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KIR3DL1 Allotype-Dependent Modulation of NK Cell Immunity against Chronic Myeloid Leukemia

Kiyotaka Izumi,* Takero Shindo,* Huong Thi Ngo,* Kaori Nakayama-Hosoya,† Koshi Akahane,‡ Minori Tamai,‡ Thao T. T. Nguyen,‡ Ai Kawana-Tachikawa,† Takeshi Inukai,‡ and Akifumi Takaori-Kondo*

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ABSTRACT

Tyrosine kinase inhibitor (TKI)–treated chronic myeloid leukemia (CML) patients with increased NK cell number have a better prognosis, and thus, NK cells may suppress CML. However, the efficacy of TKIs varies for reasons yet to be fully elucidated. As NK cell activity is modulated by interactions between their killer cell Ig-like receptors (KIRs) and HLAs of target cells, the combination of their polymorphisms may have functional significance. We previously showed that allelic polymorphisms of KIR3DL1 and HLAs were associated with the prognosis of TKI-treated CML patients. In this study, we focus on differential NK cell activity modulation through KIR3DL1 allotypes. KIR3DL1 expression levels varied according to their alleles. The combination of KIR3DL1 expression level and HLA-Bw4 motifs defined NK cell activity in response to the CML-derived K562 cell line, and Ab-mediated KIR3DL1 blocking reversed this activity. The TKI dasatinib enhanced NK cell activation and cytotoxicity in a KIR3DL1 allotype-dependent manner but did not significantly decrease effector regulatory T cells, suggesting that it directly activated NK cells. Dasatinib also enhanced NK cell cytotoxicity against K562 bearing the BCR-ABL1 T315I TKI resistance–conferring mutation, depending on KIR3DL1/HLA-Bw4 allotypes. Transduction of KIR3DL1*01502 into the NK cell line NK-92 resulted in KIR3DL1 expression and suppression of NK-92 activity by HLA-B ligation, which was reversed by anti-KIR3DL1 Ab. Finally, KIR3DL1 expression levels also defined activation patterns in CML patient–derived NK cells. Our findings raise the possibility of a novel strategy to enhance antitumor NK cell immunity against CML in a KIR3DL1 allotype-dependent manner. ImmunoHorizons, 2021, 5: 687–702.

INTRODUCTION

Although tyrosine kinase inhibitors (TKIs) have greatly improved the outcome of patients with chronic myeloid leukemia (CML) (1,2), several problems remain to be solved. Some cases acquire TKI resistance through mutation, such as T315I in the ABL kinase domain (3). Long-term TKI treatment impairs the quality of life for the majority of CML patients (4), and the cost of lifelong TKI treatment is considerable (5,6). Interestingly, a minority of cases experience long-term deep molecular remission without TKI maintenance, a status designated as treatment-free remission (7). However, the probability of treatment-free remission is...
limited, and its predictive determinants are unclear. Given that several reports show positive correlations between increased NK cell number and successful discontinuation of TKIs (8,9), antitumor NK cell immunity may underlie the CML suppression. Understanding their mode of action against CML would enable safer discontinuation of TKIs and help to overcome TKI resistance. Furthermore, elucidating determinants of antitumor NK cell immunity would help to establish a novel immunotherapy, which may also be applicable to many other malignancies.

NK cell activity is modulated through interactions between several kinds of NK cell receptors and their ligands on target cells. In particular, the killer cell Ig-like receptors (KIRs) constitute a major subset, and these stimulate or inhibit NK cell activity by ligation with specific HLA class I molecules. Ligation of inhibitory NK cell receptors with their HLA class I ligands is indispensable for educating NK cells (10), after which NK cells can attack HLA-deficient cells by recognition of “missing self” (11). Inhibitory KIRs are involved in the education process along with CD94/NKG2A, which recognizes HLA-E (12). KIR genes are encoded on human chromosome 19. Each of them shows abundant allelic polymorphism (13,14), but their functional significance is largely unknown.

KIR3DL1 is one of the inhibitory KIRs, and its known ligands are HLA-A and HLA-B molecules that contain Bw4 motifs (15). HLA-B molecules are classified into Bw4 or Bw6 subtypes based on the sequence of amino acids between motifs (15). HLA-B molecules are classified into Bw4 or Bw6 ligands are HLA-A and HLA-B molecules that contain Bw4 (5,6) and have limited, and its predictive determinants are unclear. Given that several reports show positive correlations between increased NK cell number and successful discontinuation of TKIs (8,9), antitumor NK cell immunity may underlie the CML suppression. Understanding their mode of action against CML would enable safer discontinuation of TKIs and help to overcome TKI resistance. Furthermore, elucidating determinants of antitumor NK cell immunity would help to establish a novel immunotherapy, which may also be applicable to many other malignancies.

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KIR3DL1 is one of the inhibitory KIRs, and its known ligands are HLA-A and HLA-B molecules that contain Bw4 motifs (15). HLA-B molecules are classified into Bw4 or Bw6 subtypes based on the sequence of amino acids between positions 77 and 83 in the α1 helix (16,17). According to the amino acid at position 80, Bw4 motifs are further divided into Bw4-80I (isoleucine) and Bw4-80T (threonine) motifs (18). KIR3DL1 has abundant allelic polymorphism. Its alleles are classified into high, low, and null allotypes, depending on surface expression levels (19–22). In vitro experiments have revealed some residues that vary among KIR3DL1 allotypes define their affinity for ligands and surface expression levels (23). KIR3DL1 and HLA-B allotypes differentially modulate NK cell activation (24–28). Their stronger interaction avidity was associated with greater NK cell suppression (25).

Several studies have raised the possible clinical significance of interaction avidity between KIR3DL1 and HLA-B proteins (29–31). According to these reports, interactions between KIR3DL1-low and Bw4-80T and KIR3DL1-high and Bw4-80I are defined as “strong” interactions between KIR3DL1-low and Bw4-80I, KIR3DL1-high and Bw4-80T are defined as “weak,” and any pairs that contain KIR3DL1-null or Bw6 are defined as “non.” Whereas the strong interaction showed a protective effect on disease progression in HIV-infected individuals (29), the weak or noninteractions correlated with a superior treatment response in neuroblastoma patients (30) and allogeneic transplant–treated acute myeloid leukemia (AML) patients (31). However, a recent study on AML failed to confirm this (32). Although the predicted levels of HLA class I expression on pathologic cells in different diseases may explain the difference (33), these conflicting results indicate that the clinical significance of the interaction strength still requires further verification.

We previously reported that the efficacy of TKIs in CML patients was associated with allelic polymorphisms of KIRs and HLA in a Japanese cohort (34). In accordance with the previous studies of neuroblastoma and AML (30,31), noninteracting combinations of KIR3DL1 and HLA-B predicted earlier achievement of a deep molecular response. TKIs such as imatinib and dasatinib may enhance NK cell immunity by suppressing regulatory T cells (Tregs) (35,36). However, there are no data on how these drugs regulate NK cell immunity through KIR–HLA interactions. In this study, we performed functional analysis of KIR3DL1 allotypes on NK cells in vitro. In particular, we examined how dasatinib affects NK cell activity based on KIR–HLA interaction. Furthermore, we explored the potential of KIR3DL1 blockade to enhance antitumor NK cell immunity.

**MATERIALS AND METHODS**

**Cell lines**

K562 (RCB 1897), a CML-derived cell line that lacks expression of HLA class I, was purchased from RIKEN BRC Cell Bank (Tsukuba, Japan), and human NK cell line NK-92 (CRL-2407) was purchased from American Type Culture Collection (Manassas, VA). K562 cells were cultured in RPMI-1640 medium (Nacalai Tesque, Kyoto, Japan; catalog [cat.] no. 30264-85) supplemented with 10% FBS (Life Technologies, Thermo Fisher Scientific, Waltham, MA; cat. no. 10270), and 1% penicillin–streptomycin–glutamine (Nacalai Tesque; cat. no. 198-15781), 0.2 mM folic acid (Nacalai Tesque; cat. no. 13051-82), 0.1 mM 2-ME (FUJIFILM Wako Pure Chemical, Osaka, Japan; cat. no. 198-15781), 0.02 mM myo-inositol (Nacalai Tesque; cat. no. 16050122), and 0.2 mM myo-inositol (Nacalai Tesque; cat. no. 21445-95) supplemented with 12.5% FBS, 12.5% horse serum (Life Technologies; cat. no. 16050122), 0.2 mM myo-inositol (Nacalai Tesque; cat. no. 13051-82), 0.1 mM 2-ME (FUJIFILM Wako Pure Chemical, Osaka, Japan; cat. no. 13051-82), 0.02 mM folic acid (Nacalai Tesque; cat. no. 16221-62), 1% penicillin–streptomycin (FUJIFILM Wako Pure Chemical; cat. no. 168-23191) and 100 U/ml recombinant human IL-2 (Shionogi, Osaka, Japan; cat. no. 639941ID1022).

**Generation of the T315I knock-in K562 cell line**

Introduction of the T315I mutation into K562 cells was performed by homologous recombination using the CRISPR/Cas9 system as previously described (37). In brief, single guide RNA (5′-aactcgactgatagatagaacgg-3′), single-strand oligodeoxynucleotides (5′-cgttcacctcctgccgtgctcgtctcgctcgctgggtctgag-3′), capital letters indicate mutated nucleotides), and recombinant Cas9 protein (Integrated DNA Technologies, Coralville, IA; cat. no. 1074181) were electroporated into parental cells in the presence of 10 nM DNA ligase IV inhibitor SCR7 (Xcess Biosciences, San Diego, CA; cat. no. M60082-2S). To block recutting of repaired target loci, three additional silent point mutations were simultaneously introduced. Seven days after electroporation, the cells were cultured in the presence of 0.5 μM of imatinib (Selleck Chemicals, Houston, TX; cat. no. S1026). After

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FIGURE 1. Activation of KIR3DL1+ NK cells depends on KIR3DL1 expression. (A) Representative flow cytometry contour plots of KIR3DL1+ NK cells stained with Z27 and DX9 Abs. (B) The MFI of Z27 staining was compared. The difference in MFI (delta MFI) between positive and negative populations was calculated for each donor. Boxes represent interquartile range, horizontal lines in the boxes represent the median value, and whiskers indicate the minimum and the maximum values (*p < 0.05). (C) (Continued)
~3 wk incubation in the presence of 0.5 μM of imatinib, an imatinib-resistant subline was obtained. Introduction of the T31SI mutation was confirmed by direct sequence of the PCR products between intron 5 and 6 of ABL1 gene.

**Establishment of single HLA-expressing K562 cell lines**

HLA class I alleles A*24:02, A*33:03, A*44:03, A*51:01, and B*52:01 were transduced separately into HLA-null K562 (K562-null) and K562-null with T31SI mutation cells using a ViraPower Lentiviral Expression System (Thermo Fisher Scientific; cat. no. K5330-00). After selection with 5 μg/ml blasticidin (FUJIFILM Wako Pure Chemical; cat. no. 029-18701) for 2 wk, single HLA-transduced K562 cells were stained with anti-HLA class I PE (clone W6/32, BioLegend, San Diego, CA; cat. no. 14249-24) supplemented with 1% FBS and incubated with Cells were suspended in Dulbecco PBS (Nacalai Tesque; cat. no. 12996-81) and 6 μM of PDK-1 inhibitor BX-795 (Selleck Chemicals, Houston, TX; cat. no. S1274) (38). Virus-infected cells were sorted as GFP-positive cells with a FACSria II (BD Biosciences) for use in subsequent experiments.

**PBMCs from healthy donors and patients**

Peripheral blood from healthy volunteers and CML patients was obtained with written informed consent in accordance with the guidelines of the World Medical Association’s Declaration of Helsinki. PBMCs were isolated using Lympholyte-H (Cedarlane Laboratories, Burlington, Ontario, Canada; cat. no. CL5026) following the manufacturer’s protocol. NK cells were isolated using an NK Cell Isolation Kit, human (Miltenyi Biotec, Bergisch Gladbach, Germany; cat. no. 130-092-657) following the manufacturer’s protocol. This study was approved by the institutional review board of Kyoto University Hospital (accession ID: G0697-16, https://kyoto.bvits.com/rinri/publish_document.aspx?ID=1898).

**HLA/KIR genotyping**

HLA-A and -B alleles were determined using a WAKFlow HLA typing kit (Wakunaga Pharmaceuticals, Hiroshima, Japan; cat. no. 4N705 and 4N805), which is based on PCR sequence-specific oligonucleotide probes coupled with multiple analyte profiling technology (Luminex 100 System, Luminex, Austin, TX). The primer sequences in the kit are specifically designed for each HLA locus. The HLA alleles were assigned automatically using the WAKFlow typing software (Wakunaga Pharmaceuticals). KIR3DL1 genotyping was performed by GenoDive Pharma (Atsugi, Japan). Briefly, the KIR gene region (18 KIR genes including KIR3DL1) was enriched by the capture method (captured by hybridization using an artificially synthesized DNA probe) and then sequenced by Miseq (Illumina, San Diego, CA).

**Flow cytometry**

Cells were suspended in Dulbecco PBS (Nacalai Tesque; cat. no. 14249-24) supplemented with 1% FBS and incubated with

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**Table I. HLA-A and B and KIR3DL1 alleles of healthy donors in this study**

<table>
<thead>
<tr>
<th>Donor</th>
<th>HLA-A</th>
<th>HLA-B</th>
<th>KIR3DL1</th>
<th>Z27-MFI</th>
<th>KIR3DS1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A*02:01</td>
<td>A*24:02</td>
<td>B*35:01</td>
<td>B*54:01</td>
<td>*00701</td>
</tr>
<tr>
<td>2</td>
<td>A*31:01</td>
<td>A*33:03</td>
<td>B*44:03:02</td>
<td>B*51:02</td>
<td>*01502</td>
</tr>
<tr>
<td>3</td>
<td>A*24:02</td>
<td>–</td>
<td>B*40:01</td>
<td>B*54:01</td>
<td>*01502</td>
</tr>
<tr>
<td>4</td>
<td>A*24:02</td>
<td>–</td>
<td>B*40:01</td>
<td>B*40:06</td>
<td>*00701</td>
</tr>
<tr>
<td>5</td>
<td>A*02:13</td>
<td>A*26:03:01</td>
<td>B*40:02</td>
<td>B*44:02</td>
<td>*001</td>
</tr>
<tr>
<td>6</td>
<td>A*11:01</td>
<td>A*24:02</td>
<td>B*40:02</td>
<td>B*51:01</td>
<td>*01502</td>
</tr>
<tr>
<td>7</td>
<td>A*11:01</td>
<td>A*24:02</td>
<td>B*40:01</td>
<td>B*55:02</td>
<td>*00701</td>
</tr>
</tbody>
</table>

HLA alleles with a Bw4 motif are indicated in bold.

*The last column shows the KIR3DS1 gene copy number.

---

PBMCs from a KIR3DL1-high donor were coincubated with K562-null and HLA-transduced K562 cell lines at the indicated effector to target (E:T) ratios (10:1, 3:1, and 1:1). The percentages of the CD107a low, no. 1 and no. 4; KIR3DL1-high, no. 2 and no. 3) challenged by K562-null, B*44:03, or B*52:01 transduced K562 cells at an E:T ratio of 1:1. Error bars show the SEM. *p < 0.05, **p < 0.01. ***p < 0.001. FSC, forward scatter; ns, not significant.
FIGURE 2. Anti-KIR3DL1 Ab reverses inhibition of KIR3DL1+ NK cells by HLA-B.

The levels of CD107a expression as a marker of degranulation activity of KIR3DL1+ and KIR3DL1− NK cells challenged by K562-null, B*44:03, or B*52:01 transduced K562 cells in the presence of Z27 or isotype control Abs were measured. Representative flow cytometry contour plots of a KIR3DL1-low donor (no. 1) (A) and a KIR3DL1-high donor (no. 2) (B) are presented. (C) The percentages of CD107a+ KIR3DL1+ NK cells from four donors are presented as the mean value of triplicate samples. Error bars show the SEM. *p < 0.05, **p < 0.01, ***p < 0.001. FSC, forward scatter; ns, not significant.
FIGURE 3. Dasatinib differentially activates NK cells depending on the combination of KIR3DL1 and HLA-B allotypes.

PBMCs from healthy donors were treated with 1 nM dasatinib or DMSO and then cocultured with K562-null or HLA-B–transduced K562 cells at an effector:target ratio of 1:1. (A) Representative flow cytometry contour plots showing the levels of CD107a as a marker of degranulation (Continued)
Abs. For intracellular staining of FOXP3, cells were fixed and permeabilized using a Foxp3/Transcription Factor Staining Buffer Set (eBioscience; cat. no. 00-5523-00). For intracellular staining of phosphorylated ERK1/2, cells were fixed with FIX & PERM Fixation Medium A (Invitrogen; cat. no. GAS001S100) and permeabilized with FIX & PERM Fixation Medium B (Invitrogen; cat. no. GAS002S100). The following Abs were used: anti-CD3 allophycocyanin/H7 (clone SK7, BD Biosciences; cat. no. 641397), anti-CD16 FITC (clone NKPI5, BD Biosciences; cat. no. 347523), anti-CD56 PerCP/Cy5.5 (clone B159, BD Biosciences; cat. no. 560842), anti-CD158e1 allophycocyanin (clone DX9, BD Biosciences; cat. no. 564103), anti-CD107a PE and allophycocyanin (clone H4A3, BioLegend; cat. no. 328608 and no. 328620, respectively), anti-CD158e1e2 PE (clone Z27, Beckman Coulter, Brea, CA; cat. no. IM3292), anti-HLA ABC PE (clone DX17, BD Biosciences; cat. no. 560168), anti-CD45RA PerCP/Cy5.5 (clone SK8, BD Biosciences; cat. no. 346543), anti-CD25 PE/Cy7 (clone M-A251, BD Biosciences; cat. no. 560920), anti-FOXP3 allophycocyanin (clone 236A/E7, eBioscience, Thermo Fisher Scientific; cat. no. 17-4777-42), anti-phospho-p44/42 MAPK (Erk1/2; clone D13.14.4E, Cell Signaling Technology, Danvers, MA; cat. no. 4370), and donkey anti-rabbit IgG (H + L) IgG Alexa Fluor 647 (Invitrogen; cat. no. A31573). Data were acquired using a FACSlyric (BD Biosciences) or FACSCanto II (BD Biosciences) flow cytometer and analyzed with FlowJo software (BD Biosciences).

**Measurement of NK cell activity**

The activation status of NK cells was evaluated by staining CD107a, which is expressed by lytic granules. PBMCs from healthy donors were treated with DMSO, 5 μM imatinib, or 1 nM dasatinib. The percentage of FOXP3^{high}CD45RA^{−} lymphocytes among CD4^{+} T cells were defined as effector Tregs. PBMCs from healthy donors were treated with 5 μM imatinib (Selleck Chemicals; cat. no. S1021) or DMSO (Nacalai Tesque; cat. no. 13408-64) for 24 h. Coculture with target cell lines was started after washing out the drugs.

**Measurement of effector Tregs**

FOXP3^{high}CD45RA^{−} populations among CD4^{+} T cells were defined as effector Tregs. PBMCs from healthy donors were treated with 5 μM imatinib (Selleck Chemicals; cat. no. S2475), 1 nM dasatinib, or DMSO for 24 h. The percentages of effector Tregs before and after treatment were measured by flow cytometry.

**Killing assay**

Target cells were labeled with CellTrace CFSE (Invitrogen; cat. no. C34554) or CellTrace Violet (CTV; Invitrogen; cat. no. C34557) at a final concentration of 2 μM. After incubation with effector cells for 48 h, dead target cells were stained using LIVE/DEAD Fixable Near-IR Dead Cell Stain Kit (Invitrogen; cat. no. L34975). Target cell killing was evaluated by calculating the dead cell percentage (those expressing LIVE/DEAD) among CFSE- or CTV-positive cells.

**Statistical analysis**

Student t test was used to compare mean values of two independent groups. Three or more groups were compared with one-way ANOVA test. A p value < 0.05 was considered statistically significant. All statistical analysis was performed using R software (version 3.6.3, R Foundation for Statistical Computing, Vienna, Austria).

**RESULTS**

**Activation of KIR3DL1^{+} NK cells depends on KIR3DL1 expression**

We first determined the expression levels of KIR3DL1 protein on NK cells in seven healthy donors using two different clones of anti-KIR3DL1 Abs, DX9 and Z27. The donors could be
FIGURE 4. Dasatinib enhances NK cell cytotoxicity in the context of KIR3DL1 and HLA-B noninteracting pairs. (A and B) PBMCs from healthy donors were treated with 1 nM dasatinib or DMSO and then cocultured with K562-null or HLA-B–transduced K562 cells at an effector:target ratio of 4:1. (A) Representative flow cytometry contour plots showing the dead cells after gating on (Continued)
was measured. CD107a expression after coculture with both by B-NK cells from KIR3DL1-high donors showed inhibition were inhibited by Bcells were different depending on the KIR3DL1 expression levels. NK cells from KIR3DL1-low donors showed no inhibition by B*NK cells from KIR3DL1-low donors was inhibited by B*abolition by B*(hereafter, these cell lines are described by the HLA-B alleles they express). CD107a expression as a surrogate marker of degranulation activity of KIR3DL1 and KIR3DL1− NK cells was measured by flow cytometry. Degranulation activity levels increased as the ratio of effector:target decreased (Fig. 1C). Activation of KIR3DL1− NK cells was more affected by HLA-B alleles on target K562 cells than was activation of KIR3DL1− NK cells.

Activation of KIR3DL1+ NK cells from the seven donors was measured. CD107a expression after coculture with K562-null was used as a reference, and activation of KIR3DL1+ NK cells against two HLA-B-transduced K562 cells was different depending on the KIR3DL1 expression levels. NK cells from KIR3DL1-low donors showed no inhibition by B*44:03, which has the Bw4-80T allotype, but were inhibited by B*52:01, which has the Bw4-80I allotype. NK cells from KIR3DL1-high donors showed inhibition both by B*44:03 and B*52:01 (Fig. 1D, 1E).

Anti-KIR3DL1 Ab reverses inhibition of KIR3DL1+ NK cells by HLA-B

To verify that NK cell inhibition depends on physical ligation of KIR3DL1 by HLA-B, PBMCs were first treated with anti-KIR3DL1/S1 Ab Z27 or an isotype control and then cocultured with the target K562 cells. Degranulation (as represented by CD107a expression) of KIR3DL1+ NK cells from KIR3DL1-low donors was inhibited by B*52:01, which was reversed by Z27 Ab (Fig. 2A). Similarly, Z27 Ab blocked the inhibition of KIR3DL1+ NK cells from KIR3DL1-high donors by B*44:03 and B*52:01 (Fig. 2B). The blocking effect was similar within the two groups (Fig. 2C). Z27 Ab had little effect on degranulation activity of KIR3DL1− NK cells from either group.

Dasatinib differentially activates NK cells depending on the combination of KIR3DL1 and HLA-B allotypes

We next examined the effect of dasatinib on NK cell activity. Dasatinib enhanced NK cell activity against K562-null cells at specific concentrations (Supplemental Fig. 1A, 1B). PBMCs from healthy donors were treated with 1 nM dasatinib and then cocultured with K562-null or HLA-transduced K562 cells. Dasatinib enhanced degranulation of KIR3DL1+ NK cells against K562-null cells and those transduced with B*35:01, which had the Bw6 allotype, in each of three donors (nos. 4, 5, and 3; the same applies hereafter) with different KIR3DL1 alleles (Fig. 3A, upper two rows). After dasatinib treatment, KIR3DL1+ NK cells from a KIR3DL1*007 donor showed increased degranulation activity against B*52:01 (Bw4-80I) but not against B*44:03 (Bw4-80T) cells. By contrast, KIR3DL1− NK cells from KIR3DL1*01502 and *001 donors showed increased degranulation activity against B*44:03 (Bw4-80T) but not against B*52:01 (Bw4-80I) cells (Fig. 3A, lower two rows). According to the classification of binding strength between KIR3DL1 and HLA-B, which was defined in previous clinical studies (30,31,34), the combinations in which dasatinib enhanced degranulation were weak or noninteracting pairs (Fig. 3B, 3C; the classification of KIR3DL1 and HLA-B pairs in this experiment is summarized in Supplemental Table 1).

Because the PBMCs included T cells as effectors, we examined the effects of dasatinib on Tregs. Whereas treatment with 5 μM imatinib for 1 d decreased the percentage of effector Tregs, treatment with 1 nM dasatinib did not change the effector Treg frequencies (Fig. 3D, 3E). These results indicate that dasatinib-induced NK cell activation is not associated with Treg numbers.

Dasatinib enhances NK cell cytotoxicity in the context of KIR3DL1 and HLA-B noninteracting pairs

To examine whether increased expression of CD107a as a surrogate marker of degranulation is associated with enhanced cytotoxicity, target cell lysis was evaluated by flow cytometry. PBMCs from donors with KIR3DL1*007, *01502, or *001 alleles were treated with dasatinib and then cocultured with a series of K562 cells as target cells. Dasatinib enhanced cell lysis by PBMCs from a KIR3DL1-low donor (*007) against K562-null and B*35:01 (Bw6) cells but did not change against B*44:03 (Bw4-80T) and B*52:01 (Bw4-80I) cells (Fig. 4A). In contrast, dasatinib enhanced cell lysis by PBMCs from KIR3DL1-high donors (*01502 and *001) against K562-null, B*35:01 (Bw6), B*52:01 (Bw4-80I), and B*44:03 (Bw4-80T) cells (Fig. 4B). These results indicate that dasatinib-mediated enhancement of cytotoxicity was noninteracting.

CFSE-traced target cells. The KIR3DL1 allele of each donor (nos. 4, 3, and no. 5) is indicated in parentheses. Cells positive for the LIVE/DEAD stain are dead cells. (B) The percentages of lysed target cells are presented as the mean value of triplicate samples. The interaction strength between KIR3DL1 and HLA-B is indicated in the bottom of the graph. Error bars show the SEM. *p < 0.05, **p < 0.01, ***p < 0.001. (C, D) NK cells were isolated from PBMCs of healthy donors and used as effectors. freshly isolated NK cells were treated with 1 nM dasatinib or DMSO and then cocultured with K562-null or HLA-B−transduced K562 cells at an effector:target ratio of 1:1. (C) Representative contour plots showing the dead cells among CFSE-traced target cells. The KIR3DL1 allele of each donor is indicated in parentheses. (D) The percentages of lysed target cells are presented as the mean value of triplicate samples. The interaction strength between KIR3DL1 and HLA-B is indicated in the bottom of the graph. Error bars show the SEM. *p < 0.05, **p < 0.01, ***p < 0.001. FSC, forward scatter; ns, not significant.

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FIGURE 5. Dasatinib enhances NK cell cytotoxicity against K562 cells with the BCR-ABL1 T315I mutation.

(A and B) PBMCs from healthy donors were treated with 1 nM dasatinib or DMSO and then cocultured with mutant (mt; bearing the T315I mutation) K562-null or HLA-B–transduced K562 cells at the effector:target ratio of 4:1. (A) Representative flow cytometry contour plots showing

Continued
cells were the main contributor to the lytic effects, we isolated in the KIR3DL1-high donors (Fig. 4B). To confirm that NK KIR3DL1-low donors and with non- and weak-interacting pairs enhanced only in the context of noninteracting pairs in the KIR3DL1-high donors (Fig. 4B). To confirm that NK cells were the main contributor to the lytic effects, we isolated NK cells from healthy donors and used them as effectors. The results were similar to those obtained from coculture of whole PBMCs and K562 cells (Fig. 4C, 4D).

**Dasatinib enhances NK cell cytotoxicity against K562 cells with the BCR-ABL1 T315I mutation**

To verify whether dasatinib enhances lysis of CML cells with the T315I mutation, we used T315I knock-in mutants of K562-null and HLA-B–transduced K562 cells as targets. When dasatinib-treated PBMCs from a KIR3DL1-low donor (*007) were cultured as effectors, target cell lysis was increased against mutant K562-null, mutant B*44:03 (Bw6), and mutant B*52:01 (Bw4-801) cells but unchanged against mutant B*44:03 (Bw4-801) cells (Fig. 5A). By contrast, when PBMCs from KIR3DL1-high donors (*01502 and *001) were cultured as effectors, target cell lysis was increased against mutant K562-null, B*35:01 (Bw6), and B*44:03 (Bw4-801) cells (Fig. 5A). The enhancement of target cell lysis occurred in the context of non- and weak-interacting pairs (Fig. 5B).

We next blocked interaction between KIR3DL1 and HLA-B in strong-interacting pairs using the anti-KIR3DL1 Ab Z27. NK cells from KIR3DL1-low and KIR3DL1-high donors were isolated and treated with dasatinib before coincubation with CFSE-traced target cells in the presence of Z27. Z27 addition resulted in enhanced target cell lysis by NK cells with or without dasatinib (Fig. 5C), but synergistic effects of dasatinib and Z27 were not seen against mutant K562 cells (Fig. 5D).

**NK cell line expressing KIR3DL1*01502 allotype shows an activation pattern against target cells similar to primary NK cells**

KIR3DL1+ NK cells from healthy donors may express other KIR proteins, which makes it difficult to evaluate the function of KIR3DL1 protein. Therefore, functional analysis using a KIR3DL1-negative NK-92 cell line and a KIR3DL1-transduced counterpart was performed. A KIR3DL1-high allele cDNA (KIR3DL1*01502) was introduced into NK-92 cells with a lentiviral vector along with GFP (NK-92-3DL1; Fig. 6A), and GFP-transduced NK-92 cells (NK-92-MOCK) were prepared as a control. Coculture with K562 cells resulted in activation of NK-92 cells as evaluated by flow cytometric analysis of CD107a expression as a marker of degranulation activity. Degranulation activity of NK-92-3DL1 cells was inhibited when the target K562 cells expressed B*44:03 or B*52:01, whereas the control NK-92-MOCK cells were not inhibited by these HLA-B allotypes (Fig. 6B). The inhibition by B*52:01 was stronger than by B*44:03. We also assessed intracellular signaling by measuring phosphorylation levels of ERK1/2. Phospho-ERK1/2 levels were suppressed within NK-92-3DL1 cells when challenged with B*44:03 and B*52:01 cells but not K562-null cells (Fig. 6C). The activation pattern of NK-92-3DL1 cells was consistent with that of KIR3DL1-high primary NK cells (Fig. 6D). To confirm KIR3DL1-mediated modulation of NK-92 cells, target cell lysis was evaluated in the presence of Z27 Ab. Whereas cytotoxicity of NK-92-3DL1 cells was reduced by B*44:03 and B*52:01, the addition of Z27 Ab restored cytotoxic activity (Fig. 6E, 6F).

**Activation of NK cells from CML patients depends on the expression levels of KIR3DL1**

CD107a expression was measured by flow cytometry to assess degranulation activity by KIR3DL1+ NK cells from four CML patients against K562 cells. In addition, we also determined expression levels of KIR3DL1 protein by Z27 MFI. Depending on these analyses, we examined whether the patterns of degranulation activity were consistent with those of the healthy donors. When a patient had both KIR3DL1-low and KIR3DL1-high populations, these were separately analyzed (Fig. 7A). Six populations of KIR3DL1+ NK cells from four patients were analyzed (Fig. 7B, 7C). All KIR3DL1-low populations showed inhibition only by B*52:01, and all KIR3DL1-high populations showed the inhibition by B*44:03 and B*52:01 (Fig. 7D). These results were consistent with the results obtained from healthy donors (Fig. 1D, 1E).

**DISCUSSION**

In this study, we showed that NK cell activity and cytotoxicity against K562 cells was modulated by physical interaction between KIR3DL1 and HLA-B proteins in healthy donors and CML patients (Figs. 1, 2, and 7). In addition, we showed that dasatinib enhanced NK cell activity and cytotoxicity in a KIR3DL1 allotype-dependent manner (Figs. 3 and 4), which is consistent with the findings of our previous clinical study (34). This concept was also confirmed against target cells harboring the BCR-ABL1 T315I mutation (Fig. 5). Furthermore, we proved that blocking an interaction between specific KIR3DL1
FIGURE 6. NK cell line expressing KIR3DL1*01502 allotype shows an activation pattern against target cells similar to primary NK cells.

(A) Histogram showing the expression levels of KIR3DL1 on mock-transduced (white) and KIR3DL1*01502-transduced (gray) NK-92 cells. (B and C) The activation of the NK-92 cell line after coincubation with K562-null, B*44:03, or B*52:01 transduced K562 cells was evaluated by flow cytometry measuring CD107a expression levels (B) and ERK1/2 phosphorylation levels (C). Representative contour plots are shown. (D) Levels of expression of CD107a as a marker of NK-92 activation are presented as the mean value of triplicated samples. Error bars show the SEM. *p < 0.05, **p < 0.01, ***p < 0.001. (E and F) Target cell lysis by mock-/KIR3DL1-transduced NK-92 was evaluated. Target cells were traced with CTV dye and then coincubated with NK-92 effector cells in the presence of 0.1 μg/ml isotype control or Z27 Abs. (E) Representative contour plot showing the percentage of dead cells in each condition. (F) The percentages of lysed target cells are presented as the mean value of triplicate samples. Error bars show the SEM. *p < 0.05, **p < 0.01, ***p < 0.001. FSC, forward scatter; NK-92-3DL1, KIR3DL1*01502-transduced NK-92; NK-92-MOCK, mock-transduced NK-92; ns, not significant.
FIGURE 7. NK cells from CML patients show differential activating patterns depending on the expression levels of KIR3DL1.

(A) A representative flow cytometry contour plot showing the gating for KIR3DL1-high and -low NK cell populations. CD107a expression levels as a marker of degranulation activity of each population after challenge with the K562-null target cells are shown on the right as (Continued)
and HLA proteins could enhance target cell lysis (Fig. 6). These results provide a strong rationale for KIR3DL1 allotypes as therapeutic targets for optimizing treatment of CML and possibly for use to improve treatments of other cancers.

Activation of KIR3DL1+ NK cells against K562 cells was differentially inhibited depending on HLA-B allotypes on the target cells, and this inhibition could be classified into two patterns according to the expression levels of KIR3DL1 protein on the effector cells. Furthermore, KIR3DL1 blocking by Z27 Ab reversed the inhibition. Thus, the combination of KIR3DL1 allotype and HLA-B alleles regulates activity of KIR3DL1+ NK cells. Although the combination of Bw4-80T and KIR3DL1-low allotypes is classified as a strong-interacting pair, KIR3DL1-low NK cells bearing the *007 allele were not inhibited by Bw4-80T HLA-B*44:03 (Fig. 1). However, after treatment with dasatinib, the KIR3DL1*007 and B*44:03 pairing showed no enhancement of cytotoxicity in agreement with the classification (Figs. 3, 4). The classification was originally derived from a clinical study (29), and the affinity between KIR3DL1 and HLA-B allotype molecules measured by tetramer binding assay are not always true for these definitions (39). Notably, our results indicate that modulation of NK cell immunity through KIR3DL1 and HLA can be altered in the presence of dasatinib, which has as an immunomodulatory function.

Boudreau et al. (28) showed slightly stronger binding of 3DL1*007 and *015 tetramers to B*44:03 than to B*52:01. Saunders et al. (39) reported similar results from tetramer binding assays. Results published by Foley et al. (40) demonstrated stronger inhibition of both 3DL1-high (\(*015) and -low (\(*005\)-like) NK cells by B*44:03 than by B*52:01. However, in our experiment, B*52:01 showed stronger inhibition of both 3DL1-high (\(*015) and -low (\(*007) NK cells than B*44:03. This discrepancy may be attributable to differences in target cell lines. All previous research used EBV-transformed B-LCL (721.221) cells as the target, whereas we used K562 cells. It is now assumed that 721.221 cell line does not completely lack HLA class I expression, whereas our data confirm that K562 cells completely lack HLA class I expression. Therefore, we consider that K562 cells are more suitable for analyzing KIR–HLA interactions. Tremblay-McLean et al. (41) showed that K562 cells express several ligands for activating NKG2D receptor, whereas 721.221 cells do not, which may also affect NK cell activation.

It has been reported that imatinib and dasatinib activate NK cell immunity through suppression of Tregs (35,36). Treg suppression was not apparent at the concentration of dasatinib used in our experiments, suggesting another mechanism of NK cell activation by dasatinib. Although we have not elucidated the mechanistic details, Chang et al. (42) observed downregulation of NKG2A, another inhibitory receptor on NK cells, in CML patients who took dasatinib and showed that dasatinib downregulates NKG2A through inhibition of p38 MAPK. These results imply a direct effect of dasatinib on NK cell activation and also suggests that NK cells may be more dependent on signals from KIRs in the presence of dasatinib. In addition, enhancement of NK cell immunity by dasatinib was also shown against K562 cells with the BCR-ABL T315I mutation, which is naturally resistant to dasatinib. Even when the KIR–HLA combination was a strong-interacting pair, inhibition of KIR–HLA binding resulted in enhanced killing. These results suggest a possible strategy to overcome TKI resistance in CML by targeting and modulating the KIR3DL1–HLA interaction.

This study has several limitations. First, our previous research suggested that the KIR3DL1*005 allele has a significant positive impact on the therapeutic effects of TKIs (34). However, as none of our volunteer donors carried the KIR3DL1*005 allele, we could not verify its function and specificity in vitro. We assumed that the NK cell activation pattern would be similar for those of the *005 and *007 allotypes because they share the KIR3DL1-low phenotype, although this remains to be verified. Second, as NK cell effects are an aggregate of multiple KIR–HLA interactions, our in vitro results cannot be translated directly into clinical practice, and validation in a large clinical study is needed. Third, we could not examine how dasatinib affected KIR3DL1–HLA interactions and modulated NK cell activity in detail. Despite these limitations, our results are sufficient to indicate the importance of NK cell modulation through KIR–HLA interactions during CML therapy.

Therapeutic blockade of the inhibitory KIRs on NK cells is an attractive way to enhance antitumor NK cell immunity. Its feasibility has been explored using the Ab lirilumab, which targets KIR2DL1/2/3. However, although promising results were reported in a mouse model (43), a recent phase Ib clinical trial for relapsed or refractory lymphoid malignancies failed to show additional benefits of lirilumab on PD-1 blockade (44). Intervention at the level of KIR3DL1–HLA binding could be an alternative option for a novel immune checkpoint therapy. Several studies mention that blocking inhibitory KIRs including
KIR3DL1 could enhance NK cell cytotoxicity (15, 30,39,45). Our study not only confirmed these results but also showed the significance of KIR3DL1 blocking in the presence of a TKI. Furthermore, we showed that KIR3DL1 blocking was effective even in TKI-resistant cells, providing a new treatment strategy for patients with CML that has acquired drug-resistant mutations. Notably, our results also suggest that appropriate patient selection based on the KIR3DL1 alleles could maximize the efficacy of KIR3DL1 blockade. Only a limited number of KIR–HLA pairs were examined in our study, and a more detailed analysis of KIR–HLA interactions, including validation in a large clinical study, should be conducted to optimize NK cell immunity against CML and other malignancies.

DISCLOSURES

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KIR3DL1 ALLOTYPE-DEPENDENT NK CELL MODULATION AGAINST CML


