

REF 001262U

AirLife™

Pediatric Oxygen Mask

Vinyl, Under-The-Chin Style, Medium Concentration Mask with 7 feet (2.1 m) Tubing and U/Connect-It Adapter

REF

Catalogue number

R_X ONLY

U.S. Federal law restricts this device to sales by or on the order of a physician

LOT

Lot Number

Clean, Ready to Use

Clean, Ready to Use

QTY

Quantity



Manufacturer



Not made with natural rubber latex



Use-by Date



Contains Phthalates



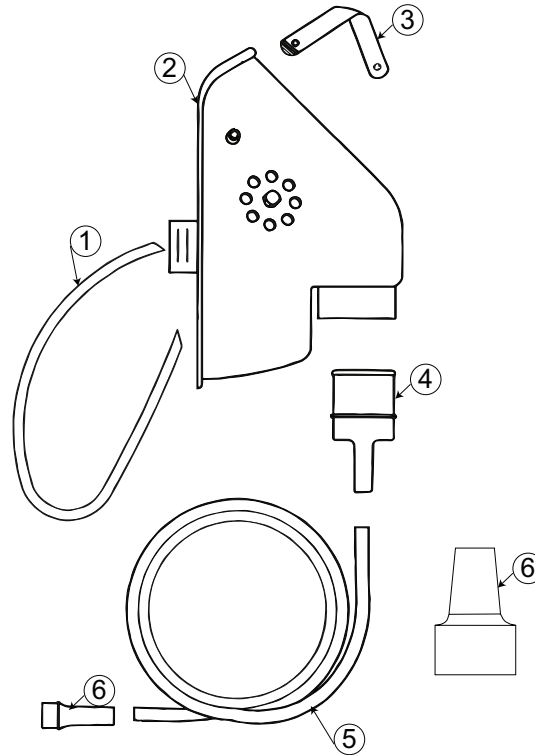
Do not re-use



Consult instructions for use



Temperature Limit



EN

English

Pediatric Oxygen Mask consists of:

- ① Elastic Strap
- ② Pediatric Oxygen Mask
- ③ Noseclip
- ④ Oxygen Connector
- ⑤ 7 feet (2.1 m) Crush Resistant Oxygen Tubing
- ⑥ U/Connect-It Adapter (Threaded Adapter)

Intended Use

Intended for oxygen to be delivered to the patient at a prescribed concentration and/or flow rate. Oxygen tubing is connected to an oxygen source on one end and a mask on the other end to deliver oxygen to the patient.

Directions for Use

1. Attach U/Connect-It Adapter of oxygen tubing directly to threaded oxygen outlet. (Separate tubing adapter may be used but it is not required.) Push and twist to seal. Adjust liter flow as prescribed by physician.

AU REP

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Made in Mexico

2. Place mask over patient's face covering both nose and mouth.
3. Slip strap over patient's head and adjust for comfort and fit.
4. Pinch nose clip to provide a seal around the nose.

Warning

- The oxygen connector in mask should not be connected directly to any endotracheal or tracheostomy tube inlet.
- Avoid kinking or occluding tubing.

Note

- Assure adequate and uninterrupted oxygen flow to patient. If oxygen flow is inadequate or interrupted, patient may not be able to inhale, or may not be able to inhale adequate volume.

Storage

Recommended normal storage temperature 0°C/+32°F to +50°C/+122°F.

Contains Phthalates (DEHP). Risks and Precautionary Measures related to Phthalates: This instruction pertains to the phthalate symbol marked on the device or its packaging. If this device is used for the treatment of children, or treatment of pregnant or nursing women; please note that the following types of procedures may increase the risk of exposure to phthalates: Exchange transfusion in neonates, total parenteral nutrition in neonates, multiple procedures in sick neonates, haemodialysis in peripuberal males, male foetus and male infant of pregnant women, and lactating women; and massive blood infusion into trauma patients. Although these procedures have the potential for increased risk of exposure, conclusive evidence of human health risks has not been established. As a precautionary measure, to reduce the potential for unnecessary exposures to phthalates, the product must be used in accordance with the instructions for use, and practitioners should refrain from using this product beyond the period of time the product is medically necessary or needed.

Dispose of all materials in accordance with local, state, and federal regulations. Decontaminate and dispose of all potentially biohazardous material.

Any serious incident that has occurred in relation to the device should be reported to the manufacturer at productquality@myAirLife.com.

AirLife®