## 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 | IMEDVGWLNV | LLLQVGLSLL |
| NMGGLGIMPV | ILLGIGIYAL | MMWDRGLGMM | SLEDVVMLNV |
| NLSNLGILPV | ILSGIGVSQV | ILEDRGFNQV | MLEDRDLFVM |
| SMLGIGIYPV | IMGNLGLIAV | LMFDRGMSLL | MLEDMSLGIM |
| TMAGIGVHVV | MAGNLGIITL | LMLDFDGSLL | SLENRGLSML |
| SMAGIGTLVV | IMGNLGLIVL | IMEDRGSLNM | ILDDGGFLLM |
| SMSGLGILPM | ILAGLGTSLL | LMNDMGFHIV | LLWNFGLLIV |
| SMAGIGIVPV | ELGGLKISTL | IMEDRGSGEM | LLFDISFLML |
| SMLGIGIVDV |  | LMWDVGLSIM | IMGDRNRNLL |
| NMAGIGMGTV |  | SMWDRGTFIM |  |
| SMLGIGILPV |  | LMLDRGSPNM |  |
| SLSGIGISAV |  | IMFDRGIGIM |  |
| DLAGLGLYPV |  | ILFDRGMNLM |  |
| NMAGIGIIQV |  | MLIDRGLSLM |  |
| NMGGLGILPV |  | IMEDRGSLIL |  |
| SMAGIGIYPV |  | LMRDYQLLQV |  |
| NLSNLGIVPV |  | LMFDRGMSVL |  |
| IMLGIGIDTL |  | LMEDIGRELV |  |
| NLSNLGIMPV |  | ILEDRGMGLL |  |
| SMLGIGIVLV |  | MMDQFNGLMM |  |
| SMAGIGVHVV |  | IMWDRDYGVM |  |
| NMAGIGILTV |  | MMWDRGFNQV |  |
| MMAGIGIVDV |  | IMSMSVSNYL |  |
| NMGGLGIVPV |  | AMGDGSYLLM |  |
| SMLGIKIVPV |  | SMWDRGMGLL |  |
| ELSGLGIQTV |  | MMENRGSGAL |  |
| SMLGIGILPM |  | LMWDSGLELM |  |
| SMAGIGILPV |  | SMWDRGLGMM |  |
| SMLGIGIVPV |  | LMWDVGWLNV |  |
|  |  | MMWDRGTFIM |  |
|  |  | MMWDRGIVPV |  |
|  |  | ILFDRGMNLM |  |

Table S1. DMF5 selection data and human target prediction.
The sequences identified from the round 3 deep-sequencing of the DMF5 10 mer 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 | ALDSRSELFL |  |
| IMTSHEVMM | GMDSRAEVFM | AMYSNSDFMV |  |
| VMESHDVFM | ALDSRSEYFL | VMDSRLEHFM |  |
| IMNSHEVMM | KMANRDENFV | SMNSRSEHFM |  |
| SMNSHEVMM | RLDGQDTKFM | SMNSKSENFL |  |
| KMNSHEVMM | LMDSRSEHFM | VLDSSSSSFL |  |
| AMQGHEYFL | IMNSRSELFL | ALDSRSENFL |  |
| AMQGHEIFL | MMNVRSELFV | ALDSKSENFL |  |
| VLQSHEVSM | TMNVRSELFV | ALDSRSEIFL |  |
| AMQSHEVTL | KMNSRSELFL | SMNSRADMFV |  |
| LMSGDYQFV | TMNVRSEHFM | SMYSRQEMMV |  |
| TMHNHEVMM | SMNSRSELFL | RMWSRSEDMV |  |
| VMHNHEVMM | KMNSRSEHFM | VLRARSDVFV |  |
| TMTGHEVFM | TMQSHDASFL | ALDSREEVFV |  |
| TMTGHEVFV | VMQGHDASFL | 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 |  |  |
|  | RMEAHSSHFV |  |  |

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 | $4^{\star 01: 01}$ | $4^{*} 01: 01$ |
|  | 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 | Vß | CDR3ß | Va | CDR3a |  |
| 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 B |  |  |  |  |  |  |
| Tumor | Healthy | Vß | CDR3ß | Va | CDR3a |  |
| 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 | CASSPSGLAGSNLGNEQFF | TRAV19 | CALSSRGSTLGRLYF |  |
| 11 | 0 | TRBV5-1 | CASSRIDSTDTQYF | TRAV4 | CLVGEVGTASKLTF |  |
| 10 | 0 | TRBV19 | CASSIPRGSSQPQHF | TRAV12-2 | CAVDSGGYNKLIF |  |
| 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.

|  |  | TCR 2A |  |  |  | TCR 3B |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Name | Peptide | Library | 2014PPM | 2017PWM | 2017DL | Library | 2014PPM | 2017PWM | 2017DL |
| 1s1 | LMDMHNGQ | 1 |  |  |  |  |  |  |  |
| 1s2 | RLDAMNGQ | 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 | VLDLFQGQ |  | 1 | 6 |  |  |  | 3 | 2 |
| 1sj2 | MMDFFNAQ |  | 2 | 1 |  |  |  | 2 | 160 |
| 1sj3 | LLNLNNGQ |  | 3 | 28 |  |  |  |  |  |
| 1sj4 | QMDYEEGQ |  | 4 | 5 |  |  |  | 1 | 100 |
| 1sj5 | GLSSQNGQ |  | 5 |  |  |  |  |  |  |
| 2sj1 | TLHYYEMH |  | 154 |  |  |  | 1 |  |  |
| DG1 | VIDFLNNQ |  |  | 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 | VVLGEQVQ |  |  |  | 5 |  |  |  |  |
| DG2 | HVDYKNVQ |  |  | 9 | 9 |  |  |  |  |
| 1sPWM4 | AISMFNGQ |  |  | 4 |  |  |  | 197 | 166 |
| DG3 | RIDPHDGQ |  |  | 8 |  |  |  | 94 | 11 |
| DG4 | SLDSYNAQ |  | 10 | 37 |  |  |  |  |  |
| 2sDL1 | TIHSYQGQ |  |  |  |  |  |  | 8 | 1 |
| 2sDL3 | SLDFYQGH |  |  |  |  |  |  | 7 | 3 |
| 2sDL4 | I IDAFKGQ |  |  |  |  |  |  |  | 4 |
| 2sDL6 | RIDDGQGQ |  |  |  |  |  |  | 365 | 6 |
| 2sDL7 | TILQYSGQ |  |  |  |  |  |  | 70 | 7 |
| 2sPWM4 | LVDFWEAQ |  |  | 40 |  |  |  | 4 |  |
| 2sPWM5 | TIAHYSGQ |  |  |  |  |  |  | 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 | AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGA |
| ILL_R | CAAGCAGAAGACGGCATACGAGATCGGTCTCGGCATTCCTGCTGAACCGCTCTTC |
| ILL_B2M_R | ATTCCTGCTGAACCGCTCTTCCGATCTNNNNNNNNCAGGATGTCTACTGTAAACTTG |
| ILL_BC_1 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTAGCATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_2 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTATGCTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_3 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTCACATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_4 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTCCAGTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_5 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTCCTCTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_6 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTGACATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_7 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTGCACTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_8 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTGCTGTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_9 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTGGGATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_10 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTGGGTTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_11 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTGGTGTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_12 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTTACCTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_13 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGGTAGTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_14 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGGTTTTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_15 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGTAACTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_16 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGTACTTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_17 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGTATCTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_18 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGTGACTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_19 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGTGATTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_20 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGTTCCTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_21 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGTTGATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_22 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTAACGTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_23 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTACACTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_24 | CTACACGACGCTCTTCCGATCTNNNNNNNNTTACCTTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_25 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGCAGTTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_26 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGCCTATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_27 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGCGTATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_28 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGCGTTTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_29 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGCTACTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_30 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGCTATTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_31 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGCTTCTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_32 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGGAAGTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_33 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGGATATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_34 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGGATGTTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_35 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGGCAATTAGCATGTTATTGCTAGCGTTTTGGCT |
| ILL_BC_36 | CTACACGACGCTCTTCCGATCTNNNNNNNNTGGCATTTAGCATGTTATTGCTAGCGTTTTGGCT |
| A2_8mer_S | TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNTGGGTGGAGGAGGTTCTGGAGGTGGTG |
| A2_9mer_S | TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNNKNTGGGTGGAGGAGGTTCTGGAGGTGGTG |
| A2_10mer_S | TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNNKNNKNTGGGTGGAGGAGGTTCTGGAGGTGGTG |
| A2_11mer_S | TTCTGTTATTGCTAGCGTTTTGGCTNNKMTGNNKNNKNNKNNKNNKNNKNNKNNKNTGGGTGGAGGAGGTTCTGGAGGTGGTG |
| A2_V5_AS | GAAGAGGATTAGGAATTGGTTTACCGGATCCAGAGGATGGTTCCCATCTCAATGTCAATG |
| A2_MYC_AS | ACCACCACCACCAGATCCACCACCACCCAAGTCTTCTTCGGAGATAAGCTTTTGTTC |
| A2_HA_AS | CGTAGTCTGGAACGTCGTATGGGTAGGATCCAGAGGATGGTTCCCATCTCAATGTCAATG |
| A2_VSV_AS | GTCTATTCATTTCAATATCAGTATAGGATCCAGAGGATGGTTCCCATCTCAATGTCAATG |
| 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.

