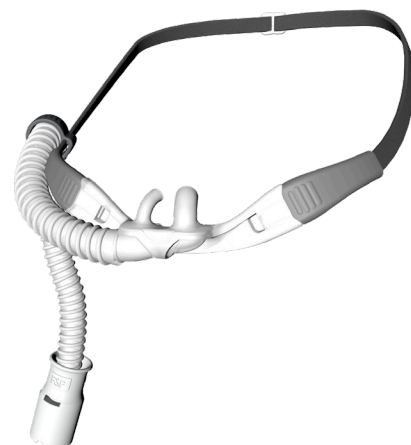


Asymmetric™ Nasal High Flow Interface

The Optiflow™+ Duet delivers asymmetric nasal high flow therapy to patients.



PRODUCT SPECIFICATIONS

| | |
|--|--|
| Product code | OPT962 (small) OPT964 (medium) OPT966 (large) |
| Quantity | Box of 20 |
| Box components | Nasal Cannula Interface Tube clip User instructions |
| Weight (Pack of 20 including box) | 1.02 kg (2.2 lb) |
| Pack dimensions | Length: 240 mm (9.4 inch) Width: 190 mm (7.5 inch) Height: 225 mm (8.9 inch) |

PERFORMANCE SPECIFICATIONS

| | |
|--------------------------------|---|
| Shelf life | 3 years |
| Flow range | With the F&P Airvo™ series: OPT962 10 – 50 L/min OPT964 10 – 60 L/min OPT966 10 – 60 L/min With the F&P 850™ system: 5 – 60 L/min With the F&P 950™ system: 5 – 70 L/min |
| Humidity settings | Up to 37 °C dew point temperature |
| Condensation management | Evaqua™ technology breathable tubing for reduction of mobile condensate |
| Usage | Single patient use |
| Duration of use | 14 days hospital use |
| Compatible humidifiers | Airvo series; MR850; F&P 950 |
| Compatible circuits | 900PT56x-series tube; or tube and chamber kit (e.g. 900PT561) RT series kits with 22 mm heated inspiratory tube and chamber (e.g. RT232) F&P 950 Adult Heated Circuit kit (e.g. 950A40) |

COMPONENTS & COMPOSITION

| | |
|------------------------------|---|
| Predominant materials | Thermoplastic elastomers, polyethylene (PE), polypropylene (PP), acetal, polyester elastomer, ABS, nylon |
| Materials not present | Not made with natural rubber latex, phthalates (DEHP, DBP, BBP), or BPA |
| Manufacturing mode | Produced in a controlled working environment |
| Disposal | Dispose of products and packaging according to local guidelines. Hospitals and long-term care facilities should discard according to their standard method for disposing of contaminated product. |

REGULATORY

| | |
|--------------------------|-------------------------------------|
| Country of origin | New Zealand or Mexico |
| Classification | AU-IIa; EU-IIa; Canada-II; USA-II |
| Notified body | TÜV SÜD Product Service GmbH CE0123 |

Please note that the information in this specifications sheet (including product information and images) is summarized and provided for illustrative purposes only. Please refer to the relevant user instructions for more information and confirm details with your local Fisher & Paykel Healthcare representative prior to placing an order. Information subject to change without notice.

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