

US Patent No 6,843,769

INSTRUCTIONS FOR USE REF: A011, A021, A031, A041, A811, A821, A831, A841

## INTENDED USE AND DESCRIPTION

Airtrag SP is a video laryngoscope to facilitate intubation. It allows visualization of the airway during intubation. It minimizes hyper extension of the neck and permits intubating from virtually any position. ET Tube is loaded into the lateral channel of the device and when advanced it is guided towards the glottis of the patient. It is a SINGLE USE medical device provided clean and ready to use. To perform its purpose Airtrag SP must have attached to its proximal end, either

- its eyecup or
- the WiFi Camera (Ref A-390) offered by the manufacturer, attached after removing its eyecup, or
- an Endoscopic camera attached to the eyecup.

### SIZES AND SPECIFICATIONS BY PRODUCT CODE

Regular: A-011

Blue. For use with ETT 7.0 – 8.5 Minimum patient mouth opening:

Maximum insertion portion width: 30.0 mm.

Working Length: 119.5 mm.

Field of View: Vertical 32 degrees; Horizontal 27 degrees Direction of View: Vertical 12 degrees; Horizontal 22 degrees

## Small: A-021

Green. For use with ETT 6.0 - 7.5

Minimum patient mouth opening: 15 mm

Maximum insertion portion width: 28.5 mm.

Working Length: 115.1 mm.

Field of View: Vertical 32 degrees; Horizontal 27 degrees Direction of View: Vertical 12 degrees; Horizontal 22 degrees

## Paediatric: A-031

Purple. For use with ETT 4.0-5.5

Minimum patient mouth opening: 12 mm.

Maximum insertion portion width: 24.6 mm.

Working Length: 75.8 mm.

Field of View: Vertical 28 degrees; Horizontal 21 degrees Direction of View: Vertical 12 degrees; Horizontal 22 degrees

#### Infant: A-041

Grey. For use with ETT 2.5-3.5

Minimum patient mouth opening: 11 mm.

Maximum insertion portion width: 22.6 mm.

Working Length: 62.7 mm.

Field of View: Vertical 27 degrees; Horizontal 21 degrees Direction of View: Vertical 12 degrees; Horizontal 22 degrees

## **USE INSIDE OF THE MRI ENVIRONMENT**

This medical device was determined to be MR conditional according to the terminology specified by ASTM Standard Practice for Marking Medical Devices and Other Items for Safety in the Magnetic Resonance Environment.

The device can be used in the MRI environment according to the following conditions:

• Static magnetic field of 3-Tesla or less

 Spatial gradient magnetic field of 720-Gauss/cm or less IMPORTANT NOTE: The device is intended for use inside of the MRI environment (e.g., in the MR system room). It will not be utilized directly inside of the MR system (e.g., inside of the bore of

the scanner), during its operation (i.e., scanning). As such, the assessment of magnetic field interactions for the device specifically involved evaluations of translational attraction in relation to exposure to a 3-Tesla MR system, only.

## **TECHNIQUE FOR USING THE AIRTRAQ SP**

#### I. PREPARATION & TEST

- Select the appropriate size Airtrag SP based on the size ETT to be
- Press orange switch located below the battery cover to Turn On the light.
- After 30 seconds check that the light is steady. If so, anti-fog system is fully activated.

NOTE: If light automatically turns off, the unit has become defective and must not be used.

- Lubricate the ETT and place it into the lateral channel of the Airtraq SP without contacting the lens.
- Align the tip of the ETT with the end of the lateral channel.

## II. AIRTRAQ SP PLACEMENT (Fig. 1)

- Insert the Airtrag SP into the midline of the patient's mouth. Take special care to avoid pushing the tongue inside the oropharynx.
- Before it reaches the vertical plane, begin looking to identify airway structures.
- Continue insertion until the epialottis is identified. Place the tip of the Airtrag SP in the vallecula. Alternatively, the tip can be placed under the epiglottis, lifting it out of the way.
- · Gently lift up the Airtrag SP to expose the vocal cords.

## III. ETT INSERTION (Fig. 2)

- · Align the vocal cords in the center of the visual field by gently moving the tip of the Airtrag SP as needed.
- · Gently advance the ETT in the lateral channel. If needed rotate counterclockwise ET tube inside the channel (corkscrew maneouver). Check insertion depth.
- Inflate the ETT cuff as normal and check for proper positioning.

midline.



Do not insert too deep



Center glottis Advance ETT

- Lift gently
- Twist Airtraq to center vocal cords

Remove midline

Corkscrew ETT

# IV. AIRTRAQ SP REMOVAL (Fig. 3) • Separate the ETT from the Airtrag SP by pulling it laterally from the ETT, while holding the ETT in position. • Remove the Airtrag SP from the patient's airway following the



## WARNINGS AND PRECAUTIONS

- This product should only be used by personnel trained in insertion of endotracheal tubes.
- Do not sterilize.
- Do not incinerate unless batteries have been removed.
- Do not submerge in liquids.
- Use only with non-flammable anesthetics.
- Do not put pressure on the teeth with this device.
- Do not force the Airtrag SP into the upper airway.

#### ISAGE TIPS

- 1. Initial experience should be gained in non-difficult airways.
- 2. Insert the Airtraq SP, avoiding the tongue, and slide it softly and
- 3. Keep the Airtraq SP in the mouth's midline.
- 4. Look before the Airtrag SP gets to the vertical plane.
- Do not insert too deep. If structures (arytenoids, epiglottis, etc.) are not clearly recognized, withdraw the Airtraq SP slightly.
- Once the tip is located at the epiglottis, either at the vallecula (Macintosh style), or under the epiglottis (Miller style), gently lift up the Airtraq SP (do not tilt or use a lever action).
- 7. Advance the ETT slowly. If needed rotate ETT inside the channel.

## **BATTERY CHARACTERISTICS**

Each Airtraq SP is equipped with two AAA batteries in serial connection that provide a voltage of 3 volts. The batteries provide power to the LED light and to the anti-fog system when the switch is turned on. They should not be replaced.

## STORAGE, TRANSPORT, SHELF LIFE AND SERVICE LIFE

The Airtraq SP should not be used, stored or transported at temperatures below  $-5^{\circ}$ C/23°F or over  $55^{\circ}$ C/131°F. The relative humidity must not exceed 95 %. The air pressure must not exceed 500 to 1060 hPa.

Airtrag SP shelf life is limited to the expiration date.

Airtraq SP service life is limited to 40 accumulative minutes, 5 minutes before its end; the Airtraq SP will start blinking for 2 seconds every 20 seconds. Once maximum operating time is reached the Airtraq SP will blink continuously until battery is depleted.

#### **DISPOSAL**

To dispose the Airtrag SP once it has been used:

- Separate the eyecup from the main body by pulling it up.
- Remove the battery cover by pulling it away from the main body (pull away from the small notches).
- Remove the batteries from the Airtraq SP and place them in an appropriate battery recycling container (dispose of them according to established recycling policies). The batteries are classified as non hazardous waste material and comply with European Directive WEEE. However, the manufacturer recommends separating them from standard trash.
- Discard the Airtraq SP as any other potentially contaminated waste.

#### MANUFACTURER'S WARRANTY

The manufacturer warrants the Airtraq SP against faulty materials or manufacturing defects for only one use or until the expiration date, whichever comes first, provided that the Airtraq SP is used in accordance with the procedures set forth in these instructions. This Warranty is applicable only if the device is purchased from an authorized distributor.

This device has not been designed to be cleaned or sterilized. Use beyond this recommendation may generate serious consequences in the product's performance and will void the Airtraq SP's warranty. The manufacturer disclaims all other warranties, whether expressed or implied, including, without limitation, the warranties of merchantability or fitness for a particular use.

#### **USER ASSISTANCE INFORMATION**

Instructions for use are available online at https://www.airtraq.com/IFU

Visit www.airtraq.com for further advice on using this device Any serious incident that occurs in relation to this device should be reported to the manufacturer and the competent Health Authority in which the user and/or patient is established. For communication with manufacturer, please email user.assistance@airtraq.com, or:

- Contact directly with Manufacturer at its address or by phone:
  USA & Canada: +1877-624-7929
  FU & Other: +34944804690
- Contact Manufacturer's Representative for your area (details below) or
- 3. Contact your Local Distributor

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Airtraq is a registered trademark.



**Rx Only** 

























#### GLOSSARY OF SYMBOLS

GLOSSAKT OF STMBOLS					
SYMBOL	SYMBOL MEANING	SYMBOL	SYMBOL MEANING	SYMBOL	SYMBOL MEANING
REF	Reference Number	QTY	Number of Units	<b></b>	Legal Manufacturer
LOT	Batch code	<u>M</u>	Manufacturing Date		Use-By date
ECREP	Authorized representative in the European Community		Importer	MD	Medical device
I	Fragile, handle with care	www.airtraq.com/fu	Consult instructions for use	2	Do not re-use
8 10/2019 Sec. (1) 2019	Temperature limit.	% 60% 10%	Humidity limitation	0 • 0 O	Atmospheric Pressure Limitation
LATEX	Not Made with Natural Rubber Latex	<b>*</b>	Keep Dry / Protect from moisture	Rx Only	Federal (USA) law restricts the use of this device to sale by or on the order of a physician
€	CE Mark, European Conformity	EAC	EAC Mark, Eurasian Conformity	UK	UKCA Mark, UK Conformity Assessment
11	This side up	<b>S</b>	Do not use if package is damaged		Distributor
<u> </u>	Separate collection for waste of electrical and electronic equipment	<b>*</b>	Type BF applied part	Australian Sponsor	Australian sponsor
MR	Magnetic resonance conditional	0	Box/packaging recyclable		