

Cardinal Health™ surgical gloves technical data sheet

Protexis™ Latex Micro

Sterile latex powder-free surgical gloves with nitrile coating

Size	5.5	6	6.5	7	7.5	8	8.5	9
Cat. No.	2D72NT55X	2D72NT60X	2D72NT65X	2D72NT70X	2D72NT75X	2D72NT80X	2D72NT85X	2D72NT90X

Technical information

Primary material	Natural rubber latex with Nitrile coating (Nitrile coating: triple-dip manufacturing process integrates latex and Nitrile—outer layer is latex; middle layer is a blend of latex and Nitrile, inner layer is Nitrile.) Powder-free: content is ≤ 2.0 milligrams/glove, in accordance with ASTM D 1624 and ISO 21171. Caution: This product contains natural rubber latex which may cause allergic reactions, including anaphylactic responses. Safe use of these gloves by or on latex-sensitized individuals has not been established.
Donning agent	Inner Nitrile Layer. Lubricated with CPC (Cetylpyridinium chloride), Darvan L and Silicone.
Color	Light Brown
Chemical additives	ZDBC (Zinc Dibutyldithiocarbamate) is used in manufacturing and residual chemical below detectable level.
Leachable protein	<50 micrograms/gram of total extractable protein, in accordance with EN 455-3 using ASTM D 5712 (Modified Lowry Protein Method).
Thickness (per ASTM D 3577 ≥ 0,10 mm)	Finger tip: 0.17 mm Palm: ≥0.14 mm Cuff: ≥0.14 mm
Length	Sizes 5.5–6.5: 11.1 in. / 282 mm Sizes 7.0–9.0: 11.6 in. / 295 mm
Force at break, before accelerated ageing (per EN 455-2 ≥ 9 N)	17N
Force at break, after accelerated ageing (per EN 455-2 ≥ 9 N) (7 days at a temperature of 70° C in an oven)	17N
Elongation at break, before accelerated ageing (per ASTM D 3577)	≥750%
Freedom from holes (per EN 455 AQL 1.5)	0.65 AQL
Design	Exclusive glove mold with an independent thumb design allows for an anatomical fit and more natural movement in the fingers, thumb and palm.
Cuff design	Proprietary interlocking, beaded cuff design reduces roll-down.
Sterilization	Gamma radiation.

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Expiration date	35 months from date of manufacture Manufacture and expiration dates are printed on packaging (YYYY-MM format).
Packaging	50 pairs per dispenser box/4 boxes per carton.
Storage recommendations	Do not store near sources of ultraviolet light, heat or x-rays.
Country of Origin	Thailand
Regulations and quality standards	Cardinal Health manufacturing facilities are certified to EN ISO 13485 by BSI. Products conform to EU Directive 93/42/EEC and CE marked. Products meet the requirements of European harmonized standards EN 455-1, -2, -3 and -4. Tested per EN 420:2003+A1:2009, EN 388:2016, EN ISO 374-1:2016, EN 374-2:2014, EN 374-4:2013, EN ISO 374-5:2016 & EN 16523-1:2015.
CE certificate number and classification	CE 553828; Class IIa.
PPE Certification number	CE 694147
Other	Single use only.

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For healthcare professionals only.

Important information: Prior to use, refer to the instructions on the dispenser box supplied with this device for indications, contraindications, side effects, suggested procedure, warnings and precautions. CARDINAL HEALTH, the Cardinal Health LOGO, ESSENTIAL TO CARE and PROTEXIS are trademarks of Cardinal Health and may be registered in the US and/or in other countries.
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