Hematopoietic Cells and Radioresistant Host Elements Influence Natural Killer Cell Differentiation

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Summary

Radioresistant host elements mediate positive selection of developing thymocytes, whereas bone marrow-derived cells induce clonal deletion of T cells with receptors that are strongly autoreactive. In contrast to T cell development, little is known about the elements governing the natural killer (NK) cell repertoire, which, similar to the T cell repertoire, differs between individuals bearing different major histocompatibility complex (MHC) phenotypes. We have used murine bone marrow transplantation models to analyze the influence of donor and host MHC on an NK cell subset. We examined the expression of Ly-49, which is strongly expressed on a subpopulation of NK cells of H-2b mice, but not by NK cells of H-2a mice, probably because of a negative effect induced by the interaction of Ly-49 with Dα. To evaluate the effect of hematopoietic cell H-2β expression on Ly-49 expression of H-2b NK cells, we prepared mixed allogeneic chimeras by administering T cell-depleted allogeneic (B10.A, H-2β) and host-type (B10, H-2b) marrow to lethally irradiated B10 mice, or by administering B10.A marrow to B10 recipients conditioned by a nonmyeloablative regimen. Expression of H-2β on bone marrow-derived cells was sufficient to downregulate Ly-49 expression on both H-2a and H-2b NK cells. This downregulation was thymus independent. To examine the effect of H-2a expressed only on radioresistant host elements, we prepared fully allogeneic chimeras by administering B10 bone marrow to lethally irradiated B10.A recipients. B10 NK cells of these fully allogeneic chimeras also showed downregulation of Ly-49 expression. The lower level of H-2a expressed on H-2b × H-2a F1 cells induced more marked downregulation of Ly-49 expression on B10 NK cells when presented on donor marrow in mixed chimeras than when expressed only on radioresistant host cells. Our studies show that differentiation of NK cells is determined by interactions with MHC molecules expressed on bone marrow-derived cells and, to a lesser extent, by MHC antigens expressed on radioresistant host elements.

T cells undergo negative and positive selection during development, resulting in self-MHC-restricted recognition of peptide antigens, and deletion of T cell clones with strong reactivity to self-antigens. Positive selection is thought to be mediated predominantly by thymic epithelial cells, whereas clonal deletion of self-reactive T cell progenitors is mediated primarily by bone marrow-derived cells populating the thymus (1, 2). Less is known about the selection processes which NK cells undergo during their differentiation. Although patterns of NK cell-mediated recognition are known to differ between MHC-disparate mouse strains (3) and human individuals (4), the environmental determinants of the NK cell repertoire have yet to be defined. We have used radiation bone marrow chimeras to address separately the influence of bone marrow-derived cells and of radioresistant host elements on NK cell expression of a surface molecule that recognizes MHC class I antigens and regulates cytolytic activity.

Ly-49 is a recently defined surface molecule expressed on a subpopulation of NK cells (5). The putative interaction of Ly-49 with Dα or Dβ class I MHC molecules on target cells globally inhibits cytolysis by Ly-49+ NK cells suggesting that Ly-49 is an inhibitory NK cell receptor specific for these MHC class I antigens (5). Presumably resulting from a negative effect induced by recognition of Dα or Dβ, Ly-49 is not detected on NK cells of animals bearing these MHC antigens (6). We took advantage of this observation to evaluate the effect of Dα expression on bone marrow or radioresistant host elements on NK cell Ly-49 expression.

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Materials and Methods

Animals. C57BL/10 (B10, H-2^s, K^bD^b), B10.A (H-2^a, K^bD^a), AK/JscNCR (A/J, H-2^d) and (B10 x B10.A)F1 (H-2^a x H-2^d) mice were purchased from Frederick Cancer Research Facility, Frederick, MD. B10.BR (H-2^s, K^bD^a) mice were purchased from the Jackson Laboratory (Bar Harbor, ME). Mice were housed in sterile microisolator cages in a viral antibody-free environment.

Bone Marrow Transplantation (BMT). Chimeras were prepared as described (7) by reconstituting 10.25 Gy whole body irradiated (WBI) (1.03 Gy/min, ^137Cs source) B10, B10.A, or (B10 x B10.A)F1 mice, or 9.5 Gy WBI A/J mice with the indicated T cell–depleted (TCD) marrow inocula. T cell depletion of bone marrow cells (BMC) was carried out as described (8). All bone marrow inocula were T cell depleted, unless otherwise indicated.

Another BMT model involved administration of 5 x 10^6 non-TCD B10 BMC to B10.BR mice conditioned with sublethal (7 Gy) total body irradiation (TBI) and pretreated with depleting doses of anti-NK1.1 mAb PK136 on days -6 and -1 before BMT. These animals showed complete donor-type reconstitution in the lymphoid and myeloid lineages.

A third BMT model involved administration of 15 x 10^6 non-TCD B10.A or A/J BMC to B10 mice pretreated with anti-CD4 and anti-CD8 mAbs, 7 Gy thymic irradiation, and 3 Gy TBI. This nonmyeloablative conditioning regimen led to induction of permanent mixed chimerism after BMT, as previously described (10).

Thymectomy. Animals were anesthetized intraperitoneally with pentobarbital and thymectomy was perfomed under a partial median sternotomy, as described (11).

mAbs and Flow Cytometry (FCM). FCM analyses were performed on a FACScan® or FACSsort® (Becton Dickinson & Co., Mountain View, CA). Chimerism was evaluated by FCM analysis of peripheral white blood cells (WBC) prepared by hypotonic lysis of RBC. WBC were stained with FITC- or biotin-conjugated (plus PE-streptavidin [PEA]) D^d-specific mouse IgG2a mAb 34-2-12 (12), K^b-specific mouse IgG2a mAb 16-11-13 (13), or K^b-specific mouse IgG2a mAb 36-7-5 (13) to distinguish H-2^d (D^d-positive, K^b-negative) from H-2^b (D^b-negative, K^b-negative) cells. Nonreactive IgG2a mAb HOPC1 was used as a negative control. Chimerism was evaluated for lymphocytes, monocytes, and granulocytes, which could be distinguished on the basis of forward angle versus 90° light scatter characteristics.

For analysis of Ly-49 expression on NK cells, PBL were prepared and labeled with anti-Ly-49 mAb A1-FITC (14) and NK1.1-specific IgG2a mAb PK136-biotin (PharMingen, San Diego, CA) plus PEA, or with PK136-PE (PharMingen). To determine the percentage of NK1.1^+ cells that were Ly-49^+, 10,000 gated PK136^+ cells (distinguished by staining above negative control mAbs HOPC1-biotin/PEA or Leu4-PE) were collected for analysis of staining with A1-FITC. The percentage of gated PK136^+ cells staining with control antibody HOPC1-FITC was subtracted from the percentage of gated PK136^+ cells staining with A1-FITC. PBL were also stained with negative control mouse IgG2a mAb HOPC1-FITC versus PK136-biotin/PEA and 36-7-5 (anti-K^b)-FITC versus PK136-biotin/PEA (or PK136-PE).

Results and Discussion

Recovery of Ly-49^+ NK Cells in Syngeneic (B10→B10) BMT Recipients. We used two-color FCM with Ly-49–specific mAb A1 to evaluate Ly-49 expression on NK1.1^+ cells of chimeras prepared in the B10.A (H-2^d, D^d)/B10 (H-2^b, D^b) strain combination. At all time points tested 7–30 wk after BMT, B10→B10 BMT recipients showed similar percentages of Ly-49^+ NK cells to those in B10 controls tested in the same assay (Figs. 1 f and 2). Although “dull” Ly-49 staining was variably detected on NK cells of D^d-bearing H-2^d B10.A, H-2^b x H-2^d (B10 x B10.A)F1, or H-2^b B10.BR mice, only Ly-49 staining producing a “dull” shoulder above background staining on B10.A NK cells had a peak fluorescence of 10 channels, compared with 41 channels for the distinct peak produced by Ly-49 staining of B10 NK cells in the same assay, brightly staining Ly-49^+ cells were not detected on normal mice of these strains, consistent with previous results (6). The reason for this variable dull Ly-49 expression is not known, and is currently under investigation.

A subpopulation of NK1.1^+ cells expresses TCR-α/β (16–18). We used three-color FCM analysis to determine the percentage of NK1.1^+ Ly-49^− cells that expressed TCR-α/β in PBL of three B10→B10 BMT recipients and a normal B10 mouse. Only 1.22–6.01% of NK1.1^+ Ly-49^− PBL expressed TCR-α/β. Thus, the vast majority of NK1.1^+ Ly-49^− cells were TCR-α/β-negative.

H-2^b Bone Marrow Downregulates Ly-49 Expression on H-2^d NK Cells. To examine the effect of hematopoietic cell H-2^d (D^d) expression on Ly-49 expression by D^b NK cells, we prepared mixed chimeras by reconstituting lethally irradiated B10 mice with a mixture of TCD allogeneic and syngeneic (host type) marrow obtained from B10 and B10.A donors (i.e., B10+B10.A→B10). These mixed chimeras show specific T cell tolerance to antigens of the donor and of the host (19). The percentage of NK1.1^+ cells in PBL of control and chimeric animals was similar (3.4 ± 1.3% SD for normals; 2.8 ± 1% SD for mixed chimeras; P = 0.1), and two-color FCM analysis showed that the proportion of NK cells of donor and host type was similar to the proportion of total PBL of each type. For example, in a chimera in which 48.5% of PBL

1 Abbreviations used in this paper: BMC, bone marrow cell; BMT, bone marrow transplantation; Cy-SA, CyChrome-streptavidin; FCM, flow cytometry; PEA, phycoerythrin-streptavidin; TBI, total body irradiation; TCD, T cell depleted.
Figure 1. Ly-49 downregulation on NK cells of mixed chimeras. Kk (a-d) and Ly-49 (e-h) staining is shown for gated NK1.1+ PBL from normal B10.A (a and e), B10 BMT recipient (b and f), B10+B10.A→B10.A (c and g), and B10+B10.A→B10 mixed chimera (d and h) 8 wk after BMT. Despite the presence of Kk+ (B10) NK cells in both types of mixed chimeras (c and d), significant Ly-49 expression was not detectable (g and h).

Figure 2. H-2k expressed on bone marrow-derived cells or on radioresistant host elements downregulates Ly-49 expression on B10 NK cells. The mean percentage ± SD of NK1.1+ PBL expressing Ly-49 at 5-9 wk after BMT is shown for lethally irradiated (10.25 Gy) B10, B10.A and (B10 × B10.A)F1 recipients of the indicated TCD BMC inocula. One group of B10.A mice received non-TCD B10 marrow, as indicated. (*) Significant difference from B10→B10 syngeneic controls (p < 0.05, Student's t test). The mean ± SD percentage of NK1.1+ PBL expressing Ly-49 was 8.3 ± 3.8 for normal B10 (n = 15), 0.5 ± 0.5 (n = 10) for normal B10.A, and 0.55 (n = 2) for normal (B10 × B10.A)F1 controls. Mean percentages of Dd-positive PBL in mixed chimeras were as follows (TCD BMC donors, left of the arrow; recipient strains, right of the arrow): B10+B10.A→B10 68.6% (range 28.2-90.7%); B10+B10.A→B10.A 45.7% (range 22.7-90.0%); B10+(B10 × B10.A)F1→B10 68.7% (range 64.4-72.0%).

Figure 3. (B10 × B10.A)F1 bone marrow-derived cells downregulate Ly-49 expression on B10 NK cells to a greater extent than do radioresistant (B10 × B10.A)F1 host cells. (Solid histograms) Staining with nonreactive control mAb HOPC-1-FITC; (empty histograms) staining with anti-Ly-49 mAb A1-FITC on gated Dd-, NK1.1+ PBL obtained from typical B10→B10, B10+(B10 × B10.A)F1→B10, and B10+(B10 × B10.A)F1 BMT recipients analyzed by three-color FCM in a single assay performed 10-23 wk after BMT. Gated Dd− NK1.1+ PBL were analyzed for (B10 × B10.A) F1→(B10 × B10.A) F1 recipients (top right) in the same assay.
TBI, 7 Gy thymic irradiation, and administration of anti-CD4 plus anti-CD8 mAbs as previously described (10), showed a similar downregulation of Ly-49 expression on B10 NK cells 3 and 20 wk after BMT (data not shown). Therefore, engraftment of H-2^a BMC without lethal irradiation was sufficient to downregulate Ly-49 expression on B10 NK cells as early as 3 wk after BMT.

Similar downregulation of Ly-49 expression was also observed in chimeras prepared by this regimen after adult thymectomy (data not shown), indicating that bone marrow-derived cells can exert their downregulatory influence on Ly-49 expression at an extrathymic site.

A previous study showed that administration of β2 microglobulin-negative, class I MHC-deficient bone marrow to class I-expressing recipients results in a pattern of NK cell function that resembles that of the class I-deficient donors (20). Our Ly-49 studies in mixed chimeras demonstrate for the first time that MHC molecules expressed on bone marrow-derived cells can influence the differentiation of NK cells not expressing these MHC molecules. This result suggests that expression of alloantigen on donor bone marrow-derived cells might tolerate host-type NK cells that could otherwise be alloreactive against donor marrow. Since mixed chimerism confers several advantages over completely allogeneic reconstitution as an approach to tolerance induction across both allogeneic and xenogeneic barriers (19, 21), the possible ability of donor marrow-derived cells to tolerate host-derived NK cells could have important implications, particularly in xenotransplantation, in which NK cell activation in recipients results in a pattern of NK cell differentiation as an approach to tolerance induction across both allogeneic and xenogeneic barriers (19, 21).

Expression of H-2^d on Radioresistant Host Elements Downregulates Ly-49 Expression on B10 NK Cells. To determine whether D^d expression on radioresistant host elements could also downregulate Ly-49 expression, we studied fully allogeneic TCD B10→B10.A (10.25 Gy WBI) chimeras, in which D^d is expressed only by the host. In these chimeras, Ly-49 expression was markedly downregulated on B10 NK cells evaluated 4–39 wk after BMT (Figs. 4f and 2).

A small proportion of host-type cells is detectable in the PBL of allogeneic chimeras of this type (22) (Fig. 4d). To rule out the possibility that these residual host-type cells are required for Ly-49 downregulation, we performed similar studies in B10.A recipients of non-TCD B10 BMC, in which no host-type lymphohematopoietic cells are detectable (22) (compare Fig. 4g to syngeneic B10→B10 control, Fig. 4a). These animals also showed marked downregulation of Ly-49 expression on B10 NK cells (c.e., Figs. 4i and 2), indicating that detectable H-2^a bone marrow-derived cells need not be present in order to induce Ly-49 downregulation.

A similar analysis was performed on fully allogeneic B10→B10.BR chimeras prepared by administration of 5 × 10^6 non-TCD B10 BMC to 7 Gy irradiated, NK cell-depleted B10.BR (H-2^a) mice. H-2^a mice normally do not contain brightly staining Ly-49^+ NK cells (6). Only 1.0 ± 0.7% of B10 NK cells from B10→B10.BR recipients expressed Ly-49 10 wk after BMT. However, unlike results in B10.A hosts, NK cells from B10→B10.BR chimeras showed an eventual increase in Ly-49 expression, with NK cells of three of four animals analyzed 30 wk after BMT showing percentages and intensities of Ly-49 expression that were below those detected in B10→B10 BMT recipients, but significantly greater than those detected in normal B10.BR controls. The fourth animal resembled a normal B10.BR control. Thus, the duration of Ly-49 downregulation induced by an H-2^a host environment was more prolonged than that induced by an H-2^k host environment with similar background genes, suggesting differences in the potency of different MHC molecules in mediating this effect. Studies performed in the B10→A/J (H-2^b→H-2^a) strain combination, however, showed only a partial reduction in the percentage of NK cells expressing Ly-49 and in the intensity of Ly-49 expression on B10 NK cells (data not shown), suggesting that host factors in addition to MHC may influence Ly-49 expression. Studies are in progress to identify these additional host influences.

To determine whether the level of H-2^a expressed on host tissues was critical in determining Ly-49 expression on B10

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Expression of H-2^a on radioresistant host elements leads to downregulated Ly-49 expression on NK1.1^+ cells. (a, d, and g) K^b expression on PBL of B10(-T) (i.e., TCD B10→B10, B10(-T)→B10.A, and B10(+T)) (i.e., non-TCD B10→B10.A) BMT recipients, respectively. A small population of surviving K^b→B10.A cells can be detected in PBL from B10.A recipients of TCD B10, but not of non-TCD B10 marrow (d vs g). (b, e, and h) Control staining with FITC-HOPC1 mAb for gated NK1.1^+ cells of these BMT recipients. (c, f, and i) Staining of the same gated NK1.1^+ populations with anti-Ly-49 mAb A1-FITC.
NK cells, we performed similar analyses in lethally irradiated (B10 × B10.A)F1 recipients of TCD B10 BMC. Although the percentage of Ly-49+ NK cells and the intensity of Ly-49 staining in these mice was significantly reduced compared with syngeneic controls (percentages summarized in Fig. 2; typical FCM histograms shown in Fig. 3), this reduction was much less marked than that observed for B10 NK cells developing in homozygous B10.A recipients, or for B10 NK cells in B10+(B10 × B10.A)F1→B10 BMT recipients (Figs. 2 and 3). These results suggest that a higher level of H-2 expression is required on the radioreistant host environment than on bone marrow-derived cells in order to produce a similar degree of Ly-49 downregulation on H-2+ NK cells.

Thus, the MHC phenotype of the host environment influences NK cell Ly-49 expression, but to a lesser degree than does the MHC of engrafted bone marrow. Preliminary results of functional studies using the RMA tumor in chimeras are consistent with those presented here i.e., an influence of bone marrow-derived cells and of the host nonlymphohematopoietic environment on NK cell function can be detected (our unpublished data). Radioreistant host elements that might influence NK cell development could include marrow stroma, since the normal marrow environment appears to play an important role in NK cell development (23). Although our FCM technique was sensitive to 0.1% contamination of H-2-disparate cells in a mixing study, and adoptive transfer studies from fully allogeneic chimeras in class I-mismatched donor-host combinations suggest that no host-type hematopoietic stem cells survive after lethal irradiation and non-TCD allogeneic BMT (our unpublished data), we have not ruled out the formal possibility that host marrow-derived cells below the level of detection in these assays survive in non-TCD B10→B10.A chimeras. However, we believe it is unlikely that such low levels of microchimerism within the lymphohematopoietic system, if present, would mediate the potent and durable effect we observed in non-TCD B10→B10.A chimeras. It is also formally possible that long-lived radioreistant host marrow-derived cells residing outside of the lymphohematopoietic system, such as host Langerhans cells that persist in the skin of long-term chimeras (our unpublished data), are the radioreistant host elements that influence the NK cell repertoire.

Previous studies suggested that the interaction of Ly-49 on NK cells with D4 expressed on target cells provides a negative signal to the NK cell, resulting in failure to kill the target (5). The reduced Ly-49 expression on NK cells of D4-bearing mice, and of chimeras expressing D4 on bone marrow-derived cells or radioreistant host elements, is consistent with the hypothesis that negative signaling through this interaction with D4 in vivo leads to either destruction of Ly-49+ NK cells or to downregulation of Ly-49 expression. We favor the possibility that Ly-49 expression is downregulated, as we have seen a continuum in the level of Ly-49 expression, with reduced anti-Ly-49 staining intensity in B10→(B10 × B10.A)F1, (B10 × B10.A)F1→B10.A/J, and long-term B10→B10.BR chimeras. Even lower, but nevertheless detectable, levels of Ly-49 expression were variably detected on normal B10.A, (B10 × B10.A)F1, and B10.BR NK cells, and in mixed chimeras involving H-2+ donors. Thus, it is possible that all of the differences between strains and various types of chimeras reflect varying intensities of Ly-49 expression, rather than the presence or absence of Ly-49+ NK cells.

Ly-49 is one of a family of structurally similar type II integral membrane proteins that is genetically linked to the NKR-P1 family of NK cell-associated genes on mouse chromosome 6 (14, 24, 25). If the interaction of such an NK-associated structure, or a combination of such structures on a single NK cell, results in a high affinity interaction with the target cell's class I MHC molecules, then perhaps the negative signal transduced to the NK cell could lead to downregulation of the interacting surface molecule(s) or deletion of that NK cell subset. The lack of Ly-49 expression on NK cells of D4+ mice or chimeras might reflect a very high affinity interaction between D4 and Ly-49, resulting in such a potent negative signal that high levels of Ly-49 expression do not persist. If, on the other hand, the overall affinity of these interactions were intermediate, then the less intense negative signal provided by interactions with self-class I MHC molecules might prevent the killing of autologous targets while allowing that NK cell subset to persist in vivo. These NK cells, on the other hand, would kill targets not expressing the relevant self-class I molecule(s), leading to a pattern of lysis consistent with the "missing self" hypothesis (26-28).

Another NK cell subset marker, recognized by mAb 5E6, is expressed on NK cells that resist engraftment of homozygous H-2+ but not of H-2+ marrow (29). Thus, interactions of the 5E6 molecule with MHC molecules could be postulated. Ly-49 expression is downregulated in mice expressing the MHC molecules to which it presumably can bind, whereas 5E6+ NK cells are detectable in animals expressing D4. This difference could be explained by our affinity/NK cell subset model. Since 5E6+ NK cells cannot reject H-2+ marrow, it could be postulated that the 5E6 molecule is an Ly-49-like recognition structure with intermediate affinity for D4. Expression of 5E6 would ensure that this subset of NK cells did not kill autologous targets in H-2+ × H-2+ mice, and would not reject homozygous H-2+ marrow. In contrast, 5E6- NK cells in F1 mice might rely on expression of an H-2+ binding Ly-49-like molecule to inhibit lysis of autologous targets, and would thus be unable to reject homozygous H-2+ marrow. The 5E6+ subset present in H-2+ homozygous mice, on the other hand, might express an additional Ly-49-like molecule that interacts with intermediate affinity to an H-2+ MHC molecule in order to prevent killing of its autologous, D4-negative cells. Thus, phenotypically distinguishable subsets of 5E6+ cells might be present in homozygous versus heterozygous H-2+ versus H-2+ mice. Coexpression of a combination of clonally distributed NK cell surface molecules has recently been reported to be associated with the inability of human NK cells to kill targets bearing HLA-Cw3 (4, 30).

In vivo studies indicate that NK cell phenotype is most readily controlled by MHC expressed on bone marrow-derived cells, but can also be regulated by radioreistant host
cells. Thus, NK cells resemble T cells in that both bone marrow-derived and radioresistant host elements determine the presence or absence of cells with certain MHC recognition structures, although bone marrow-derived cells have a more potent ability to ensure that mature cells with host-reactive receptors are not released (31, 32). This negative selection of T cells is mediated by a process of cell death, whereas it remains possible that negative selection of Ly-49–bearing NK cells is not due to permanent deletion but to dynamic regulation of Ly-49 expression by another mechanism, such as decreased transcription, translation or post-translational processing. Studies currently in progress should help to further clarify the mechanism and site of NK cell adaptive differentiation.

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