

# Standard Pol and Standard WL Pol microscopes

## Operating Instructions

	Page
Standard Pol and WL Pol transmitted-light microscopes	4
Intermediate tube with Bertrand optics and diaphragm	6
Analyzers	7
Auxiliary objects and compensators	7
Assembly of the transmitted-light polarizing microscopes	9
Adjusting the transmitted-light microscope	16
Orthoscopic image	16
Conoscopic image	18
Objectives Pol Z, Pol eyepieces, total magnification	20
Standard Pol and WL Pol incident-light microscopes	21
Assembly of the incident-light polarizing microscopes	22
Adjusting the incident-light microscope	24
Epiplan Pol objectives	25
H-PI and H-Pr Pol reflectors	26

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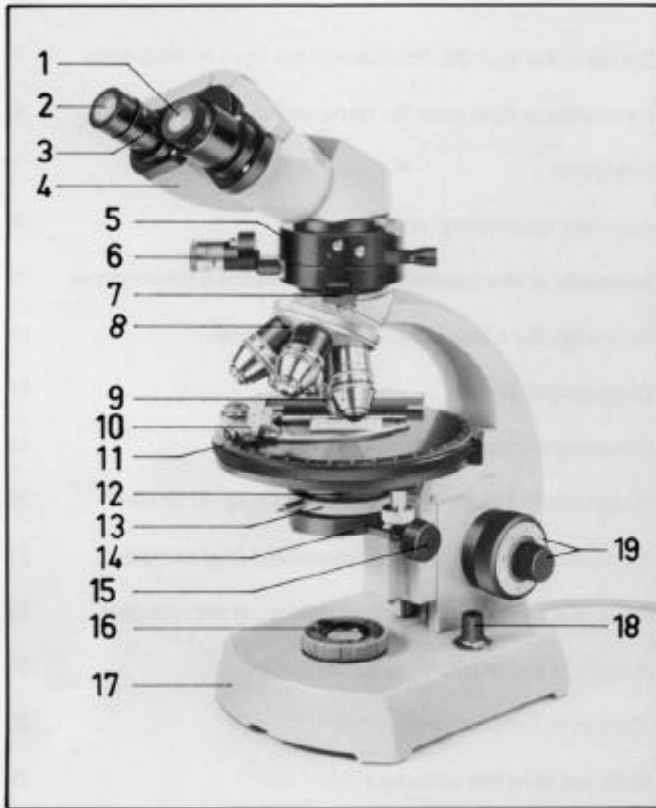
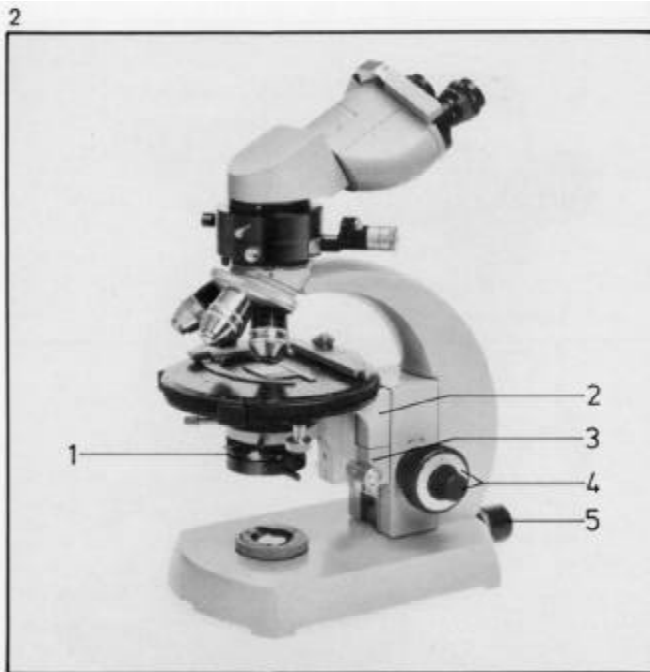


Fig. 1. Standard 18 Pol polarizing microscope (46 40 20) 1

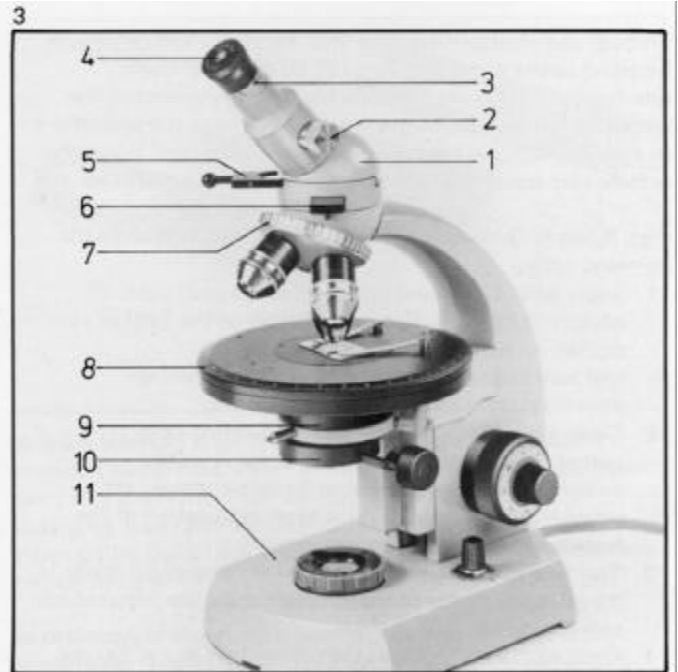
- 1 Ocular lens
- 2 Ocular tube
- 3 Binocular tube
- 4 Binocular tube S/30° Pol (47 30 36) with lever mechanism which keeps tube length and crossline orientation constant. The crossline angles are set to the user's D.
- 5 Alignment screw of Pol eyepiece engaged in a tube notch. When the alignment screw of the Pol eyepiece engages the horizontal notch in the edge of the left or right tube, the horizontal crossline bar will be aligned parallel with the oscillation direction of the polarizer (0° position/East-West); the vertical one parallel with that of the analyzer (90° position).
- 6 Nosepiece
- 7 Auxiliary object or compensator
- 8 Strain-free Pol Z objectives in centering mount
- 9 Attachable mechanical stage Pol (47 33 2b-9903) to move the specimen within a 30 x 40 mm area
- 10 Large total polarizing stage (47 34 00) with graduation, auxiliary clamp, and 100 µm scale and lockable total y rotation
- 11 Condenser 1.3 with swing-out front lens Z Pol (46 52 63) and screw-in auxiliary lens EL Pol (46 51 44)
- 12 Vertically adjustable condenser carrier with centering screws
- 13 Control for vertical condenser adjustment
- 14 Luminous-field iris diaphragm setting ring; holder for 32 mm dia. filters inside the ring
- 15 Standard base with in-base 6 V 10 W halogen illuminator and transformer
- 16 Power switch and brightness control of 6 V 10 W halogen illuminator
- 17 Coarse/fine focusing control; one interval of the fine control corresponds to 5 µm - 0.005 mm vertical stage movement.



**Fig. 2: Standard WL Pol microscope (49 15 59) for orthoscopic and conoscopic binocular observation**  
Microscope parts which are different for Standard WL Pol and Standard Pol (Fig. 1) microscopes:

- 1 Swing-in carrier with rotary polarizer and swing-out auxiliary lens (47 08 65-9902); the polarizer mount is graduated in 15° intervals with click stops at 0° and 90°. In normal position (click stop 0°) the oscillation direction is East-West
- 2 Attachable stage carrier (47 15 40) with rotary polarizing stage (47 34 66)
- 3 Attachable condenser carrier (47 15 58-9901) with centering screws
- 4 Coarse/fine focusing controls which act on the specimen stage independently of each other  
**Note: Before first use of the operating controls remove the transport lock by lifting the rack-and-pinion box with the aid of the coarse focusing controls and remove the plastic plate.**
- 5 6 V 15 W in-base illuminator

All optical components between polarizer and analyzer (condenser, objectives) contain strain-free lens elements and bear the red engraving "Pol"



**Fig. 3: Standard 16 Pol microscope for orthoscopic and conoscopic monocular observation**

- 1 Monocular polarizing tube (47 30 30-9905) with analyzer
- 2 Operating knob for Bertrand lens and diaphragm for conoscopic observation
- 3 Alignment screw of Pol eyepiece engaged in horizontal notch of tube edge (see 1.3, meaning Fig. 1, item 3)
- 4 Kpl crossline eyepiece 8x Pol (46 39 25)
- 5 Auxiliary object or compensator
- 6 Nosepiece with 5 Pol Z objectives
- 7 Rotary polarizing stage (47 34 64) with graduation, vernier and clamp  
Two stage clips (47 33 73) to secure the specimen
- 8 Condenser 0.9 Z Pol (46 52 62) with swing-out from lens and screw-in auxiliary lens EL Pol (46 51 44)
- 9 Non-rotary, swing-out polarizer on carrier (47 08 66); oscillation direction East-West
- 11 Stand base with in-base 6 V 10 W illuminator and in-base transformer

Microscope parts such as focusing controls (1 19) and knob (1.18) correspond to those of the Standard 18 Pol.

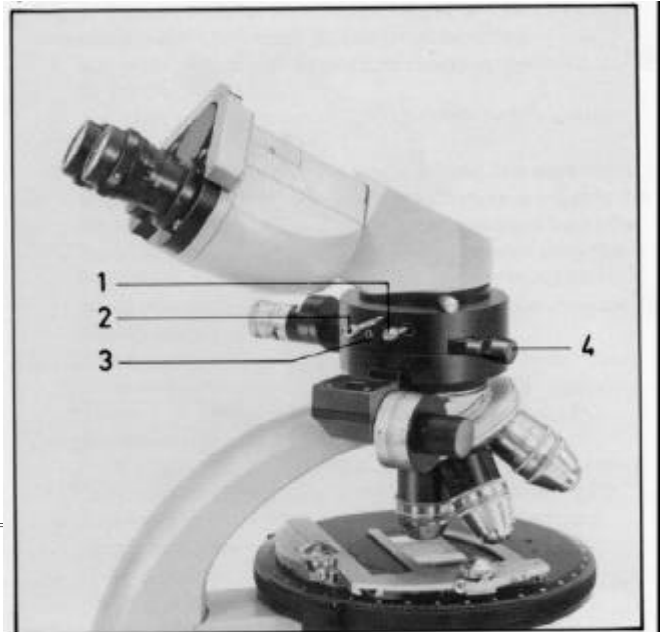
1) The 6- or 10-digit numbers in brackets are ordering numbers; they are imprinted on some items or modules.

Through the intermediate tube with focusable and centerable Bertrand optics and diaphragm (47 30 56) conoscopic interference figures are made visible in the eyepieces of the binocular Pol tube or images projected through the phototube to a photomicrographic or TV camera. Conoscopic image and orthoscopic surrounding field can be observed simultaneously.

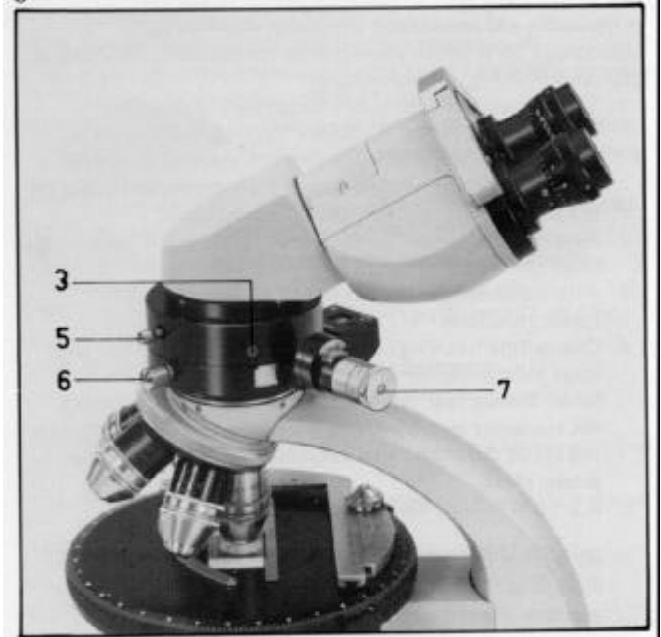
**Figs. 4 and 5: Operating controls of intermediate tube with Bertrand optics**

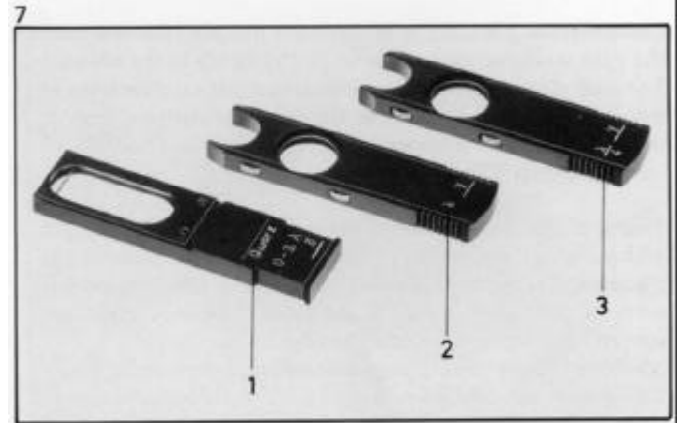
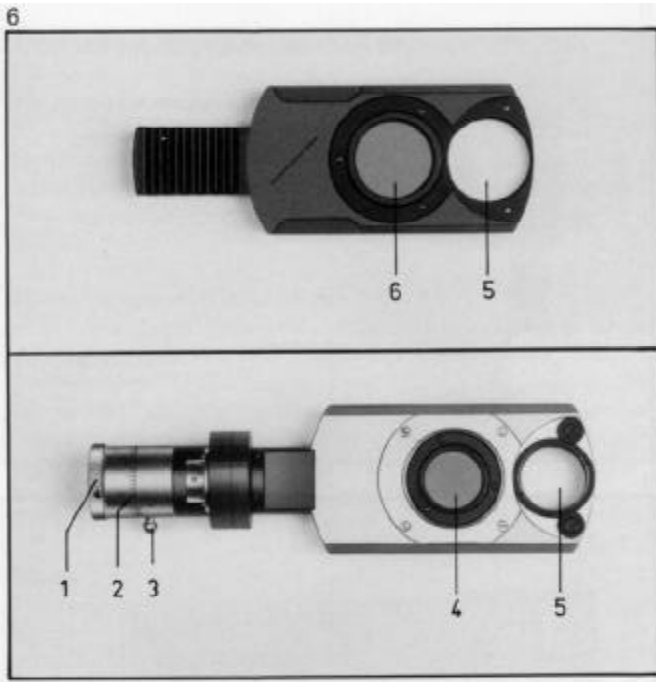
- 1 Slider with 3 Bertrand pinhole diaphragms to cut off objects of different size in the center of the field of view  
 pushed in: maximum diaphragm opening  
 half way in or out: medium diaphragm opening  
 pulled out: minimum diaphragm opening
- 2 Slider with diaphragm for the surrounding field  
 pushed in: only the conoscopic image is visible  
 pulled out: conoscopic and orthoscopic images are simultaneously visible, which eases localization of the ~~feature to be studied conoscopically~~
- 3 Two hex socket screws to center the conoscopic image in the crossline center of the eyepiece using the supplied hex socket wrench
- 4 Knob with scale to focus the conoscopic image. Certain focusing positions of the Bertrand optics can be marked with the aid of the scale
- 5 Slider to operate the Bertrand optical system  
 pushed in: Bertrand optics operative (conoscopic observation)  
 pulled out: Bertrand optics inoperative (orthoscopic observation). In this position slider (2) must also be pulled out to make the entire orthoscopic field of view visible.
- 6 Analyzer slider clamping screw prevents the slider from being pulled out accidentally and secures the set position of the slider
- 7 Analyzer

4



5





Aligned auxiliary objects and compensators are contained in sliders and are used either in the position of normal compensation or in the position of anomalous compensation. The auxiliary objects and compensators are held in place by a spring which remains in its slot.

Two different analyzers are contained in a slider. Analyzer (47 36 62-9903) (6.4) is rotary. Its angle of rotation can be read to within  $1/10^\circ$  on the  $360^\circ$  graduation with towards higher numbers, analyzer (6.4) rotates counterclockwise (mathematically positive). When set to  $90^\circ$  the oscillation direction of the analyzer (North-South) is thus crossed with that of the polarizer. Analyzer (47 36 63-9903) (6.6) is non-rotary and firmly set to  $90^\circ$  position. When the analyzer is pulled out of the beam path, a plane-parallel optical element in slider (6.5) takes its place in the beam path. It suppresses disturbing polarization effects due to beam-splitting or reflecting elements above the analyzer slider (analyzer effects), and avoids the simulation of the pleochroism which does not exist in the object.

fringent objects, i.e. to determine which of the refractive indices, the higher ( $n_\gamma'$ ) or the lower one ( $n_\alpha'$ ) belongs to which principal oscillation direction.

In **conoscopic observation** the sign of the optical character of the indicatrix of anisotropic objects (optically positive or negative) can be determined with the aid of auxiliary objects.

**Auxiliary object  $\lambda$**  (47 37 04) (7.2) enlarges or reduces by 550 nm the phase differences of the wave pairs coming from the object. **Auxiliary object  $\lambda/4$**  (47 37 14) (7.3) changes the phase difference by ca. 140 nm.

The auxiliary object has a NO-SW  $\gamma$ -direction. When the  $\gamma$ -direction of the object is parallel with the  $\gamma$ -direction of the auxiliary object, the two phase differences sum each other (addition position). The object will appear in a "higher" interference position.

subtract each other (subtraction position). The interference color of the object will be "lower" by the phase difference of the auxiliary object.

By means of the auxiliary objects, the interference colors of the object are shifted towards higher or lower orders towards higher or lower orders (addition or subtraction position).

**Compensators** are used to measure the phase differences of the light waves which are due to birefringence in the object. For such measurements the principal oscillation directions of the object must be diagonal to the crossed polarizers. For more details of the compensators see the relevant operating instructions (p. 27).

**Tilting compensators** feature either a birefringent plane plate which can be tilted about a horizontal axis, or two cemented plane-parallel plates in subtraction position. When measuring with the compensator B or E, the phase difference from the sum of both tilting angles can be directly read off the table contained in the corresponding operating instructions. The tilting angle can be read to within  $0.05^\circ$  on the graduated drum of the compensator.

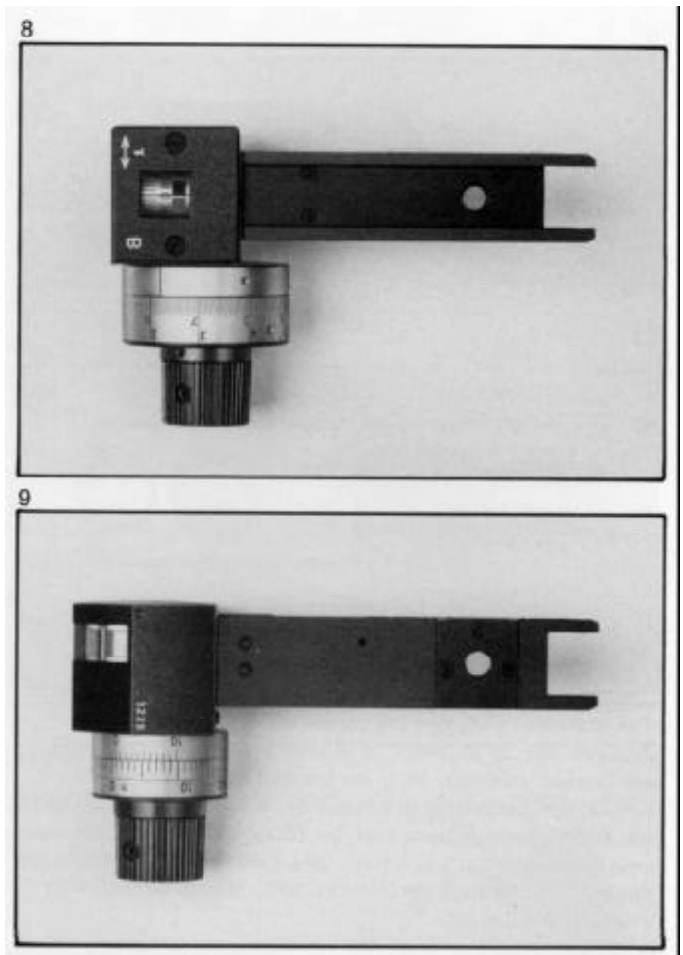
**Tilting compensator B (41 37 40) (Fig. 8)**  
to measure phase differences up to ca. 5 orders ( $5 \lambda$ )

**Tilting compensator E (41 37 41)**  
to measure phase differences up to ca. 20 orders ( $20 \lambda$ )

**Rotary compensators for small phase differences** contain a birefringent plane plate which can be rotated about the microscope axis. Angles of rotation can be read to within  $0.1^\circ$  on the knob with graduation and vernier.

**Brace-Köhler rotary compensator (41 37 31) (Fig. 9)**  
to measure phase differences up to  $\lambda/10$ . The calibration values for the wavelengths of the lines C', D, e and F' are given in the operating instructions of each compensator (designated by the number of the compensator).

**$\lambda/4$  plate (47 37 18) for Sénarmont compensation** acts in conjunction with the rotary analyzer (47 36 62-9903). Its optical element is a quarterwave plate whose principal oscillation directions are parallel with those of polarizer or analyzer. To compensate the phase difference in the object the analyzer is turned by a certain amount and the angle of rotation read off. The measuring range is  $1 \lambda$  (see operating instructions G 41-510/I).



## Exclusive microscope parts of Standard WL Pol

### Mounting condenser carrier (10.2)

Flick up lever (10.3). Place the right guide ledge of the carrier against the rail (10.1). Swing the left side in until the spring bolt snaps in behind the guide ledge. Lower condenser carrier as far as it will go. Flick down lever and tighten it slightly.

### Mounting specimen stage on change rail

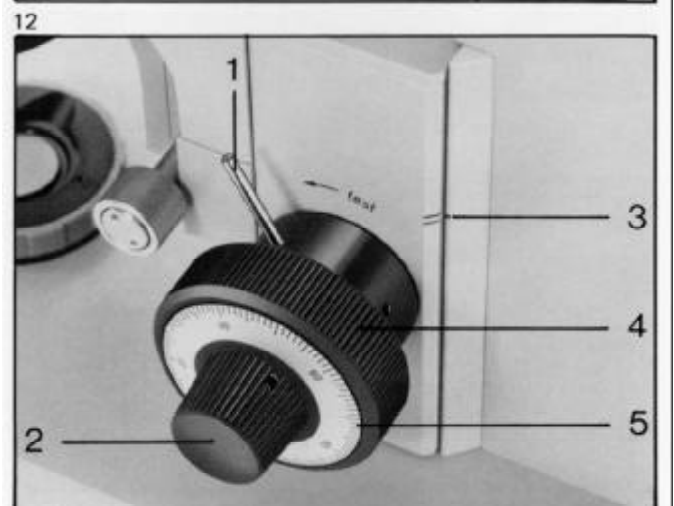
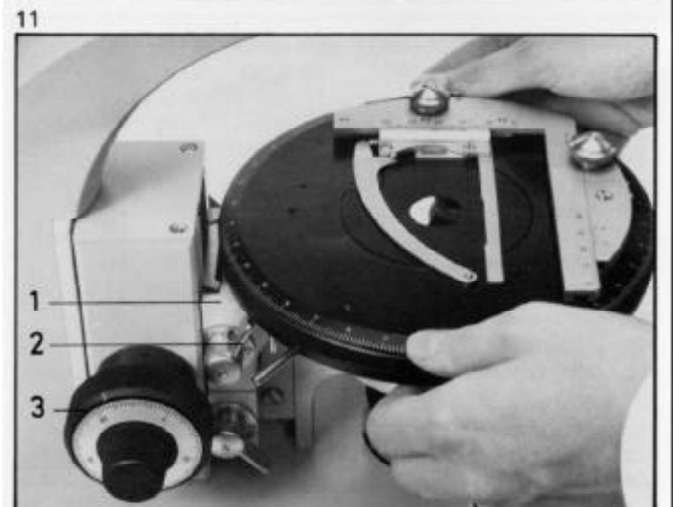
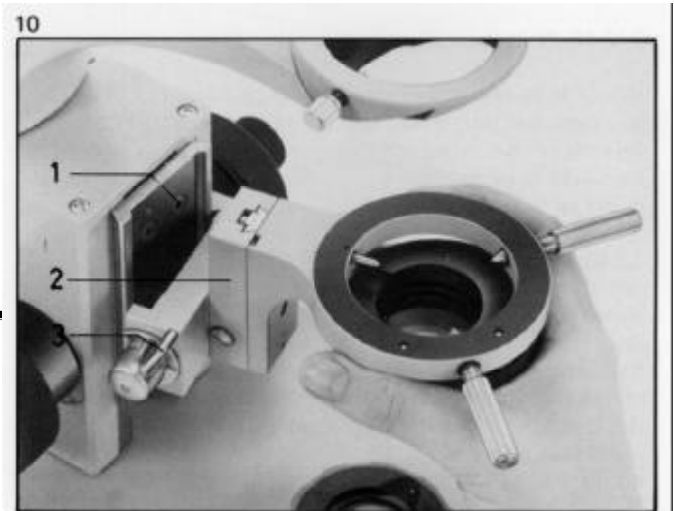
it will go.

Flick up lever (11.2) of stage carrier (11.1). Insert stage carrier from above in the change rail: let engage first the lower right projection, then the left spring bolt and at last the upper right projection. While doing so press the stage carrier against the change rail and lower it as far as the condenser carrier. Flick down lever.

### Coarse/fine focusing control

It acts on the stage carrier. The motion of the coarse focusing control (12.4) can be stiffened or smoothed. Plug the supplied metal pin (12.1) into the borehole and turn it. The motion is stiffened when turning it in the direction of the arrow. Set fine focusing control (12.2) to a medium working range by turning the fine focusing control until dot (12.3) is embraced by the two index lines. Focus on the specimen first with the coarse focusing control (12.4) then with the fine focusing control; the fine focusing control will thus have the same clearance in either direction.

One full turn of the fine focusing control corresponds to a vertical stage movement of 0.1 mm (1 interval of its graduation corresponds to 0.002 mm = 2  $\mu$ m).



## 6 V 15 W in-base illuminator

Hold 6 V 15 W filament lamp (38 00 18-1740) (13.1) with a soft cloth and plug it into socket (46 80 10-9904) (13.2) so that the red dot is opposite the red pin. Push lamp in, turn it clockwise as far as it will go and let it snap in. Remove fingerprints on the bulb.

Turn clamping ring (14.2) until its red dot is opposite the red dot of the tube (46 70 50) (14.1). Slide filament lamp in socket (14.3) into collector tube and secure with clamping ring

on the transformer corresponds to the local mains voltage. If this is not the case, set the right voltage or call our maintenance service.

Connect 6 V 15 W lamp to the mains by cable via transformer. Mains cable socket (15.3).

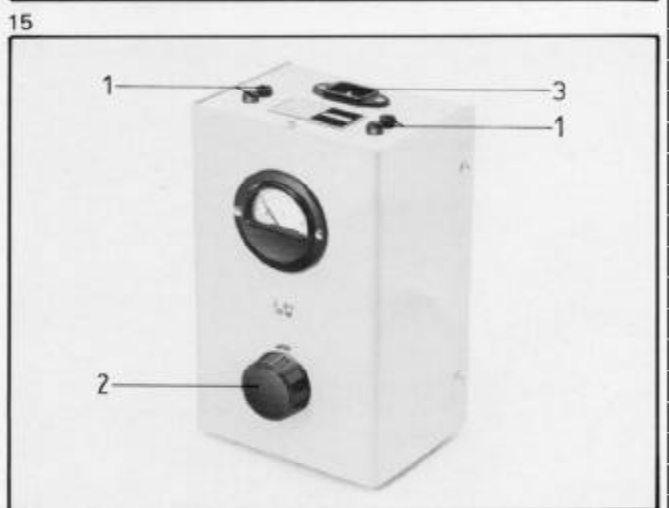
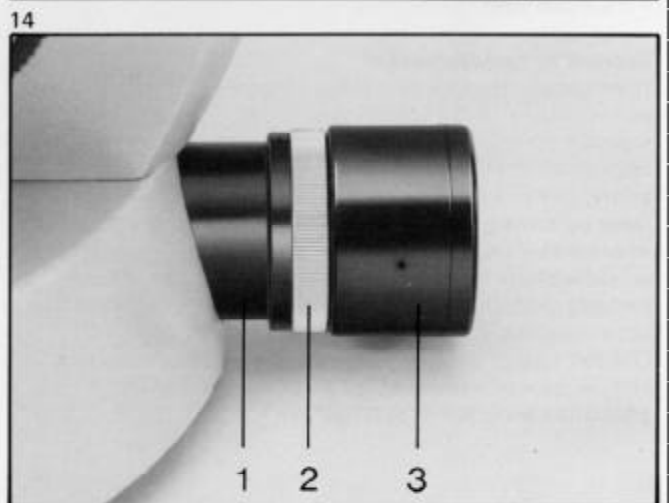
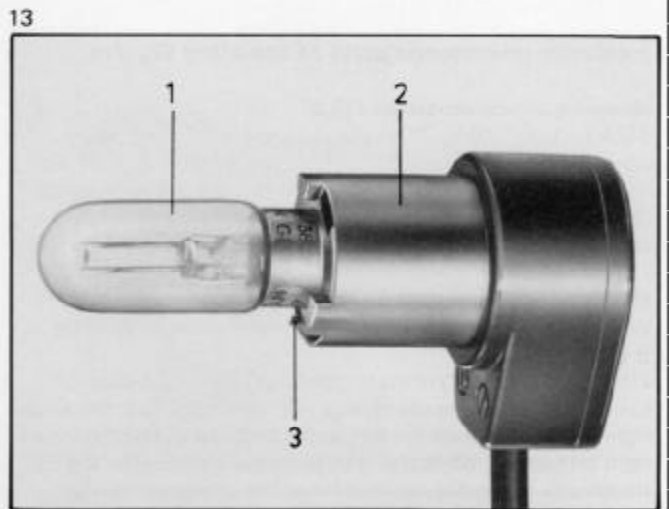
**Transformer (39 25 24-9903) with voltmeter (Fig. 15)** 100-110-115-127-220-240/2 ... 8 V/50 ... 60 Hz, power consumption 30 VA, sockets (15.1) for two 6 V 15 W illuminators.

Switch on transformer with control (15.2) and with the same control adjust the brightness.

It is generally sufficient to run the filament lamp at under-voltage which increases its life. Run the lamp at overvoltage only for short periods.

Instead of transformer (39 25 24-9903) (Fig. 15), step-up transformer (39 25 64-9903) can be used: adjusted for 100-127-220-240/3-4-5-6-7-8 V/50 ... 60 Hz, power consumption 25 VA, with CEE plug. The secondary voltage can be set in steps of 1 V. Only a 6 V 15 W lamp can be connected to its

of knob.



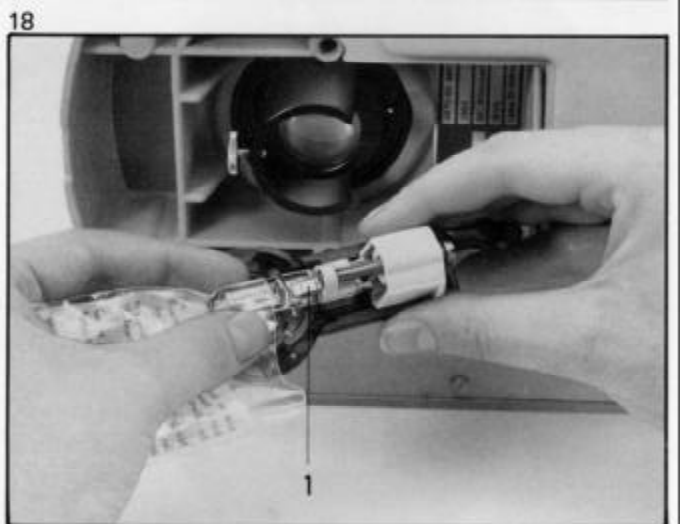
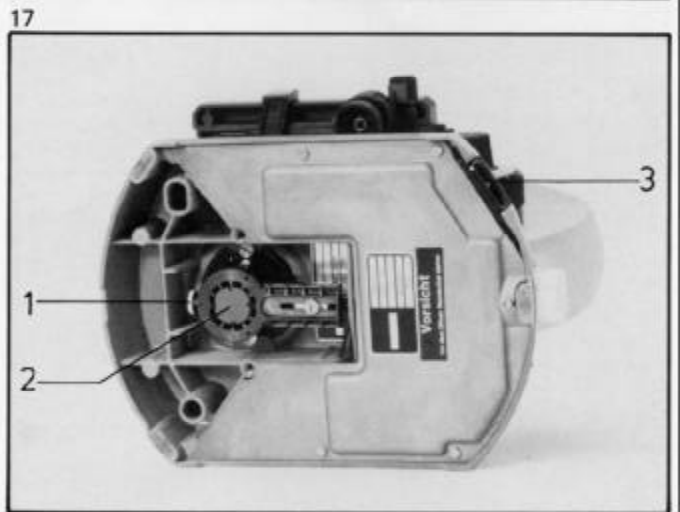
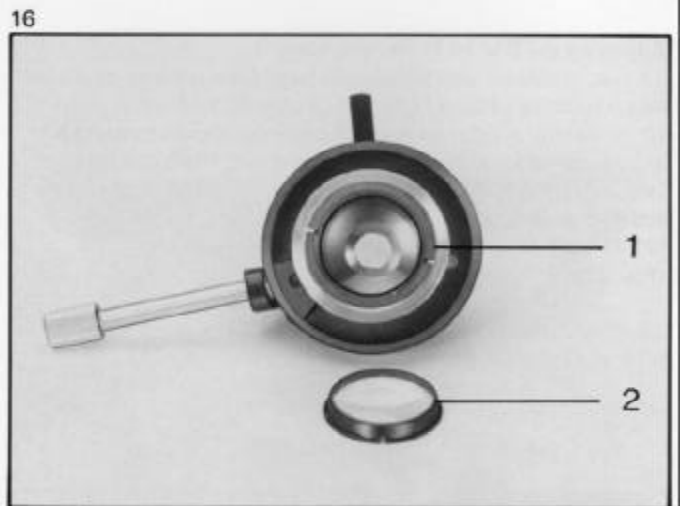
**6 V 10 W in-base halogen illuminator**  
for Standard microscopes only

Screw auxiliary lens EL Pol (46 51 44) (16.2) from below into condenser (16.1).

Socket (17.3) for mains cable. In-base transformer for 220/9 V, 50 ... 60 Hz, max. power consumption 18 VA, protection class III complying with VDE regulations or in-base transformer for 120/9 V, 50 ... 60 Hz, UL-listed.

**Fitting the 6 V 10 W halogen lamp**

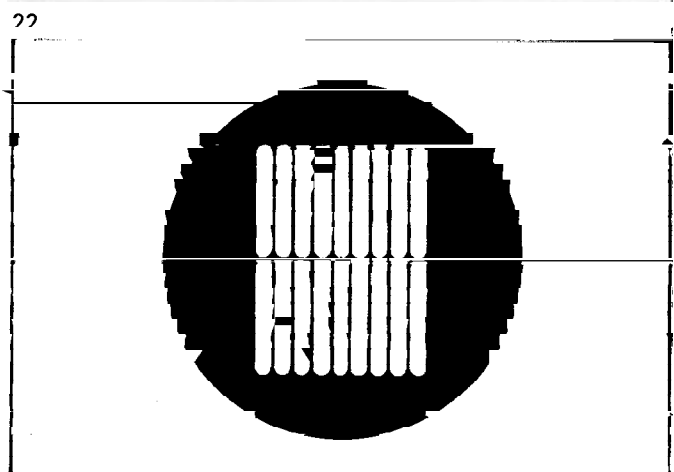
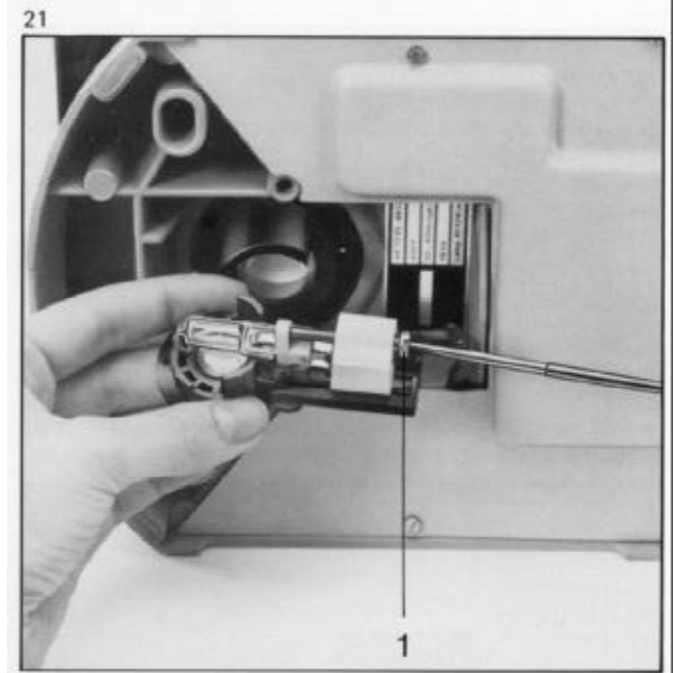
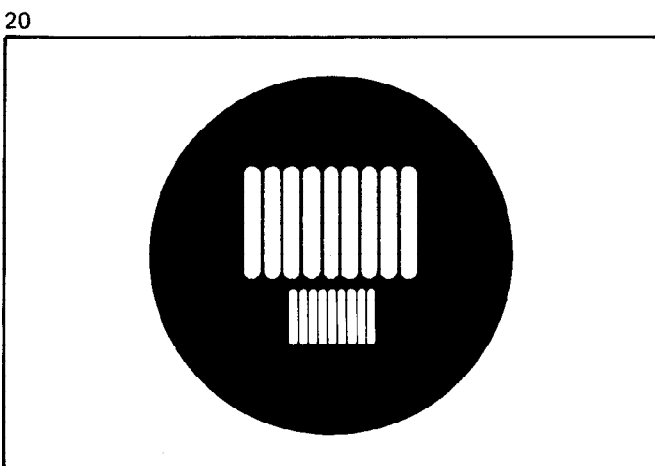
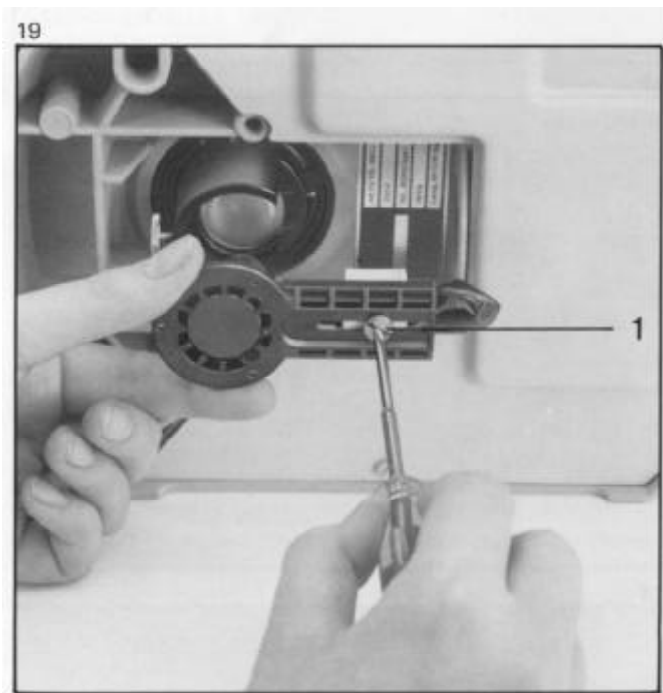
Put microscope on the side on a table. Loosen knurled screw (17.1) and pull out lamp socket (17.2). Pull defective lamp out of the two metal clamps (18.1). Hold new halogen lamp on plastic bag in which it is supplied and plug it into the



### Adjusting the 6 V 10 W halogen lamp

To ensure correct adjustment the lamp filament and its mirror image must be of equal size and lie next to each other. While watching lamp and concave mirror: loosen screw (19.1) so that the lamp socket can just be moved. Move socket with lamp until the lamp filament and its mirror image are next to each other (Fig. 20). Tighten screw (19.1). Then turn screw (21.1) until lamp filament and its mirror are of equal size (Fig. 22).

After adjustment insert lamp housing in diaphragm insert with ground glass and secure with knurled screw (17.1)



## Microscope parts of Standard Pol and WL Pol

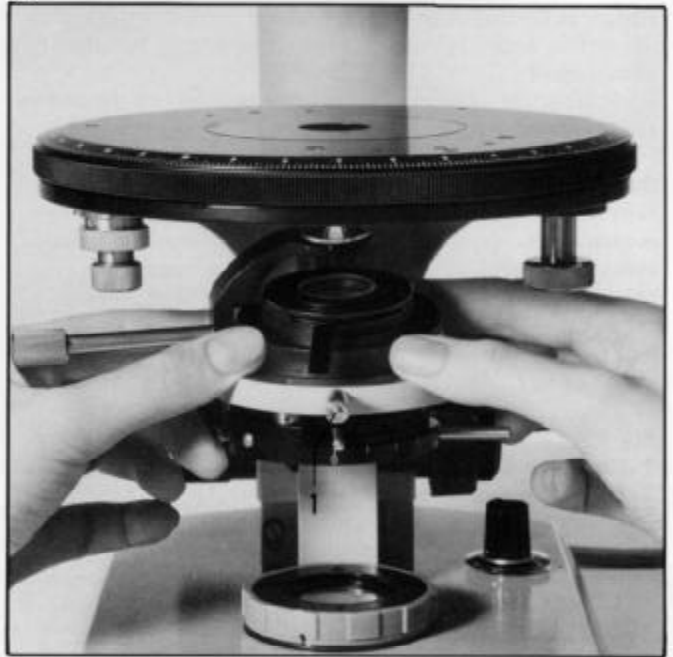
### Mounting condenser (Fig. 23)

With coarse focusing control rack up microscope stage as far as it will go and with control (1.15) set condenser carrier to lowest position to provide ample space for insertion of the condenser. Press dovetail ring of condenser against spring bolt (23.1) and insert the condenser, rack it up to topmost position with control (1.15).

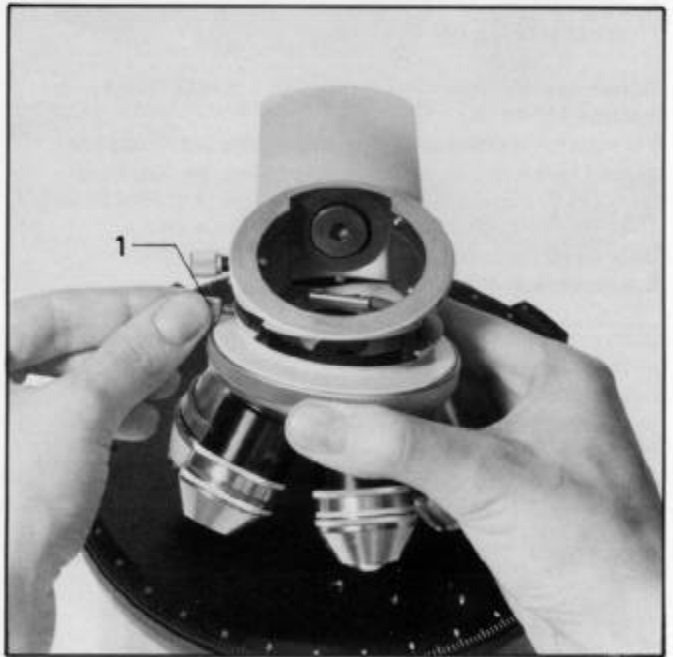
### Mounting glide nosepiece on stand head with glide mount (Fig. 24)

Attach glide nosepiece with objectives slightly tilted so that at first ca. 2/3 of the rail with the loosened clamping screw (24.1) engage the guide. Swing the other guide rail up and slide the glide nosepiece into the glide mount as far as it will go. To move the glide nosepiece pull out the spring locking pin at the end of the knurled screw after having slightly loosened clamping screw (24.1). Then tighten clamping screw again.

23



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### Rotary polarizing stage

The rotary polarizing stage is firmly mounted on Standard Pol microscopes.

On Standard WL Pol microscopes it is mounted on the change rail by way of a stage carrier (47 15 40) (25.5).

### Rotary polarizing stage (47 34 64) (3.8)

runs in precision ball bearing; dia. of stage area 150 mm; removable reducing plate. The stage features bores and tapped holes for: stage clips, attachable mechanical stage Pol (Fig. 26), heating and cooling cell (47 80 13) and universal rotary stage. Its 360° graduation with vernier allows measurements of the angle of rotation to within 1/10°. The stage can be fixed in any desired position.

**Fig. 25: Large rotary polarizing stage (47 34 66)**

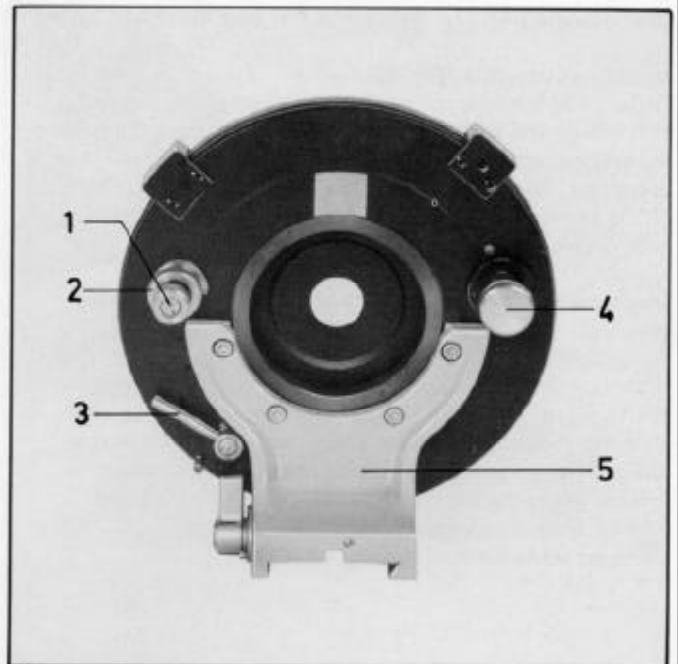
- 1 Control for stage rotation
- 2 Knob to engage and disengage control (1)
- 3 Lever to clamp the stage rotation
- 4 Knob to operate 45° click-stop system to change quickly between parallel and diagonal positions when working with auxiliary objects or compensators. It is operative when pulled out in any desired position, inoperative when snapped in at a click stop.
- 5 Stage carrier (47 15 40) to attach the specimen stage to the change rail (see Fig. 11).

### Attachable mechanical stage Pol (47 33 25-9903) and point counter (47 40 35)

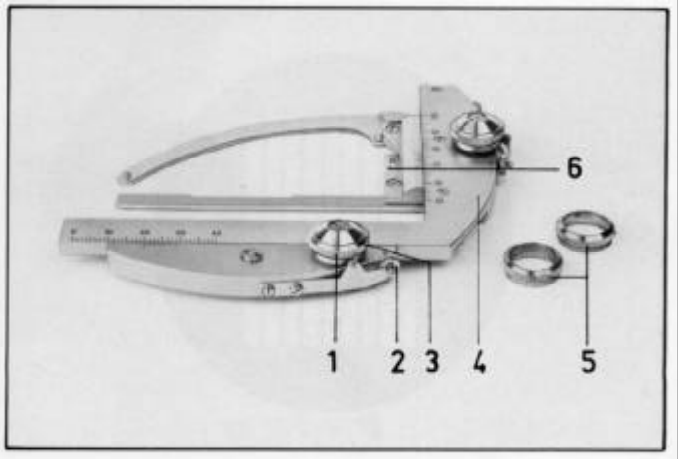
To mount the attachable mechanical stage (26.4) use one tapped hole and 2 bores of the stage plate. By operating lever (26.6) standard-sized specimen slides (e.g. 76 x 26 mm, DIN 58 884 or 46 x 26 mm) can be moved within a 30 x 40 mm area and coordinates determined to within 0.1 mm within this area.

With the **point counter** 0.2 or 0.3 mm intervals are marked as the specimen is moved. Pulling out knurled knob (26.2) overrides the click stop. To mount the click-stop rings, loosen the three screws (26.1), take off ring (26.3) and fix click-stop rings (26.5).

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27



#### Fitting tubes (Fig. 27)

The viewing tube is fitted either directly on the stand head or above the intermediate tube (Fig. 27) with its dovetails. The ring dovetails feature two 180° displaced notches (28.1) into which spring bolt (28.2) or (27.1) snaps in.

Tilt tube slightly, press its ring dovetail notch against the spring bolt of the loosened clamping screw (27.1) of the intermediate tube, and put on the entire ring dovetails until they snap in. Tighten clamping screw (27.1).

Mounting of the viewing tube either as shown in Fig. 1 or according to Fig. 2.

#### Mounting intermediate tube on stand (Fig. 28)

As with the polarizing tube press down spring bolt (28.2) with one ring dovetail notch of intermediate tube (28.1), mount the tube completely and secure it with clamping screw.

The intermediate tube is correctly aligned when the slide-in direction of the analyzer slider (30.1) is SW-NO seen through the binocular polarizing tube. The analyzer is then aligned North-South in 90° position.

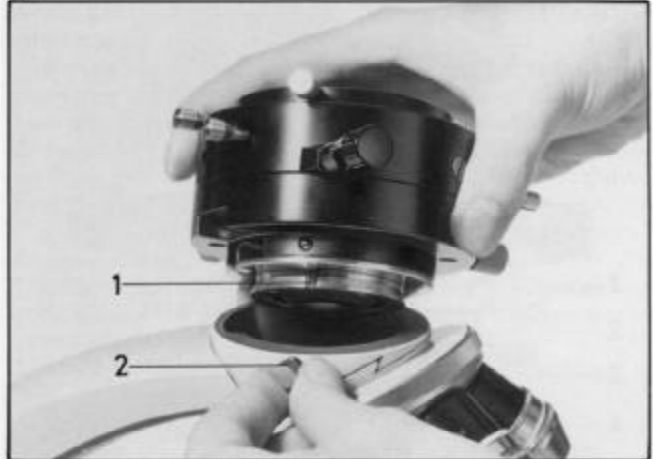
#### Focusing the eyepiece on the cross hairs

Look on a bright surface through Pol eyepiece (29.1) with reticle. Observe the cross hairs with relaxed eye. Turn diopter setting ring at first away from you and then towards you until the cross hairs are just in focus. Equip one of the tubes with Kpl eyepiece Pol (29.1) with reticle so that the alignment screw engages one of the two notches of the tube edge. Equip the other tube with a Kpl eyepiece with focusing eyelens (foc).

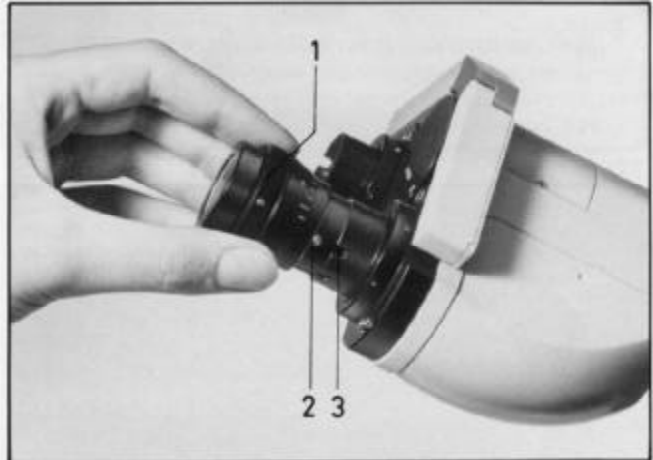
#### Slide in analyzer slider (Fig. 30)

Loosen knurled screw (30.2) and slide analyzer (30.1) into tube slot.

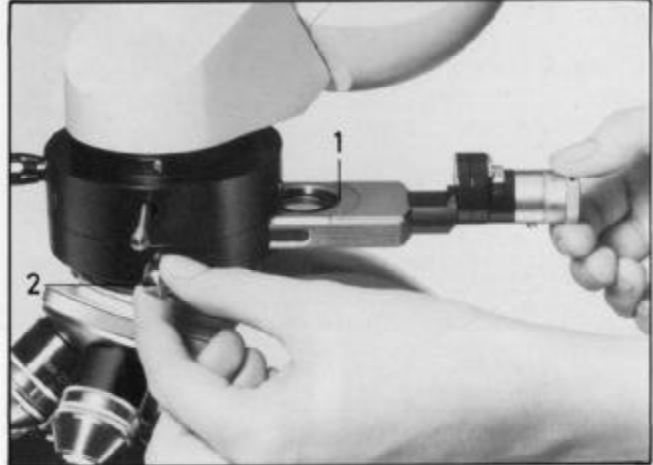
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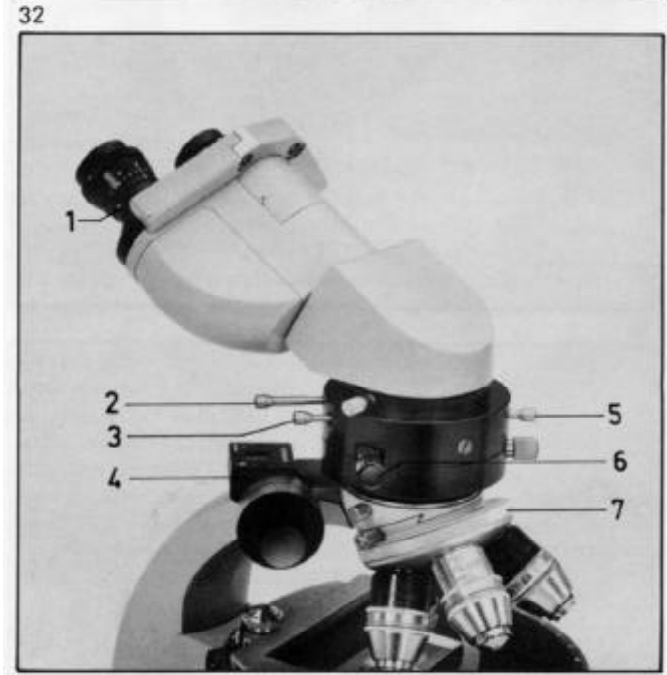
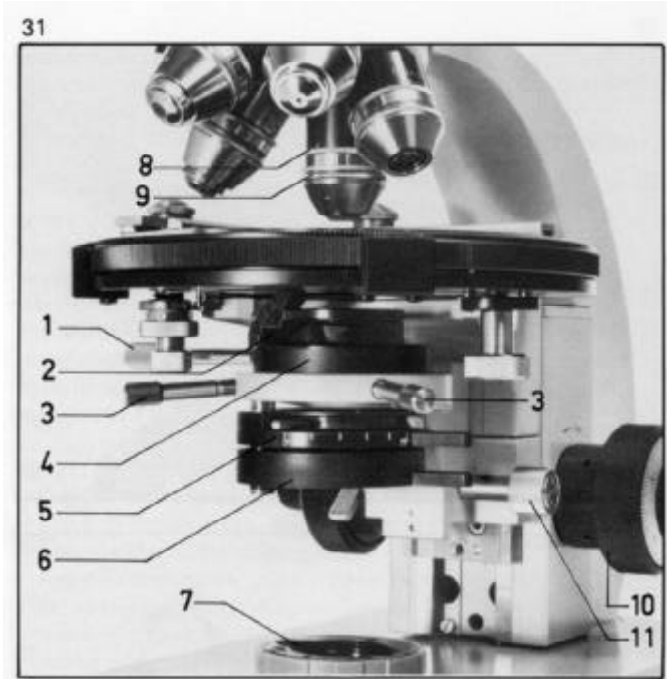


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**Centrosopic image**

1. The microscope is set up as follows: The condenser is raised to its top position. The condenser lens is swung in. The condenser lens is swung in.

2. The condenser (31.4) with control (31.1) is raised to its top position. The condenser lens is swung in. The condenser lens is swung in.

3. The condenser lens (10.1) is screwed into the condenser in the factory. On the intermediate tube pull out analyzer, auxiliary object or filter (10.2, 10.3, 10.4) and slides (10.2) and (10.3).

4. The condenser lens is swung in. The condenser lens is swung in. The condenser lens is swung in.

**Focus on the specimen**

5. While looking through the eyepiece with adjusted reticle, focus on the specimen with control (31.10). With diopter ring of the eyepiece focus for the other eye.

**Centering the Pol Z objectives (with centering rings)**

6. With the point of intersection of the eyepiece cross hairs, turn specimen stage. If a feature in the cross-hair center changes its position, the objective must be centered.

7. The attachable mechanical stage to the objective-axis.

8. To the cross-hair center by slightly turning centering rings.

9. The objective is centered.

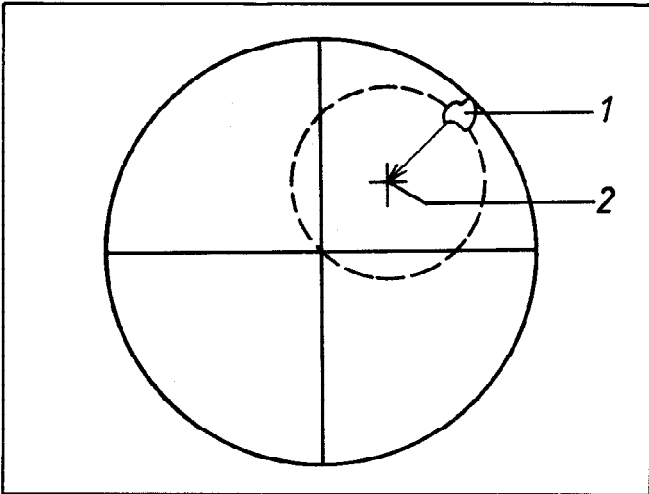
10. The objective is centered when the feature (35.2) remains in the cross-hair center while the stage is turned. All other features move on concentric circles about the cross-hair center.

11. The objective is centered when the feature (35.2) remains in the cross-hair center while the stage is turned. All other features move on concentric circles about the cross-hair center.

12. Center one objective after the other in this manner. To turn in another objective, hold the nosepiece on its knurled ring.

13. The objective is centered to avoid decentering.

33



**Adjust Köhler illumination**

While looking through the tube close luminous field diaphragm (31.7) about one half. Set condenser to focusing position with knob (31.11), i.e. focus the image of the luminous field diaphragm in the object plane (Fig. 36). Center this image with screws (31.3) (Fig. 37), open luminous field diaphragm until its image just illuminates the field of view (Fig. 38).

**Observation of the aperture diaphragm**

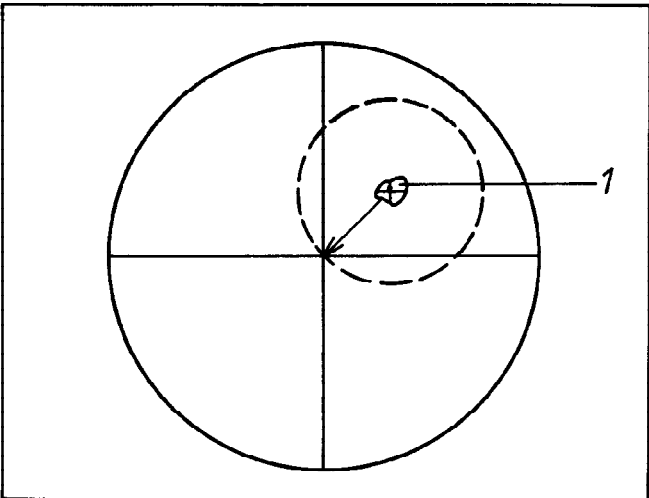
1. With diopter (46 48 89) in case of normal monocular or binocular tube. Plug the diopter into the tube instead of one eyepiece.
2. Through the swung-in Bertrand lens (3.2) in case of the monocular polarizing tube (3.1).
3. With swung-in Bertrand optics (32.5) and diaphragm (32.2) in case of intermediate tube (47 30 56) (Fig. 32). Focus the aperture diaphragm with knob (32.6).

**Adjusting the aperture diaphragm**

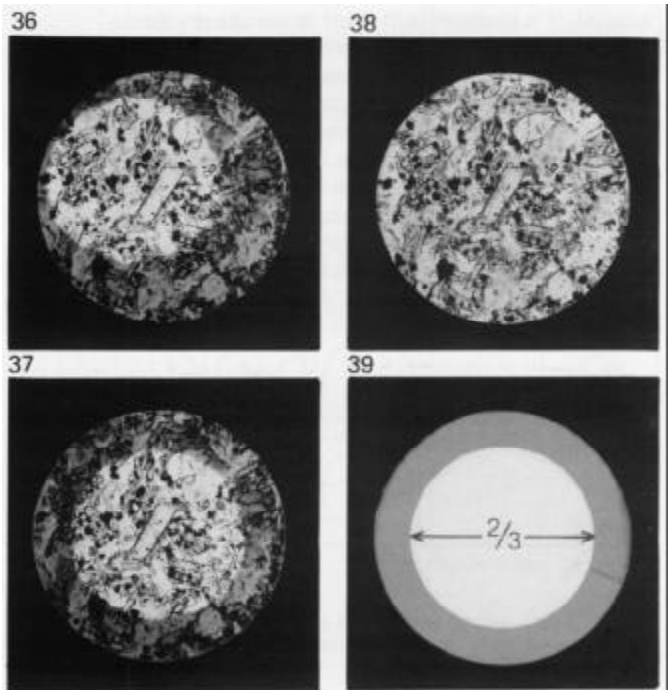
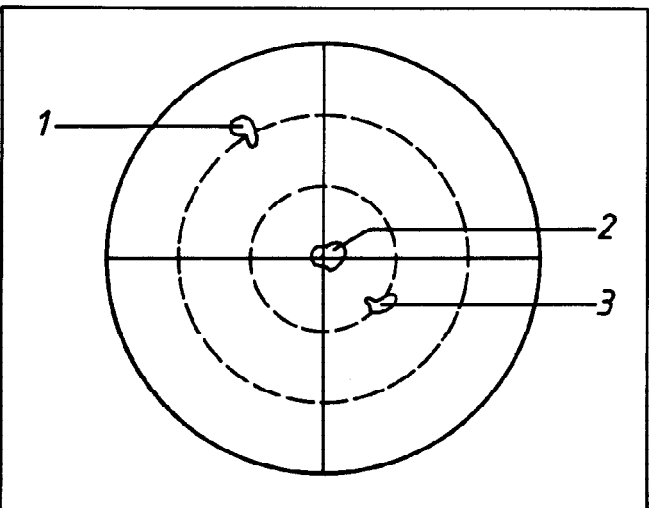
Close aperture diaphragm (31.2) until its image frees about 2/3 (Fig. 39) of the objective exit opening. Closing the diaphragm further increases the contrast but reduces resolution. It is recommended to check and/or correct the aperture-diaphragm setting after objective change.

(For a small convergence of the illuminated light bundles the aperture diaphragm must be closed further. This is necessary, for instance, to determine the extinction direction of anisotropic crystals, to observe a high-contrast Becke line, or to increase the black-and-white or color contrast between crossed polarizers.)

34



35



**Alignment of polarizer and analyzer**

The reference direction is East-West.

The pre-aligned oscillation direction of a non-rotary polarizer is East-West.

The oscillation direction of a rotary polarizer is East-West if its angle scale is set to  $0^\circ$ .

The pre-aligned oscillation direction of a fixed analyzer is perpendicular to that of the polarizer. The pre-aligned oscillation direction of a rotary analyzer (30.1) is parallel with that of the polarizer if its angle is set to  $0^\circ$  or perpendicular if it is set to  $90^\circ$ . Swing polarizer and analyzer into the beam path.

**Working with low-power objectives**

Planachromat 2.5/0.08 Pol objective (46 01 18) images a large object field; to fully illuminate this field the condenser front lens must be swung out.

Auxiliary lens (31.6) remains in the beam path. Diaphragm (31.7) acts as aperture diaphragm.

An exceptionally large object field is covered by the illumination remove the condenser and swing in auxiliary lens (31.6). Diaphragm (31.7) acts as aperture diaphragm.

**Working with oil immersion objectives**

Only when the supplied oil is used can the capabilities of the immersion objectives be fully utilized. Avoid air bubbles on the specimen during immersion, because they would impair the image. Swing objective out of the beam path and apply a drop of immersion oil to the coverglass. Turn immersion objective into beam path. Carefully focus with fine focusing control. It is generally sufficient to use the dry Pol condenser.

The immersed condenser Pol, aperture 1.3 (46 52 63) is required to cover the conoscopic image under the largest possible aperture angle. With the oiler apply a drop of immersion oil to the condenser front lens, so that front lens and specimen slide bottom are connected by the oil. After termination of the examination remove the oil from all surfaces with a clean, soft cotton cloth and optical detergent solution from the kit (43 90 07).

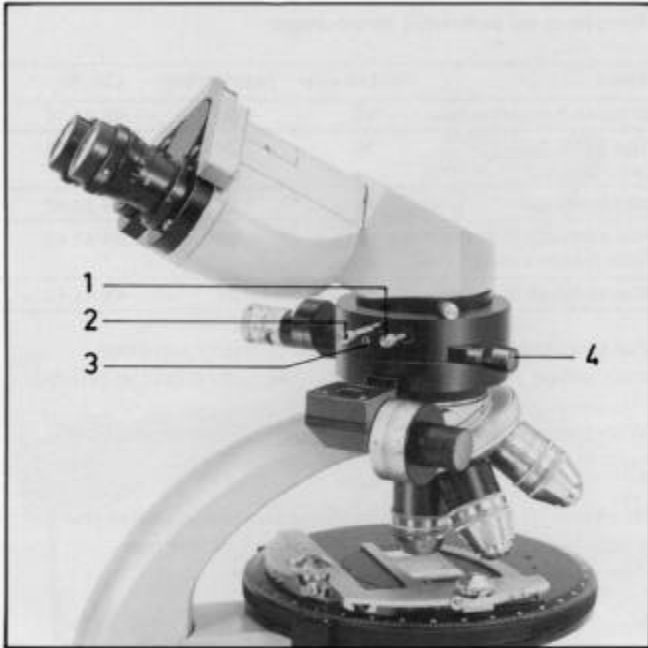
**Conoscopic image (interference figure)**

Adjust specimen for orthoscopic observation (see p. 16) and move it to the center of the cross hairs. Turn in suitable Pol objective of high aperture (25x or higher). Open luminous field diaphragm and condenser (aperture) diaphragm. Polarizer and analyzer must be in crossed position. Focus the interference figure with knob (40.4). (Write down the number of the finder scale for the corresponding objective.) Select the best pinhole diaphragm with slider (40.1). If desired, cut off orthoscopic image in the environment of the conoscopic image with slider (40.2).

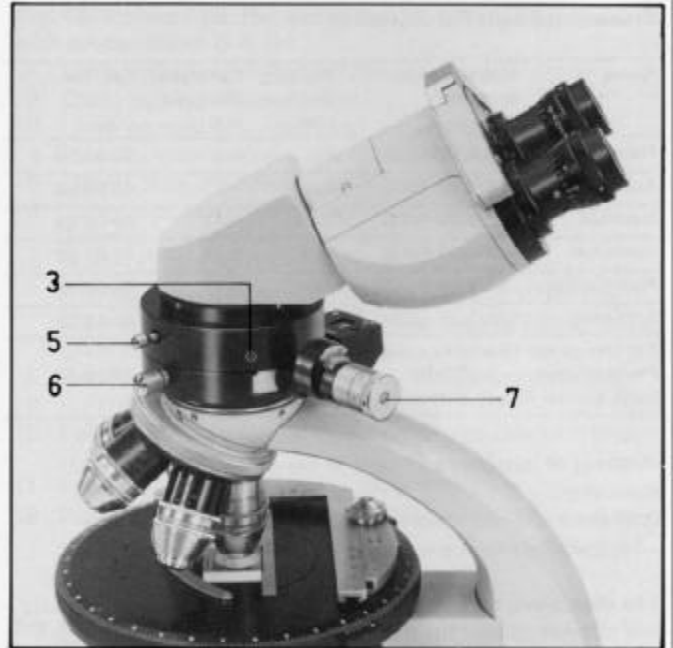
Adjust the microscope with the supplied pinhole diaphragm the reticle with the supplied hex socket wrenches and the hex socket screws (40.3) and (41.3).

On the monocular polarizing tube (3.1) swing in Bertrand lens and diaphragm with knob (3.2) and observe the interference figure.

40



41



The oscillation directions of the orthoscopic image and the character of birefringence of the conoscopic interference image are indicated in the table below

Reference directions with auxiliary objects and compensators	Conoscopic adjustment						Orthoscopic adjustment	
	optically uniaxial section perpendicular to the optical axis		optically biaxial section perpendicular to the acute bisectrix				addition position	subtraction position
			cross position		diagonal position			
	+	-	+	-	+	-		
λ plate								
During insertion of quartz wedge 0-3 λ and tilting of compensators B and E							higher	lower
Auxiliary object λ/4							interference colors	
	Displacement direction of interference fringes in the conoscopic image						η <sub>ω</sub> in the object	

\*The indicated colors of the orthoscopic image are examples which are given when first order white is visible without auxiliary object

**Transmitted-light Pol objectives**

Name	Magnification/ aperture	Working distance mm	Coverglass thickness mm	Order No.
Planachromat	2.5/0.08 Pol Z	8.5	—	46 01 18
Achromat	10/0.22 Pol Z	5.0	—	46 04 08
Neofluar	25/0.60 Pol Z	0.54	0.17	46 06 28
Plan-Neofluar	63/0.90 Pol Z	0.09	0.17	46 08 18
Achromat	100/1.25 oil Pol Z	0.09	0.17	46 19 08
For low-power observation and photomicrography:				
Planachromat	1.25/0.04	4	—	46 20 14
(with normal 45 mm parfocal length)				

Meaning of numbers and signs in the column coverglass thickness:

0.17 cover the objective by an 0.17 mm thick coverglass  
— specimen can be used with or without coverglass

The objectives designated Pol contain strain-free lens elements and do not change the polarization state between polarizer and analyzer, which is a pre-condition for exact phase-difference measurements with compensators.

The designation Pol Z refers to transmitted-light objectives in centering mount.

The Pol Z objectives 25x, 40x and 63x have exceptionally high apertures and are therefore particularly well suited for conoscopic examination.

**The total magnification of the microscope** results from the multiplication

$$M_{\text{microscope}} = M_{\text{obj}} \times M_{\text{eyep}}$$

for example: 400 = 40 × 10

Meaning of abbreviations:

$M_{\text{obj}}$  = objective magnification, here 40x

$M_{\text{eyep}}$  = eyepiece magnification, here 10x

**Eyepieces for polarizing microscopes**

Name	Magnification	Field angle	Order No.
C 8x/16 Pol with reticle	16	30°	46 39 15
Kpl 8x/18 Pol	18	33°	46 39 25
Kpl 8x/18 foc	18	33°	46 39 23
Kpl 12.5x/18 W Br Pol	18	48°	46 41 45
Kpl 12.5x/18 W Br Pol with crossline micrometer	18	48°	46 41 45

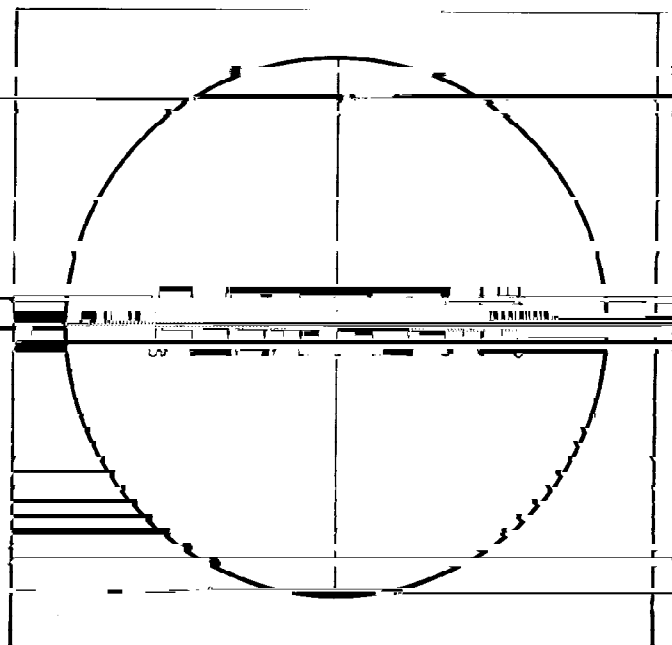
**Pol eyepieces** contain a pre-aligned reticle or crossline micrometer (Fig. 42) and a screw (32.1) to align the reticle.

With the wide fields of view of **wide-angle eyepieces (W)** large-diameter object fields can be surveyed.

**Br eyepieces** are high-eyepoint eyepieces which offer the eyeglass wearer the same wide field of view the non-eyeglass wearer enjoys.

**Eyepieces foc** come with diopter setting rings.

**Fig. 42: Eyepiece crossline micrometer**



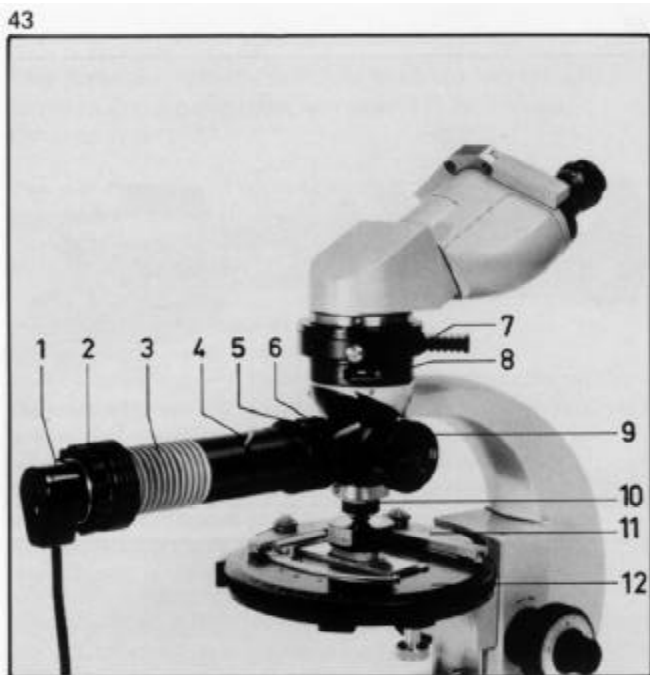


Fig. 43. Standard Pol microscope with epi-condenser II A ore

- 3 Lamp housing with clear-glass lamp condenser
- 4 Lever to open and close the aperture iris diaphragm
- 5 Slider (47 36 63-9903) with analyzer slid into large tube slot, its observation direction is North-South, when the slider is completely pulled out a quartz plate will be in the path

- 10 Epiplano Pol objective on centering change ring with thread W U.6 (48 02 86)

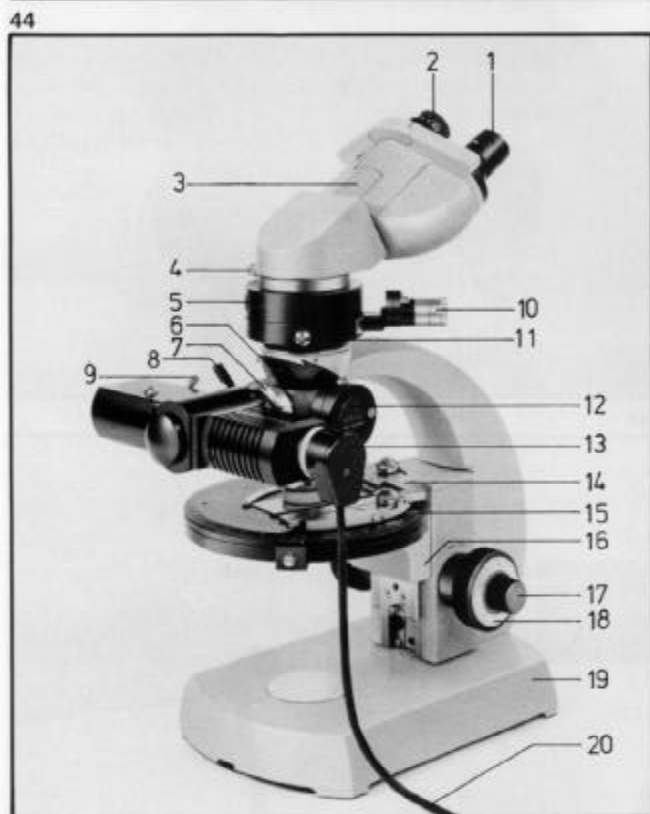
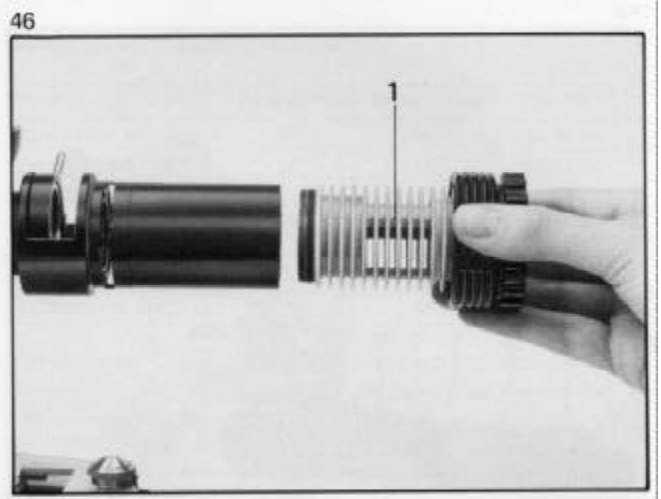
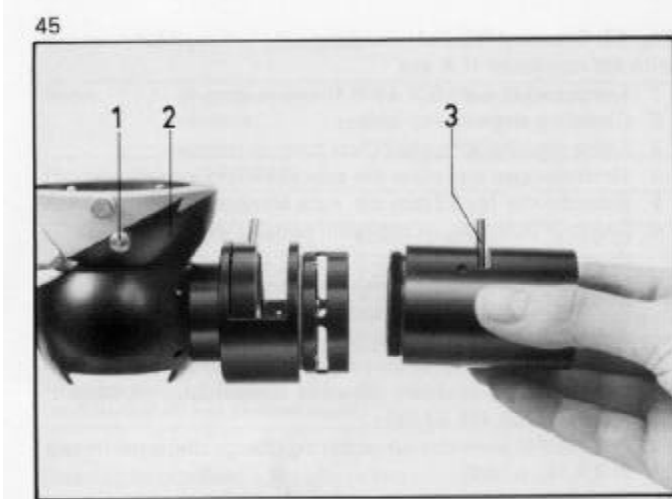


Fig. 44. Standard WL Pol microscope

- 3 Binocular tube S 30 Pol (47 30 30) with reticle erection system
- 5 Intermediate tube (47 30 39-9902)
- 6 Epi-condenser II S ore (48 64 86)
- 7 Pre-aligned, rotatable through 90°, oscillation direction East-West in 0° position, lockable
- 10 Slider with rotary analyzer (47 30 62-9903)
- 11 Clamping screw of analyzer slider
- 14 Attachable mechanical stage Pol (47 33 25-9903) to move the specimen within a 30 x 40 mm area
- 15 Rotary polarizing stage (47 34 04) with graduation and vernier and clamping device
- 19 Standard WL stand
- 20 Transformer for connection of lamp and transformer

For Standard U4 or U6 Pol microscope with incident-light accessory  
 For operating instructions see 1.1.1.



Clamp attachable stage carrier with specimen stage to change rail of Standard WL microscope (see p. 9).

**Mounting epi-condenser on stand head with glide mount (Fig. 45)**

Mount epi-condenser (45.2) slightly tilted, so that at first ca. 2/3 of the rail engage the guide with the clamping screw (45.1). Then swing in the other rail and slide the epi-condenser in as far as it will go. When sliding the epi-condenser into or out of the glide mount, pull out the spring locking pin at the end of the knurled screw after having slightly loosened clamping screw (45.1).

Secure epi-condenser with clamping screw.

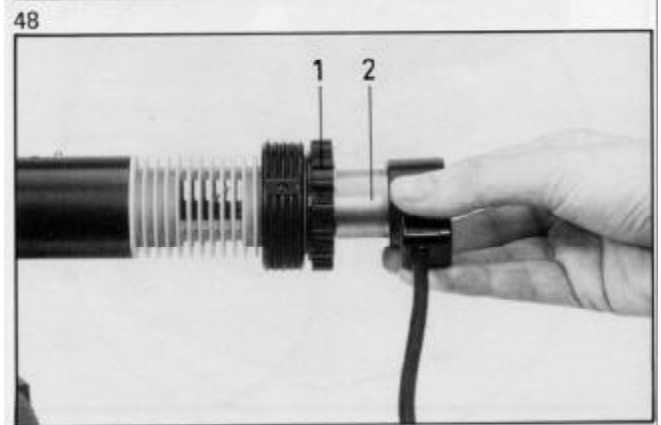
**Screw aperture diaphragm part (45.3) to epi-condenser II A**

Screw lamp housing with lamp condenser (46 72 55) (46.1) to aperture diaphragm part.

Hold **6 V 15 W filament lamp** (38 00 18-1740) (47.1) with a soft cloth and plug it into socket (46 80 10-9904) (47.3); red dot must be opposite red pin. Press down lamp, turn it clockwise as far as it will go and let it snap in. Remove fingerprints on the glass bulb. Slide sleeve (47.2) over lamp. Turn clamping ring (48.1) so that its red dot is opposite the red dot of the lamp housing with lamp condenser.

Slide filament lamp in socket (48.2) into lamp housing and secure with clamping ring.

Connect 6 V 15 W lamp to the line by cable via transformer.



**Slide in reflector (Fig. 49)**

Slide plane-glass reflector H-PI Pol (46 62 65) into epi-condenser so that pin (49.1) engages notch (49.2). Tighten clamping screw (49.3).

The H-Pr Pol prism reflector (46 62 61) is recommended for examinations of weakly anisotropic substances. Its image brightness is considerably higher than that of the plane-glass reflector (see p. 26).

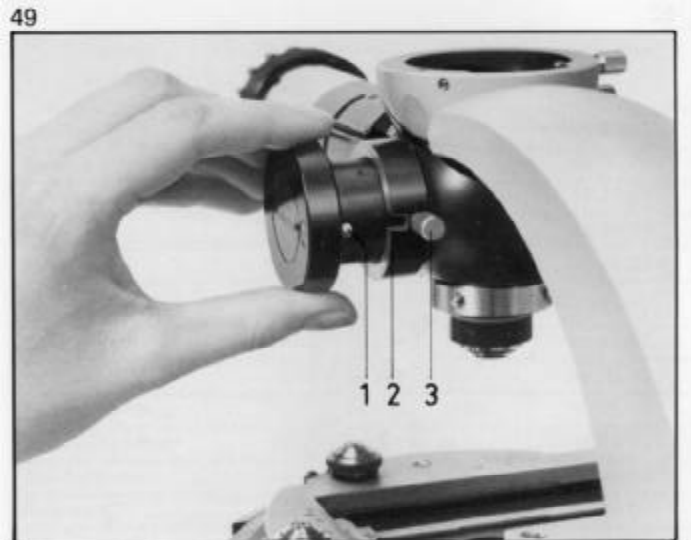
**Screw Epiplan Pol objective (50.3) into centering change ring**

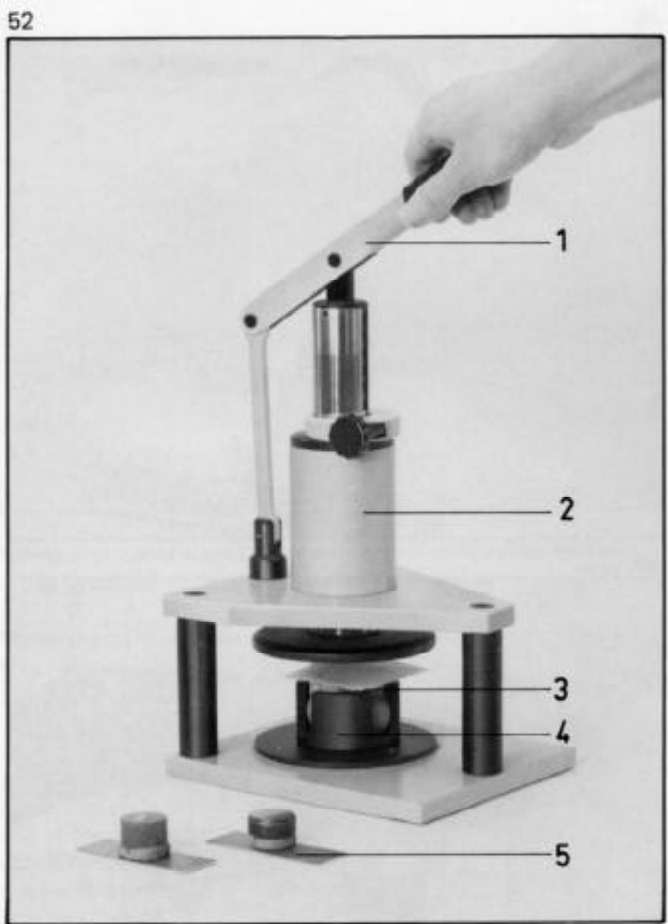
Insert change ring with Epiplan Pol objective from the side in the dovetail guide (50.1) and turn it backwards as far as it will go. For mounting of intermediate tube, binocular Pol tube, and eyepieces, and slide-in of analyzer see the corresponding instructions for the transmitted-light polarizing microscopes (p.15).

The tube slot for auxiliary objects becomes accessible when a lever is slid to the left.

Either an East-West-aligned, swing-in polarizer (46 62 89) (43.6) is firmly mounted on the epi-condenser, or a rotary polarizer (47 36 16) (51.2) inserted in the filter holder so that pin (51.1) engages notch (51.3).

Turn polarizer to 0° position and secure with knurled screw (Fig. 51); its oscillation direction is East-West. The second filter holder accepts a 32 mm dia. filter in holding ring (46 72 52).





**Preparations**

Place polished section on pot (52.4) filled with plasticine or on a specimen slide with plasticine and align it perpendicular to the microscope axis by pressing down lever (52.1) of the levelling press (47 89 62) (52.2). The polished section to be examined should be covered by a sheet of tissue paper to avoid scratches on the surface. Polished sections on normal specimen slides (52.5) can be aligned with the levelling press in the same manner. Fix specimen in the specimen holder on the specimen stage. Connect microscope illuminator with transformer. Before mains connection check whether the voltage set on the transformer coincides with the local mains voltage. If not, set the correct voltage or call our maintenance service. Switch on transformer and adjust brightness. Set polarizer to 0°, mount Epiplan Pol objective (e.g. 16x) on the epi-condenser. Swing analyzer and auxiliary object out of the beam path.

70 mm are the maximum height of specimens (distance between stage top surface and focusing plane) which can be examined under a standard VLE incident-light microscope.

Microscopes Standard 08 and 18 with slide mount on the stand head can be used with the same equipment. The maximum specimen height is 15 mm. Remove eye-piece (44.1) from the tube, hold it against a bright surface and watch the reticle, thereby leaving your focus.

The user's diopter value is then indicated by the red dot on the diopter scale.

Try to insert eye-piece into tube so that the alignment screw is visible.

**Center Epiplan Pol objective**

By rotating the objective (40.3) and center the intersection of the reticle cross hairs of the eyepiece (see p. 10).

**Adjust aperture diaphragm**

The image of the aperture diaphragm becomes visible in the field of view. The image is replaced by diopter (40 48 89), or, when with the intermediate tube, the Bertrand optical system is brought into the beam path (Fig. 32) (see p. 17).

Adjust aperture iris diaphragm of epi-condenser II A with lever (50.4).

Adjust aperture iris diaphragm of epi-condenser II A with lever (43.4).

Close aperture diaphragm so far that its image releases ca. 1/3 of the objective exit opening. Closing the diaphragm further improves the contrast but reduces the resolution. Check and/or correct the aperture diaphragm adjustment after objective exchange. Insert eyepiece in tube or swing Bertrand optical system out of the beam path.

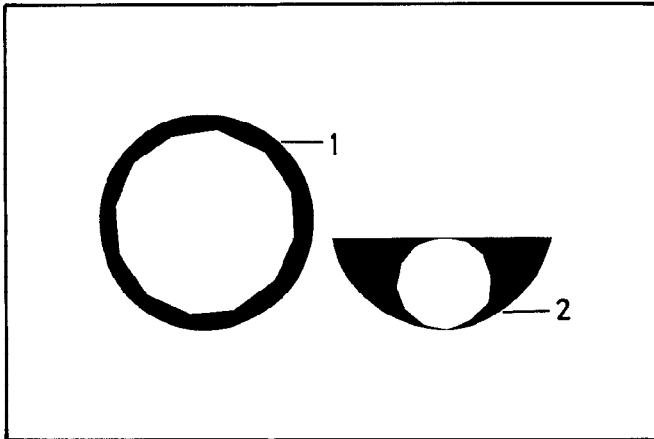


Fig. 53: Aperture-diaphragm images in objective aperture

- 1 with H-PI Pol plane-glass reflector
- 2 with H-Pr Pol prism reflector

#### Cross incident-light polarizer and analyzer

Slide in analyzer (43.7) or set rotary analyzer (44.10) to  $90^\circ$ .  
 Slide in polarizer (43.6) or set polarizer (44.7) to  $0^\circ$ . It may  
 be necessary to turn it slightly until a strongly reflecting iso-

tropic phase of the specimen, or, better a surface mirror  
 appears black. Set lamp to maximum brightness. Focus on a  
 highly anisotropic object. Four times darkness and four times  
 brightness must appear during stage rotation, and the dark and  
 bright specimen phases must re-appear exactly every  $90^\circ$  stage  
 rotation. If, for instance, only 2 bright phases are observed  
 during a stage rotation through  $360^\circ$ , the wave trains coming  
 from the polarizer are not aligned perpendicular to the  
 reflection plane of the reflector.

When a dark phase is observed, the light need not be rotated by  
 slight amounts until the above-mentioned phenomena are  
 observed.

Dry objectives	Magnification/ aperture	Working distance mm	Cat. No.
Epiplan	4/0.10 Pol	9.0	46 20 01
Epiplan	8/0.20 Pol	7.1	46 20 02
Epiplan	16/0.35 Pol	3.1	46 20 03
Epiplan	40/0.85 Pol	0.23	46 20 04
Epiplan	80/0.95 Pol	0.09	46 20 80

#### Immersion objectives

Epiplan	4/0.10 oil Pol	0.3	46 20 06
Epiplan	8/0.20 oil Pol	0.3	46 20 07
Epiplan-Neofluar	16/0.50 oil Pol	0.32	46 15 76
Epiplan-Neofluar	25/0.80 oil Pol	0.38	46 16 76
Epi-Achromat	40/0.85 oil Pol	0.5	46 20 09
Epiplan	100/1.25 oil Pol	0.25	46 20 05

#### Objectives with long working distances

LD-Epiplan	16/0.30 Pol (covergl.thickn.1.5)	4.1	46 21 23
LD-Epiplan	40/0.60 Pol	3.4	46 21 24
LD-Epiplan	100/1.25 Pol (covergl.thickn.0)	0.25	46 20 05
Cap	LD 40 Pol (covergl.thickn.0)	2.4	46 29 16

#### Epiplan Pol objectives

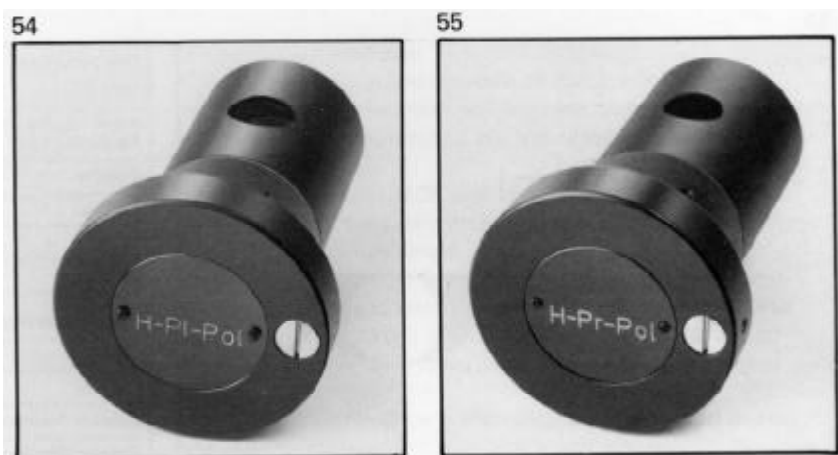
contain strain-free lens elements, which ensures that the  
 polarization state between polarizer and analyzer is not  
 changed.

LD-Epiplan objectives (coverglass thickness 1.5) have long  
 working distances. They are used with LD-caps (coverglass  
 thickness 0) or LD-caps (coverglass thickness 1.5).

LD-cap (coverglass thickness 0) of the objective on the  
 objective mount. Screw on aluminum ring.

When changing between high and low magnifications by  
 exchanging the objectives the specimen need not be created.  
 The same oil immersion is the same at all magnifications.

The characteristics of the H-PI Pol (Fig. 54) and H-Pr Pol (Fig. 55) reflectors used in epi-condensers II A (46 62 30) and II ST (46 62 15-9901) for incident-light microscopy in polarized light are listed in the table below.



**H-PI plane glass reflector,  
Vickers licence (46 62 65)**

**H-Pr Pol prism reflector  
(46 62 61)**

<b>Resolving power of the objective</b>	fully utilized	better resolution of structures normal to prism edge
<b>Illumination aperture</b>	fully utilized	up to 50 % objective aperture
<b>Specimen illumination</b>	practically perpendicular	unilateral oblique illumination
<b>Image brightness</b>	losses through beam splitting; efficiency noticeably spectrum-dependent (selective), e.g. photometry	higher
<b>Illumination of total field of view in unpolarized or singly polarized light</b>	uniform	with high-aperture objectives slight vignetting at upper and lower edges of image
<b>Extinction of entire field of view with crossed polarizer and analyzer</b>	homogeneous	homogeneous
<b>Preferred use</b>	observation in polarized light, photomicrography	weak anisotropy low reflection

K 41-003	Handbook of incident-light microscopy
G 41-119/I	Standard microscope with 6 V 10 W in-base transilluminator for transmission
<del>G 41-120/I</del>	<del>Standard and Standard WL microscopes with 6 V 15 W in-base transilluminator</del>
G 41-310/III	Microscope illuminator 100
G 41-415	MC 63 attachment camera
G 41-417	MC 63 A attachment camera
G 41-505	Standard 04 Pol microscope
G 41-510/I	Sénarmont compensators
G 41-511	Auxiliary objects $\lambda$ and $\lambda/4$
G 41-516	Brace-Köhler compensators
G 41-521	Tilting compensator B, measuring range $5 \lambda$
G 41-523	Tilting compensator E, measuring range $20 \lambda$
S 41-500.0	Michel-Lévy color chart
A 41-656.1	Antiflex Device