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Determining antigen recognition through barcoding of mhc multimers

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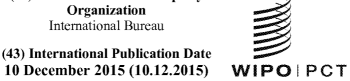
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(57) Abstract: The present invention describes the use of nucleic acid barcodes as specific labels for MHC multimers to determine the antigen responsiveness in biological samples. After cellular selection the barcode sequence will be revealed by sequencing. This technology allows for detection of multiple (potentially >1000) different antigen-specific cells in a single sample. The technology can be used for T-cell epitope mapping, immune-recognition discovery, diagnostics tests and measuring immune reactivity after vaccination or immune-related therapies.

DETERMINING ANTIGEN RECOGNITION THROUGH BARCODING OF MHC MULTIMERS

Technical field of the invention

5 The present invention relates to antigen recognition through nucleic acid labelled MHC multimers.

Background of the invention

The adaptive immune system is directed through specific interactions between immune cells and antigen-presenting cells (e.g. dendritic cells, B-cells, monocytes and macrophages) or target cells (e.g. virus infected cells, bacteria infected cells or cancer cells). In important field in immunology relates to the understanding of the molecular interaction between an immune cell and the target cell.

15 Specifically for T-lymphocytes (T-cells), this interaction is mediated through binding

- between the T-cell receptor (TCR) and the Major Histocompatibility Complex (MHC) class I or class II. The MHC molecules carries a peptide cargo, and this peptide in decisive for T-cell recognition. The understanding of T-cell recognition experienced a dramatic technological breakthrough when Atman et al. (1) in 1996 discovered that multimerization of single peptide-MHC molecules into tetramers would allow sufficient binding-strength (avidity) between the peptide-MHC molecules and the TCR to determine this interaction through a fluorescence label attached to the MHC-multimer. Such fluorescent-labelled MHC multimers (of both class I and class II molecules) are now widely used for determining the T-cell specificity. The MHC multimer associated fluorescence can be
- 25 determined by e.g. flow cytometry or microscopy, or T-cells can be selected based on this fluorescence label through e.g. flow cytometry or bead-based sorting. However, a limitation to this approach relates to the number of different fluorescence labels available, as each fluorescence label serve as a specific signature for the peptide-MHC in question.
- 30 Thus, this strategy is poorly matching the enormous diversity in T-cell recognition. For the most predominant subset of T-cells (the αβ TCR T-cells), the number of possible distinct αβ TCRs has been estimated at ~10¹⁵ (2) although the number of distinct TCRs in an individual human is probably closer to 10⁷ (3). Therefore, much effort has attempted to

expand the complexity of the T-cell determination, with the aim to enable detection of multiple different T-cell specificities in a single sample. A more recent invention relates to multiplex detection of antigen specific T-cells is the use of combinatorial encoded MHC multimers. This technique uses a combinatorial fluorescence labelling approach that allows for the detection of 28 different T-cell populations in a single sample when first published (4,5), but has later been extended through combination with novel instrumentation and heavy metal labels to allow detection of around 100 different T-cell populations in a single sample (6).

10 The requirement for new of technologies that allow a more comprehensive analysis of antigen-specific T-cell responses is underscored by the fact that several groups have tried to develop so-called MHC microarrays. In these systems, T-cell specificity is not encoded by fluorochromes, but is spatially encoded (7,8). In spite of their promise, MHC microarrays have not become widely adopted, and no documented examples for its value in the multiplexed measurement of T-cell responses, for instance epitope identification, are available.

Considering the above, there remains a need for a high-throughput method in the art of detection, isolation and/or identification of specific antigen responsive cells, such as antigen specific T-cells.

Further, there remains a need in the art, considering the often limited amounts of sample available, for methods allowing detection, isolation and/or identification of multiple species of specific antigen responsive cells, such as T-cells, in a single sample.

25

Summary of the invention

The present invention is the use of nucleic acid-barcodes for the determination and tracking of antigen specificity of immune cells.

30 In an aspect of the present invention a nucleic acid-barcode will serve as a specific label for a given peptide-MHC molecule that is multimerized to form a MHC multimer. The multimer can be composed of MHC class I, class II, CD1 or other MHC-like molecules. Thus, when the term MHC multimers is used below this includes all MHC-like molecules. The MHC multimer is formed through multimerization of peptide-MHC molecules via different backbones. The barcode will be co-attached to the multimer and serve as a

30

specific label for a particular peptide-MHC complex. In this way up to 1000 to 10.000 (or potentially even more) different peptide-MHC multimers can be mixed, allow specific interaction with T-cells from blood or other biological specimens, wash-out unbound MHC-multimers and determine the sequence of the DNA-barcodes. When selecting a cell population of interest, the sequence of barcodes present above background level, will provide a fingerprint for identification of the antigen responsive cells present in the given cell-population. The number of sequence-reads for each specific barcode will correlate with the frequency of specific T-cells, and the frequency can be estimated by comparing the frequency of reads to the input-frequency of T-cells. This strategy may expand our understanding of T-cell recognition.

The DNA-barcode serves as a specific labels for the antigen specific T-cells and can be used to determine the specificity of a T-cell after e.g. single-cell sorting, functional analyses or phenotypical assessments. In this way antigen specificity can be linked to both the T-cell receptor sequence (that can be revealed by single-cell sequencing methods) and functional and phenotypical characteristics of the antigen specific cells.

Furthermore, this strategy may allow for attachment of several different (sequence related) peptide-MHC multimers to a given T-cell – with the binding avidity of the given peptide-MHC multimer determining the relative contribution of each peptide-MHC multimer to the binding of cell-surface TCRs. By applying this feature it is possible to allow the determination of the fine-specificity/consensus recognition sequence of a given TCR by use of overlapping peptide libraries or alanine substitution peptide libraries. Such determination is not possible with current MHC multimer-based technologies.

- 25 Thus, one aspect of the invention relates to a multimeric major histocompatibility complex (MHC) comprising
 - two or more MHC's linked by a backbone molecule; and
 - At least one nucleic acid molecule linked to said backbone, wherein said nucleic acid molecule comprises a central stretch of nucleic acids (barcode region) designed to be amplified by e.g. PCR.

Another aspect of the present invention relates to a composition comprising a subset of multimeric major histocompatibility complexes (MHC's) according to the invention, wherein each set of MHC's has a different peptide decisive for T cell recognition and a unique "barcode" region in the DNA molecule.

Yet another aspect of the present invention is to provide a kit of parts comprising

- a composition according to the invention; and
- one or more sets of primers for amplifying the nucleic acid molecules.

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Still another aspect of the present invention is to provide a method for detecting antigen responsive cells in a sample comprising:

- providing one or more multimeric major histocompatibility complexes (MHC's) according to the invention or a composition according to the invention;
- 10 contacting said multimeric MHC's with said sample; and
 - detecting binding of the multimeric MHC's to said antigen responsive cells, thereby detecting cells responsive to an antigen present in a set of MHC's.

wherein said binding is detected by amplifying the barcode region of said nucleic acid molecule linked to the one or more MHC's.

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Further aspects relates to different uses.

Brief description of the figures

Fig. 1 describes the generation of barcode labelled MHC multimers.

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- Fig. 2 describes the generation of a library of barcode labelled MHC multimers.
- Fig. 3 describes the detection of antigen responsive cells in a single sample.
- 25 Fig. 4 describes the possibility of linking the antigen specificity (tracked by the barcode) to other properties.
 - Fig. 5 shows in a set of experimental data that the invention is experimentally feasible.
- 30 Fig. 6 onwards shows experimental data of examples 2 onwards.

The present invention will now be described in more detail in the following.

Detailed description of the invention

Definitions

Prior to discussing the present invention in further details, the following terms and conventions will first be defined:

5

TCR: T-cell receptor

MHC: Major Histocompatibility Complex

Multimeric MHC: Multimeric Major Histocompatibility Complex

10 Nucleic acid barcode

In the present context, a nucleic acid barcode is a unique oligo-nucleotide sequence ranging for 10 to more than 50 nucleotides. The barcode has shared amplification sequences in the 3' and 5' ends, and a unique sequence in the middle. This sequence can be revealed by sequencing and can serve as a specific barcode for a given molecule.

15

Sequencing

In the present aspect it is understood that sequencing also relates to e.g. deep-sequencing or next-generation sequencing, in which the amplified barcodes (the PCR product) is sequenced a large number of repetitive time (number of total reads, e.g. 100.000s of reads). The number of reads for the individual barcode sequence will relate to their quantitative presence in the amplification product, which again represents their quantitative presence before amplification, since all DNA-barcodes have similar amplification properties. Thus, the number of reads for a specific barcode sequences compared to the total number of reads will correlate to the presence of antigen responsive cells in the test-sample.

Referring now to the invention in more detail, Fig. 1 describes how peptide-MHC molecules, nucleic acid (DNA)-barcodes and (optional) fluorescent labels are assembled to form a library of MHC multimers each holding a DNA-barcode specific for the given peptide-MHC molecule involved. Fig 1A) the barcode is designed to have a unique sequences that can be determined through DNA sequencing. Also the barcode have shared amplification ends, enabling amplification of all DNA-barcodes simultaneously in a PCR reaction. DNA-barcodes are attached to the MHC-multimerization backbone (e.g. via a biotin linker binding to streptavidin on the multimer backbone). Fig 1B represents the multimer backbone. This may be any backbone that allow multimerization of macro-

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molecules. The backbone may (optionally) hold a fluorescence label (illustrated by the asterisk) to track the total pool of MHC multimer binding cells irrespectively of the peptide-MHC multimer specificity. Fig 1C represents the peptide-MHC molecule of interest, carrying a specific peptide cargo (horizontal line). Fig. 1D represents the assembled peptide-MHC multimers carrying the DNA barcode.

Multimeric major histocompatibility complex (MHC)

An aspect of the invention relates to a multimeric major histocompatibility complex (MHC) comprising

- 10 two or more MHC's linked by a backbone molecule; and
 - at least one nucleic acid molecule linked to said backbone, wherein said nucleic acid molecule comprises a central stretch of nucleic acids (barcode region) designed to be amplified by e.g. PCR.

Different types of backbones may be used. Thus, in an embodiment the backbone molecule is selected from the group consisting of polysaccharides, such as glucans such as dextran, a streptavidin or a streptavidin multimer. The skilled artisan may find other alternative backbones.

- The MHC's may be coupled to the backbone by different means. Thus, in an embodiment the MHC's are coupled to the backbone through a streptavidin-biotin binding or a streptavidin-avidin binding. Again other binding moieties may be used. The specific binding may use specific couplings points. In another embodiment the MHC's are linked to the backbone via the MHC heavy chain.
- The MHC consists of different elements, which may partly be expressed and purified from cell systems (such as the MHC heavy chain and the Beta-2-microglobulin element).

 Alternatively, the elements may be chemically synthesized. The specific peptide is preferably chemically synthesized.
- 30 All three elements are required for the generation of a stable MHC (complex). Thus, in an embodiment the MHC is artificially assembled.
 - The multimeric MHC may comprise different numbers of MHC's. Thus, in yet an embodiment the multimeric major histocompatibility complex (MHC) is composed of at

least four MHC's, such as at least eight, such as at least ten, 2-30, 2-20, such as 2-10 or such as 4-10 MHC's.

The nucleic acid component (preferably DNA) has a special structure. Thus, in an embodiment the at least one nucleic acid molecule is composed of at least a 5' first primer region, a central region (barcode region), and a 3' second primer region. In this way the central region (the barcode region) can be amplified by a primer set. The length of the nucleic acid molecule may also vary. Thus, in another embodiment the at least one nucleic acid molecule has a length in the range 20-100 nucleotides, such as 30-100, such as 30-80, such as 30-50 nucleotides. The coupling of the nucleic acid molecule to the backbone may also vary. Thus, in a further embodiment the at least one nucleic acid molecule is linked to said backbone via a streptavidin-biotin binding and/or streptavidin-avidin binding. Other coupling moieties may also be used.

- 15 In a further embodiment the at least one nucleic acid molecule comprises or consists of DNA, RNA, and/or artificial nucleotides such as PLA or LNA. Preferably DNA, but other nucleotides may be included to e.g. increase stability.
- Different types of MHC's may form part of the multimer. Thus, in an embodiment the MHC is selected from the group consisting of class I MHC, a class II MHC, a CD1, or a MHC-like molecule. For MHC class I the presenting peptide is a 9-11mer peptide; for MHC class II, the presenting peptide is 12-18mer peptides. For alternative MHC-molecules it may be fragments from lipids or gluco-molecules which are presented.
- 25 It may also be advantageously if it was possible to determine the complete pool of bound multimers when incubated with a sample (of cells). Thus, in a preferred embodiment, the backbone further comprises one or more linked fluorescent labels. By having such coupling better quantification can be made. Similar the labelling may be used for cell sorting.

30

Composition

Fig 2 illustrates the generation of a full barcode library. Fig 2A, this library is composed of multiple, potentially more than 1000 different peptide-MHC multimers, each with a specific DNA-barcode. Such that barcode#1 codes for peptide-MHC complex#1, barcode#2 codes for peptide-MHC complex#2, barcode#3 codes for peptide-MHC complex#3, and so on

until the possible mixture of thousands different specificities each with a specific barcode. Fig 2B represents the final reagent, which is a mixture of numerous different MHC-multimers each carrying a specific DNA barcode as a label for each peptide-MHC specificity.

5

As previously described a pool (library) of different sets of multimeric major histocompatibility complexes (MHC's) may be used to analyze an overall cell population for its specificity for peptides. Thus, another aspect of the invention relates to a composition comprising a subset of multimeric major histocompatibility complexes

10 (MHC's) according to the invention, wherein each set of MHC's has a different peptide, decisive for T cell recognition and a unique "barcode" region in the DNA molecule. In the present context, it is to be understood that each specific multimeric major histocompatibility complex is present in the composition with a certain number and that there is subset of different multimeric major histocompatibility complexes present in the composition.

Preferably all specific region for each multimeric MHC can be determined with only a few primer sets, preferably only one primer set. Thus, in an embodiment the primer regions in the DNA molecule are identical for each set of MHC's. In this way only one primer set is required. In an alternative embodiment, the multimeric MHC's are grouped by different primer sets, thereby allowing multiplication of different sets of the multimeric MHCs. In this way background noise may be limited, while also retrieving information of specific bindings. Thus, different primer set for different sets of MHC's may be used.

The number of individual sets of multimeric MHC's may vary. Thus, in an embodiment the composition comprises at least 10 different sets of multimeric MHC's such as at least 100, such as at least 500, at least 1000, at least 5000, such as in the range 10-50000, such as 10-1000 or such as 50-500 sets of MHC's.

30 Kit of parts

The composition of the invention may form part of a kit. Thus, yet an aspect of the invention relates to a kit of parts comprising

- a composition according to the invention; and
- one or more sets of primers for amplifying the DNA molecules.

Method for detecting antigen responsive cells in a sample

In Fig. 3 it is illustrated how this library can be used for staining of antigen responsive cells in a single sample. Fig 3A, cells in single cell suspension (may e.g., but not exclusive, originate from peripheral blood, tissue biopsies or other body fluids) are mixed with the 5 peptide-library represented in fig 2B. Fig 3B, after staining, cells are sequentially washed and spun to remove residual MHC multimers that are not bound to a cellular surface. Specific cell populations, e.g. T-cells (CD8 or CD4 restricted), other immune cells or specifically MHC multimer binding T-cells may be sorted by flow cytometry or others means of cell sorting/selection. Fig 3C, the DNA-barcode oligonucleotide sequences 10 isolated from the cell population is amplified by PCR. Fig 2D, this amplification product is sequenced by deep sequencing (providing 10-100.000s of reads). The sequencing will reveal the specific barcode sequence of DNA barcodes attached to cells in the specimen after selection, as these will appear more frequent than sequences associated to the background of non-specific attachment of MHC multimers. The "signal-to-noise" is 15 counteracted by the fact that any unspecific MHC multimer event will have a random association of 1/1000 different barcodes (dependent of the size of the library), making it even more sensitive than normal multimer staining.

Through analyses of barcode-sequence data, the antigen specificity of cells in the specimen can be determined. When DNA-barcode#1 is detected above background level of reads it means that peptide-MHC multimer#1 was preferentially bound to the selected cell type. Same goes for barcode no. 2, 3, 4, 5,etc. up to the potential combination of more than 1000 (nut not restricted to this particular number). When the number of input cells are known, e.g. when cell populations of interest is captured via a fluorescence signal also attached to the multimer by flow cytometry-based sorting or other means of capturing/sorting, the specific T-cell frequency can be calculated comparing the frequency of barcode-reads to the number of sorted T-cells.

Therefore, the multimeric MHC's and/or the compositions according to the invention may be used for different purposes. Thus, yet another aspect of the invention relates to a method for detecting antigen responsive cells in a sample comprising:

- providing one or more multimeric major histocompatibility complexes (MHC's) or a composition according to the invention;
- contacting said multimeric MHC's with said sample; and

- detecting binding of the multimeric MHC's to said antigen responsive cells, thereby detecting cells responsive to an antigen present in a set of MHC's.

wherein said binding is detected by amplifying the barcode region of said nucleic acid molecule linked to the one or more MHC's (through the backbone).

5

In an embodiment the method includes providing the (biological) sample.

As known to the skilled person, unbound molecules should preferably be removed. Thus, in an embodiment unbound (multimeric) MHC's are removed before amplification, e.g. by washing and/or spinning e.g. followed by removing of the supernatant.

The type of sample may also vary. In an embodiment the sample is a biological sample. In an embodiment the sample is a blood sample, such as an peripheral blood sample, a blood derived sample, a tissue biopsy or another body fluid, such as spinal fluid, or saliva.

15 The source of the sample may also vary. Thus, in a further embodiment said sample has been obtained from a mammal, such as a human, mouse, pigs, and/or horses.

It may also be advantageously to be able to sort the cells. Thus, in an embodiment the method further comprises cell sorting by e.g. flow cytometry such as FACS. This may e.g. be done if the backbone is equipped with a fluorescent marker. Thus, unbound cells may also be removed/sorted.

As also known to the skilled person, the measured values are preferably compared to a reference level. Thus, in an embodiment said binding detection includes comparing measured values to a reference level, e.g. a negative control and/or total level of response in the sample. In a further embodiment, said amplification is PCR such as QPCR.

As also previously mentioned the detection of the barcode includes sequencing of the amplified barcode regions. Thus, in an embodiment the detection of barcode regions includes sequencing of said barcode region, such as by deep sequencing or next generation sequencing.

specificities.

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Use of a multimeric major histocompatibility complex

In Fig 4, it is illustrated how this technology can be used to link different properties to the antigen specificity of a cell population. Fig 4A. illustrates how cells after binding to a barcode labeled MHC multimer library may be exposed to a certain stimuli. Cell

5 populations can be selected based on the functional response to this stimuli (e.g., but not exclusive, cytokine secretion, phosphorylation, calcium release or numerous other measures). After selecting the responsive or non-responsive population (following the steps of Fig 2), the DNA barcodes can be sequenced to decode the antigen responsiveness, and thereby determining the antigen-specificities involved in a given 10 response.

Fig 4B illustrates how cells can be selected based on phenotype, to link a certain set of phenotypic characteristics to the antigen-responsiveness.

Fig 4C represents the possibility for single-cell sorting of MHC-multimer binding cells based on the co-attached fluorescence label on the MHC multimer. Through single-cell 15 sorting the antigen-specificity of the given cell can be determined on a single cell level through sequencing of the associated barcode label. This can be linked to the TCR that can also be sequenced on a single cell level, as recently described (10). Hereby, this invention will provide a link between the TCR sequence, or other single-cell properties and the antigen specificity, and may through the use of barcode labeled MHC multimer 20 libraries enable definition of antigen-specific TCRs in a mixture of thousands different

Fig 4D illustrates the use of barcode labeled MHC multimer libraries for the quantitative assessment of MHC multimer binding to a given T-cell clone or TCR transduced/transfected cells. Since sequencing of the barcode label allow several

- 25 different labels to be determined simultaneously on the same cell population, this strategy can be used to determine the avidity of a given TCR relative to a library of related peptide-MHC multimers. The relative contribution of the different DNA-barcode sequences in the final readout is determined based on the quantitative contribution of the TCR binding for each of the different peptide-MHC multimers in the library. Via titration based analyses it
- 30 is possible to determine the quantitative binding properties of a TCR in relation to a large library of peptide-MHC multimers. All merged into a single sample. For this particular purpose the MHC multimer library may specifically hold related peptide sequences or alanine-substitution peptide libraries.

Fig 5 shows experimental data for the feasibility of attaching a DNA-barcode to a MHC 35 multimer and amplify the specific sequences following T-cell staining. Fig 5A shows the staining of cytomegalovirus (CMV) specific T-cells in a peripheral blood samples. The specific CMV-derived peptide-MHC multimers was labeled with a barcode (barcode#1) and mixed with an irrelevant/non-specific peptide-MHC multimer labeled with barcode (barcode#2) and mixed with 998 other non-barcode labeled non-specific MHC multimers.

Data here shows the feasibility for staining of CMV-specific T-cells in a mixture of 1000 other MHC multimers. Data is shown for three different staining protocols. Fig 5B shows the readout of the specific barcode sequences by quantative PCR. Barcode#1 (B#1) determining the CMV specific T-cell in detected for all three staining protocols, whereas the irrelevant/non-specific barcode signal, barcode#2 (B#2) is undetectable.

10

Overall, the multimeric MHC's or compositions comprising such sets of MHC's may find different uses. Thus, an aspect relates to the use of a multimeric major histocompatibility complex (MHC) or a composition according to the invention for the detecting of antigen responsive cells in a sample.

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Another aspect relates to the use of a multimeric major histocompatibility complex (MHC) or a composition according to the invention in the diagnosis of diseases or conditions, preferably cancer and/or infectious diseases.

20 A further aspect relates to the use of a multimeric major histocompatibility complex (MHC) or a composition according to the invention in the development of immune-therapeutics.

Yet a further aspect relates to the use of a multimeric major histocompatibility complex (MHC) or a composition according to the invention in the development of vaccines.

25

Another aspect relates to the use of a multimeric major histocompatibility complex (MHC) or a composition according to the invention for the identification of epitopes.

In sum, the advantages of the present invention include, without limitation, the possibility for detection of multiple (potentially, but exclusively, >1000) different antigen responsive cells in a single sample. The technology can be used, but is not restricted, for T-cell epitope mapping, immune-recognition discovery, diagnostics tests and measuring immune reactivity after vaccination or immune-related therapies.

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This level of complexity allow us to move from model antigens to determination of epitopespecific immune reactivity covering full organisms, viral genomes, cancer genomes, all vaccine components etc. It can be modified in a personalized fashion dependent of the individuals MHC expression and it can be used to follow immune related diseases, such 5 as diabetes, rheumatoid arthritis or similar.

Biological materials are for instance analyzed to monitor naturally occurring immune responses, such as those that can occur upon infection or cancer. In addition, biological materials are analyzed for the effect of immunotherapeutics including vaccines on 10 immune responses, Immunotherapeutics as used here is defined as active components in medical interventions that aim to enhance, suppress or modify immune responses, including vaccines, non-specific immune stimulants, immunosuppressives, cell-based immunotherapeutics and combinations thereof.

- 15 The invention can be used for, but is not restricted to, the development of diagnostic kits, where a fingerprint of immune response associated to the given disease can be determined in any biological specimen. Such diagnostic kits can be used to determining exposure to bacterial or viral infections or autoimmune diseases, e.g., but not exclusively related to tuberculosis, influenza and diabetes. Similar approach can be used for immune-20 therapeutics where immune-responsiveness may serve as a biomarker for therapeutic response. Analyses with a barcode labelled MHC multimer library allow for highthroughput assessment of large numbers of antigen responsive cells in a single sample.
- Furthermore, barcode labelled MHC-multimers can be used in combination with single-cell 25 sorting and TCR sequencing, where the specificity of the TCR can be determined by the co-attached barcode. This will enable us to identify TCR specificity for potentially 1000+ different antigen responsive T-cells in parallel from the same sample, and match the TCR sequence to the antigen specificity. The future potential of this technology relates to the ability to predict antigen responsiveness based on the TCR sequence. This would be 30 highly interesting as changes in TCR usage has been associated to immune therapy (11,12).

Further, there is a growing need for the identification of TCRs responsible for target-cell recognition (e.g., but not exclusive, in relation of cancer recognition). TCRs have been 35 successfully used in the treatment of cancer (13), and this line of clinical initiatives will be J 2015/165007 FC 1/DK2015/05015

further expanded in the future. The complexity of the barcode labeled MHC multimer libraries will allow for personalized selection of relevant TCRs in a given individual.

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Due to the barcode-sequence readout, the barcode labeled MHC multimer technology

3 allow for the interaction of several different peptide-MHC complexes on a single cell surface, while still maintaining a useful readout. When one T-cell binds multiple different peptide-MHC complexes in the library, there relative contribution to T-cell binding can be determined by the number of reads of the given sequences. Based on this feature it is possible to determine the fine-specificity/consensus sequences of a TCR. Each TCR can potential recognize large numbers of different peptide-MHC complexes, each with different affinity (14). The importance of such quantitative assessment has increased with clinical used of TCRs and lack of knowledge may have fatal consequences as recently exemplified in a clinical study where cross recognition of a sequence related peptide resulted in fatal heart failure in two cases (15,16). Thus, this particular feature for quantitative assessment of TCR binding of peptide-MHC molecules related to the present invention, can provide an efficient solution for pre-clinical testing of TCRs aimed for clinical use.

Also related to the above, this allows for determination of antigen responsiveness to
20 libraries of overlapping or to very similar peptides. Something that is not possible with
present multiplexing technologies, like the combinatorial encoding principle. This allows
for mapping of immune reactivity e.g. to mutation variant of viruses, such as, but nor
exclusive, HIV.

In broad embodiment, the present invention is the use of barcode labelled MHC multimers for high-throughput assessment of large numbers of antigen responsive cells in a single sample, the coupling of antigen responsiveness to functional and phenotypical characteristic, to TCR specificity and to determine the quantitative binding of large peptide-MHC libraries to a given TCR.

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While the foregoing written description of the invention enables one of ordinary skill to make and use what is considered presently to be the best mode thereof, those of ordinary skill will understand and appreciate the existence of variations, combinations, and equivalents of the specific embodiment, method, and examples herein. The invention should therefore not be limited by the above described embodiment, method, and

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examples, but by all embodiments and methods within the scope and spirit of the invention.

Additional items of the invention:

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- 5 Item 1: Use of barcode labelled MHC multimers for multiplex detection of different T-cell specificities in a single sample, enabling simultaneous detection of potentially more than 1000 different T-cell specificities where the specificity is revealed through sequencing of the barcode label.
- 10 Item 2: Use of barcode labelled MHC multimers in combination with single-cell sorting and TCR sequencing, where the specificity of the TCR can be determined by the coattached barcode. This will enable identification of TCRs specific for a mixture of numerous (potentially, but not restricted to >1000) different peptide-MHC multimers, and match the TCR sequence to the antigen specificity.
- Item 3: Use of barcode labelled MHC multimers for determining the affinity and binding motif of a given TCR. The barcode labelling strategy will allow for attachment of several different (sequence related) peptide-MHC multimers to a given T-cell with the binding affinity determining the relative contribution by each peptide-MHC multimer. Thereby it is possible to map the fine-specificity/consensus recognition sequence of a given TCR by

use of overlapping peptide libraries or e.g. alanine substitution libraries.

- *Item 4:* Use of barcode labelled MHC multimers to map antigen responsiveness against sequence related/similar peptides in the same libraries, e.g. mutational changes in HIV infection. This has not been possible with previous MHC multimer based techniques.
- Item 5: The use of barcode-labelled MHC multimers to couple any functional feature of a specific T-cell or pool of specific T-cells to the antigen (peptide-MHC) recognition. E.g. determine which T-cell specificities in a large pool secrete cytokines, releases Calcium or other functional measurement after a certain stimuli.

It should be noted that embodiments and features described in the context of one of the aspects of the present invention also apply to the other aspects of the invention.

All patent and non-patent references cited in the present application, are hereby incorporated by reference in their entirety.

The invention will now be described in further details in the following non-limiting examples and items.

ITEMS

- 1. A multimeric major histocompatibility complex (MHC) comprising
 - two or more MHC's linked by a backbone molecule; and
- at least one nucleic acid molecule linked to said backbone, wherein said nucleic acid molecule comprises a central stretch of nucleic acids (barcode region) designed to be amplified by e.g. PCR.
- The multimeric major histocompatibility complex according to item 1, wherein the backbone molecule is selected from the group consisting of polysaccharides, such as
 glucans such as dextran, a streptavidin or a streptavidin multimer.
 - 3. The multimeric major histocompatibility complex according to item 1 or 2, wherein the MHC's are coupled to the backbone through a streptavidin-biotin binding, streptavidin-avidin.

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- 4. The multimeric major histocompatibility complex according to any of the preceding items, wherein the MHC's are linked to the backbone via the MHC heavy chain.
- 5. The multimeric major histocompatibility complex (MHC) according to any of the preceding items, wherein the MHC is artificially assembled.
 - 6. The multimeric major histocompatibility complex (MHC) according to any of the preceding items, composed of at least four MHC's, such as at least eight, such as at least ten, 2-30, 2-20, such as 2-10 or such as 4-10 MHC's.

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7. The multimeric major histocompatibility complex (MHC) according to any of the preceding items, wherein the at least one nucleic acid molecule is composed of at least a 5' first primer region, a central region (barcode region), and a 3' second primer region.

- 8. The multimeric major histocompatibility complex (MHC) according to any of the preceding items, wherein the at least one nucleic acid molecule has a length in the range 20-100 nucleotides, such as 30-100, such as 30-80, such as 30-50 nucleotides.
- 5 9. The multimeric major histocompatibility complex (MHC) according to any of the preceding items, wherein the at least one nucleic acid molecule is linked to said backbone via a streptavidin-biotin binding and/or streptavidina-avidin binding.
- 10. The multimeric major histocompatibility complex (MHC) according to any of the
 preceding items, wherein the at least one nucleic acid molecule comprises or consists of DNA, RNA, and/or artificial nucleotides such as PLA or LNA.
- 11. The multimeric major histocompatibility complex (MHC) according to any of the preceding items, wherein the MHC is selected from the group consisting of class I MHC, a15 class II MHC, a CD1, or a MHC-like molecule.
 - 12. The multimeric major histocompatibility complex (MHC) according to any of the preceding items, wherein the backbone further comprises one or more linked fluorescent labels.

- 13. A composition comprising a subset of multimeric major histocompatibility complexes (MHC's) according to any of items 1-12, wherein each set of MHC's has a different peptide decisive for T cell recognition and a unique "barcode" region in the DNA molecule.
- 25 14. The composition according to item 13, wherein the primer regions in the DNA molecule are identical for each set of MHC's.
- 15. The composition according to item 13 or 14, comprising at least 10 different sets of MHC's such as at least 100, such as at least 500, at least 1000, at least 5000, such as in the range 10-50000, such as 10-1000 or such as 50-500 sets of MHC's.
 - 16. A kit of parts comprising
 - a composition according to any of items 13 to 15; and
 - one or more sets of primers for amplifying the nucleic acid molecules.

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- 17. A method for detecting antigen responsive cells in a sample comprising:
 - providing one or more multimeric major histocompatibility complexes (MHC's)
 according to any of items 1-12 or a composition according to any of items 13-15;
 - contacting said multimeric MHC's with said sample; and
- 5 detecting binding of the multimeric MHC's to said antigen responsive cells, thereby detecting cells responsive to an antigen present in a set of MHC's.

wherein said binding is detected by amplifying the barcode region of said nucleic acid molecule linked to the one or more MHC's.

- 10 18. The method according to item 17, wherein unbound MHC's are removed before amplification, e.g. by washing and/or spinning.
- 19. The method according to item 17 or 18, wherein the sample is a blood sample, such as an peripheral blood sample, a blood derived sample, a tissue biopsy or another body15 fluid, such as spinal fluid, or saliva.
 - 20. The method according to any of items 17-19, wherein said sample has been obtained from a mammal, such as a human, mouse, pigs, and/or horses.
- 20 21. The method according to any of item 17-20, wherein the method further comprises cell sorting by e.g. flow cytometry such as FACS.
- 22. The method according to any of items 17-21, wherein said binding detection includes comparing measured values to a reference level, e.g. a negative control and/or total level25 of response.
 - 23. The method according to any of item 17-22, wherein said amplification is PCR such as QPCR.
- 30 24. The method according to any of items 17-13, wherein the detection of barcode regions includes sequencing of said region such as deep sequencing or next generation sequencing.
- 25. Use of a multimeric major histocompatibility complex (MHC) according to any of items 1-12 or a composition according to any of items 13-16 for the

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detecting of antigen responsive cells in a sample.

26. Use of a multimeric major histocompatibility complex (MHC) according to any of items
1-12 or a composition according to any of items
13-16 in the diagnosis of diseases or
5 conditions, preferably cancer and/or infectious diseases.

27. Use of a multimeric major histocompatibility complex (MHC) according to any of items 1-12 or a composition according to any of items 13-16 in the development of immune-therapeutics.

- 28. Use of a multimeric major histocompatibility complex (MHC) according to any of items 1-12 or a composition according to any of items 13-16 in the development of vaccines.
- 29. Use of a multimeric major histocompatibility complex (MHC) according to any of items 1-12 or a composition according to any of items 13-16 for the identification of epitopes.

Examples

EXAMPLE 1

5 Figure 5 shows results that act as proof-of-principle for the claimed invention. Fig 5A, Flow cytometry data of peripheral blood mononuclear cells (PBMCs) from healthy donors.

Materials and methods

PBMCs were stained with CMV specific peptide-MHC multimers coupled to a specific 10 nucleotide-barcode. In addition to CMV peptide-MHC reagents the cells were stained in the presence of negative control reagents i.e. HIV-peptide MHC multimers coupled to another specific barcode label and the additional negative control peptide-MHC reagents (p*) not holding a barcode – all multimers were additionally labeled with a PEfluorescence label. The amounts of MHC multimers used for staining of PBMCs were 15 equivalent to the required amount for staining of 1000 different peptide-MHC specificities i.e. 1x oligo-labeled CMV specific MHC multimers, 1x oligo-labeled HIV specific MHC multimers and 998x non-labeled p*MHC multimers, so as to give an impression whether background staining will interfere with the true positive signal. Prolonged washing steps were included (either 0 min (A), 30 min (B) or 60 min (C)) after removing the MHC 20 multimers, and data from all experiments are shown. The PE-MHC-multimer positive cells were sorted by fluorescence activated cell sorting (FACS) Fig 5B, Cross threshold (Ct) values from multiplex qPCR of the sorted PE-MHC-multimer positive cells. QPCR was used to assess the feasibility of detecting certain cell specificity through barcode-labeled peptide-MHC-multimers. Reagents associated with a positive 25 control (CMV) barcode and a negative control (HIV) barcode were present during staining,

Examples of nucleic acid sequences are:

DNA-barcode oligo for CMV MHC multimer attachment:

but negative control (HIV) barcode-peptide-MHC multimers should be washed out.

DNA-barcode oligo for HIV MHC multimer attachment:

5GAGATACGTTGACCTCGTTGAANNNNNTCTATAGGTGTCTACCTCACTTAAGC TCTTGGTTGCAT

5= Biotin-TEG

Results

Results shows Ct value only detectable to the CMV peptide-MHC multimer associated barcode, whereas the HIV-peptide MHC multimer associated barcode was not detected

Conclusion

This experiment is a representative example of several similar experiment performed with other antigen specificities. Overall these data show that it is feasible to

- stain with 1000 different MHC-multimers in a single sample while still maintain a specific signal,
 - 2) attach a DNA-barcode to an MHC multimer,
 - 3) amplify the DNA-barcode after cellular selection steps,
 - 4) read the barcode with QPCR, using barcode specific probes.
- 5) obtain a specific signal corresponding to the antigen specific T cell population present in the sample, while non-specific MHC multimer barcodes are non-detectable.

Together these (and similar data available) provide proof of feasibility for the steps described in figure 1, 2, and 3.

EXAMPLE 2

This example relates to

- 25 i) the stability of DNA oligonucleotides, used in one embodiment of the invention, in blood preparations, and
 - ii) an embodiment of the invention, in which certain tagged Dextramers (detection molecules in which the binding molecule is a number of peptide-MHC complexes, and the label is a DNA oligonucleotide) are enriched for. Allowing identification of the Dextramers
- 30 with binding specificity for certain (subpopulations of) cells in the cell sample tested.

 In i) it is shown that DNA oligos are stable during handling in PBMC's and in blood for a time that will allow staining, washing and isolation of T cells and subsequent amplification

of DNA tags.

In ii) Show that a model system consisting of DNA-tagged Dextramers with MHC specificities for CMV. Flu and negative control peptide will locate to and can be

captured/sorted with relevant T cell specificities and can be identified by PCR amplification and/or sequencing.

A. Stability of single-stranded and double-stranded oligonucleotides in blood preparations

<u>DNA tag oligo design.</u> 69-nucleotide long, biotinylated TestOligo consisting of 5'primer region (22nt yellow)-random barcode region (6xN-nt)-kodon region (21nt green/underlined)-3'primer region (20nt blue) were prepared:

10 'b' = Biotin-TEG 5' modification

'h' = HEG (terminal modifications)

Forward-01 primer GAGATACGTTGACCTCGTTG

Reverse-01 primer ATGCAACCAAGAGCTTAAGT

TestOligo-01

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15 bGAGATACGTTGACCTCGTTGAANNNNNN<u>TCTATCCATTCCATCCAGCTC</u>ACTTAAGC TCTTGGTTGCAT

TestOligo-02

20 TestOligo-03

bhgagatacgttgacctcgttgaannnnnn<u>tctatccattccatccagctc</u>acttaag ctcttggttgcath

TestOligo-04

bhGAGATACGTTGACCTCGTTGAANNNNNTCTTGAACTATGAATCGTCTCACTTAAG

25 CTCTTGGTTGCATh

TestOligo-05

bhGAGATACGTTGACCTCGTTGAANNNNNN<u>TCTATAGGTGTCTACTACCTC</u>ACTTAAG CTCTTGGTTGCATh

TestOligo-06

30 bhGAGATACGTTGACCTCGTTGAANNNNNTCTTTATTGGAGAGCACGCTCACTTAAG CTCTTGGTTGCATh

Q-PCR probes for quantifying the amount of TestOligos 1-6:

+ = locked nucleic acid (LNA) modified RNA nucleotide

35 LNA-3

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The stability of oligo-tags by Q-PCR was analyzed under conditions relevant for T cell isolation:

The testOligos 1-6 were incubated in anticoagulated EDTA blood, and following incubation the amount of each of the testOligos was determined using Q-PCR using the abovementioned primers and probes. The oligo tags were quantified by QPCR with SYBR® Green JumpStart™ Taq ReadyMix™ according to manufacturer's protocol in combination with any capillary QPCR instruments (e.g. Roche LightCycler or Agilent Mx3005P).

Because of the different termini of the testOligos 1-6, this also was a test of the stability of non-modified DNA oligo tag vs HEG modified 5' and HEG modified 5' and 3' (TestOligo- 01, -02 and -03 respectively).

The results are shown in figure 6. It is concluded that the stability of the testOligos is appropriately high for all variants tested, to perform the invention.

30 B. Generation and screening of a 3 member DNA tagged MHC Dextramer library for screening of antigen specific T cells in a lymphoid cell sample.

This experiment involves the generation of 3 DNA- tagged Dextramers, each with a unique specificity, as follows:

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Dextramer 1: Flu (HLA-A*0201/GILGFVFTL/MP/Influenza)

Dextramer 2: CMV (HLA-A*0201/NLVPMVATV/pp65/CMV)

Dextramer 3: Negative (HLA-A*0201/ALIAPVHAV/Neg.Control).

5 Each of these Dextramers thus have a unique pMHC specificity (i.e. the three Dextramers have different binding molecules), and each Dextramer carries a unique label (DNA oligonucleotide) specific for that one pMHC specificity.

The library of DNA-tagged Dextramers are screened in a preparation of lymphoid cells 10 such as anticoagulated EDTA blood or preparations of peripheral blood mononucleated cells (PBMC's). Those Dextramers that bind to cells of the cell sample will be relatively more enriched than those that do not bind.

Finally, the MHC/antigen specificity of the enriched Dextramers is revealed by 15 identification of their DNA tags by Q-PCR with DNA tag-specific probes or by sequencing of the DNA tags.

- 1. Production of 3 different DNA tagged Dextramers with HLA-A*0201-peptide (pMHC) complexes.
 - a. pMHC complexes are generated and attached to dextran, along with unique DNA tags identifying each of the individual pMHC complexes, as follows.
 - i. Generation of DNA tagged Dextramers with Flu (HLA-A*0201/GILGFVFTL/MP/Influenza), CMV (HLA-A*0201/NLVPMVATV/pp65/CMV) and Negative (HLA-A*0201/ALIAPVHAV/Neg.Control).
 - 1. Dextramer stock is 160 nano molar (nM), TestOligo stock is diluted to 500 nM. Mix 10 micro liter (uL) 160 nM dextramer stock with 10 uL 500 nM TestOligo stock. Incubate 10 min at r.t. Mix with 1,5 ug pMHC complex of desired specificity. Adjust volume to 50 uL with a neutral pH buffer such as PBS or Tris pH 7,4, and store at 4 degrees Celsius. This will produce a DNA tagged Dextramer with approximately 3 oligo tags and 12 pMHC complexes, respectively, per Dextramer.

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- a. Dex-Oligo-03 = Dextramer with TestOligo-03 and HLA-A*0201/NLVPMVATV/pp65/CMV.
- b. Dex-Oligo-04 = Dextramer with TestOligo-04 and HLA-A*0201/GILGFVFTL/MP/Influenza.
- c. Dex-Oligo-05 = Dextramer with TestOligo-05 and HLA-A*0201/ALIAPVHAV/Neg.Control.

2. Preparation of cell sample for screening for antigen-specific T cells.

- a. Appropriate cell samples for identification of antigen specific T cells are preparations of lymphoid cells such as preparations of peripheral blood mononucleated cells (PBMC's) or anticoagulated blood. Such preparations of cell samples are prepared by standard techniques known by a person having ordinary skill in the art.
- b. Transfer in the range of 1E7 lymphoid cells (from PBMC or EDTA anticoagulated blood) to a 12 x 75 mm polystyrene test tube.
- c. Add 2 ml PBS containing 5% fetal calf serum, pH 7.4. Centrifuge at 300 x g for 5 min. Remove supernatant and resuspend cells in a total volume of 2,5 ml PBS containing 5% fetal calf serum, pH 7.4.

3. <u>Preparation and modification of library of DNA tagged Dextramers with three MHC/peptide specificities (from 1).</u>

- a. Mix 5 ul 10 uM biotin with 10 ul each of Dex-Oligo-03, Dex-Oligo-04 and Dex-Oligo-05.
- 4. Mixing of preparations of lymphoid cells with a library of DNA tagged MHC Dextramers.
 - a. Mix 1E7 lymphoid cells in 2,5 mL (from 2b) with 30 uL library of DNA tagged Dextramers (from 3a).
 - b. Incubate 30 min at r.t.
 - c. Centrifuge at 300 x g for 5 min. and remove the supernatant.
 - d. Resuspend pellet in 2,5 ml PBS containing 5% fetal calf serum, pH 7.4. Centrifuge at 300 x g for 5 min. and remove the supernatant.
 - e. Resuspend pellet in 2,5 ml PBS containing 5% fetal calf serum, pH 7.4

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- 5. Capture of all CD8+ antigen specific cells by magnet assisted cell sorting, performed according to Miltenyi Biotec catalog nr 130-090.878, Whole Blood CD8 MicroBead protocol.
 - a. Ad 100 uL Whole Blood C MicroBeads (Miltenyi Biotec catalog nr 130-090.878) to resuspended lymphoid cells from 4e. Mix and allow capture of CD8+ T cells for 15 min at r.t.
 - b. Place Whole Blood Column in the magnetic field of a suitable MACS Separator. For details see the Whole Blood Column Kit data sheet.
 - c. Prepare column by rinsing with 3 mL separation buffer (autoMACS Running Buffer or PBS containing 5% fetal calf serum, pH 7.4).
 - d. Apply magnetically labeled cell suspension (4e) onto the prepared Whole Blood Column. Collect flow-through containing unlabeled cells.
 - e. Wash Whole Blood Column with 3×3 mL separation buffer (autoMACS Running Buffer or PBS containing 5% fetal calf serum, pH 7.4).
 - f. Remove Whole Blood Column from the separator and place it on a new collection tube.
 - g. Capture CD8+ T cells by pipetting 5 mL Whole Blood Column Elution Buffer or PBS containing 5% fetal calf serum, pH 7.4 onto the Whole Blood Column. Immediately flush out the magnetically labeled cells by firmly pushing the plunger into the column.
 - h. Centrifuge at 300 x g for 5 min. and remove the supernatant. Resuspend the collected CD8+ cells in 50 uL and store at minus 20 degrees Celsius for subsequent analysis.
- 6. <u>Identification of Dextramers that bound significantly to antigen specific T cells of the lymphoid cell sample.</u>
 - a. Quantifying ratios of DNA oligo tags in input (3a) vs captured fraction (5h) by sequencing or alternatively quantifying by QPCR using the DNA tag specific probes LNA-3, LNA-4 and LNA-5 will reveal the relative abundance of antigen specific T cells in the lymphoid cell sample.
 - Quantifying ratios of DNA oligo tags in input (3a) vs captured fraction (5h) by QPCR using the DNA tag specific probes LNA-3, LNA-4 and LNA-5.
 - 1. Make 25 uL QPCR reactions of
 - a. input of library of DNA tagged Dextramers (3a)

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b. output of library of DNA tagged Dextramers (5h)

c. Standard curves of 10 to 1E8 TestOligo-03, TestOligo-04 and TestOligo-05 respectively.

- 2. Mix 12,5 uL JumpStart Tag ReadyMix (Sigma-Aldrich # D7440) with 0,125 uL 100 uM primer each of Forward-01 and Reverse-01, 0,625 ul 10 uM of either probe LNA-3, LNA-4 or LNA-5, 0,025 ul Reference dye (Sigma-Aldrich # R4526) and 12.5 uL of either input of library of DNA tagged Dextramers (3a), output of library of DNA tagged Dextramers (5h) or Standard curves of 10 to 1E8 TestOligo-03, TestOligo-04 and TestOligo-05 respectively.
- 3. Run two step QPCR thermal profile Cycle 1 = 5 min at 95 degrees Celsius, Cycle 2-40 = 30 sec at 95 degrees Celsius and 1 min at 60 degrees Celsius.
- 4. Estimate the relative abundance of T cells with antigen specificity against one of the three MHC Dextramers by plotting the QPC cycle time (Ct) values of the input of library of DNA tagged Dextramers (3a), the output of library of DNA tagged Dextramers (5h) in a plot of Ct values of the QPCR standard curve of TestOligo-03, TestOligo-04 and TestOligo-05 respectively.
- ii. Quantifying ratios of DNA oligo tags in input (3a) vs captured fraction (5h) by ultra-deep sequencing.
 - 1. Make 25 uL PCR reactions of
 - a. input of library of DNA tagged Dextramers (3a)
 - b. output of library of DNA tagged Dextramers (5h)
 - 2. Mix PCR reaction using any standard PCR master mix with 1,25 uL 10 uM primer each of Forward-01 and Reverse-01, and 12,5 uL of either input of library of DNA tagged Dextramers (3a) or output of library of DNA tagged Dextramers (5h). Top up to 25 uL with pure water. For example use 2x PCR Master Mix from Promega containing Taq DNA polymerase, dNTPs, MgCl2 and reaction buffers.
 - 3. Ultra Deep Sequencing of the above PCR product can be provided by a number of commercial suppliers such as for

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example Eurofins Genomics, GATC Biotech or Beckman Coulter Genomics using well established Next Generation Sequensing technologies such as Roche 454, Ion Torrent, the Illumina technology or any other high throughput sequencing technique for PCR amplicon sequencing.

4. PCR amplicon analysis of the relative abundances of the input of library of DNA tagged Dextramers (3a), the output of library of DNA tagged Dextramers (5h) will reveal the relative abundance of T cells with antigen specificity against one of the three MHC Dextramers.

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7. Predicted results and comments

a. It is expected that the relative abundance and ratios of DNA oligo tags in input of a library of DNA tagged Dextramers (3a) as estimated by QPCR or sequencing is primarily affected by three parameters namely i) the ratio in which the DNA oligo tags were supplied during the generation of the DNA tagged Dextramers (1.a.i.1), ii) how the library input was mixed (3a) and iii) how efficiently the individual DNA oligo tags are amplified in the PCR reactions.

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 In an example, the relative ratios of DNA oligo tags in input of a library of DNA tagged Dextramers as generated in 3a and as measured by QPCR or sequencing would be between 1 to 10 fold of each other.

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b. It is expected that the relative abundance and ratios of DNA oligo tags in the output of library of DNA tagged Dextramers (5h) as estimated by QPCR or sequencing, in addition to the three parameters mentioned in 7a, is primarily affected by three additional parameters namely i) the number of antigen specific T cells with specificity for one of the three MHC-peptide combinations ii) the affinity of the T cell receptor of the given T cell for the given MHC-peptide complex and finally iii) the efficiency of separating antigen-specific T cells and their associated DNA tagged MHC Dextramers from unbound DNA tagged MHC Dextramers by washing and cell capture.

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 i. In an example, the relative ratios of DNA oligo tags in output of a library of DNA tagged Dextramers as generated in 5h and as measured by QPCR or sequencing would be more than 10 fold in

favor of those DNA oligo tags coupled to an MHC Dextramer with an MHC-peptide complex for which antigen-specific T cells are present in the lymphoid cell sample.

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1. In a lymphoid cell sample from an influenza positive and CMV positive HLA-A0201 donor with antigen-specific T cells against HLA-A*0201/NLVPMVATV/pp65/CMV and HLA-A*0201/GILGFVFTL/MP/Influenza and no antigen-specific T cells against HLA-A*0201/ALIAPVHAV/Neg.Control it is expected that the relative ratios of TestOligo-03 (Dex-Oligo-03 = Dextramer with TestOligo-03 and HLA-A*0201/NLVPMVATV/pp65/CMV), TestOligo-04 (Dex-Oligo-04 = Dextramer with TestOligo-04 and HLA-A*0201/GILGFVFTL/MP/Influenza) and TestOligo-05 (Dex-Oligo-05 = Dextramer with TestOligo-05 and HLA-A*0201/ALIAPVHAV/Neg.Control) will be more than 10 fold in the favor of TestOligo-03 and TestOligo-04 over TestOligo-05. That is TestOligo-03 and TestOligo-04 is expected to be more than 10 fold more abundant or frequent than TestOligo-05 as measured by sequencing or QPCR of the output of library of DNA tagged Dextramers (5h) if they were supplied in equal amounts in the input of library of DNA tagged Dextramers (3a).

EXAMPLE 3

This is an example where the Sample was blood from one CMV positive and HIV negative donor which was modified to generate Peripheral blood mononuclear cells (PBMCs). The Backbone was a dextran conjugate with streptavidin and fluorochrome (Dextramer backbone from Immudex).

The MHC molecules were peptide-MHC (pMHC) complexes displaying either CMV

(positive antigen) or HIV (negative antigen) derived peptide-antigens. The MHC molecules were modified by biotinylation to provide a biotin capture-tag on the MHC molecule. The MHC molecule was purified by HPLC and quality controlled in terms of the formation of functional pMHC multimers for staining of a control T-cell population.

The oligonucleotide labels were synthetized by DNA Technology A/S (Denmark). The 35 label was synthetically modified with a terminal biotin capture-tag. The labels were

combined oligonucleotide label arising by annealing an A oligonucleotide (modified with biotin) to a partially complimentary B oligonucleotide label followed by enzymatic DNA polymerase extension of Oligo A and Oligo B to create a fully double stranded label.

- The MHC molecule was synthetized by attaching MHC molecules in the form of biotinylated pMHC and labels in the form of biotin-modified oligonucleotide onto a streptavidin-modified dextran backbone. The MHC molecule further contained a modification (5b) in the form of a fluorochrome. Two different MHC molecules were generated wherein the two individual MHC molecules containing different pMHC were encoded by corresponding individual oligonucleotide labels.
- 10 An amount of sample, PBMC's (1b) was incubated with an amount of mixed MHC molecules (5) under conditions (6c) that allowed binding of MHC molecules to T cells in the sample.

The cell-bound MHC molecules were separated from the non-cell bound MHC molecules (7) by first a few rounds of washing the PBMC's through centrifugation sedimentation of cells and resuspension in wash buffer followed by Fluorescence Activated Cell Sorting (FACS) of fluorochrome labeled cells. T cells that can efficiently bind MHC molecules will fluoresce because of the fluorochrome comprised within the MHC molecules; T cells that cannot bind MHC molecules will not fluoresce. FACS-sorting leads to enrichment of fluorescent cells, and hence, enrichment of the MHC molecules that bind T cells of the PBMC sample.

FACS isolated cells were subjected to quantitative PCR analysis of the oligonucleotide label associated with the MHC molecules bound to the isolated cells to reveal the identity of MHC molecules that bound to the T cells present in the sample.

This experiment thus reveal the presence of T cells in the blood expressing a T cell receptor that recognize/binds to peptide-MHC molecules comprised in the peptide-MHC multimeric library.

- Sample preparation. The cell sample used in this experiment was obtained by preparing PBMC's from blood drawn from a donor that was CMV positive as well as HIV negative as determined by conventional MHC-multimer staining.
 - a. Acquiring sample: Blood was obtained from the Danish Blood Bank
 - b. Modifying sample: Peripheral blood mononuclear cells (PBMCs) were isolated from whole blood by density gradient centrifugation. The density gradient medium, Lymphoprep (Axis-Shield), which consists of carbohydrate polymers and a dense iodine compound, facilitate separation of the individual constituents of blood. Blood samples were diluted 1:1 in RPMI (RPMI 1640, GlutaMAX, 25mM Hepes; gibco-Life technologies) and carefully layered onto the Lymphoprep. After centrifugation, 30 min, 490g,

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PBMCs together with platelets were harvested from the middle layer of cells. The isolated cells, the buffy coat (BC), was washed twice in RPMI and cryopreserved at -150 ℃ in fetal calf serum (FCS; gibco-Life technologies) containing 10% dimethyl sulfoxide (DMSO; Sigma-Aldrich). BC's used in this example are listed in table 6 together with their respective virus specificities. Their virus-specificities had been identified by conventional MHC multimer staining protocols.

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2. BackboneBackbone preparation: The backbonebackbone is a dextran molecule, to which has been attached streptavidin and fluorochromes. The streptavidin serves as attachment sites for biotinylated oligonucleotides and biotinylated pMHC complexesMHC molecule. The fluorochrome allows separation of cells bound to MHC molecules from cells not bound to MHC molecules.

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a. In this example backbones were linear and branched dextran molecules of 1000-2000 KDa with covalently attached streptavidin (5-10 per backbone) and fluorochromes (2-20 per backbone) in the form of PE. Backbones are essentially Dextramer backbone as described by Immudex. In this example the backbones are also named SA conjugate.

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3. MHC molecule preparation: The MHC molecules used in this example were two different class I MHC-peptide complexes. MHC heavy chains (HLA-A0201 and HLA-B0702) and B2M were expressed in E.coli as previously described (Hadrup et al. 2009) and each refolded with two peptide antigens. The individual specificities (peptide-MHC molecules, allele and peptide combination) were generated in the following way

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a. Synthesis: MHC molecules in this example was specific pMHC monomers that were produced from UV-exchange of selected HLA-I monomers carrying a UV-conditional 9-residue peptide-ligand (p*). When exposed to UV-light (366 nm) the conditional ligand will be cleaved and leave the binding groove empty. Due to the instability of empty MHC-I molecules, the complexes will quickly degrade if they are not rescued by replacement with another peptide that match that HLA-type. In this way specific pMHC monomers were produced by mixing excess of desired HLA ligands with p*MHC monomers. p*MHC monomers were refolded, biotinylated and purified as previously described (Hadrup et al. 2009)

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purified as previously described (Hadrup et al. 2009).

i. HIV derived peptide ILKEPVHGV from antigen HIV polymerase and

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CMV derived peptide TPRVTGGGAM from antigen pp65 TPR (Pepscan Presto, NL) were diluted in phosphate buffered saline (DPBS; Lonza) and mixed to final concentrations100μg/ml:200μM (HLA-A02: ILKEPVHGV and HLA-B07:TPRVTGGGAM). The mixtures were exposed to 366 nm UV light (UV cabinet; CAMAG) for one hour and optionally stored for up to 24 h at 4 °C.

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- b. Modification: No further modifications
- c. Purification: The panel of MHC molecules was moved to eppendorph tubes and centrifuged 5 min, 5000g, to sediment any MHC molecules not in solution, before being added to the cells.

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4. Label preparation: In this example, two different oligonucleotides, of the same length but partially different sequence, were generated. Each of the oligonuclotides became attached to a specific pMHC, and thus encoded this specific pMHC. The oligonucleotides were biotinylated, allowing easy attachment to the dextran-streptavidin conjugate backbone.

- a. Synthesis: labels were DNA oligonucleotides which were purchased from DNA Technology (Denmark) and delivered as lyophilized powder. Stock dilutions of 100 μM label were made in nuclease free water and stored at -20 °C.
 - i. The label used was named 2OS label system and was developed to increase the complexity of a limited number of oligonucleotide sequences by a combinatorial label-generation strategy to produce multiple unique labels from a more confined number of label precursors. The strategy, referred to as 2OS, involved annealing and subsequent elongation of two partially complimentary oligonucleotide-sequences (an A oligo and a B oligo) that fostered a new unique oligonucleotide-sequences that was applied as a DNA oligonucleotide label. E.g. by combining 22 unique oligonucleotide-sequences (A label precursor) that are all partly complementary to 55 other unique oligonucleotide sequences (B label precursor) a combinatorial library of 1,210 different (Ax+By) labels could be produced (e.g. with 100 in table 9).
 - 1. Partly complementary A and B oligonucleotides were annealed to produce two combined A+B oligonucleotide labels (A1+B1 to produce A1B1 and A2+B2 to produce A2B2). A and a B oligos were mixed as stated in table 3, heated to 65 ℃ for 2 min and cooled slowly to <35 ℃ in 15-30 min. The annealed A and B oligos were then elongated as stated in table 3. Components of the elongation reaction were mixed just before use. After mixing, the reaction was left 5 min at RT to allow elongation of the annealed oligonucleotides. The reagents used for annealing (left) and elongation (right) of partly complementary oligonucleotides is described in Table 3. Reagents marked in italic were from the the Sequenase Version 2.0 DNA Sequencing Kit (Affymetrix #70770).
- b. Modification: All labels were diluted to working concentrations (640nM) in nuclease free water with 0.1% Tween.
- c. Purification: No further purification of labels were performed.
- 5. MHC molecules preparation: The MHC molecules (pMHCs) and Labels (oligonucloetides) were attached to the backbone (backbone, dextranstreptavidine-fluorchrome conjugate), to form the MHC molecules, in a way so that a given pMHC is always attached to a given oligonucleotide.
 - a. Synthesis: For preparation of MHC molecules the Backbone was labeled with the Label in the form of a biotinylated AxBx oligo prior to addition of pMHC.
 - i. Creation of MHC molecules were performed by addition of label in two fold excess over backbone (2:1 label:backbone) and incubated at least 30 min, 4℃. Optionally the backbone were stored for up to 24 h at 4℃ after coupling of the label. Prior to coupling MHC molecules, pMHC monomers, these were centrifuged 5 min, 3300g. SA conjugate (Dextramer backbone, Immudex) with conjugated streptavidin (SA) and fluorochrome (PE) were aliquoted into plates according to table 1. Avoiding the precipitate, MHC molecules were

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added to the aliquoted SA conjugate and incubated 30 min, RT. Following complex formation, D-biotin (Avidity Bio200) was added together with 0.02% NaN2 in PBS to the final concentration of pMHC monomer listed in table 1, and incubated at least 30 min or up to 24h at 4°C. Assembled MHC molecules were stored up to four weeks at 4 ℃. Two sets of two MHC molecules were generated. Each set with the two specificities individually labeled. The label was inverted between the two sets as described below.

- 1. 1xCMV specific pMHCs coupled to 2OS-A1B1, 1xHIV specific pMHCs coupled to 2OS-A2B2
- 2. 1xCMV specific pMHCs coupled to 2OS-A2B2, 1xHIV specific pMHCs coupled to 2OS-A1B1.
- b. Modification: No further modifications were performed
- c. Purification: MHC molecules were centrifuged 5 min, 3300g, to sediment any MHC molecules not in solution, before being added to the sample.
- 6. Incubation of sample and MHC molecules: The cell sample and the MHC molecules were mixed in one container, to allow the MHC molecules to bind the T cells that they recognize.
 - a. Amount of sample: 1x10E6-2x10E6 cells in the form of BC's, were used.
 - b. Amount of MHC molecule: According to table 1. 1 ug/ml calculated in relation to each MHC molecule (peptide-MHC molecule) was required per incubation
 - c. Conditions: BCs were thawed in 10 ml, 37 ℃, RPMI with 10% fetal bovine serum (FBS), centrifuged 5 min, 490g, and washed twice in 10 ml RPMI with 10% FBS. All subsequent washing of cells refer to centrifugation 5 min, 490 g, with subsequent removal of supernatant. 2x10E6 cells were washed in 200ul barcode-buffer (PBS/0.5% BSA/2 mM EDTA/100 µg/ml herring DNA) and resuspended in this buffer to approximately 20µl per staining. Prior to incubation of cells with MHC molecules cells were incubated with 50 nM dasatinib, 30 min, 37 °C (Lissina et al. 2009). MHC molecules were centrifuged for 5 min, 3300g, prior to addition to cells. 1 ug/ml 5 each MHC molecule (per pMHC) was required per incubation. After adding MHC molecules, the cells were incubated 15 min, 37 ℃. The antibody mixture listed in table 2 were added together with 0.1 µl near- IRviability dye (Invitrogen L10119) that stains free amines. Antibody staining was essentially as for conventional MHC multimer staining. Cells were incubated 30 min, 4℃. Cells were then washed twice in 200 ul barcode buffer and incubated in

200 ul 1% paraformaldehyde in phosphate buffered saline (DPBS; Lonza) over night at 4℃.

- 7. Enrichment of MHC molecules with desired characteristics: In this Example, the MHC molecules were enriched by using flow cytometry, more specifically, Fluorescence-Activated-Cell-Sorting (FACS). The MHC molecules carry a fluorochrome. Hence, the cells that bind MHC molecules will fluoresce, and can, by applying a FACS sorter, be separated from the cells that do not bind MHC molecules and therefore do not fluoresce. As a result, the MHC molecules that bound to cells will be enriched for.
 - a. Apply: Cells were sorted on a BD FACSAria, equipped with three lasers (488 nm blue, 633nm red and 405 violet). The flow cytometry data

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analyses was performed using the BD FACSDiva software version 6.1.2. The following gating strategy was applied. Lymphocytes were identified in a FSC/SSC plot. Additional gating on single cells (FSC-A/FSC-H), live cells (near-IR-viability dye negative), and CD4, CD14, CD16, CD19, CD40 negative (FITC)/CD8 positive cells (PerCP) were used to define the CD8 T cell population (table 2). The cells that were bound to at MHC molecule were defined within the PerCP positive population

b. Wash: Cells were washed twice in barcode-buffer where after the cells were ready for flow cytometric acquisition. Optionally cells were fixed in 1% paraformaldehyde O.N., 4°C, and washed twice in barcode-buffer. Fixed cells were stored for up to a week at 4°C.

- c. Separate: Optionally cells were acquired up to one week after fixation in 1% paraformaldehyde. The multimer positive cells were sorted by FACS, as described in 7a, into tubes that had been pre-saturated for 2h-O.N. in 2% BSA and contained 200 µl barcode-buffer to increase the stability of the oligonucleotides that followed with the sorted cells. The sorted fluorochrome (PE) positive cells were centrifuged 5 min, 5000 g, to allow removal of all excess buffer. Cells were stored at -80 °C.
- 8. Identification of enriched MHC molecules: By identifying the Label (in this Example, the oligonucleotide label), the pMHCs that bound cells could be identified. Therefore, the oligonucleotides that were comprised within the MHC molecules that were recovered with the cells, were analyzed by quantitative PCR using Label-specific Q-PCR probes. This allowed the identification of pMHCs that bound cells of the cell sample.
 - a. Labels derived from sorted cells were analyzed by QPCR according to table 4 QPCR was performed with the kit: Brilliant II QRT_PCR Low ROX Master Mix Kit (Agilent technologies, #600837). The thermal profile is listed in table 5. PCR was run on the thermal cycler: Mx3000P qPCR system (Agilent Technologies).

Results and conclusions on Example 3

After sorting and qPCR the resultant Ct values confirmed that Labels were successfully recovered and enriched only when associated with the CMV epitope, while they were not detected when associated with the HIV epitope (figure 7).

35 Thus, it was verified that the 2OS labels were recovered after cellular interaction, sorting and qPCR only T cell recognizing the given pMHC molecule were present in the sample.

Figure 7:

Detection of a B7 CMV pp65 TPR specificity amongst negative control barcoded pMHC dextramers. A unique 2OS barcode was associated with the positive control reagents in 1., while another unique 2OS barcode was associated with the positive control reagents in 2. The spare barcode in each experiment was associated with the HIV negative control reagent. A, Representative dot plot showing the PE positive population after staining with

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the CMV and HIV pMHC multimers carrying separate 2OS-barcodes. **B**, Ct values from multiplex qPCR of the sorted PE-pMHC-dextramer positive cells. Cells were stained with 1. and 2. respectively. Reagents associated with a positive control (CMV) 2OS barcode and a negative control (HIV) 2OS barcode were present during staining, but the negative control (HIV) barcoded pMHC dextramer was evidently washed out. The results obtained from two individual experiments are presented in separate bars. Approximately 200 cells were applied in each separate PCR. QPCR was run in duplicates and Ct values are shown as mean ±range of duplicates.

10 Example 4

This is an example where the Sample (1) was blood from one CMV positive and HIV negative donor which was modified (1b) to generate Peripheral blood mononuclear cells (PBMCs).

The Backbone (2) was a dextran conjugate with streptavidin and fluorochrome (Dextramer backbone from Immudex).

- The example is similar to example 1 except that a 1000 fold excess of MHC molecules with irrelevant MHC molecules but without label were included. The MHC molecules used (3) are peptide-MHC (pMHC) complexes displaying either CMV (positive antigen) or HIV (negative antigen) derived peptide-antigens or pMHC complexes displaying irrelevant
- 20 peptide antigen. The MHC molecules were modified (3b) by biotinylation to provide a biotin capture-tag on the MHC molecule. The MHC molecules were purified (2c) by HPLC. The Labels (4) were oligonucleotides. The oligonucleotides were synthetized (4a) by DNA Technology A/S (Denmark). The labels were synthetically modified (4b) with a terminal biotin capture-tag.
- 25 The MHC molecule (5) was synthetized (5a) by attaching MHC molecules in the form of biotinylated pMHC and labels in the form of biotin-modified oligonucleotide onto a streptavidin-modified dextran backbone. The MHC molecule further contained a modification (5b) in the form of a fluorochrome. Three different MHC molecules were generated wherein the two of these individual MHC molecules containing CMV- and HIV-
- directed pMHC were encoded for by corresponding individual oligonucleotide labels. MHC molecules with irrelevant MHC molecules were not encoded for with oligonucleotide label. An amount of sample, PBMC's (1b) was incubated with an amount of mixed MHC molecules (5) in a ratio of 1:1 and in addition a 1000 fold of unlabeled p*MHC labeled backbone was included under conditions (6c) that allowed binding of MHC molecules to T
- 35 cells in the sample.

The cell-bound MHC molecules were separated from the non-cell bound MHC molecules (7) by first a few rounds of washing the PBMC's through centrifugation sedimentation of cells and resuspension in wash buffer followed by Fluorescence Activated Cell Sorting (FACS) of fluorochrome labeled cells. T cells that can efficiently bind MHC molecules will.

- 5 fluoresce because of the fluorochrome comprised within the MHC molecules; T cells that cannot bind MHC molecules will not fluoresce. FACS-sorting leads to enrichment of fluorescent cells, and hence, enrichment of the MHC molecules that bind T cells of the PBMC sample.
- FACS isolated cells were subjected to quantitative PCR analysis of the oligonucleotide 10 label associated with the MHC molecules bound to the isolated cells to reveal the identity of MHC molecules that bound to the T cells present in the sample.

This experiment thus revealed the peptide-MHC specificity of the T cell receptors of the T cells present in the blood sample. It further revealed the feasibility of enriching for T cells specific for the CMV-antigen (positive) over the HIV-antigen (negative) and an excess of

- 15 MHC molecule displaying irrelevant peptide antigens.
 - 1. Sample preparation. The cell sample used in this experiment was obtained by preparing PBMC's from blood drawn from a donor that was CMV positive as well as HIV negative as determined by conventional MHC-multimer staining.
 - a. Acquiring sample: Blood was obtained from the Danish Blood Bank
 - b. Modifying sample: Peripheral blood mononuclear cells (PBMCs) were isolated from whole blood by density gradient centrifugation. The density gradient medium, Lymphoprep (Axis-Shield), which consists of carbohydrate polymers and a dense iodine compound, facilitate separation of the individual constituents of blood. Blood samples were diluted 1:1 in RPMI (RPMI 1640, GlutaMAX, 25mM Hepes; gibco-Life technologies) and carefully layered onto the Lymphoprep. After centrifugation, 30 min, 490g, PBMCs together with platelets were harvested from the middle layer of cells. The isolated cells, the buffy coat (BC), was washed twice in RPMI and cryopreserved at -150 ℃ in fetal calf serum (FCS; gibco-Life technologies) containing 10% dimethyl sulfoxide (DMSO; Sigma-Aldrich). BC's used in this example are listed in table 6 together with their respective virus specificities. Their virus-specificities had been identified by conventional MHC multimer staining protocols.
- 2. Backbone preparation: The backbone used in this example is a dextran molecule. 35 to which has been attached streptavidin and fluorochromes. The streptavidin serves as attachment sites for biotinylated oligonucleotides (Label) and biotinylated pMHC complexes (MHC molecules). The fluorochrome allows separation of cells bound to MHC molecules and cells not bound to MHC molecules. 40
 - a. In this example backbones were linear and branched dextran molecules of 1000-2000 KDa with covalently attached streptavidin (5-10 per backbone) and fluorochromes (2-20 per backbone) in the form of PE. Backbones are

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essentially Dextramer backbone as described by Immudex. In this example the backbones are also named SA conjugate.

- 3. MHC molecules preparation: The MHC molecules used in this example were two different class I MHC-peptide complexes. MHC heavy chains (HLA-A02 and HLA-B07) and B2M were expressed in E.coli as previously described (Hadrup et al. 2009) and each refolded with two peptide antigens. The individual specificities (allele and epitope combination) were generated in the following way.
 - a. Synthesis: A in experiment 1.
 - i. As in experiment 1
 - b. Modification: No further modifications
 - c. Purification: as in experiment 1
- 4. Label preparation: In this experiment, two different oligonucleotides, of the same length but partially different sequence, were generated. Each of the oligonuclotides become attached to a specific pMHC, and thus encodes this specific pMHC. The oligonucleotides were biotinylated, allowing easy attachment to the dextran-streptavidine conjugate backbone.
 - a. Synthesis: In this example the labels were DNA oligonucleotides which were purchased from DNA Technology (Denmark) and delivered as lyophilized powder. Stock dilutions of 100 μM label were made in nuclease free water and stored at -20 °C.
 - i. As in experiment 1
 - ii. Partly complementary A and B oligonucleotides were annealed to produce two combined A+B oligonucleotide labels (A1+B1 to produce A1B1 and A2+B2 to produce A2B2). A and a B oligos were mixed as stated in table 3, heated to 65 ℃ for 2 min and cooled slowly to <35 ℃ in 15-30 min. The annealed A and B oligos were then elongated as stated in table 3. Components of the elongation reaction were mixed just before use. After mixing, the reaction was left 5 min at RT to allow elongation of the annealed oligonucleotides. The reagents used for annealing (left) and elongation (right) of partly complementary oligonucleotides is described in Table 3. Reagents marked in italic were from the the Sequenase Version 2.0 DNA Sequencing Kit (Affymetrix #70770).
 - b. Modification: All labels were diluted to working concentrations (640nM) in nuclease free water with 0.1% Tween.
 - c. Purification: No further purification of labels were performed.
- 5. MHC molecules preparation: The MHC molecules (pMHCs) and Labels (oligonucloetides) were attached to the backbone (dextran-streptavidine-fluorchrome conjugate), to form the MHC molecules, in a way so that a given pMHC is always attached to a given oligonucleotide.
 - Synthesis: For preparation of MHC molecules the Backbone was labeled with the Label in the form of a biotinylated AxBx oligo prior to addition of pMHC.
 - i. Creation of MHC molecules were performed by addition of label in two fold excess over backbone (2:1 label:backbone) and incubated 30 min, 4℃. Prior to coupling MHC molecules, pMHC monomers, these were centrifuged 5 min, 3300g. SA conjugate (Dextramer backbone, Immudex) with conjugated streptavidin (SA) and fluorochrome (PE) were aliquoted into tubes according to table 1.

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Avoiding the precipitate, MHC molecules were added to the aliquoted SA conjugate and incubated 30 min, RT. Following complex formation, D-biotin (Avidity Bio200) was added together with 0.02% NaN2 in PBS to the final concentration of pMHC monomer listed in table 1, and incubated 30 min, 4 ℃. Assembled MHC molecules were stored up to four weeks at 4°C. Two sets of two MHC molecules were generated. Each set with the two specificities individually labeled. The label was inverted between the two sets as described below.

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- 1. iv. 1xCMV specific pMHCs coupled to 2OS-A1B1, 1xHIV specific pMHCs coupled to 2OS-A2B2
- 2. v. 1xCMV specific pMHCs coupled to 2OS-A2B2, 1xHIV specific pMHCs coupled to 2OS-A1B1.
- b. Modification: No further modifications were performed

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- c. Purification: MHC molecules were centrifuged 5 min, 5000g, to sediment any MHC molecules not in solution, before being added to the sample.
- 6. Incubation of sample and MHC molecules: The cell sample and the MHC molecules were mixed in one container, to allow the MHC molecules to bind the T cells that they recognize.
 - a. Amount of sample: 1x10E6-2x10E6 cells in the form of BC's, were used.
 - b. Amount of MHC molecule: According to table 1. 5 ul of each MHC molecule was required per incubation (1 ug/ml in respect to pMHC)
 - Conditions: BCs were thawed in 10 ml, 37 °C, RPMI with 10% fetal bovine serum (FBS), centrifuged 5 min, 1500g, and washed twice in 10 ml RPMI with 10% FBS. All subsequent washing of cells refer to centrifugation 5 min, 490 g, with subsequent removal of supernatant. 1x10E6-2x10E6 cells were washed in barcode-buffer (PBS/0.5% BSA/2 mM EDTA/100 µg/ml herring DNA) and resuspended in this buffer to approximately 20µl per staining. Prior to incubation of cells with MHC molecules cells were incubated with 50nM dasatinib, 30 min, 37 °C. MHC molecules were centrifuged for 5 min, 3300g, prior to addition to cells. 5 ul of each MHC molecule was required per incubation (1 ug/ml in respect to pMHC). After adding MHC molecules, the cells were incubated 15 min, 37 ℃. The antibody mixture listed in table 2 were added together with 0.1 µl near- IRviability dye (Invitrogen L10119) that stains free amines. Antibody staining was essentially as for conventional MHC multimer staining. Cells were incubated 30 min, 4℃.

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7. Enrichment of MHC molecules with desired characteristics: In this Example, the MHC molecules were enriched by using flow cytometry, more specifically, Fluorescence-Activated-Cell-Sorter (FACS). The MHC molecules carry a fluorochrome. Hence, the cells that bind MHC molecules will fluoresce, and can be separated from the cells that do not bind MHC molecules and therefore do not fluoresce, by a FACS sorter. As a result, the MHC molecules that bound to cells will be enriched for.

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a. Apply: Two different flow cytometers were used for acquisition. A BD FACSCanto II equipped with three lasers (488 nm blue, 633nm red and 405 violet) and a BD LSR II cytometer equipped with five lasers. Only four lasers on the LSR II were used throughout this study (488 nm blue laser. 640 nm red laser, 355 nm UV laser and 405 nm violet laser). Additionally

cells were sorted on BD FACSAria and FACSAria II, equipped with three lasers (488 nm blue, 633nm red and 405 violet). All flow cytometry data analyses used the BD FACSDiva software version 6.1.2. The following gating strategy was used. All initial gatings of CD8 positive cells were performed alike. Lymphocytes were identified in a FSC/SSC plot. Additional gating on single cells (FSC-A/FSC-H), live cells (near-IR-viability dye negative), and dump-channel negative/CD8 positive cells (FITC/PerCP) were used to define the CD8 T cell population.

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b. Wash: Cells were washed twice in barcode-buffer where after the cells were ready for flow cytometric acquisition. Optionally cells were fixed in 1% paraformaldehyde O.N., 4°C, and washed twice in FACS buffer or barcode-buffer. Fixed cells were stored for up to a week at 4°C.

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- c. Separate: Optionally cells were acquired up to one week after fixation in 1% paraformaldehyde. The multimer positive cells were sorted by FACS, as described in 7a, into tubes that had been pre-saturated for 2h-O.N. in 2% BSA and contained 200 µl barcode-buffer to increase the stability of the oligonucleotides that followed with the sorted cells. The sorted fluorochrome (PE) positive cells were centrifuged 5 min, 5000 g, to allow removal of all excess buffer. Cells were stored at -80 ℃.
- 8. Identification of enriched MHC molecules: By identifying the Label (in this Example, the oligonucleotide label), the pMHCs that bound cells can be identified. Therefore, the oligonucleotides that were comprised within the MHC molecules that were recovered with the cells, were analyzed by quantitative PCR using Label-specific Q-PCR probes. This allowed the identification of pMHCs that bound cells of the cell sample.
 - a. Labels derived from sorted cells were analyzed by QPCR as in experiment

Results and conclusions on Example 4

30 After sorting and qPCR the resultant Ct values confirmed that Labels were successfully recovered and enriched for only when associated with the CMV epitope, while they were not detected when associated with the HIV epitope (figure 8).

It was verified that the 2OS labels were recovered after cellular interaction, sorting and qPCR only if they were associated with positive control reagents.

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Figure 8.

Detection of a CMV specificity amongst negative control barcoded pMHC dextramers. A unique barcode is associated with the positive control reagents in 1., while another unique barcode is associated with the positive control reagents in 2. The spare barcode in each

40 experiment is associated with the HIV negative control reagent. In addition 998x unlabeled negative control reagents are present in both 1. and 2.

- A, Ct values from multiplex qPCR of the sorted PE-pMHC-dextramer positive cells. Cells were stained with 1. and 2. respectively. Reagents associated with a positive control (CMV) barcode and a negative control (HIV) barcode were present during staining, but the negative control (HIV) barcoded pMHC dextramer was evidently washed out.
- 5 Approximatly 575 cells were analyzed in each separate qPCR. **B.** The estimated number of barcodes bound per cell relative to the obtained Ct-values. It is evident that there are some differences in the Ct values shown in B, even though the same number of cells were present in all qPCRs. This is however leveled when the values are normalized in respect to their specific probes. QPCR was run in duplicates, here showing mean ± range of dublicates.

Example 5

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This is an example where the Sample (1) was blood which was modified (1b) to generate Peripheral blood mononuclear cells (PBMCs).

- 15 The Backbone (2) was a dextran conjugate with streptavidin and fluorochrome (Dextramer backbone from Immudex).
 - The MHC molecules (3) are peptide-MHC (pMHC) complexes displaying an 8-10 amino acid peptide-antigen. The MHC molecule was modified (3b) by biotinylation to provide a biotin capture-tag on the MHC molecule. The MHC molecule was purified (2c) by HPLC.
- The Label (4) was an oligonucleotide. The oligonucleotide label was synthetized (4a) by DNA Technology A/S (Denmark) and was synthetically modified (4b) with a terminal biotin capture-tag. In parts of the example the oligonucleotide label was further modified by annealing to a partially complimentary oligonucleotide label giving rise to a combined oligonucleotide label.
- 25 The MHC molecule (5) was synthetized (5a) by attaching MHC molecules in the form of a biotinylated pMHC and labels in the form of a biotin-modified oligonucleotide onto a streptavidin-modified dextran backbone (Dextramer backbone from Immudex, Denmark). The MHC molecule further contains a modification (5b) in the form of a fluorochrome. A library of 110 different MHC molecules were generated wherein individual MHC molecules containing different pMHC were encoded by corresponding individual oligonucleotide labels.
 - An amount of sample, PBMC's (1b) was incubated with an amount of a library of MHC molecules (5) under conditions (6c) (e.g. incubation time, buffer, pH and temperature) allowing binding of MHC molecules to T cells in the sample.

The cell-bound MHC molecules were separated from the non-cell bound MHC molecules (7) by first a few rounds of washing the PBMC's through centrifugation sedimentation of cells and resuspension in wash buffer followed by Fluorescence Activated Cell Sorting (FACS) of fluorochrome labeled cells. T cells that can efficiently bind MHC molecules will fluoresce because of the fluorochrome comprised within the MHC molecules; T cells that cannot bind MHC molecules will not fluoresce. FACS-sorting leads to enrichment of fluorescent cells, and hence, enrichment of the MHC molecules that bind T cells of the PBMC sample.

FACS isolated cells were subjected to PCR amplification of the oligonucleotide label
associated with the MHC molecules bound to cells. Subsequent sequencing of individual
DNA fragments generated by the PCR reaction revealed the identity of MHC molecules
that bound to the T cells present in the sample.

This experiment thus revealed the peptide-MHC specificity of the T cell receptors of the T cells present in the blood sample.

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- 1. Sample preparation. The cells sample used in this experiment was obtained by mixing blood drawn from 2 different donors BC 260 and 171 (table 6). To provide a titration of the B0702 CMV pp65 TPR responses in a B0702 negative donor sample. 5 fold dilution of BC 260 in 171 was performed i.e. 100, 20, 5, 1, 0.2, 0.04, 0.0125, 0.0025% of BC 260 corresponding to a theoretical frequency of specific cells of 5%, 1%, 0.2%, 0.04%, 0.008%, 0.0016% and 0.00032% B0702 CMV pp65 TPR. Thus, the sensitivity of the method as well as the relevance of the results obtained in the experiment could be evaluated at the end of the experiment, by comparison with data obtained in parallel, using other methods but similar cells.
 - a. Acquiring sample: Blood was obtained from the Danish Blood Bank.
 - b. Modifying sample:
 - i. As in experiment 1
 - ii. Mixing of two blood samples as described above

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- 2. Backbone preparation: as in experiment 1
- 3. MHC molecules preparation: The MHC molecules used in this example were class I MHC-peptide complexes. The individual specifies (allele and epitope combination) were generated as described in experiment 1. Here we used a library of 110 different peptide MHC molecules corresponding to table 10.

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- a. Synthesis: As described in experiment 1
 - i. Both the MHC heavy chain and B2M was expressed in E.coli as previously described (Hadrup et al. 2009).

ii. p*MHC monomers were refolded and purified as previously described (Hadrup et al. 2009)

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 Modification: The p* UV conditional peptide-ligand was exchanged with the peptide antigens to be explored to produce specific peptide MHC monomers. 42

i. Peptides (Pepscan Presto) were diluted in phosphate buffered saline (DPBS; Lonza) and mixed to final concentrations of 100μg/ml:200μM (monomer:peptide) in individual wells of 384 well plates. Maximum volumes of 70μl were prepared in the respective well formats. The mixtures were exposed to 366 nm UV light (UV cabinet; CAMAG) for one hour and optionally stored for up to 24 h at 4 °C.

c. Purification: as in experiment 2

- 4. Label preparation: In this example, 110 different oligonucleotides, of the same length but different sequence, were generated. Each of the oligonuclotides became attached to a specific pMHC, and thus encoded this specific pMHC. The oligonucleotides were biotinylated, allowing easy attachment to the dextranstreptavidin conjugate backbone.
 - a. Synthesis: In this example the labels were DNA oligonucleotides which were purchased from DNA Technology (Denmark) and delivered as lyophilized powder. Stock dilutions of 100 μM label were made in nuclease free water and stored at -20 °C. Two types of DNA oligonucleotide labels were used and named 1OS and 2OS respectively.
 - 120 1OS labels were ordered from DNA Technology as single stranded DNA-oligonucleotides with 5' biotinylation modification. Labels were diluted to working concentrations (640nM) in nuclease free water with 0.1% Tween. See table 9 and 10 for 1OS label sequences.
 - ii. A 2OS label system was developed to increase the complexity of a limited number of oligonucleotide sequences by a combinatorial label generation strategy to produce multiple unique labels from a more confined number of label precursors. The strategy, referred to as 2OS, involved annealing and subsequent elongation of two partially complimentary oligonucleotide-sequences that fostered a new unique oligonucleotide-sequences that was applied as a DNA oligonucleotide label (table 9 + 10). E.g. by combining 20 unique oligonucleotide-sequences (A label precursor) that are all partly complementary to 60 other unique oligonucleotide sequences (B label precursor) a combinatorial library of 1,200 different (Ax+By) labels could be produced.
 - 1. Partly complementary A and B oligonucleotides were annealed to produce a combined A+B oligonucleotide label. An A and a B oligo was mixed as stated in table 3, heated to 65 °C for 2 min and cooled slowly to <35 °C in 15-30 min. The annealed A and B oligos were then elongated as stated in table 3.4. Components of the elongation reaction were mixed just before use. After mixing, the reaction was left 5 min at RT to allow elongation of the annealed oligonucleotides. The reagents used for annealing (left) and elongation (right) of partly complementary oligonucleotides is described in Table 3. Reagents marked in italic were from the the Sequenase Version 2.0 DNA Sequencing Kit (Affymetrix #70770).</p>

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- b. Modification: All labels were diluted to working concentrations (640nM) in nuclease free water with 0.1% Tween. Elongated oligonucleotide sequence 2OS labels were now treated as 1OS labels.
- 5. MHC molecules preparation: The MHC molecules (pMHCs) and Labels (oligonucloetides) were attached to the backbone (a dextran-streptavidinfluorchrome conjugate), to form the MHC molecules, in a way so that a given pMHC is always attached to a given oligonucleotide – maintaining a 1:1 relation between a pMHC and an oligonucleotide.
 - a. Synthesis: For preparation of MHC molecules the backbone was labeled in the form of a biotinylated oligonucleotide prior to addition of pMHC.
 - i. Creation of MHC molecules were, if not stated otherwise, performed by addition of label in two fold excess over backbone (2:1 label:backbone) and incubated 30 min, 4℃. Binding of label to the backbone (backbone) was always determined after titration when a new batch of backbone and or labels was used. Prior to coupling MHC molecules, pMHC monomers, these were centrifuged 5 min, 3300g. SA conjugate (Dextramer backbone, Immudex) with conjugated streptavidin (SA) and fluorochrome (PE) were aliquoted into new 96 well plates matching the peptide exchange reaction setup. Differences in the procedure for assembling PE. Avoiding the precipitate, MHC molecules were added to the aliquoted SA conjugate and incubated 30 min, RT. Following complex formation, D-biotin (Avidity Bio200) was added together with 0,02% NaN2 in PBS, and incubated 30 min, 4°C. Assembled MHC molecules were stored up to four weeks at 4℃.
 - b. Modification: When the total volume of combined panel of MHC molecules exceeded 100 µl per incubation with sample the volume was reduced.
 - i. Size exclusion spin columns Vivaspin 500, Sartorius) with a cut-off at 300 kDa were saturated by adding 500µl 2% BSA/PBS and centrifuging 5000g, until the volume had passed through. Subsequently, the columns were washed twice by adding 500µl PBS and centrifuging 5000g until no considerable volume was left in the columns. The combined panel of MHC molecules was added to the spin column and centrifuged 5000g, 4°C until the desired volume resided in the column (approximately 80 µl per incubation with sample).
 - c. Purification: The panel of MHC molecules was moved to eppendorph tubes and centrifuged 5 min, 5000g, to sediment any MHC molecules not in solution, before being added to the cells.
- 6. Incubation of sample and MHC molecules: The cell sample and the MHC 40 molecules were mixed in one container, to allow the MHC molecules to bind the T cells that they recognize.
 - a. Amount of sample: 2x10E6 cells in the form of BC's
 - b. Amount of MHC molecule
- c. Conditions: All washing of cells refer to centrifugation 5 min, 490 g, with 45 subsequent removal of supernatant. 2x10E6 cells where transferred to individual wells of 96 well plates, washed in barcode-buffer (PBS/0.5% BSA/2 mM EDTA/100 µg/ml herring DNA) and resuspended in this buffer to approximately 20_{µl} per staining. When incubating sample with MHC

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molecules the cells were incubated with 50nM dasatinib, 30 min, 37 ℃ (Lissina et al. 2009). MHC molecules were centrifuged for 5 min, 3300g, prior to addition to cells. 3 ul of each MHC molecule was required per incubation (1 ug/ml in respect to pMHC). After adding MHC molecules, the cells where incubated 15 min, 37 ℃. The antibody mixture listed in table 2 were added together with 0.1 µl near- IR-viability dye (Invitrogen L10119) that stains free amines. Antibody staining was essentially as for conventional MHC multimer staining. Cells were incubated 30 min, 4 ℃ before washing of any MHC molecules or antibodies that did not bind to the cells. Cells were subsequently fixed by adding 50 ul 1% paraformaldehyde

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7. Enrichment of MHC molecules with desired characteristics: In this Example, the MHC molecules were enriched by using flow cytometry, more specifically, Fluorescence-Activated-Cell-Sorter (FACS). The MHC molecules carry a fluorochrome. Hence, the cells that bind MHC molecules will fluoresce, and can be separated from the cell that do not bind MHC molecules and therefore do not fluoresce, by a FACS sorter. As a result, the MHC molecules that bound to cells will be enriched for.

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a. Apply: Throughout this study two different flow cytometers where used for acquisition. A BD FACSCanto II equipped with three lasers (488 nm blue, 633nm red and 405 violet) and a BD LSR II cytometer equipped with five lasers. Only four lasers on the LSR II were used throughout this study (488 nm blue laser, 640 nm red laser, 355 nm UV laser and 405 nm violet laser). Additionally cells were sorted on BD FACSAria and FACSAria II, equipped with three lasers (488 nm blue, 633nm red and 405 violet). All flow cytometry data analyses used the BD FACSDiva software version 6.1.2.

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i. The following gating strategy was used. All initial gatings of CD8 positive cells were performed alike. Lymphocytes were identified in a FSC/SSC plot. Additional gating on single cells (FSC-A/FSC-H), live cells (near-IR-viability dye negative), and dump-channel negative/CD8 positive cells (FITC/PerCP) where used to define the CD8 T cell population.

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b. Wash: Cells were washed twice in barcode-buffer where after the cells were ready for flow cytometric acquisition. Optionally cells were fixed in 1% paraformaldehyde O.N., 4°C, and washed twice in FACS buffer or barcode-buffer. Fixed cells were stored for up to a week at 4°C.

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c. Separate: Optionally cells were acquired up to one week after fixation in 1% paraformaldehyde. The multimer positive cells were sorted into tubes that had been pre-saturated for 2h-O.N. in 2% BSA and contained 200 μl barcode-buffer to increase the stability of the oligonucleotides that followed with the sorted cells. The sorted multimer positive cells were centrifuged 5 min, 5000 g, to allow removal of all excess buffer. Cells were stored at -80 °C.

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i. Gates were drawn to define the positive events from the single conjugated fluorochrome, i.e. PE or APC.

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ii. The capacity of pMHC dextramers were evaluated based on the mean fluorescent intensity (MFI) or the stain index (SI). SI is a measure of population separation, taken into account also potential effects on the negative population (background) and the spread of the background:

- 8. Identification of enriched MHC molecule: By identifying the Label (in this Example, the oligonucleotide label), the pMHCs that bound cells can be identified. Therefore, the oligonucleotides that were comprised within the MHC molecule that were recovered with the cells, were sequenced. This allowed the identification of pMHCs that bound cells of the cell sample.
 - a. Labels derived from sorted cells were amplified by PCR prior to sequencing. See table 4 for composition of the PCR. PCR was performed with the kit: Taq PCR Master Mix Kit (Qiagen, #201443). The thermal profile is listed in table 5. PCR was run on the thermal cycler: GeneAmp, PCR System 9700 (Applied Biosystem). PCR products were visualized, after gel electrophoresis on a Bio-Rad Gel Doc EZ Imager.
 - i. The forward and reverse primer included adaptors for the sequencing reaction (A-key and P1-key respectively which are compatible with Ion Torren sequencing, Life Technologies).
 - ii. Moreover the forward primer carried a sample-identification barcode (table 8). Labels on sorted cells and their associated MHC molecules derived from individual samples were amplified with primers holding a specific sample-identification sequence (Table 8). This facilitated distribution of sequence reads derived from every single sample. Additionally, the input of concentrated panels of MHC molecules (before mixing with cells) were allocated a sample-identification barcode through PCR (referred to as the panel-input). Sequencing of the panel-input would allow normalization of the analyzed sequence output.
 - iii. Positive sequence reads were aligned to sequences that read from the sample-barcode-identity at the 5'-end all the way through the pMHC-barcode-identity. The numbers of reads were normalized according to the total number of reads that mapped to the same sample-barcode-identity and according to the panel-input reads.

Deconvolute label on MHC molecule

- Sequencing of DNA oligonucleotide labels was carried out on a 314 Ion Torrent chip (GeneDx). Adaptors were introduced via primers during PCR (refer to table 8 for adaptor sequences)
 - i. A sequence database was created consisting of the possible combinations of 15 sample-identification barcodes and 358 pMHC barcodes (118 1OS + 240 2OS), together with the primer and annealing sequences from both the 1OS and 2OS systems. This accumulated to 5370 sequences that could be expected from a sequencing run. Each sequencing read was then used to search the database for alignments, using the nucleotide BLAST algorithm, with a match reward of 1, mismatch reward of -2 and a gap cost of 2 for both opening and extending a gap. In this way sequencing errors were penalized equally, whether a base was miscalled or inserted/deleted in the sequencing read compared to the actual sequence.
 - ii. Alignments were discarded by the following criteria:
 - E-value > 1e-12; insufficient length of alignment (should be greater than 60 or 102 bases for the 1OS and 2OS systems, respectively)

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- 2. Start position in subject sequence larger than 2, i.e. fewer than 5 out of 6 bases in the unique part of the sample-identification barcode was included in the alignment.
- 3. If multiple alignments could still be found for any sequencing read, only the alignment with the best percent identity was kept. Finally, the number of reads mapping to each barcode in the database was counted.

iii. Identifying overrepresented barcodes: Relative read counts were calculated by normalizing each read to the total read count mapping to the same sample-identity barcode. The relative read counts were then used to calculate the fold change per barcode compared to the control sample-barcode input (barcoded detection-molecule panel that was not mixed with cells). Significantly overrepresented barcodes were identified using a 2-sample test for equality of proportions on the raw read counts in a sample versus the control-barcode input, and p-values were corrected for multiple testing using the Benjamini-Hochberg FDR method.

Result of example 5:

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- This example shows the feasibility for detection of antigen responsive T-cell in a large mixture of different pMHC multimer (MHC molecules). We show the sensitivity of the barcode-labelled MHC multimers being at least able to detect 0.00032% of specific T-cell out of CD8 T cells. We find exact correlation withprevious described (low throughput) methods.
- Figure 9. Schematic presentations of the number of specific 1OS barcode reads mapped to seven different samples. A 5% B7 CMV pp65 TPR response (barcode 88) were spiked into a HLA-B7 negative BC in fivefold dilutions, creating seven samples (5%, 1%, 0.2%, 0.04%, 0.008%, 0.0016% and 0.00032%). This BC has a population of A11 EBV-EBNA4 specific T cell (corresponding to barcode 4). Samples were stained with the same panel comprising 110 differently 1OS barcoded-pMHC-dextramers. The bars show the total reads normalized to the input panel in each sample. Experiments were performed in duplicate. Here showing mean.
- 35 Figure 10. Schematic presentations of the number of specific 2OS barcode reads mapped to seven different samples. A 5% B7 CMV pp65 TPR response (barcode A3B18) were spiked into a HLA-B7 negative BC in fivefold dilutions, creating seven samples (5%, 1%, 0.2%, 0.04%, 0.008%, 0.0016% and 0.00032%). This BC has a population of A11 EBV-EBNA4 specific T cell (corresponding to barcode A1B4). Samples were stained with the same panel comprising 110 differently 2OS barcoded-pMHC-dextramers. The bars show

the total reads normalized to the input panel in each sample. Experiments were performed in duplicate. Here showing mean.

Example 6:

5 Examples 6 is conducted exactly as examples 5, with the only difference that we have used a different sample. Here we detect antigen responsive T-cells in 5 different donor blood samples.

Results example 6:

This example shows the feasibility to detect numerous different specificities in different donor samples using DNA barcode labelled MHC multimers. Obtained data show the feasibility for high-throughput screening of T-cell reactivity in numerous donor to assess immune reactivity associated with disease development, vaccination, infection etc.

- Figure 11. A schematic presentations of the number of specific 1OS barcode reads
 15 mapped to six different samples. Six BCs were stained with the same panel comprising
 110 differently 1OS barcoded-pMHC-dextramers. Bar charts show the total reads
 normalized to the input panel in each sample (p<0.05). Each pie chart show significant
 (p<0.01) reads mapped to that sample.
- 20 Figure 12. Schematic presentations of the number of specific 2OS barcode reads mapped to six different samples. Six BCs were stained with the same panel comprising 110 differently 2OS barcoded-pMHC-dextramers. Bar charts show the total reads normalized to the input panel in each sample (p<0.05).

25 Tables:

	SA conjugate/μl exchange	D-biotin	End: pMHC TET	Amount per staining
SA	0.092 µl (0.1 µg/ml)	28 µM	100 μg/ml	1 μΙ
PE	1.32 µl	12.6 µM	44 μg/ml	3 µl
APC	0.73 μΙ	9.78 µM	24.25 μg/ml	3 μΙ

Table 1: A listing of reagents required for production of pMHC multimers produced from 100 μ g/ml exchange reaction. The amounts of the respective reagents used for staining 1x106-2x106 cells in 100 μ l are also specified.

Table 2: A listing of the components in the antibody mixture added after incubation with MHC molecules or after staining with conventional MHC multimers.

Target	Conjugate	Amount (μl)	Source
CD8	PerCP	2	Invitrogen MHCD0831
CD4	FITC	1.25	BD bioscience 345768
CD14	FITC	3.13	BD bioscience 345784
CD16	FITC	6.25	BD bioscience 335035
CD19	FITC	2.50	BD bioscience 345776
CD40	FITC	1.56	Serotec MCA1590F

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Table 3: The reagents used for annealing (left) and elongation (right) of partly complementary oligonucleotides. Reagents marked in italic were from the the Sequenase Version 2.0 DNA Sequencing Kit (Affymetrix #70770).

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Annealing reaction	(10 µl)	Elongation reaction	(15.5 µl)
Oligo A (100 μM)	2.6 μΙ	Annealing reaction	10 μl
Oligo B (100 μM)	5.4 µl	0.1M DTT	1 μ1
Sequenase reaction buffer	2 μ1	H ₂ O	0.5 μ1
		8x diluted Sequenase polymerase	2 μl
		5x diluted Sequence extension mixture	2 μΙ
			•

Table 4: The PCR Master mix applied prior to sequencing of labels on MHC molecules associated with sorted cells. The forward and reverse primer included adaptors for the

sequencing reaction (A-key and P1-key respectively). Moreover the forward primer carried a sample-identification sequence (table 8).

Component	Volume per sample	
	(μl)	
Master Mix	25	
Forward Primer (5 µM)	3	
Reverse Primer (5 µM)	3	
Nuclease free H ₂ O	9	
Template	10	

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Table 5: The thermal profile applied for amplification of labels on MHC molecules associated with sorted cells. 36 cycles were applied if >1,000 cells were sorted while 38 cycles were applied if <1,000 cells were sorted.

Temperature (°C)	Time	No. of cycles
- 95	10 min	1
- 95	30 s	
- 60	45 s	36-38
- 72	30 s	
- 72	4 min	1
- 4	∞	

Table 6. BCs included in the experiments 3-6. The virus specificities detected by combinatorial encoding of conventional MHC multimers with 25 virus peptides. The frequency of each response is listed along with the 1OS and 2OS label numbers appointed in the experiment.

	Epitope	Freq. (%)	108	20S
BC261	A2 FLU MP 58-66 GIL	0.1249	24	A3B4
	A3 EBV EBNA 3a RLR	0.0258	60	A6B10

	A2 EBV LMP2 FLY	0.0075	27	A3B7
BC266	A1 CMV pp65 YSE	0.0859	1	A1B1
	A1 FLU BP-VSD	0.0628	3	A1B3
BC171	A11 EBV-EBNA4	0.3	4	A1B4
	A3 CMV pp150 TVY	0.015	61	A1B11
BC254	A2 FLU MP 58-66 GIL	0.0522	24	A3B4
	A2 EBV LMP2 FLY	0.014	27	A3B7
	A2 CMV pp65 NLV	1.1279	28	A3B8
BC268	A2 FLU MP 58-66 GIL	0.2523	24	A3B4
	A2 CMV pp65 NLV	05445	28	A3B8
BC260	A2 FLU MP 58-66 GIL	0.0456	24	A3B4
	A2 CMV pp65 NLV	0.134	28	A3B8
	B7 CMV pp65 TPR	4.5395	88	A3B18

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Table 7: Test Oligos with different end modifications

5 'b' = Biotin-TEG 5' modification

'h' = HEG (terminal modifications)

Forward-01

GAGATACGTTGACCTCGTTG

Reverse-01

ATGCAACCAAGAGCTTAAGT

Reverse-03

hATGCAACCAAGAGCTTAAGT

10 TestOligo-01

TestOligo-02

15 CTCTTGGTTGCAT

TestOligo-03

TestOligo-04

20 bhGAGATACGTTGACCTCGTTGAANNNNNNTCTTGAACTATGAATCGTCTCACTTAAG CTCTTGGTTGCATh TestOligo-05

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bhGAGATACGTTGACCTCGTTGAANNNNNNTCTATAGGTGTCTACCTCACTTAAG CTCTTGGTTGCATh

TestOligo-06

5 bhGAGATACGTTGACCTCGTTGAANNNNNNTCTTTATTGGAGAGCACGCTCACTTAAG CTCTTGGTTGCATh

	Probe-03 Tm 64,9	8TCTATCCATTCCATCCAGCTC7	8 = FAM; 7 = BHQ-1-plus
	Probe-04 Tm 57,3	8TCTTGAACTATGAATCGTCTC7	8 = FAM; 7 = BHQ-1-plus
10	Probe-05 Tm 58,5	9TCTATAGGTGTCTACTACCTC7	9 = HEX; 7 = BHQ-1-plus
	Probe-06 Tm 60,9	2TCTTTATTGGAGAGCACGCTC1	2 = Cy5; 1 = BHQ-2-plus

LNA-3 TCTATCCATTCCATCCAGC 8 = FAM; 7 = BHQ-1-plus LNA-4 TCT[+T][+G][+A]AC[+T][+A]TG[+A][+T]CGTC 8 = FAM; 7 = BHQ-1-plus 15 LNA-5 TCT[+A][+T][+A]GG[+T][+G]TC[+T][+A][+C]TACC 9 = HEX; 7 = BHQ-1-plus LNA-6 TCT[+T][+T][+A]TT[+G][+G]AG[+A][+G][+C]ACGC 2 = Cy5; 1 = BHQ-2-plus

Table 8:

A-Keys hold the sample identification barcode and keys for the Ion torrent sequencing.

P1-keys only holds Ion torrent sequencing key

A-Key 10S-F1-1	CCATCTCATCCCTGCGTGTCTCCGACTCAGGAAGATGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-2	CCATCTCATCCCTGCGTGTCTCCGACTCAGTCCTGAGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-3	CCATCTCATCCCTGCGTGTCTCCGACTCAGTGTGGAGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-4	CCATCTCATCCCTGCGTGTCTCCGACTCAGCATTTAGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-5	CCATCTCATCCCTGCGTGTCTCCGACTCAGTTACCCGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-6	CCATCTCATCCCTGCGTGTCTCCGACTCAGATTCTCGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-7	CCATCTCATCCCTGCGTGTCTCCGACTCAGAGACCCGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-8	CCATCTCATCCCTGCGTGTCTCCGACTCAGCGCATGGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-9	CCATCTCATCCCTGCGTGTCTCCGACTCAGTCCTCGGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-10	CCATCTCATCCCTGCGTGTCTCCGACTCAGATTCCTGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-11	CCATCTCATCCCTGCGTGTCTCCGACTCAGCGTCGAGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-12	CCATCTCATCCCTGCGTGTCTCCGACTCAGGCCAATGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-13	CCATCTCATCCCTGCGTGTCTCCGACTCAGATACGGGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-14	CCATCTCATCCCTGCGTGTCTCCGACTCAGGTCAGAGATTCTATAAACTGTGCGGTCC
A-Key 10S-F1-15	CCATCTCATCCCTGCGTGTCTCCGACTCAGCGAGTTGATTCTATAAACTGTGCGGTCC
A-Key 20S-F1-1	CCATCTCATCCCTGCGTGTCTCCGACTCAGCTGGGGGAAGTTCCAGCCAG
A-Key 2OS-F1-2	CCATCTCATCCCTGCGTGTCTCCGACTCAGCTCCACGAAGTTCCAGCCAG
A-Key 20S-F1-3	CCATCTCATCCCTGCGTGTCTCCGACTCAGCTTACCGAAGTTCCAGCCAG
A-Key 20S-F1-4	CCATCTCATCCCTGCGTGTCTCCGACTCAGTGGCAGGAAGTTCCAGCCAG
A-Key 20S-F1-5	CCATCTCATCCCTGCGTGTCTCCGACTCAGTGAGTAGAAGTTCCAGCCAG
A-Key 20S-F1-6	CCATCTCATCCCTGCGTGTCTCCGACTCAGATTCAGGAAGTTCCAGCCAG
A-Key 20S-F1-7	CCATCTCATCCCTGCGTGTCTCCGACTCAGTGAGCTGAAGTTCCAGCCAG
A-Key 20S-F1-8	CCATCTCATCCCTGCGTGTCTCCGACTCAGGGCGTGGAAGTTCCAGCCAG
A-Key 2OS-F1-9	CCATCTCATCCCTGCGTGTCTCCGACTCAGAAATTGGAAGTTCCAGCCAG
A-Key 20S-F1-10	CCATCTCATCCCTGCGTGTCTCCGACTCAGGCTGACGAAGTTCCAGCCAG
A-Key 20S-F1-11	CCATCTCATCCCTGCGTGTCTCCGACTCAGTTCTTAGAAGTTCCAGCCAG
A-Key 20S-F1-12	CCATCTCATCCCTGCGTGTCTCCGACTCAGTGGTGGGAAGTTCCAGCCAG
A-Key 20S-F1-13	CCATCTCATCCCTGCGTGTCTCCGACTCAGGCAGTCGAAGTTCCAGCCAG
A-Key 20S-F1-14	CCATCTCATCCCTGCGTGTCTCCGACTCAGTCGTGAGAAGTTCCAGCCAG
A-Key 20S-F1-15	CCATCTCATCCCTGCGTGTCTCCGACTCAGTACAGTGAAGTTCCAGCCAG

P1-key 10S-R1	CCTCTCTATGGGCAGTCGGTGATGAGTACATGATAGCGCCGTAC
P1-key 2OS-R1	CCTCTCTATGGGCAGTCGGTGATCTGTGACTATGTGAGGCTTTC

Table 9: Representative oligo's applied in the 110 member library of MHC molecules (examples 3-6)

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	5'		6xN
Oligo name	modification	Forward primer region	region
10S-1-Oligo-1	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNN
10S-1-Oligo-2	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNNN
10S-1-Oligo-3	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNNN
10S-1-Oligo-4	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNN
10S-1-Oligo-5	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNNN
10S-1-Oligo-6	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNN
10S-1-Oligo-7	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNN
10S-1-Oligo-8	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNN
10S-1-Oligo-9	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNN
10S-1-Oligo-10	Biotin-C6-	AGATTCTATAAACTGTGCGGTCCTT	NNNNN

Coding region

TATGAGGACGAATCTCCCGCTTATA
GGTCTTGACAAACGTGTGCTTGTAC
GTTTATCGGGCGTGGTGCTCGCATA
CCGATGTTGACGGACTAATCCTGAC
TAGTAGTTCAGACGCCGTTAAGCGC
CCGTACCTAGATACACTCAATTTGT
GGGGTTCCGTTTTACATTCCAGGAA
TATCCCGTGAAGCTTGAGTGGAATC
GGTATGGCACGCCTAATCTGGACAC

Coding region

Reverse primer region

GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG
GGTACGGCGCTATCATGTACTCATG

Annealing region

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20S-A

	5'		6xN
Oligo name	modification	Forward primer region	region
2OS-1-Oligo-A1	Biotin-C6-	GAAGTTCCAGCCAGCGTCACAGTTT	NNNNNN
2OS-1-Oligo-A2	Biotin-C6-	GAAGTTCCAGCCAGCGTCACAGTTT	NNNNNN
2OS-1-Oligo-A3	Biotin-C6-	GAAGTTCCAGCCAGCGTCACAGTTT	NNNNN
2OS-1-Oligo-A4	Biotin-C6-	GAAGTTCCAGCCAGCGTCACAGTTT	NNNNNN
2OS-1-Oligo-A5	Biotin-C6-	GAAGTTCCAGCCAGCGTCACAGTTT	NNNNNN
2OS-1-Oligo-A6	Biotin-C6-	GAAGTTCCAGCCAGCGTCACAGTTT	NNNNNN

CGAGGCAATGGTTAACTGACACGT GGTCAGCATCATTTCC
CAGAAAGCAGTCTCGTCGGTTCGAA GGTCAGCATCATTTCC
TAAGTAGCGGCATAATGTACGCTC GGTCAGCATCATTTCC
GGATCCAGTAAGCTACTGCGTTTAT GGTCAGCATCATTTCC

GGGCTGCGGAGCGTTTACTCTGTAT GGTCAGCATCATTTCC AAACGTATGTGCTTTGTCGGATGCC GGTCAGCATCATTTCC

2OS-B

Oligo name	5' modification	Forward (2OS-R) primer region	6xN region
2OS-1-Oligo-B1		CTGTGACTATGTGAGGCTTTCTCGA	NNNNNN
2OS-1-Oligo-B2	•	CTGTGACTATGTGAGGCTTTCTCGA	NNNNN
2OS-1-Oligo-B3		CTGTGACTATGTGAGGCTTTCTCGA	NNNNN
2OS-1-Oligo-B4		CTGTGACTATGTGAGGCTTTCTCGA	NNNNN
2OS-1-Oligo-B5		CTGTGACTATGTGAGGCTTTCTCGA	NNNNN
2OS-1-Oligo-B6		CTGTGACTATGTGAGGCTTTCTCGA	NNNNN
2OS-1-Oligo-B7		CTGTGACTATGTGAGGCTTTCTCGA	NNNNN
2OS-1-Oligo-B8		CTGTGACTATGTGAGGCTTTCTCGA	NNNNNN
2OS-1-Oligo-B9		CTGTGACTATGTGAGGCTTTCTCGA	NNNNN
2OS-1-Oligo-B10		CTGTGACTATGTGAGGCTTTCTCGA	NNNNNN

Coding region	Annealing region
GCCTGTAGTCCCACGCGATCTAACA	GGAAATGATGCTGACC
CAACCATTGATTGGGGACAACTGGG	GGAAATGATGCTGACC
ACGTTTAAGCATCTGTACTCCAGAT	GGAAATGATGCTGACC
GAATTGAAGCCATCGTTTCGCGCAA	GGAAATGATGCTGACC
CGTAGCTTTTGTAGCGTCTGAGGGC	GGAAATGATGCTGACC
AATCGTCAGTCCCTGTTTCGACATC	GGAAATGATGCTGACC
CGGTGGTAGGTGATACTTCTGTACC	GGAAATGATGCTGACC
TGACTATCGGGGCGTGACATGAGCT	GGAAATGATGCTGACC
GTTGGTGAAACTACCGACGCTTTAC	GGAAATGATGCTGACC
AATGGAGGTGCAGGAATACTCTCGT	GGAAATGATGCTGACC

Table 10: liste of labels and pMHC molecules in 110 member library (examples 3-6)

Barcode 10S	Barcode 2OS	HLA	Peptide	Sequence
1	A1B1	A1	CMV pp65 YSE	YSEHPTFTSQY
2	A1B2	A1	CMV pp50 VTE	VTEHDTLLY
3	A1B3	A1	FLU BP-VSD	VSDGGPNLY
4	A1B4	A11	EBV-EBNA4	AVFDRKSDAK
5	A1B5	A11	HCMV pp65	GPISGHVLK
6	A1B6	A11	VP1	DLQGLVLDY
7	A1B7	A11	VP1	VLGRKMTPK
8	A1B8	A11	VP1	VTLRKRWVK
9	A1B9	A11	VP1	LVLDYQTEY
10	A1B10	A11	VP1	GQEKTVYPK
11	A2B1	A11	VP1	VTFQSNQQDK
12	A2B2	A11	VP1	LKGPQKASQK
13	A2B3	A11	VP1	NVASVPKLLVK
14	A2B4	A11	VP1	TSNWYTYTY
15	A2B5	A11	VP1	LVLDYQTEYPK
16	A2B6	A11	VP1	TLRKRWVKNPY
17	A2B7	A11	VP1	AVTFQSNQQDK
18	A2B8	A11	VP1	PLKGPQKASQK
19	A2B9	A2	VP1	RIYEGSEQL
20	A2B10	A11	VP1	SLFSNLMPK
21	A3B1	A2	VP1	KLLVKGGVEV
22	A3B2	A11	VP1	SLINVHYWDMK
23	A3B3	A2	HPV E6 29-38	TIHDIILECV
24	A3B4	A2	FLU MP 58-66 GIL	GILGFVFTL
25	A3B5	A2	EBV LMP2 CLG	CLGGLLTMV
26	A3B6	A2	EBV BMF1 GLC	GLCTLVAML
27	A3B7	A2	EBV LMP2 FLY	FLYALALLL
28	A3B8	A2	CMV pp65 NLV	NLVPMVATV
29	A3B9	A2	EBV BRLF1 YVL	YVLDHLIVV
30	A3B10	A2	HPV E7 11-20	YMLDLQPETT
31	A4B1	A2	CMV IE1 VLE	VLEETSVML
32	A4B2	A2	VP1	GCCPNVASV
33	A4B3	A2	VP1	SITQIELYL
34	A4B4	A2	VP1	LQMWEAISV
35	A4B5	A2	VP1	AISVKTEVV
36	A4B6	A2	VP1	KMTPKNQGL
37	A4B7	A2	VP1	TVLQFSNTL

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38	A4B8	A2	VP1	GLFISCADI
39	A4B9	A2	VP1	LLVKGGVEVL
40	A4B10	A2	VP1	ELYLNPRMGV
41	A5B1	A2	VP1	NLPAYSVARV
42	A5B2	A2	VP1	TLQMWEAISV
43	A5B3	A2	VP1	QMWEAISVKT
44	A5B4	A2	VP1	VVGIS <u>S</u> LINV
45	A5B5	A2	VP1	SLINVHYWDM
46	A5B6	A2	VP1	HMFAIGGEPL
47	A5B7	A2	VP1	FAIGG <u>E</u> PLDL
48	A5B8	A2	VP1	NLINSLFSNL
49	A5B9	A2	VP1	FLFKTSGKMAL
50	A5B10	A2	VP1	ALHGLPRYFNV
51	A6B1	A2	VP1	NLINSLFSNLM
52	A6B2	A2	VP1	FLDKFGQEKTV
53	A6B3	A2	VP1	VKGGV <u>E</u> VLSV
54	A6B4	A24	HCMV 248-256	AYAQKIFKIL
55	A6B5	A24	EBV LMP2	IYVLVMLVL
56	A6B6	A24	EBV BRLF1	TYPVLEEMF
57	A6B7	A24	EBV BMLF1	DYNFVKQLF
58	A6B8	A3	CMV pp150 TTV	TTVYPPSSTAK
59	A6B9	A3	FLU NP 265-273 ILR	ILRGSVAHK
60	A6B10	A3	EBV EBNA 3a RLR	RLRAEAQVK
61	A1B11	A3	CMV pp150 TVY	TVYPPSSTAK
62	A1B12	A3	EBV BRLF1 148-56 RVR	RVRAYTYSK
63	A1B13	A3	VP1	ASVPKLLVK
64	A1B14	A3	VP1	CCPNVASVPK
65	A1B15	A3	VP1	ITIETVLGR
66	A1B16	A3	VP1	NTLTTVLLD
67	A1B17	A3	VP1	ALHGLPRYF
68	A1B18	A3	VP1	VASVPKLLVK
69	A1B19	A3	VP1	VSGQPMEGK
70	A1B20	A3	VP1	KASSTCKTPK
71	A2B11	A3	VP1	KTPKRQCIPK
72	A2B12	A3	VP1	YTYTYDLQPK
73	A2B13	A3	VP1	PITIETVLGR
74	A2B14	B7	VP1	SVARVSLPM
75	A2B15	A3	VP1	NSLFSNLMPK
76	A2B16	A3	VP1	KVSGQPMEGK
77	A2B17	A3	VP1	TVYPKPSVAP
78	A2B18	A3	VP1	SLINVHYWDMK
79	A2B19	A3	VP1	GVEVLSVVT
80	A2B20	A3	VP1	PLDLQGLVL
81	A3B11	A3	VP1	GLDPQAKAK
82	A3B12	A3	VP1	EVWCPDPSK
83	A3B13	A3	VP1	ADIVGFLFK
			<u> </u>	

84	A3B14	A3	VP1	KTSGKMALH
85	A3B15	A3	VP1	KMALHGLPR
86	A3B16	A3	VP1	RYFNVTLRK
87	A3B17	A3	VP1	TLRKRWVKN
88	A3B18	B7	CMV pp65 TPR	TPRVTGGGAM
89	A3B19	B7	CMV pp65 RPH-L	RPHERNGFTV
90	A3B20	B7	EBV EBNA RPP	RPPIFIRLL
91	A4B11	B7	VP1	KPGCCPNVA
92	A4B12	B7	VP1	QPIKENLPA
93	A4B13	B7	VP1	LPRYFNVTL
94	A4B14	B7	VP1	MPKVSGQPM
95	A4B15	B7	VP1	YPKPSVAPA
96	A4B16	B7	VP1	KPSVAPAAV
97	A4B17	B7	VP1	APLKGPQKA
98	A4B18	B7	VP1	APKRKASSTC
99	A4B19	B7	VP1	SVARVSLPML
100	A4B20	B7	VP1	YPKTTNGGPI
101	A5B11	87	VP1	YPKPSVAPAA
102	A5B12	B7	VP1	KPGCCPNVASV
103	A5B13	B7	VP1	NPRMGVNSPDL
104	A5B14	B7	VP1	LPAYSVARVSL
105	A5B15	B7	VP1	TPTVLQFSNTL
106	A5B16	B7	VP1	LPRYFNVTLRK
107	A5B17	B7	VP1	YPVVNLINSLF
108	A5B18	B7	VP1	YPKPSVAPAAV
109	A5B19	B7	VP1	KPSVAPAAVTF
110	A5B20	В7	VP1	APKRKASST

Table 11: Number of cells sorted in examples 3-6, using the 110 member library

Six BCs

108	CD8 cells	Sorted cells	Fraction (%)
BC171	55036	3737	6,790101025
BC254	228535	33 69	1,474172446
BC261	49227	792	1,608873179
BC266	27769	1237	4,454607656
BC268	120307	2490	2,069705005
2 0 S	CD8 cells	Sorted cells	Fraction (%)
BC171	80851	4681	5,789662466
BC254	175729	2663	1,515401556
BC261	57926	816	1,408693851
BC266	46916	2077	4,427061131
BC268	250144	4157	1,661842779

5 Table 12:

Abbreviations

1OS Single oligo system

2OS Two oligo system

AIRE Autoimmune regulator

10 APC Allophycocyanin

Barcode oligonucleotide sequence

BC Buffy coat

B cell B lymphocyte

BSA Bovine Serum albumin

15 CD Cluster of differentiation

CDR Complementary-determining regions

CMV Cytomegalovirus

Ct Cross threshold

CTL Cytotoxic T lymphocyte

20 CyTOF Cytometry by time-of-flight

DC Dendritic cells

DMSO Dimethyl sulfoxide

dT Thymidine backbone

EBV Epstein-Barr virus

EDTA Ethylenediaminetetraacetic acid

ELISPOT enzyme-linked immunospot

ER Endoplasmatic reticulum

5 FACS Fluorescence activated cell sorting

FBS Fetal Bovine Serum

FCS Fetal calf serum

FITC Fluorescein isothiocyanate

HEG Hexaethylene glycol

10 HIV Human immunodeficiency virus

HLA Human leukocyte antigen

HPLC High-performance liquid chromatography

IFN Interferon

li Invariant chain

15 IL Interleukin

MHC Major Histocompatibility Complex

N6 Randon six nucleotides

NIR Near-infrared

nt Nucleotide

20 O.N. Over night

PBMC Peripheral blood mononuclear cell

PBS Phosphate buffered saline

PCR Polymerase chain reaction v

PE R-phycoerythrin

25 PerCP Peridinin chlorophyll

p* UV-conditional peptide

PBS Phosphate buffered saline

pMHC Peptide-Major histocompatibility complex

PCR Polymerase chain reaction

30 qPCR Quantitative polymerase chain reaction

RAG1/RAG2 Recombinant activating genes

RT Room temperature

SA Streptavidin

SI Stain index

35 TAP1/TAP2 Transporter associated with antigen processing

T cell T lymphocyte
TCR T cell receptor
TEG Triethylene glycol
TET Tetramers
5 Th T helper cells
TIL Tumor Infiltrating Lymphocyte
Tm Melting temperature

TNF Tumor necrosis factor
Treg T regulatory cells vi vii

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Claims

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- 1. A multimeric major histocompatibility complex (MHC) comprising
- two or more MHC molecules linked by a backbone molecule; and
- at least one nucleic acid molecule linked to said backbone, wherein said nucleic acid molecule comprises a central stretch of nucleic acids (barcode region) designed to be amplified by e.g. PCR.
- The multimeric major histocompatibility complex according to claim 1, wherein the
 backbone molecule is selected from the group consisting of polysaccharides, such as glucans such as dextran, a streptavidin or a streptamer multimer.
 - 3. The multimeric major histocompatibility complex according to claim 1 or 2, wherein the MHC molecules are coupled to the backbone through a streptavidin-biotin binding and/or streptavidin-avidin and/or via the MHC heavy chain and/or via light chain (B2M).
 - 4. The multimeric major histocompatibility complex (MHC) according to any of the preceding claims, wherein said multimeric MHC is composed of at least four MHC molecules, such as at least eight, such as at least ten, 2-30, 2-20, such as 2-10, such as 4-10 or such as 15-30 MHC molecules.
 - 5. The multimeric major histocompatibility complex (MHC) according to any of the preceding claims, wherein the at least one nucleic acid molecule is composed of at least a 5' first primer region, a central region (barcode region), and a 3' second primer region.
 - 6. The multimeric major histocompatibility complex (MHC) according to any of the preceding claims, wherein the at least one nucleic acid molecule has a length in the range 20-200 nucleotides, such as 20-150, such as 20-100, such as 30-80, such as 30-50 nucleotides.
 - 7. The multimeric major histocompatibility complex (MHC) according to any of the preceding claims, wherein the at least one nucleic acid molecule is linked to said backbone via a streptavidin-biotin binding and/or streptavidin-avidin binding.

8. The multimeric major histocompatibility complex (MHC) according to any of the preceding claims, wherein the at least one nucleic acid molecule comprises or consists of DNA, RNA, and/or artificial nucleotides such as PNA or LNA.

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- The multimeric major histocompatibility complex (MHC) according to any of the preceding claims, wherein the MHC is selected from the group comprising class I MHC, a class II MHC, a CD1, or other MHC-like molecules.
- 10 10. The multimeric major histocompatibility complex (MHC) according to any of the preceding claims, wherein the backbone further comprises one or more linked fluorescent labels and/or His-tags, and/or metalion tags, and/or other selectable tags or labels.
- 15 11. A composition comprising:
 - i. a subset of multimeric major histocompatibility complexes (MHC molecules)
 according to any of claims 1-10, wherein each subset of MHC molecules has a
 different peptide decisive for T cell recognition and
 - ii. a nucleic acid molecule comprising a central stretch of nucleic acids (barcode region).
 - 12. The composition according to claim 11, comprising at least 10 different subsets of MHC molecules such as at least 100, such as at least 500, at least 1000, at least 5000, such as in the range 10-100000, such as 10-50000, such as 10-1000 or such as 50-500 sets of MHC molecules.
 - 13. A method for detecting antigen responsive cells in a sample comprising:
 - i) providing one or more multimeric major histocompatibility complexes (MHC molecules) according to any of claims 1-10 or a composition according to any of claims 11-12;
 - ii) contacting said multimeric MHC molecules with said sample; and
 - iii) detecting binding of the multimeric MHC molecules to said antigen responsive cells, thereby detecting cells responsive to an antigen present in a set of MHC molecules, wherein said binding is detected by amplifying

the barcode region of said nucleic acid molecule linked to the one or more MHCmolecules through the backbone molecule.

- 14. The method according to claim 13, wherein the sample is a blood sample, such as a peripheral blood sample, a blood derived sample, a tissue sample or body fluid, such as spinal fluid, or saliva.
- 15. The method according to any of claims 13-14, wherein said sample has been obtained from a mammal, such as a human, mouse, pigs, and/or horses.
- 16. The method according to any of claim 13-15, wherein the method further comprises cell selection by e.g. flow cytometry such as FACS, magnetic-bead based selection, size-exclusion, gradient centrifugation, column attachment or gel-filtration.
- 17. The method according to any of claims 13-16, wherein said binding detection includes comparing measured values to a reference level, e.g. a negative control and/or total level of response.
- 18. The method according to any of claim 13-17, wherein said amplification is PCR such as QPCR.
 - 19. The method according to any of claims 13-18, wherein the detection of barcode regions includes sequencing of said barcode region, by deep sequencing, highthroughput sequencing or next generation sequencing, or detection of said barcode region by QPCR,
 - 20. Use of a multimeric major histocompatibility complex (MHC) according to any of claims 1-10 or a composition according to any of claims 11-12 for the detecting of antigen responsive cells in a sample.
 - 21. Use of a multimeric major histocompatibility complex (MHC) according to any of claims 1-10 or a composition according to any of claims 11-12 in the diagnosis of diseases or conditions, preferably cancer and/or infectious diseases.

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22. Use of a multimeric major histocompatibility complex (MHC) according to any of claims 1-10 or a composition according to any of claims 11-12 in the development of immune-therapeutics, and/or development of vaccines, and/or identification of epitopes.

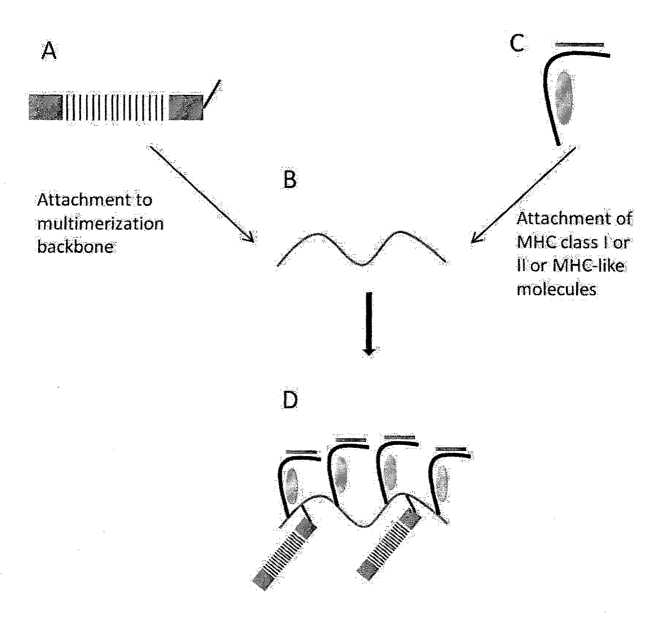
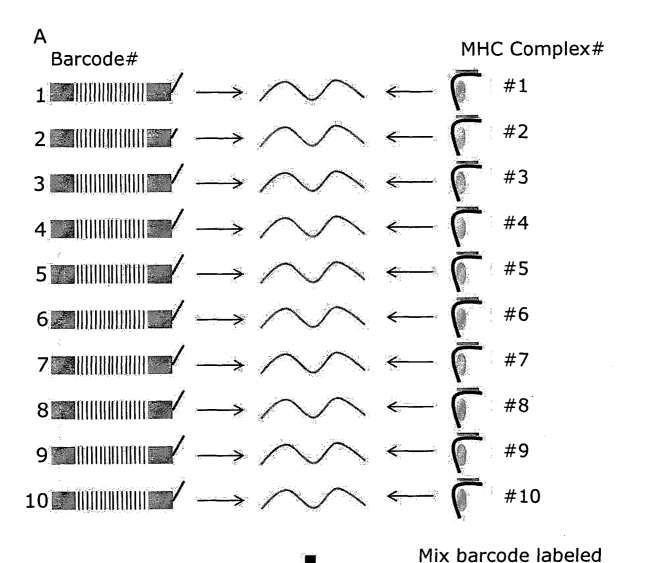
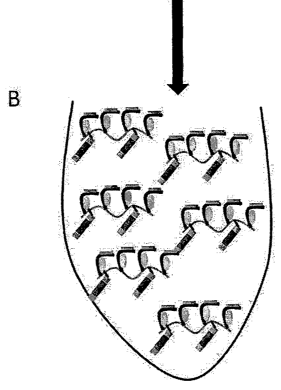


Fig. 1





peptide-MHC multimer library in one tube. Possible complexity up to 1000-10.000 different MHC multimers

Fig. 2

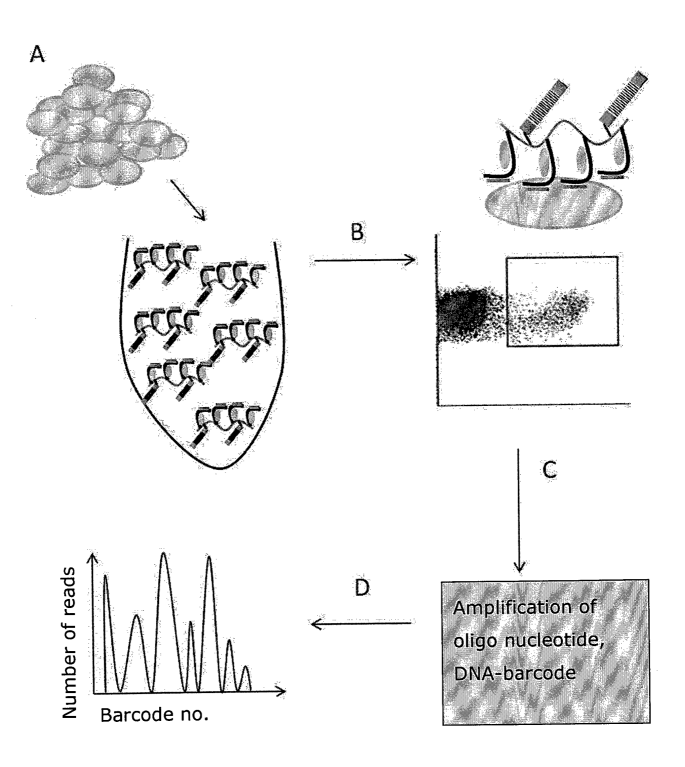


Fig. 3

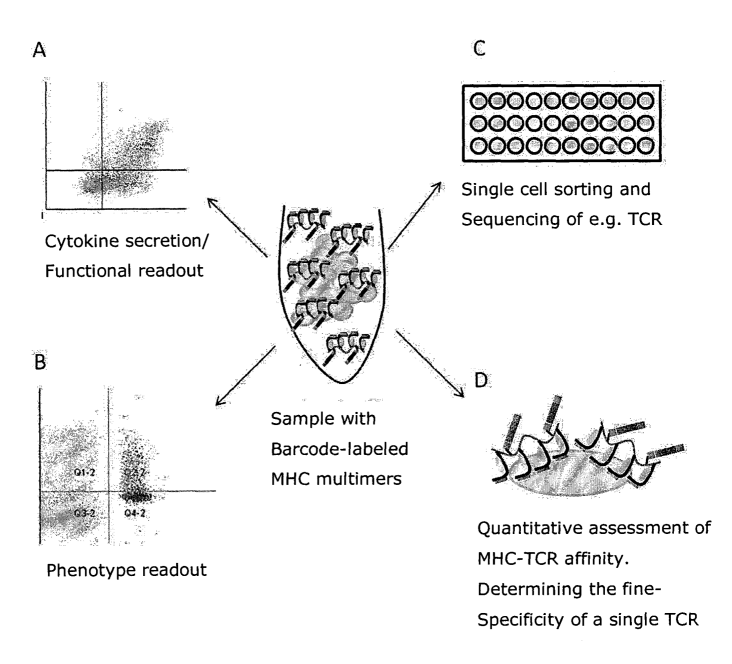


Fig. 4

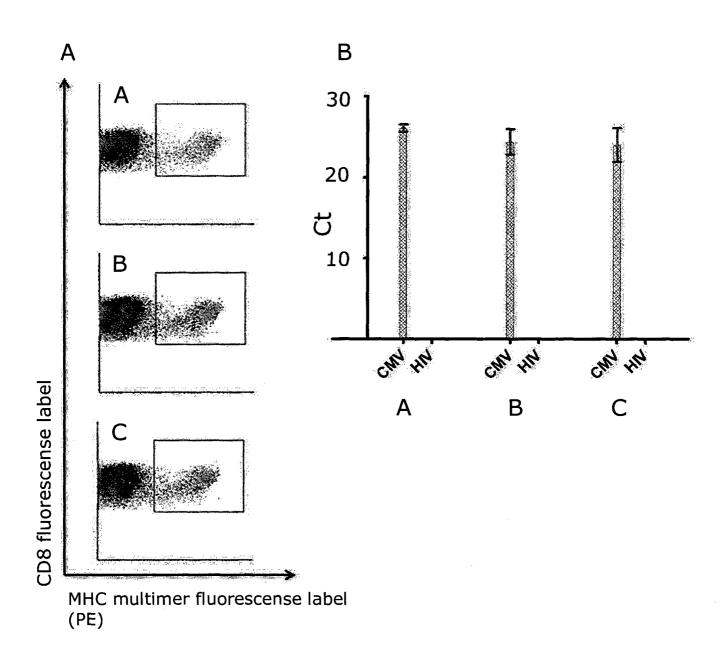
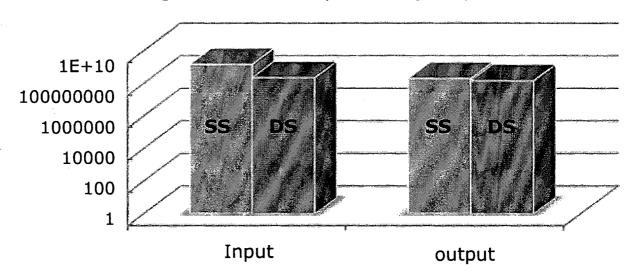


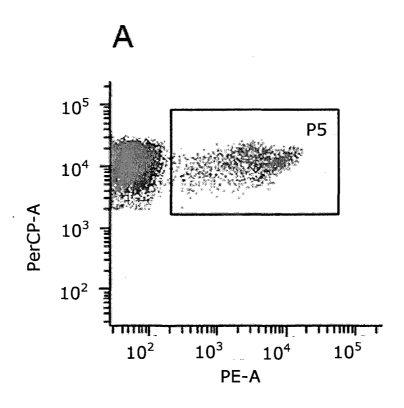
Fig. 5

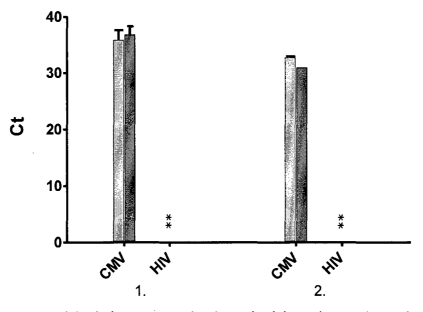
Stability of DNA oligo codons in blood (EDTA) during 24h incubation (MACS capture)



Single stranded (SS) and double stranded (DS) DNA

Fig. 6



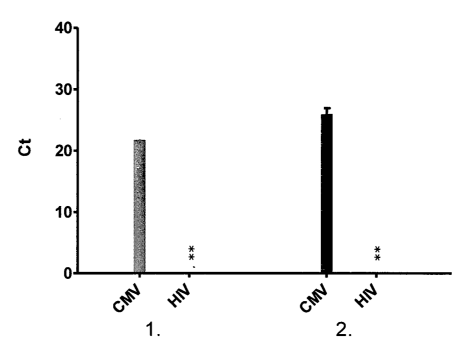


** did not reach threshold within 40 cycles

Fig 7

8 / 12

PCT/DK2015/050150



** did not reach threshold within 40 cycles

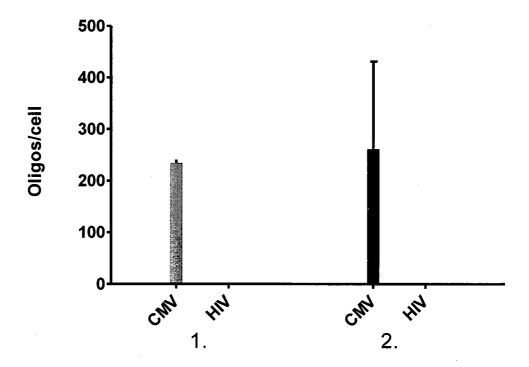
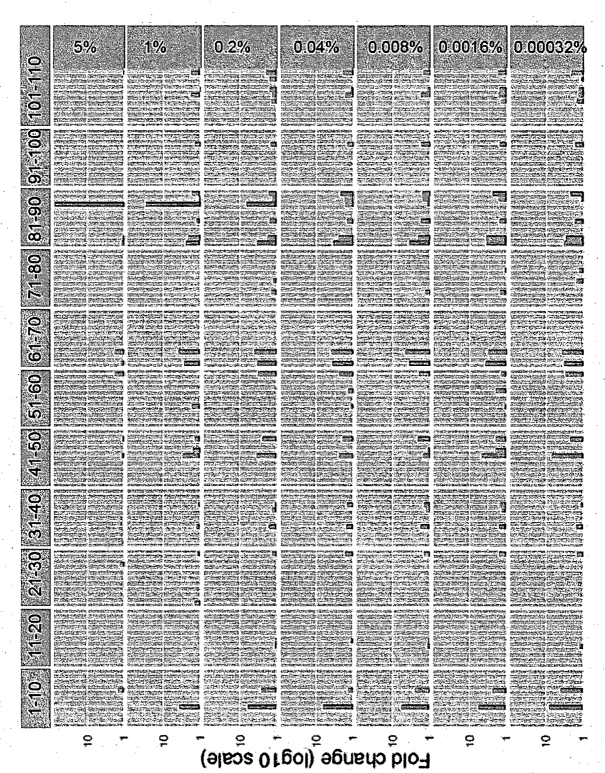
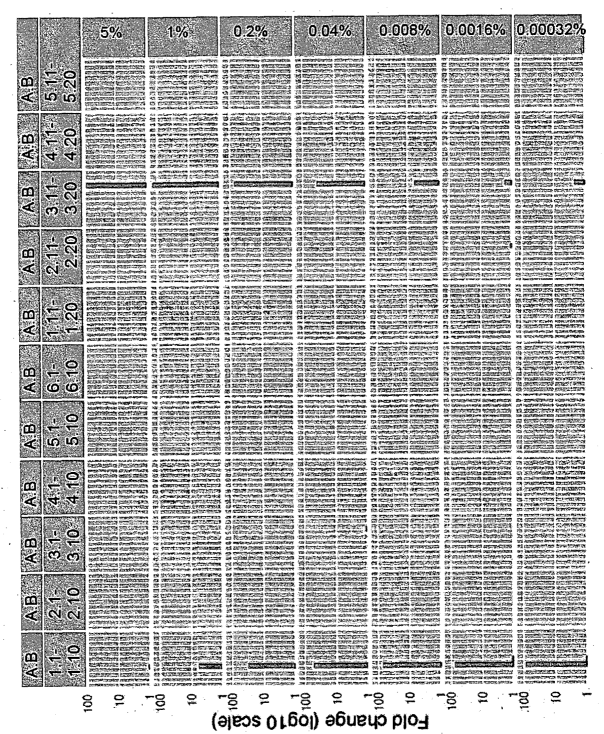


Fig 8



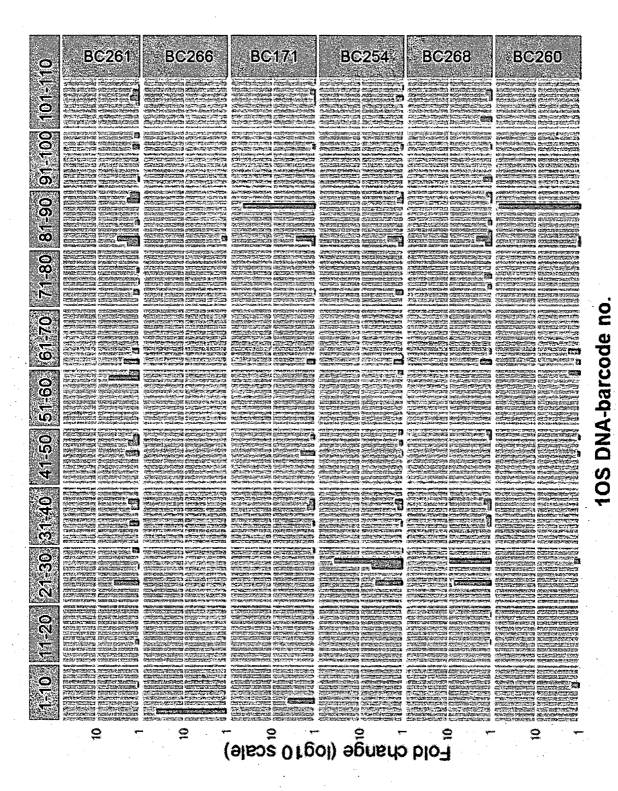
signif p < 0.05 p >= 0.05

Fig 9



signif p < 0.05 p >= 0.05

Fig 10



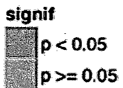


Fig 11

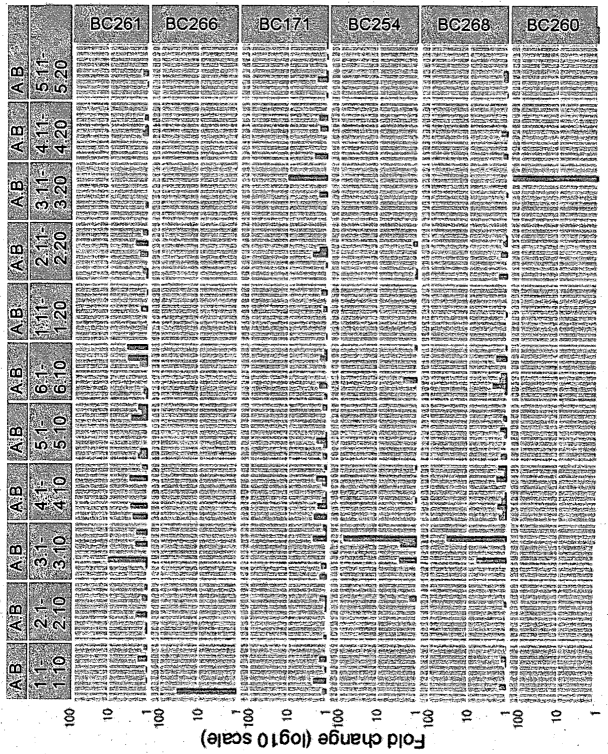


Fig 12

INTERNATIONAL SEARCH REPORT

International application No PCT/DK2015/050150

A. CLASSIFICATION OF SUBJECT MATTER INV. C07K14/74 C12Q1/68

C12N15/10

G01N33/543

G01N33/569

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

ADD.

Minimum documentation searched (classification system followed by classification symbols) $C07\,K$ C120 C12N G01N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal, WPI Data, BIOSIS, Sequence Search

WO 2009/126828 A2 (CALIFORNIA INST OF TECHN [US]; UNIV CALIFORNIA [US]; KWONG GABRIEL A [) 15 October 2009 (2009-10-15) page 19 - page 22 page 34 - page 46	1-6, 8-11,13, 16,17,20 7,12,14, 15,18,
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page 19 - page 22 page 34 - page 46	7,12,14,
figures 1-3; examples 6,12	19,21,22
WO 2010/037397 A1 (DAKO DENMARK AS [DK]; BRIX LISELOTTE [DK]; SCHOELLER JOERGEN [DK]; PED) 8 April 2010 (2010-04-08) example 9	7
US 2005/019843 A1 (CHEN DANIEL SHIN-YU [US] ET AL) 27 January 2005 (2005-01-27) page 6 - page 7	12,14, 15,21,22
	BRIX LISELOTTE [DK]; SCHOELLER JOERGEN [DK]; PED) 8 April 2010 (2010-04-08) example 9 US 2005/019843 A1 (CHEN DANIEL SHIN-YU [US] ET AL) 27 January 2005 (2005-01-27) page 6 - page 7

Further documents are listed in the continuation of Box C.	X See patent family annex.
"A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier application or patent but published on or after the international filling date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "&" document member of the same patent family
Date of the actual completion of the international search 28 September 2015	Date of mailing of the international search report $09/10/2015$
Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Authorized officer Deleu, Laurent

INTERNATIONAL SEARCH REPORT

International application No
PCT/DK2015/050150

C(Continua	ation). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Υ	WO 2012/094492 A2 (JEANSEE LLC [US]; MACULA ANTHONY J [US]) 12 July 2012 (2012-07-12) page 35 - page 36	18,19
Υ	WO 2009/077173 A2 (PHILOCHEM AG [CH]; NERI DARIO [CH]; MELKKO SAMU [CH]; MANNOCI LUCA [CH) 25 June 2009 (2009-06-25) example 4	19
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