Nikon

AX/AX R with NSPARC

# AX AX R with NSPARC

Confocal based Super Resolution Microscope

Nikon Ax

Shedding New Light On MICROSCOPY

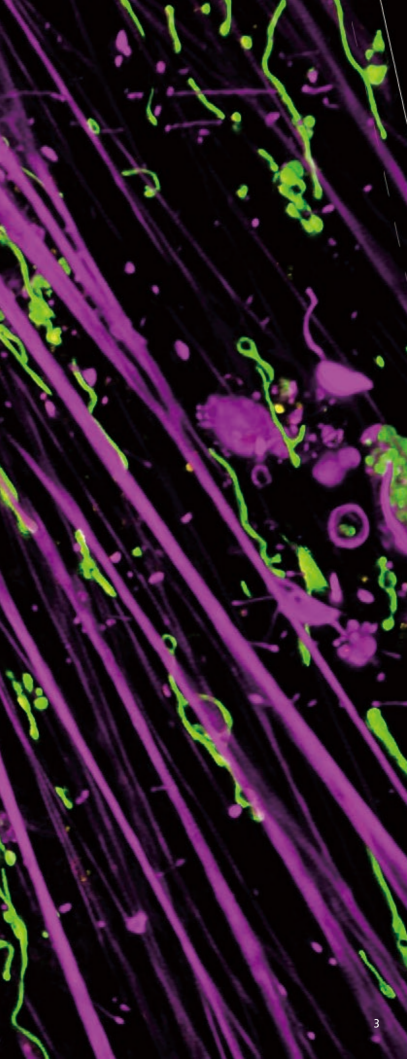
# Igniting New Confocal Imaging Potential

The AX/AX R confocal microscope provides an unprecedented field of view and improved pixel density, enabling the collection of more data over a wide area and down to deep regions in tissues. Its remarkably high sensitivity and high-speed scanning reduce phototoxicity to the specimen and minimize damage. Equipped with image acquisition/analysis functions utilizing Artificial Intelligence (AI), the AX/AX R can efficiently acquire highly reliable data.

The newly developed Nikon Spatial Array Confocal (NSPARC) detector utilizes a detector array to collect a two-dimensional image at each scanned point. It adds additional potential to imaging by providing a new level of ultra-low noise sensitive spatial detection, enabling improved lateral resolution of 1 airy unit/~230 nm\* and improved signal-to-noise ratio.

The Nikon AX series brings new solutions to confocal imaging

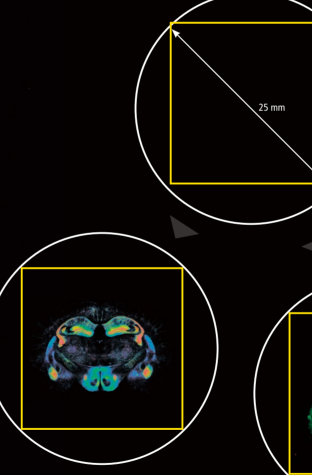
\*Lateral optical resolution as defined by the Abbe diffraction limit (0.61l / NA)



# See More of the Specimen

With the largest field-of-view on both inverted and upright microscope stands available (25 mm diagonal), more specimens fit in one FOV with more objective lens choices than ever before. Coupled with scanning sizes up to 8192 x 8192 pixels, sampling beyond the optical diffraction limit is possible even at low magnifications with the AX/AX R. Using lower magnifications with longer working distances and high numerical apertures enables more flexible specimen preparations to be used, while the large FOV allows simultaneous high resolution in one image. Collect more data in every image, and at faster rates. This high throughput is beneficial in not only academic research but also drug discovery research.

The AX/AX R has a 25 mm diagonal FOV, much larger than other confocal instruments.



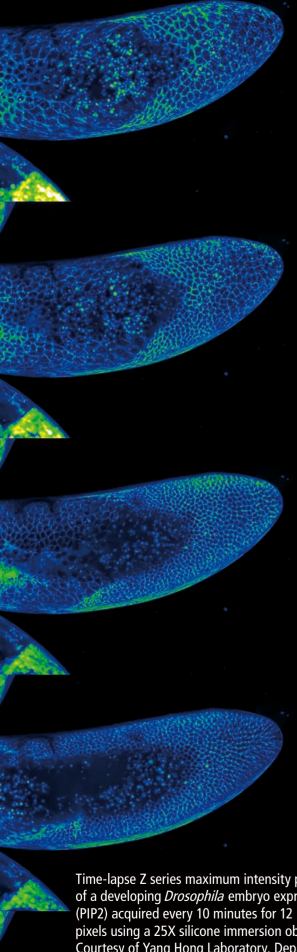
\*Cleared adult mouse brain acquired with 1X objective in one acquisition of 1 FOV

> \*Drosophila sp. embryo development easily fits within the FOV using a high NA 25X SIL 1.05 NA objective

Whole mouse bladder optically cleared with iDISCO and acquired at 8192 x 8192 pixels using a 2X Plan Apo objective, effective pixel size 0.6 µm (over 5X the spatial resolution of a typical monochrome CMOS camera). Courtesy of Dr. Gerry Apodaca, Integrative Systems Biology, Department of Medicine, University of Pittsburgh in collaboration with Dr. Alan Watson at the Center for Biological Imaging, University of Pittsburgh.

\*It would not be possible to capture this sample in one FOV or at this resolution with other commercial confocal systems

\*Danio sp. 2d+ embryo 4X



# Observe with Minimal Disturbance...

Laser scanning confocal imaging is principally challenging on specimen viability, as it applies focused laser illumination point by point on a sample.

The AX R's high speed resonant scanning, which decreases the illumination time by more than 20X typical confocal scanning times, greatly reduces biases caused by merely acquiring images.

Reducing the acquisition time also allows for extremely highspeed imaging (up to 720 fps at 2048 x 16 pixels). The result: longer time-lapse imaging and/or more frequent high-speed imaging of living samples that enables dynamic events to be captured. This is useful for imaging living samples, and dramatically increases the efficiency of such drug discovery research as multipoint assays.

Time-lapse Z series maximum intensity projection images of a developing *Drosophila* embryo expressing PLC-PH::GFP (PIP2) acquired every 10 minutes for 12 hours at 2K x 1K pixels using a 25X silicone immersion objective. Courtesy of Yang Hong Laboratory, Department of Cell Biology, University of Pittsburgh in collaboration with the Center for Biological Imaging.

# ...and Acquire more Rapidly...

Confocal imaging, notoriously slow because of its point-scanning requirement for high quality 3-dimensional imaging at high resolution, is greatly changed by fast imaging with the AX R's resonant scanning capabilities.

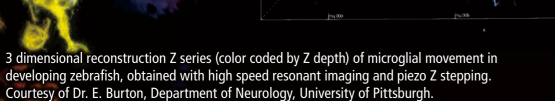
Utilizing 2048 x 2048 pixel resonant scanning and a 25 mm FOV on a large intestinal sample montage, acquiring 25 high-resolution images and merging them in under 2 minutes

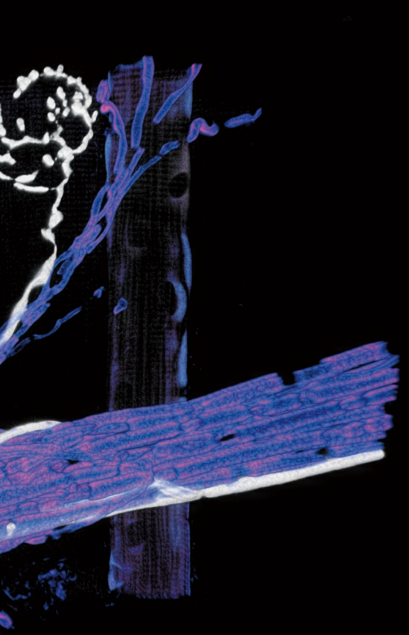
# ...and with Ultrafine Detail

With a full 25 mm FOV, up to 8192 x 8192 pixels, and the capability for supravideo frame rates, the AX/AX R allows for spectacular imaging with high resolution, at both low and high magnifications.

The entire range of a whole organism or system biology down to intracellular imaging is achievable on one instrument.

This high resolution capability is beneficial for target selection in drug discovery.





# Mouse muscle acquired with a 25X SIL immersion objective using 2048 x 2048 pixel resonant scanning

# Detectors In Tune with Labels

The AX/AX R's all new DUX-VB detector custom-tunes emission bandwidths to a library of labels and probes, and provides the freedom to fine-tune emission bands to minimize unwanted fluorescence. Simply select the number of labels in your specimen and their catalog names. Alternatively, you can define the desired emission ranges, or even simply the emission color: the AX/AX R and NIS-Elements software does the rest, including optimizing the dichroic mirror and laser excitation choices best suited for imaging. Or, acquire hyperspectral images in up to 66 emission channels for unmixing.

Optionally, the AX/AX R's base DUX-ST detector allows up to 12 discreet bandpasses of emission, upgradable to 18.

And all detector systems can be customized with high sensitivity and low noise GaAsP or Multi-alkali PMT detectors to provide the best detector for sensitivity and wavelength response requirements as well as budgets.

# Superior Optics for Confocal imaging

Controlling the manufacturing and implementation of optics from raw materials all the way to complete microscope systems brings unparalleled optical quality and performance. Complementary optical design means the confocal system, microscope, and objectives all are optimized and matched for superior quality and resolution. Nikon's CFI60 and 75 infinity corrected optical system has numerous options for magnifications, working distances, and immersion mediums, paired for use with an extremely wide variety of samples and specimen preparations.

### CFI Plan Apochromat Lambda S 25XC Sil/40XC Sil

Using silicon oil as the immersion liquid when imaging objects having a refractive index of around 1.4 enables this lens to acquire high quality images with reduced spherical aberration, even with thick samples.

### CFI Apochromat LWD Lambda S 20XC WI/40X WI



This objective corrects chromatic aberrations in a wide wavelength range from visible to near-IR light. With its high numerical aperture, as well as long working distance, it is a powerful tool for imaging thick living specimens.

**CFI Plan Apochromat Lambda D 10X** 

This objective provides superb aberration correction up to the periphery of its large 25 mm field of view, enabling sharp image acquisition of entire samples with digital cameras that have large image sensors. It also corrects chromatic aberration over a wide wavelength range.

### **CFI Plan Apochromat VC 60XC WI**

A water immersion objective that ensures clear image acquisition even within the deep areas of samples. It corrects chromatic aberrations in the shorter wavelength range and is suitable for multicolor confocal imaging.

Maximum intensity projection of Z stack images of marmoset brain acquired with a 60X 1.27 NA water immersion objective using 2048 x 2048 pixel resonant scanning and a DUX-VB detector with user-defined emission bands.

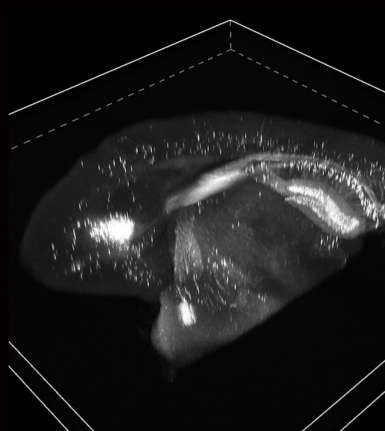
Maximum intensity projection of Z stack images, color-coded by depth, of vascular development in embryonic zebrafish acquired with a 10X 0.45 NA Plan Apo Lambda S objective using 1024 x 2048 pixel resonant scanning. Courtesy of Erika Driekorn and Dr. Beth Roman, Department of Human Genetics, University of Pittsburgh Graduate School of Public Health.

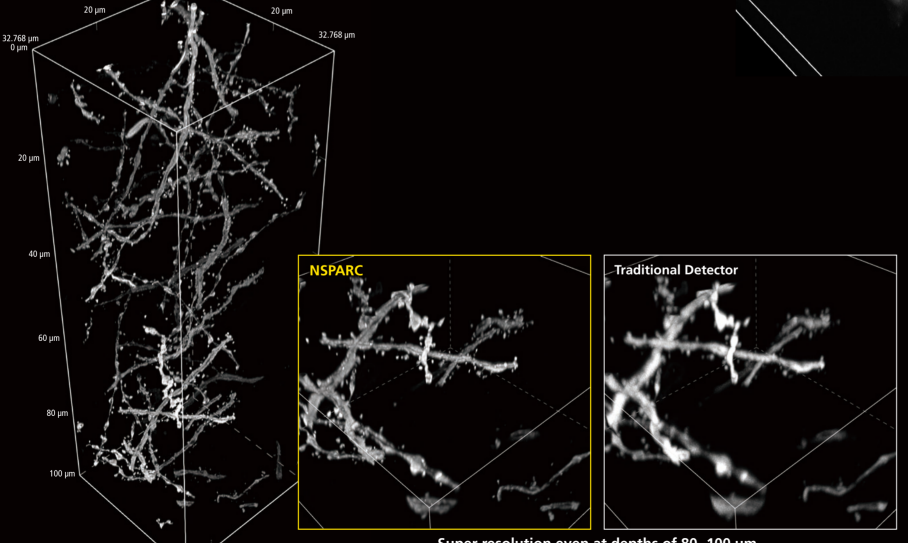
# 3D Confocal Imaging with Super Resolution

NSPARC's array detection enables simultaneous collection of multiple sampling of emission light per pixel, which is beneficial in the Z direction as well as laterally and improves axial resolution.

As a user can optically restrict the emission to the NSPARC detector, confocal imaging can be achieved with a high signal-to-noise ratio, even for 3D observation of thick samples such as cranial nerves, organoids, and organ chips.

Nikon provides various objectives to match specimens, including silicon immersion objectives to minimize refractive index mismatches. Coupled with the AX/AX R's ultra-large 25mm FOV, the system supports a wide selection of objectives capable of obtaining image data from large overviews down to extremely fine details, which can then be measured and analyzed.





Thy1-EGFP mouse neuron (optically cleared) Sample courtesy of: Lin Daniel, PhD. SunJin Lab Co.

Super-resolution even at depths of 80~100  $\mu m$ 

# AX/AX R

# Large FOV confocal



CFI Plan Apochromat Lambda D 4X

# NSPARC Spatial Array Detector Technology

Where a point scanning detector traditionally produces an intensity output only for each pixel NSPARC comprises an array of 25 detectors that operates more like an extremely sensitive camera: two-dimensional spatial information from each scanned pixel is collected by the detector.

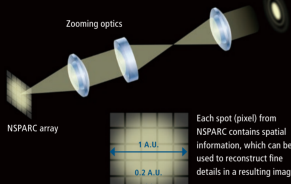
Optical lenses direct emission light to the detector, allowing it to be used with various objectives and magnifications, while simultaneously allowing the user to define the size of the illumination spot on the detector array. This enables oversampling of the conventional single airy unit emission from the confocal plane.

This two-dimensional information is immediately used to obtain ultra-fine structural information, which is lost in conventional detection.

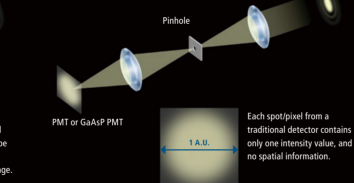
### NSPARC

### **Traditional Detector**

With NSPARC detection, the fluorescence emission light is directed through optical lenses to the detector array, where the projected light can fill the array.



With conventional PMT or GaAsP detectors, fluorescence emission light passes through a variable sized emission pinhole (usually set to 1AU).



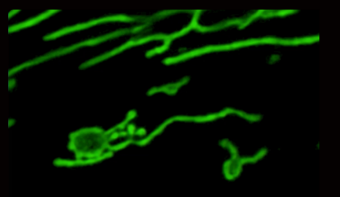
# Extremely Flexible Hardware Configurations

NSPARC detector can be integrated on its own as the only detector for the AX/AX R, or as an additional detector for a multichannel AX/AX R system with either the DUX-VB or ST detector units and an optional transmitted light detector.



# Low Noise, High Sensitivity Detection

The combination of ultra-short dwell time resonant imaging with the AX R resonant scanner and NSPARC's sensitivity allows longer, less phototoxic imaging data to be collected in live-cell assays. As an added benefit, additional spatial information is acquired for every pixel using the detector array, which can be used to further enhance resolution and brightness.

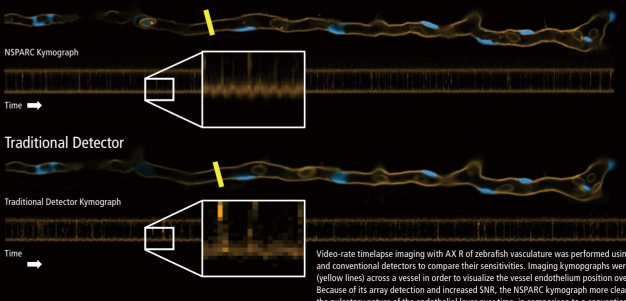


Mitochondrial dynamics captured using AX R resonant scanner coupled with a CFI Plan Apochromat Lambda 100X objective.

# High Speed Imaging with Improved Sensitivity

With the capability to detect single photons, the NSPARC detector's extremely low noise profile and exceptional sensitivity make it a perfect match for AX R confocal resonant high-speed imaging, where the pixel dwell time is as short as 200 nanoseconds, enabling image acquisition at video frame rate. More details can be extracted from images for downstream analysis and computation.

### NSPARC



Video-rate timelapse imaging with AX R of zebrafish vasculature was performed using NSPARC and conventional detectors to compare their sensitivities. Imaging kymopgraphs were created yellow lines) across a vessel in order to visualize the vessel endothelium position over time. Because of its array detection and increased SNR, the NSPARC kymograph more clearly displays the pulsatory nature of the endothelial layer over time, in comparison to a conventional GaAsP detector

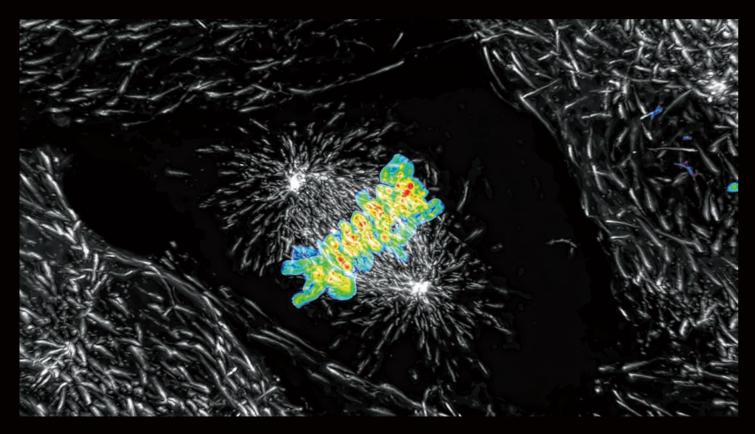
# Comprehensive Imaging N/S Elements Software

NIS-Elements Imaging Software allows integrated control of microscopes and peripheral devices, as well as confocal systems. In addition to various functions for confocal imaging, it has a wide range of AI tools that support streamlining of image analysis and optional modules that enable customization of analysis and experiment workflows.



## **ER (Extended Resolution)**

NIS-Elements ER can be used to improve confocal spatial resolution up to 120 nm (lateral)/300 nm (axial) using GPU-processing with automatic parameter settings and user-defined options.



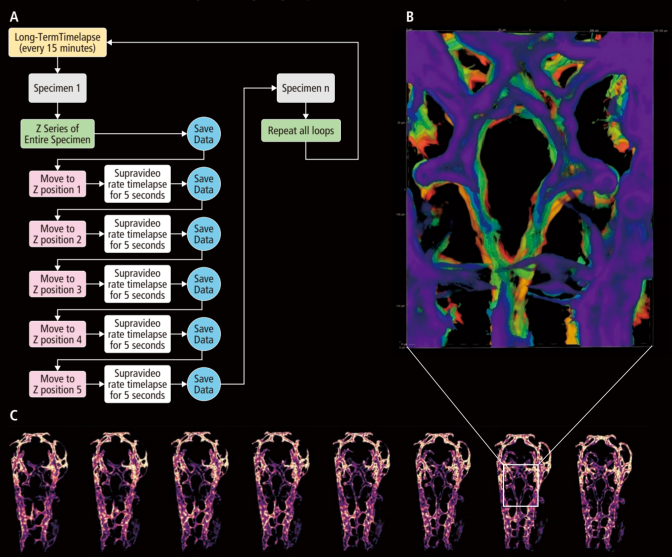
Maximum intensity projection of Z stack images of a live sample to which ER has been applied, acquired with a 60X 1.4 NA Plan Apo Lambda oil immersion objective using 2048 x 1024 AX R resonant scanning at 15 fps.



# **Customized Definition of Experiments**

NIS-Elements has built-in multidimensional (multi-XY,Z,T, multichannel) experiment capabilities. Adding the optional JOBS module allows even more customization such as setting up non-orthogonal experiments with multiple paths and dimensions. Oftentimes, experiments require customization to streamline acquisition and capture all necessary data points.

Analysis of data can be done even in real-time during the experiment, and the direction of the experiment can even be changed based on the results of the analysis. Users have ultimate flexibility in designing experiments that maximize their data output needs.



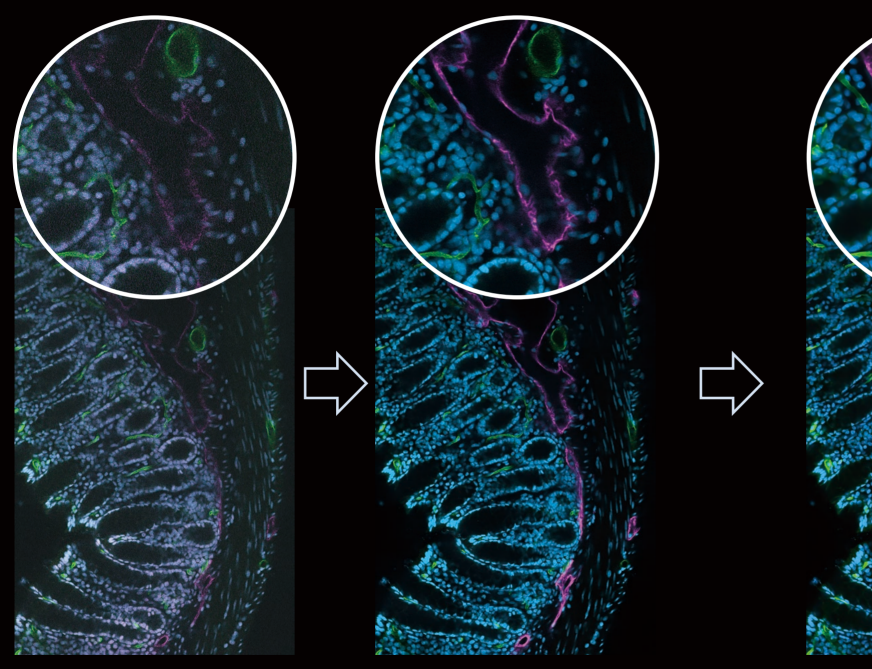
Time lapse images of Danio sp. vascular development acquired with a 25X SIL immersion objective using 1024 x 2048 AX R resonant scanning.

(A) JOBS experiment protocol shown above.

(B) Inset showing vascular development with overlay color representing time, where each color represents a different time point. (C) Maximum intensity projections of entire FOV of Z series at points in time during the progress of the experiment. Courtesy of Erika Driekorn and Dr. Beth Roman, Department of Human Genetics, University of Pittsburgh Graduate School of Public Health.

# Al Software Innovations Designed to Assist

From acquisition to analysis, Nikon's NIS-Elements software is a pioneering leader in the implementation of convolutional neural network (CNN) based deep learning for microscopy. Several AI tools are available, many targeted specifically for assisting users in acquiring, processing, and analyzing confocal data. These tools aid users in achieving adequate signal-to-noise ratio (SNR) images for image processing and analysis, and more tools for both segmentation and image enhancement or modality transformation.



## **Starting Point**

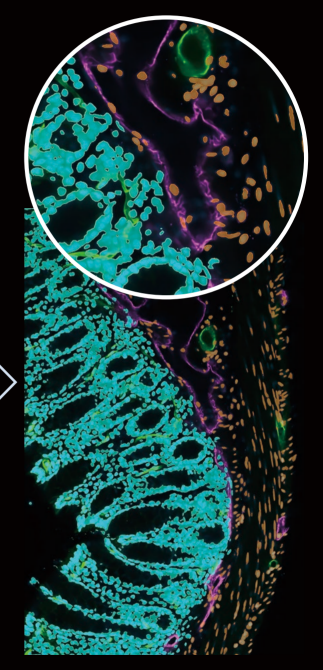
Confocal imaging has multiple variables that must be fine-tuned for the best image quality, a statistically valid signal-to-noise ratio, and longterm sample stability. NIS-Elements AI tools are designed to assist in achieving these targets.

## Autosignal.a

New for AX/AX R: Autosignal.ai can suggest the best illumination and detection settings automatically, instead of users manually attempting to find the best settings by trial and error, or while scanning live and exposing the sample unnecessarily.

### Denoise.a

Shot noise is the main noise source in confocal imaging. Denoise.ai can remove the shot noise component from confocal images, improving the image quality and assisting in downstream segmentation.



### Segment.ai

A toolbox of AI functions assist users in easy segmentation of images; after training the AI, segmentation that would take hours by traditional methods (such as samples with uniform intensity, making traditional thresholding of different morphologies nearly impossible) can be done in seconds.

# **Optional Components**



### Water Immersion Dispenser

A software-controlled, automatic water dispenser enables longterm time-lapse imaging using refractive-index matching water immersion objectives in any environment, including incubation.

### Automatic Correction Collar

Moves the objective correction collar to the optimum position for best resolution both remotely and by software control. Motorized collars allow users to adjust the correction collar without disturbing the specimen position, even in incubated enclosures or environmental chambers.

# **Multiple Modalities**

### LAPP Modular illumination system

The Ti2-E microscope supports up to 5 episcopic illumination sources, which can be used in tandem with AX/AX R confocal imaging: total internal reflection fluorescence (TIRF), point, raster or field stimulation devices, and fluorescence light sources can all be integrated onto the same microscope stand, and used in the same experiments.

### Total Internal Reflection Fluorescence (TIRF)

The incident angle of a laser and corresponding penetration depth of the evanescent field can be controlled via NIS-Elements software. When multiple TIRF modules are mounted, the penetration depth can be independently set for each wavelength.

### Photostimulation: Point and Raster Scanner

The XY galvano scanning unit can stimulate the desired area of a sample using laser point scanning. It allows simultaneous photostimulation and confocal imaging.

### Photostimulation: Digital Micromirror Device (DMD)

The DMD module enables photoactivation of userspecified patterns rather than photoactivation of a single spot. This allows stimulation of multiple points and tracking of their behavior. The DMD module can be used with either laser illumination or less phototoxic LED illumination.

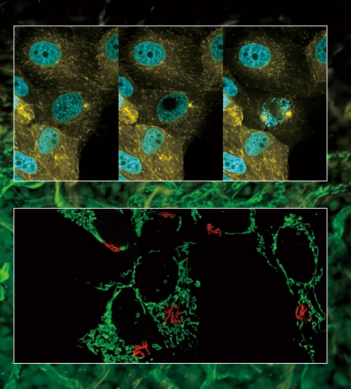
### Opti-Microscan photostimulation device

This device enables photostimulation over a 400 to 700 nm wavelength range\*, allowing photostimulation IR imaging with visible light. Simultaneous stimulation, sequential stimulation and manual stimulation are available. \* Depends on filter cube type.



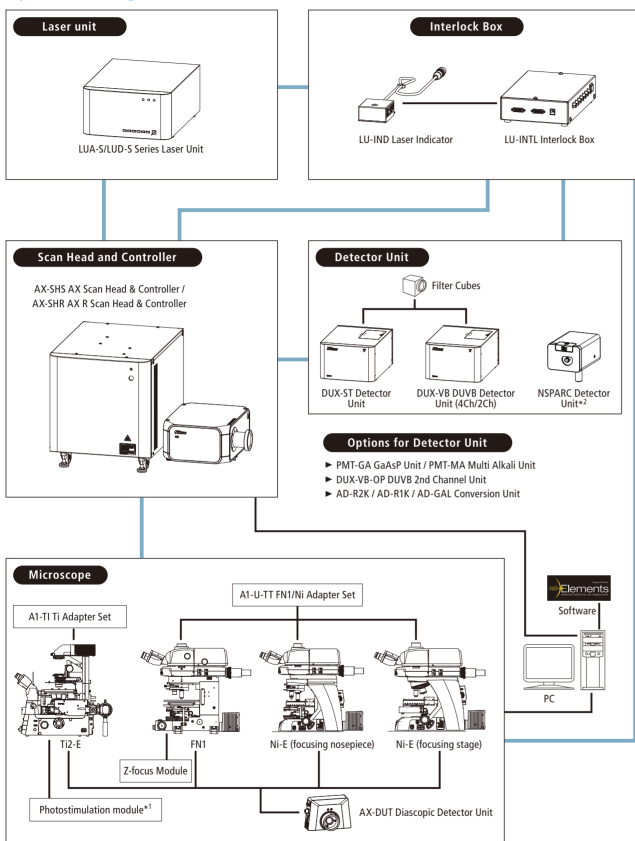








# System diagram



<sup>\*1</sup> Adapter for Ti2-LAPP system, light source, hybrid dichroic mirror for photostimulation and imaging, and control board are all required. \*2 Compatible only with TI2-E and Ni-E (cannot be configured for FN1).

# **Specifications**

-		
Scan Head	AX Galvano Scanner	25 mm FOV galvano scanner Up to 8192 x 8192 pixels Up to 10 fps at 512 x 512 pixel Pixel dwell time up to 2 millise Supports bidirectional imaging
	AX R Galvano+Resonant Scanner 2K or 1K	25 mm FOV resonant scanner 25 mm FOV galvano scanner Up to 2048 x 2048 pixels for 2 Up to 720 fps at 2048 x 16 pix 30 fps at 2048 x 512 pixels for Supports bidirectional imaging
Scan Head Input/Output Port		One FC fiber laser input port a An additional FC fiber laser inp
Dichroic Mirror		Up to 6 customizable mirrors
Pinhole		Variable aperture with 6 blade
FOV		Maximum 25 mm diameter (ci
Laser		Up to 8 visible lasers Compatible range: 405-750 nr
Detector	DUX-VB Detector Unit	2 or 4 channels Freely tunable emission bands Up to 12 bandpass filters Multi-alkali PMT or GaAsP PM
	DUX-ST Detector Unit	2 or 4 channels Up to 18 bandpass filters Multi-alkali PMT or GaAsP PM
	NSPARC Detector Unit	Lateral resolution 100 nm* <sup>1</sup> , A Equipped with SPPC (Single Pi Up to 7 barrier filters can be n 561 nm, 594 nm and 640 nm) With galvano scanner: Can be 8192 pixels With resonant scanner: Can b of 128 to 1024 pixels <sup>*2</sup>
Diascopic Detector		
	lector	Compact PMT detector
Z step		
Z step Compatible N		Ti2-E: 0.01 μm, 0.02 μm (with Ti2-E inverted microscope with
		<ul> <li>Ti2-E: 0.01 μm, 0.02 μm (with</li> <li>Ti2-E inverted microscope with</li> <li>Ni-E and FN1 upright microscope</li> <li>Photostimulation (point raster</li> <li>Fluorescence lifetime imaging</li> <li>Piezoelectric Z (or XYZ)</li> <li>Environmental chamber or end</li> </ul>
Compatible N		<ul> <li>Ti2-E: 0.01 μm, 0.02 μm (with</li> <li>Ti2-E inverted microscope with</li> <li>Ni-E and FN1 upright microscope</li> <li>Photostimulation (point raster</li> <li>Fluorescence lifetime imaging</li> </ul>
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\*1 These values were measured using 40 nm diameter beads for lateral resolution and 100 nm diameter beads for axial resolution with excitation by a 488 nm laser. Actual resolution depends on laser wavelength and optical configuration.

\*2 Resolution of 2048 pixels cannot be set.

pixels illiseconds aging and linescan imaging

ner for 2K (1024 x 1024 pixels for 1K) 16 pixels for 2K (720 fps at 1024 x 16 pixels for 1K) els for 2K (1024 x 512 pixels for 1K) aging and linescan imaging

ort and one FC fiber signal output port er input port and FC fiber signal output port can be optionally added

blades

er (circle) inscribed by a rectangle

50 nm

ands with  $\pm$  1 nm accuracy and up to 66 discrete spectral channels

PMT options

PMT options

\*<sup>1</sup>, Axial resolution 300 nm\*<sup>1</sup>

le Pixel Photon Counter) array detector

be mounted (supports excitation at 405 nm, 445 nm, 488 nm, 514 nm, ) nm)

an be used with X resolution of 64 to 8192 pixels, Y resolution of 128 to

an be used with X resolution of 256, 512 and 1024 pixels, Y resolution

with encoder control), FN1 stepping motor: 0.05 µm, Ni-E: 0.025 µm

with maximum FOV of 25 mm roscopes with maximum FOV of 25 mm

aster or digital micromirror) ging including fast FLIM

enclosure TIRF, N-STORM or N-SIM S

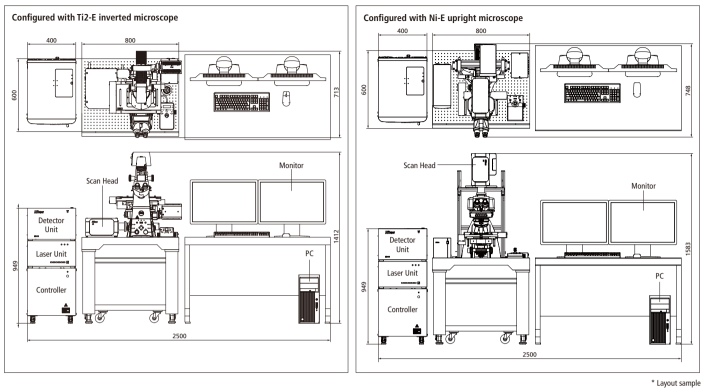
36 gray levels with fine integration) with multiple file output options

4bit Professional with GPU-accelerated graphics card

umidity 70% RH or less (no condensation)

# Layout

Unit: mm



Specifications and equipment are subject to change without any notice or obligation on the part of the manufacturer. November 2022 @2022 NIKON CORPORATION

🔥 WARNING

TO ENSURE CORRECT USAGE, READ THE CORRESPONDING MANUALS CAREFULLY BEFORE USING YOUR EQUIPMENT.

#### Monitor images are simulated.

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WARNING-LASER RADIATION AVOID EXPOSURE TO BEAM

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