

Transmissible cytotoxicity of multiple myeloma cells by cord blood-derived NK cells is mediated by vesicle trafficking

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Natural killer cells (NK) are important effectors of anti-tumor immunity, activated either by the downregulation of HLA-I molecules on tumor cells and/or the interaction of NK-activating receptors with ligands that are overexpressed on target cells upon tumor transformation (including NKG2D and NKP30). NK kill target cells by the vesicular delivery of cytolytic molecules such as Granzyme-B and Granulysin activating different cell death pathways, which can be Caspase-3 dependent or Caspase-3 independent. Multiple myeloma (MM) remains an incurable neoplastic plasma-cell disorder. However, we previously reported the encouraging observation that cord blood-derived NK (CB-NK), a new source of NK, showed anti-tumor activity in an *in vivo* murine model of MM and confirmed a correlation between high levels of NKG2D expression by MM cells and increased efficacy of CB-NK in reducing tumor burden. We aimed to characterize the mechanism of CB-NK-mediated cytotoxicity against MM cells. We show a Caspase-3- and Granzyme-B-independent cell death, and we reveal a mechanism of transmissible cell death between cells, which involves lipid–protein vesicle transfer from CB-NK to MM cells. These vesicles are secondarily transferred from recipient MM cells to neighboring MM cells amplifying the initial CB-NK cytotoxicity achieved. This indirect cytotoxicity involves the transfer of NKG2D and NKP30 and leads to lysosomal cell death and decreased levels of reactive oxygen species in MM cells. These findings suggest a novel and unique mechanism of CB-NK cytotoxicity against MM cells and highlight the importance of lipids and lipid transfer in this process. Further, these data provide a rationale for the development of CB-NK-based cellular therapies in the treatment of MM.

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Natural killer cells (NK) are important effectors of anti-tumor immunity of the immune system. They can be activated by inhibition of killer cell immunoglobulin (Ig)-like receptor (KIR) receptors on NK due to downregulation of HLA-I on tumor cells or by the interaction of NK-activating receptors with ligands that are overexpressed on target cells. These receptors include NKG2D and the family of NK cytotoxicity receptors (NKP30, NKP44, NKP46),¹ which can be on the cell surface and in the endocytic compartment² from where they traffic to tumor cells in exosomes to exert cytotoxicity.³

NK deliver cytolytic molecules such as Granzyme-B (GrB) and Granulysin to target cells. GrB induces Caspase-3-dependent apoptotic cell death with reactive oxygen species (ROS) generation.⁴ Alternatively, Granulysin activates Caspase-3-dependent cell death through ROS generation^{5,6} and Caspase-3-independent cell death via endoplasmic reticulum (ER) stress⁷ or lysosomal cell death through release of cathepsins.⁸

Multiple myeloma (MM) is a plasma-cell disorder characterized by clonal proliferation of malignant plasma-cells in the

bone marrow (BM) and monoclonal protein in the blood or urine.^{9,10} Plasma cells synthesize large quantities of Igs, which are folded in the ER. An excess of Ig synthesis causes a failure in this folding process leading to the release of unfolded/misfolded Igs.¹¹ These Igs are degraded by intracellular protein degradation pathways, including the ubiquitin–proteasome system and autophagy. Proteasome inhibitors (PIs) are potent anti-MM agents,¹² which block the protein degradation process in MM cells leading to ER stress-mediated cell death.^{13,14} An excess of PI-induced ER stress can increase autophagy^{15,16} leading to eventual refractory disease in some patients.^{17–19} Therefore new anti-MM strategies are needed.

Previously, we have demonstrated that cord blood-derived NK (CB-NK) show anti-tumor activity in an *in vivo* MM murine model²⁰ and observed that the expression of NKG2D by MM tumor cells correlated with lower tumor burden following CB-NK cellular therapy. Here we characterize the CB-NK-mediated cytotoxicity against MM cells and observe a

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Abbreviations: NK, natural killer cells; CB-NK, cord blood-derived NK cells; MM, multiple myeloma; ROS, reactive oxygen species; ER, endoplasmic reticulum; GrB, Granzyme-B

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Caspase-3- and Gr-B-independent cell death and reveal a mechanism of transmissible cell death between cells that involves lipid-protein vesicle transfer from CB-NK to MM cells. These vesicles are secondarily transferred from recipient MM cells to neighboring MM cells, thereby amplifying the initial CB-NK cytotoxicity achieved. This indirect cytotoxicity involves the transfer of NKG2D and NKP30 and leads to lysosomal cell death and reduced ROS levels in MM cells.

Results

NKG2D expression in MM cells after CB-NK treatment correlates with lower MM progression, and NKG2D and NKP30 contribute more to the cytotoxicity of MM cells than in K562 cells. We have previously shown that CB-NK exert anti-MM activity in a murine MM model.²⁰ Immunodeficient mice were injected with the ARP1 MM cell line. MM cells were identified in the BM, spleen, lymph nodes (LN) and ovaries and demonstrated higher MFI for NKG2D in the BM, LN and ovaries from CB-NK-treated mice *versus* untreated mice (Supplementary Figures S1A–D and F–I). CB-NK in CB-NK recipient mice were CD56⁺, CD3[−], NKG2D⁺ and CD16⁺ (Supplementary Figure S1J). In treated mice, MFI for NKG2D in MM cells in LN correlated inversely with tumor burden ($R = -0.886$, $P = 0.04$; Supplementary Figure S1J), suggesting a transfer of NKG2D from CB-NK to MM cells with a beneficial effect.

As the presence of NKG2D in MM cells was associated with lower tumor burden, we investigated the cytotoxic role of NKG2D. We also examined NKP30, as it has an important role in the killing of MM cell lines,²¹ MICA/B (NKG2D ligand) and the synergistic effects of these receptors. We compared our results from ARP1 cells with K562 cells, where, in the absence of HLA-I, NK cytotoxicity is mediated by KIR receptors.

After blocking these receptors, we found a higher contribution of these in the killing of ARP1 *versus* K562 cells (Figures 1a–d). The greatest impact in ARP1 cells was mediated by NKP30 followed by NKG2D (Figures 1b and d). At a 20:1 effector: target cell ratio, we observed a 33% (NKG2D blocked), 49% (NKP30 blocked) and 73% (both blocked) reduction in ARP1 killing (Figure 1d) *versus* 4% (NKG2D blocked), 3% (NKP30 blocked) and 19% (both blocked) reduction in K562 killing (Figure 1c). This effect was reproduced at high effector: target cell ratios (Figures 1c and d, $P < 0.0001$). A similar pattern was observed with primary MM CD138⁺ cells showing at 20:1 effector: target cell ratio a 26.6% (NKG2D blocked) and 51% (NKP30 blocked) reduction in CD138⁺ cell killing (Figures 1e–g). Moreover, in NK from MM patients (MM-NK) that showed lower cytotoxicity than CB-NK (Supplementary Figure S2A), NKG2D and NKP30 receptors did not impact NK-MM cytotoxicity (Supplementary Figure S2B), revealing this unique feature of CB-NK cytotoxicity and the superiority of CB-NK for MM treatment.

NKG2D and NKP30 are transferred into MM cells via lipid trafficking and the endocytic pathway. We reported an inverse correlation between NKG2D expression in MM-ARP1 cells and disease progression and demonstrated that NKG2D and NKP30 were involved in CB-NK cytotoxicity.

The trafficking of these receptors to MM cells was subsequently followed using confocal fluorescence microscopy. Cell trackers were used to label CB-NK and tumor cells. Tumor cells were labeled in blue using CMAC and CB-NK in green with bodipy-505/515. This has an affinity for neutral lipids and has a chloromethyl group that allows binding to membrane proteins, thereby labeling proteins and lipids. We observed that bodipy staining allowed us to follow the trafficking of contents from CB-NK to ARP1 cells (Figure 2, and Supplementary Movie S1).

Within 10 min of tumor cell/CB-NK co-culture, CB-NK transferred their contents to MM cells. This was via lipid-protein vesicles and larger volume vesicles (Figure 2a). Further, some of the lipid-protein vesicles trafficked between ARP1 cells after transfer from CB-NK (Figure 2b). Such transfer was not detected in the absence of CB-NK (Figure 2c). This trafficking also involved the immunological synapse, as demonstrated by actin co-staining (Figures 2a and b). Of note, MM-NK showed less intense immunological synapse *versus* MM cells and lower bodipy staining (Supplementary Figure S2C) in comparison to CB-NK supporting the finding of lower MM-NK cytotoxicity and indicating lower levels of lipids/cytotoxic vesicles in MM-NK. The stronger immunological synapses and bodipy staining, indicative of higher levels of lipids, were characteristic of more potent CB-NK.

Time-lapse *in vivo* imaging revealed that the intensity of the materials transferred to ARP1 cells by CB-NK increased with time as indicated by the increase in Bodipy (regions of interest (ROIs) 1–3 in Figures 2d and e). Also, after CB-NK transferred vesicles to ARP1 cells, these ARP1 cells could then transfer Bodipy-containing vesicles to neighboring ARP1 cells (ROIs 4 and 5 in Figures 2d, f and g and Supplementary Movie S1). This transfer was associated with membrane blebbing appearance (Figure 2g) indicative of apoptotic-like mechanisms.²² Imaging revealed that, after 20 min of co-culture, NKG2D and NKP30 were detected in ARP1 cells (Figures 3a–c), whereas they could not be detected in ARP1 cells that had not been with CB-NK (Figure 3d). After 3 days of co-culture, flow cytometry confirmed surface and intracellular presence of NKG2D and NKP30 in ARP1 cells (Figures 3e and f).

We investigated the role of the endocytic pathway in the NKG2D and NKP30 transfer to ARP1 cells. Cells uptake molecules from the extracellular environment by endocytosis. NK-activating receptors are continuously recycled and degraded by endocytic pathway.² Once produced, NK-activating receptors are delivered to early endosomes (defined by EEA1). They are then either recycled to the plasma membrane (defined by Rab11) and exocytosed, or delivered to late endosomes (defined by Rab7), to be targeted to lysosomes for degradation.^{2,23,24} We therefore analyzed whether NKG2D and NKP30 once transferred between cells entered the endocytic pathway in MM cells.

We observed that bodipy co-localized with NKG2D and NKP30 (column 3, Figure 4) confirming concomitant lipid and protein trafficking between cells. Co-localization of NKG2D and NKP30 with EEA1 (column 4, Figures 4a and d), Rab11 (column 4, Figures 4b and e) and Rab7 (column 4, Figures 4c and f) indicated active trafficking of NKG2D and NKP30 through endocytic pathway involving lipid-protein trafficking.

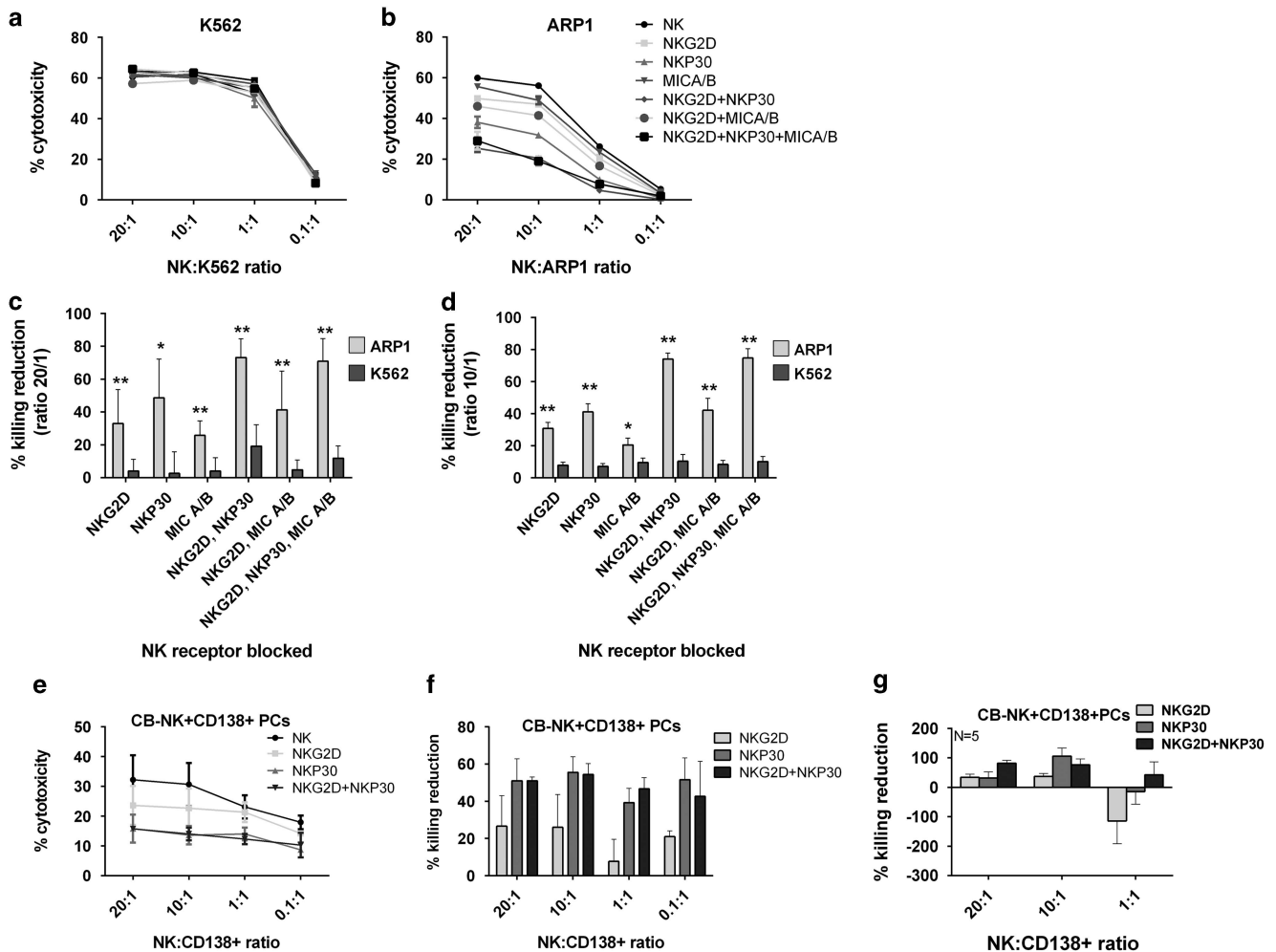


Figure 1 NKG2D and NKP30 contribute more to the cytotoxicity of MM cells than in K562 cells: (a and b) Cytotoxicity assays of CB-NK against K562 (a) and ARP1 (b) cells after blocking NKG2D, NKP30 in CB-NK and MICA/B in target cells. (c and d) Percentage of killing reduction obtained in panels a and b after blocking the receptors. Bars represent mean \pm S.E.M. values at a effector:target cell ratio 20:1 (c) and 10:1 (d) ($n=8$). (e) CB-NK cytotoxicity assay *versus* primary CD138⁺ MM cells, adding CB-NK before and after blocking NKG2D, NKP30 and both together. (f) Percentage of killing reduction obtained in panel e after blocking the receptors. (g) Mean values of experiment shown in panels e and f with CD138⁺ cells from five MM patients. Bars represent mean \pm S.E.M. * $P \leq 0.05$. ** $P \leq 0.0001$

To confirm the importance of lipid-protein trafficking in the transfer of NKG2D and NKP30 and in CB-NK cytotoxicity, we analyzed the expression levels of NKG2D and NKP30 in target cells following preincubation of CB-NK with Filipin III (a lipid raft inhibitor)^{22,25} and performed cytotoxicity assays adding U18666A (a cholesterol synthesis and transport inhibitor²⁶). Lipid rafts inhibition reduced the expression of NKG2D and NKP30 in ARP1 cells after CB-NK exposure (Figure 4g), providing evidence to support a role for lipid-protein trafficking in the transfer of NKG2D and NKP30. Cholesterol synthesis inhibition in CB-NK reduced cytotoxicity only *versus* ARP1 (Figures 4h and i); however, for K562 we observed higher cytotoxicity at 10:1 effector:target ratio, suggesting different roles for lipids in CB-NK cytotoxicity depending on the target cell. We confirmed in U266 and KMM1 MM cells the cytotoxic role of CB-NK-derived lipids (Supplementary Figure S3A). Interestingly, peripheral blood NK (PB-NK) of healthy individuals failed to show evidence of a cytotoxic role of lipids *versus* MM (Supplementary Figure S4A), suggesting another unique feature of CB-NK.

The mechanisms underlying CB-NK-mediated cytotoxicity differ between target cells. As we had evidence for different cytotoxic mechanisms depending on the target cells (MM *versus* K562) (Figure 1), other parameters were assessed. Upon interaction with target cells, NK degranulate releasing secretory lysosomes with GrB,²⁷ activating Caspase-3 in target cells with features of apoptosis, such as membrane blebbing and ROS generation.⁴ CB-NK showed less degranulation *versus* ARP1 than *versus* K562 cells (mean value: 7.6 *versus* 68%, $P=0.029$) (Figures 5a and b), which was confirmed for other MM cell lines (Supplementary Figure S3B). Further, we noted that GrB was transferred to ARP1 cells after contact (Figure 5c). Interestingly, inhibition of GrB and Caspase-3 in NK-K562 and NK-ARP1 co-cultures reduced the killing of K562 cells by CB-NK as expected but increased the killing of ARP1 cells by CB-NK (Figures 5d and e). Inhibition of GrB and Caspase-3 at a 20:1 effector:target ratio yielded a reduction of 13% in CB-NK cytotoxicity against K562 cells but an increase of 12% in killing against ARP1 cells (Figures 5e, $P < 0.05$).

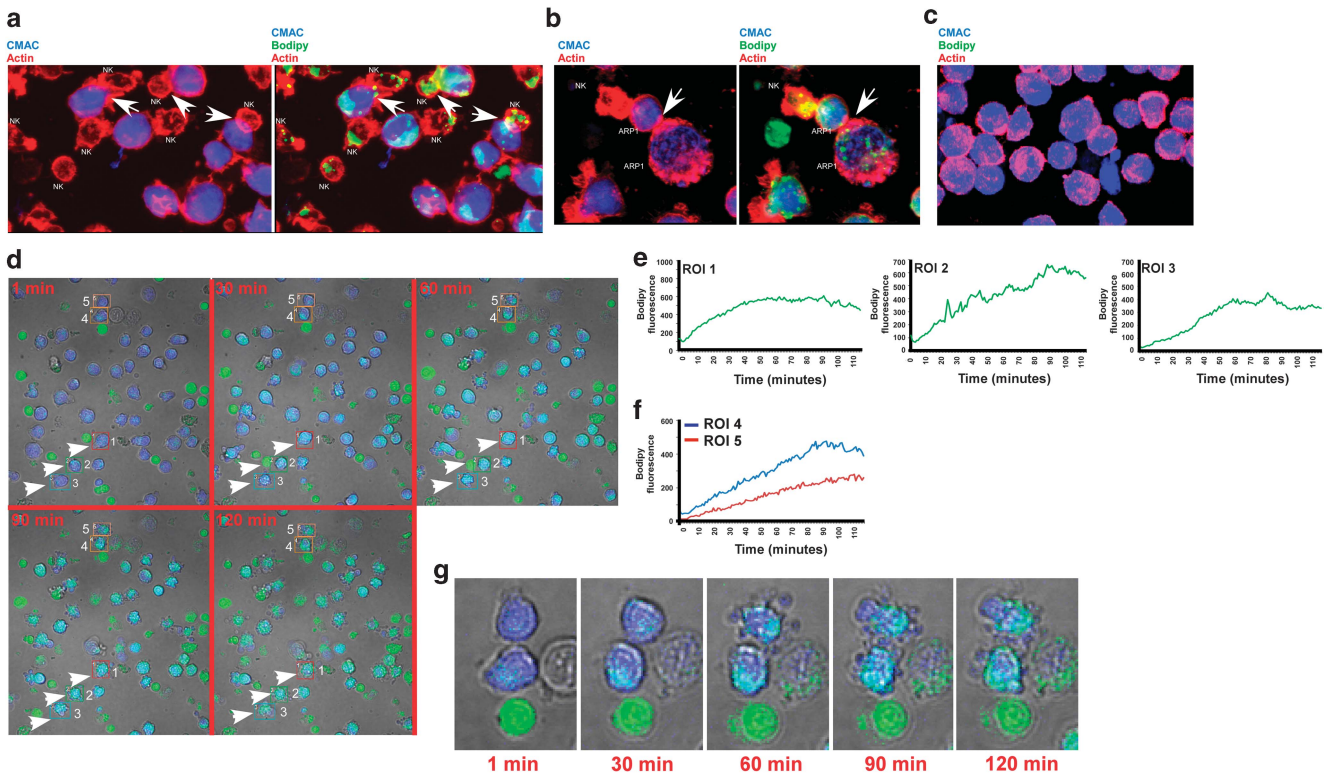


Figure 2 CB-NK contents are transferred to ARP1 cells via lipid vesicles or lipid droplets (LDs). (a) CB-NK and ARP1 cells were co-cultured in cell pellet for 10 min. The two images are the same showing ARP1 cells stained in blue (CMAC) and CB-NK in green (bodipy). Actin is shown in red. CMAC and actin are shown in the first image and the three colors together in the second image. Arrows indicate transfer of CB-NK content in green into ARP1 cells, indicating lipid trafficking. (b) Same experiment as panel a showing one CB-NK transferring content to one ARP1 cell and this ARP1 cell transferring content to a secondary ARP1 cell through the immunological synapse. (c) ARP1 cells alone as control stained with CMAC and actin in red. (d) Different time points taken from a time lapse *in vivo* imaging for 2 h (Supplementary Movie S1) where ARP1 cells were stained in blue (CMAC) and CB-NK in green (bodipy). Five different ARP1 cells are indicated in regions of interest (ROIs). ROIs 1–3 indicate three ARP1 cells that had been touched by the same CB-NK. ROI 4 indicates one ARP1 cell touched by a CB-NK and then this ARP1 cell transfers the CB-NK content into another ARP1 cell (ROI 5). (e) Analysis of the increase during time of green staining (bodipy) in ROIs 1–3 from panel d. (f) Analysis of the increase during time of green staining (bodipy) in ROIs 4 and 5 in panel d. (g) Amplified images from ROIs 4 and 5 in panel d to show ARP1 cells showing membrane blebbing. See also Supplementary Movie S1

GrB- and Caspase-3-independent CB-NK cytotoxicity was also confirmed for other MM cell lines (Supplementary Figure S3C). However, cytotoxicity of PB-NK from healthy individuals *versus* K562 and ARP1 was GrB and Caspase-3 dependent (Supplementary Figure S4B), providing another unique characteristic of CB-NK. Because CB-NK degranulation indicates release of lysosomes, we measured lysosomes in K562 and ARP1 cells after CB-NK. As controls, CD19⁺ cells from healthy individuals were used. Whereas in K562 and CD19⁺ cells there was an increase in lysosomes after CB-NK (Figure 5f, $P < 0.0001$), in ARP1 there was a decrease (Figure 5f, $P < 0.0001$), confirming that different mechanisms of CB-NK cytotoxicity against MM cells exist.

CB-NK induce lysosomal cell death in MM cells, and NKG2D and NKP30 are involved in this mechanism. We aimed to understand the mechanisms underlying CB-NK cytotoxicity *versus* MM cells. We had observed lipid trafficking between cells, cytotoxicity that was independent GrB and Caspase-3 and decreased lysosomes after CB-NK exposure (Figure 5). Granulysin can cause Caspase-3-independent cell death with ER stress and lysosomal permeabilization.^{7,8} After permeabilization, lysosomes

release cathepsins leading to a Caspase-3-independent cell killing called ‘lysosomal cell death’.⁸ We hypothesized that CB-NK induce lysosomal cell death in MM cells. We analyzed lysosome markers (Lyso-tracker and Rab7, which also defines lysosomes) and the impact of cysteine cathepsins in CB-NK cytotoxicity. After 40 min with CB-NK, ARP1 cells had reduced lysosomes levels (Figures 6a, b, and d, $P < 0.0001$). Further, cathepsins inhibition reduced CB-NK cytotoxicity *versus* ARP1 and not *versus* K562 (Figure 6e), confirming a lysosomal cell death for MM cells; which was confirmed in other MM cell lines, by detecting both a decrease in Rab7 and the impact of cathepsins in CB-NK cytotoxicity (Supplementary Figures S3D and E). Primary MM CD138⁺ cells also showed decreased lysosomes levels after CB-NK (Supplementary Figure S3F). Interestingly, cytotoxicity of PB-NK from healthy individuals *versus* K562 or ARP1 cells was not affected by cathepsins inhibition (Supplementary Figure S4C), indicating another unique feature of CB-NK.

ROS were also decreased in ARP1 cells (Figures 6c and f, $P < 0.0001$) and in primary MM CD138⁺ cells after CB-NK exposure (Supplementary Figure S3F). As GrB leads to ROS production⁴ and Caspase-3-dependent cell death, we

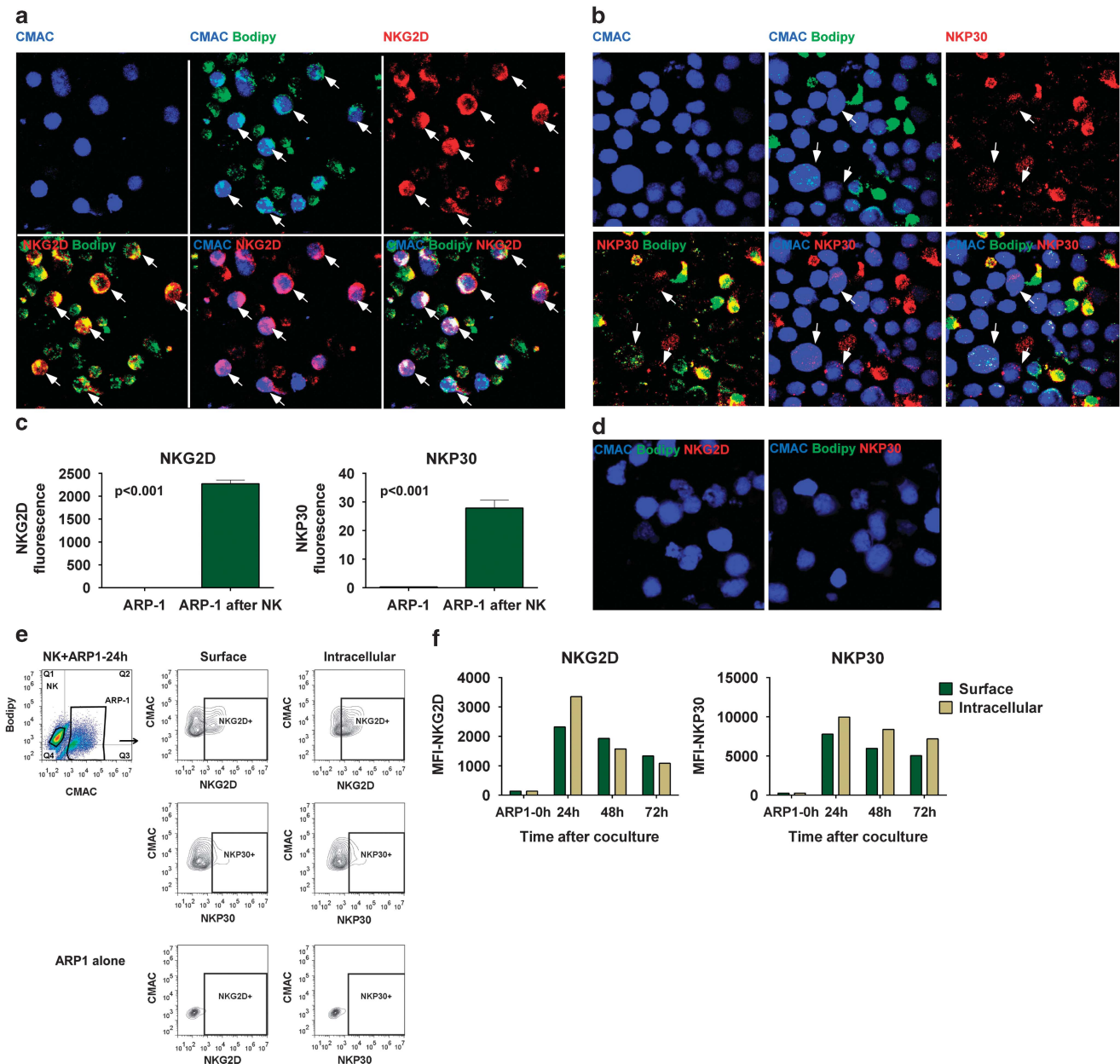
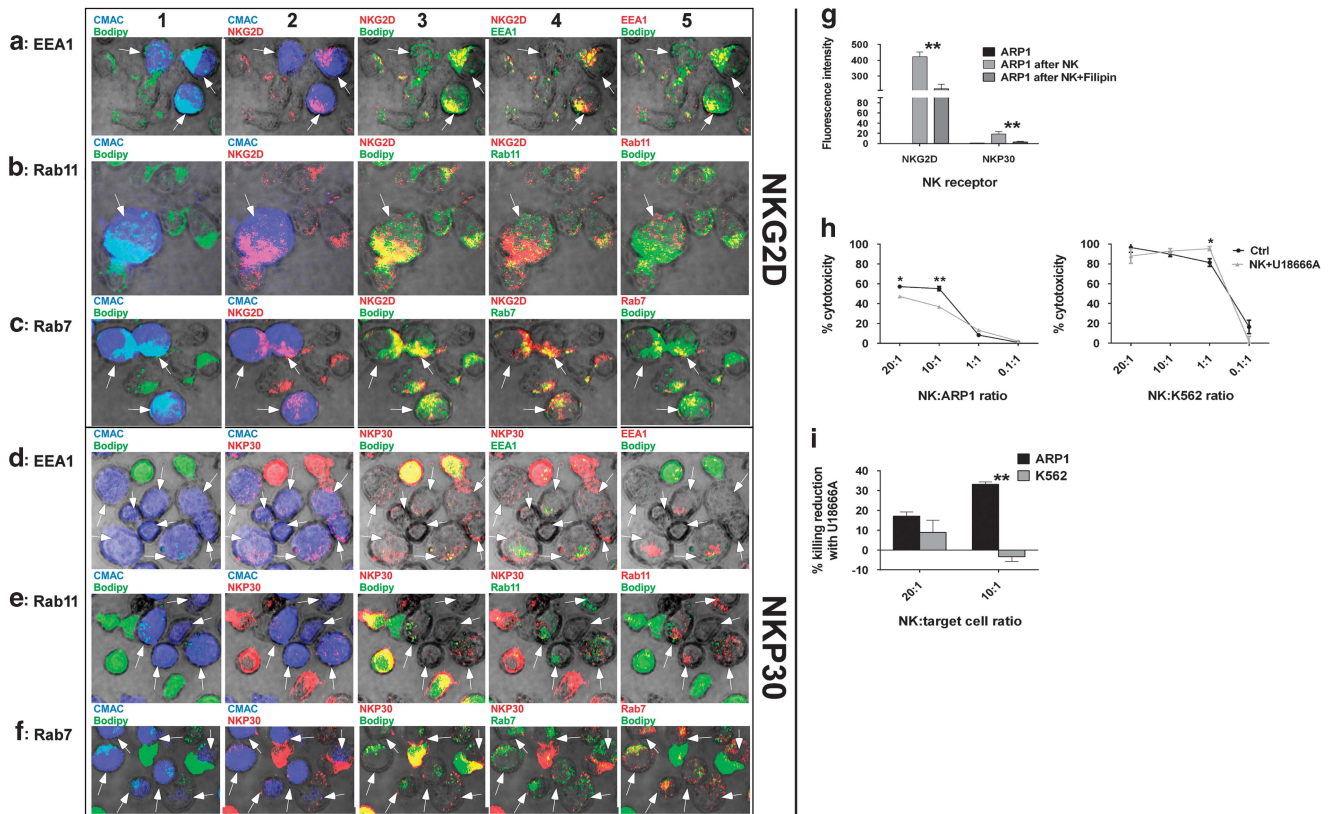


Figure 3 NKG2D and NKP30 are transferred into MM cells: (a) NKG2D and (b) NKP30 in ARP1 cells after co-culturing with CB-NK for 20 min. ARP1 cells were stained in blue (CMAC) and CB-NK in green (bodipy). NKG2D and NKP30 are indicated in red. All the images represent the same showing different combinations of the three colors. Arrows indicate ARP1 cells with NKG2D or NKP30. (c) Fluorescence values obtained from images in panels a and b. Bars represent mean \pm S.E.M. (d) As control, ARP1 cells alone were stained for NKG2D and NKP30 with the same settings than in images a and b. Results from panels a to d were confirmed in three different experiments. (e and f) Detection by flow cytometry of NKG2D and NKP30 in ARP1 cells after 24, 48 and 72 h of co-culture with CB-NK. ARP1 cells were gated based on CMAC staining. (e) Representative plot of the experiment at 24 h of co-culture. ARP1 cells unstained alone were used as control. (f) Median fluorescence intensity (MFI) values for NKG2D and NKP30 in ARP1 cells at the different times of co-culture, performing both surface and intracellular staining

investigated whether this pathway was responsible for the decreased ROS levels observed. Inhibition of GrB and Caspase-3 did not abrogate the decrease in ROS (Figure 6f), confirming that GrB and Caspase-3 in MM cells neither activate cell death nor ROS production. We investigated whether the decreased ROS after CB-NK treatment was only characteristic of MM cells. In other tumor cells analyzed (K562), there was also a decrease in ROS after CB-NK treatment (Figure 6g, $P < 0.0001$); however, CD19⁺ cells

from healthy individuals showed an increase in ROS after CB-NK (Figure 6g, $P < 0.0001$).

To determine whether NKG2D and NKP30 were involved in the decreased levels of lysosomes and ROS in MM cells after CB-NK exposure, these receptors were blocked, and lysosomes and ROS levels analyzed. Blocking either NKG2D (Figure 6h, $P < 0.0001$) or NKP30 (Figure 6i, $P < 0.0001$) abrogated the CB-NK effect on the decreased lysosome and ROS levels in MM cells after CB-NK exposure, indicating that



both receptors were involved in the cell death mediated by CB-NK.

MM stroma decreases CB-NK cytotoxicity. Because of the protective role of the stroma in MM, we analyzed the impact of MM stroma in CB-NK cytotoxicity. MM stroma reduced CB-NK cytotoxicity for MM cells (ARP1, RPM1 and KMM1, Figure 7a, $P < 0.05$). We observed that both MM cells and CB-NK adhered strongly to the stroma making difficult the formation of the immunological synapse between CB-NK and MM cells. However, we still observed vesicle transfer from CB-NK to MM cells and also secondary transfer between MM cells (Figure 7b). Because CXCL12 tethers hemopoietic cells into the stroma,²⁸ we measured CXCL12 secretion by the stroma and observed a reduced CXCL12 secretion after the addition of MM cells or MM cells and CB-NK (Figure 7c, $P < 0.05$), suggesting a possible role for CXCL12 in reducing the anti-MM cytotoxicity of CB-NK. Therefore we performed cytotoxicity assay adding AMD3100, which inhibits binding of CXCL12 to CXCR4 in hemopoietic cells. A slight recovery in CB-NK cytotoxicity versus RPM1 MM cell line was observed

($P = 0.09$, Figure 7d), providing evidence to support this possibility.

CB-NK cytotoxicity is transmissible between MM cells. We demonstrated lipid–protein trafficking between MM cells after CB-NK contact. Recipient MM cells appeared to become apoptotic with membrane blebbing (Figure 2g and Supplementary Movie S1). We examined whether NKG2D and NKP30 were secondarily transferred from ARP1 cells previously exposed to CB-NK (‘primary (1°) ARP1 cells’) to adjacent ARP1 cells (‘secondary (2°) ARP1 cells’). Bodipy-labeled CB-NK and CMAC-labeled ARP1 cells were co-cultured for 40 min; these ARP1 cells were defined as 1° ARP1 cells. Alive 1° ARP1 cells were then FACS-sorted. These sorted 1° ARP1 cells were then either left alone (1° ARP1 alone) or co-cultured (1° ARP1 with 2° ARP1) in cell pellet for another 40 min with unstained fresh 2° ARP1 (2° ARP1 after 1° ARP1) cells allowing differentiation between the two ARP1 cell populations (CMAC⁺ 1° ARP1 cells versus unstained 2° ARP1 cells). Then, in these two sets of ARP1 cells, we analyzed the expression of NKG2D and NKP30 (Figure 8a).

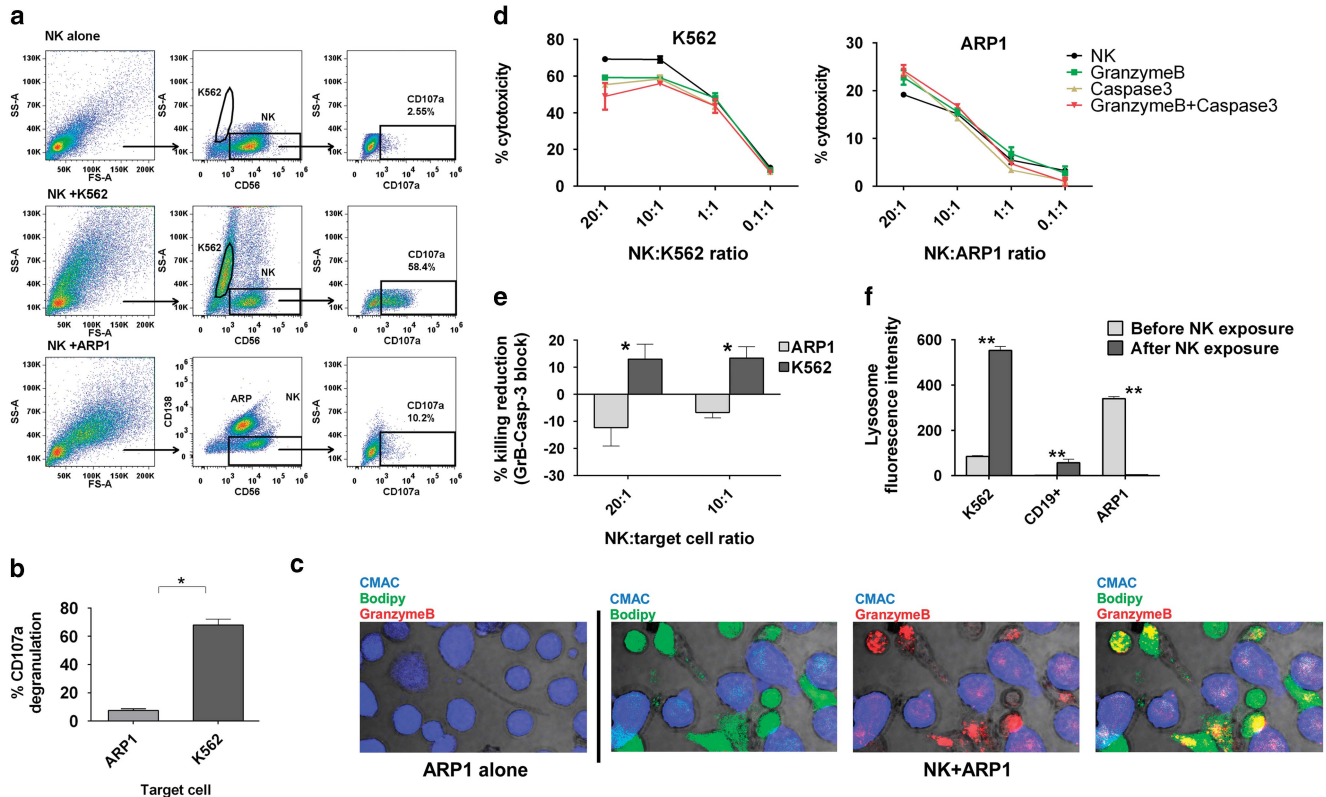


Figure 5 The mechanisms underlying CB-NK-mediated cytotoxicity differ between target cells: (a) Degranulation (CD107a expression) of CB-NK after 4 h of incubation with K562 or ARP1 cells. (b) Mean \pm S.E.M. values from a representative experiment shown in a ($n = 4$). (c) GrB expression in ARP1 cells after being co-cultured with CB-NK for 20 min. ARP1 cells were stained in blue (CMAC) and CB-NK in green (bodipy). GrB is indicated in red. (d) CB-NK cytotoxicity reduction after inhibiting GrB and Caspase-3. Killing is shown for control CB-NK, after inhibiting GrB in CB-NK, Caspase-3 in target cells and both, GrB in CB-NK and Caspase-3 in target cells. (e) Percentage of killing reduction after inhibiting GrB and Caspase-3 in four different CB units of the representative experiment shown in d. (f) Lysosome levels before and after CB-NK exposure for 40 min for K562 cells, CD19⁺ cells from a healthy donor and ARP1 cells. Bars represent mean \pm S.E.M. * $P \leq 0.05$. ** $P \leq 0.0001$

NGK2D (Figures 8b and d, $P < 0.0001$) and NKP30 (Figures 8c and e, $P < 0.0001$) were transferred secondarily to neighboring 2^o ARP1 cells (2^o ARP1 after 1^o ARP1 in Figure 8) from 1^o ARP1 cells, which had primarily synapsed with NK cells (1^o ARP1 with 2^o ARP1 in Figure 8). Co-staining for actin indicated that this transfer occurred through an immunological synapse between ARP1 cells, especially for NKG2D (Figures 8d and e).

As NKG2D and NKP30 were transferred to 2^o ARP1 cells, we also analyzed the downstream effects of this transfer on lysosomes and ROS levels and whether this effect was translated in some indirect cytotoxicity of CB-NK against these 2^o ARP1 cells. These 2^o ARP1 cells demonstrated decreased lysosome and ROS levels after contact with 1^o ARP1 cells (2^o ARP1 after 1^o ARP1: in Figures 8f and g, $P < 0.0001$), in comparison to the control sorted ARP1 cells (ARP1 ctrl: in Figures 8f and g). We also noticed a compensating effect between the two populations of ARP1 cells, as after 1^o ARP1 cells were exposed to CB-NK and then incubated with 2^o ARP1 cells, the 1^o ARP1 cells began to normalize their loss in lysosomes and ROS (1^o ARP1 alone versus 1^o ARP1 with 2^o ARP1: in Figures 8f and g, $P < 0.0001$).

To determine whether CB-NK cytotoxicity was also transmitted between ARP1 cells, we performed cytotoxicity

assay by adding 1^o ARP1-sorted cells as effectors and 2^o ARP1 cells as targets. 1^o ARP1 cells, which previously synapsed with CB-NK, demonstrated 7% cytotoxicity against 2^o ARP1 cells at a 40:1 effector:target ratio. Although this cytotoxicity decreased at lower effector:target cell ratios, it was greater than the cytotoxicity of the control ARP1 cells untouched by CB-NK (Figure 8h), indicating a transmissible cell death from CB-NK to MM cells and then from these MM cells to neighboring MM cells. Finally, we tested whether the MM stroma impacted on this indirect 'domino-effect' cytotoxicity. Cytotoxicity assay containing MM stroma reduced indirect cytotoxicity of 1^o ARP1 versus 2^o ARP1 at 20:1 effector:target cell ratio (Figure 8i), suggesting that MM stroma may reduce 'domino-effect' cytotoxicity.

Discussion

We show a novel CB-NK-mediated cytotoxicity against MM cells that is GrB and Caspase-3 independent, involves transmissible cytotoxicity between CB-NK and MM cells via lipid-protein vesicle trafficking and secondarily transferred from these MM cells to neighboring MM cells. This lipid-protein transfer involves NKG2D and NKP30 receptors, causes lysosomal cell death and reduced ROS levels. This cytotoxicity appears to be unique of CB-NK and different from

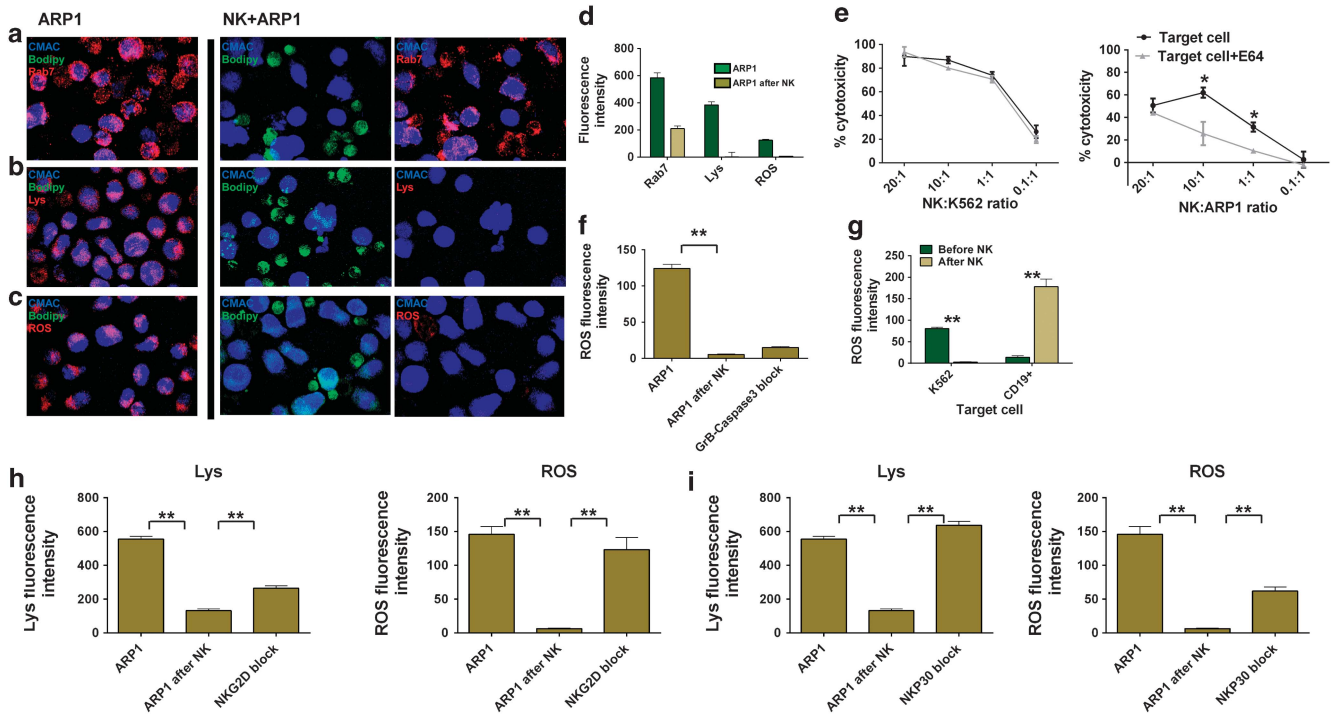


Figure 6 CB-NK induce lysosomal cell death in MM cells, and NKG2D and NKP30 are involved in this mechanism: (a–d) Analysis of lysosomes (Rab7 and Lyso-tracker: Lys) and ROS production in ARP1 cells before and after co-culture in cell pellet with CB-NK for 40 min. ARP1 cells were labeled in blue (CMAC), CB-NK in green (bodipy) and each marker is indicated in red. (a) Late endocytic and lysosome marker Rab7. (b) Lysosome marker (Lyso-tracker). (c) ROS production. (d) Mean \pm S.E.M. values of slides analyzed from a to d. Results from a to c were confirmed in three different experiments. (e) CB-NK cytotoxicity reduction after incubating target cells (K562 and ARP1) with E64 (a cysteine cathepsin inhibitor). (f) ROS variation after inhibiting GrB in CB-NK and Caspase-3 in ARP1 cells and co-culturing for 40 min. (g) ROS levels before and after CB-NK exposure for 40 min for K562 cells and CD19⁺ cells from healthy individuals. (h and i) Variation of lysosome (Lys) and ROS levels after CB-NK exposure for 40 min (ARP1 after NK) and after blocking (h) NKG2D and (i) NKP30 in CB-NK (NKG2D block and NKP30 block). Lysosomes were determined with the Lyso-tracker. Bars represent mean \pm S.E.M. ** $P \leq 0.0001$

that seen against K562 cells where NKG2D and NKP30 do not have such a relevant role.

Lysosomal cell death happens after ruptured lysosomes release cathepsins leading to Caspase-3-independent cell death, which can show membrane blebbing, and which, depending on the magnitude of lysosomal permeabilization, will activate an apoptotic-like programmed cell death or necrosis cell death.²⁹ Lipids and Granulysin are involved in this type of cell death as lipids destabilize lysosomes³⁰ and Granulysin mediates a Caspase-3-independent lysosomal cell death with membrane blebbing.⁸ These studies support our findings where we show a transmissible MM cell death mediated by CB-NK, involving lipid–protein transfer with a decrease in lysosomes, cell death dependent on Cathepsins and showing membrane blebbing in MM cells. NKG2D and NKP30, associated with the decreased lysosomes levels, were also indirectly transferred to new MM cells indicating their possible role in this indirect cytotoxicity. Although no measurement of Granulysin was made, our findings do not discard a possible role of Granulysin in this lysosomal cell death.

Cells exchange information via exosomes/lipid vesicles, and NK release cytotoxic exosomes/vesicles expressing NK-activating receptors.^{3,31} Lipids inhibit autophagy³² and can activate ER stress-associated cell death.³³ Here we show lipid trafficking between cells and that this exchange of

‘information’ in lipid structures, which contain NKG2D and NKP30, occurs between CB-NK and MM cells and afterwards between adjacent MM cells being associated with cytotoxicity against MM cells. Even the bone marrow stroma reduced this CB-NK direct and indirect cytotoxicity, it was not an impediment for this lipid transfer from CB-NK to MM cells and secondarily to neighboring MM cells.

GrB-induced NK death is usually accompanied by increased ROS production, a process which is Caspase-3 dependent.⁴ We found that even though GrB is transferred into ARP1 cells by CB-NK, the GrB and Caspase-3 pathway was not involved neither in the killing of ARP1 cells nor in the decreased ROS levels detected. By comparison, NKG2D- and NKP30-associated MM cell killing was associated with decreased ROS, providing evidence for a role for these receptors. As ROS is associated with survival in tumor cells and with a lowering of the interaction between cancer cells and the immune system,^{34,35} this CB-NK-associated ROS reduction might make tumor cells more vulnerable to cell-based immunotherapeutic strategies.

MM cells, which are susceptible to ER stress cell death, have high autophagy levels.^{18,36} Autophagy uses lysosomes for degradation,³⁷ associates to increased ROS³⁸ and contributes to chemotherapeutic resistance.¹⁷ We observed decreased lysosomes and ROS in MM cells after CB-NK

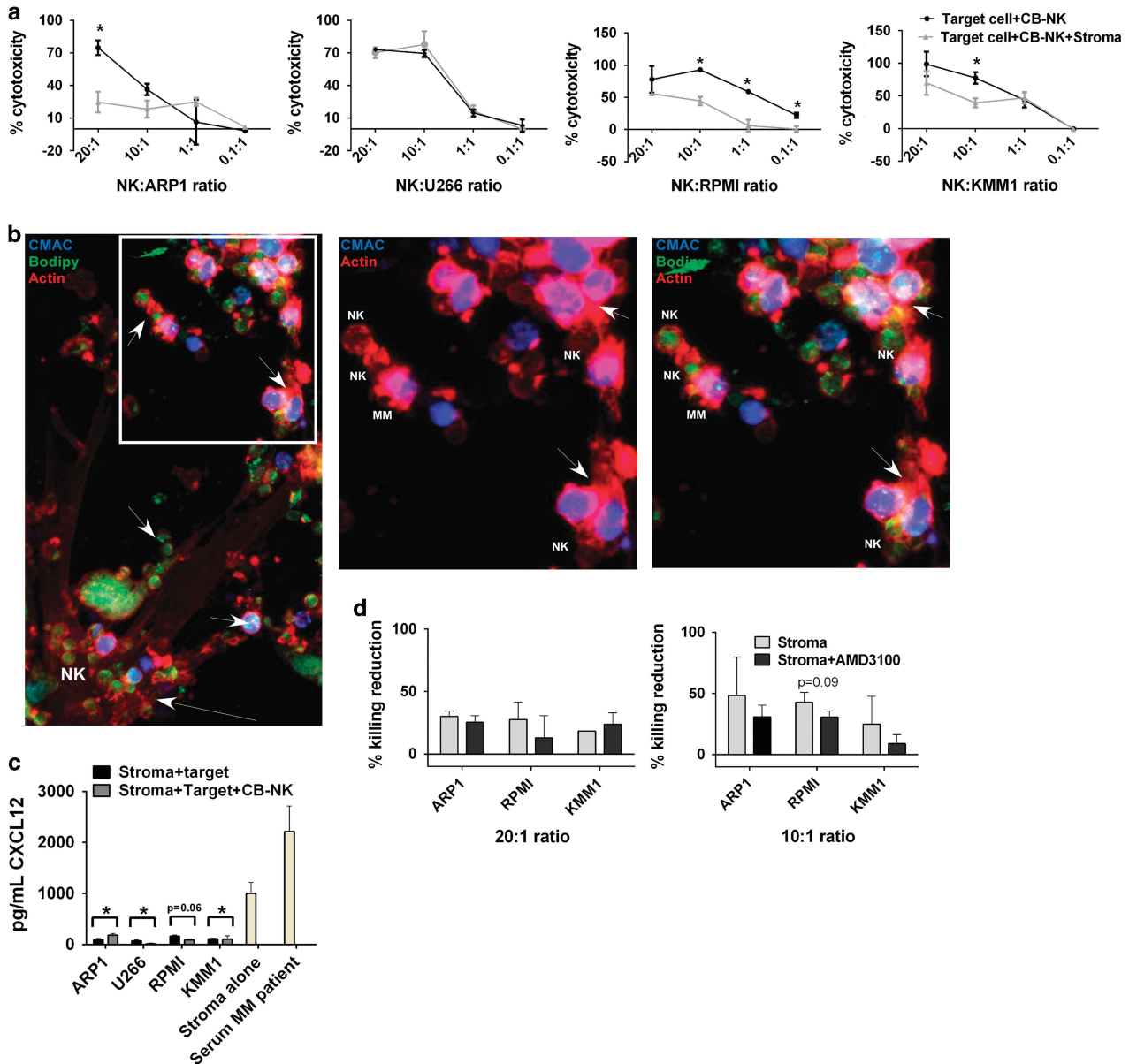


Figure 7 MM stroma decreases CB-NK cytotoxicity: (a) CB-NK cytotoxicity reduction after adding MM stromal cells into cytotoxicity assays. Cytotoxicity assays for MM cells were done in parallel either adding (Target cell + CB + NK + Stroma) or not adding (Target cell + CB-NK) MM stromal cells to the assay. (b) Confocal fluorescence image of stromal MM cells with CB-NK (in green with bodipy) and KMM1 MM cells (in blue with CMAC). Actin is shown in red. Stromal cells can be visualized by the actin filaments. Arrows in the lower part show CB-NK adhered to stromal cells. Arrows in the upper part show transfer of CB-NK content to MM cells and also secondarily between MM cells. The white square is shown amplified in the right side showing either two colors (CMAC and actin) or the three colors together (CMAC, bodipy and actin). Same images were confirmed for other MM cell lines. (c) CXCL12 levels in the supernatant of cytotoxicity assays with MM cells adding either stroma with target MM cells and CB-NK (Stroma + target + CB-NK) or stroma with target cells alone (Stroma + target). Serum MM patient was added as positive control. *P*-values represent analysis of stroma alone versus stroma + target. (d) Cytotoxicity CB-NK reduction at 20:1 or 10:1 effector:target ratio, after adding (Stroma + AMD3100) or not adding (Stroma) AMD3100 to the cytotoxicity assay with stroma, CB-NK and MM cells. **P* ≤ 0.05

exposure, which could suggest decreased autophagy levels. Although an extensive study of autophagy markers was not undertaken here, the possible autophagy reduction in MM cells by CB-NK could reduce the refractoriness, which happens in some MM patients after PI treatment.^{17,19}

In summary, we show a unique mechanism of CB-NK cytotoxicity that involves transmissible cytotoxicity between cells, where lipids appear as 'vehicles' to perform this mechanism. This opens new pathways for the use of CB-NK

as a cellular immunotherapy option and for developing therapeutic strategies using CB-NK-derived lipids.

Material and Methods

Ethics statement. All research involving human materials was approved by the MD Anderson Cancer Center (MDACC) Institutional Review Board (IRB). CB units were obtained from healthy donors who gave written informed consent. All animal work was performed under an MDACC Institutional Animal Care and Use Committee (IACUC)-approved protocol specific to this study.

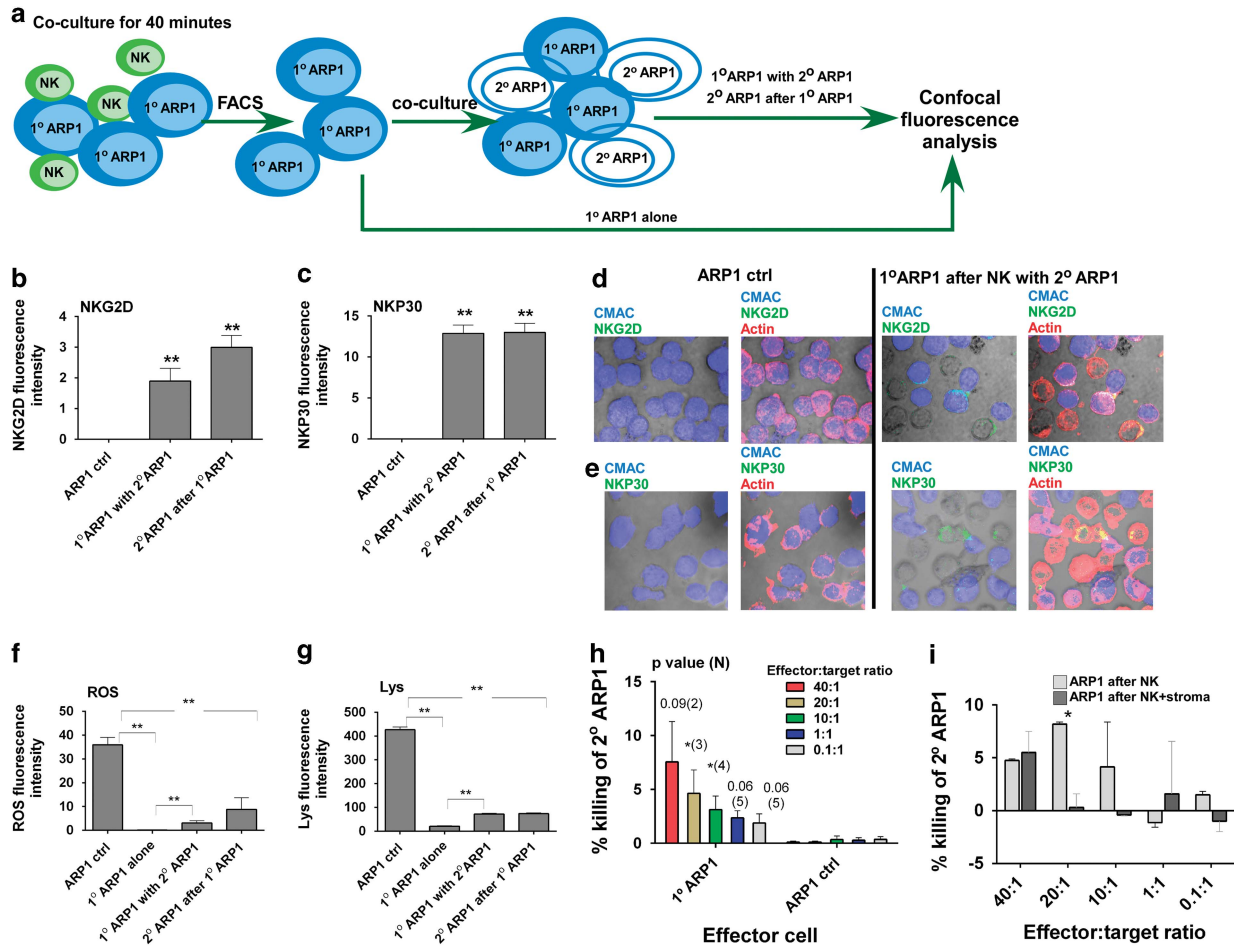


Figure 8 CB-NK cytotoxicity is transmissible between MM cells: (a) Design experiment for results from b to h. Bodipy-labeled CB-NK and CMAC-labeled ARP1 cells (1° ARP1 cells) were co-cultured in cell pellet for 40 min, then 1° ARP1 cells were FACS sorted and half of them were left alone (1° ARP1 alone), and half of them (1° ARP1 with 2° ARP1) were co-cultured in cell pellet for 40 min with fresh unstained ARP1 cells (2° ARP1 cells after 1° ARP1), then confocal fluorescence experiments and analyses were performed discriminating between the two sets of ARP1 cells based on 1° ARP1 cells labeled in blue (CMAC) and 2° ARP1 cells unstained. (b and c) NKG2D (b) and NKP30 (c) expression in 1° ARP1 cells and 2° ARP1 cells. Bars represent mean \pm S.E.M. (d and e) Representative images of NKG2D (d) and NKP30 (e) expression in green in both sets of ARP1 cells of values obtained in b and c. Actin is shown in red. Lysosome (f) and ROS (g) expression in the two sets of ARP1 cells. *ARP1 ctrl* indicates ARP1 cells alone sorted as control for the experiments. Bars represent mean \pm S.E.M. Results from b to g were confirmed in three different experiments. (h) Cytotoxicity assay performed adding as effector cells 1° ARP1 sorted cells and as targets fresh ARP1 cells. Effectors were added at different effector:target cell ratios. In parallel, ARP1 cells alone were sorted and added as effectors to the cytotoxicity assay as control. (i) Cytotoxicity reduction of 1° ARP1 (as effectors) versus fresh ARP1 cells (as targets) after adding (ARP1 after NK + stroma) or not adding (ARP1 after NK) stroma to the assay. * $P \leq 0.05$. ** $P \leq 0.0001$

Cell cultures. K562-based antigen-presenting cells expressing membrane-bound IL-21 (‘Clone 9.mblL21’) was kindly provided by Dean A Lee and Laurence JN Cooper (MDACC, Houston, TX, USA). CB-NK were obtained using methods as previously described.²⁰ Briefly, NK cells were MACS-selected (Miltenyi Biotec, San Diego, CA, USA) from CB mononuclear cells (CBMC) and co-cultured with irradiated Clone 9.mblL21 cells at a ratio of 2:1 (Clone 9: NK cells). IL-2 (Proleukin; Chiron, Emeryville, CA, USA) was added at 400 U/ml every other day. At day 14, CB-NK were harvested for all the experiments. Culture media was comprised of 45% RPMI-1640 (Cellgro, Manassas, VA, USA) and 45% Click’s media (Irvine Scientific, Santa Ana, CA, USA) supplemented with 10% AB human serum (Atlanta Biologicals, Lawrenceville, GA, USA). ARP1 cell line was kindly provided by Multiple Myeloma Research Center (Little Rock, AK, USA) to Lymphoma/Myeloma Department (M.D. Anderson Cancer Center, Houston, TX, USA). K562, U266, RPMI and KMM1 cell lines were obtained from American Type Culture Collection (ATCC, Rockville, MD, USA). K562, ARP1 and KMM1 were cultured in RPMI supplemented with 10% fetal bovine serum (FBS) and U266 with 15% FBS. CD19⁺ cells from healthy donors and CD138⁺ primary cells from MM patients were obtained by MACS selection (Miltenyi Biotec). Peripheral blood NK (PB-NK) from healthy individuals and NK from MM patients (MM-NK) were

obtained after doing Ficoll in peripheral blood and afterwards MACS selection of NK cells (Miltenyi Biotec).

Luminescence and fluorescence *in vivo* imaging. Generation of green fluorescent protein-Firefly luciferase (GFP-FFLuc)-expressing ARP1 cell line for *in vivo* experiments was previously reported.²⁰ Mice were anaesthetized and subjected to serial bioluminescence imaging as previously described²⁰ using a Xenogen IVIS 200 system (Caliper, Waltham, MA, USA). After euthanizing the mice, tumor areas were visually inspected and then visualized also under a fluorescent stereo microscope Leica M2025 FA (Leica Microsystems Inc, Buffalo Grove, IL, USA). The Lymphoma/Myeloma Department gave us advice to set up the *in vivo* MM model using ARP1 cell line.

Cytotoxicity assays. Two different techniques were used to label target cells: either with ⁵¹Cr (Amersham Pharmacia Biotech, Piscataway, NJ, USA) for 2 h or with BATDA reagent (Perkin-Elmer, Waltham, MA, USA) for 10 min, then washed and added to effector cells (NK) in triplicate into 96-well round bottom plates (Corning, Corning, NY, USA) at different NK:target cell ratios. After 4 h, supernatants were transferred either to a Luma-Plate-96 (Perkin-Elmer) for ⁵¹Cr or

to White flat bottom plates with Delfia Europium solution. ^{51}Cr or Batda-Europium release by target cells after lysis was measured either on a TOP Count NXT microplate scintillation and luminescence counter (Perkin-Elmer) for ^{51}Cr or in a fluorescence reader (Tecan, Männedorf, Switzerland). Cytotoxicity was determined by the formula: cytotoxicity = (sample value – spontaneous lysis)/(maximum lysis – spontaneous lysis) \times 100%. For blocking or inhibition assays, all reagents were added for 30 min at 37 °C either to NK or target cells before the cytotoxicity assay. U18666A was added to NK for 2 h at 37 °C previous to the cytotoxicity assay. Anti-NKG2D (clone 1498810) or anti-NKP30 (clone p30-15) were added to NK cells at 5 $\mu\text{g}/\text{ml}$. GranzymeB inhibitor Z-AAD-CMK (Calbiochem, San Diego, CA, USA) was added to NK at 600 nM. Caspase-3 inhibitor Z-DEVD-FMK (BD Biosciences, San Jose, CA, USA) was added to target cells at 20 μM . Cathepsin inhibitor E64 (Sigma-Aldrich, St. Louis, MO, USA) was added to target cells at 10 μM . The lipid raft inhibitor Filipin III (Sigma-Aldrich) was added to NK cells. Cholesterol synthesis inhibitor U18666A (Sigma-Aldrich) was added to NK cells at 5 $\mu\text{g}/\text{ml}$. The CXCL12-CXCR4 inhibitor AMD3100 (Sigma-Aldrich) was added at 40 μM . For calculation of cytotoxicity reduction after blocking NK receptors, inhibiting GrB and Caspase-3 pathway or adding Filipin III, U18666A, E64 or AMD3100, values were calculated relative to control NK cells without blocking any receptor or without inhibiting any pathway.

Stromal cells and ELISAs. BM stromal cells (BMSC) from MM patients were obtained from BM aspirate after doing Ficoll to isolate mononuclear cells, expanded *in vitro* in DMEM + 10% FBS till they were 85% confluent and then the night before to the cytotoxicity assays they were added into 96-well plates at 10 000 cells/well where the cytotoxicity assay was performed. CXCL12 levels in culture supernatant were measured with Human CXCL12/SDF-1 alpha Quantikine ELISA Kit (R&D Systems, Minneapolis, MN, USA).

Confocal fluorescence microscopy. Target cells were stained with CellTracker Blue CMAC (7-amino-4-chloromethylcoumarin) and NK were stained with CellTracker Green Bodipy dye (8-Chloromethyl-4,4-Difluoro-1,3,5,7-Tetramethyl-4-Bora-3a,4a-Diaza-S-Indacene) (Invitrogen, Life Technologies, Eugene, NY, USA) for 30 min at 37 °C in serum-free media. After washing, cells were mixed at a 1 : 1 ratio in 1.5-ml tubes, spun at low speed and allowed to co-culture in cell pellet at 37 °C for 10, 20 or 40 min, depending on the experiment. For the 24–72 h experiments, cells were added into a 24-well plate at 1 million cells/ml and then transferred onto microscope slides using a cell concentrator (Cytofuge 2, IRIS International, Chatsworth, CA, USA), fixed with 3% methanol-free formaldehyde and then permeabilized. For immunostaining, primary antibodies were incubated for 30 min at 37 °C and secondary antibodies were incubated for 20 min at 37 °C. To visualize CB-NK and MM cells with the stroma, BMSC from MM patients were added over microscope slides in well chambers the night before of the experiment to allow the stroma to adhere to the slides. Next day, CB-NK (stained in green with bodipy) and MM cells (stained in blue with CMAC) were added and allowed to co-culture for 40 min, then slides were fixed, permeabilized and immunostained for actin (Phalloidin) as described above.

Primary antibodies used in confocal fluorescence microscopy were GrB-Alexa Fluor-647, NKP30-PE and NKP30-Alexa Fluor-647 from BD Biosciences, NKG2D-Alexa Fluor-700 (R&D Systems), EEA1, Rab7 (D95F2), Rab11 (D4F5) from Cell Signaling Technology (Boston, MA, USA), Phalloidin-Alexa Fluor-594 and Phalloidin-Alexa Fluor-647 from Invitrogen (Invitrogen, Life Technologies). Secondary antibody used was Anti-Rabbit IgG-Alexa Fluor-647 from Cell Signaling. For ROS and lysosome detection, Cell ROX Deep Red Reagent and Lyso Tracker Deep Red (Molecular Probes, Invitrogen, Life Technologies) were added into the media at 3 μM and 75 nM, respectively.

Images were acquired using an Olympus IX81 microscope (Olympus Corporation, Center Valley, PA, USA). Image acquisition was performed using 405, 488 and 633 lasers for excitation, doing Z-stack acquisition images and applying the corresponding filters for each dye used. Image analysis was performed with the software FluoView Viewer FV-ASW version 04.00.02.09 (Olympus Corporation). To obtain fluorescence intensity values, four different views/slide were analyzed, quantifying 100 cells/view. Numerical values obtained were then used to perform statistical analysis with SPSS statistics, version 19 (IBM, Chicago, IL, USA).

Time-lapse *in vivo* imaging. Time-lapse *in vivo* imaging was performed by staining target and effector cells with CMAC and Bodipy as described above. Cells were added together into an eight-well plate and imaged using the 405- and 488-nm wave length lasers for 2 h performing one image acquisition/minute.

Flow cytometry. To visualize tumor cells and CB-NK in the different tissues in mice by flow cytometry and degranulation of CB-NK *in vitro*, 10^6 cells/ml were stained for 30 min at 4 °C. The following conjugated antibodies were used: NKG2D-PeCy7, CD16-PeCy5, CD138-APC, CD56-V450, CD56-PeCy5, and CD107a-V450 from BD Biosciences. Data were acquired by the Beckman Coulter Gallios device (Beckman Coulter, Inc., Brea, CA, USA). Analysis was performed using the FlowJo (Tree Star, Ashland, OR, USA) software.

FACS sorting. To FACS-sort ARP1 cells that had been primarily synapsed with CB-NK, ARP1 cells stained in blue with CMAC and CB-NK in green with Bodipy were allowed to co-culture in cell pellet for 40 min. Then, propidium iodide was added at 3 $\mu\text{g}/\text{ml}$ and cells were sorted in a BD FACSAria III cell sorter (BD Biosciences), based on FSC/SSC properties, single cells, viability (propidium iodide exclusion) and CMAC⁺ positivity. Sorted cells were used afterwards for cytotoxicity assays and confocal fluorescence microscopy experiments.

Indirect cytotoxicity. For indirect cytotoxicity experiments, sorted ARP1 cells that had been co-cultured in cell pellet with CB-NK were added as effector cells in a cytotoxicity assay with fresh ARP1 cells (never exposed to CB-NK) as target cells at different effector:target cell ratios. As a negative control, ARP1 cells untouched by CB-NK were sorted and used in parallel also as effector cells. For the stroma experiment, in parallel ARP1 cells were added to a plate with BMSC from MM patients previously added the night before. Cytotoxicity assays were performed as described above.

Statistical analysis. Pearson's correlation test was performed to analyze correlation between MFI for NKG2D and disease progression in mice. Mann–Whitney *U* test was used to analyze the comparison between groups for the confocal fluorescence images, flow cytometry and cytotoxicity assays. Paired *T*-test was performed to analyze the difference in the cytotoxicity assay for different groups in the individual experiments. Mann–Whitney *U* test was used to analyze the comparison between different groups in cytotoxicity assays for the analysis of all different experiments together. All statistical analyses were performed using either SPSS statistics, version 19 or GraphPad Prism, version 6.02 (GraphPad Software, Inc., San Diego, CA, USA).

Conflict of Interest

The authors declare no conflict of interest.

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