

Technical product specification

Product name	semperguard latex powdered	Version / Index no:
Spec code	LXF-/LXS-050NA-N-3CZ	semperguard latex powdered_Version
Date of issue	28/02/2019	G_February 2019_EN

General information

single use examination and disposable protective glove, non sterile Type

Labelling

information printed on dispenser box

Shape

ambidextrous - straight fingers

Material Colour

Natural Rubber Latex (NRL) natural white

Inside Outside pre-powdered no treatment

Cuff / surface

rolled cuff / microrough

Shelf life

5 years

Available sizes

XS (5-6) S (6-7) M (7-8) L (8-9) XL (9-10)

Dimensions, physical properties and biocompatibility

Glove length

median ≥ 240 mm (according to EN 455-2)

Minimum wall thickness

at finger

0.20 mm (double measured) / 0.10 mm (single measured) 0.16 mm (double measured) / 0.08 mm (single measured)

at palm at cuff

0.12 mm (double measured) / 0.06 mm (single measured)

Glove width

according to EN 455-2: median XS ≤ 80 mm, S 80 ± 10 mm, M 95 ± 10 mm, L 110 ± 10 mm, XL

≥ 110 mm

Force at Break

Tensile Strength

median ≥ 6 N (during shelf life according to EN 455-2) min. 14 MPa after aging (according to ASTM D3578) min. 500% after aging (according to ASTM D3578)

Elongation at Break

Residual powder / ≤ 15 mg/dm²

Powder content

Performance requirements and inspection levels

Freedom from holes (Barrier)

AQL ≤ 1.5

(as per EN 455-1, sampling in accordance with ISO 2859-1, G-1)

Dimensions and physical properties

(as per ASTM D3578, sampling in accordance with ISO 2859-1, S-2)

Standards, guidelines & quality certificates

Quality certification

regulations

ISO 9001, ISO 13485, ISO 14001

Conformity to directives and

- Medical Device Directive 93/42/EEC: Class I - PPE Regulation (EU) 2016/425: Category III

- Food Contact Materials Regulation (EC) 1935/2004

Conformity to standards

EN 420, EN ISO 374-1, EN 374-2, EN 16523-1, EN 374-4, EN ISO 374-5, EN 455 1-4, ASTM D3578 (except stress at 500% elongation), ASTM F1671



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Instructions and additional statements

Storage instruction

Store in original packaging in a dry and dark place at 10 °C to 30 °C. Refer to guidelines of storage of rubber products as described in ISO 2230:2002. Ensure that storage area is kept cool, dry and dust free, avoid ventilation and storage close to photocopy equipment. Copper-ions discolour the glove. Protect gloves against ultraviolet light sources, as sunlight and oxidizing agents. Storage above 30 °C will lead to accelerated aging and should be avoided.

Cautionary statement and ingredient information

This product contains natural rubber latex which may cause allergic reactions, including anaphylactic responses.

This product contains accelerators (Dithiocarbamate type, Zinc-mercaptobenzothiazol) not to be used in a hypersensitivity of these substances.

For further information, a list of substances contained in the glove is available upon request.

Reporting system

Medical device vigilance and reporting system

According to the official reporting criteria of the Medical Device directive, incidents caused by examination gloves must be reported immediately to our Medical Device reporting officer. E. Meil:

our Medical Device reporting officer. E-Mail:

sempermed.complaints@semperitgroup.com or Tel.: +43 2630 310 0

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Remark

Replaces all previous versions.

All standards references refer to the date of document issue.