

## Technical product specification

Product name	semperclean MC	Version / Index no:
Spec code	LFM-110NA-N-6CZ	semperclean MC_Version D_February
Date of issue	28/02/2019	2019_EN

### General information

Type	single use examination and disposable protective glove, non sterile
Labelling	information printed on packaging
Shape	anatomical
Material	Natural Rubber Latex (NRL)
Colour	natural white
Inside	polymer coated / powder free
Outside	chlorinated
Cuff / surface	rolled cuff / microrough
Shelf life	3 years
Available sizes	6 , 6.5, 7, 7.5, 8, 8.5, 9

### Dimensions, physical properties and biocompatibility

Glove length	sizes 6 - 6.5: median 270 mm, sizes 7 - 7.5: median 280 mm, sizes 8 - 8.5: median 285 mm
Minimum wall thickness	<i>at finger</i> max. 0.54 mm (double measured) <i>at palm</i> 0.42 ± 0.05 mm (double measured) <i>at cuff</i> min. 0.32 mm (double measured)
Glove width	median size 6: 77 ± 5 mm, size 6.5: 83 ± 5 mm, size 7: 89 ± 5 mm, size 7.5: 95 ± 5 mm, size 8: 102 ± 6 mm, size 8.5: 108 ± 6 mm, size 9: 114 ± 6 mm,
Force at Break	median ≥ 6 N (during shelf life according to EN 455-2)
Tensile Strength	min. 14 MPa after aging (according to ASTM D3578)
Elongation at Break	min. 500% after aging (according to ASTM D3578)
Residual powder / Powder content	≤ 2 mg (according to EN 455-3)

### 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	AQL 4.0 (as per ASTM D3578, sampling in accordance with ISO 2859-1, S-2 )

### Standards, guidelines & quality certificates

Quality certification	ISO 9001
Conformity to directives and regulations	<ul style="list-style-type: none"> <li>- Medical Device Directive 93/42/EEC: Class I</li> <li>- PPE Regulation (EU) 2016/425: Category I or III</li> <li>- Food Contact Materials Regulation (EC) 1935/2004</li> </ul>
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 F1671

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### Instructions and additional statements

<b>Storage instruction</b>	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.
<b>Cautionary statement and ingredient information</b>	<p>This product contains natural rubber latex which may cause allergic reactions, including anaphylactic responses.</p> <p>This product contains accelerators (Dithiocarbamate type, Zinc-mercaptobenzothiazol) not to be used in a hypersensitivity of these substances.</p> <p>For further information, a list of substances contained in the glove is available upon request.</p>

### Reporting system

<b>Medical device vigilance and reporting system</b>	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-Mail: <a href="mailto:sempermed.complaints@semperitgroup.com">sempermed.complaints@semperitgroup.com</a> or Tel.: +43 2630 310 0
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<b>Remark</b>	Replaces all previous versions. All standards references refer to the date of document issue.
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