

Spinning Disc microscope

Nikon Eclipse Ti

Principal:

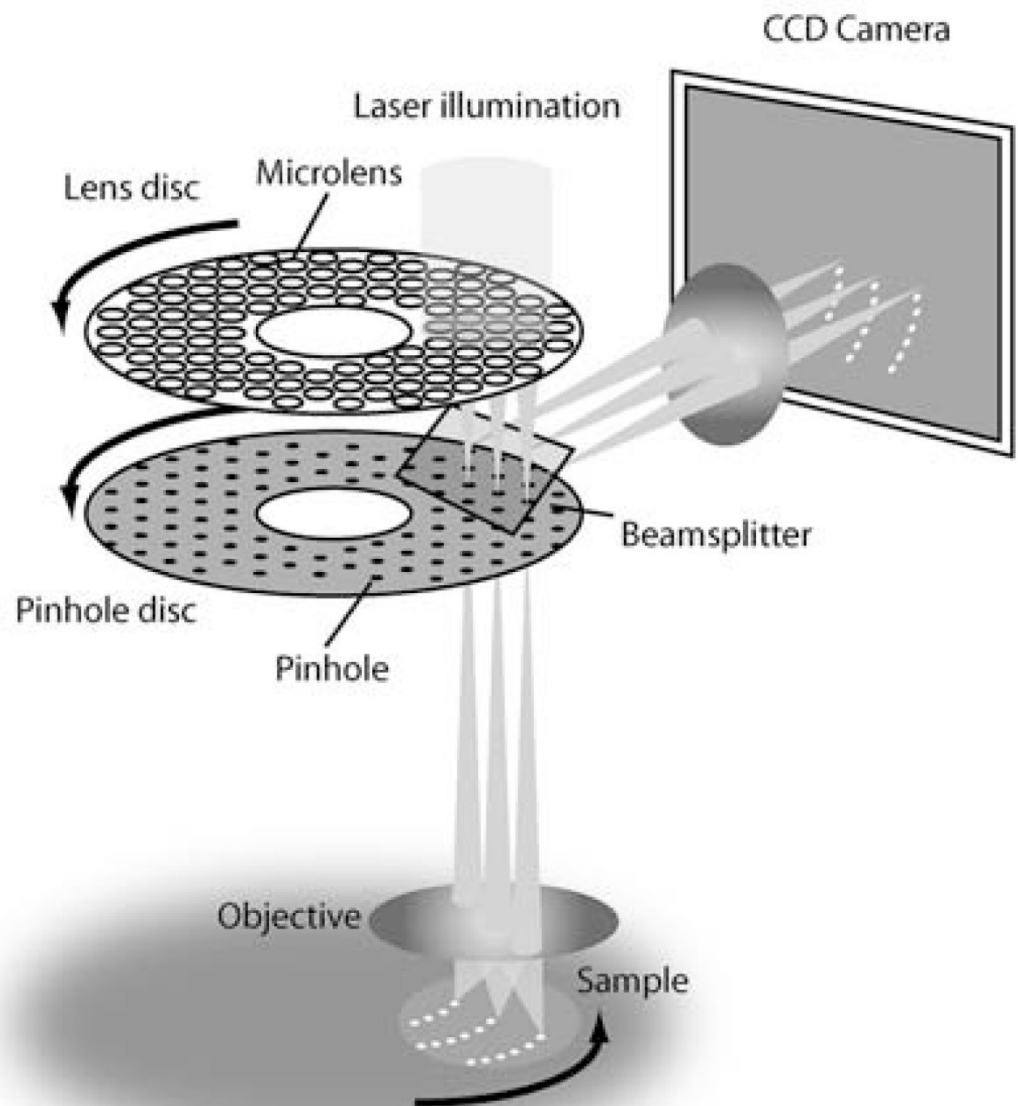
- Fast rotating discs are mounted to each other
- A Laser is projected on the first disc with micro lenses
- The focused and split beams will pass the pinholes in the next disc
- The beams rotate at high speed to excite fluorophores in the specimen in de field of view.
- Out of focus signals from your specimen will be blocked by the pinholes in the second disc
- The dichroic mirror projects the confocal-like-image on a camera

Benefits:

- High speed confocal-like imaging
- Lower bleaching rate due to speed of laser intensity

Downside:

- Fixed pinhole size



Start system

- Switch on the power by the power socket (on the floor under the table)
- Switch the mercury lamp on and then ignite it.
 - If not needed don't turn the mercury lamp on
 - The unit on the shelf closest to the door
- Switch on all devices with a button on the shelf and the table.
 - Camera and laser box are automatically enabled
- Next switch on the devices with key (Spinning disc unit and Laser control panel)
 - SPD unit has a shutter, press the little button. A red LED will go on
- Start the computer and start the program Metamorph.
 - Select suited user account.

Live Cell imaging:

- When imaging at 37°C:
- Place heating unit in microscope table before heating the device
 - Don't use any force if the unit is already heated before installing in microscope table
 - Wait till the unit is cooled down before installing the heating unit
- The heating device to maintain your cells uses water.
 - Fill reservoir with demineralized water if it runs empty
- Allow the system to warm up for ~30 minutes
- Settings for the heating unit are printed and at the wall

Finding focus

- Put the objective in the lowest position using button A
 - Lowest position is ~500 um
- Choose the objective with the preferred magnification
- Put a little drop emersion oil on the objective
 - Only if it's an oil emersion objective
- Place the metal ring in the middle of the heating chamber
 - Secure with the metal clips
- Centre you cells above the objective with the joystick
 - Button on top of joystick will change speed, S (slow) or F (fast)
- Turn focus wheel until the oil is in contact with the coverslip
- In MetaMorph choose transmission button in the Taskbar to activate transmission light
- With button D you can use the fine adjustment control knob
 - course, fine or extra fine
- Finding appropriate fluorescence cells can be done with the mercury lamp
 - Buttons are in the taskbar
 - Increase or decrease brightness by changing the ND filters
 - Located at the right back side of the microscope
 - ND 4, 8 will block intensity, they can be combined
- To change coverslips, bring objective down with lowest button A. Bring objective back to previous height with highest button A



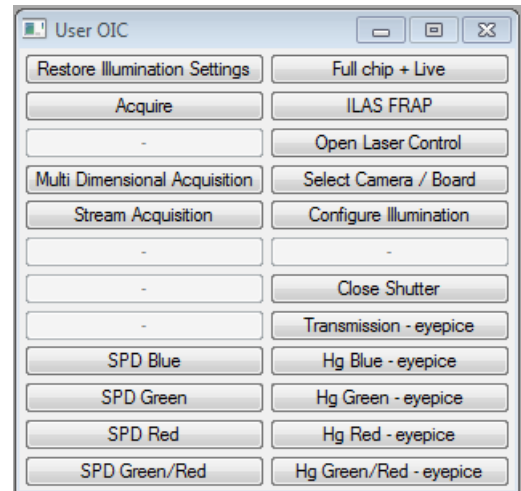
Perfect Focus System (PFS)

- If focus is found a LED (E) will turn orange
 - Focus can be kept using the Perfect Focus system
 - Press the green blinking button E
- The focus of the PFS is adjustable with an extra focus wheel module
- The turntable navigate the lens up and down as shown and with the blue button toggles between fine (pressed) and coarse (out)
- Focusing with perfect focus should be viewed through the camera



Metamorph - Taskbar

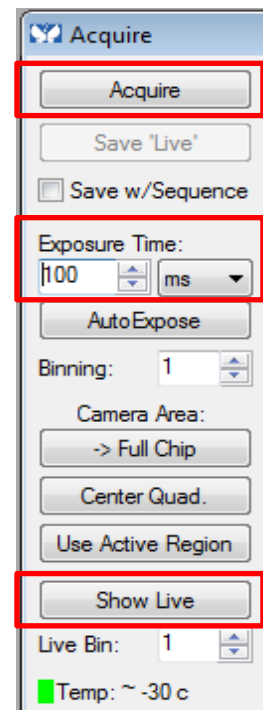
- Metamorph functions from a Taskbar and an Acquire window
- You can restore illumination settings with the button: "Restore OIC settings"
- Open the Acquire window with Acquire
- Access Acquisition modes with: MDA & Stream
- Select preferred illumination settings
 - Left side: lasers for camera,
 - right side: mercury lamp or transmission to eyepiece
- Open FRAP or laser console
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Metamorph - Acquire window

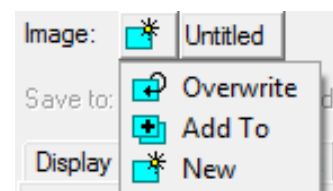
Sidebar left:

- Exposure time
 - Minimum interval time at full chip is 90ms, quad chip is 50ms
 - Exposure time can be lower than interval time
- Full Chip, Quad Chip, Use Active Region
 - Use full chip, middle quadrant, or region selected by ROI
- Live bin & bin
 - Combine pixels to increase readout speed
 - Use only for live viewing or real recording
 - Normal state = 1
- Start Live/Stop Live
 - Start/stop live mode



Acquire images

- To acquire live mode press "Acquire" button
- Separate images can be recorded as separate images or in stack



Tab: Display:

- Auto scale option
 - Enable Auto scale or change custom auto scale
 - Full bit-range is saved to file, this is only for visualization in the software
- Visualize saturated pixels

Tab Acquire:

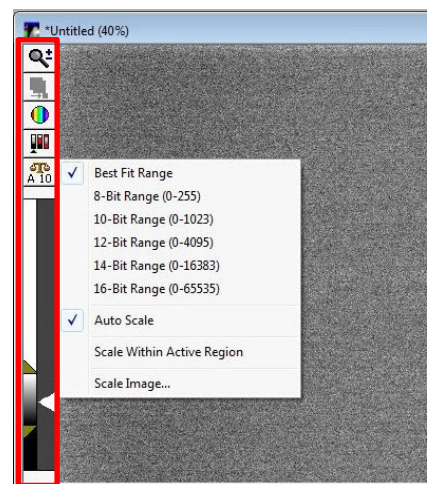
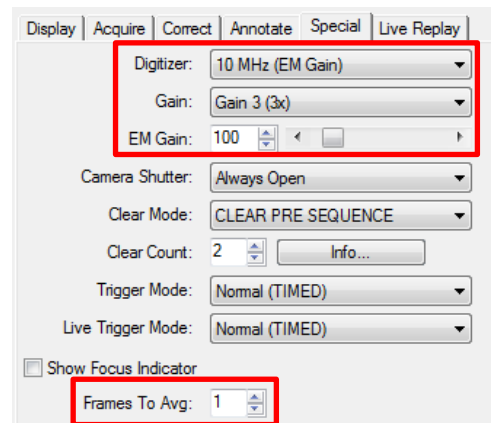
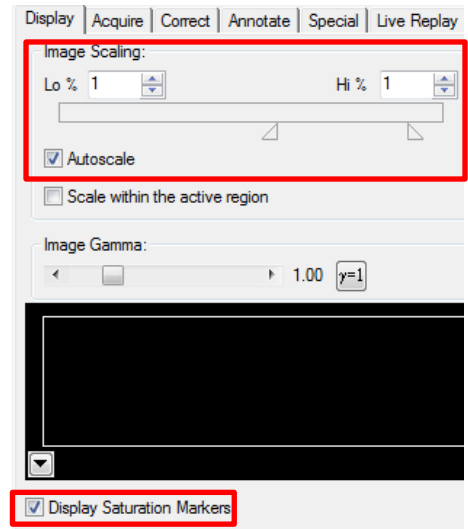
- Be sure to set the Illumination choice to "Current shutter"

Tab Special:

- Gain options:
 - 10 MHz (EM gain), gain3 3x is standard setting
 - Control gain between 0 and 1000
 - Use "Frames To Avg" option to average images / or use longer exposure time

Live image window

- Magnifying glass
- Display image in other LUT
 - Grayscale, Color by wavelength
- Histogram of intensities
 - Manual scaling can be done with the orange arrows
- Auto scaling options
 - Best fit range, 8/12/14/16 bit
 - Auto scale on/off
 - Scale within region



Streaming imaging

Open the Stream Acquisition window with the button in the taskbar

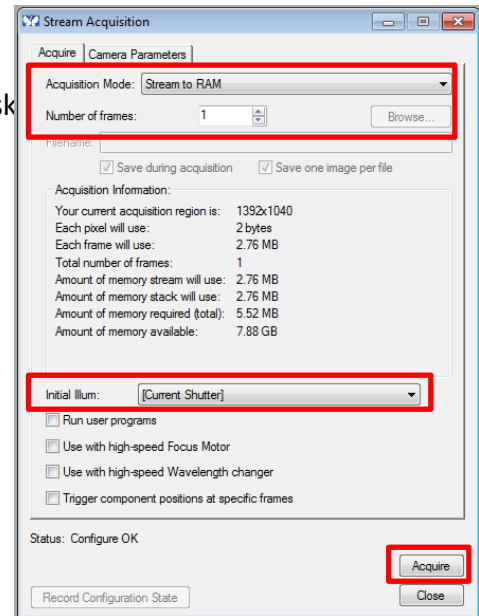
- Start recording with Acquire button
- Check if chosen configuration is ok

Tab Acquire:

- Acquisition modes:
 - RAM save later to hard disk
 - HD choose name and location to save
Save as on file or separate TIF files
- Enter the desired number of frames
- Select "Initial illumination": *Current Shutter*

Tab Camera Parameters:

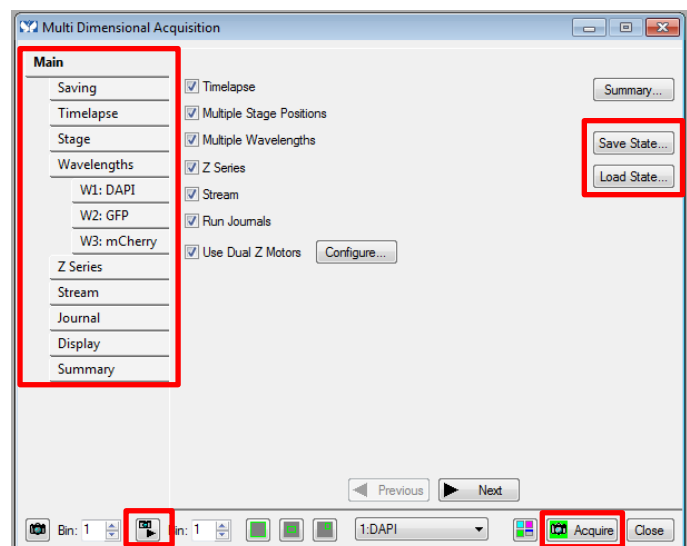
- Recording options:
 - Average frames
 - Number of frames to skip, during acquisition
 - Don't show recording (faster when exposure time < minimal interval time)
- Camera readout default options should be:
 - Camera state: HALT
 - Shutter mode: OPEN PRE SEQUENCE
 - Clear mode: CLEAR PRE SEQUENCE



Multi-Dimensional Acquisition (MDA)

Combine multiple tasks: z-stack, timelapse, streaming, multiple positions, multiple wavelengths, use of journals.

- Open the MDA window using Taskbar
- Select acquisition mode(s) in *Main*
- Continue with the tabs shown left:
 - Saving: select filename and folder
 - Select illumination settings
 - ... options for acquisition modes
- When setup is done press acquire
- Save or load settings of experiment setup, with Save/Load State buttons
- Live mode can be started from MDA window



MDA: Time lapse imaging

- Activate the time-lapse option in MDA's main window
- Set the desired interval and time
- Choose the desired illumination settings in the wavelength tab
 - Or multiple when the option multiple wavelengths are selected
- Start recording by clicking on Acquire

MDA: Z-stack

- Activate Z-stack option in MDA's main window
 - Activate use dual Z motor
 - Set Z-stack motor to: *Piezo*
 - Set all other options to: *TiZ*
- Select option: Center around current
 - Height of objective is shown between -50 μm and 50 μm
 - Not the same as display microscope , different Z-motor
- Give range, step size (in μm), number of steps
- Choose the desired illumination settings in the wavelength tab
- Start recording by clicking on Acquire
- Don't use PFS during Z-stack
 - Activate "Run Journal" in MDA's main window
 - Choose PSF off before Z-stack and PFS on after Z-stack

* For extensive multi-position, -wavelength, -z-stack experiment see the manual on the OIC website, www.erasmusoic.nl

Shut down system

- For live cell imaging:
 - Remove Tokai Hit incubator from the stage and place it on the table
 - Close the Tokai Hit incubator with a plastic well
- **Clean objective carefully**
 - Remove oil with lens paper:
 - Hold lens paper tight between your fingers and wipe over objective
 - Clean lens with 2-propanol on your lens paper
- Cover stage with lens paper box
- Save data to OIC network storage (O:\ drive)
 - To collect the data from your own pc, connect to network storage drive via the address:
<\\oic-station\oic>
User: guest, password: guest
 - When the hard disk of the microscopes pc gets full, data will be deleted
 - You are responsible for your own data, the network-drive is only for transport
 - Extended manual about network storage can be found on the OIC website
www.erasmusoic.nl Facility>Manuals>OIC network drive
- **If you are the last user of the day**
- Shut down all devices
 - Shut down mercury lamp at last
 - Camera power is switched at the adapter box at the floor
- Switch off the power socket located at the floor

Contact

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