Protective Efficacy of Cross-Reactive CD8⁺ T Cells Recognising Mutant Viral Epitopes Depends on Peptide-MHC-I Structural Interactions and T Cell Activation Threshold

Sophie A. Valkenburg¹, Stephanie Gras², Carole Guillonneau¹, Nicole L. La Gruta¹, Paul G. Thomas³, Anthony W. Purcell⁴, Jamie Rossjohn², Peter C. Doherty^{1,3}, Stephen J. Turner¹, Katherine Kedzierska^{1*}

1 Department of Microbiology and Immunology, University of Melbourne, Parkville, Australia, 2 The Protein Crystallography Unit, Department of Biochemistry and Molecular Biology, Monash University, Clayton, Australia, 3 Department of Immunology, St Jude Children's Research Hospital, Memphis, Tennessee, United States of America, 4 Department of Biochemistry and Molecular Biology, the Bio21 Molecular Science and Biotechnology Institute, University of Melbourne, Parkville, Australia

Abstract

Emergence of a new influenza strain leads to a rapid global spread of the virus due to minimal antibody immunity. Preexisting CD8⁺ T-cell immunity directed towards conserved internal viral regions can greatly ameliorate the disease. However, mutational escape within the T cell epitopes is a substantial issue for virus control and vaccine design. Although mutations can result in a loss of T cell recognition, some variants generate cross-reactive T cell responses. In this study, we used reverse genetics to modify the influenza NP₃₃₆₋₃₇₄ peptide at a partially-solvent exposed residue (N->A, NPN3A mutation) to assess the availability, effectiveness and mechanism underlying influenza-specific cross-reactive T cell responses. The engineered virus induced a diminished CD8⁺ T cell response and selected a narrowed T cell receptor (TCR) repertoire within two VB regions (V β 8.3 and V β 9). This can be partially explained by the H-2D^bNPN3A structure that showed a loss of several contacts between the NPN3A peptide and H-2D^b, including a contact with His155, a position known to play an important role in mediating TCR-pMHC-I interactions. Despite these differences, common cross-reactive TCRs were detected in both the naïve and immune NPN3A-specific TCR repertoires. However, while the NPN3A epitope primes memory T-cells that give an equivalent recall response to the mutant or wild-type (wt) virus, both are markedly lower than wt->wt challenge. Such decreased CD8⁺ responses elicited after heterologous challenge resulted in delayed viral clearance from the infected lung. Furthermore, mice first exposed to the wt virus give a poor, low avidity response following secondary infection with the mutant. Thus, the protective efficacy of cross-reactive CD8⁺ T cells recognising mutant viral epitopes depend on peptide-MHC-I structural interactions and functional avidity. Our study does not support vaccine strategies that include immunization against commonly selected cross-reactive variants with mutations at partially-solvent exposed residues that have characteristics comparable to NPN3A.

Citation: Valkenburg SA, Gras S, Guillonneau C, La Gruta NL, Thomas PG, et al. (2010) Protective Efficacy of Cross-Reactive CD8⁺ T Cells Recognising Mutant Viral Epitopes Depends on Peptide-MHC-I Structural Interactions and T Cell Activation Threshold. PLoS Pathog 6(8): e1001039. doi:10.1371/journal.ppat.1001039

Editor: Daniel C. Douek, NIH/NIAID, United States of America

Received March 15, 2010; Accepted July 13, 2010; Published August 12, 2010

Copyright: © 2010 Valkenburg et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was funded by the NHMRC Project Grants to PCD (Al454595) and KK (Al454312), a University of Melbourne Early Career Researcher Grant (to KK), NIH grant Al170251 and NHMRC program grant to PCD, SJT (Al567122). KK and NLG are NHMRC RD Wright Fellows, AWP is a NHMRC Senior Research Fellow, SJT is a Pfizer Senior Research Fellow and CG is a Marie Curie International Fellow and is supported by the 6th FP of the EU, Marie Curie #040840. JR is an ARC Federation Fellow. SAV is a recipient of the Australian Postgraduate Award. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: kkedz@unimelb.edu.au

Introduction

Virus-specific CD8⁺ T cells play a critical role in host defence via the production of antiviral cytokines, the direct killing of virusinfected cells and the establishment of immunological memory [1]. The selection of CD8⁺ T cells into an immune response requires specific interaction between the T cell receptor (TCR) and virus peptide bound to Major Histocompatibility Complex class I (pMHC-I) molecules on the surface of infected host cells. The processing of virus proteins into short fragments generates thousands of peptides that might potentially form pMHC-I epitopes, but only a few elicit CTL responses [2]. Virus escape mutants are well documented for persistent infections and constitute a major problem for $CD8^+$ T cellmediated control and vaccine design [3,4,5,6,7,8,9]. With regard to the influenza A viruses, mutational changes driven by $CD8^+$ cytotoxic T lymphocytes (CTLs) are unlikely to result in long-term persistence within the individual, as other mechanisms (particularly antibody) can ultimately mediate virus clearance [10]. Even so, the fact that such mutants can be found in nature suggests that influenza virus-specific CTLs are of protective value. Perhaps this reflects that the infection of new subjects favours the selection of mutant viruses that are more slowly controlled (and thus shed for longer), particularly in the face of a seasonal "bottleneck" where

Author Summary

Introduction of a new influenza strain into human circulation leads to a rapid global spread of the virus due to minimal antibody immunity. Established T-cell immunity towards conserved viral regions provides some protection against influenza and promotes rapid recovery. However, influenza viruses mutate to escape the protective immunity. We found that established T cell immunity can recognise influenza mutants with variations at positions that are partially involved in T cell recognition. However, an initial priming with the mutated variant decreases recognition of the original parental virus. This finding results from a markedly lower functional guality and limited structural interactions of the mutant. In terms of possible vaccination strategies for rapidly changing viruses or tumours, it appears that priming with crossreactive mutants that display such characteristics would be of no benefit as the same level of T cell immunity against such mutants can be elicited by exposure to the original virus.

much of the population is already immune [11]. In humans, influenza escape variants have been observed for CD8⁺ T cell epitopes presented in context of several HLAs, including HLA-B8, HLA-B27 and HLA-B35 [12,13,14,15,16,17,18,19]. The immunogenic peptides can be modified at an MHC anchor residue, resulting in defective binding to the MHC-I glycoprotein, or at a TCR contact site. Mutations at TCR contact residues lead to partial (cross-reactive) or total (non-cross-reactive) loss of recognition by wt CD8⁺ T cells [13], with some variants eliciting epitope-specific CD8⁺ T cell responses that are both novel and of substantial magnitude [12].

Using influenza A virus infection of B6 mice [20], we showed previously that virus variants with mutations at critical solventexposed residues that are important for TCR binding can generate effective but non-cross-reactive CD8⁺ CTL responses to what are essentially new epitopes [21,22]. This raises the possibility that it might be worthwhile to think in terms of vaccinating against likely virus escape mutants. The present analysis focuses on the crossreactive (to wt D^bNP₃₆₆) CD8⁺ T cell response to the mutant D^bNPN3A epitope [22] formed by substitution of the partially solvent exposed, and non-prominent for TCR binding, residue at position (P) 3 of the influenza NP_{366–374} peptide [21,23]. The findings have implications for vaccines to combat virus mutants and tumour escape variants, and suggest that immunisation against likely cross-reactive variants would have to be carefully evaluated to see if the strategy is worthwhile.

Results

Earlier analysis [21,22] using sequential alanine substitutions in the immunogenic NP₃₆₆₋₃₇₄ peptide established that there is some cross-reactive, though diminished, stimulation following the incubation of wt D^bNP₃₆₆-specific CD8⁺ T cells with the NP mutant peptide containing a single asparagine to alanine substitution at position (P) 3 (NPN3A mutant). Such cross-reactive CD8⁺ T cell responses after NPN3A stimulation are not surprising as the partially solvent exposed P3N residue is unlikely to play any prominent role in contacting the TCR [23]. The wt NP₃₆₆₋₃₇₄ peptide binds to H-2D^b in an extended conformation where the P2-Ser, P5-Asn and P9-Met are the anchor residues, P3-Asp is a semi-anchor (partially buried/partially exposed), and P4-Glu, P6-Met, P7-Glu and P8-Thr are solvent exposed and available for contact by the TCR (Fig. 1). This provides the basis for a defined experimental system that can be used to determine what happens when a TCR repertoire is selected to what would seem (from *in vitro* analysis) to be a suboptimal, cross-reactive mutant epitope. The interpretation of *in vitro* analysis alone should, however, be tempered with caution, as another mutant (M6A) that did not cross-react at all with the wt epitope elicited a completely novel, *in vivo* endogenous CTL response of equivalent magnitude when expressed in an infectious influenza A virus [22]. What would be the case for TCR responses elicited by influenza A viruses expressing the mutant NPN3A peptide in the native viral protein?

Consequences of 1° and 2° homologous challenge for CTL response magnitude

The NPN3A mutation was engineered into PR8 (H1N1) and HKx31 (H3N2) influenza viruses (PR8NPN3A, HKNPN3A) to allow cross-challenge experiments in the absence of antibody neutralisation. The B6 mice were immunised i.p. with the virulent PR8 mutant and wt viruses, while the HK viruses were used for primary (1°) i.n. infection of naïve mice or secondary (2°) i.n. challenge of PR8-immune (>30d previously) mice. Naïve (primary) or PR8-immune (PR8, or PR8NPN3A) mice were challenged i.n. with the homologous virus (HK, or HKNPN3A) and $CD8^+$ T cell responses were measured using the D^bNP_{366} and D^bNPN3A tetramers (Fig 2). It was immediately apparent that the splenic D^bNPN3A⁺CD8⁺ set elicited by the HKNPN3A challenge was significantly smaller on d10 (p<0.05) than the D^bNP₃₆₆⁺CD8⁺ T cell response induced by the wt virus (Fig 2B), though there was no significant difference between D^bNPN3A⁺CD8⁺ and D^bNP₃₆₆⁺CD8⁺ T cell numbers at the site of infection (BAL, Fig 2A). This has been seen before [24] and suggests that the need to clear virus from the lung results in preferential CTL localization to the site of infection when immune T cell numbers are limited. The profile of a diminished D^bNPN3A⁺CD8⁺ T cell response was maintained into memory (d28, Fig 1C; p<0.02), supporting the view that the relative size of persistent T cell pools reflects the extent of antigen driven proliferation during the acute anti-viral response.

A high proportion of the wt $D^bNP_{366}^+CD8^+$ T cells in BAL (94.0% (1⁰, d10); 93.2 (2⁰, d8); Fig 2A) and spleen (69.8% (1⁰, d10); 93.7% (2⁰, d8); Fig 2B) bound the D^bNPN3A tetramer and produced IFN- γ when stimulated by the NPN3A peptide (data not



Figure 1. A stick conformation illustrating how the influenza $NP_{366-374}$ interacts with H-2D^b. $NP_{366-374}$ peptide binds to the H-2D^b in an extended conformation. The P3-Asp, P5-Asn and P9-Met are the anchor residues, whereas the P4-Glu, P6-Met, P7-Glu and P8-Thr are solvent exposed and available for contact by the TCR. doi:10.1371/journal.ppat.1001039.g001



Figure 2. Acute and recall CD8⁺ T cell responses to D^bN3A and D^bNP₃₆₆. The magnitude of CD8⁺ T cell responses at the peak (d10 1° A,B; d8 2° D,E) or memory (d28, CF) phases following 1° (A–C) or 2° (D–F) infection. Cells were stained with the D^bNP₃₆₆-PE, D^bNPN3A-PE or D^bPA₂₂₄-PE tetramers and anti-CD8-PerCPCy5.5 mAb. The numbers of epitope specific CD8⁺ T cells were calculated from the % cells staining and the total cell counts. The wt HK or HK-NPN3A viruses were used for 1° i.n. infection and 2° i.n. challenge of i.p.-primed (PR8 or PR8-NPN3A i.p. >30d previously) B6 mice. Data are mean±SD n=5 mice per group, *=p<0.01. Memory T cell numbers were also analysed on d60, and the tetramer analysis was replicated using the ICS assay (data not shown). (G, H) Representative dot plots are shown for (G) D^bNP₃₆₆ or NPN3A tetramer staining and (H) intracellular cytokine assay after infection with either HK or HK-NPN3A for the same individual mouse.

shown). Similarly, the majority ($\geq 90\%$) of CD8⁺ T cells elicited by HKNPN3A infection bound the D^bNP₃₆₆ tetramer and produced IFN- γ (data not shown) at levels equivalent to those seen for the response to NPN3A, indicating a high level of cross-reactivity between the D^bNP₃₆₆⁺CD8⁺ and NPN3A⁺CD8⁺ T cell responses. Staining with the D^bPA₂₂₄ tetramer was included to establish that the NPN3A mutation neither diminished nor enhanced other influenza-specific CD8⁺ T cell responses (Fig 2).

Despite the decreased $D^bNPN3A^+CD8^+$ T cell numbers generated following primary infection (Fig 2A–C), the recall response was substantial following HKNPN3A challenge of PRNPN3A-immune mice (Fig 2DE) and the NP>PA immunodominance hierarchy that has long been recognized for secondary responses to wt influenza A viruses in H2^b mice [25] was maintained (Fig 2DE). Similarly, the total cell numbers for memory D^bPA₂₂₄-specific T cells on d28 were comparable for those primed and boosted within the wt and mutant virus combinations (Fig 2F), while the D^bNPN3A⁺ CD8⁺ set was 8-fold smaller than the wt D^bNP₃₆₆⁺CD8⁺ population (p<0.005). Again, the results following secondary challenge support the view that, at least in the earlier (d28) stages of memory, T cell numbers reflect the extent of clonal expansion during the acute phase [26]. In our experiments, we detected D^bNP₃₆₆⁺CD8⁺ and NPN3A⁺CD8⁺ T cells by two techniques, tetramer staining and IFN- γ ICS. Both techniques gave us comparable antigen-specific CD8⁺ T cell numbers, indicating that both tetramers accurately detected epitope-specific populations (Fig. 2GH).

Given that the NPN3A mutation was associated with a numerically diminished response following infection with either the wt or NPN3A influenza A viruses (Fig 2), we also asked if there was any effect on CD8⁺ T cell function or phenotype, particularly for markers (CD62L and CD127) that discriminate between memory T cell subsets [27,28]. More of the D^bNPN3A⁺CD8⁺ T cells remained CD62L^{hi} when sampled at the peak of the response

(d10) following primary challenge (Fig S1A), confirming the impression from the quantitative analysis (Fig 2 A-C) that there may be less clonal expansion. On the other hand, IL-7R expression was comparable for the D^bNPN3A⁺CD8⁺ and D^bNP₃₆₆⁺CD8⁺ T cell populations generated in virus-infected naïve mice (Fig S1 CD) with both being (p < 0.01) lower than the values for the D^bPA₂₂₄-specific set (Fig S1C). This suggests that IL-7R levels may be antigen dose- rather than magnitude-dependent. Neither of these differential effects was apparent for d28 memory T cells specific for the mutant or wt epitopes (Fig S1BD). Functional analysis of cytokine production based on short term (5h) stimulation with cognate peptide in the ICS assay showed no obvious differences at any stage for the D^bNP₃₆₆ and D^bNPN3Aspecifc T cells, though the usual divergence [29] from the D^bPA₂₂₄⁺CD8⁺ T cell response was observed (p<0.01) (Fig S1 E-H). These data suggest that NPN3A mutation leads to crossreactive, but diminished, CD8⁺ T cell responses with comparable cytokine production profiles.

Crystal structure and thermostability of the H2Db-NP-N3A complex

Can the smaller response to D^bNPN3A be correlated with structural constraints or any decrease in stability for the pMHC-I complex? The D^bNPN3A crystal structure containing the heavy chain of H-2D^b (residues 1–275), the β 2 microglobulin (residues 1– 99) and the 9 residues of the NPN3A peptide was determined to a 2.6 Å resolution (Fig. 3 and Table 1), with a final R_{factor} of 22.1% and an $R_{\rm free}$ of 30.4%. The structure of the H-2Db-NPN3A complex was compared (Fig. 3B, Table 1) to the wt D^bNP₃₆₆ [23]. As observed for the wt D^bNP₃₆₆, the mutant NPN3A peptide bound H-2D^b in an extended conformation, the P2-Ser, P5-Asn and P9-Met represent the anchor residues, P3-Ala semi-anchor residue, whereas the P4-Glu, P6-Met, P7-Glu and P8-Thr are solvent exposed and available for contact by the TCR (Fig. 3AB). The D^bNP₃₆₆ and D^bNPN3A structures are very similar with a root mean square deviation (r.m.s.d.) of 0.44 Å on the $\alpha 1 - \alpha 2$ domains and a r.m.s.d of 0.20 Å on the peptides. With the exception of P3-Ala, the structure of D^b-NP₃₆₆ and D^b-NPN3A superimpose well.

However, the mutation of P3-Asn to Ala leads to a loss of 35 contacts between the peptide and the MHC molecule. In comparison to the wt NP₃₆₆ that makes 636 contacts with the H-2D^b molecule, the NPN3A peptide achieves only 601 contacts. Interestingly, the Asn3 Ala mutation abolishes contacts between the P3 residue of the peptide with His155 and the Tyr156 of the MHC and eliminates one hydrogen bond between the P3-Asn^{$N\delta 2$} and the P4-Glu^O (Fig. 3CD). The loss of inter-residue bonding between 3N and 4E within the NPN3A peptide may also be important for TCR recognition as it changes the characteristics of the wt D^bNP₃₆₆ epitope (defined as type III constraint) [30] into an unconstrained D^bNPN3A epitope. This change of constraint within the epitope could affect the dynamics of the peptide upon TCR ligation, and subsequently alter TCR reactivity toward the mutated epitope. In addition, the loss of contacts with the His155 of the MHC is of interest, as position 155, termed the "gate-keeper residue" [31] is involved in contacting the TCRs in all TCRpMHC-I complexes solved to date [32]. Thus, the lack of interactions between the NPN3A peptide and the MHC through His155 may also affect recognition of the complex by D^bNP₃₆₆specific TCRs.

The loss of contacts between the peptide and the MHC molecule could lead to decreased stability of the peptide and subsequent changes in NPN3A presentation. To test this hypothesis, we performed a thermostability assay on both NP_{366}

and NPN3A bound to the H-2D^b molecule. The NP₃₆₆ and NPN3A peptides are equally effective at stabilising H-2D^b. The pMHC-I complex with the NP₃₆₆ wt peptide had a Tm of $51.8\pm0.7^{\circ}$ C and D^bNPN3A showed a comparable level of thermostability (Tm = $51.4\pm1^{\circ}$ C), irrespective of the concentrations of the complex used for the assay. This suggests that the NPN3A mutation does not modify the stability of the pMHC-I complex when compared to the cognate epitope.

Naïve precursor frequency and TCR repertoire for D^b NPN3A

Is the smaller D^bNPN3A⁺CD8⁺ T cell response a consequence of diminished naïve CTL [33]? We found (Fig 4A) similar naïve CTLp counts for D^bNPN3A (34.6±13.08) and D^bNP₃₆₆ (28.5 ± 11.0) , indicating that the smaller response to D^bNPN3A⁺CD8⁺ is not due to reduced number of naïve precursors. Furthermore, assessing the extent of V β 8.3 bias (the dominant V β for the D^bNP₃₆₆⁺CD8⁺ set) within the naïve D^bNPN3A⁺CD8⁺ population (Fig 4B) showed that the extent of V β 8.3 usage (mean 13.2% \pm 4.3) (Fig. 4B) was much the same as that determined previously [33] for naïve D^bNP₃₆₆⁺CD8⁺ CTLps $17.1\% \pm 7.4$ However, sequencing (mean the naïve D^bNPN3A⁺Vβ8.3⁺CD8⁺ TCR CDR3β regions showed a clear difference from the comparable wt-specific set. The "public TCR" dominance characteristic of $D^b NP_{366}^+ V\beta 8.3^+ CD8^+ T$ cells in both pre-immune [33] and immune [34] TCRB repertoires [34] was not a prominent feature of the D^bNPN3A-specific TCR repertoire. These public TCRs were found in only one (SGGANTGQL and SGGGNTGQL) or two (SGGSNTGQL) of the 10 mice tested (Fig. 4C). Thus, although the naïve CTLp frequencies are comparable for D^bNP₃₆₆⁺Vβ8.3⁺CD8⁺ and D^bNPN3A⁺Vβ8.3⁺CD8⁺ T cells and there is some overlap of some cross-reactive TCRs, the two repertoires are far from identical. The roughly equivalent numbers of precursors specific for the wt D^bNP₃₆₆ and mutant NPN3A peptides were unexpected considering the lower response after infection with the mutated virus. The lower magnitude of the NPN3A⁺CD8⁺ T cell response and narrower TCR β repertoire suggest that only a proportion of naïve NPN3A⁺CD8⁺ precursors are being recruited into the immune response or that, once recruited, these cells do not expand efficiently. Inefficient recruitment and/or expansion early after influenza infection, despite large naïve CTL precursor numbers, have been recently documented by our group as key determinants of subdominance for D^bPB1-F2₆₂ and K^bNS2₁₁₄-specific responses [33].

$\mathsf{D}^b\mathsf{NPN3A}$ selects distinct and more restricted TCR $\mathsf{V}\beta$ signatures

The next step was to dissect the immune $D^bNPN3A^+CD8^+$ CTL repertoire to determine how TCR diversity relates to the size of the immune $D^bNPN3A^+CD8^+$ T cell response. The D^bNPN3A specific CD8⁺ T cells were first analysed for V β usage by staining with a panel of anti-TCRV β mAbs and the D^bNPN3A tetramer. After infection with the NPN3A viruses, the strong V β 8.3 bias characteristic of responding $D^bNP_{366}^+CD8^+$ T cells [34,35] was prominent for the $D^bNPN3A^+CD8^+$ sets in only 50% of the immune mice (n = 10). The mutant $D^bNPN3A^+CD8^+$ T cells utilized a variable spectrum of TCRV β elements, with overrepresentation of V β 7, 8.1/8.2, 9, 11, and 12 after primary (Fig S2A) or V β 6, 7 and 9 following secondary (Fig S2B) NPN3A virus challenge.

We analysed TCR β clonotypes within the V β 8.3 as (i) it is still a preferred region (29.8% ±20.2) of NPN3A⁺CD8⁺ T cell response)



Figure 3. Structure of the D^bNP₃₆₆₋₃₇₄ and D^bNPN3A complexes. The H2D^b molecule is in a cartoon representation with the α 1-helix on the back and the α 2-helix removed for better clarity. The peptide is represented in stick conformation with the C terminus on the right. (A) The NPN3A is in purple and (B) the NP₃₆₆ in blue, with the p3 mutation in yellow. (B) Overlay of the peptide-binding cleft for H2D^bNP₃₆₆ and H2D^bNPN3A with the α 1-helix on top and the α 2-helix on the bottom. (C) Contacts with the H-2D^b molecule by (C) P3N in NP₃₆₆ and (D) P3A in NPN3A. The Asn3 mutation to alanine abolishes contacts between the P3 residue of the peptide with His155 and the Tyr156 of the H-2D^b. doi:10.1371/journal.ppat.1001039.g003

and (ii) clonotypes within this region could be highly relevant for cross-reactive $\mathrm{CD8}^{+}\ \mathrm{T}$ cell responses between NP_{366} and NPN3A as they are prominent in both populations. Overall, the mutant and the wt immune V β 8.3 repertoires appear different. Single-cell RT-PCR and sequencing of the CDR3ß region of tetramer⁺V β 8.3⁺CD8⁺ T cells following either HK or HK-NPN3A infections showed that (Table S1) the mutant D^bNPN3A⁺CD8⁺ T cells utilized different J β regions (primary (1⁰): J β 1.1, J β 1.3; secondary (2⁰): J β 1.1, J β 1.6, J β 2.2) in comparison to the wt D^bNP₃₆₆⁺CD8⁺ population (1⁰: J β 2.2, J β 2.4; 2⁰: J β 2.2), and showed evidence of more variable CDR3β loop lengths (8–9 aa). The reduction (relative to the wt $D^{b}NP_{366}$) in V $\beta 8.3$ usage by the D^bNPN3A⁺CD8⁺ T cells reproduces the relative loss of "public" wt TCRs found for the naïve repertoire (Fig 4B). This likely reflects that the N3A mutation has disrupted an "optimal" TCR/ pMHC fit that maximizes antigen-driven clonal expansion. Indeed, the public TCRs were not a prominent feature of the D^bNPN3A⁺CD8⁺ set, being found in only 4 of the 7 NPN3Ainfected mice at a very low frequency, namely 0% in the 1⁰ response and 22.9% in the 2^0 response (Table 2). This is in contrast to the D^bNP₃₆₆⁺CD8⁺ Vβ8.3⁺ TCR repertoire [34] that is largely (90%) comprised of high-frequency, public clonotypes found in all infected B6 mice [34].

These results also establish that the naïve $D^bNPN3A^+ CD8^+$ TCR repertoire (Fig. 4C) is predictive of the immune $D^bNPN3A^+ CD8^+$ TCR β response (Table 2). The relative lack of a "public" response translated to a profile of reduced "sharing" between individual mice, and a total increase in the number of $D^bNPN3A^+CD8^+$ clonotypes due to the 'private' nature of TCR β repertoire for each mouse (Table 2). Even so, the clonotypic diversity within individual NPN3A virus-infected mice was reduced compared to the spectrum found following wt virus-infection. This was true whether the cells were sorted using the wt D^bNP_{366} or the mutant D^bNPN3A tetramer, reflecting the significant cross-reactivity between the D^bNP_{366} and D^bNPN3A -specific populations recovered from mice infected with the NPN3A viruses. By the measure of tetramer binding, it thus seems that the NPN3A virus is selecting a less diverse repertoire than the wt virus, with the repertoire being almost completely cross-reactive with that elicited by the wt D^bNP_{366} epitope.

Similarly, when the $D^bNP_{366}^+V\beta 8.3^+CD8^+$ T cells induced by wt HK infection were sequenced, the majority of the TCR β clonotypes were detected with both the D^bNP_{366} and D^bNPN3A tetramers (Table 3). However, a switch in frequency was seen for some CDR3 β -defined clonotypes, indicating selective binding of particular TCRs by the D^bNPN3A tetramer. For example in M9 and M10 (Table 3) the 'public' SGGANTGQL CDR3 β -sequence dominated the $D^bNP_{366}^+V\beta 8.3^+CD8^+$ T cell population, whereas this sequence was less commonly detected in the same mice by the D^bNPN3A tetramer. The difference could, of course, reflect diversity in TCR α usage. Table 1. Data collection and refinement statistics.

Data Collection Statistics	H2D ^b -NPN3A
Temperature	100K
Space group	1222
Cell Dimensions (a,b,c) (Å)	93.64, 94.98, 132.24
Resolution (Å)	50.00 - 2.60 (2.70-2.60)
Total number of observations	131799 (13218)
Number of unique observations	18241 (1905)
Multiplicity	7.2 (6.9)
Data completeness (%)	98.6 (97.7)
l/σ _t	27.55 (6.37)
R _{merge} ^a (%)	6 (30.3)
Refinement Statistics	
Non-hydrogen atoms	
Protein	3040
Water	46
Resolution (Å)	2.60
R _{factor} ^b (%)	22.1
R _{free} ^b (%)	30.4
Rms deviations from ideality	
Bond lengths (Å)	0.012
Bond angles (°)	1.468
Ramachandran plot (%)	
Most Favoured Region	85.5
Allowed Region	12.5
Generously allowed region	1.7
B-factors (Å ²)	
Average main chain	40.806
Average side chain	40.807

 ${}^{a}R_{merge} = \Sigma |I_{hkl} - \langle I_{hkl} \rangle | / \Sigma I_{hkl}.$

 ${}^{b}R_{factor} = \Sigma_{hkl} ||F_o| - |F_c| |/\Sigma_{hkl}|F_o|$ for all data except \approx 5% which were used for R_{free} calculation. Values in parentheses are for the bin of highest resolution (approximate interval = 0.5 Å).

doi:10.1371/journal.ppat.1001039.t001

Following clonotype selection with the D^bNP₃₆₆ and D^bNPN3A tetramers, sequencing of the secondary $TCR\beta$ repertoire (M11-M13) induced by challenge with the homologous (PR8, then HK) viruses showed less divergence within the wt $D^{b}NP_{366}^{+}CD8^{+}T$ cell specific population. Interestingly, clonotypes like KGGS-NTGQL were enriched by the D^bNPN3A tetramer in some but not other mice (Table 3; present in M9 but not M12) suggesting, again, the likely importance of $V\beta$ - $V\alpha$ chain pairing for recognition of the native D^bNP₃₆₆⁺CD8⁺ T cells with the mutant D^bNPN3A tetramer. Thus, the analysis suggests that only a subset of the repertoire generated by wt infection is able to recognize the D^bNPN3A epitope, though this population is more diverse than that generated in response to the mutant NPN3A virus. All the statistical differences (Table 2, Table 3) were confirmed when the data were standardized to a number of sequences (data not shown).

We further analyzed the D^bNPN3A⁺V β 9⁺CD8⁺ set that was prominent in 2 of the HK-NPN3A secondarily challenged mice. An average of 1.5±1 TCR β clonotypes was found within this population (Table S2). Again, the D^bNPN3A⁺V β 9⁺CD8⁺ TCR β repertoire emerged as essentially restricted and private. However, TCR β analysis within other V β s for NPN3A⁺CD8⁺ T cells would need to be performed to compare the whole TCR repertoires.

Analysis of V α chain usage for the mutant D^bNPN3A⁺CD8⁺ and wt D^bNP₃₆₆⁺CD8⁺ T cells by PCR with a panel of V α specific primers established that those two T cell responses indeed tend to utilise different V α chains. The wt D^bNP₃₆₆⁺CD8⁺ T cell population [36] (Day EB, unpublished) tended to use V α 8 and V α 17.3 (n = 3). Conversely, the D^bNPN3A⁺CD8⁺ TCRs preferentially expressed V α 4, V α 5 and V α 11 (Table S1). While the sample size is small and there are at least 72 different V α chains, these results provide a snapshot of the mutant D^bNPN3A⁺ and wt D^bNP₃₆₆⁺ populations and suggest that there are differences in both V β and V α TCR chain usage.

Response characteristics following heterologous challenge and TCR/pMHC-I avidity

As there was substantial cross-reactivity in vitro (Fig. 2) for the D^bNP₃₆₆ and D^bNPN3A-specific responses, it was important to determine whether memory T cells that cross-react for the D^bNP₃₆₆ and D^bNPN3A epitopes are preferentially recalled by secondary infection with the heterologous virus. Mice that were primed with the PR8NPN3A and then challenged with either the wt HK or mutant HKNPN3A viruses showed equivalent recall of D^bNPN3A⁺CD8⁺ T cells during the acute phase of the secondary response. This was detected by IFN- γ production (Fig 5A) and tetramer staining (data not shown) and is consistent with the TCR $CDR3\beta$ analysis (Table 2). Conversely, when mice were firstly primed with wt PR8, then later infected i.n. with either the wt HK or mutant HK-NPN3A, the D^bNP₃₆₆⁺CD8⁺ T cells were differentially recalled indicating that (in the absence of primary selection from the naïve repertoire by the mutant epitope) only some of the D^bNP₃₆₆⁺CD8⁺ memory T cells that were expanded by heterologous challenge bind D^bNPN3A (Fig. 5A). Interestingly, wt priming and challenge $(1^{0} \text{ PR8->}2^{0} \text{ X31})$ resulted in significantly higher CD8⁺ T cells numbers (Fig. 5B) than were found for any secondary CD8⁺ T cell responses after NPN3A priming (1º PR8-NPN3A->2º X31-NPN3A and 1º PR8-NPN3A- $>2^0$ X31). These results lead to two main conclusions: (i) priming with the wt virus elicits CD8⁺ T cells that respond relatively well to a subsequent infection with cross-reactive variant (ie 1^0 PR8-> 2^0 $X31-NPN3A = 1^{\circ} PR8-NPN3A \ge 2^{\circ} X31-NPN3A$; (ii) priming with the cross-reactive variant can be detrimental as the diminished primary response may limit the full expansion of CD8⁺ T cells that are able to respond to the subsequent wt infection (ie 1° PR8-NPN3A-> 2° X31 is lower than 1° PR8-> 2° X31) and skew the TCR usage. Thus, using NPN3A for either priming or the challenge gives an equally poor response.

To determine whether such decreased CD8⁺ responses elicited after heterologous challenge would affect influenza virus clearance, we performed experiments to examine the protective efficacy of cross-reactive CD8⁺ T cell repertoires. We performed prime-andchallenge studies in µMT mice lacking B cells to ensure that antibody responses did not mask any possible inhibitory effects of "suboptimal" TCRs on viral clearance. As suggested by CD8⁺ T cell data, assessment of lung viral titres showed delayed viral clearance on d6 after the secondary infection in case of heterologous prime-and-boost (PR8->X31-NPN3A and PR8-NPN3A->X31) compared to homologous infections (PR8->X31 or PR8-NPN3A->X31-NPN3A) (Fig. 5B). These results indicate that recall of "suboptimal" TCRs for a single T cell specificity can lead to delayed viral clearance, despite the presence of other influenza CD8⁺ T cell responses (D^bPA₂₂₄⁺CD8⁺, D^bPB1₆₂⁺CD8⁺, $K^{b}NS1_{114}^{+}CD8^{+}, K^{b}PB1_{703}^{+}CD8^{+}).$



Naïve N3A⁺Vβ8.3⁺CD8⁺ T cell clonotypes

CDR3β seq	Jβ	aa	Found in #Mice	Found in #Sequences
SGGANTGQL	2.2	9	• 1	1
SGGSNTGQL	2.2	9	2	2
SGGGNTGQL	2.2	9	1	1
SGGGKTGQL	2.2	9	1	1
RRDRGGNTL	2.2	9	2	2
SDSKNTEV	1.1	8	1	1
SDARTEV	1.1	8	1	1
RTGRGANSDY	1.2	10	1	1
SEGGGRYAEQ	2.1	10	1	1
SDALQGSQNTL	2.4	10	1	1
SADSSYEQ	2.6	8	1	1
SDASQYEQ	2.6	8	1	1
Total number			. 9	14

*public D^bNP₃₆₆-specific TCR clonotypes. Experimental details are contained in Supplementary Table 3.

Figure 4. TCR repertoire of naïve D^b **NPN3A⁺CD8⁺ T cells.** Effects of the N3A substitution within NP₃₆₆₋₃₇₄ on (A) naïve T cell precursor frequency, (B) V β 8.3 bias and (C) naïve TCR β repertoire within the D^b NPN3A⁺V β 8.3⁺CD8⁺ set. (A) Naïve D^b NPN3A⁺ and D^b NP₃₆₆⁺CD8⁺ T cell precursors within lymph nodes and spleens were identified with the D^b NPN3A⁻PE and D^b NP₃₆₆⁻PE tetramers using an established tetramer enrichment protocol [53]. (B) Naïve D^b NPN3A⁺CD8⁺ T cells were assessed for the conservancy of V β 8.3 bias. (C) a CDR3 β sequences within the naïve D^b NPN3A⁺V β 8.3⁺CD8⁺ T cells were determined by a single-cell PCR, amplification of V β 8.3 transcripts and sequencing. doi:10.1371/journal.ppat.1001039.g004

These patterns of complete, or partial, cross-stimulation following *in vivo* virus challenge were reflected in the results found for *in vitro* measurements of "functional avidity" (Fig 6). Pulsing immune spleen cells recovered directly *ex vivo* with graded doses of wt or mutant peptide in the ICS assay showed comparable levels of IFN- γ induction (Fig 6AC) in every situation but one, the exposure

HK-NPN3A infection		aa	1 [°] response				2 ⁰ response									
			M1		M2		М3		M4		M5		M6		M7	
			NP	N3A	NP	N3A	NP	N3A	NP	N3A	NP	N3A	NP	N3A	NP	N3A
SGGANTGQL	252	9							3			9		2		5
SGGSNTGQL	252	9										11	1		2	3
SGGGSTGQF	252	9					11									
SGGGNTGQL	252	9							12	1.5			99	3		
SGGGNNGHL	252	9					2									
KGGGNTGQL	252	9									100	80				
SDAASTEV	1S1	8			84	88										
SDAANTEV	1S1	8			16	12			19	11						
SDAVATEV	1S1	8							62	82						
SDASSTEV	1S1	8													98	92
SDSANTEV	1S1	8							3	4.5						
RRDRGGNTL	153	9	95	100												
SGGTENSPL	156	9												95		
SDAQLYAEQ	251	9	2													
SVGGRAEQ	2S1	8								1.5						
SDWGSQNTL	2S4	9					87	100								
SDGGGTYEQ	2S5	9	2													
TOTAL sequences			44	42	32	43	64	32	34	66	59	35	72	61	50	61

Table 2. Frequency of TCR β clonotypes in D^bNPN3A⁺ V β 8.3⁺CD8⁺ T cells after 1⁰ (M1, M2) and 2⁰ (M3 to M7) mutant HK-NPN3A infection detected with either the D^bNP₃₆₆⁺ or D^bNPN3A⁺ tetramer.

M: individual mouse; NP: D^bNP₃₆₆ tetramer; N3A: D^bNPN3A tetramer.

 1^{0} responses were generated by i.n. HK-NPN3A infection of mice; 2^{0} responses were generated by priming mice with i.p. PR8-NPN3A virus then challenging with i.n. HK-NPN3A virus; $D^{b}NP_{366}$: complex of H2D^b and NP₃₆₆₋₃₇₄ peptide; $D^{b}NPN3A$: complex of H2D^b and NPN3A₃₆₆₋₃₇₄ peptide.

doi:10.1371/journal.ppat.1001039.t002

Table 3. Frequency of TCR β clonotypes in D^bNP₃₆₆⁺V β 8.3⁺CD8⁺ T cells after 1⁰ (M8–M10) and 2⁰ (M11–M13) wt influenza virus infection detected with either the D^bNP₃₆₆⁺ or D^bNPN3A⁺ tetramer.

HK infection			1 ⁰ response							2 ⁰ response						
	_		M8		M9	M9		M10		M11		M12		M13		
CDR3β	Jβ	aa	NP	N3A	NP	N3A	NP	N3A	NP	N3A	NP	N3A	NP	N3A		
SGGANTGQL	252	9	67	70	53	3	63	22	10	15	36	48	24	22		
SGGSNTGQL	252	9	22	16	2	17	27		50	13	2		50	16		
SGGGNTGQL	252	9	11	7										5		
SGGGSTGQL	252	9				17	7									
RGGSNTGQL	252	9			2						4					
RGGANTGQL	252	9							5	13	15	36	6	11		
RGGGNTGQL	252	9										4				
RGGAPTGQL	252	9							10							
KGGSNTGQL	252	9				63	3				38	2				
KGGGNTGQL	252	9							25	60	4	4		3		
SGGQGNSPL	252	9										2				
KAGGSTGQL	252	9						48								
KAGGGTGQL	252	9		5												
RALGRNTEV	151	9			2			3								
SDAGKTEV	1S1	8											6	30		
RDSANTEV	151	8										2	15	8		
SDAGAEQ	251	7		2										5		
SDWGWQNTL	254	9			40			27								
TOTAL sequences			27	43	45	30	30	60	40	47	47	44	34	37		

M: individual mouse; NP: D^bNP₃₆₆ tetramer; N3A: D^bNPN3A tetramer;

 1^{0} responses were generated by i.n. HK infection of mice; 2^{0} responses were generated by priming mice with i.p. PR8 viruses then challenging i.n. with the HK virus; $D^{b}NP_{366}$: complex of H2D^b and NP₃₆₆₋₃₇₄ peptide; $D^{b}NPN3A$: complex of H2D^b and NPN3A₃₆₆₋₃₇₄ peptide.

*Predominant: ≥15%, [#]Common: present in all mice sampled

doi:10.1371/journal.ppat.1001039.t003

of wt-primed T cells to the NPN3A peptide (Fig 6B). Thus, while the immune repertoire selected by D^bNP3A shows evidence of equivalent avidity following stimulation with the NPN3A or NP₃₆₆ peptides, D^bNP_{366} induces a response that is of higher avidity for the wt than the mutant epitope. The same effect was seen even more clearly when three (all expressing the SGGGNTGQL CDR3 β) T cell hybridoma lines [37] specific for D^bNP_{366} were stimulated with the two peptides (Fig 6DE). This result is in accord with findings from both the TCR repertoire analysis of crossreactive clonotypes, assessed by tetramer binding (Table 2), and the response magnitudes determined following homologous and heterologous virus challenge (Fig 5). Thus, priming and recall of cross-reactive CD8⁺ T cells recognising mutant viral epitopes reflects functional (defined as responsiveness to a peptide) pMHC-TCR avidity.

To determine the pMHC-I avidity of the responding $D^bNP_{366}^+CD8^+$ and NPN3A⁺CD8⁺ T cells, we additionally performed tetramer dilution (Fig. 7A–C) and tetramer dissociation (Fig. 7D–E) assays. While tetramer dissociation assay measured the "off-rate" component of pMHC-I avidity, tetramer dilution technique assessed the overall pMHC-I avidity (both the "on" and "off" rates). Furthermore, we also assessed CD8β-dependence for functional activation (a measure of low avidity CD8⁺ T cells) of both wt D^bNP₃₆₆⁺CD8⁺ and the mutant NPN3A⁺CD8⁺ T cells by anti-CD8β mAb blocking, followed by IFN- γ ICS (Fig. 7G–I). Our data obtained from those three additional measures of pMHC-I avidity confirmed the results obtained by the peptide titration

combined with ICS (functional avidity, Fig. 6) and further suggested significantly lower pMHC-I avidity of the wt $\rm D^bNP_{366}{}^+CD8^+$ (generated by the wt HK infection) for NPN3A variant.

Discussion

The P3N within the immunodominant influenza virus-specific $D^{b}NP_{366}$ epitope is a partially solvent exposed, and non-prominent for TCR binding, residue that is predominantly buried within the MHC cleft [21,23]. The NPN3A mutation leads to both decreased recruitment of CD8⁺ T cells and a narrowed clonotype selection profile within V β 8.3 and V β 9 regions. Structurally, the mutation leads to a loss of a number of contacts between the NPN3A peptide and the MHC-I molecule, including a contact with the gate-keeper residue at position 155, and unaltered stability of the H-2D^b-NPN3A complex. The fact that the NPN3A mutation affects contacts with the MHC-I at His155, known to play an important role in TCR-pMHC structures in general, is likely to indirectly compromise the TCR recognition. By loosing the bond with the Asn3, His155 may gain more flexibility and thus be inappropriately placed for the subsequent TCR ligation onto the NPN3A peptide. Alternatively, it is also possible that a small part of the solvent-exposed head group of the Asn3 residue in the wt NP_{366} peptide might, to some extent, be directly interacting with the TCR following ligation.



Figure 5. Heterologous 2° challenge with the wt HK and mutant HK-NPN3A viruses. Naïve (A) B6 or (B) μ MT mice were primed i.p with the mutant PR8-NPN3A and challenged i.n. 6 weeks later with 1×10^4 p.f.u. of either the wt HK or the mutant HK-NPN3A virus. Alternatively, naïve mice were primed i.p. with the wt PR8 virus and challenged i.n. 6 weeks later with 1×10^4 p.f.u. of the wt HK or the mutant HK-NPN3A virus. (A) The magnitude of the CD8⁺ T cell response measured 8d after 2^o infection was determined by the ICS assay for IFN- γ . Data represent mean \pm SD (n = 5) (B) Viral clearance after the secondary homologous (PR8->X31 and PR8-NP-N3A>X31-NPN3A) or heterologous (PR8->X31-NPN3A and PR8-NP-N3A>X31-NPN3A) or heterologous (PR8->X31-NPN3A and PR8-NP-N3A>X31) challenge. Lungs were sampled at days 5 and 6 after secondary infection and homogenized for virus titration by plaque assay on MDCK cell monolayers. Data represent the mean and n = 3–5 mice per group. *= p<0.01 #= p<0.05.

Surprisingly, despite the loss of several contacts between NPN3A peptide and H-2D^b, the stability of the peptide-MHC-I complex remains constant for both NP₃₆₆ and NPN3A. This suggests that the Asn3 as a secondary anchor does not play an important role in stabilizing the peptide within the MHC-I. The structural basis for the diminished recruitment of D^bNPN3A-specific CD8⁺ T cells is thus likely to rest in the way that the partially-solvent exposed residue contacts MHC-I and modifies TCR ligation.

The emerging $D^bNPN3A^+CD8^+$ T cell population was characterised by different V α and V β preference, distinct CDR3 β sequences and a lower overall TCR diversity in comparison to wt $D^bNP_{366}^+CD8^+$ T cells. These findings suggest that a very limited spectrum of CD8⁺ T cells can recognise the D^bNPN3A mutant structure. Interestingly, a similar P3 substitution in the influenza virus D^bPA₂₂₄ epitope had no affect on D^bPA₂₂₄⁺CD8⁺ T cell recognition and recruited a diverse array of TCR clones comparable to the spectrum found for the wt response [21]. This suggests that the partially-exposed P3 residue plays a greater structural and/or TCR recognition role for the "featureless" D^bNP_{366} than for the "featured" D^bPA_{224} complex reflecting, in turn, the more limited spectrum of TCRs that bind D^bNP_{366} [21]. Taken together, it appears that partially-exposed residues within viral peptides can provide important contacts with the MHC-I, which can in turn cause remote effects that modify antigenicity for the TCR-pMHC-I complex and impact on both TCR repertoire selection and the magnitude of CD8⁺ T cell responses.

Interestingly, there were no differences in function or phenotype characteristics for the $D^bNP_{366}^+CD8^+$ and NPN3A⁺CD8⁺ T cells, although those two CTL sets had a high proportion of different TCR clones. This is in accordance with previous studies showing that the simultaneous production of antiviral cytokines [29] and IL-7R [20] expression is antigen dose- rather than magnitude-related. Conversely, levels of the CD62L [38,39,40] activation marker differed for the $D^bNP_{366}^+CD8^+$ and NPN3A⁺CD8⁺ populations, indicating that response magnitude has some relationship to the activation status



Figure 6. Decreased TCR functional avidity of D^bNPN3A-primed T cells for D^bNP₃₆₆. (A–C) Responsiveness of CD8⁺ T cells to varying peptide concentrations was determined as a measure of functional TCR avidity for the cognate pMHCl complex. (A,B) D^bNP₃₆₆⁺ CD8⁺ and (A,C) D^bNPN3A CD8⁺ T cells recovered from mice 2^o-challenged with wt or NPN3A viruses were assessed for functional TCR avidity using the ICS assay. The enriched splenocytes analysed by IFN- γ ICS were stimulated *in vitro* with 4-fold dilutions of the NP₃₆₆, or NPN3A peptide. Data represent mean±SD (n = 4), * = p<0.1. (D,E) The response profiles of monoclonal D^bNP₃₆₆-specific LacZ-inducible (22) T cell hybridomas (3 distinct lines all with the "public" SGGNTGQL CDR3 β sequence) to peptide-pulsed H-2D^b splenocytes presenting the (D) wt D^bNP₃₆₆ or (E) D^bNPN3A epitopes were assessed in LacZ assay [22]. doi:10.1371/journal.ppat.1001039.g006

of CD8⁺ T cells [40,41,42] which may, perhaps, reflect the extent of CTL proliferation.

The TCR repertoires specific for D^bNP₃₆₆ and D^bNPN3A appear to be quite distinct. The response overall for wt D^bNP₃₆₆⁺CD8⁺ T cells is characterised by conserved, "public" clonotypes that constitute the majority $(83.5\% \text{ in } 1^0 \text{ response and}$ 92.3% in 2^0 response) of the selected TCR repertoire [34]. These public clonotypes are not a prominent feature of the D^bNPN3A⁺CD8⁺ set, being found only in 4/7 NPN3A-infected mice at very low frequency. Since we know that these TCR clonotypes are present in all B6 mice [34], the difference presumably reflects the lower TCR avidity for D^bNPN3A, as indicated by the T cell hybridoma analysis where they were shown to require 1000 times more NPN3A than NP366 peptide for optimal stimulation. Since the public clonotypes cannot be efficiently recruited into the immune response by the mutated N3A virus, this could have created a "hole" in TCRs capable of recognising the mutated epitope, which subsequently can lead to a reduction in T cell immunogenicity [43]. However, though both the D^bNP₃₆₆⁺CD8⁺ and D^bNPN3A⁺CD8⁺ T cell responses are characterised by quite distinct TCR repertoires, the majority are bound by both the D^bNP₃₆₆ and D^bNPN3A tetramers and can be detected by stimulation with either the NP_{366} or the NPN3A peptides, suggesting that a clonal dissection of TCR clonotypes is needed to make a valid interpretation about the truly crossreactive CD8⁺ T cell responses. These findings also raise questions concerning the true correlation between pHMC-I tetramer binding in vitro and the in vivo selection of a responding TCR repertoire.

Overall, the results indicate that a loss of a number of contacts between the NPN3A peptide and the MHC-I molecule and lower functional and structural pMHC-I avidity (for wt D^bNP_{366}) D^bNPN3A selects a narrowed TCR repertoire of "best fit" TCRs from a spectrum of naïve clonotypes that, once activated, clonally

expanded and engaged in an immune response, have sufficient avidity to be recalled by exposure to the wt D^bNP_{366} epitope. Conversely, the "better" fit D^bNP_{366} finds a sufficient spectrum of high avidity TCRs within that available naive repertoire and does not (likely because of clonal competition) select most of the TCR $\alpha\beta$ pairs that interact optimally with D^bNPN3A . Priming with the wt virus thus establishes memory for only a very limited secondary response to the mutant. Similar to our results, subtle variations within the anchor residue of H^b peptide/I-E^k also decreased peptide-MHC class II affinity and the activation of responding T cells [44].

Thinking about this in terms of possible vaccination strategies for use against rapidly changing viruses or tumor epitopes, it appears that priming with cross-reactive mutants that have characteristics comparable to NPN3A would be of no benefit (or even could be detrimental as evidenced by delayed viral clearance) as the same level of T cell immunity against such mutants can be elicited by exposure to the wt epitope. On the other hand, changes like the non-cross-reactive NP-M6A mutation [22] that induce a completely novel, high quality response might merit incorporation in an experimental vaccine or immunotherapy strategy. Overall, working out the topographical constraints that determine these differential response profiles would seem eminently worthwhile.

A further reason for defining the structural rules governing TCR cross-recognition is that similar effects have been found for different epitopes derived from unrelated viruses. Published studies provide evidence for cross-reactive CD8⁺ T cell responses between influenza A virus and Epstein-Barr virus (EBV) [45], influenza A virus and Hepatitis C virus (HCV) [46,47] or lymphocytic choriomeningitis virus (LCMV) and Pichinde virus (PV) [48]. Such heterologous cross-reactive immunity can unintentionally skew TCR recruitment, result in a narrow TCR repertoire and subsequent viral escape [48] as well as influenza disease severity



Figure 7. Decreased pMHC avidity and CD8*β***-independence for functional avidity of D^bNPN3A-primed T cells for D^bNP₃₆₆.** pMHC-TCR avidity was assessed by three measures: (A–C) the overall pMHC avidity (tetramer "on" and "off" rates) by tetramer dilution, (D–F) tetramer "off" rate by tetramer dissociation and (G–I) CD8*β*-dependence by anti-CD8*β* blocking combined with ICS. Splenocytes were obtained from mice 2^o-challenged with wt or NPN3A viruses. (A–C) D^bNP₃₆₆⁺ CD8⁺ and D^bNPN3A CD8⁺ T cells were stained with 2-fold dilutions of PE-conjugated tetramers, followed by anti-CD8 staining. (D–F) Cells were stained with either the D^bNP₃₃₆ or D^bNPN3A tetramers at the saturating concentration, then incubated at 37°C with a mAb to H2D^b to prevent rebinding of dissociated tetramer. The progressive diminution in tetramer staining was measured by flow cytometric analysis. (G–I) Lymphocytes were pre-cultured in the presence or absence of anti-CD8*β* antibody (5 µg/ml). Cells were then stimulated for 5 h with peptide, IL-2 and GolgiStop also in the presence or absence of anti-CD8*β* antibody (5 µg/ml). Following stimulation, cells blocking mAb, (H) mean fluorescence intensity (MFI) of IFN-γ staining, (I) the percentage of CD8⁺ T cells producing IFN-γ in the presence or absence of anti-CD8*β* for IFN-γ production. Tetramer staining was performed in the presence of NAAz, then washed and incubated with anti-CD8 mAb. The progressive diminution in tetramer staining was measured. The Td50 value defining the time to 50% tetramer loss. Data represent mean±SD (n=4–5 mice per group), *p<0.01; #p<0.05.

doi:10.1371/journal.ppat.1001039.g007

[45]. The topic of TCR cross-reactivity in CD8⁺ T cell responses clearly merits more attention.

Materials and Methods

Ethics statement

All animal experimentation was conducted following the Australian National Health and Medical Research Council Code of Practice for the Care and Use of Animals for Scientific Purposes guidelines for housing and care of laboratory animals and performed in accordance with Institutional regulations after pertinent review and approval by the University of Melbourne Animal Ethics Experimentation Committee in Melbourne.

Mice and viral infection

C57BL/6J (B6, H2^b) and μ MT mice were bred and housed under specific pathogen free conditions at the Department of Microbiology and Immunology, University of Melbourne. For the generation of acute influenza responses mice were lightly anaesthetised by inhalation of methoxyflurane and infected intranasally (i.n.) with 1×10^4 plaque forming units (p.f.u.) of HK-X31 (H3N2, X31) or modified HK-X31 (HK-NPN3A) influenza A viruses in 30µl of PBS. For recall responses mice were first primed intraperitoneally (i.p.) with 1.5×10^7 p.f.u. of the serologically distinct PR8 (H1N1) or modified PR8 (PR8-NPN3A) influenza A viruses, in 500 μ l of PBS. Viruses share the same internal components for NP and PA from which CD8 epitopes are derived [49]. Virus stocks were grown in the allantoic cavity of 10 day old embryonated chicken eggs, from which the viral titre was determined by plaque assay on monolayers of Madin derby canine kidney (MDCK) cells.

Generation and titration of recombinant influenza viruses

Recombinant influenza viruses with the single amino acid substitution (N3A) within the NP₃₆₆ peptide, ASNENMETM, were generated using the eight-plasmid reverse genetics system [50]. The substitution was first incorporated by site directed mutagenesis using PCR primers encoding N3A₃₆₆ peptide, ASAENMETM, and the opposite primer encoding NP. Recombinant PCR products encoding N3A₃₆₆ were digested with BsmB1 and ligated into the alkaline phosphatase treated pHW2000 vector. Recombinant viruses (HK-NPN3A and PR8-NPN3A) were rescued following transfection of MDCK-293T cell coculture with the eight plasmids encoding influenza segments. Viruses were then amplified in the allantoic cavity of 10-day old embryonated chicken eggs, and the viral titre determined by plaque assay of allantoic fluid infecting monolayers of MDCK cells. There was any evidence of altered fitness *in vivo* for the HKNPN3A virus, as the kinetics of virus growth and clearance following i.n. challenge of naïve B6 mice were found to be equivalent for the wt HK and mutant viruses (Fig S3A). Similarly, the levels of CTL activity found using target cells infected with HKNPN3A or the HK wt viruses were comparable, and the same was seen for peptide pulsed cells, suggesting that antigen presentation of NPN3A peptide remains constant (Fig S3 BC).

Determination of viral titres

Lungs taken from mice after primary viral infection (Fig. S3A) or prime-and-boost approach using homologous (PR8->X31 and PR8-NPN3A->X31-NPN3A) or heterologous (PR8->X31-NPN3A and PR8-NPN3A->X31) strategy (Fig. 5B) were homogenised and the virus-containing supernatant above the cell debris was harvested and stored at -70° C. Titres of infectious virus in the lung supernatants were determined by plaque assay on monolayers of MDCK cells.

Tissue sampling and cell preparation

Spleen and bronchoalveolar lavage (BAL) samples were recovered from mice at acute phases of the primary and secondary infections (d10 and d8 respectively), and the BAL samples were incubated on plastic petri-dishes for 1hr at 37^{0} C to remove macrophages. The spleens were disrupted and enriched for CD8⁺ T cells using goat anti-mouse IgG and IgM antibodies (Jackson ImmunoResearch Labs, West Grove, PA, USA). For assessment of naïve precursor frequency of N3A₃₆₆⁺CD8⁺ T cells, spleens and lymph nodes (inguinal, brachial, axillary, superficial cervical, and mesenteric) were collected from naïve mice and processed to single-cell suspensions.

Tetramer and phenotypic staining of CD8⁺ T cells

Lymphocytes from the BAL and spleen were stained with tetramers conjugated to Strepavidin-APC or PE (Molecular Probes, Eugene, OR, USA) at optimal staining concentrations $(10 \ \mu g/ml \ D^b NP_{366}, 40 \ \mu g/ml \ D^b N3A_{366}, and 10 \ \mu g/ml \ D^b PA_{224})$ tetramers) for 1hr at room temperature. Cells were washed twice in FACS buffer, and stained with 1 µg/ml CD8-PerCP Cy5.5, 5 µg/ml CD62L-FITC and 5 µg/ml CD127-APC mAbs (BD Biosciences) for 30 mins on ice, washed twice and analysed by flow cytometry on the FACS Calibur (BD Immunocytometry) and analysed by CellQuest Pro software (BD Immunocytometry). We titrated all the batches of all the tetramers used in this study. We used tetramers at optimal concentrations (10-40µg/ml) based on both the percentage of epitope-specific CD8⁺ T cells and the mean fluorescence intensity (MFI) of tetramer staining. A Scatchard analysis [51] based on the tetramer dilution as say (Fig. 7 A–C) was also plotted (Fig. S4) and confirmed our observations from routine tetramer titrations that the D^bNP₃₆₆ tetramer displays slightly superior pMHC binding capacities over the NPN3A tetramer at a concentration $<5\mu$ g/ml.

Tetramer dilution and tetramer dissociation analyses

CD8⁺ T cells from spleen were stained with the D^bNP₃₆₆ and D^b-NPN3A tetramers conjugated to Streptavidin-PE (Molecular Probes, Eugene, OR) for 60mins at room temperature. For a tetramer dilution assay, 2-fold dilutions of PE-conjugated tetramers were used at a range of concentrations (0.15–40 μ g/ml). For a tetramer dissociation assay, lymphocytes were stained at the optimal concentration of PE-conjugated tetramers as assessed by tetramer titration as determined by both the percentage of tetramer⁺CD8⁺ T cells and mean fluorescence intensity (MFI). Cells were washed twice in FACS buffer (10%BSA/0.02% NaAz

in PBS), stained with a FITC-conjugated mAb to CD8 α (BD Biosciences Pharmingen) for 30mins on ice, washed and analysed by flow cytometry. As a measure of pMHC avidity, splenic T cells were used in tetramer dissociation assay [29]. After staining with tetramer, T cells were washed and incubated in the presence of anti-H2D^b antibody at 5µg/ml at 37^oC to prevent tetramer rebinding. Cells were removed at intervals, stained with the FITC-conjugated mAb to CD8 α and analysed by flow cytometry. Loss of tetramer⁺CD8⁺ T cells at particular time-points was calculated in comparison to tetramer staining at t = 0mins.

Peptide stimulation and intracellular cytokine staining

Enriched T cell populations from spleen and BAL were stimulated with one of the NP₃₆₆, N3A₃₆₆, PA₂₂₄ or PB1₇₀₃ peptides (AusPep) for 5 hrs at 37°C, 5% CO₂ in the presence of 1µg/ml Golgi-Plug (BD Biosciences Pharmingen) and 10U/ml recombinant human IL-2 (Roche, Germany) (BD Biosciences). Cells were washed twice with FACS buffer, stained with CD8-PerCP Cy5.5 for 30 mins on ice, fixed, permeabilised and stained with anti-IFN- γ -FITC (5µg/ml), TNF- α -APC (2µg/ml), and IL-2-PE (2µg/ml) mAbs (Biolegend). Samples were acquired using flow cytometry, and the total cytokine production calculated by subtracting background fluorescence using no peptide controls. In selected experiments, lymphocytes were stimulated with varying concentrations of peptides, three-fold dilutions ranging from 300nM to 0.0008nM to determine the sensitivity specific peptides, defined as 'functional avidity' [52].

TCR avidity for pMHC complex by CD8β-dependence

Splenocytes were obtained from mice sampled on d6 after secondary infection. Lymphocytes were pre-cultured in the presence or absence of anti-CD8 β antibody (53.5–8) (10 µg/ml). Cells were then stimulated for 5 h with peptide, IL-2 and GolgiStop also in the presence or absence of anti-CD8 β antibody (5 µg/ml). Following stimulation, cells were analysed for CD8 and IFN γ expression. Shown is the percentage of CD8⁺ cells producing IFN- γ after stimulation in the presence of anti-CD8 β blocking mAb.

Determination of N3A₃₆₆⁺CD8⁺ T cell precursor frequency

Naïve N3A366-specific CD8⁺ T cells were identified as described [33,53]. Briefly, processed lymph nodes and spleen samples were resuspended in 100 µl of Sorter buffer, 100 µl FcR block (24G2 and CD16 culture supernatant, 1% mouse and 1% rat serum) was added, and clumps of dead cells were discarded. Tetramers at optimal staining concentrations (D^bN3A₃₆₆-PE at 40µg/ml and $D^{b}NP_{366}$ -PE at 10µg/ml) were added to the cell mix and incubated for 1 hour at room temperature in the dark. Cells were washed and resuspended in 400 µl buffer with 100 µl anti-PE microbeads (Miltenyi Biotec), and incubated at 4°C for 25 mins. Following two washes, cells were resuspended in 3 ml of buffer and cells that had bound the microbeads were purified on a magnetic LS column according to manufacturers instructions (Miltenyi Biotec). Cells eluted from the column were centrifuged $(515 \times g, 6 \text{ min}, 4^{\circ}\text{C})$, supernatant carefully aspirated to leave 90 µl buffer remaining, and 10 µl antibody cocktail was added for 30 min at 4°C. The antibody cocktail contained anti-CD8-APC Cy7 (Pharmingen, BD), anti-CD4-PE Cy7 (eBiosciences), anti-B220-FITC (Pharmingen, BD), anti-CD11b-FITC (eBiosciences), anti-CD11c-FITC (eBiosciences), anti-F4/80-FITC (eBiosciences), anti-CD62L-APC (Pharmingen, BD), and anti-CD3-PerCP Cy5.5 (Pharmingen, BD) mAbs. Cells were washed in 2 ml buffer, centrifuged $(515 \times g, 6 \text{ min}, 4^{\circ}\text{C})$, and supernatant aspirated leaving 100 µl. Resuspended cells were passed through 45 µm

sieve, and data acquired by flow cytometry on the LSR II (Becton Dickinson) and analysed with FlowJo (Treestar Inc.) software. In selected experiments, naïve NPN3A⁺V β 8.3⁺CD8⁺ T cells were single-cell sorted for TCR β analysis. Experimental details of the single cell RT-PCR designed to amplify naïve NPN3A⁺V β 8.3⁺CD8⁺ T cells using 2 different sorting strategies are listed in Table S3.

Hybridoma LacZ assay

LacZ-inducible T cell hybridomas specific for NP₃₆₆ peptide [37,54] were resuspended at 1×10^{6} cell/mL and aliquots (100µl) and dispensed into 96-well flat-bottom plates together with 5×10^{5} naïve splenocytes (APCs). Cells were cultured in the presence of NP₃₆₆ or NPN3A peptides at concentrations ranging from 10^{-15} M to 10^{-4} M for 18h at 37^{0} C. The cells were then washed with PBS, fixed with 100µl of 2% formaldehyde/0.2% glutaral-dehyde in PBS for 5min on ice, washed in PBS and incubated with 2.5mg of 5-bromo-4-chloro-3-indolyl β-D-galactoside (X-gal) for 16h at 37^{0} C. The LacZ⁺ hybridomas were then counted using a light microscope, and the number of NP₃₆₆-specific hybridomas was calculated by subtracting the "background" LacZ expression for cells cultured in the absence of the peptide.

Isolation of single cell CD8⁺ T cells, RT-PCR and sequencing

Splenocytes were stained with 10 μ g/ml D^bNP₃₆₆ or 40 μ g/ml D^bN3A₃₆₆-PE tetramer in sort buffer (PBS with 0.1% BSA) for 60 mins at room temperature, washed and stained with 1 μ g/ml anti-CD8-Allophycocyanin and 10 μ g/ml of either anti-V β 8.3 or anti-V β 9 for 30 mins on ice, washed twice with sort buffer. Single lymphocytes were isolated by sorting with a FACS Aria (BD Immunocytometry), into 80 wells of an empty 96 well twintec plate (Eppendorf). mRNA transcripts were reversed transcribed to cDNA, using a Sensiscript kit (Qiagen) according to manufacturers instructions, and the CDR3 β region amplified by a nested hot start PCR using V β primers [34]. Positive PCR products were purified using Qiagen PCR purification kit and sequenced.

Protein expression, purification, crystallisation and structure determination

H2-D^b and β 2-microglobulin molecules were expressed in *Escherichia Coli* as inclusion bodies, refolded with the NP-N3A (ASAENMETM) peptide and purified as previously described [55,56]. The H2D^bNP-N3A complex crystals were obtained at 3mg/ml by the hanging-drop vapour diffusion technique at 20°C. Crystals were grown with a reservoir containing 0.1 M sodium citrate at pH 5.7, 28% PEG 3350 (w/v), 0.2 M lithium sulphate. The crystals belong to space group **I222** and the unit cell dimensions were consistent with one molecule per asymmetric units (Table S1).

The crystals were flash frozen to a temperature of 100K before data collection in-house with a Rigaku RU-200 rotating-anode Xray generator. The data were processed and scaled with the XDS [57]. The crystal structure was solved using the molecular replacement method using the program Phaser [58] from the CCP4 suite of programs [59]. The search probe used to solve the structure was the structure of mouse MHC class I H2-D^b minus the peptide (Protein Data Bank accession number 3CPL) [22]. The progress of refinement was monitored by the R_{free} value with neither a sigma nor a low-resolution cutoff being applied to the data. The refinement protocol used includes several cycles of refinement with REFMAC [59] followed by manual model rebuilding with O program [60]. 'Translation, liberation and screw-rotation' displacement refinement was used to model anisotropic displacements of defined domains was used during the refinement process. The electron density around the NPN3A peptide was unambiguous, and all the side chains were built at full occupancy. Some mobile loops in the heavy chain of the H-2D^b molecule (residues 191–201; 220–228 and 247–254) have been removed from the final model due to missing electronic density. Final refinement statistics are summarized in Table I, the coordinates of the H2Db-NP-N3A complex have been deposited with the Protein Data Bank under accession numbers 3FTG.

Thermostability measurements of recombinant class I complexes using circular dichroism (CD)

Circular Dichroism Spectra were measured on a Jasco 815 spectropolarimeter using a thermostatically controlled cuvette. A far-UV spectra was collected from 190nm to 250nm. The UV minimum was determined as 219 nm for H2Db-NP-N3A. The measurements for the thermal melting experiments was made at the minimum for H2Db-NP-N3A, at intervals of 0.1° C at a rate of 1° C/min from 20°C to 90°C. The Jasco Spectra Manager software was used to view and smooth the traces and then the GraphPad Prism software was used to plot Temperature versus % unfolded. The midpoint of thermal denaturation (Tm) for each protein was determined as the point at which 50% unfolding was achieved. The measurements were done in duplicate at two concentrations (4 μ M and 2.2 μ M) in a solution of 10mM Tris pH 8, 150mM NaCl.

Protein Data Bank accession number

The atomic coordinates have been deposited in the Protein Data Bank, www.pdb.org (PDB ID code 3FTG).

Supporting Information

Figure S1 Phenotypic and functional characteristics of NPN3A⁺CD8⁺ T cells. (A–D) Tetramer⁺ CD8⁺ T cells were characterized for (A, B) CD62L and (C, D) IL-7R α expression following primary (A, C; d10) or secondary (B, D; d8) challenge. (E–H) The ICS assay was used to measure cytokine production following in vitro stimulation with the cognate peptide following primary (E, G) or secondary (F, H) challenge for TNF- α^+ (E, F) and IL-2⁺ within the IFN- γ^+ set (G, H). Data represent mean±SD from 5 mice per group, *=p<0.01.

Found at: doi:10.1371/journal.ppat.1001039.s001 (2.47 MB TIF)

Figure S2 TCR Vβ usage in primary and secondary NPN3A⁺ CD8⁺ and D^bNP₃₆₆⁺CD8⁺ T cell responses. Primary (A, C) or secondary (B, D) responses were generated by infection with either the (A, B) HK-NPN3A or (C, D) HK virus. Splenocytes were stained with the (A, B) D^bNPN3A or (C, D) D^bNP₃₆₆ tetramer, anti-CD8 and anti-Vβ mAbs conjugated with FITC, then the tetramer⁺CD8⁺ cells were analysed for profiles of Vβ staining. Shown are results for (A–D) individual mice (n = 4). S: spleen. Found at: doi:10.1371/journal.ppat.1001039.s002 (1.33 MB TIF)

Figure S3 N3A substitution within NP₃₆₆ does not affect antigen presentation or the rate of viral clearance. Effects of N3A substitution within NP₃₆₆ on (A) viral clearance and (B, C) antigen presentation was assessed. (A) Naïve mice were infected with either the wt HK or mutant HK-NPN3A virus. Lungs were sampled at days 3, 6, or 8 after infection and homogenized for virus titration by plaque assay on MDCK cell monolayers. Data represent the mean and n = 5 mice per group. (B, C) *ex vivo*⁵¹Cr-mediated cytotoxicity was assessed after incubation of target $(H2^b)$ EL-4 target cells either (B) infected with HK or HK-NPN3A virus or (C) pulsed with $1\mu M$ NP $_{366}$ or NPN3A peptides.

Found at: doi:10.1371/journal.ppat.1001039.s003 (1.21 MB TIF)

Figure S4 Scatchard analysis of TCR avidity for pMHC by tetramer dilution assay at the acute secondary time point (d8). Scatchard analysis of tetramer dissociation and correlation coefficient (\mathbb{R}^2) based on tetramer binding MFI (Holmberg K et al, J Immunol 171:2427, 2003) are shown. Memory mice primed with either (A, D) PR8 or (B, C) PR8-NPN3A viruses were challenged with either (A, D) HK or (B, C) mutant HK-NPN3A virus. Data represent mean of MFI/tetramer concentration (μ g/ml) versus MFI of tetramer staining, from 4 mice per group.

Found at: doi:10.1371/journal.ppat.1001039.s004 (1.96 MB TIF)

Table S1 Summary of TCR β and TCR α repertoire for D^bNP_{366} and $D^bNPN3A \ V\beta 8.3^* \ T$ cells following 1^0 and 2^0 infection with the wt or mutant NPN3A influenza virus

Found at: doi:10.1371/journal.ppat.1001039.s005 (0.05 MB DOC)

References

- Doherty PC, Turner SJ, Webby RG, Thomas PG (2006) Influenza and the challenge for immunology. Nat Immunol 7: 449–455.
- Thomas PG, Brown SA, Keating R, Yue W, Morris MY, et al. (2007) Hidden epitopes emerge in secondary influenza virus-specific CD8+ T cell responses. J Immunol 178: 3091–3098.
- Pircher H, Moskophidis D, Rohrer U, Burki K, Hengartner H, et al. (1990) Viral escape by selection of cytotoxic T cell-resistant virus variants in vivo. Nature 346: 629–633.
- Moore C, John M, James I, Christiansen F, Witt C, et al. (2002) Evidence of HIV-1 adaptation to HLA-restricted immune responses at a population level. Science 296: 1439–1443.
- Price DA, West SM, Betts MR, Ruff LE, Brenchley JM, et al. (2004) T cell receptor recognition motifs govern immune escape patterns in acute SIV infection. Immunity 21: 793–803.
- Fernandez C, Stratov I, De Rose R, Walsh K, Dale C, et al. (2005) Rapid viral escape at an immunodominant simian-human immunodeficiency virus cytotoxic T-lymphocyte epitope exacts a dramatic fitness cost. J Virol 79: 5721–5731.
- Neumann-Haefelin C, McKiernan S, Ward S, Viazov S, Spangenberg HC, et al. (2006) Dominant influence of an HLA-B27 restricted CD8+ T cell response in mediating HCV clearance and evolution. Hepatology 43: 563–572.
- Butler NS, Theodossis A, Webb AI, Dunstone MA, Nastovska R, et al. (2008) Structural and Biological Basis of CTL Escape in Coronavirus-Infected Mice. J Immunol 180: 3926–3937.
- Butler NS, Theodossis A, Webb AI, Nastovska R, Ramarathinam SH, et al. (2008) Prevention of cytotoxic T cell escape using a heteroclitic subdominant viral T cell determinant. PLoS Pathog 4: e1000186.
- Topham DJ, Doherty PC (1998) Clearance of an influenza A virus by CD4+ T cells is inefficient in the absence of B cells. J Virol 72: 882–885.
- Gog J, Rimmelzwaan G, Osterhaus A, Grenfell B (2003) Population dynamics of rapid fixation in cytotoxic T lymphocyte escape mutants of influenza A. Proc Natl Acad Sci U S A 100: 11143–11147.
- Rimmelzwaan GF, Boon AC, Voeten JT, Berkhoff EG, Fouchier RA, et al. (2004) Sequence variation in the influenza A virus nucleoprotein associated with escape from cytotoxic T lymphocytes. Virus Res 103: 97–100.
- Boon AC, de Mutsert G, Graus YM, Fouchier RA, Sintnicolaas K, et al. (2002) Sequence variation in a newly identified HLA-B35-restricted epitope in the influenza A virus nucleoprotein associated with escape from cytotoxic T lymphocytes. J Virol 76: 2567–2572.
- Voeten JT, Bestebroer TM, Nieuwkoop NJ, Fouchier RA, Osterhaus AD, et al. (2000) Antigenic drift in the influenza A virus (H3N2) nucleoprotein and escape from recognition by cytotoxic T lymphocytes. J Virol 74: 6800–6807.
- Tan PT, Heiny AT, Miotto O, Salmon J, Marques ET, et al. (2010) Conservation and diversity of influenza A H1N1 HLA-restricted T cell epitope candidates for epitope-based vaccines. PLoS One 5: e8754.
- Rimmelzwaan GF, Kreijtz JH, Bodewes R, Fouchier RA, Osterhaus AD (2009) Influenza virus CTL epitopes, remarkably conserved and remarkably variable. Vaccine 27: 6363–6365.
- Berkhoff EG, de Wit E, Geelhoed-Mieras MM, Boon AC, Symons J, et al. (2005) Functional constraints of influenza A virus epitopes limit escape from cytotoxic T lymphocytes. J Virol 79: 11239–11246.
- Wahl A, McCoy W, Schafer F, Bardet W, Buchli R, et al. (2009) T-cell tolerance for variability in an HLA class I-presented influenza A virus epitope. J Virol 83: 9206–9214.
- Wahl A, Schafer F, Bardet W, Buchli R, Air GM, et al. (2009) HLA class I molecules consistently present internal influenza epitopes. Proc Natl Acad Sci U S A 106: 540–545.

Table S2 Frequency of TCR β s in D^bNPN3A⁺ V β 9⁺CD8⁺ T cells after 1⁰ mutant HK-NPN3A infection detected with either the D^bNP₃₆₆⁺ or D^bNPN3A⁺ tetramer

Found at: doi:10.1371/journal.ppat.1001039.s006 (0.05 MB DOC)

Acknowledgments

We thank Dr John Stambas for review of the manuscript, Dr Richard Webby for help with NPN3A viruses, Ken Field and Jason Waithman for FACS sorting.

Author Contributions

Conceived and designed the experiments: SAV SG NLLG AWP JR PCD SJT KK. Performed the experiments: SAV SG CG NLLG KK. Analyzed the data: SAV SG CG PGT KK. Contributed reagents/materials/analysis tools: AWP JR KK. Wrote the paper: SAV SG CG NLLG PGT AWP JR PCD SJT KK.

- Kedzierska K, La Gruta NL, Turner SJ, Doherty PC (2006) Establishment and recall of CD8+ T cell memory in a model of localized transient infection. Immunol Rev 211: 133–145.
- Turner SJ, Kedzierska K, Komodromou H, La Gruta NL, Dunstone MA, et al. (2005) Lack of prominent peptide-major histocompatibility complex features limits repertoire diversity in virus-specific CD8(+) T cell populations. Nat Immunol 6: 382–389.
- Kedzierska K, Guillonneau C, Gras S, Hatton LA, Webby R, et al. (2008) Complete modification of TCR specificity and repertoire selection does not perturb a CD8+ T cell immunodominance hierarchy. Proc Natl Acad Sci U S A 105: 19408–19413.
- Young AC, Zhang W, Sacchettini JC, Nathenson SG (1994) The threedimensional structure of H-2Db at 2.4 A resolution: implications for antigendeterminant selection. Cell 76: 39–50.
- Belz GT, Wodarz D, Diaz G, Nowak MA, Doherty PC (2002) Compromised influenza virus-specific CD8(+)-T-cell memory in CD4(+)-T-cell-deficient mice. J Virol 76: 12388–12393.
- Belz GT, Stevenson PG, Doherty PC (2000) Contemporary analysis of MHCrelated immunodominance hierarchies in the CD8+ T cell response to influenza A viruses. J Immunol 165: 2404–2409.
- Hou S, Hyland L, Ryan KW, Portner A, Doherty PC (1994) Virus-specific CD8+ T-cell memory determined by clonal burst size. Nature 369: 652–654.
- Sallusto F, Lenig D, Forster R, Lipp M, Lanzavecchia A (1999) Two subsets of memory T lymphocytes with distinct homing potentials and effector functions. Nature 401: 708–712.
- Kaech SM, Tan JT, Wherry EJ, Konieczny BT, Surh CD, et al. (2003) Selective expression of the interleukin 7 receptor identifies effector CD8 T cells that give rise to long-lived memory cells. Nat Immunol 4: 1191–1198.
- La Gruta NL, Turner SJ, Doherty PC (2004) Hierarchies in cytokine expression profiles for acute and resolving influenza virus-specific CD8+ T cell responses: correlation of cytokine profile and TCR avidity. J Immunol 172: 5553–5560.
- Theodossis A, Guillonneau C, Welland A, Ely LK, Clements CS, et al. (2010) Constraints within major histocompatibility complex class I restricted peptides: presentation and consequences for T-cell recognition. Proc Natl Acad Sci U S A 107: 5534–5539.
- Tynan FE, Burrows SR, Buckle AM, Clements CS, Borg NA, et al. (2005) T cell receptor recognition of a 'super-bulged' major histocompatibility complex class I-bound peptide. Nat Immunol 6: 1114–1122.
- Godfrey DI, Rossjohn J, McCluskey J (2008) The fidelity, occasional promiscuity, and versatility of T cell receptor recognition. Immunity 28: 304–314.
- 33. La Gruta NL, Rothwell WT, Cukalac T, Swan NG, Valkenburg SA, et al. (2010) Primary CTL response magnitude in mice is determined by the extent of naive T cell recruitment and subsequent clonal expansion. J Clin Invest.
- Kedzierska K, Turner SJ, Doherty PC (2004) Conserved T cell receptor usage in primary and recall responses to an immunodominant influenza virus nucleoprotein epitope. Proc Natl Acad Sci U S A 101: 4942–4947.
- Kedzierska K, Day EB, Pi J, Heard SB, Doherty PC, et al. (2006) Quantification of Repertoire Diversity of Influenza-Specific Epitopes with Predominant Public or Private TCR Usage. J Immunol 177: 6705–6712.
- Zhong W, Dixit SB, Mallis RJ, Arthanari H, Lugovskoy AA, et al. (2007) CTL recognition of a protective immunodominant influenza A virus nucleoprotein epitope utilizes a highly restricted Vbeta but diverse Valpha repertoire: functional and structural implications. J Mol Biol 372: 535–548.

- Deckhut AM, Allan W, McMickle A, Eichelberger M, Blackman MA, et al. (1993) Prominent usage of V beta 8.3 T cells in the H-2Db-restricted response to an influenza A virus nucleoprotein epitope. J Immunol 151: 2658–2666.
- Wherry EJ, Teichgraber V, Becker TC, Masopust D, Kaech SM, et al. (2003) Lineage relationship and protective immunity of memory CD8 T cell subsets. Nat Immunol 4: 225–234.
- Kedzierska K, Venturi V, Field K, Davenport MP, Turner SJ, et al. (2006) Early establishment of diverse TCR profiles for influenza-specific CD62Lhi CD8+ memory T cells. Proc Natl Acad Sci U S A 103: 9184–9189.
- Kedzierska K, Venturi V, Valkenburg SA, Davenport MP, Turner SJ, et al. (2008) Homogenization of TCR Repertoires within Secondary CD62Lhigh and CD62Llow Virus-Specific CD8+ T Cell Populations. J Immunol 180: 7938–7947.
- Jabbari A, Harty JT (2006) Secondary memory CD8+ T cells are more protective but slower to acquire a central-memory phenotype. J Exp Med 203: 919–932.
- Masopust D, Ha SJ, Vezys V, Ahmed R (2006) Stimulation history dictates memory CD8 T cell phenotype: implications for prime-boost vaccination. J Immunol 177: 831–839.
- Wallace ME, Bryden M, Cose SC, Coles RM, Schumacher TN, et al. (2000) Junctional biases in the naive TCR repertoire control the CTL response to an immunodominant determinant of HSV-1. Immunity 12: 547–556.
- Kersh GJ, Miley MJ, Nelson CA, Grakoui A, Horvath S, et al. (2001) Structural and functional consequences of altering a peptide MHC anchor residue. J Immunol 166: 3345–3354.
- Clute SC, Watkin LB, Cornberg M, Naumov YN, Sullivan JL, et al. (2005) Cross-reactive influenza virus-specific CD8+ T cells contribute to lymphoproliferation in Epstein-Barr virus-associated infectious mononucleosis. J Clin Invest 115: 3602–3612.
- Wedemeyer H, Mizukoshi E, Davis AR, Bennink JR, Rehermann B (2001) Cross-reactivity between hepatitis C virus and Influenza A virus determinantspecific cytotoxic T cells. J Virol 75: 11392–11400.
- Urbani S, Amadei B, Fisicaro P, Pilli M, Missale G, et al. (2005) Heterologous T cell immunity in severe hepatitis C virus infection. J Exp Med 201: 675–680.
- Cornberg M, Chen AT, Wilkinson LA, Brehm MA, Kim SK, et al. (2006) Narrowed TCR repertoire and viral escape as a consequence of heterologous immunity. J Clin Invest 116: 1443–1456.

- Allan W, Tabi Z, Cleary A, Doherty PC (1990) Cellular events in the lymph node and lung of mice with influenza. Consequences of depleting CD4+ T cells. J Immunol 144: 3980–3986.
- Hoffmann E, Mahmood K, Yang C, Webster R, Greenberg H, et al. (2002) Rescue of influenza B virus from eight plasmids. Proc Natl Acad Sci U S A 99: 11411–11416.
- Holmberg K, Mariathasan S, Ohteki T, Ohashi PS, Gascoigne NR (2003) TCR binding kinetics measured with MHC class I tetramers reveal a positive selecting peptide with relatively high affinity for TCR. J Immunol 171: 2427–2434.
- Stifka MK, Whitton JL (2001) Functional avidity maturation of CD8(+) T cells without selection of higher affinity TCR. Nat Immunol 2: 711–717.
- Moon JJ, Chu HH, Pepper M, McSorley SJ, Jameson SC, et al. (2007) Naive CD4(+) T cell frequency varies for different epitopes and predicts repertoire diversity and response magnitude. Immunity 27: 203–213.
- Crowe SR, Turner SJ, Miller SC, Roberts AD, Rappolo RA, et al. (2003) Differential antigen presentation regulates the changing patterns of CD8+ T cell immunodominance in primary and secondary influenza virus infections. J Exp Med 198: 399–410.
- 55. Clements CS, Kjer-Nielsen L, MacDonald WA, Brooks AG, Purcell AW, et al. (2002) The production, purification and crystallization of a soluble heterodimeric form of a highly selected T-cell receptor in its unliganded and liganded state. Acta Crystallogr D Biol Crystallogr 58: 2131–2134.
- Macdonald W, Williams DS, Clements CS, Gorman JJ, Kjer-Nielsen L, et al. (2002) Identification of a dominant self-ligand bound to three HLA B44 alleles and the preliminary crystallographic analysis of recombinant forms of each complex. FEBS Lett 527: 27–32.
- Kabsch W (1993) Automatic processing of rotation diffraction data from crystals of initially unknown symmetry and cell constants. J Appl Cryst 26: 795–800.
- McCoy AJ, Grosse-Kunstleve RW, Adams PD, Winn MD, Storoni LC, et al. (2007) Phaser Crystallographic Software. J Appl Cryst 40: 658–674.
- (1994) The CCP4 suite: programs for protein crystallography. Acta Crystallogr D Biol Crystallogr 50: 760–763.
- Jones T, Zou J, Cowan S, Kjeldgaard M (1991) Improved methods for building protein models in electron density maps and the location of errors in these models. Acta Crystallogr A 1991: 110–119.