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UNC-45A Is a Nonmuscle Myosin IIA Chaperone Required for NK Cell Cytotoxicity via Control of Lytic Granule Secretion

Yoshie Iizuka,* Frank Cichocki,† Andrew Sieben,* Fabio Sforza,‡ Razaul Karim,§ Kathleen Coughlin,* Rachel Isaksson Vogel,* Riccardo Gavioli,‡ Valerie McCullar,† Todd Lenvik,† Michael Lee,§ Jeffrey Miller,† and Martina Bazzaro*

NK cell’s killing is a tightly regulated process under the control of specific cytoskeletal proteins. This includes Wiskott–Aldrich syndrome protein, Wiskott–Aldrich syndrome protein–interacting protein, cofilin, Munc13-4, and nonmuscle myosin IIA (NMIIA). These proteins play a key role in controlling NK-mediated cytotoxicity either via regulating the attachment of lytic granules to the actin-based skeleton or via promoting the cytoskeletal reorganization that is requisite for lytic granule release. UNC-45A is a highly conserved member of the UNC-45/CRO1/She4p family of proteins that act as chaperones for both conventional and nonconventional myosins. Although we and others have shown that in lower organisms and in mammalian cells NMIIA-associated functions, such as cytokinesis, cell motility, and organelle trafficking, are dependent upon the presence of UNC-45A, its role in NK-mediated functions is largely unknown. In this article, we describe UNC-45A as a key regulator of NK-mediated cell toxicity. Specifically we show that, in human NK cells, UNC-45A localizes at the NK cell immunological synapse of activated NK cells and is part of the multiprotein complex formed during NK cell activation. Furthermore, we show that UNC-45A is disposable for NK cell immunological synapse formation and lytic granules reorientation but crucial for lytic granule exocytosis. Lastly, loss of UNC-45A leads to reduced NMIIA binding to actin, suggesting that UNC-45A is a crucial component in regulating human NK cell cytotoxicity via promoting the formation of actomyosin complexes. The Journal of Immunology, 2015, 195:4760–4770.

Natural killer cells are critical for immune responses against viral infections and cancer (1). NK cell–mediated cytotoxicity begins with the formation of an active NK cell immunological synapse (NKIS) between the effector and the target cell and culminates with the release of lytic granule content for target cell killing (2–4). Unlike CTLs, which are present in small precursor frequencies and must undergo differentiation and expansion before target cell killing, NK cells are ready-to-kill cells armed with a constitutive pool of lytic granules. Thus, NK cell killing is a tightly regulated process that is particularly sensitive to cytoskeletal dynamics (4–8). A number of cytoskeletal-associated proteins including Wiskott–Aldrich syndrome protein (WASP), WASP-interacting protein, cofilin, Munc13-4, and nonmuscle myosin IIA (NMIIA) are involved in the stepwise cytoskeletal reorganization that is requisite for lytic granule release (3, 6, 9, 10). Mutations in the genes coding for these proteins severely compromise NK cell–mediated cytotoxicity and result in severe immunodeficiency (11, 12).

In lower organisms and in mammalian cells, NMIIA-assisted functions including cytokinesis, cell motility, and organelle trafficking are dependent upon the presence of its cochaperone UNC-45A (13–17). UNC-45A is a highly conserved member of the UCS (UNC-45/Cro1/She4p) protein family, which plays a crucial role in chaperone motor protein assembly (18, 19). Our group and others have recently shown that UNC-45A interacts with and affects the folding and stability of myosins including NMIIA via direct binding with myosin heads (19). This allows for its efficient binding to actin. Although high UNC-45A RNA expression levels have been reported in NK cells (20), the cellular localization and functional relevance of UNC-45 protein in NK cells has yet to be described. Given the functional dependency of NK cells on cytoskeletal dynamics in general and for NMIIA function in particular, we sought to investigate the role of the NMIIA cochaperone UNC-45A during NK cell–mediated cytotoxicity.

In this study, we show that UNC-45A protein is abundantly expressed in human NK cells, where it interacts with lytic granules and binds to NMIIA. Furthermore, we show that small hairpin RNA (shRNA)-mediated silencing of UNC-45A severely affects NK cell cytotoxicity. Lastly, our results show that impairment of NK cell cytotoxicity in UNC-45A knockdown cells is not due to an inability of these cells to form active immunological synapses, but to a deficiency in lytic granule secretion via a mechanism involving alteration of actomyosin contractility.
Materials and Methods

Isolation of cells from peripheral blood

Human subjects were used as per the Institutional Review Board approval and with their consent. Adult peripheral blood was obtained from healthy donors. PBMCs were isolated by centrifugation using a Ficoll–Paque Premium. NK cells were negatively selected from PBMCs using Clini MACS CD3 Reagent (273-01, Miltenyi Biotech), and cultured at 37°C with 5% CO₂ and a 5:1 (v/v) mix of DMEM (11995; Life Technologies) and F-12 (11765; Life Technologies) containing 10% human heat-inactivated serum (HP1022; Valley Biomedical), 25 μM 2-ME (21985-023; Invitrogen), 50 μM ethanolamine (E0135; Sigma), 1.7 μg/l sodium selenite (214485; Sigma), 20 mg/l ascorbic acid, and 1000 U/ml human IL-2 (200-02; Peprotech). Monocytes were isolated by positive magnetic selection using CD14 microbeads (Miltenyi Biotech) according to the manufacturer's instructions. B, T, and NK cells from the monocyte-depleted population were all isolated by negative magnetic bead selection according to the manufacturer's instructions.

Modulation of UNC-45A expression levels in human NK cells

For UNC-45A silencing in YT, NKL, and peripheral blood NK (p-NK) cells, scramble and UNC-45A shRNAs (5'-CGTGAAGATTACGACAAAGCA-3' #1 and 5'-CCACCTCAAGCCTGAGATTACA-3' #2) were inserted into the GFP-containing lentiviral YT cells or GFP-containing lent-PEP empty vector-expressing NKL cells. Plasmids were cotransfected with the VSVG envelope, and the delta-8.9 plasmids and infection were performed as we previously described (21). Forty-eight hours postinfection, GFP⁺ CD³⁺ CD56⁺ p-NK cells were sorted and collected using a FACSaria II (BD Biosciences). For UNC-45A overexpression in NKL cells, lent-PEP-UNC-45A was cotransfected with the VSVG envelope, and the delta-8.9 plasmids and infection were performed as we previously described (21).

Quantitative RT-PCR

Total RNA was purified using the Qiagen RNNeasy Mini kit (Qiagen). cDNA was generated by SuperScript III Transcriptase (Invitrogen) and used as template for quantitative real-time PCR using SYBR Green Master Mix (Roche) and the Applied Biosystems 7300 Real-Time PCR System. The primers used for quantitative real-time PCR were 5'-TCGGAGGCAAGAAGGGTCAC-3' and 5'-ACCCTTATCACTTGCTTCAGG-3' for UNC-45A and 5'-TGCAACCAACACTGTATTACG-3' and 5'-GGCATTGGACGTTCATGAG-3' for GAPDH.

NK cell lines

YT, NKL, 721.221, and K562 cells were cultured at 37°C with 5% CO₂ in RPMI 1640 with 10% FBS. A total of 100 U/ml human IL-2 was added for YT and NKL, 92-02 (American Type Culture Collection) were cultured in RPMI 1640 with 12.5% horse serum, 12.5% FBS, 0.1 mM 2-ME (200-02; Peprotech). Monocytes were isolated by positive magnetic selection using CD14 microbeads (Miltenyi Biotech) according to the manufacturer’s instructions. B, T, and NK cells from the monocyte-depleted population were all isolated by negative magnetic bead selection according to the manufacturer’s instructions.

Isolation of lytic granules from NK cells.

Postnuclear lysate (P NL) and crude lysosomal fraction (CLF) were obtained by centrifugation at 100,000 x g for 90 min at 4°C and pelleting of lytic granules in PNL at 180,000 x g as previously described (22). Lysosomal fractions (a-f) were obtained by Optiprep density gradient centrifugation using the Lysosomal Isolation Kit (Sigma–Aldrich) according to the manufacturer’s instructions. Fraction of 0.2 ml was harvested for subsequent analysis.

Formation and measurement of effector–target conjugates.

Scramble and UNC-45A knockdown YT cells (GFP⁺) were mixed with an equal volume of CellTracker Orange–loaded 721.221 target cells in a 5:1 polystyrene round-bottom tube at a 2:1 ratio. Cells were centrifuged at 20 × g for 1 min, incubated in 5% CO₂ at 37°C for 15 min without agitation. At the end of incubation time, the cells were vortexed for 3 s at maximum vortex speed and immediately fixed with 4% paraformaldehyde, staining with mouse anti-perforin Ab, and followed by Texas red–conjugated anti-mouse IgG and Ab and analyzed via fluorescence microscopy.

Measurement of cell viability in UNC-45A knockdowns NK cells

For measurement of NK cell viability after UNC-45A knockdown, stable YT cells expressing either scramble or UNC-45A knockdown (KD1# and KD2#) were analyzed by trypan blue to exclude dead cells.

[35S]Cr-release assay

NK cell cytotoxic activity was tested in standard 5-h [35S]Cr-release assays as previously described (2). In brief, 721.221 and K562 target cells were labeled with 0.1 μCi/10⁶ cells of Na₂[35S]CrO₄ for 90 min at 37°C and incubated with effector cells at the indicated ratio. Per each condition, the percentage specific release was calculated with ice-cold 2% paraformaldehyde and [35S]Cr-labeled target cells. Cpm were assessed by complete lysis of target cells by using 1% Triton X-100.

Western blotting and coimmunoprecipitation.

Western blot analysis, cell lysates (10–20 μg) and lysosomal fractions (50 μl) from each sample were separated by SDS-PAGE, transferred to polyvinylidene difluoride membranes, and subjected to Western blot analysis using the indicated Abs according to the manufacturer’s recommendations. For coimmunoprecipitation experiments, human NK cells were lysed into lysis buffer (50 mM Tris, pH 7.4, 150 mM NaCl, 1% Nonidet P-40, 1% protease inhibitor mixture, 1% phosphatase inhibitor mixture). The lysates were preclarified and precipitated with primary Ab and protein A/G beads. Pellets were washed three times with ice-cold lysis buffer and analyzed by SDS-PAGE. Immunofluorescence microscopy.

For analysis of UNC-45A subcellular localization, YT NK cells overexpressing UNC-45A-GFP fusion protein or primary NK cells were incubated with 721.221 or K562 target cells, respectively, at a 2:1 ratio in suspension for 30 min and adhered to poly-L-lysine–coated glass slides (Polyprep; Sigma-Aldrich) for 15 min. After fixation and permeabilization, cells were stained with the anti-myosin IA and anti-UNC45A antibodies (primary cells only) Abs, followed by FITC-conjugated donkey anti-mouse IgG or by Texas red–conjugated goat anti-rabbit IgG for UNC-45A and NMIIA, respectively, and analyzed via fluorescence microscopy. For measurement of lytic granules polarization, scramble and UNC-45A knockdown YT cells (GFP⁺) were incubated with or without 721.221 target cells at a 2:1 ratio. Cells were adhered to poly-L-lysine–coated glass slides, followed by fixation with 4% paraformaldehyde, staining with mouse anti-perforin Ab, and followed by Texas red–conjugated anti-mouse IgG Ab and analyzed via fluorescence microscopy.

Isolation of lytic granules from NK cells.

Postnuclear lysate (P NL) and crude lysosomal frac-
Granzyne A release assay

Granzyne A has an esterase activity, which can be measured using the substrate BLT ester as previously described (24). Specifically, scramble and UNC-45A knockdown NKL cells were either mock-treated (DMSO) or treated with 25 ng/ml PMA and 0.5 μg/ml ionomycin for 5 h at 37°C. Cell-free supernatant and total cell lysate obtained by repeated freeze-thaw were assayed for esterase (granzyne A) activity using BLT as a substrate. The specific release of serine esterase was measured by incubation of 45 μl cell-free supernatant with 180 μl 0.2 M Tris-HCl, pH 8.1 containing 0.2 mM BLT and 0.22 mM dithio-bis-nitrobenzoic acid. The percentage of specific enzymatic release was calculated on the basis of the total enzymatic content (total lysate). Absorbance was read using a spectrophotometer at 405/600 nm (ELISA reader 190; Molecular Devices).

IFN-γ secretion assay

Scramble and UNC-45A knockdown NK-92 cells were rested in IL-2-free medium for 16 h after stimulation with (or without) 10 ng/ml IL-12 and 100 ng/ml IL-18 for 4, 7, and 16 h. The supernatant was collected and IFN-γ secretion was measured using a commercial ELISA kit (DY285; R&D Systems).

Measurement of intracellular perforin by FACS analysis

For intracellular measurement of perforin expression levels, YT NK cells transduced with either shRNA-scramble or shRNA targeting two different UNC-45A regions (1 and 2), original YT cells, or isotype-treated (anti-IgG) YT cells were fixed, permeabilized, and stained with PE-conjugated anti-perforin Ab (556434; BD Pharmingen). At the end of the incubation period, cells were washed and analyzed using a BD LSR II flow cytometer.

Myosin II–actin functional assay

ATP-dependent myosin II–actin filament interaction was measured by adapting a recently described method for quantification of myosin II–actin binding (15). Specifically, UNC-45A knockdown (KD81 and KD2) or UNC-45A–GFP-expressing YT cells were lysed in buffer containing 20 mM Tris, pH 7.0, 0.5% Triton, 25 mM KCl, 5 mM MgCl2, 0.5 mM EGTA, 1 mM PMSE, and 1× proteinase inhibitor mixture. After elimination of unbroken cells and debris (100,000 × g for 30 min at 4°C), supernatants were further spun at 100,000 × g for 30 min at 4°C using a TL100 Beckman, and the supernatants supplemented with 30 μg/ml human platelet nonmuscle actin and incubated for 2 h on ice. Actin-bound myosin II was pelleted by ultracentrifugation (100,000 × g for 30 min at 4°C) to release functional myosin II from the pellet. The supernatant served as input for the binding reaction. A total of 5 μM Mg-ATP and 150 μg/ml actin was added to the supernatant; then the sample was transferred to a new tube (4°C) and incubated for 2 h on ice. After the incubation, 10 μl apyrase (0.1 U/μl; Sigma) was added to one of the samples for ATP minus actin binding (motor domain-dependent actin binding). Ten minutes after incubation at room temperature, the samples were centrifuged at 100,000 × g for 30 min at 4°C. The supernatants were discarded and the pellet washed once with 100 μl lysis buffer plus 5 mM ATP either treated without or with apyrase. After centrifugation, the pellets were resuspended in 30 μl SDS-PAGE sample buffer and processed for Western blot analysis.

Cell image analysis

Images were analyzed using either ImageJ or Imaris (version 8.1.2; Bitplane) software. For immunofluorescence analyses, thresholds were determined automatically using Costes’ algorithms in the Imaris program or calculated using ImageJ. Areas analyzed were either the entirety of the NK cell, defined using phase-contrast images to determine the perimeter of the NK cell, half of the cell oriented toward the NKIS, or the polarized region of a conjugated NK cell. The polarized region was defined as a triangular area with its tips located at the edge points of the NK–target cell interface and the center of the NK cell. Granules were considered polarized if >90% of granules were localized within this area. The mean fluorescent intensity (MFI) was defined as the quantified intensity of the fluorophores in the polarized area. To determine the amount of UNC-45A that localizes to the NKIS upon conjugation, we drew a line parallel to the cell-cell contact site through the center of the NK cell body. The ratio was defined as fluorescent intensity between the NKIS-oriented half of the cell and the entire cell body. Perforin localization was quantified by segregating the NK cell into four quadrants. Quadrants were defined by placing the intersection of perpendicular lines at the center of the NK cell, with the lines parallel to the sides of the field of vision. The upper left quadrant was labeled Q1, the upper right Q2, the bottom left Q3, and the bottom right Q4. Perforin puncta were then counted for each quadrant. Arbitrary units are defined as the values generated from ImageJ analyses and are relative to each respective experiment. Perforin puncta were then counted for each quadrant. Cell morphology was analyzed as cell area and circularity using ImageJ software, with the cell perimeter defined using the ImageJ drawing tool. Cell circularity was defined as the ratio between the area of the cell and its perimeter (4π × area/π×radius2). The ratio ranges from 0 to 1, with 1 indicating a perfect circle and values progressing toward 0 representing an increasingly elongated shape. For all of the quantifications, between 10 and 30 cells/condition were analyzed. Experiments were repeated a minimum of three times.

Statistical analysis

Results are reported as mean ± SD of three or more independent experiments. Unless otherwise indicated, statistical significance was assessed by two-tailed Student t test using Prism (V.4 GraphPad, San Diego, CA) and Excel. The level of significance was set at p < 0.05.

Results

UNC-45A controls NK cell-mediated cytotoxicity

Microarray analysis indicates a pattern of high UNC-45A RNA expression in human blood cells and in NK cells in particular (20). Thus, we first investigated whether UNC-45A is expressed at protein levels in human lymphocytes. To this end, lysates from CD14+ monocytes, CD19+ B cells, CD4+ T cells, CD8+ T cells, CD3–CD56+ NK (p-NK) cells, and three NK cell lines YT, NKL, and NK-92 were subjected to Western blot analysis using an mAb against UNC-45A. As shown in Fig. 1A, we found that UNC-45A is expressed at protein levels in all the cell subsets analyzed, confirming the presence of UNC-45A protein in human lymphocytes.

Next, we tested the hypothesis that UNC-45A is a binding partner of NMIIA in the NK cell context. To this end, lysates from NK cell lines and primary p-NK cells were immunoprecipitated with an anti-UNC-45A mAb, and binding to NMIIA was evaluated by Western blot. As shown in Fig. 1B, UNC-45A coimmunoprecipitates with NMIIA in both human YT and NKL NK cell lines and primary p-NK cells. To test whether UNC-45A has a potential role in controlling NK-mediated functions, we investigated UNC-45A subcellular localization in activated human NK cells. To this end, human YT NK cells expressing UNC-45A–GFP fusion protein and primary NK cells were incubated with 721.221 or K562 target cells, respectively, and the effector–target conjugates were stained with Abs against UNC-45A and/or NMIIA. Immunofluorescence microscopy revealed that UNC-45A and NMIIA colocalize at the effector–target cell interface of activated YT and p-NK cells (Fig. 1C, left and right panels, respectively). Specifically, image quantification analyses revealed that ~60% of the total UNC-45A present in the YT cells localizes at the NKIS and >70% of the total UNC-45A present in p-NK cells localizes at the NKIS (Fig. 1D, top panel). Furthermore, image quantification analyses revealed that in both YT and p-NK cells, >70% of the total UNC-45A colocalizes with NMIIA in NK cells (Fig. 1D, bottom panel). Taken together, this suggests that UNC-45A is a likely candidate for mediating NMIIA-associated functions in NK cells.

UNC-45A colocalizes with perforin in activated NK cells and associates with lytic granules

Given that UNC-45A colocalizes with and binds to NMIIA in NK cells (Fig. 1), we next wanted to evaluate whether UNC-45A associates with perforin-containing lytic granules in NK cells. To test this, YT NK cells were transduced with UNC-45A–GFP fusion protein and incubated with 721.221 target cells and conjugates stained with a mAb against perforin to allow for visualization. UNC-45A shows a ubiquitous subcellular localization and
a partial overlap with perforin granules in unengaged NK cells (Supplemental Fig. 1). In activated NK cells, however, UNC-45A is polarized at the effector–target cell interface, where it colocalizes with perforin (Fig. 2A, left panels). Specifically, image quantitation analyses revealed that ~70% of the total cellular UNC-45A colocalizes with perforin at the effector–target cell interface (Fig. 2A, right panel).

Next, we biochemically tested whether UNC-45A interacts with lytic granules. Cell lysates from YT NK cells were separated by density gradient fractionation (22) and the derived fractions, along with PNL and CLFs, were analyzed by Western blot using Abs against UNC-45A, NMIIA, and LAMP-1. UNC-45A was found specifically in the lytic granule preparation containing NMIIA and LAMP-1 (Fig. 2B). Quantification of the relative abundance of UNC-45A, NMIIA, and LAMP-1 in NK cell–derived lysosomal fractions is shown in Fig. 2C. Taken together, these results suggest that UNC-45A is a component of a multiprotein complex formed during NK cell activation and may play a role during NKIS formation and/or lytic granule exocytosis.

UNC-45A knockdown impairs the cytotoxic activity of human NK cells

Next, we wanted to test whether the presence of the NMIIA cochaperone UNC-45A is required for NK cell–mediated cytotoxicity. To this end, UNC-45A was knocked down via lentiviral delivery of shRNAs targeting multiple UNC-45A regions in NK cell lines and p-NK cells. Efficiency of UNC-45A knockdown was evaluated by Western blot and RT-PCR analysis showing a reduction in UNC-45A expression of ~90% in YT cells (Fig. 3A) and ~75% in p-NK cells (Fig. 3B). Next, we evaluated the effect of UNC-45A knockdown on the capability of NK cell lines and p-NK cells to kill target cells. YT cells infected with lentiviral particles carrying either shRNA scramble or shRNAs directed against two different UNC-45A sequences (1 and 2) were incubated with 721.221 target cells and residual specific lysis evaluated by 51Cr-release assays. UNC-45A knockdown resulted in a dramatic reduction in NK cell cytotoxicity relative to the scramble controls (Fig. 3C, left panel). A similar reduction in target cell lysis was observed with specific pharmacological inhibition of NMIIA using blebbistatin (Fig. 3C, middle panel). Quantification of the reduction in NK cell–mediated lysis as a result of UNC-45A knockdown is shown in Fig. 3C (right panel). Next, we tested whether a similar effect could be observed upon UNC-45A knockdown in primary NK cells. Primary CD3+CD56+ NK cells isolated from healthy donors were infected with lentivirus harboring scramble or UNC-45A shRNA, sorted, and incubated with K562 target cells. UNC-45A knockdown resulted in significant impairment of cytotoxic capabilities in primary NK cells in 51Cr-release assays (Fig. 3D, left and middle panels). Quantification of the reduction in primary NK cell lysis resulting from UNC-45A knockdown is shown in Fig. 3D (right panel). To exclude the possibility that the reduction in the cytotoxic capability of UNC-45A knockdown NK cells was due to impairment of cell viability, we evaluated both cell morphology and proliferative activity in scramble and UNC-45A knockdown YT cells. Our results show that UNC-45A knockdown did not affect NK cell morphology as evaluated by determination of cells’ area and circularity or proliferation rate (Supplemental Figs. 2, 3, respectively). Together, these data suggest that UNC-45A plays a critical role in regulating NK-mediated cytotoxicity without interfering with cell viability.
UNC-45A knockdown does not prevent formation of effector–target conjugates

Reduction in cytotoxic capabilities in NK cell lines and primary NK cells after UNC-45A knockdown (Fig. 3), along with its colocalization with NMIIA at the NKIS (Figs. 1, 2), suggests that the loss of UNC-45A might hinder cell killing by perturbing the formation of effector–target conjugates. To test this hypothesis, we transduced YT NK cells with lenti-pEF-shRNAmir targeting scramble sequences or two different UNC-45A regions, incubated with CellTracker Orange–loaded 721.221 target cells and the number of effector–target conjugates in control and UNC-45A knockdowns evaluated using two different approaches. In the first approach, the number of green:red (YT NK:721.221) conjugates was manually counted under the microscope and reported as the number of conjugates per field. We did not observe a significant difference in terms of ability to form conjugates in control versus UNC-45A knockdown NK cells (Fig. 4A). To confirm this result, the effect of UNC-45A silencing on conjugate formation was measured by FACS analysis. YT NK cells were transduced with scramble shRNAmir or UNC-45A shRNAmir and incubated with CellTracker Orange–loaded 721.221 target cells. After fixation, FACS analysis was performed to determine the number of conjugates, defined as the green-red double-positive population. We did not observe a difference in terms of the ability of NK cells to form conjugates after UNC-45A knockdown (Fig. 4B). Specifically, quantification of the percentage of effector–target conjugates at $t = 0$ and at $t = 10$ min revealed the absence of a statistically significant difference in UNC-45A knockdown cells versus control (Fig. 4C). Thus, the role of UNC-45A in NK cell cytotoxicity is independent of NKIS formation.

UNC-45A knockdown does not prevent lytic granule polarization and does not alter perforin levels

After NKIS formation, polarization of the lytic granules at the cell–cell contact site is a required event for NK cell–mediated cytotoxicity. Because UNC-45A knockdown did not hinder NKIS formation (Fig. 4), we next evaluated whether the reduced lytic capacity of NK cells after UNC-45A knockdown is due to inefficient lytic granule polarization. To test this hypothesis, we incubated YT NK cells transduced with scramble or UNC-45A shRNAmir with 721.221 target cells and visualized recruitment of perforin to the NKIS via fluorescence microscopy. In unengaged NK cells, perforin was localized throughout the cytoplasm (Supplemental Fig. 4), whereas in engaged NK cells, perforin-containing lytic granules were found to localize at the IS in both scramble and UNC-45A knockdown cells (Fig. 5A). Quantification of the percentage of cells with polarized lytic granules in scramble versus UNC-45A knockdown (#1, #2) is shown in Fig. 5B. Next, we addressed the question whether UNC-45A knockdown would alter intracellular levels of perforin in NK cells. To this end, we first quantified the MFI in the perforin-polarized regions in scramble versus UNC-45A knockdown YT cells. As shown in Fig. 5C, we did not observe any statistically significant difference in terms of MFI in scramble versus UNC-45A knockdown cells. To confirm this result, we next measured the intracellular levels of perforin in scramble versus UNC-45A knockdown YT cells via FACS analysis. As shown in Fig. 5D, the intracellular amount of perforin in UNC-45A knockdown cells did not differ from the one in scramble or isotype-treated controls. Taken together, this suggests that UNC-45A is not required for polarization of perforin-containing lytic granules to the NKIS nor does it alter intracellular levels of perforin in NK cells.

UNC-45A is obligate for NK cell degranulation and IFN-γ secretion

NK cell–mediated cytotoxicity occurs when, after polarization at the NKIS contact site, lytic granules are secreted into the space between NK cell and target. The lack of an observable effect on NKIS formation (Fig. 4) and lytic granule polarization (Fig. 5) in NK cells after UNC-45A knockdown suggested that the loss of UNC-45A might reduce NK cell–mediated cytotoxicity by preventing lytic granule secretion at the NKIS site. To test this...
We hypothesized that UNC-45A knockdown could hinder NK cell-mediated cytotoxicity. To test this, we transduced NKL and NK-92 cells with either scramble shRNA or shRNAs targeting UNC-45A. The efficiency of UNC-45A knockdown in NKL and NK-92 cells was evaluated by Western blot analysis showing a reduction in UNC-45A expression of ∼60% in NKL cells (Fig. 6A, left panel) and ∼90% in NK-92 cells (Fig. 6A, right panel). Cells were then either mock-treated (DMSO) or treated with 50 ng/ml PMA and 0.5 μg/ml ionomycin. The effect of PMA/ionomycin stimulation on NK cell degranulation was evaluated by measuring the increase in surface expression of the lysosome marker CD107a. Our results show that UNC-45A knockdown is associated with reduced expression of CD107a on the surface of NKL (Fig. 6B, left panel) and NK-92 (Fig. 6C, left panel) cells relative to scramble controls. Quantification of CD107a surface expression (% of

**FIGURE 3.** UNC-45A knockdown hinders NK cell–mediated cytotoxicity. (A, left panel) Western blot analysis of UNC-45A expression in YT cells transduced with either shRNA-scramble or shRNA targeting two different UNC-45A regions (1 and 2). β-Actin was used as a loading control. (Right panel) Quantification of the percentage of UNC-45A expression normalized to β-actin. (B, left panels) UNC-45A knockdown in primary CD3− lymphocytes (P4) virally infected with lenti-pEF-UNC45-shRNAmir (1 and 2). Forty-eight hours postinfection, GFP+ CD3− CD56+ cells were sorted (P5) for quantification of UNC-45A levels. (Right panel) Efficiency of UNC-45A knockdown in primary NK cells measured by quantitative RT-PCR. The mRNA values reported are relative to the control gene (GAPDH) and normalized to control-treated cells. (C, left panel) Residual specific lysis in YT cells infected with lentiviral particles carrying either shRNA scramble or shRNAs directed against two different UNC-45A sequences (1 and 2) incubated with 721.221 target cells at the indicated ratios. A representative experiment is shown. (Middle panel) Residual specific lysis in YT cells either mock- (DMSO) or blebbistatin (75 μM)-treated and incubated with 721.221 target cells at the indicated ratios. A representative experiment is shown. (Right panel) Average of five independent experiments reported as reduction in specific lysis as compared with the controls (set at 100%). (D, left and middle panels) Residual specific lysis in lenti-pEF-UNC45-shRNAmir (1 and 2) or scramble infected, CD3−/CD56+/GFP+-sorted p-NK cells derived from healthy donors (donors 1 and 2) incubated with K562 target cells at the indicated ratios. (Right panel) Average of three independent experiments reported as reduction compared with controls (set at 100%). For all experiments (conducted in triplicates), specific lysis was assessed by [51Cr]-release assay. Statistical significance was set at p = 0.05.
UNC-45A knockdown does not prevent effector–target conjugate formation. (A) YT NK cells transduced with lenti-pEF-UNC45-shRNAmir targeting either scramble or two different UNC-45A (1 and 2) regions were incubated with CellTracker Orange–loaded 721.221 target cells. The number of conjugates (effector green/target red) per condition were counted under fluorescence microscopy and reported as the number of conjugates per field. The average of three independent experiments is shown. (B) YT NK cells transduced with lenti-pEF-UNC45-shRNAmir targeting either scramble of two different UNC-45A (1 and 2) regions were incubated with CellTracker Orange–loaded 721.221 target cells. Samples were either immediately fixed (t₀) or fixed after 10 min of incubation (t₁₀). The number of conjugates was quantified by FACS analysis. The average of three independent experiments is shown. (C) Quantification of percentage of effector–target conjugates in shRNA scramble or shRNA UNC-45A (1 and 2) at t = 0 and t = 10 min.

control) for YT and NK-92 is shown in Fig. 6B and 6C (right panels), respectively.

To directly determine the impact of UNC-45A on secretion of lytic granule content, we measured the effect of UNC-45A knockdown on granzyme A secretion in both NKL and PMA/ionomycin stimulation. UNC-45A knockdown resulted in a dramatic reduction in granzyme A secretion in both NKL and NK-92 NK cells (Fig. 6D, left and right panels, respectively). Next, we evaluated the impact of modulating the UNC-45A expression levels in NK cells on IFN-γ secretion. To this end, NK-92 cells transduced with either shRNA-scramble or shRNA consisting of a pool (shRNA–UNC-45A pool) of three UNC-45A–specific shRNAs were either mock-treated or treated with 10 ng/ml IL-12 and 100 ng/ml IL-18 over a period of 4, 7, or 16 h. At the end of each incubation time, residual IFN-γ was measured by ELISA assay. As shown in Fig. 6E, UNC-45A knockdown resulted with statistically significant reduction in IFN-γ by NK cells starting as early as 4 h from treatment and observable for up to 16 h.

To test whether the absence of UNC-45A in NK cells is associated with a reduction in NMIIA–actin binding capacity. As shown in Fig. 7B (left panel), UNC-45A knockdown did not affect the total expression levels of NMIIA within NK cells. However, when lysates from either scramble or UNC-45A knockdown YT NK cells were exposed to exogenous actin in presence (+) of the ATP diphosphohydrolase apyrase, this resulted with a dramatic decrease in the NMIIA–actin interactions as compared with controls (Fig. 7B, middle and right panels). Furthermore, when we performed the complementary experiments by over-expressing GFP-tagged UNC-45A in the same cells, we observed the opposite effect: a significant in NMIIA–actin binding over controls (Fig. 7C). Quantification of actin–NMIIA binding for the earlier described experiments is given in Fig. 7D. To further confirm the requirement of UNC-45A for NMIIA and actin interaction, we visualized UNC-45A, NMIIA, and actin in UNC-45A–GFP–expressing YT NK cells conjugated with 721.221 cells. As shown in Fig. 7E, UNC-45A, NMIIA, and actin all appear to localize at the interface between effector and target cell. Taken together, our data suggest that reduced lytic granule exocytosis in NK cells lacking UNC-45A may be caused by impairments in cellular cytoskeletal dynamics.

Discussion
Perturbing of the actin-based microfilaments within NK cells has dramatic consequences on their activities including cytotoxicity.
Thus, understanding of mechanisms regulating the cytoskeletal dynamic within NK cell is fundamental for the understanding of NK cell's functions in both the physiological and the pathological context.

Exposure of NK cells to the NMIIA inhibitor blebbistatin, for instance, causes a block of the myosin heads in a state of low affinity for actin resulting with impaired NK-mediated cytotoxicity (2, 22, 27–29). Mutations in \( \text{WAS} \) (encoding WASp) on the other side are associated with reduced perforin granule polarization at the NKIS due to inefficient F-actin branching (5, 30–33). Furthermore, loss of WASp-interacting protein functions causes defects in NK cell–mediated killing via increased proteasome-mediated WASp degradation and its mislocalization (32, 34–37).

Lastly, inefficient tethering of lytic granules to the plasma membrane in NK cells carrying mutations in the \( \text{UNC13D} \) gene is associated with a complete loss of NK cell–mediated cytotoxicity (11, 38–41).

FIGURE 5. UNC-45A knockdown does not interfere with lytic granules polarization or with perforin expression levels. (A) YT NK cells transduced with either shRNA-scramble or shRNA targeting two different UNC-45A regions (1 and 2) were incubated with 721.221 target cells. After fixation, cells were stained with an anti-perforin mAb followed by Texas red goat anti-mouse IgG and analyzed by fluorescence microscopy. A representative experiment is shown. Arrows indicate NKISs. Scale bars, 5 \( \mu \)m. (B) Quantification of polarized granules per condition reported as the percentage of cells with polarized lytic granules out of the total number of conjugates. The averages of three independent experiments are shown. (C) Quantification of MFI in polarized regions in YT cell transduced with either scramble or shRNAs targeting UNC-45A regions (1 and 2). (D) Intracellular staining of perforin in YT cell transduced with either scramble of shRNAs targeting UNC-45A regions (1 and 2) or original, or treated with anti-mouse IgG as isotype control and analyzed by FACS using a PE-conjugated anti-perforin Ab.

Together our data demonstrate that UNC-45A is a component of the NK cell cytoskeleton and plays a key role during NK cell–mediated cytotoxicity.
FIGURE 6. UNC-45A knockdown prevents lytic granule secretion and IFN-γ release in NK cells. (A) UNC-45A protein expression levels in lysates of NKL (left panel) and NK-92 (right panel) cells upon transduction with either shRNA-scramble or shRNA targeting two different UNC-45A regions (1 and 2) and in NK-92 NK cells after transduction with shRNA scramble or shRNA consisting of a pool (shRNA–UNC-45A pool) of three 19- to 25-nt UNC-45A–specific shRNAs. β-Actin was used as a loading control. (B) NKL cells transduced with either shRNA-scramble or shRNA targeting two different UNC-45A regions (1 and 2) and either mock-treated (DMSO) or treated in presence of 25 ng/ml PMA and 0.5 μg/ml ionomycin. Surface CD107a expression was analyzed by FACS. A representative experiment is shown. (Right panel) Quantification of CD107a expression averaged from three independent experiments. (C, left panels) NK-92 cells transduced with either shRNA-scramble or shRNA consisting of a pool (shRNA–UNC-45A pool) of three 19- to 25-nt UNC-45A–specific shRNAs either mock-treated (DMSO) or treated in the presence of 50 ng/ml PMA and 0.5 μg/ml ionomycin. The effect of PMA/ionomycin stimulation on NK cells was evaluated by measuring the increase in surface expression of CD107a by FACS analysis. Representative experiment. (Right panel) Quantification of the CD107a expression. Average of three independent experiments. (D, left panel) NKL cells transduced with either shRNA-scramble or shRNA targeting two different UNC-45A regions (1 and 2) and either mock-treated (DMSO) or
mediated cytotoxicity in a step between approach of the lytic granules to the NKIS and their release at the effector–target cell interspace. Although the nature of this step is yet to be identified, we propose that the effect of UNC-45A in controlling lytic granule exocytosis could be either dependent or independent from its effect on regulating NMIIA activity, and that these two scenarios are not mutually exclusive. One possibility is that UNC-45A plays a role in assisting NMIIA in transporting the lytic granules through F-actin at the IS to facilitate their fusion to the cell membrane. In a separate scenario, UNC-45A acts independent of NMIIA and is part of the multisubunit complex that regulates the events of priming, docking, and fusion of lytic granules to the cell membrane leading up to target cell killing.

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FIGURE 7. Effect of UNC-45A knockdown on NMIIA expression levels and ATP-sensitive interactions of NMIIA with actin filaments. (A) Expression levels of UNC-45A, NMIIA, pNMIIA H chain (HC), and pNMIIA L chain (LC) in YT cells transduced with either shRNA-scramble or shRNA targeting two different UNC-45A regions. β-Actin was used as a loading control. (B) NMIIA–actin binding capability in YT cells transduced with either scramble or shRNA targeting two different UNC-45A regions in the absence (−) and presence (+) of the ATP diphosphohydrolase apyrase. (C) NMIIA–actin binding capability in YT cells transduced with empty vector (lenti-pEF-GFP) or lenti-pEF-UNC-45A-GFP in the absence (−) and presence (+) of the ATP diphosphohydrolase apyrase. (D) Quantification of NMIIA actin-binding capability expressed as a percentage of the control (average of two independent experiments) and obtained by comparing lines 5 and 7 and 9 and 11 of (B) and lines 4 and 6 of (C). (E) YT NK cells expressing UNC-45A–GFP fusion protein were incubated with 721.221 target cells. After fixation, effector–target conjugates were stained with an anti-NMIIA polyclonal Ab and an anti-actin mAb followed by Texas red goat anti-rabbit IgG and Alexa Fluor 647–conjugated donkey anti-mouse IgG and analyzed by fluorescence microscopy. Arrow indicates the NKIS.
Disclosures

The authors have no financial conflicts of interest.

References


