

Instructions for Use

REF

Anesthesia Screen 800-0016

INTENDED USE

Intended use is to aid anesthesiologist in viewing patient during surgery. The intended users of this device are medical professionals within hospitals and surgery centers.

INSTRUCTIONS FOR USE

Become familiar with the features of patient positioning device before use with a patient. Always practice use on a nurse, physician or appropriate volunteer prior to using clinically.



Attaching to Side Rail

- 1. Attach Schure Socket XL (sold separately) on rail. Insert mounting post into socket. Lock position by turning handle clockwise
- 2. Adjust screen to desired height range: 15" to 33" (38 cm x 84 cm). Fix desired position by turning height-adjustable lock handle clockwise.

Detaching from Side Rail

1. Unlock clamp by turning handle counterclockwise. Remove mounting post from socket.

GENERAL SPECIFICATIONS

Device Dimensions (maximum)

- Length: 23" +/- 0.5" (58 cm +/- 1 cm)
- Height: 35 3/8" +/- 0.5" (90 cm +/- 1 cm)
- Depth: 1 3/4"+/- 0.5" (4 cm +/- 1 cm)
- Device Weight: 3 +/- 0.5 lbs. (1.3 +/- .22 kg)
- Attaches to rail of surgical table anywhere along rail
- Single-person installation

COMPONENT OVERVIEW

Anesthesia Screen is a surgical table accessory that aids anesthesiologist in viewing patient during surgery.

Required product for use: 800-0134 Schure Socket XL (sold separately)

US: 0.374" x 1.122" (9.5 mm x 28.5 mm) PN# 800-0134

Denyer: 0.236" x 1.496" (6 mm x 38 mm) PN# 800-0134-DEN Europe: 0.394" x 0.984" (10 mm x 25 mm) PN# 800-0134-EU Eschmann (UK): 0.236" x 1.260" (6 mm x 32 mm) PN# 800-0134-UK Japan: 0.354" x 1.260" (9 mm x 32 mm) PN# 800-0134-JPN Swiss: 0.394" x 1.181" (10 mm x 30 mm) PN# 800-0134-SWISS

GENERAL INFORMATION

- Product not made with Natural Rubber Latex
- Product warranty covers product from manufacturing defects for period of 2 years
- If damaged in shipping, call Customer Service at (888) 724-8763 or (781) 982-7000 to obtain Return Material Authorization number. For product warranty issues, contact Customer Service.
- CE marked medical device according to MDR (EU) 2017/745
- Product is maintenance-free, check product condition before next use
- Life of device is 5 years under normal use
- Store device between -4°F to +86°F (-20°C to 30°C)

DISPOSAL

- General Prevent infection by cleaning and disinfecting product before disposal
- Packaging Dispose packaging material via household waste according to national requirements
- SchureMed accepts back used or retired products or dispose of product in accordance with national requirements

CLEANING RECOMMENDATION

Follow current Association of periOperative Registered Nurses Journal Guidelines for proper cleaning and disinfection procedures.



WARNING!

Adhere to standards for blood-borne pathogens from the Occupational Safety and Health Administration. Use recommended protective clothing, gloves, masks and eye protection to clean accessory.



PRODUCT WARNINGS

WARNING!

Do not reuse device if there are obvious signs of damage or functional issues. Consult manufacturer before reusing.

CAUTION

Strictly read/follow manufacturer's directions for cleaning fluids. DO NOT use cleaners containing phenolics.

- 1. Remove major contaminants from accessory with disposable materials. Follow appropriate bio-hazard waste disposal procedures.
- 2. Apply cleaning fluid liberally to entire accessory and wipe with clean, lint-free cloth until all moisture and cleaning fluid is removed from accessory
- 3. Let accessory dry

USER NOTICE

Any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the Member State in which the user and/or patient is established.

UDI Basic UDI-DI: 081005737F0043KX

eIFU Language Versions

To download and print the Instructions for Use, please go to http://www.schuremed.com/schuremed-eifu.

Symbol Glossary

Symbol	Title	Symbol Description
<u>l</u>	Manufacturer	Indicates the medical device manufacturer.
EC REP	Authorized Representative in the European Community	Indicates the authorized representative in the european community.
><	Use-by Date	Indicates the date after which the medical device is not to be used.
LOT	Batch Code	Indicates the manufacturer's batch code so that the batch or lot can be identified.
REF	Catalogue or Item Number	Indicates the manufacturer's catalogue or item number so that the medical device can be identified.
SN	Serial Number	Indicates the manufacturer's serial number so that a specific medical device can be identified.
\triangle	Caution	To indicate that caution is necessary when operating the device or control close to where the symbol is placed, or to indicate that the current situation needs operator awareness or operator action in order to avoid undesirable consequences.
MD	Medical Device	Indicates the item is a medical device.
[UDI]	Unique Device Identifier	Indicates a barcode as conteining unique device identifier information.
CE	CE Marking	European Conformity.
2	Single Patient Use	Indicates the item is a single patient use medical device.



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EC REP Authorized Representative

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