DP80 specifications

Item		Specifications Color / Monochrome 2CCD camera	
Camera type			
		Pixel shifting (only for color CCD)	
		Cooling system: Peltier device (max. Ta -10 degreeC)	
Image sensor		[Color]	
	Size	2/3 inch 1.45 mega pixels color CCD, RGB colors on chip filter (Bayer)	
		[Monochrome]	
		2/3 inch 1.45 mega pixels monochrome CCD	
	Scanning mode	Progressive scan	
Camera mount		B4 mount (2/3 inch Bayonet mount)	
Exposure control	Mode	Auto, SFL-Auto, Manual	
	Adjustment	±2.0 EV step: 1/3 EV	
	Exposure time	23 µs to 60 s	
Metering modes		Full image, 30%, 1%, 0.1%	
Binning		2x2, 4x4	
Live frame rate		1360 x 1024 (1x1): 15 fps, 680 x 512 (1x1) : 15 fps	
		680 x 510 (2x2): 29 fps, 340 x 250 (4x4) : 57 fps	
	Image Resolution	[Centering OFF]	[Centering ON]**
Color		4080 x 3072 (Pixel shift)	3648 x 2736 (Pixel shift)
		2040 x 1536 (Pixel shift)	1824 x 1368 (Pixel shift)
		1360 x 1024 (1x1), 680 x 512 (1x1)	1216 x 912 (1x1), 608 x 456 (1x1)
		680 x 510 (2x2), 340 x 250 (4x4)	608 x 456 (2x2), 304 x 228 (4x4)
	ISO Sensitivity	ISO200/400/800/1600 equivalent	
	A/D	14 bit *Number of effective bit: 12 bits@16 bit mode image	
	Image acquisition time *	4080 x 3072: approx. 3.3 s, 1360 x 1024: approx. 0.3 s	
	Color space	sRGB, AdobeRGB (only for color CCD)	
		[Centering OFF]	[Centering ON]
	Image Resolution	1360 x 1024 (1x1), 680 x 512 (1x1)	1216 x 912 (1x1), 608 x 456 (1x1)
Monochrome	-	680 x 510 (2x2), 340 x 250 (4x4)	608 x 456 (2x2), 304 x 228 (4x4)
	Gain	x0.5/x1/x2/x4/x8/x16	
	A/D	14 bit *Number of effective bit: 14 bits@16 bit mode image	
	Full well capacity	17000e- (Gain 0.5x)	
	Readout noise	7e-	
	Dynamic range	2300:1 (Gain 0.5x)	
	Quantum efficiency	55% (500 nm)	
	Dark current	0.4e-/pixel/s (Ta=25degere C)	
	Image acquisition speed *	1360 x 1024: approx. 7.7fps, 680 x 510 (2 x 2): approx. 14.3fps,	
		340 x 250 (4 x 4): approx. 20fps	
Image file format		File formats supported by cellSens software	
Dimensions, weight	Camera head	133 mm(W) x 130 mm(D) x 139 mm(H) /approx. 2.5 kg	
	Interface cable	Approx. 2.8 m/approx. 0.23 kg	
	External trigger cable	Approx. 0.2 m/approx. 40 g	
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DP80 system	requirement
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4 GB or more

250 W or more

CPU

RAM

HDD

Graphics

OS

Extension slot

Power supply

PC/AT compatible

(8 GB recommended for high speed image acquisition)

1280x1024 (min. 1024 x768) monitor resolution with

32-bit-video card with separate graphics memory (no integrated graphics processor with shared memory)

Windows 7 Professional/Ultimate with SP1 (64 bit)

Intel Core i5/ Intel Core i7 / Intel Xeon

Free space of 1 GB or larger

PCI-Express x1 Rev. 1.0a or later

Compatible with half size or LowProfile PCle board (106.7 mm x 174.6 mm)

Unoccupied FDD power cable, HDD

power cable (4-pin size), or Serial ATA power cable must be available * Please use computers that conform to the regulations for your region.

(at the time of installation)

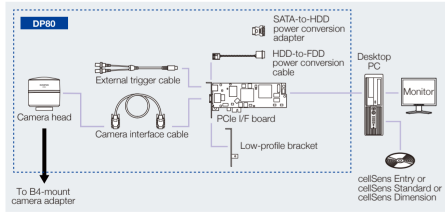


Your Vision, Our Future

A cutting-edge digital microscopy camera equipped with dual CCD sensors, providing both high sensitivity monochrome and high-quality single-shot color images.

* Image acquisition time and speed may be reduced if exposure time increases or several tasks are active in the background.
** "Centering" is a camera function which aligns the positions of the color and monochrome CCD sensors.

DP80 System Diagram



ø133 OLYMPUS Weight: approx 2.5 kg

(Unit: mm)

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DP80 Camera Head Dimension

- All company and product marines are registed of their respective owners.
 Images on the PC monitors are simulated.
- Specifications and appearances are subject to change without any notice or obligation on the part of the manufacturer.



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DP80



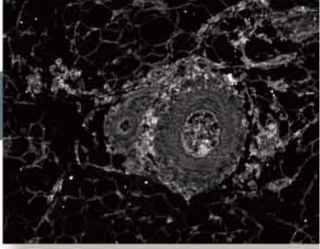




A single camera featuring color / monochrome dual CCD sensors.









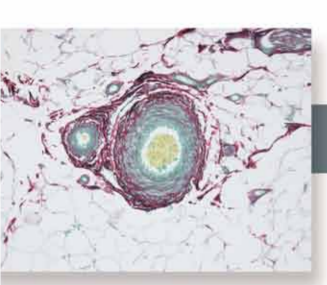
A simple switching operation allows the use of either a color or monochrome CCD

OLYMPUS **DP80**

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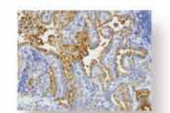


sensor on the single camera unit.



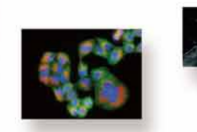


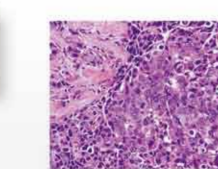
Superior bright-field images at high resolution





Clear fluorescence images with excellent sensitivity





Consider how convenient and easy it would be if both high resolution bright-field and high-sensitivity fluorescence images could be observed and acquired using a single microscope camera. The DP80 digital camera fulfills this simple yet unfulfilled requirement for two cameras in one.

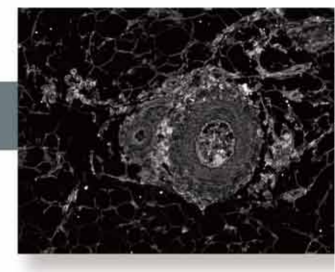
Since it is possible to rapidly switch between the monochrome CCD sensor and color CCD sensor, it is possible to easily acquire high-quality bright-field and high-sensitivity fluorescence images of the same field. Without the need to switch camera ports with a prism, the time required to align the camera sensors is eliminated.

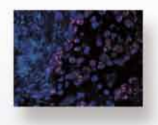
The DP80 assists you in your research activities from observation to documentation in a smooth and easy manner.

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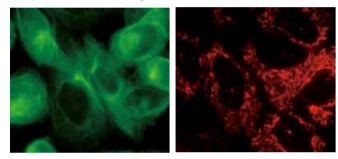


High quality images adapted to observation and imaging methods can be readily obtained.

A monochrome camera that detects and captures dim fluorescence images

Clearly observe weak fluorescence signals with the **DP80's high sensitivity**

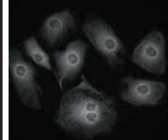
We significantly improved the DP80's fluorescence imaging performance by incorporating a separate high dynamic range monochrome CCD sensor within the body of the camera. Combined with thermo-electric cooling and high resolution capture, the DP80 meets the demands for low-light fluorescence imaging. With a high quantum efficiency along a wide spectrum, the DP80 provides exceptional fluorescence singal detection.



High-sensitivity fluorescence imaging available up to Cv7 wavelengths

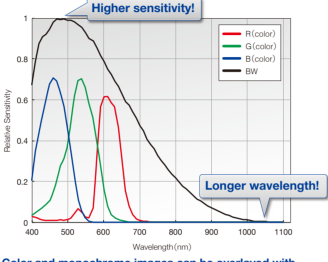
The DP80 responds to a wide range of wavelengths from visible to near infrared. Now sensitive to fluorescence signals with longer wavelengths, the DP80 captures near-IR wavelengths from samples stained with Cy7 (767 nm).





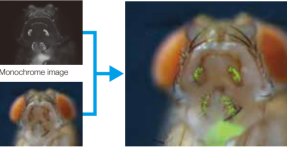
Color image (exposure time: 60 s)





Color and monochrome images can be overlayed with pixel precision

Since the camera is equipped with a centering function, which minimizes positional differences between color images and monochrome images, a combined image can be produced by precisely superimposing the monochrome images and color images that are acquired for a given observation method. Since positions accurately fixed morphology and localization can be examined by, for example, superimposing a bright-field image and a fluorescence image.



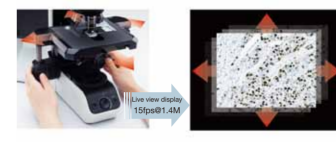
Superimposed image

Color image



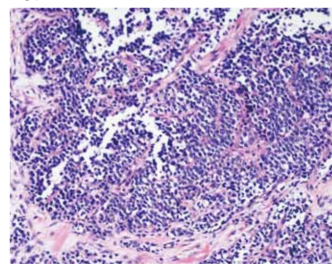
High definition uncompressed live images of 1360×1024 pixels

Live view display of high definition RGB 24-bit color images of 1360 × 1024 pixels at 15 frames per second.Distortion-free focusing or framing is provided because there is no deterioration of image quality due to non-compression, and so sample details are sharp and clear whether the sample is stationary or moving.



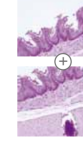
High resolution imaging up to 12.5 megapixels

The DP80 uses pixel-shifting technology to reach a maximum recording image size of 4080 × 3072, a high resolution equivalent to 12.5 megapixels



Faithful panoramic imaging, with high-quality in brightfield or fluorescence

Multiple-region capture of saved images can be easily recombined and restitched seamless into a single image. Numerous annotations and comments can be saved with images for later retrieval. These features are useful for standardization and accuracy control of inspection and research processes.



Research workflow is improved through Olympus cellSens imaging software

Efficient support of observation and experimentation

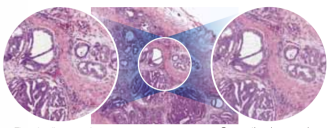
High-guality color and multi-channel imaging is automated with the DP80's switchable CCD sensors and the intuitive Olympus cellSens software interface. From complex image acquisition to image processing to report generation, the researcher can focus on research activities instead of routine labor-intensive acquisition setup and data preparation.



Fine-detail Processing that suppresses pseudo-colors and moiré artifacts

The DP80 is equipped with Fine Detail Processing, which reduces pseudo-colors and moire of ultrastructures and improves resolving power. Clear imaging of details is achieved by fully extracting the resolving power of objective lenses.



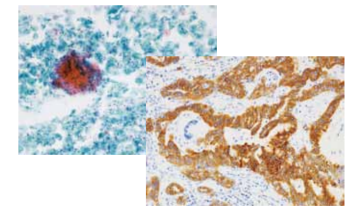


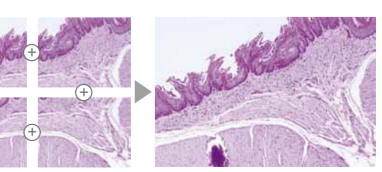
Fine-detail processing

Conventional processing

Superior color reproducibility captures fine differences in color

Subtle hue differences within colors such as brown, blue, and purple were difficult to distinguish in the past, but now, such slight differences in color can be reproduced by incorporating AdobeRGB* color space which reproduces a wide range of color tones and a new algorithm of color reproduction. Color images faithful to the original samples can be acquired. *Color reproduction fidelity depends on monitor specifications. Monitors supporting AdobeRGB are required to accurately reproduce images recorded in AdobeRGB mode





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High quality bright-field and high-sensitivity multi-channe I fluorescence imaging The DP80 alone can provide all of the images that are illustrated below.

Caputured by monochrome CCD with pseudo-colors

Caputured by color CCD

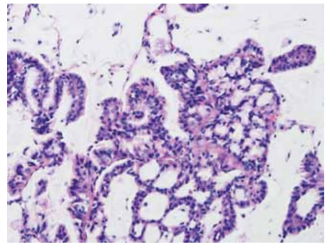


Fig.1: Histology of lung with EML4-ALK fusion gene. (HE stain).

Caputured by color CCD

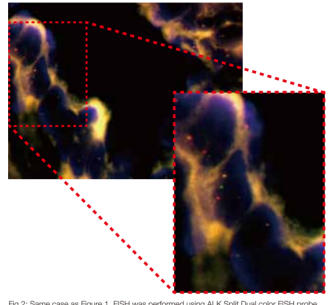


Fig.2: Same case as Figure 1. FISH was performed using ALK Split Dual color FISH probe (green = FITC and red = TexRed) (GSP Laboratory). The abnormal ALK split signal was observed as green and red colored signal, in addition to normal yellow-colored signal.

Caputured by monochrome CCD with pseudo-colors

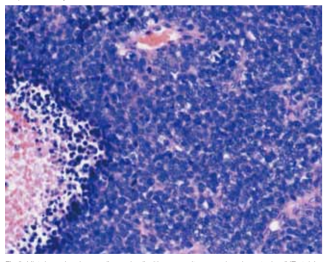


Fig.3: Histology showing small round cell with many mitoses and nuclear atypia. (HE stain)

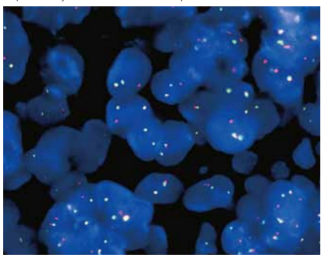


Fig.4: Same case as Figure 3. FISH was performed using EWSR1 (22q12) dual color break apart rearrangement FISH probe (green = spectrum green and orange = spectrum orange) (Vysis™, Abbott Japan). The abnormal EWSR1 split signal was observed as green and orange colored signal, in addition to normal yellow-colored signal.

Image data courtesy of : JAPANESE FOUNDATION FOR CANCER RESEARCH The Cancer Institute, Division of Pathology Noriko Motoi, M.D., Ph.D. Yuichi Ishikawa, M.D., Ph.D.

Caputured by color CCD



Surface of Drosophila melanogaster expressing fluorescence protein in periphearal sensory cells. Image data courtesy of : Institute of Molecular and Cellular Biosciences, University of Tokyo Kei Ito, Ph.D.

Caputured by color CCD

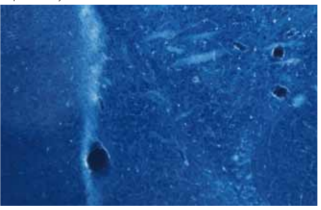
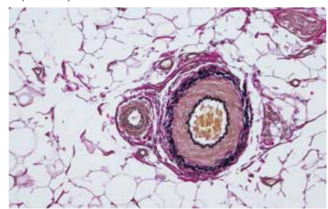


Image data courtesy of : RIKEN BRAIN SCIENCE INSTITUTE Neural Circuitry of Memory Joshua P. Johansen, Ph.D.

Caputured by color CCD

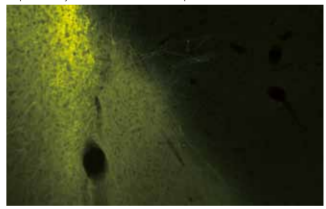


Observation of Collagen type I and type III with multicolor immuno-fluorescence staining during wound healing process Bright-field image of total collagen with Elastica van Gieson (EVG) staining (Left; DP80 color mode) and, multi-fluorescent pseudo-color image of collagen type I labeled with Cy7 and collagen type II erythrocytes and/or other tissue components.

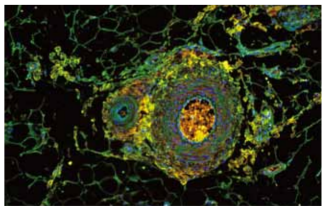
Caputured by monochrome CCD



Caputured by monochrome CCD with pseudo-colors



In the dark field images we can see the borders of the lateral amygdala, a brain region important for fearful emotions. In the fluorescence image are cells and processes expressing a fusion protein of channelrhodopsin/EYFP. Channelrhodopsin is a blue light activated non specific cation channel that is used in 'optogenetics' experiments. We can express channelrhodopsin in lateral amygdala neurons and produce emotional fear memories by activating the cells with blue light. These microscope images allow us to verify that expression of channelrhodopsin has occured in the lateral amygdala.



Caputured by monochrome CCD with pseudo-colors

labeled with Cy5 respectively (Right; DP80 monochrome mode). Location of the collagen type I and III is confirmed clearly by the long-wavelength observation without auto-fluorescence noise such as