The invention relates to a method for sorting, staining or detecting T cells of the immune system using a protein kinase inhibitor.
Use of a Protein Kinase Inhibitor to detect Immune cells, such as T cells.

The invention relates to the use of a protein kinase inhibitor to detect cells of the immune system and in particular T cells; and also to a method for the detection of cells of the immune system, and in particular T cells, comprising the use of a protein kinase inhibitor; a method for conserving reagents when detecting cells of the immune system, and in particular T cells, comprising the use of a protein kinase inhibitor; the use of a protein kinase inhibitor in cell sorting of immune cells and in particular T cells; and also a method of cell sorting immune cells and in particular T cells comprising the use of a protein kinase inhibitor.

INTRODUCTION

T-cells detect antigens in the form of peptides bound to major histocompatibility complex (MHC) molecules at the cell surface. This primary recognition event enables the orchestration of adaptive immunity and targeted destruction of transformed and pathogen-infected cells. T-cell specificity is determined by the highly variable complementarity determining regions of the T cell receptor (TCR). The TCR/peptide-MHC (pMHC) interaction is very weak and classically endures for no longer than a few seconds at physiological temperatures. However, multimerization of pMHC molecules results in cooperative interactivity at the cell surface and ensures that the binding avidity of pMHC tetramers far exceeds the sum of the contributing monomeric affinities. This avidity effect extends the binding half-life of pMHC tetramers (Laugel et al., 2005) and enables stable coherence to the surface of T-cells bearing cognate TCRs (Altman et al., 1996; Burrows et al., 2000). Consequently, pMHC tetramers have transformed the study of antigen-specific T-cells by enabling their visualization, enumeration, phenotypic characterization and isolation from ex vivo samples (Altman et al., 1996; Chattopadhyay et al., 2006). Indeed, pMHC tetramers have been used in thousands of studies in the decade since their initial inception and have spawned the formation of several commercial companies.
We have recently used a monoclonal T-cell system to examine T-cell activation and pMHC class I (pMHCI) tetramer binding with a series of altered peptide ligands that vary in their affinity for the cognate TCR by over 100-fold (Laugel et al., 2007). Importantly, cell surface topography, including TCR and CD8 density, remain constant in this system. In this controlled system, efficient staining with tetrameric pMHCI required a monomeric TCR/pMHCI affinity of $K_D <35 \mu M$; below this threshold, there was a sharp drop off in the intensity of pMHCI tetramer staining (Laugel et al., 2007). A reasonable T-cell agonist in this system bound with a $K_D \sim 250 \mu M$ and a weak agonist bound with a $K_D >500\mu M$. However, TCR/pMHCI affinities of $>200\mu M$ were not detectable by pMHCI tetramer. Thus, using normal staining procedures, pMHCI tetramers do not necessarily detect all T-cells that can respond to a particular agonist; similarly, not all agonists for a particular T-cell can be identified physically with pMHCI tetramers. These potential limitations of pMHCI tetramer staining, which likely extend across a range of multimeric valencies, have important implications for data interpretation and present a particular problem for the detection of tumour-specific or autoreactive T-cells that tend to express low affinity TCRs (Cole et al., 2007).

In this study, we demonstrate that pre-treatment with a protein kinase inhibitor (PKI) enhances multimerized MHC staining and in particular pMHCI tetramer staining of antigen-specific CD8+ and CD4+ T-cells and describe the mechanism through which these effects operate.

Protein kinases are enzymes that modify other proteins by chemically adding phosphate groups to them (phosphorylation). This involves the removal of a phosphate group from ATP and covalently attaching it to one of three amino acids that have a free hydroxyl group. Most kinases act on both serine and threonine, others act on tyrosine, and a number (dual specific) act on all three. There are also protein kinases that phosphorylate other amino acids, including histidine kinases that phosphorylate histidine residues.
Phosphorylation usually results in a functional change of the target protein or substrate by changing enzyme activity, cellular location or association with other proteins.

Protein kinase inhibitors are types of enzyme inhibitors which specifically block the action of protein kinases. There is an entire range available for purchase and the selection of a protein kinase inhibitor involves a consideration of its specificity, permeability and stability. Moreover, protein kinase inhibitors may be reversible or irreversible depending upon whether the binding action of the protein kinase inhibitor to its target or substrate can be reversed.

Statements of Invention
The invention herein described, according to a first aspect, therefore concerns the new use of at least one protein kinase inhibitor to detect cells of the immune system and in particular T cells.

Notably, the invention can be used in relation to either CD4+ or CD8+ Tcells and so reference herein to T cells includes reference to both CD4+ and CD8+ Tcells. Further, the invention has application in relation to T cells that bind, via their TCR, all forms of T cell ligands and in particular multimerized ligands.

The process of detection may, particularly but not exclusively, involve the enhanced performance of an existing system for detection such as the enhanced staining with pMHC multimers and subsequent imaging of immune cells, such as T cells. Reference herein to pMHC multimers, be they tetramers, pentamers, octomers or dextramers, or some other multimeric scaffold, includes reference to pMHC class I and pMHC class II.

Most notably still, the invention also relates to other TCR associated MHC ligands and so to other MHC-restricted T cells such as, without limitation, HLA A,
B, C or E-restricted T cells, including CD1d-restricted T cells and T cells restricted by HLA class II molecules such as HLA DR, DQ and DP alleles.

Further the invention has application in the detection or sorting of T cells from any mammalian species including, without limitation, man, mice and monkey and also the conservation of reagent when detecting or sorting cells from any of said mammalian species.

In a preferred embodiment of the invention the new use involves the enhanced detection, or staining, of cell surface markers of the said cells and in particular T cells and, more particularly, the enhanced detection or staining of protein kinase sensitive cell surface markers such as T cell surface receptors (TCR's). Yet more ideally, the new use involves the binding of more TCR's with the chosen detection or staining agent(s).

Reference herein to the term protein kinase sensitive includes reference to a cell surface marker that binds less favourably to its detecting or staining agent in the presence of protein kinase enzymes and so can be made to bind more favourably to said agent in the presence of a protein kinase inhibitor.

Whilst not wanting to be constrained or limited by any explanation of the underlying mechanism of the invention, we believe that a protein kinase inhibitor prevents the phosphorylation of cell surface markers, such as receptors, and in particular TCR's, by their relevant kinase and so prevents receptor internalization and recycling. This means more cell surface markers, or receptors, are available at the cell surface for detection or staining by a detection or staining agent, thus increasing the intensity or sensitivity of the detection or staining process.

Importantly, the benefits of the invention apply only to T-cells that express specific TCRs; PKI treatment does not result in the staining of T-cells that bear non-cognate TCRs.
In a further preferred embodiment of the invention said use comprises the step of pre-treating a population of cells to be detected with at least one protein kinase inhibitor before exposing said population to at least one agent for detecting or staining same.

Alternatively, the said use may comprise the step of treating a population of cells to be detected with at least one protein kinase inhibitor at the same time as exposing said population to at least one agent for detecting or staining same.

In either case, said protein kinase is ideally a reversible protein kinase meaning its binding action to its target site or substrate can be reversed. Ideally, said protein kinase inhibitor is a commercially available inhibitor such as, but not limited to any one or more of the following: Dasatinib, Lck inhibitor II, Wortmannin, Resveratrol, LY294002, AS604850, PI3-K alpha inhibitor 1 or PI3-K alpha inhibitor 2.

Moreover, the protein kinase inhibitor is, ideally, a tyrosine protein kinase inhibitor.

In a yet further preferred embodiment of the invention said treatment comprises the step of exposing said population of cells to said inhibitor for a period of time up to one hour. One hour may be exceeded but the beneficial effects associated with said treatment are achieved before this time interval of more than one hour expires. Indeed, the beneficial effects are achieved within seconds of exposure but we have found it advantageous to treat said population with said inhibitor for approximately 30mins at 37 °C. More preferably still, we have found it advantageous to treat said population with an amount of inhibitor up to 30OnM and most preferably less than 20OnM such as, for example only, 100, 90, 80, 70, 60, 50, 40, 30, 20, or 1OnM.
The ability of our system to detect or stain more efficiently, and so allow us to visualize cells, and in particular T cells, with either a small number of cell surface markers or TCR's; or an average number of cell surface markers or TCR's but which have low avidity for the detecting or staining agent, enables us to detect or stain cells that would otherwise be undetectable using conventional techniques. As just one example, we are now able to detect T cells bearing TCR's with low affinity for the cognate pMHC1 ligand that would otherwise be undetectable using pMHC1 monomer or tetramer staining.

Thus, in a further aspect, our invention concerns a method of detecting cells of the immune system, and in particular T cells, which comprises the use of at least one protein kinase inhibitor.

The method of detection may, particularly but not exclusively, involve the enhanced performance of an existing system for detection such as the enhanced staining with pMHC tetramers/multimers and so subsequent imaging of immune cells, such as T cells.

In a preferred embodiment of the invention the method involves the enhanced detection, or staining, of cell surface markers of the said cells and in particular T cells and, more particularly, the enhanced detection or staining of protein kinase sensitive cell surface markers such as T cell surface receptors (TCR's). Yet more ideally, the method involves the binding of more TCR's with the chosen detection or staining agent(s).

In a further preferred method of the invention said method comprises the step of pre-treating a population of cells to be detected with at least one protein kinase inhibitor before exposing said population to at least one agent for detecting or staining same.
Alternatively, the said method may comprise the step of treating a population of cells to be detected with at least one protein kinase inhibitor at the same time as exposing said population to at least one agent for detecting or staining same.

In either case, said protein kinase is ideally a reversible protein kinase meaning its binding action to its target site can be reversed. Ideally, said protein kinase inhibitor is a commercially available inhibitor such as, but not limited to any one or more of the following: Dasatinib, Lck inhibitor II, Wortmannin, Resveratrol, LY294002, AS604850, PI3-K alpha inhibitor 1 or PI3-K alpha inhibitor 2.

Moreover, the protein kinase inhibitor is, ideally, a tyrosine protein kinase inhibitor.

In a yet further preferred method of the invention said treatment comprises the step of exposing said population of cells to said inhibitor for a period of time up to one hour. One hour may be exceeded but the beneficial effects associated with said treatment are achieved before this time interval of more than one hour expires. Indeed, the beneficial effects are achieved within seconds of exposure but we have found it advantageous to treat said population with said inhibitor for approximately 30mins at 37°C. More preferably still, we have found it advantageous to treat said population with an amount of inhibitor up to 30OnM and most preferably less than 20OnM such as, for example only, 100, 90, 80, 70, 60, 50, 40, 30, 20, or 1OnM.

Using our invention we have been able to detect low avidity antigen-specific T cells. Indeed the benefits of our invention have been so striking that our invention can be used, advantageously, to conserve detecting or staining agents.
According to a yet further aspect our invention concerns a method for conserving agents when detecting cells of the immune system and in particular T cells.

In this further aspect of our invention the agents to be conserved comprise known detecting or staining agents for detecting selected cells such as T cells within a population of cells and, more specifically, antigen-specific cells such as T cells and so T cells that have TCR's specific for the agent to be used, for example, TCR's specific for an antigen presenting molecule or cell such as a pMHC: pMHC1 or pMHC1 1 or CdId etc. in the form of a monomer or multimer.

In this aspect of the invention the method involves conserving agents by the enhanced performance of an existing system for detection such as the enhanced staining with pMHC tetramers/multimers and so subsequent imaging of immune cells, such as T cells.

In a further preferred method of the invention said conserving method comprises the step of pre-treating a population of cells to be detected with at least one protein kinase inhibitor before exposing said population to at least one agent for detecting or staining same.

Alternatively, the said method may comprise the step of treating a population of cells to be detected with at least one protein kinase inhibitor at the same time as exposing said population to at least one agent for detecting or staining same.

In either case, said protein kinase is ideally a reversible protein kinase meaning its binding action to its target site can be reversed. Ideally, said protein kinase inhibitor is a commercially available inhibitor such as, but not limited to any one or more of the following: Dasatinib, Lck inhibitor II, Wortmannin, Resveratrol,LY294002, AS604850, PI3-K alpha inhibitor 1 or PI3-K alpha inhibitor 2.
Moreover, the protein kinase inhibitor is, ideally, a tyrosine protein kinase inhibitor.

In a yet further preferred method of the invention said treatment comprises the step of exposing said population of cells to said inhibitor for a period of time up to one hour. One hour may be exceeded but the beneficial effects associated with said treatment are achieved before this time interval of more than one hour expires. Indeed, the beneficial effects are achieved within seconds of exposure but we have found it advantageous to treat said population with said inhibitor for approximately 30mins at 37 °C. More preferably still, we have found it advantageous to treat said population with an amount of inhibitor up to 30OnM and most preferably less than 20OnM such as, for example only, 100, 90, 80, 70, 60, 50, 40, 30, 20, or 1OnM.

Advantageously, the enhanced detection or staining afforded by our invention only applies to cells such as T cells that express antigen specific markers, such as TCR's, this means protein kinase treatment does not result in the detection or staining of cells, such as T cells, that bear non-cognate markers or TCR's.

Indeed, we show herein that PKI treatment lowers the TCR/pMHCI affinity threshold required for pMHCI tetramer binding by as much as 5 fold, thereby allowing the binding of pMHCI tetramers to CD8+ T-cells that express TCRs with very weak affinities for pMHCI (>80-500 µM).

This simple and universally applicable procedure thereby enables the visualization of previously undetectable tumour-specific and autoreactive CD8+ T-cells with pMHCI tetramers through the preferential enhancement of low avidity interactions with TCRs at the cell surface.
Moreover, our invention comprises at least one further benefit in that it, not only increases the detection of a selected population of cells within a given population without affecting the detection or staining of cells bearing non-cognate cell surface markers, but it also reduces agent induced cell death.

In this latter respect, we have shown that the treatment with at least one protein kinase inhibitor reduces pMHC1 multimer, and in our example tetramer, induced cell death.

According to a further aspect of the invention there is therefore provided the use of at least one protein kinase inhibitor for sorting, in a population of immune cells, a viable, selected, sub-population of cells. Ideally said cells are T cells.

According to a yet further aspect of the invention there is provided a method for sorting, in a population of immune cells, a viable, selected, sub-population of cells. Ideally said cells are T cells.

In the latter two aspects of the invention the use of at least one protein kinase inhibitor helps to ensure that a selected population of cells, typically T cells, with preferred, usually high affinity, TCR's are kept viable and, following cell sorting, are available for use.

In this further aspect of the invention the method of sorting may involve the enhanced performance of an existing system for sorting such as the enhanced sorting with pMHC tetramers/multimers and so the sorting of immune cells, such as T cells.

In a further preferred method of the invention said sorting method comprises the step of pre-treating a population of cells to be sorted with at least one protein kinase inhibitor before exposing said population to at least one agent for sorting or detecting same.
Alternatively, the said method may comprise the step of treating a population of cells to be sorted with at least one protein kinase inhibitor at the same time as exposing said population to at least one agent for sorting or detecting same.

In either case, said protein kinase is ideally a reversible protein kinase meaning its binding action to its target site can be reversed. Ideally, said protein kinase inhibitor is a commercially available inhibitor such as, but not limited to any one or more of the following: Dasatinib, Lck inhibitor II, Wortmannin, Resveratrol, LY294002, AS604850, PI3-K alpha inhibitor 1 or PI3-K alpha inhibitor 2.

Moreover, the protein kinase inhibitor is, ideally, a tyrosine protein kinase inhibitor.

In a yet further preferred method of the invention said treatment comprises the step of exposing said population of cells to said inhibitor for a period of time up to one hour. One hour may be exceeded but the beneficial effects associated with said treatment are achieved before this time interval of more than one hour expires. Indeed, the beneficial effects are achieved within seconds of exposure but we have found it advantageous to treat said population with said inhibitor for approximately 30mins at 37 °C. More preferably still, we have found it advantageous to treat said population with an amount of inhibitor up to 30OnM and most preferably less than 20OnM such as, for example only, 100, 90, 80, 70, 60, 50, 40, 30, 20, or 10OnM.

Our invention will now be exemplified by reference to the following Materials and Methods and Results wherein:-

**Figure 1: Dasatinib (PKI) substantially improves pMHC tetramer staining intensity.** A. 10^5 ILA1 CTL were re-suspended in 40µl of PBS ± 5OnM dasatinib or Lck inhibitor II (PKI's) (Calbiochem), then incubated at 37°C for 30 minutes.
Cells were then stained with cognate HLA A2/ILAKFLHWL-PE (pMHC) tetramer at a final concentration of 10µg/ml for 20 minutes at 37°C, washed twice in PBS and analyzed on a FACSCalibur (BD) flow cytometer. A >10-fold increase in median fluorescence intensity (MFI) was observed after treatment with 5OnM dasatinib (Das + Tet) or Lck inhibitor II (Lck II + Tet) compared to staining without PKI pre-treatment (Tet only). B. 10^5 ILA1 CTL were treated with various concentrations of dasatinib for 30 minutes at 37°C, then stained with either HLA A2/ILAKFLHWL tetramer or the non-cognate HLA A2/ELAGIGILTV tetramer for 20 minutes at 37°C before washing with PBS. C. 10^5 ILA1 CTL were resuspended in 40µl of PBS ± the indicated concentration of dasatinib and incubated for 60 minutes at 37°C. Cells were then stained with cognate HLA A2/ILAKFLHWL-PE tetramer at a final concentration of 10µg/ml for 20 minutes at 37°C and washed twice in PBS prior to flow cytometric analysis. D. As (A), but ILA1 CTL were incubated with 5OnM dasatinib for various times prior to staining. E. As (A), but tetramer concentration was varied to stain CTL pre-treated ± 5OnM dasatinib for 30 minutes. F. 10^5 Mel13 CTL were stained with various concentrations of HLA A2/ELAGIGILTV tetramer following incubation ± 5OnM dasatinib for 30 minutes. G. 5x10^5 splenocytes from an F5 TCR transgenic Rag^+ mouse were resuspended in PBS ± 5OnM dasatinib and incubated for 30 minutes at 37°C. Cells were subsequently stained with H2-D^ß/ASNENMDAM-PE tetramer for 20 minutes at 37°C followed by anti-CD8 Cy5.5 for 30 minutes on ice prior to two washes in PBS and analysis by flow cytometry. H. 10^5 cells of the HLA DR^ß0101-restricted, influenza virus A HA_{307-319} PKYVKQNTLKLAT-specific CD4^+ clone C6 were incubated with PBS ± 5OnM dasatinib for 30 minutes at 37°C, then stained with cognate PE-conjugated tetramer for 20 minutes at 37°C. Samples were washed with PBS before flow cytometric analysis. Irrelevant tetramer was used as a negative control in all cases.

Figure 2: Dasatinib treatment preferentially increases the ability of pMHC tetramers to stain T-cells bearing low affinity TCRs. A. 10^5 ILA1 CTL were stained with 10µg/ml PE-conjugated HLA A2 tetramer folded around the 8E, 5Y,
4 L₁ index (ILAKFLHWL), 3G8T or 3G peptides for 20 minutes at 37°C following incubation ± 50nM dasatinib for 30 minutes at 37°C. For all samples, data were acquired with a FACSCalibur flow cytometer (BD) and analyzed using FlowJo software. Irrelevant tetramer was used as a negative control. B. The MFI of tetramer staining for all of the variants in the presence and absence of dasatinib displayed in (A) are plotted against the monomeric affinity of TCR/pMHCI interactions previously measured for each of these variants expressed as the dissociation constant (KD) (Table 1). Curves were fitted as described in the Materials and methods.

Figure 3: Dasatinib reduces pMHCI tetramer-induced cell death.

Increased tetramer staining in the presence of dasatinib appears to be due partly to reduced cell death. It is known that pMHCI tetramer-induced signaling can trigger cell death (Purbhoo et al., 2001; Xu et al., 2001; Guillaume et al., 2003; Cebecauer et al., 2005). Cell death induced by tetramer staining was assessed using the amine-reactive viability dye GrViD at the end of the staining procedure, which is spectrally distinct from ViViD. PBMC were stained with ViViD to identify and allow exclusion of dead and dying cells prior to the addition of pMHCI tetramer; GrViD staining was performed after pMHCI tetramer and surface antibody staining. Data were acquired on a BD LSR II flow cytometer and analyzed using FlowJo software. ViViD⁺, CD14⁺ and CD19⁺ cells were excluded from the analysis and the frequency of GrViD-positive cells was assessed in the tetramer-positive CD3⁺CD8⁺ T-cell populations. Representative flow profiles are shown here for CD8⁺ T-cells specific for the HLA A2-restricted epitopes CMV pp65₄₋₉ (NLVPMVATV) and EBV BMLF1259-267 (GLCTLVAML). The frequencies of dead cells varied depending on the tetramer used, but the frequency of GrViD-positive dead cells within the tetramer-positive population was always lower in the presence of 50 nM dasatinib. These data, together with comparable results in other systems (data not shown), suggest that the
cumulative cell death over the time course of a staining experiment could be substantially reduced by treatment with dasatinib.

**Figure 4: Dasatinib enhances the visualization of antigen-specific CD8+ T-cells in mixed cell populations.** A. Staining of HLA A2-restricted CTL lines expanded from PBMC by one round of stimulation with the influenza matrix M158-66 peptide (GILGFVFTL) or the Melan-A/Mart-1 26-35 peptide (ELAGIGILTV). Lines were stained with cognate tetramer ± pre-treatment with 50 nM dasatinib for 30 minutes at 37°C. B. Flow cytometric profiles of live CD3+ lymphocytes stained with HLA A2 tetramers folded around the EBV BMLF1259-267 (GLCTLVAML), CMV pp65 49-503 (NLVPMVATV) or Melan-A/Mart-1 26-35 (ELAGiGILTV) peptide epitopes. 2x10^6 PBMC were stained with the amine-reactive viability dye ViViD, then stained with tetramer (1 µg in minimal staining volume) ± pre-treatment with dasatinib for 30 minutes at 37°C. Cells were then stained with cell surface markers as described in the Materials and Methods; a dump channel was used to exclude dead cells, CD.14+ and CD19+ cells from the analysis. Boolean gating was carried out to exclude aggregates. Data were acquired with a BD LSR II flow cytometer and analyzed using FlowJo software.

**Figure 5: Dasatinib allows detection of autoreactive CTL.** A. CTL clone IE6, specific for the HLA A2-restricted epitope PPII15-24, was activated with either CMV pp65 49-503 or PPIis-24 peptide for 6 hours at 37°C and then assayed for TNFα production by intracellular cytokine staining as detailed in the Materials and Methods. B. Staining of CTL clone IE6 with either an irrelevant or HLA A2/PPIi 5-24 tetramer ± pre-treatment with 50 nM dasatinib for 30 minutes at 37°C. C. Representative stainings with HLA A2/PPIi 5-24 tetramer ± pre-treatment with 50 nM dasatinib for 30 minutes at 37°C. Left panels: control subject PBMCs; middle panels: type I diabetic patient PBMCs; right panels: a short-term line expanded by one round of peptide stimulation from a type 1 diabetic patient.
Figure 6: Dasatinib results in a time dependent increase in TCR and CD8 expression levels at the CTL cell surface. The ILA1 CTL clone was treated with PBS ±50nM dasatinib at 37°C and 10^5 CTL were removed from the medium at 0, 10, 30, 60, 180 and 250 minutes. CTL were subsequently stained with anti-CD8 FITC (clone SK1; BD, Pharmingen; left panel) or anti-TCR FITC (clone BMA 031; Serotec; right panel) for 30 minutes on ice, washed twice and resuspended in PBS. Data were acquired on a FACSCalibur flow cytometer (BD) and analyzed using FlowJo software.

Figure 7: Beneficial effects of dasatinib effects are not CD8-mediated. A. Melc 5 CTL were pre-treated with PBS ± 50nM dasatinib for 30 minutes at 37°C, then stained with HLA A2 DT227/8KA cognate tetramer for 20 minutes at 37°C. After washing twice, data were acquired on a FACSCalibur flow cytometer (BD) and analyzed using FlowJo software. B. Staining of HLA A2-restricted CTL lines expanded from PBMC by one round of stimulation with the Melan-A/Mart-1 26-35 peptide (ELAGIGILTV). Lines were stained with either wild type or CD8 null cognate tetramer ± pre-treatment with 50 nM dasatinib for 30 minutes at 37°C.

Figure 8: Dasatinib blocks antigen-induced TCR downregulation and tetramer internalization from the cell surface. A. MeM 3 CTL were pre-treated with PBS ± 50nM dasatinib and exposed to C1R-A2 B cells previously pulsed with 10^8M ELAGIGILTV peptide or medium alone for 4 hours at 37°C. Cells were subsequently stained with anti-TCR-FITC (clone BMA 031; Serotec) and anti-CD8-APC (clone RPA-T8; BD Pharmingen) mAbs for 30 minutes on ice, washed twice and resuspended in PBS. Data were acquired on a FACSCalibur flow cytometer (BD) and analyzed using FlowJo software. B. 10^5 ILA-1 CTL were pre-treated with PBS (i & ii) or PBS + 50 nM dasatinib (iii & iv) for 30 minutes at 37°C, then stained with 20µg/ml HLA A2/I LAKFLH WL-Alexa488 tetramer for 15 minutes at 37°C. Microscopy was performed as described in the Materials and Methods.
Figure 9: Dasatinib enhances pMHCI tetramer on-rate. Rate of HLA A2/hTERT $^{540-548}$ (ILAKFLHWL) tetramer recruitment to the cell surface of clone ILA1 is substantially enhanced following treatment of CTL with 50nM dasatinib for 30 minutes at 37°C. Subsequent to treatment with dasatinib, on-rate experiments were performed and analyzed as described previously (Laugel et al., 2007). Curves represent the following rate estimates: fast rate 0.14/min, slow rate 0.04/min (tetramer only); fast rate 0.42/min, slow rate 0.06/min (dasatinib + tetramer).

Figure 10: Dasatinib prevents down-regulation of 'empty' TCRs. Proposed model for the mechanism by which dasatinib enhances cognate tetramer staining. Dasatinib treatment prevents TCR and coreceptor down-regulation and maintains these receptors at the cell surface, thereby increasing molecular availability for further capture of pMHCI tetramer from solution.

Materials and Methods

2.1 Cells

The ILA1 CTL clone is specific for the HLA A*0201 (HLA A2 from hereon) restricted human telomerase reverse transcriptase (hTERT) epitope ILAKFLHWL (hTERT$^{540-548}$). Mel13 and Mel5. CTL clones are specific for the HLA A2 restricted Melan-A$^{26-35}$ epitope ELAGIGILTV. ILA1, Mel5 and Mel13 CD8+ cytotoxic T lymphocyte (CTL) clones were generated and re-stimulated as described previously (Whelan et al., 1999; Laugel et al., 2005). CTL were maintained in RPMI 1640 (Gibco) supplemented with 100U/ml penicillin (Gibco), 100 µg/ml streptomycin (Gibco), 10% heat inactivated fetal calf serum (FCS; Gibco), 2.5% Cellkines (Helvatica Healthcare, Geneva), 200 IU/ml IL-2 and 25ng/ml IL-15 (Peprotech). CTL lines specific for the influenza matrix protein M1$^{58-66}$ (GILGFVFTL) and Melan-A$^{26-35}$ (ELAGIGILTV) epitopes, both restricted by human leukocyte antigen (HLA) A*0201 (HLA A2 from hereon), were
generated by pulsing 6x10^6 PBMC from an HLA A*0201+ individual with cognate peptide at concentrations of 1µM and 100µM, respectively, for 1 hour at 37°C; cells were subsequently washed and resuspended in RPMI 1640 supplemented with 100U/ml penicillin (Gibco), 100µg/ml streptomycin (Gibco) and 10% heat inactivated FCS (Gibco) only. After 3 days, increasing amounts of IL-2 were added to the media reaching a maximum concentration of 20IU/ml by day 14; lines were then tested by pMHCI tetramer staining. Patient samples were collected by leukapheresis; mononuclear cells were isolated by standard Ficoll-Hypaque density gradient centrifugation and stored by cryopreservation. For autoimmune studies, blood was obtained from two HLA-A2+ patients with type 1 diabetes; both were adults, aged 27 and 31 years, and were studied within 3 months of diagnosis. Short-term lines from these diabetic patients were established as described above using the PPlι_5-24 autoantigen preproinsulin peptide (ALWGPDPAAA); this peptide bind HLA-A2 with high affinity (Arif et al., 2004). Naïve murine CTL were obtained by harvesting splenocytes from transgenic F5 Rag+ mice. A significant percentage of CD8+ T-cells within the splenic populations of these mice express the F5 TCR, which recognizes the H-2D^b-restricted influenza H17-derived nucleoprotein peptide epitope ASNENMDAM (Mamalaki et al., 1993). The HLA DR^0101-restricted CD4+ T-cell clone C6 recognizes the influenza virus A HA_{307-319} epitope (PKYVKQNTLKLAT).

2.2 Protein Kinase inhibitors
Dasatinib was synthesized as described previously (Lombardo et al., 2004). Biological activity was tested in a cell death titration assay on BA/F3 bcr-abl cells as described previously (Magnusson et al., 2002). Dasatinib was dissolved in DMSO to a concentration of 1mM and stored in aliquots at -20°C. Once thawed, these stocks of dasatinib were stored at 4°C and used within 7 days. The 1mM DMSO stock was diluted 1/10,000 in PBS on the day of experimentation to achieve a working solution of 10nM; subsequent 1/2 dilution yielded a final concentration of 50nM in cellular assays unless stated otherwise. The 10nM
working solution was always made up on the day of experimentation as the shelf-life of this solution is short (~days). Staurosporine (Biomol), Lck inhibitor II (Calbiochem), genistein (calbiochem), herbimycin A (Calbiochem), PP2 (Calbiochem) and PP3 (Calbiochem) were dissolved and stored at -20 °C in DMSO. PKIs were dissolved in PBS prior to use and tested at concentrations of 1nM, 3nM, 5nM, 10nM, 20nM, 50nM, 100nM, 250nM, 500nM and 1μM.

2.3 pMHCI tetramer manufacture
Soluble biotinylated pMHCI monomers were produced as described previously (Wooldridge et al., 2005). Tetrameric pMHCI reagents were constructed by the addition of either R-Phycoerythrin (PE)-conjugated streptavidin (Molecular Probes, Invitrogen) or Alexa 488-conjugated streptavidin (Molecular Probes, Invitrogen) at a pMHCKstreptavidin molar ratio of 4:1. Conjugated streptavidin was added to a solution of soluble pMHCI in 5 equal aliquots at 20 minute intervals and subsequently stored in the dark at 4°C.

2.4 pMHCI tetramer staining and flow cytometry: clones and splenocytes
10⁵ MeH3, Mel5 or ILA1 CTL were pre-treated at 37°C with dasatinib at a range of concentrations (0-50nM) for a series of durations up to 1 hour. Mel13/Mel5 or ILA1 were then stained with either PE-conjugated HLA A2/ELAGIGILTV or HLA A2/ILAKFLHWL tetramer, respectively, at a final concentration of 10μg/ml for 20 minutes at 37°C. The HLA DR*0101-restricted clone C6 was stained with HLA DR*0101/ PKYVKQNTLKLAT PE tetramer for 20 minutes at 37°C. After initial experiments to determine the optimal conditions of use, all subsequent experiments were performed by incubating T-cells ± 50nM dasatinib for 30 minutes at 37°C prior to tetramer staining. The HLA DR*0101-restricted clone C6 was stained with HLA DR*0101/ PKYVKQNTLKLAT PE tetramer. Subsequent to tetramer staining, CTL clones were stained with anti-human CD8-FITC (clone SK1 ; BD Pharmingen) and 7-AAD (Viaprobe; BD Pharmingen) for 30 minutes on ice then washed twice with phosphate buffered saline (PBS); the HLA DR*0101-restricted clone C6 was stained with 7-AAD (Viaprobe; BD
Pharmaingen) only. For murine CTL, 5x10^5 splenocytes were pre-treated with 5OnM dasatinib for 30 minutes at 37^0C, stained with H2-D^b/ASNENMDAM PE-conjugated tetramer for 20 minutes at 37^0C and then anti-murine CD8-Cy5.5 for 30 minutes on ice, washed twice and re-suspended in PBS. Data were acquired using a FACSCalibur flow cytometer (BD) and analyzed using FlowJo software (Treestar Inc., Ashland, OR, USA).

2.5 pMHCI tetramer staining and flow cytometry: human peripheral blood mononuclear cells

Frozen peripheral blood mononuclear cells (PBMCs) were thawed in a 37°C water bath until a small clump of ice remained and then transferred into RPMI medium containing 10% FCS, 100U/ml penicillin, 100µg/ml streptomycin, 2mM L-glutamine (Gibco) and 100U DNase/ml (Roche Diagnostics Corporation, Indianapolis, IN, USA). PBMC were washed twice in this medium and then left to rest for 2 hours at 37^0C. After 2 washes with PBS, 2 x 10^6 PBMC were stained with live/dead® fixable violet amine reactive dye (Invitrogen Corporation, Carlsbad, California, USA), washed and incubated for 30 minutes at 37^0C in PBS alone or PBS containing 5OnM dasatinib. Subsequently, PBMC in 50µl of PBS alone or PBS containing 5OnM dasatinib were stained for 20 minutes with pHLA A2 tetramers refolded around either CMV pp65_{49-503} (NLVPMVATV), EBV BMLFI_{259-267} (GLCLTVAML) or Melan-A/Mart-1_{26-35} (ELAGIGILT) peptides. After 2 washes in PBS containing 1% FCS and 0.02% sodium azide (Sigma-Aldrich, St. Louis, MO, USA), cells were stained with a selection of the following cell surface monoclonal antibodies (mAbs): (i) anti-CD3-APC-Cy7 and anti-CD8-APC (BD Biosciences, San Jose, CA, USA); (ii) anti-CD4-PE-Cy5.5 (Caltag Laboratories, purchased through Invitogen Corporation, Carlsbad, California, USA); and, (iii) anti-CD8-quantum dot (QD)705, anti-CD 14-Pacific Blue and anti-CD19-Pacific Blue, conjugated in-house according to standard protocols (http://drmr.com/abcon/index.html). The latter two mAbs were used to exclude CD14^+ monocytes and CD19^+ B cells, which can bind tetramer non-specifically,
from the analysis. Finally, cells were washed and resuspended in PBS containing 1% paraformaldehyde (PFA). Stained PBMC were acquired on a BD LSR II (Becton Dickinson Immunocytometry Systems, San Jose, CA, USA) and analyzed using FlowJo software (Tree Star, Inc., Ashland, OR, USA).

2.6 TCR downregulation and tetramer on-rate experiments
HLA A2-expressing C1R B cells (Hutchinson et al., 2003) were either pulsed with ELAGIGILTV peptide at a concentration of $10^{-6}$M for 60 minutes at 37°C or incubated in medium alone. After two washes with RPMI 1640 supplemented with 100IU/ml penicillin and 100µg/ml streptomycin, 60,000 HLA A2+ C1R cells (pulsed or unpulsed) were incubated for 4 hours at 37°C with 30,000 MeM3 CTL that had been pre-treated with PBS ± 5OnM dasatinib for 30 minutes at 37°C. Cells were then stained with anti-TCR-FITC (clone BMA 031; Serotec) and anti-CD8-APC (clone RPA-T8; BD Pharmingen) for 30 minutes on ice, washed twice and resuspended in PBS. Data were acquired using a FACSCalibur flow cytometer and analysed using FlowJo software (Treestar Inc., Ashland, OR, USA). Tetramer on-rate experiments were performed as previously described (Laugel et al., 2007).

2.7 Fluorescence Microscopy
10^5 ILA1 CTL were treated with PBS ± 5OnM dasatinib for 30 minutes at 37°C, then stained with Alexa 488-conjugated (Molecular Probes) HLA A2/ILAKFLHWL tetramer at a final concentration of 20µg/ml for 15 minutes at 37°C. Following two washes with PBS, each sample was fixed in 2% paraformaldehyde. After fixing, ILA1 CTL were re-suspended in 100µl of 2% FCS/PBS and then spun on to a microscope slide at 550rpm for 5 minutes using a cytopsin. Samples were subsequently analysed on a Leica DM LB2 (Leica Microsystems) fluorescence microscope.
2.8 IFN\textsubscript{γ} ELISpot assays

CD8\textsuperscript{+} T-cell responses to islet autoantigen were detected by IFN\textsubscript{γ} ELISpot as described previously (Chang et al., 2003) with the following modifications. PBMCs were pre-cultured at 37°C/5% CO\textsubscript{2} in single wells of 48-well plates at a density of 1 x 10\textsuperscript{6} cells in 0.5ml TC medium (RPMI 1640 supplemented with antibiotics (Invitrogen) and 10% human AB serum (PAA, Somerset, UK)) containing the test peptide at a final concentration of 10\textmu M. Control wells contained TC medium with an equivalent concentration of diluent (DMSO). After 24 hours incubation, non-adherent cells were re-suspended using pre-warmed TC medium (2% AB serum), washed, brought to a concentration of 10\textsuperscript{6} cells/300 \mu l, and then dispensed in 100\mu l aliquots into wells of 96-well ELISpot plates (Nunc Maxisorp; Merck Ltd., Poole, UK) pre-blocked with 1% bovine serum albumin in PBS and pre-coated with monoclonal anti-IFN\textsubscript{γ} (U-Cytech, Utrecht, NL). Assays were then developed according to the manufacturer's instructions; plates were dried and spots were counted using a BioReader 3000 (BioSys, Karben, Germany) and reported as total responder cells per 10\textsuperscript{6} PBMCs.

2.9 Intracellular cytokine staining assays

10\textsuperscript{6} CTL were stimulated with specific peptide at a concentration of 10\mu g/ml for 6 hours; brefeldin A (10\mu g/ml; Sigma-Aldrich) was added for the final 5 hours. Unrelated peptide (10\mu g/ml) was used as a negative control. Briefly, the cells were fixed in paraformaldehyde (2%; Sigma-Aldrich), permeabilized with saponin (0.5%; Sigma-Aldrich), and labeled with APC-conjugated anti-TNF\alpha mAb (BD Pharmingen). The cells were evaluated using a FACSCalibur flow cytometer (BD Biosciences). At least 10,000 events gated on forward and side scatter were analyzed using FlowJo software (Tree Star, Inc., Ashland, OR, USA). Corresponding isotype control antibodies were used to establish the quadrants for analysis.
2.10 Mathematical modelling of the dasatinib effect on pMHCI tetramer staining

A mathematical model that relates staining intensity to tetramer binding kinetics has been described previously (Laugel et al., 2007; van den Berg et al., 2007). Briefly, in this model, the high binding avidity of pMHCI tetramers is accounted for by assuming that tetramers can engage up to three TCR molecules, forming a cluster of one, two, or three TCRs in which association and dissociation between the TCRs and the tetramer pMHCI sites occurs at a much higher rate than diffusive escape of temporarily unbound TCRs, tending to stabilize the cluster until the tetramer happens to become disassociated with TCR molecules at all four sites. Moreover, in singlet and duplet clusters, a free TCR can associate contacts with free pMHCI sites to form, respectively, a duplet or triplet cluster. The dasatinib effect is incorporated into this model by assuming that dasatinib alters the rate at which singlet and duplet clusters recruit free TCRs. Thus, the expression for the relative staining intensity \( I \) becomes:

\[
I = I_{\text{m1}} + \Delta_1 \left( \frac{K_i/K_D}{r_0} + 6D(K_2/K_D)^3r_0^2 + \hat{\delta}_D^2(K_3/K_D)^6r_0^3 \right)
\]

where \( K_D \) is the single-site dissociation constant; \( K_i, K_2 \) and \( K_3 \) are kinetic parameters; \( I_{\text{m1}} \) and \( \Delta_1 \) relate the read-out to the number of surface bound tetramers (\( I_{\text{m1}} \) is a nuisance parameter, representing the background level); \( \hat{\delta}_D \) is the duplet recruitment enhancement factor and \( \hat{\delta}_T \) is the triplet recruitment enhancement factor. In the absence of dasatinib, we have \( \hat{\delta}_D = \hat{\delta}_T = 1 \), whereas these factors are greater than 1 if dasatinib promotes TCR recruitment. The scaled free TCR density, \( r_0 \), is implicitly defined by the scaled conservation law:

\[
I = r_0 + \left( \frac{K_i/K_D}{r_0} \right) r_0 + 2\hat{\delta}_D(K_2/K_D)^3r_0^2 + 3\hat{\delta}_T^2(K_3/K_D)^6r_0^3
\]

(from (Laugel et al., 2007)). Association kinetics has been found empirically to be described very well by the biphasic exponential model:

\[
K_i = I_{\text{m1}} + I_{\text{max, fast}} \left( 1 - \exp \left( \lambda_{\text{fast}}t \right) \right) + I_{\text{max, slow}} \left( 1 - \Theta \exp \left( \lambda_{\text{slow}}t \right) \right)
\]

where \( I(t) \) is the staining intensity at time \( t \) and \( \lambda \) are positive parameters (van den Berg et al., 2007). The goodness of fit is fair but not perfect; a better curve fit might be achieved with a more sophisticated model.
However, we have taken a minimalistic approach to modelling the dasatinib effect because of the mechanistic uncertainties surrounding its mode of action. The duplet recruitment enhancement factor $\delta_D$ was estimated to equal 6.53±2.93, whereas the triplet recruitment enhancement factor $\delta_T$ was estimated to equal 13.7±7.38 (by simultaneous non-linear least-squares). The fit of the model indicates that the data are consistent with the hypothesis that dasatinib makes free TCRs more readily available to pMHC tetramers for recruitment into duplet and triplet clusters.

RESULTS

3.1 PKI treatment enhances pMHC tetramer staining of CD8$^+$ and CD4$^+$ T-cells

Incubation of ILA1, a CTL clone specific for the HLA A2-restricted epitope hTERT$^{540-548}$ (ILAKFHWL)$^1$ with the PKIs dasatinib or 3-(2-(1H-benzo[d]imidazol-1-yl)-6-(2-morpholinoethoxy)pyrimidin-4-ylamino)-4-methylphenol (Lck inhibitor II; Calbiochem) resulted in a >10 fold increase in pMHC tetramer staining intensity (Figure 1A). Pre-incubation with the PKI PP2 (Calbiochem) resulted in a moderate increase in tetramer staining; however, no significant enhancement was observed when CTL were pre-treated with herbimycin, PP3, genestein (Calbiochem) or staurosporine (Biomol) (data not shown). Importantly, PKI inhibitor treatment did not enhance staining with non-cognate pMHC tetramer (Figure 1B). Identical results were obtained for the HLA A2-restricted Melan A$^{25-35}$ ELAGIGILTV-specific CTL clone Mel13 (data not shown).

Dasatinib is a reversible dual Src/Bcr-Abl kinase inhibitor with clinical applications that suppresses the activity of many kinases, including the Src protein kinase Lck (IC$_{50}$=0.4nM) (Shah et al., 2004; Carter et al., 2005; Weichsel et al., 2008). Furthermore, while dasatinib reversibly inhibits antigen-specific T-cell effector functions, it is not toxic to T-cells in the short term at concentrations <100 nM (Weichsel et al., 2008). Indeed, T-cell clones incubated in 50 nM
dasatinib for 24 hours were able to regain responsiveness to antigen within 1 hour of drug removal (data not shown). Dasatinib can also be used in flow cytometry-based applications without loss of cell viability (Weichsel et al., 2008). These properties prompted us to select dasatinib for further investigation in the current study.

The enhancement of pMHCI tetramer staining following dasatinib treatment was highly dose dependent (Figure 1C). Maximal effect was achieved by exposing CTL to 50 nM dasatinib for 1 hour, which resulted in an 89% increase in pMHCI tetramer staining intensity (Figure 1C). Unexpectedly, pre-incubation of CTL with 50 nM dasatinib for as little as 30 seconds resulted in a 60% increase in pMHCI tetramer staining (Figure 1D). Furthermore, incubation with 50 nM dasatinib for 30 minutes significantly enhanced the staining of both ILA1 and Mel13 CTL over a wide range of pMHCI tetramer concentrations (Figure 1E&F). Pre-incubation with dasatinib also enhanced pMHCI tetramer staining of naïve murine F5 TCR CTL directly ex vivo (Figure 1G) and pMHCII tetramer staining of a HLA DR*010-restricted CD4+ T-cell clone (Figure 1H). Thus, pre-incubation with 50 nM dasatinib for 30 minutes provides a quick and easy way to enhance pMHC tetramer staining efficiency in both human (CD4+ and CD8+ T-cells) and murine systems. These effects are highly specific; increased pMHC tetramer binding only occurs in the presence of a cognate TCR/pMHC interaction (Figure 1B&H).

3.2 Dasatinib preferentially enhances pMHCI staining of T-cells bearing low affinity TCRs

In order to dissect further the effects of dasatinib, we examined pMHCI tetramer staining using several altered peptide ligands for the ILA1 CTL clone that differ in their binding affinity for the ILA1 TCR by >100-fold (Table 1). Pre-incubation with dasatinib enhanced staining efficiency with all variant pMHCI tetramers (Figure 2A). The percentage increase in tetramer staining afforded by pre-incubation with dasatinib for 8E, 5Y, 4L, ILA index, 3G8T and 3G pMHCI tetramers was 675%, 1825%, 324%, 111%, 75% and 26%, respectively. Thus, the benefits of dasatinib pre-treatment in terms of enhanced tetramer staining
intensity are greater for peptide variants that exhibit weaker interactions with the ILA1 TCR. The intensity of pMHCI tetramer staining in the presence and absence of dasatinib treatment was plotted against the monomeric TCR/pMHCI dissociation constants and a curve fitted according to the mathematical model outlined in the Materials and Methods (Figure 2B). The data demonstrate that, in the absence of dasatinib, there is a sharp reduction in tetramer staining intensity for ligands with TCR/pMHCI $K_D > 35\mu M$; this is consistent with previous observations (Laugel et al., 2007). In the presence of dasatinib, however, the TCR/pMHCI affinity threshold for this sharp drop-off did not occur until the $K_D$ exceeded 200$\mu M$. In fact, dasatinib treatment allows detectable staining of the ILA1 clone even when the agonist TCR/pMHCI $K_D$ exceeds 500$\mu M$. Dasatinib treatment therefore enables the physical detection of CTL bearing TCRs with low affinity for the cognate pMHCI ligand that would otherwise be undetectable using pMHCI tetramer staining alone.

### 3.3 Dasatinib reduces pMHCI tetramer-induced cell death

Previous studies have shown that soluble pMHCI tetramer-induced signalling can trigger cell death (Purbhoo et al., 2001; Xu et al., 2001; Guillaume et al., 2003; Cebeauer et al., 2005). This can reduce the number of live cells that remain after pMHCI tetramer staining under normal conditions. Dasatinib blocks antigen-specific signalling and subsequent T-cell effector functions (Weischel et al., 2008). Consequently, we hypothesized that dasatinib could prevent pMHCI tetramer-induced cell death. Indeed, the percentage of tetramer-positive cells that died when PBMCs were stained directly ex vivo with pMHCI tetramers representing epitopes derived from cytomegalovirus (CMV) and Epstein-Barr virus (EBV) was reduced in the presence of dasatinib (Figure 3). Therefore, dasatinib exerts three beneficial effects: (i) it increases the intensity of pMHCI and pMHCI\textsuperscript{II} tetramer staining; (ii) it preferentially enhances pMHCI tetramer staining of T-cells bearing low affinity TCRs; and, (iii) it reduces pMHCI tetramer-induced cell death.
3.4 Substantial improvements in the detection of antigen-specific CD8+ T-cells directly ex vivo

The above findings suggest that dasatinib treatment might enable the identification of low avidity antigen-specific CD8+ T-cells directly *ex vivo* that cannot be 'seen' in the absence of the drug. To test this idea, we first examined the staining of CTL lines in the presence or absence of 50 nM dasatinib. Staining improvements were observed in three different CTL lines raised against the Melan-A/Mart-1 26-35 epitope (ELAGIGILTV) and three different CTL lines stimulated with the influenza matrix M158-66 epitope (GILGFVFTL), all derived from HLA A2+ individuals. Representative data are shown in Figure 4A. In all cases, dasatinib treatment substantially enhanced the staining intensity of cognate CD8+ T-cells without concomitant increases in the tetramer-negative population. In accordance with the results above, CD8+ T-cells that stained poorly with pMHCI tetramer exhibited the greatest benefit from dasatinib treatment. The staining intensity of all cognate CD8+ T-cells increased by at least 2-fold after dasatinib treatment, but T-cells that bound tetramer weakly exhibited increases of >20-fold in their fluorescence intensity. In many cases, larger populations of cells that stained with the corresponding pMHCI tetramer were detected after dasatinib treatment. This increase in tetramer* cells after dasatinib treatment likely reflects the combined effects of a lower detection threshold in terms of TCR/pMHCI affinity and the fact that dasatinib reduces pMHCI tetramer-induced cell death (Figure 3). Subsequently, we examined whether dasatinib could enhance pMHCI tetramer staining of cognate CD8+ T-cells in direct *ex vivo* PBMC samples and enable the detection of antigen-specific CD8+ T-cells that are 'invisible' with routine staining procedures. Indeed, a substantial increase in both pMHCI staining intensity and the percentage of antigen-specific CD8+ T-cells was observed at both 4 °C and 37°C in PBMC samples stained with HLA A2 tetramers specific for antigens derived from CMV, EBV and Melan A (Figure 4B).
3.5 pMHCI staining of functional autoimmune CTL following dasatinib treatment

We next examined pMHCI tetramer staining of IE6, a preproinsulin (PPIis-24)-specific HLA A2-restricted autoreactive CTL clone isolated from a patient with type 1 diabetes. This CTL clone produces TNFα, IFNγ and MIP1β on stimulation with target cells pulsed with cognate PPI-derived peptide antigen (Figure 5A), and does not stain with cognate pMHCI tetramer using conventional staining procedures (Figure 5B). An identical result was obtained for 2D6, a different PPI-specific CTL clone isolated from a patient with type 1 diabetes (data not shown). Dasatinib treatment allowed both CTL clones to bind cognate tetramer without affecting staining with non-cognate tetramer (Figure 5B & data not shown). In keeping with these findings, dasatinib treatment allowed the identification of a HLA A2/PPIis-24 tetramer-positive population directly ex vivo from a type 1 diabetic patient, consistent with a corresponding IFNγ ELISpot response to PPIis-24 peptide of 13 responder cells per 10⁶ PBMCs (Figure 5C & data not shown). Dasatinib did not increase direct ex vivo HLA A2/PPIis-24 tetramer staining in healthy HLA A2-matched control subjects (Figure 5C). A seven-fold increase in the percentage of autoreactive CTL was observed when short-term lines expanded from two type 1 diabetic patients were stained in the presence of dasatinib (Figure 5C & data not shown). Thus, dasatinib treatment allows the detection of functional autoreactive CTL that are otherwise undetectable with standard staining conditions.

3.6 How does dasatinib exert its beneficial effects on pMHIC tetramer staining?

Previous studies have demonstrated that incubation with Src kinase inhibitors results in enhanced TCR and CD8 expression at the cell surface (Luton et al., 1994; D’Oro et al., 1997). Consistent with these observations, we have recently demonstrated that increased levels of TCR and CD8 are seen at the cell surface following incubation with dasatinib for 4 hours (Weischel et al., 2008). Initially, therefore, we investigated this increase in TCR and CD8 levels as a possible
mechanism for the observed effects on tetramer binding. The beneficial effects of dasatinib on pMHCI tetramer staining were observed within seconds of dasatinib treatment (Figure 1D), whereas significant increases in TCR and CD8 levels were not observed until > 30 minutes (Figure 6). Therefore, this time dependent accumulation of TCR and CD8 at the cell surface cannot explain the effects of dasatinib on tetramer binding.

We next investigated whether the mechanism of PKI action operates through TCR- or CD8-mediated effects. Dasatinib treatment enhanced pMHCI tetramer staining of the HLA A2-restricted ELAGIGILTV-specific Melc5 CTL clone and a CTL line raised against the Melan-A/Mart-12635 epitope (ELAGIGILTV) with both wildtype and CD8-null (DT227/8KA) tetramers (Figure 7A&B), thereby demonstrating that PKIs can exert their effects in the absence of a pMHCI/CD8 interaction. Thus, consistent with effects on pMHCII tetramer binding (Figure 1), dasatinib does not enhance pMHCI tetramer binding via CD8-mediated effects.

TCR expression levels are not static and TCRs are constantly being down-regulated from the cell surface (Krangel, 1987). TCR internalization is thought to be mediated by three different mechanisms: (i) constitutive recycling of the TCR between intracellular compartments and the plasma membrane in resting cells by an unknown mechanism (Dietrich et al., 2002); (ii) protein kinase C activation (Minami et al., 1987; Dietrich et al., 2002); and, (iii) lck-mediated tyrosine phosphorylation following TCR ligation by specific pMHCI ligand. Dasatinib has been shown to target lck and therefore has the potential to inhibit the latter pathway of TCR down-regulation. Indeed, dasatinib treatment was found to block antigen-induced TCR downregulation from the CTL surface (Figure 8A). pMHC tetramers are rapidly internalized under normal staining conditions (Whelan et al., 1999) and therefore we reasoned that dasatinib might exert its beneficial effects by blocking this process. To this end, fluorescence microscopy was performed in the presence and absence of dasatinib (Figure 8B). HLA A2/ILAKFLHWL-Alexa488 tetramer capping and internalization was blocked in the presence of dasatinib and remained on the cell surface where it formed a
ring that was visibly brighter than tetramer that had been internalized (Figure 8B). Thus, by preventing TCR downregulation, PK inhibition acts to drive the system towards a higher number of surface TCRs and a higher number of potential productive engagements with pMHC tetramer. This has the effect of increasing tetramer on-rate, at least in pMHCI systems (Figure 9).

4. DISCUSSION

pMHC tetramer technology has revolutionized the study of antigen specific T-cells. However, one major limitation of this technique is that pMHCI, and most likely pMHCII, tetramer staining is dependent on a distinct TCR affinity threshold (Laugel et al., 2007). Consequently, pMHC tetrarmers fail to identify T-cells that express TCRs with low affinity for cognate antigen; such low affinity interactions characterize TCR/pMHCI binding in tumor-specific and autoreactive CD8+ T-cells (Cole et al., 2007). Here, we demonstrate that a short incubation with a reversible PKI such as dasatinib results in three major benefits in terms of pMHC tetramer staining. First, substantial improvements in pMHC tetramer staining intensity are observed. This effect applies to both CD4+ and CD8+ T-cells (Figure 1). Indeed, the beneficial effects are so striking even at low pMHC tetramer concentrations that dasatinib treatment could be used to conserve reagent. Second, dasatinib treatment reduces tetramer-induced cell death that has been previously reported to be an issue with pMHCI tetramer staining protocols (Purbhoo et al., 2001; Xu et al., 2001; Guillaume et al., 2003; Cebecauer et al., 2005). Third, the benefits of dasatinib treatment are greater for TCR/pMHCI interactions of weak affinity and, as a result, dasatinib enhances the detection of low avidity CD8+ T-cells; this effect increases the number of CD8+ T-cells that can be detected directly ex vivo, particularly in the setting of tumor-specific and autoreactive CD8+ T-cell populations. Such effects are also likely to apply to CD4+ T-cells, which typically bind cognate pMHCII antigens with affinities lower than those reported for pMHCI systems (Cole et al., 2007).

Importantly, no increase in background staining was seen in any of the systems tested here (Figure 1B, 1H, 5B&5C). In fact the tetramer negative background
was actually seen to decrease with dasatinib treatment in some staining experiments (Figure 4C&5C). Dasatinib proved to be a particularly powerful tool in the detection of autoreactive CTL from type I diabetic patients. No increase in the PPI tetramer positive population was observed in healthy donors and indeed staining was only ever seen if a functional response to the preproinsulin peptide was evident. Therefore benefits only apply to T-cells that express TCRs specific for the pMHCI tetramer in use. This conclusion is further strengthened by the finding that the beneficial effects are TCR mediated and do not involve the CD8 coreceptor. Therefore dasatinib facilitates the specific TCR/pMHCI interaction rather than the non-specific pMHCI/CD8 interaction.

Dasatinib prevents TCR downregulation and tetramer internalization from the cell surface. How does this effect result in faster tetramer on-rates and the beneficial effects described above? When an individual pMHCI molecule in a pMHCI tetramer engages a cell surface TCR, this engagement can be either 'productive' or 'non-productive' in terms of capturing the tetramer from solution (Figure 10). A productive engagement requires a second pMHCI in the tetramer to bind a second TCR before the first pMHCI dissociates. By preventing TCR downregulation after non-productive engagement, dasatinib treatment acts to maintain TCRs on the cell surface where they are available for future interactions. There is increasing evidence that non-triggered TCRs are internalized with engaged TCRs (Niedergang et al., 1997; San Jose et al., 2000). Thus, it is possible that the tetramer internalization that occurs after productive engagement in the absence of dasatinib treatment also results in the internalization of non-engaged TCR and coreceptor. By preventing TCR downregulation, dasatinib and other effective PKIs would act to maintain surface TCRs and therefore enable a higher number of potential productive engagements with pMHCI tetramer as indicated by the red arrows (Figure 10).

In summary, we have demonstrated that a short incubation with reversible PKIs such as dasatinib substantially improves the staining intensity of cognate T-cells with pMHCI tetramers and can expose concealed antigen-specific T-cells that
bear low affinity TCRs. These benefits are restricted to cognate T-cells and are not accompanied by concomitant increases in background staining. This simple and universally applicable technique is likely to be beneficial in all studies of antigen-specific T-cells.
REFERENCES


activation using alpha3 domain mutants of MHC class I/peptide complex. Immunity 14, 591-602.
Table One

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Table 1: Affinity measurements of the interaction between the ILA1 TCR and hTERT<sub>540</sub>-54β<sub>4</sub> pMHCI variants. Summary of the results obtained by nonlinear analysis of surface plasmon resonance binding equilibrium experiments as detailed in Laugel et al. 2007 (Laugel et al., 2007) and Melenhorst et al (Melenhorst et al., 2008). K_D values were determined by analyzing the data in nonlinear curve fittings to the equation AB=BxAB<sub>max</sub>/(K_D+B) assuming 1:1 Langmuir binding.
1. A method of detecting, within a population of cells, T cells of the immune system comprising exposing said population of cells to at least one protein kinase inhibitor either before, or whilst, exposing said population of cells to at least one detecting or staining agent that detects or stains protein kinase sensitive cell surface markers of said T cells, such as T cell surface receptors (TCR'S).

2. Use of at least one protein kinase inhibitor to treat a population of cells in order to detect, within said population of cells, T cells of the immune system via the detection or staining of protein kinase sensitive cell surface markers of said T cells, such as T cell surface receptors (TCR's), with at least one detecting or staining agent for said cell surface markers; wherein said population of cells are treated with said protein kinase inhibitor either before, or whilst, exposing said population of cells to said at least one detecting or staining agent.

3. A method of conserving agents when detecting, within a population of cells, T cells of the immune system that involves enhancing the performance of said agents by exposing said population of cells to at least one protein kinase inhibitor either before, or whilst, exposing said population of cells to at least one detecting or staining agent that detects or stains protein kinase sensitive cell surface markers of said T cells, such as T cell surface receptors (TCR's), and thereby conserving the amount of said detecting or staining agent required to detect or stain said T cells.

4. A method according to claim 3 wherein the agents to be conserved comprise known detecting or staining agents for detecting or staining T cell surface markers.
5. A method of sorting, in a population of cells, a viable, selected, sub-population of cells comprising exposing said population of cells to at least one protein kinase inhibitor either before, or whilst, exposing said population of cells to at least one sorting agent that sorts, detects or stains a selected type of cell and thereby enables said selected cells to be sorted, detected or stained and simultaneously, or subsequently, isolated from said population of cells; wherein exposure of said population of cells to said protein kinase inhibitor increases the viability of said selected cells.

6. Use of at least one protein kinase inhibitor for sorting, within a population of cells, a viable, selected, sub-population of cells by exposing said population of cells to at least one protein kinase inhibitor either before, or whilst, exposing said population to at least one sorting agent that sorts, detects or stains a selected type of cell and thereby enables said selected cells to be sorted, detected or stained and simultaneously, or subsequently, isolated from said population of cells; wherein exposure of said population of cells to said protein kinase inhibitor increases the viability of said selected cells.

7. A method or use according to any preceding claim wherein said protein kinase inhibitor is a reversible protein kinase meaning its binding action to its target site or substrate can be reversed.

8. A method or use according to any preceding claim wherein said protein kinase inhibitor is any one or more of the following: Dasatinib, Lck inhibitor II, Wortmannin, Resveratrol, LY294002, AS604850, PI3-K alpha inhibitor 1 or PI3-K alpha inhibitor 2.

9. A method or use according to any preceding claim wherein said protein kinase inhibitor is a tyrosine protein kinase inhibitor.
10. A method or use according to any preceding claim wherein said agent is a multimerized ligand.

11. A method or use according to claim 10 wherein said ligand is a peptideMHC multimer such as a tetramer, pentamer, octomer or dextramer.

12. A method or use according to claim 11 wherein said peptideMHC multimer is a peptideMHC class I and/or a peptideMHC class II multimer.

13. A method or use according to any preceding claim wherein said agent is an antigen specific marker that is specific for at least one T cell TCR.

14. A method or use according to any preceding claim wherein said treating of said population of cells comprises exposing said population of cells to said inhibitor for a period of time up to one hour.

15. A method or use according to claim 14 wherein said period is approximately 30mins.

16. A method or use according to any preceding claim wherein said protein kinase inhibitor is used at a concentration of up to 30OnM.

17. A method or use according to claim 16 wherein said concentration is less than 20OnM.
**Fig. 1A**

**Fig. 1B**
**Fig. 1C**

![Bar chart showing the relationship between MFI Tetramer and Dasatinib concentration.](image)

**Fig. 1D**

![Line graph showing the relationship between MFI Tetramer and incubation time with 50nM Dasatinib.](image)
Fig. 1E

Fig. 1F
Fig. 3
Fig. 5C
Fig. 6

TCR

CD8

MFI

Time (minutes)

Dasatinib

No Dasatinib
Fig. 9
Tetramer binding

Non-productive

TCR downregulation

dasatinib

Productive

TCR downregulation

dasatinib

dasatinib

Fig. 10