Rescue of Impaired NK Cell Activity in Hodgkin Lymphoma With Bispecific Antibodies In Vitro and in Patients

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Natural killer (NK) cells represent a key component of the innate immune system against cancer. Nevertheless, malignant diseases arise in immunocompetent individuals despite tumor immunosurveillance. Hodgkin lymphoma (HL) is characterized by CD30+ tumor cells and a massive infiltration of immune effector cells in affected lymph nodes. The latter obviously fail to eliminate the malignant cell population. Here, we tested for functional NK cell defects in HL and suggest an improvement of NK function by therapeutic means. We demonstrate that peripheral NK cells (pNK) from patients with HL fail to eliminate HL cell lines in ex vivo killing assays. Impaired NK cell function correlated with elevated serum levels of soluble ligands for NK cell receptors Nkp30 (BAG6/BAT3) and NKG2D (MICA), factors known to constrict NK cell function. In vitro, NK cell cytotoxicity could be restored by an NKG2D/Nkp30-independent bispecific antibody construct (CD30xCD16A). It artificially links the tumor receptor CD30 with the cytotoxicity NK cell receptor CD16A. Moreover, we observed that NK cells from patients treated with this construct were generally activated and displayed a restored cytotoxicity against HL target cells. These data suggest that reversible suppression of NK cell activity contributes to immune evasion in HL and can be antagonized therapeutically.

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INTRODUCTION

There is a recent clinical and experimental evidence to show that natural killer (NK) cells are critically involved in the recognition and elimination of tumor cells. The NKG2D receptor and the group of natural cytotoxicity receptors (NCRs) (NKP46, NKP44, and NKP30) are regarded as the major NK cell receptors in tumor defence.1,2 NKG2D is a member of the C-type lectin superfamily which triggers NK cell activation. Ligands of the human NKG2D receptor are the major histocompatibility complex class I (MHC I)-related molecules MICA/MICB, and the UL16-binding proteins ULBP, which are rarely expressed in healthy tissues but induced on cells upon transformation. This induction alerts the immune system to the dangerous cell.1 The ligands for NKP46 and NKP44 are still enigmatic although heparan sulfate proteoglycans, vimentin, and proliferating nuclear antigen have been described as ligand structures.3–5 A member of the B7 family, B7-H6, and the nuclear factor BAG6 (also referred to as BAT3), which is released from tumor cells under stress conditions via the exosomal pathway, were recently identified as tumor cell-associated ligands for NKP30.6,7 Silencing of NKG2D during tumor progression results from the persistent exposure of ligands expressed on the surface of target cells.9,10 Moreover, tumor cells release ligands into the environment by shedding. The soluble molecules not only block NKG2D but also induce receptor internalization and degradation.11–13 Soluble MICA molecules in addition engage NKG2D on CD4+ T cells to promote proliferation and differentiation of regulatory T cells that additionally inhibit the immune response.11

Plasma levels of soluble ligands correlate with disease progression in many hematological and solid tumors.14–17 Here, we demonstrate that peripheral NK cells from patients with Hodgkin lymphoma (HL) are functionally inactive, which is in line with former studies on NK cell function in HL.18,19 The observed NK cell dysfunction correlates to elevated serum levels for ligands engaging NKG2D (MICA) and NKP30 (BAG6/BAT3). We provide an evidence that immunotherapeutic strategies targeting NK cells are promising, because NK cell cytotoxicity could be restored in vitro and in patients by treatment with a novel human antibody construct that is designed for the treatment of HL and other CD30-expressing malignancies. The tetravalent, bispecific antibody used in this study targets CD30 on Hodgkin Reed-Sternberg (HRS) cells with two of its...
binding sites, whereas the activating receptor CD16A on NK cells (CD30xCD16A, AFM13) is targeted by the other two binding sites, thereby selectively cross-linking tumor and NK cells. CD16A (FCGR3A) is the human low-affinity IgG Fc receptor that is expressed on the surface of NK cells, macrophages, a subset of monocytes, and T cells. The engagement of CD16 triggers its interaction with both FcRI-γ and CD3-ζ immunoreceptor tyrosine-based activation motif complexes. This induces the recruitment and activation of phosphotyrosine kinases including Syk and ZAP70, finally resulting in the activation of NK cell-effector functions.

CD30, a member of the tumor necrosis factor receptor family, is highly expressed on Hes1 cells, but rarely and faintly expressed in normal tissue and thus represents an excellent target structure for immunotherapy. Although more than 80% of patients with HL are cured by combined radio- and chemotherapy, there is still a high and unmet need for both treatment options for patients who relapse or fail to respond to front-line treatment and for
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Figure 2 The surface expression of NKG2D on HL-derived pNK cells is reduced. (a) Four-color flow cytometry was performed to determine the expression of the activating NK cell receptors NKG2D, NKp30, and NKp46 on gated NK cells (CD3−/CD56+/CD16+) in HL samples. Significant differences (Mann–Whitney U test) are indicated. (b) Serum levels of MICA, BAG6, and MIF in HL samples were collected before therapy (BT), after therapy (AT), and from patients after therapy with later relapse (ATre). Significant differences are indicated (Mann–Whitney U test). (c) Histology: Tissue samples from patients with HL were stained with the BAG6-specific monoclonal antibody 3E4. Three samples from different patients are shown. ***P ≤ 0.0001; *P ≤ 0.05. HL, Hodgkin lymphoma; MIF, macrophage migration inhibitory factor; NK, natural killer.

therapies that have limited side effects.22 Our findings suggest that immunotherapeutic approaches are an effective and promising alternative to standard therapies.

RESULTS
Function and phenotype of peripheral NK cells is altered in patients with HL

It is a hallmark of HL that the malignant cells in affected lymph nodes are surrounded by immune effector cells including lymphocytes, that are unable to recognize and kill the tumor cells.23 Here, we demonstrate that the recognition and killing of the HL-derived target cell line L428 was impaired in peripheral NK (pNK) cells isolated from patients with HL (Figure 1a), although this cell line was efficiently lysed by NK cells from healthy donors. The difference between NK cell cytotoxicity from patients (samples were taken before therapy) and healthy donors was highly significant (P = 0.0001). Fluorescence activated cell sorter analysis confirmed published data24 that reported absence or very low expression of MHC I on L428 cells (Figure 1b, first panel) excluding an MHC I-mediated suppression of HL-NK cells in these assays. CD95 (APO-1/Fas) and CD262 (DR5), death receptors involved in NK cell-mediated killing and several other costimulatory adhesion molecules including ICAM-1 and ICAM-2 were expressed on L428 target cells (Figure 1b). The expression of ligands for the NCRs NKp30 and NKp46 was not detectable upon staining with recombinant receptors, whereas various ligands for NKG2D (MICA/B, members of the ULBP family) were assured using both, specific antibodies and NKG2D-Fc protein (Figure 1b). Lysis of L428 target cells by healthy NK cells was mainly dependent on NKG2D, as an NKG2D-blocking antibody was able to suppress NK cell-dependent killing (Figure 1c).

NKG2D surface expression on HL-derived pNK is reduced

We then analyzed the expression pattern of a panel of NK cell markers and receptors such as CD16, the NCRs including NKp30, NKp46, NKp44, and NKG2D. The samples were obtained from patients (mean age: 38) before (n = 40, BT), during (n = 39, DT), and on completion/after radio/chemotherapy (n = 17, AT). Surprisingly, no significant difference between the NK cells derived from healthy donors (n = 23) and those derived from untreated patients with HL was observed for CD16, NCRs, and for the activation markers CD25, CD69, and CD71, although NKp30 and NKp46 were significantly downregulated during therapy (Figure 2a and data not shown). The only distinctive feature of NK cells from patients with HL was a significantly decreased NKG2D surface expression (P = 0.0001 for healthy versus untreated patients with HL (Figure 2a, left panel)). It is well established that tumor cells release soluble ligands for NKG2D to inhibit antitumor immune responses via downregulation of the receptor.12 In fact, analysis of about 300 HL serum samples (Figure 2b) indicated that the NKG2D ligand MICA was
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Results for each sample are represented by mean fluorescence intensity (MFI). **P ≤ 0.0001; *P ≤ 0.005; *P ≤ 0.05. Significances are indicated. HL, Hodgkin lymphoma; IL-2, interleukin 2; NK, natural killer.

**HL serum derogates NK cell activity**

Consequently, to gain insight into the impact of the aberrantly expressed serum factors, we analyzed the effect of patient serum on NK cell function. Interestingly, stimulation with interleukin-2 (IL-2) resulted in the activation of pNK cells isolated from patients (HL-NK) and led to a robust target cell killing. This indicates that NK cell “anergy” is reversible (Figure 3a, right panel). However, the IL-2–mediated activation was not observed in the presence of patient serum (HL-NK/HLs), suggesting that HL serum factors suppress IL-2 function or IL-2 signaling pathways of NK cells in this disease. Supporting this conclusion, we demonstrate that HL serum was sufficient to suppress the cytotoxicity of normal (healthy donor) NK cells against target cells, whereas serum from healthy donors (NK/healthy serum) had no effect on NK cell activity (Figure 3a, left panel). In line, we observed that overnight incubation of freshly isolated NK cells with patient serum with HL (n = 5) resulted in a significant downregulation of NKG2D, NKp30, and the activation marker CD69 (Figure 3b) in comparison with control sera of healthy donors (n = 5). A significant reduction was also observed for CD16, whereas the expression of CD16 and NKp46 remained stable rendering them as promising therapeutic targets. These data correlated with the elevated serum level for BAG6 targeting NKp30 and for MICA targeting NKG2D (Figure 2b) and may explain the reduced cytotoxicity of patient peripheral NK cells and healthy NK cells upon incubation with HL sera (Figures 1a and 3a). This *ex vivo* cocultivation experiment allowed a direct estimation of the HL sera-mediated effects on a given polyclonal NK cell population. However, the results were not exactly reflected by the comparison of the NK cell phenotype of healthy donors versus patients with HL (Figure 2a), which is probably due to the pattern of heterogeneous receptor expression among the donors.

**HL-NK cells can be activated in vitro and in vivo with an CD30xCD16A**

Provided that the inhibition of HL-derived pNK cells is not intrinsic but depends on their environment—soluble serum factors—NK cells might be excellent targets for immunotherapy. Therefore, we tested the impact of a tetravalent, bispecific antibody construct CD30xCD16A (TandAb AFM13; Affime Therapeutics, Heidelberg, Germany) on HL-derived pNK cells in *vivo* and *in vitro*. AFM13 is designed for the treatment of Hodgkin lymphoma and is currently tested in a phase I clinical study (ClinicalTrials.gov identifier NCT01221571) to assess the safety in patients with HL. The construct targets CD16A on NK cells and binds simultaneously to the surface receptor CD30, a member of the tumor necrosis factor family. CD30 is overexpressed on malignant HRS cells and contributes to the proinflammatory tumor microenvironment...
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Figure 4 CD30xCD16A activates effector cells from patients with HL in vitro and in vivo. (a) The cytotoxicity of HL-PBMCs (left panel) or purified patient NK cells (right panel) was analyzed by a standard 3 hours europium release assay with L428 target cells in the presence of CD30xCD16A (AFM13) or a control α-CD30 single chain (αCD30sv control). NK cell cytotoxicity against CD30− target cells (293T) in the presence of AFM13 is indicated (AFM/CD30-target). (b) NK cells from healthy donors were incubated with either AFM13 (10 μg/ml), L428 target cells (ratio 5:1), or AFM in combination with L428 for 16 hours, and the expression of CD69 on NK cells was estimated by flow cytometry. (c) CD69 is upregulated on NK cells of patients with HL in response to CD30xCD16A therapy. Flow cytometry: The percentage of CD69+ NK cells in the peripheral blood of CD30xCD16A-treated patients (two different doses) was estimated at the following time points: before therapy (pre), 24 hours after first (d1), second (d8), third (d15), and last (d22) CD30xCD16A infusion. Each curve represents one patient. (d) Ex vivo cytotoxicity assay: NK cells were isolated from the peripheral blood of patients before start of the therapy and 24 hours after the end of first AFM13 infusion to assess the cytotoxicity against the Hodgkin lymphoma cell line L428 (left panel). Left panel: one representative of four patients; right panel: mean of six patients tested. NK cells from two out of six patients tested remained inactive upon treatment. (e) The killing assay was performed in the presence of a MHC I-blocking antibody (α-HLA I), an isotype control (iso), or without antibodies (ctr) (right panel). (f) The binding of AFM13 to peripheral NK cells was estimated before and after (24 hours and 48 hours) the first AFM13 infusion by flow cytometry. The percentage of NK cells loaded with AFM13 is indicated: d, day; HL, Hodgkin lymphoma; MFI, mean fluorescence intensity; MHC I, major histocompatibility complex class I; NK, natural killer; pre, pretreatment.

in this malignancy. Previous studies showed promising clinical results for a bivalent, bispecific monoclonal antibody targeting CD16 and CD30; however, the potency to trigger patient NK cell function and target cell killing has not been analyzed so far.

Cytotoxicity assays revealed that the bispecific protein, but not a αCD30 single chain control protein, significantly enhanced lymphocyte and HL-NK cell-dependent killing (Figure 4a) of L428 target cells. The lysis of CD30− 293T cells remained unaffected. Control experiments proved that NK cell activation was dependent on the presence of both target cells and CD30xCD16A, because neither L428 cells nor CD30xCD16A alone were sufficient to induce the expression of the activation marker CD69 on NK cells (Figure 4b). CD69 belongs to the family of C-type lectin receptors and triggers cytotoxicity and cytokine production of NK cells via pathways activating PLC and vav1. CD69 is induced after activation with IL-2, IL-12, or tumor necrosis factorα and also in response to anti-CD16 monoclonal antibodies; however, the CD30xCD16A failed to upregulate CD69 in the absence of target cells, arguing against nonspecific activity of this antibody construct (Figure 4b).

In the next step, we asked whether an activation of NK cells monitored by CD69 upregulation was detectable in study patients treated with CD30xCD16A (Figure 4c). Patient NK cells were isolated and analyzed before the start of therapy (pre) and 24 hours after each of 4 weekly infusions. After the first treatment, CD69 expression was induced in all patients analyzed. The initial induction showed a decline in response to further treatments; however, the percentage of CD69+ NK cells was still higher in five out of seven patients compared with the pretreatment level. Upregulation of CD69 suggests an enhanced cytotoxic potential of the affected NK cells. In fact, ex vivo killing assays demonstrated these functional consequences: patient NK cells isolated before therapy were inactive, whereas NK cells isolated 24 hours after a single AFM13 infusion were nearly as cytotoxic as NK cells from healthy donors in four out of six patients tested (Figure 4d). To further prove that the killing of the target cells was independent of MHC
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I-missing self-activation, any remaining inhibitory self-antigens were blocked through the addition of an MHC I-specific antibody (Figure 4e). This blocking of MHC I antigens had no impact on target cell killing; therefore, it can also be expected in the autologous patient setting. The binding of AFM13 to the NK cells was detectable on the isolated patient cells by fluorescence activated cell sorter analysis using an antibody with specificity for the construct (Figure 4f). AFM13 was thus present in the cytotoxicity assays with patient NK cells as effector cells. Thus, the high cytotoxic activity of these cells might be due to an antibody-dependent cellular cytotoxicity-like–mediated mechanism. The enhanced killing, mediated through CD30xCD16A, is most likely restricted to CD30+ target cells as shown for NK cells from healthy donors, although this could not be directly tested due to the limitation of patient cells. Taken together, the treatment with an antibody construct targeting CD16A on NK cells can overcome NK cell inhibition in vivo, despite the presence of inhibitory factors in the patient blood. The overall objective of the AFM13 study (ClinicalTrials.gov identifier NCT01221571 ongoing study, 28 patients treated so far), which is a single arm phase I dose escalation trial for patients with relapsed or refractory HL, is to evaluate safety, tolerability, pharmacokinetics, immunogenicity, antitumor activity, the maximum tolerated dose, and the optimal biological dose of AFM13. The study drug was well tolerated and adverse events were generally mild (Table 1). Three out of six patients that were heavily pretreated achieved stable disease (SD), reflecting antitumor activity of AFM13. These patients showed clear upregulation of CD69 on NK cells, and two of them showed efficient ex vivo killing of HL cells after treatment. One patient with SD but no enhanced NK cell-mediated tumor cell killing ex vivo was treated with the least dose of 0.5 mg/kg body weight of all patients tested in this setting. This dose was obviously sufficient to trigger upregulation of CD69 but might not be high enough to preactivate the NK cells above the necessary stimulus threshold with one single infusion. Thus, repeated NK cell stimulation for 4 weeks by AFM13 treatment on a weekly basis could be sufficient to reach this threshold leading to the observed efficacy of the drug in vivo. Moreover, the cytotoxicity of patient NK cells was also enhanced in two patients with SD probably indicating that NK cell activation by AFM13 is a prerequisite for the biological activity of AFM13 but alone not

Table 1 Clinical characteristics and response

| Patient no. | 1 | 2 | 3 | 4 | 5 | 6 |
|-------------|---|---|---|---|---|---|
| Age (years) | 43 | 57 | 69 | 21 | 72 | 26 |
| Sex | F | M | M | M | F | F |
| Stage at first diagnosis | II A | UNK | II BE | IV A | IV A | II A |
| Stage at start of treatment | II AE | III A | III BE | IV B | IV A | III A |
| Histology | Nodular sclerosing | Mixed cellularity | Nodular sclerosing | Nodular sclerosing | Nodular sclerosing | Nodular sclerosing |
| B-symptoms | No | No | No | No | No | No |
| Number of therapies | 6 | 6 | 3 | 3 | 3 | 4 |
| Time since initial diagnosis (month) | 73 | 540 | 3 | 12 | 16 | 250 | 148 |
| Time since last treatment (month) | 3 | 4 | 1 | 3 | 18 | 82 |
| Response to last therapy | Yes | No | No | Yes | No | Yes |
| Dosage (mg/kg bw) | 0.5 | 1.5 | 1.5 | 4.5 | 4.5 | 7 |
| Induction of cytolytic function | −5.0 | 23.0 | 45.0 | −9.0 | 61.0 | 63.0 |
| Fold induction of CD69+ NK cells | 5.8 | 2.0 | 1.1 | 0.92 | 4.5 | 3.3 |
| Response to AFM13, Cycle 1 | SD (0%) | PD | PD | PD | SD (–7%) | SD (–25%) |
| Tumor size (cm²) before AFM13 | 19.3 | 10.2 | 20.8 | 40.7 | 21.4 | 8.8 |
| Tumor size (cm²) after 1st AFM13 cycle | 19.0 | 3.0 | 9.5 | 48.6 | 19.8 | 6.3 |
| Fever | Chills (Grade 1); hypertension (Grade 3) | Chills (Grade 1); | Bronchospasm (Grade 2); | Pain lumbar vertebra worsening (moderate); | Grade 1 | Grade 1 |
| Drug related AE’s | Grade 1 | Grade 2 | Grade 2; | Ague (Grade 2); anemia worsening | Grade 1 | Grade 1 |
| Nausea, vertigo, chills, vomiting, sinus tachycardia, palpitations |

Abbreviations: AE, adverse events; bw, body weight; f, female; h, hours; m, male; NK, natural killer; PD, progressive disease; SD, stable disease. Clinical characteristics of AFM13 treated patients. The induction of the cytolytic function of the NK cells was calculated as difference between % of lysis before and 24 hours after AFM13 infusion.
sufficient for the therapy response. So far, we do not know much about homing of the activated NK cells to the tumor tissue and the influence of the HL tumor microenvironment on NK cell activity in these patients, which should be addressed in further studies. Recent data showed that the HL microenvironment might compromise immune effector functions.33 Taken together, AFM13 has demonstrated encouraging biologic activity and seems to be a new feasible targeted therapy for heavily pretreated patients with HL. These findings strongly support the high demand for therapeutic strategies aimed at restoring NK cell function to avoid tumor evasion from the patient immune system.

DISCUSSION

NK cells represent the main component within the innate immune system involved in recognition and elimination of cancer cells. Here, we report that NK cells from patients with HL have defects in target cell killing and reveal an “anergic” phenotype. The impaired cytotoxicity was attributed to an impaired NKG2D-mediated recognition. The reduced lysis of the target cells was reflected by a downregulation of the NKG2D receptor on peripheral NK cells that correlated to elevated levels of sMICA in patient serum samples. There is an experimental and clinical evidence showing that the NKG2D/NKG2D-L system represents a major part of tumor cell rejection and recent studies demonstrated that there is an impaired immunosurveillance in NKG2D-deficient mice.34 For diseases such as chronic lymphocytic leukemia, a positive correlation of soluble NKG2D ligands to disease stage and bad prognosis was reported.35 Shedding of NKG2D-L from the tumor cell surface is one of the best studied mechanisms that circumvent detection by the innate immune system through reducing activating signals from the cell surface.1,2 Moreover, the released soluble ligands for NKG2D are known to counteract immunosurveillance in two ways: they inhibit NKG2D-cytotoxicity by binding/blocking the receptor and induce its downregulation on the cell surface of T cells and NK cells.1,2 Recently, it was shown that patient leukemia sera with high sNKG2D-L levels reduce NKG2D on NK cells promoting immune evasion,17 whereas other serum factors and their effects on NK cell receptor expression were not addressed in this study. The prominent role for soluble NKG2D ligands in the evasion of NK cell recognition is widely accepted, although not reported for HL so far. Aside from that, our data suggest that not only MICA, the ligand for NKG2D, contributes to NK cell inhibition, but also the factors MIF and BAG6 reveal elevated serum levels in patients versus healthy donors. MIF, known as the MIF, reduces NK cell function also by NKG2D downregulation.25,26 High level of BAG6, the soluble ligand for NKp30, might further constrict NK cell function as incubation of healthy NK cells with HL serum resulted not only in NKG2D downregulation but also in a decreased NKp30-surface detection.

A reduced expression of NCRs on peripheral or tumor-associated NK cells has already been described for tumors such as leukemia, multiple myeloma, and breast cancer.96,37 A critical role for NKp30 for immune surveillance was already proposed but only recently demonstrated in gastrointestinal tumors38 shedding light on its potential role in antitumor immunity. Analysis of larger patient cohorts is needed to uncover potential diagnostic or prognostic value of MIF and BAG6 as novel markers for the treatment outcome in HL. However, a direct correlation of ligand serum level and receptor expression level could not be established in this study, because NK cell phenotype and ligand serum level data were only available from different patients.

The impaired NKG2D-dependent killing of target cells by cytotoxic T cells that were obtained from HL tissue was recently attributed to the release of MICA and ULBP3 in combination with TGFβ from bystander and HRS cells, which caused downregulation of NKG2D on effecter lymphocytes in the microenvironment.33 The data presented here indicate that tissue-derived factors of tumors shape the phenotype and function of peripheral immune cells, which may further inhibit antitumor immune responses.

However, the inhibition of NK cell activity of patient NK cells could be restored with IL-2, a cytokine that is well known to trigger NK cell activity via Ras/MAPK, JAK/Stat, and PI 3-kinase/Akt signaling pathways. Interestingly, IL-2 failed to activate NK cells in the presence of patient serum, suggesting that serum factors act against NK cell stimulation. In line with our data, a reversible suppression of peripheral NK cells, that promote self tolerance, was recently reported for breast cancer.37,39

Killing dysfunction of NK cells from patients with HL could be restored with AFM13, a bispecific antibody construct that targets CD16A on effector cells and binds simultaneously to the surface receptor CD30. AFM13 belongs to a new group of antibody formats that has two binding sites for each antigen as it is a homodi-meric consisting of two polypeptides pairing head-to-tail with each other. This allows tetravalent binding and leads to a comparable avidity as for IgG. In contrast to the previously used CD30-CD16 targeting antibody HRS-3/A9,40 AFM13 consist solely of variable domains and therefore avoids Fc-mediated side effects. A previous study showed promising results against xenotransplanted solid human HL for a bivalent, bispecific monoclonal antibody targeting CD16 and CD30,31 indicating that these targets are well suited for an NK cell activating approach.

CD30, a member of the tumor necrosis factor family is overexpressed on malignant HL cells and associated with constitutive CD30 signaling, known to contribute to the proinflammatory tumor microenvironment.30,41

Enhancement of patients NK cell cytotoxicity against L428 target cells using AFM13 worked in both settings, after direct addition of the construct to the cytotoxicity assay and after treatment of patients. CD16, the low-affinity IgG receptor, is responsible for antibody-dependent cellular cytotoxicity and the engagement of the receptor causes degranulation of NK cells thus triggering cell lysis.20,42 Both in vitro and in vivo studies indicate that the CD16-mediated antibody-dependent cellular cytotoxicity is the predominant mode by which an antitumor response is achieved. In vivo studies using mice with defects in Fc receptor expression demonstrated that the activity of therapeutic antibodies was dependent on the expression of Fc receptors on immune cells.43,44

The general activation of NK cells by the bispecific construct was exclusively observed in the presence of CD30+ target cells and was reflected by the induced expression of the NK cell activation marker CD69. CD69 cross-linking induces the cytotoxic activity and costimulates cytokine production via phosphotyrosine kinases of the Src family. Engagement of CD69 through either antibodies
or—under physiological conditions—with undefined ligands on target cells initiates the activation of PLC and Vav1, both involved in the development of cytotoxicity. The treatment with the bispecific tetravalent antibody CD30xCD16A that induced CD69 expression on the effector cells may thus be regarded a first step in NK cell activation that results in the target cell killing upon direct interaction of NK cells with target cells via the CD30 antibody fragment. Evidence for NK cell activation as a two-stage process by means of first an unspecific activation and second target cell specific activation step was described previously. Other groups also reported that antibody-based approaches with antibody-drug conjugates like Brentuximab vedotin or bispecific antibodies with promising results against B-cell lymphomas, confirming the principle efficiency of novel immunological therapeutics.

Taken together, the impaired function of patient NK cells that are characterized by a decreased NKG2D expression does not reflect a general cytolytic dysfunction. We were able to show that the impaired NK cell function in patients with HL can be restored: NK cells from patients with HL were activated upon treatment with a construct targeting CD16 and CD30 (ex vivo and in vivo), resulting in an improved recognition of an HL-derived target cell line. This data warrant further studies on the feasibility, toxicity, and the clinical response of patients with HL. Furthermore, the NK cell population—in contrast to the T cells—is reconstituted quickly after the end of standard therapies in HL (data not shown). Innovative immune therapies for patients following chemo/radiotherapy that target NK cells rather than T cells are therefore extremely promising.

MATERIALS AND METHODS

Patients. The research was approved by the Ethics Committee of the University Clinic of Cologne and all human participants gave written informed consent. NK cells were isolated with the NK Cell Isolation Kit and AutoMACS (Miltenyi, Bergisch-Gladbach, Germany) as described previously from healthy humans (blood or buffy coats), from patients and AutoMACS (Miltenyi, Bergisch-Gladbach, Germany) as described in informed consent. NK cells were isolated with the NK Cell Isolation Kit University Clinic of Cologne and all human participants gave written consent. NK cell purity of NK cells used (CD3−, CD56+) was evaluated by flow cytometry; Study Identifier NCT01221571 (ClinicalTrials.gov). The HD14–18 trials of the German Hodgkin Study Group, or from patients before/during/after standard chemo/radiotherapy (patients participating in the Deutsche Forschungsgemeinschaft (SFB832, TP19) to E.P.v.S. The University Clinic Cologne for support. U.R. and C.H. are employees of Affimed Therapeutics AG. We thank particularly the blood donors for their generous contribution and Birgit Cathof, Director of the Institut für Transfusionsmedizin, University Clinic Cologne for support. U.R. and C.H. are employees of Affimed Therapeutics AG. This study was supported by a grant from the Deutsche Forschungsgemeinschaft (SF8832, TP19) to E.P.v.S. The other authors declared no conflicts of interest.

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