Fluorescence Imaging with Olympus IX81 Microscope
Last updated April 26, 2022

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You must read the MIC Facility Manual before training. It covers lab and laser safety, training policy, scheduling, and biosafety requirements.

USERS MUST RECEIVE TRAINING FROM MIC STAFF. Getting trained by another user is neither sufficient nor permissible for independent operation of the microscope. In other words, without the official training, you can be in the room and watch, but you are not allowed to touch the microscope.

ACKNOWLEDGMENT POLICIES
The use of the facility must be properly acknowledged in any publication (including web pages). You can use the following statement:
“The use of the Microscopy and Imaging Center facility at Texas A&M University is acknowledged. Research Resource ID (RRID): SCR_022128. The Olympus FV1000 microscope acquisition was supported by the Office of the Vice President for Research at Texas A&M University.”

Users are also required to file a copy of any relevant publication containing the acknowledgment with the MCF administrative office.

Notes:
**Microscope slides and other types of sample carriers:**

Microscope objectives are designed to view the sample through a thin glass window, 0.17 mm thick (this is the #1.5 coverglass). On the inverted microscope, the bottom of the sample chamber must be that thin and optically clear. If your samples are in a standard plastic Petri dish or a regular multi-well culture plate, you will not be able to get the best image quality with the 10x objective, and you will not be able to focus at all with the other objectives.

There are two long-working dry objectives, 20x/0.4 and 60x/0.7, that may be used with these samples. These objectives are NOT installed on the microscope. Please consult MIC staff for specific instructions.

**Recommended types of samples:**

- Slides with a 0.17 mm thick coverglass, sealed with nail polish or other suitable sealer that has dried. Please do not get wet nail polish on the objectives, it may get expensive.
- Chambered slides where the culture chamber was removed after staining, suitable mounting medium and a coverglass was applied.
- Chambered slides with optically clear, thin bottom (e.g. Ibidi microslides, Greiner CELLview cell culture Slides)
- Imaging-type 96-well plates with optically clear, thin bottom – these MUST be specifically the type where the bottom is minimally recessed relative to the skirt of the plate. **CONSULT THE MIC STAFF IF YOU ARE NOT SURE WHICH PLATE YOU NEED.**

**NOT Recommended:**

- LabTek II chambered coverglass – the chambers are too close to the end of the coverglass, pose a risk of damaging microscope objectives by the slide holder.
- 95 well plates with deeply recessed bottom – They create a risk of damaging microscope objectives. The edge rows and columns CANNOT be imaged,
- **If you use these plates or chambers, when the objective is in focus and you move to the well on the end or edge of the plate, the metal sample holder will hit the objective and damage it. You and your PI will be responsible for objective repair/replacement ($$$).**
Lasers:
Even though no lasers are used for wide-field fluorescence imaging, the microscope is a part of the laser scanning confocal system and all users MUST take the online Laser Safety course, located at https://ehsdtraining.tamu.edu, before their first training session.
Standard operating procedure (SOP) and a copy of Texas Administrative Code §289.301 (Registration and Radiation Safety Requirements for Lasers) are located in the confocal microscope room.
The last user must be able to properly shut down the entire system, including the lasers if they were left on by the previous user.

Mercury Lamp Precautions
- The lamp emits strong UV and visible radiation. Do not look into the source or disassemble the lamp housing.
- Do not turn the lamp on if the lamp usage counter reached its expected lifetime (300hr)!
  - After turned on, it takes ~ 15 min for the lamp to reach full brightness.
  - Lamp must be ON for at least 30 min before it can be switched OFF.
- After the lamp has been switched OFF, it must cool down (~15 min) before it may be switched ON again.
Biosafety requirements and rules for work in the MIC

The Olympus microscope room has been approved as BL-1 space. In order to be able to bring the active BL-1 material to the MIC, the MIC facility and room number MUST be listed in the investigator’s IBC permit, in Section F, Agent use and Storage Locations. The investigator is required to send a copy of the IBC permit listing the relevant MIC lab and the BL1 organisms, to MIC office (microscopy@tamu.edu) BEFORE bringing the BL-1 samples.

The Olympus microscope room is not approved for BL-2 or higher samples.

### F. Agent use and storage locations.

<table>
<thead>
<tr>
<th>Location ID</th>
<th>Campus</th>
<th>Building Number</th>
<th>Room Number</th>
<th>Room Use</th>
<th>Current Bio-safety Level</th>
<th>Shared Lab?</th>
<th>Other PIs</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Texas A&amp;M</td>
<td>1530</td>
<td>1117</td>
<td>Cell and tissue culture</td>
<td>BSL-2</td>
<td>Yes</td>
<td>*</td>
</tr>
<tr>
<td>3</td>
<td>Texas A&amp;M</td>
<td>1530</td>
<td>1118</td>
<td>Confocal microscopy</td>
<td>BSL-1</td>
<td>Yes</td>
<td>*</td>
</tr>
<tr>
<td>4</td>
<td>Texas A&amp;M</td>
<td>1530</td>
<td>1121</td>
<td>Laboratory</td>
<td>BSL-1</td>
<td>Yes</td>
<td>*</td>
</tr>
<tr>
<td>5</td>
<td>Texas A&amp;M</td>
<td>1530</td>
<td>1120</td>
<td>Microscopy room</td>
<td>BSL-1</td>
<td>Yes</td>
<td>*</td>
</tr>
</tbody>
</table>

* The list of other PIs is maintained by the MIC

All users of rooms listed in table F must follow the rules. This applies even to those users that do not work with samples requiring IBC permit:

**I) For users that DO NOT work with BL-1 agents:**

- Closed toe shoes are required in the confocal microscope room. Upon exiting the microscope room, users are required to wash hands. The sink in the Bioprep lab (Rm. 1121) or in the Culture room (Rm. 1117) may be used.

**II) For users working with active BL1-agents:**

- The MIC facility and room number MUST be listed in the investigator’s IBC permit, in *Section F, Agent use and Storage Locations*; The PI is required to send a copy of the IBC permit, listing the relevant MIC lab and the BL-1 organisms, to microscopy@tamu.edu BEFORE the BL-1 samples can be brought to the MIC. Without the MIC facility listed in the PI’s permit, no BL-1 work is allowed in the MIC.
- Samples being brought to the MIC must be contained, in accordance with the operating procedure in investigator’s IBC permit, to prevent spills during transport
- Closed toe shoes must be worn
- The use of gloves should be restricted to only handling the sample to avoid contaminating general work area. Touching the microscope or the control computer keyboard and mouse with gloved hands should be avoided. Gloves may not be worn outside of the rooms listed in table F.
- The user must ensure proper disposal of the samples - taking all samples as well as contaminated microscope slides, etc., back to his/her laboratory for disposal. Samples may not be put in regular trash in the MIC.
- Any spills and contaminations must be cleaned up, decontaminated by the user and the emergency contact person notified.
- At the end of the work session, the work area, including the microscope stage must be wiped clean using a disinfectant (70% ethanol or disinf ecting wipes).
- Upon exiting the BL-1 work area, users must wash hands; the sink is the Bioprep lab (Rm. 1121).
Biological Spill Response: BL1 Laboratory

The following procedures are provided as a guideline for biohazard spill clean up in a BL1 laboratory. Although the biological material in a BL1 level spill should not be a significant health hazard, you have the obligation to minimize the release of recombinant organisms and biohazardous material from the laboratory.

In the event of a spill:

- If a biohazardous material spills on you, remove any contaminated clothing and wash any exposed body parts.
- If a biohazardous material gets in your eyes, flush at the nearest eyewash station.
- If the spill area is large or in a common use area, mark, label, or otherwise denote the area so others may avoid it.
- Using materials from your spill kit:
  - Put on gloves, lab coat, and eye protection.
  - Cover the spill with absorbent material.
  - Pour disinfectant over the entire area. Allow the area to soak for 30 minutes.
- If warranted, contact the principal investigator, assess the magnitude of the spill, and formulate further plans of action.
- Safely pick up any broken glass with forceps or sweep into a dustpan, and dispose the residue into a broken glass/sharps container.
- Place spill materials into an autoclave bag.
- Make sure area is cleaned and disinfected thoroughly.
- Soak any contaminated clothes and shoes in a tray with disinfectant.
- If the spill is greater than 25 ml or contains recombinant DNA, report the spill to the Office of Research Compliance and Biosafety at 979.862.4549.
STARTUP PROCEDURE

1. Check in the microscope room if the instrument seems ready for use
2. Start your iLab kiosk session
3. Fill out the log book, turn ON the following components
4. Mercury Lamp Power Supply. If the time counter exceeds 300 hours, do not turn it on, ask MIC Staff to replace the Hg lamp
5. Microscope Control Box
6. Prior Optiscan stage controller
7. Photometrics Camera
8. Computer
9. Log in the computer: User = Fluoview Password =
10. Open the software:
    “Micromanager”. When prompted for the name of the configuration file, choose the default “MMConfig_MIC_Prime.cfg” or select your own config file, located in the D:/User_Data/Camera Imaging/
11. The software now tries to take control of the microscope and other hardware. If you get error messages, try re-loading the hardware configuration (Tools - Load Hardware Configuration). If the Photometrics Prime camera was not detected, you will have to turn the camera on and wait ~30 s while the camera is initialized – when the light next to the power switch on the camera stops blinking, the camera is ready, you can re-load the HW configuration.
12. You can change/modify the configuration file as needed and save it with a different name (do NOT overwrite the default MIC config file).
Using the Olympus IX81 microscope

Adjusting the oculars for optimal viewing
1. Adjust the interpupillary distance of the binoculars so that you can see with both eyes
2. Set both oculars to “0” focal correction (see scale on the side of the ocular tube)
3. Use the microscope focus knob to bring the specimen in focus for your left eye.
4. Now, take your hand off the focus knob. Turning the ocular focus correction, bring the specimen in focus for your right eye.

Focusing
Turning the microscope focus wheel towards you will move the objectives upwards, closer towards the specimen (= focus into the specimen). Turning the focus knob away from you will lower the objective, i.e., move it away from the specimen.
**Setting up Köhler illumination**

Köhler illumination produces an even field of illumination and also avoid projecting images of dust onto the specimen.

1. Place specimen on the stage, and switch to the DIC filter (in software). Make sure the microscope is set to visual observation, not the camera port (top left button on the front of the microscope).
2. Focus the specimen (Fig. 2A). From now on, do not touch the focus knob.
3. Completely open the Condenser Aperture and completely close the Field Aperture (Fig. 2B).
4. Focus the edges of the Field Aperture by cranking the condenser up or down using the condenser focus knob. Double check that the specimen and the Field Aperture are in focus simultaneously (Fig. 2C).
5. If the Field Diaphragm is off center, use the two adjusting screws to center the condenser so that the diaphragm image is exactly centered in the field of view (Fig. 2D).
6. Open the Field Diaphragm until its shadow just disappears from the field of view (Fig. 2E). Opening the Field Diaphragm more than that causes extra glare and decreases image contrast.

**Figure 2**: Köhler illumination – Focusing and centering the condenser Field Diaphragm

- A: Focus the specimen
- B: Field Diaphragm closed down, not focused, not centered
- C: Field Diaphragm focused, not centered
- D: Field Diaphragm focused and centered
- E: Field Diaphragm opened just enough to disappear from view
**Differential Interference Contrast (DIC, a.k.a. Nomarski) observations**

Set up Kohler illumination first.

Make sure the polarizer in the condenser is engaged

Check that the Filter set selected in the software is DIC.

The slider with Nomarski prism below the objective needs to be in the optical path (pushed in). The “hockey stick” handle on the slider should be set according to the objective used: pulled out for dry objectives, pushed in for immersion objectives. *(Explanation: This handle controls the distance of the Nomarski prism from the flange of the objective. Dry and immersion objectives have their back focal plane at a different position. The Nomarski prism must be close to the BFP for best performance.)*

If the objectives are switched using the software’s pre-set group, the correct Nomarski prism is automatically engaged (and the objective turret retracts before turning to prevent damage or contamination of the objectives).

*If you use the buttons on the keypad and on the microscope to switch objectives, you have to manually switch to the correct Nomarski prism in the condenser turret.: Each objective requires a different condenser prism for DIC to work. See table 1 below. Use the control keypad to turn the condenser turret (two buttons on the keypad, labeled “GFP” and “Cy3”).*

Table 1: Condenser turret positions for DIC observation with individual objectives

<table>
<thead>
<tr>
<th>Objective</th>
<th>Condenser turret position</th>
<th>“hockey stick” handle position on the DIC objective prism</th>
</tr>
</thead>
<tbody>
<tr>
<td>UPLSAPO 10x/0.4 dry</td>
<td>5</td>
<td>OUT</td>
</tr>
<tr>
<td>UPLSAPO 20x/0.7 dry</td>
<td>5</td>
<td>OUT</td>
</tr>
<tr>
<td>UPLSAPO 40x/0.9 dry</td>
<td>2</td>
<td>OUT</td>
</tr>
<tr>
<td>UPLSAPO 60x/1.2 water imm</td>
<td>4</td>
<td>IN</td>
</tr>
<tr>
<td>UPLSAPO 100x/1.4 oil imm</td>
<td>6</td>
<td>IN</td>
</tr>
<tr>
<td>UPLSAPO 20x/0.85 oil imm.</td>
<td>5</td>
<td>IN</td>
</tr>
</tbody>
</table>
Using the 20x and 40x dry objectives - how to keep them clean:
If the slide is clean and does not have any immersion oil, you may switch to the 20x or 40x dry objective. To do so, either switch the objectives via software, or press the "ESC" button on the microscope (green button behind the focusing wheel) - this retracts the objective turret to prevent damage or contamination of the objectives. Then use the control keypad to switch to the desired objective and push the "Esc" button again to return to the focus position. You should need only minimal re-focusing (1/4 or 1/2 turn of the focus wheel).

Beware that if your slide already has immersion oil on it, switching to the 20x or 40x objectives will get oil on these dry lenses and will require their careful cleaning by the MIC staff. If this happens, please let the MIC staff know and do not try to clean the lenses yourself. The front lens elements are very delicate and prone to scratching, which would ruin the optics.

Using the oil immersion objectives:
- Remove the sample from the microscope
- Select the “100x oil” or “20x oil” objective in the software
- Apply a small drop of immersion oil on the objective
- Press the ESC button on the microscope to lower the objective
- Put the sample (slide) on the microscope stage in the same orientation as it was before.
- Press the “ESC” button to bring the objective into focus

CLEANING
- NEVER USE KIMWIPES OR OTHER TISSUE PAPER TO CLEAN OBJECTIVES. USE ONLY LENS PAPER (e.g. Fisher 11-996)
- Using clean lens paper gently blot off the oil from the lens. Do NOT drag the paper across the lens, just dab off the oil. The front lens of the objective is very delicate and must be protected from scratching.
- Wipe off any oil from the objective barrel.
- Thorough cleaning of the oil immersion objectives is performed by the MIC staff only.
Using the 60x/1.2 Water Immersion objective

- Remove the sample from the microscope. Make sure the coverslip on your slide is clean and dry, with absolutely no oil
- Select the 60x/1.2 water immersion objective in the software
- Set the adjustment collar on the objective to the thickness of the coverslip on your specimen (Coverslips #1.5 are approximately 0.17 mm thick)
- Apply a small drop of clean ultra-filtered water on the objective, using a plastic Pasteur pipette
- Press the “ESC” button on the microscope to lower the objective turret
- Place the sample on the microscope stage in the same orientation as it was before
- Press the “ESC” button to bring the objective into focus

CLEANING the 60x/1.2 Water Immersion Objective

- NEVER USE KIMWIPES OR OTHER TISSUE PAPER TO CLEAN OBJECTIVES. USE ONLY LENS PAPER (e.g. Fisher 11-996)
- Using clean lens paper gently blot off the water from the lens. Do NOT drag the paper across the lens, just dab off the water. The front lens of the objective is very delicate and must be protected from scratching.
- Thorough cleaning of the oil immersion objectives is performed by the MIC staff only.

The importance of coverslip thickness for image quality

Use coverslip #1.5 (~0.17 mm thickness) for best image contrast and resolution. The microscope objectives work best with those coverslips. The 40x dry and 60x water immersion objectives have correction collars for coverslip thickness compensation, but they are hard to reach. It is best to leave them adjustment at 0.17 mm and use the correct coverslip.
**Fluorescence microscopy**

FILTER SETS:
There are six filter sets (filter cubes) available, four of them are installed in the motorized filter turret. Users who need to use a filter set not currently installed will need to swap the filter cubes, as shown in the section “Replacing filter cubes for fluorescence imaging”. Filters are described by a central wavelength and bandwidth, or LP for “longpass”. Thus, 350/50 denotes 325-375 nm bandpass.

<table>
<thead>
<tr>
<th>name</th>
<th>Chroma Cat. #</th>
<th>Excitation</th>
<th>Beamsplitter</th>
<th>Emission</th>
</tr>
</thead>
<tbody>
<tr>
<td>DAPI/Hoechst/AMCA</td>
<td>31000v2</td>
<td>350/50</td>
<td>400dclp</td>
<td>460/50</td>
</tr>
<tr>
<td>Cyan GFP v2 (CFP)</td>
<td>31044v2</td>
<td>436/20</td>
<td>455dclp</td>
<td>480/40</td>
</tr>
<tr>
<td>Endow GFP</td>
<td>41017</td>
<td>470/40</td>
<td>495lp</td>
<td>525/50</td>
</tr>
<tr>
<td>Yellow GFP (YFP)</td>
<td>41028</td>
<td>500/20</td>
<td>515lp</td>
<td>535/30</td>
</tr>
<tr>
<td>Cy3</td>
<td>41007a</td>
<td>545/30</td>
<td>570lp</td>
<td>610/75</td>
</tr>
<tr>
<td>Texas Red</td>
<td>41004</td>
<td>560/55</td>
<td>595lp</td>
<td>645/75</td>
</tr>
</tbody>
</table>

- Focus on the specimen using transmitted light (DIC), if possible
- Switch to the desired filter in the software (e.g., GFP)
- Switch from the camera port to eyepiece observation (top left button on the front of the microscope)
- Open the mechanical shutter below the objective turret. Illumination light can be attenuated using a lever on the mercury lamp housing
- Find and focus the specimen
- If you will not be acquiring images right away, close the mechanical shutter or switch back to DIC filter set in the software
- Beware that the Nomarski prism below the objective degrades the fluorescence image. Thus, if high resolution imaging is needed, pull the Nomarski prism out of the optical path
Camera imaging

- Select the desired filter set in the FILTER menu.
- Select the desired objective in the OBJECTIVE menu. Take care not to get oil on dry objectives or on the water immersion 60x objective.
- Run "Live View" and set the Exposure to get a good image. You may need to adjust the histogram to display the image properly (Click "Auto", then tweak the Histogram Min and Max manually)
- You may also adjust the display gamma by pulling the intensity line up or down.
- Switch to the next filter and optimize exposure time as needed.
- To set up Multidimensional Acquisition, click the “Multi-D Acq” button
Multidimensional Acquisition
Set parameters as needed:

- **Time Points**
- **Multiple Positions (XY):** you can mark XY positions or define a matrix of XY coordinates
- **Slices (Z):** if doing z-stacks; set z-start, z-end and z-step (see Table 2 for optimal z-steps).
- **Channels:** set the filter, exposure time, z-offset (to correct for focus shift between different colors) and check whether to acquire the channel for every slice in a z-stack or just once. You can also set to skip a number of frames in a time lapse series.
- **Assign a name to the experiments (directory and name prefix).** Make sure “Save image files to acquisition directory” is checked.
- **Select the acquisition order (e.g., Channels, slices, time points)**
- **Check the “Save Image” option, choose the destination folder and the file name.** Data should be saved to the data disk drive to your folder in “D:Camera_Imaging”. Each time you click “Acquire”: the software will create a new folder with the base name and a sequential number, for example “my-great-sample_1”, “my-great-sample_2”, “my-great-sample_3”, … The folder then contains your image stack file or individual images, as well as the metadata.
- **Usually, saving images as a single multi-image tiff is preferred.**
**XY Pixel size, Recommended Z-step, Sampling and Nyquist Criterion**

The Photometrics Prime sCMOS camera has 2048 x 2048 pixels (6.5μm pixel size). To capture the image at full resolution, the smallest resolved detail in the primary image should be represented by at least two pixels ("Nyquist criterion").

The default configuration file contains spatial calibration for each objective when used without the additional 1.6x magnification. As long as you switch the objectives using the button in the software, rather than using the keypad by the microscope, the pixel size is known to the software and saved with the acquired image. When additional magnification is introduced by engaging a 1.6x field lens (the handle on the microscope is pulled out), the spatial calibration is no longer valid. You could change the spatial calibration (pixel size for each of the objectives) in your config file to the 1.6x field lens values. See table 2 below. You should then save your own copy of your configuration file. It may be easier to just indicate the additional magnification in the image name, and apply the correct spatial calibration later.

If the 1x tube lens magnification provides correct Nyquist sampling, there is usually no good reason to use the 1.6x additional magnification. You would be getting a dimmer image of a smaller field of view, without any gain in resolution.

**Table 2. Pixel size for Photometrics Prime sCMOS camera.** Resolution is calculated for 500 nm wavelength. **Green fill indicates that the XY pixel size satisfies the Nyquist criterion.** All other conditions result in **undersampled images.** XY resolution is calculated as $D_{XY} = 0.61 \cdot \lambda / NA$, Z resolution is calculated as $D_z = 1.4 \cdot \lambda \cdot \eta / (NA^2)$, where NA is the numerical aperture of the objective, $\lambda$ is the wavelength of signal (here, assumed 500 nm, i.e., 0.5 μm) and $\eta$ is the refractive index of the immersion medium (air, 1.0; water, 1.33; oil, 1.515).

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>10x/0.4 dry</td>
<td>1x</td>
<td>0.6470</td>
<td>0.763</td>
<td>4.38</td>
<td>1.90</td>
</tr>
<tr>
<td>10x/0.4 dry</td>
<td>1.6x</td>
<td>0.4034</td>
<td>0.763</td>
<td>4.38</td>
<td>1.90</td>
</tr>
<tr>
<td>20x/0.7 dry</td>
<td>1x</td>
<td>0.3227</td>
<td>0.436</td>
<td>1.43</td>
<td>0.62</td>
</tr>
<tr>
<td>20x/0.7 dry</td>
<td>1.6x</td>
<td>0.2019</td>
<td>0.436</td>
<td>1.43</td>
<td>0.62</td>
</tr>
<tr>
<td>40x/0.9 dry</td>
<td>1x</td>
<td>0.1596</td>
<td>0.339</td>
<td>0.86</td>
<td>0.38</td>
</tr>
<tr>
<td>40x/0.9 dry</td>
<td>1.6x</td>
<td>0.0996</td>
<td>0.339</td>
<td>0.86</td>
<td>0.38</td>
</tr>
<tr>
<td>60x/1.2 WI</td>
<td>1x</td>
<td>0.1080</td>
<td>0.254</td>
<td>0.65</td>
<td>0.28</td>
</tr>
<tr>
<td>60x/1.2 WI</td>
<td>1.6x</td>
<td>0.0675</td>
<td>0.254</td>
<td>0.65</td>
<td>0.28</td>
</tr>
<tr>
<td>20x/0.85 oil</td>
<td>1x</td>
<td>0.3205</td>
<td>0.359</td>
<td>1.47</td>
<td>0.64</td>
</tr>
<tr>
<td>20x/0.85 oil</td>
<td>1.6x</td>
<td>0.2004</td>
<td>0.359</td>
<td>1.47</td>
<td>0.64</td>
</tr>
<tr>
<td>100x/1.4 oil</td>
<td>1x</td>
<td>0.0654</td>
<td>0.218</td>
<td>0.54</td>
<td>0.24</td>
</tr>
<tr>
<td>60X/0.7 dry, LWD</td>
<td>1x</td>
<td>0.1121</td>
<td>0.436</td>
<td>1.43</td>
<td>0.62</td>
</tr>
</tbody>
</table>
Photometric Prime sCMOS Gain and Noise
Camera S/N AF16F202005  Tested 6/27/2016
Dark current, measured at sensor temperature 10 °C below ambient: \(0.013839 \text{e}^{-}/\text{pix/s}\)

<table>
<thead>
<tr>
<th>Readout Speed/data bits</th>
<th>Gain Setting</th>
<th>Gain (electrons/ADU)</th>
<th>Read Noise (e(^{-}))</th>
<th>Non-linearity (%)</th>
<th>Full Well (ke(^{-}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>250MHz/16bit</td>
<td>1</td>
<td>0.47</td>
<td>1.2</td>
<td>1.1</td>
<td>0.72</td>
</tr>
</tbody>
</table>

Data management and conversion

Storing your data
- The imaging computer is not for long-term data storage. After copying your files to your storage medium and confirming they are OK, erase the files from the computer. Old data files will be purged periodically.
- Always create two copies of the data files, store them in different locations, e.g. one in the lab, the other one at home.

Viewing and processing the image files
Micromanager allow saving multi-dimensional data (time-lapse, z-stack, multi-channel) either as individual images (one tiff file for each channel, z-position, time-point) or as a single multi-image TIFF file. This multi-image TIFF file also contains all the metadata in the file header. The best way to open these images is to download and install MicroManager on your computer (http://www.micro-manager.org).
The multi-file data can also be opened using the freeware ImageJ or Fiji (http://fiji.sc).
Replacing filter cubes for fluorescence imaging

The filter turret holds six filter cubes. Cube #1 is for laser scanning confocal imaging, cube #6 is for transmitted light observation (brightfield and DIC), so there are only four positions left for fluorescence imaging. Most of the time, there is DAPI, GFP, Cy3 and Texas Red cubes installed. There are additional filter cubes in the storage cabinet for CFP and YFP. Users received instructions during the initial training how to do the swap.

1. If the mercury lamp is on, make sure the light is blocked: pull the adjustment lever on the lamp housing all the way towards you.

2) Loosen the locking screw by ~1/2 turn (access hole from the right side of the microscope)

3) Pull the filter turret from the left side of the microscope and lay it in top of the camera. Loosen the screw in the center of the turret cover and remove the cover. Find the filter cube you want to remove (NEVER remove the LSM cube #1) – read the label on the side of the cube. Loosen the locking screw for the cube, remove the cube, replace with the desired filter cube and gently tighten the screw.
4. Replace the filter turret cover. Pay attention to the precise orientation of the cover. The small screw head protruding from the side of the cover must go in the notch next to the turret motor.

5. Insert the turret back into the microscope, gently tighten the turret lock screw (access hole from the right side of the microscope)

6. Attach a temporary label above the corresponding filter button on the keypad to indicate that the filter was replaced.
SHUTDOWN PROCEDURE

- Remove specimen from the stage

- Using **lens paper** only, gently blot excess oil from the oil immersion objective. Do not wipe the lens or rub the lens with the paper. NEVER USE KIMWIPES or other kinds of paper on the objective lens. Thorough cleaning of the objectives is done by the MIC staff only.

- Switch to the lowest magnification objective (10x).

- Clean the work area, wipe the table surfaces with 70% ethanol.

- Turn off all control boxes and the computer, the Mercury lamp power supply is the last to be turned off.

**TURN OFF THE CAMERA**

- Replace the plastic cover. Only cover the microscope itself, NOT the mercury lamp or the camera or surrounding area.

- If you are the last user for the day, and if the lasers are still running, it is your responsibility to **turn the lasers off!**

**Laser turn off procedure:**

1. 405 nm diode lasers: turn the key to OFF, turn the power switch to OFF.
2. Red and green HeNe lasers – turn the knob to OFF **ONLY IF YOU ARE THE LAST USER FOR THE DAY**
3. Argon laser: TURN OFF **ONLY IF YOU ARE THE LAST USER FOR THE DAY. IF NOT, LEAVE THE ARGON LASER RUNNING.**
   
   i. **Turn the key to OFF position**
   
   ii. **WAIT!! The laser must cool down.** The fan to stop automatically when the laser unit has cooled down. It takes several minutes
   
   iii. Set the power switch to OFF.

- Sign off in the log book.

- Turn the lights off; make sure the door is closed when you leave.

- Finish your Kiosk session