

Product Information Version 1.0

# ZEISS LSM 880 with Airyscan

Revolutionize Your Confocal Imaging



### **Revolutionize Your Confocal Imaging with Airyscan**

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Your samples tend either to be very small, move very fast or bleach very quickly. Or do all of that at once. To get unbiased data from live cells or other weakly abelled samples, there's no such thing as too much sensitivity, resolution or peed. Each photon of emission light is precious.

Now you can use multicolor samples with any label and get image quality like you've never seen before. With Airyscan you are always able to select the optimal acquisition strategy for your sample: Simply decide whether you want to gain 1.7x higher resolution in all three dimensions – resulting in a 5x smaller confocal volume. Or push the sensitivity beyond the limits of all conventional confocals. Or use the increase in signal-to-noise ratio to speed up your image acquisition. The choice is yours.



HeLa cells, Actin stained with Phalloidin-Alexa 546,
 AP3 with Alexa 488, DAPI. Courtesy of S. Traikov, BIOTEC,
 TU Dresden, Germany



See for yourself how Airyscan gives you better data than ever before. Book a hands-on demonstration in one of our ZEISS Microscopy Labs now. >> www.zeiss.com/Ism880

### Simpler. More Intelligent. More Integrated.

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# Airyscan: Enter a New World of Confocal Performance

Imagine a true confocal that could give you high sensitivity, improved resolution in x, y and z, and high speed – all that in a single system.

With Airyscan you will be increasing the resolution of your imaging far beyond the resolution of a classic confocal point scanning microscope.

You can resolve 140 nm laterally and 400 nm axially, at 488 nm – without sacrificing sensitivity or speed. In the optional Virtual Pinhole Mode, you can decide even after the acquisition, which pinhole size best suits your application.

### Perform Quantitative Imaging

Scientific results depend on unbiased data. LSM 880 guarantees gentle imaging of your sample through homogeneous illumination, combining linear scanning with a sensitive detection infrastructure.

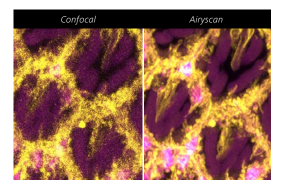
Working with identical pixel times and continuous scanner monitoring, you can perform quantitative imaging at all speeds and scan modes.

Get robust and reliable results, even from your most demanding single molecule imaging and analysis.

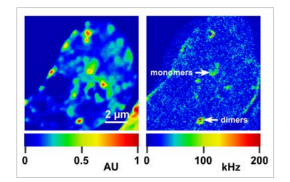
#### **Increase Your Productivity**

Save time on investigations into localization and interaction of proteins that require multiple fluorescent labels. LSM 880 collects all these signals in one go, with high speed and high sensitivity. You perform simultaneous spectral detection in a single scan with the highest number of descanned or non-descanned channels – featuring GaAsP technology, too.

LSM 880 lets you take full advantage of large fields of view and the highest speed of any linear scanning confocal – up to 13 fps.



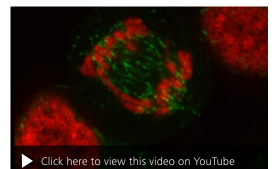
Drosophila retina, labelled for actin (AF 488) shown in magenta and Na,K-ATPase (GAR-CF 633) shown in yellow. Maximum projection of a 10 µm Z-stack, Plan-APOCHROMAT 20x/0.8



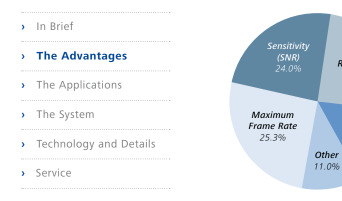
Heterochromatin protein 1 (HP-1) fused to GFP and expressed in the nucleus of a human Hep G2 cell.

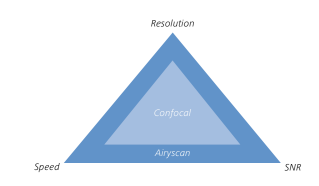
*Left panel* shows the distribution of *HP-1* between Euchromatin and denser Heterochromatin areas.

**Right panel** represents a brightness map demonstrating dimerization of HP-1 within heterochomatic regions. Sample: courtesy of P. Hemmerich, Leibniz-Institute for Age Research (FLI), Jena, Germany



You Mitosis in HeLa-Kyoto cell line, Histone 2B (H2B, red, mCherry) and microtubule end binding protein 3 (EB3, green, EGFP) during mitosis. Sample: courtesy of J. Ellenberg, EMBL, Heidelberg, Germany



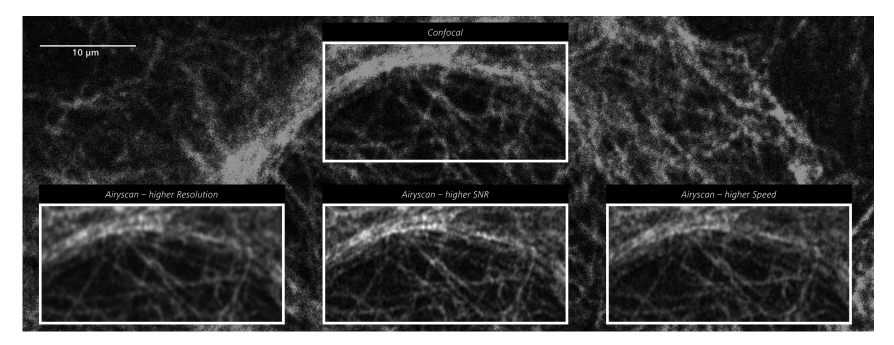


A survey among 250 researchers working with confocal microscopes revealed that their imaging would benefit most from an increase in sensitivity, resolution and speed.

Resolution in Z

24.7%

Airyscan extends exactly those parameters for your experiments and even allows combinations, such as increased resolution and signal-to-noise-ratio (SNR) at a given speed.



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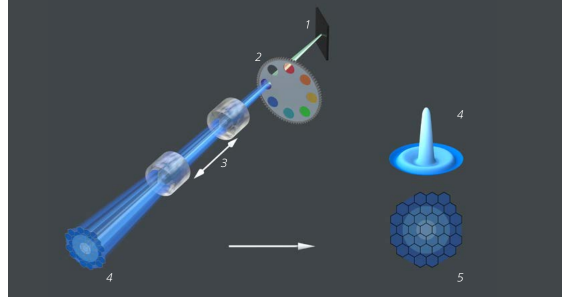
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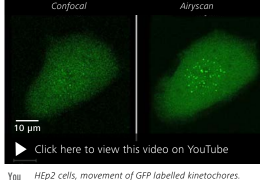
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### The Airyscan Principle

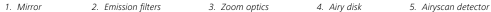
| The beampath of every microscope transforms the emission signal of even an infinitesimally small spot into a complex three-dimensional structure called an Airy disk |
|--|
| or Airy pattern. This transformation is known as point spread function (PSF).  |

A classic confocal microscope illuminates one spot on your sample to detect the emitted fluorescence signal. Out-of-focus emission light is rejected at a pinhole, the size of which determines how much of the Airy disk reaches the detector. You can increase the resolution by making the pinhole smaller, but signal-to-noise drops significantly since less valuable emission light is passing through. With Airyscan ZEISS introduces a new concept. Instead of throwing light away at the pinhole, a 32 channel area detector collects all light of an Airy disk simultaneously. Each detector element functions as a single, very small pinhole. Knowing the beampath and the spatial distribution of each Airy disk enables a very light efficient imaging: you can now use all of the photons that your objective collected. It's up to you whether you use the additional information from your sample to get better signal-to-noise, resolution or speed. In the optional Virtual Pinhole Mode, you can decide even after the acquisition, which pinhole size best suits your application.

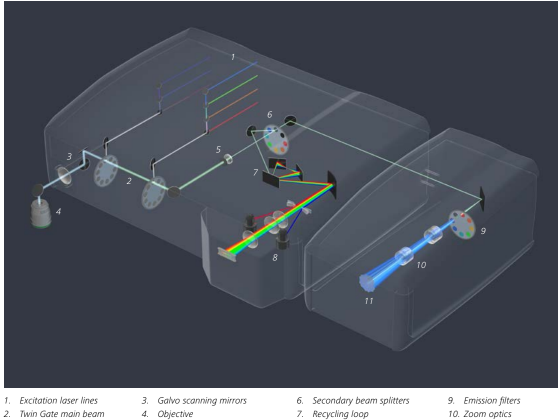




Tube Courtesy of V. Döring, Leibniz-Institute for Age Research (FLI), Jena, Germany



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- splitters
- 5. Pinhole and pinhole optics
- 8. Quasar detection unit
  - 11. Airyscan detector

### Beampath of LSM 880 with Airyscan

Emission light travels through the Twin Gate main dichroic beam splitter with its very efficient laser suppression to deliver supreme contrast. Then, at the secondary beam splitter, all emission light either travels via the recycling loop to the internal spectral detection unit (Quasar) with up to 34 channels. Or, light is sent to the revolutionary Airyscan detector with GaAsP technology.

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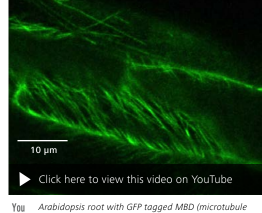
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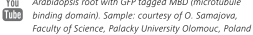
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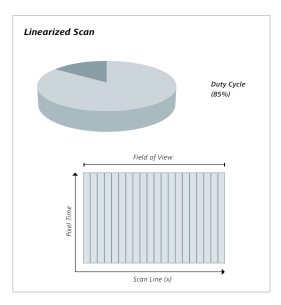
# To fully resolve the movement of labeled proteins in dynamic cellular and subcellular processes you often need to image at around 10 frames per second. Now, with LSM 880, you can achieve up to 13 frames per second at 512 x 512 pixels.

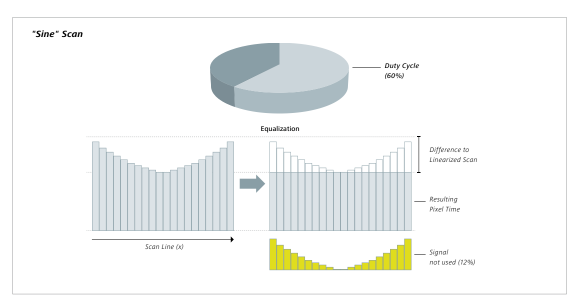
Fast and Linear Scanning - Your Powerful Combination

While you're performing unidirectional or bidirectional scanning, LSM 880 is constantly monitoring and calibrating the scanner position. This guarantees a stable field of view and equal pixel integration times over the whole field of view. Linear scanning is an essential prerequisite for your quantitative and correlative imaging. It gives you a constant signal-to-noise level and uniform exposure to the illuminating laser through-out the scanned area, including manipulated regions of interest. Unlike traditional sine scanning confocals, LSM 880 uses more than 80% of the scanning time for data acquisition. That means you will enjoy a 29% improvement of signal-to-noise ratio because of longer pixel integration times at a defined frame rate. You can't always influence the orientation of your structure of interest as regards to the detection optics, but with LSM 880 you can always adapt the scanfield and rotate it freely to best suit your sample.









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### Parallel Acquisition of Multiple Channels

It takes multiple labels to analyze interactions between different cellular or subcellular structures, but you can achieve the highest timing precision and speed up your imaging time by recording their intensities simultaneously. LSM 880 lets you acquire the entire spectrum – and all your labels – in just one scan with 32 channels, 512 x 512 pixels at 5 frames per second.

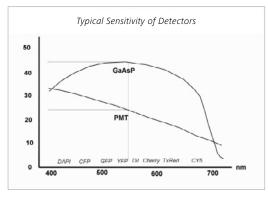
Set up 10 channels for multichannel spectral imaging and then add the transmission detector. You can now image all fluorescent dyes and the additional oblique contrast in a single scan. This protects your sample and also saves you time.

Especially for your demanding multiphoton experiments, you will profit from having this fundamental capability: up to 12 non-descanned detectors can be read out in parallel.

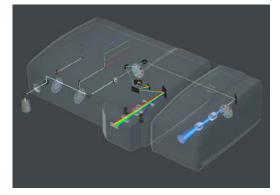
GaAsP detectors have proven to be the best choice for confocal imaging of weak fluorescent signals. In photon counting mode you can use them for single molecule techniques such as fluorescence correlation spectroscopy (FCS) and cross correlation spectroscopy.

### Benefit from the Most Spectral Confocal

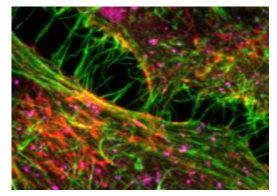
Investigations into localization and interaction of proteins often require multiple fluorescent labels. Now you can save time and collect all these signals in one go, with high speed and high sensitivity. LSM 880 lets you perform spectral detection with any number of descanned or non-descanned channels in a single scan.



Typical spectral quantum efficiency (QE) of PMT and GaAsP detectors.



Beampath LSM 880 with Airyscan



HeLa cells, Actin stained with Phalloidin-Alexa 546, AP3 with Alexa 488, DAPI. Courtesy of S. Traikov, BIOTEC, TU Dresden, Germany

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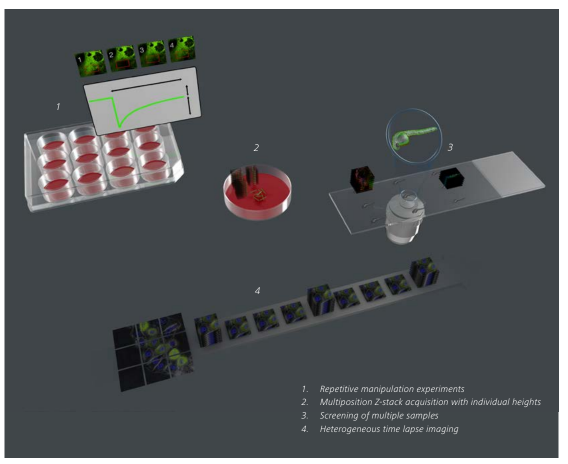
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### Experiment Designer: Your Smart Automation Module for Enhanced Productivity

Sometimes your applications require complex acquisition strategies. Especially for statistical analysis, repetitive imaging of a large number of samples with different imaging set ups comes into play. Experiment Designer is an easy-to-use module for ZEN imaging software that sets up your imaging for multiple positions, using the large number of imaging modalities of LSM 880. It is complemented by a number of hardware and software options so your sample always stays in focus, even during the most demanding long-term time lapse experiments.





With the ZEN software module Experiment Designer you can set up complex imaging routines consisting of freely defined and repeatable experiment blocks with multi-position tile scans of multichannel Z-stacks.

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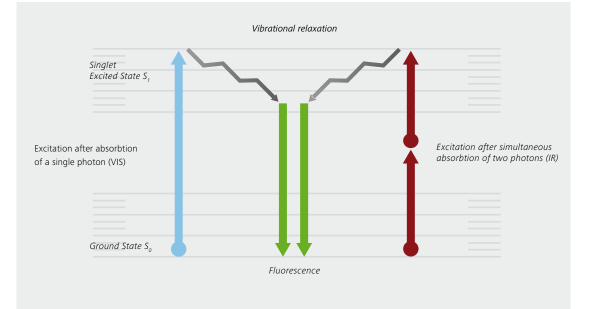
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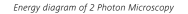
### **Multiphoton Microscopy**

Multiphoton microscopy lets you acquire optical sections of deep tissue layers. This imaging method makes use of three basic principles:

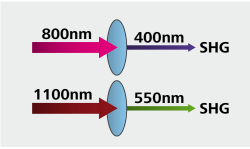
- The longer the wavelength of light, the less it is scattered when entering tissue. Therefore, excitation light in the near-infrared range penetrates deeper into biological samples than visible light can.
- Light of a wavelength between 600 to 1300 nm experiences the lowest absorption in tissue, making it nearly transparent in this spectral range.
- A fluorescent dye with an excitation maximum at 500 nm can be excited with one photon of the same wavelength. Or with two photons of the doubled wavelength – 1000 nm – that arrive simultaneously.

A powerful pulsed laser at wavelengths of 700 to 1300 nm makes sure that enough photons arrive simultaneously to excite the fluorescent dye. Outside the focal plane, the laser intensity drops exponentially and produces no excitation. Another effect allows you to visualize non-stained tissue structures when excitation light of very high intensity interacts with tissue by doubling its frequency.





The nonlinear effect of frequency doubling (SHG) on non-centrosymmetric molecules with predominantly periodic alignment occurs, for example, in striated muscle and collagen.



Simplified mechanism of second harmonic generation.

### > In Brief

### Superresolution Microscopy

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LSM 880 is the only confocal laser scanning microscope that can be combined with three complementary superresolution techniques in one system, delivering true multimodal imaging of your samples. Your LSM 880 with Airyscan can deliver resolutions down to 140 nm laterally and 400 nm axially even in thicker and denser samples. Acquire even smaller structures by combining it with ELYRA, the dedicated superresolution system for structured illumination (SR-SIM) and photoactivated localization microscopy (PALM). SR-SIM works best with thinner, less scattering samples and delivers large fields of view with a resolution down to 120 nm laterally and 350 nm axially. PALM uses photo-switchable fluorescent molecules, recording them over time and then superimposing these data. This lets you achieve resolutions down to 20 nm laterally and 50 nm axially.



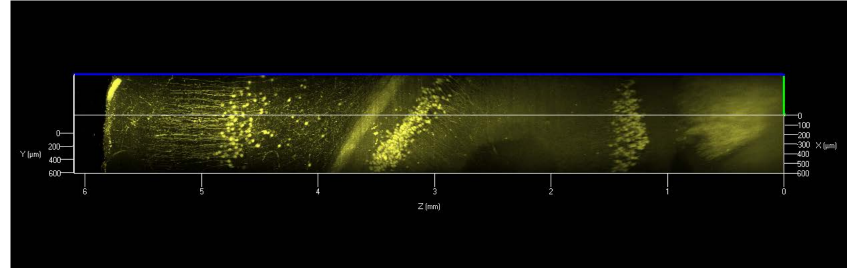
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### **Clearing Methods**

Tissue clearing opens up a new dimension of optical penetration depth into biological samples such as tissue sections, mouse brains, embryos, organs, spheroids or biopsies.

With Axio Examiner and LD Plan-APOCHROMAT 20x/1.0, you can look deep into tissue that has been treated with the Scale method which has a refractive index of 1.38. That lets you image up to six times deeper than with a multiphoton microscope and up to 60 times deeper than with a conventional laser scanning microscope. Get ready to be impressed by the quality of structural information you retrieve from the deepest layers: expect a big push forward, especially in basic neurobiological research and mapping of neuron networks.



Maximum intensity projection, brain of 7-week old YFP-H mouse, fixed and cleared with Scale clearing technique (Hama et al, Nat Neurosci. 2011). Courtesy of H. Hama, F. Ishidate, A. Miyawaki, RIKEN BSI, Wako, Japan

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As your needs grow, LSM 880 grows with you, forming the basis for a number of enhancements. Like every system from ZEISS, open interfaces and a modular architecture guarantee the seamless interaction of all components now and in the future. These include:



Combine Axio Observer with incubation to get the best tool for long term live cell imaging with stable temperature conditions.



Axio Examiner is your microscope stand for LSM 880 multiphoton systems. Combine the system with Airyscan or incubation for your demanding experiments with living specimens.



The upright research microscope Axio Imager can be combined with LSM 880, Airyscan and incubation, too.



Additional accessories such as a Z piezo stage and a leveling insert are recommended for superresolution Airyscanning.



Use BiG.2 with its two GaAsP detectors for FCS, photon counting experiments and FLIM.



BiG.2 works perfectly as a non-descanned detector, also providing a highly sensitive direct coupled detector for FLIM.

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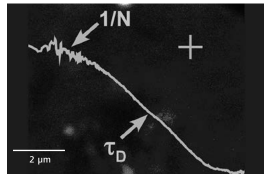
The GaAsP NDD two channels with flexible filter settings completes the ensemble of non-descanned detectors for Axio Examiner.



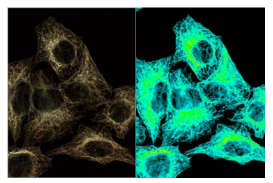
Airyscan can be added to any of the LSM 880 system configurations, including the version with BiG.2 detector.



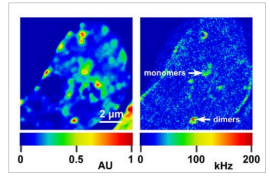
You can add a choice of cameras from the Axiocam series to LSM 880 for widefield imaging experiments – also in combination with LSM imaging.



Fluorescence correlation spectroscopy is the established method mainly for kinetic studies on the single molecule level – it works easily with GaAsP detectors.



Förster Resonance Energy Transfer (FRET) and Fluorescence Recovery After Photobleaching (FRAP) are methods to study molecule interaction and motion.



With the Number&Brightness analysis tool you evaluate the connection between the intensity in your sample and the number of molecules responsible for this signal intensity.

# **Tailored Precisely to Your Applications**

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| Typical Applications, Typical Samples  | Task  | ZEISS LSM 880 offers   |
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| Antibody stained tissue slices   | Document morphological relations of structures with a resolution of 140 nm (xy) / 400 nm (z) at 488 nm excitation | Airyscan with GaAsP detector for superresolution imaging   |
| Tissue cleared with scale solution   | Image cleared tissue with up to 5.6 mm in Z   | Special objective corrected for immersion medium of refractive<br>index 1.38 working with confocal or multiphton imaging on<br>Axio Examiner |
| Live cell culture  | Study the motility of vesicles and organelles   | Up to 13 frames per second time lapse imaging  |
|  | Document the kinetics of endo and exocytosis  | Mixed mode imaging with LSM and superresolution based on photoactivated localization   |
|  | Screen and document cells expressing the desired fluorescent label<br>in response to pharmacological treatment    | Widefield imaging using Axiocam  |
| Live cell culture with two labels  | Study the motility of subcellular structures  | Airyscan with GaAsP detector to image 2 colors with time lapse imaging in 2D or 3D at 2.5 frames per second                                  |
|  | Explore the interaction of two proteins with fluorescent lifetime microscopy                                      | BiG.2 as detector for FLIM and third party electronics and software  |
|  | Explore the interaction of two proteins exploiting the Förster Resonance Energy Transfer effect                   | FRET analysis tool   |
| Live cells with multiple labels  | Image over long time in an automated way  | Experiment Designer software tool combined with spectral imaging   |
| Fixed cell culture specimens   | Document cellular structures in superresolution in 3D with about 2x the resolution of a confocal                  | Structured illumination with ELYRA   |
| Live or fixed cells with multiple labels<br>and overlapping emission signals | Examine the interplay of multiple proteins  | Parallel acquisition of all signals with spectral imaging<br>at 5 full frames per second and online or post processed<br>unmixing            |
| Cellular structures with weak labels   | Image subcelluar structures at physiological expression levels  | Airyscan with GaAsP detector or LSM 880<br>with GaAsP detector   |
| Living organisms/animals   | See the interaction of cells within living tissue   | Multiphoton extension of LSM 880   |
|  | Imaging of live tissue with cells expressing multiple different fluorescent proteins                              | Multiphoton extension of LSM 880 with OPO  |

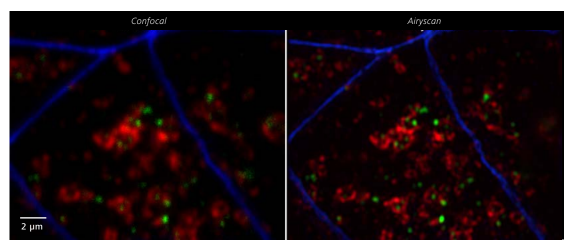
# **Tailored Precisely to Your Applications**

| > In Brief                                 | Typical Applications, Typical Samples                                  | Task  | ZEISS LSM 880 offers  |
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| > The Advantages                           | Plant roots  | Follow the changes of subcellular structures over time with a high resolution                   | Airyscan with GaAsP detector for superresolution imaging<br>beyond 40 µm deep into tissue with up to 5 full frames per        |
| > The Applications                         | Model organisms, e.g. Zebrafish, Drosophila or C. elegans              | See fine details of the organisation and dynamics   | second (512 x 512) Airyscan with GaAsP detector for superresolution imaging   |
| > The System                               |  | of endogeneously expressed FP proteins  | beyond 40 µm deep into tissue   |
| <ul> <li>Technology and Details</li> </ul> | Live samples with varying labelling intensities over the field of view | Collect all image information and decide on the way to present the best image in contrast later | Airyscan with GaAsP detector for imaging with 3 Airy units<br>and the abiltiy to virtually close the pinhole post-acquisition |

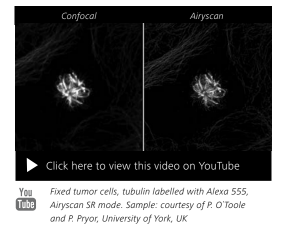
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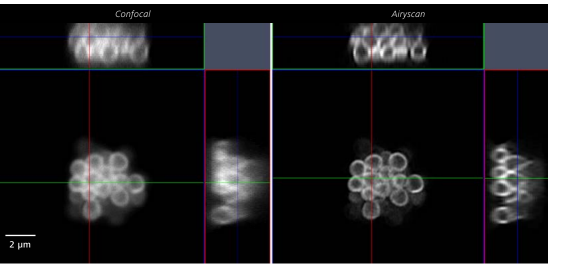
# ZEISS LSM 880 at Work

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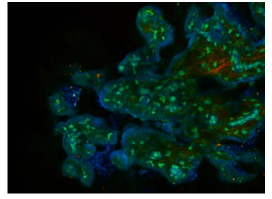


Human RPE cells, ZO1 (tight junction marker) in blue, photoreceptor outer segments stained with FITC in green, EEA1 (endosomal marker) in red. Courtesy of S. Almewadar, CRTD, TU Dresden, Germany





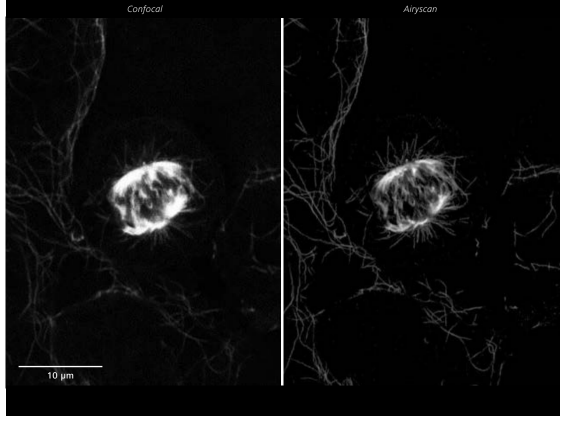
Fluorescent 1µm ringbeads imaged at 488 nm



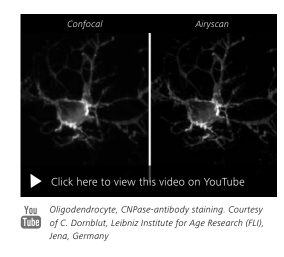
Skin tissue from pig labelled with Ethyleneblue. The unfixed sample was excited with 1100 nm using an OPO (optical parametric oscillator). Fluorescent lifetime measurement was performed using the detector module BiG.2 connected to the TCSPC electronics from Becker&Hickl. The color coded image shows the variation in lifetime within different types of skin cells.

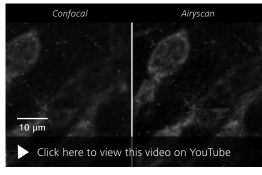
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Fixed tumor cells, tubulin labelled with Alexa 555, Airyscan SR mode. Sample: courtesy of P. O'Toole and P. Pryor, University of York, UK





You Slice of mouse brain, CNPase-antibody staining, imaged with 10x objective. Courtesy of C. Dornblut, Leibniz Institute for Age Research (FLI), Jena, Germany

### ZEISS LSM 880 at Work

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The cells are IMR90 human diploid lung fibroblasts. The DNA has been stained with DAPI, the telomeric G strand (leading strand) in green with a Peptide Nucleic Acid probe and Alexa 488 and the telomeric C strand (lagging strand) in red with a Peptide Nucleic Acid probe and Alexa-546. Prior to their harvest the cells have been treated with siRNAs targeting RTEL1 (2). RTEL1 is a helicase that is essential for telomere replication, and lack of the protein leads to stalled forks at telomeres and telomere breakage. This can be seen by individual telomeres that appear as more than one dot, as highlighted in the images. Airyscan resolves multiple telomere dots, thereby allowing an accurate quantification of telomere replication problems. Sample: Courtesy of J. Karlseder (Molecular and Cell Biology Laboratory) and J. Fitzpatrick (Waitt Advanced Biophotonics Core), The Salk Institute, La Jolla, USA.

# **ZEISS LSM 880: Your Flexible Choice of Components**

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### 1 Microscope

- Inverted stand: Axio Observer
- Upright stand: Axio Examiner, Axio Imager
- Port for coupling of ELYRA
- Camera port
- Manual or motorized stages
- Incubation solutions
- Fast Z piezo inserts
- Definite Focus

### 2 Objectives

- C-APOCHROMAT
- Plan-APOCHROMAT
- W Plan-APOCHROMAT, LD Plan-APOCHROMAT
- EC Plan-NEOFLUAR

### 3 Illumination

- UV laser: 355 nm, 405 nm
- VIS laser: 440 nm, 458 nm, 488 nm, 514 nm, 543 nm, 561 nm, 594 nm, 633 nm
- NIR laser for multiphoton imaging: Ti:Sa, OPO

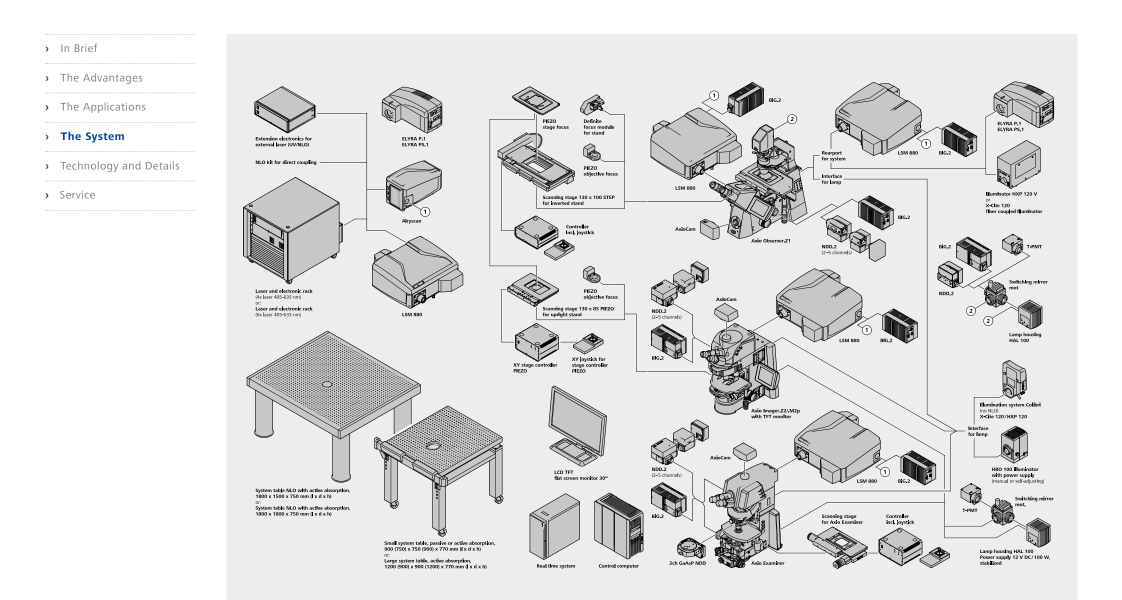
### 4 Detection

- 3 or 34 descanned spectral channels (GaAsP and/or PMT)
- Airyscan detector
- 2 additional GaAsP channels (BiG.2)
- Up to 6 non-descanned GaAsP detectors
- Up to 12 non-descanned GaAsP or PMT detectors total
- Transmitted light detector (T-PMT)

### 5 Software

 ZEN (black edition), recommended modules: Tiles & Positions, Experiment Designer, FRAP, FRET, RICS, FCS, Deconvolution, 3D VisArt

# **ZEISS LSM 880: System Overview**



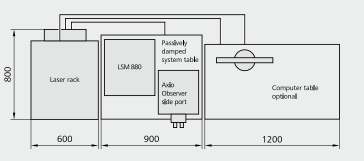


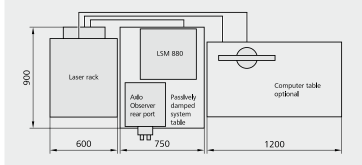
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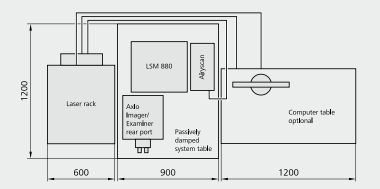
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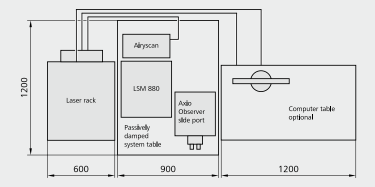
LSM 880 on Small System Table

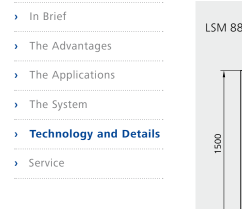




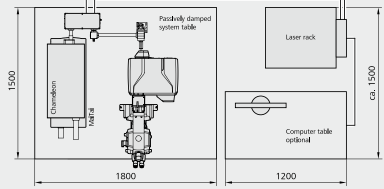
### LSM 880 with Airyscan on Large System Table



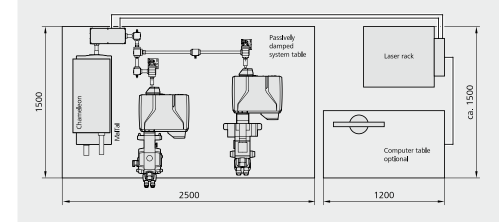




LSM 880 Equipped with Two Photon Laser (NLO) for Single Stand

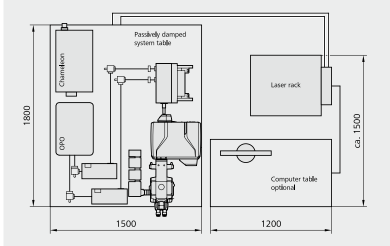


LSM 880 Equipped with Two Photon Laser (NLO) for Dual Stand

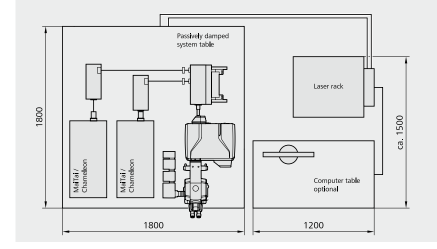


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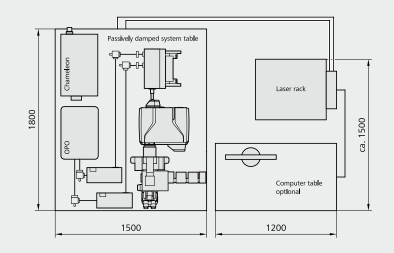
LSM 880 NLO on Axio Examiner Equipped with NLO Laser and OPO



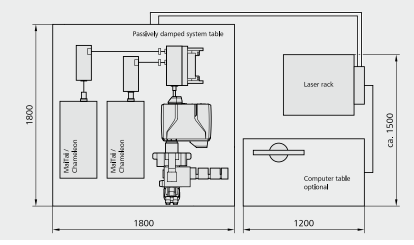
### LSM 880 NLO on Axio Examiner Equipped with Two NLO Lasers



### LSM 880 NLO on Axio Observer Equipped with NLO Laser and OPO



### LSM 880 NLO on Axio Observer Equipped with Two NLO Lasers



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| Physical Dimensions   | Length (cm) | Width (cm) | Height (cm) | Weight (kg) |
|---|-------------|------------|-------------|-------------|
| Small passively damped system table                                   | 90          | 75         | 77          | 80          |
| Small actively damped system table                                    | 90          | 75         | 77          | 90          |
| Large actively damped system table                                    | 120         | 90         | 77          | 120         |
| Active anti-vibration table (NLO)<br>for Mai Tai Laser or Chameleon   | 180         | 150        | 75          | 200         |
| Active anti-vibration table (NLO)<br>for two-microscope configuration | 250         | 150        | 75          | 400         |
| Periscope for two NLO lasers or NLO laser with OPO                    | 40          | 37         | 70          | 40          |
| Active anti-vibration table (NLO) for Laser Chameleon with OPO        | 180         | 180        | 75          | 410         |
| Scanning Module LSM 880   | 50          | 45         | 22          | 27          |
| Microscope  | 50          | 35         | 50          | 20          |
| Electronic Rack with laser units                                      | 80          | 60         | 65          | 80          |
| Plug-in unit external laser   | 70          | 55         | 25          | 10          |
| Laser module UV   | 80          | 60         | 65          | 40          |
| Airyscan  | 40          | 20         | 24          | 12          |
| Fiber optic cable, UV   | 200         |            |             |             |
| Fiber optic cable, VIS(ible)  | 250         |            |             |             |
| Cables  | 250         |            |             |             |

### Microscopes

| Stands              | Upright: Axio Imager.Z2, Axio Examiner.Z1<br>Inverse: Axio Observer.Z1 with side port or rear port  |
|---------------------|---|
| Z Drive             | Smallest increment Axio Imager.Z2: <25 nm;<br>Axio Observer.Z1: <25 nm;<br>Axio Examiner: <30 nm;<br>fast piezo objective or stage focus available; Definite Focus for Axio Observer.Z1 |
| XY Stage (optional) | Motorized XY scanning stage, for Mark & Find function (xyz) as well as Tile Scan (Mosaic Scan);<br>smallest increment of 1 µm (Axio Observer) or 0.2 µm (Axio Imager)                   |

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| Scanning module       |   |
|-----------------------|---|
| Scanner               | Two independent, galvanometric scanning mirrors with ultrashort line and frame flyback  |
| Scanning Resolution   | $4 \times 1$ to 8192 x 8192 pixels, also for multiple channels, continuously adjustable   |
| Scanning Speed        | 19 x 2 speed levels; up to 13 images/sec. with 512 x 512 pixels (max. 430 images/sec 512 x 16), up to 6875 lines per second   |
| Scanning Zoom         | 0.6× to 40×; digitally adjustable in increments of 0.1 (Axio Examiner: 0.67× to 40×)  |
| Scanning Rotation     | Can be rotated freely (360 degrees), adjustable in increments of one degree, freely adjustable xy offset  |
| Scanning Field        | 20 mm field diagonal (max, 18 mm for Axio Examiner) in the intermediate image plane, with full pupil illumination   |
| Pinholes              | Master pinhole with preset size and position; can be adjusted as desired for multitracking and short wavelengths (such as 405 nm)   |
| Beam Path             | Exchangeable Twin Gate beamsplitter with up to 100 combinations of excitation wavelengths and outstanding laser line suppression;<br>manual interface port for external detection modules (such as BiG.2, Airyscan, third party detectors, internal detection<br>with spectral signal separation and signal recycling loop for compensation of polarization effects |
| Detection options     |   |
| Detectors             | 3 or 34 spectral detection channels, GaAsP and/or PMT (QE 45% typical for GaAsP)  |
|                       | 2 additional GaAsP detection channels (BiG.2)   |
|                       | Airyscan module with spatial detection (32 channels GaAsP)  |
|                       | Up to 12 non-descanned detection channels (PMT and/or GaAsP)  |
|                       | Transmitted light detector (PMT)  |
| Spectral Detection    | 3 or 34 simultaneous, confocal reflected-light channels, GaAsP and / or PMT based   |
|                       | freely adjustable spectral detection area (resolution down to 3 nm)   |
| Data Depth            | 8 bit, 12 bit or 16 bit available; up to 35 channels simultaneously detectable  |
| Real-Time Electronics | Microscope, laser, scanning module and additional accessory control;<br>data acquisition and synchronization management through real-time electronics;<br>oversampling read-out logic   |

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| ZEN Imaging Software             |   |
|----------------------------------|---|
| System Configurations            | Workspace to conveniently configure all of the motorized functions of the scanning module, laser and microscope;<br>save and restore application configurations (Re-Use)  |
| System Self-Test                 | Calibration and testing tool to automatically test and calibrate the system   |
| Recording Modes,<br>Smart Setup  | Spot, Line/Spline, Frame, Tiles, Z Stack, Lambda Stack, Time Series and all combinations (xyz, lambda, t), online calculation and visualization of ratio images, average and summation (by line/image, adjustable), Step Scan (for higher image frame rates); quick set up of imaging conditions using Smart Setup by simply selecting the labelling dye          |
| Crop Function                    | Easily select scanning areas (simultaneously select zoom, offset, rotation)   |
| Real ROI Scan,<br>Spline Scan    | Scans of up to 99 designated ROIs (regions of interest) as desired and pixel-by-pixel laser blanking; scan along a freely defined line  |
| ROI Bleaching                    | Localized bleaching in up to 99 bleach ROIs for applications such as FRAP (fluorescence recovery after photobleaching) or uncaging;<br>use of different speeds for bleaching and imaging, use of different laser lines for different ROIs   |
| Multitracking                    | Rapidly change excitation lines when recording multiple fluorescences for the purpose of minimizing signal crosstalk and increasing dynamic range   |
| Lambda Scan                      | Parallel or sequential acquisition of image stacks with spectral information for every pixel  |
| Linear Unmixing                  | Acquisition of crosstalk-free, multiple fluorescence images using simultaneous excitation; online or offline and automatic or interactive unmixing. Advanced unmixing logic with indication of reliability  |
| Visualization                    | XY, orthogonal (XY, XZ, YZ), Cut (3D section); 2.5D for time series of line scans, projections (maximum intensity); animations;<br>Depth coding (inverse colors), brightness, gamma and contrast settings; color table selection and modification (LUT), character functions  |
| Image Analysis and<br>Operations | Co-localization and histogram analysis with individual parameters, number & brightness analysis, profile measurement along user-defined<br>lines, measurement of lengths, angles, areas, intensities and much more; operations: addition, subtraction, multiplication, division, ratio, shift<br>filters (low-pass, median, high-pass, etc., also user-definable) |
| Image Management                 | Features for managing images and the corresponding imaging parameters; multiprint feature   |

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| Optional Software        |   |
|--------------------------|---|
| 3D VisArt                | Rapid 3D and 4D reconstructions and animations (available modes: shadow projections, transparency projection, surface rendering)                                    |
| Deconvolution            | 3D image restoration based on calculated point-spread functions (modes: nearest neighbor, maximum likelyhood, constrained iterative)                                |
| ROI-HDR                  | Imaging mode: High Dynamic Range, intelligent, local improvement of the dynamic signal range, user-selectable gain and laser power                                  |
| Physiology               | Comprehensive evaluation software for online and offline calibration of ion concentrations  |
| FRET                     | Acquisition of FRET (Förster resonance energy transfer) image data with subsequent evaluation;<br>Acceptor Photobleaching and Sensitized Emission methods supported |
| FRAP efficiency analysis | Acquisition of FRAP (fluorescence recovery after photobleaching) experiments with subsequent evaluation of intensity kinetics                                       |
| Visual Macro Editor      | Creation and editing of macros using representative symbols to program routine workflows  |
| RICS Image Correlation   | Single molecule imaging and analysis using multialkali or GaAsP PMT detectors (publ. v. Gratton)  |
| Experiment Designer      | Defintion of customized imaging configurations and procedures   |
| Macro Environment        | VBA Macro recording and editing   |

| n Brief              | Lasers  |  |  |  |
|----------------------|---|--|--|--|
| he Advantages        | Laser Insert RGB<br>(pigtailed; 458, 488, 514, 543, 561, 594, 633 nm) | Single-mode polarization preserving fiber            |  |  |
|                      | (pigtaileu, 400, 400, 514, 545, 501, 534, 655 1111)                   | Laser beam attenuation for all lasers by VIS-AOTF    |  |  |
| e Applications       |   | Ar laser (458, 488, 514 nm, 25 or 35 mW)             |  |  |
| ne System            |   | HeNe laser (543 nm, 1 mW) DPSS laser (561 nm, 20 mW) |  |  |
| e system             |   |  |  |  |
| chnology and Details |   | HeNe laser (594 nm, 2 mW)                            |  |  |
|                      |   | HeNe laser (633 nm, 5 mW)                            |  |  |
| Service              | Laser Insert V (pigtailed; 405, 440 nm)                               | Single-mode polarization preserving fiber            |  |  |
|                      |   | Diode Laser pulsed/cw (405 nm, 30 mW)                |  |  |
|                      |   | cw mode  | max power ca. 15 mW at fiber out                                   |  |
|                      |   |  | range 0.6 – 15 mW w/o attenuator,<br>attenuation by a factor of 25 |  |
|                      |   | pulsed mode  | repetition rate 20 – 50 – 80 MHz                                   |  |
|                      |   |  |  |  |
|                      |   |  | peak power: 50 – 300 mW  |  |
|                      |   |  | pulse width: 50 – 90 ps  |  |
|                      |   | jitter < 20 ps                                       |  |  |
|                      |   |  |  |  |
|                      |   | Diode Laser pulsed/cw (440 nm, 25 mW)                |  |  |
|                      |   | cw mode  | max power ca. 15 mW at fiber out                                   |  |
|                      |   |  | range 0.6 – 15 mW w/o attenuator,<br>attenuation by a factor of 25 |  |
|                      |   | pulsed mode  | repetition rate 20 – 50 – 80 MHz                                   |  |
|                      |   |  |  |  |
|                      |   |  | peak power: 50 – 300 mW  |  |
|                      |   |  | pulse width: 50 – 90 ps  |  |
|                      |   |  | jitter < 20 ps   |  |

| Advantages            | Laser Module UV (355 nm)   | Single-mode polarization preserving fiber                   |                                  |  |
|-----------------------|--|---|----------------------------------|--|
| e Auvantayes          |  | Laser beam attenuation by AOM<br>DPSS laser (355 nm, 60 mW) |                                  |  |
| ne Applications       |  |   |                                  |  |
| ne System             | Power Requirements   |   |                                  |  |
| echnology and Details | LSM 880 has a main power supply cord and plug, either CEE red (3/N/PE 400/230V/16A), or NEMA L 14-30P (2/N/Ground 120/240V/30A), and the matching mains socket outlet.<br>The mains socket outlet must be equipped with a fuse having minimum tripping characteristic C according to IEC/EN 60898. |   |                                  |  |
| Service               | Line Voltage   | 3/N/PE 400/230 V AC (±10 %)                                 | 2/N/Ground 240/120 V AC          |  |
|                       | Line Frequency   | 5060 Hz   | 5060 Hz                          |  |
|                       | ZEISS LSM incl. VIS Laser  |   | 2 4 4 25 4                       |  |
|                       | 7ELSE LEM incl. VIS Locar  |   |                                  |  |
|                       | Max. Current   | 3 phases at 16 A  | 2 phases at 25 A                 |  |
|                       | Power  | Phase 1 = 600 VA max.                                       | Phase 1 = $800 \text{ VA max}$ . |  |
|                       |  | Phase $2 = 500$ VA max.                                     | Phase $2 = 1600VA$ max.          |  |
|                       |  | Phase 3 = 1500 VA max.                                      |                                  |  |
|                       | Power Consumption  | 2100 VA max.  | 2100 VA max.                     |  |
|                       | Multiphoton Laser  |   |                                  |  |
|                       | Power Consumption  |   |                                  |  |
|                       | Ti:Sa laser 1  | 800 VA max.   | 800 VA max.                      |  |
|                       | Ti:Sa laser 2  | 800 VA max.   | 800 VA max.                      |  |
|                       | OPO  | 600 VA max.   | 600 VA max.                      |  |
|                       | Heat emission<br>without Ti:Sa or OPO laser)   | 2000 W max.   | 2000 W max.                      |  |
|                       | EMC test   |   |                                  |  |
|                       | according to DIN EN 61326-1 (05/2010)  |   |                                  |  |
|                       | 1. Noise emission according to CISPR 11 / DIN EN 55011 (04/2011)   |   |                                  |  |

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| For operation the system has to be placed in a closed room       |  |
|--|--|
| 1. Operation, specified performance                              | T = 22 °C $\pm$ 3 °C without interruption (24 h a day independently whether system is operated or switched-off)<br>It has to be ensured that the air-flow of the air-conditioning is not directed at the system. |
| 2. Operation, reduced performance                                | T = 15 °C to 35 °C, any conditions different from item 1. and 5.   |
| 3. Storage, less than 16 h                                       | T = -20 °C to 55 °C  |
| 4. Storage, less than 6 h  | T = -20 °C to 55 °C  |
| 5. Temperature gradient  | ±0.5 °C/h  |
| 6. Warm up time  | 1 h, for high-precision and/or long-term measurements $\geq$ 3 h   |
| 7. Relative humidity   | <65 % at 30 °C   |
| 8. Operation altitude  | max. 2000 m  |
| 9. Loss of heat (without Ti:Sa or OPO laser)                     | 2 kW   |
| 10. Vibrations under operation conditions<br>(with system table) | 5 μm pp at 5 Hz<br>10 μm pp at 10 Hz<br>10 μm pp at 20 Hz  |
| 11. Shipping shock (LSM 880 box)                                 | 10 g   |











LSM 880 meets the requirements according to IEC 60825-1:2007

### Count on Service in the True Sense of the Word

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Because the ZEISS microscope system is one of your most important tools, we make sure it is always ready to perform. What's more, we'll see to it that you are employing all the options that get the best from your microscope. You can choose from a range of service products, each delivered by highly qualified ZEISS specialists who will support you long beyond the purchase of your system. Our aim is to enable you to experience those special moments that inspire your work.

#### Repair. Maintain. Optimize.

Attain maximum uptime with your microscope. A ZEISS Protect Service Agreement lets you budget for operating costs, all the while reducing costly downtime and achieving the best results through the improved performance of your system. Choose from service agreements designed to give you a range of options and control levels. We'll work with you to select the service program that addresses your system needs and usage requirements, in line with your organization's standard practices.

Our service on-demand also brings you distinct advantages. ZEISS service staff will analyze issues at hand and resolve them – whether using remote maintenance software or working on site.

#### Enhance Your Microscope System.

Your ZEISS microscope system is designed for a variety of updates: open interfaces allow you to maintain a high technological level at all times. As a result you'll work more efficiently now, while extending the productive lifetime of your microscope as new update possibilities come on stream.







Profit from the optimized performance of your microscope system with services from ZEISS – now and for years to come.

>> www.zeiss.com/microservice

# The moment your data change scientific minds. This is the moment we work for.

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Carl Zeiss Microscopy GmbH 07745 Jena, Germany microscopy@zeiss.com www.zeiss.com/lsm880



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