• LERUPSSP®

HLA-B*27 unit dose single well Product Insert

101.911-96 – including *Taq* **pol.**, IFU-01 **101.911-96u – without** *Taq* **pol.**, IFU-02

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Olerup SSP® HLA-B*27 unit dose single well¹

Product number: 101.911-96 – including *Taq* polymerase

101.911-96u – without *Taq* **polymerase**

Lot number: 7D4

Expiry date: 2018-09-01

Number of tests: 96 Number of wells per test: 1

Storage - pre-aliquoted primers: dark at -20°C

- PCR Master Mix: -20°C - Control DNAs: -20°C

This Product Description is only valid for Lot No. 7D4.

Complete product documentation consists of generic Instructions for Use (IFU), lot specific Product Insert, Worksheet and Certificate.

CHANGES COMPARED TO THE PREVIOUS *OLERUP* SSP® HLA-B*27 UNIT DOSE SINGLE WELL LOT (56Y)

The format of the Product Insert and Worksheet have been changed.

The HLA-B*27 unit dose single well specificity and interpretation tables have been updated for the HLA-B alleles described since the previous *Olerup* SSP® HLA-B*27 lot (Lot No. 56Y) was made. The kit design is based on IMGT/HLA database 3.22.0.

As of lot series V, the Specificity Table is included in the lot-specific Product Insert, and the Interpretation Table is included in the Worksheet.

The HLA-B*27 unit dose is unchanged compared to the previous *Olerup* SSP® HLA-B*27 unit dose single well lot.

1. The primers in this kit do <u>not</u> amplify the B*27:05:23, B*27:23, B*27:29, B*27:75, B*27:77, B*27:85, B*27:129 and B*27:140 alleles.



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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*27:01 to B*27:145. The primers in this kit do <u>not</u> amplify the B*27:05:23, B*27:23, B*27:29, B*27:75, B*27:77, B*27:85, B*27:129 and B*27:140 alleles.

Positive and negative control DNAs are included in the kit.

DNA 1; a B*27-positive DNA as a positive control, IHW 9315, CML, B*08:01.27:05:02.

DNA 2; a B*73-positive DNA as a negative control, **IHW 9280, LK707, B*52:01:01,73:01.** (A B*73:01-positive DNA was chosen as negative control, as this is most similar to the B*27 group of alleles in the primer matching regions.)

We recommend including one positive and one negative control DNA in each test set up. The kit contains enough control DNAs to perform 16 test set ups. If more than 16 test set ups per kit are run other positive and negative DNA samples can be used as controls (e.g. positive and negative samples from previous tests).

PLATE LAYOUT

Each 8 well strip is intended for testing 8 sample DNAs.

1 1 1 1 1 1 1 1

The cut PCR plate is marked with 'B27' in silver/gray ink.

Well No. 1 is marked with the Lot No. '7D4'.

A faint row of numbers is seen between wells 1 and 2 or wells 7 and 8 of the PCR trays. These stem from the manufacture of the trays, and should be disregarded.

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

Please note: When removing each 8 well PCR plate, make sure that the remaining plates stay covered. Use a scalpel or a similar instrument to carefully cut the foil between the plates.

INTERPRETATION

Due to the sharing of sequence motifs between HLA-B alleles the B*07:197, B*15:342, B*44:97 and B*49:26 is amplified by the B*27 unit dose single well primer mix. Thus, the interpretation of B*27 unit dose single well SSP subtypings will only be influenced by these alleles, and not by other HLA Class I alleles. The primer mix in this kit does <u>not</u> amplify the B*27:05:23, B*27:23, B*27:29, B*27:75, B*27:77, B*27:85, B*27:129 and B*27:140 alleles.

UNIQUELY IDENTIFIED ALLELES

All the B*27 alleles except the B*27:05:23, B*27:23, B*27:29, B*27:75, B*27:77, B*27:85, B*27:129 and B*27:140 alleles, i.e. B*27:01 to B*27:145, recognized by

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¹HLA-B alleles listed on the IMGT/HLA web page 2015-October-10, release 3.22.0, www.ebi.ac.uk/imgt/hla.

²Alleles that have been deleted from or renamed in the official WHO HLA Nomenclature up to and including the last IMGT/HLA database release can be retrieved from web page http://hla.alleles.org/alleles/deleted.html.

HLA-B*27 unit dose single well

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PROTOCOL

DNA EXTRACTION

Extracted, highly pure DNA is needed for SSP typings. DNA samples to be used for PCR-SSP HLA typing should be re-suspended in dH_2O . The A260/A280 ratio should be 1.6 – 2.0 by UV spectrophotometry for optimal band visualization during electrophoresis.

We recommend automated DNA extraction with the QIAGEN EZ1 DSP DNA Blood System. ACD blood should be used as starting material.

Alternatively, the DNA can be extracted by any preferred method yielding pure DNA. When using alternative methods, the DNA concentration should be adjusted to 30 ng/ μ l. **Do not use heparinised blood with these methods.**

Recommended DNA concentration using:

EZ1-extracted DNA, 15 ng/μl.

DNA extracted by other methods, 30 ng/µl.

Concentrations 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. If necessary, dilute the extracted DNA in dH₂O.

PCR AMPLIFICATION

101.911-96 - including Tag polymerase

For one HLA-B*27 typing add at room temperature to a PCR well:

2 μl sample or control DNA (30 ng/μl)

 $3~\mu l$ PCR Master Mix complete with Taq – mix well before taking your aliquot

5 ul dH₂O

101.911-96u – without Tag polymerase

For one HLA-B*27 typing add at room temperature to a PCR well:

2 μl sample or control DNA (30 ng/μl)

 $3 \mu I PCR Master Mix without \textit{Taq}$ polymerase – mix well before taking your aliquot

0.1 μl *Tag* polymerase (5 units/μl)

 $4.9~\mu l~dH_2O$

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The 8 well PCR plate is marked with the lot number. Cover the primer tray(s) with the provided adhesive PCR seals. Check that all reaction wells are completely covered to prevent evaporative loss during PCR amplification. The Olerup SSP® Compression Pad (Product No. 103.505-06) can be applied on top of the adhesive PCR seals to prevent evaporation during thermal cycling.

PCR cycling parameters:

	, J.			
1.	1 cycle	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
4.	End - hold	RT		if less than 8 hours
		4°C		if longer than 8 hours

Total reaction volume in each well, 10 µl.

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 μ l per 100 ml gel solution. For maximal ease of handling use our ethidium bromide dropper bottles (Product No. 103.301-10). Note: Ethidium bromide is a carcinogen. Handle with appropriate personal protective equipment.

Load the PCR products, preferably using an 8-channel pipette. Load a DNA size marker (100 base pair ladder, DNA Size Marker Product No. 103.202-100 or DNA Size Marker for short gel runs 103.203-100) 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*.

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Expected results:

Dependent on the presence or absence and the specificity of the B*27 alleles in the sample the following results are possible (B*27 alleles not amplified by the kit and co-amplified non-B27 alleles are excluded):

Mix	Result	
+	B*27 positive	
-	B*27 negative	

Positive control: The positive control mix has to be positive for the B*27 specific product as defined in the lot-specific Interpretation and Specificity Tables. Absence of B*27 specific bands in one or both wells might indicate failure of the test.

The negative control DNA must only give rise to the internal control bands of 430 or 515 base pairs respectively and no B*27 specific bands. Additional bands might indicate inappropriate test conditions or contamination.

PCR MASTER MIX

The PCR Master Mix including *Taq* polymerase contains:

Taq polymerase 0.4 unit per 10 μl SSP reaction

 $\begin{array}{ll} \text{nucleotides} & \text{final concentration of each dNTP is 200 } \mu\text{M} \\ \text{PCR buffer} & \text{final concentrations: 50 mM KCI, 1.5 mM MgCI}_2, \end{array}$

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 µg/ml

The same PCR Master Mix is used for all Olerup SSP kits including Taq polymerase.

The PCR Master Mix without *Taq* contains:

nucleotides final concentration of each dNTP is 200 µM PCR buffer final concentrations: 50 mM KCl, 1.5 mM MgCl₂,

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 µg/ml

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



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SPECIFICITY TABLE

HLA-B*27 SSP typing

Specificity and size of the PCR product of the HLA-B*27 unit dose single well SSP typing.

Primer	Size of spec.	Size of control band ²	Amplified HLA-	Other amplified
Mix	PCR product ¹		B*27 alleles ³	HLA-B alleles ⁴
1	145 bp, 205 bp	430 bp	*27:01-27:05:22, 27:05:24-27:21, 27:24-27:28, 27:30- 27:74, 27:76, 27:78-27:84, 27:86- 27:128, 27:130- 27:139, 27:141- 27:145	*07:197, 15:342 ^w , 44:97, 49:26

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 HLA-specific PCR products of more than one length this is indicated if the size difference is more than 20 base pairs. Size differences of 20 base pairs or less are not given. For high resolution SSP kits, the alleles listed are specified according to amplicon length.

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 control primer pair gives rise to a band of 430 base pairs.

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

 3 The primers in this kit do <u>not</u> amplify the B*27:05:23, B*27:23, B*27:29, B*27:75, B*27:77, B*27:85, B*27:129 and B*27:140 alleles.

⁴Due to the sharing of sequence motifs between HLA-B alleles the B*07:197, B*15:342^w, B*44:97 and B*49:26 is amplified by the B*27 unit dose single well primer mix. 'w', may be weakly amplified.

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PRIMER SPECIFICATION

Well No.	1
Length of spec.	145
PCR product	205
Length of int.	430
pos. control ¹	
5'-primer(s) ²	167
	^{5'} -gCT ^{3'}
	363
	363 ^{5'} -AAT ^{3'}
3'-primer(s) ³	272
	^{5'} -TgC ^{3'}
	272
	^{5'} -TgC ^{3'}
	527
	5' -CCT 3'
Well No.	1

¹The internal positive control primer pairs amplify segments of the human growth hormone gene. The control primer pair gives rise to a band of 430 base pairs.

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

²The nucleotide position 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.

³The nucleotide position 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.

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CE	CELL LINE VALIDATION SHEET						
HLA-B*27 unit dose SSP kit ²							
				Well			
					1		
				Production No.	201665801		
				nci	365		
				Prod No.	216		
				ΔZ	2(
	IH	WC cell line ¹		A-B			
1	9001		*07:02		-		
2		LK707	*52:01	*73:01	-		
3		E4181324	*52:01				
4		GU373	*15:10	*53:01	_		
5		KAS011	*37:01		-		
6	9353		*39:01	*51:01	_		
7	9020		*18:01		-		
8	9025		*35:01		-		
9		YAR	*38:01		-		
10		LKT3	*54:01		-		
11		PITOUT	*44:03		-		
12	9052		*57:01		_		
13		JESTHOM	*27:05		+		
14		OLGA	*15:01	*15:20	-		
15	9075		*40:01		-		
16		SWEIG007	*40:02		-		
17		CTM3953540	*08:01	*55:01	-		
18		32367	*14:01	*56:01	_		
19		BM16	*18:01		-		
20		SLE005	*40:01		_		
21		AMALA	*15:01				
22		KOSE	*35:03	+=0.00			
23	9124		*40:02	*56:02	_		
24		JBUSH	*38:01		_		
25		IBW9	*14:02				
26		WT49	*58:01	*54.04			
27		CH1007	*07:05	*51:01 *44:03			
28		BEL5GB	*44:02	44:03	_		
29		MOU	*44:03		_		
30	9021		*42:01		븬		
31		DUCA F HAG	*18:01 *41:02				
32 33		MT14B	*40:01				
34	9098		*38:01				
35		SSTO	*44:02		\vdash		
36		KT17	*15:01	*35:01	\vdash		
37		HHKB	*07:02	33.01	\vdash		
38	9099		*15:01		\vdash		
39	9315		*08:01	*27:05	+		
40		WHONP199	*13:02	*46:01			
41		H0301	*14:02	70.01	\vdash		
42		TAB089	*46:01				
43		T7526	*46:01				
44	9057		*38:01				
45		SHJO	*42:01	*50:01			
46		SCHU	*07:02	00.01			
47		TUBO	*51:01				
48		TER-ND	*35:01	*44:03			
70	3303	ILIVIND	55.01	77.00	انسا		



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¹The provided cell line HLA specificities are retrieved from the http://www.ihwg.org/hla web site. The specificity of an individual cell line may thus be subject to change.

²The specificity of each primer solution in the kit has been tested against 48 well characterized cell line DNAs and where applicable, additional cell line DNAs.

One additional 3'-primer was tested by separately adding one additional 5'-primer.

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