Sustained NKG2D engagement induces cross-tolerance of multiple distinct NK cell activation pathways

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NKG2D is a multisubunit activation receptor that allows natural killer (NK) cells to detect and eliminate stressed, infected, and transformed host cells. However, the chronic exposure of NK cells to cell-bound NKG2D ligands has been shown to impair NKG2D function both in vitro and in vivo. Here we have tested whether continuous NKG2D engagement selectively impacted NKG2D function or whether heterologous NK cell activation pathways were also affected. We found that sustained NKG2D engagement induced cross-tolerance of several unrelated NK cell activation receptors. We show that receptors that activate NK cells via the DAP12/KARAP and DAP10 signaling adaptors, such as murine NKG2D and Ly49D, cross-tolerize preferentially NK cell activation pathways that function independent of DAP10/12, such as antibody-dependent cell-mediated cytotoxicity and missing-self recognition. Conversely, DAP10/12-independent pathways are unable to cross-tolerize unrelated NK cell activation receptors such as NKG2D or Ly49D. These data define a class of NK cell activation receptors that can tolerate mature NK cells. The reversible suppression of the NK cells’ cytolytic function probably reduces the NK cells’ efficacy to control endogenous and exogenous stress yet may be needed to limit tissue damage. (Blood. 2008;111:3571-3578) © 2008 by The American Society of Hematology

Introduction

Using a number of distinct recognition strategies and receptors, natural killer (NK) cells can detect host cells that are infected or that undergo malignant transformation. A limited number of NK cell receptors in the mouse directly recognize “nonself” ligands expressed on infected cells. Ly49H recognizes mouse cytomegalovirus (MCMV) m157,1 and NKp46 has been reported to recognize influenza virus hemagglutinin.2 Further, Ly49D recognizes xenogeneic major histocompatibility complex (MHC) class I molecules.3

In general, however, NK cells react to alterations in the expression of endogenous “self” ligands. Accordingly, a second set of activation receptors mediates “induced-self” recognition, that is, NK cell reactions to host cells that (over-) express specific endogenous self ligands due to infection or stress. Induced-self receptors include NKG2D (which interacts with RAE-1α-ε, H60, and MULT1),4 CD28 (B7.1),5 CD226 (Necl-5, nectin-2), and CRTAM (Necl-2).6 Additional activation receptors are specific for (not completely defined) ligands that are constitutively expressed on normal, that is, noninfected, nonstressed host cells. To prevent the lysis of normal cells, NK cells express inhibitory receptors, many of which are specific for MHC class I (MHC-I) molecules.7,9 Viral infection or transformation frequently leads to the down-regulation of MHC-I molecules, which allows diseased cells to escape recognition by cytolytic T cells.10 However, MHC-I low host cells provide insufficient NK cell inhibition signals, leading to NK cell-mediated lysis. This type of NK cell reactivity is known as “missing-self” recognition.11 Inhibitory receptors implicated in missing-self recognition include several Ly49 receptors and CD94/NKG2A (interacting with MHC-Ia and Ib, respectively),12 mouse 2B4 (CD48),13 NKRP1 (Crb, Oci1),14,15 and KLRG1 (cadherin).16,17 Relevant activation receptors include NKp46.2 Finally, NK cells can recognize and kill antibody-coated host cells (also termed ADCC for antibody-dependent cell-mediated cytotoxicity) using the low-affinity receptor for the Fc portion of IgG (FcRyIII, CD16). Thus, NK cells use multiple distinct recognition strategies and a multitude of surface receptors to scan the expression of cell-surface molecules in order to identify diseased host cells.

In general, activating NK cell receptors form multisubunit receptor complexes, whereby specialized adaptor molecules transduce signals into the cell. A first set of adaptors used by NK cells signals via ITAMs (immunoreceptor tyrosine-based activation motifs); these are DAP12/ KARAP (associated with NKG2D-S, Ly49D and Ly49H),18,19 FcRy (CD16 [ADCC] and NK1.1),20,21 as well as CD3ζ and/or FcRy (NKp46).22,23 In addition, NK cells also use DAP10, which lacks an ITAM but instead contains a YINM sequence that closely resembles a motif present in costimulatory molecules such as CD28.24 In NK cells, where DAP10 is associated with NKG2D, this adaptor mediates primary activating function rather than costimulation. Upon receptor engagement, the tyrosines in the ITAM or in YINM are phosphorylated by src family kinases, which generates docking sites for Syk family kinases and PI3K/Grb2, respectively.25 This mediates calcium influx and the activation of extracellular signal-regulated kinases (ERK), which are necessary events for inducing the release of cytotoxic granules and, consequently, target cell lysis.

In contrast to normal cells, where NKG2D ligands are expressed transiently, many primary tumors and established tumor cell lines express NKG2D ligands constitutively.5 Consequently, NKG2D engagement confers efficient NK cell reactivity to NKG2D ligand—expressing tumor cells.26,27 However, the sustained exposure of NKG2D with tumor cell-bound ligand in vitro results in
surface modulation of the receptor and a gradual reduction of receptor function.\textsuperscript{28,29} Eventually, the NKG2D receptor is completely uncoupled from the mobilization of calcium influx and the excretion of cell-mediated cytosis.\textsuperscript{29} Impaired receptor function correlates with a reduced association of NKG2D with the relevant signaling adaptors DAP10 and DAP12.\textsuperscript{29} The enforced constitutive expression of NKG2D ligands (Rae-1a, Rae-1e or MICA) as transgenes in mice has confirmed that sustained encounter of NKG2D ligands impairs NKG2D functions in vivo.\textsuperscript{30,32}

As there is evidence for functional cross-talk between certain NK cell activation receptors,\textsuperscript{33} we have addressed whether prolonged NKG2D engagement selectively impairs NKG2D function or whether unrelated NK cell activation receptors or pathways are also affected. We find that sustained NKG2D engagement cross-tolerizes several unrelated NK cell activation receptors. Indeed, DAP10/12-dependent NK cell activation pathways, such as NKG2D and Ly49D, are found to preferentially cross-tolerize DAP10/12-independent receptors or pathways. Conversely, DAP10/12-independent receptors or pathways are unable to induce cross-tolerance. These data identify a class of NK cell activation receptors that can tolerate mature NK cells, highlighting a novel adaptive feature of innate immune effector cells.

**Methods**

**Mice**

C57BL/6 (B6) (H-2\textsuperscript{b}) mice were purchased from Harlan (Horst, The Netherlands). Mice expressing truncated, nonfunctional DAP12 (DAP12ki) have been described before.\textsuperscript{34} All mice were older than 6 weeks when used for experiments.

**Cell lines and transfectants**

Several cell lines used here were described before.\textsuperscript{29} RMA m157 were generated following polyomavirus chain reaction (PCR) amplifying m157 from MCMV DNA and cloning of the product into a pEF BOS IRES puro expression vector. Stable transfectants were established after electroporation and selection with puromycin. The Chinese hamster ovary (CHO) Pro5 cell line was kindly provided by Wayne Yokoyama, Washington University, St Louis, MO.

**NK cell culture**

NK cell culture conditions and chronic exposure to tumor cells has been described.\textsuperscript{29} Briefly, B6 splenocytes were depleted of B and T cells using nylon wool and mAb 17A2 (anti-CD3e) followed by anti-\textsuperscript{rat} IgG M450 Dynabeads (Dynal, Oslo, Norway). Carboxyfluorescein diacetate succinimidyl ester (CFSE, Molecular Probes, Eugene, OR)–labeled and irradiated (3000 rad) tumor cells (Dynal, Oslo, Norway). Carboxyfluorescein diacetate succinimidyl ester (CFSE, Molecular Probes, Eugene, OR)–labeled and irradiated (3000 rad) tumor cells were added at a 1:1 ratio to NK cells (10\textsuperscript{6}/mL). After 3 days of coculture in the presence of interleukin-2 (IL-2; 500 ng/mL), the remaining tumor cells were exposed NK cells to RMA tumor cells that were transfected with the NKG2D ligand H60 (Table 1). The coculture of NK cells with parental RMA cells had no effect on NKG2D function as compared with NK cells that were cultured alone (data not shown). In contrast, the exposure of NK cells to RMA-H60 tumor cells almost completely blunted NKG2D function (Figure 1A). Thus, NK cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules. RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules.

**Cytotoxicity assays**

Standard (4 hour) \textsuperscript{51}Cr release assays were performed as described before.\textsuperscript{29} Immunoprecipitation and Western blot

**Results**

Sustained NKG2D engagement impairs multiple NK cell activation pathways

To probe the impact of prolonged NKG2D engagement, we exposed NK cells to RMA tumor cells that were transfected with the NKG2D ligand H60 (Table 1). The coculture of NK cells with parental RMA cells had no effect on NKG2D function as compared with NK cells that were cultured alone (data not shown). In contrast, the exposure of NK cells to RMA-H60 tumor cells almost completely blunted NKG2D function (Figure 1A). This effect required cell-cell contact and could not be induced with recombinant (soluble or immobilized) H60.\textsuperscript{29}

We next tested whether sustained NKG2D engagement affected the function of additional, unrelated NK cell activation pathways. To address this we used various NK cell–target cell lines (Table 1). RMA and RMA/S cells activate NK cells through unknown receptors. Unlike RMA, RMA/S cells are killed since they express low levels of MHC-I molecules and thus fail to engage inhibitory Ly49 receptors (“missing-self” recognition). Remarkably, the prolonged exposure to RMA-H60 cells prevented the NK cell–mediated lysis of RMA/S cells (Figure 1A). Coating of RMA cells with a mAb to Thy1 (\textsuperscript{aThy1}) can overcome NK cell inhibition and mediates ADCC through the engagement of the receptor for the constant portion of IgG (Fc\textsuperscript{RIII}) (CD16). Antibody-directed lysis of RMA cells was also defective when NK cells were previously exposed to

**Antibodies and flow cytometry**

One million cells were incubated with mAb 2.4G2 (anti-CDF16/32) to reduce background before staining with the following mAbs (obtained from BD PharMingen [San Diego, CA] unless indicated otherwise): PK136 (anti-NK1.1), anti-NKG2D (R&D Systems, Oxon, United Kingdom), 4E5 (anti-Ly49D), 3D10 (anti-Ly49H kindly provided by W. Yokoyama), 2.4G2 (anti-CDF16/32, LIR1). Rat anti-NKG2D and NKp46 antisera were generated using soluble NKG2D and NKp46 as the immunogen, respectively.\textsuperscript{35} Intracellular interferon (IFN\textgamma) detection and calcium ([Ca\textsuperscript{2\textplus}]\textsuperscript{2+}) mobilization was determined as described.\textsuperscript{29}
RMA-H60 cells (Figure 1A). In contrast, we observed significant, albeit reduced, lysis of RMA cells that express the MCMV m157 protein (Figure 1A). While Ly49H-independent lysis was impaired, Ly49H-dependent lysis of RMA m157 cells was near normal (Figure 1A). This is similar to the Ly49D receptor, which also retained significant activity after prolonged NKG2D triggering.29

Table 1. Expression of activating NK cell receptors after chronic stimulation

<table>
<thead>
<tr>
<th>Stimulator cell</th>
<th>Cognate receptor</th>
<th>NK1.1</th>
<th>NKG2D</th>
<th>CD16</th>
<th>Ly49D</th>
<th>Ly49H</th>
<th>Nkp46</th>
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<tbody>
<tr>
<td>RMA*</td>
<td>Unknown</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>RMA H60†</td>
<td>NKG2D</td>
<td>95 ± 13</td>
<td>18 ± 8**</td>
<td>57 ± 14</td>
<td>93 ± 14</td>
<td>95 ± 23</td>
<td>92 ± 7</td>
</tr>
<tr>
<td>RMA+/Thy1‡</td>
<td>CD16</td>
<td>89 ± 18</td>
<td>97 ± 14</td>
<td>51 ± 17</td>
<td>106 ± 20</td>
<td>100 ± 1</td>
<td>nd††</td>
</tr>
<tr>
<td>RMA/S§</td>
<td>Unknown</td>
<td>108 ± 8</td>
<td>122 ± 17</td>
<td>95 ± 14</td>
<td>100 ± 8</td>
<td>105 ± 7</td>
<td>nd</td>
</tr>
<tr>
<td>CHO</td>
<td>Ly49D</td>
<td>64 ± 18</td>
<td>92 ± 38</td>
<td>66 ± 15</td>
<td>41 ± 10</td>
<td>85 ± 1</td>
<td>nd</td>
</tr>
<tr>
<td>RMA m157¶</td>
<td>Ly49H</td>
<td>nd</td>
<td>nd</td>
<td>nd</td>
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<td>nd</td>
<td>nd</td>
</tr>
</tbody>
</table>

*RMA cells activate NK cells through unknown activation receptors. Lysis is prevented by inhibitory Ly49 receptors specific for MHC class I molecules. Consequently, RMA cells are resistant to NK cell–mediated lysis.
†Transfection of RMA cells with H60 overcomes MHC-I–dependent inhibition due to an excess of NK cell activation signals through NKG2D engagement.
‡Coating of RMA cells with anti-Thy1 mAb (α-Thy1) results in ADCC through CD16 engagement.
§Like the parental RMA, RMA/S cells stimulate NK cells through unknown receptors. Lysis occurs since RMA/S express very low levels of MHC-I molecules such that NK cells are not inhibited. This is known as “missing-self recognition.”
¶Transfection of RMA cells with MCMV m157 overcomes MHC-I–dependent inhibition due to an excess of NK cell activation signals through Ly49H engagement.
#The mean fluorescence intensity of staining of the indicated receptor in NK cells exposed to control RMA cells was arbitrarily set to 100.
**Significant deviations from 100 are indicated in bold face.
††nd indicates not determined.
A substantial fraction (10%-15%) of chronically stimulated NK cells was intracellular IFN-γ even in the absence of any restimulation. Restimulation through NKG2D using mAb or H60-transfected RMA cells failed to further increase the fraction of iIFN-γ NK cells, while anti-Ly49D mAb or CHO cells readily increased IFN-γ production. In contrast to Ly49D, restimulation using anti-NK1.1 mAb or RMA/S cells was unable to improve IFN-γ production (Figure 1B). Thus, sustained NKG2D engagement results in an unexpectedly broad, yet nevertheless selective, dysfunction of several NK cell activation receptors.

Upon the separation of NK cells from RMA-H60 cells and further culture in the presence of IL-2 for 18 hours, NKG2D function made a full recovery (Figure 1C and Coudert et al29). Similarly, ADCC recovered significantly, albeit not completely (Figure 1C). In contrast, missing-self recognition remained defective (Figure 1C). When NK cell function was analyzed 42 hours after the removal of RMA-H60 cells, also missing-self recognition had recovered significantly, albeit not completely (Figure 1D). Thus, even though the recovery of missing self-recognition is significantly delayed, cross-tolerance eventually reverses.

Basis for defective NK cell function

NK cell-mediated target lysis is dependent on intracellular mobilization of calcium ([Ca^{2+}]_i). Upon the cross-linking of NKG2D, control RMA exposed NK cells showed a robust ([Ca^{2+}]_i) mobilization response. In contrast, ([Ca^{2+}]_i) mobilization was essentially absent in RMA-H60–exposed NK cells (Figure 2), which is in agreement with our previous observations. This result may be related in part to the 3- to 5-fold reduced NKG2D levels on RMA-H60–exposed NK cells (Table 1 and Coudert et al29). However, RMA-H60–exposed NK cells also displayed deficient ([Ca^{2+}]_i) mobilization in response to CD16 cross-linking (Figure 2). In these cases receptor expression levels were normal (Table 1 and Coudert et al29). Notably, ([Ca^{2+}]_i) mobilization was observed upon Ly49D cross-linking although the percent of responding cells was reduced (Figure 2). Thus, chronic NKG2D engagement impairs ([Ca^{2+}]_i) mobilization by multiple unrelated, but not all, NK cell activation receptors. These data, together with the above lysis and IFNγ assays, show that chronic NKG2D stimulation disrupts missing-self recognition and the function of the CD16 (ADCC), NK1.1, and NKp46 activation receptors. In contrast, the function of the Ly49D and Ly49H receptors is to a significant extent preserved.

The role of DAP12 for NKG2D-mediated NK cell inactivation

In short-term cytokine-stimulated NK cells, NKG2D signals via DAP12 and DAP10 adaptors. We addressed whether both adaptors are required for the induction of cross-tolerance. Similar to wild-type NK cells, the chronic NKG2D stimulation of NK cells that lack a functional DAP12 adaptor (DAP12ki) resulted in complete NKG2D dysfunction (Figure 3 and Coudert et al29). This indicates that DAP10 is sufficient to disable NKG2D function. In contrast, missing-self recognition and ADCC were only partly disabled when DAP12ki NK cells were chronically stimulated through NKG2D (Figure 3). This suggests that DAP10 participates in NK cell cross-tolerization. However, profound and broad NK cell inactivation via NKG2D requires that sustained NKG2D signaling occurs via both DAP10 and DAP12 adaptors.

Association of distinct NK cell activation receptors with signaling adaptors

Chronic NKG2D stimulation strongly depletes NK cells of the signaling adaptors DAP10 and DAP12 as well as CD3ζ but not of FcRγ (data not shown and Coudert et al29). According to the literature CD16 and NK1.1 are associated with FcRγ and thus function independent of DAP10 and DAP12. Nevertheless, CD16
and NK1.1 function (as well as missing-self recognition) is impaired following chronic NKG2D stimulation. Conversely, Ly49D and Ly49H, which share DAP12 with NKG2D, remain at least in part functional.

We verified some of the reported associations of activation receptors with signaling adaptors. In agreement with available data, Ly49D and Ly49H were not only associated with DAP12, but also with DAP10, albeit at lower levels (Figure 4A). Unexpectedly, cell surface expression of Ly49H was not only improved upon cotransfection of DAP12 but also upon cotransfecting DAP10 (Figure 4A). Substantial Ly49D cell surface expression was observed in the absence of adaptor cotransfection. Nevertheless, similar to Ly49H, Ly49D expression further increased upon cotransfection with either DAP12 or DAP10 (Figure 4A). Corresponding improvements of cell surface expression were not apparent for NKRP1C (NK1.1) (Figure 4A), indicating that NK1.1 does neither associate with DAP10 nor DAP12.

In agreement with these findings, endogenous DAP12 and DAP10 did not communoprecipitate with NK1.1 (or CD16 or NKp46) from lysates of (day 7) cytokine-cultured NK cells (Figure 4B). All these receptors are associated with FcRγ (Figure 4B). In contrast, endogenous FcRγ did not associate with NKG2D, Ly49D, or Ly49H. Rather, NKG2D is linked to DAP10 and to DAP12 in agreement with published data.47 Note that NKG2D, which associates with DAP-12, is expressed at low levels upon prolonged culture of NK cells in IL-2.47 In agreement with the cotransfection experiments, Ly49D and Ly49H were not only associated with DAP12 but also with DAP10, albeit at lower levels (Figure 4B; Figure S1, available on the Blood website; see the Supplemental Materials link at the top of the online article).

Consistent with an association of DAP10 with Ly49D and Ly49H, significant receptor cell surface expression was detected on NK cells from DAP12 null mice.38 While we obtained no evidence for Ly49D function when using long-term cultured ( > 6 days) NK cells from DAP12ki mice (Figure S1, in agreement with published data34,38), we noted significant residual Ly49D-dependent lysis of CHO cells when short-term (3-day) cultures of DAP12ki NK cells were used (Figure 4C). These data indicate that Ly49D is partially and transiently functional in the absence of DAP12. In contrast, CD16 function (Figure 3) and missing-self recognition (Figure 3)34,38 occurred entirely independent from DAP12 function.

Thus the NKG2D, Ly49D, and Ly49H receptors share the signaling adaptors DAP10 and DAP12 while they do not use FcRγ. Conversely, the CD16, NK1.1, and NKp46 receptors share FcRγ, but they do not associate with DAP10 or DAP12. Thus, chronic NKG2D stimulation impacts the function of the DAP10/12-independent CD16, NK1.1, and NKp46 receptors possibly downstream of FcRγ.

**Cross-tolerization is induced via specific NK cell activation pathways**

To test whether NKG2D had a unique capacity to induce broad NK cell dysfunction, we chronically stimulated mature NK cells via additional activation receptors or pathways. Sustained exposure to antibody-coated RMA cells (ADCC) prevented subsequent ADCC but left all other NK cell activation pathways unperturbed (Figure 5A). CD16 cell surface expression was reduced approximately 50% (Table 1), which is unlikely to account for a complete loss of ADCC. Prolonged exposure to RMA/S cells (missing-self recognition) had no negative effect on NK cell function. Rather, subsequent missing-self reactivity was actually somewhat increased (Figure 5B). Finally, we tested the effects of chronic Ly49D engagement. As compared with RMA controls, the exposure of purified Ly49D + NK cells to CHO led to an almost complete loss of Ly49D function (Figure 5C). Such NK cells further showed impaired NKG2D function, defective ADCC (CD16), and missing-self recognition (Figure 5C). These defects were due to chronic Ly49D triggering, as the 3 pathways were functional in Ly49D- NK cells cocultured with CHO cells (Figure 5C). Thus, broad NK cell dysfunction can be induced via the NKG2D as well as the Ly49D receptor, which both signal through DAP10 and DAP12. In contrast, DAP10/12-independent NK cell activation pathways are not competent to induce cross-tolerization.

**Discussion**

The chronic engagement of NK cells with NKG2D ligand-expressing tumor cells in vitro and in vivo results in impaired NKG2D function. Here, we showed that sustained engagement of NKG2D could cross-tolerize a set of unrelated NK cell activation receptors. This, together with a corresponding capacity of Ly49D, suggests that DAP10/12-dependent NK cell activation pathways...
the experiments using transgenic mice cannot discriminate whether the effect on missing-self recognition was due to an impact on NK cell development or whether tolerance induction occurs in functionally mature NK cells. As we used mature NK cells, we conclude that peripheral NK cells are susceptible to tolerance induction. This is of importance with respect to the possible impact of NKG2D ligand-expressing tumor cells on mature NK cells in situ.

The prolonged exposure to RMA/S cells (missing-self recognition) had no negative effect on the reactivity of mature NK cells. On the contrary, subsequent missing-self reactivity was actually somewhat more efficient (Figure 5B). A similar effect was previously reported based on in vivo experiments. A possible explanation is that the receptors that trigger cytotoxicity in response to RMA/S target cells induce the production of cytokines, which may then act in combination with IL-2 to further enhance the cytotoxic activity of NK cells. This putative feedback loop does not seem to operate when inhibitory Ly49 NK cell receptors are co-engaged by MHC-I–expressing RMA cells. This model remains to be tested.

Despite the fact that chronic missing-self recognition did not induce tolerance in vitro, our data may nevertheless be of relevance to understand NK cell tolerance in MHC-I mosaic mice and/or in MHC-I–deficient patients or mice. In the appropriate (developmental) context in vivo, the continuous exposure to MHC-deficient host cells and thus chronic NK cell activation signals (due to a failure to receive MHC-I–dependent inhibition signals) may eventually lead to NK cell tolerance and cross-tolerance induction, as described herein.

Cross-tolerization has been noted in macrophages stimulated through Toll-like receptors (TLRs). Macrophages pretreated with the TLR-4 ligand LPS (lipopolysaccharide) show no or reduced production of inflammatory cytokines in response to a second stimulation with LPS. This is referred to as LPS or endotoxin tolerance. LPS tolerance is based at least in part on the modulation of membrane proximal cytoplasmic signal transduction such as a decrease of TLR tyrosine phosphorylation. Interestingly, the initial exposure of TLR-2 to lipoteichoic acid ligand induced cross-tolerance against subsequent LPS (TLR-4) or CpG DNA (TLR-9) stimulation. These data illustrate that innate immune cells such as NK cells and macrophages do not mount invariant responses to cognate receptor stimulation. Rather, the response of such cells is influenced by prior receptor stimulation. In the case of NK cell cross-tolerance, the transient shutdown of the cytolytic response of NK cells may serve to limit tissue damage during infection.

NK cells chronically stimulated through NKG2D suffer from a number of defects. NKG2D expression is reduced, and NK cells are almost devoid of the signaling adaptors DAP10 and DAP12 (as well as CD3ζ), which can readily account for NKG2D dysfunction. Here we determined the function of NK cell activation pathways, which signal independently of DAP10 and DAP12. CD16-mediated ADCC and [Ca²⁺]i flux was defective even though NK cells had normal amounts of the CD16 signaling adaptor FcRγ. Similarly, IFNγ production and ([Ca²⁺]i) flux induced by NK1.1 cross-linking, which similarly signals via FcRγ, was also defective. These data suggest that prolonged NKG2D stimulation modulates cytoplasmic signaling downstream of FcRγ. The fact that Ly49D cross-linking still triggered ([Ca²⁺]i) flux suggested that the defect was at or upstream of Phospholipase C (PLC)γ. Indeed, Ly49D-dependent killing is impaired in PLCγ2-deficient NK cells. Overall, the observed defects of NKG2D ligand–expressed NK cells correspond well to those of NK cells lacking the 3 guanine nucleotide exchange factors Vav-1, -2, and -3. Vav-1 is required for NK cell–mediated lysis downstream of NKG2D/DAP10 and likely other receptors that activate PI3K.
Vav-2 and -3 are required for NK cell-mediated lysis induced via DAP12 and other ITAM-bearing adaptors.\textsuperscript{51} Missing-self recognition occurs independent from ITAM-bearing adaptors but requires PI3K and src family kinase activity.\textsuperscript{35,52,53} Indeed, missing-self recognition is deficient in the absence of Vav-1 and also impaired upon chronic NKG2D engagement. Conversely, Ly49D functions in the absence of Vav-1, -2, and -3\textsuperscript{51} and retains partial functionality after constitutive NKG2D triggering.\textsuperscript{29} Future investigations will thus address whether vav proteins play a role in tolerance and cross-tolerance induction in NK cells.

We have also shown that chronic NKG2D stimulation reduces the cellular size and the expansion of NK cells that are cultured in IL-2,\textsuperscript{29} suggesting a connection between NKG2D and IL-2/15 receptor signaling. In agreement with these observations, while this manuscript was under revision, Horng et al\textsuperscript{54} reported a direct link between the NKG2D-DAP10 and the IL-2/15 receptor signaling pathways in NK cells. These data, together with the data shown herein, suggest that NKG2D is connected not only with a critical cytokine receptor but also with multiple distinct NK cell activation receptors or pathways. It will thus be interesting to see whether cross-tolerance induction is related to a dysfunction of the IL-2/15 receptor.

Irrespective of the molecular basis, tumor-infiltrating NK cells have been shown to exert reduced functions of NKG2D-dependent and -independent activation pathways when tested ex vivo.\textsuperscript{4} Broad NK cell tolerance can now be accounted for by the chronic stimulation of NK cells by NKG2D ligand-expressing tumor cells. To exploit NKG2D function for tumor immunity, it will be critical to identify the factors and conditions that shift NKG2D function from NK cell activation to NK cell tolerization.

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Authorship

Contribution: J.C. designed research, performed research, and analyzed data; L.S. and F.G. performed research and analyzed data; E.V. provided reagent and discussed data; W.H. designed research and wrote the paper.

Figure 5. Selectivity of NK cell cross-tolerance induction. B6-derived NK cells were exposed to anti-Thy1 mAb (α-Thy1)–coated RMA cells to induce chronic ADCC (▲) (A) or exposed to MHC class I–deficient RMA/S cells (missing-self recognition, ⋄) (B). Ly49D− (●) and Ly49D+ NK cells (□) were exposed to CHO cells (C). As controls, NK cells were exposed to RMA cells (○) or to RMA-H60 cells (●). After 3 days of coculture residual tumor cells were removed and the cytolytic activity of NK cells toward RMA-H60, anti-Thy1 mAb (α-Thy1)–coated RMA, RMA/S, and CHO target cells was determined. The Ly49D–dependence of CHO lysis was ensured by blocking with anti-Ly49D mAb (■). Error bars are SD.
Conflict-of-interest disclosure: E.V. is a cofounder and shareholder of Innate-Pharma. The remaining authors declare no competing financial interests.

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