The Ly-49D Receptor Activates Murine Natural Killer Cells
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Summary
Proteins encoded by members of the Ly-49 gene family are predominantly expressed on murine natural killer (NK) cells. Several members of this gene family have been demonstrated to inhibit NK cell lysis upon recognizing their class I ligands on target cells. In this report, we present data supporting that not all Ly-49 proteins inhibit NK cell function. Our laboratory has generated and characterized a monoclonal antibody (mAb) (12A8) that can be used to recognize the Ly-49D subset of murine NK cells. Transfection of Cos-7 cells with known members of the Ly-49 gene family revealed that 12A8 recognizes Ly-49D, but also cross-reacts with the Ly-49A protein on B6 NK cells. In addition, 12A8 demonstrates reactivity by both immunoprecipitation and two-color flow cytometry analysis with an NK cell subset that is distinct from those expressing Ly-49A, C, or G2. An Ly-49D+ subset of NK cells that did not express Ly-49A, C, and G2 was isolated and examined for their functional capabilities. Tumor targets and concanavalin A (ConA) lymphoblasts from a variety of H2 haplotypes were examined for their susceptibility to lysis by Ly-49D+ NK cells. None of the major histocompatibility complex class I–bearing targets inhibited lysis of Ly-49D+ NK cells. More importantly, we demonstrate that the addition of mAb 12A8 to Ly-49D+ NK cells can augment lysis of FcγR+ target cells in a reverse antibody-dependent cellular cytotoxicity–type assay and induces apoptosis in Ly-49D+ NK cells. Furthermore, the cytoplasmic domain of Ly-49D does not contain the V/Ix-YxxL immunoreceptor tyrosine-based inhibitory motif found in Ly-49A, C, or G2 that has been characterized in the human p58 killer inhibitory receptors. Therefore, Ly-49D is the first member of the Ly-49 family characterized as transmitting positive signals to NK cells, rather than inhibiting NK cell function.

Members of the Ly-49 gene family encode type II integral transmembrane proteins and are primarily expressed on the surface of murine NK cells. Several members of the Ly-49 family of proteins can bind to class I MHC and transmit inhibitory signals to NK cells. When expressed on target cells, selected class I proteins can prevent NK cells from delivering their lethal hit. Recognition of class I molecules by Ly-49+ NK cells has been proposed as a regulatory mechanism to prevent lysis of normal host cells. However, NK cell lysis can proceed upon downregulation of host class I after transformation or viral infection (1). Recent studies have identified eight Ly-49 gene family members in NK cells from B6 mice (2, 3). The prototypic member of the Ly-49 family, Ly-49A, has been shown to recognize the class I molecules H-2Dd and H-2Dk (4, 5). The interaction of Ly-49A with H-2Dd has been postulated to transmit a negative signal to the NK cell. This hypothesis has been formulated because Ly-49A+ NK cells are apparently not capable of mediating antibody-dependent cellular cytotoxicity or lectin-induced cytotoxicity against H-2Dd–expressing target cells. Upon addition of mAb A1, which recognizes Ly-49A+ NK cells (6), enhanced lysis of target cells that is not FcR dependent is observed (4). Studies have also shown that Ly-49A can recognize carbohydrate expressed on the surface of target cells, which may contribute to the interaction of Ly-49A and class I proteins (7).

The Ly-49G2 subset of NK cells also has been shown to be inhibited by target cells expressing H-2Dd and/or H-2Ld (8). Studies with Ly-49G2+ NK cells have relied primarily on the reversal of target cell inhibition by mAb 4D11 (anti–Ly-49G2) and mAb specific for H-2Dd and H-2Ld.
The Ly-49C⁺ subset of NK cells has been shown to bind the class I molecules of the H-2b, H-2d, H-2k, and H-2 haplotypes (9). Recent data from Yu et al. (10) demonstrates that Ly-49C⁺ NK cells from BALB/c and BALB.B mice are inhibited by H-2d and H-2k class I antigens. The authors in this study concluded, however, that not all Ly-49C⁺ NK cells function the same way in all mouse strains, and suggest that allelic differences may regulate class I recognition by these cells (10). Previous data by members of this group have shown that 5E6⁺ NK cells can reject bone marrow grafts expressing H-2b but not H-2b (11). These results suggest that Ly-49C binding to its H-2d ligand may not be inhibitory in the strains studied. In H-2d strain mice, Ly-49C⁺ NK cells may be responsible for promoting hematopoiesis through the upregulation of GM-CSF, as demonstrated by Murphy et al. (12), implying further that some Ly-49 family members may upregulate NK cell function.

The Ly-49 gene family now consists of eight distinct molecules in a single inbred strain. The original Ly-49 gene has been renamed Ly-49A, and the others have been designated Ly-49B-H (2, 3, 9). mAb specific for the Ly-49A, C, and G2 molecules have helped provide significant information on their functional attributes. Functional characterization of other Ly-49 family members has been hampered by the lack of antibodies that specifically recognize each molecule. Ly-49D is of particular interest because it has a cytoplasmic domain that is significantly different from other Ly-49 family members. Ly-49D is of particular interest because it has a cytoplasmic domain that is significantly different from other Ly-49 family members. The original Ly-49 gene family may upregulate NK cell function.

Materials and Methods

Mice and Rats. All mice and rats were obtained from the Animal Production Facility, FCRDC. Mice were between 8 and 20 wk old when killed.

Generation of mAb 12A8. IL-2–cultured B6 NK cells were used to immunize Fischer 344 rats. Approximately 5–10 × 10⁶ cells were injected subcutaneously into rats and boosted every 2 wk for a total of four boosts. After the final boost, the rat spleen was harvested and spleen cells were fused with the SP2/0 hybridoma by PEG. Supernatants were first screened for rat IgG by ELISA, and G2 molecules have helped provide significant information on their functional attributes. Functional characterization of other Ly-49 family members has been hampered by the lack of antibodies that specifically recognize each molecule. Ly-49D is of particular interest because it has a cytoplasmic domain that is significantly different from other Ly-49 family members. The original Ly-49 gene family may upregulate NK cell function.

Flow Cytometry Analysis. NK cells were stained as previously described and were analyzed on either a FACScan® or FACSsort® (Becton Dickinson & Co., Mountain View, CA). Cell sorting was performed on either an Epics 750 (Coulter Electronics, Hialeah, FL) or a FACSsort® (Becton Dickinson). NK cells were either stained with the primary antibody followed by an isotype-specific, FITC-conjugated secondary reagent, or directly with a FITC-labeled primary antibody. Cells sorted for mAb 12A8 were stained with biotinylated 12A8, followed by Streptavidin PE (Becton Dickinson). Antibodies used in these experiments included 2.4G2 (FcyRII/III), PK136 (NK-1.1), A1 (Ly-49A), SW5E6 (Ly-49C), and 4D11 (Ly-49G2). All antibodies were semi-purified by salt fractionation (2X) of ascites fluid. RM21 antibody against murine CD2 was affinity purified before use.

NK Cell Isolation. Murine splenic NK cells were isolated as described previously (13). Essentically, nylon wool–nonadherent (NWNAD)⁺ cells were obtained and subjected to mAb plus complement (mAb + c⁻) depletion of T cells using anti-CD4 and CD8 mAbs. Partial depletion of Ly-49A, C, and G2⁺ NK cells was obtained by including mAb SW5E6 and 4D11 to the cocktail of anti-CD4 and CD8 mAbs. Lysis of NK cells with mAb 5E6 and 4D11 accomplished two goals: the first was to reduce the number of NK cells expressing known Ly-49 receptors such as C and G2 (and coincidentally, those cells coexpressing Ly-49A), and the second was to enrich the NK cell population for cells expressing 12A8 (Ly-49D) to make sorting these cells less time consuming. Treatment of NWNAD cells with this cocktail resulted in an NK cell population that was enriched for 12A8⁺ (Ly-49D⁺) NK cells, but generally <10% Ly-49A, C, and G2⁺. The remaining B cells were further removed by immunoabsorbance to anti–mouse IgG–coated petri plates. After culture for 6–12 d in 1,000 U/ml IL-2, the resulting cell populations were >95% NK-1.1⁻.

Tumor Targets and Con A Blasts. Tumor targets were maintained in culture as previously described (13). Splenic Con A lymphoblasts were essentially prepared by the method of Chadwick and Miller (14). Blasts were prepared and frozen at ~70°C before labeling.

Cytotoxicity Assays. Tumor targets and Con A lymphoblasts were labeled with ⁵¹Cr and used in 4-h cytotoxicity assays, as described previously (13). Assays involving mAb included the specific antibodies at a concentration of ~2 µg/well for the duration of the cytotoxicity assay. Data are either presented as lytic units per 10⁷ cells or the percent of specific lysis.

Immunoprecipitation. IL-2–cultured NK cells were ³²P-labeled with lactoperoxidase and lysed in 0.5% Triton X-100 containing protease inhibitors, as described previously (13). A single lysate was subjected to sequential immunoprecipitation using mAb 12A8, followed by the live cell ELISA assay against IL-2–propagated B6 NK cells, as previously described (13). mAb 12A8 was isolated and the hybridoma was cloned by limiting dilution. mAb 12A8 is of the rat IgG2a isotype.

Transfection of Cos-7. cDNA plasmids encoding Ly-49A, D, E, F, G1, and G2 were initially transfected into Cos-7 cells using the DEAE-dextran method (15). Cells were harvested with versene (GIBCO BRL, Gaithersburg, MD) 3 d after transfection and were analyzed by flow cytometry analysis (FCA) for binding of mAb 12A8. Reactivity was confirmed as shown in Fig. 2 by using Cos-7 cells transfected with Lipofectamine (Life Technologies, Gaithersburg, MD). A 1:60 dilution of Lipofectamine and 5 µg of Ly-49 plasmid cDNA was used according to the manufacturer’s instructions. Transfection was performed for 6 h at 37°C.
followed by washing, trypsinization, and replating cells in a T-150 flask. After 72 h of culture at 37°C, the cells were removed using HBSS without Ca\(^{2+}\) and Mg\(^{2+}\) and with 1 mM EDTA. FCA was performed as described previously.

**Apoptosis.** Highly enriched populations of NK cells were obtained from C57BL/6 splenocytes as described previously. The methods used to measure apoptosis have been developed by Ortaldo et al. (15a). 10\(^6\) NK cells were cultured for 24 h in RPMI 1640 + 10% FBS + 10\(^3\) U/ml IL-2 in 24-well plates. NK cells were collected, and viable cells were obtained after passage over Lympholyte-M (Cedarlane Labs., Ontario, Canada). Cells were then cultured in 48-well plates at 10\(^6\) cells/well, to which a variety of mAb were added, including A1, 12A8, 4D11, 5E6, and PK136 at a concentration of 40 \(\mu\)g/well in a total volume of 100 \(\mu\)l. After a 15-min incubation at room temperature, 40 \(\mu\)g of a secondary goat anti–mouse or anti–rat antibody was added to each well. Plates were incubated at 37°C for 6 and 12 h. NK cells were harvested and stained with propidium iodide for FCA. Flow cytometry was performed on a FACSort\textsuperscript{TM} using Lysis II software (Becton Dickinson), which analyzed the percentage of cells incorporating propidium iodide vs. forward scatter.

**Results**

**mAb 12A8 Identifies a Unique Subset of NK Cells.** mAb 12A8 was generated by fusing spleen cells from Fischer 344 rats that had been immunized with C57BL/6–IL-2–cultured NK cells, with the SP2/0 myeloma. Screening of lymphoid cells by FCA revealed that mAb 12A8 did not react with thymocytes, T cells, or B cells from B6 mice (data not shown). mAb 12A8 did react, however, with a unique subset of B6 NK cells. Fig. 1 A shows the staining profile of mAb 12A8 on highly enriched populations of freshly isolated NK cells. Two-color FCA of 12A8 vs. NK–1.1 demonstrated that virtually all 12A8\(^+\) cells were NK–1.1\(^+\), and that 12A8\(^+\) cells represented ~63% of the NK–1.1\(^+\) cells. The staining profiles and reactivity of mAb 12A8 on freshly isolated and IL-2–cultured NK cells indicated that this antibody may be similar to the previously characterized mAbs, 4D11 (13) and 5E6 (11). Two-color FCA was used to examine this possibility. Fig. 1 presents the staining profiles of mAb 12A8 vs mAb A1 (Ly–49A; Fig. 1 B), mAb 4D11 (Ly–49G2; Fig. 1 C), and mAb 5E6 (Ly–49C; Fig. 1 D). These staining profiles revealed that mAb 12A8 reacted with a unique subset of B6 NK cells, and that it did not demonstrate reactivity similar to mAb 5E6 or 4D11. However, an interesting staining profile was observed on NK cells when costained with mAb 12A8 vs. A1 (Fig. 1 B). It was apparent that all cells that stained with mAb A1 were also 12A8\(^+\). One explanation for the expression of mAb

![Figure 1](mAb 12A8 identifies a distinct subset of C57BL/6 NK cells. Freshly isolated C57BL/6 splenic NK cells were enriched by passage over nylon wool columns, depleted of T and B cells, and prepared for FCA as follows: (A) PK136 plus biotinylated 12A8 followed by FITC–goat anti–mouse IgG2a and Streptavidin-PE, (B) FITC-A1 plus biotinylated 12A8 followed by Streptavidin PE, (C) FITC-4D11 plus biotinylated 12A8 followed by Streptavidin PE, and (D) FITC-5E6 plus biotinylated 12A8 followed by Streptavidin PE.)
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12A8 on all A1\(^+\) cells could simply be that all A1\(^+\) cells express both Ly-49A and another molecule recognized by mAb 12A8. An alternative explanation would be that mAb 12A8 cross-reacts with both Ly-49A and another highly homologous member of the Ly-49 gene family. Blocking studies were performed with mAbs 12A8 and A1 to determine if binding of one mAb could block the reactivity of the alternate mAb. Binding of mAb 12A8 to NK cells does reduce the staining intensity (mean channel fluorescence) of FITC A1; however, mAb A1 does not block binding of FITC 12A8 (data not shown). Control blocking studies revealed that mAb 12A8 did not affect staining by mAb 4D11 or 5E6. The results of our FCA data suggested that mAb 12A8 identified a unique molecule on the surface of NK cells with possible cross-reactivity to Ly-49A.

mAb 12A8 Immunoprecipitates a 50-kD Homodimer on NK Cells. Considering the unique phenotypic characterization of the 12A8\(^+\) subset of NK cells and its reactivity with Ly-49D, we decided to examine the biochemical characteristics of the antigen recognized by this antibody. IL-2–cultured NK cells from B6 mice were surface labeled with \(^{125}\)I, lysed, and subjected to immunoprecipitation with an mAb linked to protein G–Sepharose. Preliminary experiments demonstrated that mAb 12A8 immunoprecipitated a glycoprotein of \(~100\) kD (nonreduced) and \(~50\) kD (reduced). To confirm our transfection data and to determine if mAb 12A8 could immunoprecipitate a unique Ly-49 glycoprotein, we performed sequential immunoprecipitation (IP). A single lysate from \(^{125}\)I-labeled NK cells was subjected to IP, first with an isotype control mAb, followed by mAbs A1, 5E6, 4D11, and 12A8, respectively. IP with mAb A1 was performed with an excess of protein G–Sepharose–
lymphocytes, and Ly-49C and Ly-49D cells. mAb 12A8 recognizes a unique glycoprotein on B6 NK cells. Data on both NK cells and transfected Cos-7 cells, that minimally higher than those seen with mAb A1.5E6 IP, resulted in bands under both nonreducing and reducing conditions that were consistent with previously published reports. mAb 12A8 immunoprecipitation yielded proteins with noticeably higher molecular weight bands than those seen with mAb 4D11 and pE6, respectively. Approximately equal CPMs were applied to a 10% SDS-PAGE gel and electrophoresed under both nonreduced (A) and reduced (B) conditions.

Figure 3. Immunoprecipitation using mAb 12A8 reveals a unique disulfide-linked homodimer. B6 NK cells were cultured for 7 d in high dose IL-2. Cells were radiolabeled with 125I and lysed in 0.5% Triton X-100. Sequential IP was performed on a single lysate by the following mAb cross-linked to protein G-Sepharose: Control rat IgG2A, A1, 5E6, 4D11, and 12A8, respectively. Data from these IP experiments are presented in Figure 3A and B, respectively. Coexpression of Ly-49A on NK cells, Con A blasts of varied H-2 haplotypes. In further attempts to establish any possible class I restriction of the lytic capacity of Ly-49D+ NK cells, Con A lymphoblasts from seven different H-2 haplotypes were generated. Ly-49D+ cells were isolated as described above and expanded in IL-2 for 6-10 d. Splenic Con A lymphoblasts were generated for 48 h and labeled with either 51Cr or 111In, and were used as targets in a 4-h cytotoxicity assay.

The purified NK cell subsets were examined for their cytolytic properties against various class I target cells. A panel of tumor target cells was examined for susceptibility to lysis by Ly-49D+/A− and Ly-49D−/A− NK cells. Table 1 presents the lytic data on the ability of the input population (Ly-49A, C, and G2 reduced), Ly-49D+/A−, and Ly-49D−/A− NK cells to lyse tumor target cells expressing different class I phenotypes. It is apparent from this data that Ly-49D+/A− NK cells are capable of lysing tumor target cells expressing H-2a, H-2b, H-2d, and H-2k. Lysis by Ly-49D+ NK cells was equivalent to, if not greater than, Ly-49D−/A− NK cells and the input population. These results indicate that lytic activity of Ly-49D+ cells is not inhibited by these class I-bearing tumor targets. Previous studies in our laboratory have demonstrated that both P815 and WEHI-164 (H-2d targets) are not lysed efficiently by Ly-49A+ or G2+ NK cells when compared to the Ly-49A− and G2− subsets (data not shown).

Table 1. Ly-49D+ NK Cells Lyse Tumor Targets That Express Different Class I Phenotypes

<table>
<thead>
<tr>
<th>Target</th>
<th>Ly-49D+/A− (%)</th>
<th>Ly-49D−/A− (%)</th>
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<tbody>
<tr>
<td>YAC H2a</td>
<td>11%</td>
<td>1%</td>
</tr>
<tr>
<td>P815 H2d</td>
<td>12%</td>
<td>4%</td>
</tr>
<tr>
<td>EL-4 H2b</td>
<td>5%</td>
<td>4%</td>
</tr>
<tr>
<td>SST H2k</td>
<td>38%</td>
<td>77%</td>
</tr>
<tr>
<td>WEHI-164 H2d</td>
<td>7%</td>
<td>4%</td>
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*Results are presented as lytic units at 30% per 10⁶ cells. The data are from one of two similar experiments performed.
pressing H-2a, H-2b, H-2d, H-2k, H-2s, and H-2q are all lysed by Ly-49D+ NK cells. In general, the patterns of lysis observed indicated that Ly-49D+ NK cells are comparable to both the Ly-49D− and input cells against the majority of the blast targets. These results support the lytic data using tumor targets that Ly-49D+ NK cells do not appear to be inhibited by any of the class I-expressing target cells tested to date. One conclusion from this data is that the glycoprotein encoded by the Ly-49 gene may not interact with class I MHC. If recognition does occur between Ly-49D-encoded proteins and class I, signals that inhibit NK lysis are not generated.

Ly-49D+ NK Cells Mediate Reverse Antibody-dependent Cellular Cytotoxicity (RADCC) in the Presence of mAb 12A8. Since Ly-49D+ NK cells failed to exhibit any pattern of class I restriction, we decided to examine mAb 12A8 for its ability to activate the lytic potential of Ly-49D+ NK cells. A panel of different tumor target cells, including both FcγRII/III (17), and can be blocked by the addition of mAb 2.4G2, which is specific for FcγRII/III (18). As a further control in these experiments, it can be seen that mAb PK136 is capable of augmenting the lysis of both the Ly-49D+ and D− subsets (>95% NK-1.1+) against the FcγRI targets P815 and WEHI-3. The Ly-49D− subset, however, is not augmented in the presence of mAb 12A8 to lyse these same targets. These results also reveal that the augmented lysis observed in the presence of mAb 12A8 can be abrogated upon addition of mAb 2.4G2 (identical to that seen using PK136 + 2.4G2). Similar results have been obtained upon pretreatment and washing of Ly-49D+ NK cells with mAb 12A8 before performing cytotoxicity assays (data not shown). FcγRI target cells such as SST (Fig. 5 C) and WEHI 164 (Fig. 5 D) do not demonstrate any significant augmentation of lysis by Ly-49D+ NK cells in the presence of mAb 12A8. The data shown in Fig. 5 (C) shows that mAb 12A8 weakly augments killing by Ly-49D+ NK cells against the SST target. This finding has not been reproducible in other experiments using this target and was not considered significant. Based on the above experiments, we conclude that mAb 12A8 engagement of the Ly-49D molecule augments the lytic function of NK cells. These results establish for the first time that a member of the Ly-49 gene family may upregulate NK lytic function upon binding the appropriate ligand.

mAb 12A8 Induces Apoptosis of Ly-49D+ NK Cells. Since mAb 12A8 to mediate RADCC in Ly-49D+ NK cells presents functional data for involvement of the Ly-49D molecule in activating the lytic mechanism of NK cells. We also have obtained evidence that the Ly-49D molecule may be involved in triggering apoptotic events in NK cells that lead to cell death. The results shown in Fig. 6 demonstrate that mAb 12A8 induces apoptosis in ~40% of bulk NK cells by 6 h. Although some variability was observed in different experiments, none of the other Ly-49-specific antibodies induced any consistent or potent apoptosis. Since mAb 12A8 stains ~60% of bulk NK cells, of which up to 20% may represent cross-reacting Ly-49A+ cells, it appears that most Ly-49D+ NK cells are susceptible to mAb 12A8–induced apoptosis. Of interest, mAb PK136 that recognizes the NK-1.1 antigen (mouse NKR-P1) was not very effective at inducing apoptosis. Only 10–12% of NK cells demonstrated PI uptake after the addition of PK136. These results support the hypothesis that Ly-49D represents an activation molecule on NK cells. Although...
Ly-49A, C, and G2 have been demonstrated to be class I inhibitory receptors on NK cells, cross-linking of their specific receptors does not induce apoptotic associated events in NK cells. Therefore, intracellular signaling events mediated by Ly-49D appear to be different from those mediated by Ly-49A, C, and G2. Apoptotic-associated events have also been observed using the CellFit program (Becton Dickinson) to measure the DNA index of murine NK cells after appropriate stimulation and cross-linking of the Ly-49 mAb. Using this methodology, we have observed increases in abnormal DNA content in NK cells treated with mAb 12A8, but not with 5E6 or 4D11 (data not shown).

Discussion

Eight distinct members of the Ly-49 gene family have been identified (Ly-49A–H), although specific mAbs that recognize Ly-49A (A1/YE1-32 and 48), Ly-49C (5E6), and Ly-49G2 (4D11) are the only ones available. The availability of these mAbs has helped to further define the functional role of NK cells that express these receptors. The importance of generating mAbs against the remaining, undefined members of the Ly-49 gene family is paramount to understanding the functional role that these molecules play in NK cell biology. Investigation of the proposed class I inhibitory properties mediated by members of the Ly-49 gene family may be facilitated by acquiring such reagents. The prototype molecule, Ly-49A, when expressed on NK cells, has been shown to bind the class I molecules H2-Dd and Dk. This recognition of class I ligand by the Ly-49A receptor has been shown to inhibit the lytic properties of Ly-49A+ NK cells. A similar recognition process has been proposed for Ly-49G2+ NK cells. Inhibition of lysis by Ly-49G2+ NK cells, however, has only been observed against selected target cells that appear to express high levels of H2-Dd and Dk. This recognition of class I ligand by the Ly-49A receptor has been shown to inhibit the lytic properties of Ly-49A+ NK cells. A similar recognition process has been proposed for Ly-49G2+ NK cells. Inhibition of lysis by Ly-49G2+ NK cells, however, has only been observed against selected target cells that appear to express high levels of H2-Dd and/or Ld (8, 18). Experiments to determine if Ly-49G2 specifically binds to H2-Dd and/or Ld are now in progress. Brennan et al. (9) have observed binding of Ly-
49C+ cells to multiple H-2 haplotypes, including H-2d, H-2k, H-2a, and H-2b. Furthermore, Ly-49C+ NK cells have been demonstrated to be the subset of NK cells that is responsible for rejecting H-2d, but not H-2b, bone marrow allografts (11). These experiments imply that Ly-49C+ NK cells also may be capable of upregulating NK cell function upon appropriate receptor/ligand interaction. A recent report by Stoneman et al. (19) demonstrates that glycosylation of the Ly-49C core protein varies in different mouse strains. They have implied that the extent to which the Ly-49C molecule is glycosylated may explain the functional discrepancies observed by Ly-49C+ NK cells from different mouse strains. They have implied that the extent to which the Ly-49C molecule is glycosylated may explain the functional discrepancies observed by Ly-49C+ NK cells from different mouse strains. Of further interest is the recent data from Yu et al. that describes the role of Ly-49A and C+ NK cells in mediating hybrid resistance (10). These authors conclude that class I inhibition of Ly-49A and C+ NK cells support the “missing self” and not the “hemopoietic histocompatibility antigen” hypothesis for hybrid resistance. The data from this group also suggested that allelic differences in Ly-49C+ NK cells may control which class I molecules inhibit their lytic function.

Our laboratory has generated an mAb designated 12A8 that recognizes the Ly-49D molecule expressed on the surface of B6 NK cells. Functional analysis of Ly-49D+ NK cells demonstrate that they do not appear to be restricted in their lytic capacity against either tumor targets or Con A blasts of different H-2 haplotypes. More significantly, we have shown that the protein receptor encoded by Ly-49D is capable of enhancing the lytic potential of NK cells. In the presence of FcyR+ target cells, mAb 12A8 triggers Ly-49D+ NK cells to lyse these targets in a RADCC fashion similar to that which occurs with the mAb PK136 against the NK-1.1 (NKR-P1) molecule (17). Furthermore, cross-linking of mAb 12A8 on the surface of NK cells using a secondary anti-rat antibody induces IL-2–activated NK cells to undergo apoptotic events resulting in programmed cell death. Studies with both T cells (20, 21) and human NK cells (22) have revealed that IL-2–activated cells can undergo apoptosis when stimulated through activating surface receptors such as CD3 and CD16, respectively. The significance of this finding is that mAb specific for the Ly-49A, C, and G2 proteins did not induce positive signaling events leading to apoptosis. Therefore, our data provide strong evidence that Ly-49D is not an inhibitory receptor on NK cells as are Ly-49A, C, and G2. Ly-49D appears capable of enhancing lytic function, and possibly signals NK cells through alternate pathways.

A question arises as to whether or not the observed RADCC results obtained with mAb 12A8 on Ly-49D+ NK cells could be an intrinsic function of the mAb itself, rather than that of the Ly-49D receptor. It would be tempting to determine if Ly-49A+ NK cells could be induced to mediate RADCC by mAb 12A8 because of its cross-reactivity. The results of such an experiment could not be interpreted, however, because the percentage of Ly-49A+ NK cells that coexpress Ly-49D is unknown. Until an antibody is produced that specifically reacts with Ly-49D, we must at least entertain the possibility that mAb 12A8 could have some intrinsic capacity to activate Ly-49D molecules.

Because of the unique activating properties of Ly-49D+ NK cells compared to the inhibitory properties observed in Ly-49A, C, and G2+ NK cells, we compared the amino acid homology of these proteins. Fig. 7 A is a schematic comparison of Ly-49D vs. Ly-49A, C, and G2, based on the proposed amino acid sequence of these molecules (3). There is obviously a high degree of homology between Ly-49D and A that is highest in the extracellular domain (86%) and least homologous in the cytoplasmic domain (51%). The structural homology between Ly-49A and D in their extracellular domains probably accounts for the cross-reactivity that is observed with mAb 12A8, since it can bind to both Ly-49D and Ly-49A. Furthermore, we can also speculate that the lack of homology in the cytoplasmic region of these molecules accounts for the different signaling properties of these proteins. The cytoplasmic domains of Ly-49A, C, and G2 all share at least two potential serine/threonine phosphorylation sites (P). These three molecules also share similar sites for tyrosine phosphorylation (Y) in their intracellular domains. A significant divergence in phosphorylation motifs is seen in the cytoplasmic domain of Ly-49D. Ly-49D contains only one potential serine/threonine phosphorylation motif, and none of the tyrosine phosphorylation motifs found in either Ly-49A, C, or G2. Fig. 7 B presents a 13–amino acid consensus sequence comparing the Ly-49A, C, G2 and D molecules with a similar cytoplasmic region of the human p58 inhibitory receptors. Long et al. (23) have recently described a V/IxYxxL consensus sequence in p58 proteins that may represent an immunoreceptor tyrosine-based inhibitory motif (ITIM). This V/IxYxxL motif is present in the Ly-49 inhibitory receptors A, C, and G2. However, the Ly-49D receptor does not contain this ITIM motif, which further supports our functional data that Ly-49D is an activation receptor on murine NK cells.

Figure 6. mAb 12A8 induces apoptosis in Ly-49D+ NK cells. This graph presents the data obtained from a representative experiment in which percent cell death (PI uptake) is calculated after 6 and 12 h of antibody treatment. Cell death is calculated over that observed with media alone (33, 35 & 38% respectively). Maximum cell death was observed after 6 h of treatment with mAb 12A8 and cross-linking. This is a representative experiment of eight similar experiments performed.
In summary, the present study describes the first member of the Ly-49 family of NK receptors that appears to positively regulate NK function, rather than to inhibit it. The detailed structure/function relationship of the ITIMs of Ly-49 family members will greatly aid in the understanding of the mechanism of action of these novel NK cell receptors.

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