Olerup SSP[®] DNA Size Marker for short gel runs

Product number:	103.203-100
Lot number:	69M
Volume:	2 x 500 μl
Concentration:	20 ng/µl
Format:	Ready-to-load, 10 µl per gel lane
Expiry date:	2013-August-01
Storage:	2-8°C

DESCRIPTION 2 vials, 500 μl each. DNA ladder, 4 bands: 50, **200**, 500 and 1 000 bp. Double intensity of 200 bp band. 20 ng/μl in 10 mM Tris-HCl (pH 9.0), 50 mM KCl, 5% (v/v) glycerol (99.5%), cresol red 100 ng/μl.

The DNA ladder contains 4 fragments, double intensity of 200 bp band:

Number of base pairs
1000
500
200
50

- PRODUCT USE The DNA Size marker can be resolved well in standard LE agarose gels of 1-2%, in NuSieve 3:1 or Metaphor agarose gels up to 4%. The DNA Size Marker for short gel runs is supplied in a *ready-to-load format*. Typically load 10 μl per gel lane of the ready-to-load DNA Size Marker for short gel runs, equals 200 ng of DNA Size Marker for short gel runs.
- **STORAGE** Store at 2-8°C.
- **SHIPPING** The DNA Size Marker for short gel runs is shipped at ambient temperature.
- **STABILITY** The DNA Size Marker for short gel runs is stable for 24 months when stored at 2-8°C.

CERTIFICATE OF ANALYSIS

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10 μI DNA Size Marker was size-separated on a 2% agarose gel stained by ethidium bromide.

Results: Four fragments of 50, 200, 500 and 1000 bp were visible on a UV transilluminator. Double intensity of 200 bp fragment.

Date of approval: 2011-August-19

Approved by:

Quality Control, Supervisor

Declaration of Conformity

Product name: Product number: Lot number:	DNA Size Marker for short gel runs 103.203-100 69M
Intended use:	DNA Size Marker for short gel runs
Manufacturer:	<i>Olerup</i> SSP AB Franzengatan 5 SE-112 51 Stockholm, Sweden <i>Phone:</i> +46-8-717 88 27 <i>Fax:</i> +46-8-717 88 18

We, *Olerup* SSP AB, hereby declare that this product, to which this Declaration of Conformity relates is in conformity with the following Standard(s) and other normative document(s) ISO 9001:2008 and ISO 13485:2003, following the provisions of the 98/79/EC Directive on *in vitro* diagnostic medical devices, Annex III, as transposed into the national laws of the Member States of the European Union.

The Technical Documentation File is maintained at *Olerup* SSP AB, Franzengatan 5, SE-112 51 Stockholm, Sweden.

The Authorized Representative located within the Community is: *Olerup* SSP AB.

Stockholm, Sweden 2011-August-19

Ann-Cathrin Jareman Head of QA and Regulatory Affairs

ADDRESSES:

Manufacturer:

Olerup SSP AB, Franzengatan 5, SE-112 51 Stockholm, Sweden. Tel: +46-8-717 88 27 Fax: +46-8-717 88 18 E-mail: info-ssp@olerup.com Web page: http://www.olerup-ssp.com

Distributed by: Olerup GmbH, Löwengasse 47 / 6, AT-1030 Vienna, Austria. *Tel:* +43-1-710 15 00 *Fax:* +43-1-710 15 00 10 *E-mail:* support-at@olerup.com *Web page:* http://www.olerup.com

Olerup Inc., 901 S. Bolmar St., Suite R, West Chester, PA 19382 *Tel:* 1-877-OLERUP1 *Fax:* 610-344-7989 *E-mail:* info.us@olerup.com *Web page:* http://www.olerup.com

For information on Olerup SSP distributors worldwide, contact Olerup GmbH.