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Supplemental Information

Antigen Identification for Orphan T Cell Receptors Expressed on Tumor-Infiltrating Lymphocytes

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Cluster 1 Peptides	Cluster 1 Predictions	Cluster 2 Peptides	Top 10 Cluster 2 Predictions
SMLGIGIVPV	EAAGIGILTV	MMWDRGMGLL	MLWDVQSGQM
SMAGIGIVDV	TLGGIGLVTV	IMEDVGWLVN	LLLQVGLSLL
NMGGLGIMPV	ILLGIGIYAL	MMWDRGLGMM	SLEDVVMLNV
NLSNLGILPV	ILSGIGVSQV	I LEDRGFNQV	MLEDRLDFVM
SMLGIGIYPV	IMGNLGLIAV	LMFDRGMSLL	MLEDMSLGIM
TMAGIGVHVV	MAGNLGIITL	LMLDFDGSLL	SLENRGLSML
SMAGIGTLVV	IMGNLGLIVL	IMEDRGSLNM	I LDDGGFLLM
SMSGILGILPM	ILAGLGTSL	LMNDMGFHV	LLWNFGLLIV
SMAGIGIVPV	ELGGLKISTL	IMEDRGSGEM	LLFDISFLML
SMLGIGIVDV		LMWDVGLSIM	IMGDRNRNLL
NMAGIGMGTV		SMWDRGTFIM	
SMLGIGILPV		LMLDRGSPNM	
SLSGIGISAV		IMFDRGIGIM	
DLAGLGLYPV		I LFD RGMNLM	
NMAGIGIIQV		MLLDRGLSLM	
NMGGLGILPV		IMEDRGSLIL	
SMAGIGIYPV		LMRDYQLLQV	
NLSNLGIVPV		LMFDRGMSVL	
IMLGIGIDTL		LMEDIGRELV	
NLSNLGIMPV		I LEDRGMGLL	
SMLGIGIVLV		MMDQFNGLMM	
SMAGIGVHVV		IMWDRDYGVM	
NMAGIGILTV		MMWDRGFNQV	
MMAGIGIVDV		IMSMSVSNYL	
NMGGLGIVPV		AMGDGSYLLM	
SMLGIKIVPV		SMWDRGMGLL	
ELSGLGIQTV		MMENRGSGAL	
SMLGIGILPM		LMWDSGLELM	
SMAGIGILPV		SMWDRGLGMM	
SMLGIGIVPV		LMWDVGWLVN	
		MMWDRGTFIM	
		MMWDRGIVPV	
		I LFD RGMNLM	

Table S1. DMF5 selection data and human target prediction.

The sequences identified from the round 3 deep-sequencing of the DMF5 10mer library selections after clustering by reverse hamming distance. Using these clusters, predictions were made on the Uniprot database using 2014PPM. The 9 predictions for the 'GIG' cluster and top 10 predictions for the 'DRG' cluster are listed.

Related to Figure 2 and Figure S1.

NKI2 9mers	NKI2 10mers		NKI2 11mers
VMISHENFM	VMNGDSGTFL	TLMSRSDLFL	ILSNRGHEVFV
TMQSHEVML	YMAVRSENFM	ILNSRDEAMM	ILSNRGHENFM
TMQSHENFM	RMPNKQENFV	ALNSRDEAMM	ILSNRGHDVFM
VMQSHEVML	IMDSKSEHFM	ALDSRLEFFV	ILSNRGHEIFL
VMISHEIFL	IMDSREEVFV	VMDSRLEFFV	ILSNRGHEYFL
IMTSHEVML	IMDSRSEHFM	ALDSRSELF	
IMTSHEVMM	GMDSRAEVFM	AMYSNSDFMV	
VMESHDFVM	ALDSRSEYFL	VMDSRLEHFM	
IMNSHEVMM	KMANRDENFV	SMNSRSEHFM	
SMNSHEVMM	RLDGQDTKFM	SMNSKSENFL	
KMNSHEVMM	LMDSRSEHFM	VLDSSSSSFL	
AMQGHEYFL	IMNSRSELF	ALDSRSENFL	
AMQGHEIFL	MMNVRSELFV	ALDSKSENFL	
VLQSHEVSM	TMNVRSELFV	ALDSRSEIFL	
AMQSHEVTL	KMNSRSELF	SMNSRADMFV	
LMSGDYQFV	TMNVRSEHFM	SMYSRQEMMV	
TMHNHEVMM	SMNSRSELF	RMWSRSEDMV	
VMHNHEVMM	KMNSRSEHFM	VLRARSDVFV	
TMTGHEVFM	TMQSHDASFL	ALDSREEVFV	
TMTGHEVFV	VMQGHASFL	SMNSREEIFL	
VMQGHESFL	KMNSHSGTFL	SMSGFSESFV	
VMISHEVML	KMNGKSEDFM		
TMTGHEVML	DMDNRLDRDM		
SMVGMEHSM	IMDSKSEIFL		
AMQGHEHFM	SMNSHSGTFL		
VMEGDYWFL	SMNSREEHFM		
SMQSHEWML	IMNSHSGTFL		
YMQTHESFM	IMDSKSENFL		
	AMDSKSENFL		
	IMDSRADMFV		
	SMNSREEVFV		
	KMNSREEVFV		
	ALDSRSEHFM		
	AMDSRSEHFM		
	AMDSRADMFV		
	LMDSRSQIFV		
	GMTSRSDYMV		
	VMNSRSEHFM		
	VMNSRSDWFL		
	YMNSHDPYTV		
	RMDSRSQDFV		
	RMEAHSSHVF		

Table S2. NKI2 selection data by peptide length.

The sequences identified from the round 3 deep-sequencing of the NKI2 library selections listed by peptide length. Related to Figure 3.

HLA	Patient A	Patient B
A	2:01	2:01
	2:01	2:06
B	7:02	15:01
	15:01	35:01:00
C	ND	ND
	ND	ND
DRB1	1:01	4:07
	4:04	4:07
DRB345	4*01:01	4*01:01
	ND	4*01:01
DQA	1:01	3:01
	3:01	3:01
DQB	3:02	3:02
	5:01	3:02

Table S3. Patient HLA typing results.

The HLA type of two patients with colorectal adenocarcinoma. ND = Not determined. Related to Figure 4.

Patient A					
Tumor	Healthy	V β	CDR3 β	V α	CDR3 α
23	12	TRBV7-2	CASSLGLEQFF	TRAV8-3	CAGGGGADGLTF
6	0	TRBV7-3	CASSLGGGHTEAFF	TRAV19	CALSEAEAAGNKLTF
5	0	TRBV7-9	CASSLVNGLGYTF	TRAV19	CALSEAGMDSNYQLIW
4	0	TRBV15	CATSRDRGQDEKLFF	TRAV14/DV4	CAMREGRYSGAGSYQLTF
4	0	TRBV9	CASSADTGVNQPQHF	TRAV10	CVVTETNAGKSTF
4	0	TRBV10-1	CASSRDTVNTEAFF	TRAV19	CALSEARGGATNKLIF
1	0	TRBV20-1	CSARDYQGSQPQHF	TRAV12-2	CAVNSGNTGKLIF
1	0	TRBV20-1	CSARDYQGSQPQHF	TRAV20	CAVPFLYNQGGKLIF
1	0	TRBV9	CASSADTGVNQPQHF	TRAV12-2	CAVNDFNKFYF
Patient B					
Tumor	Healthy	V β	CDR3 β	V α	CDR3 α
35	0	TRBV11-2	CASSQGVGQFKNTQYF	TRAV12-2	CAVETSNTGKLIF
23	0	TRBV7-2	CASSLSGRQGGSYEQYF	TRAV29/DV5	CAASSTGNQFYF
21	0	TRBV9	CASSSSGGLVDTQYF	TRAV19	CALSAGASGAGSYQLTF
20	0	TRBV2	CASMGRSYGYTF	TRAV39	CALMNYGGATNKLIF
16	0	TRBV11-3	CASSLETGTAIYEQYF	TRAV13-1	CAADNNNARLMF
12	0	TRBV11-3	CASSPSGLAGSNLGNQFF	TRAV19	CALSSRGSTLGRLYF
11	0	TRBV5-1	CASSRIDSTDTQYF	TRAV4	CLVGEVGTASKLTF
10	0	TRBV19	CASSIPRGSSQPQHF	TRAV12-2	CAVDSSGGYNKLIF
8	0	TRBV10-3	CAIKGGDRGVNTEAFF	TRAV14/DV4	CAMREPNNAGNMLTF
4	3	TRBV20-1	CSARLASYNEQFF	TRAV12-2	CAVRRATDSWGKLQF
1	1	TRBV10-1	CASSRDFVSNEQYF	TRAV19	CALSEARGGATNKLIF

Table S4. Orphan TCRs screened on the HLA-A*02:01 library.

The TCRs screened on the HLA-A*02:01 library. TCR sequences were chosen based on clonality in the tumor, phenotypic profile, exclusivity to the tumor, and additionally by related TCR sequences. The number beneath tumor and healthy labels indicate the number of times a paired TCR sequence was seen from this tissue.

Related to Figures 5 and 6.

		TCR1A			
Name	Sequence	Library	2014PPM	2017PWM	2017DL
91	SMGVTYEM	1			
92	YMGVSYEM	2			
93	YMGVVYEM	3			
94	KMGVTYEM	4			
95	FMGVTYEM	6			
9DL1	NMEVTYEI			1	1
9DL2	FITVTEEI			82	2
9DL3	HIQVTNEI				3
9DL4	HLIVSYEL			27	4
9DL5	HLGVTKEL			5	5
9PWM1	RLGVTYFV			2	
9PWM2	YLPVTYHI			3	60
9PWM3	GLGQTYEI			4	19
Ex1	EYGVSYEW				
Ex2	EYGVQNYV				
9EX5	EMGVSYEM				

Table S5. Peptide prediction results for TCR 1A.

Peptides tested to stimulate TCR 1A cells. Peptides are derived from the library, 2014PPM, 2017PWM and 2017DL predictions and the rank of the peptide from the predictions or library selections are shown. Cells are left intentionally blank. Related to Figure 7, S3, and S4.

Name	TCR 2A							TCR 3B						
	Peptide	Library	2014PPM	2017PWM	2017DL	Library	2014PPM	2017PWM	2017DL	Library	2014PPM	2017PWM	2017DL	
1s1	LMDMHNGQ	1												
1s2	RIIDAMNGQ	2												
1s3	RMDYNNMQ	3												
1s4	SMDTFQGQ	4				9								
1s5	GMDYHNGH	5												
2s1	TMDFYQGQ	41				1								
2s2	KMDYFSGQ	42				2								
2s3	SMDWFQGQ					3								
2s4	LMDYWQGQ					4								
2s5	NMMWFQGQ					5								
1sj1	VLDLJFQGQ		1	6							3	2		
1sj2	MMDFFNAQ		2	1							2	160		
1sj3	LINLNGQ		3	28										
1sj4	QMDYEEGQ		4	5							1	100		
1sj5	GLSSQNGQ		5											
2sj1	TLHYEYEMH		154							1				
DG1	VIDFLNQ			21	10						44			
1sDL1	VIDQLNGQ			3	1						24	154		
1sDL2	VVDFLKGQ			31	2						8	16		
1sDL3	QVDGESVQ				3									
1sDL4	HIDYFNNQ			2	4						32	49		
1sDL5	VVIGEQQ				5									
DG2	HVDYKNVQ			9	9									
1sPWM4	AISMFGQ			4							197	166		
DG3	RIDPHDGQ			8							94	11		
DG4	SLDSYNAQ		10	37										
2sDL1	TIHSYQGQ										8	1		
2sDL3	SLDFYQGH										7	3		
2sDL4	IIDAFKGQ											4		
2sDL6	RIDDGQGQ										365	6		
2sDL7	TILQYSGQ										70	7		
2sPWM4	LVDFWEAQ			40							4			
2sPWM5	TIAHYSQ										5	30		
2sPWM6	KVDFPQDQ										6	97		
DG6	TIRELQGQ										148	13		
DG7	FIDFVQRQ										130	14		

Table S6. Peptide prediction results for TCRs 2A and 3B.

Peptides tested to stimulate TCRs 2A and 3B cells. Peptides are derived from the library, 2014PPM, 2017PWM, and 2017DL predictions and the rank of the peptide from the predictions or library selections are shown. Cells are left intentionally blank. Related to Figure 7, S3, and S4.

		TCR4B			
Name	Peptide	Library	2014PPM	2017PWM	2017DL
B9_1	RMEQVDWTV	1			
B9_2	KLEFMDWRL	2			
B9_3	WLDNFELCL	3			
B9_4	TLEYMDWLV	4			
B9_5	EMMLFDWKV	5			
B9DL1	KLEQLDWTV			2	1
B9DL2	TMETIDWKV			1	2
B9DL3	DLEQMEQTV			59	3
B9DL4	TLEELDWCL			102	4
B9DL5	TLEDMAWRL			75	5
B9DL6	NVEEMDWLI			4	6
B9DL7	NVEEMDWMV			3	7
B9DL8	LLEDLDWDV				8
B9DL9	TLEAMNTTV			914	9
B9DL10	VLEEVDWLI			122	10
B9DL11	WLEDVEWQV			18	11
B9PWM1	KMENFDKTV			5	50
B9PWM2	NMEYMTWDV			6	36
B9PWM3	FVENVEWRV			7	399
B9PWM4	NVEYYDIKL			8	
B9PWM5	HLEQVDKAV			9	34
B9PWM6	EMEQVDAVV			10	287

Table S7. Peptide prediction results for TCR 4B.

Peptides tested to stimulate TCR 4B cells. Peptides are derived from the library, 2014PPM, 2017PWM, and 2017DL predictions and the rank of the peptide from the predictions or library selections are shown. Cells are left intentionally blank. Related to Figure 7, S3, and S4.

Name	Primer Sequence
ILL_F	AATGATACGGCGACCACCGAGATCTACACTCTTCCCTACACGACGCTCTCCGA
ILL_R	CAAGCAGAAGACGGCATACGAGATCGGTCTCGGCACTTCTGCTGAACCGCTCTTC
ILL_B2M_R	ATTCTGCTGAACCGCTCCCGATCTNNNNNNNNCAGGATGCTACTGTAAACTTG
ILL_BC_1	CTACACGACGCTCTCCGATCTNNNNNNNTAGCATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_2	CTACACGACGCTCTCCGATCTNNNNNNNTTATGCTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_3	CTACACGACGCTCTCCGATCTNNNNNNNTTACATAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_4	CTACACGACGCTCTCCGATCTNNNNNNNTCCAGTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_5	CTACACGACGCTCTCCGATCTNNNNNNNTTCTCTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_6	CTACACGACGCTCTCCGATCTNNNNNNNTTGACATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_7	CTACACGACGCTCTCCGATCTNNNNNNNTTGACATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_8	CTACACGACGCTCTCCGATCTNNNNNNNTTGGTGTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_9	CTACACGACGCTCTCCGATCTNNNNNNNTTGGATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_10	CTACACGACGCTCTCCGATCTNNNNNNNTTGGTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_11	CTACACGACGCTCTCCGATCTNNNNNNNTTGGTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_12	CTACACGACGCTCTCCGATCTNNNNNNNTTACCTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_13	CTACACGACGCTCTCCGATCTNNNNNNNTGTAGTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_14	CTACACGACGCTCTCCGATCTNNNNNNNTTGTAGTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_15	CTACACGACGCTCTCCGATCTNNNNNNNTTAACTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_16	CTACACGACGCTCTCCGATCTNNNNNNNTTACTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_17	CTACACGACGCTCTCCGATCTNNNNNNNTTACTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_18	CTACACGACGCTCTCCGATCTNNNNNNNTTGACTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_19	CTACACGACGCTCTCCGATCTNNNNNNNTTGATTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_20	CTACACGACGCTCTCCGATCTNNNNNNNTTCCCTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_21	CTACACGACGCTCTCCGATCTNNNNNNNTTGGTATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_22	CTACACGACGCTCTCCGATCTNNNNNNNTTACGTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_23	CTACACGACGCTCTCCGATCTNNNNNNNTTACCTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_24	CTACACGACGCTCTCCGATCTNNNNNNNTTACCTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_25	CTACACGACGCTCTCCGATCTNNNNNNNTGCACTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_26	CTACACGACGCTCTCCGATCTNNNNNNNTGCACTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_27	CTACACGACGCTCTCCGATCTNNNNNNNTGCGTATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_28	CTACACGACGCTCTCCGATCTNNNNNNNTGCGTATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_29	CTACACGACGCTCTCCGATCTNNNNNNNTGCTACTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_30	CTACACGACGCTCTCCGATCTNNNNNNNTGCTATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_31	CTACACGACGCTCTCCGATCTNNNNNNNTGCTCTTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_32	CTACACGACGCTCTCCGATCTNNNNNNNTTGAAGTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_33	CTACACGACGCTCTCCGATCTNNNNNNNTTGGATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_34	CTACACGACGCTCTCCGATCTNNNNNNNTTGGATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_35	CTACACGACGCTCTCCGATCTNNNNNNNTTGGCAATTAGCATGTATTGCTAGCGTTTTGGCT
ILL_BC_36	CTACACGACGCTCTCCGATCTNNNNNNNTTGGCATTAGCATGTATTGCTAGCGTTTTGGCT
A2_8mer_S	TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNTGGTGGAGGTTCTGGAGGTGGTG
A2_9mer_S	TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNTGGTGGAGGTTCTGGAGGTGGTG
A2_10mer_S	TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNTGGTGGAGGTTCTGGAGGTGGTG
A2_11mer_S	TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNTGGTGGAGGTTCTGGAGGTGGTG
A2_V5_AS	GAAGAGGATTAGGAATGGTTACCGGATCCAGGATGGTCCCCTCAATGTCAATG
A2_MYC_AS	ACCACACCCACAGATCCACACCCCAAGCTTCTCCGGAGATAAGCTTTTGTTC
A2_HA_AS	CGTAGTCTGGAACGTCGTATGGTAGGATCCAGAGGATGTTCCCATCTCAATGTCAATG
A2_VSV_AS	GTCTATTCAATTCAGTATAGGATCCAGAGGATGTTCCCATCTCAATGTCAATG
A2_Vector	TTTTCAATTAAGATGCAGTTACTTCGCTGTTTTTCAATATTTTCTGTTATTGCTAGCGTTTTGGCT

Table S8. Primers used to generate and deep-sequence pHLA libraries.

Letter codes for nucleotides follow the IUPAC naming system. Related to Figure 1.