

PI

Sterile
Polyisoprene



NOT MADE WITH
NATURAL RUBBER LATEX

A multipurpose solution

- Engineered to **protect** in a wide array of clinical cases
- A **multipurpose solution** that offers tactile response with barrier protection
- Interlocking, beaded cuff design helps to **prevent roll-down**
- Proprietary hand mold with an independent thumb design allows for an **anatomical fit and natural movement**



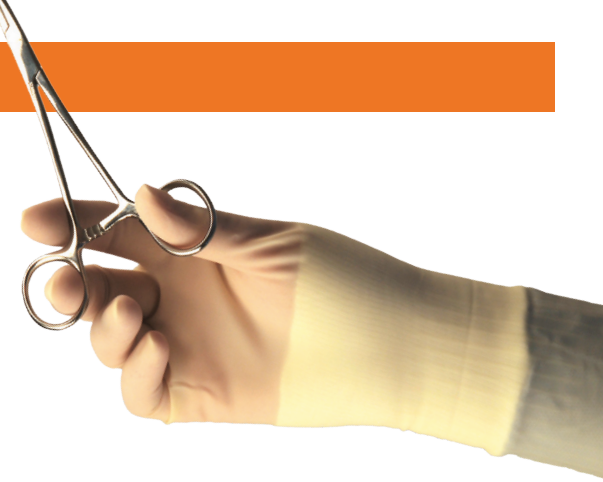
Backed by the expertise of Cardinal Health, Protexis™ PI Surgical Gloves meet all relevant FDA and ASTM standards, including also the requirements of European harmonized standards EN 455-1, -2, -3 and-4. They include those for physical dimensions, physical properties and freedom from holes. Documentation is available upon request.

Properties (before aging)

Force at break	19N ¹
Elongation at break (elasticity)	≥ 650% ²
Puncture resistance (cuff)	Min 5N ³
Freedom from holes	0.65 AQL ⁴

Chemotherapy agent permeation⁶

	Agent	Minimum breakthrough detection time in minutes (0.01 µg/cm ² /minute)
1	Carmustine (3.3 mg/mL)	15.26 (Do not use)
2	Cisplatin (1.0 mg/mL)	> 240
3	Cyclophosphamide (20 mg/mL)	> 240
4	Doxorubicin HCL (2.0 mg/mL)	> 240
5	Etoposide (20 mg/mL)	> 240
6	Fluorouracil (50 mg/mL)	> 240
7	Ifosfamide (50 mg/mL)	> 240
8	Methotrexate (25 mg/mL)	> 240
9	Mitomycin C (0.5 mg/mL)	> 240
10	Mitoxantrone (2.0 mg/mL)	> 240
11	Paclitaxel (6.0 mg/mL)	> 240
12	Thiotepa (10 mg/mL)	16.04 (Do not use)
13	Vincristine Sulfate (1.0 mg/mL)	> 240



When chemotherapy drugs are present, glove selection should be based on the specific type(s) of chemicals used. Users must refer to the instructions for use supplied with the box, for the chemicals being used to determine an adequate level of protection.

1. Force at Break tested in accordance with EN 455-2
2. Elongation at Break tested in accordance with ASTM D 3577
3. Puncture resistance (cuff) tested in accordance with AS/NZS 4179
4. Freedom from holes tested in accordance with EN 455-1
5. Thickness tested in accordance with ASTM D 3577
6. Chemotherapy agent permeation tested in accordance with ASTM D 6978-05
7. 35% reduction of materials used as compared to previous Cardinal Health packaging design. Cardinal Health Technical Council Conference 2011, presentation on file

Catalog no.	Size	Length	Thickness ⁵			Material	Color	Cuff type	Qty/ bx	Qty/ cs
			Finger	Palm	Cuff					
2D72PT55X	5.5	287 mm	0.23 mm	0.17 mm	0.17 mm	Synthetic polyisoprene (PI)	Cream	Beaded/ Rolled	50	200
2D72PT60X	6									
2D72PT65X	6.5									
2D72PT70X	7	300 mm								
2D72PT75X	7.5									
2D72PT80X	8									
2D72PT85X	8.5									
2D72PT90X	9									

THE CARDINAL HEALTH™ PROTEXIS™ SURGICAL GLOVES PROMISE:

We protect so you can perform.

Cardinal Health is dedicated to providing protection, performance and expertise so wearers can perform confidently and focus on their patients.



Help maximize storage space: Half-fold packaging design reduces packaging material⁷.



Storage recommendations: Do not store near sources of ultraviolet light, heat or x-rays. Store in a cool, dry place, store away from direct sunlight.



Expiration: 35 months from date of manufacture. Expiration date is printed on packaging.

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. As part of its continuous product development policy, Cardinal Health reserves the right to change product specifications without prior notification. Please contact your Cardinal Health representative for additional product availability information.



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