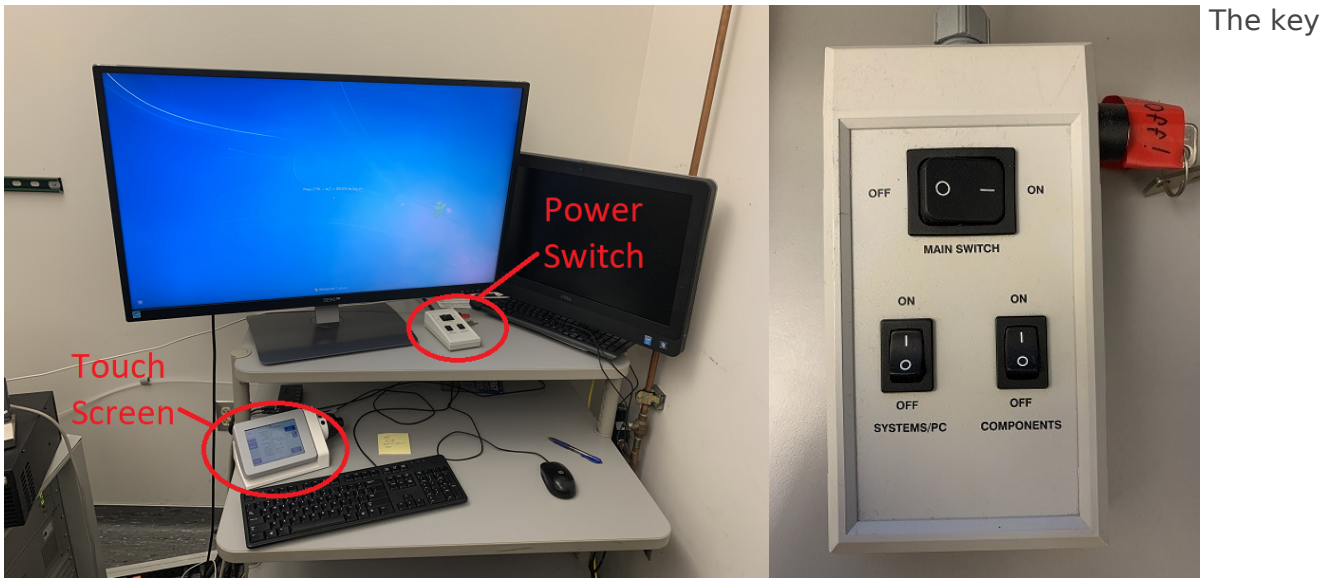


# LSM 710 Basic Operation

## Turning On the System



Turn on the X-Cite lamp next to the microscope. Turn on the Argon ion laser. Turn the key to the horizontal position and flip the small metal switch up. If the light is red lower the current control knob until it is green.

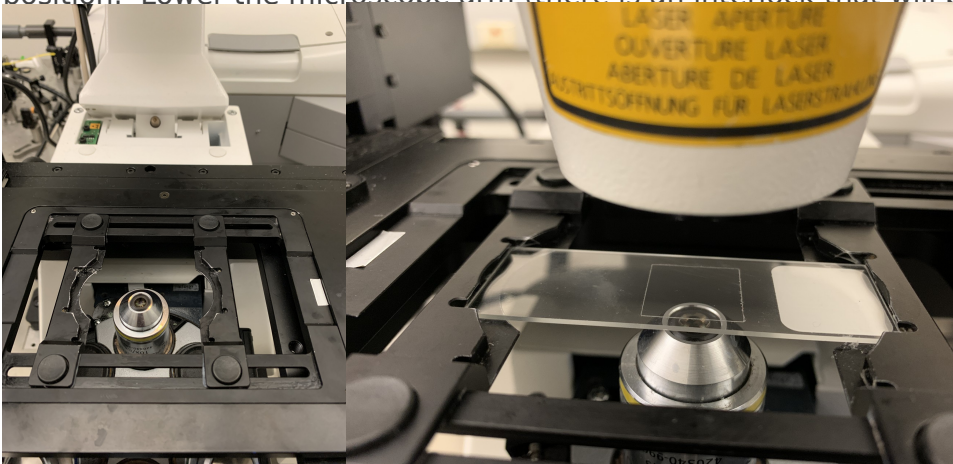


## Loading Your Sample

To load your sample, push the arm of the microscope back until it holds in place (don't grab it by the focus knob to lower the

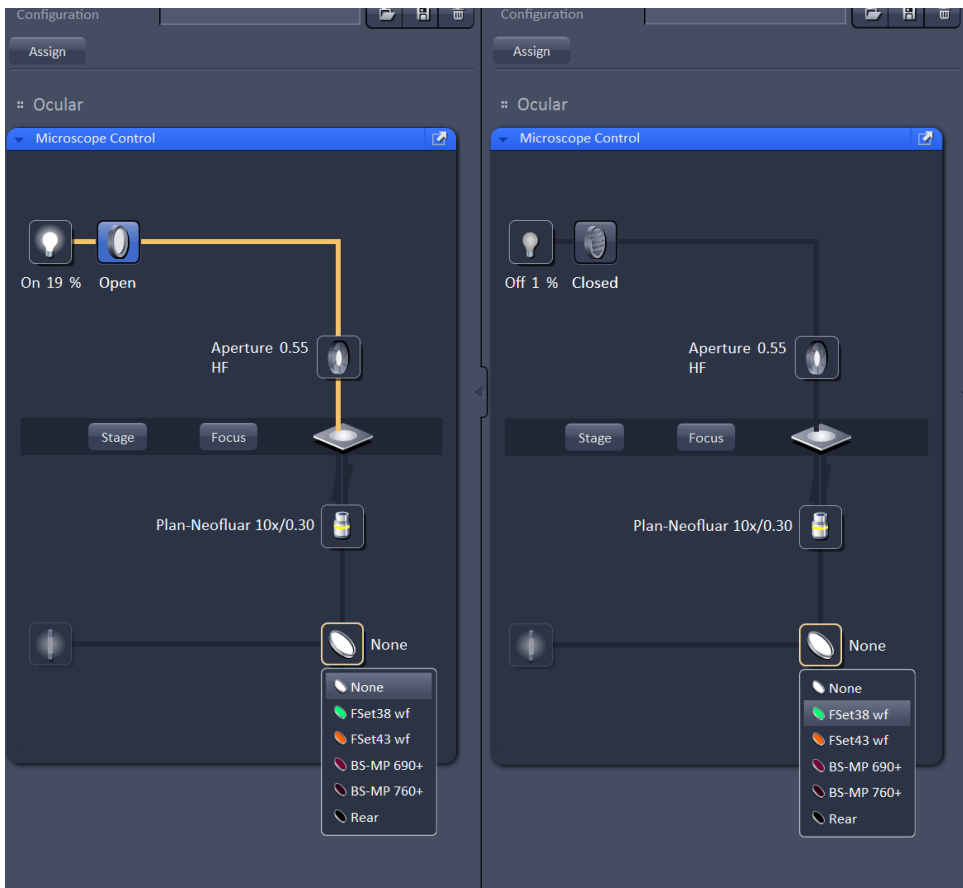


Move the sample to holder to the correct position for either a slide or a 35mm dish. Make sure to load the sample with the coverslip facing down (towards the objective). If you need to use the 40x or 63x objective add a drop of water or immersion oil to the objective lens. After placing the sample on the holder, press the upper button on the focus knob to return the objective to the working position. Lower the microscope arm (there is an interlock that will prevent the laser from turning



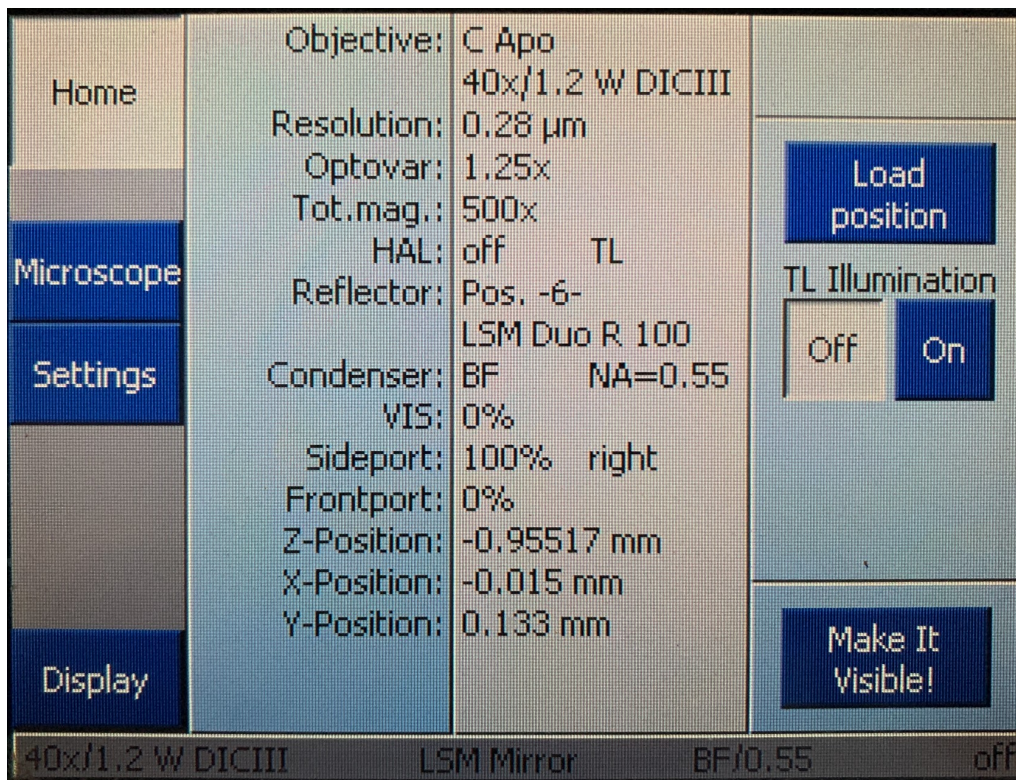
## Locating Your Sample

On the computer, login with your IGB account and password, and open the Zen program. Select "Start System" from the options it presents you. Wait for Zen to load all the components and connect to the microscope. Zen will present you with four tabs at the top: Locate, Acquisition, Processing, and Maintain. It will start you in the Locate tab; this will allow you to find the sample in the eyepiece. To find the sample in bright field, turn on the lamp, transmitted light (TL) shutter as shown below in the image on the left. You will also need to choose the empty filter from the filter turret. After those items are selected, You can look in the eyepiece and adjust the course or fine focus knob to find your sample. To find your sample with the 488 or 561 filter, close the TL shutter and switch to the green or red filter as shown in the picture on the right. If you don't see blue or green light coming from the the objective, check that the manual shutter on the X-cite lamp is open. Use the focus knob to find your sample in the eyepiece.



You can also use the touch screen to control these settings and view the sample in the eyepiece. The main menu on the touch screen will display information about the objective and stage position. There is also a button labeled "Make It Visible" that will put the microscope directly into a bright field configuration so you can view the sample.

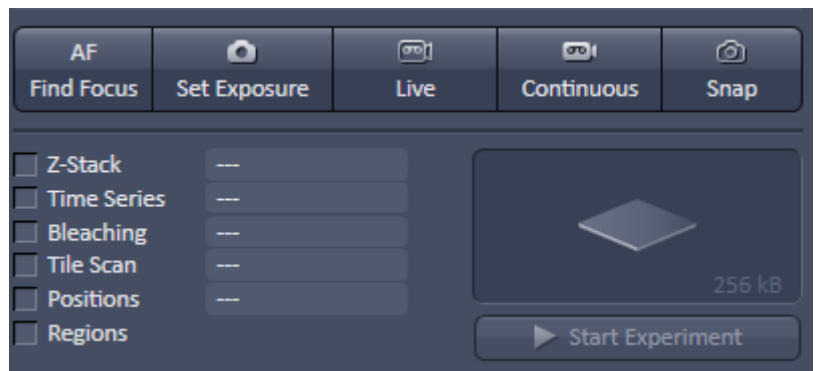




If you press the microscope button you'll be able to select the objective and reflector. Once you have found your sample in the eyepiece, you can move on to setting up your scan.

## Setting Up Your Scan

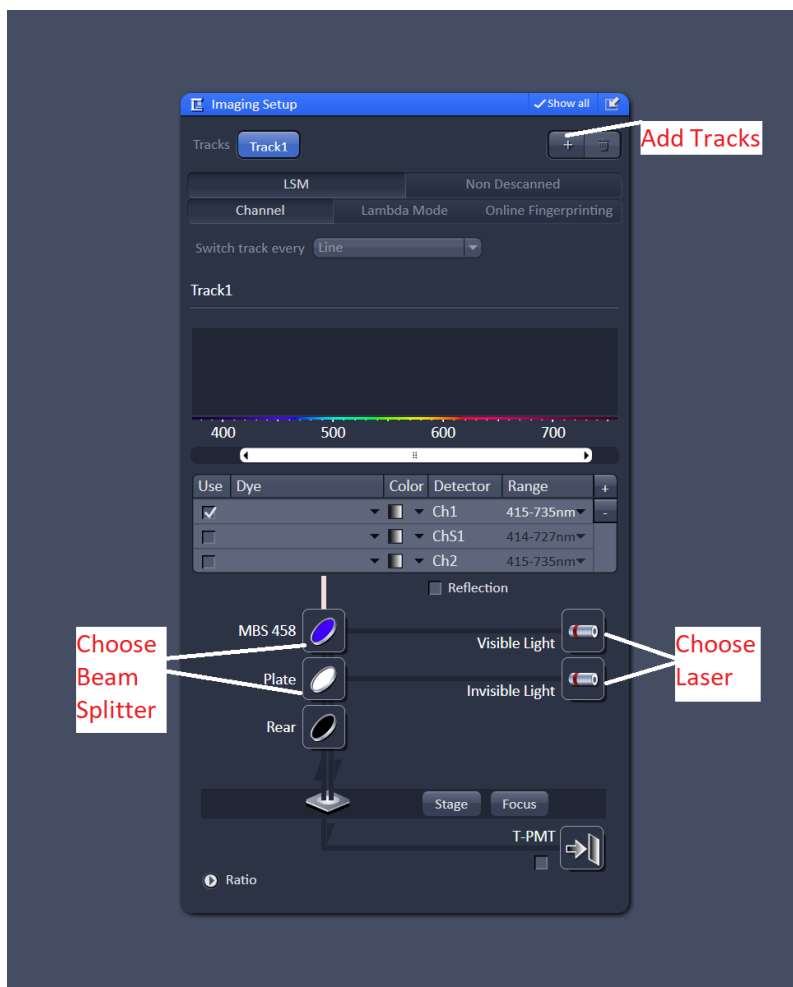
Once you have located your sample in the eyepiece, click the acquisition tab (not the menu item at the top). This is where you will setup the parameters for your image. Zen will show you a series of menus, the four most relevant (Lasers, Imaging Setup, Acquisition Mode, and Channels) will be open and the rest will be collapsed. We will go through the four menus needed to setup a scan. Also, we will be using the live and snap buttons at the top of the screen as shown below.



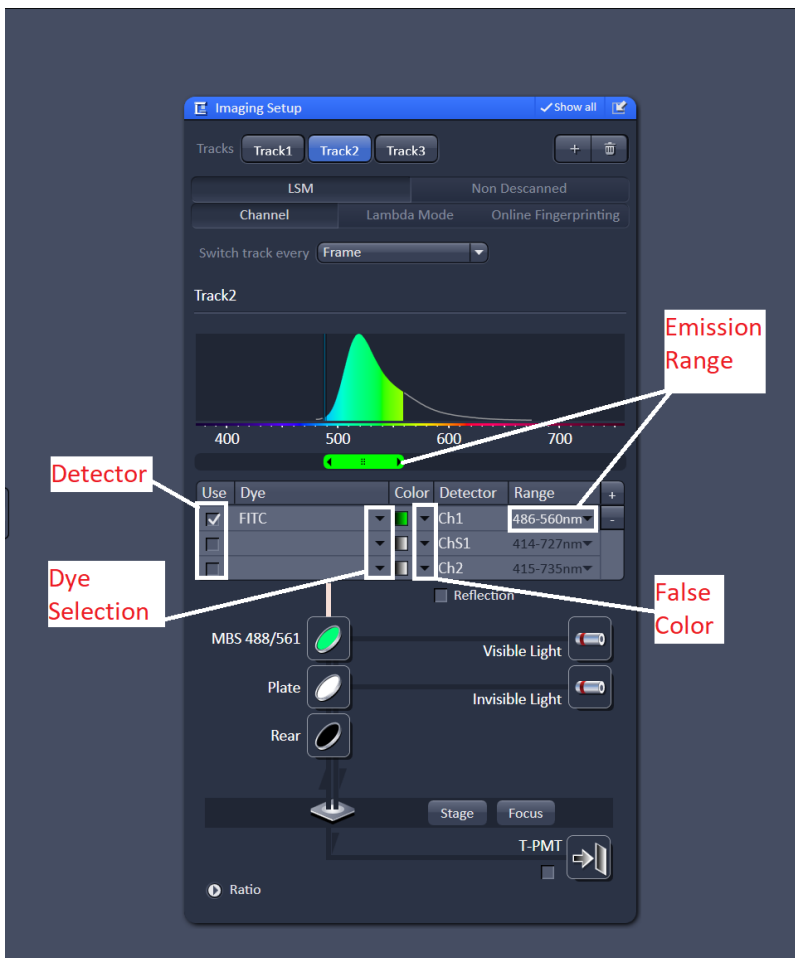
In the laser menu, turn on the 561 laser if you need it. The laser properties menu will show you the status of the laser, it should say either "connected" or "warming up". If it says "inaccessible" please contact the core staff.



Once the lasers are on. You can proceed to the "Imaging Setup" menu. Press the "+" button to add a track, you'll want to add a channel for each dye in your sample. You can have up to four tracks. I'd also recommend changing the "switch track" option from line to frame. Switching the track every line requires that all of the optics in the light path be fixed, if you aren't careful this can be frustrating.



For each track choose a laser. Then choose a beam splitter (dichroic filter) that matches the laser line. For example, if you are using the 488nm laser, choose one of the beam splitters labeled for 488nm.

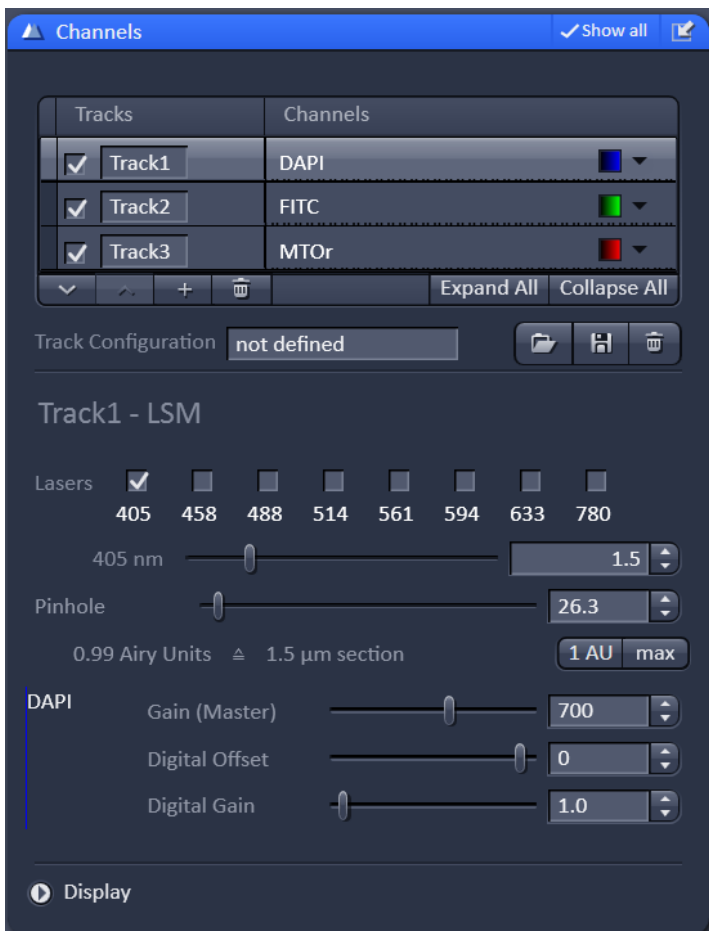


Next, select the detector you want to use. By default channel 1 is selected as the detector. As long as "switch track every frame" is selected you can just leave the detector as channel 1. Choose a dye from the database, this will display the emission spectrum of the dye. Adjust the emission range so that you are collecting most of the light from the emission spectrum. Repeat this for every track - one track for every dye in the sample.

Once the tracks are all setup, move on to the channels menu. At this point it is worth noting that "tracks" refer to the light path setup and "channels" refer to detectors. There are cases where you might want to do more than one channel per track, but for now we will just focus on using one channel for every track.

One important user interface note, the check mark box can be unchecked and the computer will ignore that track. This is different than highlighting a track/channel which will display the parameters for that channel.

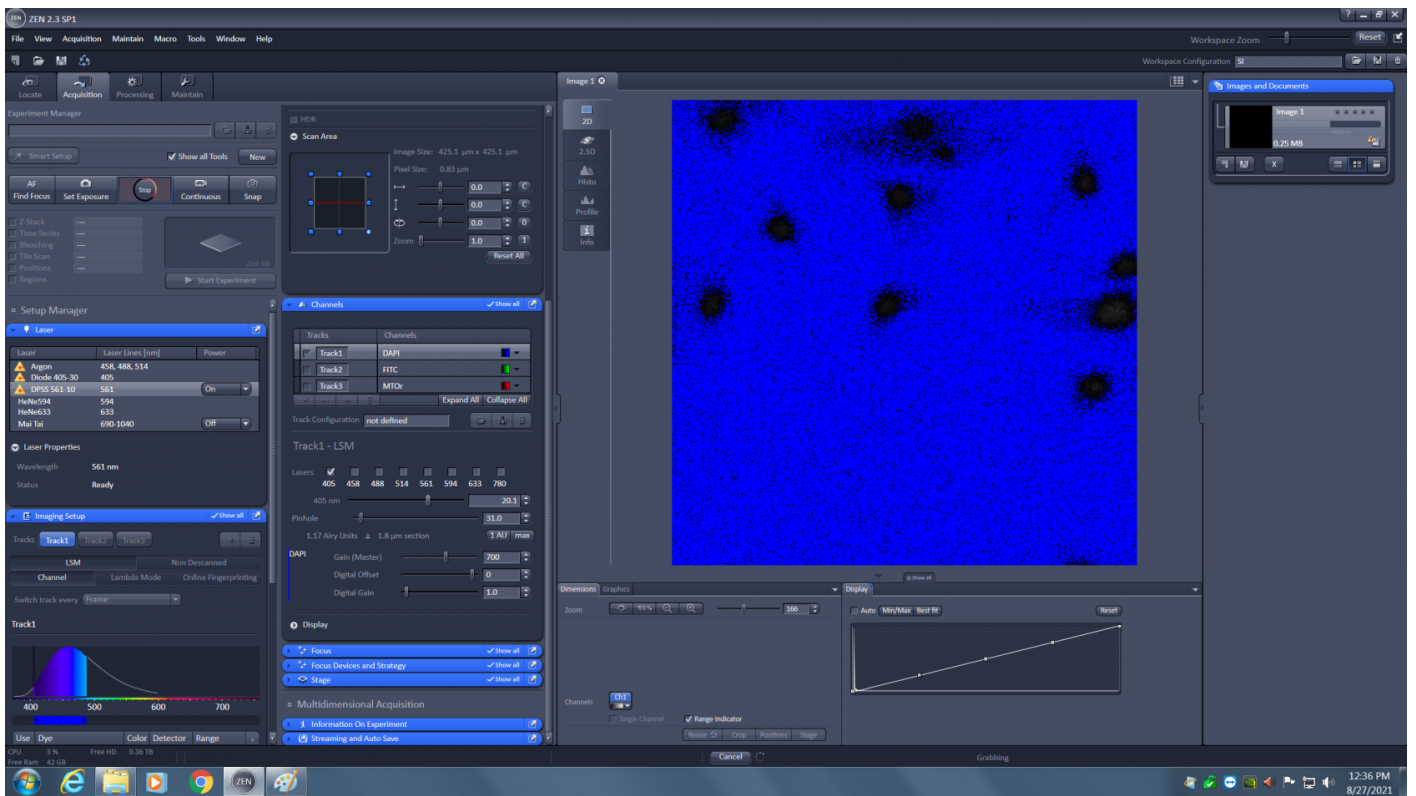




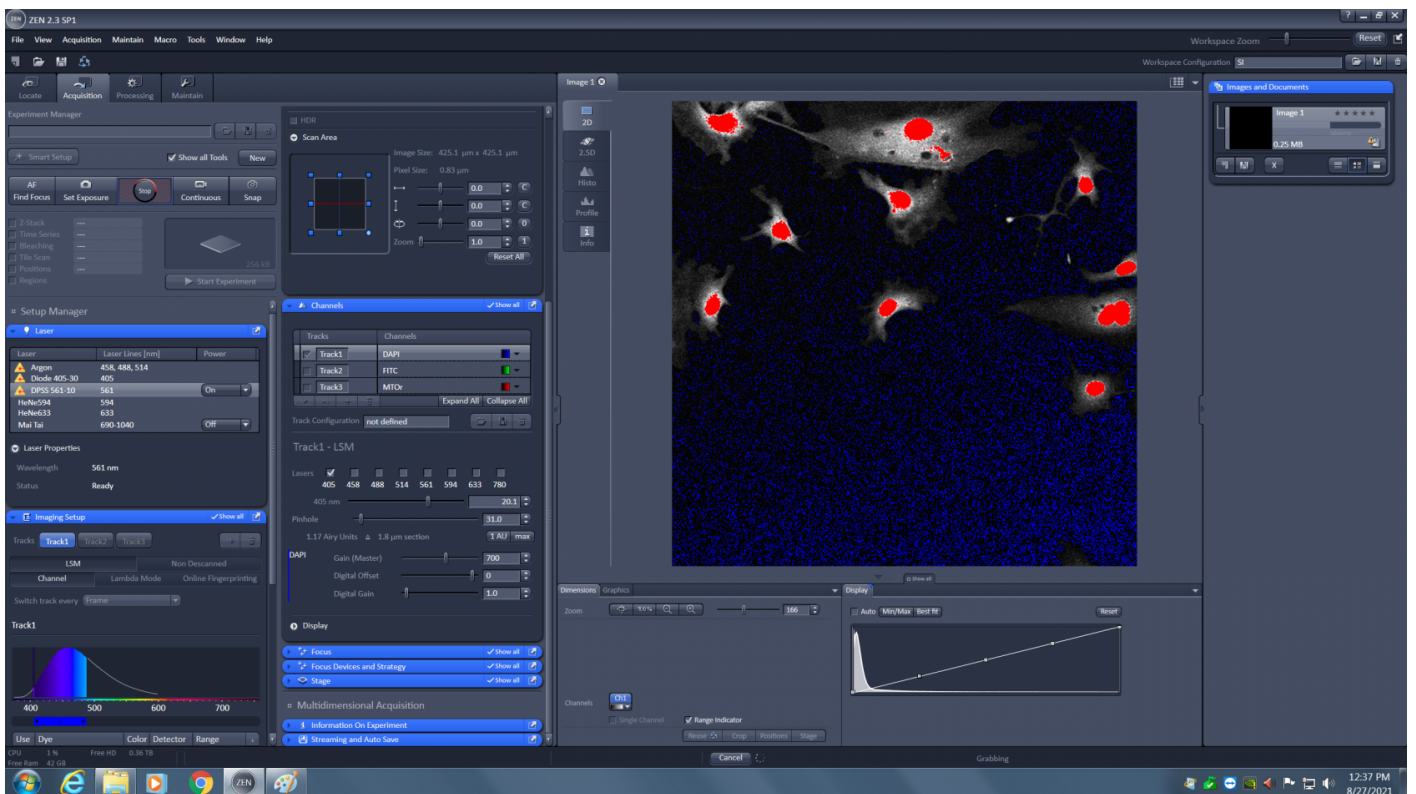
For each channel you will need to adjust the laser power, gain, and pinhole size. Start by setting the gain to 700. Raising the gain will increase your signal but will also increase your noise. Lowering the gain will lower the signal and noise, requiring you to use a higher laser power. We have found that 700 gain is a good starting point.

For the pinhole, start with the middle channel (or the most important channel). Press the 1AU button. This will set the pinhole to one airy unit (the diameter of the airy disk). Next to 1AU or 0.99 AU is an approximation of the optical section thickness, 1.5 microns in this case. The section thickness will increase as you open the pinhole. It is important to keep the pinhole diameter (26.3 in this example) the same across all channels.

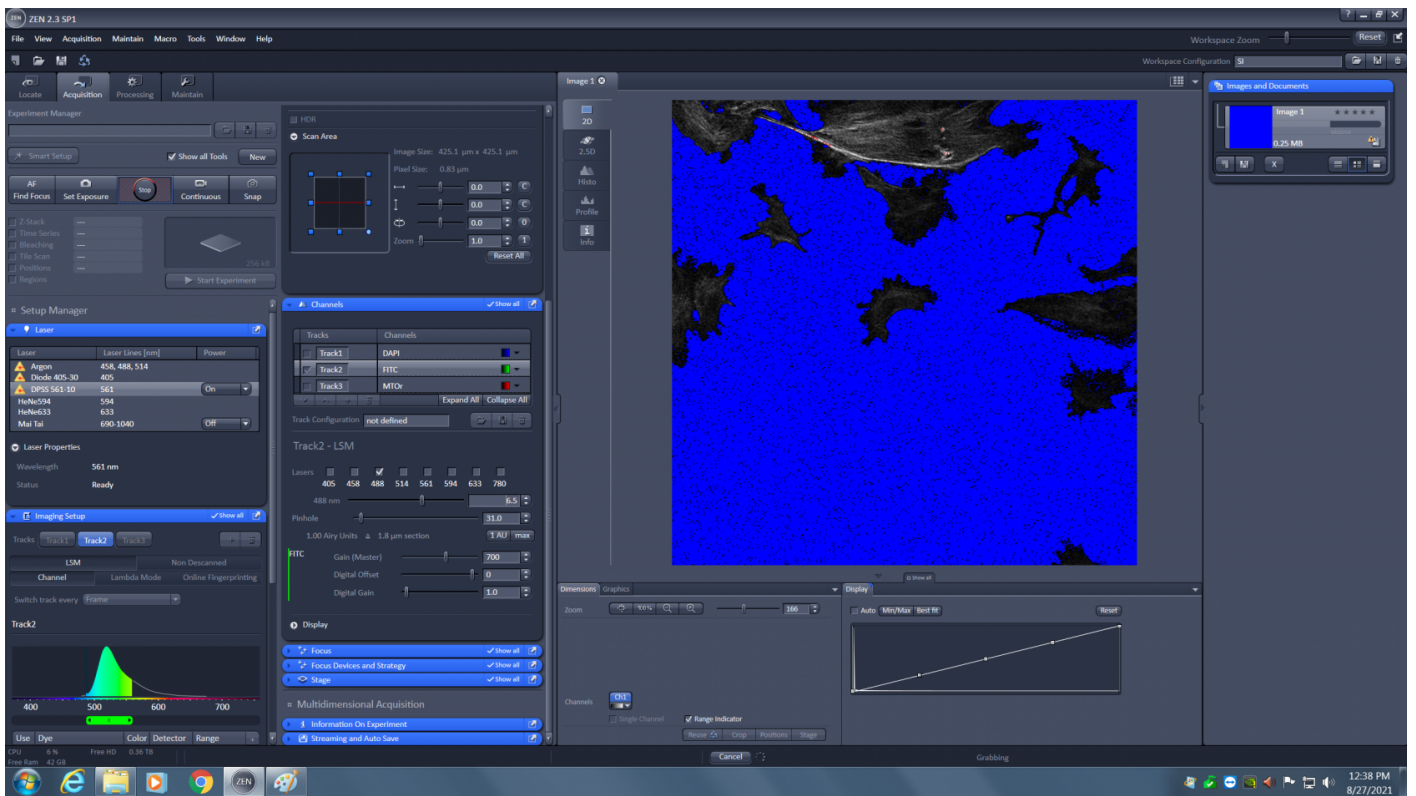
Now you need to adjust the laser power (just above the pinhole). It starts at 0.20% and can be raised to 100%. You will need visual feedback to correctly adjust this. Uncheck all but one track, and click on the the "live" live button at the top of the screen. At first you might not see anything, it is very likely the sample is not exactly in focus. Click on the "range indicator" button at the bottom of the screen. This button will turn off the false color and display the live image in a diagnostic color scheme. Blue is zero, black to white is signal, and red is saturation.



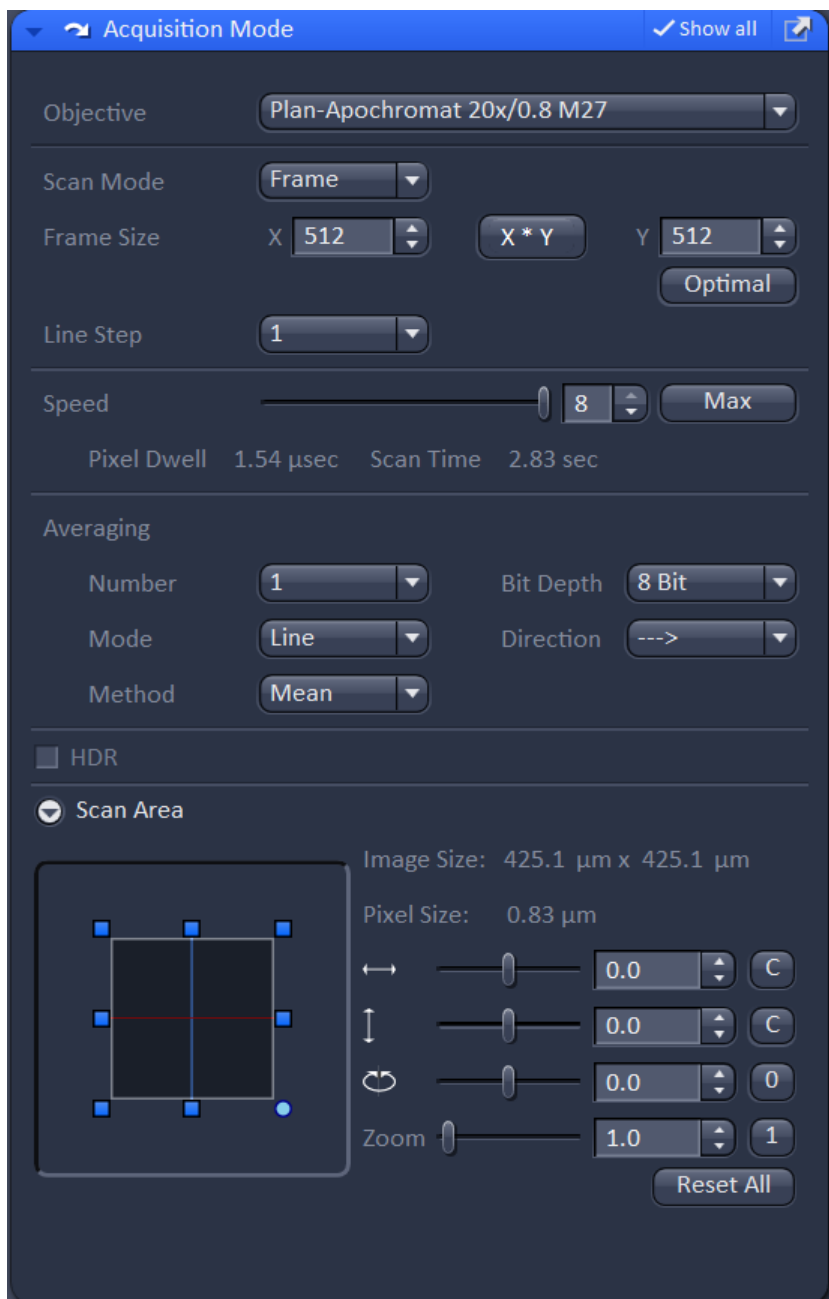
You can see in this image that there is signal, but it is weak. First adjust focus until the signal gets brighter (black to grey to white to red)..



Maximize the amount of red to make sure your sample is in focus. Then lower the laser power to that only a few red pixels are visible. You want to use as much of the detector's dynamic range as possible without saturating the detector. A few red pixels are okay, you just don't want big patches of them. The correctly adjusted image should look as shown below.



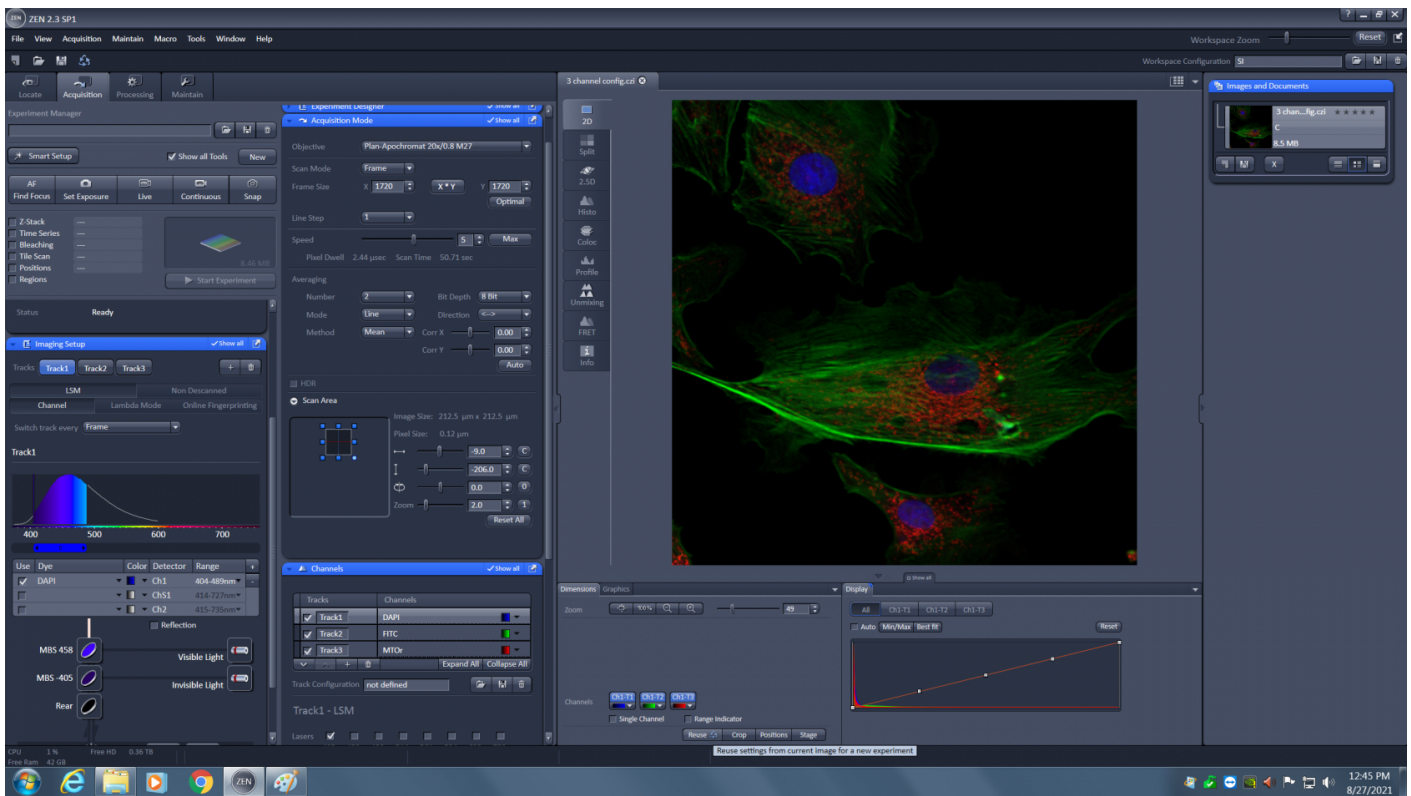
Press the "stop" button, and repeat this procedure for the other channels. Once you are done setting up all the channels we will setup the scan parameters in the acquisition menu



Start with the scan area, you can adjust the size, position, and orientation of the scan area. Adjust the size and position to frame the sample. There are two important numbers displayed next to the scan area. The "Image Size" is physical size of the image, and the pixel size is the image size divided by the the frame size. Click on the XY button to choose a frame size. If the frame size is too small, then the pixel size will be too big and your resolution will suffer. Click on the "optimal" button just below the XY button and Zen will select the correct frame size.

You'll need to choose a scan speed. This will affect the scan time and pixel dwell time. The pixel dwell time is how long the detector collects light at each pixel. I recommend not going below 1 microsecond, as this will result in a noisier image. You can choose an averaging number to average lines/frames together in order to improve the signal to noise ratio. In general keep the bit depth at 8-bit. Once the settings are correct select the "Snap" button to start the scan. One note, "snap" will use all the settings in the acquisition menu. The "Live" button with always use a 512 x 512 frame size and scan as fast as possible.





Once you have taken the image, save it as a CZI file on the D drive in the "User Files Deleted Anytime" folder. We don't guarantee files saved here are kept forever but we don't go out of our way to delete them.

The next time you use the LSM 710, you can open this CZI file and click the "Reuse" button just below the range indicator, this will reapply all the settings used to take the image. You'll still want to check the laser power settings with the range indicator as this will change from sample to sample.

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