# Olerup SSP® DNA Size Marker for short gel runs

Product number: 103.203-100/500

Lot number: 29R

Volume: Product No. 103.203-100 – 2 x 500 μl

Product No. 103.203-500 – 10 x 500 μl

Concentration: 20 ng/µl

Format: Ready-to-load, 10 µl per gel lane

Expiry date: 2014-August-01

Storage: 2-8°C

**DESCRIPTION** 103.203-100: 2 vials, 500 μl each.

103.203-500: 10 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:

<u>Fragment</u>	Number of base pairs
1	1000
2	500
3	200
4	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  $\mu$ l 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**

Olerup SSP® DNA Size Marker for short gel runs

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10  $\mu$ l 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: 2012-August-27

Approved by:

**Production Quality Control** 

## **Declaration of Conformity**

Product name: DNA Size Marker for short gel runs

Product number: 103.203-100/500

29R Lot number:

Intended use: DNA Size Marker for short gel runs

Manufacturer: Olerup SSP AB

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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.

Stockholm, Sweden 2012-August-27

Ann-Cathrin Jareman Head of QA and Regulatory Affairs

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