Coordinate Expression and Trans Presentation of Interleukin (IL)-15Rα and IL-15 Supports Natural Killer Cell and Memory CD8+ T Cell Homeostasis

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Abstract

The high affinity interleukin (IL)-15 receptor, IL-15Rα, is essential for supporting lymphoid homeostasis. To assess whether IL-15Rα’s role in vivo is to trans present IL-15, we generated mixed bone marrow chimera from IL-15Rα– and IL-2/15Rβ–deficient mice. We find that IL-15Rα–competent, IL-2/15Rβ–deficient cells are able to support IL-15Rα–deficient natural killer (NK) and memory CD8+ T cells, thus ruling out secondary signals on these cells and demonstrating that IL-15Rα–mediated presentation of IL-15 in trans is the primary mechanism by which IL-15Rα functions in vivo. Surprisingly, using IL-15– and IL-15Rα–deficient mixed chimera, we also find that IL-15 and IL-15Rα must be expressed by the same cells to present IL-15 in trans, indicating that IL-15Rα is required on a cellular level for the elaboration of IL-15. These studies indicate that IL-15Rα defines homeostatic niches for NK and memory CD8+ T cells by controlling both the production and the presentation of IL-15 in trans to NK and CD8+ memory T cells.

Key words: intracellular cytokine receptor • IL-15/IL-15Rα preassociation • mixed chimera • IL-2Rβ

Introduction

NK cells and memory CD8+ T cells both play vital roles in protecting their host from intracellular pathogens. Understanding how the survival and maintenance of these populations is regulated in vivo has recently been a field of intensive investigation. Memory CD8+ T cell maintenance is a dynamic process that is critically dependent upon two common γ chain (γc)-dependent cytokines, IL-7 and IL-15. Whereas IL-7 promotes the survival of both naive and memory CD8+ T cells, IL-15 uniquely supports basal memory CD8+ T cell proliferation (1–3). Thus, in the absence of proliferative IL-15 signals, memory CD8+ T cells undergo a slow atrophy in number, until they become essentially undetectable (2–4). In addition to maintaining memory CD8+ T cells, IL-15 and IL-15Rα are also critical for the maintenance of peripheral NK cells (5, 6). Whereas the maintenance of memory CD8+ T cells by IL-15 is mediated by both proliferation and survival, NK cell numbers are primarily maintained by regulating survival (5, 6). Thus, IL-15 plays important, nonredundant roles in maintaining the numbers of both memory CD8+ T cells and NK cells in the periphery.

Earlier studies suggested that IL-15 mediates its biological effects by binding to a high affinity, heterotrimeric receptor complex comprised of IL-15Rα, IL-2/15Rβ, and γc. IL-15Rα, which uniquely binds IL-15, is widely expressed by both hematopoietic and parenchymal cell types and has a high affinity for IL-15 (Kd ~10^{-11} M; reference 7). Although IL-15Rα may play a role in intracellular signal transduction in certain cell types, other studies suggest that the cytoplasmic tail of IL-15Rα is not critically necessary for enhancing IL-15–induced proliferation (8–11). By contrast, IL-2/15Rβ and γc heterodimers exhibit lower affinity binding for IL-15 (Kd ~10^{-9} M) in the absence of IL-15Rα, but are clearly essential for transducing IL-15–induced intracellular signals (12, 13). Gene targeting experiments have further demonstrated that all three receptor chains are required to support IL-15–dependent cell populations in vivo (14–16). Importantly, the phenotypes of IL-15– (IL-15–/–) and IL-15Rα– (IL-15Rα–/–) deficient mice are indistin-
guishable, suggesting that physiologically relevant IL-15 signals require IL-15Rα (16, 17).

Although the studies above are consistent with the idea that soluble IL-15 binds to heterotrimeric IL-15Rα, IL-2/15Rβ, and γc receptors on responsive lymphocytes (e.g., NK cells and memory CD8+ T cells) and stimulates their survival and proliferation, more recent studies have demonstrated that IL-15Rα is required in a non-cell–autonomous manner, i.e., not on responding lymphocytes, but rather on a variety of accessory cell types in the mouse (4, 5, 18). IL-15Rα expression on hematopoietic cells other than the CD8+ T cell is required for CD8+ T cell bystander proliferation and preferentially supports the basal maintenance of memory CD8+ T cells (4, 18). Moreover, IL-15Rα expression by both radiation-sensitive and radiation-resistant cells, but not by responding NK cells, is required for the peripheral survival of NK cells (5 and unpublished data). Finally, IL-15Rα expression by radiation-resistant cells, likely intestinal epithelial cells, is critical for the development of TCR-γδ intraepithelial lymphocytes (19). Therefore, as far as the development and subsequent support of several distinct IL-15–dependent cell types is concerned, the critical in vivo functions of IL-15Rα do not appear to be mediated by IL-15Rα expression on IL-15–dependent cell types.

IL-15Rα’s non-cell–autonomous role in supporting NK and memory CD8+ T cells is consistent with multiple indirect mechanisms by which IL-15 might signal through IL-15Rα on accessory cells to induce the production of proteins that subsequently support NK and memory CD8+ T cells. It is also consistent with a novel mechanism described in vitro by which IL-15Rα on accessory cells can present IL-15 in trans to IL-2/15Rβγc– bearing lymphocytes (20). Which of these multiple mechanisms is physiologically relevant has not been addressed in vivo, and the molecular mechanisms underlying this novel cellular physiology have not been investigated. Accordingly, we have used a variety of mixed radiation chimera to examine the in vivo roles of IL-15 and its various receptor chains in supporting NK and memory CD8+ T cell homeostasis.

Materials and Methods

Mice, Adoptive Transfers, and Immunization. C57BL/6j IL-15Rα and congenic Ly5.2 C57BL/6j/SJL IL-15Rα mice, and OT-1 RAG-1 and IL-15Rα OT-1 RAG-1 mice were generated

![Figure 1](image-url)

**Figure 1.** IL-15Rα expression by RAG-1–independent hematopoietic cells is sufficient to maintain NK and memory CD8+ T cells. IL-15Rα−/− mice were lethally irradiated and reconstituted with bone marrow from either IL-15Rα−/− (Wt) or IL-15Rα−/− (RαKO) mice, or a 1:1 mixture of IL-15Rα−/− and IL-15Rα−/− bone marrow (Wt/RαKO), or a 1:1 mixture of RAG-1−/− and IL-15Rα−/− bone marrow (RAG/RαKO). (A) Flow cytometric analyses of NK cell reconstitution in chimeric mice. The percentage of total lymphocytes (defined by forward and side scatter as the R1 gate) that are NK1.1+ CD3− cells is indicated in the top plots. The bottom plots are gated on NK1.1+ CD3− cells. The percentage of total lymphocytes expressing either Ly5.1 or Ly5.2 is indicated in the top right corner of the bottom plots. Note that RAG-1−/− hematopoietic cells support peripheral NK cell development. (B) Graphic representation of percentages of H2Kb-OVA+ CD8+ T cells in immunized chimera mice. 8 wk after irradiation and reconstitution, OT-1+ CD8+ T cells were adoptively transferred into the indicated chimera mice, after which mice were immunized with OVA and poly I:C. The percentage of total lymphocytes that are H2Kb-OVA+ CD8+ T cells after immunization was tracked via serial peripheral blood analyses. Note that RAG-1−/− hematopoietic cells support memory CD8+ T cell homeostasis. Data represent mean ± SEM of at least three mice per group.
IL-15Rα Expression by RAG-1-independent Hematopoietic Cells Supports NK Cell Development. As IL-15Rα is expressed by many types of hematopoietic cells, including macrophages and dendritic cells, we investigated whether IL-15Rα expression by RAG-1-independent cell types could support NK cell development. Therefore, radiation chimera were generated by reconstituting lethally irradiated IL-15Rα mice with a mixture of congenic bone marrow stem cells from RAG-1−/− and IL-15Rα−/− mice (RAG/RaKO→RaKO), or WT and IL-15Rα mice (WT/RaKO→RaKO). After 8 wk, analyses of these mixed chimera revealed that similar numbers of NK cells were present in WT/RaKO→RaKO compared with RAG/RaKO→RaKO chimera (Fig. 1A, top). Moreover, NK cells in RAG/RaKO→RaKO chimera were derived from both RAG-1−/− and IL-15Rα−/− bone marrow progenitors in approximately equal proportions (Fig. 1A, bottom right). Thus, RAG-1-independent hematopoietic cells support NK cell development as well as WT hematopoietic cells.

IL-15Rα Expression by RAG-1-independent Cells Supports Development and Maintenance of Memory CD8+ T Cells. Recent studies indicated that non-cell-autonomous expression of IL-15Rα is important for the maintenance of memory CD8+ T cells (4). To further define the hematopoietic cells that perform this function, we assessed whether IL-15Rα expression by RAG-1-independent cells is sufficient to support memory CD8+ T cells. At least 8 wk after irradiation and reconstitution, we adoptively transferred transgenic OT-1−/−CD8+ T cells into WT→RaKO, RaKO→RaKO, WT/RaKO→RaKO, and RAG/RaKO→RaKO chimera. 2 d after adoptive transfer of OT-1+ CD8+ T cells, these chimera were immunized with OVA and poly I:C, and the initial expansion, memory generation, and maintenance of OT-1+ CD8+ T cells were quantitated by analyzing the numbers of H2Kb-OVA+–reactive CD8+ T cells in peripheral blood samples. At all time points examined after immunization, WT→RaKO, WT/RaKO→RaKO, and RAG/RaKO→RaKO chimera possessed similar frequencies of H2Kb-OVA+ CD8+ T cells (Fig. 1B). In contrast, despite similar primary responses (i.e., day 4 after immunization), the population of H-2Kb-OVA+ CD8+ T cells declined progressively after 30 d in RaKO→RaKO chimera (Fig. 1B). Moreover, normal frequencies of functional memory H-2Kb-OVA+ CD8+ T cells were observed in the spleens and lymph nodes of RAG/RaKO→RaKO, but not RaKO→RaKO, chimera up to 90 d after immunization (not depicted). Therefore, IL-15Rα expression on RAG-1-independent hematopoietic cells supports memory CD8+ T cell generation and maintenance.

IL-2/15Rβ−/− Hematopoietic Cells Support IL-15−dependent Cell Types in Trans; In Vivo Evidence for Trans Presentation as the Exclusive Mechanism of IL-15Rα-mediated Lymphoid Homeostasis. Homeostatic maintenance of NK cells and memory CD8+ T cells requires IL-15Rα expression in a non-cell–autonomous fashion, and in vitro studies suggest that IL-15Rα may function by presenting IL-15 in trans to these cells (4, 5, 19, 20). Taken together, these studies sug-
The frequency of IL-15R expression is required by hematopoietic cells to support NK cell development and survival. (A) Model illustrating two distinct mechanisms by which non-cell-autonomous IL-15Rα expression is required to support IL-15–dependent cell types. Mechanism 1 requires IL-2Rβ expression on accessory cells to mediate signal transduction, whereas mechanism 2 (trans presentation) does not. (B) Flow cytometric analysis of NK cell reconstitution in indicated chimeric mice depends upon IL-15Rα−, but not IL-2/15Rβ−, competent cells. IL-15Rα−/− mice were lethally irradiated and reconstituted with bone marrow from either IL-15Rα+/− or IL-15Rα−/− mice, or with various ratios of IL-15Rα+/− and IL-15Rα−/− bone marrow, or assorted ratios of IL-2/15Rβ−/− and IL-15Rα−/− bone marrow. Mice were bled and NK cell populations in chimeric mice were assessed 8 wk after reconstitution. The percentages of total lymphocytes that are NK cells (NK1.1+ CD3− cells) are indicated in each of the top panels. The bottom plots are gated on NK1.1+ CD3− cells in WT/RaKO and RβKO/RaKO chimeras. The percentages of total lymphocytes expressing either Ly5.1 or Ly5.2 in WT/RaKO and RβKO/RaKO chimeras shown in the bottom right corner.

Figure 2. IL-15Rα, but not IL-2/15Rβ, expression is required by hematopoietic cells to support NK cell development and survival. (A) Model illustrating two distinct mechanisms by which non-cell-autonomous IL-15Rα expression is required to support IL-15–dependent cell types. Mechanism 1 requires IL-2Rβ expression on accessory cells to mediate signal transduction, whereas mechanism 2 (trans presentation) does not. (B) Flow cytometric analysis of NK cell reconstitution in indicated chimeric mice depends upon IL-15Rα−, but not IL-2/15Rβ−, competent cells. IL-15Rα−/− mice were lethally irradiated and reconstituted with bone marrow from either IL-15Rα+/− or IL-15Rα−/− mice, or with various ratios of IL-15Rα+/− and IL-15Rα−/− bone marrow, or assorted ratios of IL-2/15Rβ−/− and IL-15Rα−/− bone marrow. Mice were bled and NK cell populations in chimeric mice were assessed 8 wk after reconstitution. The percentage of total lymphocytes that are NK1.1+ CD3− cells is shown. Plots are representative of at least two mice per condition, and all experiments were performed three times with similar results.

The frequency of IL-15Rα−competent cells controls the frequency of peripheral NK cells. IL-15Rα−/− mice were lethally irradiated and reconstituted with bone marrow from either IL-15Rα+/− or IL-15Rα−/− mice, or with various ratios of IL-15Rα+/− and IL-15Rα−/− bone marrow, or assorted ratios of IL-2/15Rβ−/− and IL-15Rα−/− bone marrow. Mice were bled and NK cell populations in chimeric mice were assessed 8 wk after reconstitution. The percentages of total lymphocytes expressing either Ly5.1 or Ly5.2 in WT/RaKO and RβKO/RaKO chimeras shown in the bottom right corner.

We then investigated the cell-autonomous role of IL-2/15Rβ expression in supporting NK cells by examining the genotype of surviving NK cells in WT/RaKO→RαKO and RβKO/RaKO→RαKO chimera via congenic markers. Consistent with the data above, the percentages of NK cells derived from IL-15Rα+/− and IL-15Rα−/− bone marrow stem cells were similar to the percentages of non–IL-15Rα−dependent lymphocytes (B and CD4+ T cells) in WT/RaKO→RαKO chimera (Fig. 2 B, bottom left). By contrast, the percentage of total NK cells that were derived from Ly5.1+ IL-2Rβ−/− cells (8.4%) was dramatically reduced when compared with the percentage of other Ly5.1+ lymphocytes (41%) reconstituted in RβKO/RaKO→RαKO chimera (Fig. 2 B, bottom right). These data indicate that IL-2/15Rβ expression is required on NK cells for their development and maintenance.

Finally, as both WT/RaKO→RαKO and RβKO/RaKO→RαKO chimera generally contained reduced percentages of NK cells compared with WT/RaKO chimera, we hypothesized that the percentage of peripheral NK cells might be a function of the relative percentage of IL-15Rα−competent hematopoietic cells. To investigate this possibil-
ity, we reconstituted IL-15Rα−/− mice with either 1:1 or 1:4 mixtures of IL-15Rα−/− and either WT or IL-2/15Rβ−/− bone marrow stem cells. Examination of the percentage of NK cells in these chimeras revealed that the numbers of NK cells decreased as the proportion of IL-15Rα–competent (either WT or IL-2/15Rβ−/−) bone marrow stem cells decreased (Fig. 2 C). These findings suggest that the relative frequency of IL-15Rα–competent hematopoietic cells regulates the size of the peripheral NK cell pool.

**IL-2/15Rβ−/− Hematopoietic Cells Support Memory CD8+ T Cells.** Memory phenotype CD8+ T cells are dependent upon both IL-15 and IL-15Rα for their development and peripheral survival (16, 17, 22). To investigate whether IL-2/15Rβ−/− cells could support memory phenotype CD8+ T cells in a non-cell–autonomous fashion, we examined the reconstitution of this population in RβKO/RαKO chimera. CD44hi IL-2/15Rβhi CD8+ T cells were readily observed in both WT→RαKO and RβKO/RαKO→RαKO chimera, but not in RαKO→RαKO chimera (Fig. 3 A). Notably, there was no obvious population of IL-2/15Rβ−/− CD44hi CD8+ T cells present in RβKO/RαKO→RαKO chimera, suggesting that IL-2/15Rβ, but not IL-15Rα, expression by CD44hi CD8+ memory phenotype cells is critical for their peripheral maintenance (Fig. 3 A and not depicted).

CD44hi IL-2/15Rβhi CD8+ T cells include antigen-experienced memory cells as well as cells that may have been activated via alternate mechanisms (e.g., homeostatic expansion). To directly assess the ability of IL-2/15Rβ−/− hematopoietic cells to support antigen-experienced memory CD8+ T cells, we adoptively transferred naive OT-1+ RAG-1−/− CD8+ T cells into WT→RαKO, RαKO→RαKO, WT/RαKO→RαKO, or RβKO/RαKO→RαKO chimera 8 wk after reconstitution. These chimeras were then immunized with OVA and poly I:C, and the frequency of OT-1+ CD8+ T cells was serially examined as described above. Similar primary responses of OT-1+ CD8+ T cells were observed in all types of chimera during the first 20–30 d after immunization. However, after ~50 d, progressive loss of these cells was noted in RαKO→RαKO chimera, but not in the other types of chimera, all of which contained IL-15Rα–competent hematopoietic cells (Fig. 3 B). Thus, in parallel to our findings with NK cells, IL-2/15Rβ−/− hematopoietic cells are capable of supporting memory CD8+ T cells in a non-cell–autonomous fashion, and this finding suggests that hematopoietic cells use IL-15Rα to support memory CD8+ T cells exclusively by a trans presentation mechanism (Fig. 2 A).

**Coordinate Expression of IL-15 and IL-15Rα by Hematopoietic Cells Is Required for Supporting NK Cells In Vivo.** The findings described above support a model whereby RAG-1–independent hematopoietic cells use IL-15Rα, but not IL-2/15Rβ, to present IL-15 in trans to IL-2/15Rβ−, but not IL-15Rα−, dependent receptors on NK and memory CD8+ T cells. However, it remained unclear why NK and memory CD8+ T cells are able to respond to soluble or plate-bound IL-15 in vitro, but apparently fail to receive IL-15 signals in mice that are IL-15Rα deficient but IL-15 competent. One possible explanation would be that IL-15Rα is

![Figure 3](image-url)
not only required on accessory cells for trans presenting IL-15, but is also required for making IL-15 bioavailable for trans presentation. In this scenario, IL-15 would not be freely available in the serum of mice to be bound by IL-15Rα-presenting cells, but might need to be produced by the same cells that produce IL-15Rα. To test this hypothesis, we investigated whether coordinate expression of IL-15 and IL-15Rα was required to support NK and memory CD8+ T cells in vivo (Fig. 4). Lethally irradiated IL-15Rα−/− mice were reconstituted with WT (WT→RαKO), IL-15Rα−/− (RαKO→RαKO), IL-15−/− (15KO→RαKO), a mixture of WT and IL-15Rα−/− (WT/RαKO→RαKO), or a mixture of IL-15−/− and IL-15Rα−/− bone marrow stem cells (15KO/RαKO→RαKO). Roughly half of the hematopoietic cells in 15KO/RαKO→RαKO chimera should express IL-15Rα, but not IL-15, whereas the other half of hematopoietic cells and all residual stromal cells should express IL-15, but not IL-15Rα. Thus, if IL-15 can be secreted from IL-15Rα−/− cells to bind to IL-15Rα expressed on IL-15−/− cells, then 15KO/RαKO→RαKO chimera should support IL-15-dependent lymphocytes as well as WT/RαKO→RαKO chimera. Alternatively, if IL-15Rα is required for IL-15 elaboration, then 15KO/RαKO→RαKO chimera would not be able to support NK and CD8+ memory T cells. Analyses of NK cells in these various chimera 8 wk after reconstitution revealed that WT→RαKO and WT/RαKO→RαKO chimera contained significant numbers of peripheral NK cells, whereas

Figure 4. Two models for the relationship of IL-15-producing and -presenting cells. In model 1, IL-15 is secreted by a distinct IL-15-producing cell, and subsequently bound and presented by an IL-15Rα–competent cell. In model 2, coordinate expression of IL-15 and IL-15Rα is required for efficient presentation of IL-15 to responding cells.

Figure 5. Coordinate expression of IL-15 and IL-15Rα is required for NK cell development and maintenance. IL-15Rα−/− mice were lethally irradiated and reconstituted with bone marrow cells from either IL-15Rα−/− (Wt), IL-15Rα−/− (RαKO), or IL-15−/− (15KO), mice, or with 1:1 mixtures of either IL-15Rα−/− and IL-15Rα−/− bone marrow cells (Wt/RαKO), or IL-15Rα−/− and IL-15−/− bone marrow cells (15KO/RαKO). (A) Flow cytometric analyses of endogenous NK cells in bone marrow (BM) and spleens (SPL) of chimera mice 8 wk after reconstitution with the indicated bone marrow genotypes. Note that endogenous NK cells are not supported in chimera generated from a mixture of IL-15−/− and IL-15Rα−/− bone marrows. Data are representative of at least four mice per group. (B) Flow cytometric analysis of IL-15Rα expression on LPS-stimulated, bone marrow–derived dendritic cells of the indicated genotypes. Bone marrow–derived dendritic cells were stimulated with LPS for 24 h in vitro and stained for IL-15Rα expression. (C) Graphic representation of survival of adoptively transferred NK cells after transfer into the indicated mixed chimera. The percentage of total lymphocytes that were adoptively transferred NK cells (i.e., CFSE− NK1.1+ CD3+ cells) is indicated at each time point. Note that adoptively transferred NK cells are not supported in chimera generated from a mixture of IL-15−/− and IL-15Rα−/− bone marrows. Data represent mean ± SEM of at least two mice per group.
15KO→RaKO, RaKO→RaKO, and 15KO/RaKO→RaKO chimera revealed that transferred NK cells persisted for >2 d in WT→RaKO and WT/RaKO→RaKO chimera, but not in RaKO→RaKO, 15KO→RaKO, or most notably, 15KO/RaKO→RaKO chimera (Fig. 5 C). Therefore, coordinate expression of IL-15 and IL-15α is required to support peripheral NK cell survival.

Coordinate Expression of IL-15 and IL-15α by Hematopoietic Cells Is Required for Supporting Memory CD8+ T Cells In Vivo. Next, we investigated whether coordinate expression of IL-15Rα and IL-15 is required for the development and maintenance of memory phenotype CD8+ T cells. Analyses of tissues from various chimera revealed that IL-2/15Rβ/CD8+ T cells were readily observed in WT→RaKO chimera, but not in RaKO→RaKO, 15KO→RaKO, or 15KO/RaKO→RaKO chimera (Fig. 6 A). Therefore, coordinate expression of both IL-15Rα and IL-15 is necessary for maintenance of memory phenotype CD8+ T cells.

Finally, we examined whether coordinate expression of IL-15 and IL-15Rα is required for the generation and maintenance of antigen-experienced memory CD8+ T cells. We adoptively transferred OT-1+ CD8+ T cells into WT→RaKO, WT/RaKO→RaKO, RaKO→RaKO, 15KO→RaKO, and 15KO/RaKO→RaKO chimera, immunized these mice with poly I:C and OVA 2 d later, and examined the kinetics of transgenic T cell responses. Although the primary expansions of these cells were similar 4 d after immunization in the various chimera, memory OT-1+ CD8+ T cells were subsequently maintained in WT→RaKO and WT/RaKO→RaKO chimera, but not in RaKO→RaKO, 15KO→RaKO, and 15KO/RaKO→RaKO chimera (Fig. 6 B and not depicted). This selective loss of memory OT-1+ CD8+ T cells in 15KO→RaKO and 15KO/RaKO→RaKO chimera was particularly evident when the numbers of functional memory OT-1+ CD8+ T cells were assessed by analyzing IFN-γ production in response to the cognate peptide SIINFEKL. Although WT→RaKO chimera had significant numbers of IFN-γ+ CD44hi CD8+ T cells 90 d after immunization, RaKO→RaKO, 15KO→RaKO, and 15KO/RaKO→RaKO chimera possessed negligible numbers of SIINFEKL-responsive cells (Fig. 6 C). Thus, like NK cells, functional memory CD8+ T cells require coordinate expression of IL-15Rα and IL-15 for their maintenance.

Discussion

RAG-1–independent, IL-15Rα–competent Cells Define Homeostatic Space for NK Cell Survival and Memory CD8+ T Cell Homeostasis. IL-15 regulates the homeostasis of NK and memory CD8+ T cells, and the high affinity IL-15R, IL-
15Rα, is critical for mediating IL-15’s functions in vivo. These observations suggest that the bioavailability of IL-15 and IL-15Rα defines a homeostatic space that regulates the numbers of these lymphocytes that an organism possesses at any one time. However, the cellular and molecular bases of these homeostatic interactions are poorly understood. In this work, we have examined the cellular mechanisms by which IL-15Rα supports lymphoid homeostasis in vivo. Our experiments indicate that RAG-1–independent hematopoietic cells comprise the predominant cell types that provide IL-15Rα–dependent homeostatic support. In this regard, myeloid cells such as macrophages and dendritic cells express low levels of both IL-15 and IL-15Rα constitutively, and express higher levels in response to proinflammatory stimuli. We have also found that the proportion of IL-15Rα–competent hematopoietic cells in WT/RαKO→RαKO mixed chimera correlates directly with the number of NK cells maintained in these mice. Thus, the number of IL-15Rα–competent accessory cells might be a limiting resource for NK and CD8+ T cells in resting animals. We have separately examined both the survival of peripheral NK cells and the generation and maintenance of memory CD8+ T cells in various mixed chimera, and found similar IL-15Rα requirements for these distinct populations. As the peripheral homeostasis of NK cells is largely maintained by cell survival, while memory CD8+ T cell homeostasis is supported by both survival and proliferation, these results suggest that IL-15Rα regulates multiple cellular processes in a cell-type– and context-dependent fashion. Taken together, these experiments help define the nature of “homeostatic space” available to IL-15–responsive lymphocytes.

Trans Presentation as the Dominant Physiological Mechanism by Which IL-15Rα Supports NK and CD8+ Memory T Cells In Vivo. Previous studies indicated that IL-15Rα supports NK cell and CD8+ memory T cell homeostasis in a non-cell–autonomous fashion in vivo (4, 5, 18). The non-cell–autonomous mechanism(s) by which IL-15Rα–competent hematopoietic cells support NK and CD8+ T cells could occur via two nonexclusive mechanisms. First, IL-15Rα–competent accessory cells could transduce signals through their heterotrimeric IL-15Rs and synthesize secondary proteins that support NK and CD8+ T cells. Alternatively, IL-15Rα–competent accessory cells could use IL-15Rα to directly present IL-15 in trans to NK and CD8+ T cells. Previous studies have shown that IL-2/15Rβ expression is required for IL-15–induced proliferative responses (7, 12). Moreover, IL-2/15Rβ−/−, but not IL-2Rα−/−, mice lack NK cells, similar to both IL-15−/− and IL-15Rα−/− mice. Taken together, these data suggest that IL-2/15Rβ is critical for IL-15 responses. Hence, our finding that IL-2/15Rβ−/− hematopoietic cells perform as well as WT hematopoietic cells in supporting NK and CD8+ T cells indicates that accessory cells mediate this function without transducing IL-15 signals themselves, and without producing secondary proteins that in turn support lymphocytes. Therefore, these accessory cells use IL-15Rα exclusively to directly present IL-15 in trans to NK cells and CD8+ memory T cells. This finding is consistent with recent findings that IL-2/15Rβ−/− hematopoietic cells can support intraepithelial lymphocyte homeostasis (19). Thus, trans presentation is likely to be the exclusive physiological mechanism by which IL-15Rα supports NK and CD8+ T cells in vivo.

Our studies also shed light on the cell–autonomous requirements for IL-15Rα signaling in lymphoid homeostasis. Analyses of congenic NK cells recovered from RβKO/RαKO→RαKO mixed chimera indicate that IL-2/15Rβ expression by IL-15–responsive NK and memory CD8+ T cells is required for their homeostasis. This finding confirms previous suggestions that high expression levels of IL-2/15Rβ on these cells correlates with their sensitivity to IL-15–dependent signals in vivo (23, 24). Meanwhile, analyses of congenic NK cells recovered from our WT/RαKO→RαKO mixed chimera indicate that IL-15Rα on IL-15–dependent NK and memory CD8+ T cells is entirely dispensable for their homeostasis in vivo. These in vivo data are consistent with the fact that IL-15Rα+/+ and IL-15Rα−/−/− memory CD8+ T cells respond similarly to a given dose of IL-15 in vitro, regardless of whether it is provided as a plate-bound IL-15/IL-15Rα–γc complex or as a soluble cytokine (unpublished data). Hence, despite the fact that IL-15Rα can augment signaling responses to soluble IL-15 in transfected cells, it is unlikely that IL-15Rα on NK and memory CD8+ T cells facilitates binding of IL-15 to IL-2/15Rβ and γc receptor chains on these cells in vivo. Similarly, it is unlikely that IL-15Rα on accessory cells transfers IL-15 to IL-15Rα on responding NK and CD8+ memory T cells. Taken together, these studies indicate that IL-2/15Rβ receptors are critical, whereas IL-15Rα receptors are entirely dispensable on NK and memory CD8+ T cells for their homeostasis.

Trans presentation is a novel mechanism by which cytokine signals are transduced. Although a previous report suggested that IL-2Rα could present IL-2 in trans (25), IL-2Rα alone binds IL-2 with low affinity (Kd ~10^-8 M) and in vivo studies with IL-2Rα−/−/− T cells indicated that IL-2Rα plays a cell-autonomous role in supporting T cells (26–28). Thus, trans presentation is unlikely to be the physiological mechanism by which IL-2Rα supports T cells. Signaling through the IL-6 receptors, IL-6Rα and gp130, more closely resembles IL-15R signaling. Soluble IL-6Rα is produced by both proteolytic cleavage of IL-6Rα and alternative splicing. Soluble IL-6Rα binds IL-6 in solution and IL-6–sIL-6Rα complexes then bind to gp130 receptors on cell surfaces to initiate signal transduction events (29, 30). Nevertheless, there might be fundamental differences between IL-6Rα– and IL-15Rα–mediated signaling. Specifically, in contrast to IL-6Rα, it is unclear if IL-15 and IL-15Rα can form soluble complexes that can signal to IL-2/15Rβ−/− receptors on responding cells (11, 31). Therefore, the ability of IL-15Rα on the surface of accessory hematopoietic cells to present IL-15 in trans to IL-2/15Rβ and γc low affinity dimeric receptors on NK and memory CD8+ T cells in vivo may represent a novel
mechanism of cytokine signaling that may involve cell to cell contact.

Coordinate Expression of IL-15 and IL-15Ra by Trans Presenting Accessory Cells. Our studies with 15KO/RaKO→RaKO mixed chimera indicate that IL-15 and IL-15Ra must be expressed by the same accessory cells to support both NK and memory CD8+ T cells in vivo. As IL-15−/− cells express normal levels of cell surface IL-15Ra (Fig. 5B), and as IL-15Ra−/− cells express normal levels of IL-15 mRNA (5, 18), the inability of 15KO/RaKO→RaKO chimera to support NK and memory CD8+ T cells suggests that IL-15Ra−/− cells may not elaborate IL-15 protein. This surprising finding would explain why IL-15Ra−competent lymphocytes respond to heterologous IL-15 in vitro, but fail to respond to IL-15 elaborated from IL-15–competent cells in vivo. Thus, IL-15Ra might be essential for either the translation of IL-15 mRNA or the trafficking of IL-15 protein to the cell surface. With regards to the latter possibility, it is known that the signal sequences of IL-15 mediate protein secretion poorly (32). As IL-15Ra associates with IL-15 with high affinity, one intriguing possibility is that IL-15Ra may bind to IL-15 intracellularly and facilitate trafficking of IL-15/IL-15Ra to the surface of accessory cells. Intracellular association of IL-15 and IL-15Ra has recently been described in several contexts, including endosomal recycling of internalized IL-15Ra–IL-15 complexes that follow binding of extracellular IL-15 to surface IL-15Ra (20, 33, 34). By contrast, our current findings suggest that the critical interactions between IL-15 and IL-15Ra occur within IL-15-producing cells, before IL-15’s emergence on the plasma membrane. In addition, as soluble IL-15 has been difficult to document in mice, it is possible that IL-15Ra recognizes and binds to IL-15 exclusively within cells that synthesize both proteins. Therefore, these experiments indicate novel cell biological requirements for the regulation of IL-15, and also provide a compelling explanation for why trans presentation is the physiological mechanism by which IL-15 supports lymphoid homeostasis.

In summary, we have examined the mechanism by which IL-15Ra supports NK cell survival and CD8+ memory T cell proliferation in vivo. Our findings indicate that myeloid accessory cells do not use IL-15Ra to transduce signals leading to the elaboration of secondary proteins that support lymphoid homeostasis. Instead, these cells use IL-15Ra to present IL-15 in trans to IL-2/15Rβ–bearing receptors on the surface of NK and CD8+ memory T cells. These accessory cells must coordinate synthesis of IL-15 and IL-15Ra to present IL-15 in trans. Therefore, the critical events regulating homeostatic niches for NK and CD8+ memory T cells in vivo can be focused upon the production and trans presentation of IL-15 by IL-15Ra–expressing myeloid cells.

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