The novel immunotoxin HM1.24-ETA' induces apoptosis in multiple myeloma cells

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Despite new treatment modalities, the clinical outcome in a substantial number of patients with multiple myeloma (MM) has yet to be improved. Antibody-based targeted therapies for myeloma patients could make use of the HM1.24 antigen (CD317), a surface molecule overexpressed on malignant plasma cells and efficiently internalized. Here, a novel immunotoxin, HM1.24-ETA', is described. HM1.24-ETA' was generated by genetic fusion of a CD317-specific single-chain Fv (scFv) antibody and a truncated variant of Pseudomonas aeruginosa exotoxin A (ETA'). HM1.24-ETA' inhibited growth of interleukin 6 (IL-6)-dependent and -independent myeloma cell lines. Half-maximal growth inhibition was observed at concentrations as low as 0.3 nM. Target cell killing occurred via induction of apoptosis and was unaffected in co-culture experiments with bone marrow stromal cells. HM1.24-ETA' efficiently triggered apoptosis of freshly isolated/cryopreserved cells of patients with plasma cell leukemia and MM and was active in a preclinical severe combined immunodeficiency (SCID) mouse xenograft model. Importantly, HM1.24-ETA' was not cytotoxic against CD317-positive cells from healthy tissue (monocytes, human umbilical vein endothelial cells). These results indicate that CD317 may represent a promising target structure for specific and efficient immunotoxin therapy for patients with plasma cell tumors.
cytotoxic activity in MM cells in vitro as well as in vivo and may represent an interesting approach for the treatment of MM.

MATERIALS AND METHODS

Isolation of primary patient MM cells

Fresh plasma cell leukemia and MM cells were isolated from blood or bone marrow aspirates drawn from patients after obtaining informed consent in accordance with the Declaration of Helsinki. Briefly, citrate- or heparin-anticoagulated blood or bone marrow was layered over a discontinuous gradient consisting of 70 and 62% Percoll (Biochrom, Berlin, Germany), respectively. After centrifugation, mononuclear cells (MNC) were collected from the serum/Percoll interface. CD138+ cells were enriched from MNC using CD138 MicroBeads (Miltenyi, Bergisch Gladbach, Germany) according to the manufacturer’s protocols. The cells were either used directly for the cytotoxicity assays or cryopreserved before further analysis. Experiments reported here were approved by the Ethics Committee of the Christian-Albrechts-University (Kiel, Germany).

Isolation of monocytes from healthy donors

Monocytes were enriched from MNC of healthy donors by magnetic activated cell sorting using ‘Monocyte Isolation Kit II’ (Miltenyi) according to the manufacturer’s protocol.

Cell lines

RPMI-8226, L363, Jurkat, Raji and CEM cells were obtained from the German Collection of Microorganisms and Cell Cultures (DSMZ, Braunschweig, Germany). JK-6 and INA-6 cells were established in our laboratory. All cell lines were cultured in RPMI-1640-Glutamax-I medium (Invitrogen, Karlsruhe, Germany) containing 10% fetal calf serum, penicillin and streptomycin (R10). The cloned sequences were confirmed by Sanger sequencing.

Generation of HM1.24-ETA+

The CD317-scFv was derived by Sfi digestion of pSEC-HM1.24xdsCD16, and ligated to Sfi-digested pet27b(+)ETA carrying a variant of Pseudomonas exotoxin A codon optimized for the expression in Escherichia coli and resulting in pet27b(+)-HM1.24-ETA. The cloned sequences were confirmed by Sanger sequencing.

Expression and purification of immunotoxins

The scFv-ETA fusion proteins were expressed under osmotic stress as previously described. Briefly, arctic express (DE3) RP cells were transformed with the expression construct and overnight cultures in 2xYT medium (Carl Roth GmbH, Karlsruhe, Germany) (supplemented with 20 µg/ml gentamycin, 75 µg/ml streptomycin, 50 µg/ml kanamycin and 1% glucose) were incubated at 37 °C with shaking. The culture was diluted to an OD560 below 0.1 and further incubated at 37 °C in a shaker incubator until OD560 reached 1.0. Osmotic stress was induced as described previously and the incubation temperature was reduced to 28 °C. After 1 h of incubation with agitation, the temperature was further reduced to 13 °C and IPTG (1 mM final concentration) was added. Induced cultures were harvested 16–20 h after induction. The bacterial pellet from 1 l culture was resuspended in 20 ml of extraction buffer (0.5 M sucrose, 0.1 M Tris, 1 mM EDTA, pH 8.0). The suspension was stirred for 30 min at 4 °C and sonicated with 4 bursts of 30 s followed by intervals of 30 s for cooling. The extract was cleared by centrifugation for 40 min at 20 000 g and 4 °C. The scFv-ETA fusion proteins were enriched by affinity chromatography using streptactin agarose matrix (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. Aggregates and/or higher molecular weight contaminants were removed by gel filtration chromatography as described earlier.

SDS–polyacrylamide gel electrophoresis and western blot analysis

SDS–polyacrylamide gel electrophoresis was performed using standard procedures. Proteins were stained with Coomassie brilliant blue R250 (Sigma-Aldrich Chemie GmbH, Munich, Germany) or transferred to Immuno-Blot PVDF membrane (Bio-Rad, Hercules, CA, USA). Recombinant immunotoxins were detected with mouse anti-penta-His antibody (Qiagen). Cleavage of poly(ADP-ribose) polymerase (PARP) was analyzed using whole-cell protein extracts prepared from 1 × 10⁶ cells as previously published. Full-length PARP and its specific cleaved product were detected using mouse anti-human PARP antibody (Cell Signaling Technology, Danvers, MA, USA). Horseradish peroxidase-conjugated goat anti-mouse antibodies (Dianova, Hamburg, Germany) were used as secondary antibodies. Detection of bound antibody was performed with SuperSignal West Dura Extended Duration Substrate (Pierce, Rockford, IL, USA).

Capillary electrophoresis

Quantitative analysis of purified immunotoxins was performed using capillary electrophoresis on an Experion system (Bio-Rad) according to the manufacturer’s protocol.

Flow cytometric analyses

For immunofluorescence staining, 3 × 10⁵ cells were washed in phosphate-buffered saline (PBS) supplemented with 1% bovine serum albumin (Sigma-Aldrich, Munich, Germany) and 0.1% sodium-azide (PBA buffer). To analyze immunotoxin binding, cells were incubated with HM1.24-ETA+ or control proteins at indicated concentrations for 30 min on ice. After washing twice with 500 µl PBA buffer, cells were stained with Alexa-Fluor-488 coupled mouse anti-penta-His antibody (Qiagen) or rabbit anti-exotoxin A polyclonal antibodies and fluorescein isothiocyanate (FITC)-labeled Fab1 fragments of polyclonal goat anti-rabbit antibodies (Sigma-Aldrich).

The purity of CD138+–enriched cells was analyzed by staining 3 × 10⁵ cells with FITC-conjugated CD38 and PE-conjugated CD138 antibodies (Beckman Coulter, Fullerton, CA, USA) according to the manufacturer’s protocol using appropriate controls. CD317 expression on CD138+ cells was determined by staining 3 × 10⁵ cells with a humanized HM1.24-IgG antibody (kindly provided by Chugai Pharmaceuticals Inc., Tokyo, Japan) or irrelevant IgG1 control, and FITC-conjugated goat anti-human IgG Fab1 fragments (Beckman Coulter) as secondary antibody. Cells were analyzed on a flow cytometer (FCS300, Beckman Coulter, Brea, CA, USA).

The surface expression level of CD317 was quantified by indirect immunofluorescence analysis using the QIFKIT (Dako, Glostrup, Denmark) and mouse CD317 monoclonal antibody (clone 26F8, E Bioscience, San Diego, CA, USA) as primary antibody according to the manufacturer’s protocol.

Measurement of cytotoxic effects of immunotoxin

For evaluation of cytotoxic effects of the immunotoxin, cells were seeded at 2 × 10⁶ cells per 200 µl in 96-well plates. HM1.24-ETA+ or a control toxin was added at the indicated concentrations. After 3 days, vital cell mass was measured using colorimetric tetrazolium (MTT/MTS)-based assays (Cell Proliferation Kit I; Roche, Mannheim, Germany; Promega, Madison, WI, USA). For the detection of early stages of apoptosis and cell death, cells were seeded at 2 × 10⁵ cells per ml in 24-well plates with increasing immunotoxin concentrations. For analyzing the kinetics of apoptosis induction, the immunotoxin was used at 100 ng/ml. Cells were stained with FITC-conjugated annexin V and 7-aminoactinomycin D (7-AAD; Beckman Coulter, Fullerton, CA, USA) according to the manufacturer’s protocol, and subsequently analyzed by flow cytometry. For blocking experiments, a 50-fold molar excess of parental antibody was added 30 min before adding immunotoxin.

MM/bone marrow stromal cells (BMSC) co-culture

BMSC were obtained from MNC isolated by ficoll density centrifugation of patient-derived bone marrow aspirates and subsequently cultured in R10+. For co-culture experiments, stromal cells were trypsinized and transferred to 96-well plates (0.5 × 10⁵ cells/well), and MM cells (INA-6, 2 × 10⁵ per well) were added the following day. Cells were treated with HM1.24-ETA+ at the concentrations indicated. For blocking experiments, the parental HM1.24-IgG1 antibody or a control antibody were added in molar excess. After 3 days of culture, stromal cell viability was analyzed by a colorimetric MTS-based assay (Promega). DNA synthesis of INA-6 cells under co-culture conditions was measured by [3H]-thymidine uptake. In brief, cells were pulsed with [3H]-thymidine (Tdr, 1 µCi/well; specific activity, 5.0 Ci/mmol; Hartmann Analytic, Braunschweig, Germany).
for 6 h. Subsequently, DNA was transferred onto glassfiber filters and counted in a β-scintillation counter (Perkin Elmer, Rodgau, Germany).

INA-6 xenograft tumor model
Seven-week-old female severe combined immunodeficiency (SCID) beige mice (Charles River, Sulzfeld, Germany) were maintained under pathogen-free conditions and injected intraperitoneally with $2.5 \times 10^5$ INA-6.Tu1 cells (these INA-6 cells have already been passaged in SCID mice) in 1 ml PBS. Starting on day 2 until day 12 every other day, and on days 20, 27 and 34, mice were treated intraperitoneally with HM1.24-ETA' (total of 9 injections, 15 μg toxin each) or PBS as vehicle control. Tumor engraftment was monitored twice per week. Mice were killed when suffering from tumor burden, ascites, when paraplegia was observed or at the end of the experiment (day 125). All animal experiments and care were performed in accordance with legal regulations and the guidelines provided by the Federation of European Laboratory Animal Science Associations (FELASA) as well as approval of institutional authorities.

Statistical analyses and data processing
Graphical and statistical analyses were performed using GraphPad Prism 5.0 software (GraphPad Software, La Jolla, CA, USA). Curve fits were calculated assuming a sigmoidal dose response with variable slope. P-values were calculated using Student’s t-test, one- or two-way analysis of variance and the null hypothesis was rejected when $P < 0.05$. Survival curves were analyzed using the log rank test.

Homology modeling
A homology model was calculated for HM1.24-ETA’ by modeling the scFv (template: 1H8N) and ETA’ (template: 1IKP) separately using the YASARA Structure software (YASARA Biosciences, Graz, Austria). The individual domains were manually fused using the YASARA Structure software. Ribbon drawings were performed using Discovery Studio 2.0 Visualize software (Accelrys Inc., San Diego, CA, USA).

RESULTS
Generation and antigen-specific binding of HM1.24-ETA’
HM1.24-ETA’ was generated by fusing a CD317-specific scFv34 to a sequence optimized version of a truncated Pseudomonas exotoxin A lacking the receptor binding domain I (ETA’, Figure 1a). As controls, immunotoxins using a CD64-specific

Figure 1. Design, purification and antigen-specific binding of HM1.24-ETA’. (a, left) Design of the recombinant immunotoxin HM1.24-ETA’. ETA’, truncated ETA fragment consisting of domains II and III of Pseudomonas exotoxin A; KDEL, endoplasmic reticulum retention motif; S, pelB secretion leader fused to STREP-II-6xHis-tag; Sfl, restriction site; T7, T7 promoter; VL, VH, variable regions of the light and heavy chain. (a, right) Calculated homology model of HM1.24-ETA’. (b) HM1.24-ETA’ purified by affinity chromatography was analyzed by capillary electrophoresis under reducing conditions. SP, system peaks. (c) Higher molecular mass contaminants were removed by gelfiltration. Graph shows reanalysis after final purification step. (d) Dose-dependent binding of HM1.24-ETA’ or control immunotoxin CD64-ETA’ was analyzed on L363 cells. Mean values ± s.e.m. from three experiments are shown. (e) Competition binding experiments were performed to demonstrate specificity of HM1.24-ETA’ binding. Cells were stained with HM1.24-ETA’ and subjected to flow cytometry (black). Binding of HM1.24-ETA’ was inhibited with a molar excess of parental antibody (gray). Unstained cells (white). Data from one representative experiment out of three are shown.
The immunotoxins were expressed in E. coli and purified from periplasmatic extracts by affinity chromatography and gel filtration (Figures 1b and c). HM1.24-ETA specifically and dose dependently bound to CD317-positive L363 MM cells (Figure 1d). Binding of the immunotoxin was completely inhibited by preincubation of the cells with a molar excess of parental antibody, demonstrating that antigen specificity was not altered by fusing the toxin component (Figure 1e).

Specific inhibition of myeloma cell line growth
To test whether the recombinant immunotoxin was biologically active, proliferation assays were performed. Proliferation of CD317-expressing myeloma cell lines L363, RPMI-8226, JK-6 and the IL-6-dependent plasmacytoma cell line INA-6 was significantly inhibited by HM1.24-ETA. CD317-negative CEM (T-ALL) and Raji (Burkitt’s lymphoma) cells were not compromised (Figure 2a). A control immunotoxin did not show this inhibitory effect (Figure 2b). Half-maximal inhibition was achieved at concentrations of 30–50 ng/ml corresponding to ~0.3–0.5 nM.

HM1.24-ETA is not cytotoxic against CD317-positive nonmalignant monocytes and HUVECs
Besides its expression on MM cells (Supplementary Figure 1), CD317 expression has also been detected on normal/nonmalignant tissue. For example, nonmalignant subepithelial plasma cells expressed CD317 as evidenced by immunohistochemistry of tissue sections (Figure 3a). In line with recently published data by Erikson et al.31 who reported ‘high level’ expression of CD317 on monocytes, flow cytometric analysis revealed significant staining of monocytes (Figure 3b). In addition, using immunohistochemistry staining and RNA profiling, Erikson et al.31 reported CD317 expression on different normal tissues, including endothelial cells from blood vessels. Importantly, although CD317 expression on different normal tissues, including nonmalignant cell types (Figures 3b and c). This unexpected finding may be related to a significantly lower antigen density expressed on healthy tissue vs MM cell lines (Supplementary Figure 1), differences in the kinetics of internalization, the cellular activation state or, as suggested in recent reports, because of different functions of CD317 in selected cell types.39,40 In line with these findings, induction of low levels of surface-expressed CD317 on Jurkat T cell leukemia cells (CD317-inducible, CD7-positive) by interferon-γ (IFN-γ) (Supplementary Figure 2) did not result in HM1.24-ETA-triggered cell death (Supplementary Figure 2). Importantly, Jurkat cells were significantly inhibited by CD7-ETA, a similarly constructed immunotoxin directed against CD7, demonstrating that lack of response to HM1.24-ETA was not because of a general resistance to ETA-based conjugates (Supplementary Figure 2).

As the kinetics of internalization may impact the amount of delivered payload, the properties of monocytes and endothelial cells to internalize HM1.24 in comparison with plasma cell leukemia cells, a time-dependent clearance of a HM1.24-specific antibody bound to the cell surface, was analyzed by flow cytometry (Supplementary Figure 3A). After incubation at 37°C, the amount of surface-bound CD317 antibody on monocytes was reduced to 60% compared with monocytes incubated at 4°C (Supplementary Figures 3A and B). Therefore, monocytes displayed an internalization rate nearly comparable to the plasma cell leukemia cell line L363 (Supplementary Figures 3A and B). On endothelial cells (HUVEC), no differences in surface-bound antibody were detected after 4 h (Supplementary Figures 3A and B). This reduced internalization of antibody may explain the insusceptibility of CD317-positive endothelial cells (HUVECs) for the induction of apoptosis by the immunotoxin HM1.24-ETA.

For CD317-positive monocytes displaying comparable internalization of CD317 without being affected by the delivery of HM1.24-ETA immunotoxin, other cellular characteristics may account for the ‘resistance’ of the cells. As the catalytic ETA domain abrogates protein synthesis, resting cells with low metabolical activity, for example isolated monocytes, may be less affected by this mode of action in contrast to malignant plasma cells that are highly active in protein synthesis.

In line with this hypothesis, G28–5 scFv-PE40, an immunotoxin targeting CD40, efficiently killed CD40-positive malignant B cells, whereas monocytes and endothelial cells expressing CD40 were resistant to the immunotoxin.41 Activation of these cell populations by IFN-γ (monocytes) or a combination of IFN-γ and tumor necrosis factor-α (endothelial cells) sensitized these cells for the action of the immunotoxin. To analyze the influence of IFN-γ-mediated activation of monocytes on their susceptibility to HM1.24-ETA-induced apoptosis, isolated monocytes were treated with the immunotoxin for 3 days with or without IFN-γ treatment. Cell viability was reduced in a dose-dependent manner by HM1.24-ETA only in the presence of IFN-γ (Supplementary Figure 3C). Thus, cytokine-mediated activation of monocytes sensitized these cells for HM1.24-ETA-induced cell death, whereas
nonactivated monocytes were almost unaffected. Interestingly, the surface level of CD317 on monocytes was not altered by IFN-\(\gamma\) treatment (Supplementary Figure 3D). Therefore, IFN-\(\gamma\)-induced activation rather than altered surface expression levels of CD317 may account for the enhanced potency of the immunotoxin HM1.24-ETA on IFN-\(\gamma\)-activated monocytes.

Together, these data suggest that a certain extent of surface expression on healthy tissue may not necessarily result in toxic side effects against nonmalignant tissue and the metabolic state/activation status of a target cell may govern its susceptibility to immunotoxin action.

Antigen-specific induction of apoptosis by HM1.24-ETA

To analyze whether the inhibition of MM cell growth induced by HM1.24-ETA was antigen-specific and occurred via induction
of apoptosis, JK-6 cells were treated with a single dose of HM1.24-ETA' (100 ng/ml) for 24 h and analyzed by annexin V/7-AAD staining and flow cytometry. Preincubation of the cells with a molar excess of the parental antibody HM1.24-IgG1, but not with an irrelevant IgG1, blocked induction of apoptosis. Detection of cleaved PARP confirmed antigen-specific induction of apoptosis by HM1.24-ETA'. Data show one representative experiment out of three performed.

Dose- and time-dependent induction of apoptosis by HM1.24-ETA'

To analyze dose-dependent induction of apoptosis, L363, JK-6, INA-6 and Raji were treated with varying toxin concentrations (Figure 5a). A single dose of 100 ng/ml of HM1.24-ETA' induced apoptosis in almost all JK-6 and L363 cells within 24 h. Half-maximal induction of apoptosis occurred at concentrations as low as 20 and 30 ng/ml, respectively. In contrast, INA-6 cells were less sensitive to immunotoxin treatment, although INA-6 cells expressed higher levels of CD317 than JK-6 or L363 (Supplementary Figure 1), suggesting that additional cellular factors govern extent of immunotoxin-induced cell death.

To investigate the kinetics of apoptosis induction by HM1.24-ETA', the cell lines were incubated with a single dose of toxin that was found to be sufficient to induce apoptosis within 24 h of treatment. Cells were analyzed by annexin V/7-AAD staining at different time points (Figure 5b). Half-maximal induction of apoptosis was detected after 6.5 h. Approximately 80% of JK-6 and L363 cells were found to be annexin V/7-AAD positive already after 8 h, indicating that toxin uptake and processing was fast and efficient. Raji cells that served as controls were unaffected by immunotoxin treatment (Figures 5a and b).

Stromal cells do not protect plasma cells from HM1.24-ETA'-induced apoptosis

BMSC have been reported to protect myeloma cells from the cytotoxic effects of therapeutic drugs. To address whether bone marrow stroma mediates resistance to HM1.24-ETA', INA-6 cells were co-cultured with stromal cells obtained from patient bone marrow aspirates. Addition of a single dose of HM1.24-ETA' (30 ng/ml) completely inhibited INA-6 proliferation in the presence of BMSC (Figure 6a), whereas the unconjugated HM1.24-IgG1 antibody had no inhibitory effect on myeloma cell growth. In contrast, the viability of BMSC was not affected, even at higher HM1.24-ETA' concentrations up to 300 ng/ml (Figure 6b), indicating that HM1.24-ETA' was directly active against myeloma cells and not via affecting stromal cell viability. HM1.24-ETA'-mediated killing during co-culture was antigen specific, as indicated by blocking experiments using the parental antibody. Together, these data indicate that bone marrow stroma has no protective activity against HM1.24-ETA'-mediated induction of apoptosis.

HM1.24-ETA' induces apoptosis in freshly isolated/cryopreserved plasma cell leukemia and MM cells

To investigate the cytotoxic activity of HM1.24-ETA' on freshly isolated tumor cells, malignant plasma cells were enriched from MNC obtained from bone marrow aspirates, pleural effusion or peripheral blood samples of seven patients with MM or plasma cell leukemia (Supplementary Table 1). Purity of CD138+...
selected plasma cells from patients was ~95% (Figure 7a).

Importantly, isolated CD138⁺ cells from all patients homogeneously expressed CD317 on the cell surface (Figure 7a and Supplementary Figure 1), although at lower levels compared with MM cell lines (Supplementary Figure 1).

HM1.24-ETA’ induced apoptosis in primary malignant plasma cells in a dose-dependent manner. Approximately 75% of plasma cells (patient no. 1) treated with a single dose of 1 μg/ml for 24 h were found to be in early apoptotic state or dead as measured by annexin V/7-AAD staining (half-maximal effective concentration: 250 ng/ml, Figure 7b). The percentage of apoptotic cells after treatment with 100 ng/ml of immunotoxin increased from 25% after 24 h to 47% after 72 h, showing time-dependent induction of apoptosis at low immunotoxin concentrations (Figure 7b).

To further confirm the potency of HM1.24-ETA’ on patient-derived tumor cells, primary malignant plasma cells of six additional patients (2–7) were subjected to immunotoxin treatment (Figures 7c and d). HM1.24-ETA’ induced antigen-dependent apoptosis in all samples investigated (Figures 7c and d). The percentage of apoptotic cells obtained after 24 h of treatment with a suboptimal single dose of HM1.24-ETA’ (100 ng/ml) was variable between the different patients, indicating varying sensitivity or kinetics of immunotoxin uptake. Blocking experiments further confirmed the antigen specificity of the immunotoxin (Figures 7c and d). When mean values were calculated from all experiments, significant killing was observed between the nontreated and HM1.24-ETA’-treated group (Figure 7d, left panel), or between the specifically blocked and irrelevantly blocked sample group (Figure 7d, right panel). Thus, the induction of antigen-dependent killing was demonstrated in a panel of samples (Figures 7c and d) that were derived from patients differing in disease stage and subtype, as well as cytogenetic abnormalities (Supplementary Table 1).

HM1.24-ETA’ inhibits INA-6 tumors in SCID mice

To evaluate whether HM1.24-ETA’ was also active in vivo, 2.5 × 10⁷ INA-6 cells that demonstrated the lowest sensitivity to HM1.24-ETA’ in vitro were injected intraperitoneally into 7-week-old SCID beige mice. Starting on day 2 after tumor cell injection, mice were treated with 9 doses of HM1.24-ETA’ (total dose: 135 mg) or vehicle (PBS). HM1.24-ETA’ prevented plasmacytoma engraftment after the observation time of 125 days in 8/10 mice treated with HM1.24-ETA’. The difference in survival of HM1.24-ETA’-treated vs control group reached statistical significance (Figure 8; P<0.02).

DISCUSSION

The novel single-chain immunotoxin HM1.24-ETA’ targeting the CD317 antigen, a surface receptor overexpressed on malignant plasma cells, was designed to advance therapeutic options for patients with MM, and potent antmyeloma activity was demonstrated in vitro and in a preclinical in vivo model.

HM1.24-ETA’ specifically triggered apoptosis in malignant plasma cells at low nM concentrations, comparable to potent ETA-based immunotoxins, such as BL22 (CAT-3888).
Interestingly, induction of apoptosis in cell lines was detected within a few hours after treatment, indicating that uptake and intracellular processing of the toxin occurred efficiently and rapidly. This may be because of CD317 involvement in cell signaling as well as in endocytosis. Earlier studies suggested the transmembrane and GPI-linked protein CD317 is a highly internalized antigen that is not dramatically down-modulated after internalization. Even in the absence of ligand or antibody binding, CD317 is internalized from lipid rafts in a clathrin-dependent manner. It is mainly transported to recycling

Figure 7. HM1.24-ETA induces apoptosis in malignant plasma cells isolated from patients with multiple myeloma or plasma cell leukemia. (a) Mononuclear cells from blood of a patient (p#1) with plasma cell leukemia were analyzed for expression of CD317 on CD138+/CD38+ malignant plasma cells (black) and isotype control (gray). (b) CD138 magnetic activated cell sorting (MACS)-sorted primary tumor cells were treated with HM1.24-ETA. Immunotoxin-induced apoptosis was measured by annexin V/7-AAD staining. Mean values of single experiments set up in duplicates are shown. (c) Primary tumor cells of two patients were incubated with a single dose of 100 ng/ml HM1.24-ETA for 24 h. Annexin V/7-AAD staining indicated that HM1.24-ETA induced apoptosis. The cytotoxic effect mediated by HM1.24-ETA is blocked by preincubation of the tumor cells with a molar excess of parental antibody HM1.24-IgG1 but not by an irrelevant antibody. (d) Comparison of untreated sample group with treated sample group. (d, right) The corresponding data with HM1.24-IgG1 blocking antibody or irrelevant blocking antibody. HM1.24-ETA = 100 ng/ml, 24 h treatment; ●, untreated; ■, HM1.24-ETA treatment; □, HM1.24-ETA + irrelevant IgG1; ○, HM1.24-ETA + HM1.24-IgG1.
endosomes and/or the trans-Golgi network without being delivered to late endosomes or lysosomes for degradation. CD317 is located at the cell surface, but also exists in intracellular stores and shuttles between the trans-Golgi network and the cell surface.\textsuperscript{46} Thus, an efficient 'transport cycle' for the immunotoxin of the receptor may account for the observed potency.\textsuperscript{26,39} Besides these highly dynamic characteristics of CD317 in cells active in secretion or internalization, recent reports suggested that in polarized endothelial and epithelial cells, CD317 exhibits a more structural function. CD317 is engaged in adhesion of myeloid cells to CD317-expressing endothelial cells.\textsuperscript{26} In polarized epithelial cells a complex of RICH2 and CD317 is involved in the organization of the subapical actin cytoskeleton.\textsuperscript{41} In these cells no intracellular depots of CD317 were detectable, whereas CD317 was only internalized inefficiently. These reports are consistent with our surface retention studies of a CD317-specific antibody (Supplementary Figure 3). Clathrin-mediated endocytosis depends on the interaction of the AP2 adaptor complex and the corresponding binding motive within the cytoplasmic domain of CD317.\textsuperscript{47} This motive shared by the AP2 adaptor complex and the corresponding binding motive within the respective function in a given cellular background, the level of tumor targeting.\textsuperscript{31} The data presented here with human monococytes and HUVEC clearly demonstrate that surface expression of CD317 on healthy tissue does not inevitably result in killing of CD317-positive cells. Antigen surface expression density, the respective function in a given cellular background, the level of internalization and, as shown for monocyes, the activation status may govern susceptibility to HM1.24-ETA and other immunotoxins.\textsuperscript{39-41} A potent Fc-engineered CD317-specific IgG1 antibody was successfully tested in cynomolgus monkeys, demonstrating a significant and specific reduction of normal plasma cells in blood as well as bone marrow. Importantly, treatment was well tolerated and immunohistochemistry studies on normal tissues showed a similar staining pattern in humans and cynomolgus monkey.\textsuperscript{49} Furthermore, a native CD317 antibody has been tested in patients without adverse events, underlining that therapeutic administration of CD317-targeting agents is feasible.\textsuperscript{57} Interestingly, elevated surface expression levels of CD317 have also been reported on the tumor vasculature of B-cell lymphomas,\textsuperscript{55} on primary lung cancer cells\textsuperscript{57} and glioblastoma cells.\textsuperscript{28} Furthermore, gene expression profiling revealed elevated CD317 mRNA levels in endometrial cancer.\textsuperscript{59} These data indicate that CD317 may also represent a target structure for therapy in indications other than MM and that, in special situations, it may be possible to target both tumor and tumor vasculature.\textsuperscript{28} The HM1.24-ETA\textsuperscript{0} immunotoxin reported here represents an interesting immunotherapeutic option in patients with plasma cell disorders. Even in patients with a high tumor load, a suppressed immune system or limited hematopoiesis, including situations early after stem cell transplantation, such an approach can be used. In these settings, the action of immunotoxins in contrast to unconjugated or bispecific antibodies may be less dependent on functionally active immune effector cells. Furthermore, efficient responses to immunotoxin treatment most likely will not be influenced by FcR polymorphisms, as reported for unconjugated antibodies.\textsuperscript{8} Thus, the CD317 antigen may represent an interesting receptor for antigen-specific delivery of cytotoxic compounds, suggesting that HM1.24-ETA\textsuperscript{0} could serve as a potent immunonjugate for treatment of plasma cell disorders and possibly other CD317-positive tumors.\textsuperscript{20}
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