

DR low resolution screening Certificates 101.103.48/12 – including *Taq* polymerase, IFU-01 101.103.48/12u – without *Taq* polymerase, IFU-02

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Visit <u>www.labproducts.caredx.com</u> for "Instructions for Use" (IFU)

Lot No.: **3L6** 

**Lot-specific Information** 

## CERTIFICATE OF ANALYSIS

Olerup SSP® DR low resolution screening

Product number: 101.103-48/12 – including *Tag* pol.

101.103-48u/12u - without *Taq* pol.

Lot number: 3L6

Expiry date: 2024-07-01

Number of tests: 48 tests – Product No. 101.103-48/48u

12 tests - Product No. 101.103-12/12u

Number of wells per test: 23 + 1

Well specifications:

Well No.	<b>Production No.</b>	Well No.	Production No.	Well No.	<b>Production No.</b>
1	2020-179-01	9	2018-942-09	17	2018-942-17
2	2020-202-02	10	2020-179-10	18	2020-179-18
3	2020-179-03	11	2020-179-11	19	2020-179-19
4	2020-179-04	12	2020-179-12	20	2020-179-20
5	2020-179-05	13	2020-179-13	21	2020-179-29
6	2020-218-06	14	2020-179-14	22	2020-179-30
7	2020-179-07	15	2020-179-15	23	2020-179-31
8	2020-179-08	16	2020-179-16		

The negative control primer pairs, **Production No. 2020-205-01**, can detect contamination with PCR products diluted 10<sup>-7</sup>.

Results of Quality Control: No false positive or false negative amplifications

obtained.

Date of approval: 2020 -08 -26

Approved by:

**Production Quality Control** 

**C€** 0197 For to Vii

For In Vitro Diagnostic Use MA100 v04 CoA\_DoC IVD Annex II List B Date: August 2020, Rev. No: 00



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## **Declaration of Conformity**

Product name:

Olerup SSP® DR low resolution screening

Product number:

101.103-48/48u, -12/12u

Lot number:

3L6

Intended use:

DRB1 low resolution histocompatibility testing

Manufacturer:

CareDx AB

Franzengatan 5

SE-112 51 Stockholm, Sweden **Phone:** +46-8-508 939 00 **Fax:** +46-8-717 88 18

We, CareDx 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) EN ISO 13485:2016, following the provisions of the 98/79/EC Directive on *in vitro* diagnostic medical devices, Annex II List B, conformity assessed using Annex IV, as transposed into the national laws of the Member States of the European Union.

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

The Authorized Representative located within the Community is: CareDx AB.

Notified Body: TÜV Rheinland LGA products, Tillystrasse 2, D-90431 Nürnberg, Germany. (Notified Body number: 0197.)

Stockholm, Sweden

Date:

7020-08-27

Quality Assurance

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