



Product Main Number 1000006807

ABENA *Excellent*

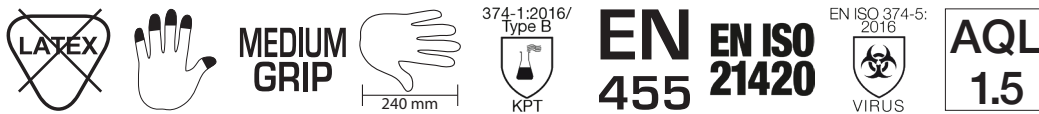
Examination gloves, accelerator-free, ABENA Excellent, XL, dark blue, nitrile, powder-free, accelerator-free

- ✓ Elastic
- ✓ Fits to hand
- ✓ No latex proteins
- ✓ Accelerator-free



Product Description

Elastic examination glove, no rubber chemicals/accelerators. Can be used for a variety of tasks such as cleaning, care, food handling, laboratory, medication handling, cytostatics and hormone creams. Contains neither latex proteins nor accelerators. Safe for use in the event of latex allergy and contact allergy to rubber chemicals.



Specifications

Base name	Examination gloves, accelerator-free
Brand	ABENA
Sub-Brand	Excellent
Size	XL
Secondary color	Dark blue
Properties	Accelerator free, finger textured, latex free, medium grip, rolled cuff
Features	Powder-free, accelerator-free
Single or multiple use	Single use
Material	Nitrile
Ingredients / Composition	Nitrile NBR, Zinc oxide, Potassium Hydroxide, Titanium Dioxide, Blue pigment
Ingredient list	Nitrile NBR, Zinc oxide, Potassium Hydroxide, Titanium Dioxide, Blue pigment
Length/depth	240 mm
Width	115 mm
Thickness	Min. 0,05 mm
Certificates	CE. Food contact materials. CAT III. MD.
CE Category (Personal Protective Equipment)	CAT III
CE Class (Medical Devices)	Class I
Product or test standards	EN ISO 374-1:2016 Type B KPT, EN 455, EN ISO 21420:2020, EN ISO 374-2:2019, EN ISO 374-4:2019, EN ISO 374-5:2016 Virus, AQL 1.5
Directives, regulations and acts	(EC) No 1935/2004, (EC) No 2023/2006, (EU) 2016/425, BEK nr 681 af 25/05/2020, MDR (EU) 2017/745
Shelf Life	3 years
Storage Instructions	Store dry, clean and at room temperature.
Product Disposal Instructions	Can be disposed of with normal household waste sorted according to local regulations.
Packaging Disposal Instructions	Can be disposed of with normal household waste sorted according to local regulations.

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Instructions for use/application

Examine the gloves for flaws and defects before use.

Packaging data

Unit	Contains	Length	Width	Height	EAN
cll	2000 pcs	37 cm	25 cm	25 cm	5703538056765
pck	200 pcs	24 cm	12.4 cm	7.5 cm	5703538056758
pair	2 pcs				
pcs	1 pcs				



Regulation (EU) 2017/745 is a regulation of the European Union on the clinical investigation and sale of medical devices for human use. It repeals Directive 93/42/EEC, which concerns medical devices, and Directive 90/385/EEC, which concerns active implantable medical devices, on 26 May 2021.



The CE mark guarantees that a product is safe to use and complies with all safety precautions. CE stands for Conformité Européenne (European Conformity) and is mainly found on electronic equipment, safety equipment, construction products and medical equipment.



Third-party type approval is required for all Category III personal protective equipment (PPE) products. The showing of the CE logo for chemical protective gloves requires that tests are carried out in accordance with test standards specified in EN ISO 374-1: 2016 + A1: 2018 - such as EN 16523-1: 2015 + A1: 2018 to determine the resistance to the permeation of chemicals. The results of this test determine the relevant pictogram symbols that can be used on the packaging and labeling.



The glass fork symbol guarantees that products have been tested in accordance with European legislation and approved for food contact. The symbol is mandatory on products used for food contact.



The product does not contain latex.



The glove has a finger textured surface.



The glove has a medium grip.



The glove is 240 mm long.



The standard specifies the requirements for protective gloves against chemicals and microorganisms. Type B has been tested for a permeation time of at least 30 minutes with the chemicals sodium hydroxide 40%, hydrogen peroxide 30% and formaldehyde 37%.



EN 455 consists of four standards that gloves must be tested against in order to be a medical glove. This standard specifies the requirements for disposable medical gloves.



This standard specifies the general requirements and relevant test procedures for glove design and construction, innocuousness, comfort and efficiency, as well as the marking and information supplied by the manufacturer applicable to all protective gloves.



The standard specifies the requirements for protective gloves against hazardous chemicals and microorganisms. This part of the standard describes the glove's resistance to bacteria, fungi and viruses.

