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Aging promotes acquisition of naive-like CD8⁺ memory T cell traits and enhanced functionalities

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Protective T cell memory is an acquired trait that is contingent upon the preservation of its constituents and therefore vulnerable to the potentially deleterious effects of organismal aging. Here, however, we have found that long-term T cell memory in a natural murine host-pathogen system can substantially improve over time. Comprehensive molecular, phenotypic, and functional profiling of aging antiviral CD8⁺ memory T cells (CD8⁺ T_M) revealed a pervasive remodeling process that promotes the gradual acquisition of distinct molecular signatures, of increasingly homogeneous phenotypes, and of diversified functionalities that combine to confer a CD8⁺ T_M-autonomous capacity for enhanced recall responses and immune protection. Notably, the process of CD8⁺ T_M aging is characterized by a progressive harmonization of memory and naive T cell traits, is broadly amenable to experimental acceleration or retardation, and serves as a constitutional component for the “rebound model” of memory T cell maturation. By casting CD8⁺ T_M populations within the temporal framework of their slowly evolving properties, this model establishes a simple ontogenetic perspective on the principal organization of CD8⁺ T cell memory that may directly inform the development of improved diagnostic, prophylactic, and therapeutic modalities.

Introduction

The complex of aging, memory, and protection has long been recognized as a theme central to the immunological sciences. In at least one respect, though, this topic has not yet been explored in detail: how aging shapes and transforms the cardinal properties of T cell memory established early in life in response to acute infectious insults. The principal component of protective T cell memory is the population of pathogen-specific CD8⁺ memory T cells (CD8⁺ T_M) that emerges, following resolution of acute disease, within several weeks through a process that combines the preferential survival of defined CD8⁺ effector T cell (CD8⁺ T_E) subsets with their gradual maturation. The subsequent maintenance of specific CD8⁺ T_M over extended time periods is integral to the organism's capacity to curtail secondary (IIP) infections, minimize clinical disease, and avert potential death (1–4). Conversely, any measures that modulate CD8⁺ T_M homeostasis and preservation, including the process of aging, may also subvert effective immune protection. Indeed, a substantial body of literature demonstrates that aging can exert a wide-spread and often deleterious influence on the efficient coordination of T cell responses by compromising or distorting relevant organ systems (thymic involution), the distribution of T cell subsets (loss of naive T cells [T_N], oligoclonal expansions of memory-phenotype T cells [T_{MP}]) and specificities (reduced TCR diversity), as well as altering T cell phenotypes,

signal transduction, metabolism, telomere biology and functionalities (diminished responsiveness, exhaustion, replicative senescence, etc.) (5–10). Although the available evidence suggests that CD8⁺ T_M populations maintained under ideal conditions (i.e., in the absence of persisting or deliberately introduced antigen) are less adversely affected by the ravages of age (3, 10), pathogen-specific CD8⁺ T_M appear to slowly evolve, as documented by their gradual though seemingly limited phenotypic and functional conversion (11–17). At a minimum, these observations emphasize the dynamic nature of CD8⁺ T_M maturation, but they may also insinuate more extensive adaptations that CD8⁺ T_M accrue over time and that may ultimately modify the efficacy of immune protection. In fact, a recent report by Martin et al. provided the first evidence for enhanced antiviral protection exerted by late (>8 months after challenge) as compared with early (day 30–45) CD8⁺ T_M (17).

To better define the nature, scope, and consequences of pathogen-specific CD8⁺ T_M aging, we employed a natural host-pathogen system — the acute infection of mice with lymphocytic choriomeningitis virus (LCMV) — and delineated the progressive accumulation of distinctive molecular, phenotypic, and functional properties of CD8⁺ T_M populations maintained for life. Our results demonstrate that aging of antiviral CD8⁺ T_M is defined by a prolonged and pronounced molecular remodeling process that is associated with the acquisition of a unique, more homogeneous phenotype emulating the appearance of naive CD8⁺ T cells (CD8⁺ T_N). Furthermore, this remodeling process imparts surprisingly favorable traits onto aged CD8⁺ T_M (diversified effector functionalities, enhanced IIP reactivity, and improved protective capacity) and can be accelerated or

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restrained as a whole by modulating the extent and/or speed of primary (I°) $CD8^{+} T_{E}$ differentiation.

Beyond the context of aging, however, the primacy of temporal relationships between multiple $CD8^{+} T_{M}$ subsets outlined here permits the development of a potentially novel perspective on the generation and maintenance of $CD8^{+} T$ cell memory that correlates protective potential with a progressive alignment of $CD8^{+} T_{M}$ and T_{N} properties, that posits a dedifferentiation process as a core feature of $CD8^{+} T$ cell memory in general, and that we have organized in a modification of the decreasing potential model (1, 4) termed the “rebound model” of $CD8^{+} T_{M}$ maturation. The notion that dedifferentiated $CD8^{+} T_{M}$ with defined molecular traits, phenotypic signatures, and functional properties can provide enhanced protection will likely have practical implications for public health initiatives such as the design of immunization strategies, monitoring of vaccine efficacy, and the development of improved treatment modalities for infectious diseases.

Results

Enhanced I° reactivity of aged $CD8^{+} T_{M}$. The LCMV model of acute viral infection permits the generation of specific $CD8^{+} T_{M}$ that are maintained for life and poised to be effectively recruited into a I° response (11). To assess the recall potential of aging $CD8^{+} T_{M}$ populations in greater detail, we utilized an in vivo system in which I° expansions of both young and old $CD8^{+} T_{M}$ populations could be monitored within the same host. In brief, cohorts of young adult B6 and B6-congenic mice were challenged with LCMV in a staggered fashion and allowed to age for up to ~2 years. Young (d58 after infection) and old (d850) $CD8^{+} T_{M}$ were then isolated, mixed at a ratio of 1:1, and adoptively transferred (AT) into naive congenic recipients that were challenged with LCMV (Figure 1A). Despite their advanced age, old $CD8^{+} T_{M}$ consistently gave rise to greater $I^{\circ} CD8^{+} T_{E}$ expansions, a phenomenon that is observed in lymphatic and nonlymphoid tissues alike and independent of the particular choice of congenic T cell marker (Figures 1, A and B, and Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI88546DS1). Similar findings were originally reported for the Sendai virus system (14) and more recently also for *Listeria monocytogenes* and LCMV models (16, 17), providing evidence for the more general occurrence of this phenomenon and a rationale for further investigations into the nature of $CD8^{+} T_{M}$ recall responses that, paradoxically, improve with age.

Differentiating LCMV-specific $CD8^{+} T$ cell responses according to 7 MHC-I-restricted epitopes, we found the age-associated augmentation of $I^{\circ} CD8^{+} T_{E}$ expansions to be independent of immunodominant determinants and functional avidities, an observation that was also reflected in the comparable frequencies of specific $I^{\circ} CD8^{+} T_{E}$ within the respective young and old donor $CD8^{+} T$ cell compartments (Figure 1C and Supplemental Figure 1B). Indeed, as determined by granzyme B (GZMB) and CCL5 content (12, 18), the entire pool of expanded donor $CD8^{+} T$ cells bore hallmarks of specific activation, and both young and old $I^{\circ} CD8^{+} T_{E}$, regardless of their differential expansion, exhibited a comparable spectrum of inducible functionalities (Supplemental Figure 1, C and D). The improved accumulation of $I^{\circ} CD8^{+} T_{E}$ derived from old $CD8^{+} T_{M}$ must be grounded in their enhanced proliferation and/or survival, and to distinguish between these possibilities

we determined expansion kinetics, proliferation rates, and survival of $CD8^{+} T_{E}$ throughout the I° response. In the experimental setting employed here, $I^{\circ} CD8^{+} T_{E}$ typically start to divide only more than 48 hours after LCMV infection (19), and the comparable proliferation indices and $I^{\circ} CD8^{+} T_{E}$ numbers found in the spleen 60 hours after challenge demonstrated that young and old $CD8^{+} T_{M}$ were recruited into the peripheral I° response with similar kinetics (Figure 1D). Differential expansion of $I^{\circ} CD8^{+} T_{E}$, however, became apparent in blood and spleen on day 5 after AT/rechallenge, was further amplified on subsequent days (Figure 1, D and E), and primarily emerged through the better survival of old $I^{\circ} CD8^{+} T_{E}$ rather than their faster or prolonged proliferation (Supplemental Figure 1, E-G).

Kinetics of early $I^{\circ} CD8^{+} T_{E}$ expansions in draining and non-draining lymph nodes (LNs). In order to visualize the earliest events of the I° response in vivo, we resorted to the p14 chimera system: purified naive p14 cells (TCRtg $CD8^{+} T$ cells specific for the LCMV-GP₃₃ determinant) were transferred into B6 recipients that were inoculated with LCMV and subsequently served as donors for p14⁺ T_{M} . Within 48 hours after p14⁺ T_{M} AT/rechallenge, old but not young I° p14⁺ T_{E} largely disappeared from the circulation (Figure 1F), suggesting their sequestration to other tissues, most likely LNs (20). In the ensuing 16 hours, old I° p14⁺ T_{E} returned to the blood, and both young and old I° p14⁺ T_{E} began to proliferate at a comparable pace in the spleen (Figure 1G). At the same time, however, old and young I° p14⁺ T_{E} responses already began to diverge in the LNs, especially the draining mediastinal LNs (MedLNs) (21) where enhanced proliferation rates of aged I° p14⁺ T_{E} correlated with their significantly increased numbers (Figure 1H). These observations could also be extended to the B6 system and the first emergence of differential $I^{\circ} CD8^{+} T_{E}$ expansions on day 5 after challenge: here, old outnumbered young I° D^bNP₃₉₆ $CD8^{+} T_{E}$ by a factor of ~3 in the MedLNs but by less than 2-fold in blood, spleen, and other LNs (Figure 1, E and I). Thus, greater I° expansions of aged $I^{\circ} CD8^{+} T_{E}$, driven from day 5 onwards primarily by a survival advantage, were precipitated between days 2 and 4 by an earlier onset of proliferation in LNs, in particular the MedLNs.

Improved protective capacity of aged $CD8^{+} T_{M}$ as a T_{M} -intrinsic property. Taken together, our results suggest that aging $CD8^{+} T_{M}$ acquire certain intrinsic properties, although the potential influence of several variables such as recipient age and adaptive changes of transferred $CD8^{+} T_{M}$, contemporaneous I° responses generated by the host, and a direct competition between concurrent $I^{\circ} CD8^{+} T_{E}$ responses has not yet been ascertained. To determine if the acquisition of enhanced I° reactivity by aging $CD8^{+} T_{M}$ simply constituted a natural adaptation that could be reversed upon prolonged exposure to a young host environment or rather was restricted to that particular environment, we conducted complementary mixed AT/rechallenge experiments and found that neither extended parking of transferred cells prior to infection nor advanced host age affected the differential expansion of old versus young $I^{\circ} CD8^{+} T_{E}$ (Figure 2, A and B). Since the AT of low $CD8^{+} T_{M}$ numbers does not preclude the development of simultaneous host T cell responses directed against the same LCMV determinants, we also monitored $I^{\circ} CD8^{+} T_{E}$ populations generated by young and old recipients. Interestingly, and consistent with the notion that aged mice have a reduced capacity to mount de novo antiviral $CD8^{+} T_{E}$ responses (22), the advanced

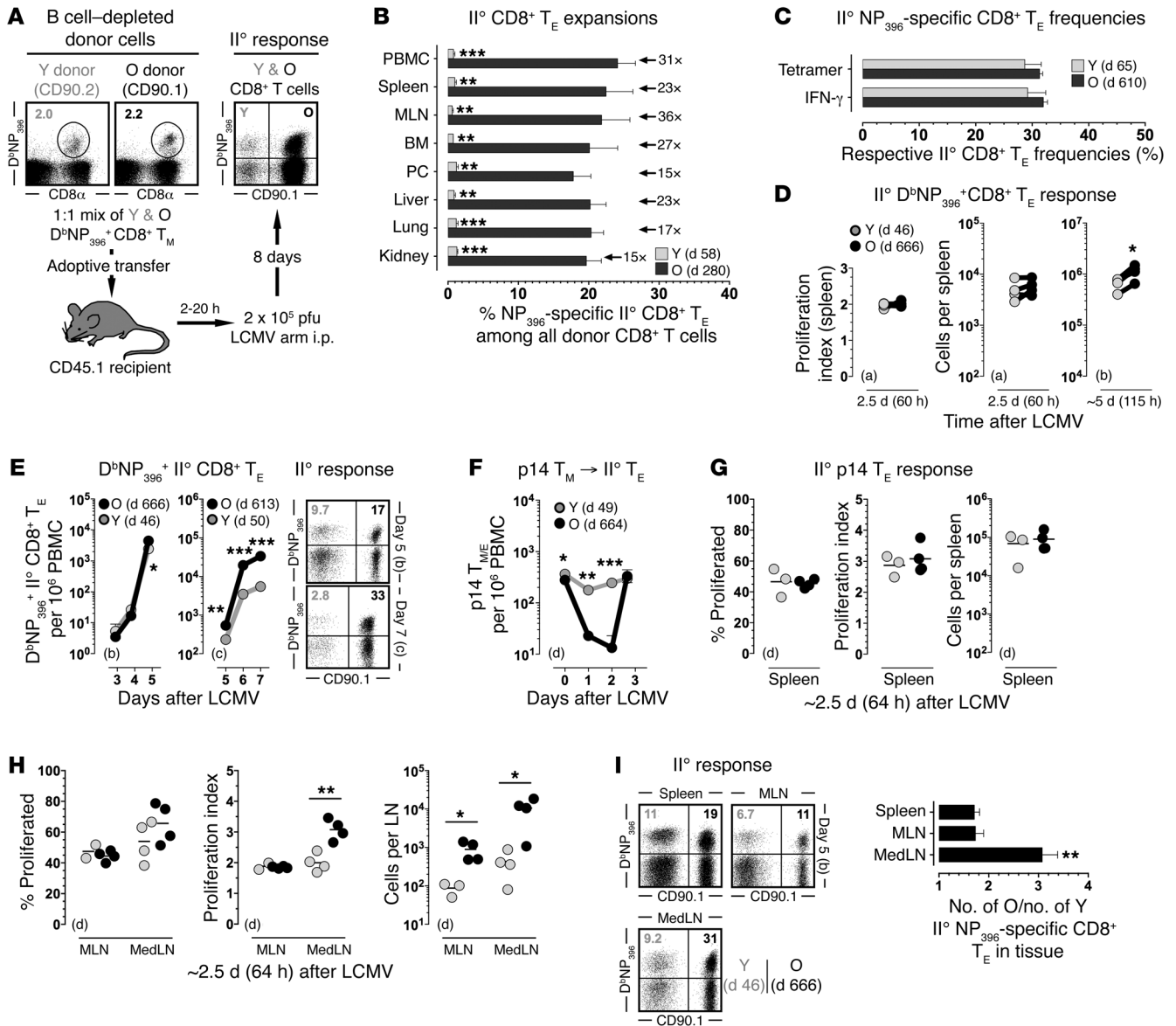


Figure 1. Enhanced II° reactivity of aged CD8+ T_M. (A) Outline for mixed AT/rechallenge experiments. Unless noted otherwise, CD8+ T cell-enriched populations containing 2 × 10³ congenic young (Y) and old (O) D^bNP₃₉₆+CD8+ T_M each were transferred i.v. into congenic recipients prior to i.p. LCMV challenge and analysis of II° CD8+ T_E expansions on day 8. (B) Relative frequencies of II° D^bNP₃₉₆+CD8+ T_E among all donor CD8+ T cells; donor age (time after I° LCMV challenge), factor of differential II° expansion and statistical significance are indicated. (C) Respective frequencies of NP₃₉₆-specific II° CD8+ T_E in young and old CD8+ donor T cell compartments as assessed by tetramer or IFN-γ staining. (D) Mixed AT/rechallenge experiments were conducted with CFSE-labeled donor cells to calculate proliferation indices; corresponding data points from individual recipients are connected by a line. (E) Expansion kinetics of II° CD8+ T_E in peripheral blood (dot plots gated on all donor CD8+ T cells). (F–H) AT/rechallenge experiments were performed with CFSE-labeled p14 T_M. (I) II° D^bNP₃₉₆+CD8+ T_E responses analyzed on day 5. To visualize earlier stages of the II° response in panels D–I, escalating numbers of CD8+ T cell-enriched donor populations were transferred containing (a) 2 × 10⁵, (b) 4 × 10⁴ or (c) 5 × 10³ young and old D^bNP₃₉₆+CD8+ T_M each, or (d) 1 × 10⁶ p14 T_M. Data are the mean ± 1 SEM or feature individual data points with n ≥ 3 mice for multiple independent experiments. *P < 0.05, **P < 0.01, and ***P < 0.001 by 1 way ANOVA or Student's t test. BM, bone marrow; MedLN, mediastinal lymph nodes; MLN, mesenteric lymph nodes; PBMC, peripheral blood mononuclear cells; PC, peritoneal cavity.

age of recipient mice significantly compromised their I° CD8+ T_E responses despite the fact that it readily supported enhanced II° expansions of aged CD8+ T_M (Figure 2B). While these data suggest an at least partially independent regulation of I° and II° CD8+ T cell immunity, a direct evaluation of this contention required a model system in which II° CD8+ T cell immunity could be studied in the complete absence of concurrent I° T cell responses: OT-I mice, due to their skewed TCR specificity for ovalbumin, cannot generate

LCMV-specific T cell responses, but otherwise provide a T cell-replete in vivo environment that eschews artifacts associated with T cell deficiency (23). As shown in Figure 2C, the disparate reactivity of old versus young CD8+ T_M was preserved in OT-I recipients, indicating that their differential II° reactivity is not affected by contemporaneous I° CD8+ T_E responses.

We next addressed the issues of potential competition among II° CD8+ T_E populations and immune protection by

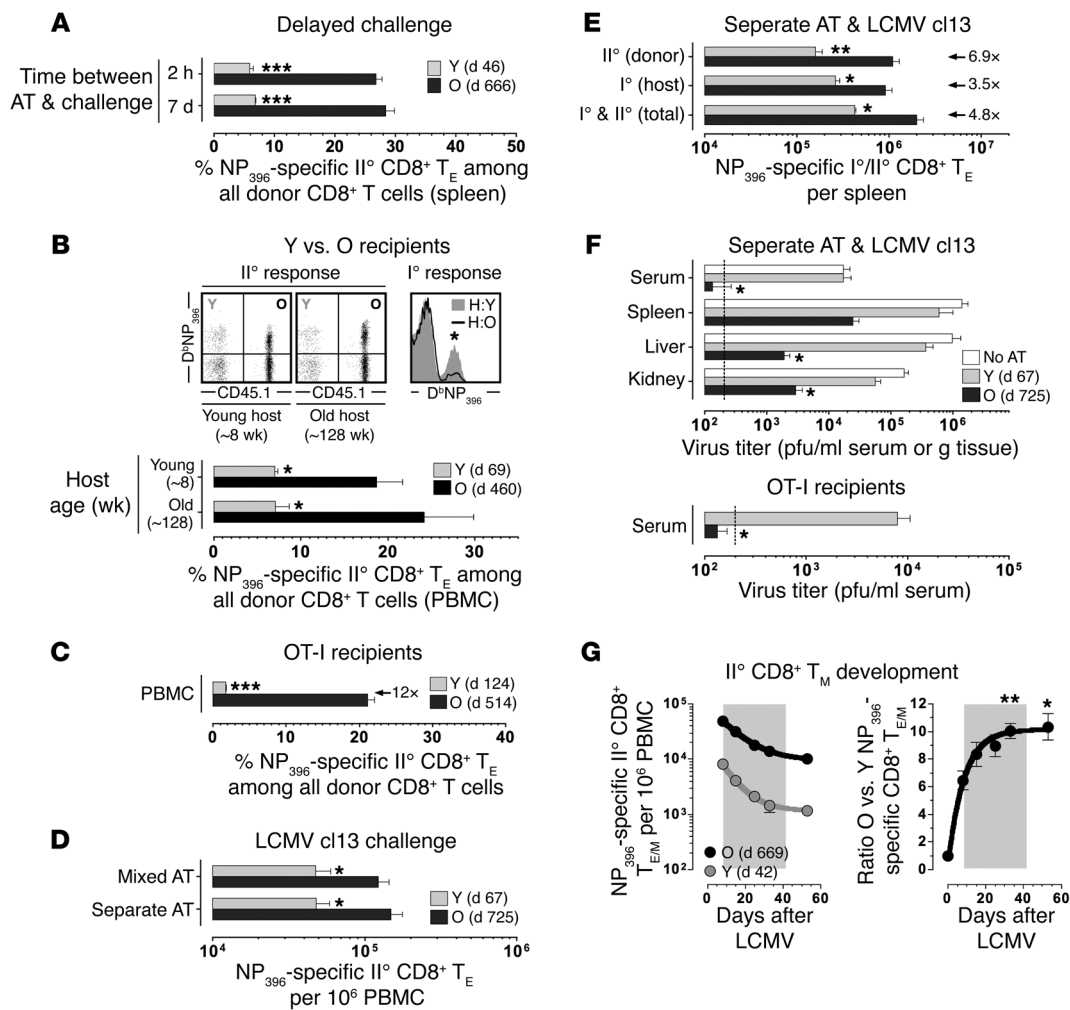


Figure 2. Improved protective capacity of aged CD8⁺ T_M as a T_M-intrinsic property. (A) Recipients of mixed young (Y) and old (O) D^bNP₃₉₆⁺CD8⁺ T_M populations (5 × 10³ each) were challenged 2 hours or 7 days after AT. (B) II^o expansions of young vs. old CD8⁺ T_M transferred into naive young vs. old recipients. Upper right: concurrent I^o D^bNP₃₉₆⁺CD8⁺ T_E responses generated by young (~8 weeks) vs. old (128 weeks) hosts. (C) Mixed AT/rechallenge experiments using OT-I recipients. (D) Young and old donor cells depleted of B and CD4⁺ T cells were combined or transferred separately prior to LCMV cl13 challenge. (E) Quantification of concurrent I^o and II^o CD8⁺ T_E responses after separate AT/LCMV cl13 rechallenge as performed in panel D. (F) Top, virus titers in tissues (day 8) after separate AT/LCMV cl13 rechallenge; bottom, AT of 5 × 10³ D^bNP₃₉₆⁺CD8⁺ T_M, OT-I recipients and determination of viral titers on day 6. (G) Contraction kinetics of II^o CD8⁺ T_E populations after LCMV Armstrong challenge; asterisks indicate a significant rise of the O:Y ratio as compared with day 8 (gray background = contraction phase); data generated with n ≥ 3 mice/group in multiple independent experiments. *P < 0.05, **P < 0.01, and ***P < 0.001 by 1 way ANOVA or Student's *t* test. PBMC, peripheral blood mononuclear cells.

use of mixed versus separate AT experiments and rechallenge with high-dose LCMV clone13 (cl13). Here, LCMV cl13-infected recipients of old and young CD8⁺ T_M, whether transferred separately or in combination, supported the enhanced II^o proliferation of aged CD8⁺ T_M in an identical fashion, ruling out that differential expansion resulted from direct competition between II^o CD8⁺ T_E populations (Figure 2D). We also note that the heightened II^o expansion of aged CD8⁺ T_M in fact boosted I^o LCMV-specific CD8⁺ T_E reactivity, a likely consequence of accelerated virus control (Figure 2E). Indeed, as determined by infectious virus titers in the separate AT/LCMV cl13 rechallenge experiments and similar to the findings of Martin et al. (17), aged CD8⁺ T_M populations mediated significantly enhanced immune protection (Figure 2F), a phenomenon also observed in OT-I recipients and thus in an experimental scenario where

LCMV-specific T cell responses are exclusively derived from the donor population (Figure 2F).

We also monitored the generation of II^o CD8⁺ T_M cells using the mixed AT/LCMV Armstrong rechallenge protocol and confirmed the attenuated contraction of II^o in relation to I^o CD8⁺ T_E populations (24) (Figure 2G and not shown). Importantly, a lesser contraction of aged as compared with young II^o CD8⁺ T_E further amplified their numerical differences and promoted a pronounced disparity of respective II^o CD8⁺ T_M pool sizes (Figure 2G). Altogether, these observations demonstrate that antiviral CD8⁺ T_M are subject to an extended maturation process that results in the acquisition of CD8⁺ T_M-intrinsic properties that foster enhanced II^o reactivity, immune protection, and II^o CD8⁺ T_M potential.

Global phenotypic and molecular alterations of aging antiviral CD8⁺ T_M. Given the relative dearth of knowledge about the mod-

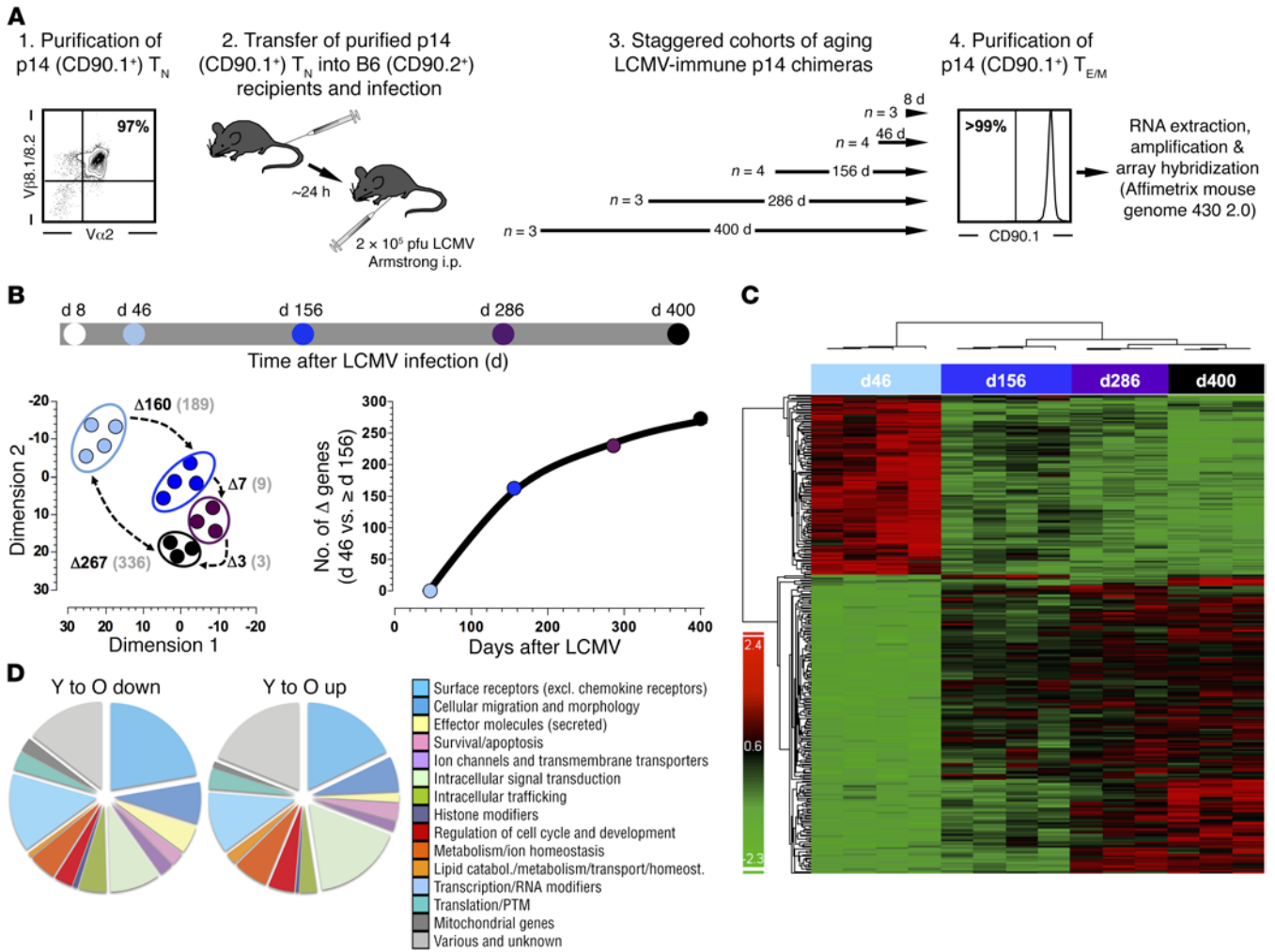


Figure 3. Molecular remodeling of aging antiviral CD8⁺ T_M. (A) Generation of LCMV-immune p14 chimeras (B6 mice transduced with 5 × 10⁴ purified congenic p14⁺ T_N prior to LCMV infection) was performed in a staggered fashion such that isolation of splenic p14⁺ T_E (day 8) and p14⁺ T_M (days 46, 156, 286, and 400) could be performed at the same time. (B) Top, timeline/color code for microarray analyses of purified p14⁺ T_{E/M}; bottom left, MDS analysis of DEGs (P < 0.05, FDR < 5%); numbers represent DEGs (black) and individual probe sets (gray) comparing the indicated p14⁺ T_M populations (P < 0.05, fold change > 2.0); bottom right, numbers of DEGs comparing day-46 and older p14⁺ T_M. (C) Heat-map clustering of p14⁺ T_M DEGs. (D) Distribution of p14⁺ T_M DEGs (day 46 [Y] vs. day 400 [O]; see Supplemental Table 1) across 15 major gene ontology categories.

ulation of pathogen-specific CD8⁺ T_M maintained for extended periods of time, we performed an initial survey of aging LCMV-immune mice to detail the temporal regulation of major antigens expressed by specific CD8⁺ T_M. Perhaps surprisingly, none of these markers nor cellular size or granularity/complexity were maintained at stable levels. Ranging from the marginal (cell size and granularity/complexity) to the modest (CD44⁺CD45⁺TCRVβ⁺CD3ε⁺CD8α) and more pronounced (MHC-I⁺CD90⁺), specific CD8⁺ T_M properties were subject to an extended maturation process that lasted for at least ~2 years and introduced significant increases among all parameters evaluated (Supplemental Figure 2).

To delineate the extent of molecular remodeling in aging antiviral CD8⁺ T_M, we generated LCMV-immune chimeras in a staggered fashion followed by contemporaneous purification of I⁺ p14⁺ T_E (d8) and T_M (d46, d156, d286, and d400) from individual mice, and processing for microarray hybridization (Figure 3A). Basic microarray analyses were conducted using both PGS (Fig-

ure 3, B and C) and Genespring (Figure 3D) software and yielded essentially similar results: multidimensional scaling (MDS) of differentially expressed genes (DEGs) in individual p14⁺ T_M populations demonstrated distinctive clustering according to age; molecular differences accrued with a diminishing pace over time; and unsupervised hierarchical clustering identified 2 clusters of genes, the expression of which progressively decreased (117 genes, ~2/5 DEGs) or increased (185 genes, ~3/5 DEGs) with age (Figure 3, B and C, and Supplemental Table 1).

Manual classification of DEGs into broad gene ontology-inspired categories revealed a relative dominance of cell surface receptors/ligands, migration- and morphology-associated genes, and intracellular signaling molecules, suggesting that CD8⁺ T_M aging may be accompanied by alterations of their phenotypes and reactivities (Figure 3D). Indeed, ingenuity pathway analysis (IPA) identified several canonical signaling pathways that included cytokines, chemokines/receptors, major costimulatory

Table 1. Rank-ordered listing of genes exhibiting >5-fold downregulation in old (day 400) vs. young (day 46) p14⁺ T_M and corresponding differential protein expression by D^bNP₃₉₆⁺CD8⁺ T_M

Rank	Genes downregulated > 5-fold by aging p14 ⁺ T _M			Proteins downregulated by aging D ^b NP ₃₉₆ ⁺ CD8 ⁺ T _M		
	Gene ID	Gene	FC ^a	FC	Figure	
1	14938	<i>Gzma</i>	-88.2	-10.4	7A	
2	16634/40	<i>Klra3/Klra9</i>	-45.2	-3.4	4A	
3	241447	<i>Lass6</i>	-32.5			
4	20308	<i>Ccl9</i>	-27.8	-1.7	Unpublished ^b	
5	11658	<i>Alcam (CD166)</i>	-25.8	-2.0	S10A	
6	13051	<i>Cx3cr1</i>	-20.7	-7.0	10A	
7	15277	<i>Hk2</i>	-15.7			
8	16364	<i>Irf4</i>	-12.4			
9	243655	<i>Klre1</i>	-12.2	not translated	4A	
10	17059	<i>Klrb1c</i>	-11.7	-3.5	4A	
11	103012	<i>Firre</i>	-10.4			
12	433470	<i>AA467197 (Nmes1)</i>	-9.5			
13	14425	<i>Galnt3</i>	-8.0			
14	57916	<i>Tnfrsf13b (TACI)</i>	-7.6	not translated	S7	
15	74103	<i>Nebi</i>	-7.5			
16	20947	<i>Swap70</i>	-7.0			
17	50928	<i>Klrg1</i>	-7.0	-3.0	4A	
18	19713	<i>Ret</i>	-6.9	-1.8	S10C	
19	20305	<i>Ccl6</i>	-6.1	not translated	Unpublished ^b	
20	72661	<i>Serp2</i>	-5.8			
21	77647	<i>Trat1 (TRIM)</i>	-5.5	-1.1	S8B	
22	21413	<i>Tcf4</i>	-5.3			
23	27218	<i>Slamf1 (CD150)</i>	-5.3	not translated	S9A	
24	16639	<i>Klra8</i>	-5.2	not translated	4A	

^aAll fold-changes (FC) are significant (genes, adjusted $P < 0.0025$; proteins, $P < 0.05$) (1-way ANOVA).

^bUnpublished observations (JE and DH).

ry components, and TLRs (Supplemental Table 2). Of particular prominence were IL-6, IL-10, and TGF- β signaling pathways, all of which have been implicated in the regulation of CD8⁺ T cell immunity under conditions of chronic LCMV infection (25–27), and we note the potential relevance of altered Wnt/ β -catenin signaling, a pathway that promotes the generation of CD8⁺ stem cell memory T cells (CD8⁺ T_{SCM}) (28) (Supplemental Table 2). Lastly, higher order network analyses revealed 16 networks containing at least 10 focus molecules, a moderate degree of connectivity, and predominant signature functions of cell death, cell-to-cell signaling/interactions, and cellular movement (Supplemental Table 3). Thus, the process of aging introduced a series of substantial and diverse molecular changes into a clonotypic CD8⁺ T_M population that collectively point towards altered phenotypic and functional traits that may directly shape evolving aspects of CD8⁺ T_M homeostasis and II^o reactivity.

Coordinated regulation of related gene and protein expression by aging CD8⁺ T_M: the C-type lectins. Based on the parallel observations of progressive phenotypic and molecular CD8⁺ T_M conversion processes, we initiated a systematic survey of corresponding gene and protein expression dynamics. Focusing at first on a subset of genes exhibiting a particularly pronounced expression decline (>5-fold downregulation in old vs. young p14⁺ T_M), we found that nearly half of these transcripts (11 of 24) coded for cell surface receptors/ligands, and that approximately half of the latter molecules

belonged to the same subgroup of C-type lectins, the KLR family (Table 1). We therefore broadened our analytical strategy to investigate the temporal regulation of all murine KLR members. Here, among the 11 of 45 KLR genes present in p14⁺ T_M, 10 demonstrated a progressive and significant downmodulation in aging p14⁺ T_M (Supplemental Table 4). Corresponding analyses of KLR gene products conducted with polyclonal aging D^bNP₃₉₆⁺CD8⁺ T_M demonstrated an excellent correlation between the temporal regulation of mRNA (p14 chimeric system) and protein (B6 system) expression: while some KLR genes were not translated (*Klra8* and *Klre1*), the levels of other KLR proteins declined significantly with advancing age; only *Klrl1* deviated from this pattern due to stable expression by practically all specific CD8⁺ T_M (Figure 4A and Supplemental Table 4).

The global decline of KLR gene and protein expression is noteworthy since it stands in apparent contrast with the notion that elevated expression of human

KLR and KIR (functional ortholog of the murine KLR subset of Ly49 genes) proteins is a hallmark of presenescent and senescent T cells in aged populations (9). To reconcile these discrepancies, we determined KLR expression by aging CD8⁺ T cells of undefined specificity in unmanipulated B6 mice. In confirmation and extension of previous reports (29), we observed a gradual acquisition of KLR⁺ phenotypes, with some receptors (*Klrc1*, *Klrd1*, and *Klrl1*) eventually expressed by more than 50% of old CD8⁺ T cells (Supplemental Figure 3). The fact that aging CD8⁺ T cells at large are subject to progressive alterations that are diametrically opposed to those observed for virus-specific CD8⁺ T_M emphasizes the importance of analytical CD8⁺ T cell-subset differentiation, and if enhanced T cell-expressed KLRs are considered an indicator for possible senescence, the steady loss of KLR⁺ phenotypes by aging antiviral CD8⁺ T_M may be viewed as a reversal thereof.

In light of the good correspondence between mRNA and protein expression data, we extended our analyses to the entire family of C-type lectins including the prominent members *Cd62l* and *Cd69* (Figure 4B, Supplemental Figure 4, and Supplemental Table 4). As expected (12), CD62L expression by aging CD8⁺ T_M increased substantially, but the abundance and subtle rise of *Cd69* mRNA was unexpected since CD69 protein is not detectable in resting splenic CD8⁺ T_M populations (Figure 4B). Rather, CD69 expression is a hallmark of tissue-resident CD8⁺ T_M and LN-populating subsets, where it contributes to the limitation of T cell

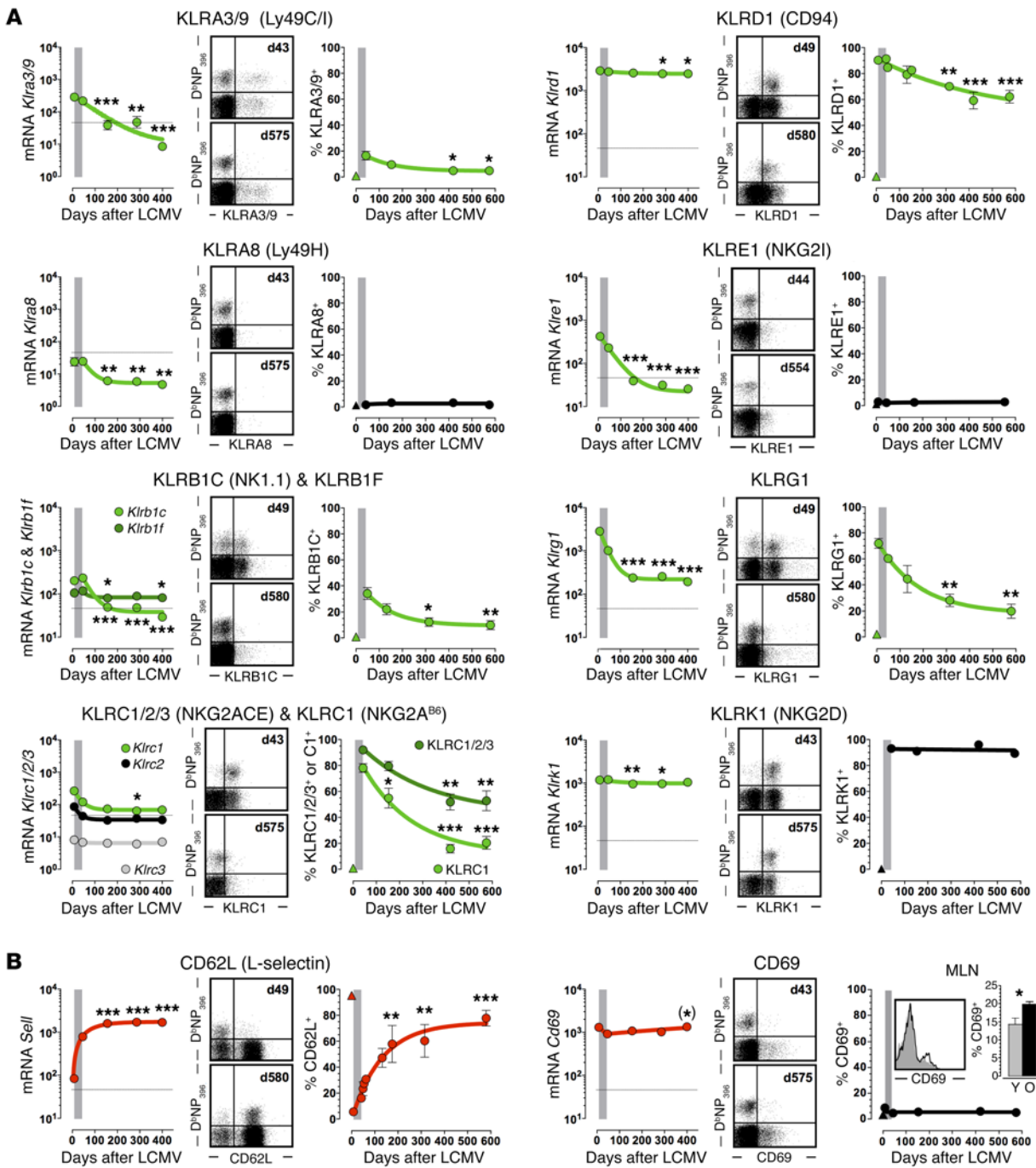


Figure 4. Temporal regulation of C-type lectin expression by virus-specific CD8⁺ T_{E/M}. The figure is organized into modules consisting of 3 panels each: left, mRNA expression (p14⁺ T_{E/M} microarrays); middle, representative dot plots of young (top) and old (bottom) splenic CD8⁺ T cells unless indicated otherwise; right, temporal regulation of D^bNP₃₉₆-CD8⁺ T_{E/M} properties. The gray bars demarcate the transition from T_E to early T_M stage (day 42), and the triangle symbols refer to CD44^{lo}CD8⁺ T_N. All values are the mean ± 1 SEM with n ≥ 3 individual mice per time point; *P < 0.05, **P < 0.01, and ***P < 0.001 comparing young and older (> day 100) p14⁺ T_M or D^bNP₃₉₆-CD8⁺ T_M, respectively, using 1-way ANOVA. Significant differences emerging during the memory phase are highlighted in red (upregulation) or green (downregulation). This outline also serves as a template for Supplemental Figures 4–14 and the data are further summarized and expanded to related genes in Supplemental Table 4. (A) Temporal regulation of KLR expression. (B) Temporal regulation of CD62L and CD69 expression. (*) indicates statistical significance determined by Student's *t* test rather than ANOVA. The insert in the CD69 module depicts young (gray, day 67) and old (black, day 725) D^bNP₃₉₆-CD8⁺ T_M recovered from MLNs and a corresponding data summary (Y, day 66; O, day 563). MLN insert bar diagram analysis in (B) determined by Student's *t* test.

gress (30). Notably, the fraction of LCMV-specific CD69⁺CD8⁺ T_M isolated from LNs was slightly elevated among aged populations (Figure 4B), perhaps a contributing factor to altered trafficking patterns of aged CD8⁺ T_M (JE, BD, and DH, unpublished

observations). Expression of 3 additional gene products (CD72b/c, CD93, and CD205) was minimal or absent, and mRNA species for the remaining C-type and other lectins were mostly maintained at stable levels (Supplemental Figure 4 and Supplemental Table 4).

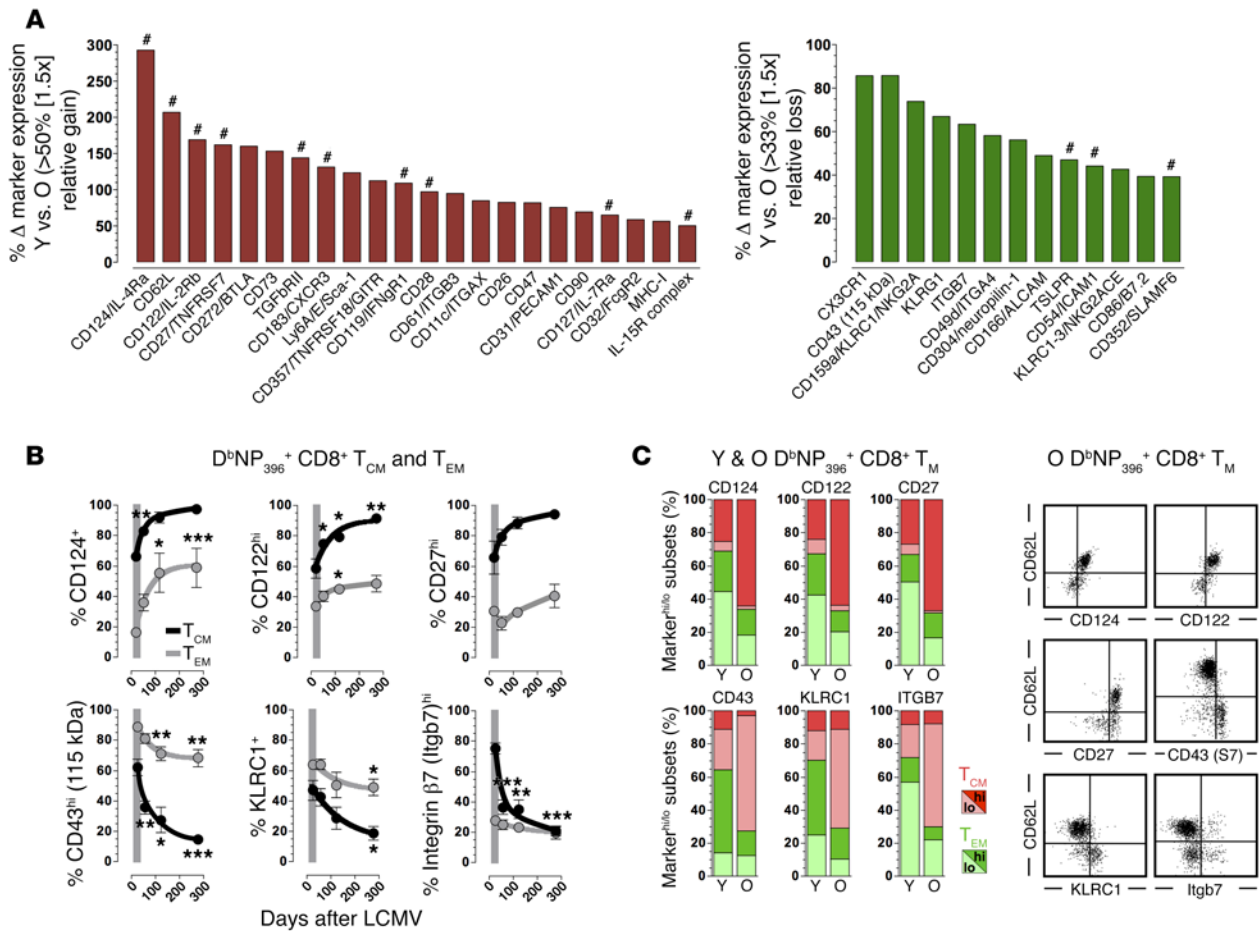


Figure 5. Gradual phenotypic conversion of aging antiviral CD8⁺ T_M populations. (A) Ranking of major D^bNP₃₉₆⁺ CD8⁺ T_M-expressed surface ligands/receptors according to differential expression between young (6–7 weeks) and old (>80 weeks) populations; primary expression data for all markers are found in Figure 4 and Supplemental Figures 2 and 4–14. Inclusion criteria for present ranking: differential expression observed in > 25% of D^bNP₃₉₆⁺ CD8⁺ T_M as well as > 1.5-fold up- or downregulation; #, molecular pathways interrogated in related work (JE, BD, and DH, unpublished observation). (B) Phenotypic conversion of blood-borne D^bNP₃₉₆⁺ CD8⁺ T_M stratified according to T_{EM} (CD62L^{lo}) and T_{CM} (CD62L^{hi}) subsets. (C) Relative distribution of young (days 52–58) vs. old (days 570–578) D^bNP₃₉₆⁺ CD8⁺ T_M phenotypes (marker^{hi}, dark red or green; marker^{lo}, light red or green) in relation to T_{CM} (red) and T_{EM} (green) subsets; dot plots gated on aged D^bNP₃₉₆⁺ CD8⁺ T_M; data generated with $n \geq 3$ mice/group in multiple independent experiments. * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$ by 1 way ANOVA. Y, young; O, old.

Molecular and phenotypic remodeling of aging antiviral CD8⁺ T_M; beyond the T_{EM}/T_{CM} paradigm. Using a similar approach to gene selection and ontology as employed above, we allocated ~25% of DEGs in Supplemental Table 1 to 30 structural or functional families that were subsequently investigated for the presence and temporal regulation of all known family members. With an emphasis on surface receptors/ligands, survival/apoptosis, effector activities, and signaling, we analyzed ~1,570 genes, ~520 of which were expressed by aging p14⁺ T_M. In the course of the extended memory phase (d46–d400), more than one third of expressed genes exhibited a significant increase or decrease, and for greater than 80% of these DEGs the changes constituted a continuation of dynamic adaptations already discernible in the transition from effector memory (T_{EM}) to early memory stage (Figure 4, Supplemental Table 4, and Supplemental Figure 4–14). The pronounced molecular reconfiguration evident during the relatively brief contraction phase of p14⁺ T_E largely confirmed previous observations (31); similarly, the slower transcriptomic

alterations accrued by aging p14⁺ T_M corresponded remarkably well to the molecular differences very recently reported for early and late p14⁺ central memory T cells (p14⁺ T_{CM}) (17). However, our observation that the dynamic signatures of aging p14⁺ T_M exhibited a prominent and slowly fading echo of these changes indicates that the continued molecular remodeling of p14⁺ T_M is a fundamental property associated with CD8⁺ T_M longevity.

To correlate the molecular evolution of p14⁺ T_M with temporal alterations at the protein level, we evaluated expression of ~140 cell surface antigens by D^bNP₃₉₆⁺ CD8⁺ T_M in aging LCMV-immune B6 mice (Figure 4, Supplemental Table 4, and Supplemental Figures 2 and 4–14); note that these analyses also included antigens not differentially expressed at the mRNA level, and were expanded in some cases to other LCMV-specific CD8⁺ T_M subsets (not shown). Although expression levels of the various phenotypic markers expectedly differed substantially, ~80% of all antigens analyzed were present on specific CD8⁺ T_M subsets, and of these, ~65% were progressively increased or decreased in a statistically signif-

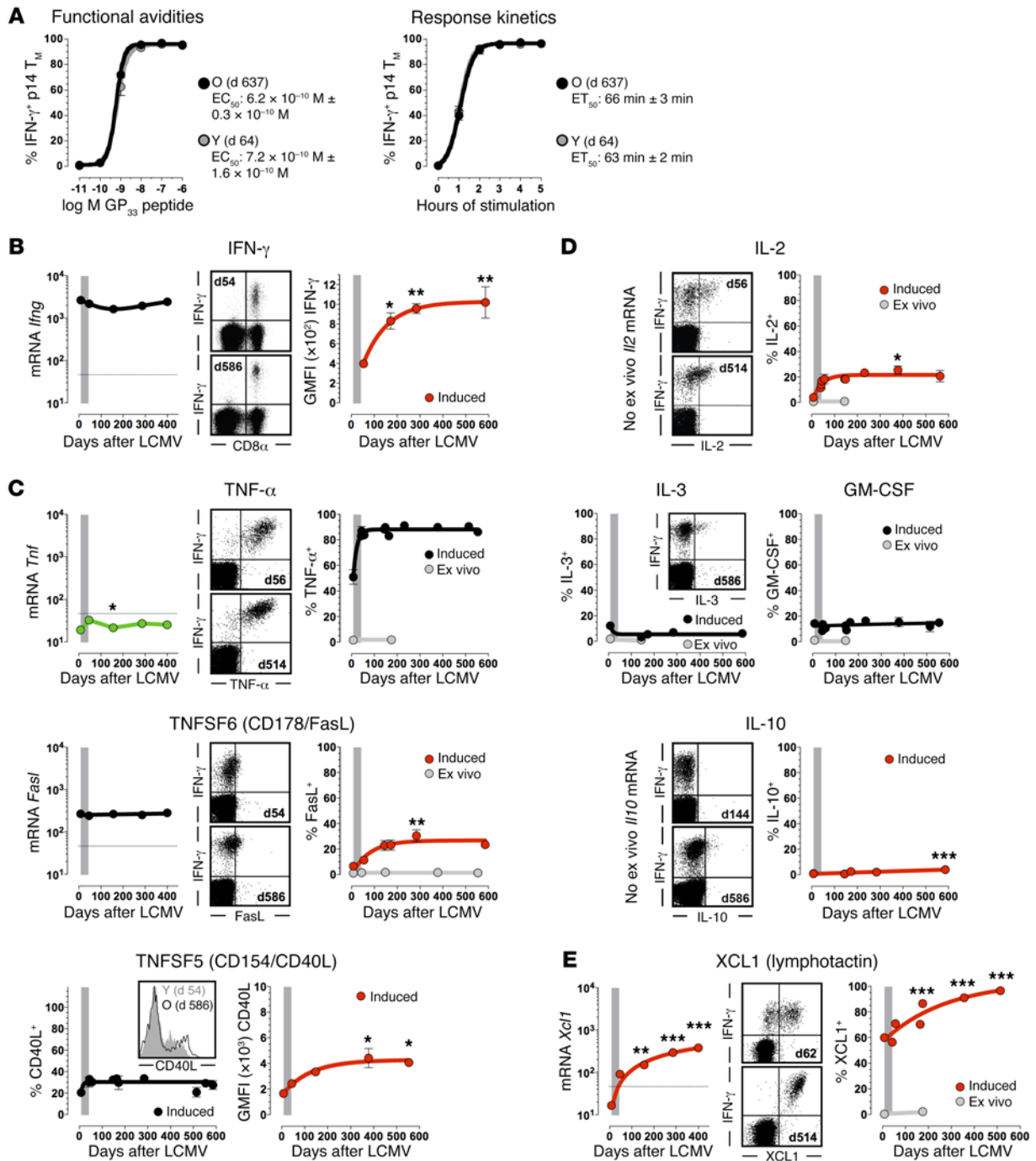


Figure 6. Evolving functionalities of aging antiviral CD8⁺ T_M: cytokines, TNFSFs, chemokines. (A) Functional avidities (EC₅₀ values) and response kinetics (ET₅₀ values) were calculated by plotting the fraction of IFN- γ ⁺ p14 T_M as a function of GP₃₃ peptide concentration (5-hour stimulation) or stimulation time (10⁻⁶ M peptide) and nonlinear regression analysis. (B–E) Temporal regulation of IFN- γ , TNFSF, cytokine, and chemokine mRNA (p14 T_{E/M} microarray data, ex vivo) as well as protein expression (NP₃₉₆-specific CD8⁺ T_{E/M} ex vivo [gray] and induced [black/red]); panels are organized in modules as detailed in Figure 4, dot plots are gated on NP₃₉₆ peptide-stimulated CD8⁺ T cells. **P* < 0.05, ***P* < 0.01, and ****P* < 0.001 by 1-way ANOVA and asterisks indicate statistical differences between young (~days 42–65) and older CD8⁺ T_M. Note that the amount of induced IFN- γ and CD40L content (geometric mean fluorescence intensity values) rather than percentage of IFN- γ - or CD40L-producing CD8⁺ T_M increased with age; data generated with *n* ≥ 3 mice/group in multiple independent experiments.

icant manner. Adding another ~10% of cell surface antigens that exhibited discernible yet nonsignificant trends towards altered expression levels, we conclude that ~75% of phenotypic CD8⁺ T_M markers, many of which have been implicated in the active regula-

tion of immune responses (32), are subject to continued modulation during the long-term maintenance of T cell memory (a rank-ordered listing of major differentially expressed surface receptors/ligands is displayed in Figure 5A). Of further importance is

the apparent translational regulation captured in our phenotypic analyses: for several surface receptors/ligands subject to particularly pronounced expression changes in aging CD8⁺ T_M (Figure 5A), corresponding mRNA levels did not significantly change over time (TGFβ-RII [Supplemental Figure 6], glucocorticoid-induced TNFR family-related gene [GTR; Supplemental Figure 7], CD28 [Supplemental Figure 8], CD31, CD32, CD47 [Supplemental Figure 10, A and B], CD49d, CD61, integrin β, [Supplemental Figure 11], CXCR3 [Supplemental Figure 12A], CD26, and Ly6A [Supplemental Figure 13, A and B]); in an unusual case we even found divergent gene and protein regulation (CD11c; Supplemental Figure 11). These observations underscore the importance of complementary molecular and phenotypic analyses for an inclusive characterization of CD8⁺ T_M aging.

Collectively, our findings suggest that phenotypic conversion rather than stability constitutes a hallmark of specific CD8⁺ T cell memory, and also may provide an adjusted perspective on the popular T_{EM}/T_{CM} paradigm (33): our longitudinal analyses of respective CD62L^{lo}CD62L^{hi} subsets (Figure 5B) corroborate the observation that both populations accrue phenotypic changes over time and that in turn align with their respectively increasing recall potentials (14, 17). As a net result of these interrelated maturation dynamics, T_{CM} exhibiting cardinal properties of aged CD8⁺ T_M (e.g., CD124⁺CD122^{hi}CD27^{hi}CD43^{lo}KLRC1 integrin β₇^{lo}) eventually come to dominate the pool of aging LCMV-specific CD8⁺ T_M (Figure 5C). Here, the expanded context of dynamic expression patterns largely (albeit not completely) shared by CD62L and multiple other surface receptors/ligands supports the general utility of T_{EM}/T_{CM} distinction, but also reveals the essentially arbitrary and limiting nature of CD62L expression levels as denominators for major CD8⁺ T_M subsets.

Evolving functionalities of aging antiviral CD8⁺ T_M. The expedient elaboration of various effector activities by specific CD8⁺ T_M constitutes a cardinal component of immune protection, yet the extent to which age-associated adaptations may alter kinetics, quantity, and/or quality of CD8⁺ T_M functions remain incompletely defined. Using induced IFN-γ production as a principal readout, young and old p14⁺ T_M exhibited identical functional avidities and response kinetics and thus a complete preservation of determinants that shape the rapid CD8⁺ T_M response to cognate antigens (Figure 6A). The amount of CD8⁺ T_M-produced IFN-γ, however, increased with age, suggesting that older CD8⁺ T_M may become more potent I^o CD8⁺ T_E (Figure 6B). To assess this possibility at large, we first defined the spectrum of young CD8⁺ T_M activities by quantifying constitutive and TCR-induced expression of ~200 p14⁺ T_M genes comprising perforin/granzymes, interferons, cytokines/chemokines, TNF superfamily (TNFSF), TGF-βSF, PDGF, and MMP members (Supplemental Figure 15 and data not shown). Then, for ~90% of transcripts demonstrating a significant TCR-induced upregulation, we analyzed corresponding protein expression in aging LCMV-immune B6 mice. Overall, we observed a strict conservation of CD8⁺ T_M functionalities (inducible TNF-α, GM-CSF, IL-3, and chemokine [CCL1, -3, -4, -5] production) in the absence of obvious immune deviation by aged CD8⁺ T_M (no acquisition of rapid IL-4, IL-13, IL-17A, IL-21, TGF-β, or TNFSF1b, -3, -7, -8, -9, -10, -11 production capacity; Figure 6, C and D, Supplemental Table 4, and data not shown).

Five additional effector functions, however, improved over time, including a growing proportion of CD8⁺ T_M capable of induced FasL expression, a modest increase of IL-2 production potential as shown previously (12, 17), and a more pronounced augmentation of inducible XCL1 chemokine synthesis (Figure 6, C-E). At the single-cell level, aging CD8⁺ T_M also made more CD40L and a very small subset acquired the capacity to produce IL-10 (Figure 6, C and D), perhaps through enhanced IL-27 responsiveness as a consequence of their increasing CD130 (also known as IL-6 signal transducer) expression (Supplemental Figure 6A); the limited IL-10 production capacity of aged CD8⁺ T_M, however, was lost upon rechallenge since all blood-borne and splenic I^o CD8⁺ T_E failed to synthesize IL-10 (data not shown). In summary, aging of CD8⁺ T_M confers enhanced functionality, and although autocrine IL-2 production can promote I^o CD8⁺ T_E expansions (34), it apparently combines with other functional gains to advance more effective pathogen control (JE, BD, and DH, unpublished observations).

Lastly, granule exocytosis is central to the cytolytic activity of CD8⁺ T_E and presumably also CD8⁺ T_M, although diverging temporal expression patterns of components within this pathway and limited knowledge about their precise contribution to in vivo CD8⁺ T_M killing (35) complicate any predictions about evolving CD8⁺ T_M cytotoxic T lymphocyte (CTL) functions. For example, several relevant mRNA species are downregulated in aging p14⁺ T_M (Supplemental Figure 15, A and B) and the precipitous decline of *Gzma* message constitutes the most pronounced molecular difference between young and old p14⁺ T_M (Table 1 and Figure 7A). Corresponding protein expression analyses demonstrated ex vivo-detectable GZMA in a major CD8⁺ T_E subset but also revealed, in contrast with the rapid downregulation of GZMB (12, 24), an attenuated decline that resulted in complete GZMA loss only ~1 year after infection (Figure 7A), a pattern similar to that of human CD8⁺ T_M analyzed after smallpox vaccination (36). Further evidence for a possible reduction of cytotoxic CD8⁺ T_M function may be derived from the marked decline of CD8⁺ T_M-expressed CX3CR1 (Table 1 and Figure 5A), a chemokine receptor that phenotypically (though not functionally) correlates with CD8⁺ T_M CTL potential (37). On the other hand, we noted a subtle trend among mRNA expression patterns shared by several genes associated with CD8⁺ T_{E/M} effector functions but not evident in other gene families: after a slight (*Ifng*, *Prfl*, *Tnf*) or more prominent (*Gzmb/k/m*, *Ccl3/4*, *S1pr5*) decline of message that lasted ~5 months, mRNA levels began to steadily rise again (Figures 6B and 7A, Supplemental Figure 15A, Supplemental Figure 12B, and data not shown). In addition, aging CD8⁺ T_M exhibited a small increase in degranulation capacity (Figure 7B).

To ascertain how these expression patterns are integrated into actual killing capacity, we conducted an in vivo CTL assay (35) with young and old LCMV-immune mice. As shown in Figure 7C, young CD8⁺ T_M exhibited faster and more pronounced early CTL activity, but by 6 hours both young and old CD8⁺ T_M eliminated the majority (≥80%) of target cells in the spleen. In a modified assay employing AT of both CD8⁺ T_M and targets into naive recipients to achieve an in vivo effector:target (E:T) ratio of 1:1 (38), CTL functions of young and old CD8⁺ T_M proved indistinguishable (Figure 7D). Therefore, aging CD8⁺ T_M, in spite of decreasing killing kinetics, largely preserve their capacity for effective target cell destruc-

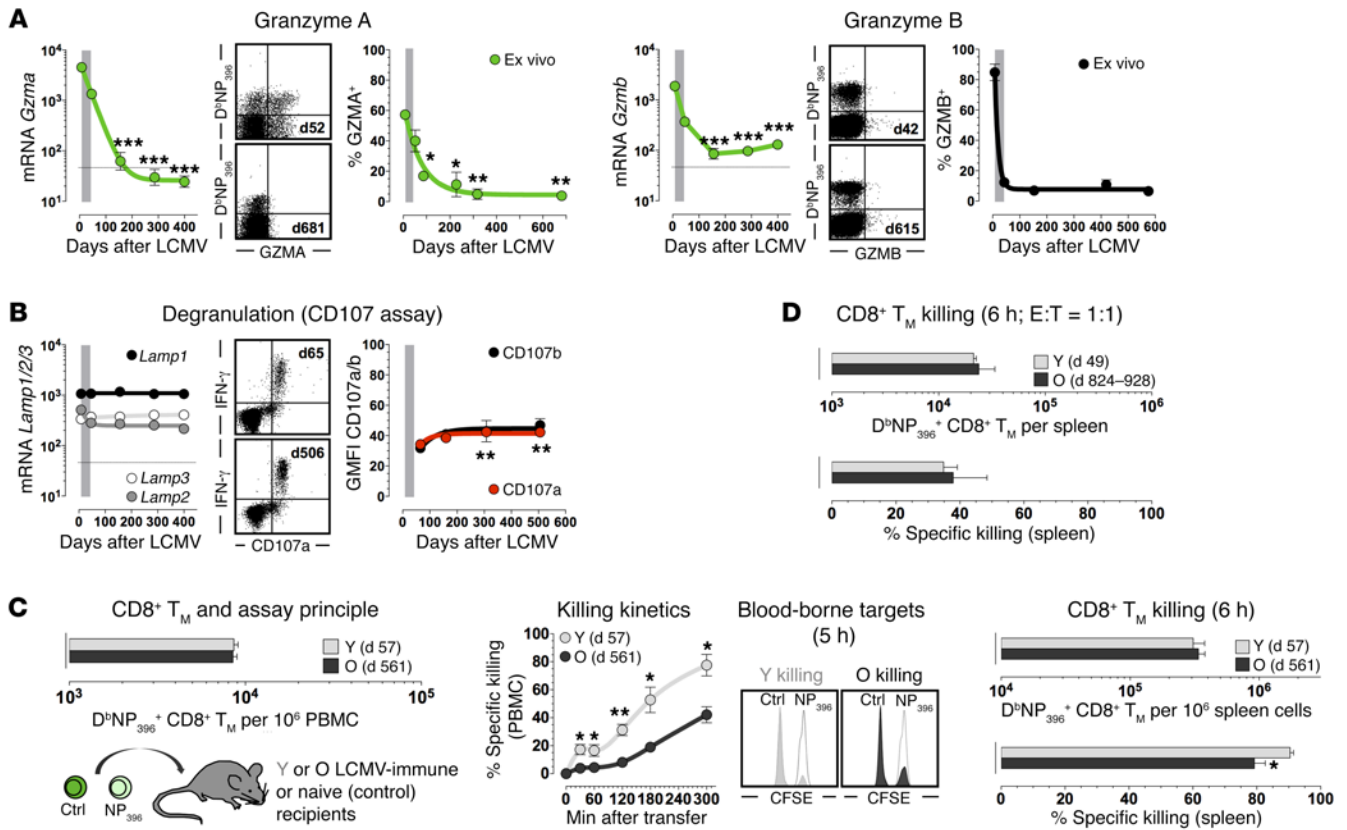


Figure 7. Evolving functionalities of aging antiviral CD8⁺ T_M: granzymes, degranulation, and CTL activity. (A) Temporal regulation of granzyme mRNA (p14⁺ T_{E/M} microarray data) and protein (D^bNP₃₉₆ + CD8⁺ T_{E/M}) expression analyzed directly ex vivo. (B) *Lamp1*, -2, and -3 mRNA expression and degranulation capacity of aging p14⁺ T_{E/M} and NP₃₉₆-specific CD8⁺ T_M respectively. (C) Left, young (Y) and old (O) LCMV-immune mice harboring identical numbers of blood-borne CD8⁺ T_M were used to assess CD8⁺ T_M CTL activities in vivo; middle, killing kinetics determined in blood after AT of 4.5 × 10⁶ differentially CFSE-labeled control and NP₃₉₆ peptide-coated target cells each (histograms gated on target cells retrieved 5 hours after AT into LCMV-immune [young, gray; old, black] or naive [open] mice); right, splenic D^bNP₃₉₆ + CD8⁺ T_M numbers at conclusion of assay and extent of specific killing. (D) D^bNP₃₉₆ + CD8⁺ T_M abundance and killing activity in spleen after AT of 4.5 × 10⁵ purified young or old D^bNP₃₉₆ + CD8⁺ T_M into naive recipients followed by transfer of 4.5 × 10⁵ sensitized and control target cells each; data generated with n ≥ 3 mice/group in multiple independent experiments. *P < 0.05, **P < 0.01, and ***P < 0.001 by 1-way ANOVA or Student's *t* test. PBMC, peripheral blood mononuclear cells.

tion. We also note that target killing by CD8⁺ T_M appears to proceed according to a sigmoidal association (Figure 7C) rather than the exponential association observed for CD8⁺ T_E (35), suggesting that CD8⁺ T_M CTLs may be less reliant on immediate cytotoxicity as conferred, for example, by perforin release from CD8⁺ T_E (38).

Accelerating the maturation process of CD8⁺ T_M: p14-titration chimeras. A notable feature of the extended CD8⁺ T_M conversion process is its prolonged yet asynchronous nature (Figure 4, Supplemental Figure 2, and Supplemental Figures 4-14); i.e., the pace and extent of conversion is distinct for different gene products and thus likely contributes to the great phenotypic variety among CD8⁺ T_M populations (1). The origins of phenotypic heterogeneity are rooted in the earliest stages of the T cell response, since diverse CD8⁺ T_E populations can be derived from a single naive CD8⁺ T cell, possibly through asymmetric division but certainly as a consequence of distinct priming interactions and division histories (39). That the degree of pathogen replication, accompanying inflammation, and effective pathogen elimination decisively shape CD8⁺ T_E differentiation is further documented in multiple studies employing chimeras generated with titrated numbers of TCRtg CD8⁺ T_N as well as their transfer at different times after

pathogen challenge and/or experimental reduction of attendant inflammation or antigen presentation. Importantly, an increase of TCRtg CD8⁺ T_N input numbers and/or decrease of inflammation/antigen presentation can accelerate CD8⁺ T_M generation as determined by expansion/contraction kinetics, CD8⁺ T_M cellularity, and/or limited phenotypic and functional analyses (2, 40-43).

Here, we revisited these concepts with an expanded focus on the kinetics of progressive CD8⁺ T_M maturation and associated in vivo functionalities as a consequence of their early ontogeny; i.e., we constructed p14 chimeras with graded numbers of p14⁺ T_N (2 × 10² to 2 × 10⁵, referred to as p14^{2e2}-p14^{2e5} chimeras) to determine the relative speed with which subsequently established p14⁺ T_M populations would acquire properties resembling endogenously generated aged CD8⁺ T_M. In most tissues, initial numerical expansion of I^o p14⁺ T_E does not occur until ~72 hours after LCMV challenge, but proliferation starts somewhat earlier in the MedLNs (19, 21). Enumerating p14⁺ T_E in the MedLNs on day 3 or 4 after infection, we found that their expansions in p14^{2e2}-p14^{2e5} chimeras tracked exactly with original p14⁺ T_N input numbers, indicating that the early kinetics of p14⁺ T_E population growth are largely impervious to p14⁺ T_N precursor number (Fig-

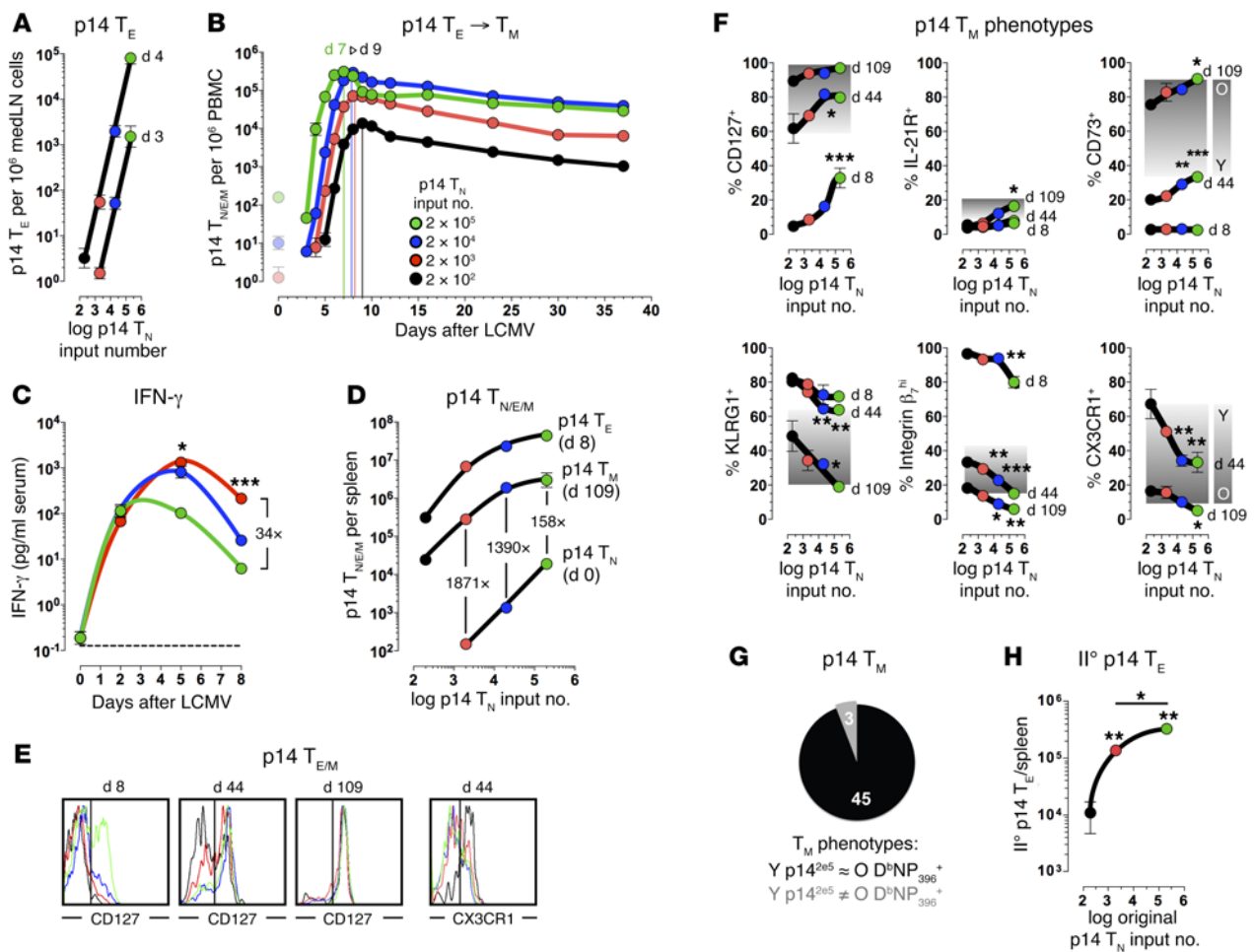


Figure 8. Accelerating the maturation process of CD8⁺ T_M: p14-titration chimeras. (A) Chimeras were constructed with graded p14⁺ T_N numbers (2 × 10² to 2 × 10⁵; color legend in panel B), and p14⁺ T_E expansions were quantified in the MedLNs on days 3–4 (connected data points, respectively). (B) Kinetics of p14⁺ T_E expansion/contraction and p14⁺ T_M development in blood (5 combined experiments; n = 3–6); data for day 0 indicate p14⁺ T_N retrieved in the absence of infection, and vertical colored lines indicate respective p14⁺ T_E peak expansions. (C) Serum IFN-γ as a function of p14⁺ T_N input number. (D) Absolute numbers of transferred p14⁺ T_N (day 0), T_E (day 8) and T_M (day 109) in spleen; values indicate fold difference between p14⁺ T_N and T_M cellularity. (E) CD127 and CX3CR1 expression by splenic p14⁺ T_{E/M} recovered from respective chimeras on days 8, 44, and 109. (F) Kinetics of phenotypic p14⁺ T_{E/M} differentiation as function of p14⁺ T_N precursor frequency. Analyses of splenic p14⁺ T_E (day 8) and T_M (days 44 and 109) were conducted as above; data points from p14⁺ T_{E/M} analyzed at the same time point are connected by a line, and statistics compare p14^{2e2} vs. p14^{2e3}-p14^{2e5} chimeras. For comparative purposes, the graded background shading demarcates the spread of respective marker expression from young (Y, light) to old (O, dark) D^bNP₃₉₆⁺CD8⁺ T_M in B6 mice (Figure 4 and Supplemental Figures 5–14). (G) Summary of preceding analyses comprising 48 markers categorized according to similarity between p14⁺ T_M from young p14^{2e5} chimeras (~7 weeks) and old D^bNP₃₉₆⁺CD8⁺ T_M (≥80 weeks). (H) AT/rechallenge experiments were performed with 2 × 10³ p14⁺ T_M purified from respective p14-titration chimeras infected ~7 weeks earlier, AT, LCMV Armstrong challenge and quantification of II° p14⁺ T_E expansions 8 days later (n = 3; representative of 3 similar experiments). *P < 0.05, **P < 0.01, and ***P < 0.001 by 1-way ANOVA or Student's t test.

ure 8A). We next monitored the dynamics of the p14⁺ T_E response in peripheral blood. Following LCMV challenge and an initial drop of blood-borne p14⁺ T cells similar to that reported for other tissues (19) (and also observed for aged II° p14⁺ T_E in peripheral blood; Figure 1F), p14⁺ T_E started to expand after more than 72 hours and, consistent with the early p14⁺ T_E expansion kinetics in the MedLNs, further analysis of population dynamics throughout the I° response revealed no significant differences among respective population growth rates (Figure 8B). As expected (2, 40), an increase of p14⁺ T_N numbers both enhanced and expedited the peak expansions of p14⁺ T_E (from d9 in p14^{2e2} to d7 in p14^{2e5} chimeras) (Figure 8B), but we note that a 1,000-fold difference in p14⁺ T_N input numbers produced an only 23-fold difference between

respective p14⁺ T_E burst sizes on d7–d9 (Figure 8B). Thus, higher numbers of naive precursors restrain the I° response by earlier termination of proliferative CD8⁺ T_E expansion and corresponding curbing of the relative burst size.

Nevertheless, greater p14⁺ T_E expansions in the higher input chimeras exerted better inflammation control at later stages (d5–d8) of the effector response (Figure 8C), and brought about a correspondingly larger p14⁺ T_M population (Figure 8D). A comparison of p14^{2e2} versus p14^{2e5} chimeras, for example, showed that the relative difference between splenic p14⁺ T_E cellularities (d8, 140-fold) was largely maintained in the memory phase (d109, 123-fold) (Figure 8D), but these calculations do not take into account the precise timing of p14⁺ T_E peak expansions. We therefore compared the numbers of

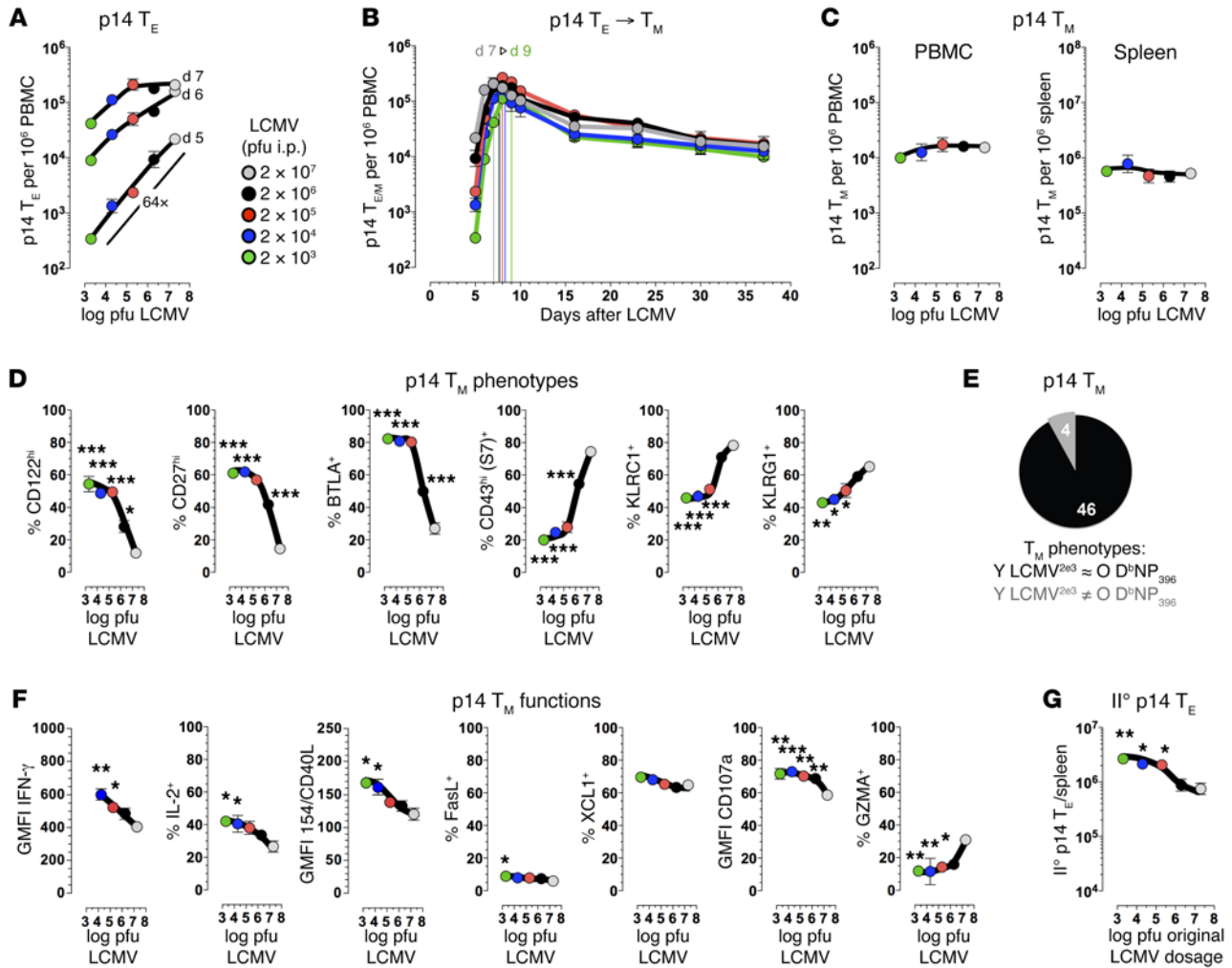


Figure 9. Delaying the maturation process of CD8⁺ T_M; virus-titration chimeras. (A) Virus-titration chimeras were constructed with 10⁴ p14⁺ T_N, challenged with graded dosages of LCMV Armstrong (2 × 10³ to 2 × 10⁷ PFU), and early p14⁺ T_E expansions were quantified in blood (days 5–7); note the 64-fold difference (P = 0.006) between low-dose (LCMV^{2e3}) and high-dose (LCMV^{2e7}) virus-titration chimeras on day 5. (B) Kinetics of p14⁺ T_E and T_M development in peripheral blood of virus-titration chimeras; vertical colored lines indicate time of respective peak expansions. (C) p14⁺ T_M numbers in blood (day 37) and spleen (day 42). (D) Relative kinetics of phenotypic p14⁺ T_M differentiation as a function of virus challenge dosage were determined on days 42–49; data are displayed as in Figure 8F, and statistics compare p14⁺ T_M properties of high-dose (2 × 10⁷ PFU) vs. lower dose (≤2 × 10⁶ PFU) virus-titration chimeras. (E) Pie chart summary of preceding analyses comprising 50 markers stratified according to similarity between young p14⁺ T_M recovered from LCMV^{2e3} virus-titration chimeras (6–7 weeks old) and old D^hNP₃₉₆-CD8⁺ T_M (≥80 weeks old). (F) Functional properties of p14⁺ T_M (day 42) were determined after a 5-hour peptide stimulation (IFN-γ, IL-2, CD40L, FasL, XCL1, CD107a), or directly ex vivo (GZMA). (G) AT/rechallenge experiments were performed with 2 × 10³ p14⁺ T_M purified from virus-titration chimeras (day 42) and Il^o p14⁺ T_E expansion analyses on day 8. All summary data obtained with n ≥ 3 mice/group in multiple independent experiments. *P < 0.05, **P < 0.01, and ***P < 0.001 by 1-way ANOVA. PBMC, peripheral blood mononuclear cells.

splenic p14⁺ T_N retrieved in the absence of infection with p14⁺ T_M (d109) and calculated that p14^{2e5} chimeras generated a p14⁺ T_M pool that was 158-fold greater than the starting population; p14^{2e3} chimeras, in contrast, produced a 1,871-fold bigger p14⁺ T_M compartment (Figure 8D). We conclude that the relative extent to which clonal expansion populates the subsequent CD8⁺ T_M compartment is curtailed by higher CD8⁺ T_N frequencies, likely through restricted CD8⁺ T_E differentiation that favors accelerated CD8⁺ T_M development (2).

To assess the extent of phenotypic CD8⁺ T_M alterations acquired as a function of increasing p14⁺ T_N input numbers, we first evaluated the regulation of IL-7Ra/CD127 expression. As shown previously (2), re-expression of CD127 by p14⁺ T_E and T_M proceeded with kinetics proportional to the numbers of available

p14⁺ T_N, but by ~3 months after challenge, practically all p14⁺ T_M, irrespective of p14⁺ T_N precursor frequency, expressed high levels of CD127; conversely, downmodulation of p14⁺ T_M-expressed CX3CR1 was progressively delayed by limiting p14⁺ T_N input numbers (Figure 8E). Although a similar experimental modulation of CD8⁺ T_M maturation kinetics has been demonstrated for additional surface receptors/ligands (2, 17), the actual extent of phenotypic alterations in this experimental scenario remains unknown. Our comprehensive characterization of p14^{2e2}-p14^{2e5} chimeras now reveals a striking correlation between p14⁺ T_N precursor frequencies and multiple dozens of p14⁺ T_M phenotypes: for ~94% of markers analyzed (45 of 48), an increase of p14⁺ T_N accelerated the conversion towards an aged CD8⁺ T_M phenotype, a pattern that

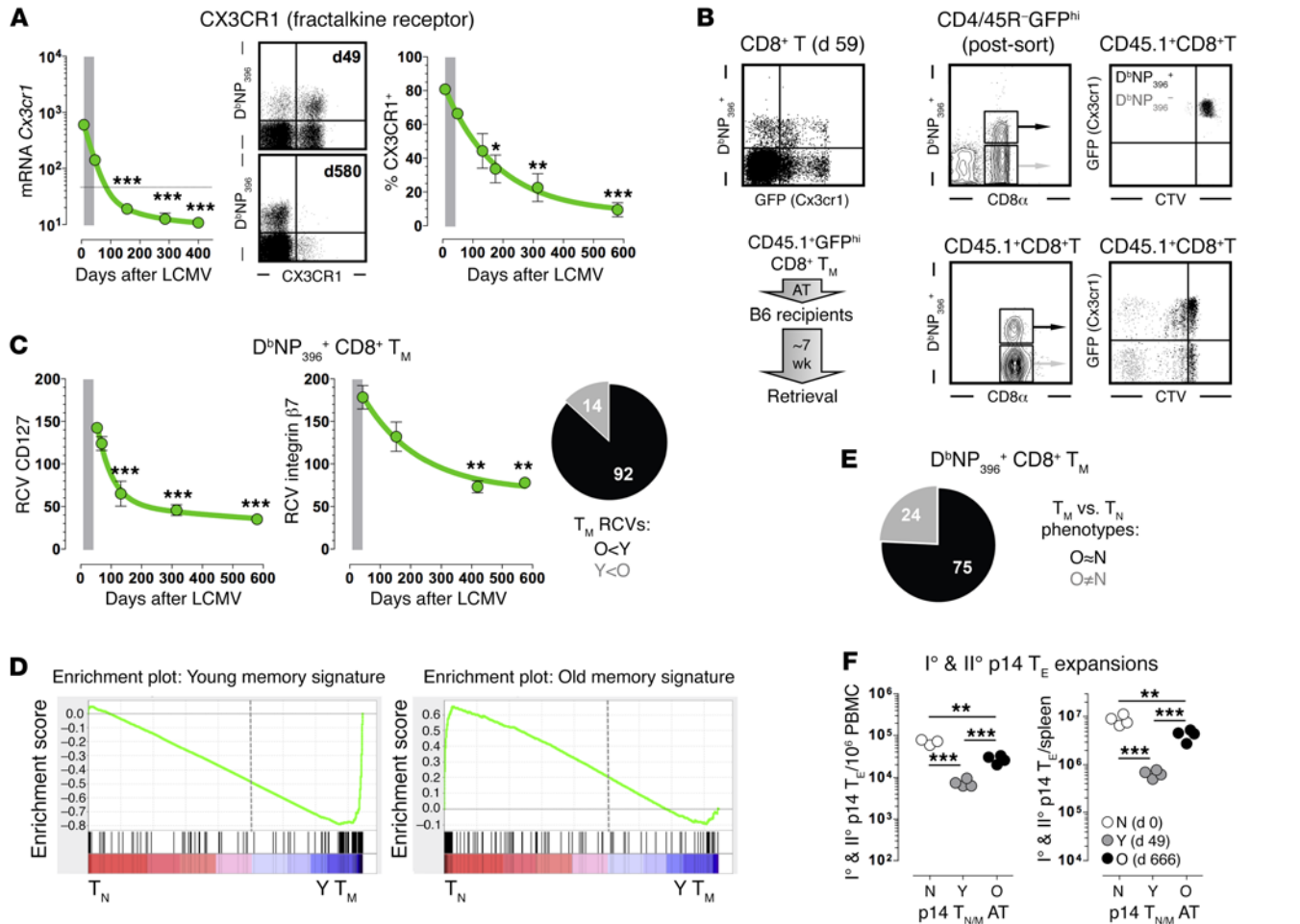


Figure 10. Progressive conversion of aging virus-specific CD8⁺ T_M towards homogeneity and naiveté. (A) Temporal regulation of *Cx3cr1* mRNA and protein expression. (B) Top, splenic CD8⁺ T cells from LCMV-immune *Cx3cr1*^{+/Gfp} mice were analyzed for GFP expression, or loaded with CellTrace Violet (see Supplemental Methods for more information) and sorted according to GFP^{hi} status; bottom, experimental outline and properties of retrieved CD8⁺ T cells. (C) Progressive acquisition of more homogeneous CD8⁺ T_M phenotypes as determined by declining RCV values; the pie chart summarizes RCV analyses of 106 markers expressed by young (6–7 weeks old) and old (>80 weeks old) D^bNP₃₉₆⁺CD8⁺ T_M. O, old; Y, young (D) The top 180 genes overexpressed by young or old p14⁺ T_M (signatures) were investigated by GSEA using datasets for polyclonal CD8⁺ T_N (T_N) and LCMV-GP₃₃-specific CD8⁺ T_M (day 30 [Y T_M]) (see Supplemental Figure 18); note respective enrichments of young p14⁺ T_M signatures in Y T_M and old p14⁺ T_M in T_N ($P < 0.0001$). (E) Expression of 99 markers was compared between aging D^bNP₃₉₆⁺CD8⁺ T_M and CD8⁺ T_N and categorized according to similarity (O≈N, conversion towards CD8⁺ T_M phenotypes; O≠N, contrasting expression). (F) AT/rechallenge experiments conducted with 2×10^3 purified p14⁺ T_N or young/old T_M, and analyzed for I^o (N) and II^o (Y vs. O) p14⁺ T_E expansions in blood (day 7.5) and spleen (day 8). * $P < 0.05$, ** $P < 0.01$, and *** $P < 0.001$ by 1-way ANOVA. PBMC, peripheral blood mononuclear cells.

often became discernible already at the T_E stage (Figure 8, F and G). Perhaps most importantly, the relative preponderance of aged p14⁺ T_M phenotypes in the higher input chimeras also correlated with improved II^o p14⁺ T_E reactivities (Figure 8H).

Delaying the maturation process of CD8⁺ T_M: virus-titration chimeras. If the above contentions are correct, CD8⁺ T_M maturation at large should also be amenable to modulation under conditions of fixed CD8⁺ T_N precursor frequencies, i.e., by experimental variation of the viral burden and degree of inflammation. We therefore challenged p14 chimeras harboring identical numbers of p14⁺ T_N with graded dosages of LCMV ranging across 4 orders of magnitude. Interestingly, the kinetics of I^o p14⁺ T_E expansion and p14⁺ T_M generation in these virus-titration chimeras were distinctly different from those in the p14-titration chimeras: an escalation of the virus challenge dose accelerated the timing of p14⁺ T_E peak expansions (2×10^3 PFU on d9 vs. 2×10^7 PFU on d7), yet neither

respective burst sizes nor contraction kinetics or p14⁺ T_M numbers were greatly affected by the different infection protocols (Figure 9, A–C). At the same time, the phenotypic disparity of p14⁺ T_M populations in the virus-titration chimeras was pronounced: for 92% of cell surface receptors/ligands analyzed (46 of 50), high-dose challenge readily delayed the acquisition of aged CD8⁺ T_M phenotypes (Figure 9, D and E; expression patterns of 49 individual markers are shown in Supplemental Figure 16). Thus, the population growth rates of I^o CD8⁺ T_E directly determine the kinetics of subsequent phenotypic CD8⁺ T_M maturation.

This conclusion is further supported by an analysis of p14⁺ T_M effector activities. An increase of LCMV inoculation dosage depressed inducible p14⁺ T_M functions (IFN- γ , IL-2, CD40L, and degranulation) and also delayed the downmodulation of GZMA (the impact on restrained FasL and XCL1 synthesis was less pronounced, although it followed the same trend), and p14⁺ T_M popu-

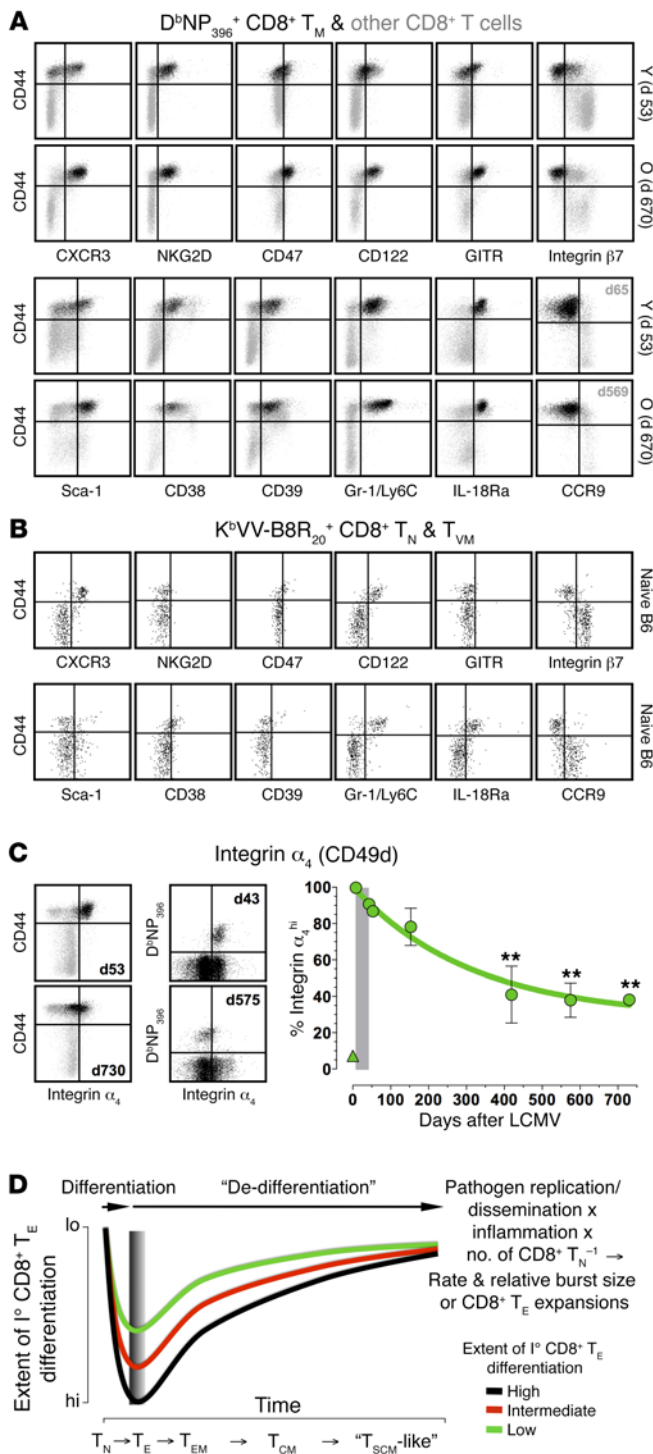


Figure 11. Memory markers old and new: the “rebound model” of T_M maturation. (A) Phenotypic properties of young and old splenic CD8⁺ T_M (CCR9⁻; PBMC d65/d569); dot plots distinguish D^bNP₃₉₆⁺CD8⁺ T_M (black) and other CD8⁺ T cells (gray). (B) Phenotypic properties of virus-specific CD8⁺ T_N enriched from naive B6 mice; dot plots are gated on K^bVV-B8R₂₀⁺CD8⁺ T_N with CD44 expression distinguishing genuine CD44^{lo}CD8⁺ T_N and CD44^{hi}CD8⁺ T_{VM}. (C) Kinetics of integrin α₄/CD49d expression by D^bNP₃₉₆⁺CD8⁺ T_{E/M}. Dot plots are gated on CD8⁺ T cells with left-hand plots distinguishing D^bNP₃₉₆⁺CD8⁺ T_M (black) from other CD8⁺ T cells (gray); triangle symbol: CD44^{lo}CD8⁺ T_N. (D) The “rebound model” of T_M maturation: kinetics of progressive conversion towards a mature and naive-like CD8⁺ T_M phenotype (dedifferentiation) are accelerated or delayed according to the extent of initial T_E differentiation (integrated metric for the product of pathogen replication/dissemination, inflammation and reciprocal CD8⁺ T_N precursor frequency that ultimately determines rate and relative burst size of CD8⁺ T_E expansions). Colored curves refer to CD8⁺ T_{E/M} populations that proceed through lesser (green), intermediate (red), or advanced (black) stages of I^o T_E differentiation, and the gray-shaded area demarcates the variable timing of peak CD8⁺ T_E expansions according to experimental conditions (earlier, high pathogen challenge dosage/inflammation or CD8⁺ T_N precursor frequencies; later, reduced infection/inflammation or CD8⁺ T_N numbers). **P < 0.01 by 1-way ANOVA.

position CD8⁺ T_M to exert improved immune protection, but also impinge on the subsequent development of II^o CD8⁺ T cell memory (Figure 2, F and G). To further explore the impact of I^o CD8⁺ T_M age on II^o CD8⁺ T_M generation, we conducted contemporaneous analyses 2–3 months after performing original mixed AT/rechallenge experiments and found that II^o CD8⁺ T_M derived from old as opposed to young I^o CD8⁺ T_M exhibited more mature phenotypes and diversified functional capacities consistent with an overall accelerated reacquisition of aged CD8⁺ T_M features (Supplemental Figure 17, A–D); in addition, these properties directly corresponded with a greater recruitment of old II^o CD8⁺ T_M into a III^o response (Supplemental Figure 17, A and E). It is noteworthy that the precipitated II^o memory formation from aged I^o CD8⁺ T_M proceeds in the wake of larger II^o expansions (i.e., increased relative burst size) and therefore potentially more pronounced effector differentiation. However, enhanced II^o CD8⁺ T_E responses primarily emerge as a consequence of better II^o CD8⁺ T_E survival and not their faster or prolonged proliferation (Figure 1, D–G, and Supplemental Figure 1, E–G), suggesting that these cells may better resist advanced effector differentiation. Given that the distinctive properties accrued in the process of CD8⁺ T_M aging can be efficiently passed on to II^o CD8⁺ T_M generations, we conclude that the original priming history shapes both I^o and II^o CD8⁺ T cell memory formation as well as III^o reactivity.

Extended phenotypic and molecular CD8⁺ T_M conversion: towards homogeneity and naiveté. The changing phenotypic composition of the virus-specific CD8⁺ T_M compartment could emerge from the progressive molecular/phenotypic conversion of individual CD8⁺ T_M (4) or an outgrowth of CD8⁺ T_M subsets with a competitive advantage (41). To distinguish between these possibilities, we built on our observation that aging CD8⁺ T_M substantially downregulated transcription/translation of CX3CR1 (Figure 10A). Using a mouse model in which *Cx3cr1* is replaced with a GFP reporter (44), GFP^{hi}CD8⁺ T_M were enriched from LCMV-immune heterozygous *Cx3cr1*^{+/Gfp} mice, labeled with a proliferation dye, and transferred into naive recipients (Figure 10B). Within ~7 weeks after AT, ~15%–

lations with less mature phenotypes and decreased functionalities also exhibited lower II^o expansions in vivo (Figure 9, F and G). Taken together, the p14 titration and virus titration experiments demonstrate that CD8⁺ T_M maturation in general is precipitated by original priming conditions that restrain CD8⁺ T_E differentiation through paced or abortive proliferative expansion (low challenge dosage or high CD8⁺ T_N precursor frequency), reduced inflammation, and/or expedited pathogen control (not shown).

Differential relay of phenotypic and functional properties from I^o to II^o CD8⁺ T_M populations. Superior recall capacities not only

20% of $D^bNP_{396}^+CD8^+T_M$ lost GFP expression without exhibiting signs of proliferation (Figure 10B), providing evidence that specific $CD8^+T_M$ are subject to a direct molecular conversion process, a conclusion consistent with earlier observations about $p14^+$ or polyclonal $CD8^+T_M$ conversion at the level of CD62L re-expression (12, 40). Although the notions of $CD8^+T_M$ conversion and subset outgrowth are not mutually exclusive (40), the transformation of aging $CD8^+T_M$ phenotypes according to different rates and degrees (Figure 4, and Supplemental Figures 2 and 4-14) may be more effectively accounted for by the conversion model (experiments providing a more definitive answer to this question are in progress).

Despite preservation of some phenotypic heterogeneity even at very late memory stages, the long-term maintenance of $CD8^+T_M$ populations appeared to result in the acquisition of more homogeneous appearances (Figure 4 and Supplemental Figures 5-14), an observation also made by Martin et al., who found fewer phenotypic subsets identified by 5-parameter Boolean gating in late versus early $p14^+T_{CM}$ populations (17). To evaluate this notion in our model, we determined the robust coefficient of variation (RCV) of marker expression and found that nearly 90% of $CD8^+T_M$ -expressed surface antigens (92 of 106) exhibited lower RCV values in old as compared with young $CD8^+T_M$, demonstrating that aging indeed fosters increased phenotypic homogeneity (Figure 10C). At the molecular level, our definition of unique young and old $p14^+T_M$ signatures also permitted their investigation by gene set enrichment analyses using published $CD8^+$ T cell datasets comprising assorted specificities (TCRtg/tetramer $^+T_{MP}$), differentiation stages (T_N/T_E ; $I^o/II^o/III^oT_M$), subsets ($T_{EM}/T_{CM}/T_{SCM}$), ages (young/old), operational states (resting/activated/exhausted), and species (murine/human). Altogether, we conducted 28 pairwise comparisons that confirmed the expected similarity of young $p14^+T_M$ to various resting $CD8^+T_M$ subsets but also revealed a notable enrichment of old $p14^+T_M$ signatures in both murine and human naive $CD8^+T_N$ (Figure 10D, Supplemental Figure 18, and data not shown). This conclusion was further supported by corresponding phenotypic analyses: relating the conversion trajectories of individual $CD8^+T_M$ markers to respective expression patterns of $CD8^+T_N$, we found that aging $CD8^+T_M$ gradually adopted an ever more naive phenotype (Figure 10E).

Given the unexpected constellation of increasing age, improved functionalities, and mostly naive $CD8^+T_M$ phenotypes, we revisited the notion of differential I^o versus II^o reactivity by directly comparing the proliferative capacity of $CD8^+T_N$ with that of young and old $CD8^+T_M$. Using the AT of limiting $p14^+T_{N/M}$ numbers as an experimental tool to gauge their maximal proliferative potential, the II^o reactivity of old versus young $p14^+T_M$ was significantly enhanced as anticipated, but remarkably, even aged $p14^+T_M$ did not expand to the same extent as $p14^+T_N$ (Figure 10F). Thus, the evolution of aging $CD8^+T_M$ phenotypes and proliferative potential is marked by a progressive harmonization with $CD8^+T_N$ properties.

Differentiating $CD8^+T_M$ subsets: memory markers old and new. What then are the distinctive phenotypic properties of mature $CD8^+T_M$ and $CD8^+T_N$ populations? To some extent, the answer depends on the precise definition of T cell naiveté. Using the $CD44^{lo}$ status as a reference, the phenotypic conversion of aging $CD8^+T_M$ did not produce populations identical to the $CD44^{lo}CD8^+T_N$ subset. Indeed, the analyses summarized in Figure 10E define multiple

markers that distinguish mature $CD8^+T_M$ from $CD44^{lo}CD8^+T_N$, including several antigens not previously considered as memory markers (e.g., CD38, CD39, and CD47) (Supplemental Table 5 and Figure 11A). Alternatively, T cell naiveté can be defined as lack of prior exposure to specific antigens, and we reported earlier that up to 30% of the virus-specific $CD8^+$ T cell repertoire in noninfected wild-type mice is in fact $CD44^{hi}$, a $CD8^+T_N$ subset termed “virtual memory” cells ($CD8^+T_{VM}$) (45). As shown here, $CD8^+T_{VM}$ and T_{MP} exhibited largely shared memory marker signatures, and a comparison of virus-specific and bulk $CD44^{lo}CD8^+T_N$ also revealed essentially identical phenotypes (Figure 11, A and B). While $CD8^+T_{VM}$ and T_{MP} were somewhat similar to $CD44^{lo}CD8^+T_N$ (CD39 $^-$ Sca-1 lo NKG2D $^-$ GITR lo) or of a mixed phenotype (CD47 int CD38 int), they also exhibited key features of aged $CD8^+T_M$ (CXCR3 $^+CD122^{hi}$ integrin β_7^{lo} Gr-1/Ly6C hi IL-18Ra $^+CCR9^+$) (Figure 11, A and B). Furthermore, expression of integrin α_9 , widely used as a genuine memory marker and found at low levels on $CD8^+T_{VM}$ (45), was in fact gradually downmodulated by aging $CD8^+T_M$ (Figure 11C). Taking into account the residual phenotypic heterogeneity of aged $CD8^+T_M$ populations and the constitutive diversity of $CD8^+T_N$ comprising both $CD44^{lo}$ and $CD44^{hi}$ subsets, their overall similarity therefore appears even more striking.

$CD8^+T_M$ aging as a dedifferentiation process: the rebound model of T_M maturation. From the perspective of the extended and cumulative $CD8^+T_M$ maturation process, our results suggest a simple amendment to the decreasing potential model of T_M generation (1, 4): improved II^o $CD8^+T_M$ reactivity and associated protective capacity develop with a probability and at a pace inversely proportional to the extent of I^o $CD8^+T_E$ differentiation. This contention, based on a reevaluation of earlier observations (2, 40) in the expanded context of $CD8^+T_M$ aging developed here, is further supported by our in vivo titration experiments demonstrating that the $CD8^+T_M$ aging process at large can be accelerated or delayed by simply manipulating the conditions of initial $CD8^+T_E$ priming. Accordingly, the degree of I^o $CD8^+T_E$ differentiation, primarily a product of pathogen replication/dissemination, inflammation, and reciprocal $CD8^+T_N$ precursor/dissem frequency that together control the speed and relative extent of proliferative $CD8^+T_E$ expansion, directly determines the odds and kinetics with which, over a period of months to years, $CD8^+T_M$ and T_N properties are progressively aligned (Figure 11D).

The notion of shared features between $CD8^+T_M$ and T_N is also supported by global transcriptome analyses of murine and human $CD8^+$ T cell subsets that overall reveal a substantial degree of similarity (~95%) (46), and that position $CD8^+T_{MP}$, T_{CM} , or T_{SCM} populations, respectively, in increasing proximity to $CD8^+T_N$ (31, 47-49). And although $CD8^+T_M$ are traditionally regarded as more potent, recent studies have identified several advantageous properties of $CD8^+T_N$ in comparison with $CD8^+T_{CM}$ subsets, such as their reduced antigen threshold requirements (50), their greater capacity for generation of $CD8^+T_E$ with superior proliferative and antitumor potential (51), or the better expansion of OT-I $^+$ and $p14^+T_N$ in relation to various $CD8^+T_M$ populations (total $T_M/T_{EM}/T_{CM}$) following bacterial or LCMV cl13 challenge (16, 52). If $CD8^+T_{MP}$ in comparison with $CD8^+T_N$, therefore can be said to have traded potential (i.e., the entire range of possible features to be acquired within the context of a I^o response) for potency (i.e., the more limited spectrum

of actual CD8⁺ T_M properties), the transcriptional, phenotypic, and functional alterations accrued during their prolonged maturation not only suggest a partial reacquisition of the potential inherent to CD8⁺ T_N, but in fact a process of dedifferentiation we refer to as the “rebound model” of T_M maturation (Figure 11D).

However, the acquisition of certain features clearly marks aged CD8⁺ T_M as distinct from CD8⁺ T_N (Figure 11A), and the peculiar blend of naive and memory properties intimates a pronounced similarity to T_{SCM}: with the exception of CD44 expression, aged CD8⁺ T_M not only conform to the murine CD8⁺ T_{SCM} phenotype (28), but bear an extraordinary resemblance to the more extensively characterized human CD8⁺ T_{SCM} population (Supplemental Table 6) with whom they also share superior proliferative and protective potential (49). However, while Gattinoni et al. proposed a scheme that positions T_{SCM} as the least differentiated CD8⁺ T_M subset in a developmental progression that casts the acquisition of stem cell-like properties as the result of arrested T cell differentiation (T_N→T_{SCM}→T_{CM}→T_{EM}→T_E), the “rebound model” posits that T_{SCM}-like CD8⁺ T_M populations constitute a late, advanced, perhaps even “terminally dedifferentiated” stage in the extended CD8⁺ T_M maturation process (T_N→T_E→T_{EM}→T_{CM}→T_{SCM}-like) (Figure 11D).

Discussion

The aging immune system is subject to a confluence of complex adaptations that collectively can compromise specific T cell immunity in particular, immune homeostasis and protection in general, and therefore health and longevity at large (5–10). In apparent contrast, we demonstrate that prolonged preservation of CD8⁺ T_M populations specific for an acute viral pathogen promotes a series of extensive adaptations that impart enhanced II° reactivity and improved protective capacity unto aging CD8⁺ T_M. With the rebound model of CD8⁺ T_M maturation, we have further developed a conceptual and actionable framework that integrates these manifold dynamics with earlier findings about slowly changing CD8⁺ T_M properties including a most recent distinction between early and late CD8⁺ T_{CM} attributes, as documented by Martin et al. (11–17). The precise correlation of proliferative potential and protection, however, is more complex since it is shaped by the nature of the infection (pathogen, route, dosage), the number of available CD8⁺ T_M, their II° reactivity and access to the sites of infection, and the coordinated deployment of various effector functions. In some settings, CD8⁺ T_M populations with more effector-like properties were shown to provide better protection despite poor II° expansion (20, 53). Such observations certainly underscore the challenges associated with the definition of general principles applicable to T cell-mediated immune protection, but it should also be noted that clinically relevant differences (e.g., the level of infectious pathogen) may emerge at times only under very specific experimental conditions (e.g., the “right” combination of transferred CD8⁺ T_M numbers, pathogen challenge dosage, and timing of analyses). Therefore, a combinatorial analysis of such experimental parameters will be particularly informative (52).

The prolonged nature of the CD8⁺ T_M maturation process described here reinforces the notion of early I° immunization and delayed booster regimens as a preferred vaccination strategy (3), an approach that can be refined through heterologous prime-boost (HPB) vaccinations of intact hosts that promote stepwise enlarge-

ments of specific CD8⁺ T_M pools and thus can precipitate improved higher-order recall responses and CD8⁺ T_M formation (54). However, in agreement with the rebound model, HPB vaccination fails to generate superior CD8⁺ T_M populations in experimental settings that limit available CD8⁺ T_M numbers (54), emphasizing that the maturation kinetics of CD8⁺ T_M populations, and with that effective immune protection, are contingent on the specific conditions of their generation. Of particular note here is the precise temporal development of the I° CD8⁺ T_E response, i.e., the combination of I° CD8⁺ T_E population growth rates and relative CD8⁺ T_E burst sizes that determines the pace at which mature phenotypic and functional CD8⁺ T_M properties are acquired at large (Figure 11D). In the case of recall responses, the speed of II° CD8⁺ T cell memory formation is further contingent on the relative survival of II° CD8⁺ T_E populations, itself a function of I° CD8⁺ T_M maturation stage, such that accelerated II° CD8⁺ T_M development may occur despite greater II° CD8⁺ T_E expansion; similar considerations likely apply to the establishment of III° and higher-order memory, though the respective CD8⁺ T_M dedifferentiation processes are progressively delayed or rendered abortive (55).

Our results further indicate that age-associated deteriorations of T cell immunity may be restricted to particular subsets (e.g., T_N or T_M specific for persistent pathogens) (5, 8, 10, 22) while others, as shown here for bona fide pathogen-specific CD8⁺ T_M, are not only spared but apparently improve over time. The extent to which similar processes are operative in humans remains to be investigated by an expansion of longitudinal analyses such as those of Miller et al. (56) or the recent cross-sectional survey of major T cell subsets in multiple tissues across 6 decades of life by Thome et al. (57). The conceptualization of T cell memory as an extended dedifferentiation process in the wake of a defined stimulatory event, however, will have ramifications beyond the context of aging: the introduction of time as an organizing principle for the astonishing phenotypic and functional heterogeneity of the T_M compartment (58) provides a simple ontogenetic perspective that can account for the relational complexity of numerous CD8⁺ T_M properties, and that may conceive of altered T cell modalities observed under conditions of repeated pathogen exposure, persistent infection, absence of CD4⁺ T cell help or age-associated formation of clonal expansions as instances of delayed, aborted, impaired, or divergent dedifferentiation. That the rebound model contributes to an integrated and facilitated understanding of the organization of T cell memory, to the efficient identification of relevant molecular pathways in control of changing homeostatic and recall CD8⁺ T_M properties, and to the improved practical coordination of diagnostic, prophylactic, and therapeutic interventions will be a condition of its utility.

Methods

Mice, viruses, and challenge protocols. C57BL6/J, congenic B6.CD90.1 and B6.CD45.1, and B6.CD45.1.CX3CR1^{Gfp/Gfp}, and OT-I mice were purchased from The Jackson Laboratory; p14 mice were obtained on a B6.CD90.1 background from M. Oldstone (The Scripps Research Institute, La Jolla, California, USA). We only used male mice in this study to avoid potential artifacts due to gender-mismatched AT settings. LCMV Armstrong (clone 53b) and cl13 were provided by M. Oldstone and grown and titered as described (11, 35). Eight- to ten-week-old mice were inoculated i.p. with 2×10^5 PFU LCMV Armstrong; in

some cases, graded dosages (2×10^3 to 2×10^7 PFU) of LCMV Armstrong were used (Figure 9). For II° challenges, recipients of $\text{CD8}^+ \text{T}_{\text{M}}$ were inoculated i.p. with 2×10^5 PFU LCMV Armstrong or 2×10^6 PFU LCMV cl13 i.v. Aging LCMV-immune mice were excluded if they presented with the following conditions: (a) gross physical abnormalities (lesions, emaciation, weight loss); (b) lymphatic tumors at time of necropsy; or (3) T cell clonal expansions in the $\text{D}^{\text{b}}\text{NP}_{396}^+$, $\text{D}^{\text{b}}\text{GP}_{33}^+$, or $\text{D}^{\text{b}}\text{GP}_{276}^+ \text{CD8}^+ \text{T}_{\text{M}}$ compartment. According to these criteria, up to ~30% of aging mice were excluded from the study.

Tissue processing, cell purification, adoptive transfers, and flow cytometry. Standard protocols and procedures were employed as detailed in Supplemental Methods.

Microarray hybridization and analysis. $\text{p14}^+ \text{T}_{\text{E/M}}$ were generated as shown in Figure 3A, purified to greater than 99%, and processed for hybridization to Affymetrix M430.2 arrays as detailed in Supplemental Methods; the data can be retrieved from the NCBI's Gene Expression Omnibus (GEO GSE38462).

Statistics. Data handling, analysis, and graphic representation was performed using Prism 6.0 (GraphPad Software). All data summarized in bar and line diagrams are expressed as the mean \pm 1 SEM and asterisks indicate statistical differences calculated, where applicable, by 1-way ANOVA with Dunnett's multiple comparisons or Student's *t* test (2-tailed unpaired or paired); a *P* value of < 0.05 was considered significant ($*P < 0.05$, $**P < 0.01$, and $***P < 0.001$).

Study approval. All procedures involving laboratory animals were conducted in accordance with the recommendations in the NIH "Guide for the Care and Use of Laboratory Animals", protocols were approved by Institutional Animal Care and Use Committees of the University of Colorado (70205604[05]1F, 70205607[05]4F, B-70210[05]1E) and

Icahn School of Medicine at Mount Sinai (IACUC-2014-0170), and all efforts were made to minimize suffering of animals.

Author contributions

JE, BD, and DH designed the study and individual experiments; JE, BD, TTN, FV, KH, KJ, RMK, ETC, and DH conducted experiments and acquired data; JE, BD, AKF, LEH, RMK, ETC, and DH analyzed data; RMK provided reagents; and DH wrote the manuscript.

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