abbelight instrument CAPTURE THE EVANESCENCE Your upgradable nanoscope for single-molecule imaging





abbelight instrument

abbelight offers two modules adaptable to most inverted microscopes: SAFe 180 and SAFe 360.



Illumination on large field of view 2D STORM nanoscopy Data analysis Image: Description of the second o

With these instruments, researchers can perform any type of single-molecule localization imaging.

SAFe 360



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Simultaneous multicolor















From microscopy to nanoscopy

Revealing structures and dynamics at the nanoscale

Standard **fluorescence microscopy** techniques (widefield, confocal,...) operate in the resolution range of 200–300 nm laterally and 500–800 nm axially. However, biological structures and processes that occur at a lower scale require superior resolution.

Among recent techniques that break the diffraction limit, *i.e.* super-resolution techniques, **nanoscopy** retrieves structural or dynamic quantitative information with the highest resolution achievable in light microscopy.



Stefan Hell, Eric Betzig, and William Moerner were awarded the chemistry Nobel Prize in 2014 for their work on nanoscopy techniques.



Single-molecule Localization Microscopy (SMLM) is the nanoscopy technique that retrieves structural or dynamic quantitative information with the highest precision achievable.

SMLM principle

SMLM relies on the ability to randomly activate only a subset of fluorescent molecules in order to distinguish them spatially.

By repeating the process in consecutive image acquisitions, accumulated raw data are processed to detect single molecules with a nanometric precision (down to 10 nm).

Data guantification and analysis are then performed to resolve either structures or dynamics at the nanoscale level.



microscopy

The uniqueness of SMLM is that it gives rise not only to highly resolved images, but also to the 3D coordinates of single molecules, opening up new avenues for spatial and temporal quantitative analysis.



Localizing molecules in 2D

In order to reconstruct a nanoscopy image, each molecule is detected and localized by specialized algorithms.



To determine the x and y positions of each molecule, a commonly used localization algorithm is Gaussian fitting.



Because images are now obtained at the nanoscale level, new challenges arise. Effects that were negligible at the microscopy level now need to be taken into account.



Resolution = 2,35 × (Localization precision) \otimes (labeling density, drift...)



SMLM approaches... from structure to dynamics

Current SMLM approaches only differ in how the fluorophores activation-inactivation is induced. Among them, STORM, PALM and PAINT resolve spatial structures with nanometric precision, while SPT and smFRET reveal temporal dynamic processes in living cells.

Structures

- STORM (STochastic Optical Reconstruction Microscopy)
 - Standard organic fluorescent dyes (cyanines, rhodamines, oxazines...)
 - Specific imaging buffer (containing a reducer, which induces the transition to a dark state, and an oxygen scavenging system to stabilize this state before returning to the ground state)
- PALM (Photoactivated Localization Microscopy)
 - Photo-activatable or -convertible fluorescent proteins (mEos3,2, Dendra2, PA-mCherry, mMaple,...);
 - No specific buffer, live-cell compatible
- PAINT (Point Accumulation for Imaging in Nanoscale Topography)
 - Specific fluorophores that have the ability to emit fluorescence only upon binding to their biological target (ex: Nile Red, which fluoresces only when interacting with membranes)
 - No specific buffer, live-cell compatible



		Contraction of the
STORM	PALM	PAINT
NHS-AF-647	RNAP-mEos3.2 *	Nile Red-membrane

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Dynamics

 sptSMLM combines Single Particle Tracking with SMLM (PALM or STORM) to obtain spatially and temporally highly resolved diffusion maps of single molecules

SPT



Raw data



- Planck Institute, Marburg (Single molecule Fluorescence
- SmFRET (Single molecule Fluorescence Resonance Energy Transfer) provides distance measurements in single-molecule reaction trajectories
 High FRET dyes are close to each other



DAISY technology

DAISY is the combination of two complementary strategies developed by abbelight and used to extract the "Z" position of a particle. (Cabriel et al. *BioRxiv 2018*)

The first approach is called **Magnified Astigmatism** and is inspired by the astigmatic PSF shaping published by Huang et al. (*Science 2008*) and commonly used in commercial systems.

The second approach exploits the near field information encoded in every single emitter, developed and published under the name **DONALD** by Bourg et al. (*Nature Photonics 2015*)

	abbeligh	t instrument		Nanoscopy market
3D method	Magnified astigmatism (in dual-view system)	DONALD	DAISY	Standard astigmatism
Imaging depth	5-10 µm	0,5 µm	5-10 µm	5-10 μm
Capture range	1,2 µm	0,5 µm	1,2 µm	800 nm
Lateral loc. Precision*	10 nm	10 nm *	10 nm	10 nm
Lateral resolution**	23 nm	23 nm	23 nm	23 nm
Axial loc. Precision*	13 nm	13 nm *	13 nm	22 nm
Axial resolution**	> 30 nm + focus & drift dependence	30 nm	30 nm	> 50 nm +focus & drift dependence
Axial drift	Degrade axial resolution above	Not sensitive	Not sensitive	Degrade axial resolution above

* Mean value, for dSTORM imaging using AF647 and abbelight buffer

** Mean value, resolution = 2,35 × (Localization precision) ⊗ (labeling density, drift...)



Magnified Astigmatism

ASTIGMATIC PSF SHAPING is a very efficient way to extract the relative position of a single particle regarding the focus plane of the objective. Using an astigmatic lens, a controlled aberration can be induced, measured, and related to the distance between the objective's focal plane and the emitting particle.

The stronger the aberration is, the better the axial precision is. However, the lateral resolution is also degraded. Therefore, for conventional setups using a single camera, a compromise has to be found between astigmatism strength and X,Y localization precision.

Using a dual camera system, the astigmatic deformation can be enhanced while preserving the best lateral resolution. This is what we call Magnified Astigmatism.

Key features

- + Capture range of 1200 nm
- + Enhanced astigmatic lens for better Z precision
- + No loss of lateral resolution
- + Two controls for false positive detections
- Relative to the focal plane: sensitive to axial drift
- Amplified chromatic aberrations

Camera 1



Camera 2







DONALD

Any single emitter is a dipole, emitting two components of fluorescence:

The far-field emission is a propagative wave, always collected in the low angles of the objective (UAF) and commonly used for any fluorescence microscopy technique.

Although it is often forgotten, dyes also have a near-field emission, which can also be collected in the objective - if the dye is close enough to the coverslip - but in the high angles of the objective (SAF).

Since UAF is constant and SAF decays exponentially, a simple ratio of intensities, for each dye, determines its absolute distance to the coverslip.

Besides its simplicity, the strength of this photophysical measurement is its insensitivity to axial drift or aberration.

Key features

- + Absolute measurement: insensitive to drift
- + No loss of lateral resolution
- + No chromatic aberration
- + Compatible with PSF shaping methods
- Loss of axial precision above 300 nm
- _ Capture range limited to 600 nm above coverslip

Far field emission



DAISY technology

DAISY



Absolute VS Relative measurement

DRIFT-FREE

Since the theoretical localization precision is almost the same between DONALD and Magnified Astigmatism, the final resolution of the image will be different because of the axial drift. Even with the best focus control system, small oscillations and large drift of the focal plane position can occur, directly impacting the resolution of the final image.

STATISTICAL ANALYSIS

An absolute axial measurement enables straightforward statistical 3D multicolor analysis, since the reference is always the coverslip surface. It is now possible to compare thousands of acquisitions, on different samples, for different proteins, and easily compile the data*.

*Bouissou et al. ACSnano 2017



SAFe 360 optical path

The SAFe 360 module has been designed to integrate and use both Magnified Astigmatism and **DONALD** at the same time, and therefore DAISY.

The optical cubes and specific lenses can be easily replaced to tune the system depending on the user's needs.

- 1. Fluorescence from the microscope
- Beam splitter 50/50 2.
- Camera 1 for 2D and SAF detection 3.
- SAF physical filtering for DONALD 4.
- 5. Strong cylindrical lens to induce magnified astigmatism
- 6. Camera 2 for 3D astigmatic detection





SAFe Light

abbelight **SAFe Light** technology offers the **largest field of view** in nanoscopy and requires **lower laser power**.

- ✤ 200x200 µm² field of view with 300 mW laser power
- More intensity on the sample with lower laser power
- Illumination adaptable to the sample
- TIRF, HiLo or EPI illumination modes
- 16-times more quantitative data
- Homogenous illumination, no interference patterns in the image

-	

Scanning for homogenous illumination faster than the camera acquisition time (200 fps for the largest field of view)

Widefield microscopy

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Widefield microscopy



Nanoscopy

TIRF microscopy



Standard field of view in nanoscopy market: 50x50 µm²

COS7 cells – mitochondria AF647 Sample provided by N. Bourg abbelight

20 µm

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Illumination adapted to the sample







	EPI	HiLo	TIRF
Name	Epifluorescence	Highly inclined and laminated optical sheet	Total internal reflection fluorescence
Type of illumination	In-depth illumination, higher background	Limited background, not restricted to coverslip	Illumination close to the coverslip, removal of in-depth background
Examples of biological structures	Structures far from the coverslip: nucleus, thick cells, tissues	Slightly in-depth samples	Structures close to the coverslip: membranes, cytoskeleton, in vitro surfaces



Simultaneous multicolor

Multicolor imaging is a powerful way to assess colocalization between different biological structures.

- Method 1: acquire different colors sequentially
- Method 2: use dichroic cubes to allow simultaneous multicolor nanoscopy.

Both these methods are possible with SAFe 360.

However, they both require excitation with several lasers and compatibility of imaging buffers, and they can lead to chromatic aberrations.



abbelight implemented a technology to perform simultaneous multicolor single-molecule imaging with only one excitation laser: spectral demixing (Winterflood et al. Bophys 2015).

This method is **compatible** with a vast array of cell lines and standard commercial fluorophores.



NEO software

Abbelight's NEO SAFe software provides **a user-friendly all-in-one workspace** for acquisition, processing, and analysis of nanoscopy data.

Feature	NEO SAFe software
Control of instrument	Laser power and illumination angle (EPI, HiLo, TIRF)
	Cameras
Control of acquisition parameters	Region of interest size
	Exposure time (down to 10 ms per frame)
	Frame number
Live reconstruction of nanoscopy data	Choice of localization parameters: Localization algorithms (center of mass, Gaussian fitting, phasor) Intensity threshold Background substraction method
	Live visualization and reporting
Live drift correction	Cross-correlation
Decision-making tools to guide acquisition	Real-time SQUIRREL algorithm, Culley et al. 2017



Nanoscopy data, unlike standard microscopy images, are **coordinate-based** rather than pixelbased, opening up new avenues for in-depth data analysis.

NEO offers a variety of tools for nanoscopy data visualization and analysis.

Feature	NEO SAFe software
Visualization	3D visualization
	Region of interest crops
	Mutlicolor visualization
	Possibility to export images in TIFF format compatible with commonly used softwares
Descriptive spatial statistics	Localization distribution
	Measuring tools
Clustering analysis	K-Ripley function, DBSCAN, Voronoi tessellation
	Centroid, density and volume measurements
	Possibility to export quantifications in format compatible with commonly used softwares





© 2019 abbelight contact@abbelight.com | www.abbelight.com abbelight, 6 rue Jean Calvin, 75005 Paris, France