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Expansion of allogeneic NK cells with efficient antibody-dependent cell cytotoxicity against multiple tumor cells

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Running title: Production of NK cells with ADCC
Abstract

Monoclonal antibodies (mAbs) have significantly improved the treatment of certain cancers. However, in general mAbs alone have limited clinical activity. One of their main mechanisms of action is to induce antibody-dependent cell-mediated cytotoxicity (ADCC), which is mediated by natural killer (NK) cells. However, most cancer patients present severe immune dysfunctions affecting NK activity. This could be circumvented by the injection of allogeneic, expanded, NK cells, which is safe. Nevertheless, despite the strong cytolytic potential of expanded NK cells against different tumors, clinical results have been poor.

Methods: We thus propose the combination of allogeneic NK cells and mAbs to improve cancer treatment. We generated expanded NK cells (e-NK) with strong in vitro and in vivo ADCC response against different tumors and mediated by different therapeutic mAbs, i.e. rituximab, obinutuzumab, daratumumab, cetuximab and trastuzumab.

Results: Remarkably, e-NK cells can be stored frozen and after thawing armed with mAbs. They mediate ADCC through degranulation dependent and independent mechanisms. Furthermore, they overcome some of the anti-apoptotic mechanism found in leukemic cells.

Conclusion: In summary, we have established a new protocol for activation/expansion of NK cells with high ADCC activity. The use of mAbs in combination with e-NK cells could potentially improve cancer treatment.
**Introduction**

Recent progresses in cancer treatment are mainly related to the development of new-targeted therapies [1]. These require the identification of suitable targets that are mainly expressed by the tumor cell population and/or playing a critical role for neoplastic cell development. In this context, monoclonal antibodies (mAbs) have improved the treatment of certain cancers. Indeed, rituximab (RTX), an IgG1 mAb directed against CD20 antigen, has now become the backbone for the treatment of most B-cells chronic lymphocytic leukemia (B-CLL) and B-cells non-Hodgkin’s lymphoma (B-NHL) patients. The combination of RTX with conventional chemotherapy has shown better efficacy in randomized clinical trials. This success has also been found with other cytotoxic mAbs such as trastuzumab in breast cancer or cetuximab in colorectal carcinoma and squamous cell carcinoma of the head and neck [2, 3]. However, it is remarkable to note that the mAbs themselves have modest clinical activity. For example, RTX or obinutuzumab (OBZ; previously GA101, Roche, Genentech) when used as monotherapy in patients with relapsed follicular lymphoma (FL) have demonstrated short progression-free survival (PFS) [4]. These data indicate that there is a need to optimize their use in co-therapy. RTX success is related to its capacity to induce Fc-antibody-dependent cell-mediated cytotoxicity (ADCC). One receptor for human IgG1 is FcγRIIIa (CD16a), which is expressed on natural killer (NK) cells and macrophages. The influence of FcγRIIIa-158VF polymorphism on RTX clinical response strongly suggests that ADCC is critical [5]. Based on these results, there has been an attempt to produce new anti-CD20 mAbs that exhibit higher affinity for FcγRIIIa either by Fc mutations or by glycoengineering [4, 6]. This later strategy, leading to low fucose content of the N-glycan, is currently under clinical investigations in B-cell malignancies with the mAb OBZ, which shows stronger ADCC *in vitro* and in a lymphoma xenograft mouse model compared to RTX and improved clinical activity for treating B-CLL [4]. This clinical benefit has been observed in other B-cell malignancies [4]. OBZ is approved for first-line B-CLL in association with chlorambucil and in combination with bendamustine for the treatment of patients with FL who relapse or are refractory to RTX-containing regimen [4]. Initial results show that lenalidomide, which stimulates NK cell activity [7], activates NK cells in OBZ-treated patients [8].

Hematological cancer patients possess antitumor NK cells that are unable to control disease [9, 10]. Blood-borne tumor cells use different mechanisms for immune escape [11, 12], e.g. inducing NK cell dysfunction [7, 13]. This mechanism has also been observed in a variety of solid tumor patients [3]. In addition, NK cell differentiation may be inhibited by the presence of tumor cells e.g. acute myeloid leukemia (AML) cells infiltrating bone marrow [14, 15].
Therefore, the failure of mAb as monotherapy could be related to impaired NK cell function and hence, there is a clinical interest to reactivate or replace patient NK cells [16]. Clinical-grade production of allogeneic NK cells is efficient and NK cell–mediated therapy after hematopoietic stem cell transplantation (HSCT) seems safe [15, 17, 18]. However, despite the strong cytolytic potential of expanded NK cells against different tumors, clinical results have been very limited [15, 17, 18].

The combination of allogeneic NK cells with mAb could improve cancer treatment by replacing the defective effector immune cells. In addition, mAbs would effectively guide these effectors to their tumor targets. Several groups have tried this combination with miscellaneous results that could be due to deficient CD16 expression or lack of proper activation of expanded NK [19-22]. In addition, these works did not include a systematic study of the effect of these cells in combination with several mAbs on different tumors, and also they did not include primary tumor cells.

The aim of this work is to generate allogeneic NK cells with strong ADCC response against different tumors and mediated by different therapeutic mAbs. In addition, NK cell production should be easily scaled up and developed with good manufacturing practices (GMP). We have produced umbilical cord blood (UCB)-derived NK cells because UCB are rapidly available, present low risk of viral transmission and have less strict requirements for HLA matching and lower risk of graft-versus-host disease (GvHD) [17]. For NK cell expansion we used Epstein–Barr virus (EBV)-transformed lymphoblastoid B cell lines as accessory cells, which induce a unique NK cell genetic reprogramming [23], generating effectors that overcome the anti-apoptotic mechanism of leukemic cells [24] and that are able to eliminate tumor cells from patients with poor prognosis [25]. We show that NK cells obtained with our protocol are able to perform ADCC \textit{in vitro} and \textit{in vivo}. The ADCC response was induced by different therapeutic antibodies and against multiple target cells.
Materials and Methods

Ethical statement

Experimental procedures were conducted according to the European guidelines for animal welfare (2010/63/EU). Protocols were approved by the Animal Care and Use Committee “Languedoc-Roussillon” (approval number: CEEA-LR-12163). The use of human specimens for scientific purposes was approved by the French National Ethics Committee. All methods were carried out in accordance with the approved guidelines and regulations of this committee. Written informed consent was obtained from each patient prior to surgery.

Chemicals

The D1D2 peptide has been previously described [26]. IL-2 and IL-15 were obtained from Miltenyi Biotec. To produce deglycosylated cetuximab, a commercial cetuximab solution was treated overnight with PNGaseF (Promega) at 37°C in 50 mM sodium bicarbonate buffer (pH 7.8) at 125 U/mg of cetuximab. Deglycosilated cetuximab was purified by gel filtration using a Sephadex 75 column in PBS, and sterilized by filtration. The AF 647 Goat F(AB’)2 anti human IgG (H+L) min x (BOV,HRS,MS) was from Interchim.

B-CLL patients

Data and samples from patients were collected at the Clinical Hematology Department of the CHU Montpellier, France, after patient’s written consent and following French regulations. Patients were enrolled in two independent clinical programs approved by the “Comités de Protection des Personnes Sud Méditerranée I”: ref 1324 and HEMODIAG_2020 (ID-RCB: 2011-A00924-37). Samples were collected at diagnosis and kept by the CHU Montpellier [10, 27]. For analysis peripheral blood mononuclear cells (PBMCs) were obtained by ficoll gradient and stored frozen in liquid nitrogen until use. The percentage of CD19⁺CD5⁺ cells was always higher than 90%. Other samples from hematological cancer patients were obtained from the same collections.

Cell lines

The (EBV)-transformed lymphoblastoid B cell line PLH, the hematopoietic cell lines HL60 and MV4-11 (acute myeloid leukemia), Daudi and Raji (Burkitt’s lymphoma derived, CD20⁺), K562 (erythroleukemia), MM.1S and U266 (multiple myeloma), MEC1 (B-chronic lymphocytic leukemia) and its variants MEC-1-BCL-XL and MEC-1-MCL-1 were cultured in 10% FBS RPMI medium. The adherent cell lines Calu-1 (lung cancer), A549 (lung
adenocarcinoma), SK-OV-3 (ovarian carcinoma) and SKBR3 (breast adenocarcinoma) were cultured in 10% FBS DMEM medium.

**PBMC and UCBMC purification**
UCBs were obtained from healthy donors from the CHU Montpellier. PBMC and UCB mononuclear cells (UCBMC) were respectively collected from peripheral blood samples and UCB units using Histopaque -1077 (Sigma). Briefly, 13 ml Histopaque were added to 50 ml centrifugation tubes and 30 ml of 1/2 diluted blood in RPMI, (Invitrogen) were slowly added at the top. Tubes were centrifuged at 1600 rpm for 30 minutes at 20 °C without break. Mononuclear cells were collected from the interlayer white ring, washed in RPMI and suspended in RPMI medium supplemented with 10% FBS (Invitrogen).

**Isolation and activation of human NK cells**
Frozen UCBMCs were depleted of T cells by using EasySep™ CD3 Positive Selection Kit (STEMCELL technologies). Cells were cultured for 10 to 20 days with γ-irradiated PLH cells at 1:1 NK cell:accessory cell ratio in presence of IL-2 (100 U/ml) and IL-15 (5 ng/ml), or with IL-2 alone (1000 U/ml). PLH cells were added every four days and fresh cytokines every two days. At the end of the process NK cell purity (CD56+/CD3-) was always higher than 90%.

**FACS analysis**
For phenotype analysis, cells were stained with 7AAD (Beckman) to identify viable cells and antibodies against the surface markers CD25-FITC, CD45RO-FITC, CD69-PE, CD62L-PE, CD19-PE, CD3-PE, CD19-ECD, CD56-PECy7, CD56-APC, CD3-APC, CD45-APCAlexaFluor750, CD45RA-APCAlexaFluor750, CD16-PacificBlue, CD57-PacificBlue, CD45-KromeOrange, CD16-KromeOrange (Beckman), CD158b-FITC, CD158a-PE, CD107a-HV500 (BD Biosciences), CD158e-Vioblue (Miltenyi). 1x10^5–3x10^5 cells were incubated for 20-30 minutes at 4 °C with different antibodies in PBS containing 2.5% FBS. Cells were then washed and suspended in 200-250 µl of the same media. Staining was analyzed on a Gallios flow cytometer (Beckman) using the Kaluza software. Alive lymphocytes were gated using FSC/SSC and 7AAD staining. B lymphocytes (CD19+), T lymphocytes (CD3+CD56-) and NK cells (CD56+CD3-) were distinguished using respectively CD19, CD3 and CD56 antibodies.

**NK cell mediated cytotoxicity**
Fresh or frozen (stored in liquid nitrogen) NK cells were labeled with 3 µM of CellTracker™ Violet BMQC Dye (Life Technologies) and incubated overnight with target cells at different E:T ratios. Subsequently, phosphatidylserine (PS) translocation and membrane damage were analyzed in the violet fluorescence negative target cell population by flow cytometry using Annexin V-FITC (Immunostep) and 7AAD (BD Biosciencies) or propidium iodide (PI) as previously described [24, 28]. We consider all cells positive for annexin-V and/or PI (or 7-ADD) as death (or dying).

In ADCC experiments, we incubated target cells with the relevant antibodies (RTX and OBZ at 10 µg/ml; daratumumab, cetixumab and trastuzumab at 5 µg/ml) for 30 minutes at 37°C. When we armed NK cells, we incubated them at the same concentration of antibodies before washing and incubation with target cells. EGTA was used at 1 mM to block the granular exocytosis pathway and MgCl₂ at 1.5 mM to maintain the osmotic pressure.

We used the tetrazolium dye MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazonium bromide) to determine cellular viability. We added 10 µL of MTT (5 mg/ml) to the adherent cells (100 µL of medium after 2 washes with PBS) and we incubated them for 1 h at 37°C, and added 100 µL of 0.05 M HCL in isopropanol to dissolve the crystals and quantified color at 550 nm in a spectrophotometer.

In all experiments we calculated the basal cell death in the absence and presence of the different mAbs. These values were subtracted from those obtained after NK cell or NK cell+mAb treatments to generate the specific natural cytotoxicity or specific ADCC, respectively. All mAbs gave very low levels (<3%) of cytotoxicity in presence of heat inactivated serum media.

**Evaluation of RTX-armed e-NK**

e-NK cells (2x10⁵) were incubated for 1 h with 10 µg/ml RTX, washed and incubated with 1:800 solution of a goat F(ab')2 anti human IgG (H+L) for 30 min at 4°C. After incubation, NK cells were washed with PBS and RTX binding was analyzed by FACs. As a control cells were only stained with the a goat F(ab')2 anti human IgG.

**NK degranulation assay**

Briefly, 50x10³ target cells per well were placed in RPMI, 10% FBS, IL-2 100 U/ml with monensin (BD Biosciencies) in a 96-well V-bottom plate. NK and target cells were incubated overnight at 37°C in 5% CO₂ and living cells were counted using a Muse cytometer (Millipore) with the count and viability kit (Millipore). As a control, NK cells were incubated
without targets. CD107a+ NK cells were analyzed on a Gallios flow cytometer (Beckman Coulter) using 7AAD, CD45RO-FITC, CD19-PE, CD56-PECy7, CD3-APC, CD45RA-APCAlexaFluor750, CD16-KromeOrange and CD107a-HV500 (BD Biosciences). Results were analyzed using Kaluza software.

**In vivo experiments**

*In vivo* experiments were carried out using 6 to 8 weeks/old male NOD scid gamma (NSG) mice. Mice were bred and housed in pathogen-free conditions in the animal facility of the European Institute of Oncology–Italian Foundation for Cancer Research (FIRC), Institute of Molecular Oncology (Milan, Italy). For engraftment of human cells, mice were subcutaneously engrafted with 5x10^6 BCL-P2 or 10x10^6 LNH1 primary tumor cells derived from a BCL (P2) patient or a diffuse large B-cell lymphoma (DLBCL) patient (LNH1). At day 4, we engrafted 15 (BCL-P2) or 10 (LNH1) million e-NK cells and at day 6, mice were treated i.p. with RTX (in saline medium) 3 mg/kg once a week x 3 weeks; or with a combination of both. Tumor growth was monitored at least once a week using a digital calliper, and tumor volume was calculated according to the formula: \( L \times W^2/2 = \text{mm}^3 \), where \( W \) represents the width and \( L \) the length of the tumor mass.
Results

Costimulation with the EBV lymphoblastoid PLH cell line is more efficient to expand UCB NK cells for clinical use than high IL-2 stimulation

Cytokines and encounter with target cells induce dissimilar gene expression on NK cells [23]. We used umbilical cord blood (UCB) cells and compared two NK cell activation/expansion protocols: one using a high dose of the cytokine IL-2 (1000 U/ml) and the other one by cell costimulation. The costimulation protocol was performed with the EBV cell line PLH together with low concentrations of IL-2 (100 U/ml) and IL-15 (5 ng/ml) [29]. NK cell expansions are jeopardized by T cells, therefore we depleted them from UCB before expansion. We observed that NK cell cultures underwent massive cell death at day 12 in IL-2-driven expansion (data not shown). Hence, we compared at day 10 different parameters that reflect NK activity.

Proliferation. Costimulation-drive expansion was largely more efficient (Fig. 1A).

Cytotoxicity. IL-2-drive expansion induced NK cells with superior natural cytotoxicity against K562 (Fig. 1B), Daudi (supplementary. Fig. 1A) and primary CD20⁺ B cell lymphoma cells (BCL P2; Fig. 1C). Moreover, they also showed higher ADCC activity with RTX (Fig. 1C and supplementary Fig. S1A). However, natural cytotoxicity gradually increased with the costimulation protocol when cells were activated for longer periods of time (Fig. 1D). This correlated with a notable large-scale expansion of cells (Median+/−STDV, IL-2 10 d (16.8 +/- 22.2), costimulation 10 d (140.5 +/- 235.8) and costimulation 20 d (260.9 +/- 141.2), n=10).

Activation markers. Both protocols increased the expression of the activation marker CD69 (supplementary Fig. S2B) and decreased of CD45RA expression to originate CD45RA<sub>dim</sub> cells (supplementary. Fig. 2B), which was associated to increase in the activation marker CD45RO as previously published [9]. Costimulation kept higher CD16 expression (supplementary Fig. S2B).

Exhaustion markers. We investigated the expression of two markers that could suppress NK cell-mediated cytotoxicity: TIM-3 [30] and PD-1 [31]. Both protocols did not affected their expression PD-1 expression (supplementary Fig. S2B).

Maturation and homing markers. UCB NK cells showed low percentage of CD62L<sup>+</sup> cells (18.1+/−6.7, average+/−STD, n=3) that increased 10 d after IL-2 treatment (68.2+/−13.5) and costimulation (56.7+/−11.9). However at day 20 post-costimulation, CD62L expression was lost (1.6+/−0.4). IL-2-stimulated cells did not survive so long; hence we could not observe a
possible decrease in CD62L. CD62L is a "homing receptor" facilitating naive lymphocytes to enter secondary lymphoid tissues. Mature NK cells express low CD62L, which favors their peripheral trafficking [32].

In agreement with others [33], few naïve UCB NK cells expressed CD57 (1.2 +/- 1.3, average +/- STD, n=3). IL-2 stimulation barely increased expression (7.3 +/- 3.9) and costimulation did not change it (0.6 +/- 0.6). Longer costimulation, i.e. 20 days, did not alter it (1.6 +/- 0.5). However, the lack of CD57 expression did not impair NK cell cytolytic activity (see below).

In summary, costimulation showed higher number of activated and functional NK cells with higher CD16 expression. This prompted us to exclusively perform costimulation for the next experiments. However IL-2 induced higher and faster cytotoxicity and could be the best option for autologous NK cell grafts.

**Frozen/thawed NK cells keep their cytolytic activity**

For clinical purposes, it will be advantageous to store and have a bank of cryopreserved expanded NK cells that will be “ready to use” [34]. Compared with fresh expanded NK cells, frozen/thawed NK cells roughly lost on average 50% of their cytolytic activity (Fig. 1E). As shown in Fig. 1D, 20 day-activation showed higher cytolytic activity than shorter expansions. For next experiments we used 20-21 days costimulation-induced expansion of UCB-derived NK cells containing more than 90% of NK cells and kept frozen until use. Hereafter, we called them e-NK.

**e-NK cells mediate ADCC against target cells with diverse CD20 levels**

We observed that e-NK performed ADCC similarly on Raji and Daudi cells, which express high CD20 levels (supplementary Fig. S2A), than on primary tumor cells (supplementary Fig. S2B). P2 cells probably express more CD20 than P148, however, they were slightly less sensitive to RTX-mediated ADCC. In fact, e-NK performed ADCC even if their natural cytotoxicity against some patient cells was low or absent (Fig. 2A supplementary Fig. S2B-E). Hence, e-NK show strong ADCC with RTX independently of their natural cytotoxicity and with lower variability (Fig. 2A-B). The glycoengeenering mAb OBZ [4, 35] induced higher ADCC than RTX (Fig.2C and supplementary Fig. S2B-C).

**e-NK cells can be “armed” with mAbs to facilitate treatment**
We next used e-NK cells coated with anti-CD20 mAb (“armed” e-NK cells) as an alternative to antibody-coated target cells. “CD20-armed” e-NK showed similar results than opsonizing tumor cells with anti-CD20 (supplementary Fig. S3A). The presence of RTX after e-NK “arming” was visualized by using a fluorescent anti-IgG (supplementary Fig. S3B). “OBZ-armed” e-NK were also efficient to generate ADCC (supplementary Fig. S4).

Statistical analysis of several e-NK productions on 5 CLL patients did not show any differences between opsonizing targets or “arming” e-NK (Fig. 3A). Moreover, the analysis of these 4 e-NK expansions on the CLL targets showed that all productions could be armed (Fig. 3B). Combining all these results statistically showed the significant ADCC mediated by e-NK (Fig. 3C).

Cytotoxicity requires degranulation and cell interaction by ICAM

Primary human NK cell cytotoxicity is largely independent of degranulation [36] and resides in death receptor binding on tumor cells by death receptors ligands expressed by NK cells. In agreement, e-NK natural cytotoxicity was only partially diminished by the degranulation inhibitor EGTA (Fig. 4A). In contrast, EGTA largely blocked ADCC.

The interaction of NK cell expressed LFA-1 with its target cell ligand ICAM modulates NK cell cytotoxicity [37]. Blocking this interaction with the D1D2 peptide [26] partly reduced natural cytotoxicity and almost completely abolished ADCC (Fig. 4B). Therefore, our e-NK keep similar mechanisms of eliminating target cells than primary human cells.

e-NK mediate ADCC with daratumumab

Next we tested if e-NK produced ADCC with the anti-CD38 daratumumab [38]. We used 3 target cells that expressed CD38 (MM.1S, MV4-11 and BCL-P2) and observed that three different e-NK preparations showed ADCC with daratumumab (supplementary Fig. S5A-C). In contrast daratumumab failed to induce ADCC against U266 that did not express CD38 (Supplementary Fig. 5D). Statistical analysis of several e-NK productions on MM.1S and P2 showed that e-NK efficiently performed ADCC with daratumumab (Fig. 4C).

e-NK mediate ADCC with cetuximab

Next we analyzed if e-NK cells mediated ADCC with other mAbs used to treat solid tumors. We used the cell lines Calu-1 and A549. Calu-1 cells express more epidermal growth factor receptor (EGFR) than A549 (data not shown). Both lines are targets of the anti-EGFR cetuximab, which has been proposed for using with adoptively transferred expanded
allogeneic NK cells in clinical trials for cervical cancer [39]. In fact, *in vitro* and clinical data suggest that cetuximab mediates ADCC through NK cells [22, 34]. We observed a relatively large diversity of the natural cytotoxicity of the different e-NK donors versus these target cells. The decreased on cell viability, measured by MTT formation, was low (supplementary Fig. S6A). However, the increased in cell death, measured by annexin-V/7-ADD staining, was higher. This showed that e-NK had induced on targets the initial steps of apoptosis (annexin-V staining) but longer times were required to evaluate cell viability with MTT. Cetuximab increased early apoptosis and accelerated the process of cell death, decreasing viability. Statistical analysis of several e-NK productions on CALU-1 and A549 showed that e-NK efficiently performed ADCC with cetuximab (Fig. 5A).

EGTA diminished ADCC but not significantly (supplementary Fig. S6B-C). This suggested that the mechanism of action only partly involved degranulation, indicating a possible participation of death ligand-induced apoptosis.

e-NK mediate ADCC with trastuzumab

We next tested the anti-HER2 mAb trastuzumab on SK-BR-3, which express high HER2, and A549, which express low levels, cells. e-NK showed ADCC with trastuzumab in both cell lines by decreasing viability or increasing apoptosis (supplementary Fig. S7A). Natural cytotoxicity as well as ADCC heavily depends on degranulation because EGTA largely decreased both (supplementary Fig. S7A-B). Statistical analysis of several e-NK productions on SK-BR-3 and A549 showed that e-NK efficiently performed ADCC with trastuzumab (Fig. 5B). Finally, we extended this study to the ovarian cell line SK-OV-3 that was mainly resistant to both, natural cytotoxicity and ADCC, during short treatment (data not shown). However, 6-day treatment revealed that NK cells, mainly with trastuzumab, efficiently killed these cells (Fig. 5C).

e-NK mediated ADCC in vivo

We next evaluated e-NK activity *in vivo* by engrafting primary tumor cells from a BCL patient (P2) or from a DLBCL patient (LNH1) into NSG mice. Four days later mouse were engrafted with e-NK and 2 days later treated with RTX. e-NK, and mainly RTX, decreased tumor growth (Fig. 6A). But co-treatment was more effective, totally protecting mice from BCL cells and 4 out of 5 mice from DLBCL cells.

e-NK cells showed ADCC against chemoresistant cells
EBV-activated NK cells overcome anti-apoptotic mechanisms of chemoresistant cells [24, 25]. Overexpression of BCL-X<sub>L</sub> and MCL1 are common features of several hematological cancers [40]. Jurkat cells over-expressing BCL-X<sub>L</sub> are resistant to doxorubicin and to soluble TRAIL [41]. MCL1 over-expression protects from ibrutinib cytotoxicity (O. Gonzalo and I. Marzo, Doctoral Thesis, University of Zaragoza, 2017). e-NK killed the CD20<sup>+</sup> MEC-1 B-CLL cell line overexpressing BCL-X<sub>L</sub> or MCL1, which gave them low protection against ADCC compared to the wild type cell line (supplementary Fig. S8). Statistical analysis of several e-NK productions showed that they kill chemoresistant cell lines (Fig. 6B). However, BCL-X<sub>L</sub> overexpression significantly decreases ADCC at high E:T ratios. This suggests that chemoresistance could partially protect tumor cells from e-NK-mediated ADCC, but not from e-NK natural cytotoxicity.
Discussion

High numbers of tumor-infiltrating NK correlates with a better prognosis in some tumors [15]. NK are the first lymphocytes to recover after HSCT including after UCBT. The speed of recovery correlates with the prognostic [17]. Despite these findings, NK cell adoptive immunotherapy has given low clinical benefit. Perhaps current expansion protocols fail to produce enough NK cells to support clinical success or generate cells with impaired activity [15]. We focused on developing a protocol to obtain NK cells in sufficient number and with high ADCC activity that could be mediated by different therapeutic mAbs. From UCB-derived NK we produce e-NK that only partially decrease ADCC function after cryopreservation (Fig. 1), preserving ADCC in vivo (Fig. 6). Interestingly, we have recently described that expression of the CD45 isoform CD45RO identifies anti-tumor NK cells in human patients [10]. e-NK decrease CD45RA and increase CD45RO expression suggesting that they would show elevated antitumor natural cytotoxicity as we have confirmed here. e-NK do not require relatively high Ag levels to perform ADCC because they were effective with cells expressing variables Ag levels in different cell lines.

Coupling mAbs with e-NK should improve the use of both treatments in several clinical contexts. First, most cancer patients possess impaired NK cells [3, 15, 17]. This should diminish the clinical activity of mAb. e-NKs should overpass this hitch by increasing mAb-induced ADCC in patients with immune defects. Second, the clinical activity of NK cells in solid cancers is uncertain [15, 17]. Probably these effectors scarcely recognize these targets in vivo and/or fail to infiltrate these tumors e.g., in some tumors NK have been detected in the tumor stroma but not within the tumor lesion [42-44]. Moreover, the adoptive transfer of autologous NK cells in patients as single therapy maintains high levels of circulating NK cells but did not mediate tumor regression [3, 17, 45]. mAbs should bring e-NK to the selected targets and can also facilitate target elimination by favoring the recognition of opsonized cells. In fact, haploidentical NK cells plus anti-GD2 mAb therapy has promising antitumor activity in pediatric recurrent/refractory neuroblastoma [46, 47]. Third, e-NKs overcome the anti-apoptotic mechanism of leukemic cells (Figure 7 and [24]) allowing elimination of tumor cells from poor prognosis patients [25].

An inconvenient of allogeneic expanded NK cell engraftment is that they survive few days in vivo [48]. The persistence of ex vivo haploidentical IL-2-activated and -expanded NK-DLIs reaches a maximum of 7 days in lymphoma patients [49]. This leaves grafted NK cells short time to eliminate their targets. The advantage is that NK will be less prompted to generate clinical problems as those found with CAR-T cells, which produce some chronic effects.
related to their long-term persistence ([50]; http://www.medscape.com/viewarticle/876591). One of the main concerns for the use of allogeneic immune cells is the incidence of GVHD. Allogeneic NK cells infusion is well tolerated in cancer patients [3, 17] and the severity of aGVHD correlates with impaired reconstitution of the NK cell compartment [51]. To our knowledge, engraftment of NK cells have been linked to GVHD only when combined to HLA-matched, T-cell–depleted nonmyeloablative peripheral blood stem cell transplantation [52]. NK cells likely contributed to GVHD in this setting by augmenting underlying T-cell alloreactivity [52].

An interesting alternative to allogeneic NK are KIR/KIRL blocking antibodies that activate endogenous NK cells [53]. This approach has the inconvenient that cancer patient NK cells are hyporeactive [3, 15, 17], suggesting that they could be inefficient to totally eliminate tumors. Moreover, recent clinical data suggest that such antibodies modify the endogenous NK repertoire making KIR-expressing NK cells, which are those with higher cytolytic activity, hyporeactive [54]. This raises concerns to the clinical use of these antibodies. There are other ways to activate endogenous NK cells such as the use of lenalidomide (LEN) [7, 8]. Preliminary results from the Phase Ib/II clinical trial GALEN suggest that LEN could facilitate OBZ-mediated NK cell activation [8] as was observed with RTX [55].

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Figure legends.

**Figure 1. Optimization of NK cell expansion protocol.** A) Comparison of costimulation (PLH accessory cells + IL-2 100 U/ml + IL-15 5 ng/ml) and IL-2 (1000 U/ml) expansion protocols using three UCB donors (2903, 3464, 2928). B) K562 cells were incubated overnight with costimulation- (circles) or IL-2-activated (triangles) NK cells from two different donors at different effector:target (E:T) ratios. Cell death was analyzed by 7-AAD staining. C) BCL Patient 2 cells were incubated overnight with costimulation- or IL-2-activated NK cells from three different donors, in presence (black symbols) or absence (white symbols) of RTX (10 µg/ml). D) NK cells were expanded by costimulation for different days from 2 donors. At these days NK cells were frozen. They were thawed at the same time and tested for cytotoxicity against the cell lines. E) NK cells from 2 donors were expanded for 20 days and frozen or kept fresh before testing their cytotoxicity.

**Figure 2. e-NK mediate ADCC with anti-CD20 mAbs.** A) PBMCs from CLL patient 148 were incubated for 1h with 10 µg/ml of RTX and overnight with e-NK cells obtained from nine different donors at 1:1 E:T ratio. Cell death was analyzed by 7AAD staining. The right panel shows the statistical analysis. B) 7 e-NKs were tested against 13 CD20+ target samples at different E:T ratios as described in (A). C) 5 e-NKs were tested against 6 CD20+ target samples at different 1:1 E:T ratio as described in (A).

**Figure 3. Anti-CD20-armed e-NK show ADCC activity.** PBMCs from CLL samples were incubated for 1h with 10 µg/ml of RTX and overnight with donor e-NK cells at 3:1 E:T ratio (antibody-coated target cells condition). Alternatively, e-NK cells were incubated for 1 h with 10 µg/ml of RTX before incubating them overnight with target cells (antibody-armed NK cell condition). B-CLL cell death was analyzed by 7-AAD.

**Figure 4. ADCC requires degranulation and LFA-1/ICAM interaction.** A-B) Daudi cells were incubated overnight with e-NK cells from two different donors and/or RTX (10 µg/ml) as described in Fig. 2. Cytotoxic assays were performed also in the presence of 1 mM EGTA (A) or 15 µg/ml D1D2 protein (B). Cell death was analyzed by 7-AAD staining. C) e-NK produce ADCC with daratumumab. Three different e-NK cell productions were tested against the CD38+ cell line MM.1S and BCL-P2 cells that expressed CD38. Target cells were pre-
incubated for 1 h with 5 µg/ml daratumumab before overnight incubation at different E:T ratios with e-NK. Cell death was analyzed by 7-AAD staining.

**Figure 5.** e-NK perform ADCC with cetuximab and trastuzumab against EGFR and HER2 positive cell lines, respectively. A) Tumor cells were incubated with 5 µg/ml cetuximab for 1 h and overnight with 4 e-NK preparations at 3:1 E:T ratio. Subsequently, we measured cell viability (MTT) and cell death (annexin-V/PI). B) Tumor cells were incubated with 5 µg/ml trastuzumab for 1 h and overnight with e-NK cells from 4 different donors at 3:1 E:T ratio. Subsequently, we measured cell viability (MTT). C) 1x10^4 SK-OV-3 cells were plated and 24h later treated with trastuzumab and cultured with 5x10^5 NK cells. After 6 days, medium was removed and cells were fixed and stained. The right graph shows the statistical analysis comparing cells incubated with e-NK alone or with trastuzumab. Two areas of 2 different experiments were counted and the average of cells/area depicted in the graphic.

**Figure 6.** e-NK show ADCC in vivo and overcome mechanisms of drug resistance. A) 5 NSG mice/group were subcutaneously engrafted with BCLP2 (left) or 10x10^6 (right) cells and treated with e-NK and/or RTX. B) e-NK cell-induced ADCC overcome anti-apoptotic mechanisms of drug resistance. CD20^+ MEC-1 cells overexpressing BCL-X_L and MCL1 were incubated with of RTX (10 µg/ml). After 1 hour, e-NK cells from 3 different donors were added overnight. Cell death was analyzed by 7AAD/annexin-V labeling.
Figure 2

A

B

C

---

* * *

RTX

control + RTX

* * *

RTX OBZ

% Cell death

% Cell death

% Cell death
Figure 3

A

B

C

- Antibody-coated target cells
- Antibody-armed NK cells

NK cells

CLL P421  CLL P520  CLL P424  CLL P570  CLL P576

NK 3168  NK 3533  NK 2903  NK 3464

% Cell death
Figure 5

A

- **CALU-1**
- **A549**

B

- **SKBR3**
- **A549**

C

- Control
- Trastuzumab

NK cells

**n° cell**
Figure 6

A

5x10^6 BCL P2

Days from tumor injection

Tumor volume (mm³)

B

MEC-1

MEC-1 BCL-XL

MEC-1 MCL-1

% Cell death

E:T

+RTX

CONTROL

**

*

**
Graphical Abstract

Antigen

Target cell

mAb

NK cell

CD16

NK Expansion

Antibody-coated target cells

Antibody-armed NK

Degranulation

Tumor cell death