

Operating Manual **Axioscope 5/7/Vario**



Upright Microscope for Routine and Entry-level Research

We make it visible.

Knowledge of this manual is essential for the operation of the instrument. Please familiarize yourself with the contents of this manual and pay special attention to instructions concerning safe operation of the instrument.

The specifications are subject to change; the manual is not covered by an update service.

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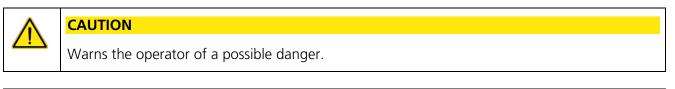
1.1 Notes on instrument safety

The Axioscope 5/7/Vario microscopes were engineered, manufactured and tested in accordance with the DIN EN 61010-1 (IEC 61010-1) and IEC 61010-2-101 safety standards for electrical measuring, control and laboratory equipment.

The Axioscope 5/7/Vario microscopes meet the requirement of Low Voltage Directive 2014/35/EU, EMC(Electromagnetic Compatibility) Directive 2014/30/EU, and carry the **C C** conformity marking.

The present manual contains information and safety warnings with which the operator must comply.

The following warning, instructional and informational symbols are used in the manual:



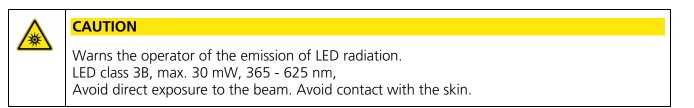
CAUTION

Warns the operator of hot surfaces.



CAUTION

Warns the operator of the emission of UV radiation.





CAUTION

Warns the operator of crushing hazard.



CAUTION

Warns the operator to disconnect the instrument from the power supply before opening or doing any work on the instrument.



ATTENTION

This symbol indicates the risk of damage to the instrument or the system.

NOTE

This symbol indicates information or instructions which must be followed especially carefully.

The Axioscope 5/7/Vario microscopes and their original accessories may be used only for the microscopy procedures described in this operating manual.

The following guidelines are especially important:



CAUTION

The microscope may only be plugged into an electrical outlet equipped with a safety contact. The safety feature must not be disabled by using an extension cord which does not have a protective ground conductor.



CAUTION

Whenever it becomes apparent that any of the safety mechanisms are out of order, the microscope must be switched off and protected from any inadvertent use. Please contact the ZEISS Service Department or the Carl Zeiss Microscopy Service before switching the microscope on again.

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<u>!\</u>	

CAUTION

The stands are equipped with integrated power supply units. These units adapt to a voltage range of 100 to 240 V and a frequency range of 50 to 60 Hz. No further voltage adjustment is necessary on the microscope itself.

The 12 V DC 100 W external power supply unit for HAL 100 covers a voltage range from 100 to 127 V AC and 200 to 240 V AC \pm 10%, and a frequency range of 50 to 60 Hz. It adapts automatically to the applied voltage.

The 24 V DC 30 W external power supply unit for Axioscope 5 Vario covers a voltage range from 100 V to 240 V and a frequency range of 50 to 60 Hz.

The power supply units for the HBO 100 adapt to a voltage range from 100 to 240 V AC and a frequency range of 50 to 60 Hz. They adapt automatically to the applied voltage. No further voltage adjustment is necessary.

Before you switch on the instrument, please verify that the voltage you are about to use is suitable.

	CAUTION
	Always disconnect the microscope from the power supply before you open it or change a fuse!
	Make sure that the fuses are suitable for the applied nominal current. Never use any makeshift fuses and do not short-circuit the fuse holders.



CAUTION

The microscopes do not have any special safety devices to protect users from acid or from potentially infectious, toxic, radioactive or other samples that may be hazardous to your health. Compliance with all statutory requirements, especially national accident prevention regulations, is required when handling such samples.



CAUTION

Only authorized personnel are permitted to operate the instruments. The personnel must have been instructed about and be aware of the risks involved in using the microscope. The Axioscope is a precision instrument whose functionality can easily be damaged or even destroyed when handled improperly.



CAUTION

Operating the instrument in an area with a potentially explosive atmosphere is prohibited. It may be operated only on stable, non-flammable surfaces.

Specimens must be disposed of appropriately in accordance with applicable statutory regulations and internal work instructions.



CAUTION

Gas discharge lamps, such as the HBO 100, emit ultraviolet radiation which may cause injuries to your eyes and skin. Never look directly into the beam and avoid exposing your skin directly to the beam. Never operate the microscope without the necessary safety equipment (e.g. special damping filters or the fluorescence protection screen). Gas discharge lamps have a high internal pressure when hot. Never change the lamps while they are still hot and always use protective gloves and masks when changing them.



CAUTION

Avoid any contact with the hot lamp case. Before you change a lamp, disconnect the instrument from the power supply and wait 15 minutes for the lamp to cool down.



CAUTION

When the lamp module is equipped with appropriate LED modules, or when a white light source coupled by a fluid light conductor is used, ultraviolet radiation may leak which can cause injury to your eyes and skin. Never look directly into the light and avoid exposing your skin directly to the light. Always use the appropriate safety equipment when operating the microscope.



ZEISS

CAUTION

Never look into the light beam of the illumination equipment - with or without optical instruments - not even if you just want to look at a specimen. Irreparable eye damage may occur!



CAUTION

Do not hold any flammable or easily combustible materials into the light beam.



CAUTION

The immersion fluid Immersol 518 N[®] is irritating to the skin. Avoid any contact with skin, eyes and clothes. Read the safety data sheets on Immersol 518 N[®], Immersol 518 F[®] and Immersol W[®] carefully. If skin contact occurs, wash off with lots of water and soap. If eye contact occurs, flush with water immediately for at least 5 minutes. Seek medical assistance if irritation continues.



CAUTION

Dispose of immersion fluid Immersol 518 $N^{\rm \tiny (8)}$ appropriately: Do not allow to contaminate surface water or get into drains.



CAUTION

For stands equipped with motorized focusing drives, there is a risk of fingers being crushed in the working area when the microscope stage is lowered. Do not reach into the area between the microscope stage/stage carrier/condenser carrier and the base of the stand while the stage is being lowered.



ATTENTION

The manufacturer assumes no liability for any use of the microscope, its assemblies or individual parts if these are used in in a manner or for any purpose other than those described in this operating manual. This also applies to any maintenance or repair work not performed by authorized maintenance and repair personnel. Moreover, the warranty and any claims thereunder will be voided.



ATTENTION

Microscope lamps radiate a lot of heat which could damage the heat-sensitive fluorescence filters. Do not remove the heat protection filter when using a fluorescence filter; otherwise the fluorescence filter may not function properly.

	ATTENTION
	Dirt and dust may affect the microscope's performance. Protect it by using a dust cover when it is not in use. Always make sure that the instrument is switched off before covering it up.
	ATTENTION
Ŀ	Closing or covering the ventilation slits may result in heat accumulation which could damage the instrument and even start a fire. Ensure that the ventilation slits are always kept open, uncovered and clear. Do not insert or drop anything into them or let anything fall into them.
	ATTENTION
Ċ	Always position equipment in a manner that allows easy disconnection from the power supply.
	ATTENTION
	Do not use a power supply cord with an inadequate rating.
_	
	ATTENTION
Ŀ	When using the HAL or HBO illuminators, a base plate for the Axioscope (000000-2202-526) is mandatory.
	ATTENTION
	Do not dispose of defective microscopes in the household waste. Comply with the applicable statutory regulations for their disposal. Specimens must also be disposed of appropriately in accordance with applicable statutory regulations and internal work instructions.

1.2 Warning labels on the microscopes

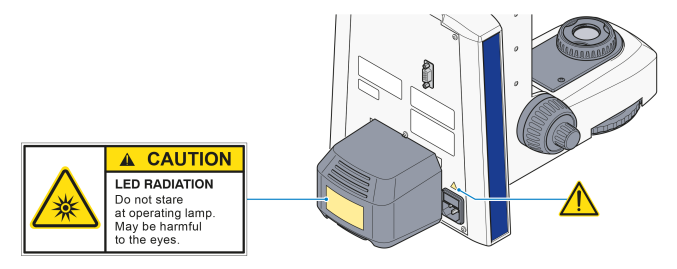


Fig. 1-1 Warning labels on microscopes with LED light source for transmitted light

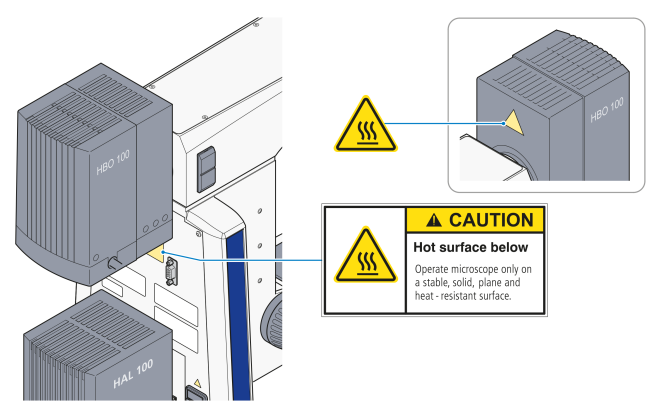


Fig. 1-2 Warning labels on microscopes with HAL 100 for transmitted light and HBO 100 for reflected light

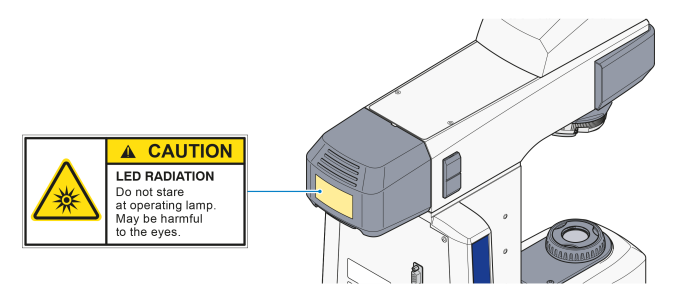


Fig. 1-3 Warning label on microscopes with LED light source for reflected light

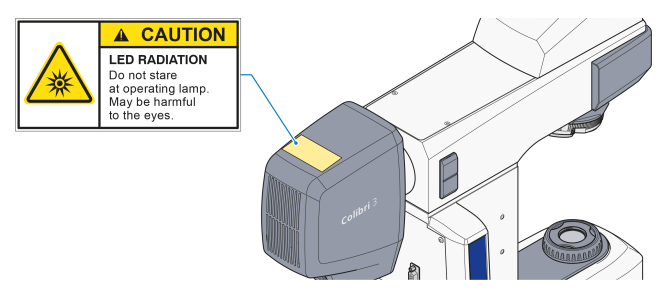
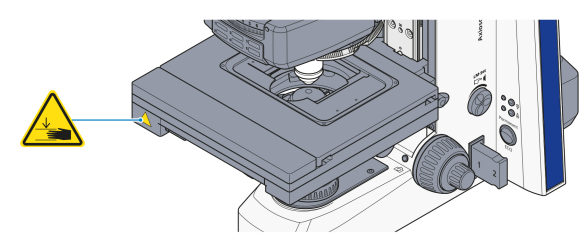


Fig. 1-4 Warning label on microscopes with Colibri 3 light source for reflected light





1.3 Notes on the warranty

The manufacturer guarantees that the instrument is free of any material and workmanship defects upon delivery. If you become aware of any deficiencies, please contact us immediately and take all necessary precautions in order to avoid further damage. Upon notice of deficiencies it is up to the manufacturer's discretion to correct the deficiencies or to deliver a defect-free instrument. Defects due to ordinary wear and tear (especially on wearing parts) and to improper handling are not covered by our warranty.

The manufacturer is not liable for damage to the instrument due to incorrect operation, negligence or any other manipulation of the instrument, in particular due to the removal or replacement of instrument parts or due to the use of accessories from other manufacturers. This will immediately void the warranty.

No maintenance or repair work, except for the instances mentioned in the manual, may be performed on the microscopes. Only ZEISS service personnel or personnel authorized especially by ZEISS may perform repair work on the microscopes. In the event of malfunction of your instrument, please contact the ZEISS Microscopy Service Department (see page 147) or the ZEISS agency assigned to your country.

2 DESCRIPTION OF THE INSTRUMENT

2.1 Intended use

The Axioscope 5/7/Vario microscopes are microscopes designed for biological and medical applications as well as materials analyses.

Depending on the configuration of the microscope stand, they may be used with transmitted light only or with a combination of transmitted light and reflected light.

Typical areas of use for the Axioscope 5/7/Vario in biomedical applications include:

- medical analysis in laboratories, hospitals, physicians' practices,
- science and research (colleges, universities) in medical and biological fields,
- industrial applications (pharmacology, food technology),
- analysis of blood and tissue samples of the human body.

Typical fields of application of the Axioscope 5/7/Vario in Materials applications include:

- routine research in aerospace and metallographic laboratories,
- quality assurance and control in the automobile and mechanical engineering industries
- failure analysis in the electronics and semiconductor industries,
- The analysis of optical properties of minerals in geoscience studies, and
- morphology studies in the chemical, polymer and mineral exploration industries.

Depending on the accessories with which each instrument is equipped, the following microscopy and contrasting techniques can be performed:

Transmitted light

- Bright-field (BF)
- Dark-field (DF)
- Phase contrast (PhC)
- Differential interference contrast (DIC)
- PlasDIC
- Polarization (Pol): orthoscopy and conoscopy

Reflected light

- Bright-field (BF)
- Dark-field (DF)
- Differential interference contrast (DIC)
- Circular differential interference contrast/total interference contrast (C-DIC/TIC)
- Fluorescence
 - Polarization (Pol)

The binocular photo tubes can be connected via a special adapter with a microscopy camera, a reflex camera or a digital/video camera for documentation purposes.

Using a ZEISS Axiocam 202 mono or Axiocam 208 color camera, the camera can be connected to an external HD monitor via HDMI connection or to a PC/laptop via USB connection to control camera functionality. Advanced analysis functions for images are available if the ZEN software is installed on the PC. Furthermore, the camera functions can be controlled via WiFi connection by tablet PC or smartphone using the Labscope/Matscope app.

2.2 Technical data

Dimensions (length x width x height)

Microscope stand for Axioscope 5/7	approx. 293.5 mm x 240 mm x 367.5 mm
Microscope stand for Axioscope Vario	approx. 429 mm x 458.5 mm x 700 mm

Weight

Microscope stand for Axioscope 5/7	
(depending on stand configurations and accessories)	approx. 14 to 20 kg
Microscope stand for Axioscope Vario	approx. 32 kg

Ambient conditions

Shipping (in packaging):

Permissible ambient temperature	40 to +70 °C
Permissible humidity (no condensation)	max. 75% at 35 °C
Storage:	
Permissible ambient temperature	+10 to +40 °C
Permissible humidity (no condensation)	max. 75% at 35 °C
Operation:	
Permissible ambient temperature	+10 to +40 °C
Permissible relative humidity (no condensation)	max. 75% at 35 °C
Highest permitted altitude of use	max. 2000 m
Air pressure	800 hPa to 1060 hPa

Operational specifications

Operational area	Closed rooms
Protective class	
Protection type	IP 20
Electrical safety	in accordance with DIN EN 61010-1 (IEC 61010-1)
	in conjunction with CSA and UL regulations
Overvoltage category	
	conforming to EN 55011 Class B
	conforming to DIN EN 61326/A1
Mains voltage for the Axioscope 5/7	-
with internal power supply unit	
Mains voltage for the Axioscope Vario	
with external power supply unit	
Mains frequency	
Power consumption of the Axioscope 5	
with internal power supply unit	
Power consumption of the Axioscope 7	
· · ·	
Power consumption of the Axioscope Vario	
with external power supply unit	

HBO 100 W power supply unit

Range of application	indoor
Protection class	
Protection type	IP 20
Mains voltage	100 VAC 240 VAC
	No voltage adjustment required
Mains frequency	50 to 60 Hz
Power consumption with HBO 103 in operation	155 VA

Fuses in accordance with IEC 127

Axioscope 5/7 microscope stand for LED illumination in transmitted light:	2 T 3.	15 A/H,	5x20 mm
Axioscope 5/7 microscope stand for HAL 50 illumination in transmitted light:	2 T 3.	15 A/H,	5x20 mm
HBO 100 W power supply unit	T 2	.0 A/H,	5x20 mm
12 V DC 100 W external power supply unit	2 T 5	.0 A/H,	5x20 mm

Light sources

LED illumination transmitted light/reflected light	
Power consumption	max. 10 W
Adjustment of light source	continuous approx. 10 to 800 mA
Halogen lamp	
Adjustment of light source	infinitely variable from approx. 3 to 12 V
Halogen lamp	
Adjustment of light source	infinitely variable from approx. 3 to 12 V
Mercury vapor short arc lamp	HBO 103 W/2
Power consumption for HBO 103 W/2	

Axioscope 5/7/Vario

Stand with manual/motorized stage focusing	
Coarse drive	approx. 4 mm/rotation
Fine drive	approx. 0.4 mm/rotation; approx. 4 µm scale intervals
Lifting range	approx. 25 mm
Height stop	mechanically variable
Condenser 0.9/1.25 H with optional	
modulator disk	for bright-field, dark-field and
	phase contrast 1, 2, 3 or PlasDIC
Manual objective change	via nosepiece, 6-positions, M27, coded
Manual reflector module change	via reflector slider, 2-positions coded, reflector turret 4-positions coded or 6-positions coded

2.3 Interface diagram

The following figure shows a diagram of the microscope's hardware interfaces. The Axioscope 5 stand TL/RL with LED/HAL illumination serves as the example.

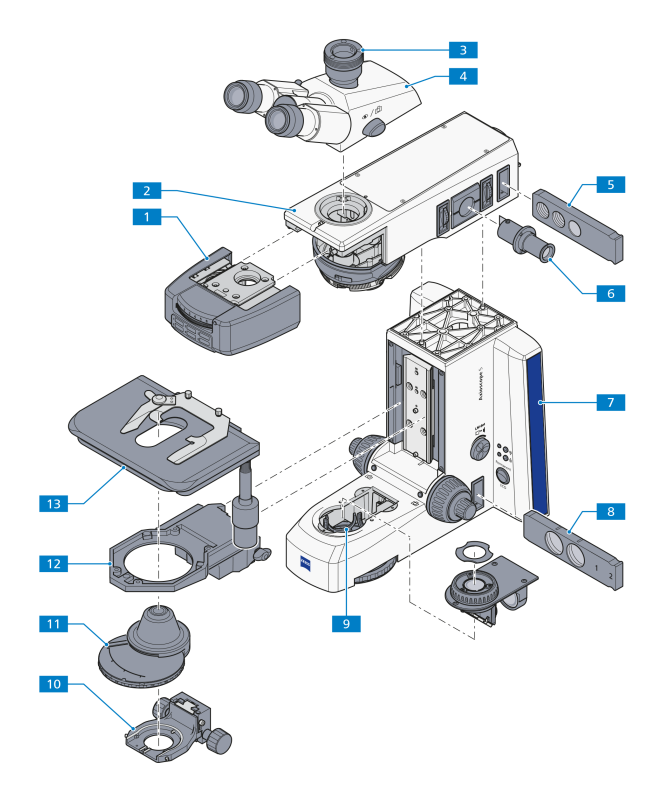


Fig. 2-1 Interface diagram (example: Axioscope 5 stand, TL/RL)

Key to Fig. 2-1:

- 1 Reflector turret
- 2 Upper part of the stand
- 3 Camera interface 60N on the photo tube
- Tube/photo tube 4
- Filter slider for reflected light 5
- 6 Adjustment tool
- 7 Lower part of the stand
- Filter slider for transmitted light 8
- Filter wheel for transmitted light 9 Condenser carrier
- 10
- Condenser 11
- 12 Stage carrier
- Stage 13

Controls and functional elements on the microscope 2.4

2.4.1 Stand models

Nine stand models are available in the delivery program:

1. Axioscope 5 stand HAL 50	(430035-9032-000)
2. Axioscope 5 stand, Bio-TL	(430035-9201-000)
3. Axioscope 5 stand, Bio-TL/RL	(430035-9061-000)
4. Axioscope 5 stand, Pol-TL	(430035-9261-000)
5. Axioscope 5 stand, Pol-TL/RL	(430035-9291-000)
6. Axioscope 5 stand, Mat-TL/RL	(430035-9121-000)
7. Axioscope 7 stand, Mat-TL/RL, motorized	(430035-9330-000)
8. Axioscope 5 Vario material stand	(430035-9150-000)
9. Axioscope 5 Mat-RL	(430035-9091-000)

2.4.2 Axioscope 5 stands, Bio-TL

Transmitted light stands for bioscience equipped with the following main components:

- 1. Version 1 (430035-9032-000): HAL 50 halogen illumination Version 2 (430035-9201-000): LED 10 W-TL illumination or optionally HAL 100 halogen illumination using an external power supply
- 2. Nosepiece 6-positions M 27 HF, coded
- 3. Mechanical stage 75x50 R
- 4. Binocular photo tube 30x/23 (50:50), reversed image
- 5. Eyepiece E-PL 10x/23 GW foc.
- 6. Condenser 0.9/1.25H
- 7. Coded changeable reflector turrets and reflector slider

Key to Fig. 2-2:

- 1 Eyepieces
- 2 Binocular tube part
- **3** Binocular tube/photo tube (with mounted Axiocam 208, optionally)
- 4 Transmitted light stand (consisting of upper part and lower part)
- **5** Transmitted light illuminator
- 6 Intensity/LM knob for light intensity and Light Manager function (LM)
- 7 Indicator light for transmitted light
- 8 Permanent/ECO mode switch
- **9** Filter slider transmitted light
- **10** Focusing drive fine adjustment (left & right)
- **11** Focusing drive coarse adjustment (left & right)
- 12 Coaxial knurled knob for adjusting the mechanical stage in direction Y
- **13** Coaxial knurled knob for adjusting the mechanical stage in direction X
- **14** Snap buttons
- **15** Nosepiece
- **16** 6-position filter wheel (operable from left & right)
- 17 Field diaphragm
- 18 Condenser with aperture diaphragm and centering screws (with optional modulator disk)
- 19 Knurled knob for adjusting the condenser vertically (left & right)
- 20 Stage carrier
- 21 Condenser carrier
- 22 Release lever for height stop on focusing drive
- 23 Main switch I/O
- 24 Mechanical stage with dual specimen holder

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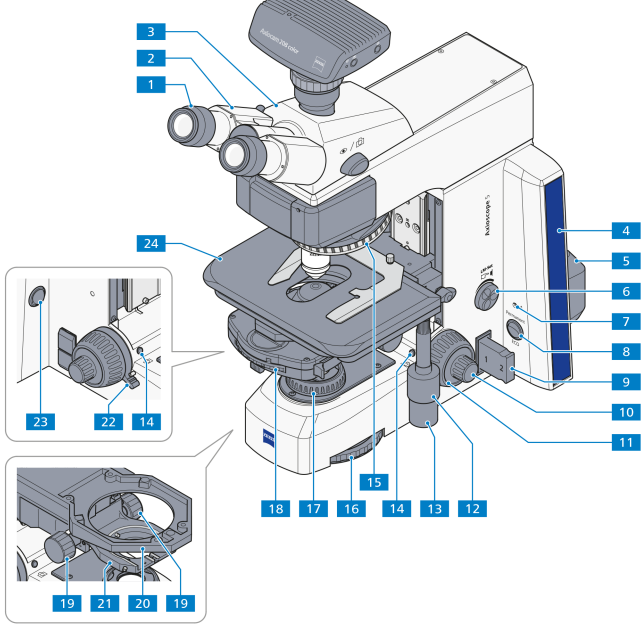


Fig. 2-2 Controls and functional elements of the Axioscope 5 stands, Bio-TL

2.4.3 Axioscope 5 stand, Bio-TL/RL

Transmitted and reflected light fluorescence stand (430035-9061-000) for bioscience equipped with the following main components:

- 1. LED 10 W-TL illumination or optionally with HAL 100 halogen illumination using an external power supply for transmitted light applications
- 2. Colibri 3, HBO 100, HXP illumination for reflected light fluorescence applications
- 3. Nosepiece 6-positions M27 HF, DIC, coded
- 4. Mechanical stage 75x50 R
- 5. Binocular photo tube 30x/23 (50:50) reversed image
- 6. Eyepiece E-PL 10x/23 GW foc.
- 7. Condenser 0.9/1.25H
- 8. Coded changeable reflector turret or reflector slider

Key to Fig. 2-3:

- 1 Eyepieces
- 2 Binocular tube part
- 3 Binocular tube/photo tube (with mounted Axiocam 208 or Axiocam 202, optionally)
- 4 Field diaphragm
- **5** Adjustment tool
- 6 Aperture diaphragm or FL attenuator
- **7** Filter slider for reflected light
- 8 Reflected light illuminator
- 9 Transmitted and reflected light stand (consisting of upper part and lower part)
- 10 Intensity/LM knob for light intensity, Light Manager Function (LM) and switching between fluorescence channel
- **11** Transmitted light illuminator
- **12** Indicator light for reflected light
- **13** Reflected light (RL) button
- 14 Transmitted light (TL) button
- **15** Indicator light for transmitted light
- 16 Permanent/ECO mode switch
- **17** Filter slider transmitted light
- **18** Focusing drive fine adjustment (left & right)
- **19** Focusing drive coarse adjustment (left & right)
- 20 Coaxial knurled knob for adjusting the mechanical stage in direction Y
- 21 Coaxial knurled knob for adjusting the mechanical stage in direction X
- 22 Snap buttons
- 23 Nosepiece
- 24 6-position filter wheel (operable from left & right)
- 25 Field diaphragm
- 26 Condenser with aperture diaphragm (with optional modulator disk)
- 27 Knurled knob for adjusting the condenser vertically (left & right)
- 28 Stage carrier
- **29** Condenser carrier with centering screws
- **30** Release lever for height stop on focusing drive
- 31 Main switch I/O
- **32** Mechanical stage with specimen holder
- **33** Knurled wheel for switching the reflector turret position
- **34** Reflector turret (for replaceable reflector modules)

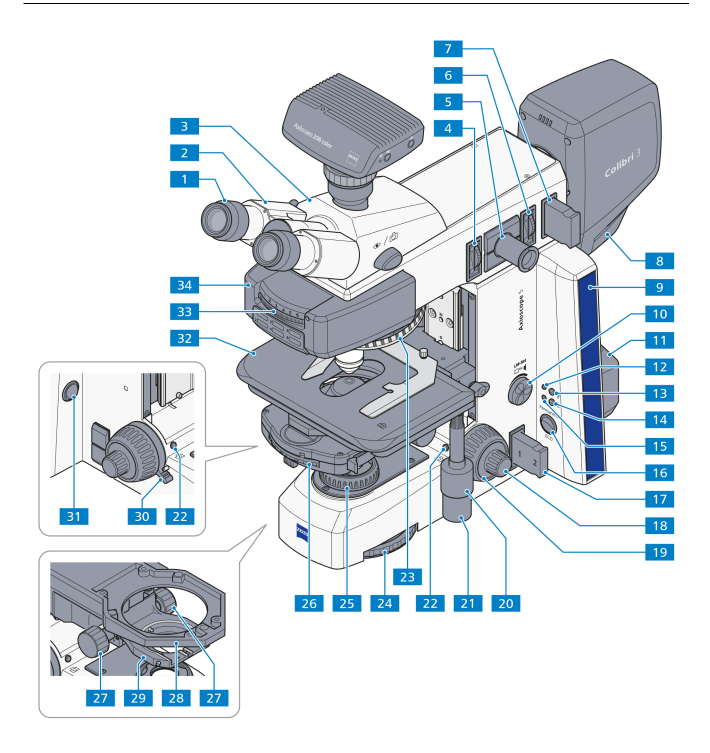


Fig. 2-3 Controls and functional elements of the Axioscope 5 stand, Bio-TL/RL

2.4.4 Axioscope 5 stand, Pol-TL/RL

Transmitted and reflected light stand for polarization (430035-9291-000) equipped with the following main components:

- 1. LED 10 W-TL illumination or optionally with HAL 100 halogen illumination using an external power supply for transmitted light applications
- 2. LED 10 W-RL illumination for reflected light applications
- 3. Nosepiece 6-positions M27: 5-positions HF/DF/Pol, centerable and 1x HF/DF/DIC fixed, coded
- 4. Pol rotary stage, 360° with 45° click stops (freely definable)
- 5. Binocular photo tube 30x/23 (100:0/0:100), reversed image
- 6. E-PL 10x/23 GW foc. Pol Eyepiece with reticle
- 7. Pol condenser 0.9 H
- 8. Mount for polarization slider A 14x40 mm, d = 36 mm
- 9. Polarizer D, 90° rotatable, removable
- 10. Quartz depolarizer with tube-lens for Axioscope tubes
- 11. Coded changeable reflector turret or reflector slider

Key to Fig. 2-4:

- 1 Eyepieces
- Binocular tube part 2
- 3 Binocular tube/photo tube
- 4 Slot for polarizer slider A 60x30 mm
- 5 Field diaphragm
- 6 Aperture diaphragm
- 7 Filter slider for reflected light
- Reflected light illuminator 8
- Transmitted and reflected light stand (consisting of upper part and lower part) 9
- Intensity/LM knob for light intensity and Light Manager function (LM) 10
- Transmitted light illuminator 11
- Indicator light for reflected light 12
- 13 Reflected light (RL) button
- Transmitted light (TL) button 14
- Indicator light for transmitted light 15 Permanent/ECO mode switch 16
- Filter slider transmitted light
- 17
- Focusing drive fine adjustment (left & right) 18 Focusing drive – coarse adjustment (left & right)
- 19 Snap buttons 20
- 21 Nosepiece
- 6-position filter wheel (operable from left & right) 22
- 23 Field diaphragm
- Condenser with aperture diaphragm (with optional modulator disk) 24
- 25 Knurled knob for adjusting the condenser vertically (left & right)
- 26 Polarizer
- Condenser carrier with centering screws 27
- 28 Stage carrier
- Release lever for height stop on focusing drive 29
- 30 Main switch I/O
- 31 Pol rotary stage
- 32 Knurled wheel for switching the reflector turret position
- 33 Reflector turret (for replaceable reflector modules)

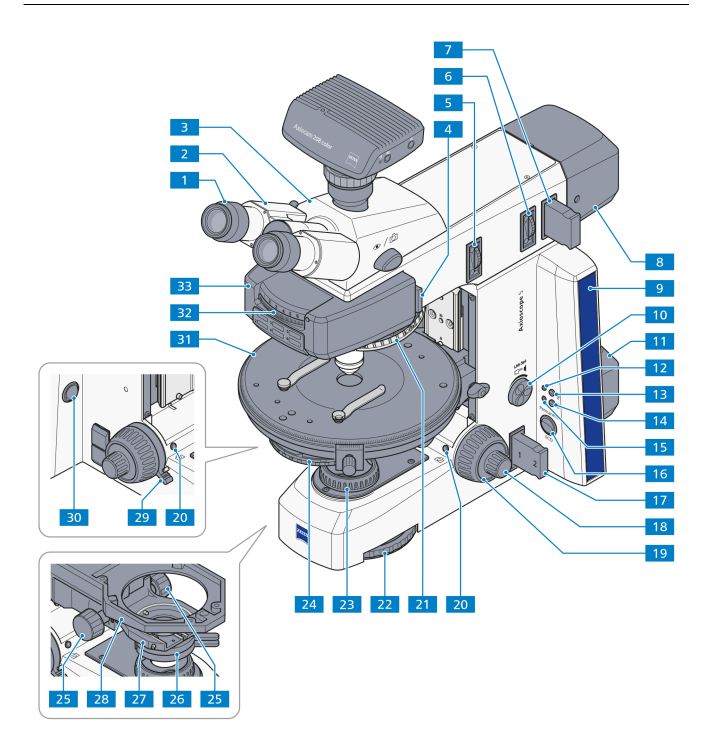


Fig. 2-4 Controls and functional elements of the Axioscope 5 stand, Pol-TL/RL

2.4.5 Axioscope 5 stand, Mat-TL/RL

Transmitted and reflected light stand for material (430035-9121-000) equipped with the following main components:

- 1. LED 10 W-TL illumination or optionally with HAL 100 halogen illumination using an external power supply for transmitted light applications
- 2. LED 10 W-RL illumination for reflected light applications
- 3. Nosepiece 6-positions M27 HF/DF/DIC, coded
- 4. Mechanical stage 75x50 R
- 5. Binocular photo tube 20x/23 (100:0/0:100), upright image with sliding prism
- 6. Eyepiece E-PL 10x/23 GW foc.
- 7. Condenser 0.9/1.25H
- 8. Mount for polarization slider A 14x40 mm d = 36 mm
- 9. Coded changeable reflector turret or reflector slider

Key to Fig. 2-5:

- 1 Eyepieces
- 2 Binocular tube part
- **3** Binocular tube/photo tube
- 4 Slot for polarizer slider A 60x30 mm
- **5** Field diaphragm
- 6 Aperture diaphragm
- 7 Filter slider for reflected light
- 8 Reflected light illuminator
- 9 Transmitted and reflected light stand (consisting of upper part and lower part)
- **10** Intensity/LM knob for light intensity and Light Manager function (LM)
- **11** Transmitted light illuminator
- **12** Indicator light for reflected light
- **13** Reflected light (RL) button
- **14** Transmitted light (TL) button
- **15** Indicator light for transmitted light
- **16** Permanent/ECO mode switch
- **17** Filter slider transmitted light
- **18** Focusing drive fine adjustment (left & right)
- **19** Focusing drive coarse adjustment (left & right)
- 20 Coaxial knurled knob for adjusting the mechanical stage in direction Y
- **21** Coaxial knurled knob for adjusting the mechanical stage in direction X
- 22 Snap buttons
- 23 Nosepiece
- 24 6-position filter wheel (operable from left & right)
- 25 Field diaphragm
- 26 Condenser with aperture diaphragm (with optional modulator disk)
- 27 Knurled knob for adjusting the condenser vertically (left & right)
- 28 Stage carrier
- **29** Condenser carrier with centering screws
- **30** Release lever for height stop on focusing drive
- 31 Main switch I/O

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- 32 Mechanical stage with specimen holder
- **33** Knurled wheel for switching the reflector turret position
- 34 Reflector turret (for replaceable reflector modules)

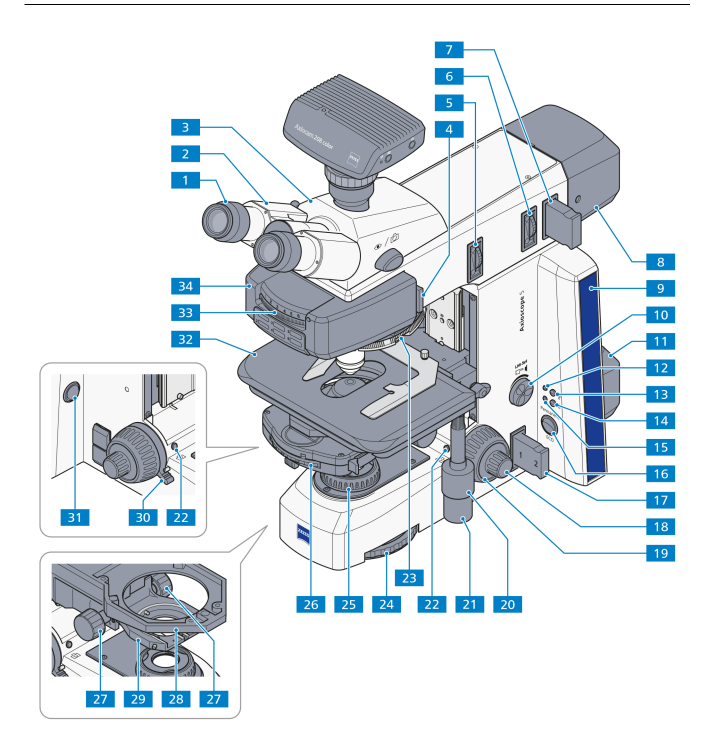


Fig. 2-5 Controls and functional elements of the Axioscope 5 stand, Mat-TL/RL

2.4.6 Axioscope 7 stand, Mat-TL/RL mot.

Transmitted and reflected light stand for material (430035-9330-000) equipped with the following main components:

- 1. LED 10 W-TL illumination or optionally with HAL 100 halogen illumination using an external power supply for transmitted light applications
- 2. LED 10 W-RL illumination for reflected light applications
- 3. Nosepiece 6-positions M27 HF/DF/DIC, coded
- 4. Motorized 80x60 mechanical stage
- 5. Binocular photo tube 20x/23 (100:0/0:100), upright image with sliding prism
- 6. Eyepiece E-PL 10x/23 GW foc.
- 7. Condenser 0.9/1.25H
- 8. Mount for polarization slider A 14x40 mm d = 36 mm
- 9. Coded changeable reflector turret or reflector slider

Key to Fig. 2-6:

- **1** Eyepieces
- 2 Binocular tube part
- **3** Binocular tube/photo tube
- 4 Slot for polarizer slider A 60x30 mm
- 5 Field diaphragm
- 6 Aperture diaphragm
- 7 Filter slider for reflected light
- 8 Reflected light illuminator
- 9 Transmitted and reflected light stand (consisting of upper part and lower part)
- **10** Intensity/LM knob for light intensity and Light Manager function (LM)
- **11** Transmitted light illuminator
- **12** Indicator light for reflected light
- **13** Reflected light (RL) button
- **14** Transmitted light (TL) button
- **15** Indicator light for transmitted light
- 16 Permanent/ECO mode switch17 Filter slider for transmitted light
- 17 Filter slider for transmitted light18 Focusing drive fine adjustment (left & right)
- Focusing drive line adjustment (left & right)
 Focusing drive coarse adjustment (left & right)
- 20 Snap button
- 20 Shap butto 21 Nosepiece
- **22** 6-position filter wheel (operable from left & right)
- 23 Field diaphragm
- 24 Condenser with aperture diaphragm (with optional modulator disk)
- 25 Knurled knob for adjusting the condenser vertically (left & right)
- 26 Stage carrier
- 27 Condenser carrier with centering screws
- 28 Stage control button
- 29 Main switch I/O
- **30** Mechanical stage 80x60 mot. with specimen holder
- **31** Knurled wheel for switching the reflector turret position
- **32** Reflector turret (for replaceable reflector modules)

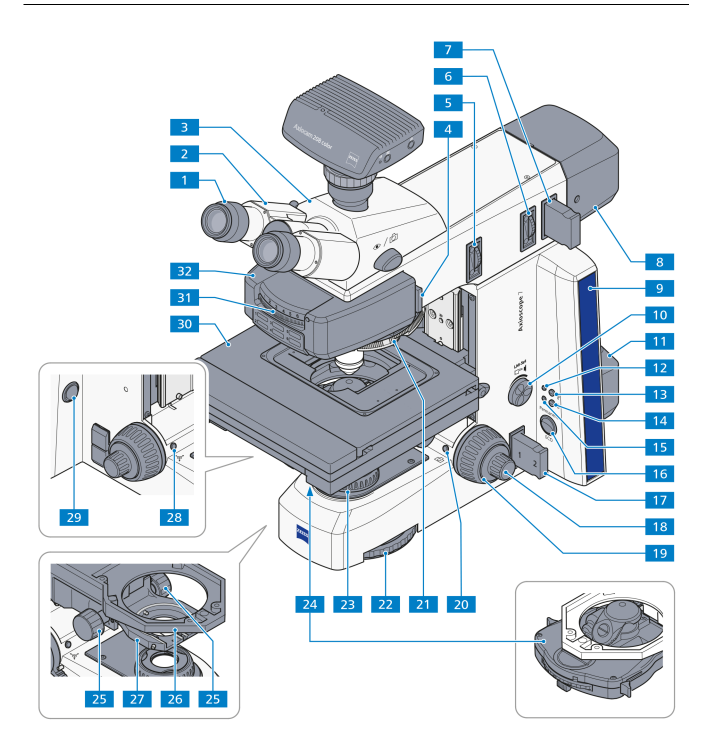


Fig. 2-6 Controls and functional elements of the Axioscope 7 stand, Mat-TL/RL mot.

2.4.7 Axioscope 5 Vario material stand

Transmitted and reflected light stand for material (430035-9150-000) equipped with the following main components:

- 1. Upper stand part including reflected-light illumination LED 10 W, with interface for HAL / HBO / HXP
- 2. Focusing gear box Axioscope Vario, 15 mm focus lift
- 3. Nosepiece 6-positions HF/DF/DIC, coded
- 4. Binocular photo tube 20x/23 (100:0/0:100), upright image with sliding prism
- 5. Eyepiece E-PL 10x/23 GW foc.
- 6. Coded changeable reflector turret or reflector slider
- 7. Stand column 560 mm with crank for vertical adjustment
- 8. Stage carrier and base plate

Key to Fig. 2-7:

- 1 Eyepieces
- 2 Binocular tube part
- **3** Binocular tube/photo tube
- 4 Release lever for vertical adjustment
- **5** Stand column Axioscope Vario 560 mm
- 6 Hand wheel for vertical adjustment7 Slot for polarizer slider A 60x30 mm
- 8 Field diaphragm
- 9 Aperture diaphragm
- **10** Filter slider for reflected light
- **11** Reflected light illuminator
- 12 Intensity/LM knob for light intensity and Light Manager function (LM)
- **13** Focusing drive coarse adjustment
- **14** Focusing drive fine adjustment
- **15** Snap button
- **16** Indicator light for reflected light
- 17 Permanent/ECO mode switch
- **18** Base plate
- **19** Stage carrier, H=140 mm
- 20 Nosepiece
- 21 Knurled wheel for switching the reflector turret position
- 22 Reflector turret (for replaceable reflector modules)

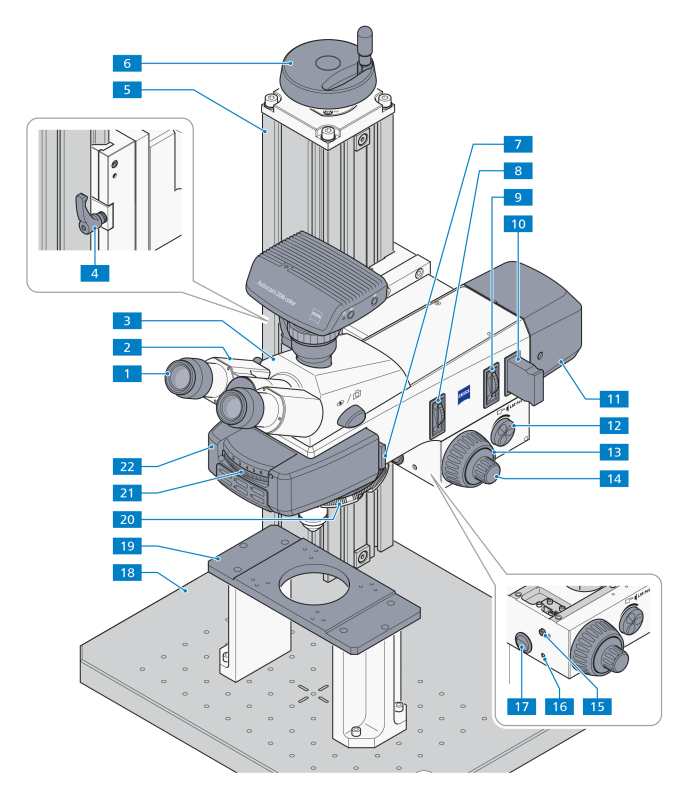


Fig. 2-7 Controls and functional elements of the Axioscope 5 Vario material stand

Functions of stands keys and display elements 2.4.8

For the location of the keys and display elements on your stand, refer to sections 2.4.2 - 2.4.7.

R NOTE Each time the microscope is switched on, the first setting of parfocal value for any nosepiece position will clear the parfocal settings for ALL nosepiece positions.

	Availability					
Кеу	Axioscope 7 mot.	Axioscope 5	Axioscope Vario	Action	Functionality/Description	
Main switch I/O (see Fig. 2-6/29)	Yes	Yes	Yes	I = on; O = off	Switches the microscope on/off	
Permanent/ ECO switch (see Fig. 2-6/16)	Yes	Yes	Yes	Switch:	 Switches between Permanent (continuous) mode and ECO mode of the microscope illumination: Permanent mode active: illumination is continuously switched on. ECO mode active: illumination switches off after 15 minutes without action. Note: Don't use ECO mode for experiments involving time-lapse or video recording. 	
Indicator light (see Fig. 2-6/12 and 15)	Yes	Yes	Yes	Blinking ***:	Indicates microscope function (normal/setting)	
RL/TL button (see Fig. 2-6/ 13 and 14)	Yes	Option	No	Short press *:	Switches RL/TL illumination alternately on/off. The respective indicator light is continuously illuminated. Second press to the RL/TL button to turn off/on the illumination (the indicator light not affected).	
	Yes	Yes	Yes	Turn:	Controls the light intensity of the active light source	
Intensity/ LM knob (see Fig. 2-6/ 10)				Short press *:	Repeated short pressing switches a single LED or all LEDs of the fluorescence light source together on or off.	
				Long press **:	Saves the set light intensity. During this action, the indicator light blinks twice in GREEN and the image background appears BLACK for 300 ms (this does not apply to halogen illumination).	
				Long press for 20 s:	Activates the factory default settings (enables Light Manager, sets light intensity to the initial value, clears all saved parfocal positions). When the knob is pressed, the indicator light starts blinking in RED after 3 s until 20 s is reached. After 20 s, the indicator light blinks in GREEN. Then release the knob. The indicator light turns to GREEN permanently if the system reset is done. After factory default reset, re-power the system.	

Short press: hold less than 1 second, then release.

Long press: hold at least 1.5 seconds. BLINK: the indicator light alternately goes on/off at 500 ms intervals. ***

	Availability					
Кеу	Axioscope 7 mot.	Axioscope 5	Axioscope Vario	Action	Functionality/Description	
Left Snap				Short press *:	Snaps an image	
button (see Fig. 2-5/ 22) (if ZEISS Axiocam attached)	No	Yes	No	Long press **:	Starts video recording; another short press is required to stop recording. When recording is finished, the monitor display turns BLACK for 300 ms.	
Right Snap button (see Fig. 2-6/ 20) (if ZEISS Axiocam attached)		Yes	Yes	Short press:	Snaps an image; when the snap is completed, the monitor display turns BLACK for 50 ms.	
	Yes			Long press **:	Starts video recording; another short press is required to stop recording. When recording is finished, the monitor display turns BLACK for 300 ms.	
Snap button + Intensity/LM knob (see Fig. 2-6/20+10)	Yes	Yes	Yes	Long press **:	Enables/disables the Light Manager: - Disabling: The indicator light blinks GREEN / ORANGE / GREEN in sequence. - Enabling: The indicator light blinks GREEN / GREEN / GREEN in sequence.	
Stage control button on the left (see Fig. 2-6/ 28)	Yes	No	No	Short press *:	 Switches between XY stage control and Z axis control via focus drives If Z axis control is active: the indicator light lights in GREEN permanently the left or right fine focusing drive knobs are controlled for slow Z-movement (focusing) the left and right coarse focusing drive knobs are controlled for fast Z-movement (focusing) If XY stage control is active: the indicator light blinks GREEN the indicator light blinks GREEN the left focus drive knobs (fine or coarse) control Y movement (slow or fast) of the stage the right focus drive knobs (fine or coarse) controls X movement of (slow or fast) the stage 	
				Long press **:	 Records the parfocal position. 1. the LED shuts off for 300 ms (darkness indicates action to user) For halogen illumination: Do NOT shut off the light. 2. the indicator light blinks twice in GREEN 	
Right snap button + Stage				Press simultaneously	Load/unload	
control button on left (see Fig. 2-6/20+28)	Yes	No	No	Press simultaneously again	Load/unload	

Short press: hold less than 1 second, then release.
 Long press: hold at least 1.5 seconds.
 BLINK: the indicator light alternately turns on/off at 500 ms intervals

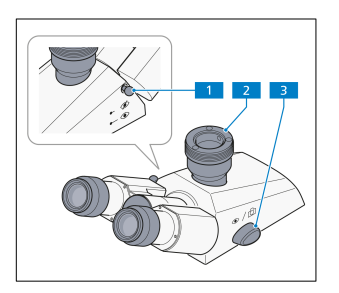


Fig. 2-8 Binocular photo tube

- Left push-pull rod (Fig. 2-8/1) pushed in: Eyepiece shutter is closed.
- Left push-pull rod (Fig. 2-8/1) pulled out: Eyepiece shutter is open.

2.5 Controls and functional elements on microscope components

2.5.1 Binocular photo tube 30°/23 (100:0/0:100)

The photo port (Fig. 2-8/**2**) of the binocular tube can be used to mount a reflex camera, a microscope camera or a video camera with the aid of appropriate adapters.

Using the shift knob (or the push-pull rod on the right side of other photo tubes), the light can be directed to the eyepieces or to the mounted camera.

- Shift knob (Fig. 2-8/3) in forward position (eye symbol):
 - 100% light to the eyepieces
- Shift knob (Fig. 2-8/3) in back position (camera symbol): 100% light to the camera
- Whenever you want to take photos with a long exposure time, we recommend that you block any ambient light from entering the eyepieces by using the eyepiece shutter or the eyepiece cover (included in the dust protection kit). If neither is available, remove the eyepiece and place the included dust protection caps over the eyepiece barrels.
- Vary eyepiece distance by moving the eyepiece barrels apart or together.

NOTE NOTE

For polarization microscopy, we recommend using the Pol photo tube with upright image and one eyepiece reticle (graticule).

2.5.2 Binocular ergo tube/ergo photo tube 20°/23 and ergo photo tube 15°/23, each with continuous vertical adjustment

These ergonomic tubes are designed for a field of view of 23 mm. The viewing angle is 20° or 15°.

The ergo photo tubes are equipped with one camera port each (Fig. 2-9/1 and Fig. 2-10/2).

The camera ports can be used to mount a reflex camera, a microscope camera or a video camera with the aid of appropriate adapters.

The camera port can carry a maximum weight of 2.5 kg (camera plus cable). For a fee, the ZEISS service personnel will equip your instrument with a higher maximum load, if needed.

The ergo tubes permit a height adjustment range of 44 mm.

A larger range can be obtained by swiveling the binocular section from the lower to the upper observation position (depending on the interpupillary distance).

- Continuous vertical adjustment by means of the rotary knobs (Fig. 2-9/3 and Fig. 2-10/1)
- The adjustment range of the ergo photo tube 20°/23 can be read off the vertical scale (Fig. 2-9/4).
- The binocular part of the ergo photo tube 15°/23 is also continuously horizontally retractable up to 50 mm.

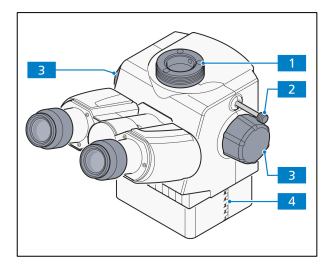


Fig. 2-9 Binocular ergo photo tube 20°/23 with vertical adjustment

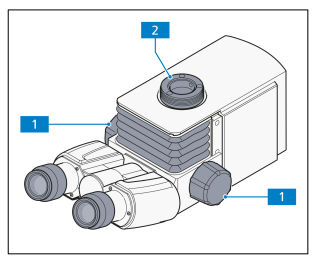
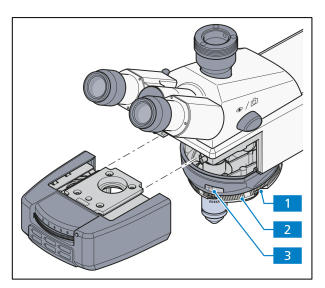


Fig. 2-10 Binocular ergo photo tube 15°/23, telescopic with vertical adjustment

The **ergo photo tube 20°/23** has two switch positions (scale: 100:0/0:100):

- Push-pull rod (Fig. 2-9/2) pushed in: 100% to the eyepieces
- Push-pull rod (Fig. 2-9/2) pulled out: 100% to camera port

The ergo photo tube 15°/23 (Fig. 2-10) is available only with an upright image and fixed scale (50:50).



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Fig. 2-11 Nosepiece with DIN slot for compensators

2.5.3 Nosepiece with objectives

- Equipped with DIN slot (Fig. 2-11/3) for compensators or fluorescence protection shield
- M27 mounting thread for six objectives
- Equipped with three, six or no DIC positions (Fig. 2-11/1) depending on the configuration
- Turning the knurled wheel of the nosepiece (Fig. 2-11/2) enables you to change the objectives quickly

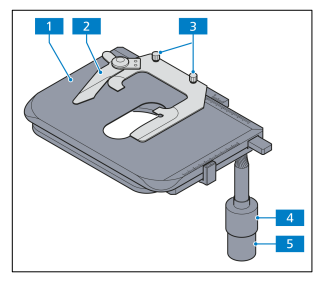


Fig. 2-12 Mechanical stage with specimen holder

2.5.4 Mechanical stage 75x50

- Mechanical stage (Fig. 2-12/1) with specimen holder (Fig. 2-12/2) for holding and positioning the samples
- Two clamping screws (Fig. 2-12/3) for attaching and removing the specimen holder
- Coaxial knurled knobs for X (Fig. 2-12/5) and Y adjustment (Fig. 2-12/4)

2.5.5 Low-power system for 2.5x/4x objectives mounted on the condenser carrier

The low-power system enables full illumination of the object field when using a low-magnification objective (2.5x-4x) in combination with the condenser 0.9/1.25 H.

It can be centered and remains in the beam path throughout the use of the corresponding objective.

- Swing the low-power system (Fig. 2-13/2) in or out of the beam path using the handle (Fig. 2-13/1). Make sure the module locks in securely.
- Do not use the low-power system with LED illumination for transmitted light.

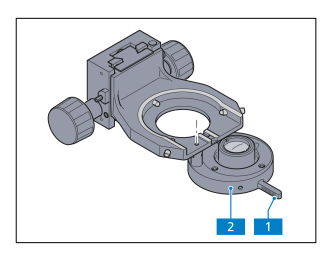


Fig. 2-13 Low-power system mounted on the condenser carrier

ATTENTION

If the condenser carrier with mounted low-power system is moved too far down, the low-power system may collide with the field diaphragm and damage it!

2.5.6 0.9/1.25 BF condenser and 0.9 Pol condenser

The 0.9/1.25 BF condenser (Fig. 2-14/**1**) with aperture diaphragm (Fig. 2-14/**2**) can be optionally equipped with a modulator disk (Fig. 2-14/**3**) for:

- bright-field BF
- dark-field DF
- phase contrast PhC 1, PhC 2, PhC 3
- PlasDIC (optional: To use PlasDIC, the ring diaphragm must be replaced with a 2 mm or 4.5 mm slit diaphragm for PlasDIC)

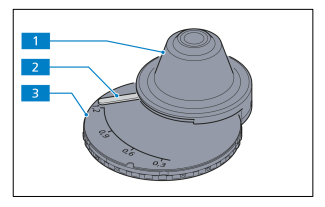


Fig. 2-14 0.9/1.25 BF condenser with modulator disk



ATTENTION

Do not clean the front optics of the condensers Pol with acetone.

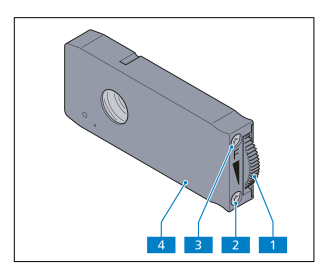


Fig. 2-15 14x40 mm stop slider

2.5.7 14x40 mm stop sliders for aperture and field diaphragms

To adjust the reflected light beam path using the KÖHLER method, one 14x40 mm stop slider (Fig. 2-15/4) mounted in the upper part of the stand is required to function as a field diaphragm (**F**) and another to function as the aperture diaphragm (**A**).

- To open the diaphragm, turn the knurled wheel (Fig. 2-15/1) up.
- To close the diaphragm, turn the knurled wheel (Fig. 2-15/1) down.
- To center the diaphragm to the beam path, adjust the centering screws (Fig. 2-15/2 and 3) with an Allen wrench (AF 3).

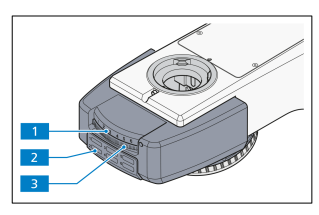
F	NOTE
	When applying fluorescent light, you may want to mount an FL attenuator (if not pre- installed) on the level of the aperture diaphragm in order to attenuate the stimulation intensity.

2.5.8 Reflector turrets with 4 or 6 coded positions

These reflector turrets are equipped with four or six coded push-and-click (P&C) reflector positions.

Turn the knurled wheel to adjust the reflector position. The marking indicates which reflector position is currently in the beam path.

For easy identification we advise that you write down the appropriate data for each position of the mounted reflector modules on the adhesive labels provided with the instrument. Then attach the labels at the appropriate points.

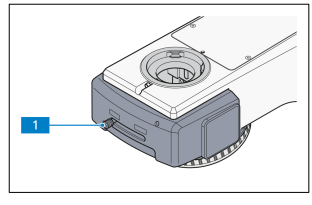


- 1 Display showing the number of the active reflector position
- 2 Spaces for labels with position data3 Knurled wheel with position marks

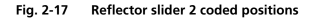
Fig. 2-16 Reflector turret with 6 coded positions

2.5.9 Reflector slider with 2 coded positions

The reflector slider with two coded positions is equipped with two individually loadable reflector positions for P&C modules which can be slid into the beam path.



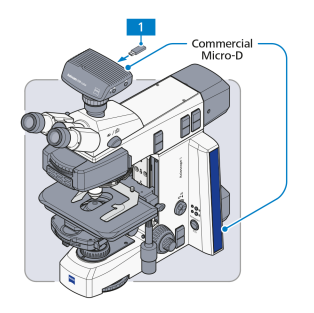
1 Slider



2.5.10 Microscope operating modes

2.5.10.1 Using the microscope imaging system works on as a standalone system

The microscope with Axiocam 208/202 can be used in standalone mode. The camera acts as the control interface and is powered by the microscope via a USB (Commercial Micro-D power) cable. A USB Type-C drive is included in the package and can be inserted in the USB slot on the back of the camera for storing data. Then images are recorded and saved on the USB drive. Functions of the microscope stand such as the Light Manager and encoding are automatically launched. The camera is equipped with image enhancement functions such as true color and noise reduction.



Functionality:

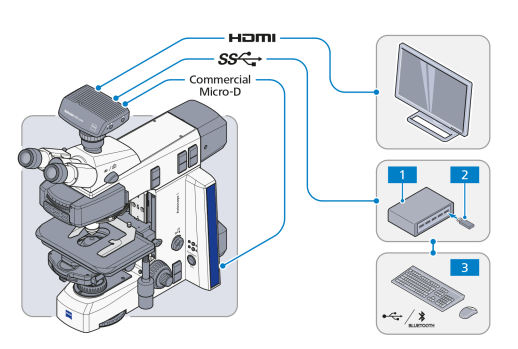
- Light Manager
- Coded components
- Image enhancement (true color, noise reduction)
- Record and save image on the USB drive
- Record and save video on the USB drive

1 USB Type-C drive is included in the package.

R ³	NOTE
	The snap button on the stand works only when a USB drive is detected by the camera port.

2.5.10.2 Connecting the microscope to an HD monitor, TV or projector via an HDMI cable

A monitor can be connected to the camera via an HDMI cable. The camera is powered by the microscope via a USB (Commercial Micro-D power) cable. A USB hub can be connected via the USB port on the camera. A wireless or wired mouse and keyboard can be connected to the camera via the USB hub, which together with the monitor, function as the control interface. Functions such as the Light Manager, encoding and image enhancement are automatically launched. Live images can be viewed on the monitor display and advanced features are available in the on-screen display (OSD). When the Axioscope is operated together with the Colibri 3 illuminator, the one-key fluorescence function can be used. Images can be snapped and saved on the USB Type-C drive, which is connected via the USB hub.

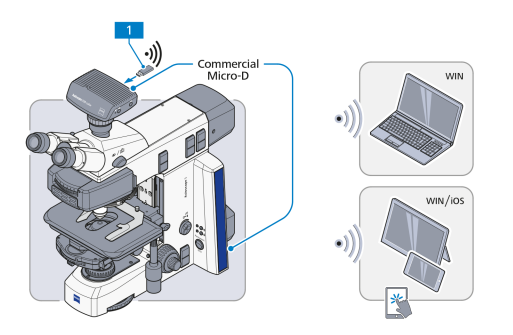


- Light Manager
- Coded components
- Image enhancement
- Observe live image on display
- Snap and save image on the USB drive
- Record and save video on the USB drive
- One-key fluorescence*
- Advanced features in OSD
- *: Works only when the Axioscope is used with Colibri 3

- **1** USB hub (input type C to output type A)
- 2 USB Type-C drive included in package
- 3 Mouse keyboard

2.5.10.3 Using the microscope imaging system with Labscope/Matscope via a Wi-Fi dongle connection

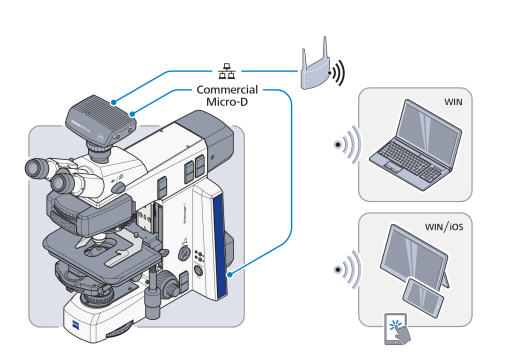
The camera is powered by the microscope via a USB (Commercial Micro-D power) cable. An optional monitor can be connected to the camera via an HDMI cable. The recommended USB Wi-Fi dongle can be connected to the camera via the USB hub. The control interface can be a PC or portable electronic devices that uses Wi-Fi. Functions such as the light manager, encoding, ECO mode and image enhancement are automatically launched. When a monitor is connected, live images can be viewed on the monitor display. Live images can also be viewed on a PC or portable devices and advanced features in Labscope/Matscope are available. The one-key fluorescence function can be used with the Axioscope in conjunction with the Colibri 3 illuminator.



- Light Manager
- Coded components
- ECO mode
- Image enhancement
- Observe live image
- Snap and save image via software
- One-key fluorescence*
- Advanced features in Labscope/ Matscope
- *: Only with specific microscopy configuration
- **1** USB Wi-Fi dongle (please see the ZEISS website for the recommended model)

2.5.10.4 Using the microscope imaging system with Labscope/Matscope via a WLAN router connection

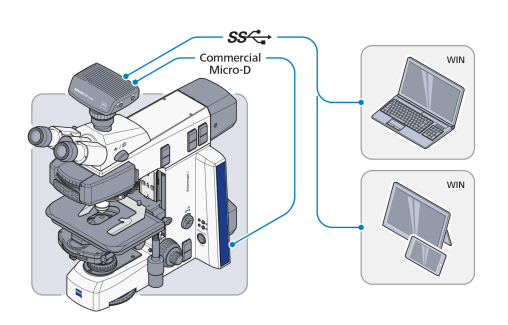
The camera is powered by the microscope via an USB (Commercial Micro-D power) cable. An optional monitor can be connected to the camera via an HDMI cable. A router is connected to the camera via Ethernet. The control interface can be a PC or portable electronic devise controlled via Ethernet or Wi-Fi. Functions such as the Light Manager, encoding, ECO mode and image enhancement are automatically launched. When a monitor is connected, live images can be viewed on the monitor display. Live images can also be viewed on PC or portable devices and advanced features in Labscope/Matscope are available. With Axioscope in connection with Colibri 3 illuminator, the one-key fluorescence function can be used.



- Light Manager
- Coded components
- ECO mode
- Image enhancement
- Observe live image
- Snap and save image via software
- One-key fluorescence*
- Advanced features in Labscope/ Matscope
- *: Only with specific microscopy configuration

2.5.10.5 Using the microscope imaging system with Labscope/Matscope via a USB connection

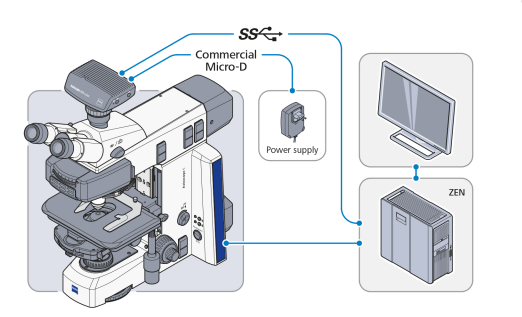
The camera is powered by the microscope via an USB (Commercial Micro-D power) cable. An optional monitor can be connected to the camera via an HDMI cable. A PC or Windows Surface can be connected to the camera via USB cable. Functions such as the Light Manager, encoding, ECO mode and image enhancement are automatically launched. When a monitor is connected, live images can be viewed on the monitor display. Live images can also be viewed on a PC or Surface computer and advanced features in Labscope/Matscope are available. With Axioscope in connection with Colibri 3 illuminator, the one-key fluorescence function can be used.



- Light Manager
- Coded components
- ECO mode
- Image enhancement
- Observe live image
- Snap and save image via software
- One-key fluorescence*
- Advanced features in Labscope/ Matscope
- *: Only with specific microscopy configuration

2.5.10.6 Using the microscope imaging system with ZEN software via a USB connection

The camera is powered via an Commercial Micro-D power cable connected to an external power socket. A workstation can be connected to the camera and the microscope stand via USB cables at the same time. Functions such as the Light Manager, encoding, ECO mode and image enhancement are automatically launched. With a monitor is connected, live images can be viewed on the monitor display and advanced features are available in OSD (on-screen display). Live images can be viewed on the workstation and basic features in ZEN are available.



- Light Manager
- Coded components
- ECO mode
- Image enhancement
- Observe live image
- Snap and save image via software
- Basic features in ZEN

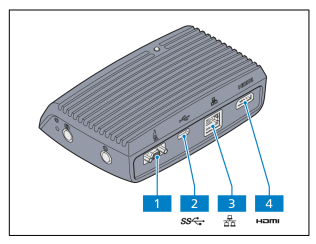


Fig. 2-18 Camera connector panel (rear side)

2.5.11 Axiocam 202 mono/208 color controls and connectors

The camera connection panel contains the following connectors (see Fig. 2-18):

- 1 port for power supply and communication to the microscope stand (via Commercial Micro-D cable)
- 2 port for camera control and image transfer (USB 3.0)
- **3** Gigabit Ethernet port (RJ45) for communication and image transfer
- 4 HDMI port for image data transfer to a monitor, TV or projector

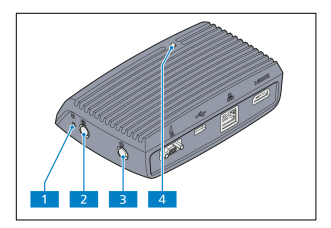


Fig. 2-19 Camera operator panel (right side)

The camera operator panel contains the following elements (see Fig. 2-19):

- 1 camera factory reset button
- 2 image / video capture button
- **3** OSD menu button
- 4 LED function indicator

NOTE NOTE

For more information about OSD see the Axiocam 202/208 User Guide.

2.5.12 OSD functionality with Axiocam 202 mono/208 color

The On Screen Display menu (OSD menu) is shown on whatever display the camera is connected to via an HDMI cable.

• Press the **OSD menu** button (Fig. 2-19/**4**) to open the menu (see Fig. 2-20).

R3	NOTE
	For more information about OSD functionality, see the Axiocam 202/208 User Guide.



Fig. 2-20 OSD menu, Home

3 START-UP

The Axioscope microscopes can be installed, modified and started up by the customer himself.

On request, the ZEISS Service installs or modifies the microscope for an extra charge.

R	NOTE
	Before installing and starting-up the microscope, read the Notes on instrument safety (see section 1.1) carefully and thoroughly.

The assembly activities described on the following section are illustrated for the most part by way of examples for one microscope stand type. However, they apply analogously to other microscope stand models. Special features are described separately.

The same activities apply when assembling a microscope stand with an upper stand part and the stand column Vario – special activities for the stand column Vario are described separately.

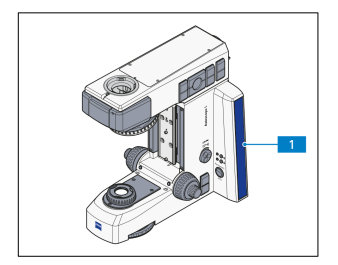


Fig. 3-1 Setting up the microscope

3.1 Mounting standard components

3.1.1 Unpacking and setting up microscope stand

- Remove all components from the packaging and check that all components described on the delivery note are present.
- Set up the microscope stand (Fig. 3-1/1) on a vibration-free, level, hard and non-combustible surface.
- Keep the original packaging for storage or for returning the instrument to the manufacturer, or dispose of it properly.

3.1.2 Mounting the Axioscope 5 Vario material upper stand part on the stand column

ATTENTION

Equipment or parts having a mass of 18 kg or more shall be provided with means for careful lifting and carrying.

If you are going to use the Axioscope 5 Vario material stand consisting of the Vario upper stand part (Fig. 3-2/2) combined with gear box (Fig. 3-2/3) and the Vario stand column (Fig. 3-2/1), proceed as follows:

- Unpack the Vario upper stand part with gear box and the Vario stand column.
- Position the Vario upper stand part (Fig. 3-2/2) on the mounting plate (Fig. 3-2/5) of the Vario stand column and tighten it with the 6 Allen screws (Fig. 3-2/4).
- In order to adjust the microscope vertically, loosen the release lever (Fig. 3-2/6) on the stand column and adjust the height (according to the size of the specimen) with the hand wheel (Fig. 3-2/7).
- Then tighten the release lever again.

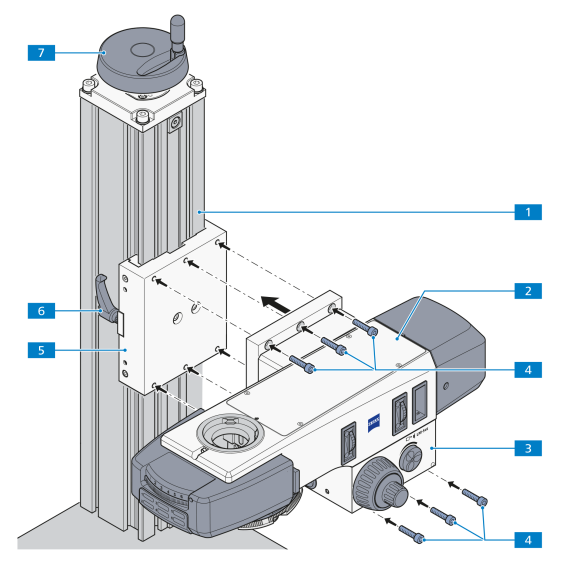
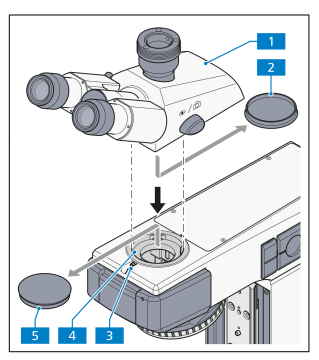


Fig. 3-2 Mounting the Vario upper stand part on the stand column



ZEISS

Fig. 3-3 Attaching the binocular tube

3.1.3 Mounting the binocular tube/photo tube

- Loosen the screws (Fig. 3-3/3) with the Allen wrench (AF 3).
- Remove the dust cover caps (Fig. 3-3/2 and 5) from the bottom of the tube and the dovetail ring socket on the stand.
- Hold the binocular tube (Fig. 3-3/1) or the binocular photo tube at an angle and insert it with the dovetail ring into the stand's dovetail ring socket (Fig. 3-3/4).
- Then push it into a horizontal position. Turn the binocular tube into the desired viewing position and tighten the Allen screws with the Allen wrench.

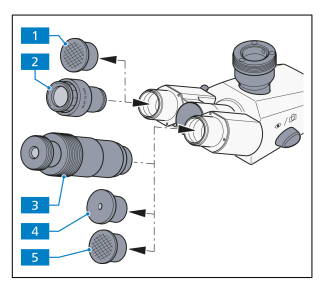


Fig. 3-4 Inserting eyepieces

3.1.4 Inserting eyepieces or auxiliary microscope or pinhole diaphragm

- Remove both dust cover caps (Fig. 3-4/1 and 5) from the binocular tube.
- Remove both eyepieces (Fig. 3-4/2) from the box and insert them into the binocular tube to the stop.
- Instead of an eyepiece you may insert an auxiliary microscope (Fig. 3-4/**3**) into one of the binocular eyepiece tubes in order to observe aperture, phase and darkfield diaphragms and to center phase and darkfield diaphragms. These diaphragms can be focused with the adjustable eye lens of the auxiliary microscope.
- The auxiliary microscope (Fig. 3-4/**3**) or the diopter (Fig. 3-4/**4**) can be used to observe conoscopic images.

The PL 10x/23 GW foc. Pol eyepiece contains a reticle firmly glued into it (cannot be changed), which is of defined orientation. When changing the interpu-pillary distance on the binocular photo tube, the two eyepiece tubes follow this rotary motion synchronously, so that the position of the orientation grooves in the eyepiece tubes remains unchanged.

• Insert the eyepiece into the binocular photo tube, fitting the locating screw (Fig. 3-5/2) into the tube's orientation groove (Fig. 3-5/1).

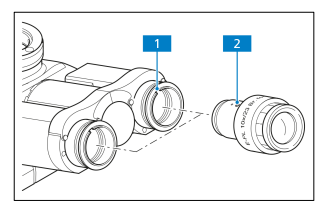


Fig. 3-5 Inserting the E-PL 10x/23 GW foc. Pol eyepieces into the binocular photo tube with upright image

The PL 10x/23 GW foc. Pol eyepiece (444038-9900-000) can be combined with a PL 10x/23 GW foc. eyepiece (444036-9000-000).

Inserting the eyepiece reticle

Eyepiece reticles (Fig. 3-6/**5**) can be used with eyepieces (Fig. 3-6/**3**) marked with a red dot (Fig. 3-6/**R**).

- Unscrew mounting stop (Fig. 3-6/6) from the eyepiece (Fig. 3-6/3).
- Pull locking ring (Fig. 3-6/4) out.
- Insert eyepiece reticle (Fig. 3-6/**5**) into the mounting stop and fix it with the locking ring.
- Screw mounting stop (Fig. 3-6/6) into the eyepiece (Fig. 3-6/3).

Make sure that the side of the reticle (Fig. 3-6/**5**) with the line pattern on it is facing the mounting stop in the tube (Fig. 3-6/**6**).

After insertion into the beam path of the microscope it is readable true to side.

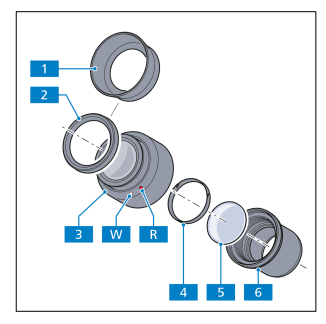


Fig. 3-6 Inserting the eyepiece reticle

The slight image shift caused by the additional path through glass is taken into account on the diopter scale by the fact that the zero point position is indicated not by the white dot (Fig. 3-6/W), but the red dot (Fig. 3-6/R).

NOTE NOTE

The eyepiece reticles must be inserted under dust-free conditions. This should be carried out only by ZEISS Service.

Inserting reversible eyecups

The eyepieces have rubber protection rings to avoid scratches on eyeglasses. These may be replaced by reversible eyecups as desired.

• To do this, remove the eyeglass protection rings (Fig. 3-6/2) from the eyepieces and mount the eyecups (Fig. 3-6/1).

Sometimes the eyeglass protection rings are seated very tightly in the eyepiece groove, so you may need a blunt object (wooden stick) to prod them off.

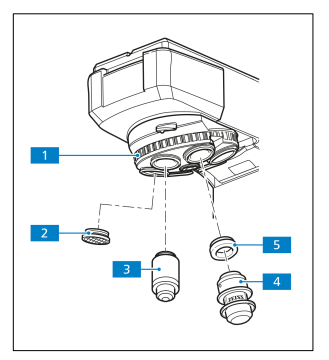


Fig. 3-7 Screwing in objectives

3.1.5 Screwing in objectives

- Lower the mechanical stage with the stage carrier to the bottom stop or lower the stage carrier with the dovetail guide.
- Remove the dust protection caps (Fig. 3-7/2) from the appropriate nosepiece openings.
- Take the objective (Fig. 3-7/3) out of its box and mount it on the nosepiece (Fig. 3-7/1), starting with the smallest magnification factor (set up clockwise).
- Instead of an objective, you may mount an object marker (Fig. 3-7/4) on any position of the nosepiece with the aid of a W0.8/M27 adapter (Fig. 3-7/5). Don't forget to recap the object marker when not in use to prevent it from drying out.

r S	NOTE
	Always place dust protection caps over the positions in the nosepiece which are currently not in use.
F	NOTE

The W0.8/M27 adapter must be used with the W0.8 objective.

3.1.6 Inserting and removing push-and-click (P&C) reflector modules into the reflector insert

To install or uninstall the modules, the reflector insert (reflector turret with 4 or 6 coded positions, reflector slider with 2 coded positions) must be removed from the upper part of the stand.

Inserting a module:

- Remove the reflector insert (Fig. 3-8/1) from the upper part of the stand (section 3.1.7) and put it aside with the top side facing down.
- Insert the module (Fig. 3-8/3) (with the top side facing down) with the aid of its mounting elements on its left and right (Fig. 3-8/4) at a slant from the top into the lower spring clips (Fig. 3-8/2) on the reflector socket.
- Press the module against the upper spring clips of the reflector turret until it engages firmly.

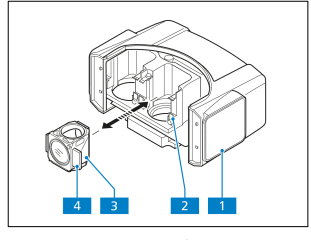


Fig. 3-8 Changing the reflector module in the reflector insert

Removing a module:

- Disengage the module from the upper spring elements by tilting it forward, then lift it off the lower spring elements and remove the module.
- After installing or uninstalling the reflector module, don't forget to replace the reflector insert or the cover cap.

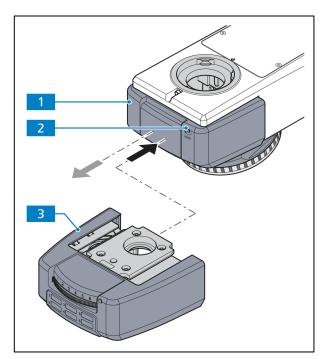


Fig. 3-9 Mounting the reflector insert



- First, remove the cover cap (Fig. 3-9/1) by inserting the Allen wrench (AF 3) into the slot (Fig. 3-9/2). Turn the locking screw counterclockwise and pull the cover cap forward.
- Push the reflector insert (Fig. 3-9/**3**) with the reflector modules P&C (e.g. reflector turret with 6 coded positions) into the upper part of the stand until it stops, hold it and tighten the locking screw again.

NOTE

When the reflector insert is removed, the reflected light path is automatically blocked so that the glare will not affect the operator.

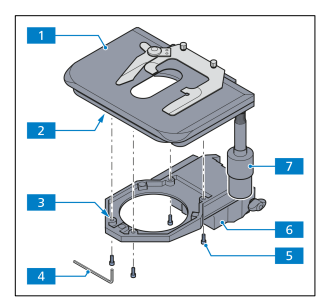


Fig. 3-10 Changing the mechanical stage, 75x50 R

3.1.8 Mounting the mechanical stage

Depending on the type, the mechanical stages can either be stable or they can be rotated and centered. The shift range is 75 mm in the Xdirection and 50 mm in the Y-direction.

Stages are available with a coaxial gear knob (Fig. 3-10/**7**) located on the right or the left side of the stage.

To assemble/disassemble a stage, remove the stage carrier, then reattach the stage carrier with the mounted stage to the stand (see section 3.1.15 on page 67).

(1) Stable mechanical stage, 75x50 R

a) How to remove the stage

- Remove the four mounting screws (Fig. 3-10/5) on the stage carrier (Fig. 3-10/6) using the Allen wrench (AF 3) (Fig. 3-10/4).
- Remove the stage (Fig. 3-10/1) from the stage carrier. This is easier if you turn the stage and its carrier over.

b) How to mount the stage

- Fit the stage (Fig. 3-10/1) onto the stage carrier (Fig. 3-10/6) so that the threaded holes in the bottom of the stage (Fig. 3-10/2) are on top of the corresponding holes in the stage carrier (Fig. 3-10/3).
- Insert the four mounting screws (Fig. 3-10/5) from below through the holes in the stage carrier and screw them into the bottom of the stage.
- Tighten the screws.

F	NOTE
	After long hours of use, the stage guide may be loose and the stage travel range may be decreased. However, this is not a malfunction and can be corrected easily as described below.

c) Recovering the stage travel range in X-axis

After long hours of use, the X direction range will gradually become smaller. This is not a quality issue and can be easily reset.

• Hold the specimen holder's two screws and move it to the left and right until hits the end stops, the reset is complete (Fig. 3-11/1, 2).

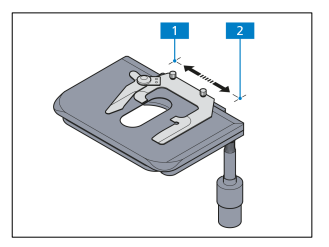


Fig. 3-11 Recovering the stage travel range in X-axis

(2) Rotatable mechanical stage 75x50/240° R

NOTE

Rotatable stages can be used only with the stage carrier for rotatable stages (430710-9011-000).

a) How to remove the stage

- Loosen the screw cap (Fig. 3-12/2) of the spring pin with approx. 3 turns.
- Press the stage forward against the spring pin (Fig. 3-12/1), lift the back side off the stage carrier (Fig. 3-12/3) and remove it by lifting it upward.
- Tighten the screw cap (Fig. 3-12/2).

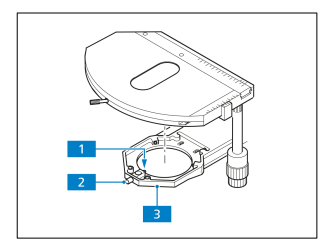


Fig. 3-12 Changing the rotatable mechanical stage

b) How to mount the stage

- Loosen the screw caps (Fig. 3-12/2) of the spring box with approx. 3 turns.
- Put the stage with the dovetail ring notch onto the spring pin (Fig. 3-12/1).
- Press the stage forward against the spring pin and lower its back side into the stage carrier (Fig. 3-12/3).
- Tighten the screw caps (Fig. 3-12/2).

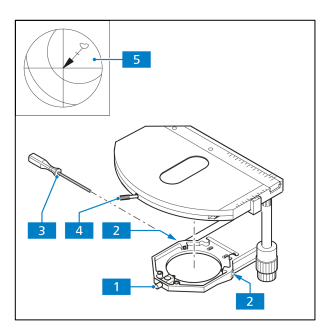


Fig. 3-13 Centering the rotatable mechanical stage

c) How to center the stage

When using objectives with a high magnification, only one selected objective can be centered exactly at a time.

All stages are delivered pre-centered, i.e. when the stage is turned, the detail of the specimen in the focus remains in the center of the image. If the detail shifts out of the center when the stage is turned (Fig. 3-13/5), then it becomes necessary to re-center the stage.

- Loosen the clamping screws of the stage (Fig. 3-13/4) and the screw cap of the stage carriers (Fig. 3-13/1).
- Determine the maximal shift of the specimen (Fig. 3-13/**5**, arrow point) to the cross line in the eyepiece by turning the stage.
- Shift the specimen detail towards the center of the cross line by turning each the two centering screws on the stage carrier (Fig. 3-13/2) with an Allen wrench (AF 1.5) (Fig. 3-13/3) half the length of the arrow. Repeat the procedure if the specimen detail shifts out of the center again when turning the stage.
- After finishing the centering procedure tighten the screw caps (Fig. 3-13/1) again.

The stage can be turned by 240° within a shifting range of $y \le 27$ mm. No turning of the stage is possible outside of this range.

(3) Adjusting the drive length on the ergonomic drives

An axial shift of the coaxial knurled knobs by a **maximum of 15 mm** can extend the drive length of the **X** and **Y** shift on the mechanical stages with ergonomic drives.

(4) Removing and mounting the additional sleeves

Both coaxial knurled knobs for the stages are equipped with additional sleeves for an even more sensitive adjustment of the specimen position. These sleeves can be removed in those cases when it is important to shift the specimen more quickly.

- Loosen both clamping screws (Fig. 3-14/4) on the lower additional sleeve (Fig. 3-14/3) and remove them by pulling them down. Then loosen both clamping screws (Fig. 3-14/2) on the upper additional sleeves (Fig. 3-14/1) and remove them by pulling them down as well.
- Re-mount the additional sleeves on the coaxial knurled knobs by following the above steps in reverse order. Tighten the clamping screws when you are done.

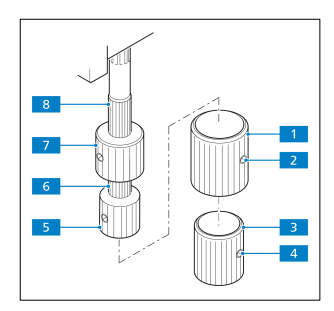


Fig. 3-14 Adjusting ergonometric drive

(5) Adjusting the friction (smoothness of operation) on both coaxial knurled knobs of the ergonomic drives

The smoothness of ergonomic operation is factory pre-set to a medium level. The operator can change the friction as follows:

a) How to adjust the X-direction

- Remove the additional sleeves (Fig. 3-14/1 and 3) from the coaxial knurled knobs by loosening the clamping screws.
- Shift the X-coaxial knurled knob (Fig. 3-14/5) downward and the Y-coaxial knurled knob (Fig. 3-14/7) upward.
- Hold the X-coaxial knurled knob (Fig. 3-14/5) and turn the light-colored knurled ring above it (Fig. 3-14/6) to the right (increased smoothness) or left (decreased smoothness) until you reach the desired level.

b) How to adjust the Y-direction

- Hold the Y-coaxial knurled knob (Fig. 3-14/7) and turn the light-colored knurled sleeve above it (Fig. 3-14/8) to the right (increased smoothness) or the left (decreased smoothness) until you reach the desired level.
- Replace the additional sleeves and tighten the clamping screws.

R	NOTE
•	You can increase the service life of the stage by cleaning the specimen holder regularly and avoiding particulates from the mechanical guide system of the X-motion.

3.1.9 Mechanical stages with friction adjustment

The drive length and friction for the X- and Y-adjustment can be adjusted individually on the standard mechanical stage (432035-9002-000) and the mechanical stage 432035-9070-000.

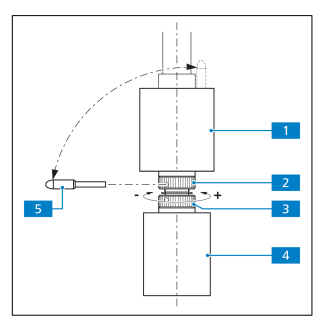


Fig. 3-15 Adjusting the frictional momentum

(1) Adjusting the drive length on the stage drive

The drive length of the X- and Y-drive can be adjusted by an axial shift of the coaxial knurled knob (Fig. 3-15/**4** and **1**) within a range of approx. 15 mm.

(2) Adjusting the frictional torque of the coaxial knurled knobs for the X-/Y-adjustment of the mechanical stage

The frictional torque of the coaxial knurled knobs is factory-preset to a medium level which can be changed as follows if necessary:

a) X-drive

- Slide the coaxial knurled knob for the X-adjustment (Fig. 3-15/**4**) all the way down.
- Take the enclosed adjustment pin (Fig. 3-15/5) out of the coaxial knurled knob for the Y-adjustment (Fig. 3-15/1) and stick it into one of the holes on the lower hole nut (Fig. 3-15/3).
- Hold the coaxial knurled knob for the X-adjustment (Fig. 3-15/4) and turn the hole nut with the adjustment pin clockwise (lower frictional torque: –) or counterclockwise (higher frictional torque: +) until you reach the desired level of friction (see Fig. 3-15).
- The adjustment should not be more than **one** rotation.

b) Y-drive

- Slide the coaxial knurled knob for the Y-adjustment (Fig. 3-15/1) all the way up.
- Stick the enclosed adjustment pin (Fig. 3-15/5) into the hole on the upper hole nut (Fig. 3-15/2).
- Hold the coaxial knurled knob for the Y-adjustment (Fig. 3-15/1) and turn the hole nut with the adjustment pin clockwise (lower frictional torque: –) or counterclockwise (higher frictional torque: +) until you reach the desired level of friction.
- The adjustment should not be more than **one** rotation.
- Replace the adjustment pin in the coaxial knurled knob for the Y-adjustment (Fig. 3-15/1).

3.1.10 Attaching the Pol rotary stage

Two rotary stage models are available. The mounting procedure is the same for each model.

3.1.10.1 Removing the Pol rotary stage

- Loosen the screw cap (Fig. 3-16/8) of spring pin with about 3 turns.
- Push the Pol rotary stage (Fig. 3-16/5) to the front against the spring-loaded pin (Fig. 3-16/9), lift it off the stage carrier (Fig. 3-16/7) at the back and remove it upwards.
- Retighten the screw cap (Fig. 3-16/8).

3.1.10.2 Attaching the Pol rotary stage

- If necessary, loosen the screw cap (Fig. 3-16/8) of the spring pin with about 3 turns.
- Place the Pol rotary stage with the groove of the dovetail (stage bottom) on the spring-loaded pin (Fig. 3-16/9).

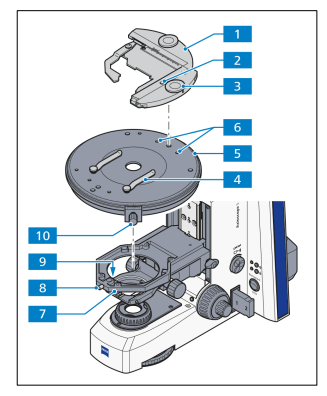


Fig. 3-16 Changing the Pol rotary stage with click stop, attachable Pol object guide and stage clips

- Attach the rotary stage (432035-9191-000) with the clamping screw (Fig. 3-16/10) pointing to the front right.
- Push the Pol rotary stage to the front against the spring-loaded pin and lower it at the back into the stage carrier (Fig. 3-16/7), then release it.
- Retighten the screw cap (Fig. 3-16/8).

F	NOTE
	When you are using the rotary stage (432035-9131-000), the click stop must point to the front.

3.1.10.3 Removing the attachable Pol object guide and mounting stage clips

- Turn the control wheel (Fig. 3-16/3) of the Pol object guide (Fig. 3-16/1) until the clamping screw becomes visible in the mounting hole (Fig. 3-16/2).
- Use the Allen wrench to loosen the clamping screw. Remove the Pol object guide Pol upwards.
- Insert the stage clips (Fig. 3-16/4) into the holes on the stage provided for this purpose.
- Attach the Pol object guide (453560-0000-000) for the large rotary stage (453550-0000-000) in the same way.
- Attach the D Pol specimen holder for transmitted light specimens or the A Pol specimen holder for reflected light specimens and mount the balance weight on the stage.

3.1.10.4 Removing the stage clips and attaching the Pol attachable object guide

- Remove stage clips (Fig. 3-16/4) from rotary stage Pol.
- Insert the Pol object guide (Fig. 3-16/1), introducing the two cylindrical pins on its underside into the respective holes (Fig. 3-16/6) and tighten the clamping screw firmly (Fig. 3-16/2). Where required, turn the control first to make the clamping screw accessible.

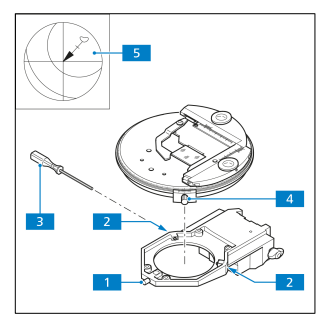


Fig. 3-17 Centering the Pol rotary stage

3.1.10.5 Centering the Pol rotary stage

All stages are factory-pre-centered, i.e. when rotating the stage the specimen feature set to the center of the field of view will remain in the center. If the specimen feature moves off the center of the field of view (Fig. 3-17/**5**) while you are rotating the stage, the stage should be re-centered as follows.

- Before centering the stage, you must adjust the KÖHLER illumination on the microscope (see section 4.2.1).
- For stage centering, use a contrast-rich specimen and an eyepiece with reticle.
- Loosen the stage click-stop (Fig. 3-17/4) or clamp (on the Pol rotary stage with clamping device) and unscrew the stage carrier cap (Fig. 3-17/1).
- Rotate the stage to determine the position of maximum offset of the specimen feature (Fig. 3-17/**5**, origin of arrow) from the center of the eyepiece reticle.
- Turn the two centering screws on the stage carrier (Fig. 3-17/2) with one Allen wrench (AF 1.5) (Fig. 3-17/3) each to move the specimen feature by half the arrow length towards the reticle center. Rotate the stage again to check if the specimen feature moves off. Repeat the centering procedure, if necessary.
- After centering, retighten the screw cap (Fig. 3-17/1).

3.1.10.6 Centering objectives of polarizing microscope stands

The nosepiece with 5-positions HF/DF/Pol and 1-position HF/DF/DIC is equipped with five centerable objective mounts (without DIC slots) as well as one objective mount with DIC slot (noncenterable). Accordingly, all objectives can be centered relative to the rotary stage.

Stage centering is necessary to ensure that a specimen feature located in the center of the field of view does not move out while you are rotating the stage. Centering all objectives ensures that the specimen feature remains in the center of the field of view even when the objective is changed.

- Before centering the stage, you must adjust the KÖHLER illumination on the microscope (see section 4.2.1).
- For stage centering, use a contrast-rich specimen and an eyepiece with reticle.

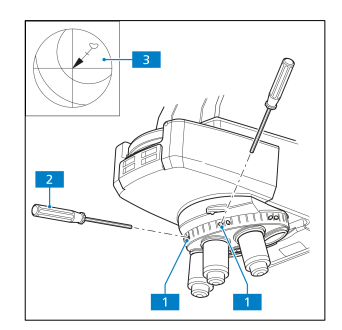


Fig. 3-18 Centering objectives

- First, turn the nosepiece to swing the non-centerable objective mount (mount with DIC slot) into the light path. Center the rotary stage for the non-centerable objective mount as described under 3.1.10.5.
- Turn the nosepiece to move a centerable objective mount into the light path.
- Rotate the stage to determine the position of maximum offset of the specimen feature (Fig. 3-18/3, origin of arrow) from the center of the eyepiece reticle.
- Turn the two centering screws on the nosepiece (Fig. 3-18/1) using one AF 1.5 Allen wrench (Fig. 3-18/2) each to move the specimen feature by half the arrow length towards the reticle center. Rotate the stage again to check whether the specimen feature moves off. Repeat the centering procedure, if necessary.
- Center the other four objectives in the same manner.

₽	NOTE
	To preserve this centered state it is imperative to change the objectives only by turning the nosepiece by its knurled ring, without touching the objectives.

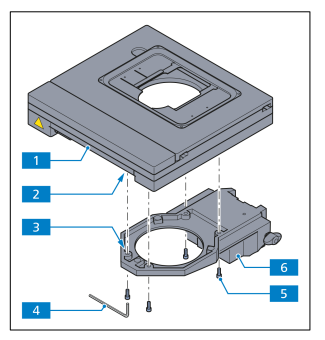


Fig. 3-19 Changing the motorized mechanical stage

3.1.11 Mounting/removing the 80x60 motorized mechanical stage on the Axioscope 7 motorized material stand

The motorized stage can only be mounted on/removed from the stand as an entire unit with the stage carrier (see section 3.1.15 on page 67).

How to remove the stage

- Unplug the power cable connector (Fig. 3-20/1) of the stage from the socket on the stand (Fig. 3-20/2).
- Turn the stage (Fig. 3-19/1) and stage carrier (Fig. 3-19/6) upside down and remove the four mounting screws (Fig. 3-19/5) with the Allen wrench (AF 3) (Fig. 3-19/4).
- Carefully remove the stage from the carrier.

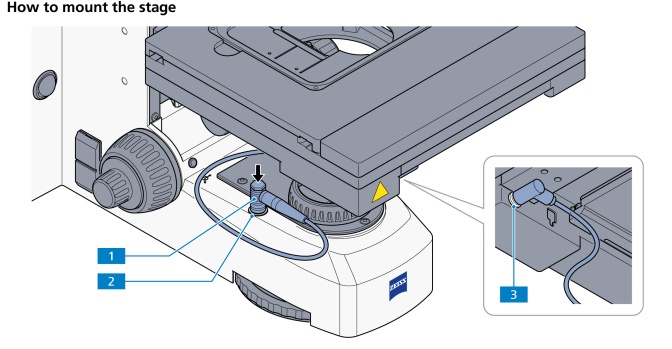


Fig. 3-20 Plugging in the power cable of the motorized mechanical stage

- Turn stage (Fig. 3-19/1) and stage carrier (Fig. 3-19/6) upside down carefully and match the threaded holes in the bottom of the stage (Fig. 3-19/2) to the corresponding holes in the carrier (Fig. 3-19/3).
- Insert the four mounting screws (Fig. 3-19/5) into the holes.
- Align the stage in XY-direction and tighten the screws with the Allen wrench (AF 3).
- Plug the power cable connectors (Fig. 3-20/1) into the sockets on the stage (Fig. 3-20/3) and on the stand (Fig. 3-20/2).

3.1.12 Mounting the condenser carrier

- Adjust the guide of the condenser carrier (Fig. 3-21/**3**) using the coaxial knurled knob (Fig. 3-21/**2**) until both screws (Fig. 3-21/**4**) become accessible.
- Mount the condenser carrier on the stage carrier (Fig. 3-21/1), slide it firmly and straight up to the upper stop and tighten both screws.

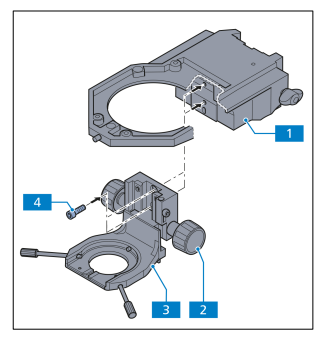


Fig. 3-21 Mounting the condenser carrier

3.1.13 Mounting the condenser

- Slide the stage carrier with the focusing drive to the upper stop.
- Unscrew both centering screws (Fig. 3-22/**5**) on the condenser carrier until you cannot see the end parts anymore.
- Unscrew the clamping screw (Fig. 3-22/9) on the right side of the condenser carrier until the maximum vertical adjusting range is usable.
- Slide the condenser carrier (Fig. 3-22/3) with the knurled knob for vertical adjustment (Fig. 3-22/2) all the way down.
- Open the front lens (if switchable) on the condenser using the lever (Fig. 3-22/7).
- Insert the condenser (Fig. 3-22/8) between the condenser carrier (Fig. 3-22/3) and the stage carrier (Fig. 3-22/1). Adjust the guide pin on the bottom of the condenser in the direction of the slot (Fig. 3-22/6).

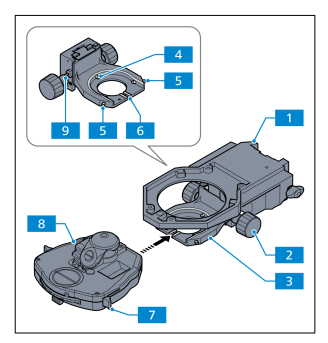


Fig. 3-22 Mounting the condenser

- Push the condenser with the dovetail ring against the spring box (Fig. 3-22/4) of the condenser carrier until the condenser can be placed on the condenser carrier horizontally.
- Release the condenser and its pin screw will slide into the front slot (Fig. 3-22/6).
- Tighten the centering screws until they engage in the condenser's dovetail ring.
- Screw in the clamping screw (Fig. 3-22/9) without clamping the vertical drive.

F	NOTE
	Follow the same procedure when mounting other condensers.
L F	NOTE

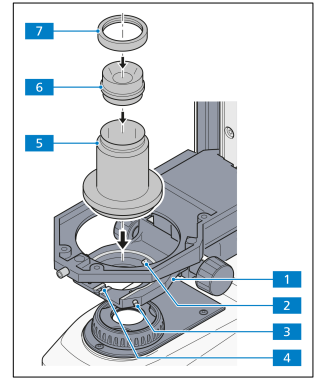


Fig. 3-23 Mounting the darkfield condenser

3.1.14 Mounting the darkfield condenser

NOTE NOTE

The condenser holder Z for darkfield is required to place darkfield condensers in the condenser carrier.

- Insert the darkfield condenser (Fig. 3-23/6) in the condenser holder Z (Fig. 3-23/5) and screw on the fastening ring (Fig. 3-23/7).
- Push the condenser holder Z (Fig. 3-23/5) against the spring box (Fig. 3-23/2) into the condenser carrier (Fig. 3-23/1).
- Tighten both centering screws (Fig. 3-23/**3**) on the condenser carrier (Fig. 3-23/**1**) until they grip the dovetail ring of the condenser holder Z (Fig. 3-23/**5**).

3.1.15 Mounting the stage carrier

- Screw the shoulder screw (Fig. 3-24/**3**) into the appropriate opening:
 - Opening with mark 0 (Fig. 3-24/3):
 No sample space extension is mounted.
 - Opening with mark **30** (Fig. 3-24/**2**): The sample space extension for **30 mm** is mounted.
 - Opening with mark 60 (Fig. 3-24/1): The sample space extension for 60 mm (430030-9111-000) is mounted.
- Insert the stage carrier (Fig. 3-24/**5**) at a slight angle (beneath the shoulder screw) from the left into the guide, then push it in straight and tighten the wing screw (Fig. 3-24/**4**) slightly.
- Push the stage carrier along the guide upward until it engages in the shoulder screw. Tighten the wing screw.
- Check to ensure that the guide is accurately positioned.

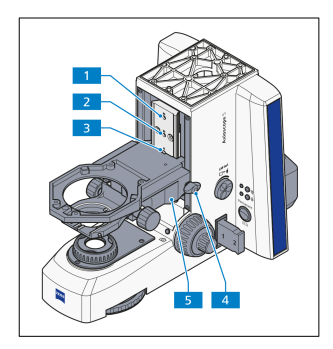
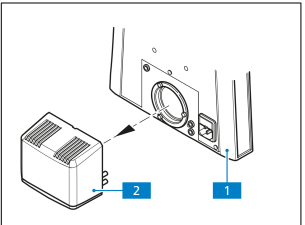


Fig. 3-24 Mounting the stage carrier



Removing the HAL 50 halogen Fig. 3-25 illuminator from the stand

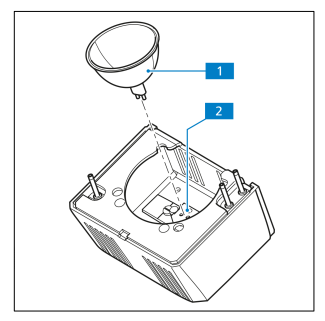


Fig. 3-26 Inserting halogen lamp 12 V, 50 W

3.1.16 Changing the 12 V, 50 W halogen lamp of the HAL 50 halogen illuminator

- Switch off the microscope and disconnect it from the power supply. Wait at least 15 minutes for the halogen lamp to cool down.
- Remove the HAL 50 halogen illuminator (Fig. 3-25/2) from the back of the stand (Fig. 3-25/1) and put it down with the open side facing up.
- Remove the used 12 V, 50 W halogen lamp (if necessary) by pulling it out of the cap (Fig. 3-26/2) in an upward motion.
- Push the new lamp (Fig. 3-26/1) with its two • cap pins gently and carefully into the lamp cap (Fig. 3-26/2). Be careful not to touch the protective glass of the lamp. To avoid bending its cap pins or damaging the lamp holder, the lamp must not be tilted.
- Place the HAL 50 halogen illuminator with the connecting pins into the back side of the microscope and push until the lamp engages securely.
- Reconnect the microscope to the power supply.

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3.1.17 Mounting the illuminator TL LED10/RL LED10

- Using the Allen wrench (AF 3) (Fig. 3-27/7), loosen the clamping screw of the TL illumination connector (Fig. 3-27/6) or of the RL illumination connector (Fig. 3-27/2) at the back of the stand.
- Connect the illuminator's power plug (Fig. 3-27/4) to the socket (Fig. 3-27/3) on the lower part of the stand (or the upper part of the stand).
- Match the illuminator's (Fig. 3-33/5) dovetail ring to the connection piece on the stand. Ensure that the illumination cable is not being pinched or clamped.
- Tighten the clamping screw (Fig. 3-33/6 or 2) with the Allen wrench (AF 3).

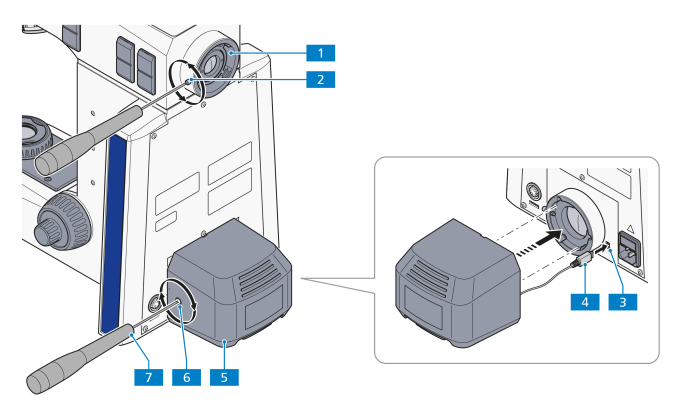


Fig. 3-27 Mounting the TL LED10 illuminator or RL LED10 illuminator

3.1.18 Mounting the Axioscope base plate on the stand

L	NOTE
	When using the HAL or HBO illuminators, a base plate for Axioscope (000000-2202-526) is mandatory.

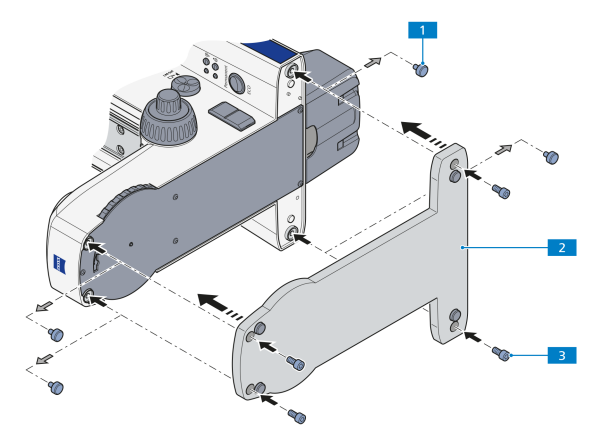


Fig. 3-28 Mounting the base plate

- Remove the four rubber feet (Fig. 3-28/1) from the bottom of the stand.
- Align the base plate (Fig. 3-28/**2**) with the stand and insert a hexagonal socket screw (M6) (Fig. 3-28/**3**) into each of the four holes in the base plate.
- Tighten the screws (M6) with an Allen wrench to fix the base plate to the stand.

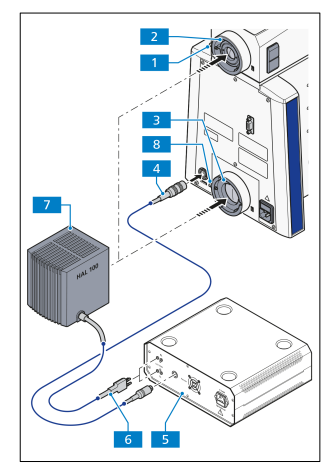
3.1.19 Mounting and adjusting the HAL 100 halogen illuminator

The HAL 100 halogen illuminator may serve as a light source for transmitted light and reflected light procedures (except fluorescence light) as long as either the lower or upper part of the stand is equipped with the appropriate connection piece. When mounting the halogen illuminator, no difference is made between reflected light and transmitted light procedures.

3.1.19.1 Mounting the HAL 100

ATTENTION Before switching on the halogen illuminator, remove the tools (Fig. 3-31/7) included in the illuminator case to prevent them from being damaged by heat.

 With the Allen wrench (AF 3) loosen the clamping screw of the respective illumination connector (lower for TL (Fig. 3-29/8), upper for RL (Fig. 3-29/1)) on the back of the stand.





• Insert the lamp case (Fig. 3-29/7) with the dovetail ring into the connector (Fig. 3-29/2 or Fig. 3-29/3). Now fixate it by tightening the clamping screw (Fig. 3-29/1 or Fig. 3-29/8) with the Allen wrench.

F	NOTE
	Due to the lack of ground clearance it is important to avoid tilting the illuminator when inserting it into the lower stand part. Check for straight and correct fit.

- Connect the 3-pin lamp plug (Fig. 3-29/6) to the respective socket on the back of the external power supply unit (suited for two lamps) (Fig. 3-29/5): plug the TL illuminator into the lower TL socket, plug the RL illuminator into the upper RL socket.
- Connect the cable (Fig. 3-29/4) for illumination intensity control (see section 3.3.2).

F	NOTE
	If this cable is connected to the stand, the illumination intensity of the HAL 100 in use can be varied by rotating the Intensity/LM knob (Fig. 2-4/ 10).
	If this cable is not connected to the stand, the illumination intensity of the HAL 100 in use can be varied by rotating the adjusting wheel of the external power supply unit.
	To switch between the illuminators for reflected and for transmitted light, flip the toggle switch of the power supply unit to TL for transmitted light or to RL for reflected light (see section 3.3).

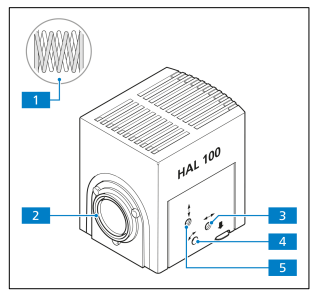


Fig. 3-30 Adjusting the HAL 100

3.1.19.2 Adjusting the HAL 100

Coarse setting

- After loosening the clamping screw (Fig. 3-29/1 or Fig. 3-29/8), remove the ready-to-operate halogen illuminator (Fig. 3-30/2) from the microscope stand.
- Switch on the external power supply unit of the HAL 100 (see section 3.3.3).
- Direct the light beam towards a projecting surface (wall) which is at least 3 m away.



Never look directly into the lightemitting aperture of the illuminator!

- Adjust the adjustment screw (Fig. 3-30/**3**) using an Allen wrench (AF 3) until both images of the lamp filament appear as sharp as possible on the projection surface.
- Adjust the adjustment screws (Fig. 3-30/4 and 5) so that the lamp filament of one image exactly fills the gaps of the reflected image (Fig. 3-30/1).

Fine setting

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- Mount the illuminator on the microscope stand again and tighten the clamping screw to lock it.
- If applicable, remove the filter from the beam path.
- Focus the objective ≤ 40x on the flat specular surface of a sample and go to a free area of the specimen.
- Remove the eyepiece and center the lamp filament and its reflection in the eye pupil image with the adjustment screws (Fig. 3-30/4 and 5).
- Optimize the even illumination of the pupil image with the adjustment screws (Fig. 3-30/3).
- If applicable, replace the filter in the beam path.

3.1.19.3 Changing the halogen lamp 12 V, 100 W

	CAUTION		
	Hot surface! Danger of burns!		
	Always let the illuminator case cool off for at least 15 min before touching it!		
r) B	NOTE		
	It is not necessary to remove the illuminator case from the stand when		

• Switch off the external power supply unit of the HAL 100 (see section 3.3.3). Disconnect the 3-pin plug from the HAL 100.

replacing the halogen lamp.

- Press the release button (Fig. 3-31/3) of the halogen illuminator HAL 100 (Fig. 3-31/1). Pull the lamp carrier (Fig. 3-31/2) out completely and put it aside.
- Press both spring levers downwards (Fig. 3-31/**5**) and pull out the old halogen lamp (Fig. 3-31/**6**) in an upward motion.
- Press both spring levers downwards, insert the new lamp into the lamp cap (Fig. 3-31/4) and release the spring levers. Use only the included special tools (Fig. 3-31/7) when holding/ grabbing the halogen lamp. Traces of skin grease on the halogen lamp can impair its life span.

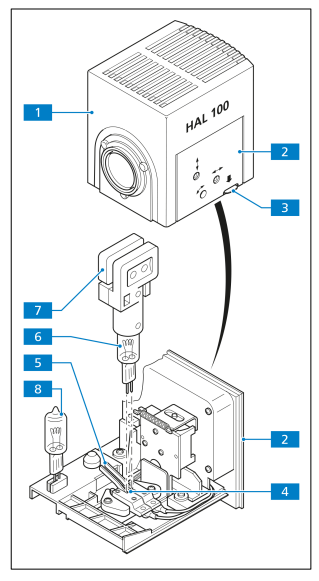


Fig. 3-31 Changing the HAL 100

- Press the spring levers down again briefly. Follow the adjustment steps described in section 3.1.19.2.
- Replace the lamp carrier and slide it in until it engages fully.
- Re-connect the 3-pin plug of HAL 100.



ATTENTION

Before switching on the halogen illuminator, remove the tools (Fig. 3-31/**7**) included in the illuminator case to prevent them from being damaged by heat. The replacement lamp (Fig. 3-31/**8**) may remain in the illuminator case.

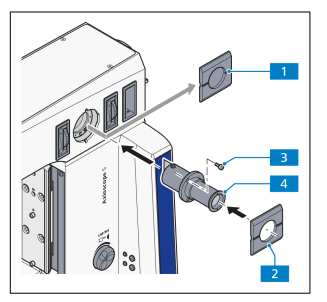


Fig. 3-32 Inserting the adjustment tool

3.1.20 Inserting the adjustment tool of HBO 100 into the stand Axioscope 5 TL/RL

- Remove the cover (Fig. 3-32/1) from the mounting aperture of the adjustment tool.
- Insert the adjustment tool (Fig. 3-32/4) and tighten the three included screws (Fig. 3-32/3).
- Attach the cap (Fig. 3-32/**2**) to the opening. Make sure it locks.
- Slide in the movable connecting piece of the adjustment tool.

The adjustment tool is used to adjust the HBO 100 illuminator; see section 3.1.21.3.

3.1.21 Mounting and adjusting the HBO 100 illuminator

3.1.21.1 Inserting the HBO 103 W/2 mercury vapor short arc lamp

For safety reasons, the HBO 100 illuminator and the HBO 103 W/2 mercury vapor short arc lamp are packed separately.

L	NOTE
	Please read the manual included with the unit before inserting or replacing the HBO 103 W/2 lamp.

• Insert the HBO 103 W/2 lamp into the illuminator as described in the accompanying manual.

	ATTENTION
<u>!</u>	Please use an FL attenuator (if it is not installed continuously) to modify the light intensity. Gray filters are not resistant enough on the long run.

NOTE

3.1.21.2 Mounting the HBO 100 illuminator

- Remove the cover from the reflected light connection piece (Fig. 3-33/1).
- Insert the illuminator case (Fig. 3-33/3) with the dovetail ring in the reflected light connection piece (Fig. 3-33/1) on the back of the unit. Tighten the clamping screw (Fig. 3-33/2) with the Allen wrench (AF 3).
- Connect the HBO 100 lamp cable to the power supply unit (Fig. 3-34/1). Secure with retaining ring.
- First, plug the cord into the outlet (Fig. 3-34/**2**) of the power supply unit, then insert the power plug into the electrical outlet.

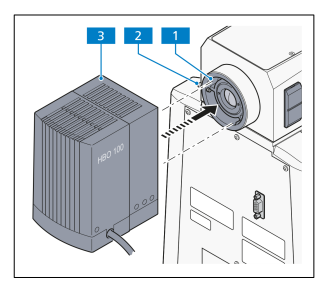


Fig. 3-33 Mounting the HBO 100

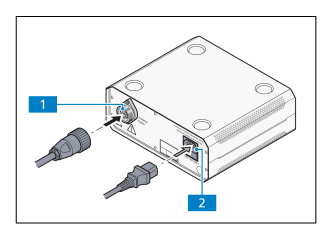


Fig. 3-34 Power supply unit for HBO 100

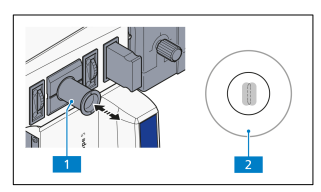


Fig. 3-35 Adjustment tool

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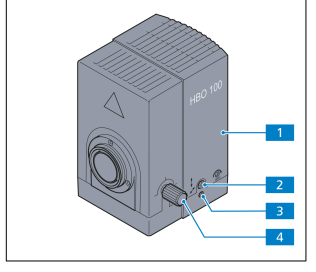


Fig. 3-36 Adjusting the HBO 100

3.1.21.3 Adjusting the HBO 100 illuminator

The HBO 100 illuminator is available in two versions (manual and automatic adjustment).

The self-adjusting HBO 100 (423011-9901-000) adjusts automatically when the power supply unit is turned on.

Follow the instructions below to adjust the HBO 100 illuminator manually (423010-0000-000).

NOTE

If the FL attenuator is in the reflected light beam path, make sure that it is set to 100% transmission.

- Turn on the HBO 100 illuminator (Fig. 3-36/1) on the power supply unit for the HBO 100 (Fig. 3-63/1) and wait until it reaches its operational temperature.
- Pull the adjustment tool (Fig. 3-35/1) out of the microscope stand. The lighter colored focal point of the lamp HBO 103 W/2 and its slightly darker reflection become visible in the black glass window of the adjustment tool.
- Use the knurled button (Fig. 3-36/**4**) for collector adjustment to focus the brighter focal point.
- Use the adjustment screws (Fig. 3-36/2 and 3) to focus the darker focal point (reflection) as you would for the focal point image (Fig. 3-35/2) in the inscribed adjustment circle.

NOTE The two focal points of the HBO 103 W/2 lamp should be close to each other in the adjustment circle of the adjustment tool!

• Replace the adjustment tool in its original position.

3.1.22 Mounting the Colibri 3 illumination system and changing the LED modules

3.1.22.1 Mounting the Colibri 3 illumination system on the upper part of the Axioscope stand



CAUTION

Before removing the Colibri 3 from the microscope stand, please make sure the main power is switched off. Never look directly into the light-emitting aperture of the lamp.

- With the Allen wrench (AF 3) (Fig. 3-37/6) loosen the clamping screw (Fig. 3-37/5) of the RL illumination connector on the upper back of the stand.
- Align the illuminator's (Fig. 3-37/1) dovetail ring with the connection piece (Fig. 3-37/2) at the stand.
- Tighten the clamping screw (Fig. 3-37/5) with the Allen wrench (AF 3).
- Connect the illuminator plug (Fig. 3-37/4) to the stand socket (Fig. 3-37/3).
- Tighten the fixing screws on the plug.

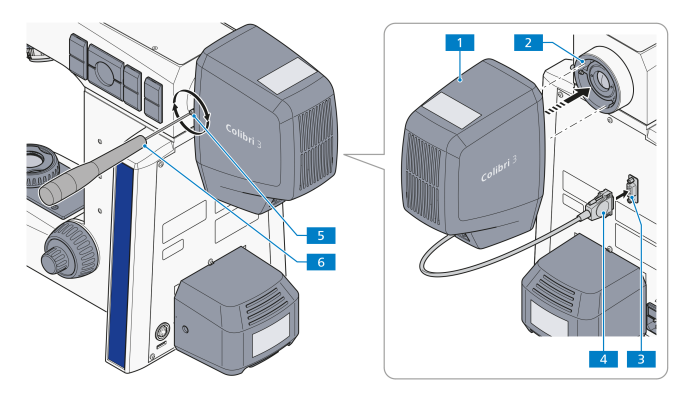


Fig. 3-37 Mounting the illumination system Colibri 3

3.1.22.2 Changing the LED modules in the Colibri 3 illumination system

How to change an LED module, i.e. the LED module for LED slot 1 (Fig. 3-38/):

- Remove the Colibri 3 from the stand: Follow the instructions of section 3.1.22.1 in reverse order.
- Loosen the four captive clamping screws (Fig. 3-38/5) on the front side with an Allen wrench (AF3).
- Remove the back cover (Fig. 3-38/7).
- Disconnect the LED module power cable connector (Fig. 3-38/1.1) from the PCBA (Fig. 3-38/1.6).
- Loosen the knurled knob (Fig. 3-38/1.3) by hand and remove the old LED module (Fig. 3-38/1.4).
- Select the right LED module with matched LED-specific labels (Fig. 3-38/1.2 and 1.5) and insert it in the correct slot (Fig. 3-38/①).

RF R

NOTE

One label on the LED slot (Fig. 3-38/**1.5**) indicates the wavelength range for this position; the other label on the LED module housing (Fig. 3-38/**1.2**) indicates the wavelength of the LED module. See the table below as well.

- Tighten the knurled knob (Fig. 3-38/**1.3**) again.
- Push the LED module power cable connector (Fig. 3-38/1.1) back to the PCBA (Fig. 3-38/1.6).
- If required, replace the LED modules of LED slots 2, 3 and 4 (Fig. 3-38/2), (3) and (4) in the same way.
- Mount the back cover (Fig. 3-38/7) and fasten the four anti-loss clamping screws (Fig. 3-38/5) with an Allen wrench (AF3).
- Re-mount the Colibri 3 on the stand again (see section 3.1.22.1).

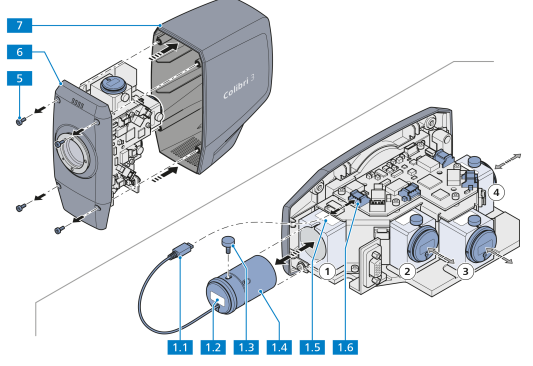


Fig. 3-38 Changing the LED modules in the Colibri 3 illumination system

Usability of LED modules for Colibri 3

Position	Wavelength range	385 nm LED 423052-9593-000	470 nm LED 423052-9573-000	505 nm LED 423052-9562-000	565 nm LED 423052-9602-000	625 nm LED 423052-9522-000
Slot 1	450~480 nm	Х	0	Х	Х	Х
Slot 2	350~415 nm	0	Х	Х	Х	Х
Slot 3	594~660 nm	Х	Х	Х	Х	0
Slot 4	508~565 nm	Х	Х	0	0	Х

0 X Usable

Not usable

Mounting the external illumination fixture HXP 120 3.1.23

Please refer to the separate operation manual of HXP 120 for instructions on installing and operating this system.

3.1.24 Mounting the Axiocam 202 mono or Axiocam 208 color

- Mount the C-mount camera adapter (Fig. 3-39/2) on the Axiocam (Fig. 3-39/1).
- Attach the Axiocam with the adapter to the camera port (Fig. 3-39/4) of the tube.
- Orient the camera to the stand and fix the position by tightening the ring nut (Fig. 3-39/3).
- Connect the camera to the stand (Fig. 3-39/7) via the USB (Commercial Micro-D) cable (USB 2.0) (Fig. 3-39/5).
- Connect the camera to an external monitor via HDMI cable (Fig. 3-39/6).
- Alternatively connect the camera to a WLAN router or PC.

NOTE NOTE

Please refer to the microscope operating system modes in section 2.5.10.

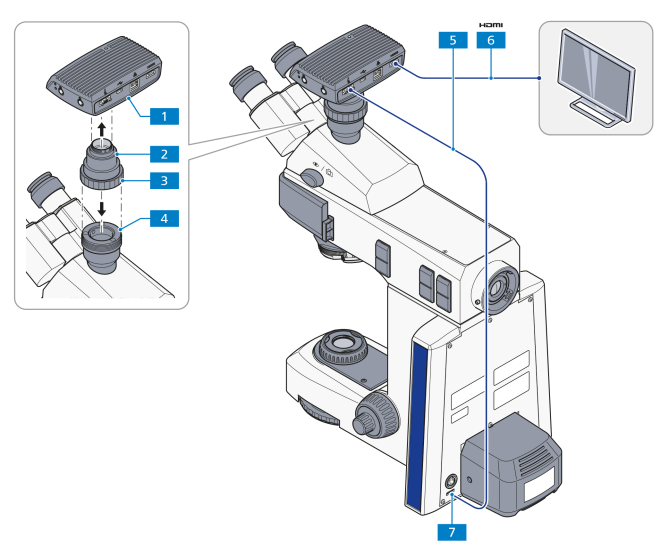


Fig. 3-39 Mounting the Axiocam 202 mono or Axiocam 208 color

CAUTION

Always disconnect the stand and the external power supply units from the power mains before beginning any mounting work.

3.2.1 Mounting the intermediate plate for Analyzer Slider

- Loosen the screw (Fig. 3-40/**5**) with the Allen wrench (AF 3) and lift off the tube (Fig. 3-40/**1**).
- Use the included ring tool to unscrew the tube lens (accessible from the bottom) and keep it free of dust.
- Screw the tube lens included with the intermediate plate into the tube.
- Insert the intermediate plate (Fig. 3-40/2) with the dovetail ring into the stand's dovetail ring socket (Fig. 3-40/6), adjust it, then fasten it with the Allen screw (Fig. 3-40/5).
- Insert the analyzer slider into the slider slot (Fig. 3-40/**3**) if applicable.
- Slightly incline the binocular tube (Fig. 3-40/1) when inserting it into the intermediate plate, adjust it and fasten it with the Allen screw (Fig. 3-40/4).

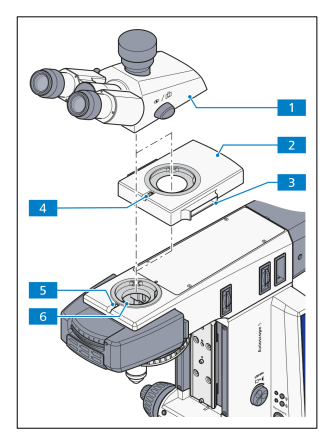


Fig. 3-40 Mounting the binocular tube with intermediate plate

3.2.2 Mounting the tube lens turret

Mounting the tube lens turret is done similarly to mounting the intermediate plate for analyzer sliders. See section 3.2.1.

F	NOTE
	Before using the tube lens turret, please remove the tube lens from the used tube.

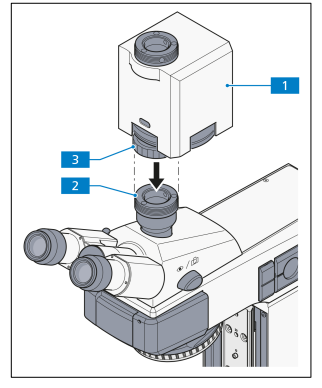


Fig. 3-41 Mounting the magnification changer

3.2.3 Mounting the magnification changer

- Remove the camera adapter or the dust protection cover from the camera port (Fig. 3-41/**2**) of the photo tube.
- Mount the magnification changer (Fig. 3-41/1) on the camera port, adjust it and fasten it with the retainer nut (Fig. 3-41/3).
- Mount the camera with the appropriate adapter on the camera port of the magnification changer.

The locking power for the magnification modules stop position can be adjusted with the screw located on the bottom of the magnification changer housing.

The screw is labeled with a white circle.

3.2.4 Changing the filters in the reflector module FL P&C

- Remove the reflector module FL P&C (Fig. 3-42/**3**) from the reflector turret (see also section 3.1.6).
- Unscrew the adapter ring (Fig. 3-42/1) with the mounting plate included with the tools (Fig. 3-42/6).
- Turn over the reflector module and let the filter (Fig. 3-42/2 or 5) drop onto a soft surface.
- Insert the emission filter (Fig. 3-42/2) or the excitation filter (Fig. 3-42/5) respectively at the position shown in Fig. 3-42.
- Fix the filters with the corresponding adapter ring (Fig. 3-42/1).

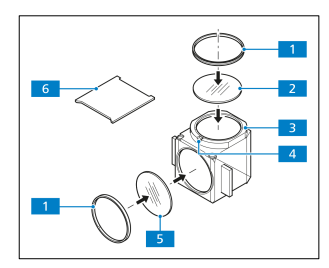


Fig. 3-42 Changing the filters in the reflector module FL P&C

NOTEWhen LED modules (423052-9593-000 for 385 nm, 423052-9573-000 for 470 nm, 423052-
9562-000 for 505 nm, 423052-9602-000 for 565 nm and 423052-9522-000 for 625 nm) are
used for fluorescence, the respective excitation filter is pre-installed in the LED module instead
of the reflector module.

The emission filter and excitation filter usually have a name and an arrow on their rims. The arrow indicates the direction of insertion and must always point to the inside of the reflector module (see arrows in Fig. 3-42). To minimize the image offset in multiple fluorescence images, the emission filter may have another mark to level the point angle.

When mounting the appropriate emission filter in the according reflector module, the mark should be aligned with the orientation groove (Fig. 3-42/**4**). This guarantees that the point angle in the reflector modules acts equally and compensates or minimizes the image offset – already low in ZEISS filter sets – even further.

When mounting a filter without any orientation marks (arrow), we recommend proceeding as follows:

Filters with reflective, di-electrical coatings are mounted with the reflective coating (Fig. 3-43/**6**) facing outward (in relation to the reflector module) in excitation filters (Fig. 3-43/**5**) and facing inward in emission filters (Fig. 3-43/**2**).

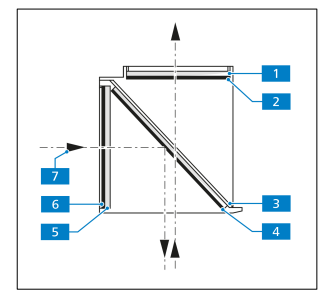


Fig. 3-43 Mounting the filters and the color splitter

The reflective coating (Fig. 3-43/4) of the color splitter (Fig. 3-43/3) faces down when the splitter is mounted.

The arrows (Fig. 3-43/7) in the figure mark the path of the illumination beam or the imaging beam.

3.2.5 Changing the color splitter in the reflector module FL P&C

!

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ATTENTION

Be extremely careful when installing the filters and the color splitter to avoid damaging and soiling the optical components.

Fig. 3-44 Opening the module

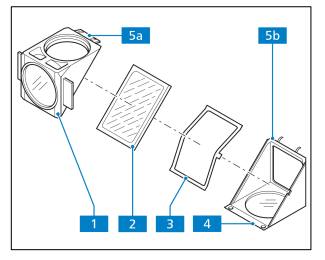


Fig. 3-45 Changing the color filter

We recommend ordering completely equipped reflector modules FL P&C, as changing the color splitter is very challenging.

When changing the color filter, please proceed as follows:

- Remove the reflector module FL P&C from the reflector turret (see also section 3.1.6).
- Loosen both slit screws (Fig. 3-44/1) with a screwdriver.
- Hold both parts of the reflector module (the **emission** part (Fig. 3-44/**2**) and the **excitation** part (Fig. 3-44/**3**) together, turn them against the mounting position and put them down.
- Now tilt the excitation part lying on top (Fig. 3-45/1) upward and lift the emission part (Fig. 3-45/4) from the holding fixtures (Fig. 3-45/5b).
- Remove color splitter (Fig. 3-45/**2**) and spring box (Fig. 3-45/**3**) from the bottom part of the module.
- Remove the old color splitter and carefully put the new splitter with the reflective coating facing upwards onto the spring box (Fig. 3-45/**3**). Put both pieces together into the bottom part of the module. Make sure that the side latch of the spring box is in the appropriate recess of the bottom part of the module.

The reflective (coated) side (Fig. 3-46/**3**) of the color splitter has a tapered edge (Fig. 3-46/**1**) or corner (Fig. 3-46/**2**).

- Place the excitation part of the module (Fig. 3-45/1) on the emission part (Fig. 3-45/4)

 holding the fixtures (Fig. 3-45/5b) and eyelet (Fig. 3-45/5a) latches together. Hold both parts together and turn them back into mounting position.
- Replace the slit screws and tighten them.
- Finally, attach the label with the name of the filter combination to the side of the module.

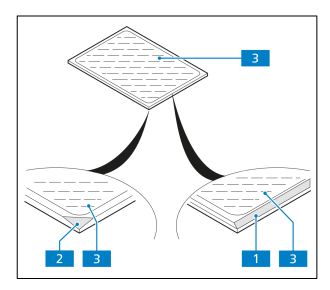


Fig. 3-46 Labeling the color splitter

3.2.6 Mounting the polarizer D or the filter holder

- For easy assembly, remove the condenser carrier (Fig. 3-47/**3**) fully, with the stage carrier.
- If necessary, unscrew stop bolt and bracket bolt with low-power system from the condenser carrier.
- Hold the polarizer or filter holder (Fig. 3-47/6) parallel to the bottom of the condenser carrier (Fig. 3-47/3) and screw the bracket bolt (Fig. 3-47/4) of the polarizer (Fig. 3-47/6) with the angled adjustment lever (Fig. 3-47/5) into the front threaded opening at the left below the condenser carrier (Fig. 3-47/3) until it stops.
- Screw the stop bolt (Fig. 3-47/2) with the adjustment lever (Fig. 3-47/1) to the stop into the rear threaded opening of the condenser carrier (Fig. 3-47/3).

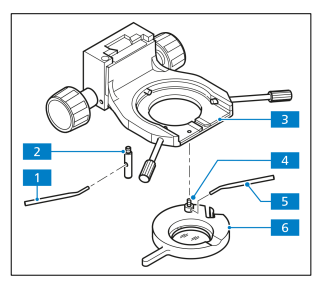


Fig. 3-47 Mounting the polarizer D

F	NOTE
	Proceed similarly when mounting the other components of polarization filter carrier.

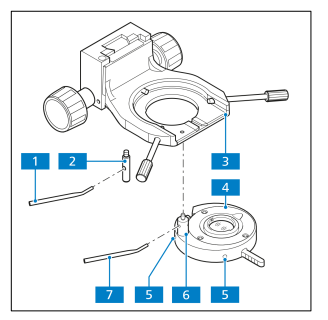


Fig. 3-48 Mounting the low-power system

3.2.7 Mounting and centering the lowpower system for 2.5x/4x objectives

- If necessary, remove the polarizer or filter holder from the condenser carrier.
- Hold the low-power system (Fig. 3-48/4) parallel to the bottom of the condenser carrier (Fig. 3-48/3) and screw the bracket bolt (Fig. 3-48/6) of the low-power system with the angled adjustment lever (Fig. 3-48/7) into the front threaded opening to the left below the condenser carrier (Fig. 3-48/3) until it stops.
- With the adjustment lever, (Fig. 3-48/1) screw the stop bolt (Fig. 3-48/2) into the rear threaded opening of the condenser barrier (Fig. 3-48/3) to the stop.
- Swing the low-power system into the beam path. Make sure that the fixture is securely engaged.
- Open aperture diaphragm and field diaphragm completely.
- Adjust both adjustment screws (Fig. 3-48/5) with the aid of two Allen wrenches (AF 1.5) until the field of view is well lit.

F	NOTE
	The low-power system is useful in only combination with the 0.9/1.25 condenser.

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3.2.8 Inserting the modulator disk in the 0.9/1.25 BF condenser

- Remove the condenser (Fig. 3-49/1) from the condenser carrier (see section 3.1.13). If the condenser cannot be lowered sufficiently, e.g. with mounted low-power system, then it might become necessary to remove it with the stage carrier, then lower it to the stop and remove the condenser.
- Loosen the clamping screw (Fig. 3-49/5) of the condenser's dial segment (Fig. 3-49/3) with the Allen wrench (AF 3) and pull out the dial segment.
- Slide the modulator disk (Fig. 3-49/**4**) with its two-pronged forked opening pointing forward into the condenser opening (Fig. 3-49/**2**). Make sure that the disk engages in the guide on both inner sides of the condenser. The guide serves as a stop for the modulator disk. The pin of the disk's clamping screw must slide into the orientation groove of the condenser.
- Tighten the disk's clamping screw with the Allen wrench (AF 3).
- Replace the condenser into its carrier (see section 3.1.13).

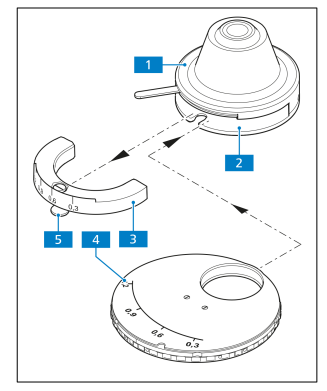


Fig. 3-49 Inserting the modulator disk in the 0.9/1.25 H condenser

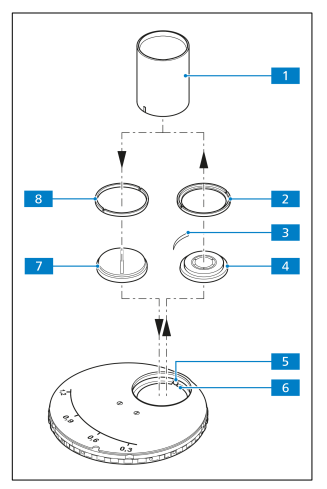


Fig. 3-50 Mounting the slit-diaphragm

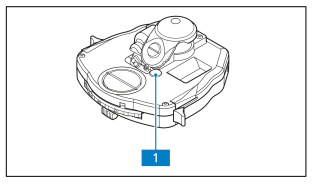


Fig. 3-51 Changing the PhC-DIC-PlasDIC diaphragm

3.2.9 Changing the PlasDIC diaphragm

How to mount the slit diaphragm (426715-9000-000):

- Turn the PhC diaphragm that needs to be replaced into the free aperture of the modulator disk.
- Screw the modulator disk's centering screws (Fig. 3-50/**5**) with the aid of an Allen wrench back until they stop.
- Unscrew the adapter ring (Fig. 3-50/**2**) of the PhC diaphragm with the included tool (Fig. 3-50/**1**).
- Remove the PhC diaphragm (Fig. 3-50/4) and spring (Fig. 3-50/3).
- With the aid of the tool (Fig. 3-50/**1**), insert the slitdiaphragm (Fig. 3-50/**7**) with the cams into the orientation grooves (Fig. 3-50/**6**).
- Screw in the adapter ring (Fig. 3-50/8) included with the slit diaphragm with the aid of the appropriate tool (Fig. 3-50/1).

3.2.10 Changing the PhC-DIC-PlasDIC diaphragm on the 0.9 BF DF PhC DIC achromaticaplanatic condenser

The diaphragm is changed according to the directions given in sections 3.2.8 and 3.2.9.

ATTENTION

After inserting the DIC module, do not tighten the condenser's centering screw again.

How to deactivate the pre-set diaphragm mechanism which automatically opens the diaphragm when phase contrast occurs:

• Remove the cover (Fig. 3-51/1). Turn the screw (Fig. 3-50/5) with the Allen wrench (AF 1.5) counterclockwise to the stop. You can now close the aperture diaphragm for contrasting procedures in DIC.

3.2.11 Changing the filter in the filter wheel transmitted light

- Remove the stage carrier (see section 3.1.12).
- Unscrew three screws (Fig. 3-52/**3**) of the field diaphragm socket (Fig. 3-52/**4**) and pull the socket out of the stand base.
- If necessary, pull the filter to be changed (Fig. 3-52/1) out of the corresponding position of the filter wheel (Fig. 3-52/2).
- Put the new filter into the position.

r P	NOTE
	One filter position is always designed as firmly installed shutter position.

- Repeat the procedure for all filter wheel positions.
- Put in the field diaphragm socket and screw it in.
- Mount the stage carrier (see section 3.1.12).

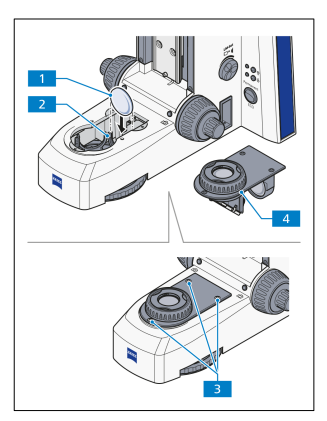


Fig. 3-52 Changing the filter in the filter wheel transmitted light

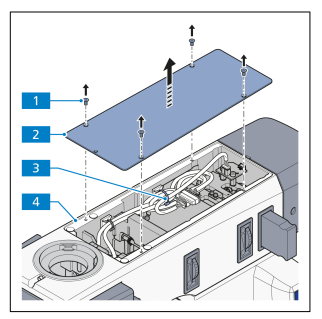


Fig. 3-53 Removing the cover of the upper stand part

3.2.12 Mounting the sample space extension, 60 mm

- Remove the binocular tube/photo tube, see section 3.1.3 (in reverse order).
- Unscrew the 4 screws (Fig. 3-53/1) of the cover and remove the cover (Fig. 3-53/2) from the upper stand part (Fig. 3-53/4).
- Cut and remove the one cable tie (Fig. 3-53/3).

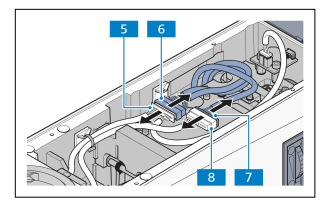


Fig. 3-54 Disconnecting the cable connections

- Disconnect the cable connection between the main control board (Fig. 3-54/7) of the lower stand part and the nosepiece sensor of the upper stand part (Fig. 3-54/8).
- Disconnect also the cable connection between the main control board (Fig. 3-54/**6**) and the turret sensor/RL LED interface (Fig. 3-54/**5**).

NOTE

Pure transmitted light stands have no reflector turret sensor / RL LED interface. Accordingly, no connection cables for this purpose are available.

- Hold the upper stand part (Fig. 3-55/4) and unscrew the 6 Allen screws (Fig. 3-55/9).
- Carefully, remove the upper stand part upward and pull the cables simultaneously downward out of the upper stand part.

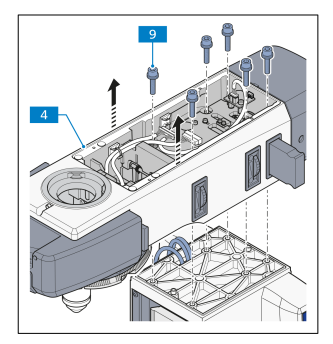


Fig. 3-55 Removing the upper stand part

- Carefully, thread the cables (Fig. 3-56/6 and 7) from the bottom through the long hole of the sample space extension (Fig. 3-56/11).
- Place the sample space extension on the lower stand part (Fig. 3-56/10) and tighten it with the 6 spacer bolts (Fig. 3-56/12) of the sample space extension.

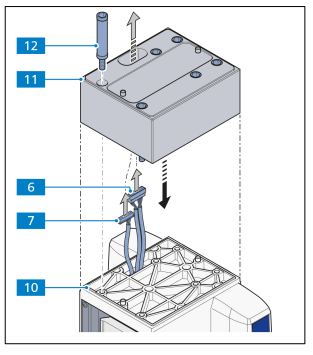
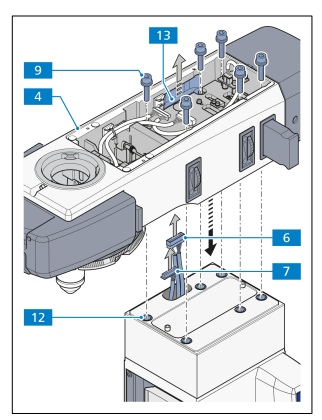


Fig. 3-56 Mounting the sample space extension



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Fig. 3-57 Mounting the upper stand part on the sample space extension

- Carefully, thread the cables (Fig. 3-57/6 and 7) from the bottom through the lateral opening (Fig. 3-57/13) of the upper stand part (Fig. 3-57/4).
- Place the upper stand part on the sample space extension (Fig. 3-57/12) and tighten it with the 6 Allen screws (Fig. 3-57/9) of the upper stand part.

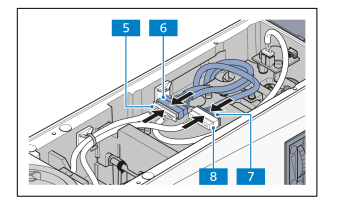


Fig. 3-58 Establish the cable connections

- Establish the cable connection between the main control board (Fig. 3-58/7) of the lower stand part and the nosepiece sensor of the upper stand part (Fig. 3-58/8).
- Establish the cable connection between the main control board (Fig. 3-58/6) and the turret sensor/RL LED interface (Fig. 3-58/5).
- Make sure the plugs are securely connected.

R)	NOTE
	Pure transmitted light stands have no reflector turret sensor / RL LED interface. Accordingly, no connection cables for this purpose are available.

- Place the cover (Fig. 3-59/2) on the upper stand (Fig. 3-59/4) part and tighten it with the 4 screws (Fig. 3-59/1).
- Secure the cables with one cable tie (Fig. 3-59/**3**).
- Mount the binocular tube/photo tube, see section 3.1.3.

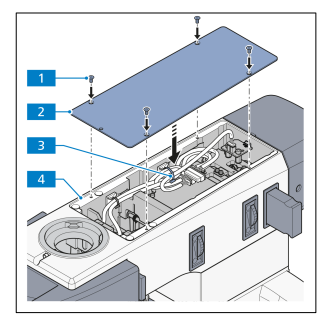


Fig. 3-59 Mounting the cover on the upper stand part

3.3 Connecting to the power supply and switching the microscope on/off

3.3.1 Microscope stand

ZEISS

• Connect the microscope's power supply (Fig. 3-60/2) with its power cord to an electrical outlet.

R3	NOTE
	The microscope can adjust to power, voltage and frequency ranges of 100 to 240 V AC, 50-60 Hz. The power supply unit adjusts automatically to the available voltage.

• Switch the microscope on or off with the main switch (Fig. 3-60/1) on the left side of the stand.

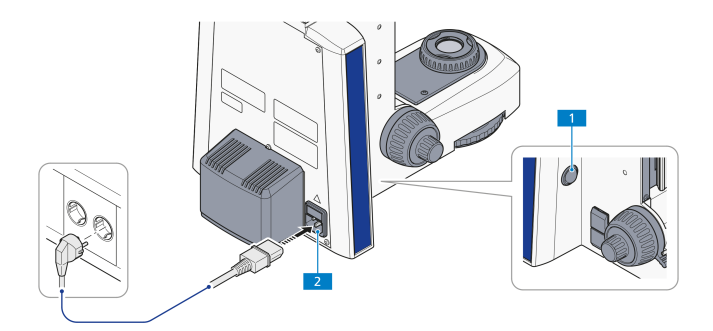


Fig. 3-60 Lower stand part for HAL 50 illumination (proceed similarly for LED illumination)

3.3.2 Connecting the HAL 100 halogen illuminator for transmitted light

NOTE
When using the HAL illuminator, a base plate for Axioscope (000000-2202-526) is mandatory.

- Mount the HAL 100 (Fig. 3-61/**4**) on the transmitted light connector of the stand (see also section 3.1.19.1).
- Connect the 3-pole lamp plug of the HAL 100 to the port for transmitted light **TL** (Fig. 3-62/**3**) of the separate power supply unit for two lamps (Fig. 3-61/**2**).
- Connect the cable for illumination intensity control (Fig. 3-61/1) to the corresponding port (**Remote**) on the back of the stand or to that of the power supply unit (Fig. 3-62/4).
- Switch the toggle switch (Fig. 3-61/**3**) for reflected or transmitted light to the **TL** position (transmitted light).

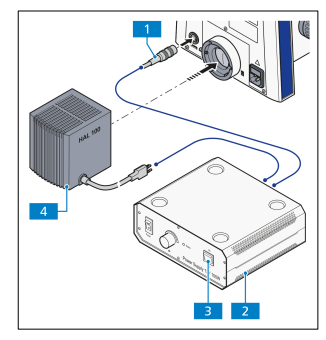


Fig. 3-61 Lower stand part for HAL 100 illumination

3.3.3 Connecting the HAL 100 halogen illuminator for reflected light

- Mount the HAL 100 on the reflected light connector of the stand (see section 3.1.19.1).
- Connect the 3-pole lamp plug of the HAL 100 to the port for reflected light **RL** (Fig. 3-62/**2**) of the separate power supply unit for two lamps.
- Connect the power cord of the external power supply unit (Fig. 3-62/**5**) to an electrical outlet.
- Switch the toggle switch for reflected or transmitted light to the **RL** position (reflected light).

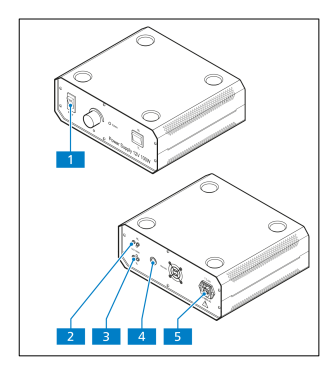


Fig. 3-62 External power supply unit for two lamps (front and back)

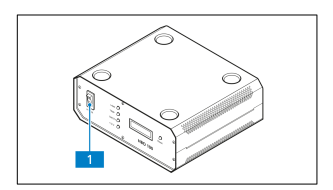


Fig. 3-63 Power supply unit for HBO 100 (front and back)

3.3.4 Switching the illumination on/off

- Turn the power switch (Fig. 3-60/1) of the microscope with the lower part of the stand for LED or HAL 50 illumination either on or off.
- halogen illuminator for Switch on the transmitted light (lower stand part for HAL 100 illumination) and reflected light (if in use) at its separate supply external power unit (Fig. 3-62/1) (also refer to the separate operation manual of the external power supply unit). Depending on whether you want to use the illuminator for reflected light or the illuminator for transmitted light, switch the togale switch (Fig. 3-61/3) for reflected or transmitted light to the TL position (for transmitted light) or to the RL position (for reflected light.
- Switch the HBO 100 illuminator for overhead fluorescence light (if in use) on or off with the power switch of the power supply unit for HBO 100 (Fig. 3-63/**1**).

F	NOTE
	If the control cable (Remote) is connected (see section 3.3.2) to the stand, the illumination intensity of the HAL 100 illuminator for transmitted or reflected light, depending on which is switched on, can be varied by rotating the Intensity/LM knob on the lower part of the stand.
∎ >>⊃	1075

R	NOTE
	If the control cable is not connected to the stand, the illumination intensity of the HAL 100 illuminator in use can be varied by rotating the knob of the external power supply unit.

3.4 Using the Light Manager function

The Light Manager (LM) function saves the ratios of the set light intensities between different combinations of objective and reflector turret positions for a given light source. When changing the light intensity of one objective/reflector combination, the light intensities of other combinations will also change according to the set ratios. This ensures that users don't need to repeatedly set up light intensities for each objective/reflector combination when switching between samples which require different illumination intensity.

After switching on the microscope, the previous setting of the Light Manager will be restored.

Disable/enable the LM function

To disable the LM function proceed as follows:

• Press one of the **Snap** buttons and the **Intensity/LM** knob simultaneously for at least 1.5 seconds.

The indicator light blinks in the following sequence: GREEN / ORANGE / GREEN

To enable the use of the LM function proceed as follows:

• Press one of the **Snap** buttons **and** the **Intensity/LM** knob simultaneously for at least 1.5 seconds.

The indicator light blinks in the following sequence: GREEN / GREEN / GREEN

Save light intensity ratios using the LM function

- 1. Switch to the first objective and/or reflector positions of interest.
- 2. Set the desired light intensity.
- 3. Press the Intensity/LM knob for at least 1.5 seconds.
- 4. The light intensity for this objective/reflector combination is then saved. When using LED as light source, after the light intensity is saved, LED is switched off for 300 ms. This is visible through eyepieces and serves as an indicator for the user.
- 5. Proceed by switching to the second objective/reflector combination, and press the Intensity/LM knob for at least 1.5 seconds. Now a ratio between the first and the second objective/reflector combinations is established.
- 6. Repeat step 5 to set light intensity ratios for more objective/reflector combinations.

3.5 Default factory settings of the microscope

The default factory settings are:

- Light Manager enabled, but no light intensity values saved
- Light intensity set to initial minimum value
- all configuration stored will be cleared

To reset the microscope to default factory settings, proceed as follows:

• Press and hold down the Intensity/LM knob for 20 seconds.

While the knob is held down from 3 s to 20 s, the indicator light blinks in RED.

It blinks GREEN after 20 seconds. When the indicator stops blinking and remains GREEN, the reset to the default factory setting is successful.



ATTENTION

Please use this function with caution as it will reset **all** existing configurations.

ZEISS

4 OPERATION

4.1 Default setting of the microscope

4.1.1 Setting the inter-pupillary distance on the binocular tube

• Adjust the individual inter-pupillary distance by symmetrically swiveling of the two eyepiece tubes towards each other (Fig. 4-1).

The correct inter-pupillary distance has been set when the observer sees only **one** round image when looking into the eyepieces!

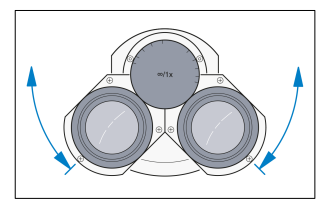


Fig. 4-1 Setting the inter-pupillary distance on the binocular tube

4.1.2 Setting the viewing height

• Adjust the viewing height to your individual requirements by swiveling the eyepiece tubes up (Fig. 4-2/**A**) or down (Fig. 4-2/**B**).

The binocular ergonomic (photo) tubes (425511-0000-000, 425512-0000-000, 445514-0000-000 and 425520-9050-000) are equipped with a continuous height adjustment in the range from 44 mm to 50 mm.

The binocular piece of the ergo tube 425520-9050-000 can also be pulled out horizontally by 50 mm.

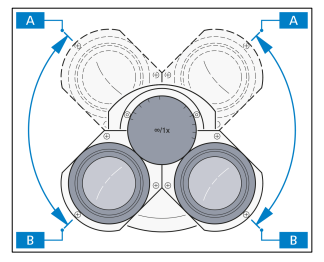


Fig. 4-2 Setting the viewing height on the binocular tube

4.1.3 Adjusting for ametropia (user's visual impairment) when using eyepiece reticles

The prerequisite for the correct use of an eyepiece reticle is two adjustable eyepieces (e.g. PL 10x/23 GW foc.) to compensate for different degrees of ametropia of the user.

To adjust the eyepieces for different degrees of ametropia, follow the subsequent steps:

- Focus on the line figure of the eyepiece reticle with the focusable eye lens of the adjustable eyepiece.
- Focus on the microscopic image of a loaded specimen with the focusing drive while observing with the eyepiece containing the eyepiece reticle.
- When both the microscopic image and the eyepiece reticle are in focus, the image for the second eye is brought into focus with the focusable eye lens of the second eyepiece.

Both microscopic images including the eyepiece reticle are thus in focus. From this point, use only the focusing drive to focus images.

4.2 Illumination and contrast methods in transmitted light microscopy

4.2.1 Setting up transmitted light brightfield microscopy using the KÖHLER method

(1) General principle of operation

Of all the optical microscopy methods, transmitted light brightfield microscopy is the one most commonly used. High-contrast or tinted (stained) samples (e.g. a blood smear) can be examined easily and quickly.

For an imaging result which is as true to the specimen as possible, we must not only consider the socalled direct bundled beams but the indirect ones as well, i.e. the beams which diffract and scatter on the sample details. According to ABBE, the image is truer to the specimen when the fraction of the **cone** of light is as large as possible.

The best performance of the microscope, and especially its objective, is achieved when the condenser, field diaphragm and aperture diaphragm are adjusted in accordance with the KÖHLER illumination principles. These fundamental basic rules for adjusting a microscope are explained in detail in section 4.2.1 (3) "Setting up transmitted light brightfield microscopy using the KÖHLER method".

(2) Instrumentation for transmitted light brightfield microscopy

Every microscope (except the one with the Vario stand column) is configured to work with the transmitted light brightfield method.

All available condensers (except special condensers like darkfield condensers) can be used for the transmitted light brightfield method.

(3) Setting up transmitted light brightfield microscopy using the KÖHLER method

- The Axioscope has been put into operation appropriately (section 2.5.10).
- The Axioscope is turned on.
- Adjust the image brightness using the Intensity/LM knob (Fig. 4-3/1) on the microscope stand.
- Put a high-contrast sample into the specimen holder of the mechanical stage.
- If condensers with a swiveling front lens are used, swivel these into the beam path with $\geq 10x$ objectives and turn the knurled knob for vertical adjustment of the condenser (Fig. 4-3/3 and Fig. 4-4/2) to the top stop. Make sure the stop is adjusted to prevent the condenser from lifting out the sample (adjusting the condenser stop, section 4.2.1 (4)).
- On condensers with revolver/modulator disks: turn the knurled ring (Fig. 4-4/3) to position H (or **BF** = brightfield).
- Bring the 10x objective into position on the nosepiece (Fig. 4-3/6) and focus the sample using the focus drive (Fig. 4-3/2).
- Close the field diaphragm (Fig. 4-3/5) enough to make it visible in field of view (even if blurred) (Fig. 4-3/A).
- Lower the condenser using the knurled knob for vertical adjustment until the edge of the field diaphragm appears sharp (Fig. 4-3/**B**).

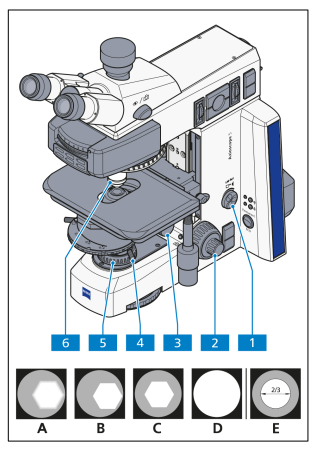


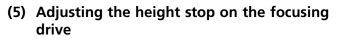
Fig. 4-3 Microscope adjustment in transmitted light brightfield microscopy

- Center the field diaphragm image with both centering screws (Fig. 4-3/4) on the condenser carrier (Fig. 4-3/C).
- Open the field diaphragm enough to make the edge of the diaphragm disappear from the field of view (Fig. 4-3/**D**).
- To adjust the aperture (contrast), take an eyepiece from the tube barrel. Looking through the barrel adjust the aperture with the adjusting lever (Fig. 4-4/**4**) of the aperture diaphragm to the size of approx. 2/3 4/5 of the diameter of the objective exit pupil (Fig. 4-3/**E**). In most cases this aperture gives the best contrast at almost full resolution and is thus the best compromise for the human eye.
- Replace the eyepiece in the tube barrel.

r S	NOTE
	Each objective changes the size of the object field and the objective aperture. The centering might also vary slightly from objective to objective, so that an adjustment of field diaphragm and aperture might become necessary for optimal results.
	With objectives < 10x, the front lens of the condenser (if swivelable) must be swiveled out of the beam path and the aperture opened fully. In order to achieve a better contrast, the field diaphragm can be pulled closer when working with large object fields. To do this you must reduce the opening by a certain range. Avoid closing it too much to guarantee a consistent illumination of the field of view.

(4) Adjusting the height stop on the condenser carrier

- Loosen the setscrew (Fig. 4-4/1) on the height stop with the Allen wrench (AF 3).
- Focus the sample with the focusing drive.
- Close the field diaphragm and adjust the condenser vertically (Fig. 4-4/2) until you get a sharp image.
- Perform the vertical adjustment on the condenser by small increments only to avoid lifting out the sample.
- Tighten the setscrew (Fig. 4-4/1) on the height stop.



- Turn the clamping lever (Fig. 4-5/1) of the stop towards the pin stop.
- Move the stage with the aid of the focusing drive (Fig. 4-5/2) to the highest position allowable without risking colliding with specimen holder or objectives.
- Now press the clamping lever down until the stop engages again.

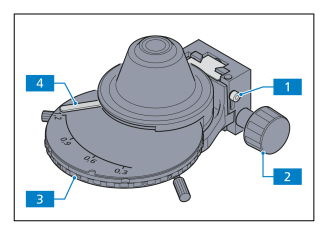


Fig. 4-4 Adjusting the height stop on the condenser carrier

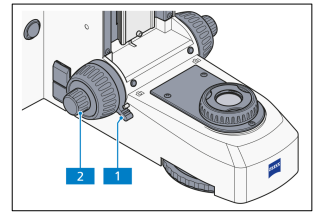


Fig. 4-5 Adjusting the height stop on the focusing drive

4.2.2 Setting up transmitted light darkfield microscopy using the KÖHLER method

(1) General principle of operation

In the transmitted light darkfield microscopy you basically illuminate the sample with an illumination aperture which is higher than the one of the objective you are using.

In darkfield microscopy, only the diffracted and scattered light portions which are important for the imaging procedure get into the objective, whereas the indirect unaffected light beams are directed past the objective. Thus a resolution of fine structures can be achieved which is below the resolution capacity of a light microscope. The fine structures now appear bright and incandescent on a dark background.

(2) Instrumentation

Condenser with transmitted light stop in position **D** e.g.:

- Condenser 0.9/1.25 H with modulator disk BF, DF, PhC 1, PhC 2, PhC 3,
- Condenser, achrom.-aplan. 0.9 BF DF PhC DIC,
- Darkfield condenser with dry darkfield (465505-0000-000 applicable aperture from 0.6 0.75),
- Ultra-condenser (465500-0000-000 applicable aperture from 0.75 1).

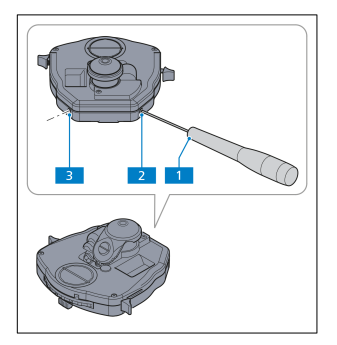


Fig. 4-6 Centering the darkfield diaphragm on the condenser, achromaticaplanatic 0.9 H D PhC DIC

(3) Setting up transmitted light darkfield microscopy

- Adjust the KÖHLER illumination as for transmitted light brightfield microscopy. Instead of the 10x objective it is necessary to use an objective with the highest possible aperture. The aperture must not exceed the aperture of the condenser being used.
- Set the revolver/modulator disks to position **D** and bring the condenser front lens into position (if applicable).
- Take the eyepiece out of the tube (or replace it with the auxiliary microscope) and check the centering of the darkfield diaphragm in the objective exit pupil. If the DF central darkfield diaphragm in the universal condenser is partially outside or is not centered to the objective exit pupil and the pupil does not appear consistently dark, it is necessary to re-adjust the centering of the darkfield diaphragm.
- If the darkfield diaphragm needs to be centered, use two Allen wrenches (AF 1.5) (Fig. 4-6/1) and adjust the two centering screws (Fig. 4-6/2 and 3) until the objective exit pupil appears consistently dark. Remove the Allen wrenches (AF 1.5) from the condenser after finishing the centering procedure.

F	NOTE
	Objectives with built-in iris apertures have apertures which are too high for the transmitted light darkfield microscopy. Stop the iris aperture down at least until it reaches the border
	aperture for the darkfield of the condenser in use.

Performance criterion for the darkfield method is always the darkest possible background for the field of view.

- Replace the eyepiece in the tube.
- Lighter areas which might still be visible in the field of view are reduced when the darkfield condenser is adjusted correctly and delicately. You now have an approximately sharp eye diaphragm image.
- Now adjust the field diaphragm diameter to the size of the field of view.

Darkfield samples need to be kept impeccably clean, more so than samples for any other method. A fingerprint, dust or any dirt particle can have a negative effect, as they brighten the background and reduce the contrast of the object image.

(4) Setting darkfield contrast with a dry darkfield condenser

- If necessary, swivel out the low-power system, polarizer or λ plate.
- Move the condenser carrier down until the end stop.
- Place the dry darkfield condenser in the condenser holder Z (see section 3.1.14).
- Place the condenser holder Z in the condenser carrier and center it roughly, so the condenser fits into the opening of the mechanical stage without contact, when moving upwards.
- Move the condenser up until the end stop. Place the specimen and adjust the illumination intensity to bright enough.
- Swivel in an objective with small magnification (e.g. 5x or 10x) and focus the specimen, using the focusing drive.
- Place a specimen as its details are distributed evenly in the field of view. This will make the image of the field diaphragm easier to identify.
- Close the field diaphragm until the end stop.
- Lower condenser until edge of field diaphragm appears sharp (luminous-field diaphragm focus level). There will be an increasing or decreasing light ring visible, when moving the focus upwards or downwards from the field diaphragm focus level (so called circular "breathing" of the field diaphragm depiction).
- Center the field diaphragm image with both centering screws on the condenser carrier.
- Swivel in the desired objective.
- Focus the specimen with the focusing drive, if necessary.
- Focus the field diaphragm, using the condenser drive. Open the field diaphragm enough to make the edge of the diaphragm disappear from the field of view.
- Optimize the contrast with the aperture diaphragm on the condenser, if necessary.

(5) Setting darkfield contrast with immersion oil darkfield condenser

- If necessary, swivel out the low-power system, polarizer or λ plate.
- Move the condenser carrier down until the end stop.
- Place the immersion oil darkfield condenser in the condenser holder Z (see section 3.1.14).
- Place the condenser holder Z in the condenser carrier and center it roughly, so the condenser fits into the opening of the mechanical stage without contact, when moving upwards.
- Move the condenser up until the end stop.
- Place a drop of immersion oil (without bubbles, if possible) on the center of the condenser front lens.
- Place a specimen. The immersion oil will disperse between condenser and specimen holder.
- Slightly move the mechanical stage back and forth to dissipate any air bubbles in the immersion oil.
- Adjust the illumination intensity to bright enough and open the field diaphragm completely.
- Swivel in an objective with small magnification (e.g. 10x) and focus the specimen, using the focusing drive.
- Center the field diaphragm on the condenser carrier with the adjustment screws and focus the image with the condenser drive.
- Place a drop of immersion oil on the specimen holder, swivel in immersion oil objective and focus the specimen.
- Close the field diaphragm until the end stop.
- Lower condenser until edge of field diaphragm appears sharp.
- Center the field diaphragm image with both centering screws on the condenser carrier.

The luminous field diaphragm appears only as a circle segment on the edge of the viewing field due to the high magnification of the immersion oil objective. As a result, the focusing and centering of the field diaphragm must be repeated. If necessary, the luminous-field objective should be opened slightly if the light intensity is too low.

The field diaphragm is centered properly when the edge of the luminous field diaphragm is centered or equidistant from the viewing field edge.

- For a sharply focused specimen, open the sharply set field diaphragm enough to make the edge of the diaphragm disappear from the field of view.
- You can improve the contrast of the microscope image by slightly adjusting the focus level of the condenser with the condenser drive.
- The background of the eyepiece image should appear equally dark.
- For immersion oil objectives with an iris diaphragm, the contrast can be further optimized by turning the adjustment of the iris diaphragm.

4.2.3 Setting up transmitted light phase contrast microscopy

(1) General principle of operation

The phase contrast method is ideal for examining thin uncolored samples, e.g. individual cells of cell cultures. Generally, the human eye cannot detect phase differences (variations in refractive index or thickness) within the different cell components.

The phase contrast method uses the optical modulators "annular phase diaphragm" and "phase ring" to convert the small phase differences in intensity differences which are visible to the human eye. The interference of different beams in the intermediate image is important for the generation of such images.

With the aid of the optically defined ring channel "annular phase diaphragm and phase ring", the bright direct light portions are attenuated and provided with a constant phase shift. The indirect light portions, however, which are diffracted by different cell particles, bypass this optical channel and their phase is affected by the difference in the sample's refractive index and thickness.

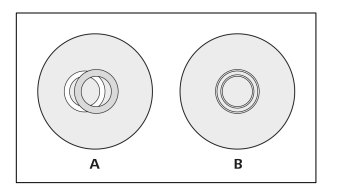
In the intermediate image plane, the partial beams are thus differently affected and achieve interference and strengthen or weaken each other (constructive and destructive interference) – depending on their phase. As a result, these interferences create image contents with intensity differences visible to the human eye.

(2) Instrumentation

- Phase contrast objectives with the phase rings PhC 1, PhC 2 or PhC 3 for low, medium and high numeral apertures which can also be used in the brightfield.
- Condenser with revolver/modulator disk which contains centerable ring diaphragms PhC 1, PhC 2 and PhC 3 for low, medium and high numeral apertures.
- The phase annular diaphragm used on the condenser must be consistent with the labeling on the objective which is used, e.g. PhC 1.

(3) Setting up transmitted light phase contrast microscopy

- Swivel the phase contrast objective, e.g. labeled Ph 1, into the beam path.
- Switch on the annular phase diaphragm on the condenser's revolver disk which has the same label as the phase contrast objective, e.g. 1.
- In order to check the centering and the overlap of the lighter annular diaphragm (in the condenser) with the darker phase ring (in the objective), remove an eyepiece from the tube and replace it with an auxiliary microscope. Use the adjusting fixture on the auxiliary microscope to focus the annular diaphragm and the phase ring in the objective exit pupil.



- Fig. 4-7 Centering the annular phase diaphragm (light-colored, in the condenser) and the phase ring (dark-colored, in the object)
- If the overlap is not exact (Fig. 4-7/A), the lighter annular diaphragm must be re-centered with the aid of two Allen wrenches (AF 1.5) (Fig. 4-6/1). Adjust the two centering screws (Fig. 4-6/2 and 3) so that you achieve a full overlap with the darker phase ring (Fig. 4-7/B).
- Remove the auxiliary microscope from the tube and replace it with the eyepiece.

To increase the image contrast, a green 32×4 interference broadband filter may be mounted on the field diaphragm or inserted into the color glass carrier (if available).

A complete phase contrast can only be achieved when the light-colored annular diaphragm (in the condenser) overlaps exactly with the dark-colored phase ring (in the objective) in the illumination beam path (Fig. 4-7/**B**).

4.2.4 Setting up transmitted light differential interference contrast (DIC)microscopy

(1) General principle of operation

The transmitted light DIC method allows for a high-contrast vivid display of transparent sample details.

The light is linearly polarized by a polarizer and is separated into two beams in a birefringent prism. The two beams penetrate two parts of the sample which are closely separated. They experience different path differences due to the different refractive indices or different sample thickness. Both beams are then reunited in a second double refractive prism and have after passing the analyzer the same polarization. Therefore both beams can interfere in the intermediate image and the path differences are thus converted into intensity differences represented by a gray scale. A compensator or λ -plate may be used for a consecutive conversion of the gray scale in a color scale.

(2) Instrumentation

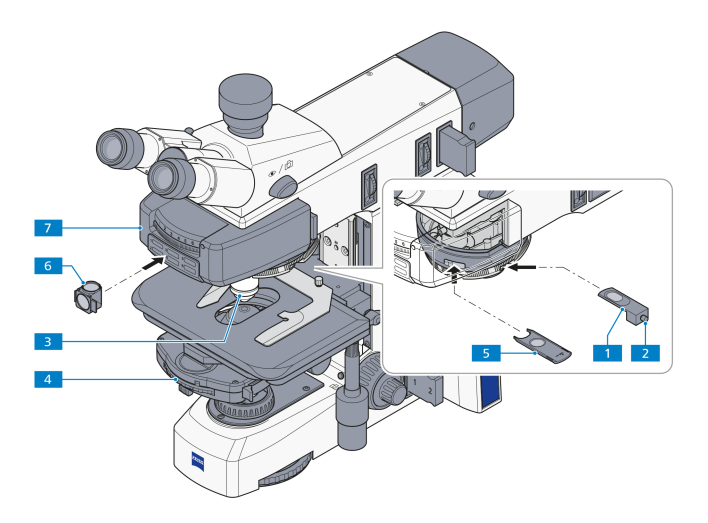
- Objectives equipped with DIC fixtures, e.g. EC Plan-NEOFLUAR objectives
- Nosepiece with slot for DIC slider
- DIC slider, compatible with the objectives in use
- Condenser with turret disk containing DIC prisms (e.g. condenser, achromatic-aplanatic 0.9 H D PhC DIC)
- Analyzer module ACR P&C for transmitted light in the reflector turret/slider or analyzer slider D/A fixed or rotatable in conjunction with a mounted intermediate plate for analyzer slider 12x46
- Rotatable mechanical stage, if necessary

(3) Setting transmitted light DIC

- Put the DIC compatible objective (Fig. 4-8/**3**) in place on the nosepiece. Slide the according DIC slider (Fig. 4-8/**1**) into the slit of the appropriate objective position. Make sure that the slider engages firmly.
- Swing in the analyzer module (Fig. 4-8/6) on the reflector turret (Fig. 4-8/7) (or slide the analyzer slider into the intermediate plate for analyzer sliders).
- Swing in the DIC prism (use position **Ph** on the condenser) (Fig. 4-8/4).
- Adjust field diaphragm and aperture diaphragm according to the KÖHLER method.
- Adjust the optimal contrast on the DIC slider (Fig. 4-8/1) with the knurled screw (Fig. 4-8/2). Symmetrical adjustment of the DIC slider along its middle position lets the sample details appear as if they were elevated or deepened.
- If desired, the compensator λ (Fig. 4-8/5) can be put into the opening above the nosepiece in order to create a chromatic DIC contrast.

R3	NOTE
	The DIC method functions with polarized light. It is disturbed when birefringent elements, e.g. foils, are put between polarizer and analyzer, as is sometimes done when doing a histological incision. The same situation occurs with Petri dishes or specimen holders which have a plastic base. In these cases we recommend using PlasDIC.

OPERATION Illumination and contrast methods in transmitted light ... Axioscope 5/7/Vario



- DIC slider 1
- Knurled screw 2
- 3 Objective on the nosepiece
- 4 Condenser with DIC prism
- 5
- Compensator λ Analyzer module Reflector turret 6
- 7

Components for the transmitted light DIC method Fig. 4-8

4.2.5 Setting up transmitted light PlasDIC contrast microscopy

(1) General principle of operation

PlasDIC can be used independently from the material of the specimen holder.

The contrast method gives a relief-like image and is especially well suited for thicker objects. The contrast is adjustable. It is possible to contrast the cavities of microtiter plates up to the edge. It is not necessary to use cultivation holders with a glass base.

(2) Instrumentation

- Abbe condenser with modulator disk and objective-dependent 2 mm slit diaphragm (A-Plan 10x and LD A-Plan 20x) or 4.5 mm slit diaphragm (in all other cases)
- Objectives:
 A-Plan 10x, 20x, 40x;
 LD A-Plan 20x, 32x, 40x;
 LD Plan-Neofluar 20x, 40x, 63x
- PlasDIC slider depending on the objective
- ACR P&C analyzer module for transmitted light and reflector turret /slider or D/A fixed or rotatable analyzer slider in connection with mounted intermediate plate for analyzer slider 12x46

(3) Setting PlasDIC

- Fully open the aperture on the condenser.
- Place the sample on the stage.
- Swing the position with the 2 or 4.5 mm slit diaphragm for PlasDIC of the condenser into the beam path. Increase brightness when switching from brightfield to PlasDIC.
- Swing the analyzer module into the beam path or slide analyzer slider in beam path in intermediate plate.
- Swing in the objective for PlasDIC.
- Slide the PlasDIC slider on the nosepiece into the DIC slit of the objective in use.
- Adjust the contrast with the knurled screw of the PlasDIC slider. You can now see the structures in relief or in pseudo-darkfield. The best result is achieved in the relief mode.

4.2.6 Setting transmitted light polarization

4.2.6.1 Detecting birefringence

(1) Application

The transmitted light polarization method is used for samples which change the polarization of the light. Such samples are called birefringent. Examples include crystals, minerals or polymers. If such birefringent substances are observed between crossed polarizers, the birefringent portion of the sample appears bright while its surroundings remain dark.

A birefringent substance can be recognized by turning the sample by 360° between crossed polarizers. The sample should show four bright and four dark appearances during the turning procedure. During the turning procedure, interference colors appear that range from gray (mostly for biological samples) through white, yellow and red until blue, depending on birefringence, thickness as well as sample orientation. The interference colors may be of the first or a higher order.

(2) Instrumentation

- Strain-free objectives
- Pol rotary stage
- D Polarizer (rotatable or fixed), if no polarizer is already integrated in the condenser.
- ACR P&C analyzer module Pol for transmitted light in the reflector turret/slider or D analyzer slider fixed or with lambda-plate.
- Depolarizer for avoiding unwanted polarizing effects

A depolarizer (quartz depolarizer) should be incorporated in all microscopes used to examine mineralogical/geological specimens.

A depolarizer extinguishes undesirable polarization effects possibly (e.g. false or pseudo-pleochroism) occurring behind the analyzer (e.g. on prism surfaces in the tube), or shifts them to higher orders.

• Insert the quartz depolarizer with tube lens 115 (428106-9010-000) into the tube

or

• insert the quartz depolarizer with tube lens 130 in combination with intermediate plate (428106-9000-000) into the tube.

(3) Setting the microscope

- Adjust the microscope as described in section 4.2.1 (3) for transmitted light brightfield microscopy using the KÖHLER method.
- Center the rotary stage Pol (Fig. 4-9/1) (see section 3.1.10.5) and objectives (see section 3.1.10.6).
- Swing the polarizer (Fig. 4-9/3) into the beam path and position it to 0° if you are using a rotatable polarizer.
- Put the analyzer (Fig. 4-9/2) into the slit for the compensator or swing analyzer module on the reflector turret/slider. Because of the crossed polarizers the field of view now appears dark.

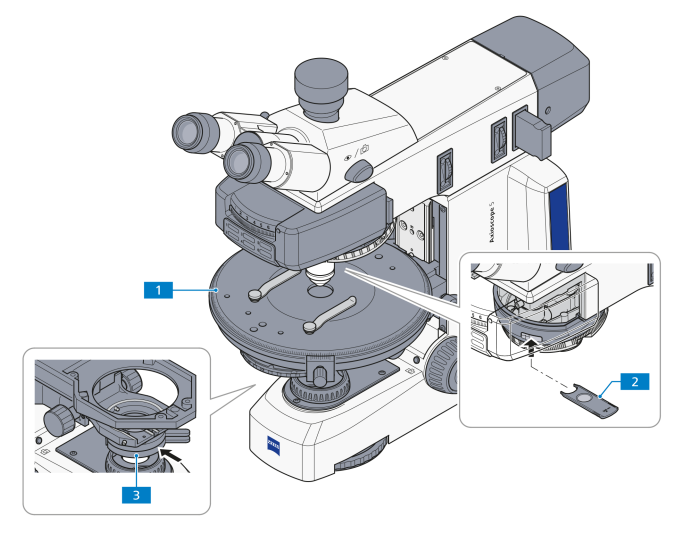


Fig. 4-9 Components for transmitted light polarization

NOTE NOTE

If an analyzer slider is going to be used for working with the Axioscope, either the reflector module H P&C or an empty position must be swiveled in on the reflector turret/slider.

• Bring the specimen to be examined into the field of view and turn it with the rotary stage. Normally, birefringent (anisotropic) objects will now show the same color and intensity variations as described above during rotation between crossed polarizers. Optically anisotropic substances may remain dark when an isotropic direction, e.g. from optically single-axle or double-axle crystals, is placed parallel to the observation direction.

4.2.6.2 Determination of the polarization direction nγ'

(1) Application

The determination of the polarization direction of n_{γ} or $n_{\gamma'}$ respectively (polarization direction with the absolute or relative largest index of refraction) and n_{α} or $n_{\alpha'}$ respectively (polarization direction with the absolute or relative smallest index of refraction) relative to the morphological directions, e.g. of crystal surfaces, crystal needles or fibers, provide an important signature of the material. This method is also used in the diagnosis of bio-crystals (e.g. gout and pseudo-gout).

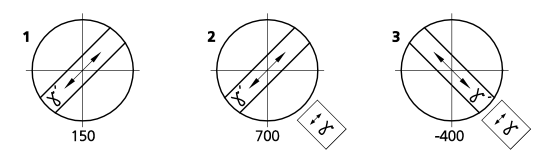


Fig. 4-10 Determination of the polarization direction $n_{\gamma'}$ in a synthetic fiber

(2) Instrumentation

- Eyepiece with reticle
- Strain-free objectives
- Rotary stage (Fig. 4-9/1)
- D polarizer (rotatable or fixed), if not using a condenser with an integrated polarizer
- ACR P&C analyzer module Pol for transmitted light in the reflector turret/slider or analyzer slider D fixed or with lambda plate
- Pol adjusting tool sample for polarization microscopy (453679-0000-000)

(3) Setting the microscope

- Adjust the microscope as described in section 4.2.1 (3) for transmitted light brightfield microscopy. Make sure the inter-pupillary distance is adjusted correctly on the binocular tube (see section 4.1.1).
- Center the rotary stage Pol (Fig. 4-9/1) and objectives (see sections 3.1.10.5 and 3.1.10.6).
- Swivel the polarizer (Fig. 4-9/3) into the beam path and position it to 0° if you are using a rotatable polarizer.
- Swivel in the analyzer module on the reflector turret (Fig. 4-9/2) (or insert the analyzer slider into the intermediate plate). Because of the crossed polarizers the field of view now appears dark.
- Place the Pol adjusting tool sample on the microscope stage and turn until the sample appears dark.
- Remove the analyzer from the beam path and align the reticle along the split cracks of the sample.

- Now insert the analyzer back into the beam path and remove the adjustment sample. The forward direction of the polarizer and analyzer is now parallel to the cross hair in the reticle (polarizer east-west, analyzer north-south).
- Turn the Pol rotary stage with the sample, e.g. a synthetic fiber, so that the sample reaches maximal darkness. The fiber is now parallel to one of the two lines in the cross hair orientation. If the deflection is significant (5° and more) you will need to recalibrate your microscope for polarization.

RF .	NOTE
	Do not change the eyepiece distance on the binocular tube any further to avoid shifting the angular position of the reticle to the fiber.

- Now turn the stage by approx. 45° until the longitudinal axis of the fiber is pointing in northeastsouthwest direction (Fig. 4-10/1). The sample now shows the strongest brightness (diagonal position). The sample can have any color in this position.
- Slide in the λ compensator.

Like the sample, the λ compensator is a birefringent object, but it has a defined path difference of 550 nm and a maximum oscillation direction n_{γ} pointing strongly to NE-SW.

When the compensator λ is put in, the sample changes its color depending on its orientation (NE-SW or NW-SE).

The changes in color are based on optical interference. It is necessary to compare the interference colors (path differences) in both diagonal positions (NE-SW and NW-SE).

The path difference results from the interference of the polarization of the sample and the polarization of the λ compensator.

The largest path difference occurs when the polarization direction of the sample or the absolute or relative largest index of refraction (n_{γ} or $n_{\gamma'}$) is parallel to the largest polarization direction of the λ compensator. The sample appears then e.g. in blue-green (Fig. 4-10/**2**).

The smallest path difference occurs when the direction of polarization of the sample with the absolute or relative smallest index of refraction (n_{α} or $n_{\alpha'}$) is perpendicular to the polarization direction of the compensator λ . The sample then appears e.g. yellow (Fig. 4-10/**3**).

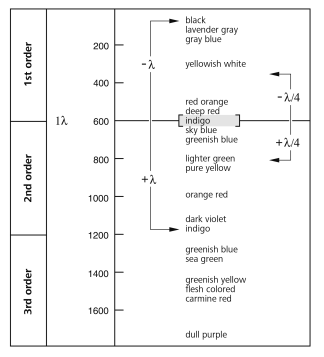


Fig. 4-11 Diagram of the Michel-Lévy color tables

(4) Conclusions

According to the Michel-Lévy color table (Fig. 4-11), the gray-white color which occurs in the bright position as discussed in the example above (Fig. 4-10/**1**) corresponds to a path difference of 150 nm.

After insertion of the λ compensator the surroundings of the artificial fiber which is not birefringent show a bright red which corresponds to a path difference of the compensator of 550 nm (the first order interference color for the path difference of 550 nm corresponds to 1 λ).

If the polarization direction of the birefringence sample to be investigated is parallel to the largest polarization direction of the compensator λ , i.e. in the NE-SW direction, the path difference of the sample (i.e. gray-white: 150 nm) and the path difference of the λ compensator (red: 550 nm) add up. This leads to a change in color of the sample from gray-white to blue-green (resulting path difference = 700 nm).

If the polarization direction of the birefringence sample to be investigated is perpendicular to the largest polarization direction of the λ compensator, i.e. in the NW-SE direction, the path difference of the sample (e.g. gray-white: 150 nm) is subtracted from the path difference of the λ compensator (red: 550 nm). This leads to a change in color of the sample from gray-white to orange (resulting path difference = 400 nm).

F	NOTE
	Michel-Lévy color tables can be ordered under literature catalog no. 42-312.

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4.2.6.3 Measuring path differences

Measuring compensators are required for the accurate measurement of path differences,. These compensators reset, i.e. compensate the path difference produced by the specimen to zero (first order black).

While the addition position as well as the subtraction position are of interest for the methods described above, **only** the subtraction position is relevant for measurements.

Path differences in the specimen can assume very small values (1/50 λ or 10 nm) and very large values (greater than 10 λ or approx. 5500 nm and more) and thus determine the compensator appropriate for the measurement.

To find the appropriate compensator, proceed as follows:

- Set the microscope as for transmitted light brightfield microscopy (see also section 4.2.1), taking particular care to set the inter-pupillary distance correctly on the binocular tube (see also section 3.1.4).
- Position the specimen feature to be examined directly above the center of the reticle.
- Restrict the size of the aperture to a value of about 0.2.
- Rotate the Pol rotary stage until the specimen is in extinction position, i.e. it appears **entirely dark**. In this position, activate the 45° click stop.
- Rotate the stage **once** (by 45°), so that the specimen is in diagonal position (sample becomes bright).

The intensity of the interference or color produced by the specimen allows the following conclusion to be drawn:

– If more or less strong interference colors appear on the object, the path difference ranges approximately between 1/2 λ and about 5 λ . The suitable compensator is:

Tilting compensator B 0-5 λ (accessory 000000-1115-700).

- If the object-side color changes from light gray/white to a strong interference color, when a λ compensator (473704-0000-000) is inserted in the compensator slot, the path difference is (1/4 ... 1/2) λ .

L	NOTE
	The occurrence of the color change effect may require the assessment of two specimen positions lying 90° apart. For this purpose, rotate the centered stage (by 2 click stops).

The suitable compensator is:

Tilting compensator B 0-5 λ (accessory 000000-1115-700).

- After insertion of the λ compensator and rotation of the specimen by 90° (2 click stops), the interference color remains to be white, in this case, however, it is a "higher-order white" and thus the path difference is > 5 λ .

The compensator appropriate for this application:

Tilting compensator K 0-30 λ (accessory 000000-1115-698).

- A dark gray appearing as the interference color suggests very small path differences (λ /10 or 54.6 nm).

The compensator appropriate for this application:

Rotary compensator Brace–Köhler λ/10 (Accessory 000000-1115-703).

• Push the compensator into the slot as far as it will go.

For measurement preparation and measurement procedure, use the enclosed operating instructions.

4.2.6.4 Circular polarization contrast

(1) Application

Unlike standard polarization contrast, circular polarization contrast does not show any dark (extinction) positions that depend on the angle of rotation (azimuth) of the specimen relative to polarizer or analyzer. This means that, while you are rotating the stage, the image will always look the same, as there are no bright and dark positions. With optical anisotropy all transparent specimens show the interference colors that are characteristic to them.

(2) Microscope configuration

- Strain-free objectives
- Pol rotary stage
- DF circular polarizer (no polarizers may be mounted on the condenser), including the corresponding $\lambda/4$ plate
- analyzer slider or D analyzer module Pol in the reflector turret

(3) Setting the microscope

- Set the microscope as for transmitted light brightfield microscopy using the KÖHLER method (see also section 4.2.1).
- Center the Pol rotary stage and objectives (if not yet done see sections 3.1.10 and 3.1.10.6).
- For the further settings, **do not** use a specimen for the time being.
- Place the analyzer in position.
- Swivel the lower part of the D circular polarizer (Fig. 4-12/3) into the light path up to the click stop and swing in the analyzer module on the reflector turret or insert the analyzer slider into the intermediate plate (if applicable). At full light intensity, assess the extinction (darkening) of the field of view without specimen.

If the extinction is not optimal, correct the vibration direction of the polarizer by slightly and carefully turning the polarizer mount using a small screwdriver (for that, use the adjusting slots (Fig. 4-12/4) on the bottom of the polarizer) until you obtain maximum extinction. Normally, however, this is not necessary because the vibration direction of the polarizer is factory-adjusted.

• Push the 6x20 mm slider with the $\lambda/4$ plate (Fig. 4-12/5) belonging to the polarizer into the compensator slot above the nosepiece as far as it will go.

- Then, swivel the top part of the D circular polarizer (Fig. 4-12/2) into the light path.
- Rotate the lever of the $\lambda/4$ plate of the D circular polarizer (Fig. 4-12/1) until obtaining maximum extinction (dark-gray field of view) (lever points 45° to the right).

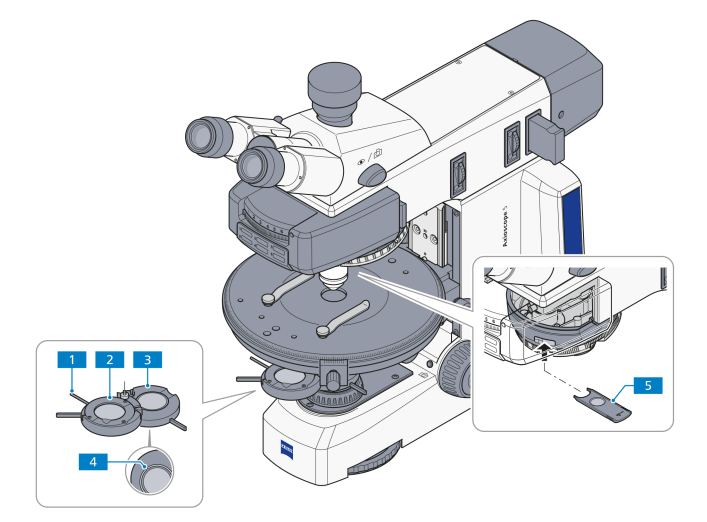


Fig. 4-12 Components for circular polarization contrast

- Do not view an anisotropic specimen unless the above adjustment has been done.
- Place the specimen to be examined on the stage.

The specimens appear constantly and independently of the stage rotation in their specific interference color, which depends on the material, specimen thickness and orientation.

F	NOTE
	To obtain a contrast-rich image at higher objective magnifications (from approx. 20x), reduce the illumination aperture to a value of between 0.15 and 0.20, i.e. close the aperture diaphragm accordingly. This creates an almost vertical (orthongonal) illumination.
	The effect of the $\lambda/4$ plate (Fig. 4-12/ 2) can be undone by either swiveling it out of the light path or turning it with lever (Fig. 4-12/ 1) into one of its two click-stop positions.

4.2.7 Setting transmitted light polarization for conoscopic observation - determining the optical character of crystals

For the classification (and thus identification) of crystalline matter, the examination of the interference image in the objective pupil delivers more valuable information than that obtained by viewing the specimen itself. The interference image becomes visible in the eyepiece if an additional optical system (fixed or focusing Bertrand lens or, in the basic version, the auxiliary microscope or diopter) is used.

In contrast to orthoscopy, this technique is called conoscopy, because here the specimen is ideally illuminated through a wide-open cone. In practical microscopic work, this means that the condenser front lens (0.9) must be in the light path, the aperture diaphragm fully open, and the objective, too, should be a high-aperture type.

4.2.7.1 Application

The determination of the optical character of transparent and weakly absorbent crystals is used to diagnose crystals. This method is also termed conoscopy.

Its main application is classical mineral microscopy. However, it also facilitates the identification and characterization of synthetic crystals, industrial minerals and plastics (e.g. films).

4.2.7.2 Simple conoscopy using the auxiliary microscope or the diopter

(1) Microscope configuration

- Recommended strain-free objectives:

N-Achroplan 50x/0.9 Pol objective or EC Plan-Neofluar 40x/0.9 Pol objective

- Axioscope stand for polarization with auxiliary microscope or diopter
- Pol rotary stage
- Pol binocular photo tube

eyepiece with crossline micrometer 14:140 (454060-0000-000) and

adjustment aid for polarization microscopy (453679-0000-000)

- Achromatic-aplanatic 0.9 H Pol condenser or 0.9 Pol condenser
- D polarizer (rotatable or fixed)
- analyzer slider or D Pol analyzer module in the reflector turret or reflector slider

(2) Setting the microscope for simple conoscopy using the diopter or the auxiliary microscope

- Set the microscope as for transmitted light brightfield microscopy using the KÖHLER method (see also section 4.2.1).
- Put the specimen onto the stage and focus on it.
- Move the polarizer and the analyzer into the light path.
- If you do not use the Pol binocular photo tube, carry out the following two steps:
 - first, align the crossline micrometer 14:140 or the eyepiece reticle to the vibration direction of the polarizer using the Pol adjustment aid. Please refer to section 4.2.6.2.
 - remove the Pol adjustment aid.
- Move a selected crystal to the center of the reticle. Only crystals above a defined size can be observed.
- Swivel in the front lens on the condenser, if necessary.
- For conoscopy of small crystals, close the luminous-field diaphragm, if necessary, to prevent the axial figure of the examined crystal from being superimposed by the axial figures of adjacent crystals.
- Swivel the 40x or 50x objective into the light path and focus with the focusing drive.
- After removing an eyepiece from the tube, you can view the axial figure in the eyepiece tube.
- For an improved assessment of the axial figure, you can (if available) insert a diopter or an auxiliary microscope in the eyepiece tube.

4.2.7.3 Conoscopy with Bertrand system module

(1) Microscope configuration

- Recommended strain-free objectives: N-Achroplan 50x/0.8 Pol objective or EC Plan-Neofluar 40x/0.9 Pol objective
- Axioscope stand for polarization with the Pol P&C Bertrand system module
- Pol rotary stage
- Pol binocular photo tube or eyepiece with crossline micrometer 14:140 (454060-0000-000) and adjustment aid for polarization microscopy (453679-0000-000)
- Achromatic-aplanatic 0.9 BF Pol condenser or 0.9 Pol condenser
- Bertrand system module in reflector turret
- D polarizer (rotatable or fixed)

(2) Setting the microscope for conoscopy using the Bertrand system module

- Set the microscope as for transmitted light brightfield microscopy using the KÖHLER method(see also section 4.2.1).
- Put the specimen onto the stage and focus on it.
- Move the polarizer into the light path.
- If you do not use the Pol binocular photo tube, carry out the next two steps:
 - Align the crossline micrometer 14:140 or the eyepiece reticle to the vibration direction of the polarizer using the Pol adjustment tool. Please refer to section 4.2.6.2.

NOTE NOTE

If you do not use the Pol binocular photo tube for your work, you must realign the crossline micrometer or eyepiece reticle after changing the inter-pupillary distance on the binocular tube.

- Remove the Pol adjustment tool.
- Put the specimen onto the stage and focus on it.

In the case of uniaxial crystals, the most favorable orientation for conoscopic viewing is obtained with those specimen features (e.g. of a thin section) that in orthoscopic viewing change the brightness as little as possible upon rotating the stage. In this case, the direction of viewing and the optical axis are \pm parallel. The same refers also to biaxial crystals, if they are viewed along or approximately in the direction of one of the two optical axes.

- Move a selected crystal to the center of the reticle.
- Swivel the front lens on the condenser into the light path.
- Swivel in the N-Achroplan 50x/0.8 Pol objective or the EC Plan-Neofluar 40x/0,9 Pol objective and focus with the focusing drive.
- Close the luminous-field diaphragm as much as is necessary to prevent the axial figure from being superimposed by the axial figures of adjacent crystals. The smallest crystal extension that can be masked out is 4 µm.
- Swivel in the Pol Bertrand system module on the reflector turret. Afterwards, the axial figure appears in the field of view.

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4.2.7.4 Conoscopy with intermediate plate and Bertrand lens slider

(1) Microscope configuration

- Recommended strain-free, high-quality objectives:
 - EC Plan-Neofluar 40x/0.9 Pol objective or
 - EC Plan-Neofluar 100x/1.30 Oil Pol objective or
 - EC Epiplan-Neofluar 50x/0.8 Pol objective or
 - EC Epiplan-Neofluar 100x/0.9 Pol objective
- Axioscope stand for polarization with intermediate plate
- Pol rotary stage
- Pol binocular photo tube
 - or

eyepiece with crossline micrometer 14:140 (454060-0000-000) and

adjustment aid for polarization microscopy (453679-0000-000)

- Achromatic-aplanatic 0.9 BF Pol condenser or 0.9 Pol condenser
- Bertrand lens slider in intermediate plate
- D Pol analyzer module in reflector turret
- D polarizer (rotatable or fixed)

(2) Setting the microscope for conoscopy using the intermediate plate and the Bertrand lens slider

- Set the microscope as for transmitted light brightfield microscopy using the KÖHLER method (see also section 4.2.1).
- Put the specimen onto the stage and focus on it.
- Move the polarizer and the D Pol analyzer module into the light path.
- First, align the crossline micrometer 14:140 or the eyepiece reticle to the vibration direction of the polarizer using the Pol adjusting tool sample. Please refer to section 4.2.6.2.

NOTE

If you do not use the Pol binocular photo tube for your work, you must realign the reticle after changing the inter-pupillary distance on the binocular tube.

- Remove the Pol adjustment tool.
- Put the specimen onto the stage and focus on it.

In the case of uniaxial crystals, the most favorable orientation for conoscopic viewing is obtained with those specimen features (e.g. of a thin section) that in orthoscopic viewing change the brightness as little as possible upon rotating the stage. In this case, the direction of viewing and the optical axis are \pm parallel. The same refers also to biaxial crystals, if they are viewed along or approximately in the direction of one of the two optical axes.

- Move a selected crystal to the center of the reticle.
- Swivel the front lens on the condenser into the light path.

- Swivel in the recommended objective and focus with the focusing drive.
- Close the luminous-field diaphragm as far as necessary to prevent the axial figures of adjacent crystals from being superimposed on the axial figure. The smallest crystal extension that can be masked out is 4 μm.
- Push the Bertrand lens slider incorporated in the intermediate plate into its active position. When you have done this, the axial figure appears in the field of view.
- Focus the axial figure by shifting the lever of the Bertrand lens slider.

4.2.7.5 Evaluation

Crystalline anisotropic specimens can be classified in optically uniaxial and biaxial specimens, each with an "optically positive" or "optically negative" character.

Uniaxial crystals show a **black cross** if the optical axis runs parallel to the viewing direction. **Depending on the degree of birefringence and the specimen thickness,** concentrically arranged colored **interference fringes (the so-called isochromates)** might appear (see also Fig. 4-13, second row).

The lines of this black cross remain closed when you rotate the stage. It can be located inside or outside the imaged objective pupil, depending on the position of the section.

With **optically biaxial** crystals, the cross resolves into two **dark hyperbola branches (the so-called isogyres) depending on stage rotation**, which are surrounded by colored interference patterns depending on the amount of birefringence and specimen thickness (suggestive of the figure "8").

When inserting a λ compensator (473704-0000-000) or $\lambda/4$ (473714-0000-000) or a 0-4 λ wedge compensator (000000-1140-663) into the compensator slot, with the initial state of the axial figure as illustrated in Fig. 4-13, the schematically shown changes in color (blue and yellow areas) will result in the axial figure, thus allowing the differentiation in "optically positive" and "optically negative".

	Optically uniaxial		iaxial Optically biaxial		
	Positive	Negative	Positive	Negative	
$\lambda \text{ plate}$ (white \rightarrow blue \rightarrow yellow)	++	+ - +		+	+ = Blue - = Yellow
Quartz wedge (Direction of movement when inserted)					Direction ofmovement
λ/4 plate (position of black spots)					

Fig. 4-13 Determining the optical character

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If the positions of the section are less favorable, i.e. the center of the cross of optically uniaxial or the isogyre of optically biaxial specimens is outside the objective pupil, the optical character can be assessed as follows:

- If the black isogyres are **straight** and run (related to the reticle) parallel through the pupil, the specimen is **optically uniaxial**.
- If the black isogyres are **curved lines**, traveling across the pupil on a circular path, the specimen is **optically biaxial**.

F	NOTE
	Axial figures can often be better represented with circular polarization. In particular, the axial angle of optically biaxial specimens (more or less the space between isogyres) can be determined more precisely and with much less ambiguity. The optical character can also be determined by inserting the λ compensator (6x20 mm) into the compensator slot.

4.3 Illumination and contrast methods in reflected light microscopy

4.3.1 Setting up reflected light brightfield microscopy using the KÖHLER method

(1) Application

Reflected light brightfield microscopy is the easiest and most commonly used RL-microscopy method. It is used to examine optically opaque specimens or samples as e.g. cut material or wafers.

For an imaging result which is as true to the specimen as possible, we must not only consider the socalled direct bundled beams but the indirect ones as well, i.e. the beams which diffract and scatter on the sample details. According to ABBE, the image is truer to the specimen when the fraction of the indirect bundled beams is larger.

The cone of light emerging from the reflected light illuminator is reflected on a color-neutral beam splitter before it passes through the objective which is focused on the specimen surface (so-called condenser function). The objective collects the light reflected on the specimen and creates, with the tube lens, the microscopic intermediate image. This image can then be examined visually or documented using a camera.

(2) Instrumentation

- Axioscope with pre-installed LED 10 W illuminator or optionally used with HAL 100 halogen lamp mounted at the back of the upper part of the stand
- ACR P&C brightfield reflector module for reflected light in the reflector turret
- Upper part of the stand with HAL 100/HBO 6-positions HD, DIC with aperture and field diaphragm sliders or upper part of the stand with HAL 100/HBO 6-positions HD, DIC and 2x 14x40 mm stop slider

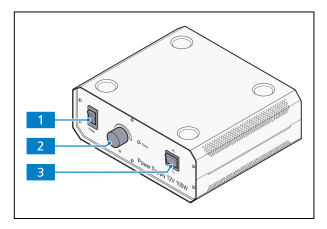


Fig. 4-14 External power supply unit for HAL 100

(3) Setting the reflected light brightfield

- The microscope is in the correct operational mode according to section 2.5.10.
- The microscope is switched on.
- When the halogen illuminator is used: Switch on the HAL 100 halogen illuminator on the external power supply unit (Fig. 4-14/**1**).
- Move the toggle switch (Fig. 4-14/3) to the **RL** position for reflected light.
- Adjust the light intensity with the control button on the external power supply unit (Fig. 4-14/**2**).
- Put a high-contrast reflected light sample on the microscope stage.
- Swing in the 10x objective on the nosepiece (Fig. 4-15/6).
- Focus the sample with the focusing drive (Fig. 4-15/5). Try to focus away from the sample to avoid any collision between the objective and sample.

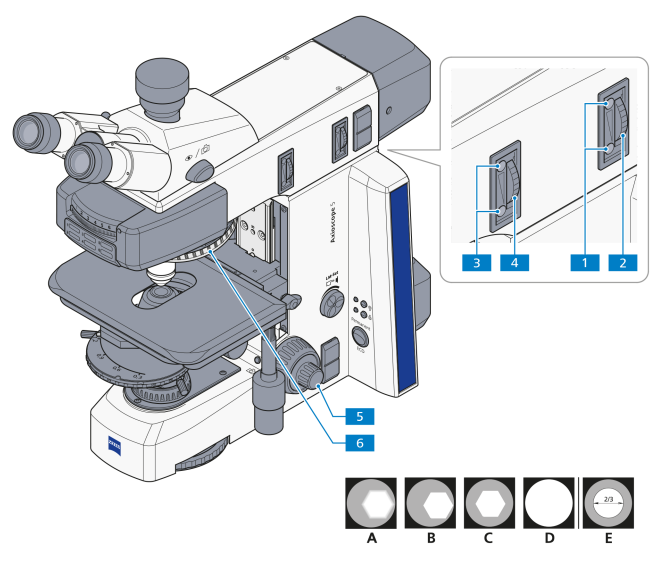
- Turn the knurled button of the aperture diaphragm (Fig. 4-15/2) to a medium position (approx. half open and half closed).
- Adjust the knurled knob on the field diaphragm (Fig. 4-15/4) so that the field diaphragm becomes visible in the field of view (Fig. 4-15/A).
- Readjust the focus on the edge of the field diaphragm with the focusing drive (Fig. 4-15/**B**) and center the field diaphragm on the edge of the field of view (Fig. 4-15/**C**) with the centering screws (Fig. 4-15/**3**).
- Now open the field diaphragm enough to make it disappear just past the edge of the field of view (Fig. 4-15/**D**).
- To adjust the aperture diaphragm (image contrast), remove an eyepiece from the tube barrel and look into the barrel or mount an auxiliary microscope instead of the eyepiece.
- Center the aperture diaphragm with the centering screws (Fig. 4-15/1) and adjust the knurled knob (Fig. 4-15/2) to about 2/3 to 4/5 of the diameter of the objective exit pupil (Fig. 4-15/E) for samples with a medium contrast quality.

In most cases this position of the aperture diaphragm gives the best contrast at almost full resolution, which is the best compromise for the human eye.

• Replace the eyepiece. Re-adjust the focus using the coarse and fine focusing drives and set the image brightness according to the reflected light sample. Readjust aperture stop diameter after each objective change.

NOTE
Never use the aperture diaphragm to regulate the image brightness! Use the control knob on the external power supply unit to adjust the light intensity!

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- Centering screw of the aperture diaphragm Knurled button of the aperture diaphragm 1
- 2
- Centering screw of the field diaphragm 3
- Knurled wheel of the field diaphragm 4
- 5 Focusing drive
- 6 Nosepiece

Setting the microscope in the reflected light brightfield Fig. 4-15

4.3.2 Setting the reflected light darkfield

(1) Application

The reflected light darkfield method is applied when samples are examined which do not have areas with different reflectivity (ideal brightfield samples), but which show deflections (as scratches, cracks, dust particles etc.) on the plane surface. All such light-scattering details appear bright in the darkfield while the reflective plane areas remain dark.

(2) Instrumentation

- Axioscope with pre-installed LED 10 W illuminator or optionally used with the HAL 100 halogen lamp mounted at the back of the upper part of the stand
- Objectives: Epiplan-Neofluar HD, EC Epiplan-Neofluar HD, Epiplan HD
- Darkfield ACR P&C reflector module for reflected light in the reflector turret/slider

(3) Setting the reflected light darkfield

- Adjust the microscope as described in section 4.3.1 for reflected light brightfield microscopy. The field diaphragm image should lie just barely outside of the edge of the field of view to avoid reflections.
- Swing the darkfield ACR P&C reflector module for reflected light on the reflector turret into the beam path.
- Remove the 6x20 mm compensator slider if mounted.
- Swing in the objective position with the darkfield objective (HD) on the nosepiece.
- If necessary, swivel in the darkfield reflector module on the reflector turret.
- Open the aperture diaphragm fully and switch off or remove neutral filters if applicable.
- Place the sample on the stage and adjust the focusing if necessary.

4.3.3 Setting reflected light DIC and reflected light C-DIC

(1) Application

The reflected light DIC and the reflected light C-DIC methods (DIC = Differential Interference Contrast; C-DIC = Differential Interference Contrast in circular polarized light) are used for the high-contrast imaging of small height differences on the surface of opaque specimen.

(2) Instrumentation

- Axioscope with pre-installed LED 10 W illuminator or optionally used with the HAL 100 halogen lamp mounted at the back of the upper part of the stand
- Objectives: EC Epiplan-Neofluar, Epiplan with the additional label "DIC" or "Pol"
- DIC slider, fitted to the objectives engraved on the top surface of the slider with amplification and aperture (in connection with the DIC P&C reflector module) or 6x20 mm C-DIC slider (in connection with the DIC P&C reflector module, compensator slot 6x20 mm)

(3) Reflected light DIC, preferably for objectives Epiplan 5x to 100x and LD-Epiplan 10x to 100x

- Adjust the microscope for reflected light brightfield microscopy as described in section 4.3.1. Open the field diaphragm enough so that the edge of the diaphragm lies just barely outside of the edge of the field of view to avoid reflections.
- Swing the DIC P&C reflector module on the reflector turret/slider into the beam path.
- Swing in the objective position with the DIC position on the nosepiece.
- Slide the DIC slider into the slot of the nosepiece (above the objective).
- Place the sample on the stage, then focus and turn the mechanical stage so the structure of interest is visible with maximum contrast.
- The contrast can be optimized with the knurled button on the DIC slider.

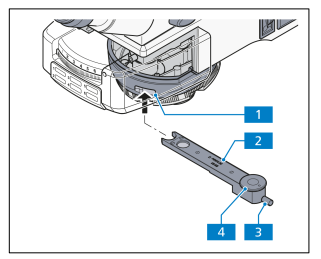


Fig. 4-16 Compensator slot 6x20 mm with C-DIC slider 6x20 mm

(4) Reflected light C-DIC

- Adjust the microscope for reflected light brightfield microscopy.
- Swing the C-DIC P&C reflector module into the beam path.
- Slide the C-DIC slider (Fig. 4-16/**2**) into the 6x20 mm compensator slot (Fig. 4-16/**1**).
- Place the sample on the stage, focus and turn the control knob (Fig. 4-16/**4**) on the C-DIC slider so the structure of interest is visible with maximum contrast. No further stage rotation is necessary.
- The contrast can be optimized by adjusting the control knob (Fig. 4-16/**3**) on the C-DIC slider.

4.3.4 Setting reflected light TIC

(1) Application

The reflected light TIC method (Micro-interferometry; TIC = Total Interference Contrast in the circular polarized light) is used in imaging and measuring specimen structures which are on hand in different azimuths.

(2) Instrumentation

- Axioscope with adjusted HAL 100 halogen illuminator mounted on the reflected light barrel
- Objectives: EC Epiplan-Neofluar, Epiplan with additional label "DIC" or "Pol"
- 6x20 mm compensator slot
- TIC slider 6x20 mm with appropriate C DIC P&C stepped reflector module

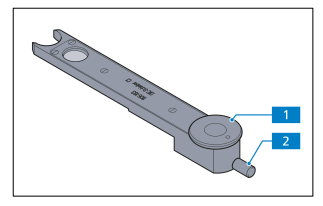


Fig. 4-17 TIC slider 6x20 mm

(3) Setting reflected light TIC

- Place the sample (e.g. a stepped object) on the stage and adjust the microscope for reflected light brightfield microscopy as described in section 4.3.1.
- Swing the C-DIC P&C reflector module on the reflector turret into the beam path.
- Slide the 6x20 mm TIC slider in the compensator slot 6x20 mm (Fig. 4-16/1). You will see chromatic interference stripes in the field of view. Using the control knob (Fig. 4-17/2) of the TIC slider, move the black interference stripe by sight to the middle of the field of view.
- To choose the structure to be measured, turn the knob (Fig. 4-17/1) on the TIC slider until the interference stripes are vertical to the direction in which the specimen is broken down (see Fig. 4-18). The interference stripes can be shifted with the aid of the control button (Fig. 4-17/2).

The step height SH is then determined with the following formula:

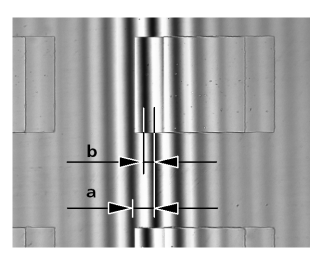
$$SH = \frac{n\Delta}{2} = \frac{\lambda b}{2a}$$

Where SH = step-height in nm

n = refractive index of the environment, mostly air (n = 1)

 Δ = phase difference

- a = distance between interference stripes
- b = offset of the interference stripes along the step
- λ = wave length of the illumination in nm



The values a and b (see Fig. 4-18) are determined with the aid of the eyepiece reticle micrometer or with the micrometer eyepiece.

If you are working with white light (without an interference filter), you must set $\lambda = 550$ nm. When interference filters are used, it is important to apply the focal point of their wave lengths.

The measured path difference depends on the aperture and increases with the illumination aperture.

Fig. 4-18 Interference stripes

Objective	Correction factor k
5x/0.15	1.0057
10x/0.25	1.0161
10x/0.30	1.0236
20x/0.4	1.0436
20x/0.50 and 50x/0.75	1.0718
50x/0.60	1.1111
50x/0.75 and 100x/0.75	1.2038
50x/0.80	1.2500
50x/0.90 and 100x/0.90	1.3929
100x/0.95	1.5241

The following correction values must be considered depending on the objective used:

Table 1: Correction depending on aperture

Example:

a = 11 mm	b = 5 mm
$\lambda = 550 \text{ nm}$	objective 20x/0.50

 $SH = \frac{\lambda \cdot b \cdot k}{2a} = \frac{550 \ nm \ \cdot \ 5 \ mm \ \cdot \ 1.0718}{22 \ mm} = 134 \ nm$

Attention:

- If the step and its surroundings are made from different materials, the phase jumps characteristic for the material must be considered. For all non-conducting materials, the phase jump is 180°, and for all semi-conductors only slightly different from 180°. Consequently, errors in the step-height determination may be neglected. However, if metals on top of glass are investigated, the results may become erroneous. The phase jumps given in table 2 were calculated for vertical light incidence and compact materials. They can serve as approximate values, since the phase jumps depend on the layer-thickness and the angle of incidence of the light. An accurate determination of the layer thickness is possible only when the complete specimen is covered with a homogeneous layer and the path differences are measured.
- If the layers and the steps are transparent, as with silicon dioxide on silicon, for example, the interference stripes can change their colors, so that the determination of the order of the interference may become problematic. This complication can be avoided if the sample is covered with a homogeneous layer.

Material	Phase jump φ
Copper	140.0°
Gold	142.5°
Silver	151.0°
Bismuth	151.0°
Nickel	157.0°
Iron	157.5°
Zinc	159.0°
Platinum	160.0°
Aluminum	160.0°
Tin	160.5°
Chrome	165.0°
Coal	160.0°
Graphite	165.0°
Silicon	177.0°
Glass	180.0°

Table 2:	Calculated phase jumps for
	compact material and
	vertical incidence of light

For a thickness measurement (step height), half the difference of the phase jump at the respective interface must be considered:

$$SH = \frac{\Delta}{2} - \frac{\delta\phi}{2}$$

Example: extreme case of copper on glass

 $\Phi_{\it copper}$ =140°, $\Phi_{\it glass}$ =180°, consequently, for the additional thickness due to the phase jump we obtain

$$\frac{\delta\phi}{2} = 20^\circ \text{ or } \frac{\lambda}{18} = 30 \ nm$$

Without consideration of the phase jump at the respective interfaces, the thickness value would be too large by 30 nm.

4.3.5 Setting reflected light polarization – proof of bireflectance and reflection Pleochroism

(1) Application

Reflected light polarization is a contrasting method suited for cut surfaces of mineral ore, coal, ceramics, special metals and alloys. Depending on the orientation of the crystals and the specimen details, the cut surfaces often react differently when reflected in linearly polarized light.

The illumination light is polarized by the polarizer before passing through the objective onto the specimen surface where it is reflected. Then the beam parts experience path differences depending on the structure and polarization of optical rotations which, when passing through the analyzer, are represented by different shades of gray. With the aid of a compensator with a λ -plate the gray contrast can be converted into a color contrast. **Caution! Do not use a 6x20 mm compensator for this conversion!**

Even when examining "dark" specimen surfaces, a rotatable $\lambda/4$ plate in front of the objective (antireflective cap) helps eliminate the reflections which are inevitable when working with objectives with very low magnification.

(2) Instrumentation

- Axioscope with pre-installed LED 10 W illuminator or optionally used with HAL 100 halogen lamp mounted at the back of the upper part of the stand
- Pol rotary stage
- Objectives: Epiplan-Neofluar Pol, EC Epiplan-Neofluar Pol, Epiplan Pol
- DIC P&C or DIC Rot I P&C reflector module in the reflector turret/slider or Pol P&C reflector module plus analyzer slider or analyzer slider plus polarizer slider

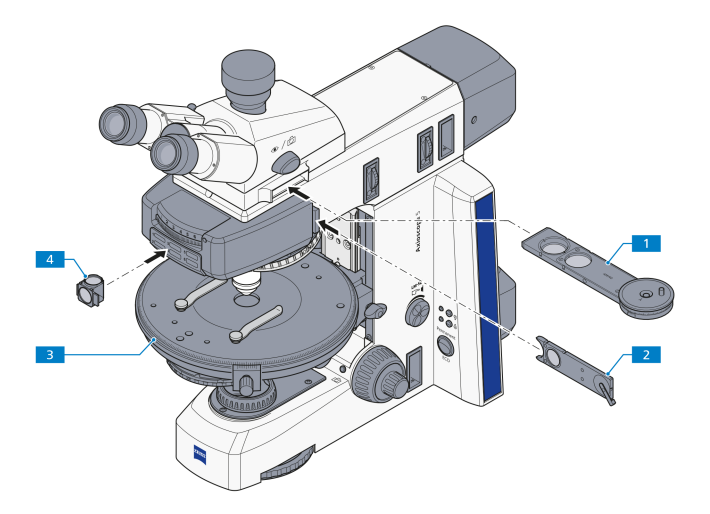
(3) Setting reflected light polarization

- Adjust the microscope for reflected light brightfield microscopy as described in section 4.3.1.
- If using the objective position with a DIC position, remove the DIC slider, if necessary.
- Swing the DIC P&C reflector module (Fig. 4-19/4) on the reflector turret into the beam path, swing in the Pol P&C reflector module and slide the analyzer slider into the compartment; or slide the analyzer slider (Fig. 4-19/1) and the polarizer slider (Fig. 4-19/2) into their compartments, if applicable. When using the fixed sliders, the polarizer must be aligned in the EAST-WEST direction and the analyzer in the NORTH-SOUTH direction.
- Place the sample on the stage, adjust the desired magnification, focus, and examine the sample in the polarization contrast which is now present while turning the Pol rotary stage (Fig. 4-19/3).

A sample is bireflectant when the sample details show differences in brightness and color which change when the stage rotates.

For samples with low bireflectance we recommend using the analyzer with a rotatable lambda plate.

Pleochroism is present when the color of the sample changes as soon as the stage rotates (overhead polarizer is turned on, analyzer is turned off).



- 1 Analyzer slider TL/RL
- 2 Polarizer slider RL
- **3** Rotary stage Pol
- 4 Reflector module in the reflector turret/slider

Fig. 4-19 Components for reflected light polarization

4.3.6 Setting up reflected light fluorescence

(1) General principle of operation

The reflected light fluorescence method is used to show fluorescent substances in typical fluorescent colors in high contrast. The light originating from a high-performance illuminator in a reflected light fluorescence microscope passes through a heat protection filter onto an excitation filter (bandpass). The filtered short-wave excitation radiation is reflected by a Dichroic Beam splitter and is focused on the sample through the objective. The sample absorbs the short-wave radiation before emitting longer-wave fluorescence radiation (Stokes' Law). This radiation is then captured from the image side by the objective and passes through the Dichroic Beam splitter. Last, the beams pass through a emission filter (longpass/bandpass) and only the long-wave radiation emitted by the sample passes.

The spectra of the excitation and the emission filter must match very closely. They must be inserted in a FL P&C reflector module together with the according Dichroic Beam splitter.

(2) Instrumentation

- Appropriate objectives, e.g. EC Plan-Neofluar or Fluar (UV-excitation)
- FL P&C reflector module and safety shutter in the upper part (additional shutter function with the filter slider is optional)
- HBO 100 or Colibri 3 or HXP 120 illuminator for excitation illumination
- Fluorescence protection shield

Image: Note The mercury vapor short arc lamp must be adjusted before starting the reflected light fluorescence method. Re-adjusting the adjustment might become necessary depending on the length of operating time.

(3) Setting up reflected light fluorescence

Setting the reflected light fluorescence is facilitated when starting with an objective of average degree of magnification, e.g. EC Plan-Neofluar 20x/0.50, and a sample of high fluorescence. The use of demonstration samples is also advisable for the start-up.

F	NOTE
	If necessary, remove the λ compensator used for the transmitted light polarization method from its compensator slot before setting the reflected light fluorescence.

- Slide the fluorescence protection shield in the compensator slot above the nosepiece.
- Switch on the objective EC Plan-Neofluar 20x/0.50.
- Turn the condenser turret to the BF transmitted light brightfield position first (or choose phase contrast if using a PhC objective) and go to the sample detail to be examined.

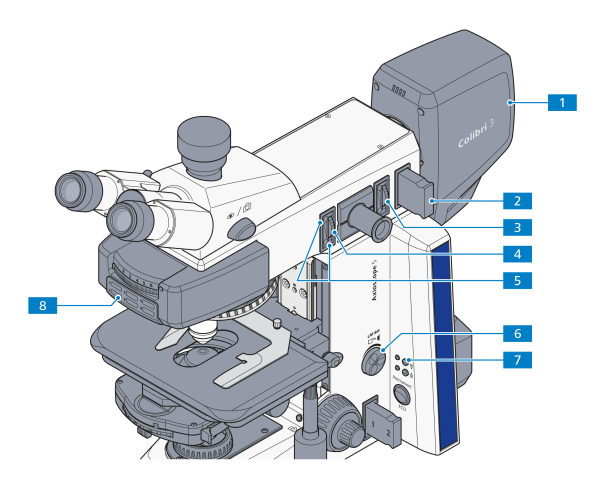


Fig. 4-20 Components for reflected light fluorescence

- Keep the light path in the reflected light illuminator blocked via the blocking position of the filter slider for reflected light (Fig. 4-20/2).
- Switch on the Colibri 3 illuminator (Fig. 4-20/1) by pushing the **RL** button (Fig. 4-20/7).
- Press the **Intensity/LM** button (Fig. 4-20/**6**) briefly for less than 1.5 seconds repeatedly to activate the required LED module or all LED modules of Colibri 3 together. The indicator light of the respective LED module on the Colibri 3 lights up when this module is switched on.

L	NOTE		
	If using the HBO 100 illuminator, switch on the external power supply and let it warm up to operational temperature for about 15 minutes.		

- On the reflector turret/slider (Fig. 4-20/8), choose the FL P&C reflector module with the desired fluorescence filter combination (depending on the excitation mode) and switch it on.
- Now unblock the light path in the reflected light illuminator with the filter slider for reflected light (Fig. 4-20/2).
- If necessary, adjust the FL attenuator to 100% transmission in order to facilitate locating fluorescence signals. Reduce the transmission later to preserve the sample.

- Remove an eyepiece from the tube and adjust the aperture diaphragm by sight.
- Open the aperture diaphragm (Fig. 4-20/3) enough to see the whole objective exit pupil.
- Replace the eyepiece in the tube and close the field diaphragm (Fig. 4-20/4) enough to make it visible in the field of view.
- Using both centering screws (Fig. 4-20/5), center the field diaphragm on the edge of the field of view.
- Open the field diaphragm enough to make it just disappear behind the edge of the field of view, or, if you are using a sample which might bleach out, reduce the field diaphragm for the field of view.
- Focus the sample again and optimize the collector position of the HBO 100 (if used). Adjust the collector so that the reflector module of the short-wave excitation illuminates the field of view evenly. A correction of the collector position is not necessary in modules with longer-wave excitation.

5 CARE, FUSE REPLACEMENT AND SERVICE

5.1 Instrument care

Maintenance of the Axioscope is limited to the following activities:

- Turn the instrument off after each use and cover it with the appropriate protection cover (dust and humidity protection).
- Do not use or keep the instrument in a humid environment (maximum humidity \leq 75%).
- Cover any open tubes with the dust protection caps.
- Remove dust or dirt from the visible optical surfaces by using a soft brush, blower brush, cotton swab, optical paper or cotton cloth.
- Remove water-soluble stains (coffee, soft drinks, etc.) with a dust free cotton cloth, which can be slightly moistened with water only or with a mild detergent diluted with water.
- Remove oil or grease stains (immersion oil, fingerprints) with a cotton swab or a dust- and lint-free free cotton cloth and the special optical cleanser L. This cleanser is made of 90% petroleum ether by volume and 10% isopropyl alcohol (IPA) by volume. The ingredients are also known by the following names: Petroleum ether: surgical spirit, benzine Isopropyl alcohol: 2-propyl alcohol, dimethylcarbinol, 2-hydroxypropane.

Clean the optical surfaces in a circular motion going from the middle towards the edge. Do not press too hard.



05/2019

ATTENTION

Do not use acetone to clean the lenses of the Pol condensers.

Please comply with the following instructions if you are going to use the microscope in a hot and humid climate:

• Keep the instrument in a bright, dry and well-aired room; humidity ≤ 75%; especially delicate parts like objectives and eyepieces should be kept in special dry closets.

Precision optical instruments may be damaged by mold if they are kept and used under the following conditions:

- If the relative humidity > 75% longer than three days at a temperature of between +15 $^\circ\!C$ and +35 $^\circ\!C$.
- If they are set up in dark rooms without sufficient ventilation.
- If there are dust and finger prints on optical surfaces.

5.2 Instrument maintenance

5.2.1 Checking the instrument

- Be sure to comply with the required voltage (e.g. on the power supply unit for HBO 100).
- Check the power cable and the power plug for damage.
- As soon as you become aware of any damage, turn off the instrument and secure. Call in a qualified professional to fix the problem.
- Check to make sure that the maximum operational hours of the mercury vapor short arc lamp are not exceeded.

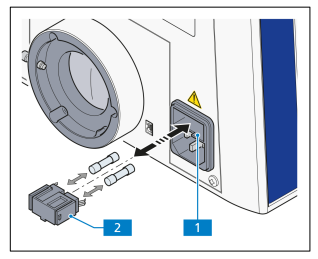


Fig. 5-1 Replacing the fuses in the stand

5.2.2 Replacing the fuses in the stand

CAUTION

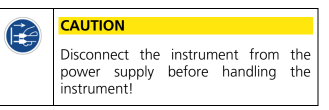
Always disconnect the instrument from the power supply before changing any fuses.

If fuses blow, the cause must be identified and any technical defect must be properly eliminated.

The fuse compartment is on the back of the microscope. It is combined with the plug and contains two fuses type **T 3.15 A/H /250 V**.

- Disconnect the plug.
- Pull the fuse retainer (Fig. 5-1/**2**) out to the front with the aid of a small screwdriver, if necessary.
- Remove the fuses from their container and replace them with new fuses.
- Push the fuse retainer back into the compartment (Fig. 5-1/1) to the stop.
- Reconnect the instrument to the power supply.

5.2.3 Replacing the fuses in the 12 VDC 100 W power supply unit



If fuses blow, the cause must be identified and any technical defect must be properly repaired.

The fuse compartment is on the back of the power supply unit and contains two fuses of the type **T 5 A/H 250 V, 5x20 mm**.

- Disconnect the power plug.
- Pull the fuse retainer (Fig. 5-2/**2**) out to the front with the aid of a small screwdriver, if necessary.
- Remove the fuses from their retainer and replace them with new fuses.
- Push the fuse retainer back into the compartment (Fig. 5-2/1) to the stop.
- Insert the power plug again.

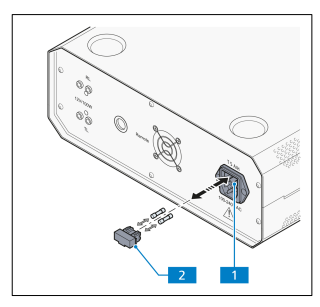


Fig. 5-2 Replacing fuses in the power supply unit

5.3 Troubleshooting

5.3.1 Microscope

Problem	Cause	Solution
No illumination light after switching on the microscope.	Nosepiece and/or reflector turret are not engaged to defined positions.	Move the nosepiece and/or reflector turret to the left or right to engage the nosepiece and/or reflector turret to defined positions. Then restart the microscope.
Shading or brightness irregularities in the field of view of the microscope; the field of	The vis/phot push-pull rod/shift knob on the photo tube is not in correct functional position (in-between position).	Move the vis/phot push-pull rod/shift knob to the correct functional position (end position).
view is not fully visible.	Nosepiece with objective is not fully engaged in its locking position.	Engage the nosepiece with the objective in its locking position.
	Condenser is not adjusted correctly.	Adjust the condenser correctly (adjustment, centering), see p. 101 ff.
	Aperture diaphragm is not adjusted correctly.	Adjust the aperture diaphragm correctly (centering, opening), see p. 101 ff.
	Field diaphragm is not adjusted correctly.	Adjust the field diaphragm correctly (centering, opening), see p. 101 ff.
	Filter is not correctly inserted in its slot.	Insert the filter correctly.
Low resolution and poor contrast.	Opening of the aperture diaphragm is not adjusted correctly.	Adjust the opening of the aperture diaphragm according to the 2/3-rule and the texture of the sample you are using, see p. 101 ff.
	Condenser is not focused correctly and front lens is not connected correctly.	Focus the condenser and turn the front lens correctly on or off, see p. 101 ff.
	Wrong thickness of the frame glass when using a transmitted light objective with a frame glass thickness correction of 0.17 mm.	Use standardized frame glasses with a thickness of 0.17 mm.
	Specimen holder is not inserted correctly.	Turn the specimen holder over, the sample side shows up.
	No immersion oil or an unspecified immersion oil is used with immersion objectives.	Use immersion oil 518 N or 518 F by ZEISS
	Air bubbles in the immersion oil.	Repeat the oiling procedure with fresh oil.
	Immersion oil on the front lens of a dry objective.	Clean the lens
	Correction setting is not set to the proper thickness of the frame glass.	Adjust the correction setting to the correct thickness of the frame glass
	Dirt or dust on the optical surfaces of objectives, eyepieces, condensers or filters.	Clean the soiled optical component

Problem	Cause	Solution
Parfocal performance not good on Axioscope 7	The focal plane was adjusted using the low-magnification objective which has a larger depth of focus than the high- magnification objective.	Determine the focal plane using the high-magnification objective.
	There is a backlash of Z-axis drive.	Adjust the focal plane from the same direction for all the nosepieces.
No light in eyepiece	The system is in ECO mode.	Turn the Intensity/LM knob clockwise to wake up the system.
	The light intensity is too low.	Turn the Intensity/LM knob clockwise to increase the light.
	The light was turned off by another pressing of the respective RL/TL button.	Press the RL or TL button according to the corresponding indicator in green color.
	LED connector is loose (when using built- in LED10 illumination).	Unmount the LED10 lamp case from the microscope stand, unplug and re- insert the connector to the socket. Check again.
	The reflector module is incorrectly installed or absent.	Check the reflect turret and make sure the correct reflector is in use.
	The field diaphragm is closed.	Check and, if necessary, open the field diaphragm.
XY stage stops at wrong position after initialization on Axioscope 7	The XY stage initialization failed.	Restart the microscope, if the issue still exists please contact ZEISS Service.
Cannot focus on the sample under high- magnification nosepiece with Axioscope 7	The Z-axis resolution was not configured with the magnification of nosepiece.	Configure the system with correct nosepiece information with MTB Configuration.
Asymmetric image sharpness, e.g. one side is sharp, one is	Condenser is not adjusted properly.	Re-adjust the condenser, see p. 101 ff.
side blurred.	Nosepiece is not engaged in its locking position.	Engage the nosepiece in its locking position (click-diaphragm).
	Sample is not fixed correctly on the mechanical stage.	Insert and fix the sample correctly in the specimen holder.
Distinct focus differences when changing the objective.	Focusable eyepieces are not adjusted correctly.	Adjust the focusable eyepieces according to the user's vision defect, see p. 99.
	Objective is not screwed in all the way.	Screw the objective in to the stop.
	Tube lens is not mounted, or it is mounted unnecessarily.	Mount the tube lens or remove it, according to the situation.

Problem	Cause	Solution
The left and the right field of view cannot be brought together in one image.	Distance of the eyepiece (distance of the pupils) is not adjusted correctly.	Re-adjust the distance of the eyepiece, see p. 99.
	Focusable eyepieces are not adjusted correctly.	Adjust focusable eyepieces according to the user's vision defect, see p. 99.
Using the microscope fatigues the eyes.	Distance of the eyepiece (distance of the pupils) is not adjusted correctly on the binocular tube.	Re-adjust the distance of the eyepiece, see p. 99.
	Focusable eyepieces are not properly adjusted.	Adjust the focusable eyepieces according to the user's vision defect, see p. 99.
	Image brightness is unacceptable.	Adjust the lamp voltage or insert a conversion filter.
	Binocular tube is misaligned optically, mechanically.	Call in service personnel for check-up/ repair.
Dirt or dust in the field of view.	Condenser is not focused properly and front lens is not in the correct on or off mode.	Focus the condenser and turn the front lens either on or off appropriately, see p. 101 ff.
	Opening of the aperture diaphragm is too small.	Adjust the opening of the aperture diaphragm according to the 2/3-rule or according to the texture of the sample, see p. 101 ff.
	Dirt of dust on optical surfaces of objectives, eyepieces, condensers, filters or samples.	Clean the optical surfaces of the soiled components. see p. 139.
Halogen lamp 12 V, 50 W does not glow although the switch in the "On" position.	Power plug is not plugged into the outlet.	Insert the plug into the outlet. Make sure outlet and instrument are adjusted to the correct voltage.
	Halogen lamp 12 V, 50 W is not mounted.	Insert halogen lamp 12 V, 50 W, see p. 68.
	Halogen lamp 12 V, 50 W is defective.	Exchange the halogen lamp 12 V, 50 W, see p. 68.
	Fuses are defective.	Exchange the fuses, see p. 140.
	The installed electrical equipment may be defective.	Call in the service personnel to check or exchange the components, if necessary, see p. 147.
	No voltage in the power socket.	Use another power socket.

Problem	Cause	Solution
Halogen lamp 12 V, 50 W flickers, illumination intensity is not stable.	Halogen lamp 12 V, 50 W is reaching the end of its life span.	Exchange the halogen lamp 12 V, 50 W, see p. 68.
	Power cable is not installed properly or is damaged.	Install the power cable properly or exchange it.
	The pins of the halogen lamp 12 V, 50 W are not properly inserted in the socket.	Insert the pins of the halogen lamp 12 V, 50 W correctly, see p. 68.

5.3.2 Axiocam 202/208

When the microscope is in use with Axiocam 202/208

Problem	Cause	Solution
LED indicator is off.	The camera is not drawing power via the USB (Commercial Micro-D) cable.	Make sure the microscope is powered on and connect the USB (Commercial Micro-D) cable to the stand.
	USB cable is not connected to a certified power supply unit.	Connect the camera to a running PC. Requirements: 5 V DC with at least 1000 mA at output.
	USB cable is not suitable.	Use the USB cable provided in the original package.
Firmware update does not function.	For an update, a USB stick must be inserted and the firmware to be updated must be saved to the root folder on the USB stick.	Insert a formatted USB stick with firmware in the root folder with at least 200 MB free memory space. Be sure to carefully follow the instructions enclosed with the firmware update.
Date/Time is wrong on the camera.	Date/time is not set correctly.	Date/time can be set in OSD under Operating System setting.
	The buffer battery has no charge.	Contact service
The image has severe noise.	The amplification (gain) is set too high.	Open 3D noise reduction; reduce the gain.
	The exposure time is set too high.	Open 3D noise reduction; reduce the exposure time.
	The light intensity is set too low.	Open 3D noise reduction; increase the light intensity.
The image is too dark or too bright.	Automatic exposure time has not been activated.	Activate auto exposure or manually adjust the exposure time and gain so that the settings are suitable for the current light situation.

Problem	Cause	Solution
The camera forgets the settings (e.g. manual white balance, resolution) if the power supply is disrupted.	Settings are not stored.	5 seconds is required for the settings to be automatically stored.
Monitor connected via HDMI does not display an image.	The camera is not delivering a signal, or signal is not compatible with the monitor.	Ensure that the camera has been switched on for at least 30 seconds and the LED indicator is green. Check the plug connections on the camera and monitor.
The camera cannot be recognized by the PC.	Camera is not recognized by Windows drive.	Make sure camera is powered on and re-connect the USB cable to the PC.
The image appears distorted on the full screen monitor.	The image aspect ratio is 16:9. The monitor may be set to a different aspect ratio and causes the distortion.	Set the monitor aspect ratio to 16:9.
The image is blurred on the screen but the sample is in focus through eyepieces.	Focus plane of the camera is different from that of the eyepieces.	Make sure the sample is focused correctly through eyepieces and adjust the camera adapter until image is in focus on the monitor.
		Or, if no adjustable adapter available, focus on monitor and adjust eyepiece focus through diopter setting.
The camera otherwise behaves abnormally.	The camera may have been put into an unintended state.	Press the factory reset button on the camera.

5.4 Maintenance and repair work

Maintenance and repair work on the mechanical, optical and electronic parts inside the instrument and on the electrical equipment of the Axioscope microscopes may be performed only by the ZEISS Service Department or specially **authorized** technical personnel.

To ensure that your microscope remains in optimal and faultless working order for the longest possible time, we recommend that you conclude a service/maintenance contract with ZEISS.

If you want to place an additional order or if a maintenance/repair incident occurs, please contact your ZEISS representative.

To place a maintenance/repair call, please contact the representative responsible for your area or the main quarters at

Carl Zeiss Microscopy GmbH Carl-Zeiss-Promenade 10 07745 Jena, Germany

microscopy@zeiss.com www.zeiss.com/microscopy



Carl Zeiss Suzhou Co., Ltd. Modern Industrial Square 3-B, No.333 XingPu Road SIP 215126 Suzhou, China

5.5 Firmware update

For updating the firmware, please use the following link for downloading the latest version:

https://www.zeiss.com/microscopy/us/downloads.html

6 ANNEX

6.1 List of abbreviations

AC Alternating Current ACR Automatic Component Recognition AF Width across flats Achromatic objectives with improved flatness of field (ICS optical products) A-Plan B/BF Bright field CCD Charge-coupled Device CSA Canadian Standards Association C-DIC Differential Interference Contrast in circular polarized light D/CGT Cover glass thickness D/DF Dark field d Diameter (e.g. of filters) Direct Current DC DIC **Differential Interference Contrast** DIN Deutsches Institut für Normung (German Institute for Standardization) ЕC **European Community** ЕC Enhanced Contrast ΕN Euronorm (European Standard) Ergo **Ergonomic/Ergonomics** FL Fluorescence foc. focusable GW Suitable for glasses wearers HAL Halogen lamp (illuminator) HBO Mercury Vapor Short Arc Lamp for Fluorescence (illuminator) HDMI High Definition Multimedia Interface ICS Infinity Color-Corrected System IEC International Electrotechnical Commission IP Internal Protection ISCP Infinity Space Camera Port ISO International Organization for Standardization LED Light Emitting Diode Light Manager LM Refractive Index for the D-line (sodium) n_D

- OSD On-screen display
- PCBA Printed Circuit Board Assembly

phot	photographic
Ph/PhC	Phase Contrast
PL	Plan
PlasDIC	Differential Interference Contrast for Plastic Receptacles
Pol	Polarization
P&C	Push&Click
R	Right-hand (coaxial knurled knob on the right-hand side of the mechanical stage)
RL	Reflected light
SLR	Single Lens Reflex
Т	Slow (type of fuse)
TIC	Total Interference contrast in circular polarized light
TL	Transmitted light
UL	Underwriter Laboratories
UV	Ultraviolet
VAC	Volt AC
vis	visual

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Bertrand lens	
Bertrand lens slider	
Bertrand system	
Bertrand system module	
Binocular photo tube	
Binocular tube	
Binocular tube part	
Birefringence	
Brightfield	
-	

C

, 22, 24, 26, 28, 30, 39, 65, 87, 101, 104

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Diaphragm	
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Ergo tube	
Ergonomics	
Evecups	
Eyeglass protection ring	
Eyepiece reticle	
Eyepieces	

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Field diaphragm Filter	
Filter holder	
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Ρ	
P&C Path difference Phase contrast Photo tube Pinhole diaphragm PlasDIC contrast Pol adjustment aid Polarization Polarization direction Polarizer	

Power supply

R

Reflected light Reflected light (RL) button	24, 26, 28, 30
Reflected light brightfield Reflected light fluorescence	
Reflected light illuminator	
Reflector insert	
Reflector module	
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6.3 Industrial property rights

The instruments, their components and the methods described in this manual are protected by the following patents:

- US 6154282
- US 6392796
- US 6595661
- US 7038847
- EP 1359453
- US 7046436
- CN 206479707 U

6.4 System overview

See next pages.

