Oxygen Tubing – Clear, 3-Channel, Ribbed Connectors

SINGLE PATIENT USE | DISPOSABLE

Device Manufacturing Specifications		
Reference Number	2001, 2002, 2004, 2014, 2025, 2030, 2035, 2040, 2050	
Manufacturer	SunMed LLC/Salter Labs	
FDA Product Code	Class I Medical Device	
Classification – EU	Class IIa, Rule 2	
CE Mark/Notified Body	CE 2797	
EU Authorized Representative	Mt Promedt Consulting GmbH	1
Classification – Canada	Class II	1
EMDN Code	R03010204 Oxygen Administration Tubing	1
GMDN Code	12875 Oxygen Administration, Tubing	1
UMDNS Code	12875 Tubing, Oxygen Connecting	1
Made In	Mexico	1
Usage	Disposable, Single Patient Use	1
Sterile	Nonsterile	1
Patient Population	Neonate, Infant, Pediatric and Adult	1
Packaging	Individually Packaged, 20/Case, 25/Case or 50/Case	
Shelf Life	5 Years	
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Description: Clear, 3-Channel safety oxygen supply tubing with ribbed connectors used as a conduit to delivery oxygen from the source gas to the patient.

Intended Purpose: To extend the length of oxygen tubing as accessory to nasal cannulas, oxygen masks and other breathing products.

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Specification	
5° C to 40° C	
-20° C to 50° C	
3/16"	
≤ 15 LPM	
3-Channel	
Ribbed Universal Push-On	
Fits 4 mm to 6 mm	
1' (0.3 m), 4' (1.2 m), 7' (2.1 m), 14' (4.2 m), 25' (7.6 m), 35' (10.7 m) 50' (15.2 m)	

Device Material				
Component	Material			
Tubing	Polyvinyl Chloride, Clear			
End Piece	Polyvinyl Chloride			
Part Number	UOM	GTIN		
2001-1	Each	00607411200007		
2001-1-50	Case	10607411200004		
2002-7	Each	00607411200014		
2002-7-50	Case	10607411200011		
2004-4	Each	00607411200038		
2004-4-50	Case	10607411200035		
2014-14	Each	00607411200106		
2014-14-50	Case	10607411200103		
2025-25	Each	00607411200168		
2025-20-25	Case	10607411200165		
2030-30	Each	00607411200175		
2035-30-20	Case	10607411200172		
2050-35	Each	00607411200182		
2050-35-20	Case	10607411200189		
2035-40	Each	00607411200199		
2035-40-20	Case	10607411200196		
2050-50	Each	00607411200236		
2050-50-20	Case	10607411200233		



Latex: SunMed[®] does not utilize latex or latex-containing material to manufacture products. To the best of our knowledge latex does not contact components or finished goods during the manufacturing process.

Phthalates: The selected materials are Non-DEHP; DEHP was not intentionally added as part of the manufacturing process.

Mercury-Lead: The selected materials do not contain lead or mercury. Biocompatible: Per device classification in ISO 10993.

SunMed manufactures product from medical grade materials in compliance with Good Manufacturing Practices (GMPs) as listed in 21 C.F.R. (U.S. Code of Federal Regulations).



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