Early Priming Minimizes the Age-Related Immune Compromise of CD8⁺ T Cell Diversity and Function

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Abstract
The elderly are particularly susceptible to influenza A virus infections, with increased occurrence, disease severity and reduced vaccine efficacy attributed to declining immunity. Experimentally, the age-dependent decline in influenza-specific CD8⁺ T cell responsiveness reflects both functional compromise and the emergence of ‘repertoire holes’ arising from the loss of low frequency clonotypes. In this study, we asked whether early priming limits the time-related attrition of immune competence. Though primary responses in aged mice were compromised, animals vaccinated at 6 weeks then challenged >20 months later had T-cell responses that were normal in magnitude. Both functional quality and the persistence of ‘preferred’ TCR clonotypes that expand in a characteristic immunodominance hierarchy were maintained following early priming. Similar to the early priming, vaccination at 22 months followed by challenge retained a response magnitude equivalent to young mice. However, late priming resulted in reduced TCRβ diversity in comparison with vaccination earlier in life. Thus, early priming was critical to maintaining individual and population-wide TCRβ diversity. In summary, early exposure leads to the long-term maintenance of memory T cells and thus preserves optimal, influenza-specific CD8⁺ T-cell responsiveness and protects against the age-related attrition of naïve T-cell precursors. Our study supports development of vaccines that prime CD8⁺ T-cells early in life to elicit the broadest possible spectrum of CD8⁺ T-cell memory and preserve the magnitude, functionality and TCR usage of responding populations. In addition, our study provides the most comprehensive analysis of the aged (primary, secondary primed-early and secondary primed-late) TCR repertoires published to date.

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Introduction
The elderly population is particularly susceptible to novel infections, especially the annual, seasonal epidemics caused by influenza A viruses [1,2], with increased occurrence, severity of infection and reduced vaccine efficacy being attributed to age-related decline in immune capacity [3–6]. The ageing effect on the immune system is considered to be multifactorial, arising from the diminished thymic export of naïve precursors due to thymic involution [7,8], the impaired recruitment [9,10] of naïve CD8⁺ T cell precursors and the replicative senescence of memory cells [11–14]. Ageing can also be associated with abnormal cellular functions such as distorted cytokine secretion (IL-2, IL-4 and IFN-γ) profiles [15–17], decreased granzyme B production [18,19] and reduced proliferative capacity due to the loss of CD28 expression [20]. Perturbations in the naive TCR repertoire have also been reported, with abnormal TCR spectratype (CDR3β length) patterns in aged mice reflecting the massive, antigen-independent expansion, of a few clonotypes [21]. Naïve T cell attrition has also been inferred from observed reductions in the diversity of antigen-specific TCR repertoires in aged mice [5,22].

Previous mouse studies have established that ageing can be associated with diminished CD8⁺ T cell efficacy and delayed influenza virus clearance [23–25]. Recent evidence has further shown that the selective loss of primary, influenza-specific CD8⁺ T cell responsiveness in older mice is characterized by a narrowing in the spectrum of TCR usage and is seen predominantly for low frequency populations, with this effect being best characterized for the prominent D b NP366⁺CD8⁺ T cell set [5,26]. Overall, the findings so far suggest that the capacity to respond effectively to new influenza infections in aged mice requires the maintenance of a diverse pool of functional peripheral T cells.

As CD8⁺ T cells tend to be specific for peptides derived from more conserved proteins that are internal to the virus, priming effective CD8⁺ T cell memory has obvious potential for countering newly emerged seasonal or pandemic influenza strains. The importance of long-lived, antigen-specific memory CD8⁺ T cells capable of rapid recall following the secondary infection has been well documented for the respiratory viruses in mice [27,28] and...
Author Summary

The elderly population is particularly susceptible to novel infections, especially the annual, seasonal epidemics caused by influenza viruses. Established T cell immunity directed at conserved viral regions provides some protection against influenza infection and promotes more rapid recovery, thus leading to better clinical outcomes. We asked whether priming early in life limits the time-related attrition of immune competence. We found that although influenza-specific T cell responses are compromised in the aged mice, vaccination with influenza early (but not late) in life ‘locks’ optimal T-cell responsiveness, maintains functional quality, persistence of preferred clones and a characteristic T cell hierarchy. Overall, our study supports development of vaccines that prime T cells early in life to elicit the broadest possible spectrum of pre-existing T cell memory and preserve the magnitude, functionality and clonal usage of responding populations for life-long immunity against influenza viruses.

Humans [29,30]. Such long-term maintenance of memory T cells leading to enhanced secondary response forms the basis for vaccination strategies based on priming CD8+ T cell memory to promote early virus clearance and decreased morbidity. The question is whether such CD8+ T cell memory can be effectively recalled in the elderly.

A recent study [6] suggested that infecting mice with LCMV or influenza at an extreme age (18–20 months) leads to defective CD8+ T cell memory and diminished recall responses following virus challenge. What happens, though, if CD8+ T cell memory is established when the mice are young? The analysis reported here compares the CD8+ T cell response profiles for young (<3 months) and aged (22 month) mice, with the latter cohort being first exposed to immunogenic influenza epitopes early or late in life. The results suggest that designing influenza vaccines which promote as broad as possible spectrum of CD8+ T cell memory in adolescence could be beneficial, even if such benefit emerges long after the subject has first been given the protective immunogen.

Results

To validate the previous studies [5,31] and determine the ageing effect on primary immune responsiveness (Figure 1A) for immunodominant DγNP366 and DPA224+ CD8+ pools, we infected young (<3 month) and old (>22 month) mice intranasally (i.n.) with 1×10^5 pfu of an infectious (H3N2, HK) influenza A virus. More importantly, as a main question of the present study, we asked whether any age-related compromise of CD8+ T cell function and diversity might be modified by priming early (at 2 months) or late (at 22 months) i.p. with 1.5×10^6 pfu of the serologically distinct PR8 (H1N1) virus that has the same immunogenic CD8+ T cell peptides as HK.

We used the i.p. priming route with the influenza virus as it does not lead to a productive viral replication (similarly to the current i.m. human influenza vaccines), but gives one-stop growth cycle with full protein production. Such non-productive immunisation with the whole virus results in priming of antigen-specific effector T cells and establishment of long-term T cell memory for subsequent challenge (Figure S1), comparable to those observed after the natural (i.n.) influenza infection [32–34]. Importantly, the i.p. priming does not elicit the whole cascade of detrimental inflammatory responses in the virally-infected lung [35] and thus avoids double pathology at the site of infection. The i.p. route of influenza priming is equivalent to the current i.m. vaccination approaches used in humans with respect to the non-productive viral immunisation.

CD8+ T cell responsiveness following early versus late infection of aged mice

The comparison of the HK-induced CD8+ T cell responses utilized young or old mice that were either immunologically naive (primary, 1^st^ Figure 1A) or had been primed at 2 months of age with the PR8 virus and challenged 20 months later (secondary, 2^nd^ Figure 1B). Immunodominant and subdominant CD8+ T cell responses were measured in the spleen (Figure 1CD) by the ex vivo IFN-γ ICS assay. Following 1^st^ challenge, the size of the low precursor frequency DγNP366+ CD8+ set in the spleen (Figure 1C) was markedly diminished in the aged animals relative to the young controls as previously observed [3,31].

Conversely, any age-related effects on CD8+ T cell numbers were not significant for DPA224+ CD8+ T cell responses are intriguing, as the naive CD8+ T cell frequencies [36] found for DPA224+ specific T cells in young mice are significantly higher than those detected for DγNP366 (≥72 versus <40 per individual, respectively), suggesting that a larger naive CD8+ T cell pool size minimizes the extent of age-related attrition and, as a consequence, the effect on primary CD8+ T cell response magnitude (Figure 1C).

Reduced magnitude of the immunodominant DγNP366-specific CD8+ T cell response that was detected for the primary, influenza-specific CD8+ T cell response in older mice (Figure 1D), was not sustained following secondary HK challenge of mice that had been primed early with the PR8 virus (Figure 1D). The numbers of DPA224+CD8+ T cells were significantly diminished across combined experiments but, otherwise, the recall responses for memory T cell pools in young or old mice primed at <2 months (at least 20 months previously) were not obviously different, emphasizing the durability of virus-specific CD8+ T cell memory [37]. In particular, the overdominance of the DγNP366-specific set that is characteristic of the secondary response to these viruses [38] was still apparent in the aged mice (Figure 1D).

The beneficial effect of the early CD8+ priming on the immunodominant low-precursor responses like the DγNP366-specific population following influenza infection at the extreme age was most striking when the relative contributions of particular antigen-specific CD8+ T cells were analysed based on total cell numbers (Figure 2, calculations based on Figure 1 for immunodominant DγNP366+CD8+ and DPA224+CD8+ pools, and data not shown for subdominant DFB1705+CD8+ and KβPB1-F262+CD8+ populations). In the aged mice, the primary CD8+ T cell responses showed a shift in the typical immunodominance hierarchy (Figure 2B), with the contribution of the immunodominant DγNP366+CD8+ population being significantly lower in the aged mice (9.4±3.6%) in comparison to young animals (43.4±15%; p<0.01; Figure 2A). The differential immunodominance hierarchy resulted mainly from significantly increased contribution of KβPB1705+CD8+ T cells (Figure 2). This led to major modifications in response hierarchy following primary influenza virus infection of aged mice KβPB1705>DPA224>DFB1705>F262>DγNP366, with the comparable profile for young mice being DγNP366>DPA224=KβPB1705>DFB1705-F262.

Conversely, recall of CD8+ T cells primed at a young age preserved the overall contribution of T cell specificities and retained the immunodominance hierarchy in aged mice primed early at 6 weeks (Figure 2D), reflecting the characteristic immunodominance hierarchy in young controls (Figure 2C).

These findings show clearly that priming the CD8+ T cell compartment at an early age leads to subsequent preservation of
CD8\(^+\) T cell numbers and immunodominance hierarchies for influenza infection in the elderly.

Age-related effects on cytokine polyfunctionality and activation status

One measure of CD8\(^+\) T cell function is the capacity to produce multiple cytokines simultaneously [39] following in vitro stimulation with peptide in the standard, 5 h ICS assay. For the primary DbPA224\(^+\) CD8\(^+\) T cell population that remained relatively constant in numbers with age (Figure 1C), the frequencies of double (IFN-\(\gamma\)/TNF-\(\alpha\)) and triple-producers (IFN-\(\gamma\)/TNF-\(\alpha\)/IL-2) were significantly lower in comparison with the young mice (Figure 3AB). Furthermore, taking mean fluorescence intensity (MFI), which represents the intensity and therefore amount of cytokine production, it also seems that the DbPA224\(^+\) CD8\(^+\) population tended to produce less TNF-\(\alpha\), though this diminution effect was not apparent for either IFN-\(\gamma\) or IL-2 (Figure 4A).

Taking the prevalence and MFI data together (Figure 3 and Figure 4), there appears to be a general decrease in cytokine polyfunctionality for the primary DbPA224\(^+\) CD8\(^+\) response.

Figure 1. Effect of age and early priming on 1\(^o\) and 2\(^o\) CD8\(^+\) T cell responses. (A) For the primary responses, naïve mice were infected i.n. with 1 x 10\(^4\) pfu of the HK (H3N2) influenza A virus either at a young (<3 months; mo) or extreme (22 mo) age. Analysis of CD8\(^+\) T cell responses was performed on d10 after the primary infection. (B) For the secondary responses of the early-primed mice, animals were primed at <2 mo i.p. with 1.5 x 10\(^7\) pfu the PR8 (H1N1) influenza A virus, then challenged 6 weeks (young) or >22 mo (aged) later i.n. with 1 x 10\(^4\) pfu of the HK virus. Analysis of CD8\(^+\) T cell responses was performed on d8 after the secondary infection. (C, D) Numbers of epitope-specific CD8\(^+\) T cells in the spleens recovered from young (filled symbols) or aged (>22 month, open symbols) B6 mice on d10 (1\(^o\), C) or d8 (2\(^o\), D) following primary (1\(^o\)) or secondary (primed young) (2\(^o\)) i.n. infection with the HK (H3N2) influenza A virus. Memory mice had been injected i.p. with the PR8 (H1N1) influenza A virus at <2 mo and were challenged. Lymphocyte populations were stimulated with the NP\(_{366}\), or PA224 peptides in the presence of Brefeldin A for 5 hrs, then stained with the anti-CD8PerCPCy5.5 mAb, fixed/permeabilised and stained with anti-IFN-\(\gamma\)-FITC mAb. Cytokine (IFN-\(\gamma\)) production was calculated by subtracting background fluorescence for the no-peptide controls, and the numbers of IFN-\(\gamma\)-specific CD8\(^+\) T cells were determined from the % cells staining and the total cell counts. Data represent individual mice (symbols) and the mean (line). Experiments were performed at least twice. * = p<0.05.

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CD8\(^+\) T cell numbers and immunodominance hierarchies for influenza infection in the elderly.

Figure 2. Immunodominance hierarchies in aged mice after 1\(^o\) infection or 2\(^o\) challenge of primed-early mice. The relative prevalence of the immunodominant D\(^\text{NP}_{366}\)/CD8\(^+\) and D\(^\text{PA}_{224}\)/CD8\(^+\) T cell population over the subdominant D\(^\text{PB1}_{1703}\)/CD8\(^+\) and K\(^\text{PB1-F}_{262}\)/CD8\(^+\) sets. Results are shown for (A, B) 1\(^o\) and (C, D) 2\(^o\) HK infection in (A, C) young and (B, D) aged mice. The relative contributions of particular antigen-specific CD8\(^+\) T cells were analysed based on total cell responses (Figure 1 for D\(^\text{NP}_{366}\)/CD8\(^+\) and D\(^\text{PA}_{224}\)/CD8\(^+\) and data not shown for D\(^\text{PB1}_{1703}\)/CD8\(^+\) and K\(^\text{PB1-F}_{262}\)/CD8\(^+\)). Data represent the mean proportion of a particular peptide-specific CD8\(^+\) population. * = p<0.01 shows a difference between young and aged animals. Experimental outline as in Figure 1AB.

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Conservation of CD8\(^+\) T Cell Memory with Age
Conversely, analysis of aged mice primed early showed that functional characteristics appear to be locked-in early and maintained in the long-term for memory T cell populations (Figure 3CDEF).

Can we detect other evidence of enduring functional change? Given that the influenza-specific CD8+ T cells generated following primary infection of aged mice were either of suboptimal functional quality (DbPA224+CD8+; Figure 3, Figure 4) or reduced in number (DbNP366+CD8+; Figure 1), the further question was whether there was any effect on cell surface activation phenotype [34,40–42]. Comparison of phenotypic markers associated with activation, trafficking and memory potential: CD62L vs. IL-7Rα (CD127), CD27 vs. CD43, and IL-7Rα vs. KLRG-1 for the D'Pa224+CD8+ and D'np366+CD8+ sets (Figure S2) showed that, with the exception of a decrease in the relative prevalence of the least activated CD27loCD43loCD8+ cells in the older mice (Figure S2AD), there were no significant differences in phenotype with age.

Aged memory T cells have a young-type TCRβ usage profile

Previous studies have found a significant skewing in TCR Vβ usage (mAb staining) and CDR3β length (spectratyping) for CD8+ T cell responses developed from naïve and memory populations by
the infection of aged versus young mice [13,21,43]. Thus, we
looked more closely at the expansion and maintenance of
responder T cell clonotypes [44,45]. As our earlier analysis of
influenza-specific CD8+ TCR clonotype diversity has focused on
the prominent Vβ8.3*DγNP366* and Vβ7*DγPA224* sets
[47], we first assessed the Vβ mAb-staining profiles to determine
whether these characteristic TCRs were also selected following
primary or secondary challenge of aged mice. Indeed for both
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primary or secondary challenge of aged mice. Indeed for both
consistent with the DγNP366* CD8+ and DγPA224* CD8+ T cell responses,
the characteristic Vβ8.3 and Vβ7 usage was observed (Figure 5),
though additional Vβ6, Vβ7 and Vβ9 biases were variously
detected in individual, older mice for the primary DγNP366*CD8+
population (Figure 5C), possibly due to the recruitment of low
frequency alternate DγNP366*CD8+ T cell responses. Despite the
absence of a prominent Vβ8.1/8.2*DγNP366* set in one of the
early-primed, secondarily-challenged at 22 month mice, the bias
was generally to Vβ8.3 suggesting that the characteristic
DγNP366*CD8+ TCRβ usage profile is retained in the persistent
memory population. The DγPA224* set was characterised across
individuals by Vβ7 TCR usage (Figure 5BDF), which was more
consistent than the DγNP366*Vβ8.3 usage, possibly reflecting the
higher number of precursors with Vβ7 surviving within the 22
month old mice.

Priming at an extreme age does not impair the recall
response magnitude

Since priming at a young age led to the typical magnitude and
quality of influenza-specific CD8+ T cell responses following viral
infection in the aged mice, we asked whether priming the mice via
a non-replicative route (i.p. priming with 1.5×107 pfu of PR8) at
extreme age (22 months) would also be beneficial for the
subsequent influenza virus infection. Since the reduced primary
DγNP366*CD8+ T cell responses in aged mice has been attributed
to the lower naïve precursors in young mice [5], this experiment
would determine whether old naïve mice could be primed at an
extreme age (at 22 months) and subsequently challenged i.n.
with 1×104 pfu of the HK influenza strain (at ~24 months;
Figure 6A) to mount an effective recall response after the
attrition had occurred. Surprisingly, despite the reduced primary
DγNP366*CD8+ T cell responses (Figure 1C) and lower magnitude
of secondary DγPA224*CD8+ sets (Figure 1E) in the spleens of aged
animals, the recall of influenza-specific CD8+ T cells was robust
and equivalent in magnitude to the young controls (Figure 6). The
numbers of both immunodominant DγNP366*CD8+ and
DγPA224*CD8+ populations were normal (Figure 6B). This
resulted in the maintained contribution of each of the T cell
specificities to influenza-specific responses (Figure 6E). Conversely,
the polyfunctionality of those secondary CD8+ T cell populations
in mice primed at the extreme age did not always resemble
effectiveness of influenza-specific CD8+ T cells recruited in young
individuals (Figure 6C). Perturbations in the TCR usage with
extreme age were evident macroscopically in the TCR Vβ usage
for DγPA224*CD8+ (Figure 6G) and especially the DγNP366*CD8+
(Figure 6F) responses, with the usage of alternate Vβ8.1/8.2 for
DγPA224*CD8+ and Vβ7 and Vβ8.1/8.2 for DγNP366*CD8+
populations. The characteristic Vβ8.3 usage for DγNP366*CD8+
was only dominant in 1 of 4 mice (Figure 6F), reflecting narrowing
of the naive DγNP366*CD8+ set with extreme age that initially
limited the primary response (Figure 1C) and/or the clonal
expansions characteristic for the aged animals as previously
reported [13,14].

Early but not late priming preserves TCRβ usage of
‘preferred’ clonotypes in the aged mice

A substantial body of work from previous studies has defined the
young B6 CDR3β TCR usage at high resolution [44,47], therefore
using these data sets from young mice we were able to compare
the spectrum of clonotype prevalence in aged mice using single-
cell RT-PCR and sequencing of the CDR3β region to determine
the spectrum of TCRβ diversity. Analysis of 1489 CDR3β

Figure 4. Impaired polyfunctionality of DγPA224-specific CD8+ T cells in the aged mice during primary but not secondary influenza
infection. (A) Primary or (B) secondary (primed young) influenza-specific CD8+ T cell responses were assessed for simultaneous production of IFN-γ,
TNF-α and IL-2 in the spleen of aged (22 months old) and young (6–8 weeks) mice. Compiled data (n = 3–5, mean±SD) are shown for the mean
fluorescence intensity (MFI) of IFN-γ, IFN-γ and TNF-α as well as IFN-γ and IL-2 staining. * = p<0.05. Experimental outline as in Figure 1AB.
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sequences for primary and secondary (primed-young and primed-old) responses from the 22 month old mice (Tables 1 and 2) showed that the dominant Jb regions and CDR3b loop lengths in the aged animals (Tables S1, S2, S3, S4, S5, S6) were comparable to those found early in life (Figures 7 and 8 for comparison with young animals). However, more inter-individual variation in the primary responses was observed in the older group (Figure S3).

While 83% of each of the TCRb repertoires involved in the primary responses to DbNP366 in young mice utilized Jb2.2 and a CDR3 length of 9 amino acids (aa), this profile was substantially diminished to 57% of the TCRb repertoire for 2/7 aged mice.

Similarly, Jb1.1, Jb1.5, and Jb2.6 collectively dominated the primary DbPA224+CD8+ responses for 7/7 young mice, while Jb2.1 and Jb2.3 emerged strongly (>55% each) for 2 of the older mice. While the primary DPa224+CD8+ repertoires in individual young mice mostly featured diverse CDR3 lengths of 5, 6, and 7 aa, 94% of the primary DPa224+CD8+ T cell repertoires in two of the aged mice could be attributed to one particular CDR3 length (i.e. 6 aa in one mouse and 7 aa in the other mouse).

Age-associated changes in TCRb repertoire usage were investigated for the DNP366+CD8+ and DPA224+CD8+ populations by sequencing individual CDR3b TCR signatures (Tables 1 and 2, Tables S1, S2, S3, S4, S5, S6) and the extent of TCRb repertoire diversity was then assessed using both the number of different aa-defined clonotypes and Simpson’s diversity index, which accounts for the clonal dominance hierarchy. These measures of diversity were estimated for a standard 22 TCRb sequences per epitope per mouse to adjust for differences in total number of sequences obtained per mouse [48]. The primary DPa224+CD8+ TCRb repertoires were found to be significantly less diverse in aged versus young mice, with a lower number of clonotypes per individual (median: 8 vs. 14, p = 0.005; Figure 7C) and a decreased Simpson’s diversity index (median: 0.72 vs 0.94, p = 0.007; Figure 7G), despite there being no significant change in the DPa224-specific CD8+ T cell response magnitude (Figure 1A). Some age-related contraction in the number of different DPa224+CD8+ TCRb clonotypes was also found following secondary infection (early priming) (median: 10 vs. 12, p = 0.007; Figure 7D), though the difference was not as large as in the primary response, largely due to the increased median diversity for the recall response in older mice. Interestingly, when mice were primed at 22 months of age and then challenged (primed old), similar results were obtained as early priming, however there appeared to be substantial increase in the similarity between some pairs of mice (Figure 7P). Surprisingly, the reduced diversity seen in the DPa224+CD8+ primary response (Figure 7CG), from which the late priming response is derived, was not carried over to the primed-old recall TCRb repertoire (Figure 7DH). This suggests that priming plays a positive role in preserving a broader spectrum of clonotype availability within the inherently diverse DPa224+CD8+ T cell repertoire, due to enhanced response magnitude.

In contrast, despite the greatly diminished magnitude of the primary DNP366+CD8+ T cell response in older mice (Figure 1A), the extent of TCRb repertoire diversity analysed at the aa level was not significantly different for young and old mice (Figure 7AE, Table 1, Table S1). The public DNP366+Vβ8.3 clonotypes can be encoded by up to 10 different nucleotide (n.t.) sequences each, with as many as 4 distinct n.t.-defined variants being present in an individual young mouse [44]. Following primary exposure
of aged animals or when mice were primed late, the three main public V\textsubscript{b}8.3\textsuperscript{+}DbNP366+ clonotypes: SGGANTGQL, SGGGNTGQL, SGGSNTGQL \cite{44} were encoded by a total of 10 and 9 distinct n.t. sequences respectively (Tables S1 and S5), in contrast to the 16 different clonotypes detected in the secondary-infected (primed early), aged mice (Table S3). As a consequence, priming early or late prior to challenge preserved a mean of 2.9 ± 0.6 and 3.0 ± 0.7 n.t.-defined public clonotypes in comparison to the 1.7 ± 1.1 public n.t. sequences detected following infection of old, naive mice. Such reduced availability of n.t.-defined public clonotypes of aged animals or when mice were primed late, the three main public V\textsubscript{b}8.3\textsuperscript{+}DbNP366+ clonotypes: SGGANTGQL, SGGGNTGQL, SGGSNTGQL \cite{44} were encoded by a total of 10 and 9 distinct n.t. sequences respectively (Tables S1 and S5), in contrast to the 16 different clonotypes detected in the secondary-infected (primed early), aged mice (Table S3). As a consequence, priming early or late prior to challenge preserved a mean of 2.9 ± 0.6 and 3.0 ± 0.7 n.t.-defined public clonotypes in comparison to the 1.7 ± 1.1 public n.t. sequences detected following infection of old, naive mice. Such reduced availability of n.t.-defined public clonotypes of aged animals or when mice were primed late, the three main public V\textsubscript{b}8.3\textsuperscript{+}DbNP366+ clonotypes: SGGANTGQL, SGGGNTGQL, SGGSNTGQL \cite{44} were encoded by a total of 10 and 9 distinct n.t. sequences respectively (Tables S1 and S5), in contrast to the 16 different clonotypes detected in the secondary-infected (primed early), aged mice (Table S3). As a consequence, priming early or late prior to challenge preserved a mean of 2.9 ± 0.6 and 3.0 ± 0.7 n.t.-defined public clonotypes in comparison to the 1.7 ± 1.1 public n.t. sequences detected following infection of old, naive mice. Such reduced availability of n.t.-defined public clonotypes of aged animals or when mice were primed late, the three main public V\textsubscript{b}8.3\textsuperscript{+}DbNP366+ clonotypes: SGGANTGQL, SGGGNTGQL, SGGSNTGQL \cite{44} were encoded by a total of 10 and 9 distinct n.t. sequences respectively (Tables S1 and S5), in contrast to the 16 different clonotypes detected in the secondary-infected (primed early), aged mice (Table S3). As a consequence, priming early or late prior to challenge preserved a mean of 2.9 ± 0.6 and 3.0 ± 0.7 n.t.-defined public clonotypes in comparison to the 1.7 ± 1.1 public n.t. sequences detected following infection of old, naive mice. Such reduced availability of n.t.-defined public

**Table 1.** CDR3\textsubscript{b} TCR repertoire of D\textsuperscript{b}NP\textsubscript{366} V\textsubscript{b}8.3\textsuperscript{+}CD8\textsuperscript{+} T cells at the acute phase of primary and secondary influenza virus infection of young and aged mice.

|                      | Primary                   | Secondary                  |
|----------------------|---------------------------|----------------------------|
|                      | Young         | Aged         | Young          | Aged primed young | Aged primed old |
| Mice analysed        | 5             | 7             | 6              | 7               | 5               |
| TCRs sequenced       | 287           | 284           | 383            | 358             | 153             |
| Different clonotypes (aa) | 24         | 29            | 22             | 15              | 9               |
| Different clonotypes (nt) | 37         | 37            | 41             | 29              | 16              |
| Clonotypes per mouse (aa) | 7.0 ± 5.1  | 5.1 ± 3.5     | 6.5 ± 2.3      | 3.6 ± 1.8       | 3.2 ± 1.5       |
| Clonotypes per mouse (nt) | 8.6 ± 5.6  | 5.6 ± 3.8     | 8.8 ± 4.2      | 4.7 ± 1.4       | 4.4 ± 1.1       |

\footnotesize*Predominant when found in more than 15% of mice.*

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sequences in the primary aged mice resulted in a loss of one of the major public clonotypes SGGGNTGQL (Figure 8A) in all 7 animals tested following primary virus challenge (Table S1). This was associated in turn with a markedly greater contribution of the SGGANTGQL clonotypes (37% versus 23%) in primarily-infected aged animals in comparison to those that were secondary challenged (Figure 8B). It is interesting to note that previously the SGGANTGQL clonotype has been associated with low pMHC avidity [49]. Thus, although the D^bNP366^CD8^+ repertoire is dominated by public TCRs encoded by multiple distinct n.t. sequences, due to codon redundancy the selective, age-related exclusion of one n.t.-defined clonotype does not necessarily equate to the disappearance of any given aa clonotype from the naive pool. However, it is still possible that the prominent TCR signatures (like SGGGNTGQL) can be lost or significantly decreased with ageing.

Significantly higher inter-individual similarity of D^bNP366^CD8^+ clonotypes per mouse (aa) 23.9±6.8 10.0±4.8 21.8±3.8 13.3±3.7 11.0±2.4

Table 2. CDR3β TCR repertoire of D^bPA224^Vβ7^CD8^+ T cells at the acute phase of primary and secondary influenza virus infection of young and aged mice.

|            | Primary | Secondary |
|------------|---------|-----------|
|            | Young   | Aged      | Young   | Aged primed young | Aged primed old |
| Mice analysed | 7      | 6         | 6       | 6                 | 6               |
| TCRs sequenced | 373    | 277       | 347     | 249               | 168             |
| Different clonotypes (aa) | 115   | 55        | 105     | 66                | 54              |
| Different clonotypes (nt) | 150   | 58        | 128     | 79                | 55              |
| Clonotypes per mouse (aa) | 23.9±6.8 | 10.0±4.8 | 21.8±3.8 | 13.3±3.7 | 11.0±2.4 |
| Clonotypes per mouse (nt) | 26.0±8.1 | 10.2±5.0 | 23.5±4.0 | 13.8±4.2 | 11.0±2.4 |

1Repeated (shared) when sequence found in more than 80% of mice.

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Discussion

The present analysis establishes the importance of priming the CD8^+ T cell compartment early in life in order to preserve CD8^+ T cell numbers, functional quality and preferential profiles of TCR usage for influenza-specific CD8^+ effector T cell responses in the elderly. In contrast, primary CD8^+ T cell responses in aged animals tended to show alterations in the typical CD8^+ T cell immunodominance hierarchy, with T cell responses to some epitopes being reduced in magnitude, a decrease in the capacity to make multiple cytokines, and changes in the extent of TCRβ repertoire diversity as a consequence of the diminished availability of naive clonotypes. These effects were minimal for the recall responses generated from memory T cell populations that were generated early, and then recalled by virus challenge more than 18 months later. Overall, the results emphasize both the durability and constancy of immune memory.

The response hierarchy following primary influenza virus infection of aged mice was K^b^PB1-F262 > D^bNP366^PA224 > D^bPB1- F262 > D^bNP366, with the comparable profile for young mice being D^bNP366 > D^bPA224 = K^b^PB1-F262 > PB1-F262. Typically subdominant epitopes accounted for 59% of the response in aged naïve mice challenged with virus compared with a 34% (Figure 2A) contribution in the young. Thus, immunodominance hierarchies may be relative to age, an idea that is clearly more relevant to the situation in long-lived humans than in mice. In contrast, the typical hierarchy [36] was maintained for both young and old mice that were primed early, with a relative contribution by subdominant epitopes of 10% and 12% (Figure 2D) respectively. Whereas when mice were primed at an extreme age subdominant epitopes contributed 26% of the anti-influenza CD8^+ T cell response and, therefore, the immunodominance hierarchy was perturbed (Figure 6E), to a lesser extent than the primary response in aged mice.

The difference in naïve precursor frequency for the D^bNP366^CD8^+ and D^bPA224^CD8^+ T cell sets is only two-fold (36 vs 79 naive precursors, respectively) [36], yet any age-related diminution in magnitude for the primary response to D^bPA224 was less apparent, suggesting that expanding CD8^+ T cell precursors prevalence by an estimate of 2–4 fold may protect immune
capacity in the long term. As all the naïve, endogenous and non-transgenic D\textsuperscript{b}NP\textsubscript{366} + \textsuperscript{CD8} + and D\textsuperscript{b}PA\textsubscript{224} + \textsuperscript{CD8} + T cells are recruited into the primary immune response [36], there would be no naïve precursors left to mount a primary \textsuperscript{CD8} + T cell responses after re-challenge for these three sets of influenza-specific \textsuperscript{CD8} + T cell populations, unless new precursors had emerged subsequently from the thymus.

With age, the relative loss in magnitude for the normally prominent D\textsuperscript{b}NP\textsubscript{366} specific response can be most likely attributed to the loss of naïve precursors with time as previously suggested [5]. Despite multiple attempts to repeat the naïve \textsuperscript{CD8} + T cell analysis for aged (22 mo) B6 mice, we were unable to recover viable tetramer + \textsuperscript{CD8} + populations (data not shown) following the application of the rigorous magnetic separation procedure that is required to recover very small numbers of antigen-specific cells from the total, peripheral \textsuperscript{CD8} + T cell pool [36,50] in the aged mice comparing to normal precursor frequencies in the young controls. This could reflect diminished structural integrity due, for

Figure 7. Comparison of TCR\textsubscript{B} diversity, and inter-individual sharing and similarity for the D\textsuperscript{b}NP\textsubscript{366} V\textsubscript{b}8.3 + \textsuperscript{CD8} + and D\textsuperscript{b}PA\textsubscript{224} V\textsubscript{b}7 + \textsuperscript{CD8} + repertoires. Shown are the relative measures of TCR\textsubscript{B} repertoire diversity, (A–D) the number of different clonotypes and (E–H) Simpson’s diversity index, and (I–L) % of repertoire comprised of shared clonotypes, and (M–P) inter-individual TCR repertoire similarity, as measured by the Morisita-Horn similarity index. The Simpson’s diversity and Morisita-Horn similarity indices account for the clonal dominance hierarchy among the different clonotypes and vary between 0 (minimum diversity/similarity) and 1 (maximum diversity/similarity). Each of the diversity, inter-individual sharing and similarity measures were estimated for a standard sample size of 22 TCR sequences per individual mouse repertoire. The repertoire diversities were calculated for each mouse per age/priming group for primary (A, E) and secondary (B, F) D\textsuperscript{b}NP\textsubscript{366} V\textsubscript{b}8.3 \textsuperscript{CD8} + TCR repertoires and for primary (C, G) and secondary (D, H) D\textsuperscript{b}PA\textsubscript{224} V\textsubscript{b}7 \textsuperscript{CD8} + TCR repertoires. The repertoire similarities were assessed between pairs of primary (M) and secondary (N) D\textsuperscript{b}NP\textsubscript{366}’V\textsubscript{b}8.3’\textsuperscript{CD8} + TCR repertoires and between pairs of primary (O) and secondary (P) D\textsuperscript{b}PA\textsubscript{224}’V\textsubscript{b}7’\textsuperscript{CD8} + TCR repertoires within the same age/priming group. To evaluate TCR sharing, clonotypes were first defined as shared or non-shared across all D\textsuperscript{b}NP\textsubscript{366}’specific or D\textsuperscript{b}PA\textsubscript{224}’specific TCR\textsubscript{B} repertoires. The proportions of the 22 TCR\textsubscript{B} sequences per D\textsuperscript{b}NP\textsubscript{366}’V\textsubscript{b}8.3’\textsuperscript{CD8} + TCR repertoire (I, J) or D\textsuperscript{b}PA\textsubscript{224}’V\textsubscript{b}7’\textsuperscript{CD8} + TCR repertoire (K, L) that were comprised of shared clonotypes were then estimated. A Mann-Whitney test was used to compare young and aged mice for the primary responses and between young mice, aged mice primed young and aged mice primed old for the secondary responses. For the comparison between age/priming groups for the secondary responses, the statistical significance for each pairwise comparison was determined at p<0.0167 (*), using Bonferroni correction for multiple pairwise comparisons.

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instance, to senescence-associated changes in membrane lipids [51]. Thus, at this time we were unable to compare naive influenza-specific CD8\(^+\) T cell precursor frequencies of aged mice to established precursor frequencies in the young controls, but rather infer results from the immunodominance hierarchy of the aged primary responses.

The comparable sizes and immunodominance hierarchies of influenza-specific CD8\(^+\) T cell responses in young and elderly following recall reflects the stability of long-term-memory pools, which has also been evidenced by earlier data showing stable memory numbers for both D\(^b\)NP\(_{366}\) V\(\gamma\)8.3\(^+\) and D\(^b\)PA224 V\(\beta\)8.3\(^+\) T cells until at least d575 after primary infection [32]. Together with the present analysis, evidence for the preservation of Vaccinia virus-specific memory populations in humans primed more than 20 years previously [52] reinforces the view that early antigen encounter minimizes the attrition of CD8\(^+\) T cell responses in the elderly. Furthermore, analysis of the 2009 H1N1 (swine-origin influenza) response in human populations showed that this newly emerged pandemic virus shared immunogenic peptides with the catastrophic 1918 H1N1 strain [53], emphasizing the likely value of establishing effective CD8\(^+\) T cell memory to all known influenza epitopes.

Early priming of the CD8\(^+\) T cell compartment also preserves CD8\(^+\) T cell functionality in the very long term. In contrast to the suboptimal peptide-induced, polynucleotide cytokine profiles expressed by CD8\(^+\) T cells generated from naive CD8\(^+\) T cells in aged animals, the recall of influenza-specific CD8\(^+\) T cell memory in the elderly is associated with functional profiles comparable to those found in the young. Since polyfunctionality (simultaneous IFN-\(\gamma\), TNF-\(\alpha\) and IL-2 production) of CD8\(^+\) T cells is thought to correlate with protective efficacy [54,55,56], establishing optimal cytokine profiles early may provide a clear advantage for virus-specific CD8\(^+\) T cell responses in the elderly.

Ageing is often associated with the attrition of the peripheral TCR repertoire, reflecting the loss of some T cell clonotypes and the large expansion of others [5,14,21]. Our study provides the most comprehensive analysis of the aged (primary, secondary primed-early and secondary primed-late) TCR repertoire published to date. The present, unbiased single-cell RT-PCR analysis of CDR3\(\beta\) usage in the elderly showed a diminished number of clonotypes during the aged primary D\(^b\)PA224 CD8\(^+\) responses when compared with the normal profiles for young individuals [44,47,57]. As naive D\(^b\)NP\(_{366}\) CD8\(^+\) and D\(^b\)PA224 CD8\(^+\) T precursors are efficiently recruited into the primary immune response [36], this primary repertoire analysis can be considered to reflect the loss of a substantial proportion of naive TCRs with ageing. Whilst a previous study [42] suggested that age-related clonal TCR attrition is more prevalent for the low precursor frequency D\(^b\)NP\(_{366}\) CD8\(^+\) repertoire, we found a greater reduction in the numbers of D\(^b\)PA224 V\(\beta\)8.3\(^+\) versus D\(^b\)NP\(_{366}\) CD8\(^+\) (down 34.9\% -specific nucleotide clonotypes per mouse recovered following primary infection of older mice (Tables 1 and 2). This is likely to reflect that there are a greater variety of n.t. types encoding public D\(^b\)NP\(_{366}\)-specific aa clonotypes across all mice than for D\(^b\)PA224-specific aa clonotypes, which potentially makes D\(^b\)PA224 CD8\(^+\) aa-defined clonotypes more vulnerable to total clonotype loss and thus reduced diversity.

The public, aa-defined D\(^b\)NP\(_{366}\) CD8\(^+\) CDR3\(\beta\) clonotypes can be encoded by up to 10 different n.t. sequences [44], meaning that the loss of one n.t.-defined public TCR may not necessary result in the elimination of that particular CDR3\(\beta\) aa sequence. Thus, it is not surprising that the D\(^b\)NP\(_{366}\) CD8\(^+\) CDR3\(\beta\) clonotypes in the aged mice following primary infection are encoded by a limited number of n.t. sequences (1.7 ± 1.1 per mouse) inferring a loss of D\(^b\)NP\(_{366}\)-specific CD8\(^+\) T cells. This was associated with the

Figure 8. Comparison between aged and young mice of the dominance of shared D\(^b\)NP\(_{366}\) CD8\(^+\) V\(\gamma\)8.3\(^+\) and D\(^b\)PA224 CD8\(^+\) V\(\beta\)7\(^+\) TCR clonotypes during primary and secondary (primed-young and primed-old) infections. Shown are the percentages of the D\(^b\)NP\(_{366}\) CD8\(^+\) V\(\gamma\)8.3\(^+\) (A, B) and D\(^b\)PA224 CD8\(^+\) V\(\beta\)7\(^+\) (C, D) TCR repertoires per mouse that are comprised of aa clonotypes shared between a particular number of mice (indicated by colour-coding) during primary (A, C) and secondary (B, D) infections. The number of mice sharing a TCR clonotype was determined across young and aged mice and both primary and secondary challenges. For example, the public D\(^b\)NP\(_{366}\)CD8\(^+\)V\(\gamma\)8.3\(^+\) TCR clonotype SGGANTGQL (red) was observed in 23 out of 30 mice (A, B). This clonotype contributed to 95% of the primary D\(^b\)NP\(_{366}\)CD8\(^+\)V\(\gamma\)8.3\(^+\) TCR repertoire of aged mouse M4 (A). There were five D\(^b\)NP\(_{366}\)CD8\(^+\)V\(\gamma\)8.3\(^+\) TCR clonotypes that were each observed in a number of mice ranging between 4 and 7 mice (beige). Three of these five clonotypes contributed to the D\(^b\)NP\(_{366}\)CD8\(^+\)V\(\gamma\)8.3\(^+\) TCR repertoire responding to secondary infection in young mouse M7 (B, as indicated the three beige segments). Multiple unshared clonotypes (light grey), which were observed in only one mouse, contributed to the TCR repertoires of many of the mice (as indicated by multiple light grey segments per column).

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Conservation of CD8\(^+\) T Cell Memory with Age
decreased contribution of two main public clonotypes (SGGGNTGQL and SGGNTGQL) and the increased prominence of one public clonotype (SGGANTGQL) in aged mice following primary influenza virus challenge. Similar epitope-specific TCR repertoire homogenisation across a population of aged mice has been recently observed for CD8+ T cell responses to H5N1 [22]. As SGGANTGQL is of lower pMHC avidity [49], the dominance of this clonotype in the aged repertoire may be one reason for the lower functional quality of DNP366+CD8+ T cell responses in the elderly.

The real advantage of priming CD8+ T cell responses early in life is reinforced by the demonstration that n.t.-defined clonotype diversity is preserved for the public DNP366+CD8+ T cell response, resulting in more equal contribution of the 3 main public clonotypes (SGGGNTGQL, SGGGGNTGQL and SGGNSGTGQL), which was not seen when mice were primed later in life (where SGGNTGQL alone dominated). Similarly, the secondary DPA224+CD8+ response in aged mice is slightly more diverse than that generated following primary virus challenge. Thus, early priming of the CD8+ T cell compartment induces a more diverse, aged repertoire by promoting the survival of public DNP366+CD8+ clonotypes. This may in turn reflect the selection of “best-fit” TCRs. Maintaining TCR repertoire diversity can enhance the efficacy of CD8+ T cell-mediated immunity [58], diminish the likelihood that mutated pathogens ‘escape’ immune recognition [59] and lead to more cross-reactive CD8+ T cell responses [53,60]. Preserving a greater breadth of responding TCRs is thus likely to be favorable for the elderly population. Taken together, our study supports the evolution of vaccine strategies to prime CD8+ T cells early in life in order to preserve the magnitude, functionality, TCR repertoire diversity and preferential TCR usage of responding populations.

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 influenza virus infection

Female C57BL/6j (B6, H2b) mice were bred and housed under specific pathogen free (SPF) conditions at the Department of Microbiology and Immunology, University of Melbourne. Primary responses: For generation of acute primary influenza CD8+ T cell responses, mice were lightly anaesthetised by inhalation of metohfluane and infected intranasally (i.n.) with 1 x 105 plaque forming units (pfu) of H3N2 (HK) influenza A viruses in 30 μl of PBS. Young mice were infected at 6–8 weeks, while aged mice were infected at 22 months of age. Secondary responses: To study the effects of early priming on aged CD8+ T cell responses, mice were first primed intraperitoneally (i.p.) at 6 weeks of age with 1.5 x 105 pfu of H1N1 PR8 influenza A virus and subsequently challenged with the serologically distinct H3N2 HK virus at extreme age of 22 months (6 weeks->22 months; primed young->challenged old). To study the effects of late priming on aged CD8+ T cell responses, mice were first primed i.p. with PR8 at 22 months and challenged 6 weeks later with HK (22 months ->23.5 months; primed old->challenged old). Control young animals were primed at 6 weeks, then challenged at 12 weeks of age (6 weeks->12 weeks; primed young->challenged young). The aged cohort of mice were held for up to 24 months in SPF conditions, monitored for signs of infection, weight loss and spontaneous tumor growth.

Tissue sampling and cell preparation

Spleens were recovered from mice at acute phases of the primary and secondary infections (day [d] 10 and d8, respectively). Spleens were depleted of B cells by incubation on 2 μg/G/ IgM coated plates (Jackson ImmunoResearch Labs) for 45 mins at 37°C, and uncold cells harvested.

Tetramer and phenotypic staining of CD8+ T cells

Enriched lymphocytes from the spleen were stained with DNP366 and DPA224 tetramers conjugated to Streptavidin-APC. For Vβ usage analysis, tetramer-stained cells were incubated a panel of FTTC conjugated anti-Vβ mAbs (2, 3, 4, 5.1/5.2, 6, 7, 8.1/8.2, 8.3, 9, 10, 12, 13, 14 and 17) [61] at 5 μg/μl, and 1 μg/ml anti-CD8-PerCP/Cy5.5. Cells were stained for 30 mins on ice, washed twice and analyzed by flow cytometry using a FACS Calibur (BD Biosciences) and Flowjo software (Treestar).

Peptide stimulation and intracellular cytokine staining

Splenocytes were stimulated with 1 μM NPE66 or PA24 peptides (AusPep) for 5 hrs at 37°C, 5% CO2 in the presence of 1 μg/ml Golgi-Plug (BD Biosciences) and 10 U/ml recombinant human IL-2 (Roche). Cells were washed twice with FACS buffer, stained with 1 μg/ml anti-CD8-PerCP/Cy5.5 mAb for 30 mins on ice, fixed, permeabilised using the BD Cytofix/Cytoperm kit and stained with 5 μg/ml anti-IFN-γ-FTTC, 2 μg/ml anti-TNF-α-APC, and 2 μg/ml anti-IL-2-PE mAbs. Samples were acquired by flow cytometry using a FACS Calibur and analysed by Flowjo. The total cytokine production was calculated by subtracting background fluorescence using no peptide controls.

Isolation of single-cell tetramer-specific CD8+ T cells, RT-PCR and CDR3β sequencing

Splenocytes were stained with DNP366-PE or DPA224-PE tetramers in sort buffer (PBS with 0.1% BSA) for 1 hr at room temperature, washed and stained with 1 μg/ml anti-CD8-APC and 5 μg/ml of either anti-Vβ8.3 or -PE-Vβ7-FTTC 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 twin-tec plate (Eppendorf). mRNA transcripts were reversed transcribed to cDNA, using a Sensiscript kit (Qiagen) according to manufacturer’s instructions, and the CDR3β region amplified by a nested PCR using Vβ primers [44,47,57]. Positive PCR products were purified using QIAGEN PCR purification kit and sequenced.

Statistical analysis

Magnitude, phenotype and function were compared between experimental aged and young groups by an unpaired Student’s t test. Clonotypic diversity was quantified using both the number of different clonotypes and Simpson’s diversity index. The overlap of...
TCRβ repertoires between mice was quantified using both the proportion of the TCRβ repertoires per mouse comprised of shared clonotypes and the Morisita-Horn similarity index. The Simpson’s diversity and Morisita-Horn similarity indices account for both the variety of distinct clonotypes (defined either at the level of the amino acid or nucleotide sequence) and the clone size (number of copies) of each clonotype involved in the epitope-specific response within each mouse [48,62]. The Simpson’s diversity and Morisita-Horn similarity indices vary between 0 (minimum diversity/similarity) and 1 (maximum diversity/similarity). The diversity and similarity measures were calculated in conjunction with a randomization procedure to correct for differences in sample sizes between mice [48,62], and were estimated for a subsample of 22 TCRβ sequences. To estimate the proportion of the TCRβ repertoires per mouse comprised of shared clonotypes, clonotypes were pre-defined as shared based on their presence in more than one mouse prior to the random subsampling of 22 sequences. A Mann-Whitney test was used to compare, between pairs of groups, the diversity (and similarity) between the aged and young groups of mice in primary responses and between young and aged (primed-young) and aged (primed-old) in secondary responses. Bonferroni correction for multiple pairwise comparisons was applied for the comparisons between the three secondary response groups (i.e. each pairwise test was assessed at the significance level of \( p = 0.05/3 = 0.0167 \)).

All statistical analyses were performed using GraphPad Prism version 5.04 (GraphPad Software Inc, San Diego, CA).

Supporting Information

Figure S1  Analysis of acute and memory CD8+ T cell responses elicited by i.p. priming. Naïve B6 mice were i.p. primed with 1.5 \times 10^7 pfu of the PR8 virus. Influenza-specific CD8+ T cell responses were analysed in the spleen at the acute (d10), early memory (d23) and late memory (10 mths) phases of infection. (A) Total numbers of tetramer+ CD8+ T cells are shown for immunodominant DNP366+ CD8+ and DPAPA224+ CD8+ T cell responses. (B) The contribution of immunodominant DNP366+ CD8+, DPAPA224+CD8+ T cell responses in comparison with subdominant DPB1707+CD8+ and KPB1-F262+CD8+ sets were calculated based on the proportions of IFN-γ+CD8+ populations. (C, D) Polyclonality of influenza-specific CD8+ T cell responses was assessed by the size of clonotypes [57,58].

Figure S2 The expression of activation markers on influenza-specific CD8+ T cells in young and aged mice. Phenotypic analysis of (A, D) CD27 vs CD43, (B, E) KLRG1 vs IL-7R, and (C, F) CD69 vs IL-7-R was determined at the acute day 8 secondary time-point for (A–C) DNP366+ and (D–F) DPAPA224 splenocytes from aged mice primed at 3 months and challenged at 22 months in comparison to young animals. Similar phenotypic data were obtained when aged mice were either primed at 22 months (primary response) or primed when young (at 6 weeks) and challenged at 22 months (secondary response). Data represent the mean ± SD of 3–5 mice per group, * = p<0.05.

Figure S3 Comparison between aged and young mice of the characteristics of the DNP366+CD8+V8.3+ and DPAPA224+CD8+V77+ TCR repertoires during primary and secondary (primed-young and primed-old) infections. The distributions of Jβ gene usage (A, B, E, F) and CDR3β length (C, D, G, H) among all DNP366+CD8+V8.3+ TCR sequences during primary (A, C) and secondary (B, D) infections and all DPAPA224+CD8+V77+ TCR sequences during primary (E, G) and secondary (F, H) infections.

Table S1 Nucleotide and amino acid CDR3β diversity profiles for primary DNP366+V8.3+CD8+ T cells in the aged (22months) mice.

Table S2 CDR3β diversity profiles for primary DPAPA224+V77+CD8+ T cells in the aged (22months) mice.

Table S3 Nucleotide and amino acid CDR3β diversity profiles for secondary DNP366+V8.3+CD8+ T cells in the aged (primed at 2months) mice.

Table S4 CDR3β diversity profiles for secondary DPAPA224+V77+CD8+ T cells in the aged (primed at 2months) mice.

Table S5 Nucleotide and amino acid CDR3β diversity profiles for secondary DNP366+V8.3+CD8+ T cells in the aged (primed at 2months) mice.

Table S6 Amino acid CDR3β diversity profiles for secondary DPAPA224+V77+CD8+ T cells in the aged (primed at 2months) mice.

Author Contributions

Conceived and designed the experiments: KK SAV MPD VV PCD. Performed the experiments: SAV MPD VV PCD. Analyzed the data: SAV KK VV THYD NLB. Wrote the paper: SAV KK VV THYD NLB MPD SJT PCD.

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