



BS-5062 Series

Polarizing Microscope Instruction Manual

To ensure the safety and obtain satisfactory performance, please study this instruction manual thoroughly before your operation.

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1. Purpose

BS-5062 Series polarizing microscope is for the field of metallurgy, geology and minerals.

BS-5062 Series polarizing microscope is with gypsum(λ), mica ($\lambda/4$) sample, quartz wedge and attachable mechanical stage. It is an ideal instrument that has perfect function and quality.

BS-5062BTR/TTR transmitting & reflecting polarizing microscope is perfect in optical and mechanical quality, and it can be used in observing even, non-even, transparent, non-transparent mineral sample.

2. Specification

2.1 Total Magnification

2.1.1 Transmitting

| Objective \ Eyepiece | 4X | 10X | 20X | 40X | 60X | 100X |
|----------------------|-----|------|------|------|------|-------|
| 10X | 40X | 100X | 200X | 400X | 600X | 1000X |

2.1.2 Reflecting

| Objective \ Eyepiece | 5X | 10X | 20X | 50X | 100X |
|----------------------|-----|------|------|------|-------|
| 10X | 50X | 100X | 200X | 500X | 1000X |

2.2 Objectives

| Objective | N.A. | Working distance (mm) | |
|--|------------------|-----------------------|-------|
| Non-stress Infinity Plan Objective (Transmitting) | PLAN 4X | 0.10 | 12.1 |
| | PLAN 10X | 0.25 | 4.64 |
| | PLAN 20X(S) | 0.40 | 2.41 |
| | PLAN 40X(S) | 0.66 | 0.65 |
| | PLAN 60X(S) | 0.80 | 0.33 |
| | PLAN 100X(S,Oil) | 1.25 | 0.19 |
| Non-stress LWD Infinity Plan Objective (Reflecting) | LPL 5X | 0.13 | 16.04 |
| | LPL 10X | 0.25 | 18.48 |
| | LPL 20X | 0.40 | 8.35 |
| | LPL 50X(S) | 0.70 | 1.95 |
| | PLAN 100X(S,Dry) | 0.90 | 1.10 |

2.3 Eyepiece

| Magnification | Type | View Field Diameter(mm) |
|---------------|------------------|-------------------------|
| 10X | High eye point | 20/22 |
| 10X | Reticule (0.1mm) | 20/22 |

2.4 Mechanical Tube Length: ∞

2.5 Head: Seidentopf binocular (trinocular) head 30°,

Interpupillary adjustable distance is 48-76mm.

Diopter adjustable range ± 5 ,

Anti-fungal systems.

2.6 Intermediate: 360°part division for analyzer, 2°30'per scale, lock system

Bertrand Lens (center adjusting)

Gypsum(λ), mica ($\lambda/4$)sample, quartz wedge

2.7 Nosepiece: Quadplex or quintuple nosepiece (center adjusting), nosepiece spanner.

2.8 Revolving Round Stage: Diameter $\Phi 174$ mm, 360° part scale, 6'per scale.

2.9 Focusing System: Coaxial coarse and fine focusing knobs, coarse stroke 22mm.

Fine division 2 μ m, condenser up-down range 22mm

2.10 Condenser: Abbe condenser, N.A.0.9/0.13 swing out condenser, adjustable aperture, aperture center can be adjustable.

360°part division for polarizer, 5° per scale, lock system

2.11 Electric components: Input voltage AC100-240V, 50/60Hz

Output voltage DC1.2-6V

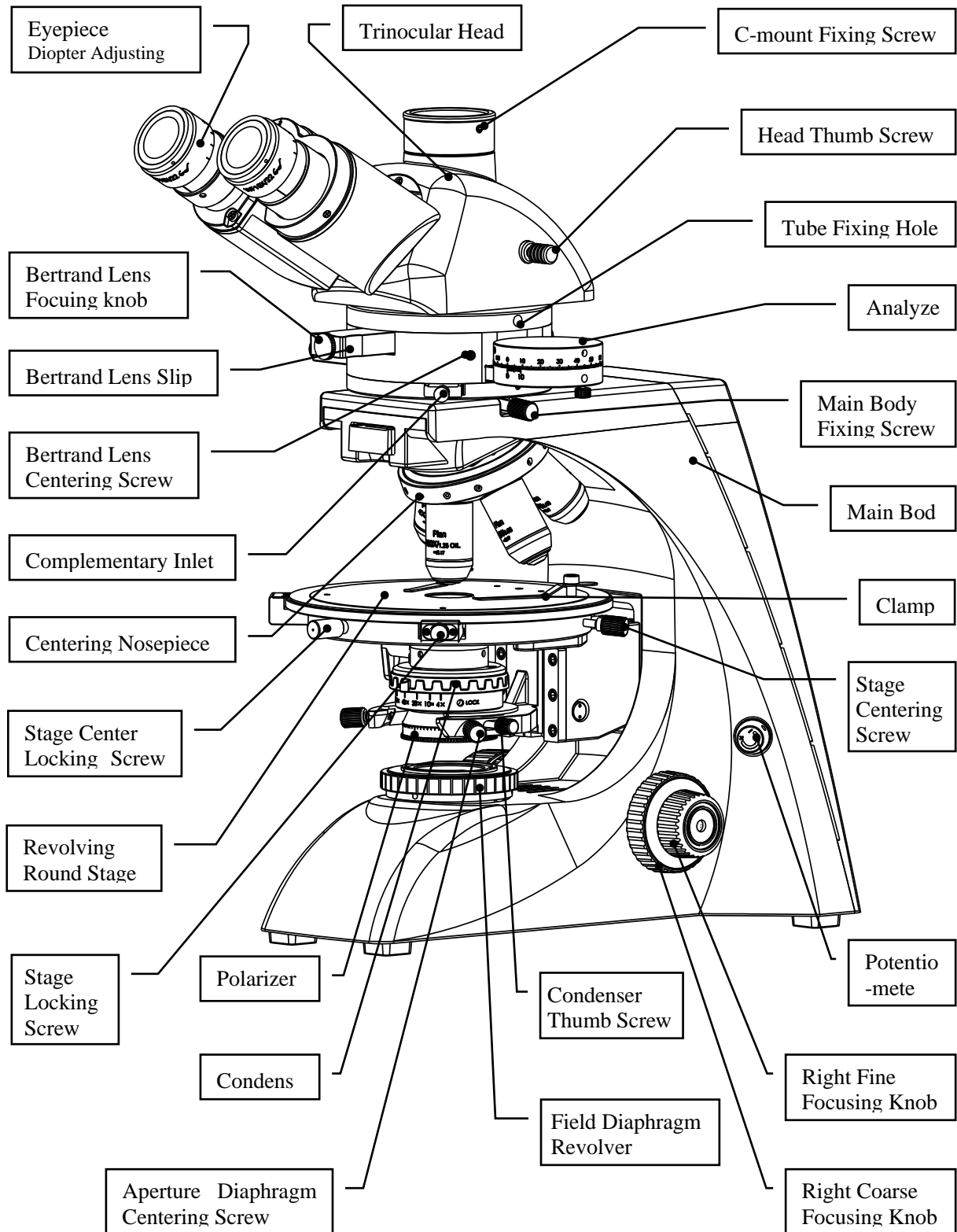
6V/30W halogen lamp

Rotation potentiometer with power switch

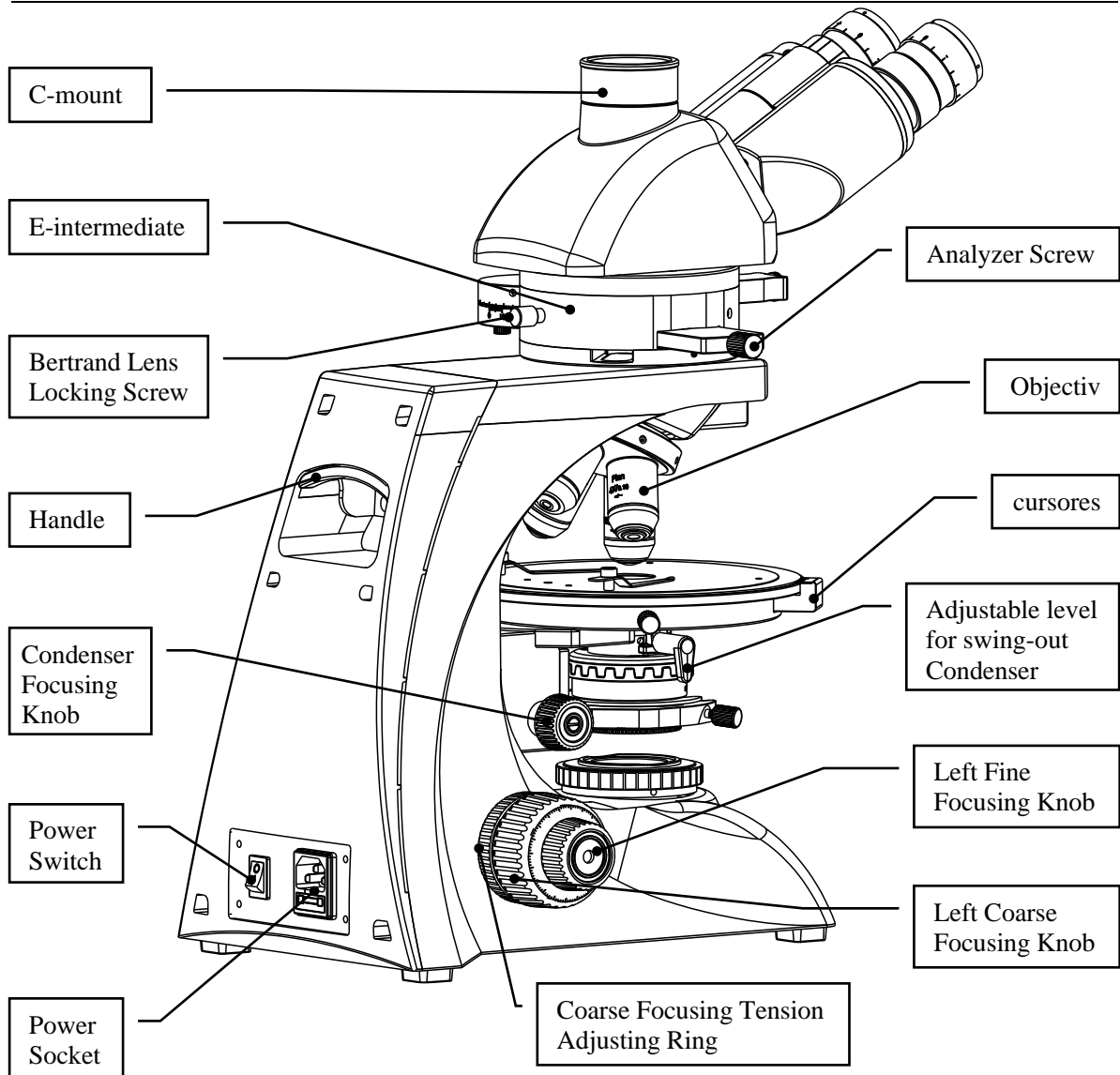
Fuse 5A $\phi 5 \times 20$

2.12 Filter: Blue (Amber, green, neutral filter optional)

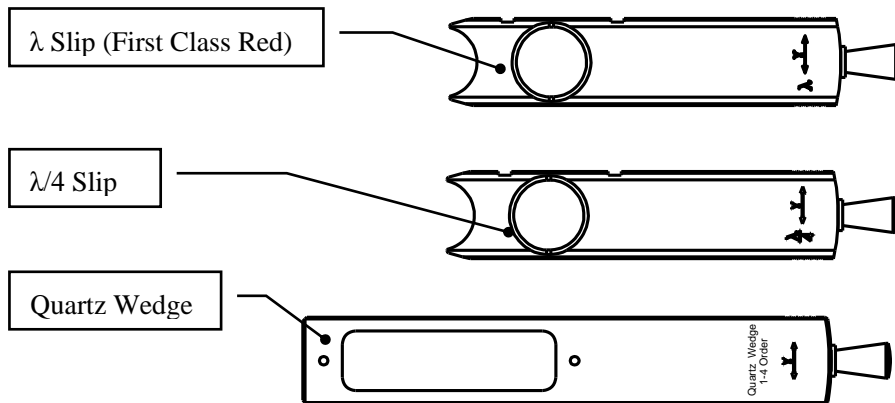
3. Parts Name

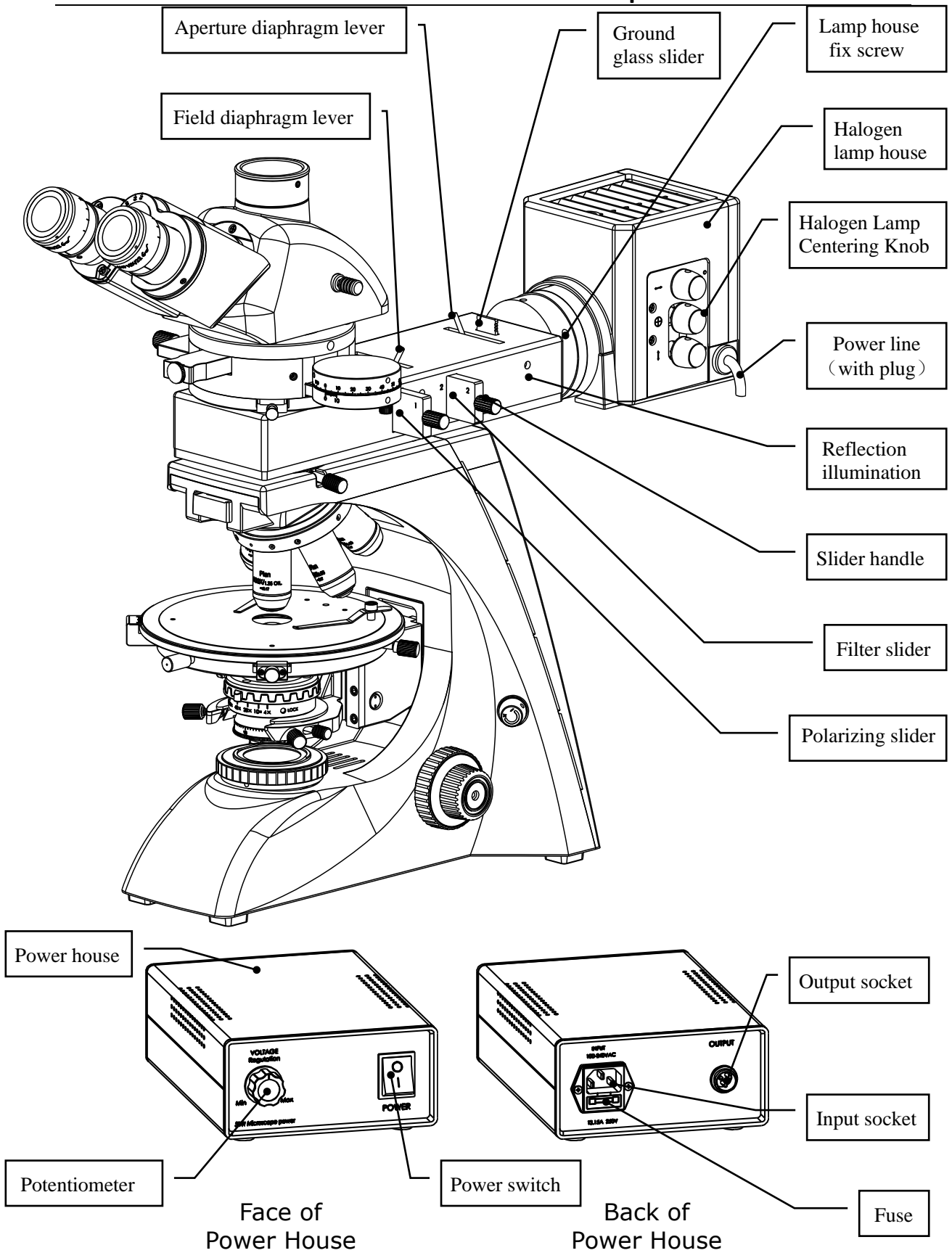


BS-5062T Transmitting Polarizer Microscope (1)

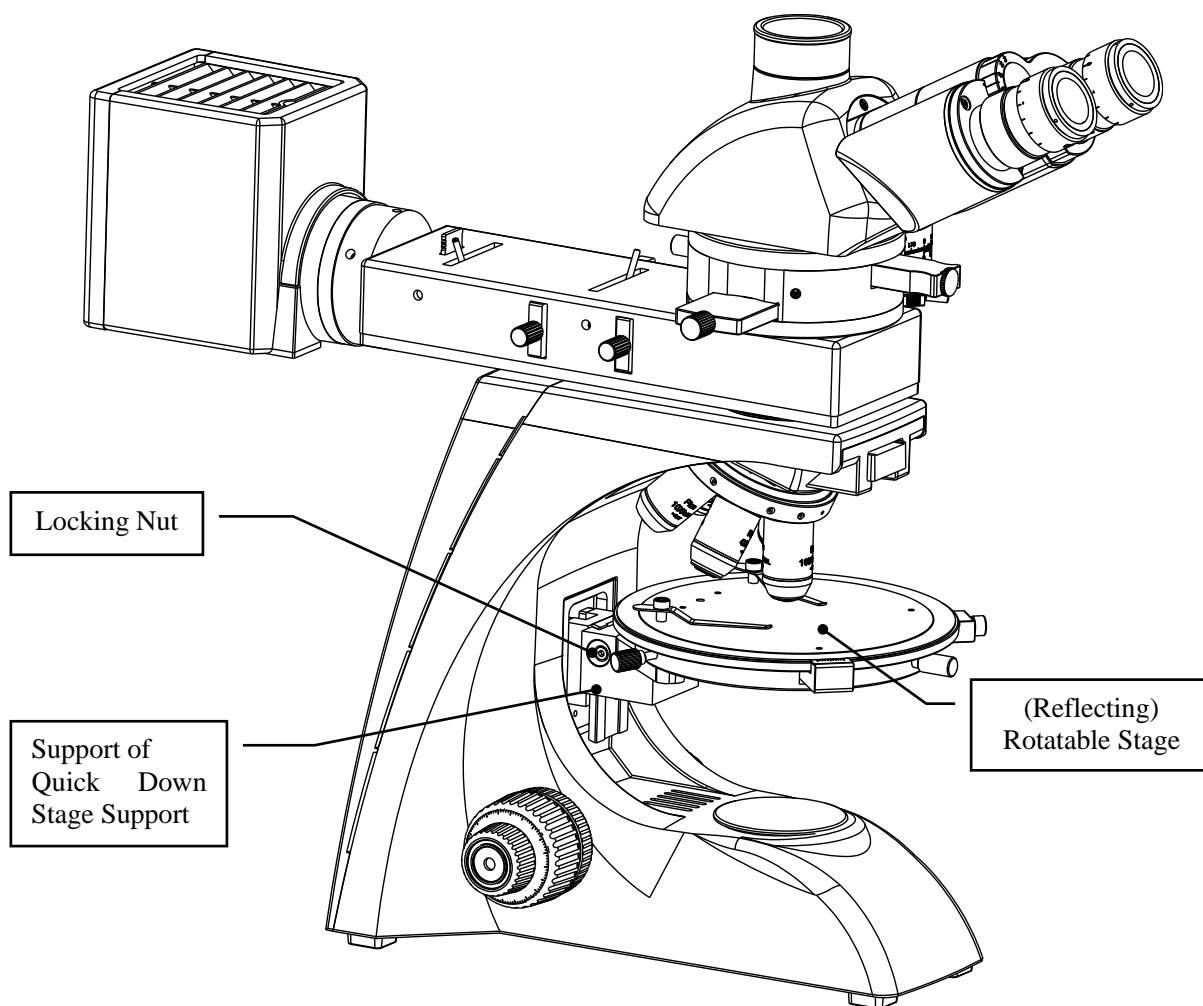


BS-5062T Transmitting Polarizer Microscope





BS-5062TTR Transmitting & Reflecting Polarizer Microscope



BS-5062TR Reflecting Polarizing Microscope

4. Installation

4.1 Installation Condition

4.1.1 The required input voltage: **100V-240V**, 50/60HZ

4.1.2 Alcohol, gasoline and paper all are burnt early, please take them away from the lamp.

4.1.3 The halogen lamp: **6V/30W**

4.1.4 The microscope should be used in environment of indoor temperature 0°-40°C and maximum relative humidity 85%.

4.1.5 Please pay attention to prevent microscope from violent shake and vibration in application and in carrying. Don't drag it on the surface of worktable to avoid damage to microscope and worktable.

4.2 Installation

4.2.1 Please confirm the installation condition meets 4.1;

4.2.2 Put out the main body and place it in table, and loose the main body fixing screw, then put the cost cover out;

4.2.3 Put out the intermediate, fix it into the main body, then tighten the main body fixing screw; If it is Transmitting & Reflecting Polarizer Microscope, put out the reflecting illumination unit to fix it.

Installation of reflecting illumination unit: Put out the halogen lamp house, fix it onto the back of reflecting illumination unit by the fixed screw, connect the power wire of halogen lamp house into the power output socket of power house.

4.2.4 Fix the head into the intermediate;

4.2.5 Insert the eyepieces into the tubes;

4.2.6 Put out the dust cover of the nosepiece, and turn the coarse focusing knob to lower the stage, and find the objective hole with yellow mark in the nosepiece. Fix the 10X objective into the hole, and turn the nosepiece clockwise, fix the other objectives as per the power.

4.2.7 Loose the condenser lock thumb and fix the condenser.

5. Operation

5.1 Turn the power switch and the potentiometer to adjust light to be available;

5.2 Fix the sample on the stage, and move it into the path;

5.3 Turn the nosepiece to put 10X objective into light path, and turn the focus knobs to get clear image.

5.4 Confirm the polarizing vibrancy direction

The polarizing vibrancy direction has set to be at west-east in factory when the scale of the polarizing is 0°.

5.5 Check polarizing and analyzer

The field should be dark completely (when there is no sample) when the scale of polarizer and analyzer is 0°. Please check the position of the polarizer and the analyzer if not so.

5.6 Choose the complementary slip as per the sample, then insert into the slip.

5.7 Put into the Bertrand Lens Slip in the condition of polarizing;

5.8 Adjust the center of the stage.

5.9 Adjustment for round stage rotating center

The rotating axis of the round stage should be in the same line with the axis of optical system. When rotating the stage, the sample's center would coincide with view field's cross division line intersection point, and image should move around cross division line intersection point in circular motion, image should always in the view field. If not, accurate data can't be got, and the operation can't be done correctly, especially for high power objective.

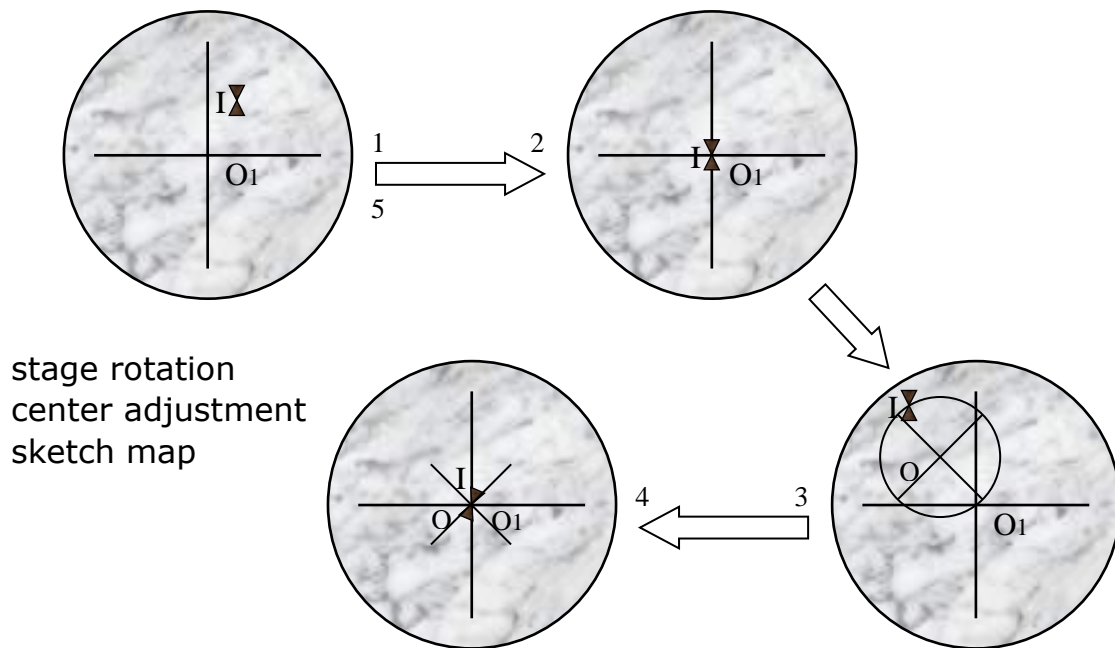
Adjusting Details:

5.9.1 After getting clear image, choose one image point (I) at the observational position, then moving mineral sample , making I in cross division line's point, and it is the center of field (mark: **O₁**).

5.9.2 Stage must rotate at least one circle, if actual rotating center (mark: **O**) isn't coincided with field center **O₁**, image point I will do circular motion around stage rotating center **O** .

5.9.3 The angle from rotating stage to initial position is 180°, point I will move from **O₁** to the full distance position , the symmetric point to point O. Adjusting the stage focusing screw, point I will move to point **O**, then move mineral sample gently, make point I coincided with field center **O₁** .

5.9.4 Rotating stage one circle again, if image point I isn't off point **O₁**, it means point **O** has coincided with point **O₁**, then the center adjustment is finished. If not, please repeat step2 , 3, 4.



- △ To confirm polarized light's vibration direction, choose a cleavage black mica, putting it in field center, then rotating the stage until the black mica color getting the darkest, the black mica cleavage direction is the polarized light's vibration direction.

6. Maintenance

6.1 Clean microscope

6.1.1 Don't touch the lens with hand, Dust on lens should be cleaned by soft brush or absorbent cotton or cleaned by absorbent cotton, lens paper with the mixture of alcohol and ether (proportion 1:4).

6.1.2 Alcohol and ether all are burnt early, please take them away from fire. Be careful for turn on and off power.

6.1.3 Don't clean painted metal and galvanizing metal with organic solvent such as alcohol, ether or the mixture of the both. Silicon cloth or soft cleaning preparation is suggested to clean it.

6.1.4 Plastic should be cleaned by soft cloth with clear water.

6.2 Environment of using and placing

6.2.1 Microscope should be used and placed in a cool, dry, non-dust, non-shake and non-corrosive gases environment.

6.2.2 Microscope should be used in environment of indoor temperature 0~40°C and maximum relative humidity 85%.

6.2.3 Removing equipment is suggested to be installed when microscope used in heavy humidity area to avoid fungus and mist damage instrument.

6.2.4 Please pay attention to prevent microscope from violent shake and vibration in application and in carrying. Don't drag it on the surface of worktable to avoid damage to microscope and worktable.

6.3 Replacement of bulb

6.3.1 Turn off power, and pull out plug.

6.3.2 Wait the bulb become cool.

▲ Please be sure that the bulb is cool, then follow by the next operations.

6.3.3 Lay aside the microscope reliably, unscrew the knurled thumb screw of the lamp housing cover on the underside of base.

6.3.4 Pull over the lamp housing cover.

6.3.5 Pull out the bulb should be replaced, hold a new bulb with silk cloth to avoid fingerprint and dust affect bulb brightness and service life, and insert fully the contact pins into the bulb socket.

6.3.6 Close the lamp housing cover, and screw the knurled thumb screw.

▲After working for above 10 hours continuously, better cut off the microscope about 30 minutes.

6.4 Replacement of fuse

6.4.1 Cut off power of microscope, and pull out the plug.

6.4.2 Unscrew fuse cap in the back of base, take out old fuse.

6.4.3 Replace a new fuse, then screw the fuse cap.

6.5 Stop to use microscope, please cut off power, cover the dust cover, and place it in a cool and dry environment.

7. Troubleshooting

In the period of using BS-5062 series microscope, if there is any trouble occurs, please referring to the following sheet listed some common troubleshooting resolve them.

| Trouble | Causation | Remedy |
|--|--|--|
| Switch on but bulb dark | Plug is unreliable | Plug in again |
| | Bulb is broken | Change bulb |
| | Fuse is broken | Change fuse |
| Bulb is flickering or brightness is unsteady | Bulb is unstable | Insert it again |
| | Bulb is broken | Replacing bulb |
| Brightness of view field isn't enough or is Uneven | Bulb specification doesn't meet the requirement | Replacing bulb |
| | Brightness isn't adjusted correctly | Adjust rotation potentiometer |
| | Objective isn't in correct position | Make the objective in correct position |
| | The size of iris aperture is too small | Adjust the size of iris aperture |
| Brightness of view field isn't enough or is Uneven | Lens (objective, eyepiece, condenser, light collector) has dust | Clean it |
| | Position of condenser is too low | Higher condenser |
| Image isn't clear (contrast or definition isn't enough) | Cover glass of specimen doesn't meet the requirement | Use required thickness cover glass (0.17mm) |
| | Cover glass of specimen isn't in up direction | Place specimen correctly |
| | Surface of objective lens is dirty (especially it is easy for the front lens of 40X objective to dip in immersion oil) | Clean it |
| | Immersion oil isn't used for 100X objective (oil) | Use immersion oil |
| | Immersion oil doesn't meet the requirement | Use immersion oil supplied by us |
| | There is bubble in immersion oil | Clear the bubble way |
| | Size of iris aperture isn't proper | Adjust the size of iris aperture |
| | Position of condenser is too low | Readjust the position of condenser |
| One side of image is dark or image is moving as focusing | Objective isn't in correct position | Make the objective in correct position |
| | Specimen isn't placed correctly | Place specimen levelly on stage and clip it with clamp |
| Objective touches specimen as changing low times objective to high times objective | Cover glass of specimen isn't in up direction | Place specimen correctly |
| | Cover glass doesn't meet the requirement | Use required thickness cover glass (0.17mm) |
| Image observed by two eyes aren't in superposition entirely. | Interpupillary distance isn't adjusted correctly | Adjust interpupillary distance according to two eyes |
| It is easy for eyes to be tired during observing | Diopter isn't adjusted correctly | Readjust diopter |