

dCODE Dextramer® (HiT) Package Insert

Products

dCODE Dextramer® (HiT) (Gold/Explore), Cat. No. WBxxxxxdHG /

UWBxxxxxHEP

CD1d dCODE Dextramer® (HiT) (Gold/Explore), Cat. No. XDxxxxxdHG /

YDxxxxxdHG

MR1 dCODE Dextramer® (HiT) (Gold/Explore), Cat. No. ZAxxxxxdHG

HLA-E dCODE Dextramer® (10x), Cat. No. URxxxxxdXC HLA-G dCODE Dextramer® (10x), Cat. No. USxxxxxdXG

Collectively denominated dCODE® (HiT) reagents.

Recommended use

Profiling and quantitation of antigen-specific T cells, NKT, MAIT, or NK cells in cell samples^{1,2,3}.

For research use only. Not for use in diagnostic or therapeutic procedures.

Materials Provided

dCODE® (HiT) reagents consisting of a dextran polymer backbone carrying multiple MHC, CD1d, MR1-, HLA-E, or HLA-G-antigen complexes, a corresponding unique DNA Barcode oligo, and R-phycoerythrin (PE) for sorting of dCODE® (HiT) positive cells.

- dCODE® (HiT) reagents are provided at a concentration of 160 nM in PBS buffer, containing 1% bovine serum albumin (BSA) and 15 mM NaN₃, pH 7.2.
- 2 μL (one test) is recommended for staining of 1-3 x 10⁶ PBMCs.
- Each dCODE® (HiT) reagent is uniquely identified by its allele / Peptide / DNA Barcode.

The unique DNA Barcode oligo comprises:

- Forward and reverse primer handle sequences for amplification of DNA Barcode
- Unique Molecule Identifier (UMI) sequence
- DNA Barcode sequence that specifies the MHC-peptide specificity

Sizes

dCODE® (HiT) - Gold: Single reagents of 25 tests (50 μ L), 50 tests (100 μ L) or 150 tests (300 μ L) each.

dCODE® (HiT) - Explore: Reagent Panels of 16, 32, 48, 64, 80, or 96 for 10 tests (20 μ L), 25 tests (50 μ L) or 50 tests (100 μ L) each.

Storage

dCODE $^{\otimes}$ (HiT) reagents should be stored at 2-8°C in the dark – the plastic vial only partially protects the reagents against light.

Precautions

Contains sodium azide (NaN₃), a chemical highly toxic in pure form. At product concentrations, though not classified as hazardous, sodium azide may react with lead and copper plumbing to form highly explosive buildups of metal azides. Upon disposal, flush with large volumes of water to prevent metal azide build-up in plumbing.

References

- Bentzen, A. et al., Large-scale detection of antigen-specific T cells using peptide-MHC-I multimers labeled with DNA barcodes. Nat. Biotechnol 2016. Doi: 10.1038/nbt.3662
- Bentzen, A. et al., Next-generation detection of antigen-responsive T cells using DNA barcode-labeled peptide-major histocompatibility complex I multimers. European Journal of Immunology, 2016. <u>Doi: 10.1002/eji.201670200</u>



3. Lyngaa, R. B et al., High-throughput discovery of T cell epitopes in type 1 diabetes using DNA barcode labelled peptide-MHC multimers. Eu Jour. of Immunol., 2016. DOI: 10.1038/nbt.3662

Patents

The dCODE® technology is disclosed in granted and pending patents within the WO 2015/185067 and WO 2015/188839 patent families including US11402373, US11585806, US11668705, EP3152232, EP3155426, HK1236546 B, AU2015271324, AU2019264685, AU2021204496, CA2951325, SG11201610177U, JP6956632 and JP7271465.

Symbols

See www.immudex.com/symbols for explanation of symbols.

Technical support

E-mail: customer@immudex.com

Telephone: +45 3110 9292 (Denmark)

Manufacturer

Immudex, Bredevej 2A, DK-2830 Virum, Denmark

Materials Required (Not Provided) Stain buffer: PBS, pH 7.4 cont. 1-5% serum and 0.1 g/L Herring sperm

DNA

Wash buffer: PBS, pH 7.4 cont. 1-5% serum Sorting buffer: PBS, pH 7.4 cont. 50% serum

100 μ M d-Biotin in PBS, pH 7.4

Antibodies identifying relevant cell surface markers (e.g. CD3, CD4, CD8) Forward and reverse amplification primers. For design see Appendix A

qPCR reaction components QIAquick® PCR Purification Kit

Procedure

- 1. Prepare PBMC sample and resuspend 1-3 x 10 6 PBMC in 50-100 μL stain buffer.
- 2. Centrifuge dCODE® (HiT) at 10,000 x g for 1 min.
- 3. Preparation of dCODE® (HiT) reagent pool:
 - a. Add 0.2 μ L 100 μ M d-Biotin per dCODE® (HiT) specificity into an empty tube
 - b. Add 2 µL of each dCODE® (HiT) specificity and mix
- 4. From this mixture, take 1 μ L and dilute with 99 μ L wash buffer. Save this for qPCR amplification as input sample (step 13)
- 5. Add the pool of dCODE® (HiT) reagents to the cell sample and mix thoroughly
- 6. Incubate in the dark at room temperature:
 - a. MHC I, CD1d, MR1, HLA-E^A, or HLA-G dCODE Dextramer[®] (HiT) pool: 10 min. incubation^B
 - b. MHC II dCODE Dextramer® (HiT): 30 min. incubation^B
 - c. dCODE® (HiT) pool comprised of a. and b.: 30 min. incubation^B
- 1. Add relevant antibodies in the volume/concentration recommended by provider. Incubate at room temperature in the dark for 20 min.
- 7. Washing:
 - a. If staining in 4 mL tubes, add 2 mL wash buffer. Centrifuge at 300 x g for 5 min. and remove the supernatant. Repeat washing with another 2 mL wash buffer



- b. If staining in 96-well microtiter plates, make 4 sequential washes using 200 μ L wash buffer per well. Centrifuge at 300 x g for 5 min. between each wash and remove supernatant
- 8. Resuspend cells in adequate volume of wash buffer
- 9. Proceed to FACS following the guidelines and practices of your sorting facility
- 10. Make a two-way sort by sorting the dCODE® (HiT)-PE-positive population and the dCODE® (HiT) PE-negative population separately
- 11. Collect sorted cells directly into tubes containing 500 μL sorting buffer.
- 12. Centrifuge the sorted cell samples at 300 x g for 10 min. Add 1 mL pure PBS and spin down for an additional 5 min. at 300 x g. Discard supernatant and resuspend cell pellet in 40 μ L PBS (Optional: Store at -20°C for up to 72h or proceed to the next step)
- 13. For each cell sample (dCODE® (HiT)-PE-positive and dCODE ® (HiT)-PE-negative), use 20 μL cell suspension to prepare a 40 μL final PCR reaction volume. For dCODE® (HiT) input mixture, use 1 μL of the diluted sample as template in a 40 μL total PCR reaction volume
- 14. Purify DNA from each PCR reaction (dCODE®-PE-positive, dCODE®-PE-negative, and input mixture) using QIAquick® PCR Purification Kit following the manufacturer's instructions. Elute each sample in 20 μL water
- 15. To prepare the final sample for sequencing, mix 10 μ L of each of the purified dCODE®-PE-positive and dCODE®-PE-negative PCR products. Add just 1 μ L PCR product from the input mixture. Remaining purified PCR products can be stored at -20°C as backup
- Send sample for sequencing.

Procedural Notes

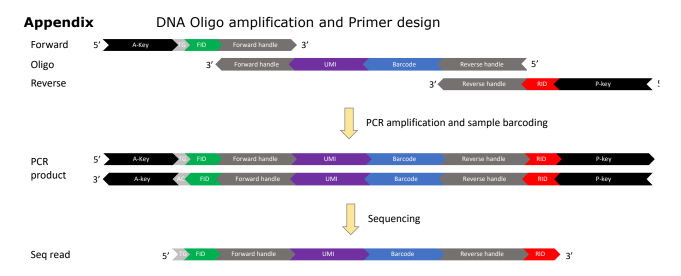
- A. HLA-E dCODE Dextramer® should be kept at 2-8°C or on ice during general handling of the reagent.
- B. Incubation time may be increased when using a high number of reagents in pool staining and requires optimization.

Data Analysis

Demultiplex the sequencing data according to sample (primer barcodes) and $dCODE^{\$}$ (HiT) reagents (oligo barcodes). Calculate the apparent enrichment (AE) for each reagent by dividing its read count in the $dCODE^{\$}$ (HiT)-PE-positive sample with the corresponding read count in the $dCODE^{\$}$ (HiT)-PE-negative sample. To get the specific enrichment, divide each AE with the (median of the) AE of the control $dCODE^{\$}$ (HiT) reagents.

Sequencing reads from the dCODE® (HiT) input mixture serve to validate the presence of all dCODE® (HiT) reagents.





Note: If the amplified product is sequenced using the A-key, the sequence read corresponds to the reverse complement of the oligo.

Amplification primers:

- Amplification primers are used in the PCR reaction to amplify the target region of the DNA barcode oligo.
- Each primer consists of a region for annealing to the oligo (forward or reverse handle), a unique sample ID sequence to individual label samples for multiplex sequencing, and a sequencing handle (A-key and P-key).
- The A-key and P-key must be matched to the sequencing platform (examples below are for Ion Torrent) and can be adapted as necessary.

P-key RID (8) Reverse handle Reverse 5'-CCTCTCTATGGGCAGTCGGTGAT + Sample ID + CTGTGACTATGTGAGGCTTTC-3'

Table 1: Examples of primer sequences for Ion Torrent sequencing (color-coding as above):

Forward Primer	Reverse Primer
CCATCTCATCCCTGCGTGTCTCCGACTCAG	CCTCTCTATGGGCAGTCGGTGATATTGCGCCC
TGCTGGGGTGGAAGTTCCAGCCAGCGTC	TGTGACTATGTGAGGCTTTC
CCATCTCATCCCTGCGTGTCTCCGACTCAG	CCTCTCTATGGGCAGTCGGTGATGACCCGTAC
TGCTCCACACGAAGTTCCAGCCAGCGTC	TGTGACTATGTGAGGCTTTC
CCATCTCATCCCTGCGTGTCTCCGACTCAG	CCTCTCTATGGGCAGTCGGTGATTGGCGTACC
TGCTTACCTGGAAGTTCCAGCCAGCGTC	TGTGACTATGTGAGGCTTTC
CCATCTCATCCCTGCGTGTCTCCGACTCAG	CCTCTCTATGGGCAGTCGGTGATTTATATGTC
TGTGGCAGCAGAAGTTCCAGCCAGCGTC	TGTGACTATGTGAGGCTTTC
CCATCTCATCCCTGCGTGTCTCCGACTCAG	CCTCTCTATGGGCAGTCGGTGATTGTTCTGCC
TGTGAGTAACGAAGTTCCAGCCAGCGTC	TGTGACTATGTGAGGCTTTC