Surgical Face Mask

EN 14683:2019 + AC:2019 TYPE IIR REF: 1094 | Anti-Fog















GIBSONMEDICAL.SE

SURGICAL FACE MASK

TYPE IIR, TIE STRAPS, ANTI-FOG



PRODUCT: Surgical Face Mask Type IIR SIZE: 17.5 x 9.5 cm MODEL: Tie Straps, Anti-Fog RANGE OF APPLICATION: Fluid resistant, disposable medical device that creates a physical barrier between the mouth and nose of the wearer and the immediate environment. Designed to prevent spreading of infections by catching pathogens shed in liquid droplets and particles from the wearer's mouth and nose. Also protects the wearer from potentially contaminated fluid splash. STRUCTURE AND COMPOSITION: Mask body made of three layer polypropylene non-woven fabric (spunbond-meltblown-spunbond) with anti-fog foam. Spunbond tie straps and aluminium nose piece. Free from latex, PVC and nickel. PRODUCT PERFORMANCE: EN 14683:2019 + AC:2019 Type IIR.

INSTRUCTIONS: Clean and disinfect hands before use. 1. Open the package and take one mask out. Avoid touching the internal side of the mask. The coloured side is the external side. 2. With the nose strip facing upwards, attach the mask to your nose and stretch the mask around your chin. 3. 4. 5. Tie the upper straps around the back of the head and lower straps around the neck. 6. Use the index finger of both hands to pinch the nose strip so it takes the shape of your nose bridge. Make sure the mask fits your face shape and that it is comfortable. NOTE: Do not touch the mask when in use. Only touch the tie straps when removing it. Discard into a waste container with a lid, or according to medical waste regulations in medical environments. Clean and disinfect hands after use.

CAUTION: Check whether the package is intact before use. Do not use it if the package is damaged. Confirm the label, production date and use-before-date. Use within the validity period. This product is disposable and should not be reused. Discard into a waste container with a lid. In a medical environment it should be discarded according to medical waste regulations.



Manufacturer - Indicates the manufacturer of the medical device



Date of manufacture - Indicates the date when the medical device was manufactured



Expiration date - Indicates the date after which the medical device should not be used



Do not expose to direct sunlight - Indicates a medical device that should not be exposed to direct sunlight



Store dry - Indicates that the medical device should be protected against moisture



Do not reuse - Indicates that the medical device is intended for single use or for an individual patient during a single procedure



Read the instruction manual - Indicates that it is necessary for the user to read the instruction manual List number - Indicates the manufacturer's list number so that the medical



batch can be identified

device can be identified Batch code - Indicates the manufacturer's batch code so that the



CE marking - Indicates that products sold in the EEA have been assessed to meet high health-, safety- and environmental protection requirements



EC | REP | Authorized Representative - Indicates the Authorized Representative of the European Community



Financial contribution towards recovery of packaging - Indicates that a financial contribution has been made towards recovery of packaging

STORAGE REQUIREMENTS: Store protected from sunlight in a dry, clean and well-ventilated environment. **RECYCLING:** Packaging is sorted as paper. Face mask as residual waste.





