Research Article

Impact of the NK Cell Receptor LIR-1 (ILT-2/CD85j/LILRB1) on Cytotoxicity against Multiple Myeloma

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The role of different receptors in natural-killer- (NK-) cell-mediated cytotoxicity against multiple myeloma (MM) cells is unknown. We investigated if an enhancement of NK-cell-mediated cytotoxicity against MM could be reached by blocking of the inhibitory leukocyte immunoglobulin-like receptor 1 (LIR-1). Our investigations revealed high levels of LIR-1 expression not only on the NK cell line NK-92, but also on myeloma cells (MOLP-8, RPMI8226) as well as on a lymphoblastoid cell line (LBCL; IM-9). Subsequent cytotoxicity assays were designed to show the isolated effects of LIR-1 blocking on either the effector or the tumor side to rule out receptor-receptor interactions. Although NK-92 was shown to be capable of myeloma cell lysis, inhibition of LIR-1 on NK-92 did not enhance cytotoxicity. Targeting the receptor on MM and LBCL did not also alter NK-92-mediated lysis. We come to the conclusion that LIR-1 alone does not directly influence NK-cell-mediated cytotoxicity against myeloma. To our knowledge, this work provides the first investigation of the inhibitory capability of LIR-1 in NK-92-mediated cytotoxicity against MM and the first functional evaluation of LIR-1 on MM and LBCL.

1. Introduction

Understanding of NK cell function has undergone a long process since their identification in 1975 [1]. NK cells have initially been regarded as part of the innate immune system, not allowing any modulation of action with respect to their changing microenvironment. Their pattern of inhibitory and activating receptors was considered to be sufficient to adequately detect tumor cells by the lack of human leukocyte antigen (HLA) class I molecules. Those tumor cells were killed instantly and without any obvious need of coactivation by other cells of the immune system [2]. This unique feature among lymphocytes has now been understood to be only the basic function of response, which is completed by diverse interactions with especially dendritic cells (DC) and T cells [3]. NK cells do extensively communicate with their surroundings, and their still-not-fully-deciphered set of receptors detects changes in the normal surface pattern on all types of tissues.

NK cell receptors are functionally divided into activating and inhibitory receptors. Their main ligands are major histocompatibility complex I (MHC-I) molecules, while some of the receptors can directly recognize specific antigens on bacteria or damaged cells. Mainly three different subclasses of NK-cell receptors (NKRs) can be distinguished.

LIR and killer immunoglobulin-like receptors (KIRs) are type I transmembrane proteins of the immunoglobulin-like receptor superfamily (IgSF). Both recognize classical HLA class I molecules, while LIR can also interact with nonclassical HLA class I and bacteria with low binding affinities [2, 4–6]. The second group of natural cytotoxicity receptors (NCRs) also belongs to type I transmembrane proteins but has poorly defined ligands. Type II transmembrane proteins of the C-lectin type superfamily include natural killer cell
Lectin-like receptor group 2 (NKG2) receptors that form heterodimers with CD94 [2].

LIRs are expressed on subsets of NK cells and T cells, as well as on monocytes, B cells, and DC, with the widest distribution for LIR-1 [7–10].

LIR-1 is an inhibitory receptor also known as immunoglobulin-like transcript 2 (ILT-2)/CD85j or leucocyte immunoglobulin-like receptor, subfamily B member 1 (LILRB1) [7]. It has first been detected in searching for the counterpart of UL18, a cytomegalovirus encoded HLA class I homolog that is expressed on infected cells [8, 11, 12].

MM is an incurable disease that is characterized by the clonal proliferation of terminally differentiated plasma cells [13, 14]. Stem cell transplantation (SCT) is so far the only option to achieve long time remission of the disease [15]. To improve the outcome of MM patients, approaches like immunomodulation and cellular therapy are under investigation. NK cells are an attractive candidate for immune therapy. They kill tumor cells without antigen-specific priming [2] and are the predominant lymphocyte subset within the first 90 days after transplantation [16–19]. LIR-1 is one of the main inhibitory NK cell receptors in this early phase after SCT [10, 16, 20].

We therefore investigated the influence of LIR-1 on MM and on other tumor entities, taking into account its broad distribution among tissues.

2. Material and Methods

2.1. Cells. Unless otherwise stated, all media and supplements were obtained from Life Technologies. Natural killer cell line NK-92 was cultured in alpha-MEM supplemented with Earl’s Salts and L-Glutamine, 12.5% equine serum, 0.2 mM inositol (Sigma-Aldrich), 0.15 M sodium chloride, 0.1% Tween 20) containing 3% nonfat dry milk. Incubation with the anti-LIR-1 (VMP55, Santa Cruz) was done overnight at 4°C under gentle agitation. After extensive washing with PBS-T, secondary one-step horseradish-peroxidase-conjugated goat anti-mouse IgG (R&D Systems) was added for 1 h. Membranes were stained with enhanced chemiluminescence agent (GE Healthcare) and exposed to X-ray film (GE Healthcare). To confirm equal loading of all gel chambers, membranes were stripped from the specific antibody using Re-Blot solution (Millipore), followed by additional staining of β-Actin (ACTB, C4, Santa Cruz).

2.2. Flow Cytometry. Monoclonal antibodies (mAb) were phycoerythrin- (PE-) conjugated CD2 (RPA-2.10, BDP harmingen), CD159a (Z199, Beckman Coulter), CD85j (HP-F1, Beckman Coulter); Pacific Blue-conjugated CD16 (MOPC-21, BD Pharmingen); fluorescein isothiocyanate- (FITC-) conjugated CD25 (B1.49.9, Beckman Coulter) and anti-IgG (goat polyclonal anti-mouse IgG, Abcam); allophycocyanin (APC)-stained CD56 (B159, BD Pharmingen) as well as appropriate isotype controls. Unconjugated anti-HLA-1 (HP-1F7) was obtained from Santa Cruz, anti-HLA-G (MEM-G/09) and -E (MEM-E/08) were obtained from Abcam. 7-Amino-Actinomycin D (7AAD, BD Pharmingen) was used to analyze dead cells. 20 μL Anti-A, B reagent was used in each sample to block unspecific bindings (OrthoClinical Diagnostics). Cells were incubated with the antibody or isotype control for 30 minutes at 4°C, washed with PBS and, if appropriate, stained with a secondary antibody followed by an additional washing step. All samples were additionally stained with 7AAD. Fluorescence was measured on a BD FACSCanto II flow cytometer and BD FACSDIVA Software v.6.1.3 (Becton Dickinson) was used for data analysis.

2.3. Transfection of COS-7 Cells with LIR-1. As a positive control for western blot experiments, COS-7 cells were transfected with pCMV6-AC vector encoding for LIR-1 (OriGene) or pCMV6-XL5 as a mock control (OriGene), using FuGene HD Transfection Reagent (Promega) [21, 22]. All steps were performed according to the manufacturer’s instructions. The vector was multiplied by transformation of E.coli cells (One Shot TOP10/P3 competent cells) with subsequent purification of the plasmids (Qiagen EndoFree Plasmid Maxi Kit).

2.4. Western Blot Analysis. Western blot was used to analyze expressed surface molecules of effector and target cells [23, 24]. Proteins were separated by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) under reducing conditions. Gels (Life Technologies) were blotted onto nitrocellulose membranes (Whatman). Membranes were blocked for one hour with TBS-T buffer (0.05 M Tris-HCL, 0.15 M sodium chloride, 0.1% Tween 20) containing 3% nonfat dry milk. Incubation with the anti-LIR-1 (VMP55, Santa Cruz) was done overnight at 4°C under gentle agitation. After extensive washing with TBS-T, secondary one-hour incubation with horseradish-peroxidase-conjugated goat anti-mouse IgG (R&D Systems) was completed with additional washing. Membranes were stained with enhanced chemiluminescence agent (GE Healthcare) and exposed to X-ray film (GE Healthcare). To confirm equal loading of all gel chambers, membranes were stripped from the specific antibody using Re-Blot solution (Millipore), followed by additional staining of β-Actin (ACTB, C4, Santa Cruz).

2.5. Cytotoxicity Assays. Cytolysis was determined in 4-hour chromium-release assays (CRAs) according to standard protocols [25–27]. Briefly, NK-92 and target cells were seeded out in fresh medium one day before functional assays. The next day target cells were labeled with 100 μCi sodium-51-chromate (51Cr) for 1.5 hours at 37°C in a humidified incubator with 5% CO₂. Two washing steps were performed with PBS (Life Technologies) and assay medium (RPMI1640, 10% FCS, 1% PenStrep; Life Technologies), respectively. Cells were resuspended to a dilution of 5 × 10⁴ cells/100 μL. NK-92 and tumor cells were coincubated at various effector: target (E:T) ratios in U-bottom microtiter
Figure 1: Expression of surface antigens. (a). Flow cytometric phenotyping of all cell lines was done with mAbs against LIR-1, CD56, CD16, CNKG2A, CD2, and CD25 as shown for NK-92. (b). Surface antigen expression on NK-92 and tumor cell lines. Values given as % positive staining after subtraction of isotype control.
plates at a total of 200 µL assay medium. Maximum lysis or spontaneous release (SR) of 51Cr were induced by adding 5% Triton-X or assay medium to 100 µL target cells, respectively. All samples were plated out in triplicates. Prior to the 4-hour incubation period, plates were carefully centrifuged to facilitate E:T contact. After incubation and additional centrifugation, 25 µL supernatant were transferred to a 96-well plate (Isoplate 96, PerkinElmer). To each well, 150 µL scintillation liquid were added (Rotiszint eco plus, Carl Roth). Plates were closed with Viewseal foils (Greiner Bio-one). Suspension was mixed thoroughly for 15 minutes at 19°C on an Eppendorf thermomixer and then measured (19, Release 19.0.0).

Results are shown as the mean of at least three independent experiments. In all experiments, SR was <20% [28].

2.8. Blocking of Tumor Cells. Target cells were incubated with 1 µg/mL of the respective mAb at a cell density of 5 × 10^4 cells/mL. Procedure and incubation times were the same as described for NK cells. F(ab')2 was used at a concentration of 1.7 µg/mL for all samples, inclusively the “untreated” control. Tumor cell incubation with mAb led to SR of <25% for MOLP-8 and RPMI8226 and up to 11% for all other cells.

2.9. Statistics. To control for indirect effects, statistical interpretation was done by multivariate Analysis of Variance (ANOVA). In all calculations, specific lysis was defined as the dependent variable. Antibodies, concentrations of the mAb, the targets and the use or no use of F(ab')2 were defined as independent variables. Wherever appropriate, interdependencies between the variables were taken into account. All calculations were done by SPSS (IBM SPSS Statistics Version 19, Release 19.0.0).

3. Results

3.1. Characterization of NK-92 and Tumor Cell Lines. NK-92 was found to express high levels of LIR-1 based on flow cytometric analysis (Figure 1). We confirmed a high expression of NKG2A and CD25, as well as small amounts of FcyRIII (CD16) [10]. Hereby, but not regarding LIR-1 expression, NK-92 cells share important similarities with the CD56^bright subset of NK cells [32].

LIR-1 was present on myeloma cells as well as on IM-9, but no target cell line expressed NKG2A. Western blot analysis confirmed the pattern of LIR-1 expression, and the strength of band representation reflected the staining intensity detected by flow cytometry (Figure 2). LIR-1 or mock-transfected as well as naïve COS-7 served as positive and negative controls, respectively. For evaluation of HLA class I expression, NK-92 cells were shared with mAb concentrations of 0.1, 1 and 10 µg/mL, respectively. Target cells were prepared as described above. NK-92 cells were washed twice to avoid interactions of the mAb with the later on coincubated target cell line. Cells were adjusted for a fix E:T ratio of 1.25:1, using 5 × 10^3 target cells/100 µL as before. In some of the experiments, this step was followed by additional preincubation with 11 µg/mL F(ab')2 15 minutes prior to coincubation of NK and tumor cells. In samples classified as “untreated,” no F(ab')2 was used [30, 31]. SR was always <20% for all cell lines except from MOLP-8, which showed a constantly high SR up to 36% [28]. To rule out toxicity of the mAb, in four experiments NK-92 were radioactively labeled (51CrNK) and treated as the unlabeled cells in a parallel series to the blocking assays. 51CrNK showed SR < 6% for all conditions with a standard deviation (SD) < 3%. No differences related to parameters F(ab')2, antibody or concentration could be detected within statistical analysis [30].

2.6. Blocking Experiments. Blocking antibodies were anti-NKG2A (CD159a, IgG2b, Z199 BeckmanCoulter) anti-LIR-1 (CD85j, IgG1, Santa Cruz), which blocks HLA-A, -B, -C, -E, and -G engagement [10, 29, 30]. IgG1 (CD85j, 292319, IgG2b, R&D Systems) and anti-HLA-I NKG2A (CD159a, IgG2b, Z199 BeckmanCoulter) anti-LIR-1 were used as isotype controls. As also shown by flow cytometric analysis, NK-92 as well as IM-9 and MOLP-8 express high levels of LIR-1, whereas no detection of LIR-1 was possible on HL60 and K562 cells.

2.7. Blocking of Effector Cells. CRA were performed as described above. Heat inactivated human serum (HS) was obtained after informed consent from healthy volunteers. NK-92 cells were preincubated in RPMI1640 with 1% PenStrep and 10% HS for 30 minutes and kept within the same medium during additional 30 minutes of incubation with mAb concentrations of 0.1, 1 and 10 µg/mL, respectively. Target cells were prepared as described above. NK-92 cells were washed twice to avoid interactions of the mAb with the later on coincubated target cell line. Cells were adjusted for a fix E:T ratio of 1.25:1, using 5 × 10^3 target cells/100 µL as before. In some of the experiments, this step was followed by additional preincubation with 11 µg/mL F(ab')2 15 minutes prior to coincubation of NK and tumor cells. In samples classified as “untreated,” no F(ab')2 was used [30, 31]. SR was always <20% for all cell lines except from MOLP-8, which showed a constantly high SR up to 36% [28]. To rule out toxicity of the mAb, in four experiments NK-92 were radioactively labeled (51CrNK) and treated as the unlabeled cells in a parallel series to the blocking assays. 51CrNK showed SR < 6% for all conditions with a standard deviation (SD) < 3%. No differences related to parameters F(ab')2, antibody or concentration could be detected within statistical analysis [30].
Figure 3: Continued.
Figure 3: HLA expression. (a). Representative flow cytometric gating strategy for analysis of HLA expression on all used cell lines (shown for RPMI8226). All cells were stained with anti-HLA-I/-E/-G IgG1, followed by secondary goat anti-mouse IgG1 FITC. (b). HLA class I molecules were detectable on all cell lines except for negative control K562. JEG-3 were strongly positive for HLA-6, while detection on NK92 and HL60 was marginal. HLA-E as the ligand for NK62A lacked on all cell lines except on minor staining on IM-9.
stimulated with 200 U/ml IL-2 the day before the assay. NK-92 cells were almost resistant to NK-92 mediated killing. NK-92 cells were the table below are given as % specific lysis. AML cell line HL60 has been described before to lack inhibitory KIR molecules [37, 38]. Furthermore, NK-92 blocking assays, for no target cell line expressed the only inhibitory capacities of LIR-1. No increase of cytotoxicity instead of primary cells allowed an isolated view on the binding partners.

The choice of NK-92 and those distinct tumor cell lines instead of primary cells allowed an isolated view on the inhibitory capacities of LIR-1. No increase of cytotoxicity due to blockade of NKG2A could be expected in subsequent blocking assays, for no target cell line expressed the only known NKG2A ligand HLA-E [36]. Furthermore, NK-92 has been described before to lack inhibitory KIR molecules [37, 38].

Other inhibitory LIRs that can be found on NK cells are only LIR-3 (ILT5) and LIR-8 as well as soluble LIR-4 for which the ligands are not yet detected [6].

LIR-1 could therefore be considered to be the only known major inhibitory receptor in this context and was expressed at high levels (Figures 1 and 2). Influence of so far unknown inhibitory receptors was ruled out by selective blockade of LIR-1.

Due to these findings, we considered the use of NK-92 and the chosen tumor cell lines as an ideal system to study the discrete influence of LIR-1 on modulation of NK-cell cytotoxicity.

3.2. Myeloma Cells Are Highly Susceptible to NK-92 Mediated Killing. Cytotoxicity of NK-92 against a panel of tumor cell lines was investigated in CRA at different E:T ratios (Figure 4) [28]. MM cell lines and IM-9 were efficiently lysed by NK-92, with highest results for IM-9 (E:T 10:1; specific lysis 69.9 ± 6.3%) followed by MOLP-8 (29.8 ± 3.9%), K652 (22.6 ± 7.6%), and RPMI8226 (21.3 ± 3.6%). HL60 was almost resistant to lysis (4.5 ± 2.5%).

Figure 4: NK-92 mediated killing of target cell lines is dependent on applied E:T ratio. CRA revealed a clearly E:T-dependent level of NK-92 mediated cytotoxicity. Results of CRA are represented as means of at least three independent experiments ± SD. Values in the table below are given as % specific lysis. AML cell line HL60 was almost resistant to NK-92 mediated killing. NK-92 cells were stimulated with 200 U/ml IL-2 the day before the assay.

3.3. Blocking of LIR-1 on NK-92 Does Not Increase Target Cell Lysis. To evaluate the influence of LIR-1 in myeloma cell lines, mAbs were used to block LIR-1 receptor-ligand interactions. As target cells lacked the expression of the HLA-E molecule, blocking of NKG2A was not expected to alter the results but was conducted as a negative control. No significant increase of tumor cell lysis could be achieved by any of the mAbs despite high concentrations (Figure 5).

As specific lysis of target cells by NK-92 was found to be independent from mAb concentration and type, results are presented as means of the used concentrations (0.1/1/10 μg/mL) or in a separate bar as means of concentration and mAbs (CD85j, CD159a, and IgG2b) (Figure 6). For MOLP-8, RPMI8226 and HL60, a significant influence of F(ab\(^{-}\))\(_2\) towards a decreased lysis seemed to be relevant, but it could not be taken into account.

What first might appear as a protective effect of the applied F(ab\(^{-}\))\(_2\) towards a reduced lysis of target cells could also be observed in the untreated sample and must therefore be considered to be a side effect caused by cell culture procedure. Only experiments with K562 were performed at the same day with and without F(ab\(^{-}\))\(_2\) and the observed specific lysis of K562 was the same for both experimental rows. This confirms the thesis of culture side effect to be responsible for significant changes in experimental results.

It is possible that LIR-1 influence could not be measured, if a maximum level of NK-activation had already been achieved before blocking of the inhibitory receptor [38, 39]. Interleukin (IL)-2 requirement during cell culture, induction of the high potential activating receptor Nkp44 by IL-2 [40], and origin of NK-92 from rapidly progressive NK cell lymphoma [38, 41] favor a preactivated condition.

3.4. Blocking of Neither LIR-1 Nor HLA Class I on Target Cells Increases Target Cell Lysis. As expected, HLA-A, -B, -C, -E, and -G blockade on tumor cells did not show any influence on lysis (Figure 7) since blocking of LIR-1 as the only relevant inhibitory NK cell receptor on NK-92 had already not modulated cytotoxicity. We also decided to selectively block LIR-1 receptors on the tumor cells before coincubation with NK-92. Though not being likely to directly change NK-cell properties, LIR-1 expression might contribute to MM resistance in so far unknown ways as its role in immune regulations is still fairly unknown (discussed below). As LIR-1 surface expression increases during B-cell and DC maturation [42], it might be directly involved in cell-cell interactions that promote survival and growth. In our experiments, LIR-1 expression on target cells seemed not responsible for a resistance to lysis (Figure 7).

3.5. mAb or F(ab\(^{-}\))\(_2\) Have No Toxic Effects on NK-92 under Experimental Conditions. To rule out toxic effects of mAb, NK-92 were labeled with 51Cr and incubated with the respective mAb in parallel to the conducted experiments.
In four independent experiments, no harmful effect of anti-LIR-1, anti-NKG2A or F(ab′)2 could be observed (Figure 8).

3.6. F(ab′)2 Stabilize Pattern of Tumor Lysis. Although not significantly affecting tumor lysis, there seemed to be an important influence of the F(ab′)2 fragments (Figure 9). They stabilized the results even though prior evaluation of surface molecules did only show very low amounts of CD16 (see above). Apart from outliers, use of F(ab′)2 seemed to even out mAb effects in the blocking experiments, leading to values that oscillate close to the origin in both directions (a). Sparing those fragments decreased relative lysis, predominantly relevant for MOLP-8 and RPMI8226 cells. Only K562 cells rendered more susceptible to NK mediated lysis with a lack of F(ab′)2 though not at a significant level (b).

4. Discussion

The aim of the present investigations was to evaluate if LIR-1 on NK cells inhibits NK-92 mediated cytotoxicity
against different tumor cell lines. Secondly, presence of LIR-1 on the target cells was validated concerning its influence on NK cell mediated lysis. LIR-1 was assumed to be the only receptor for bacterial detection [5, 8]. Data about the specific role of this receptor in “normal” action of lymphocytes, especially NK cells, are more conflicting and rare. While LIR-1 inhibition of T cell and monocyte activation has less refer to experimental settings in which HLA-G was not present on the target cell [46–48]. Available publications emphasize data that confirm an important inhibitory effect of LIR-1, but a more questioning view might refer to the number of experiments mentioned that did not deliver the pronounced outcome. We will give a short overview of available functional studies of LIR-1 on donor-derived NK cells and NK cell lines in order to demonstrate the conflicting knowledge that is available.

Godal et al. made an attempt in revealing the influence of LIR-1 on dNK in cytotoxicity against HLA-G negative AML and ALL blasts. Their results indicate that LIR-1 does only serve as a weak inhibitory NK cell receptor in the absence of HLA-G on tumor cells, but might be relevant in situations with low KIR expression as seen within the first months after stem cell transplantation (SCT) [10].

Other authors used merely HLA-transfected 721.221, murine cells (P815) or immature dendritic cells (iDC) as target cells, but often had the benefit of comparing results from NK cell lines to the performance of NK cells derived from healthy donors. In 2008, Yawata et al. conducted a series of degranulation assays of dNK against HLA class I deficient 721.221 in order to investigate the involvement of distinct receptors in “missing-self” recognition and were not able to identify any involvement of LIR-1 [49]. More successful LIR-1 mediated inhibition was achieved by Morel and Bellon who

**Figure 6:** Use of F(ab')2 does not significantly influence cytotoxicity of NK-92 cells. Statistical analysis (ANOVA) reveals a seemingly protective effect by F(ab')2 fragments against NK-92 mediated lysis. As influence of the type of antibody and the used concentration were excluded, experimental data were merged. Results were highly significant (untreated ± F(ab')2/Ab use ± F(ab')2): IM-9 P = 0.317/P = 0.211; MOLP-8 P = 0.017/P = 0.002; RPMI8226 P = 0.052/P < 0.001; HL60 P = 0.009/P < 0.001; K562 P = 0.692/P = 0.771). The same settings were used throughout all experiments. In the experimental row, K652 tests were performed at the same day and within the same panel. For all other target cells, tests including F(ab')2 were performed first and were followed by those without additional treatment 1 week later. Though IM-9 renders more susceptible to lysis to a nonsignificant degree, time is the factor most likely to be taken into account, for targets incubated with “untreated” NK cells showed the same changes in sensitivity towards lysis.

**Figure 7:** Blocking of LIR-1 and HLA on tumor cells does not lead to increased lysis by NK-92 cells. LIR-1 (CD85j) and HLA class I molecules of the tumor cell lines were blocked as before on the NK-92 cells. After incubation and extended washing, standard CRA were performed. Neither LIR-1 nor HLA blocking showed significant increase of tumor cell lysis. All experiments were performed with the use of F(ab')2 to prevent ADCC. Statistical analysis performed by ANOVA excluded any significant effect of either isotype or Ab on target cell lysis, compared to untreated samples. Discrete investigation of K562 treatment, performed as a comparison of cumulated values for all Ab-incubated samples with or without F(ab')2, excluded any influence of additional F(ab')2 on the results (P = 0.218).
tested the cytotoxicity of dNK and NK-92 against HLA-G and -B transfected 721.221 in CRA with subsequent use of blocking antibodies [40]. Their pre-studies found only 15% of LIR-1 positive dNK to be stimulated by target contact and these responsive dNK were chosen for the subsequent studies. Comparative analysis of NK-92 and dNK in different types of assays were conflicting.

No such conflict was observed by Favier et al., who incubated dNK and the NK cell line NKL with 721.221-G1 and used blocking Abs against LIR-1, NKG2A, or IgG2b. Isotype control with (+) or without (−) F(ab')2 is under the same conditions as the unlabeled effector cells in the blocking experiments. SR is given as mean of four independent experiments for untreated NK cells and as mean of an experimental series with concentration of 10, 1 and 0.1 μg/mL. "All Ab mean" integrates those results. F(ab')2 concentration was 11 μg/mL. Spontaneous release was <6%. As this control panel was performed in parallel to regular NK-92 blocking experiments, interferences by reagents, temperature, incubation period, cell viability, or technical equipment can be excluded.

Moreover, in reverse ADCC (rADCC) assays with dNK against P815, only few clones were inhibited by LIR-1. Here again, NK cell clones have shown to exhibit less predictable outcomes than cell lines [21].

Summarizing the available data, NK cell lines seem to be more reliable than dNK concerning LIR-1 mediated downregulation of cytotoxicity. The results for polyclonal dNK show high variances between the different clones that are mostly not characterized in detail. Successful LIR-1 mediated inhibition by HLA-transfected 721.221 has abundantly been shown. Involvement of additional undetected receptors could not always be excluded, and different types of assays performed with the same effector and target cells could lead to highly diverging results. Investigations of cytotoxicity of dNK and donor-derived tumor cells are rare and no sufficient information is available about the role of LIR-1 in the absence of HLA-G. Current opinion about the inhibitory influence of LIR-1 is predominantly based on investigations at the fetomaternal interface or has been gained from settings in which only a single HLA molecule was present on the target cells—mostly HLA-G or -B on 721.221. The performed experiments do not sufficiently cover the extensive binding capacities of LIR-1 to HLA class I.

Although we are aware of problems concerning the comparability of NK cell lines to donor-derived NK cells (dNK), we have chosen an experimental setting that allows studying isolated influence of LIR-1 on cytotoxicity. Our major goal was to provide a model system to overcome the common practice of using transfected target cell lines in cytotoxicity assays. Being aware of the necessity of future efforts to confirm the present findings in a brighter panel of cells and cell lines, these results might provide a first step towards a new understanding of LIR-1.

Available data do not sufficiently support the direct implication of LIR-1 in NK cell inhibition. Uptregulation of the receptor does not necessarily favor immunosuppression but might correlate with the acquisition of memory. LIR-1 surface expression increases during B cell and DC maturation [42] as well as during cytomegalovirus infection [52] and acquisition of T-cell memory [53, 54]. We suppose that an increasing LIR-1 expression also correlates with the acquisition of NK cell memory [55], supported by the surveillance that a high LIR-1 level on NK cells leads to an effective lysis of HIV-infected DC [56].

LIR-1 seems to have a high impact on regulating the balance between activation and inhibition during immune responses [42]. It might do so by cooperation with other receptors like KIRs, which favor a clustering of MHC-I [57]. As HLA-G is the only LIR-1 ligand that generates covalent dimers and trimers at the cell surface, aggregation might be a precondition to receptor’s activation. It has been proposed that these complexes increase the avidity towards LIR-1, explaining the discrepancy between the relatively low affinity of LIR-1 to its ligand and the high relevance within the context of fetal tolerance [58]. Homo- or heterotypic complex formation of the LIR-1 receptor and its ligands might be a key factor that regulates the degree of NK cell inhibition. By this, LIR-1 might work as a rheostat of NK cell activation. This hypothesis would fit the capability of LIR-1
Figure 9: Use of F(ab’)2 after blocking NK-92 receptors seems to balance lysis pattern. Values show the % relative increase or decrease of target cells lysis in a standard 4-hour CRA, calculated as follows: % relative change of specific lysis = (% specific lysis “antibody”) – (% specific lysis “untreated”) / (% specific lysis “untreated”). Though no concentration of the used mAbs showed a significant change in tumor lysis, interesting differences in the resulting lysis-patterns could be detected. (a) Apart from outliers, use of F(ab’)2 evens out mAb effects in the blocking experiments, leading to values that oscillate close to the origin in both directions. (b) Sparing F(ab’)2 decreases specific lysis, predominantly for MOLP-8 and RPMI8226 cells.

5. Conclusions

Within the present study, no alteration of NK mediated cytotoxicity against MM was observed after blockade of LIR-1. Being the only functional inhibitory receptor within this setting, major known side effects by, for example, KIR have been ruled out. This unexpected outcome opens the door to fruitful discussions about the complexity of LIR-1 interactions and its potential role within tumor defense. It is very likely that the present presumptions about functions of LIR-1 within immune regulation are far behind the real impact. We hypothesize that LIR-1 has a key role as a rheostat of NK cell modulation and is strongly involved in the acquisition of NK cell memory.

Conflict of Interests

The authors have no financial conflict of interests.

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