

DISPOS-A-GLOVE® Sterile Pairs

Essential Series

Packed as a pair, these synthetic medical exam gloves are ideal for brief, sterile applications

- **Sterile:** Packaged as a pair of sterile gloves per pouch, these powder-free gloves are suitable for sterile medical exam (non-surgical) procedures
- **Reduced allergy risk:** This synthetic glove is made from copolymer materials (ethylene vinyl acetate film), making it suitable for use by individuals with allergies to natural rubber latex (Type I allergies)
- **Heightened sensitivity:** These gloves also offer good tactile sensitivity to the wearer
- **Double-gloving friendly:** DISPOS-A-GLOVE® can suitably be used as a glove liner for increased protection
- **Cost-effective:** This synthetic alternative for sterile use is offered at a competitive price point
- **Easy donning:** These gloves are packaged on backing paper, making them easy to don

Recommended For

- Sterile procedures outside of the O.R.
- Brief, low-risk examination procedures
- Protection from Type I latex allergy in HCW's or patients



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PRODUCT INFORMATION

Material	EVA (Ethylene Vinyl Acetate)	Not Made From Natural Rubber Latex	Yes
Color	Clear	Cuff Length	Standard
Powder Content	Powder-Free	External Glove Surface	Smooth
Freedom from Holes (Inspection level I)	1.5 AQL	Palm Thickness (mm/mil)	0.03 / 1.2
Finger Thickness (mm/mil)	0.03 / 1.2	Allergy Prevention	Latex (Type I)
Available Sizes	S (6.5 - 7), M (7.5 - 8), L (8.5 - 9)	Tested For Use With Chemotherapy Drugs	No
Sterile	Yes	Antistatic	Not Tested
Glove Length (mm/inches)	278 / 10.9	Product Segmentation	Essential Series

PRODUCT INFORMATION

Size	S (6.5 - 7)	M (7.5 - 8)	L (8.5 - 9)
Product Code	MDG651	MDG751	MDG851

PACKAGING AND STORAGE

Packaging	1 pair/paper pouch 50 pouches per box 14 boxes per carton/case
Shelf Life	3 Years
Storage Instructions	Keep out of direct sunlight; store in a cool and dry place. Keep away from sources of ozone or ignition.

STANDARDS AND CERTIFICATIONS

CE Number, EN 455 Part 1, EN 455 Part 3, ISO 13485, ISO 10993-10:2002, Medical Device Regulation (EU) 2017/745 Class I



Contact your Ansell representative for ordering or more information.

North America
 US Tel: 800 952 9916
 CA Tel: 1-844-494-7854
 Email: insidesalesus@ansell.com

Central & South America
 Tel: +52(442) 296 20 50
 Email: cslac@ansell.com

Latin America & Caribbean Region
 BR +55 (11) 3356-3100
 latam.medical@ansell.com
 MX: +52 442 296 2050
 latam.medical@ansell.com
 CO: +57 1 288 3247
 latam.medical@ansell.com

Europe, Middle East & Africa/Asia Pacific
 Tel: +32 (0) 2 528 74 00
 Email: info@ansell.eu

MY +603 8310 6688
 apac.medical@ansell.com
 CN: +86 21 38275000
 infochina@ansell.com
 JP: +813 5549 8151
 info.medical.jp@ansell.com

Australia & New Zealand
 Tel: +61 3 9270 7270
 Email: protection@ap.ansell.com