Competition-Based Cell Assay Employing Soluble T Cell Receptors to Assess MHC Class II Antigen Processing and Presentation

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Abstract. Accurate assessment of antigen-specific immune responses is critical in the development of safe and efficacious biotherapeutics and vaccines. Endosomal processing of a protein antigen followed by presentation on major histocompatibility complex (MHC) class II constitute necessary steps in the induction of CD4+ T cell immune responses. Current preclinical methods for assessing immunogenicity risk consist of in vitro cell-based assays and computational prediction tools. Cell-based assays are time and labor-intensive while in silico methodologies have limitations. Here, we propose a novel cell-based assay capable of investigating an antigen’s endosomal processing and MHC class II presentation capabilities. This novel assay relies on competition between epitopes for MHC class II binding and employs labeled soluble T cell receptors (sTCRs) as detectors of epitope presentation.

KEY WORDS: antigen presentation; APC; HLA-DR; MHC; TCR.

INTRODUCTION

Development of antigen-specific immunological memory is critical in establishing long-lived protection against infections. Immunological memory, including the generation of high affinity matured antibodies, is a multi-step process dependent on antigen-specific T cell help (1). To initiate this process, antigen presenting cells (APCs) uptake and digest protein antigen into linear epitopes. These epitopes are loaded on to major histocompatibility complex (MHC) class II and presented to T cell receptors (TCRs) (2). TCR recognition by antigen-specific CD4+ helper T (Th) cells leads to cellular activation and expansion (1, 3). Activated Th cells provide help to antigen-specific B cells resulting in the generation of antibody secreting plasma cells and long-lived memory B cells (1, 4).

Antibody-specific immunity is critical in combating infection and developing efficacious vaccines. In unwanted cases, antigen-specific immunity can develop in response to protein therapeutics leading to the generation of anti-drug antibodies (ADA) that can impact both drug efficacy and tolerability (5, 6). To assess the likelihood of generating an antigen-specific immune response, several analytical tools that employ primary human immune cells have been developed. Dendritic cell (DC)-T cell assays measure CD4+ T cell proliferation in response to antigen pulsed matured autologous DCs and peripheral blood mononuclear cell (PBMC) assays quantify cytokine response and CD4+ T cell activation in antigen pulsed heterogeneous cell cultures. These assays can detect antigen-specific CD4+ T cell responses in naïve donors which may correlate with drug immunogenicity. For robustness and wide HLA coverage, multiple human blood donors are needed for analysis, making these assays both labor intensive and time consuming (7, 8). Additional limitations are the inability of these assays to easily distinguish immunogenic epitopes within a full protein.

To specifically determine epitopes presented on MHC, an LC-MS/MS approach referred as MHC-associated peptide proteomics (MAPPs) assay is utilized. In this approach, APCs are pulsed with test antigen, and MHC II-associated peptides are immunoprecipitated. Peptides subsequently eluted from MHC II molecules are then sequenced by LC-MS/MS (9, 10). While MAPPs can accurately identify epitopes in the context of a full protein, sensitivity varies depending on the protocol (11). Use of primary human cells requires the inclusion of many donors for diverse HLA coverage and further complicates outcomes. These assays also require high number cells which impose practical limits on number conditions analyzed for each compound (9).

To address the inability to test large numbers of candidates/compounds in cell-based assays, high-throughput in silico algorithms are utilized. These predictive methods are trained on large sets of MHC class II binding data and can specifically target areas of high immunogenicity risk within the protein sequence (12). Despite the evolving robustness of in silico methods, many of these tools are unable to predict cellular processing and trafficking of peptides especially for
MHC-II epitopes. Integration of MAPPs data and antigen cleavage data has improved the accuracy of these tools (13–15). However, predictions are limited by the breadth and accuracy of data sources used for training these tools.

Here we propose the development of a novel cell-based assay that takes advantage of the competition between epitopes to bind MHC class II and employs labeled soluble T cell receptors (sTCRs) to detect peptide-MHC complexes (pMHC) (16, 17). Soluble TCRs have been proposed as a novel immunotherapy engineered for targeted delivery of oncolytics and immunomodulatory agents (18, 19). However, this is the first example where sTCRs are employed to assess antigen presentation. To overcome limitations of MHC-II heterogeneity in primary cell-based assays, B lymphoblastoid cells (B-LCL) mono-allelic for the HLA-DRB1 locus are employed as APCs. This assay will determine MHC class II presentation based on the concept that epitopes restricted by the same MHC molecule compete for presentation (16). Mono-allelic APCs will be pulsed with a reference epitope alone or in the presence of a test compound. Labeled sTCRs recognizing the reference epitope/MHC complex (e:MHC) will be used to detect reference epitope presentation on MHC class II. Decreased sTCR/e:MHC binding signal in the presence of a test compound will indicate test compound presentation on MHC class II. We demonstrate that assay-detected epitopes from test compounds correlated with in silico predictions and standard MAPPs assay analysis. The assay proposed here has diverse applications ranging from vaccine development, neoantigen evaluation, and immunogenicity analysis and de-risking.

MATERIALS AND METHODS

Soluble T Cell Receptor Expression and Purification

Expression and purification of soluble T cell receptors were performed by GenScript Biotech (Piscataway, NJ) using the following conditions.

Preparation of Influenza HA sTCR. Expi293F cells were transfected with pUC57 plasmid containing HA sTCR using ExpiFectamine™ 293 reagent and cultured in serum-free Expi293™ Expression Medium (Thermo Fisher Scientific) at 37°C at 8% CO2 on an orbital shaker. Cell culture supernatants were collected on day 6 for purification.

Preparation of Stable Cell Lines Expressing HIV gag p24 sTCR. CHO-K1SP cells were transfected with pUC57 plasmid containing p24 sTCR twice and expression was detected 48 h post-transfection by Western blot. Transfected cells maintained in selection medium (CD CHO medium, 25 μM MSX, 200 μM anti-clumping agent) were divided into pools and screened by Dot-Blot and SDS-PAGE. Single clones from the top 12 cell pools were obtained by limiting dilution. The top 10 clones were maintained in Dynamis medium with 200x anti-clumping agent and incubated at 37°C, 5% CO2. Feed B solution was added on day 4, 6, 8, 10, and 12 and glucose was added to maintain the concentration range from 4 to 6 g/L. Supernatant from the top producing clone was harvested on the last day for SDS-PAGE analysis and purification.

Purification. Filtered cell culture media were loaded on to an affinity purification column and two-step purification was performed using Anti-DYKDDDDK G1 affinity resin (GenScript cat# L00432) and HiLoad16/600 Superdex200pg. The eluted fractions were pooled and exchanged to the final formulation buffer (PBS, pH 7.2). The purified protein was analyzed by SDS-PAGE, Western blot, and SEC-HPLC analysis to determine molecular weight and purity. The concentration was determined by Bradford assay with a BSA standard (Thermo Fisher cat# 23236). For all lots, purity of the sTCR was above 80% as measured by SEC-HPLC, and endotoxin level below 2.1EU/mg as measured by LAL Endotoxin Assay Kit (GenScript cat# L00350).

Synthesis of Omomyc Dimer

The Omomyc protein sequence was assembled by solid phase synthesis on a Microwave Liberty Blue synthesizer (CEM, Matthews, NC). Released and deprotection of Omomyc from its solid support were achieved by stirring the peptidyl-resin in TFA (trifluoroacetic acid)/water/TIS (triisopropylsilane)/DTT (dithiothreitol) (93:3:3:1, v:v:v:w; 10 mL/g of peptidyl-resin) for 2–3 h at room temperature. The crude protein product was precipitated and washed with diethyl ether. It was then purified to > 90% purity using preparative mass-triggered HPLC on C18 Reverse Phase columns and linear gradients of acetonitrile in water, both buffered with 0.1% TFA. The HPLC fractions containing a pure protein product were pooled and lyophilized to yield Omomyc as a white solid in TFA salt form. Conversion of the TFA to acetate salt of the protein was performed using a HiTrap Q HP strong anion exchange column (5 mL; GE Healthcare). Purified Omomyc monomer was dimerized via disulide formation by stirring it in a solution of Water (48.0 mL) and 10X TBS (pH 7.4) (48 mL), adjusted to pH 7.8 by addition of 1.2 mL of 5 M NaOH, followed by addition of DMSO (21.0 mL). HPLC fractions containing a pure protein product were pooled and lyophilized to yield Omomyc dimer as a white solid in TFA salt form. Conversion of the TFA to acetate salt form of the protein was performed using a Fast Flow Q Sepharose gel (42 mL, 0.2 mmol/mL; acetate form) (strong anion exchange, GE Healthcare).

B Lymphoblastoid Cell Culture

Mono-allelic B lymphoblastoid cells (B-LCL) were purchased from the Fred Hutchinson Cancer Research Center Cell Bank (Seattle, WA). The following cell lines were used: IHW ID IHW01124 HLA-DRB1*01:01; IHW01152 HLA-DRB1*03:01; IHW01111 HLA-DRB1*04:01; IHW09050 HLA-DRB1*07:01; IHW09036 HLA-DRB1*11:01; and IHW01055*15:01. Cell lines were maintained in RPMI-1640 media (Sigma R0883)
supplemented with 2 mM L-Glutamine (Sigma G7513), 1 mM sodium pyruvate (Sigma S8636), and 15% fetal bovine serum (FBS) (Sigma F4135) at 37°C in a 5% CO2 incubator.

**B Lymphoblastoid Antigen Pulse**

B-LCLs were cultured in X-Vivo 15 media (Lonza cat# 04-7440Q) supplemented with 10% human AB serum (HS) (Sigma cat# H3667) and 5 μg/mL of the human TLR9 ligand CpG ODN 2006 (InvivoGen cat# trl-2006) for 24 h. The following day antigens at the indicated concentrations were added to cells plated to a 6-well dish at 0.5 × 10^6 cells in 2 mL of X-Vivo 15 media with 1% human Ab serum and 5 μg/mL of CpG ODN 2006. After 24-h incubation, cells were collected for staining with soluble T cell receptors.

Peptides for antigen pulse were generated as a custom reagent by Thermo Fisher Scientific (Waltham, MA). Influenza A H5N1 HA protein (A/Vietnam/1194/2004) was purchased from Sino Biological (cat# 11062-V08H1).

**Primary Dendritic Cell Differentiation and Antigen Pulse**

Two hundred milliliters of whole blood was collected in BD Vacutainer CPT tubes (cat# 362753) from healthy human donors in the MSD Volunteer Research Donor Program. CD14+ cells were isolated from PBMCs by positive selection using CD14 Microbeads (Miltenyi Biotec cat# 130-050-201). CD14+ cells were plated to non-adherent 6 well plates at 3 × 10^6 cells/mL in 3 mL of X-Vivo 15 media, 10% autologous human serum, 1% PennStrep (Sigma cat# P4333), IL-4 (1000 IU/mL, R&D cat# 204-IL-010/CF), and GM-CSF (1000 IU/mL, R&D cat# 7954-GM-050/CF). Cytokines were replenished on days 2 and 4. Antigens were added to cells on day 4, and DCs were matured with a pulse of LPS (1 μg/mL, Sigma cat# L2880). Cells were harvested on day 5 for analysis, and markers for mature DCs were validated by flow cytometry.

**Soluble T Cell Receptor Staining and Flow Cytometry Analysis**

Cells were washed with 5 mL of FACS wash buffer (PBS, 1% HS, 2% FBS) and transferred to a 96-well plate for staining. sTCRs at a final concentration of 500 nM diluted in FACS wash buffer were added to cells for 40 min at 37°C in a final volume of 100 μL. Following sTCR incubation, cells were washed with 0.5 mL of PBS once. BioLegend Zombie Aqua cell viability stain (cat# 423102) diluted 1:500 in PBS was incubated with cells for 10 min at room temperature. The cell viability stain was not washed off to allow for detection of loss of cell viability during staining. sTCR staining buffer containing mouse anti-His PE secondary antibody (clone J095G46, BioLegend cat# 362603) at a 1:20 dilution and FeR blocking reagent at a 1:100 dilution (Miltenyi Biotec cat# 130-059-901), and cells were incubated at room temperature for an additional 20 min. Following secondary antibody incubation, cells were washed with 0.5 mL of FACS wash buffer and cells were resuspended in a final volume of 300 μL of FACS wash buffer for analysis. Cells were analyzed immediately using a BD-LSR Fortessa. **Staining of APC Markers**. Following 24 h antigen pulse cell surface marker expression of HLA-DR, CD80, and CD86 was analyzed by flow cytometry. Cells were blocked for 15 min at 4°C in FACS wash buffer containing a 1:100 dilution of FeR blocking reagent. Mouse anti-human CD86 (BV421) (clone 2331, BD Biosciences cat# 562433) at a 1:100 dilution, mouse anti-human CD80 (PE) (clone L307.4, BD Biosciences cat# 557227) at a 1:20 dilution, and mouse anti-human HLA-DR (APC-H7) (clone G46-6, BD Biosciences cat# 561358) at a 1:100 dilution were added to cells in a final volume of 100 μL for 20 min at 4°C in FACS wash buffer with FeR blocking reagent. As a control, cells were incubated with isotype control Abs at the same dilution: BV421 mouse IgG1, k (clone X40, BD Biosciences cat# 562438), PE mouse IgG1, k (clone MOPC-21, BD Biosciences cat# 555749), and APC-H7 Mouse IgG2a, k (clone G155-187, BD Biosciences cat# 560897). Following staining, cells were washed with FACS wash buffer and resuspended 300 μL of FACS wash buffer containing 7AAD viability dye at 1:100 dilution for analysis (ebioscience cat# 00-6993). Cells were not fixed and analyzed immediately using a BD-LSR Fortessa.

Data was analyzed using FlowJo (version 10). Cells were gated according to their forward and side scatter properties and single cells were selected by comparison of FSC-A to FSC-H. Dead cells were excluded from analysis. A representative gating strategy can be found in S2 Fig.

**Chloroquine Inhibition**

Chloroquine (Sigma cat# C6628) at 80 μM was added to cells 1 h prior to addition of antigens. Cells were cultured in the presence of chloroquine and antigens for 24 h before sTCR binding analysis. As the effects of chloroquine are quickly reversible, chloroquine at 20 μM was added to the FACS wash buffer in treated cells.

**MAPPs Assay**

A total of 1 × 10^7 cells per sample were submitted to Caprion Biosciences Inc. (Montreal, Quebec, CA) for MAPPs analysis. MHC class II immunoprecipitation and LS/MS-MS analysis were performed by Caprion Biosciences Inc. according to the method previously published by Mommen et al. (9). Primary DCs and B-LCLs were pulsed with Omomyc (10 μg/mL, 0.5 μM) under conditions for antigen pulse described above. Mock (vehicle control) cells were included as a control.

**In Silico Predictions**

The 90AA long Omomyc sequence was divided into 9 peptides of 25AA in length with 10AA overlap. MHC Class II binding affinity of these peptides to HLA-DRB1*11:01 or HLA-DRB1*15:01 was analyzed using the following algorithms: IEDBcons (20), Epimatrix (Epivax) (21), NN_Align (22), or NetMHC3.2 (23). The number of unique core binding epitopes within the top 5% binding rank for Epimatrix and top 10% binding rank for IEDBcons, NN_Align, and NetMHC3.2 was considered as HLA-DRB1 binders.
Data Presentation and Statistical Analysis

Graphpad prism (version 8) was used for analysis. In Fig. 2, data is expressed as the fold MFI of sTCR fluorescence in peptide-treated cells of mock-treated cells: MFI_{(reference epitope) / median MFI_{mock)}. One-way ANOVA with multiple comparisons was used to determine significance, $p > 0.05$ not significant ($ns$). In Figs. 3 and 4b, data is expressed as the percent sTCR fluorescence signal (MFI) relative to cells pulsed with reference epitope: MFI_{(test compound + reference epitope) / median MFI_{(reference epitope alone)}. In Fig. 5, data is expressed as the percent sTCR positive relative to cells pulsed with reference epitope: $\%$sTCR+_{(test compound + reference epitope) / median $\%$sTCR+_{(reference epitope alone)}. One-way ANOVA with multiple comparisons was used to compare reference peptide to test compound + reference peptide $p > 0.05$ not significant ($ns$). Bars represent median and IQR of triplicate samples.

RESULTS

A Novel In Vitro MHC Class II Presentation Assay Employing Soluble T Cell Receptors

We describe a novel cell-based assay for MHC class II epitope processing and presentation that employs fluorescently labeled sTCRs for detection of pMHC. Antigen-specific sTCRs have been developed as novel therapeutics for immune modulation and targeted drug delivery (18, 19, 24). We employed the method established by Walseng et al. for the generation of soluble T cell receptors in mammalian cells (17). For expression, a vector containing TCR alpha and beta chains with the transmembrane and cytoplasmic domain removed was transfected into mammalian cells (17) (Fig. 1a). Production of mammalian cells offers benefits over bacteria in allowing for post translational modifications. A picornavirus ribosomal skipping sequence 2A was included between TCR alpha and beta chains allowing for equimolar production of both chains and higher efficiency of pairing (25–27).

Two soluble TCRs were generated. Sequences of the variable and constant domain were utilized from the F24 TCR binding the HIV-1 capsid epitope Gag293 (p24) in the context of HLA-DRB1*01:01, HLA-DRB1*11:01, and HLA-DRB1*15:01 (28). A second TCR (F11) recognizing the influenza hemagglutinin epitope HA306 (HA) in the context of HLA-DRB1*01:01 was also expressed (29). Expression of TCR construct in mammalian cell lines produced a secreted protein dimer that separated under reducing conditions (S1 Fig). Peptides of the HIV-capsid epitope Gag293 (p24) and influenza hemagglutinin epitope HA306 (HA) will serve as reference epitopes since they can be detected by the F24 and F11 sTCRs respectively when presented by MHC-II.

To assess if any peptide or protein contains epitopes, defined as sequences presented on a specific HLA-DR allele, B lymphoblastoid cell lines (B-LCL) mono-allelic for the HLA-DRB1 locus were pulsed with the peptide or protein of interest (referred as test-peptide and test-compound respectively) and a reference epitope (Fig. 1b). A labeled sTCR specific to the eMHC was used to quantitate reference epitope presentation by flow cytometry. Since epitopes restricted by the same MHC molecule compete for binding (16), efficient presentation of the test-peptide will reduce the level of eMHC detected by the labeled sTCR (Fig. 1b). Comparing the signal obtained when cells are pulsed with a reference epitope alone versus cells pulsed with a reference epitope and a test-peptide will indicate if the test-peptide contains epitopes presented on the HLA-DR allele (Fig. 1b).

sTCRs Specifically Bind the Reference Epitope in Context of Its Cognate MHC Class II

To test if the sTCRs can bind the eMHC complex, B-LCLs were incubated with increasing concentrations of influenza HA reference epitope or HIV p24 reference epitope for 24 h. Fluorescently labeled sTCRs were employed to assess epitope presentation. Relative to mock (vehicle control), a dose-dependent increase in bound F24 p24 sTCR was observed in HLA-DRB1*11:01 and *15:01 when pulsed with the p24 epitope (Fig. 2a). Cross-recognition of the p24 epitope by F24 TCR in the context of multiple HLA-DR alleles has been previously demonstrated (28). Similarly, the F11 HA sTCR bound HLA-DRB1*01:01 and *04:01 cells when pulsed with the HA epitope (Fig. 2b). Even though F11 is reported to bind the HA epitope in the context of HLA-DRB1*01:01 only (29), F11 cross-recognized the HA epitope when presented on HLA-DRB*04:01 (Fig. 2a). Similar to cross-recognition of the F24 sTCR in the context of multiple HLA types, cross-recognition of HA epitope in the context of HLA-DRB1*01:01 and *04:01 by the HA1.7 TCR has also been observed (30). It is therefore not surprising that the HA-specific F11 sTCR was able to recognize the HA epitope in the context of both HLA-DRB*01:01 and *04:01 in our assay. Because mono-allelic cell lines are used, cross-recognition will not confound the output of this assay. Furthermore, the F11 sTCR was also able to detect the HA epitope presentation on HLA-DRB1*01:01 expressing primary DCs (S3A Fig).

To investigate the specificity of sTCR/eMHC binding, B-LCL were pulsed with either the p24 epitope or the HA epitope and stained with sTCRs (Fig. 2c and d). F24 sTCR stained HLA-DRB1*11:01 and HLA-DRB1*15:01 when pulsed with the p24 epitope only (Fig. 2c). F11 sTCR stained HLA-DRB1*01:01 and *04:01 expressing cells when pulsed with the HA epitope only (Fig. 2d). A low signal was detected when F24 sTCR was used to stain HA-epitope pulsed HLA-DRB1*15:01 expressing cells. However, the F24 was significantly better at staining p24-epitope HLA-DRB1*15:01 expressing cells (Fig. 2c). Both the p24 and the HA epitopes are promiscuous epitopes, able to bind multiple HLA-DR1 alleles, and therefore can be presented on all HLA types tested (28, 30).

The concentration of peptide input required for significant sTCR binding varied between cell lines, ranging from a
minimum detection limit of 1 μM for p24 in the HLA-DRB1*11:01 line to 400 μM HA in the HLA-DRB1*01:01 line (Fig. 2a and b). F24 TCR binds with the highest affinity to HLA-DRB1*11:01 expressing cells, followed by HLA-DRB1*15:01 and HLA-DRB1*01:01 expressing cells (28). Our analysis supports these findings, with a median 21-fold increase in binding relative to mock in HLA-DRB1*11:01 expressing cells when pulsed with 100 μM of the p24 epitope, versus a 2.5-fold increase in binding relative to mock observed when HLA-DRB1*15:01 expressing cells were pulsed with the same amount of the p24 epitope (Fig. 2a). Furthermore, HLA-DR expression levels were consistent between the different cell lines and were not affected when these cells were pulsed with the different epitopes (S4 Fig). As TCR/MHC and peptide affinity differs between HLA types, assay conditions such as concentration of the reference epitope and the sTCR will need to be optimized for each HLA allele and sTCR.

**Competition-Mediated Detection of Test Epitope Presentation on MHC-II**

To investigate the assay’s ability to detect MHC class II presentation of a test antigen by competing out the binding of the reference epitope, APCs were pulsed with the reference epitope alone or in the presence of an increasing concentration of test compound. APCs were subsequently stained with sTCR specific for the eMHC (Fig. 3a). F24 was used to stain B-LCLs pulsed with the p24 epitope as reference epitope alone or with an increasing concentration of the HA epitope as test peptide (Fig. 3a). Likewise, F11 was used to stain B-LCLs and primary human DCs pulsed with the HA epitope as a reference epitope alone or with an increasing concentration of p24 epitope as test peptide (Fig. 3b and S3BFig). A significant dose-dependent reduction in reference epitope detection was observed indicating successful presentation of the test peptide (Fig. 3a and b). This is expected since both epitopes (p24 and HA) employed in this experiment are able to bind multiple HLA-DR alleles (28, 30).

We next investigated if this assay could detect epitopes in a context of a full protein antigen. Omomyc a 90AA dimer (31) or influenza H5N1 HA a 522AA trimer (32) was used as test compounds in an experiment employing the p24 epitope as a reference epitope and the F24 sTCR (Fig. 3c and d). Both Omomyc and H5N1 HA significantly reduced sTCR detection of the p24 reference antigen (Fig. 3c and d). This indicates that both proteins contain epitopes able to bind the specific HLA-DR alleles expressed by BLC-Ls, and these cell lines are capable of processing and presenting epitopes in the context of full proteins and not just peptides.
Following the observation that the Omomyc protein antigen contains HLA-DR binding epitopes (Fig. 3b), a strategy of overlapping peptides was devised to map these epitopes. Nine peptides of 25AA in length with 10AA overlap were generated to cover the length of the Omomyc protein (Fig.4a). B-LCLs were pulsed with p24 reference epitope alone or in combination with each of the 9 overlapping Omomyc peptides and stained with the F24 sTCR (Fig. 4b). Reduced sTCR/e rMHC binding was observed in the presence of peptides spanning AA 27–70 in HLA-DRB1*11:01 expressing cells, and AA 36–61 in HLA-DRB1*15:01 expressing cells indicating the presence of MHC class II binding epitopes (Fig.4b).

To compare the output of this assay with in silico epitope prediction tools, we analyzed the Omomyc sequence for peptide/MHC class II binding using four MHC-II binding prediction algorithms: IEDBcons (33), Epimatrix (21), NN_Align (22), and NetMHC3.2 (23). The number of unique core binding epitopes was quantified using the best practices for each algorithm and indicated as “hits” which ranged from 0 to 4 epitopes (Fig. 4c). When comparing outputs, consensus between all 4 prediction algorithms was observed only for peptides detected by the in vitro sTCR assay (Fig. 4c). In silico tools predicted the presence of additional epitopes not presented in the in vitro sTCR assay, this could be due to overprediction associated with in silico immunogenicity tools as reported before (34). Next, we compared epitope presentation delineated by the sTCR-based assay versus the standard MAPPs in vitro APC assay. To perform this analysis, four B-LCL lines expressing distinct HLA-DRB1 alleles (HLA-DRB1*01:01; *03:01, *07:01, or *15:01) were pulsed with full length Omomyc for 24 h (Fig. 4d). Peptide sequences bound to MHC were eluted and sequences were determined by mass spectrometry. MHC class II bound sequences determined by MAPPs were compared with Omomyc peptides detected to bind in the sTCR assay (Fig. 4b). Epitopes were identified by MAPPs spanning AA position 29–77. This region overlapped with peptides detected by the sTCR assay which covered AA position 27–70 for HLA-DRB1*11:01 and AA 36–61 for HLA-DRB1*15:01 (Fig. 4d). Although HLA types varied between assays, the detected core binding epitopes were consistent. These core epitopes located at AA...
positions 43, 46, and 53 represent promiscuous epitopes, binding with high affinity to all included HLA types (Fig. 4b and d). MAPPs detected an epitope spanning AA position 70–77 in HLA-DRB1*03:01; *07:01; and *15:01 cells (Fig. 4d). A non-significant trend of decreased e, MHC/sTCR binding was observed when HLA-DRB1*15:01 expressing cells were pulsed with peptides spanning AA region 63–90 (Fig. 4b). Finally, we compared MAPPs detected epitopes between B-LCLs and primary DCs when pulsed with the full Omomyc protein. Both B-LCLs and primary DCs presented

Fig. 3. Competition-mediated detection of test epitope presentation on MHC-II. a F24 staining of 1055 HLA-DRB1*15:01 and 9036 *11:01 pulsed with p24 (reference epitope) alone or with an increasing concentration of HA test antigen (test peptide). b F11 staining of 1111 HLA-DRB1*04:01 and 1124 *01:01 cells pulsed with HA (reference epitope) alone or with an increasing concentration p24 test antigen (test peptide) (c, d). F24 staining of 9036 HLA-DRB1*11:01 and 1055 *15:01 cells was pulsed with p24 alone or in the presence of Omomyc (e) or influenza A H5N1 HA protein (d). The data on Y-axis is expressed as the percent sTCR fluorescence signal (MFI) relative to cells pulsed with reference epitope: MFI_{test compound} / median MFI_{reference epitope alone}. The one-way ANOVA with multiple comparisons was used to determine significance, p > 0.05 not significant (ns). For statistical comparisons in a and b, samples treated with test peptide were compared to no test epitope; for c and d, samples treated with reference peptide + protein were compared to the reference epitope alone.
peptides with the same core binding epitope centered around AA position 46 that was detected by the sTCR assay (Fig. 4b).

Minimum Peptide Length Required for Endosomal Processing

A recent study has highlighted discrepancies in the detection of MHC-II epitopes when APCs were pulsed with peptides versus a full protein in the context of MAPPs assays (35). This discrepancy was associated with endosomal processing which may not be required for an epitope to bind MHC-II in the context of a small peptide resulting in a false positive. Cells were treated with chloroquine, which inhibits endosomal activity by increasing endosomal/lysosomal pH and inhibiting the activity of acidic proteases (36). Treatment of B-LCLs with chloroquine inhibited epitope competition of the p24 reference epitope when B-LCLs were pulsed with full length Omomyc whereas in the absence of inhibitor, Omomyc-associated epitopes competed p24 binding to HLA-DRB1*11:01 and *15:01 (Fig. 5a). Treatment of B-LCLs with chloroquine did not decrease p24 presentation and detection of the p24/HLA-DR complex by the F24 sTCR (S6 Fig).

To determine the minimum length of peptide that is inhibited by chloroquine, B-LCLs were pulsed with the p24 reference epitope and Omomyc peptides of increasing lengths (15AA to 90AA) in the presence or absence of chloroquine.
Chloroquine had no impact on the ability of Omomyc peptides to compete with the reference epitope when the competing peptide length was shorter than 70AA (Fig. 5b). For peptide length ranging from 80 to 90AA, chloroquine was able to inhibit the ability of the encoded epitopes to compete the reference epitope (Fig. 5b compare top and bottom panel). In the absence of chloroquine, 80–90mer Omomyc peptides reduced sTCR binding to p24/HLA-DR1 complex by 50–60% (Fig. 5b top panel), where in the presence of chloroquine, no significant reduction of sTCR binding to p24/HLA-DR1 was observed.

**DISCUSSION**

Here we present a proof of concept for a novel *in vitro* assay capable of assessing MHC-II antigen processing and presentation and promises an accurate, fast, and cost-effective turnaround. This assay employs fluorescently labeled soluble TCRs recognizing MHC class II/epitope complex as a novel approach for assessing antigen presentation. In addition, epitope competition for MHC binding minimizes the number reagents required for this assay, as only a single sTCR and reference epitope are needed for each MHC supertype. We propose to employ APCs mono-allelic for the MHC locus to eliminate the ambiguity that arises from the co-expression of multiple MHC alleles; however, this assay was also able to assess antigen presentation in multi-allelic human primary DCs (S3A Fig).

An integral component of this assay is the concept of epitope competition for MHC antigen presentation (16). Peptide sequences as small as 15AA in length and full proteins up to 522AA were able to compete with the reference antigen for MHC class II presentation (Fig. 3a–c). The concept of competition was described by Adorini et al., who detected decreased antigen-specific T cell activation when hen egg-white lysozyme-transfected APCs were cultured with exogenous peptides (16). Sensitivity of sTCR in detecting reference antigen was highly dependent on the affinity of the TCR/eMHC complex. The detection limit for the high affinity F24 TCR was 1 μM of p24 when HLA-DRB1*11:01 expressing cells were used; on the other hand, F11 sTCR required 400 μM of HA peptide in HLA-DRB1*01:01 expressing cells. Low sensitivity can be easily overcome by selecting sTCR with a higher affinity for eMHC complex or multimerizing the sTCR to increase the avidity of eMHC/sTCR interaction (37, 38). Alternatively, high affinity TCR mimic antibodies or TCR engagement reporter cells can be used (39–43). The competition-based assay described here enabled the detection of competing epitopes, despite differences of sTCR sensitivity to the reference antigen. These analyses were able to consistently detect competition by high affinity antigens and epitopes. Furthermore, since the strength of MHC peptide binding affinity was correlated with immunogenicity (44, 45), the ability to detect high affinity epitopes is integral in identifying immunogenicity risks.

When comparing epitope presentation on B-LCLs versus primary DCs, we observed high alignment of presented epitopes between the sTCR assay and the standard MAPPs assay (Figs. 4b and S3). The application of B-LCLs as APCs capable of inducing antigen-specific CD4+ T cell activation has been widely established (46–50). While B-LCLs are utilized as the model APC in our study, the assay described here can be easily adapted to other APCs. Macrophage and DC cell lines as well as primary human APCs could be easily employed as epitope competition and detection by sTCR binding are not B-LCL dependent.

Similar binding specificities enabled clustering of HLA-DR alleles into eight super types (44). In this study, we employed four cell lines that represent four HLA II super types: HLA-DRB1*01:01, HLA-DRB1*04:01, HLA-DRB1*11:01, and HLA-DRB1*15:01. To achieve broad coverage, inclusion of additional HLA-DR super types as well as HLA-DQ and HLA-DP alleles will be required. The
The sTCR assay described here is not allele nor class specific, and the binding profile of any HLA can be determined as long as a sTCR specific for a reference epitope can be generated.

The B-LCLs employed here were capable of presenting epitopes from full-length proteins (Fig. 3b). Antigen processing by B-LCLs was inhibited by chloroquine, an inhibitor of endosomal proteases used as a correlate for endosomal processing (Fig. 5) (36). To map the specific position of epitopes within a protein sequence, a strategy of overlapping peptides is required. The sTCR assay enabled an overlapping peptide strategy spanning the entire Omomyc length to map the position of epitopes within the Omomyc protein (Fig. 4a and b). The sTCR assay detected epitopes which overlapped with those detected standard MAPPs assay (Fig. 4d). Furthermore, the sTCR assay was able to identify overprediction implied by in silico tools (Fig. 4e). MAPPs detected presentation of Omomyc AA 61–77 in 1055 HLA-DRB1*15:01 which was not detected by the sTCR assay. This discrepancy could be attributed to weaker affinity to the sTCR/peptide complex to p24/HLA-DRB1*15:01 leading to decreased sensitivity of detection. Methods for optimizing sTCR/reference peptide affinity could also further increase accuracy.

A caveat with employing peptides in cell-based antigen presentation assays is highlighted by Sorde et al., where a MAPPs assay identified an epitope when presented in the context of a peptide but not the full parent protein (35). This observation implies that unlike full length proteins, endosomal processing is not required for peptide encoded epitopes. To overcome this limitation, we identified the minimal peptide size requiring endosomal processing prior to epitope binding in the B-LCLs. Chloroquine decreased the ability of full length Omomyc to compete with the reference epitope (Fig. 5). Chloroquine treatment increased the presentation of the reference p24 epitopes which is a 15mer peptide (S6 Fig). When cells were pulsed with peptides of increasing length, pretreatment of cells with chloroquine affected the presentation of epitope encoded in peptide of 80–90AA in length (Fig. 5b). This indicates that at least for the B-LCLs used in this study, peptides with a minimum of 80AA in length should be employed to ensure endosomal processing.

CONCLUSIONS

This work describes soluble T cell receptors that can recognize an epitope/MHC class II complex in the context of four major MHC classes: HLA-DRB1*01:01, *04:01, *11:01, and *15:01. In summary, we describe a novel in vitro assay for analysis of antigen processing and presentation in the context of a specific MHC class II molecule. The described assay has the potential for detecting antigen presentation by multiple APC types within the context of both MHC classes I and II.

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AUTHOR CONTRIBUTIONS

CT and JP performed the experiments. CT, VJ, and JM contributed to experimental design, data discussion, and manuscript preparation. CT and JM oversaw the entire project. All authors read and approved the final manuscript.

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SUPPLEMENTARY INFORMATION

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