Confocal Microscopy

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Acousto-optic products

AA OPTO-ELECTRONIC QUANTA TECH

Introduction

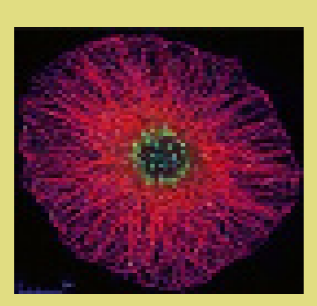
Confocal microscopy is an imaging technique used to increase micrograph contrast and/or to reconstruct three-dimensional images by using a spatial pinhole to eliminate out-of-focus light or flare in specimens that are thicker than the focal plane. This technique has been gaining popularity in the scientific and industrial communities. Typical applications include life sciences and semiconductor inspection.

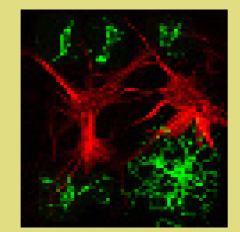
Basic concept

The principle of confocal imaging was patented by Marvin Minsky in 1961. In a conventional (i.e., widefield) fluorescence microscope, the entire specimen is flooded in light from a light source. Due to the conservation of light intensity transportation, all parts of specimen throughout the optical path will be excited and the fluorescence detected by a photodetector or a camera. In contrast, a confocal microscope uses point illumination and a pinhole in an optically conjugate plane in front of the detector to eliminate out-of-focus information. Only the light within the focal plane can be detected, so the image quality is much better than that of wide-field images. As only one point is illuminated at a time in confocal microscopy, 2D or 3D imaging requires scanning over a regular raster (i.e. a rectangular pattern of parallel scanning lines) in the specimen. The thickness of the focal plane is defined mostly by the square of the numerical aperture of the objective lens, and also by the optical properties of the specimen and the ambient index of refraction.

Different principles

Three types of confocal microscopes are commercially available: Confocal laser scanning microscopes, spinning-disk (Nipkow disk) confocal microscopes and Programmable Array Microscopes (PAM). Generally speaking, confocal laser scanning microscopy yields better image quality but the imaging frame rate is very slow (less than 3 frames/second); spinning-disk confocal microscopes can achieve video rate imaging-desired for dynamic observations.





Confocal Laser scanning Microscopy

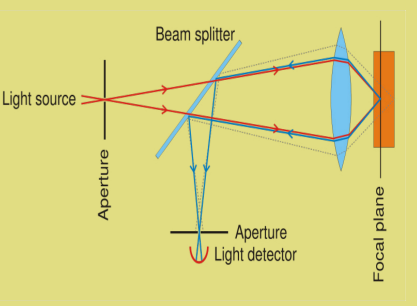
Confocal laser scanning microscopy (CLSM or LSCM) is a valuable tool for obtaining high resolution images and 3-D reconstructions. The key feature of confocal microscopy is its ability to produce blur-free images of thick specimens at various depths. Images are taken point-by-point and reconstructed with a computer, rather than projected through an eyepiece. The principle for this special kind of microscopy was developed by Marvin Minsky in 1953, but it took another thirty years and the development of lasers for confocal microscopy to become a standard technique toward the end of the 1980s.

Image formation

In a laser scanning confocal microscope a laser of detected fluorescent light. The beam is scanned across the sample in the horizontal plane using one beam passes a light source aperture and then is focused by an objective lens into a small (ideally difor more (servo-controlled) oscillating mirrors. This fraction-limited) focal volume within a fluorescent scanning method usually has a low reaction latency specimen. A mixture of emitted fluorescent light as and the scan speed can be varied. Slower scans prowell as reflected laser light from the illuminated spot vide a better signal to noise ratio resulting in better is then recollected by the objective lens. A beam contrast and higher resolution. Information can be collected from different focal planes by raising or splitter separates the light mixture by allowing only the laser light to pass through and reflecting the lowering the microscope stage. The computer can generate a three-dimensional picture of a specimen fluorescent light into the detection apparatus. After passing a pinhole the fluorescent light is detected by assembling a stack of these two-dimensional imaby a photo-detection device (photomultiplier tube ges from successive focal planes. (PMT) or avalanche photodiode) transforming the light signal into an electrical one which is recorded In addition, confocal microscopy provides a signifiby a computer. cant improvement in lateral resolution and the capa-

city for direct, non-invasive serial optical sectioning The detector aperture obstructs the light that is not of intact, thick living specimens with an absolute coming from the focal point, as shown by the dotminimum of sample preparation. As laser scanning ted grey line in the image. The out-of-focus points confocal microscopy depends on fluorescence, a are thus suppressed: most of their returning light is sample usually needs to be treated with fluorescent blocked by the pinhole. This results in sharper imadyes to make things visible. However, the actual dye ges compared to conventional fluoresence microsconcentration can be very low so that the disturbancopy techniques and permits one to obtain images ce of biological systems is kept to a minimum. Some of various z axis planes (z-stacks) of the sample. instruments are capable of tracking single fluorescent molecules. Additionally transgenic techniques The detected light originating from an illuminated vocan create organisms which produce their own lume element within the specimen represents one fluorescent chimeric molecules. (such as a fusion of pixel in the resulting image. As the laser scans over GFP, Green fluorescent protein with the protein of the plane of interest a whole image is obtained pixel interest).

by pixel and line by line, while the brightness of a resulting image pixel corresponds to the relative intensity



Resolution enhancement by the confocal principle

Laser scanning confocal microscopy (LSCM) is a Unfortunately, the probability decrease in creation of scanning imaging technique in which the resolution detectable photons has a bad effect on the signal to obtained is best explained by comparing it with ano- noise ratio. This can be compensated by using more ther scanning technique like Scanning electron mi- sensitive photo-detectors or by increasing the intencroscope (SEM). Not to be confused with phonogra- sity of the illuminating laser point source. Increasing ph-like imaging—AFM or STM, for example, where the intensity of illumination latter risks excessive the image is obtained by scanning with an atomic tip over a conducting surface.

In LSCM a fluorescent specimen is illuminated by a point laser source, and each volume element is associated with a discrete fluorescence intensity. Here, the size of the scanning volume is determined by the spot size (close to diffraction limit) of the optical system. This is due to the fact that the image of the scanning laser is not an infinitely small point but a three-dimensional diffraction pattern.

The size of this diffraction pattern and the focal volume it defines is controlled by the numerical aperture of the system's objective lens and the wavelength of the laser used. This can be seen as the classical resolution limit of conventional optical microscopes using wide-field illumination. However, with confocal microscopy it is even possible to overcome this resolution limit of wide-field illuminating techniques as only light generated in a small volume element is detected at a time. Here it is very important to note that the effective volume of light generation is usually smaller than the volume of illumination; that is, the diffraction pattern of detectable light creation is sharper and smaller than the diffraction pattern of illumination.

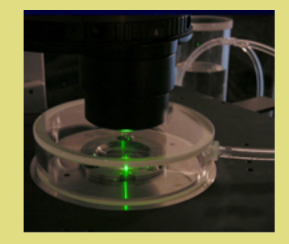
In other words, the resolution limit in confocal microscopy depends not only on the probability of illumination but also on the probability of creating enough detectable photons, so that the actual addressable volume being associated with a generated light intensity is smaller than the illuminated volume. Depending on the fluorescence properties of the used dyes, there is a more or less subtle improvement in lateral resolution compared to conventional microscopes. However, by using light creation processes with much lower probabilities of occurrence such as second harmonic generation (SHG), the volume of addressing is reduced to a small region of highest laser illumination intensity resulting in a significant improvement in lateral resolution.

bleaching or other damage to the specimen of interest, especially for experiments in which comparison of fluorescence brightness is required.

Uses

Confocal microscopy is clinically used in the evaluation of various eve diseases. It is particularly useful for imaging, qualitative analysis and quantitafication of endothelial cells of the cornea. It is used for localising and identifying presence of filamentary fungal elements in the corneal stroma in cases of keratomycosis, enabling rapid diagnosis and thereby early institution of definitive therapy.

Confocal microscopy is also used as the data retrieval mechanism in some 3D optical data storage systems and has helped determine the age of the Magdalen papyrus



Polychromatic Modulation Systems

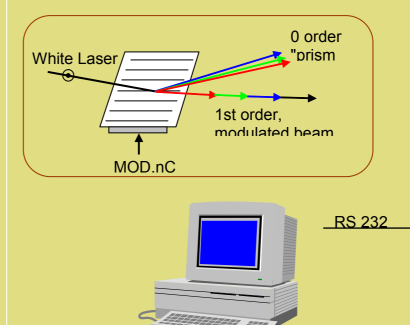
The AOTF.nC is a special acousto-optic tunable filter which uses the anisotropic interaction inside a tellurium dioxide crystal to control independently or simultaneously different lines from an incoming laser light (White laser, Ar+, Kr+, HeNe, DPSS, Dye...). Up to 12 distinct lines can be mixed and separately modulated in order to generate different colorimetric patterns.

The specific crystal cut of the AOTF.nC produces good diffraction efficiency (> 90%), narrow resolution (1-2 nm), a low cross-talk between lines, and high extinction ratio.

The large separation angle between 0 and 1st orders, as well as the excellent output chromatic colinearity (<0.2 mrd over 450-700 nm) make this AOTF a powerful tool for free space or fiber pigtailed applications.

Its associated thermal stabilisation maintains stable computer control). diffraction efficiency and reduces dramatically beam All parameters are stored in an EEPROM and are drift with single mode fiber pigtailing. This is a major automatically loaded after each switch on. advantage for high sensitivity applications.

The associated driver MOD.nC, based on PLLs (Phaby a blanking signal which produces smooth effects se Locked Loop), has been specially designed in orwithout modifying the colorimetric balance. der to exploit the best of the AOTF.nC features. The combination of the modulation input and blan-Its compact design with single power supply, low RF king signals provides the best extinction ratio perforemissions and ease of use will satisfy the most demance (> 140 dB). manding of applications, where accuracy and flexibi-



Computer

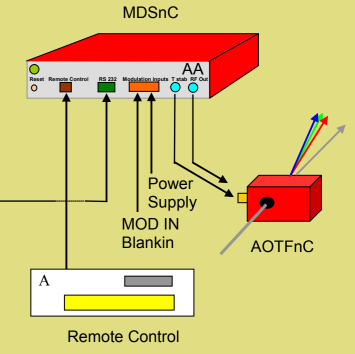


lity are key requirements.

Thanks to its complete numerical design and integrated microcontroller setting up is fast, simple and repeatable.

Access to and adjustments of functions is simple with either a bright LCD display (with remote control adjustment) or through a RS232 serial link (with

Each line is externally controlled by a distinct modulation input signal which can be TTL or analog. Additionally, all lines can be simultaneously controlled



Polychromatic Modulators

AOTFnC

Number of channels / Lines Acoustic velocity (nom) Optical wavelength range Transmission AO interaction type Selected order Input Light polarization Output Light polarization Drive frequency range Active aperture Spectral resolution (FWHM)

Separation angle (orders 0-1) Chromatic colinearity (order 1) Temperature stabilization **AO Efficiency Rise time** Max accepted RF power Electrical impedance VSWR Size **Operating temperature**

UV 4 675 m/s 350-430 nm > 80 % -nom 90% Birefringent +1 Linear parallel Linear orthogonal 110-180 MHz 2 x 2 mm² > 4.2 degrees < 0.2 mrd @351+363 nm < 0.2 mrd T or TN >=90%

980 ns / mm < 1 W all lines 50 Ohms < 2/1

Linear parallel 80-153 MHz 3 x 3 mm² nom 1-2 nm > 4.6 degrees T or TN >= 90 % /line 1010 ns / mm < 1 W all lines 50 ohms

VIS

650 m/s

> 95 %

-1

450-700 nm

Birefringent

8

< 2/1 70 x 36.6 x 35.8 mm3 10 to 40 °C

VIS 8 660 m/s 400-650 nm > 90 % Birefringent -1 Linear orthogonal Linear orthogonal Linear parallel 74-158 MHz 3 x 3 mm² nom 1-4 nm > 4 degrees < 0.3 mrd

T or TN >= 90 % /line 1000 ns /mm nom 1 W all lines 50 ohms < 2/1

Polychromatic Drivers / Digital versions

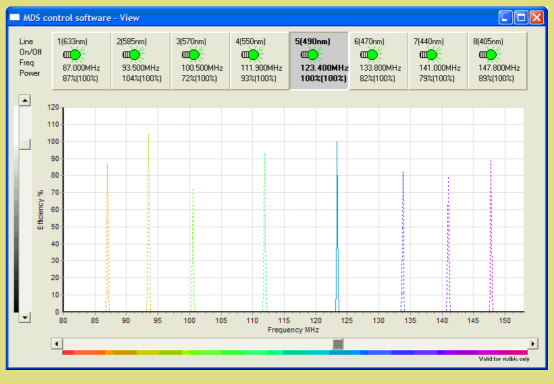
These drivers based on Direct Digital Synthesizers (DDS), produce multiple fixed stable and accurate RF frequency signals for polychromatic modulators. Their brand new design with "on the edge" technology offers unique performance in term of accuracy, speed and stability (single/multi-line), thanks to their internal temperature correction and high linearity design.

The built in amplifier delivers the necessary RF power to drive the acousto-optic device, with reduced power consumption (AA "COLD DESIGN").

The RF output power per channel can be individually modulated (MOD IN signals) or simultaneously modulated (BLANKING signal). AA focussed on a ultra low crosstalk version with superior fast and fall time.

The adjustments of the driver (Frequency & Power) can be done with a remote control, USB or through RS 232 communication to allow user flexibility in power control or frequency scanning.

USB Software





MDS control software								
Line Frequency Power	-6.7 dBm			Send Send and Store Query all			C RS232	
On/Off Lines status Select	Frequency		Powe			Status	On/Off	IMode © Internal
 Line 1 Line 2 Line 3 	141.500 133.500 128.699	MHz MHz MHz	19.0	dBm dBm dBm		ON ON OFF	<u>।</u>	C External VMode C 5V
C Line 4 C Line 5 C Line 6	124.500 117.300 105.099	MHz MHz MHz	19.0	dBm dBm dBm		OFF OFF OFF		© 10V
C Line 7 C Line 8	89.199 84.699	MHz MHz		dBm dBm	-	OFF OFF		
						A		



Number of channels Frequency range Frequency stability Frequency accuracy Frequency step Frequency control Power Supply Laboratory version Rise Time / Fall time (10-90 %) Modulation Input Control Blanking Input Control Extinction ratio @ 125 MHz

Output RF power Output Impedance V.S.W.R. Input / Output connectors Size Laboratory version Weight Laboratory version Heat exchange Laboratory version Operating temperature Maximum case temperature Option Up to 8 Octave or above in 20-180MHz - will be adapted to AO +/-2 ppm/°C Nom 1 KHz Nom 1 KHz Remote Control or USB, Option : RS232 OEM version : 24 VDC - nom 0.85A 110/230 VAC - 50-60 Hz < 50ns Analog 0-5 V or 0-10 V / 10 KOhms Analog 0-5 V or 0-10 V / 10 KOhms MOD IN > 80 dB typ 90dB typ 80dB BLK > 70 dBMOD IN + BLK > 90 dB typ 100 dB 22 dBm per channel 50 Ohms Nom < 1.5/1DB25 / SMA (DB9 for RS232) OEM version : 207 x 127 x 20.2 mm3 Rack 19", 1U OEM version : nom 1 kg nom 4 kg **OEM** version : Conduction stand alone 10 to 40 °C OEM version : 50 °C Cover with Fan