

# SLIT LAMP

**OPERATOR'S MANUAL** 



Original instructions

# NIDEK CO., LTD.

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# SAFETY PRECAUTION

# 1.1 For Safe Use

BEFORE USE, READ THIS MANUAL.

Be sure to read the operator's manual prior to operation of the device to understand the safety precautions and operating procedures thoroughly.

♥ "1.3 Usage Precautions" (page 4)

Keep this manual handy for reference.

# 1.2 Safety Signal Words

In this manual, signal words are used to designate the degree or level of safety alerting. The definitions are as follows.

# 

This signal word indicates a potentially hazardous situation which, if not avoided, may result in death or serious injury.

# 

This signal word indicates a potentially hazardous situation which, if not avoided, may result in minor or moderate injury or property damage accident.

Even situations indicated by A CAUTION may result in serious injury under certain conditions. Safety precautions must be strictly followed at all times.

# **1.3 Usage Precautions**

#### Before use

#### 

- If any serious device-related incident occurs, report it to Nidek and the competent authority in the country where the user or patient, or both reside.
- Connect the power plug to a grounded outlet.
   Electric shock or fire may occur in the event of malfunction or power leakage.
- Never modify the device. Electric shock or malfunction may result.

# 

• Do not use this device for any purposes other than the intended purpose.

NIDEK is not responsible for accidents or malfunctions caused by misuse.

♥ Intended use" (page 11)

• Be sure to read the operator's manual prior to operation of the device to understand the safety precautions and operating procedures thoroughly. Use the accessories specified by NIDEK only. Use of the device outside the scope of this manual may cause unanticipated malfunctions or

adverse events.

• Use of this device is limited to doctors or health care providers authorized by a doctor.

Unanticipated malfunctions or adverse events may result. NIDEK is not responsible for accidents or malfunctions caused by misuse.

- Do not touch the interior of the device.
  - There are no parts within the device that requires servicing by the user.
- Only personnel qualified by NIDEK or an authorized distributor can install the device.
- Install the device on a level surface with sufficient space for assembly.
  - Fingers may be caught in the microscope arm or illumination unit arm if the device shifts unexpectedly.
- Use a stable optical table that does not topple easily.

The table may topple causing injury or damage to the device.

Install the device in an environment that meets the specifications.

The observation accuracy may be impaired, and malfunction may result. In addition, there is a possibility of injury if the device receives shock and falls down.

" Environmental conditions (during use)" (page 42)

- Install the device in an environment that meets the conditions listed below.
  - · A location not exposed to water
  - · A level and stable surface free from vibration and shock
  - A location with little external light
  - · A location with low dust
  - A location where condensation may not occur
- Always pay attention to the following points when operating the device.
  - Set the light intensity as low as possible.
  - Minimize the illumination area size as much as possible (slit width, slit length).
  - When projecting the illumination light, maximize the angle between it and the visual axis as much as possible.
  - Use the color filter, and in special cases, the diffuser.

• Before transporting the device, secure the fastening knobs to secure the main body, microscope arm, and illumination unit arm. Also, maximize the slit width.

Vibration or shock may cause the main body or arms to shift unexpectedly, resulting injury or malfunction of the device. In addition, if the device is transported with the slit closed, the slit may become unable to completely close during use.

· Observe the following instructions about the power supply.

Malfunction, electric shock, or fire may result.

- Be sure to use a (HOSPITAL GRADE) power outlet which meets the power specification requirements.
- Fully insert the power plug into a power outlet.
- Install the device so that the outlet that the power plug is inserted into is easily accessible during use. In addition, ensure that the power plug can be disconnected without the use of any tool.
- Before transporting the device, disconnect the power cord from the power inlet.
- · Do not place heavy objects on the power cord.
- Never use a power strip or extension cord to supply the device with power.
- Be sure to use the provided power cord. Do not connect the provided power cord to any other device.
- Before connecting the cords, turn off the power switch and disconnect the power cord from the power outlet.

Malfunction of the device may result.

• Insert the plug into the connector according to the proper indication and orientation and do not apply undue force to make the connections.

### **During use**

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• Before use, perform visual and operation checks. If abnormal conditions are encountered, stop using the device.

If the device is used under abnormal conditions, intended results may not be achieved. Also unanticipated malfunctions or health hazards may occur due to improper diagnosis.

- Do not perform servicing or maintenance on the device during use.
- Before and after use of the device, and before observing each patient, clean the forehead rest, chinrest, and grips (optional) with clean gauze or absorbent cotton. If necessary, dampen a cloth with rubbing alcohol and gently wipe them off.

If chinrest paper is used, remove one piece after each patient.

- · Wipe up any spills on the device immediately.
- When moving the device, take care that it does not come into contact with the patient.
- Take care not to catch hands or fingers in gaps between moving parts (such as between the main unit and table, between the main unit and base plate unit, near the microscope arm and illumination unit arm, and the lower end of the chinrest elevation control). Take care not to allow the patient to unnecessarily touch the device.

Hands or fingers may be caught in gaps resulting in injury.

• When the patient comes off from the device after observation, instruct the patient not to hold the chinrest support.

The device may topple over resulting in injury.

• Keep the optical parts such as lenses free of fingerprints and dust. Also confirm that they are not dirty before use.

The observation accuracy may decrease substantially.

- Do not turn forcefully the knobs beyond the range of movement. Malfunction of the device may result.
- Do not bump the device against the end of its movement range. Malfunction of the device may result.
- Be sure to set the light intensity to the minimum level at the beginning, and raise it as necessary when projecting light on the patient's eye. Be sure to return the light intensity to the minimum level after every examination.

If a high-intensity light is unnecessarily projected on the patient's eye, damage to the patient's retina may occur.

• Pay particular attention when projecting the illumination light into the eyes of infants, aphakic patients, and patients with eye disease.

These patients are more susceptible to adverse reaction to illumination light.

• In the event of smoke or strange odors, immediately turn off the device and disconnect the power plug from the power outlet. Check that the smoke ceases to be produced and contact NIDEK or your authorized distributor.

Use of the device under such abnormal conditions may cause fire or electric shock. In case of fire, use a dry chemical (ABC) extinguisher to extinguish the fire.

- If any of the following abnormalities in the power cord should be found, replace it with a new one immediately.
- Electric shock or fire may result.
  - The inside wires of the cord are exposed.
  - The power is intermittent when moving the cord.
  - The cord is excessively hot.
- After a long period of disuse, check for any abnormality before use.
- Should the device fail, disconnect the power cord from the power outlet and contact NIDEK or your authorized distributor without touching the interior of the device.
- Do not use an applanation tonometer other than the one specified for this device. Specified applanation tonometer: CSO Z800

#### Patient environment

The patient environment is the volume of space in which contact can occur between the patient and any part of the device or between the patient and any other person(s) touching the device.

Use devices that comply with IEC60601-1 in the patient environment.

If any device that does not comply with IEC 60601-1 is to be used, use an isolation transformer or common protective grounding.



#### After use

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- When transporting the device, lift it together with the optical table. Damage or malfunction of the device may result.
- When the device is not in use, turn off the power switch and place the dust cover over the device. Dust may affect the observation accuracy.
- If the device is not to be used for a long time, disconnect the power cord from the power outlet. Fire may result.
- When transporting the device, pack the device in the provided shipping carton. Excessive vibration or shock may cause malfunction of the device.
- Maintain the surrounding temperature and humidity in the specified environmental conditions during transport and storage of the device.

↔ "Environmental conditions (during transport)" (page 42), "Environmental conditions (during storage)" (page 42)

# Maintenance

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- Only service personnel trained by NIDEK can repair the device.
  - NIDEK will not be responsible for accidents caused by improper servicing.
- Before performing maintenance, wipe the surfaces of the device (especially, the areas that come into contact with the patient) with a clean cloth dampened with rubbing alcohol.
- When the device is sent back to NIDEK for repair, wipe the surfaces of the device (especially, the areas that come into contact with the patient) with a clean cloth dampened with rubbing alcohol.
- Do not use the device beyond its service life.
  - Even with proper maintenance and check, after time, the device reliability or safety may become degraded and fail to achieve the target values.
- To ensure the continued safe use of the device, it is recommended that the manager of this device make sure that maintenance and preventive inspection are performed at least once a year.

For details of maintenance and preventive inspection, ask NIDEK or your authorized distributor. If the manager of this device cannot perform the maintenance and preventive inspection, contact NIDEK or your authorized distributor.

# Disposal

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 Follow local governing ordinances and recycling plans regarding disposal or recycling of device components.

It is recommended to entrust the disposal to a designated industrial waste disposal contractor. Inappropriate disposal may contaminate the environment.

• When disposing of packing materials, sort them by material and follow local ordinances and recycling regulations.

Inappropriate disposal may contaminate the environment.

# 1.4 Labels and Symbols

To call attention to users, labels and indications are provided on the device. If labels are peeling off, characters are fading, or otherwise becoming illegible, contact NIDEK or your authorized distributor.

	This symbol indicates that the operator is advised to refer to the related instructions in the opera- tor's manual.
	☆ "5.1 Specifications" (page 41)
-	This symbol indicates that the degree of protection against electric shock is of a Type B Applied
<b>.</b>	Part. The applied parts are the forehead rest, chinrest, and grips (optional)
Λ	↔ * Base plate unit" (page 14)
$\sim$	This symbol indicates that the device must be supplied only with alternating current.
	This symbol indicates that the connection point is suitable for direct current only.
I	This symbol indicates the state of the power switch. When this symbol side of the switch is pressed down, power is supplied to the device.
(	This symbol indicates the state of the power switch. When this symbol side of the switch is
0	pressed down, power is not supplied to the device.
п	
M	This symbol indicates the year of manufacture.
	This symbol indicates the manufacturer
	This symbol indicates that this product is to be disposed of in separate collection of electrical and
X	electronic equipment in EU.
MD	Medical device
EC REP	EU Authorized Representative
-ਲ਼ੑੑ੶	This symbol indicates the illumination control that adjusts the intensity of the illumination light.
<b>つ</b>	This symbol indicates the adjustment amount of illumination light.
-'ਲ਼ੑੑ੶ BL	This symbol indicates the illumination control of the optional external illumination.
	This symbol indicates the adjustment amount of the optional external illumination light.



INTRODUCTION

# 2.1 Device Outline

The NIDEK SLIT LAMP SL-2000 is used to magnify the eyeball, eyelid, and eyelash of patients for observation, using slit illumination light.

# Intended use

The SLIT LAMP SL-2000 is intended for use in eye examination of the anterior eye segment, from the corneal epithelium to the posterior capsule. This device is used to aid in the diagnosis of diseases or trauma which affect the structural properties of the anterior eye segment.

# Intended patient population

• Age

All ages except babies and infants

• Weight

The maximum weight bearing 1,350 N (estimated from the forehead rest and chinrest load capacity [100 N] with an assumption that the head weighs 7.4% of human body)

Health condition

Able to undergo an examination while seated

• Conditions - Visual function

One or both eyes are normal or have disease. Eyes with visual impairment are excluded.

# Intended user profile

Ophthalmologists, optometrists (OD), orthoptists (ORT)

# 2.2 Principles

Slit light is projected at an angle onto the ocular media such as the cornea or crystalline lens to optically section the eyeball. Minute lesions are highlighted by the light diffusion in the tissue. The stereo microscope magnifies any lesions for observation.

# 2.3 Configuration and Functions

# Main unit



#### 1 Filter insertion rod

The filter insertion rod is used to insert or remove the yellow filter into or from the observation optical path.

#### **2** Binoculars

The binoculars enable stereoscopic observation of the patient's eye. Adjust the pupillary distance to suit the operator.

#### **3** Diopter adjustment rings

The diopter adjustment rings are used to adjust the diopter.

#### 4 Eyecups

The cups slide forward and backward. Usually, use the eyepieces with the cups pulled out. For eyeglasses wearers, push in the cups.

#### 5 Magnification changer

The magnification changer is used to select the total magnification of the microscope.

#### 6 Filter changer

The filter changer is used to insert a filter into the illumination optical path.

#### 7 Slit rotation control

The slit rotation control is used to rotate the slit. A click is felt in the horizontal and vertical axes.

#### 8 Slit width control

The slit width control is used to adjust the slit width of the illumination light.

#### 9 Slit length control

The slit length control is used to adjust the aperture diameter or slit length of the illumination light.

#### 10 Joystick

The joystick is used to adjust the position of the main unit. By manipulating the joystick right, left, forward, and backward, the main unit moves right, left, forward, and backward. By tilting the joystick, the main unit slightly moves to the direction of tilt. By rotating the joystick, the main unit moves up and down.



#### **11** Illumination control

The illumination control is used to adjust continuously the intensity of the illumination light. When the illumination control is turned fully counterclockwise, a click is felt. Turning it until a click is felt turns off the illumination light.

#### 12 Prism head

The prism head is used to project the illumination light on the patient's eye.

#### 13 Diffuser

The diffuser is used to diffuse the illumination light throughout the visual field.

#### 14 Focusing rod

The focusing rod is inserted into the mounting hole on the arm, and used to adjust eyepiece diopter and pupillary distance.

#### 15 Scanning knob

The scanning knob is used to move the illumination light right and left. A click is felt in the middle.

#### 16 Cap

#### 17 Axis angle scale

The axis angle scale indicates the angle between the microscope arm and illumination unit arm. A click is felt at the angle of  $0^{\circ}$ .

#### 18 Illumination unit arm fastening knob

#### **19** Microscope arm fastening knob

#### 20 Main unit power cord

The main unit power cord supplies power to the main unit.

#### 21 Main unit fastening knob

### Base plate unit



#### 1 Fixation lamp

The fixation lamp is used to steady the patient's visual axis by asking the patient to focus on the lamp. The position of the fixation lamp can be adjusted by the flexible arm.

#### 2 Forehead rest

#### 3 Chinrest

#### 4 Chinrest elevation control

The chinrest elevation control is used to adjust the height of the chinrest.

#### 5 Slide plate

The slide plate reduces friction to enable the device to smoothly move along all axes.

#### 6 Eye level marker

The patient's eyes are aligned to this height.

#### 7 Fixing pins for chinrest paper

The fixing pins are used to attach the stack of chinrest paper onto the chinrest.

#### 8 Fixation lamp cord

The fixation lamp cord supplies power to the fixation lamp.

# Power supply box



#### 1 Pilot lamp

The pilot lamp illuminates when the power switch is on (

#### 2 Power switch

The power switch is used to turn on  $( \ )$  or off  $( \bigcirc )$  power to the slit lamp.

#### **3** Power inlet

The power inlet is used to connect the female plug of the power cord.

#### 4 Fixation lamp connector

The fixation lamp connector is used to connect the fixation lamp cord.

#### 5 Main unit power connector

The main unit power connector is used to connect the main unit power cord.

# External illumination (optional)



#### **1** External illumination

#### 2 External illumination control

The external illumination control adjusts continuously the intensity of the external illumination light. A click is felt when it is turned fully counterclockwise. When it is turned until a click is felt, the external illumination is turned off.

#### **3** External illumination button

Pressing this button turns on the external illumination. Pressing this button again turns off the external illumination.

When the external illumination control is at the position where a click is felt, the external illumination is not turned on even when the button is pressed.

# 2.4 Packed Contents

Part name	Quantity	Appearance
Focusing rod	1 unit	ET. D
Сар	1 unit	
Chinrest paper	2 packs	
Power cord	1 unit	
Hexagonal wrench (2 mm)	1 unit	
Phillips screwdriver	2 units	
Round head wood screws (4.1 x 13, 3.8 x 20)	2 units each	OTHE OTHER OF THE OTHER
Dust cover	1 unit	
Operator's manual	1 volume	

The following are included in the standard configuration. Check the contents before use.

# 2.5 Attachment Instructions

# 2.5.1 Attachment of accessories

# Prism head

**1** Insert the prism head <sup>(\*A)</sup> into the groove of the lamp housing <sup>(\*B)</sup>.



**2** Tighten the fastening screw <sup>(\*C)</sup> of the prism head with the Phillips screwdriver to secure the prism head.



# Binoculars

🥢 Note

binoculars (\*C).

**1** Using the hexagonal wrench, insert the fastening screw <sup>(\*A)</sup> into the microscope mount <sup>(\*B)</sup> without allowing the screw to protrude into the interior.

cannot be mounted on the microscope mount.



• If the fastening screw protrudes through the microscope mount, the binoculars

**3** Using the hexagonal wrench, tighten the fastening screw to secure the binoculars.

**2** Into the frame of the microscope mount, insert the







# Cords

Connect the cords to the power supply box.

- **1** Connect the power cord  $^{(*D)}$  to the power inlet  $^{(*A)}$ .
- **2** Connect the fixation lamp cord <sup>(\*E)</sup> to the fixation lamp connector <sup>(\*B)</sup>.
- **3** Connect the main unit power cord <sup>(\*F)</sup> to the main unit power connector <sup>(\*C)</sup>.
- **4** Confirm that the power switch is turned off  $(\bigcirc)$  and plug the power cord into the power outlet.



# 

• Connect the power plug to a grounded outlet. Electric shock or fire may occur in the event of malfunction or power leakage.

# 2.5.2 Attachment of optional accessories

Only personnel qualified by NIDEK or an authorized distributor can attach the optional accessories. The general attachment instructions are described below.

# 16x eyepieces

- **1** Remove the magnification changer <sup>(\*A)</sup>.
  - 1) Turn the magnification changer to the "32" position.

- 2) Remove the cap on the screw hole.
- 3) Remove the screw with the Phillips screwdriver to remove the magnification changer.
- **2** Mount the magnification changer for the 16x eyepieces.
  - 1) Turn the magnification changer for the 16x eyepieces to the position between the "32" and "40" positions.
  - 2) Mount it in the reverse order of how it was removed.

White characters: Magnifications when 12.5x eyepieces are attached

Green characters: Magnifications when 16x eyepieces are attached

# 🥢 Note

• Properly mount the magnification changer as shown in the illustration.

The magnification indications on the magnification changers are mirrored. If the right and left magnification changers are confused, the magnification indications do not correspond to the actual magnifications.

**3** Mount the other magnification changer in the same manner.





2



**4** Change the two eyepieces <sup>(\*B)</sup>.

- 1) Hold the base of the eyepieces and pull them out.
- 2) Completely insert the 16x eyepieces.



# Grip

**1** Remove the cap <sup>(\*A)</sup> on the mounting hole for the grip.

Using a thin rod such as a hexagonal wrench, push out the cap from the interior.





# Beam splitter

**1** Using the hexagonal wrench, loosen the fastening screw <sup>(\*A)</sup> to remove the binoculars <sup>(\*B)</sup>.

- **2** Mount the beam splitter on the microscope mount.
  - 1) Into the frame of the microscope mount <sup>(\*C)</sup>, insert the beam splitter <sup>(\*D)</sup>.
  - 2) Using the hexagonal wrench, tighten the fastening screw to secure the beam splitter.







**3** Mount the binoculars on the beam splitter.

Into the frame of the beam splitter, insert the binoculars (\*E).



2) Tighten the fastening screw (\*F) of the beam splitter to secure the binoculars.



# C-mount camera adapter

- 1 Mount the beam splitter.
  ★ \*• Beam splitter" (page 23)
- **2** Mount the camera <sup>(\*A)</sup> on the C-mount camera adapter <sup>(\*B)</sup>.

Align the camera connection part with the C-mount camera adapter connection part. Secure the C-mount camera adapter to the camera by rotating the C-mount camera adapter.

- **3** Mount the C-mount camera adapter on the beam splitter.
  - 1) Unscrew the cap <sup>(\*C)</sup> attached to the mounting hole on the beam splitter.
  - 2) Mount the C-mount camera adapter.

Align the C-mount camera adapter connection part with the mounting hole on the beam splitter. Rotate the securing ring  $^{(*D)}$  to secure the adapter.





- **4** While viewing the monitor connected to the camera, adjust the focus and orientation of the slit image or adjust the position of the slit image.
  - Focus and orientation of the slit image
    - Move the camera connection part <sup>(\*E)</sup> of the C-mount camera adapter vertically to adjust the focus of the slit image. Rotate the camera connection part to adjust the orientation of the slit image.
    - Using the provided hexagonal wrench (1.5 mm), tighten the three fastening screws <sup>(\*F)</sup> to secure the camera connection part.



Position of the slit image

Using the provided hexagonal wrench (1.27 mm), turn the three adjustment screws <sup>(\*G)</sup> to adjust the internal mirror by changing the tightness of the adjustment screws.

Adjust the slit image so that it appears in the center of the monitor.



# Co-observation scope

- 1 Mount the beam splitter.
  ★ \* Beam splitter" (page 23)
- **2** Mount the co-observation scope on the beam splitter.
  - Unscrew the cap (\*A) attached to the beam splitter mount.
  - 2) Mount the co-observation scope.

Align the co-observation scope connection part with the mounting hole on the beam splitter. Rotate the securing ring <sup>(\*B)</sup> to secure the co-observation scope.



# 2.5.3 Attachment of applanation tonometer

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• Do not use an applanation tonometer other than the one specified for this device. Specified applanation tonometer: CSO Z800

- **1** Rest the tonometer mount <sup>(\*A)</sup> on the microscope mount and tighten the fastening screw <sup>(\*B)</sup> to secure the tonometer mount.
- **2** Mount the applanation tonometer <sup>(\*C)</sup> on the tonometer mount support.



**3** Adjust the height of the applanation tonometer.

1) While looking into the eyepieces, adjust the height so that the cone <sup>(\*D)</sup> of the applanation tonometer is aligned to the center of the visual field.

#### 🥢 Note

- Projecting the illumination light on the cone of the applanation tonometer makes it easier to check the position of the cone.
- Using the provided hexagonal wrench, tighten the two adjustment screws <sup>(\*E)</sup> to secure the applanation tonometer.



\* For the operating procedure, refer to the operator's manual of the applanation tonometer.

# 2.6 Beep Sound

Device operations and errors are indicated by beep sounds.

For procedure for changing the beep sound volume setting, contact NIDEK or your authorized distributor.

• At device start-up

After the power switch is turned on ( ), a beep sounds when the device is ready.

• When the upper or lower limit of the main unit movement is detected

If the main unit has reached the upper or lower limit of its movement range when the joystick is manipulated, a beep sounds.

• When abnormalities are detected

A beep sounds repeatedly when errors occur in the device.

Turn off  $(\bigcirc)$  the power switch, then contact NIDEK or your authorized distributor.



# **OPERATING PROCEDURE**

# 3.1 Operation Flow

Device power ON

*"3.2 Preparation for Observation" (page 30)* Turn on power to the device and change the settings of the device if necessary. Conduct patient preparation.

Observation

"3.3 Operating Procedure" (page 32)

Device power OFF

"3.4 End of Operation" (page 35)

# 3.2 Preparation for Observation

**1** Minimize the intensity of the illumination light.

Turn the illumination control fully counterclockwise.

• For devices with the optional external illumination

Turn the external illumination control fully counterclockwise and turn off the external illumination button.

**2** Turn on (|) the power switch <sup>(\*A)</sup>.

Confirm that the pilot lamp (\*B) illuminates.

# 

• Do not turn on ( ) the power switch while the illumination control is turned fully clockwise.

**3** Loosen the fastening knobs to allow the secured parts of the device to move freely.

There are three types of fastening knobs: the illumination unit arm fastening knob <sup>(\*C)</sup>, microscope arm fastening knob <sup>(\*D)</sup>, and main unit fastening knob <sup>(\*E)</sup>.







### **4** Perform checks before use.

Perform the following checks before use.

- The optical parts such as lenses are clean.
- The main unit moves smoothly using the joystick.
- The illumination light is projected.
- The fixation lamp illuminates.
- The optional external illumination lights up / turns off. If any abnormality is found, stop using the device, then see *"4.1 Troubleshooting" (page 37)*.

- **5** Insert the focusing rod.
  - 1) Remove the cap on the mounting hole  $({}^{*}F)$ .
  - 2) Insert the focusing rod into the mounting hole  $({}^{*}F)_{-}$







- Turn the focusing rod so that its flat surface faces the microscope.
- 2) Project the illumination light on the focusing rod.
- Fully turn the diopter adjustment rings (\*G) to the + side and look into the eyepieces.
- 4) Observe the slit image with one eye.
- 5) Slowly turn the diopter adjustment ring <sup>(\*G)</sup> from the + side to the side until the slit image is focused sharply.

• Be sure to adjust the eyepiece diopter for each eye and do not turn the diopter adjustment ring from the - side to the + side.

The eyepiece diopter cannot be properly adjusted.

**7** Adjust the pupillary distance of the binoculars.





Before adjustment

After adjustment



**8** After the adjustment, remove the focusing rod and attach the cap.

# 3.3 Operating Procedure

# **1** Clean the areas that come into contact with the patient.

Wipe the forehead rest <sup>(\*A)</sup>, chinrest <sup>(\*B)</sup>, and grips (optional) with clean absorbent cotton or gauze dampened with rubbing alcohol.

If chinrest paper is used, remove one piece after each patient.

To attach the chinrest paper, see 4.3 *Chinrest Paper Attachment*" (page 39).

**2** Conduct patient preparation.

- 1) Instruct the patient to remove their glasses or contact lenses and sit on the chair.
- Adjust the height of the optical table and patient's chair so that the patient can easily rest their chin on the chinrest.
- Instruct the patient to place their chin on the chinrest as far forward as possible with their forehead resting gently on the forehead rest.
- 4) Rotate the chinrest elevation control <sup>(\*C)</sup> to align the level of the patient's eye with the eye level marker <sup>(\*D)</sup>.
- 5) If necessary, adjust the position of the patient's eye using the fixation lamp <sup>(\*E)</sup>.
- **3** Manipulate the joystick to project the illumination light on the patient's eye.

By manipulating the joystick, the main unit moves right, left, forward, and backward. By rotating the upper part of the joystick, the main unit moves up and down.







# 

• When moving the device, take care that it does not come into contact with the patient.

• Do not bump the device against the end of its movement range. Malfunction of the device may result.

- **4** Adjust the intensity of the illumination light by turning the illumination control.



If necessary, conduct the following procedures.

1) Change the magnification of the microscope with the magnification changer (\*F).

Microscope magnifica- tion	Diameter of real field of view
5x	40.7 mm
8x	25.7 mm
12.5x	16.1 mm
20x	10.1 mm
32x	6.4 mm

- 2) Move the slit image to the right or left with the scanning knob (\*G) or rotate the slight image with the slit rotation control  $(^{*H})$ .
- \*J \*K 3) Select the desired filter with the filter changer

(*I) <sub>.</sub>		
Index	Filter name	Effect
Blue	Blue filter	Fluorescence and staining reaction
Green	Red-free filter	Red-free
Gray	ND filter	Light reduction
White	(Free aperture)	_

\*G

\*I

\*H

4) Adjust the slit width with the slit width control (\*J).

Adjustment range: 0 to 14 mm (continuously variable)

#### 🥢 Note

- As a guide, the slit width control has scales marked at 1, 2, 5, and 10 mm positions.
- The maximum slit width is the same as the maximum aperture diameter. Therefore, when the slit width is 14 mm (the maximum width), the shape of the illumination light becomes a circle.
- 5) Change the aperture diameter and slit length with the slit length control (\*K). Adjustment range: ø0.2, 1, 3, 5, 9, 14 mm



6) Lift the diffuser <sup>(\*L)</sup> to mount it on the illumination light aperture.

Using the diffuser diffuses the illumination light throughout the visual field.

7) Insert the yellow filter.

Pulling up the filter insertion rod  $^{(^{\ast}\mathrm{M})}$  inserts the yellow filter.

 For devices with the optional external illumination, turn on the external illumination button (<sup>\*N)</sup> to turn on the illumination light.

Adjust the intensity of the external illumination light by turning the external illumination control  $(^{*O})$  clockwise. It is used to illuminate the background for image capture with a camera.







**6** Minimize the intensity of the illumination light after the observation is complete.

Turn the illumination control fully counterclockwise.

For devices with the optional external illumination
 Turn the external illumination control fully counterclockwise.

**7** To observe the eyes of the next patient, repeat from Step 1.

# 3.4 End of Operation

**1** Minimize the intensity of the illumination light.

Turn the illumination control fully counterclockwise.

- · For devices with the optional external illumination
- Turn the external illumination control fully counterclockwise and turn off the external illumination button.
- **2** Turn off  $(\bigcirc)$  the power switch.
- **3** Clean the areas that come into contact with the patient (forehead rest, chinrest, and optional grips).

For cleaning, see Step 1 in "3.3 Operating Procedure" (page 32).

- **4** Move the device to the center of the base and secure the fastening knobs.
- **5** Place the dust cover on the device.



• When the device is not in use, place the dust cover over the device.



MAINTENANCE

# 4.1 Troubleshooting

Should the device function improperly, attempt to correct the problem according to the following table before contacting NIDEK or your authorized distributor.

When	Remedy		
The pilot lamp does not illumi- nate even though the power switch is on ( ).	<ul> <li>The power cord may not be correctly connected. Reconnect it securely.</li> <li>Check whether proper voltage is applied to the power outlet.</li> </ul>		
The illumination light is not pro- jected.	<ul> <li>The main unit power cord may not be correctly connected to the power supply box. Reconnect it securely.</li> <li>When the slit is closed, the illumination light is not projected. Use the slit width control to open the slit.</li> <li>The illumination control may be turned to the position where the illumination light is turned off. Turn the illumination control clockwise.</li> </ul>		
The fixation lamp does not illu- minate.	<ul> <li>The fixation lamp cord may not be correctly connected to the power supply box.</li> <li>Reconnect it securely.</li> </ul>		
The device does not move up and down.	<ul> <li>The device has reached the upper or lower limit of its movement range. Rotate the joystick in an opposite direction.</li> <li>When the device has reached the upper or lower limit of its movement range, a beep sounds.</li> </ul>		
The optional external illumina- tion does not turn on.	<ul> <li>The external illumination button may be turned off. Turn on the external illumination button.</li> <li>The external illumination control may be turned to the position where the illumination light is turned off (turned fully counterclockwise). Turn the external illumination control clockwise.</li> </ul>		

If the symptom cannot be corrected by the above actions, contact NIDEK or your authorized distributor.

# 4.2 Cleaning

# 

- Never use an organic solvent such as paint thinner or abrasive solvents. It may damage the surface of the device.
- Never use a sponge or cloth soaked in water. The water may leak into the inside of the device and cause device failure.
- When wiping the prism head with a sponge or cloth, do not scrub the prism cover <sup>(\*A)</sup> forcibly.

The coating may be scratched. If the coating is dirty or scratched, stray light may enter the microscope visual field causing adverse effects on observation.

# Device exterior

Wipe the device exterior except for optical parts with a soft, dry cloth.

For severe stains, soak the cloth in a neutral detergent, wring well, and wipe. Finally dry with a soft, dry cloth.

\*A

# Optical parts such as lenses

First, use a blower to blow off any dust from the optical parts such as lenses. For severe stains, gently wipe the part with clean lens cleaning paper dampened with alcohol while taking care not to scratch the surface.

If the parts are not cleaned satisfactorily, repeat the cleaning procedure with new cleaning paper until they become clean.

# 4.3 Chinrest Paper Attachment

- **1** Remove the two fixing pins from the chinrest.
- **2** Remove a suitable number of chinrest papers from the pack. An entire pack of chinrest paper cannot be attached. Use 20 to 30-sheet stacks of chinrest paper.
- **3** Pass the fixing pins through the chinrest paper stack. Pass the fixing pins through the holes on either side of the stack of chinrest paper.



**4** Attach the stack of chinrest paper onto the chinrest.

- 1) Insert the pin into the hole of the chinrest while holding both fixing pin and stack of paper.
- 2) Push the other pin into the hole of the chinrest.



# SPECIFICATIONS AND TECHNICAL INFORMATION

# 5.1 Specifications

Device specifications			
Microscope	Microscope type	Galilean converging binocular, five-level magnification (rotating drum)	
	Objective lens focal length	125 mm	
	Eyepieces	12.5x	
	<ul> <li>Total magnification / diameter of real field of view</li> </ul>	5x / 40.7 mm, 8x / 25.7 mm, 12.5x / 16.1 mm, 20x / 10.1 mm, 32x / 6.4 mm	
	Interpupillary adjust- ment range	50 to 78 mm	
	Stereoscopic angle	6°	
	Diopter adjustment range	±8 D	
	Yellow filter	For fluorescent observation (the filter can be inserted or removed using the filter insertion rod)	
Base	Right and left movement distance	110 mm	
	Forward and backward movement distance	100 mm	
	• Forward and backward, right and left fine move- ment distance	20 mm (when the joystick is tilted)	
	Up and down movement distance	30 mm (motorized)	
	Chinrest up and down movement distance	80 mm	
	Fixation lamp	LED lamp (green)	

Device specifications				
Lamp housing	Projection magnification	1x		
	Slit width	0 to 14 mm (continuously variable)		
	Aperture diameter / slit length	0.2, 1, 3, 5, 9, 14 mm		
	Slit rotation	±90°		
	• Filter	Blue filter: Fluorescence and staining reaction Red-free filter: Red-free ND filter: Light reduction		
	Illumination lamp	LED lamp (white)		
	Lighting control	Minimum to maximum (continuously variable)		
External illumination	Illumination lamp	LED lamp (white): 2 units		
(optional)	Lighting control	Minimum to maximum (continuously variable)		
Power supply	Input voltage	AC 100 to 240 V 50/60 Hz		
	Power consumption	75 VA		
Dimensions and	Dimensions	346 (W) × 433 (D) × 567 (H) mm		
mass	• Mass	12 kg		
Environmental condi-	• Temperature	10 to 35°C (50 to 95°F)		
tions (during use)	Humidity	30 to 90% (non-condensing)		
	Atmospheric pressure	800 to 1,060 hPa		
Environmental condi-	• Temperature	-30 to 60°C (-22 to 140°F)		
tions (during trans- port)	Humidity	10 to 95% (non-condensing)		
	Atmospheric pressure	700 to 1,060 hPa		
Environmental condi-	• Temperature	-10 to 55°C (14 to 131°F)		
tions (during storage)	Humidity	10 to 95% (non-condensing)		
	Atmospheric pressure	700 to 1,060 hPa		
Others	<ul> <li>Expected service life (defined by manufac- turer)</li> </ul>	8 years from the date of the initial operation <ul> <li>Proper maintenance is necessary.</li> </ul>		
	Packing unit	1 unit		
Classifications	<ul> <li>Form of protection against electric shock: Class I</li> <li>Form of protection against electric shock (applied parts): Type B applied part</li> <li>Degree of protection against harmful ingress of water or particulate matter: IPXI</li> <li>Degree of safety application in the presence of flammable anesthetics and/or fla mable cleaning agents: Device which should be used in an environment where r flammable anesthetics and/or flammable cleaning agents are present</li> <li>Degree of suitability for use in an oxygen rich environment: Device which is not intended for use in an oxygen rich environment</li> <li>Mode of operation: Continuous operating device</li> </ul>			

Accessories	
Standard accesso- ries	<ul> <li>Focusing rod, cap, chinrest paper (2 packs), power cord, hexagonal wrench (2 mm), Phillips screwdriver (2 units), round head wood screws (4.1 x 13, 3.8 x 20), dust cover, Operator's Manual</li> </ul>
Optional accessories	• 1 pair of 16x eyepieces, 1 pair of grips (for patients), beam splitter, C-mount camera adapter, tonometer mount <sup>*1</sup> , co-observation scope, external illumination

\*1. The following applanation tonometer can be attached to the tonometer mount.

• CSO: Z800

For details, refer to the operator's manual of the applanation tonometer.

# 5.2 Light Hazard

Provision of information on the avoidance of light hazard from the optical device is required in ISO 15004-2:2007 "Ophthalmic instruments - Fundamental requirements and test methods -".

#### 

• The light emitted from this device is potentially hazardous. The longer the exposure time is, the greater the risk of ocular damage becomes.

Exposure to light from this device when operated at maximum intensity <sup>(\*A)</sup> will exceed the safety guideline after the following duration.

- Illumination light Normal use: 200 seconds
   With a 90 D lens: 1,300 seconds
   Illuminance: 0 to 200,000 lx
- External illumination light (optional external illumination) 924 seconds Illuminance: 0 to 5,000 lx



#### • Relative spectrum



# **5.3 EMC (Electromagnetic Compatibility)**

The device is suitable for use in hospitals except for near active HF surgical equipment and RF shielded rooms with an ME system for magnetic resonance imaging, where the intensity of electromagnetic disturbances is high.

# 

• Do not use the device near, on, or under other electronic equipment. Otherwise, it could result in improper operation. If such use is necessary, the device and the other equipment should be observed to verify that they are operating normally.

- Use of accessories, cables other than those specified or provided by the manufacturer of this equipment could result in increased electromagnetic emissions or decreased electromagnetic immunity of this equipment and result improper operation.
- Portable RF communications equipment (including peripherals such as antenna cables and external antennas) should be used no closer than 30 cm (12 inches) to any part of the device, including the specified or provided cables.

Otherwise, degradation of the performance of this equipment could result.

· Electromagnetic disturbance could blink or turn off the slit light.

# Specified cable

Part name	Connector Shielded	Cable Shielded	Ferrite Core	Length (m)
Power cord for the SL- 2000 (HG) <sup>*1</sup>	No	No	No	2.5
Fixation lamp cord	Yes	Yes	No	0.7
Main unit power cord	Yes	Yes	No	0.9
Power cord for the SL- 2000 (CE) <sup>*2</sup>	No	No	No	2.5

\*1: Test voltage AC 100 V 50 Hz

\*2: Test voltage AC 230 V 50/60 Hz, AC 240 V 50 Hz

# Essential performance

- · Slit illumination light projection function
- Observation function using converging binocular microscope

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#### Guidance and manufacturer's declaration - electromagnetic emissions

The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environment.

Emissions test	Compliance	Electromagnetic environment - guidance
RF emissions CISPR 11	Group 1	The device uses RF energy only for its internal function. Therefore, its RF emissions are very low and are not likely to cause any interference in nearby electronic equipment.
RF emissions CISPR 11	Class B	The device is suitable for use in all establishments, including domestic
Harmonic emissions IEC 61000-3-2	*1	establishments and those directly connected to the public low-voltage power supply network that supplies buildings used for domestic purposes.
Voltage fluctuations/ Flicker emissions IEC 61000-3-3	*2	

\* 1 For the regions where the rated voltage is 220 V or greater, this device complies with class A. For the regions where the rated voltage is 127 V or less, this standard is not applicable.

\* 2 For the regions where the rated voltage is 220 V or greater, this device complies with this standard. For the regions where the rated voltage is 127 V or less, this standard is not applicable.

Guidance and manufacturer's declaration - electromagnetic immunity				
The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environment.				
Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance	
Electrostatic Discharge (ESD) IEC 61000-4-2	±8 kV contact ±2 kV, ±4 kV, ±8 kV, ±15 kV air	±8 kV contact ±2 kV, ±4 kV, ±8 kV, ±15 kV air	Floor should be wood, concrete or ceramic tile. If floors are covered with synthetic material, the relative humidity should be at least 30%.	
Electrical fast transient/burst IEC 61000-4-4	±2 kV for power supply lines ±1 kV for input/output lines 100 kHz repetition frequency	±2 kV for power supply lines ±1 kV for input/output lines 100 kHz repetition frequency	Mains power quality should be that of a typical commercial or hospital environment.	
Surge IEC 61000-4-5	±1 kV differential mode ±2 kV common mode	±1 kV differential mode ±2 kV common mode	Mains power quality should be that of a typical commercial or hospital environment.	
Voltage, dips, short interruptions and voltage variations on power supply input lines IEC 61000-4-11	Dips: $0\% U_{T}$ in 0.5 cycle at 0°, $45^{\circ}$ , 90°, 135°, 180°, $225^{\circ}$ , 270°, and 315° $0\% U_{T}$ in 1 cycle and 70% $U_{T}$ in 25/30 cycles in single phase (at 0°) Short interruptions: $0\% U_{T}$ in 250/300 cycle	Dips: $0\% U_{T}$ in 0.5 cycle at 0°, $45^{\circ}$ , 90°, 135°, 180°, $225^{\circ}$ , 270°, and 315° $0\% U_{T}$ in 1 cycle and 70% $U_{T}$ in 25/30 cycles in single phase (at 0°) Short interruptions: $0\% U_{T}$ in 250/300 cycle	Mains power quality should be that of a typical commercial or hospital environment. If the user of the device requires continued operation during power mains interruptions, it is recommended that the device be powered from an uninterruptible power supply or a battery.	
Power frequency (50/ 60 Hz) magnetic field IEC 61000-4-8	30 A/m 50 Hz or 60 Hz	30 A/m 50 Hz or 60 Hz	Power frequency magnetic fields should be at levels characteristic of a typical location in a typical commercial or hospital environment.	
NOTE $U_{\rm T}$ is the a.c. mains voltage prior to application of the test level.				

Guidance and manufacturer's declaration - electromagnetic immunity				
The device is intended for use in the electromagnetic environment specified below. The customer or the user of the device should assure that it is used in such an environment.				
Immunity test	IEC 60601 test level	Compliance level	Electromagnetic environment - guidance	
Conducted RF IEC 61000-4-6	3 V 0.15 MHz to 80 MHz	3 V 0.15 MHz to 80 MHz	Portable and mobile RF communications equipment should be used no closer than 30 cm to any part of the device, including cables. <sup>a</sup>	
	6 V in ISM bands between 0.15 MHz and 80 MHz 80% AM at 1 kHz	6 V in ISM bands between 0.15 MHz and 80 MHz 80% AM at 1 kHz		
Radiated RF IEC 61000-4-3	3 V/m 80 MHz to 2.7 GHz 80% AM at 1 kHz	3 V/m 80 MHz to 2.7 GHz 80% AM at 1 kHz		
Proximity field from RF wireless communications equipment IEC 61000-4-3	See "Test specifications for enclosure port immunity to RF wireless communications equipment".	See "Test specifications for enclosure port immunity to RF wireless communications equipment".		

NOTE 1 At 80 MHz and 800 MHz, the higher frequency range applies.

NOTE 2 These guidelines may not apply in all situations. Electromagnetic propagation is affected by absorption and reflection from structures, objects and people.

а Field strengths from fixed transmitters, such as base stations for radio (cellular/cordless) telephones and land mobile radios, amateur radio, AM and FM radio broadcast and TV broadcast cannot be predicted theoretically with accuracy. To assess the electromagnetic environment due to fixed RF transmitters, an electromagnetic site survey should be considered. If the measured field strength in the location in which the device is used exceeds the applicable RF compliance level above, the device should be observed to verify normal operation. If abnormal performance is observed, additional measures may be necessary, such as reorienting or relocating the device.

Test specifications for enclosure port immunity to RF wireless communications equipment												
Test frequency (MHz)	Band <sup>a</sup> (MHz)	Service <sup>a</sup>	Maximum power (W)	Distance (m)	Immunity test level (V/m)							
385	380 - 390	TETRA 400	1.8	0.3	27							
450	430 - 470	GMRS 460, FRS 460	2	0.3	28							
710												
745	704 - 787	LTE Band 13, 17	0.2	0.3	9							
780												
810												
870	800 - 960	GSM 800/900, TETRA 800, iDEN 820, CDMA 850, LTE Band 5	2	0.3	28							
930												
1,720												
1,845	1,700 - 1,990	GSM 1800, CDMA 1900, GSM 1900, DECT LTE Band 1, 3, 4, 25: UMTS	2	0.3	28							
1,970												
2,450	2,400 - 2,570	Bluetooth, WLAN, 802.11: b/g/n, RFID 2450, LTE Band 7	2	0.3	28							
5,240		N/LAN 000 44										
5,500	5,100 - 5,800	wlan 802.11 a/n	0.2	0.3	9							
5,785		Corri										
NOTE: If nece system may b	essary to achieve e reduced to 1 n	e the immunity test level, the distance be n. The 1 m test distance is permitted by I	tween the transmitting EC 61000-4-3.	antenna and the ME	equipment or ME							

For some services, only the uplink frequencies are included.



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13.1.This Agreement constitutes the entire agreement between you and NIDEK concerning the Software, and supersedes any prior written or oral agreement between you and NIDEK. No modification of this Agreement will be binding unless otherwise agreed in writing.

#### 14. NO WAIVER

14.1.The failure of NIDEK to enforce at any time or for any period the provisions hereof in accordance with its terms will not be construed to be a waiver of such provisions or of the rights thereafter to enforce each and every provision.

#### **15. NO THIRD PARTY RIGHTS**

15.1.This Agreement is intended to be solely for the benefit of you and NIDEK and is not intended to confer any benefits upon or create any rights in favor of any person other than you and NIDEK.

#### 16. HEADINGS

16.1.All headings are for convenience only and will not affect the meaning of any provision of this Agreement.

#### 17. LANGUAGE

- 17.1.The license agreement for the Software may be provided in multiple languages. In such event, unless otherwise agreed in writing, the following shall apply:
- (i) If you use the Software in any countries outside Japan, the license agreement for the Software shall be executed and delivered in a text using the English language. The text using the English language shall prevail and control; and
- (ii) If you use the Software in Japan, the license agreement for the Software shall be executed and delivered in a text using Japanese language. The text using the Japanese language shall prevail and control.