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Blocking expression of inhibitory receptor NKG2A overcomes tumor resistance to NK cells

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A key mechanism of tumor resistance to immune cells is mediated by expression of peptide-loaded HLA class I molecule (HLA-E) in tumor cells, which suppresses NK cell activity via ligation of the NK inhibitory receptor CD94/NK group 2 member A (NKG2A). Gene expression data from approximately 10,000 tumor samples showed widespread HLA-E expression, with levels correlating with those of KLRC1 (NKG2A) and KLRD1 (CD94). To bypass HLA-E inhibition, we developed a way to generate highly functional NK cells lacking NKG2A. Constructs containing a single-chain variable fragment derived from an anti-NKG2A antibody were linked to endoplasmic reticulum–retention domains. After retroviral transduction in human peripheral blood NK cells, these NKG2A protein expression blockers (PEBLs) abrogated NKG2A expression. The resulting NKG2A−/− NK cells had higher cytotoxicity against HLA-E–expressing tumor cells. Transduction of anti-NKG2A PEBL produced more potent cytotoxicity than interference with an anti-NKG2A antibody and prevented de novo NKG2A expression without affecting NK cell proliferation. In immunodeficient mice, NKG2A−/− NK cells were substantially more powerful than NKG2A+ NK cells against HLA-E–expressing tumors. Thus, NKG2A downregulation evades the HLA-E cancer immune checkpoint and increases the antitumor activity of NK cell infusions. Because this strategy is easily adaptable to current protocols for clinical-grade immune cell processing, its clinical testing is feasible and warranted.

Introduction

Immunotherapy has become a mainstay of contemporary oncology. Major clinical responses achieved with T cell immune-checkpoint inhibitors and chimeric antigen receptor–directed (CAR-directed) T lymphocytes (1, 2) encourage the evaluation of other immunotherapeutic approaches. To this end, the anticancer potential of NK cell infusions has not yet been fully explored. In patients with acute myeloid leukemia (AML), responses associated with NK cells have been reported after allogeneic hematopoietic stem cell transplantation (3–5) and after infusion of NK cells activated ex vivo by cytokines (4, 6). Clinical activity, however, remains difficult to predict, responses are often not durable, and therapeutic potential against tumors beyond AML is unclear (7, 8).

NK cells have the inherent capacity to identify, and specifically eliminate, virally infected or transformed cells (7–9). NK activating and inhibitory receptors, ligated to varying degrees by molecules expressed on candidate target cells, regulate these functions (8, 9). The CD94/NKG2A heterodimeric receptor is one of the most prominent NK inhibitory receptors (10). It binds to a nonclassical minimally polymorphic HLA class I molecule (HLA-E), which presents peptides derived from leader peptide sequences of other HLA class I molecules, such as HLA-G (11–15). Upon ligation by peptide-loaded HLA-E, NKG2A transduces inhibitory signaling through 2 inhibitory immune-receptor tyrosine-based inhibition motifs, thus suppressing NK cytokine secretion and cytotoxicity (16–18).

Although HLA-E transcripts are ubiquitously expressed (19), surface expression of HLA-E is often weak or undetectable in tumor cell lines (20). During antitumor immune responses, however, IFN-γ secreted by immune cells upregulates HLA-E expression in tumor cells (21–23). Stabilized by peptide loading (24), the HLA-E/peptide complex dampens initial NK cell responses by binding CD94/NKG2A. HLA-E is indeed overexpressed in several tumors (25–33), and a correlation between higher HLA-E and poorer prognosis has been observed (25,27). Moreover, NKG2A+ NK cells are reportedly predominant in the tumor microenvironment. Thus, intratumoral NK cells in patients with non–small cell lung cancer have higher expression of NKG2A than those obtained from nontumoral distant sites (34, 35). Likewise, NKG2A expression is higher in NK cells infiltrating breast cancer (BRCA) tumors than in those isolated from symmetric normal breast tissue (36) and in peripheral blood NK cells of patients with AML compared with NK cells of age-matched controls (37). Because of the strong evidence for the suppressive effect of NKG2A on NK cell activity, this immune checkpoint has been targeted with antibodies, currently in clinical development, that block its interaction with HLA-E, currently in clinical development (38–41).

In this study, we determined HLA-E expression in approximately 10,000 tumor samples and assessed its relation with that of KLRC1 (NKG2A) and KLRD1 (CD94). Then we sought to develop NK cells that were impervious to the inhibitory effects of HLA-E.

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To this end, we designed constructs, named protein expression blockers (PEBLs), that prevent the transport of NKG2A to the cell-surface membrane of NK cells. Finally, we tested to determine whether downregulation of NKG2A could overcome the HLA-E-mediated mechanism of tumor resistance to NK cell killing.

Results
HLAE expression in tumor samples, relation with immune cell infiltration, and KLRC1/KLRD1 expression. We performed an unbiased analysis of HLAE expression in human tumors using data from 10,375 tumor samples, representing 33 tumor types, made available by The Cancer Genome Atlas (TCGA) Research Network. We found considerable inter- and intratumor heterogeneity in HLAE expression (Figure 1A). Tumor types with more than 100 samples whose median HLAE expression exceeded the median value (log10 transcripts per kilobase million of 3.92) included kidney renal clear cell carcinoma (KIRC), head and neck squamous cell carcinoma (HNSC), lung adenocarcinoma (LUAD), skin cutaneous melanoma (SKCM), prostate adenocarcinoma (PRAD), stomach adenocarcinoma (STAD), colon adenocarcinoma (COAD), cervical squamous cell carcinoma and endocervical adenocarcinoma (CESC), lung squamous cell carcinoma (LUSC), and rectum adenocarcinoma (READ).

When all tumors were analyzed collectively, there was a direct relation between levels of HLAE expression and those of KLRC1 (NKG2A) (Pearson’s correlation coefficient \( r = 0.48 \)) and KLRD1 (CD94) (\( r = 0.57 \)), while there was no relation with the activating member of the NKG2 family KLRC2 (NKG2C) (\( r = 0.07 \)) (Figure 1, B and C). The correlation between HLAE and KLRC1 was particularly evident in some tumors, such as bladder urothelial carcinoma (BLCA), sarcoma (SARC), CESC, HNSC, and PRAD (Figure 1D).

We further assessed the correlation between HLAE and KLRC1 using the Tumor Immune Estimation Resource (TIMER; https://cistrome.shinyapps.io/timer/), which allows estimates of immune-cell infiltration (42). Among tumors with more than 100 samples, the highest correlation values were observed in BLCA, thyroid carcinoma (THCA), thymoma (THYM), HNSC, and BRCA; the lowest were in low-grade glioma (LGG), glioblastoma multiforme (GBM), KIRC, and pheochromocytoma and paraganglioma (PCPG) (Supplemental Table 1; supplemental material available online with...
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linked to 4 different endoplasmic reticulum (ER) retention domains (Figure 2A). PEBL1 contains a (GGGGS)4AEKDEL domain; it binds the KDEL (lysine, aspartic acid, glutamic acid, leucine) peptide receptor, linked to the coat protein complex I (COPI) vesicular transport, which mediates protein traffic between ER and Golgi (44, 45). PEBLs 2–4 contain different peptides, including KKMP (lysine, lysine, methionine, proline) domains, together with CD8α hinge and transmembrane domains; these allow direct binding to COPI (45, 46).

To test the capacity of PEBLs to downregulate NKG2A, we inserted the constructs into a murine stem cell virus (MSCV) retroviral vector containing GFP and used it to express the anti-NKG2A PEBLs in the NKG2A+ cell line NK92. NKG2A expression was essentially abrogated in all GFP+ NK92 cells (Figure 2B). PEBLs were not detectable on the cell surface, whereas a surface membrane-bound anti-NKG2A scFv used as a control was highly expressed (Supplemental Figure 4), indicating that PEBLs were confined to the intracellular space.

To determine whether anti-NKG2A PEBLs could also downregulate NKG2A expression in primary human NK cells, we expanded peripheral blood NK cells (n = 10 from 9 donors) by 5- to 6-day stimulation with the K562-mb15-41BBL cell line (47, 48). We then purified NKG2A+ NK cells by magnetic bead–positive selection and transduced them with anti-NKG2A-PEBL2. The percentage of NKG2A+ cells before purification was 69.8% ± 11.8%, and this was increased to 98.8% ± 1.6% after purification. Transduction of anti-NKG2A PEBL markedly reduced NKG2A. Among GFP+ cells, representing 49.1% to 82.0% (median, 68.2%) of all NK cells, NKG2A+ cells were 7.5% ± 5.9%, while in cells transduced with a vector containing only GFP, NKG2A expression remained high (97.6% ± 3.2%; P < 0.0001) (Figure 2, C and D).

Cell markers, gene expression, and functional features of NKG2A+ NK cells. Anti-NKG2A PEBL transduction resulted in downregulation of CD94 surface expression (Figure 3, A and B). In the 7 donors tested, residual CD94+ cells represented 21.9%–45.1% (median, 37.8%) of NK cells transduced with PEBL versus 89.9%–99.7% (median, 96.8%) of NK cells transduced with GFP alone. After staining the cells with the anti-NKG2A antibody and testing for CD94 expression after cell permeabilization.
of NK cells transduced with GFP alone. We noted that expression of CD25 was lower in NK cells transduced with anti-NKG2A PEBL than in those transduced with GFP alone (Figure 3D). Nevertheless, PEBL-transduced NK cells could expand as well as control NK cells with IL-2 (Figure 3E). Likewise, the IL-2-dependent cell line NK92 continued to proliferate normally after downregulation of NKG2A (Supplemental Figure 6B).

We applied RNA-Seq to examine the expression of 44 genes encoding NK activating and inhibitory receptors, cytokines and cytokine receptors, and cytolytic molecules. There were no differences in mRNA expression for NKG2A, CD94, and NKG2C, and none of the other 41 genes was marked as a “discovery” after controlling for false discovery rate (Supplemental Table 2).

To further examine the functionality of NKG2A<sup>-null</sup> NK cells, we tested their capacity to lyse HLA-E-negative K562 leukemia cells. In both short- and long-term cultures, anti-NKG2A PEBL-transduced NK cells were as powerful as control NK cells (Figure 3, F and G). In line with these results, NKG2A<sup>-null</sup> cells and control NK cells had equal capacities to exocytose cytotoxic granules in the presence of target cells, as shown by CD107α staining (Supplemental Figure 6C). IFN-γ secretion after target stimulation was also maintained (Supplemental Figure 6D). Hence, downregulation of NKG2A by PEBL did not impair NK cell function.

**NKG2A<sup>-null</sup> NK cells avert suppression by HLA-E-expressing tumor cells.** To determine whether NKG2A downregulation released NK cells from HLA-E-mediated inhibition, we generated tumor cells with strong NKG2A-binding potential. For this purpose, we transduced the ligand of NKG2A, i.e., HLA-E plus HLA-G signal peptide (GpHLA-E) (49, 50), in tumor cell lines derived from AML (K562), osteosarcoma (U2OS), and Ewing’s sarcoma (ES8 and EW8) (Supplemental Figure 7A). These cell lines express a variable profile of inhibitory and activating NK ligands: the inhibitory HLA-class I is absent or dim in K562 and ES8, but highly expressed in U2OS and EW8; the activating NKG2D ligands MICA/B and ULBP1-3 and the DNAM-1 ligands CD112 and CD155.
Antibody binding to NKG2A may reduce its NK inhibitory activity (38, 39, 41, 55). To compare the effect of anti-NKG2A PEBL to that of antibody interference with NKG2A, we preincubated NKG2A+ cells with either the anti-NKG2A antibody Z199 or an isotype-matched nonreactive antibody. We then tested their capacity to kill GpHLA-E–transduced K562, U2OS, and ES8 cells. Although Z199 significantly improved the cytotoxicity of control NK cells against all 3 targets, the same NK cells transduced with anti-NKG2A PEBL were consistently more cytotoxic (Figure 4C).

We used these targets to test the function of NKG2A null NK cells. In 4-hour cytotoxicity assays, NKG2Anull NK cells had a markedly higher killing capacity than control NK cells (P < 0.0001 for all comparisons; Figure 4A). Exocytosis of lytic granules marked by CD107a expression was not confined to NKG2C+ cells; in fact, most CD107a+ cells lacked both NKG2A and NKG2C expression (Figure 4B). Importantly, NKG2A downregulation did not increase cytotoxicity against autologous activated CD4+ T cells (Supplemental Figure 8A), an effect that NKG2A had been previously reported to suppress (52, 53).

CD155 are expressed in all 4 lines, but at generally low levels in ES8 and EW8 (51). Regardless, GpHLA-E expression markedly inhibited CD107a secretion by NKG2A+ NK cells and NK cytotoxicity overall (Supplemental Figure 7, B and C).

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To further test the gains in antitumor activity afforded by NKG2A downregulation, we performed long-term cytotoxicity assays at low (1:2–1:8) effector/target cell (E/T) ratios against GpHLA-E–transduced cell lines. As shown in Figure 4D, tumor cell killing was again markedly superior with the anti-NKG2A PEBL cells. These results were corroborated in experiments in which U2OS-GpHLA-E was grown as a spheroid. As shown in Figure 4E and Supplemental Figure 9, NKG2A<sup>null</sup> cells were clearly more powerful.

Finally, we determined whether antibody-dependent cell cytotoxicity (ADCC) was affected by the downregulation of NKG2A with anti-NKG2A PEBL. For this purpose, we transduced the HER2<sup>+</sup> BRCA cell line SK-BR-3 with GpHLA-E (Figure 5A) and used these cells as targets for the anti-Her2 antibody trastuzumab. As shown in Figure 5B, NKG2A<sup>null</sup> cells were more powerful against GpHLA-E-SK-BR-3 cells in the absence of antibody (P = 0.0001) and maintained superior cytotoxicity in the presence of trastuzumab (P = 0.018).

NKG2A downregulation increases NK cell killing of tumor cells exposed to IFN-γ and suppresses NKG2A upregulation by IL-12. The above experiments showed that downregulation of NKG2A dramatically reduced the inhibition exerted by transduced GpHLA-E. We identified cell lines with endogenous surface HLA-E expression (Supplemental Figure 10) and tested their relative susceptibility to cell killing by NK cells with or without NKG2A downregulation. As shown in Figure 6A, even after only 4 hours of coculture, NKG2A<sup>null</sup> NK cells exerted significantly higher cytotoxicity in 3 of the 4 cell lines (P < 0.01 for the AML cell line U937, and P < 0.0001 for the hepatocellular carcinoma cell line PLC/PRF/5 and the acute lymphoblastic leukemia cell line OP-1; the remaining cell line, the Ewing’s sarcoma cell line EW8, 4-hour cell killing was higher with NKG2A<sup>null</sup> cells (76.5% ± 1.3% vs. 66.4 ± 6.2%; n = 9), but the difference was not statistically significant. When tested against a spheroid model of the U2OS cell line not transduced with GpHLA-E (this cell line expresses low surface HLA-E; Supplemental Figure 7A), NKG2A<sup>null</sup> cells were significantly more powerful than their NKG2A<sup>+</sup> counterparts (P < 0.01 at 6 and 24 hours; P < 0.001 at 12 and 18 hours; Figure 6B).

It is known that IFN-γ secreted by immune cells reactive to tumors increases expression of HLA-E, which, in turn, promotes resistance to NK cells via NKG2A ligation (23). We determined whether improvements in tumor cell killing brought about by NKG2A downregulation with PEBL extended to tumor cells exposed to IFN-γ. After 12 hours of exposure to IFN-γ (300 ng/ml), EW8, PLC/PRF/5, and U937 overexpressed HLA-E (Supplemental Figure 10). These cells were consistently more susceptible to killing by NKG2A<sup>null</sup> NK cells than by control NK cells (P < 0.01 for PLC/PRF/5; P < 0.0001 for both EW8 and U937) (Figure 6C).

Because NK cells secrete IFN-γ during activation (Supplemental Figure 6), we postulated that exposure to the supernatant from cocultures of NK cells and target cells would also induce NK cell resistance in tumor cells. To test this notion, we cultured NK cells with EW8 or U937 cells for 24 hours, collected and filtered the supernatant, and added it to fresh EW8 and U937. After 12 hours, we compared NK cells with and without NKG2A downregulation in 4-hour cytotoxicity assays. As shown in Figure 6D, exposure to the NK-conditioned medium induced considerable resistance to NKG2A<sup>+</sup> cells, but NKG2A<sup>null</sup> cytotoxicity remained high (for EW8, P < 0.0001 at 2:1 and 1:1 and P < 0.01 at 1:2; for U937, P < 0.0001 at 2:1 and P < 0.01 at 1:1 and 1:2).

Previous studies have shown that AML cells from patients overexpress HLA-E after exposure to IFN-γ, suggesting that this might favor the growth of NK-resistant leukemic cells (33). We obtained 4 bone marrow specimens from patients with AML collected at diagnosis (n = 3) or relapse (n = 1) and containing over 90% AML blasts. We exposed the cells to IFN-γ (300 ng/ml for 12 hours) and used them as targets of NKG2A<sup>+</sup> NK cells that had been transduced with either anti-NKG2D PEBL or GFP alone. As shown in Figure 6E, NKG2A<sup>null</sup> cells exerted a significantly higher cytotoxicity (P < 0.0001) in 2 of the 4 samples. Interestingly, the 2 sensitive samples were also those expressing the highest levels of HLA-E (MFI: 17171 and 12267 versus 8595 and 6390).

IL-12, secreted by macrophages and dendritic cells, is an important activator of immune responses against tumor cells, including the generation of NK cells with memory-like properties (56, 57). However, exposure of NK cells to IL-12 also increases expression of CD94/NKG2A (58–60). In the context of enhanced HLA-E expression in tumor cells caused by IFN-γ secretion, this is likely to further dampen NK cell responses. We purified NKG2A<sup>-</sup>cells from peripheral blood NK cells from 3 donors and transduced them with either GFP alone or anti-NKG2A PEBL. We noticed that coculture with K562-mb15-41BBL cells during the transduction procedure induced an increase in NKG2A expression in control NK cells, while NKG2A<sup>-</sup> remained undetectable in PEBL-transduced cells (Figure 6F). Exposure to IL-12 (20 ng/ml for 5 days) further increased NKG2A expression in control NK cells, but not in PEBL-transduced NK cells (Figure 6F).

Thus, the PEBL strategy not only downregulated NKG2A expression in NKG2A<sup>-</sup>positive NK cells, but also suppressed NKG2A expression after stimulation and IL-12 exposure. When the cyto-
The toxicity of PEBL-transduced NK cells exposed to IL-12 was tested using GpHLA-E K562 cells, there was a clear improvement in cytotoxicity over that of control NK cells (*P < 0.0001 at all E/T ratios) (Figure 6G).

Antitumor capacity of NKG2A\textsuperscript{null} NK cells in vivo. The above results indicated that NKG2A downregulation markedly enhanced the antitumor capacity of NK cells. To further challenge this finding, we engrafted the Ewing’s sarcoma cell line ES8 transduced with GpHLA-E in immunodeficient mice. As shown in Figure 7, treatment of mice with 2 infusions of control NK cells (transduced with GFP alone and expressing NKG2A) on days 1 and 5 after tumor injection only delayed tumor development. Median survival for the 2 groups was 22 and 36 days from tumor infusion. In contrast, the same NK cells with downregulated NKG2A by PEBL induced profound responses, which resulted in long-term survival for most mice (median survival was not reached after more than 269 days follow-up; *P < 0.001, log rank test compared with control NK cells). In a second model, we engrafted the osteosarcoma line U2OS expressing GpHLA-E and treated mice with 2 infusions of NK cells 3 and 10 days after tumor injection (Supplemental Figure 11). Again, control NK cells only slightly delayed relapse (median survival, 25 and 32 days, respectively). Also, in

Figure 6. Cytotoxicity of NKG2A\textsuperscript{null} NK cells against tumor cells with endogenous HLA-E expression. (A) Four-hour cytotoxicity of NK cells transduced with anti-NKG2A PEBL or GFP only (Control) against cell lines expressing endogenous HLA-E (see Supplemental Figure 10). EW8 and PLC/PRF/5 were transduced with luciferase. BrightGlo was added after 4 hours of coculture, and luminescence was measured using a Flx 800 plate reader. Cytotoxicity of U937 and OP-1 was measured by flow cytometry. Box (25th–75th percentile, median) and whiskers (minimum–maximum) plots from 3 experiments with cells from 3 donors (EW8, PLC/PRF/5), and 6 with cell lines from 2 donors (U937, OP-1) in triplicate, at 2:1, 1:1, or 1:2 E/T. (B) Spheroid tumors of U2OS-mCherry were cocultured with NK cells at 1:2 E/T in triplicate and analyzed with IncuCyte Zoom System. Data are shown as mean (± SD) red calibrated unit (RCU)/μM². Representative images at end of culture are shown. Scale bars: 300 μm. (C) Four-hour cytotoxicity against cell lines exposed to IFN-γ (300 ng/ml; 12 hours). Plots are from 3 experiments with NK cells from 3 donors (EW8, PLC/PRF/5), and 6 with NK cells from 2 donors (U937) in triplicate at 2:1, 1:1, or 1:2 E/T. (D) Similar experiments targeting cells exposed for 12 hours to conditioned medium (C.M.) from 24-hour cocultures of NK cells with the respective cell lines. Four-hour cytotoxicity was compared with that against cells not exposed to conditioned medium. (E) Four-hour cytotoxicity against primary AML cells from 4 patients, exposed to IFN-γ (300 ng/ml; 12 hours). Data are from 4 experiments with NK cells from 2 donors in triplicate at 2:1 and 1:1 E/T. (F) NKG2A-negative NK cells from 3 donors were stimulated with K562-mb15-41BBL for 7 days, transduced with anti-NKG2A PEBL or GFP alone, and then exposed to IL-12 (20 ng/ml) for 5 days. Percentage of NKG2A\textsuperscript{null} cells at each stage is shown. (G) PEBL-transduced and control NK cells were exposed to IL-12 and tested in 4-hour cytotoxicity assays against K562-GpHLA-E cells. Data are shown as mean (± SD) of triplicate measurements at each E/T. **p < 0.01; ***p < 0.001; ****p < 0.0001, t test.
CD94. The resulting NK cells not only retained their general function, but they were substantially more cytotoxic than control NK cells against target cells expressing HLA-E molecules loaded with the HLA-G signal peptide. NKG2A downregulation by PEBL also produced higher antitumor activity than interference with an anti-NKG2A antibody, while cell killing of nontransformed cells did not increase. The gains in cytotoxicity observed in short- and long-term in vitro cultures were confirmed in xenograft models, where NKG2A null cells induced durable tumor remission. These results indicate that NKG2A downregulation is an effective way to increase the antitumor activity of NK cell therapies.

In tumor cells, downregulation of classical HLA class I molecules decreases susceptibility to T cell recognition and cytotoxicity (61). At the same time, overexpression of nonclassical HLA class I molecules, such as HLA-E and HLA-G, generates resistance to NK cells through binding of CD94/NKG2A (40). CD94 has no signaling capacity, and as we showed in our study, its expression on the cell surface of NKG2A+ NK cells is largely dependent on the presence of NKG2A. CD94 can also form heterodimers with NKG2C, CD94. This model NKG2A<sup>null</sup> cells had clearly superior antitumor activity to that of their NKG2A<sup>+</sup> counterparts (median survival, not reached after 60 days follow-up; \( P < 0.01 \)). Of note, no evidence of xenoreactivity (weight loss, ruffled fur, abnormal posture, or decreased activity) was observed.

**Discussion**

The unique capacity of NK cells to recognize and kill tumor cells (8, 9) has been corroborated by reports of major clinical activity in some patients (6, 7), suggesting that NK cell infusions could have considerable potential for the treatment of cancer. Tumors, however, develop mechanisms of resistance that can nullify the NK cell effect. In this study, we developed a way to overcome the potent NK resistance mechanism mediated by HLA-E ligation of the inhibitory CD94/NKG2A receptor. We sought to abrogate its impact by removing NKG2A from the surface of NK cells and achieved this goal with an approach based on constructs designed to prevent protein surface expression. Anti-NKG2A PEBLs stopped NKG2A surface expression together with that of its coreceptor, CD94. The resulting NK cells not only retained their general function, but they were substantially more cytotoxic than control NK cells against target cells expressing HLA-E molecules loaded with the HLA-G signal peptide. NKG2A downregulation by PEBL also produced higher antitumor activity than interference with an anti-NKG2A antibody, while cell killing of nontransformed cells did not increase. The gains in cytotoxicity observed in short- and long-term in vitro cultures were confirmed in xenograft models, where NKG2A<sup>null</sup> cells induced durable tumor remission. These results indicate that NKG2A downregulation is an effective way to increase the antitumor activity of NK cell therapies.
which activates NK cells via DAP12 (10, 62). Although HLA-E can bind both inhibitory CD94/NKG2A and activating CD94/ NKG2C, it has a lower affinity for the latter (50, 63). Expression of HLA-E, and hence resistance to NK cells, can be further enhanced by exposure to IFN-γ secreted by tumor-reactive immune cells (21–23). In our experiments, we recapitulated this scenario by exposing tumor cells to IFN-γ or to supernatant collected from cocultures of NK cells and tumor cells or by enforcing expression of GpHLA-E (49, 50, 64). In all these settings, resistance to NK cells increased, but downregulation of NKG2A on NK cells overcame resistance and markedly increased cytotoxicity. Another potential mechanism of resistance during immune response to tumors is expression of NKG2A generated by exposure of NK cells (including cells originally lacking surface NKG2A expression) to higher levels of IL-12 in the tumor microenvironment (58–60). Transduction of NK cells with anti-NKG2A PEBL blocked such upregulation and restored cytotoxicity.

The importance of the CD94/NKG2A-HLA-E immune checkpoint in regulating NK cell activity against tumors has led to the ongoing clinical testing of an anti-NKG2A inhibitory antibody (39, 40). We used the sequence of the heavy and light chain regions of an anti-NKG2D antibody as a basis for developing an anti-NKG2A scFv. We found that when the scFv was coupled to selected ER/ Golgi retention domains, downregulation of NKG2A was rapid and nearly absolute. Importantly, NKG2A+ NK cells with NKG2A downregulation had more powerful cytotoxicity than those exposed to the anti-NKG2A antibody. This could be explained by the fact that CD94/NKG2A recycles between cell surface and cytoplasm independently of HLA-E ligation (65), yielding a continuous supply of new receptors on the cell surface that may eventually escape antibody blockade. NKG2A is a major NK inhibitory receptor, but there are other receptors that also regulate NK cell function. To this end, we have used PEBLs to successfully downregulate the NK inhibitory receptors KIR2DL1 and KIR2DL2/3 in primary NK cells (T. Kamiya and D. Campana, unpublished results). The relative antitumor capacity of these cells, NKG2A\textsuperscript{null} cells, and those lacking multiple inhibitory receptors remains to be investigated.

Downregulation of NKG2A might have also been achieved by using modern gene-editing methodologies based on meganucleases, TALEN, or CRISPR/Cas9, which have already been applied to genetically modify T lymphocytes (66–69). As shown in this study, PEBL-mediated downregulation of NKG2A is highly efficient and specific. It also has several potential practical advantages. First, anti-NKG2A PEBL expression can be accomplished with viral vectors identical to those currently used for expression of CAR in T lymphocytes. Therefore, the PEBL method can be readily adapted to current protocols for clinical-grade processing of genetically modified immune cells. Second, PEBL composition is similar to that of currently used CARs and should not pose the risk of nonspecific gene editing. Therefore, the introduction of PEBLs in clinical cell processing should not elicit additional regulatory concerns. Finally, anti-NKG2A PEBL can be combined with activating receptors to further enhance NK cell cytotoxicity. PEBL constructs can be assembled in multicistronic vectors that allow simultaneous expression of PEBLs and activating receptors. We have used this approach to effectively downregulate surface molecules in T cells while simultaneously expressing CARs (70, 71).

Expression of PEBL in NK cells relied on a culture system that has already been translated into a GMP-grade protocol or infusion of autologous or allogeneic NK cells in patients with cancer (8). Several approaches are being tested clinically to improve the anticancer potential of NK cell infusions, such as ex vivo expansion (8, 72, 73), activation with cytokines (4, 6), and infusion of antibodies against inhibitory receptors (39, 40, 74). Other approaches, such as CAR expression (47, 75), expression of activating receptors (76), and expression of membrane-bound cytokines (77), suggest possibilities to further enhance NK cell potency that warrant clinical exploration. The NKG2A\textsuperscript{null} cells generated in this study can evade a central mechanism of tumor resistance to NK cells. Importantly, NKG2A is also expressed in a subset of activated CD3+ T lymphocytes and its ligation can inhibit their function (23, 78, 79). In mice, loss of function of Qa-1b (equivalent to HLA-E in humans) resulting from in vivo CRISPR screening increased sensitivity of tumor cells to immune cells (80). Correlative studies in ovarian and cervical carcinomas have shown a predominant infiltration of CD8+ lymphocytes expressing NKG2A (26, 81), and recently published studies indicate that an anti-NKG2A antibody can augment T cell antitumor activity (41, 82). Thus, the approach described here could potentially be extended to protocols of adoptive T cell therapy of cancer.

Methods

Gene expression studies. RNAseqv2 expression data from 10,375 individual tumor annotated samples were downloaded from the TCGA web portal. Thirty-three tumor types, as defined by TCGA codes, were included (Figure 1). Pairwise gene expression correlation values and associated P values were calculated in R. Gene expression box plots were generated using R graphics package GGPlot2 (83). Ordered column plots, scatter plots, and linear regression lines were plotted in Microsoft Excel or GraphPad. Analyses were also performed using TIMER (42).

Cells. The human cell lines NK92, K562, SK-BR-3, PLC/PRF/5, U2OS, and U937 were obtained from ATCC. ES8 and EW8 were from the tissue repository of St. Jude Children’s Research Hospital (Memphis, Tennessee, USA). OP-1 was developed in our laboratory (84). We transduced K562, SK-BR-3, PLC/PRF/5, U2OS, ES8, EW8, and U937 cells with a Mscv-IRES-GFP (where IRES indicates internal ribosome entry site) retroviral vector containing the firefly luciferase gene (48). K562, U2OS, and ES8 cells were also transduced with Mscv-IRES-mCherry vector. Transduced cells were selected for their expression of GFP or mCherry with a MoFlo (Beckman Coulter) or a FACSaria (BD Biosciences) cell sorter. Bone marrow–derived mesenchymal stromal cells immortalized by expression of human telomerase reverse transcriptase were previously developed in our laboratory (54). NK92 cells were cultured in RPMI 1640 (Thermo Fisher Scientific) with 12.5% FBS, 12.5% horse serum, 200 IU/ml IL-2, and antibiotics; other cell lines were maintained in RPMI 1640 or in DMEM (GE Healthcare); U2OS and PLC/PRF/5 with 10% FBS (GE Healthcare) and antibiotics. Tumor cell lines were periodically validated by DNA fingerprinting analysis (DSMZ).

Mononucleated cells from peripheral blood of healthy donors, and from peripheral blood or bone marrow of patients with AML were separated by centrifugation on a Lymphoprep Density Step (Axis-Shield) and washed twice in RPMI 1640.
NK cell expansion and NKG2A selection. To expand NK cells, peripheral blood mononucleated cells were cocultured with the genetically modified K562-mb15-41BBL cell line in SCGM medium (CellGenix) containing 10% FBS and 40 IU/ml IL-2 (Proleukin, Novartis) (8, 47, 48, 51, 76, 77, 85). After 5 to 7 days of culture, residual T cells were removed using Dynabeads CD3 (Thermo Fisher). NK cells were labeled with allophycocyanin-conjugated (APC-conjugated) anti-CD159a (NKG2A) antibody, followed by anti-APC Microbeads (Miltenyi Biotec); the LS and LD columns (Miltenyi Biotec) were used to select or remove NKG2A+ cells, respectively. Selected NK cells were restimulated with K562-mb15-41BBL cells and 400 IU/ml IL-2. Recombinant human IL-12 (R&D Systems; 20 ng/ml) was used in some experiments.

PEBL constructs. We designed a scFv containing the variable regions of the heavy and light chains of the murine anti-human NKG2A monoclonal antibody Z199 (17, 43) and a 20-amino acid linker. To construct the anti-NKG2A PEBLs, the nucleotide sequence was joined to the CD8a signal peptide and sequences encoding ER/Golgi retention peptides (GGGGS)4AEKDEL (PEBL1), or CD8a hinge and transmembrane domain followed by LYYKKSRRSFIEEKKMP (PEBL2), LYKKSRRSFIDEKKMP (PEBL3), or LYCNKYSRRSFIEEKKMP (PEBL4; Figure 1A). Constructs were subcloned into the MSCV-derived retroviral vector containing IRES and GFP. Preparation of retroviral supernatant and transduction were performed as previously described (85). Briefly, MSCV retroviral vector–conditioned medium was added to polypropylene tubes coated with RetroNectin (Takara); after centrifugation and removal of the supernatant, NKG2A-selected and restimulated NK cells were added to the tubes and left at 37°C for 12 hours; fresh viral supernatant was added on 2 other successive days. Transduced NK cells were maintained in SCGM medium with FBS, antibiotics, and 400 IU/ml of IL-2 until the time of experiments.

Cell marker and mRNA expression. Expression of PEBLs was detected using a biotin-conjugated goat anti-mouse F(ab′)2, antibody (Jackson ImmunoResearch), followed by streptavidin conjugated to APC (Jackson ImmunoResearch). Other antibodies used to determine cell-marker expression are listed in Supplemental Table 3. Intracellular staining for CD94 was performed after surface staining with anti-NKG2A and permeabilization with 8E, a reagent developed in our laboratory. Cell staining was analyzed using Accuri C6 or Fortessa flow cytometers (BD Bioscience), with Diva (BD Biosciences) or FlowJo software.

Whole-transcriptome mRNA-Seq library preparation of NK cells with or without NKG2A downregulation was performed using the TruSeq Stranded mRNA Library Prep Kit (Illumina) and sequenced on a NextSeq 500 platform with a read length of 2 × 151 bp. The reads were mapped to the human genome reference hg19 using STAR (86). Reads aligned to each gene were counted using featureCounts (87), and the expression levels were calculated as fragments per kilobase of transcript per million mapped reads (FPKM). Expression levels were log transformed, followed by batch effect correction using ComBat of FPKM. All original mRNA-Seq data were deposited in the NCBI’s Gene Expression Omnibus database (GEO GSE127081).

Cytokine production, CD107a staining, and cytotoxicity assays. To measure IFN-γ production, NK cells were cocultured at 37°C 5% CO2, with target cells for 8 hours in the presence of 0.1% Brefeldin A (GolgiPlug, BD Biosciences); cells were then labeled with phycoerythrin-conjugated (PE-conjugated) anti-IFN-γ (clone 25723.11, BD Biosciences) after cell membrane permeabilization with 8E (a permeabilization reagent developed in our laboratory). To measure exocytosis of lytic granules, target and effector cells were cocultured in the presence of PE-Cyanin 7–conjugated (Cy7-conjugated) anti-human CD107a antibody (H4A3; BioLegend). After 1 hour, monensin (GolgiStop, BD Biosciences) was added to the plate, and cultures were prolonged for another 3 hours.

To test cytotoxicity, NK cells were cocultured with target cells labeled with calcein red-orange AM (Thermo Fisher Scientific). After 4 hours, the number of viable target cells was counted by flow cytometry (76). To test cytotoxicity against luciferase-labeled target cells, BrightGlo (Promega) was added after 4 hours of coculture, and luminescence was measured using a Fx800 plate reader (BioTek). To test ADCC, trastuzumab (10 μg/ml; Herceptin, Roche) was added to SK-BR-3 cells expressing GhpH-L-E prior to coculture with NK cells. For long-term cytotoxicity assay, NK cells were cocultured with mCherry-transduced target cells, and mCherry-expressing cells were cultured with the IncuCyte Zoom System (Essen BioScience). To generate tumor spheroids, U2OS cells expressing mCherry cells were seeded into ultra-low attachment 96-well round bottom plates (Corning) at a concentration of 500 cells per 100 μl of RPMI 1640. After centrifugation, cultures were maintained for 7 days. Plates were then transferred into the IncuCyte Zoom System. On day 8, NK cells were added. Images were captured with the IncuCyte Zoom System and a x10 objective lens at 6-hour intervals. Tumor cell growth was quantified based on the fluorescence intensity of mCherry.

To interfere with NKG2A function, we used a purified nonconjugated anti-NKG2A antibody (Z199, Beckman Coulter); an iso-type-matched nonreactive immunoglobulin (R&D) was used as a control. In some experiments, we used as targets cell lines (K562, ES8, EW8, U2OS, and SK-BR-3) transduced with a construct encoding HLA-E with its signal peptide (MVVMAPRTLFLLLSGALT) replaced with the HLA-G signal peptide (MVVMAPRTLFLLLSGALTLPT) (49, 50). The nucleotide sequence was cloned into the MSCV vector containing IRES and GFP and transduced in cells that were previously transduced with firefly luciferase and GFP. To detect HLA-E expression, transduced cells were labeled with APC-conjugated anti–HLA-E antibody (Thermo Fisher Scientific) and sorted with the MoFlo cell sorter. In some experiments, target cells were cultured for 12 hours with either IFN-γ (300 ng/ml; Sigma-Aldrich) or culture supernatant from cocultures of NK cells and tumor cells before the cytotoxicity assays. The culture supernatant was collected from 24-hour cocultures, centrifuged at high speed, and filtered before adding it to fresh tumor cells.

Xenograft models. ES8 cells expressing luciferase and GhpH-L-E were injected i.p. into NOD.Cg-Pkdcreid IL2rgtm16/j/Sjz (NOD/SCID IL2Rγnull) mice (Jackson Laboratory) at 2 × 104 per mouse. NK cells were expanded for 5 days. After CD3 depletion and NKG2A+ selection, NK cells were restimulated with K562-mb15-41BBL cells and transduced with anti-NKG2A PEBL or GFP only. One and five days after ES8 cell injection, PEBL-transduced and control NK cells were administered at 1 × 105 cells per mouse (7 mice per group). Another group of 9 mice received RPMI 1640 with 10% FBS instead of NK cells. All mice received i.p. injections of IL-2 (20,000 IU each) 3 times per week. Similar experiments were performed with U2OS-GphH-L-E cells injected i.p. in 15 mice, with the difference being that NK cells were adminis-
tered on days 3 and 10 after tumor injection. Bioluminescence was measured with a Xenogen IVIS Spectrum system (Caliper Life Sciences) after i.p. injection of d-luciferin potassium salt (PerkinElmer; 150 μg/g body weight). Luminescence was emitted with Living Image 3.0 software. Mice were euthanized when the sum of ventral and dorsal bioluminescence signal reached 2 to 5 × 10^10 photons per second or earlier if physical signs warranting euthanasia appeared. Mice were distributed in each group to achieve an even representation of tumor engraftment based on the luminescence signal recorded before NK cell administration. The only inclusion criteria were tumor engraftment by luminescence before NK cell infusion.

Statistics. For RNA-Seq analysis of NK cells, paired t test was used to compare expression levels, with statistical analysis performed using R. Controlling for false discovery rate was done using the 2-stage linear step-up procedure of Benjamini, Krieger, and Yekutieli, with a Q of 1% using GraphPad. Comparisons of NK cell function were analyzed using t test; P < 0.05 was considered significant. Mouse survival was analyzed by log-rank test.

Study approval. Peripheral blood samples were obtained from discarded anonymized by-products of platelet donations from healthy adult donors at the National University Hospital or the Health Science Authority Blood Banks, Singapore. Samples from patients with AML were obtained from banked surplus material. Studies were performed with approval from the Institutional Review Board, National University of Singapore. All mouse procedures were approved by the Institutional Animal Care and Use Committee and conducted in accordance with current guidelines.

Author contributions
TK developed PEBLs, performed experiments, and analyzed data. SVS and DW performed experiments and analyzed data. MR analyzed gene expression data. DC designed the study, analyzed data, and wrote the manuscript with TK, SVS, DW, and MR.

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