sharing Center, check the Ethernet connection; it should indicated Unidentified Network. If Network Cable Unplugged is shown despite the Ethernet cable being plugged in and the driver being turned on, disable and re-enable the Ethernet connection.
7. Ensure Network & sharing is properly configured at 1 Gbps by double-clicking the Ethernet connection and checking the Speed.
8. When the 1-color Microscope Driver is activated, the On/Off Switch should blink blue while initializing. If the light is sustained without any blinking when first turned on, restart the Microscope Driver.
9. Certain Intel Ethernet cards must be activated in Slave Mode to function. This mode can be found in the same menu as the Jumbo Frames (see section 3.3.2).

How can I stop the software lagging and/or dropping frames?\textsuperscript{1}

1. Deactivate all internet using programs that can conflict with the Doric Neuroscience Studio (IE Skype, Firewall, etc.)
2. Use a computer with the recommended specifications:
   - Operating System: Windows 10
   - CPU: Quad Core i7 3.46 GHz
   - RAM: 16 Gb
   - Dedicated Graphics Card
   - Desktop computer recommended

\textsuperscript{1}Dropped frames are black frames that occur when an image is lost in communication. They can easily be spotted in the Average Intensity in ROI trace if the value descends to 0.
How can I visualize recorded frames?

1. All images will appear black in Window Image Preview/Traditional Image Viewers as they are special 16 bit .tif files. Use dedicated software such as the Doric Neuroscience Studio Image Analyzer or ImageJ.
2. Due to library usage conflicts, the Dell Backup & Recovery application interferes with the loading of images in the Doric Neuroscience Studio. Uninstall the Dell Backup & Recovery application from the computer.

Can I use a USB to Ethernet adapter to connect the driver?

1. The microscope driver must be connected to a computer ethernet port.
2. Should a USB to Ethernet adapter be used for other function, such as internet access, the adapter must be disabled during the first initialization of the microscope.

7.0.2 Hardware

How do I prevent instability in the Assisted Opto-electric Rotary Joint?

1. Ensure the optical fiber Patch Cord is of equal or shorter length than the microscope Electrical Cable when connector to the Assisted Opto-Electric Rotary Joint. Even if the cable is looped, the distance from rotary joint to patch-cord connector should be shorter than the length of the electrical cable.

How do I prevent the Cannula turning in the Protrusion Adjustment Ring?

1. These two components are meant to be glued together after installation. If they have not been glued during installation, add a drop of quick-drying glue on the border between the Cannula and Protrusion Adjustment Ring.

How to protect the Cannula when the Input Protective Cap does not stay inside?

1. Fill the interior of the Cannula with KWIK-CAST (WPI) to act as a cap. After removal of the dried sealant, clean the Rod Lens outer surface using a cotton swab lightly dipped in isopropyl alcohol.

How do I stabilize the Cannula-Microscope Body connection?

1. Ensure that the Microscope Barrel is sufficiently loose (see Microscope Surgical Procedure Application Note Section 2.1 or the Instructional Video).

How to insert the Microscope Body into the Cannula with minimal force?

1. Ensure that the Microscope Clamps are sufficiently loose (screw in the barrel).
2. Ensure that the Cannula and Microscope Body are properly aligned.
3. Verify the Cannula installation instructions in Section 2.1 of the Microscope Surgical Procedure Application Note.

7.0.3 Biology

Why can't I see any individual cells?

1. It generally takes 3 to 8 weeks before the area surrounding the tip of the Rod Lens has healed enough for sharp imaging of neurons.
8.1 Warranty

This product is under warranty for a period of 12 months. Contact Doric Lenses for return instructions. This warranty will not be applicable if the unit is damaged or needs to be repaired as a result of improper use or operation outside the conditions stated in this manual.

8.2 Contact us

For any questions or comments, do not hesitate to contact us by:

Phone  1-418-877-5600

Email  sales@doricleses.com