Cancer-associated fibroblast suppresses killing activity of natural killer cells through downregulation of poliovirus receptor (PVR/CD155), a ligand of activating NK receptor

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Abstract. Cancer-associated fibroblasts (CAFs) play an important role in cancer expansion and progression in tumor microenvironment (TME), via both direct and indirect interactions. Natural killer (NK) cells play a crucial role in anticancer immunity. We investigated the inhibitory effects of CAFs on NK cell activity. CAFs were isolated from endometrial cancer tissue, while normal endometrial fibroblasts (NEFs) were obtained from normal endometrium with no pathological abnormality. NK cells were obtained from allogenic healthy volunteers. CAFs or NEFs were co-cultured at an NK/fibroblast ratio of 1:1 with or without inserted membrane. For NK cell activity, K562 cells were cultured as target cells. NK cell-killing activity was determined by calculating the ratio of PI-positive K562 cells in the presence of NK cells co-cultured with fibroblasts versus NK cells alone. To examine whether NK cell activity was suppressed by IDO pathway, we inhibited IDO activity using the IDO inhibitor 1-MT. We demonstrated that CAFs derived from endometrial cancer induced greater suppression of the killing activity of allogenic NK cells compared with normal endometrial fibroblasts (NEFs). The suppression of NK cell activity by CAFs was inhibited when a membrane was inserted between the CAFs and NK cells, but not by 1-MT, an inhibitor of IDO.

Introduction

Cancer-associated fibroblasts (CAFs) regulate not only carcinogenesis, but also the immune evasion of cancer in the tumor microenvironment (TME), which facilitates cancer cell proliferation, expansion, and metastasis (1-5). The TME is comprised of various types of cells, including microvascular endothelial cells, immune cells, and CAFs, and these cells interact directly and indirectly with each other maintaining homeostasis of the TME. Among these cells, CAFs are important regulators of the immunological microenvironment of the tumor. Sanz-Moreno et al reported that CAFs regulate immune evasion in the TME by various mechanisms, including the use of cytokines and cell attachment (6). They demonstrated that the ROCK and JAK1 signaling pathway generates a contractile force in stromal fibroblasts, allowing remodeling of the extracellular matrix and the creation of tracks for the collective migration of squamous carcinoma cells. Furthermore, Gaggioli et al demonstrated that the generation of these tracks by fibroblasts was sufficient in enabling collective invasion of squamous cell carcinoma cells (7).

NK cells play an important role in cancer immunity in the TME. A review by Chan et al identified several well-known ligands of NK paired or activating receptors that are expressed on the cell surface of target cells, including malignant cells...
Isolation of NK cells and NK cell/fibroblast co-cultures. 2-8 passages were used before the experiments. NK cells were obtained from healthy volunteers after Ficoll-Paque isolation kit from Miltenyi Biotec. NK cell purity was >95% as evaluated by flow cytometry. They were cultured in RPMI with 10% FBS, 100 IU/ml penicillin, 100 µg/ml streptomycin, at 37°C in humidified 5% CO2 atmosphere, and stimulated 1 ng/ml IL15 (R&D) for 48 h, before co-culture.

For co-culture experiments, CAFs or NEFs were seeded at 5x10^4 in 24-well plates with 500 µl of medium and cultured for 48 h. NK cells were added at 1x10^5 well, at an NK/fibroblasts ratio of 1:1, with or without 1-µm pore of cell-culture-insert. NK cells alone were also cultured in the absence of fibroblasts. After 24-h incubation, NK cells were harvested and analyzed. We confirmed that allogenic NK cells did not kill these fibroblasts during co-culture (data not shown).

**NK cell killing assay.** K562 cells (obtained from ATCC; American Type Culture Collection, VA, USA) were cultured in RPMI-1640 supplemented with 10% FBS containing 100 U/ml penicillin and 100 µg/ml streptomycin. K562 cells were resuspended at 1-2x10^5 cells/ml in 0.1% FBS/PBS and added CFSE, using CFSE Cell Division Assay kit from Invitrogen, to a final concentration of 2.5 µM CFSE staining solution, incubated cells at 37°C for 30 min. After centrifugation of cells at 300 g for 10 min, supernatant was discarded and the cells were resuspended into RPMI medium and incubated at 37°C for 30 min. The cells were washed with 1% FBS/PBS three times, and 2x10^4 of stained K562 cells in 200 µl medium were added to tubes. Harvested NK cells were diluted at 1x10^6 cells/ml and prepare dilution series 2x10^5 cells per 200 µl. After incubation the tubes at 37°C for 4 h, 5 µl of PI (propidium iodide solution, Biolegend, San Diego, CA, USA) was added for dead cell count. Percentage of PI-positive dead K562 cells (% K562 PI-positive cell) was evaluated by flow cytometry. In some experiments, allogenic NK cells were incubated for 24 h with NEFs, CAFs or no fibroblasts followed by exposure to target (K562) cells. NK cell killing activities were indicated by ratio of % K562 PI positive cells under each condition against that of NK cells only.

**Immunocytochemistry.** CAFs were grown on coverslips, then serum deprived and fixed in 4% paraformaldehyde, permeabilized with 0.1% Triton X-100, washed twice with PBS, and incubated for 60 min at room temperature with an anti-α-SMA antibody (mouse clone 1A4, ab7817, Abcam MA, USA), labeled with Alexa Flour 488 (zenon). After incubation, the slides were washed with PBS and fixed with 4% paraformaldehyde. The cells were counterstained with 4',6-diamidino-2-phenylindole dihydrochloride (DAPI).

**Western blotting.** CAFs and NEFs grown in chamber slides were fixed and each of them were collected and total proteins were extracted. Immunoblotted with anti-beta Actin antibody (rabbit polyclonal antibody, Abcam), and α-SMA (mouse clone 1A4, ab7817, Abcam).

**Flow cytometry.** Cultured cells were harvested and incubated with PerCP Cy5.5-conjugated CD90 (mouse clone Thy1, BioLegend), FITC-conjugated Vimentin (mouse clone RV202, ab8978, Abcam), PE-conjugated PVR (mouse clone SK114, BioLegend) or appropriate isotype control at 4°C for 30 min, washed twice, and analyzed using a BD FACS Calibur cytometer. The results were analyzed using Kaluza software.

**Materials and methods**

**Patients and establishment of fibroblasts.** Tumor samples were obtained from the patients with endometrial carcinoma, and normal endometrium were collected from those without pathology in uterine endometrium, undergoing surgical resection in our hospital. All women gave written informed consent and the Research Ethics Committee of the University of Tokyo approved all aspects of the study.

CAFs were isolated from the cancer tissues of endometrial cancer while NEFs were from the normal endometrium with no pathological abnormality. The tissues were minced and digested in DMEM/F12 medium, (Gibco, Japan), supplemented with 100 IU/ml penicillin, 100 µg/ml streptomycin, 1 mg/ml collagenase type I (Wako, Tokyo, Japan), and 25 ng/ml DNase, Roche Diagnostics GmbH at 37°C for 60 min, filtered with 100 and 70 µm cell strainers, BD Falcon, and centrifuged at 1,500 rpm for 5 min and washed with D-PBS, Wako. They were resuspended in DMEM/F12 with 10% FBS, 100 IU/ml penicillin, 100 µg/ml streptomycin, and cultured at 37°C in humidified 5% CO2 environment. Fibroblasts passed for 2-8 passages were used before the experiments.

**Isolation of NK cells and NK cell/fibroblast co-cultures.** NK cells were obtained from healthy volunteers after Ficoll-Paque gradient and negative magnetic selection, using human NK cell isolation kit from Miltenyi Biotec. NK cell purity was >95% as evaluated by flow cytometry. They were cultured in RPMI with 10% FBS, 100 IU/ml penicillin, 100 µg/ml streptomycin, at 37°C in humidified 5% CO2 atmosphere, and stimulated 1 ng/ml IL15 (R&D) for 48 h, before co-culture.

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siRNA experiments. NEFs were transfected with 100 pmol of siRNA with Lipofectamine RNAiMAX. The complexes of RNAi and 1.5 µl Lipofectamine RNAiMAX in 140 µl Opti-MEM I medium without serum. Then the complexes were added in the cultured cells at 4x10^4 in 24-well plates, and 500 µl of complete medium without antibiotics was gently added in each well, and incubated for 12 h at 37˚C in a CO₂ incubator to be ready for the next assay of gene knockdown. To knock down PVR, siRNA by R&D was used. The sequences were as follows: human PVR; sense, 5’-rCrArGrCuUrAuUr CrGrGrArCuUrCrArArATT; antisense, 5’-UUArGrGrArCuUr CrGrGrArAuUrGrCuUrGTT. The negative siRNA controls were obtained from Life Technologies.

Statistical analysis. Data are presented as means ± SEM. Statistical analyses were carried out using Student's t-test or Dunnett analysis using JMP software. A value of p<0.05 was considered significant. Asterisks indicate those comparisons with statistical significance (p<0.05).

Results

Isolation of CAFs from endometrial cancer. To investigate the difference in effect on NK cell activity between NEFs and CAFs, we isolated fibroblasts derived from normal endometrium (NEFs) and endometrial cancer tissue (CAFs). Uterine endometrium is composed of many glands and abundant stroma. The endometrial stroma is enriched in fibroblasts, and surrounds the endometrial glandular epithelium. Endometrial cancer can transform normal fibroblasts in the stroma into CAFs. Therefore, the comparison between endometrial CAFs and NEFs was thought to be suitable for the investigation of CAF activity. Fibroblasts were isolated from cancer tissue or normal endometrium by standard isolation methods and identified by immunostaining and western blotting. Fibroblasts isolated from endometrial cancer expressed the fibroblastic markers vimentin and CD90 (Fig. 1A). To distinguish the cancer-derived fibroblasts from the NEFs, presence of α-SMA was assessed by immunocytochemistry (Fig. 1B), and expression level of α-SMA compared with NEFs was assessed by western blotting (Fig. 1C). α-SMA localizes to the cytoplasm in cancer-derived fibroblasts (Fig. 1B). Western blotting demonstrated increased expression levels of α-SMA in two cancer-derived fibroblast samples compared with two NEF samples. Therefore, the cancer-derived fibroblasts isolated in this study possessed typical CAF characteristics, and were confirmed to be CAFs.

Suppression of NK cell-killing activity by CAFs. We assessed the effect of CAFs on the killing activity of NK cells to investigate the mechanism of CAF-mediated immune evasion. Assessment of NK cell activity is often performed by measuring allogenic NK cell killing activity against K562 cells (14). Using this NK cell-killing assay, we first examined whether CAFs isolated from endometrial cancer suppressed the killing activity of allogenic NK cells (Fig. 2). NK cells co-cultured with CAFs were assessed for killing activity by changing the E:T (NK cell: K562 cell) ratio. The proportion of
dead K562 cells increased in an E:T ratio-dependent manner in the presence of NK cells alone, or with NK cells co-cultured with CAFs, indicating that the target (K562) cells were killed by allogenic effector (NK) cells (Fig. 2). The killing activity of NK cells co-cultured with CAFs was significantly reduced, to less than one third the level of NK cells alone (Fig. 2). Next, NK cell activity was assessed when co-cultured with either NEFs or CAFs (Fig. 3). Allogenic NK cells were incubated for 24 hours with NEFs, CAFs, or no fibroblasts, and then exposed to K562 cells. NK cell-killing activity was determined by calculating the ratio of PI-positive K562 cells in the presence of NK cells co-cultured with fibroblasts versus NK cells that had been cultured alone. There was no significant difference in killing activity observed between NK cells only and NK cells co-cultured with NEFs. Again, the killing activity of NK cells co-cultured with CAFs was significantly decreased, to approximately one-third of the activity of NK cells cultured with no fibroblasts. However, NK cell activity was significantly decreased when co-cultured with CAFs, compared with NEFs (Fig. 3).

The CAF-induced decrease in NK cell activity was not rescued by indoleamine 2,3-dioxygenase inhibitor. Some studies have shown that malignant cells suppress NK cell activity via production of IDO (15,16). IDO expression was barely detected in non-stimulated CAFs, whereas it was clearly detected in CAFs co-cultured with NK cells (data not shown). Previous reports have demonstrated that IDO expression in CAFs is likely caused by NK cell-derived IFNγ (17). Therefore, we inhibited IDO activity using the IDO inhibitor 1-MT, to investigate whether IDO was involved in the suppression of NK cell activity by CAFs. NK cells and CAFs were co-cultured in the presence or absence of 1-MT, followed by assessment of NK cell-killing activities (Fig. 4). While NK cell activity was...
decreased by co-culture with CAFs, this suppression was not rescued by 1-MT treatment.

Cell-to-cell interaction was critical for the CAF-induced decrease in NK cell activity. To examine whether direct cell-to-cell interaction between NK cells and CAFs was required for the suppression of NK cell activity, NK cells and CAFs were cultured in a chamber with an inserted membrane separating these cells (Fig. 5). The suppression of NK cell activity by CAFs was completely abolished by the physical blockage of cell-to-cell interaction using the inserted membrane. These data strongly suggested that CAFs suppressed NK cell activity via cell-to-cell interaction rather than IDO, or other cytokine, production.

Cell-surface expression of PVR is decreased in CAFs compared with NEFs. We next focused on the cell-surface ligands expressed on CAFs that interact with activating NK cell receptors. Several ligands of paired or activating NK receptors have been previously demonstrated to be expressed on the cell surface of target cells, including malignant cells (18). However, few reports have investigated the cell-surface expression of NK receptor ligands on CAFs. We investigated the expression of several NK receptor ligands in CAFs. Due to the difference between CAFs and NEFs in their ability to induce NK cell activity, we hypothesized that the expression of NK receptor ligands may differ between CAFs and NEFs. Expression of poliovirus receptor (PVR/CD155), a ligand of paired NK receptors DNAM-1 and TIGIT, was observed to differ between CAFs and NEFs (Fig. 6). Flow cytometry analysis revealed that the cell-surface expression of PVR was decreased in CAFs compared with NEFs. This suggested that the downregulation of PVR in CAFs may mediate the suppression of DNAM-1 signaling.

PVR knockdown downregulates NK cell activity. To confirm whether PVR downregulation results in the decrease of NK cell-killing activity, PVR expression in NEFs was knocked down using siRNA against PVR (PVRsi) (Fig. 7A). PVRsi-transfected NEFs showed decreased cell-surface PVR expression when compared with control siRNA-transfected NEFs. The median flow cytometry index (MFI) of PVR expression in PVR knockdown cells was similar to that of NEFs. Asterisk indicates p-value of <0.05.
CAFs (Fig. 7A). Interestingly, NK cell-killing activity of NK cells co-cultured with PVRsi-transfected NEFs was decreased to approximately one third of control si-transfected NEFs (Fig. 7B). The decreases in both PVR expression and effect on NK cell activity were approximately equivalent to those of CAFs. These data suggested that the level of PVR cell-surface expression in CAFs was critical for the killing activity of NK cells.

Discussion

We demonstrated that CAFs showed increased suppression of NK cell-killing activity compared with NEFs, due to decreased PVR cell surface expression, a ligand of an NK activating receptor. In this study, human CAFs and NEFs were isolated from the stroma of endometrial cancer and normal endometrium, respectively, and their interactions with NK cells were compared. Uterine endometrium is composed of many glands and abundant stroma. The endometrial stroma is enriched in fibroblasts, and surrounds the endometrial glandular epithelium. Endometrial cancer cells can transform normal fibroblasts in the stroma into CAFs. In the cancer microenvironment, CAFs acquire cancer-specific characteristics in addition to their fibroblastic background. Therefore, we believe that the comparison between endometrial CAFs and normal endometrial stromal fibroblasts was suitable for investigation of CAF characteristics. We observed an increased α-SMA expression level in CAFs compared with NEFs, in confirmation with previous studies demonstrating the expressing of α-SMA and vimentin in active CAFs (19). CD90 is a fibroblastic marker that is also expressed in CAFs of a variety of cancers, such as lung, prostate, pancreas, and breast cancers (20-22). CAFs isolated from endometrial cancer in this study were fibroblasts showing typical CAF characteristics.

Many previous studies have demonstrated the suppression of NK cell activity by cancer cells by measuring the killing activity of allogenic NK cell against K562 cells (23). Some studies have shown that malignant cells suppress NK cell activity via the production of IDO. In this study, the suppression of NK cell-killing activity by CAFs was not rescued by exposure to 1-MT (IDO inhibitor), suggesting suppression of NK cell activity is not due to IDO production by CAFs. Prevention of cell-to-cell interaction between NK cells and CAFs completely abolished the suppression of NK cell activity by CAFs, indicating direct interaction between NK cells and CAFs was critical for the CAF-induced suppression of NK cell activity. Several studies have demonstrated that exosomes derived from CAFs may interact with NK cells to suppress NK activity. However, exosomes should be able to penetrate the transwell membrane used in this study, suggesting that the suppression of NK cell activity was not dependent on CAF exosome exposure.
Our data demonstrated cell-surface expression of PVR was reduced in CAFs, compared with NEFs. The MFI of PVR in CAFs was half that in NEFs. NK activating receptors include NKp30, NKp44, NKp46, NKG2D, DNAM-1, and LFA-1 (8). Among the ligands of these receptors, PVR, a ligand of DNAM-1, was observed in this study to be ubiquitously expressed on the cell-surface of CAFs. It is known that DNAM-1 activates NK cell activity. We observed that co-culture of NK cells and CAFs did not alter DNAM-1 expression in NK cells (data not shown). In contrast, PVR interacts with NK paired receptors, DNAM-1 (activating) and TIGIT (inhibiting). Interestingly, some studies demonstrated that PVR interacts with higher affinity to TIGIT than DNAM-1. Our data suggested that the reduced expression of PVR in CAFs resulted in decreased interaction with DNAM-1, leading to suppression of NK killing activity. Furthermore, it is possible that the high affinity of PVR to TIGIT allowed even low expression of cell-surface PVR in CAFs to bind to TIGIT, thus inhibiting NK cell activity. We also demonstrated that the NK killing activity of PVR-knockdown NEFs decreased to levels comparable with that of CAFs. The data suggested that the reduction of PVR expression in CAFs is critical for CAF-induced suppression of NK cell activity. Many studies have reported PVR involvement in NK cell-associated immune-evasion by malignant cells, and that low expression of PVR is associated with poor prognosis (24). However, this study is the first to demonstrate that the suppression of NK killing activity by CAFs was mediated by downregulation of cell-surface PVR in CAFs; however, the mechanism of PVR downregulation remains to be clarified. Recently, a study by Gong et al reported that ER stress induces the downregulation of PVR in hepatoma cells via the ATF6 and IRE1α pathways (25). ER stress, such as that induced by hypoxia in the TME, may result in downregulation of PVR in CAFs; therefore a low-level of PVR expression in fibroblasts may be a characteristic of CAFs. Further study is warranted to bring to light the mechanism by which CAFs acquire such features in the TME.

CAFs may assist malignant cells in a similar fashion to maintain the immunosuppressive microenvironment of the tumor. Additionally, immune evasion may be influenced by both CAFs and malignant cells with regard to NK cell-mediated killing activity. Therefore, soluble PVR may be used as a potential agent to activate NK cell activity in the TME. These data may provide a novel strategy for inhibiting the immune evasion system in the TME.

In conclusion, this is the first report to demonstrate that CAF-mediated suppression of NK killing activity is due to downregulation of PVR cell-surface expression in CAFs. We discovered that CAFs suppressed NK cell function via a receptor-ligand interaction, aiding cancer progression. Soluble PVR may be used as a potential agent to activate NK cell activity in the TME. These data may provide a novel strategy for inhibiting the immune evasion system in the TME.

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