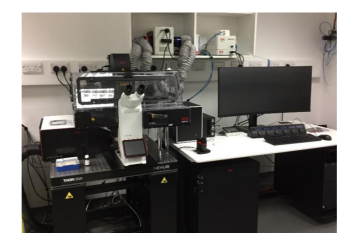
Imperial College London



Observing Life As It Happens

QUICKSTARTGUIDE: HCF2 - CONFOCAL LEICA Stellaris 8 inverted (ICTEM 314)



Contents

Startup Procedure	2
Software Setup	2
"Configuration" tab	
Re-Using settings	3
Microscope Controls	3
Changing Objectives	3
Setting the Focal plane	4
Microscope stand controls	5
Microscope chamber	5
Köhler illumination	6
Software / Image Acquisition	6
Channel Setup – Loading an existing setup	6
Channel Setup - Automatic using the "Dye Assistant"	6
Channel Setup – New manual setup	7
Setting Intensity	7
Image Format	8
Z-Stacks	
Capturing Images	8
Saving Data	9
Setting Environmental Control	9
Sample Holders	10
Changing Sample Holders	11
Shutdown Procedure	11

Startup Procedure

- Start PC
- On the central power unit Switch on:
 - Power
 - o Laser
 - turn on Laser Emission Key
- IMPORTANT III The microscope stage will auto calibrate (move) – to avoid trapped fingers DO NOT LOAD A SAMPLE UNTIL THE MICROSCOPE HAS FINISHD STARTING UP
- Login (IC network account)
- Wait until the TFT screen on the front of the microscope has finished booting
- Start LEICA Application Software "LAS X" on desktop

Software Setup

There are two possible hardware configurations for confocal: with or without environmental control

- In the start-up window from the drop-down menu select either:
 - o machine.xlhw use without environmental control
 - the environmental control can be run manually without software control
 - machine-Climate-Control.xlhw with software environmental control (NB Heating unit must be on before starting)

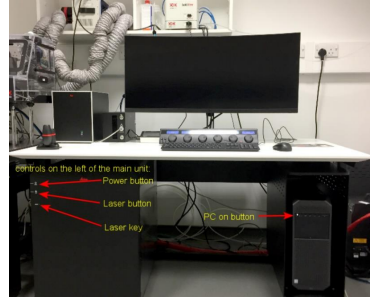
"Configuration" tab

STELLARIS 5 🗘	Configuration	Acquire	Proce	ss	Quantify	>	Analysis)

Select "Hardware" and set the required Bit Depth – 16 Bit

	ø	Hardware Settings	u L	4	User Configuration LAS X Version: 4.1.1.23273 Copyrgit © 3020 Lata Microspitens CMS Grid	bit
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		ctor Auto Selection	waat unto stante is ready		Data handling at Ki M 💿 Folders 🔘 Properts	Settings used for current mage A applied to insign (f motilate) Cabital settings from above) A could be applied to mapping them above Activate Access Management Activate Access Management

Change the range indicator lookup table to Glow Over in the "User Configuration"



eica Application Suite X 3.0.24308	MICROSYSTEMS
Configuration : Microscope :	machine.xlhw ≑ DMI8 ≑
Load settings at startup :	
	OK Cancel

• Select "Laser Config" and turn on the 405 Diode and WLL (white light) lasers



Re-Using settings

In the **Configuration** tab:

- Select IPS
- Select Load
- Select All_Users.xml from the E:\ drive
- Then applying or loading experimental settings will load all image settings

Mask Filter	*	Select Mask	
Confocal Load & Apply		IPS Masks :	م
Confocal Live Data Mode		z-Use-Mode	,
		Begin / End	
Confocal Sequential Frame / Sta	ack	Stack Config	
		Scan-Mode	
		Scan-Format	
		Time Config	
		Lambda Config	
		LUTs	
		Shutter	
Load		Scan-Direction	
Reset to Default		Resolution	

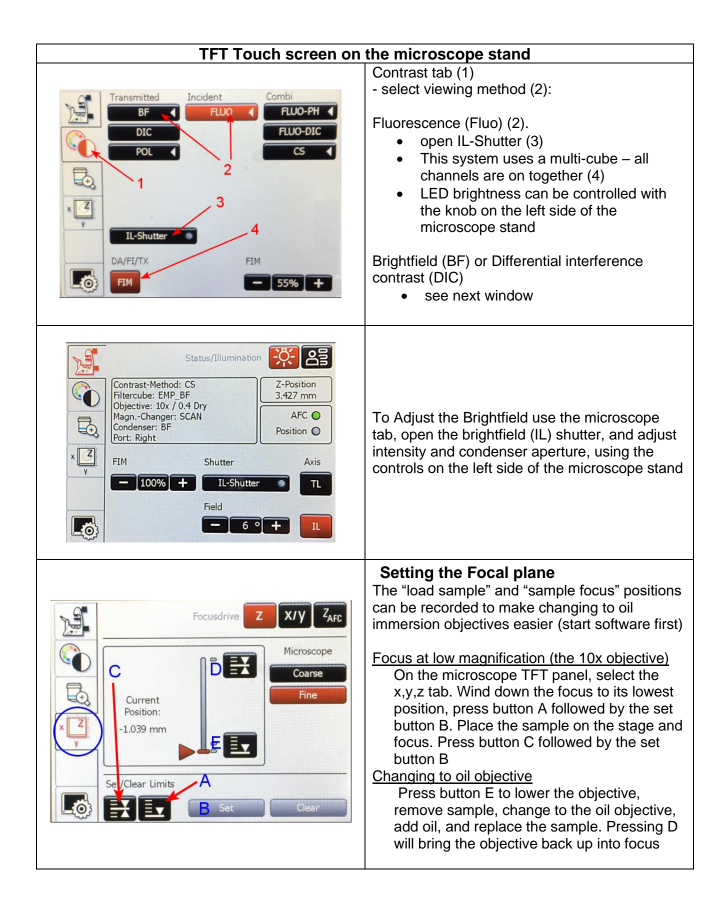
Microscope Controls

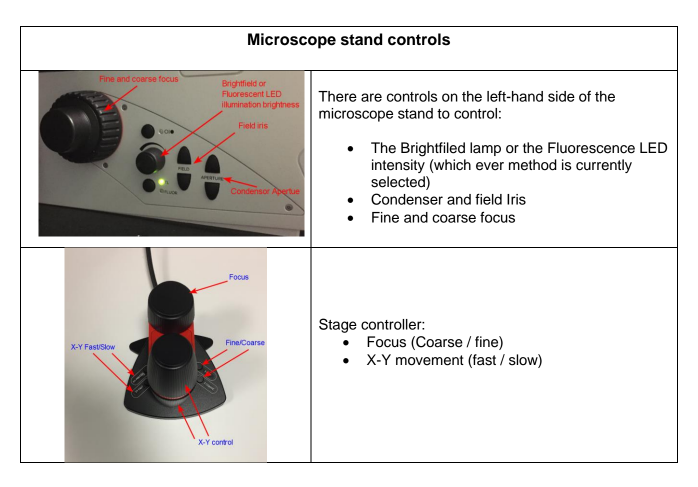
Changing Objectives

Select the "Acquire" tab and in the main setup window:



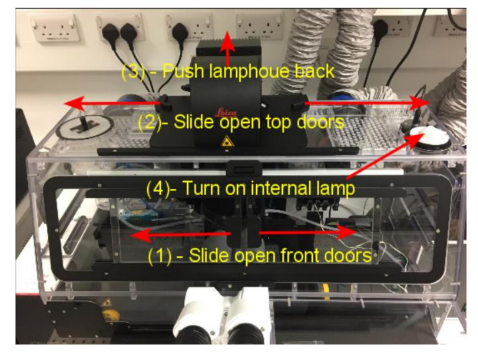
- Click on the objective and a list will appear allowing you to select the correct objective
- This will also bring up another window if changing to or from oil or water immersion
- You will need to click yes on this window to move the objective before adding oil/water
- NB clean the oil objective and sample before clicking yes and moving to the dry objectives





Microscope chamber

Loading and changing samples:



The chamber light can also be controlled with a footswitch once turned on

Köhler illumination

- Select a Low power objective (x10 or x20)
- Switch on brightfield
- Look through the eyepiece, if necessary, adjust the eyepieces for your eyes
- Adjust the brightness if necessary (TFT panel / left side of microscope stand)
- Focus on your sample
- Open the condenser aperture iris (TFT panel)
- Fully close the field iris above the condenser (1)
- Looking down the eyepiece, focus the black edges of the field iris with the large silver focus wheel (2)
- If necessary, centre the field iris with the Köhler screws (3)
 Allen keys to adjust them are found on the back of the condenser (right side)
- Re-open the illumination iris so that the black ring just disappears from your field of view

Software / Image Acquisition

Select "Acquire" tab and select the desired acquisition mode eg XYZ

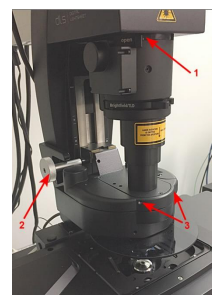
Channel Setup – Loading an existing setup

Load settings (Acquire" tab):

- Either, from a previously saved Image. Open the saved image/library in the "projects" tab and select the image. Then click on the APPLY icon on the menu above (NB. The objective, pixel number, bit depth and zoom and averaging are not re-loaded and may need re-setting).
- Or use the Load channel setup option from the main setup window to open a previously created and saved channel setup

Channel Setup - Automatic using the "Dye Assistant"

• select the Dye Assistant setup button

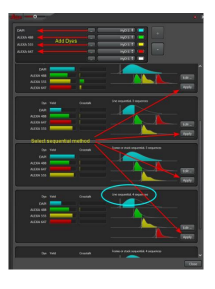








- a new window will open allowing channel selection by fluorophore.
- clicking on the "..." button will load a search window allowing dye to be selected
- repeat adding channels as necessary (a new line will automatically appear for each new channel)
- Select the sequential scan method of choice (usually Line • sequential) by pressing apply. This will setup the channels and detectors automatically



Channel Setup – New manual setup

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Searching for fluorophores, and dragging/dropping into channels

- 1. Type in dye
- 2. Select sequential method (Stack between stacks, Frame - between frames, Lines between lines)
- 3. Drag dye onto a detector (it will auto select the detectors)
- 4. Adjust detector range if required

Add further tracks/dyes as required using the (+) symbol

Setting Intensity

- Start preview scanning by clicking on either the LIVE button or FAST LIVE (bottom of the screen)
 - Live scans all channels at scan capture settings (size, averaging, etc can be quite slow).
 - Fast Live scans only the currently active channel (512 format and no averaging). 0 This is good for setting up.
- Select a channel in the Channel setup window by ٠ clicking on the "Setting" panel and use the control panel to optimally set the Smart Gain, Smart Intensity, Zoom and Focus. Repeat for each channel.
- Use the "Range Indicator" LUT to with this.

help



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Image Format

Use the XY panel to Set

- Image format (1) 1024 x 1024 is typical but depends on image requirements
- zoom factor (2) (If required, move the Zoom Area using the arrow icons, NB Zooms above 4 results in empty magnification at 1024 x 1024 pixels)
- Averaging (3) required to give you sufficient image quality:
 - o line averaging for live imaging
 - line or frame averaging for fixed cells
- Pinhole (4) is preset at 1AU but may be adjusted to change Z volume

Z-Stacks

- Adjust focus in "live" mode to start of Z- stacck and select Begin (1)
- Adjust focus in "live" mode to end of Z- stacck and select End (2)
- Set step size the "optical section" size (z) can be read from the X-Y panel or the "+" button will allow Nyquist settings to be applied

When finished with Z stack mode there is a "Trash" button to remove the stack settings





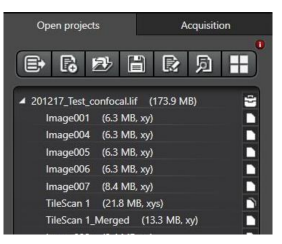
Capturing Images

Once the setup is compplete the image is recoreded using either "Capture Image" for a single image or use "Start" for an image sequence (Z-stack, tiling or time series)



Saving Data

- Images are stored in a library (.lif file format)
- Every time Capture Image or Start is pressed the image is added (but NOT SAVED) to the Library in the "Open projects" tab.
- Right clicking on the individual image names allows renaming or deleting.
- To save the images click the "Save" icon above.
- The first time it will prompt for a location and the Library name.



• Then every time you capture an image press the "Save" icon to update the library or you could lose your data.

Setting Environmental Control

This unit can be controlled either manually (independent of the software) or through the software using machine-Climate-Control.xlhw option when starting LAS-X, however, the unit needs to be turned on before running this software option.

Switching On

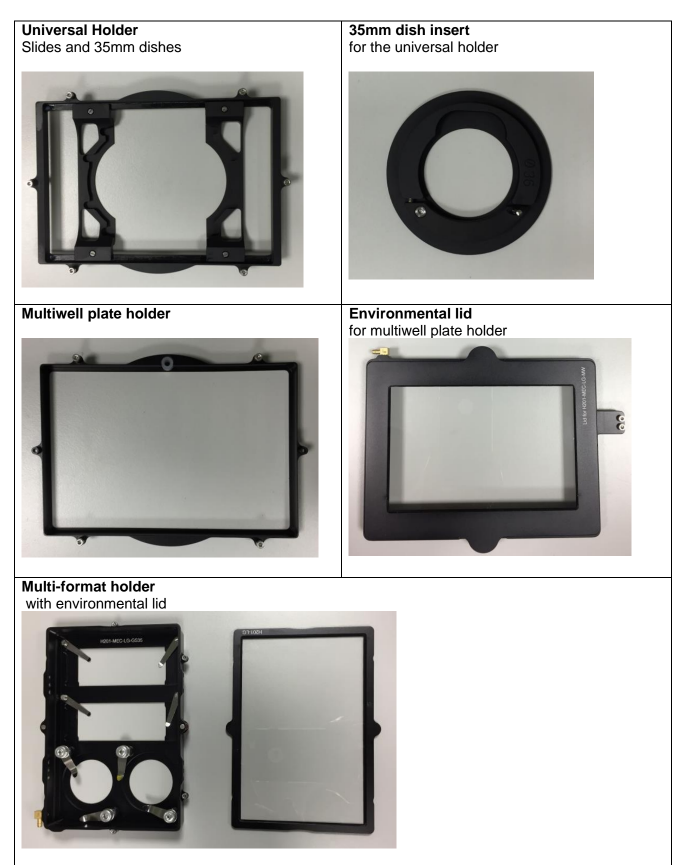
- Make sure chamber is closed
- Turn on the controller on the desk by pressing and holding the "On" button for about 3 seconds
- Press the T^oC button to adjust the temperature
- Press the CO₂% button to set CO₂



Turning off

• Turn off the controller on the desk by pressing and holding the "On" button for about 3 seconds

Sample Holders



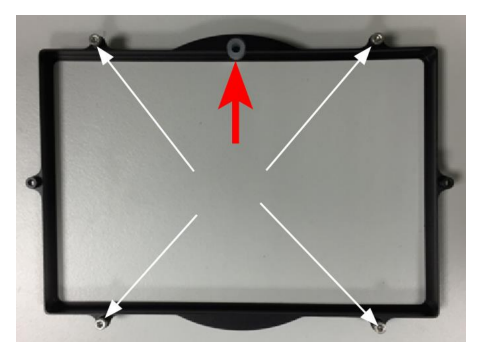
Changing Sample Holders

Slide Holders

The slide sample holders are not retained by screws but just drop in place and can be lifted in and out as required

Plate Holder

However, the multi-well plate holder is retained by a series of small screws, 2 on each side.



- To remove this sample holder, loosen these screws (4 turns) with the Allen key provided. Important, do not completely unscrew these screws Just 4-5 turns will release the stage and allow its removal with part of the screws still attached in their threads. Be careful not to lose these screws!
- To fit the holder, drop it into place with the rubber support (red Arrow) to the back of the microscope. Gently tighten the screws DO NOT OVER TIGHTEN SCREWS

Shutdown Procedure

- In the Configuration tab and laser config, switch off lasers
- Close LASX
- Update booking if necessary.
- Remove your samples & clean objective lenses with fresh lens tissue
- Clear up the desk
- Save files onto the server
- On the central power unit Switch off:
 - o Laser Key
 - o Laser
 - Power
- Shut down PC or sign out for next user