●LERUPSSP®HLA-A*80 Product Insert Page 1 of 12

101.434-06 – including *Taq* **polymerase**, IFU-01 **101.434-06u – without** *Taq* **polymerase**, IFU-02

Visit <u>www.olerup-ssp.com</u> for "Instructions for Use" (IFU)

Lot No.: **09X** Lot-specific information

Olerup SSP® HLA-A*80

Product number: 101.434-06 – including *Taq* polymerase

101.434-06u - without *Taq* polymerase

Lot number: 09X

Expiry date: 2017-January-01

Number of tests: 6
Number of wells per test: 4+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. 09X.

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-A*80 LOT (03S)

The HLA-A*80 kit is updated for new alleles to enable separation of:

- Confirmed¹ alleles as listed in the IMGT/HLA database
- Polymorphisms in exons outside of the region encoding the peptide binding domain
- Null and Alternatively expressed alleles

A well containing Negative Control primer pairs has been added.

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

Two wells have been added to HLA-A*80, wells 4 to 5.

The HLA-A*80 primer set, specificity and interpretation tables have been updated for the HLA-A alleles described since the previous *Olerup* SSP[®] HLA-A*80 lot was made (Lot No. 03S). The kit design is based on IMGT/HLA database 3.16.0.

July 2014 Rev. No.: 00 **008**

¹As described in section Uniquely Identified Alleles.

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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 primers of the wells detailed below have been exchanged, added or modified compared to the previous lot.

Well	5'-primer	3'-primer	rationale
4	New	New	New primer pair for the A*80:03 allele.
5	-	-	Negative Control.

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Well **5** contains <u>Negative Control primer pairs</u>, that will amplify more than 95% of the *Olerup* SSP[®] HLA Class I, DRB, DQB1, DPB1 and DQA1 amplicons as well as all the amplicons generated by the control primer pairs matching the human growth hormone gene.

HLA-specific PCR product sizes range from 75 to 200 base pairs. The PCR product generated by the positive control primer pair is 430 base pairs.

Length of PCR	105	200	105	80	75	80	85
product							
5'-primer ¹	164	340	440	45	45	43	36
-	5'-CAC3'	5'-Agg ^{3'}	^{5'} -TTA3'	⁵ '-Tgg ³ '	⁵ '-Tgg ³ '	⁵ '-Tgg ³ '	5'-TAC3'
							36
							^{5'} -TAT ^{3'}
3'-primer ²	231	2 nd I	507	59	58	57	47
-	⁵ '-TgC ³ '	^{5'} -AAA ^{3'}	^{5'} -TTg ^{3'}	5'-CTC3'	^{5'} -ggC ^{3'}	5'-CTC3'	5'-ACA3'
							48
							^{5'} -gCA ^{3'}
							48
							^{5'} -gCC ^{3'}
							52
							5'-TgT ^{3'}
A*	+	+	+				
B*	+	+	+				
C*	+	+	+				
DRB1				+	+		
DRB3				+	+		
DRB5				+			
DQB1					+		
DPB1						+	
DQA1							+

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

²The nucleotide position for HLA class I genes and the codon for HLA class II genes, in the 2nd or 3rd exon or the 2nd intron, matching the specificity-determining 3'-end of the primer is given in the anti-sense direction. Nucleotide and codon 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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Lot No.: **09X** Lot-specific information

PRODUCT DESCRIPTION

HLA-A*80 SSP subtyping

CONTENT

The primer set contains 5'- and 3'-primers for identifying the HLA-A*80:01 to A*80:03 alleles.

PLATE LAYOUT

Each test consists of 5 PCR reactions in an 8 well cut PCR plate. Wells 6 to 8 are empty.

1 2 3 4 NC empty empty empty

The 8 well cut PCR plate is marked with 'A80' in silver/gray ink.

Well No. 1 is marked with the Lot No. '09X'.

Wells 1 to 4 – HLA-A*80 high resolution primers.

Well 5 – Negative Control (NC).

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 heat-sealed with a PCR-compatible foil.

Please note: When removing each 8 well PCR plate, make sure that the remaining plates stay sealed. 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-A alleles non-HLA-A*80 alleles will be amplified by primer mixes 1, 3 and 4.

For further details see Specificity Table.

UNIQUELY IDENTIFIED ALLELES

The HLA-A*80 alleles, i.e. **HLA-A*80:01** and **HLA-A*80:03**, recognized by the HLA Nomenclature Committee in April 2014^{1,2} will be amplified by the primers in the HLA-A*80 subtyping kit³.

The HLA-A*80kit enables separation of the confirmed HLA-A*80 alleles as listed in the IMGT/HLA database. An HLA allele is listed as confirmed by IMGT/HLA if it has been sequenced by more than a single laboratory or from multiple sources. Current allele confirmation status for HLA-A*80 alleles is listed below.

The HLA-A*80 kit also enables identification of polymorphisms in exons outside of the region encoding the peptide binding domain and of null and alternatively expressed alleles

The HLA-A*80 subtyping kit cannot distinguish the silent mutations in the A*80:01:01:01 and 80:01:01:02 alleles.

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ALLELE CONFIRMATION STATUS

Allele	Status ¹			
A*80:01:01:01	Confirmed			
A*80:01:01:02	Unconfirmed			
A*80:02	Unconfirmed			
A*80:03	Unconfirmed			

¹Allele status "confirmed" or "unconfirmed" as listed on the IMGT/HLA web page 2014-April-17, release 3.16.0, www.ebi.ac.uk/imgt/hla.

RESOLUTION IN HOMO- AND HETEROZYGOTES

Results file with resolution in HLA-A*80 homo- and heterozygotes is available upon request.

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¹HLA-A alleles listed on the IMGT/HLA web page 2014-April-14, release 3.16.0, www.ebi.ac.uk/imgt/hla.

Alleles 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.

³The A*80:02 and the A*11:166, 30:56 and 31:85 alleles give rise to identical amplification patterns with the HLA-A*80 primer set. These alleles can be distinguished by the HLA-A low resolution and/or the HLA-A*11, HLA-A*30 and HLA-A*31 kits.

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SPECIFICITY TABLE

HLA-A*80 SSP subtyping

Specificities and sizes of the PCR products of the 4+1 primer mixes used for HLA-A*80 SSP subtyping

Primer Mix	Size of spec. PCR product ¹	Size of control band ²	Amplified HLA-A*80 alleles	Other amplified HLA Class I alleles ³
1	135 bp	800 bp	*80:01:01:01-80:03	*11:166, 30:56, 31:85
2	155 bp	1070 bp	*80:01:01:01- 80:01:01:02, 80:03	
3	165 bp	1070 bp	*80:02	*11:166, 30:56, 31:85
4	235 bp	1070 bp	*80:03	*01:12, 01:14, 01:19, 03:97, 03:122, 03:167, 11:27, 24:92, 30:45, 30:75
5 ⁴	-	-	Negative Control	

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-A*80 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 internal positive control bands are 1070 or 800 base pairs respectively, well distribution as outlined in the table. Well number 1 contains the shorter, 800 bp, internal positive control band. The well distribution of the internal controls can help in orientation of the kit on gel photo, as well as allow for kit identification. In the presence of a specific amplification the intensity of the control band often decreases.

³Due to the sharing of sequence motifs between HLA-A alleles non-HLA-A*80 alleles will be amplified by primer mixes 1, 3 and 4.

⁴Primer mix 5 contains a negative control, which will amplify more than 95% of HLA amplicons as well as the amplicons generated by control primer pairs. PCR product sizes range from 75 to 200 base pairs. The PCR product generated by the control primer pair is 430 base pairs.

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101.434-06 – including *Taq* **polymerase**, IFU-01 **101.434-06u – without** *Taq* **polymerase**, IFU-02

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Lot No.: **09X**

Lot-specific information

PRIMER SPECIFICATION

Well No.	1	2	3	4
Length of spec.	135	155	165	235
PCR product				
Length of int.	800	1070	1070	1070
pos. control ¹				
5'-primer(s) ²	176	176	176	363
	^{5'} -gCA ^{3'}	^{5'} -gCA ^{3'}	^{5'} -gCA ^{3'}	^{5'} -ATA ^{3'}
3'-primer(s) ³	270	292	299	559
	^{5'} -ACA ^{3'}	^{5'} -gTT ^{3'}	5' -CCA 3'	^{5'} -CgT ^{3'}
Well No.	1	2	3	4

¹The internal positive control primer pairs amplify segments of the human growth hormone gene. The internal positive control bands are 1070 or 800 base pairs respectively, well distribution as outlined in the table. Well number 1 contains the shorter, 800 bp, internal positive control band. The well distribution of the internal controls can help in orientation of the kit on gel photo, as well as allow for kit identification. 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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CELL LINE VALIDATION SHEET									
		HLA-A	*80 SSI	P kit⁴					
					Well				
					1	2	3	4	
				Lot No.:	201072301	201072302	201315303	201438604	
	IHV	/C cell line ¹	A *	A*					
1	9001		*24:02		-	-	-	-	
2	9280	LK707	*02:01		-	-	-	-	
3	9011	E4181324	*01:01		-	-	-	-	
4	9275	GU373	*30:01		-	-	-	-	
5	9009	KAS011	*01:01		-	-	-	-	
6	9353	SM	*02:01	*26:03	-	-	-	-	
7	9020	QBL	*26:01		-	-	-	-	
8	9025	DEU	*31:01		-	-	-	-	
9	9026	YAR	*26:01		-	-	-	-	
10	9107	LKT3	*24:02		-	-	-	-	
11	9051	PITOUT	*29:02		-	-	-	-	
12	9052	DBB	*02:01		-	-	-	-	
13	9004	JESTHOM	*02:01		-	-	-	-	
14	9071	OLGA	*31:01		-	-	-	-	
15	9075	DKB	*24:02		-	-	-	-	
16	9037	SWEIG007	*29:02		-	-	-	-	
17	9282	CTM3953540	*03:01	*80:01	+	+	-	-	
18	9257	32367	*33:03	*74:01	-	-	-	-	
19		BM16	*02:01		-	-	-	-	
20		SLE005	*02:01		-	-	-	-	
21		AMALA	*02:17		-	-	-	-	
22		KOSE	*02:01		-	-	-	-	
23	9124		*02:01	*34:01	-	-	-	-	
24		JBUSH	*32:01		-	-	-	-	
25		IBW9	*33:01		-	-	-	-	
26		WT49	*02:05		-	-	-	-	
27		CH1007	*24:10	*29:01	-	-	-	-	
28		BEL5GB	*02:01	*29:02	-	-	-	-	
29	9050		*29:02	20.02	_	_		-	
30	9021		*30:01	*68:02	_	-	-	-	
31		DUCAF	*30:02	00.02	H	-	Ť	Ť	
32		HAG	*02:01			-	Ė	Ē	
33		MT14B	*31:01			_	Ē	Ē	
34	9104		*31:01			_	Ė	Ē	
35		SSTO	*32:01		-	-			
36		KT17	*02:06	*11:01	-	_	Ë	Ē	
37		HHKB	*03:01	11.01		_	Ė	-	
38	9099		*02:17		-	-	Ė	-	
38 39	9099		*01:01	*03:01		-	Ë	-	
39 40		WHONP199	*02:07	*30:01		_	Ë	Ė	
40 41				30.01	Ė	-	÷	-	
		H0301	*03:01		-	-	-	-	
42		TAB089 T7526	*02:07	*00.07	-	-	Ŀ	-	
43			*02:06	*02:07	_	-	-	-	
44	9057		*66:01	*0.4.00	_	-	-	-	
45		SHJO	*23:01	*24:02	-	-	-	-	
46		SCHU	*03:01	*00 O 1	-	-	-	-	
47		TUBO	*02:16	*03:01	-	-	-	-	
48	9303	TER-ND	*02:01	*11:01	-	-		-	



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101.434-06 - including Taq polymerase, IFU-01 101.434-06u - without *Taq* polymerase, IFU-02

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Lot No.: **09X Lot-specific information**

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.

The 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.

No DNAs carrying the alleles to be amplified by primer solutions 3 and 4 were available. The specificities of the primers in primer solutions 3 and 4 were tested by separately adding one additional 5'-primer, respectively one additional 3'-primer.



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101.434-06 – including *Taq* **polymerase**, IFU-01 **101.434-06u – without** *Taq* **polymerase**, IFU-02

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101.434-06 – including *Taq* **polymerase**, IFU-01 **101.434-06u – without** *Taq* **polymerase**, IFU-02

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101.434-06 – including *Taq* **polymerase**, IFU-01 **101.434-06u – without** *Taq* **polymerase**, IFU-02

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Lot No.: **09X** Lot-specific information

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