Lot No.: **38G**

Lot-specific information

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Olerup SSP[®] HLA-B*27 – bulk

Product number:	101.532-48u – without <i>Taq</i> polymerase
Lot number:	38G
Expiry date:	2011-October-01
Number of tests:	48
Number of wells per test:	2
Storage - primer vials:	dark at -20°C
- PCR Master Mix:	-20°C
- Control DNAs:	-20°C
- Product Insert	RT

This Product Description is only valid for Lot No. 38G.

CHANGES COMPARED TO THE PREVIOUS OLERUP SSP® HLA-B*27 LOT

The HLA-B*27 specificity and interpretation tables has been updated for the HLA-B alleles described since the previous *Olerup* SSP[®] HLA-B*27 lot (Lot No. 54F) was made.

The HLA-B*27 bulk primer set is unchanged compared to the previous lot.



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PRODUCT DESCRIPTION

HLA-B*27 SSP typing

CONTENT

The primer set contains 5'- and 3'-primers for identifying the HLA-B27 specificity, B*2701 to B*2759N.

The primer solutions consist of specific primer mixes, i.e. group-specific primers as well as a *control primer pair* matching non-allelic sequences.

Positive and negative control DNAs are included in the kit.

DNA 1; a B*27-positive DNA as a positive control, IHW 9067, BTB, B*270502.

DNA 2; a B*73-positive DNA as a negative control, IHW 9280, LK707,

B*520101,7301. (A B*7301-positive DNA was chosen as negative control, as this is most similar to the B*27 group of alleles in the primer matching regions.)

PCR Master Mix complete with Taq, Taq polymerase, nucleotides, buffer, glycerol and cresol red, is included in the kit without Taq polymerase.

PCR Master Mix without Taq, nucleotides, buffer, glycerol and cresol red, is included in the kit without *Taq* polymerase.

2 PCR reactions with a reaction volume of 10 μ l is performed per sample.

INTERPRETATION

In addition to the HLA-B*27 alleles, the B*3702, B*4704 and B*4705 will be amplified by primer mix 2 of the HLA-B*27 kit.

UNIQUELY IDENTIFIED ALLELES

All the HLA-B*27 alleles, i.e. **B*2701 to B*2759N**, recognized by the HLA Nomenclature Committee in October 2009¹ are identified by the primers in the HLA-B*27 SSP kit.

In addition, the B*3702, B*4704 and B*4705 alleles are amplified by primer mix 2 of the HLA-B*27 kit.

¹HLA-B alleles listed on the IMGT/HLA web page 2009-10-19, release 2.27.0, <u>www.ebi.ac.uk/imgt/hla</u>.



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PROTOCOL

DNA EXTRACTION

Extracted, highly pure DNA is needed for SSP typings. We recommend isolation of DNA using GenoPrep B200 or GenoPrep B350 cartridges on the GenoMTM-6 robotic workstation (GenoVision Europe *Tel:* +43 1 710 15 00 or GenoVision Inc. USA *Tel:* +1 610 430 88 41; <u>http://www.genovision.com</u>). Using GenoMTM-6-extracted DNA ACD, EDTA and heparinised blood can be used as starting material. Because of its high purity, GenoMTM-6-extracted DNA can be diluted when used in combination with *Olerup* SSPTM products. The recommended DNA concentration is 15 ng/ul.

Alternatively – BUT DO NOT USE HEPARINISED BLOOD WITH THESE METHODS - the DNA can be extracted using trimethylammoniumbromide salts (DTAB/CTAB) or by salting out. Dissolve the extracted DNA in dH₂O.

IMPORTANT:

Optimal DNA concentration using: GenoMTM-6-extracted DNA, 15 ng/ μ l. DNA extracted by other methods, 30 ng/ μ l.

Concentration exceeding 50 ng/ μ l will increase the risk for nonspecific amplifications and weak extra bands, especially for HLA Class I high resolution SSP typings.

PCR AMPLIFICATION

101.532-48 – without Taq polymerase

For one HLA-B*27 bulk typing, dispense 5 μ l of each of the 2 HLA-B*27 bulk primer solutions into an 8 tube strip of 0.2 ml PCR tubes; primer solution 1 into well 1, primer solution 2 into well 2.

For one HLA-B*27 typing, add at room temperature in a 0.5 ml tube:

 $4 \times 2 \mu l = 8 \mu l DNA (30 ng/\mu l)$

4 x 3 μ l = 12 μ l PCR Master Mix complete with *Taq* – mix well before taking your aliquot

Mix well, dispense 5 μ l of the DNA-PCR Master Mix mixture into each of the 2 wells of an HLA-B*27 typing. Close the 8 tube PCR strip with an 8 strip lid.

101.532-48u – without Taq polymerase

For one HLA-B*27 bulk typing, dispense 5 μ l of each of the 2 HLA-B*27 bulk primer solutions into an 8 tube strip of 0.2 ml PCR tubes; primer solution 1 into well 1, primer solution 2 into well 2.

For one HLA-B*27 typing, add at room temperature in a 0.5 ml tube:

 $4 \times 2 \mu l = 8 \mu l DNA (30 ng/\mu l)$

4 x 3 μ l = 12 – 0.3 = 11.7 μ l PCR Master Mix complete with *Taq* – mix well before taking your aliquot

0.3 μl *Taq* polymerase (5 units/μl)

Mix well, dispense 5 μ l of the DNA-PCR Master Mix mixture into each of the 2 wells of an HLA-B*27 typing. Close the 8 tube PCR strip with an 8 strip lid.

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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 1. 1 cycle	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° 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 specific PCR products. The length of the specific PCR product is helpful in the interpretation of the results.

Record the presence of the internal positive control bands.

Lanes without either control band or specific PCR products should be repeated.

Interpret the typings with the *lot-specific Interpretation and Specificity Tables*.



Lot No.: 38G	Lot-specific information
PCR MASTER MIXES	
The PCR Master Mix com	nplete with <i>Taq</i> contains:
<i>Taq</i> polymerase	0.4 unit per 10 μl SSP reaction
nucleotides	final concentration of each dNTP is 200 μ M
PCR buffer	final concentrations: 50 mM KCl, 1.5 mM MgCl ₂ , 10 mM Tris-HCl pH 8.3, 0.001% w/v gelatin
glycerol	final concentration of glycerol is 5%
cresol red	final concentration of cresol red is 100 μ g/ml
	used for all Olerup SSP kits without Taq polymerase.
The PCR Master Mix with	
nucleotides	final concentration of each dNTP is 200 μ M
PCR buffer	final concentrations: 50 mM KCl, 1.5 mM MgCl ₂ , 10 mM Tris-HCl pH 8.3, 0.001% w/v gelatin
glycerol	final concentration of glycerol is 5%

The same PCR Master Mix is used for all Olerup SSP kits without Taq polymerase.

The PCR Master Mix complete with *Taq* and the PCR Master Mix without *Taq* can be shipped at ambient temperature.

final concentration of cresol red is 100 µg/ml

When stored at -20° C, the PCR Master Mix complete with *Taq* and the PCR Master Mix without *Taq* are stable for 24 months from the date of manufacture. Vials with the PCR Master Mixes can be kept at $+4^{\circ}$ C for 4 weeks, but the PCR Master Mixes are then no longer stable for 24 months.

cresol red



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SPECIFICITY TABLE

HLA-B*27 SSP typing

Specificity and size of the PCR product of the two primer mixes used for HLA-B*27 SSP typing.

Primer Mix	Size of spec. PCR product ¹	Size of control band ²	Amplified HLA-B*27 alleles	Other amplified HLA-B alleles ³
1	145 bp	430 bp	*2701-270508, 270510-2711, 2713- 2715, 2717, 2719- 2721, 2724, 2725, 2727, 2728, 2730, 2732-2759N	
24	95 bp	515 bp	*2701-270512, 2708, 2710, 2712, 2713, 2715-2718, 2723, 2725, 2726, 2728, 2729, 2731, 2736- 2740, 2742, 2744, 2745, 2747-2759N	*3702, 4704, 4705

¹Alleles are assigned by the presence of specific PCR product(s). However, the sizes of the specific PCR products may be helpful in the interpretation of HLA-B*27 SSP typings.

When the primers in a primer mix can give rise to specific PCR products of more than one length this is indicated if the size difference is 20 base pairs or more. Size differences shorter than 20 base pairs are not given. For high resolution SSP kits the respective lengths of the specific PCR product(s) of the alleles amplified by these primer mixes are given.

Nonspecific amplifications, i.e. a ladder or a smear of bands, may sometimes be seen. GC-rich primers have a higher tendency of giving rise to nonspecific amplifications than other primers.

PCR fragments longer than the control bands may sometimes be observed. Such bands should be disregarded and do not influence the interpretation of the SSP typings.

PCR fragments migrating faster than the control bands, but slower than a 400 bp fragment may be seen in some gel read-outs. Such bands can be disregarded and do not influence the interpretation of the SSP typings.

Some primers may give rise to primer oligomer artifacts. Sometimes this phenomenon is an inherit feature of the primer pair(s) of a primer mix. More often it is due to other factors such as too low amount of DNA in the PCR reactions, taking too long time in setting up the PCR reactions, working at elevated room temperature or using thermal cyclers that are not pre-heated

²The internal positive control primer pairs amplify segments of the human growth hormone gene. The two different control primer pairs give rise to either an internal positive control band of 430 base pairs or a band of 515 base pairs.

Well number 1 contains the primer pair giving rise to the shorter, 430 bp, internal positive control band in order to help in the correct orientation of the HLA-B*27 typing.

In the presence of a specific amplification the intensity of the control band often decreases.

³Due to the sharing of sequence motifs between HLA-B alleles three non-HLA-B*27 alleles will be amplified by primer mix 2.

⁴Short specific PCR fragments are less intense and not as sharp as longer specific bands.



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INTERPRETATION TABLE			
HLA-E	3*27 \$	SSP t	yping
Amplification patter	n of th	<u>e B*27</u>	01 to 2759N alleles ¹
		ell	
	1	2	
Length of spec.	145	95	Length of spec.
PCR product			PCR product
Length of int.	430	515	Length of int.
pos. control ²			pos. control
5'-primer ³	167	363	5'-primer ³
	^{5'} -gCT 3'	^{5'} -AAT ^{3'}	
3'-primer ⁴	272	418	3'-primer ⁴
	^{5'} -TgC ^{3'}	^{5'} -gTC ^{3'}	
Well No.	1	2	Well No.
HLA-B allele ⁵			HLA-B allele ⁵
*2701-270508, 270510-			*2701-270508, 270510-
270512, 2708, 2710, 2713,			270512, 2708, 2710, 2713,
2715, 2717, 2725, 2728, 2736-	1	2	2715, 2717, 2725, 2728, 2736-
2740, 2742, 2744, 2745, 2747-			2740, 2742, 2744, 2745, 2747-
2759N			2759N
*270509, 2712, 2716, 2718,			*270509, 2712, 2716, 2718,
2723, 2726, 2729, 2731, 3702,		2	2723, 2726, 2729, 2731, 3702,
4704, 4705			4704, 4705
*2706, 2707, 2709, 2711, 2714,			*2706, 2707, 2709, 2711, 2714,
2719-2721, 2724, 2727, 2730,	1		2719-2721, 2724, 2727, 2730,
2732-2735, 2741, 2743, 2746			2732-2735, 2741, 2743, 2746
HLA-B allele ⁵			HLA-B allele ⁵
Well No.	1	2	Well No.

¹Due to the sharing of sequence motifs between HLA-B alleles three non-HLA-B*27 alleles will

be amplified by primer mix 2; B*3702, B*4704, B*4705. ²The internal positive control primer pairs amplify segments of the human growth hormone gene. The two different control primer pairs give rise to either an internal positive control band of 430 base pairs or a band of 515 base pairs.

Well number 1 contains the primer pair giving rise to the shorter, 430 bp, internal positive control band in order to help in the correct orientation of the HLA-B*27 typing.

In the presence of a specific amplification the intensity of the control band often decreases.

³The nucleotide position, in the 2nd and 3rd exons, matching the specificity-determining 3'-end of the primer is given. Nucleotide numbering as on the www.ebi.ac.uk/imgt/hla web site. The sequence of the 3 terminal nucleotides of the primer is given. ⁴The nucleotide position, in the 2nd and 3rd exons, matching the specificity-determining 3'-end of

the primer is given in the anti-sense direction. Nucleotide numbering as on the www.ebi.ac.uk/imgt/hla web site. The sequence of the 3 terminal nucleotides of the primer is given. ⁵The sequence of the B*270501 allele has been shown to be identical to B*270502.

The B*2722 sequence shown to be identical to the corrected B*2706 sequence.



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CELL LINE VALIDATION SHEET						
HLA-B*27 unit dose SSP kit						
				We)	
					1	2
				Production No.	200964601	200064602
	IF	WC cell line	HL	A-B		
1	9001	SA	*0702		-	-
2	9280	LK707	*5201	*7301	-	-
3	9011	E4181324	*52011		-	-
4	9275	GU373	*1510	*5301	-	-
5	9009	KAS011	*3701		-	-
6	9353		*3901	*5101	-	-
7	9020		*1801		-	-
8	9025		*3501		-	-
9	9026		*3801		-	-
10		LKT3	*5401		-	-
11		PITOUT	*4403	1	-	-
12	9052		*5701		-	-
13		JESTHOM	*2705		+	+
14		OLGA	*1501	*1520	-	-
15	9075		*4001	1020	-	-
16		SWEIG007	*4001		-	
				*5504	-	-
17		CTM3953540	*0801	*5501	-	-
18		32367	*1401	*5601	-	-
19		BM16	*1801		-	-
20		SLE005	*4001		-	-
21		AMALA	*1501		-	-
22		KOSE	*3503		-	-
23	9124		*4002	*5602	-	-
24	9035	JBUSH	*3801		-	-
25	9049	IBW9	*1402		-	-
26	9285	WT49	*5801		-	-
27	9191	CH1007	*0705	*5101	-	-
28	9320	BEL5GB	*4402	*4403	-	-
29		MOU	*4403		-	-
30	9021		*4201		-	-
31		DUCAF	*1801		-	-
32		HAG	*4102		-	-
33		MT14B	*4001		-	-
34		DHIF	*3801		-	-
35		SSTO	*4402		-	-
_		KT17	*1501	*3501	1	-
36				3301	l -	-
37		HHKB	*0702		-	-
38	9099		*1501	*0705	-	-
39	9315		*0801	*2705	+	+
40		WHONP199	*1302	*4601	-	-
41		H0301	*1402	ļ	-	-
42		TAB089	*4601		-	-
43		T7526	*4601		-	-
44	9057		*3801		-	-
45		SHJO	*4201	*5001	-	-
46	9013	SCHU	*0702		-	-
47	9045	TUBO	*5101		-	-
48	0202	TER-ND	*3501	*4403	-	-



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CERTIFICATE OF ANALYSIS

Olerup SSP [®] HLA-B*27 SSP -	- bulk
Product number:	101.532-48u – without <i>Taq</i> polymerase
Lot number:	38G
Expiry date:	2011-October-01
Number of tests:	48
Number of wells per test:	2

Well specifications:

Well No.	Production No.
1	2009-646-01
2	2009-646-02

The specificity of the primer solutions of the kit has been tested against 48 well characterized IHWC cell line DNAs.

Results: No false positive or false negative amplifications were obtained.

Date of approval: 2009-November-08

Approved by:

Quality Control, Supervisor



Lot No.: 38G	Lot-specific information	
	Declaration of Conformity	
Product name: Product number: Lot number:	<i>Olerup</i> SSP [®] HLA-B*27 - bulk 101.532-48u 38G	
Intended use:	HLA-B*27 low resolution histocompatibility testing	
Manufacturer:	<i>Olerup</i> SSP AB Hasselstigen 1 SE-133 33 Saltsjöbaden, Sweden Phone: +46-8-717 88 27	

Fax: +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 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 *Olerup* SSP AB, Hasselstigen 1, SE-133 33 Saltsjöbaden, Sweden.

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

Notified Body: Lloyd's Register Quality Assurance Limited, Hiramford, Middlemarch Office Village, Siskin Drive, Coventry CV3 4FJ, United Kingdom. (Notified Body number: 0088.)

Saltsjöbaden, Sweden 2009-November-08

Olle Olerup Managing Director



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Lot-specific information

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