Identification and Molecular Characterization of NKp30, a Novel Triggering Receptor Involved in Natural Cytotoxicity Mediated by Human Natural Killer Cells

By Daniela Pende,* Silvia Parolini,‡ Anna Pessino,§ Simona Sivori,§ Raffaella Augugliaro,§ Luigia Morelli,§ Emanuela Marcenaro,§ Laura Accame,* Angela M alaspinia,* R oberto Biassoni,* Cristina Bottino,* Lorenzo Moretta,* and Alessandro Moretta§

From the *Istituto Nazionale per la Ricerca sul Cancro, 16132 Genova, Italy; ‡Dipartimento di Scienze Biomediche e Biotecnologie, Università di Brescia, 25100 Brescia, Italy; and the §Dipartimento di Medicina Sperimentale, Università di Genova, 16132 Genova, Italy

Summary

Two major receptors involved in human natural cytotoxicity, NKp46 and NKp44, have recently been identified. However, experimental evidence suggested the existence of additional such receptor(s). In this study, by the generation of monoclonal antibodies (mAbs), we identified NKp30, a novel 30-kD triggering receptor selectively expressed by all resting and activated human natural killer (NK) cells. Although mAb-mediated cross-linking of NKp30 induces strong NK cell activation, mAb-mediated masking inhibits the NK cytotoxicity against normal or tumor target cells. NKp30 cooperates with NKp46 and/or NKp44 in the induction of NK-mediated cytotoxicity against the majority of target cells, whereas it represents the major triggering receptor in the killing of certain tumors. This novel receptor is associated with CD3ζ chains that become tyrosine phosphorylated upon sodium pervanadate treatment of NK cells. Molecular cloning of NKp30 cDNA revealed a member of the immunoglobulin superfamily, characterized by a single V-type domain and a charged residue in the transmembrane portion. Moreover, we show that NKp30 is encoded by the previously identified 1C7 gene, for which the function and the cellular distribution of the putative product were not identified in previous studies.

Key words: natural killer cells • triggering receptor • natural cytotoxicity

Natural killer (NK) cells provide an efficient effector mechanism by which immunosurveillance eliminates tumor or virally infected cells (1–3). A well-defined characteristic of NK cells is their ability to lyse target cells deficient in expression of MHC class I molecules (4). This observation has been the basis for identification of different inhibitory receptors expressed by NK cells upon binding to MHC class I molecules expressed on target cells, these receptors deliver inhibitory signals that down-regulate cytolytic functions (1–3, 5, 6). In humans, recognition of HLA class I molecules is mediated by two types of receptors: those belonging to the Ig superfamily (Ig-SF),¹ which includes both killer inhibitory receptor (KIR; 7–14) and LIR-1/ILT-2 proteins (15, 16) whose ligands are represented by various groups of HLA-A, -B, and -C alleles; and the lectin-like CD94/NKG2A receptor complex (17), which recognizes HLA-E molecules (18, 19). The expression of these inhibitory receptors explains how NK cells can distinguish between HLA-deficient and normal cells (20, 21). However, limited information existed on the activating NK receptors responsible for triggering natural cytotoxicity. Only recently, two distinct NK-specific receptors have been identified that play an important role in the NK cell-mediated recognition and killing of HLA class I–defective target cells. These receptors, termed NKp46 and NKp44 (22, 23), are members of the Ig-SF (24, 25). Their cross-linking, induced by specific mAbs, leads to a strong NK cell activation resulting in increased intracellular Ca²⁺ levels, triggering of cytotoxicity, and lymphokine release (22, 23). Importantly, mAb-mediated masking of NKp46 and/or NKp44 resulted in inhibition of NK cytotoxicity against most, but not all, target cells. These findings, while providing evidence for a central role of NKp46 and NKp44 in natural cytotoxicity, also implied the existence of additional receptors (23, 24, 26).

¹Abbreviations used in this paper: GAM, goat anti–mouse antiserum; HRPO, horseradish peroxidase; Ig-SF, Ig superfamily; ITAM, immunoreceptor tyrosine-based activating motif; KIR, killer inhibitory receptor; NCR, natural cytotoxicity receptor; ORF, open reading frame; RT, reverse transcriptase.
In this study, we identified and characterized NKP30, a third triggering receptor involved in NK cell–mediated recognition and killing of target cells. NKP30 is a member of the IgSF characterized by a single V-type domain that is selectively expressed on the surface of human NK cells.

**Materials and Methods**

The following mAbs were produced in our lab: JT3A (IgG2a, anti-CD3); BAB281 (22) and KL247 (IgG1 and IgM, respectively, anti-NKp46); Z231 (23) and K338 (IgG1 and IgM, respectively, anti-NKp44); KD1 and c127 (IgG2a and IgG1, respectively, anti-CD16); c218 and GPR165 (IgG1 and IgG2a, respectively, anti-CD56); A6-136 (IgM, anti-HLA class I [6]); GL183 (IgG1, anti-p58.2 [7]); EB6 (IgG1, anti-p58.1 [8]); and Z199 (IgG2b, anti-NKG2A [17]).

D1.12 (IgG2a, anti–HLA-DR) mAb was provided by Dr. R. S. Accolla (University of Insibria, Varese, Italy). H2P2.6 (IgG2a, anti-CD4) mAb was provided by Dr. P. Sanchez-Madrid (Hospital de la Princesa, Madrid, Spain).

The novel mAbs were derived by immunizing 5-wk-old Balb/C mice with activated (CD3+, CD56+, and CD16+) NK cells, either NK clones (EC1 and SA260 for A76 and Z25 mAbs, respectively) or a polyclonal NK cell population (for AZ20 mAb). After different cell fusions, the mAbs were selected for the ability to induce lysis in redirected killing assays against the FcγR-I+ P815 target cells.

Purification of PBLs and Generation of Polyclonal or Clonal NK Cell Populations. PBLs were derived from healthy donors by Ficoll-Hypaque gradients and depletion of plastic-adherent cells. To obtain enriched NK cells, PBLs were incubated with anti-CD3 (JT3A), anti-CD4 (HP2.6), and anti–HLA-DR (D1.12) mAbs (30 min at 4°C), followed by goat anti–mouse-coated Dynabeads (Dynal; 30 min at 4°C) and immunomagnetic depletion (21-23). CD3+CD4+DR- cells were used in cytolytic assays or cultured on irradiated feeder cells in the presence of 100 U/ml IL-2 (Proleukin; Chiron Corporation) and 1.5 ng/ml PHA (GIBCO BRL) to obtain polyclonal NK cell populations or, after limiting dilution (27), NK cell clones.

Flow Cytometeric Analysis. Cells were stained with the appropriate mAb followed by PE- or FITC-conjugated isotype-specific goat anti–mouse second reagent (Southern Biotechnology Associates Inc.). Samples were analyzed by one- or two-color cytometric analysis (FACScan®; Becton Dickinson), as described previously (7).

Cell Lines and Cytolytic Assays. The following FcγR-negative targets were used: MEL15 (Mel15392, human melanoma [21]); M14 (human melanoma [24]); SM MC (human hepatocarcinoma [26]); A549 (human lung adenocarcinoma, no. CCL-185.1; American Type Culture Collection). FO-1 and 1174 (melanoma) were provided by Dr. S. Ferrone (Roswell Park Cancer Center, Buffalo, N.Y.); AUMA (human melanoma) was provided by Dr. P. Couille (Catholic University of Louvain, Brussels, Belgium). The FcγR-positive target was used was P815 (murine mastocytoma). PHA blasts, used as target cells, were obtained by culturing PBLs with 1.5 ng/ml PHA (GIBCO BRL).

Cells were tested for cytolytic activity in a 4-h 51Cr-release assay as described previously (8, 26), either in the absence or presence of various mAbs. The concentrations of the various mAbs were 10 μg/ml for the masking experiments and 0.5 μg/ml for the redirected killing experiments. The E/T ratios are indicated in the text.

Determination of Intraocular Free Cadmium [Ca2+]. Increase. Determination of [Ca2+], was performed as described previously (28). Fura-2-labeled NK cells were incubated for 30 min at 4°C with saturating amounts of anti-NKp30 mAb (AZ 20) or medium alone. Cross-linking of this receptor was obtained by adding into the cuvette 20 μg/ml of affinity-purified goat anti–mouse antibody (GAM; ICN Biomedicals, Inc.).

Biological Characterization of NKP30 Molecules. Integral NK cell membrane proteins (29) were prepared as follows: 25 × 106 cells were lysed in 100 μl TX buffer (20 mM sodium phosphate buffer, 1% Triton X-114, 10 mM EDTA, pH 8) for 30 min at 4°C, and centrifuged (5 min, 10,000 rpm). The supernatant was left for 10 min at 37°C, centrifuged, and the lower phase was resuspended 1:2 in TX buffer and left for 10 min at 4°C in order to clarify the lysates. The suspension was then left for 10 min at 37°C, centrifuged, and the lower phase was resuspended 1:3 in EB (0.0625 M Tris, pH 6.8, 10% glycerol, 2.3% SDS). Samples were analyzed in discontinuous SDS-PAGE, transferred to Immobilon P (Millipore Corp.,), and probed with AZ20 mAb following rabbit anti–mouse horseradish peroxidase (HRPO; DAKO), or NKP30-specific antiserum followed by donkey anti-rabbit HRPO (Nycamed Amersham plc). The Renaissance Chemiluminescence kit (NEN Life Science Products) was used for detection.

NKP30 Polyclonal Antibodies. A 2.5-kg HY/Cr male rabbit (Charles River Laboratories) was immunized with 100 μg/100 μl of the 15 amino acid peptide WVSQPPEIRTLEGSC (Eurogentec) conjugated with KLH (13). Four weekly treatments were performed, the first in association with 100 μl CFA and all others with 100 μl IFA, 1 wk after the last treatment, 10 ml of blood was drawn, and serum was tested and titered by ELISA against the immunizing and irrelevant peptides.

Analysis of the NKP30 Signal Transduction Complex. NK cells (107) were stimulated or not with 100 μM sodium pervanadate (25), and 1% digitonin lysates were preincubated five times with sepharose-protein A–coupled KD1 (anti-CD16) mAb. Lysates were then immunoprecipitated with sepharose-CNBr–coupled Z231 and BAB281 mAbs, or with sepharose protein A–coupled NKp30-specific rabbit antiserum and preimmune rabbit serum. Samples were analyzed in a 15% SDS-PAGE under reducing conditions (5% 2-ME), transferred to Immobilon P (Millipore Corp.), and probed with antiphosphotyrosine mAb (PY20-HRPO; Transduction Laboratories) or anti-CD3ζ mAb (2H2; Immunotech), followed by rabbit anti–mouse HRPO (DAKO). The Renaissance Chemiluminescence kit (NEN Life Science Products) was used for detection.

Library Screening by cDNA Expression in COS-7 Cells. The expression cDNA library was prepared in VR 1012 plasmid (Vical Inc.) using RNA extracted from IL-2–activated polyclonal NK cells obtained from two healthy donors as described previously (24, 25). The library screening procedure was as described (24, 25, 30). In brief, cDNA library was transiently transfected in COS-7 cells, and selection of positive pools was performed by immunocytochemical staining using the specific anti-NKp30 mAb A76 and sib-selection.

DNA Sequencing. DNA sequencing was performed using d-R hodaamine Terminator Cycle Sequencing kit and a 377 ABI Automatic Sequencer (Applied Biosystems/Perkin-Elmer).

Transient Transfections. COS-7 cells (5 × 105/plate) were transfected with VR 1012-NK-A1 (clone 5C) with the vector alone by the DEAE-dextran or electroporation methods as described (13). After 48 h, transfected cells were used for cytotoxicity and analysis of NKp30 TRAP expression by Northern Blotting. To analyze NKP30 transcript expression in different cell lines of
hemopoietic origin, RNA was size fractionated by denaturing agarose gel electrophoresis and transferred onto a positively charged nylon membrane (ENLife Science Products). In particular, 10 μg of total RNA prepared using CsCl gradient, or 2 μg of poly A+ RNA prepared using oligo(dT) magnetic bead separation (Dynal AS), was loaded on each lane. Northern blots were performed under stringent conditions as described (31). The NKp30 421-bp cDNA probe was obtained by PCR amplification performed with 25 pmol of each primer for 30 cycles (30 s at 94°C, 30 s at 60°C, 30 s at 72°C), followed by a 7-min incubation at 72°C. The sequences of the primers are: 5′ CAG GGC ATC TCG AGT TTC CGA CAT GGC CTG GAT GCT GTT G (NK-A1 up) and 5′ GAT TTA TTG GGG TCT TTT GAA G (A76-38 reverse). Amplification was performed with 25 pmol of each primer for 30 cycles (30 s at 94°C, 30 s at 60°C, and 30 s at 72°C), followed by a 7-min incubation at 72°C. The amplification products were subcloned in pCR2.1 by TOPO-TA Cloning kit (Invitrogen), and subsequently sequenced.

Zoo-Blot Analysis. A clone of cross-species conservation of NKp30 gene was performed using a Zoo-Blot (CLONTECH). The Southern blot contained genomic DNA from humans, Rhesus monkey, Sprague-Dawley rat, BALB/c mouse, dog, cow, rabbit, chicken, and Saccharomyces cerevisiae yeast. The hybridization probe was the same 421-bp cDNA fragment used to hybridize the Northern blot. Washes were carried out at low stringency conditions as described (32).

**Results**

Identification of a Novel NK-specific Triggering Surface Molecule. Mice were immunized with CD3+, CD16+, CD56+ NK cell clones or bulk populations. mAbs from different fusions were first selected according to their ability to induce lysis of the FcyR+ P815 target cells in a redirected killing assay using polyclonal NK cell populations or clones as effector cells. Three mAbs, A76, AZ20, and Z25 (all of IgG1 isotype), were selected that induced a strong cytolytic activity (Fig. 1 A) similar to that elicited by other mAbs specific for known triggering NK receptors, including CD16, NKp46, and NKp44 (22, 23, 26). In Fig. 1 B, the NK cell cytotoxicity induced by graded amounts of AZ20 mAb is compared with that of isotype-matched anti-CD16 or anti-CD56 mAbs. The cytolytic response to AZ20 mAb paralleled that induced by anti-CD16 mAb, whereas anti-CD56 mAb had no effect. Moreover, as shown in Fig. 1 C, a sharp [Ca2+]i increase was detected in the representative clone 3M16 after stimulation with AZ20 mAb. Notably, [Ca2+]i increments induced by this Ab occurred only in the presence of a goat anti-mouse secondary reagent, allowing efficient cross-linking of the activating receptor.

Analysis of the cell surface distribution of the molecule(s) recognized by A76, AZ20, and Z25 mAbs, performed by indirect immunofluorescence and FACS® analysis, revealed reactivity with various activated polyclonal or clonal NK cell populations derived from different donors (see below). These also included the infrequent CD16− NK cell clones. On the contrary, no mAb reactivity was detected with PHA-induced polyclonal T cell populations or TCR-α/β and -γδ T cell clones (derived from different donors). Neither was any reactivity detected with EBV-induced B cell lines, monocytic and dendritic cell lines, and different hemopoietic and nonhemopoietic tumor cell lines, including HL60, U937, E01/A3, THP-1, Daudi, Jurkat, IGR-OV, and all the various tumor cell lines used as target cells in this study (data not shown).

We recently showed that polyclonal NK cell populations from some donors were characterized by a bimodal distribution of fluorescence intensity of NKp46 molecules (NKp46high and NKp46dim), and that NK clones derived from these individuals expressed a stable NKp46high or

**Figure 1.** Triggering of NK-mediated cytolytic activity induced by three new mAbs. (A) A representative polyclonal NK cell population was analyzed for cytolytic activity in a redirected killing assay against the FcyR+ P815 target cell line in the absence or presence of c127 (anti-CD16), BAB281 (anti-NKp46), Z231 (anti-NKp44), AZ20, A76, Z25, and c218 (anti-CD56) mAbs. The E/T ratio used was 1:1. (B) The representative NK clone 3M16 was analyzed in a redirected killing assay against P815 target cells (E/T ratio 1:1) in the presence of graded amounts of AZ20 ( ), c127 (anti-CD16: △), or c218 (anti-CD56: ○) mAbs. All the mAbs used are of the IgG1 isotype. (C) Clone 3M16 was analyzed for [Ca2+]i mobilization in the presence of AZ20 mAb, followed by GAM®. The negative control is represented by cells treated with GAM alone.
NKp46<sup>dull</sup> phenotype (26). Importantly, the cytolytic activity of NK cell clones against NK-susceptible target cells strictly correlated with their NKp46 phenotype (26). We then analyzed the reactivity of the new mAbs on polyclonal NK cell populations and NK cell clones derived from individuals displaying different patterns of NKp46 expression. As shown in Fig. 2 A, the polyclonal NK cell population derived from the representative donor AM displayed a homogeneously bright phenotype when stained by either AZ20 or anti-NKp46 mAbs. On the contrary, in the polyclonal NK cells derived from donor CB, staining with the same mAbs resulted in a bimodal distribution of fluorescence. Notably, in donor CB, the same pattern of fluorescence intensity was also detectable in fresh purified NK cells (Fig. 2 A). Moreover, the analysis of several clones derived from donor CB revealed that NKp46<sup>bright</sup> clones were
consistently AZ20bright, whereas NKp46dull clones always displayed an AZ20dull phenotype (Fig. 2 B).

To further define the pattern of reactivity of the new mAbs in freshly isolated lymphocytes, PBLs derived from different individuals were assessed by double fluorescence analysis using informative mAbs. A representative donor is shown in Fig. 3 A: the surface molecule recognized by AZ20 mAb was selectively expressed on CD56+ cells. Moreover, most AZ20+ cells coexpressed CD16 molecules. However, AZ20 mAb did not stain CD3+ T lymphocytes or HLA-DR+B lymphocytes. It is of note that the CD56+ AZ20– cell population detected in this donor also expressed surface CD3 molecules (not shown). Therefore, also in freshly derived lymphocytes, the reactivity of AZ20 mAb overlaps with that of anti-NKp46 mAb. A direct comparative analysis of the surface expression of NKp46 and AZ20 mAb–reactive molecules is shown in Fig. 3 A. The two molecules were clearly coexpressed by the same cell subset. However, no diagonal distribution could be detected in cells stained by AZ20 and anti-NKp46 mAbs, whereas this type of fluorescence distribution occurred when cells were stained simultaneously by two anti-NKp46 mAbs of different isotype.

Notably, results identical to those described for AZ20 mAb were obtained with A76 and Z25 mAbs. These data suggested that the molecule recognized by the new mAbs may be distinct from NKp46. To directly evaluate this possibility, COS-7 cells transiently transfected with NKp46 cDNA (24) were analyzed for their reactivity with AZ20, A76, and Z25 mAbs. Cell transfectants, although reacting with different anti-NKp46 mAbs, were not stained by AZ20, A76, and Z25 mAbs (data not shown). Taken together, these data strongly suggest that A76, AZ20, and Z25 mAbs are specific for a novel surface molecule that defines all mature human NK cells, but is distinct from NKp46.

To analyze the biochemical characteristics of the surface molecules recognized by AZ20, A76, and Z25 mAbs, NK populations were surface labeled with 125I or biotin, immunoprecipitated with one or another mAb, and analyzed by SDS-PAGE. Under these conditions, no specific bands could be detected. Thus, integral membrane proteins were prepared from NK cells to further analyze a possible reactivity of the various mAbs in Western blot. As shown in Fig. 3 B, AZ20 mAb specifically reacted with an ~30-kD molecule, thereafter termed NKp30. Under the same conditions, both A76 and Z25 mAbs displayed a poor reactivity (data not shown).

Cross-linking of NKp30 Induces Cytolytic Activity in Freshly Derived NK Cells. Since NKp30 molecule, like NKp46, was expressed on fresh NK cells, we analyzed whether it could trigger the cytolytic activity of these cells as demonstrated previously for NKp46. As shown in Fig. 4 A, AZ20, A76, and Z25 mAbs induced a strong increase of cytolytic activity against P815 target cells, whereas the isotypespecific anti-CD56 mAb had no effect. This triggering effect was comparable to that obtained with anti-NKp46 mAb. Moreover, in these experiments, the use of AZ20 F(ab')2 fragments did not induce triggering of cytolytic activity, indicating that mAb-dependent NKp30 stimulation requires efficient cross-linking mediated by FcγR on target cells (data not shown).

Involvement of NKp30 in the Induction of Natural Cytotoxicity against Normal or Tumor Cells. Previous data showed...
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that mAb-mediated masking of NKp46 or NKp44 inhibited the non–M HC-restricted tumor cell lysis by activated NK cells (23, 24, 26). Moreover, masking of NKp46 also inhibited the natural cytotoxicity mediated by freshly isolated peripheral blood NK cells (26). We then evaluated whether masking of NKp30 could affect the cytolytic activity mediated by freshly derived NK cells or NK clones against a panel of FcγR-negative tumor target cells. As shown in Fig. 4 B, anti-NKp30 mAb, but not the isotype-matched anti-CD56 mAb, inhibited natural cytotoxicity mediated by fresh NK cells against the HLA class I-negative 1174 mel, AUMA, and FO-1 melanoma cell lines. In addition, a greater inhibitory effect occurred when anti-NKp30 mAb was used in combination with anti-NKp46 mAb. This suggests that NKp30 and NKp46 may represent receptors that act synergistically in triggering the natural cytotoxicity of fresh NK cells.

In view of these data, we further analyzed the effect of mAb-mediated masking of NKp30 on the tumor cell killing by activated NK cells. Fig. 5 shows three representative NK cell clones analyzed in a cytolytic assay against different tumor targets, including two melanomas (MEL15 and M14), a hepatocarcinoma (SMC), and a lung adenocarcinoma (A549). In previous studies, we showed that the cytolytic activity against the M14 melanoma was confined to NK clones displaying the NKp46bright phenotype and could be inhibited by mAb-mediated masking of NKp46 receptor. On the other hand, NKp46bright clones also killed MEL15. However, neither masking of NKp46 nor of NKp44 significantly inhibited their cytolytic activity (26). These data strongly suggested the existence in these clones of additional triggering receptors responsible for the cytotoxicity against MEL15 target cells. As illustrated above, NKp30 is brightly expressed in NKp46bright clones. Therefore, it is conceivable that it may play a role in the killing of MEL15 target cells. Indeed, as shown in Fig. 5, anti-NKp30 mAb sharply inhibited the NK-mediated lysis of MEL15 cells (>50% of inhibition). Anti-NKp46 mAb exerted a minor effect, whereas an isotype-matched anti-CD56 mAb had no effect (data not shown). On the contrary, lysis of M14 melanoma was inhibited by anti-NKp46 mAb, whereas anti-NKp30 mAb had virtually no effect. Thus, while NKp46 appears as the major receptor involved in lysis of M 14, NKp30 plays a central role in the killing of MEL15.
Analysis of the same NK clones in cytolytic assays against other tumor target cells such as SMCC and A549 (Fig. 5) revealed a balanced contribution of NKp46 and NKp30 to the induction of cytotoxicity. Indeed, while mAb-mediated masking of NKp46 or NKp30 alone had a moderate inhibitory effect, the simultaneous masking of the two molecules resulted in a significant inhibition. These results indicate that the two receptors may exert an additive effect in the induction of cytotoxicity against certain target cells. Cooperation in NK cell triggering was previously demonstrated for NKp46 and NKp44 (23). Further analysis revealed that NKp30 could exert an additive effect in the induction of NK-mediated cytotoxicity, not only with NKp46, but also with NKp44. Fig. 6A shows the cytolytic activity of the representative NK clone MIL69 against FO-1 or A549 tumor cells. Target cell lysis was only partially inhibited by mAb-mediated masking of NKp30, NKp44, or NKp46 receptors. However, the combined masking of two receptors resulted in a higher inhibitory effect, whereas the simultaneous masking of the three receptors gave the maximal inhibition. Isotype-matched anti-CD56 mAb had no inhibitory effect either when used alone or in combination with other mAbs (data not shown).

We further analyzed the role of NKp30 alone or in combination with other receptors in cytolytic assays using PHA-induced T cell blasts as a source of normal target cells. In these experiments, lysis of autologous cells by NK cell clones was obtained by mAb-mediated masking of HLA class I molecules on target cells to disrupt the interaction with the HLA class I–specific inhibitory receptors expressed on NK cells. Also, under these experimental conditions, the mAb-mediated masking of single receptors had only a partial inhibitory effect on cytotoxicity (Fig. 6B).

Figure 6. NKp30 cooperates with NKp46 and NKp44 in the induction of NK-mediated cytotoxicity against tumor or normal autologous target cells. (A) The representative NK clone MIL69 was analyzed for cytolytic activity against FO-1 or A549 FcγR-negative target cell lines either in the absence or presence of mAbs to the indicated molecules. The following mAbs were used: KL247 (anti-NKp46), AZ20 (anti-NKp30), and KS38 (anti-NKp44). The E/T ratios were 2.1 to 3.1 (A549). (B) Two NK cell clones (MX361 and P9) were analyzed for cytolytic activity against autologous PHA blasts in the absence (white bars) or presence of mAbs to the indicated molecules (black bars). The mAbs used were A6-136 (anti-HLA class I) KL247 (anti-NKp46), KS38 (anti-NKp44), and AZ20 (anti-NKp30). The E/T ratio was 10:1.
ingle open reading frame (O R F) of 573 bp. T ransfection of COS-7 cells with clone 5C cDNA construct resulted in the surface expression of a molecule that was recognized by all the various anti-NKp30 mAbs (Fig. 7 A), but not by anti-NKp46 mAbs as assessed by cytofluorimetric analysis. A s shown in Fig. 7 B, clone 5C ORF encoded a putative 190 amino acid polypeptide belonging to the Ig-SF, characterized by a signal peptide of 18 amino acids and by an extracellular region of 120 amino acids forming an Ig-like domain of the V type. The extracellular portion contains two potential N-linked glycosylation sites and no consensus sequences for O-linked glycosylation. A region rich in hydrophobic amino acids, potentially involved in protein-protein interactions, is connecting the Ig V-like domain with the transmembrane region. The 19 amino acid transmembrane region contains the positively charged amino acid, Arg, and the 33 amino acid cytoplasmic portion lacks typical immunoreceptor tyrosine-based activating motif (ITAM) consensus sequences. The presence of a charged amino acid in the transmembrane domain is a feature common to other triggering receptors expressed on NK cells (24, 25, 34–37). T hese charged residues are usually thought to be involved in the association with ITAM-containing signaling polypeptides.

S earching E M B L / G e n B a n k / D D B J databases revealed that the clone 5C cDNA was identical to a previously identified alternatively spliced form of the 1C7 gene (available under accession no. AF031138). T his gene has been mapped on human chromosome 6, in the T NF cluster of the MHC gene complex (38). T o date, however, owing to the lack of specific mAb, neither the function nor the surface distribution of the putative product of 1C7 gene could be identified. M oreover, the 1C7 transcript could not be revealed by Northern blot on different tissues and cell lines. O n the other hand, by reverse transcriptase (R T)-P C R, the 1C7 transcript could be amplified by RNA isolated from spleen (but not from other tissues) or certain lymphoid and myeloid cell lines. T hese data suggested that 1C7 transcripts could be poorly represented, or could be expressed at substantial levels only in a narrow range of cell types (39). O ur present analysis of NKp30 expression by Northern blotting revealed a mRNA of ~1 kb in polyclonal NK cell populations and NK cell lines, including NKL and NK3.3. O n the contrary, consistent with the lack of reactivity with anti-NKp30 mAbs, no NKp30 mRNA could be detected in human monocytes or cell lines of different histotype, including U937, Jurkat, HL60, and LCL 721.221 cells (Fig. 8 A). I n some of these cell lines that were negative for mRNA expression by Northern blot (and for anti-NKp30 mAb surface staining), it has been possible to detect transcripts when analyzed by the R T -P C R technique. T his finding is likely to reflect a low level of NKp30 transcription resulting in lack of NKp30 surface expression. M oreover, Northern blot analysis of multiple human tissues showed selective expression of NKp30 transcript only in spleen cells (data not shown). A ltogether, these data are consistent with the notion that NKp30 expression is largely NK specific.

F inally, the human NKp30 cDNA probe hybridized with genomic DNA from monkey, rat, mouse, dog, cow, and rabbit. T hese data suggest that the NKp30-encoding gene may be conserved in different species (Fig. 8 B).

B ioc hemical C haracterization of the NKp30 C omplex. A n NKp30-specific antiserum was generated by immunizing rabbits with an N H2-terminal NKp30 peptide. A s shown in Fig. 9 A, the antiserum recognized in Western blot a molecule identical to that previously detected by AZ20 mAb. U nlike the AZ20 mAb, the antiserum immunoprecipitates NKp30 molecules from polyclonal NK cell populations labeled with biotin (data not shown). T hus, a polyclonal NK cell population, treated or not with sodium pervanadate, was immunoprecipitated with the NKp30-specific antiserum and probed with antiphosphotyrosine mAb. T o avoid nonspecific binding of rabbit Ig to CD16 molecules, cell lysates were extensively precleared with anti-CD16 mAb. M oreover, in all experiments, preimmune rabbit serum was used as negative control. I n these experiments, no tyrosine phosphorylation of NKp30 receptor could be detected (data not shown). O n the other hand, NKp30 receptor associated with a molecule that became tyrosine phosphorylated upon sodium pervanadate treatment (Fig. 9 B) and comigrated with the NKp46-associated CD3ζ chain. T he
identity between the NKp30-associated molecule and CD3ζ polypeptides was directly demonstrated by its reactivity with anti-CD3ζ mAb (Fig. 9 B).

Thus, NKp30, similar to other NK triggering receptors including CD16 (34, 35) and NKp46 (23), can transduce activating signals via association with the ITAM-containing CD3ζ polypeptides. These data are in agreement with the lack of ITAM in the NKp30 cytoplasmic tail, and with the presence of a charged residue in its transmembrane portion.

Discussion

In this study, because of the generation of specific mAbs, we identified and characterized NKp30, a novel triggering receptor that plays an important role in the natural cytotoxicity of both resting and activated human NK cells. Similar to NKp46, NKp30 is selectively expressed by all NK cells, both freshly isolated and cultured in IL-2, thus representing an optimal marker for NK cell identification. Although it belongs to the Ig-SF, NKp30 does not display any substantial homology with previously identified NK receptors.

In many respects, NKp30 appeared similar to NKp46. Indeed, their parallel expression on all NK cells (including the rare CD16- cells), the existence for both of a high or low density pattern of surface expression, together with their similar functional characteristics, led to the thought that the surface molecule recognized by the new mAbs could be identical or strictly related to NKp46. However, NKp30 and NKp46 displayed different molecular masses and, functionally, appeared to play a complementary role in the induction of natural cytotoxicity. Moreover, molecular cloning revealed that NKp30 is a protein with very limited homology with NKp46, as the two molecules display only 13% identity and 15% similarity, and are encoded by genes located on different chromosomes.

The receptors responsible for the NK cell triggering during natural cytotoxicity and tumor cell lysis have remained elusive until recently. Available data were consistent with the hypothesis of the existence of multiple triggering NK receptors involved in natural cytotoxicity. In this context,
we recently identified NKp46 and NKp44, two receptors involved in recognition and lysis of a variety of tumor targets. Both belong to the Ig-SF, but neither displays significant identity. They appear to be involved in different signal transducing polypeptides (CD3ε/FcεRIγ and KARAP/DAP12, respectively) that become tyrosine phosphorylated upon NK cell activation. NKp46 and NKp44 were shown to cooperate in the process of tumor cell lysis by human NK cells. However, lysis of certain target cells was only marginally NKp46 and/or NKp44 dependent, since mAb-mediated masking of these molecules did not significantly interfere with cytotoxicity. This finding strongly suggested the existence of additional triggering receptors that could induce cytotoxicity against these target cells. Moreover, although clearly NKp46 and/or NKp44 dependent, the cytolytic activity against other tumor cell lines could not be abrogated by mAb-mediated masking of both molecules, suggesting again the existence of additional receptor(s) cooperating with NKp46 and NKp44. Indeed, we show here that NKp30 represents a receptor that may cooperate with NKp46 and NKp44 in the induction of cytotoxicity against a variety of target cells. Perhaps more importantly, NKp30 represents the major receptor in inducing NK-mediated killing of certain tumor target cells, the lysis of which is largely NKp46/NKp44 independent. Remarkably, NKp30, similar to NKp46, is also involved in NK cell activation and target cell killing by fresh NK cells.

As discussed above, the surface expression of NKp30 parallels that of NKp46. Indeed, NK cells displaying an NKp46null or an NKp46bright phenotype were also characterized by NKp30null or NKp30bright fluorescence. We showed previously that NK cell clones characterized by an NKp46null phenotype consistently express low amounts of NKp44. The finding that NK cells express parallel densities of different triggering receptors may explain the existence of NK cell subsets displaying different “natural” cytolytic activity. For example, it was difficult to understand why the cytolytic activity against some target cells (such as MEL15), although largely NKp46 independent, was essentially confined to NK clones expressing the NKp46bright phenotype. These results can now be explained by the finding that only NKp46bright cells express a high density of NKp30 receptor. Thus, the previous demonstration of major differences in cytolytic activity of NKp46null and NKp46bright cells can now be applied also to NK cells displaying different NKp30 phenotypes. Along this line, the cytolytic activity of NKp30null NK cell clones was markedly reduced compared with NKp30bright clones (data not shown).

NKp30, similar to NKp46, associates with CD3ε, which is most likely involved in signaling via the receptor complex. However, CD3ε does not appear to be necessary for the surface expression of both receptors, at least in COS-7 cells (this report, and reference 24). Molecular cloning revealed that NKp30 is the product of IC7, a gene previously mapped on human chromosome 6 in the HLA class III region (38, 39). However, neither the function nor the cellular distribution of the putative product of IC7 gene was known, and no indications existed of its role in natural cytotoxicity. In addition, the analysis of IC7 transcript expression was limited to RT-PCR, whereas no detection had been possible by Northern blot analysis (39). It should also be stressed that no correlation between transcript and surface expression could be established because of the lack of specific mAbs. In this study, we show that a precise correlation exists between the surface expression of NKp30, as determined by staining with three different mAbs, and mRNA expression, as assessed by Northern blot. On the contrary, the detection of IC7 transcripts by RT-PCR does not allow prediction of the surface expression of the IC7/NKp30 molecule.

In conclusion, the NKp30 molecule represents a third member of an emerging family of receptors, termed natural cytotoxicity receptors (NCRs; 40), that are involved in NK cell triggering upon recognition of non-HLA ligands. These receptors appear to complement each other in the induction of target cell lysis by NK cells. The relative contribution of each receptor is likely to reflect the expression/density of their specific ligands on target cells. It has recently been shown that CD16 is also involved in natural cytotoxicity, thus suggesting that in addition to Fc-binding and antibody-dependent cell-mediated cytotoxicity, CD16 may play a role in the regulation of NK cell function (41). Besides CD16 and the different NCRs, several other surface molecules that can mediate NK cell triggering have been identified in humans and rodents. These include CD2 (42, 43), CD69 (44), CD28 (45), 2B4 (46), and NKR-P1 (47). However, their actual role in natural cytotoxicity has yet to be clarified, since in most instances these activating structures are not NK restricted.

Finally, although the identification of different NCRs constitutes a major step forward in our understanding of the NK cell physiology, both the nature and the distribution of the NCR ligands on target cells remain to be determined. Based on the available data, it is possible to envisage a novel mechanism of tumor escape consisting in the downregulation (on tumor cells) of ligand molecules specifically recognized by NK-specific triggering receptors. Thus, the identification of such ligands will allow the analysis of their distribution in normal versus tumor cells, and define whether a correlation exists between ligand expression and susceptibility to NK-mediated lysis by different tumor cells.

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Address correspondence to Lorenzo Moretta, Laboratorio di Immunopatologia, Centro Biotecnologie.

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