

101.624-12 – including *Taq* polymerase, IFU-01
101.624-12u – without *Taq* polymerase, IFU-02Visit <https://labproducts.caredx.com> for
“Instructions for Use” (IFU)Lot No.: **6L7**

Lot-specific Information

CERTIFICATE OF ANALYSIS**Olerup SSP® HLA-C*12**

Product number: 101.624-12 – including *Taq* polymerase
101.624-12u – without *Taq* polymerase

Lot number: 6L7

Expiry date: 2024-10-01

Number of tests: 12

Number of wells per test: 47+1

Well specifications:

Well No.	Production No.	Well No.	Production No.	Well No.	Production No.
1	2020-146-01	17	2020-146-17	33	2020-146-33
2	2020-146-02	18	2020-146-18	34	2020-146-34
3	2020-146-03	19	2020-146-19	35	2020-146-35
4	2020-146-04	20	2020-146-20	36	2020-146-36
5	2020-146-05	21	2020-146-21	37	2020-146-37
6	2020-146-06	22	2020-146-22	38	2020-146-38
7	2020-146-07	23	2020-146-23	39	2020-146-39
8	2020-146-08	24	2020-146-24	40	2020-146-40
9	2020-146-09	25	2020-146-25	41	2020-146-41
10	2020-146-10	26	2020-146-26	42	2020-146-42
11	2020-146-11	27	2020-146-27	43	2020-146-43
12	2020-146-12	28	2020-146-28	44	2020-146-44
13	2020-146-13	29	2020-146-29	45	2020-146-45
14	2020-146-14	30	2020-146-30	46	2020-146-46
15	2020-146-15	31	2020-146-31	47	2020-146-47
16	2020-146-16	32	2020-146-32		

The negative control primer pairs, **Production No. 2020-205-01**, can detect contamination with PCR products diluted 10^{-7} .

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

Date of approval: 2020-11-19

Approved by:



Production Quality Control



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Lot No.: **6L7**

Lot-specific Information

Declaration of Conformity

Product name: *Olerup* SSP® HLA-C*12
Product number: 101.624-12/12u
Lot number: 6L7

Intended use: HLA-C*12 high resolution histocompatibility testing

Manufacturer: *CareDx* AB
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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 III, as transposed into the national laws of the Member States of the European Union.

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

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

Stockholm, Sweden

Date:

2020-11-26

Quality Assurance

