Lot-specific Information

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# *Olerup* SSP<sup>®</sup> HLA - Negative Control SSP

Product number:	102.102-01 – including <i>Taq</i> polymerase
Lot number:	05G
Expiry date:	2011-June-01
Number of tests:	96
Number of wells per test:	1
Storage - pre-aliquoted primers:	dark at -20°C
- PCR Master Mix:	-20°C
- Adhesive PCR seals	RT
- Product Insert	RT

### This Product Description is only valid for Lot No. 05G.

#### **GENERAL DESCRIPTION**

The Olerup SSP<sup>®</sup> HLA– Negative Control is intended to be used as a negative control in Olerup SSP<sup>®</sup> typings.

The primer set contains Negative Control primer pairs, that will amplify more than 95% of the *Olerup* SSP<sup>®</sup> HLA Class I, DRB, DQB1 and DPB1 amplicons as well as all the amplicons generated by the control primer pairs matching the human growth hormone gene.

The *Olerup* SSP<sup>®</sup> HLA – Negative Control has the sensitivity to detect approximately 50 copies of DNA template.

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## **PRODUCT DESCRIPTION**

## HLA - Negative Control SSP

#### CONTENT

The primer set contains Negative Control primer pairs, that will amplify more than 95% of the *Olerup* SSP<sup>®</sup> HLA Class I, DRB, DQB1 and DPB1 amplicons as well as all the amplicons generated by the control primer pairs matching the human growth hormone gene.

PCR product sizes range from 75 to 430 base pairs. The PCR product generated by the control primer pair is 430 base pairs.

Length of PCR	105	200	105	80	75	80
product						
5'-primer <sup>1</sup>	164	340	440	45	45	43
	<sup>5'</sup> -CAC <sup>3'</sup>	<sup>5'</sup> -Agg <sup>3'</sup>	<sup>5'</sup> -TTA <sup>3'</sup>	<sup>5'</sup> -Tg g <sup>3'</sup>	<sup>5'</sup> -Tg g <sup>3'</sup>	<sup>5'</sup> -Tg g <sup>3'</sup>
3'-primer <sup>2</sup>	231	2 <sup>nd</sup> I	507	59	58	57
	<sup>5</sup> '-TgC <sup>3'</sup>	<sup>5'</sup> -AAA <sup>3'</sup>	<sup>5'</sup> -TTg <sup>3'</sup>	<sup>5</sup> '-CTC <sup>3'</sup>	<sup>5'</sup> -ggC <sup>3'</sup>	<sup>5</sup> '-CTC <sup>3'</sup>
A*	+	+	+			
B*	+	+	+			
Cw*	+	+	+			
DRB1				+	+	
DRB3				+	+	
DRB5				+		
DRBJ						
DQB1					+	

<sup>1</sup>The nucleotide position for HLA class I genes and the codon for HLA class II genes, in the 2<sup>nd</sup> or 3<sup>rd</sup> exon, matching the specificity-determining 3'-end of the primer is given. Nucleotide and codon numbering as on the <u>www.ebi.ac.uk/imgt/hla</u> web site. The sequence of the 3 terminal nucleotides of the primer is given.

<sup>2</sup>The nucleotide position for HLA class I genes and the codon for HLA class II genes, in the 2<sup>nd</sup> or 3<sup>rd</sup> exon or the 2<sup>nd</sup> intron, matching the specificity-determining 3'-end of the primer is given in the anti-sense direction. Nucleotide and codon numbering as on the <u>www.ebi.ac.uk/imgt/hla</u> web site. The sequence of the 3 terminal nucleotides of the primer is given.

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The primer solution is pre-aliquoted into 0.2 ml PCR wells. Each well contains the same dried primer solution.

**PCR Master Mix complete with Taq,** Taq polymerase, nucleotides, buffer, glycerol and cresol red, as well as PCR lids are included in the kit including Taq polymerase.

**PCR Master Mix without Taq,** nucleotides, buffer, glycerol and cresol red, as well as PCR lids are included in the kit without *Taq* polymerase.

1 PCR reaction with a reaction volume of 10  $\mu$ l is performed per test.

### PLATE LAYOUT

Each test consists of 1 PCR reaction. Each well of the 8 well PCR plates contains the same primer mix.

	1	1	1	1	1	1	1	1
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The 8 well cut PCR plate is marked with 'NC' in silver/gray ink.

Well No. 1 is marked with the Lot No. '05G'.

The PCR plates are covered with a PCR-compatible foil.

**Please note:** When removing each PCR well, make sure that the remaining plates/wells stay covered. Use a scalpel or a similar instrument to carefully cut the foil between the plates/wells.

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## PROTOCOL

#### PCR AMPLIFICATION

For users of Olerup SSP<sup>®</sup> kits including Taq polymerase

Cut off one well from the 8 well PCR plate.

Add 2  $\mu$ l dH<sub>2</sub>O to the negative control well.

Add 8  $\mu$ l of the PCR Master Mix complete with *Taq*-H<sub>2</sub>O mixture to the negative control well, i.e. before the sample DNA is added to the PCR Master Mix complete with *Taq*-H<sub>2</sub>O mixture.

Add the sample DNA to the PCR Master Mix complete with Taq -H<sub>2</sub>O mixture, mix well and dispense 10 µl of the DNA-PCR Master Mix complete with Taq-H<sub>2</sub>O mixture into each of the wells of the SSP typing, but not into the negative control well.

The same PCR Master Mix Complete with Taq and the same dH<sub>2</sub>O that is used for the typings should be used in the negative control well. (The PCR Master Mix complete with Taq supplied with the Negative Control kit is intended to replace the PCR Master Mix used from the typing kits including Taq polymerase.)

#### For users of Olerup SSP<sup>®</sup> kits without Taq polymerase

Cut off one well from the 8 well PCR plate.

Add 2  $\mu$ l dH<sub>2</sub>O to the negative control well.

Add 8  $\mu$ l of the PCR Master Mix-*Taq*-H<sub>2</sub>O mixture to the negative control well, i.e. before the sample DNA is added to the PCR Master Mix-*Taq*-H<sub>2</sub>O mixture.

Add the sample DNA to the PCR Master Mix-*Taq*-H<sub>2</sub>O mixture, mix well and dispense 10  $\mu$ l of the DNA-PCR Master Mix-*Taq*-H<sub>2</sub>O mixture into each of the wells of the SSP typing, but not into the negative control well.

The same PCR Master Mix without Taq, Taq polymerase and dH<sub>2</sub>O that is used for the typings should be used in the negative control well. (The PCR Master Mix without Taq supplied with the Negative Control kit is intended to replace the PCR Master Mix used from the typing kits without Taq polymerase.)

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Use a 96 well thermal cycler with a heated lid. The temperature gradient across the heating block should be  $< 1^{\circ}$ C.

PCR cycling param	eters: 94°C	2 min	denaturation
2. 10 cycles	94°C	10 sec.	denaturation
	65°C	60 sec.	annealing and extension
3. 20 cycles	94°C	10 sec.	denaturation
	61°C	50 sec.	annealing
	72°C	30 sec.	extension

The same PCR cycling parameters are used for all the Olerup SSP kits.

#### AGAROSE GEL ELECTROPHORESIS

Prepare a 2% (w/v) agarose gel in 0.5 x TBE buffer. Dissolve the agarose by boiling in a microwave oven. Let the gel solution cool to  $60^{\circ}$ C. Stain the gel prior to casting with ethidium bromide (10 mg/ml), 5 µl per 100 ml gel solution. For maximal ease of handling use our ethidium bromide dropper bottles (Product No. 103.301-10), 1 drop of ethidium bromide solution per 50-75 ml of gel. <u>Note:</u> Ethidium bromide is a powerful carcinogen.

Load the PCR products, preferably using an 8-channel pipette. Load a DNA size marker (100 base pair ladder, Product No. 103.201-100) in one well per row.

Run the gel in 0.5 x TBE buffer, without re-circulation of the buffer, for 15-20 minutes at 8-10 V/cm.

#### **DOCUMENTATION AND INTERPRETATION**

Put the gel on a UV transilluminator and document by photography. Record the presence and absence of PCR products.

In the negative control well no PCR product should be seen. The presence of PCR product(s) indicates contamination<sup>1</sup>.

If contamination is detected, wipe test and testing of all reagents should be performed in order to detect the source of contamination

<sup>1</sup>Primer oligomer artifacts may be seen. This does not represent contamination.

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# **CERTIFICATE OF ANALYSIS**

# **Olerup SSP<sup>®</sup> HLA - Negative Control SSP**

Product number: 102.102-01 – including *Taq* polymerase Lot number: 05G 2011-June-01 Expiry date: Number of tests: 96 Number of wells per test: 1

#### Well specification:

Well No.	Production No.
1	2009-614-01

The negative control primer solution has been tested in a dilution series of the corresponding PCR products, 1 to  $10^3$  down to 1 to  $10^9$ .

Results: The negative control primer pairs can detect contamination with the corresponding PCR products diluted 1 to  $10^7$ .

Date of approval: 2009-July-01

Approved by:

Quality Control, Supervisor

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## **Declaration of Conformity**

Product name: Product number: Lot number:	<i>Olerup</i> SSP <sup>®</sup> HLA - Negative Control 102.101-01 05G
Intended use:	Negative Control in Olerup SSP <sup>®</sup> HLA typings.
Manufacturer:	<i>Olerup</i> SSP AB Hasselstigen 1 SE-133 33 Saltsjöbaden, 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 9001:2000 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, Hasselstigen 1, SE-133 33 Saltsjöbaden, Sweden.

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

Saltsjöbaden, Sweden 2009-July-01

Olle Olerup Managing Director

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**A**DDRESSES:

Manufacturer: *Olerup* SSP AB, Hasselstigen 1, SE-133 33 Saltsjöbaden, Sweden. *Tel:* +46-8-717 88 27 *Fax:* +46-8-717 88 18 *E-mail:* info-ssp@olerup.com *Web page:* http://www.olerup.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:* <u>support-at@olerup.com</u> *Web page:* <u>http://www.olerup.com</u>

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**.