IL-12 Production and Subsequent Natural Killer Cell Activation by Necrotic Tumor Cell-loaded Dendritic Cells in Therapeutic Vaccinations

Aeyung Kim1, Kwang Dong Kim1, Seung-Chul Choi1, Moon-Jin Jeong2, Hee Gu Lee1, Yong-Kyung Choe1, Sang-Gi Paik3 and Jong-Seok Lim1

1Laboratory of Cell Biology, Korea Research Institute of Bioscience and Biotechnology, Daejeon, 2Department of Oral Histology, College of Dentistry, Chosun University, Gwangju, 3Department of Biology, Chungnam National University, Daejeon, Korea

ABSTRACT

Background: Immunization of dendritic cells (DCs) pulsed with tumor antigen can activate tumor-specific cytotoxic T lymphocytes (CTL) that are responsible for protection and regression. In this study, we examined whether the uptake of necrotic tumor cells could modulate DC phenotypes and whether the immunization of necrotic tumor cell-loaded DCs could elicit efficient tumor specific immune responses followed by a regression of established tumor burdens. Methods: We prepared necrotic tumor cell-pulsed DCs for the therapeutic vaccination and investigated their phenotypic characteristics, the immune responses induced by these DCs, and therapeutic vaccine efficacy against colon carcinoma in vivo. Several parameters including phagocytosis of tumor cells, surface antigen expression, chemokine receptor expression, IL-12 production, and NK as well as CTL activation were assessed to characterize the immune response. Results: DCs derived from mouse bone marrow efficiently phagocytosed necrotic tumor cells and after the uptake, they produced remarkably increased levels of IL-12. A decreased CCR1 and increased CCR7 expression on DCs was also observed after the tumor uptake, suggesting that antigen uptake could induce DC maturation. Furthermore, co-culturing of DCs with NK cells in vitro enhanced IL-12 production in DCs and IFN-γ production in NK cells, which was significantly dependent on IL-12 production and cell-to-cell contact. Immunization of necrotic tumor cell-loaded DCs induced cytotoxic T lymphocytes as well as NK activation, and protected mice against subsequent tumor challenge. In addition, intratumoral or contra-lateral immunization of these DCs not only inhibited the growth of established tumors, but also eradicated tumors in more than 60% of tumor-bearing mice. Conclusion: Our data indicate that production of IL-12, chemokine receptor expression and NK as well as CTL activation may serve as major parameters in assessing the effect of tumor cell-pulsed DC vaccine. Therefore, DCs loaded with necrotic tumor cells offer a rational strategy to treat tumors and eventually lead to prolonged survival. (Immune Network 2003;3(3):188-200)

Key Words: Dendritic cells (DC), IL-12, NK cells, cytotoxic T lymphocytes (CTL), IFN-γ

Abbreviations: DC, dendritic cells; CCR, CC chemokine receptor; NKT, natural killer T; MIP-1α, macrophage-inflammatory protein-1α; SLC, secondary lymphoid-tissue chemokine.

Introduction

Dendritic cells (DCs) are highly effective antigen-presenting cells with the unique capability of inducing primary immune responses against tumor-associated antigens (1). DCs pulsed with synthetic tumor-derived MHC class I-restricted peptides, tumor lysates or tumor cell-derived RNA induced tumor-specific T
cell activation, host-protective and therapeutic antitumor immunity in mice and humans (2-5). The use of tumor-derived antigens in the form of tumor cell lysates or whole tumor cells has advantages in comparison with vaccinations against a single antigen, because so far only a few human tumor-associated antigens have been identified and immunity against a single antigen may be ineffective in tumors with heterogeneous cell populations. In addition, the high polymorphism of the human HLA system has presented problems in identifying and applying tumor-associated markers to be used as vaccines for cancer immunotherapy. In contrast, given that the induction of stronger cytotoxic T lymphocytes (CTL) responses appears to be a major goal of current cancer vaccine strategies, lysates-loaded DCs containing multiple known as well as unknown antigens that can be presented to T cells by both MHC class I- and class II-pathways provide the potential to induce efficient antitumor immune responses (6).

Recently, it has been shown that phagocytosis of apoptotic/necrotic tumor cells induced the maturation of DCs (7,8). DC maturation is characterized by down-regulation of antigen acquisition, increased expression of MHC and co-stimulatory molecules, production of proinflammatory cytokines, and altered expression of chemokine receptors (1,9). As they mature, DCs migrate to the T-cell areas of lymphoid organs, where antigen is presented to naive CD4+ and CD8+ T cells. Although a number of reports demonstrated that apoptotic/necrotic tumor cells might be a good source of tumor antigens for presentation to DCs in vitro and in vivo (7,10-12), the causal relation between the phenotypic characteristics of DCs that phagocytosed tumor cells, the immune responses induced by these DCs and their therapeutic vaccine efficacy against an established tumor have not been well studied. Importantly, it seems clearly not enough to phagocytose killed tumor cells to induce immune stimulatory signals leading to the breaking of tolerance to tumor antigens in vivo. Furthermore, there is a great need in controlled clinical trials for the establishment of key experimental parameters of the immune response by DC-based tumor vaccine, especially DCs loaded with necrotic tumor cells.

We have previously reported that vaccination of DC co-cultured with tumor cells in mice elicits an efficient protective antitumor activity against a challenge by colon cancer CT-26 cells (13). In this study, we prepared necrotic tumor cell-pulsed DCs for the therapeutic vaccination and investigated their phenotypic characteristics, the immune responses induced by these DCs, and therapeutic vaccine efficacy against colon carcinoma in vivo. Several parameters including phagocytosis of tumor cells, surface antigen expression, chemokine receptor expression, IL-12 production, and NK as well as CTL activation were assessed to characterize the immune response. Our data indicate that production of IL-12, chemokine receptor expression and NK as well as CTL activation may serve as major parameters to assess the effect of tumor cell-pulsed DC vaccine.

Materials and Methods

**Mice and tumor cell lines.** Five-to six-week-old female Balb/c mice purchased from the Genetic Resources Center, KRIIBB (Daejeon, Korea) were maintained under specific pathogen-free conditions until the experiments. The experiments employing the mice were performed in accordance with institutional guidelines. CT-26 cells were obtained from Seoul National University Hospital (Seoul, Korea). CT-26/NP cells, stable cell line of CT-26 cells transfected with influenza type A NP, were kindly provided from Dr. Y. C. Sung at Pohang University of Science and Technology (Pohang, Korea). Renca, renal adenocarcinoma cell line was obtained from Samsung Medical Center (Seoul, Korea). All tumor cells were cultured in RPMI containing 10% heat inactivated FBS, 100 U/ml penicillin, 100 µg/ml streptomycin, and 2 mM L-glutamine (Gibco BRL, Grand Island, NY, USA) at 37°C in 5% CO2. They were routinely checked for the absence of Mycoplasma contamination by using ELISA detection kit (Roche Diagnostics GmbH, Mannheim, Germany).

**Generation of dendritic cells from mouse bone marrow cells.** Dendritic cells were generated from bone marrow cells by following Dr. Inaba’s method with slight modification (14). In brief, bone marrow cells obtained from femurs and tibias of Balb/c mice were incubated with an antibody cocktail containing J1j.10 (anti-CD90), J11d (anti-CD11b), 3.168 (anti-CD8), GK1.5 (anti-CD4), RA3-3A1 (anti-B220), and M5/114.15.2 (anti-I-Ab,d,q & I-Ed,k) at 4 oC for 1 h. They were subjected to rabbit complement (Low-ToxR-M, Cedarlane, Onta-rio, Canada) according to the manufacturer’s instruction. To eliminate dead cells, cells were subjected to density centrifugation on Histopaque 1077 (Sigma, Saint Louis, MO, USA), and then washed twice with RPMI 1640 medium without serum. They (5×105 cells/well) were further incubated in culture medium supplemented with 10 ng/ml recombinant murine GM-CSF and IL-4 (Endogen, Woburn, MA, USA) in 24-well plates (Costar, Cambridge, USA). On days 2 and 4, non-adherent cells were discarded and culture media were replaced with fresh DC medium. On day 7, the non-adherent cells that had acquired typical dendritic cell morphology as identified on the phase contrast microscope were harvested by a gentle
swirling, and used in the subsequent experiments.

**Induction of tumor necrosis and uptake of necrotic tumor cells.** Tumor necrosis was induced by exposing the cells to one cycle of rapid freezing (liquid nitrogen) and thawing at 37°C in serum-free medium. Upon treatment all the cells became trypan blue positive. During incubation, these necrotic cells were degraded into fragments. Supernatants were obtained from necrotic cells by spinning at 1,500 rpm for 10 min. To analyze uptake of necrotic tumor cells by DCs, tumor cells were dyed green with PKH-67 (Sigma) according to the manufacturer’s protocol. Briefly, tumor cells and DCs were incubated with 2×10^6 M PKH-26 and PKH-67 at room temperature for 5 min, respectively, then stopped with the same volume of fetal bovine serum, and rinsed extensively with culture medium. PKH-26-labeled DCs at the ratio of 1:1 for 12–40 h at 4°C or 37°C. Phagocytosis of necrotic cells by DCs was defined by the percentage of double positive cells by flow cytometry. For fluorescence images, coverslips containing DCs stained with PKH-67 were mounted on glass slides after pulsing PKH-26-labeled necrotic tumor cells for 24 h and subjected to necrosis as described above, and then co-cultured with PKH-67-labeled tumor cells were subjected to necrosis as described above, then co-cultured with PKH-67-labeled DCs at the ratio of 1:1 for 12–40 h at 4°C or 37°C. Phagocytosis of necrotic cells by DCs was performed in 24-well transwell culture chamber (Costar) with 5.0 μm pore size as described previously (15). Briefly, after DCs were co-cultured with necrotic tumor cells at the ratio of 1:1 for 24 h, cells (5×10^5) were added to the top chamber in assay medium at a final volume of 100 μl, and recombinant chemokines (MIP-1α and MIP-3β; R & D Systems, Inc, Minneapolis, MN, USA) were diluted in the bottom chamber with assay medium to a final volume of 600 μl at appropriate concentrations. After the plates were incubated at 37°C in 5% CO₂ for 4 h, the cells in the bottom chamber were collected.

**Flow cytometric analysis of cell surface antigens.** To determine expression of chemokine receptors and cytokines using RT-PCR. Total RNA was isolated from each sample by the acid guanidinium thiocyanate phenol chloroform extraction method. cDNA was synthesized from 10 μg total RNA using ProSTAR™ kit (Stratagene, La Jolla, CA, USA), and used as the template for PCR (CCR1 and CCR7; 95°C for 1 min, 57°C for 1 min, and 72°C for 1 min, IL-12 p35, p40 and IFN-γ; 94°C for 1 min, 55°C for 1 min and 72°C for 1 min) using specific primers. To ensure the quality of the procedure, RT-PCR was performed on the same samples using specific primers for β-actin. PCR products were harvested and resolved on a 1.2% agarose gel containing ethidium bromide. They were quantified with Quantity One software supplemented with Gel Doc 2000 system (Biorad, Hercules, CA, USA). To compare relative expression, relative values were calculated as follows; (mean concentration×area) of CCR1, CCR7 or IFN-γ/(mean concentration×area) of β-actin. The sequence of each primer was as follows: mCCR1 (forward: 5'-TCTAGTGTTTCATCATTGGAGTGTG; reverse: 5'-GACGCACCGCGTTTGACTCTTCTTC), mCCR7 (forward: 5'-ACA-GCGCCCTCCAAGAAAGACCCG; reverse: 5'-T-GACGTCATAGGGGACATTTGACGGCTG), mIL-12 p35 (forward: 5'-TGGATCAATGAGACATCCACA; reverse: 5'-GATTCGAGACTGCTACACGTG), mIL-12 p40 (forward: 5'-ATGTTGTCTCCAGAAGCTTAAA; reverse: 5'-GCAAATTGCTACACGTTCC), and mIFN-γ (forward: 5'-AACGCTACACACTGATCT; reverse: 5'-TGTCATTGTAATGTGCGG). In vitro chemotaxis assay. An in vitro chemotaxis assay was performed in 24-well transwell culture chamber (Costar) with 5.0 μm pore size as described previously (15). Briefly, after DCs were co-cultured with necrotic tumor cells at the ratio of 1:1 for 24 h, cells (5×10^5) were added to the top chamber in assay medium at a final volume of 100 μl, and recombinant chemokines (MIP-1α and MIP-3β; R & D Systems, Inc, Minneapolis, MN, USA) were diluted in the bottom chamber with assay medium to a final volume of 600 μl at appropriate concentrations. After the plates were incubated at 37°C in 5% CO₂ for 4 h, the cells in the bottom chamber were collected.
The migrating cells were stained with trypan blue and counted five times. Each assay was performed in triplicate.

**Determination of cytotoxic activity.** Groups of BALB/c mice at 6–8 wks age were immunized subcutaneously (s.c.) in the right flank with 0.5 × 10^6 DCs, necrotic tumor cell-loaded DCs, necrotic tumor cells, or PBS. To examine CTL activity, splenocytes were harvested, stimulated in vitro with irradiated DCs pulsed with necrotic tumor cells for 3–4 days, and then used as effector cells. As effector cells for NK assay, splenocytes incubated for 12 h in the presence of IL-2 (10 ng/ml) were used. The target cells, such as CT-26, Renca, and YAC-1 were incubated with Na^51CrO_4 (100 Ci per 1 × 10^6, NEN-DuPont) for 90 min at 37 °C with shaking every 15 min. Radioisotope-labeled target cells were combined with various numbers of effector cells in 96-well U-bottomed culture plates (Corning, NY, USA) in triplicate. Wells containing only culture medium and target cells were used as the spontaneous ^51Cr release control, whereas those containing 1% SDS and target cells were used as the maximum release. After incubation for 4 h at 37 °C in 5% CO_2, supernatants (100 μl) were collected from each well, and the ^51Cr release was measured in a gamma counter (Wallac Inc., Gaithersburg, MD, USA). The percentage of specific lysis was calculated as follows:

\[
\text{percentage of specific lysis} = \left( \frac{\text{cpm of test} - \text{cpm of spontaneous release}}{\text{cpm of maximum release} - \text{cpm of spontaneous release}} \right) \times 100.
\]

Data shown are the mean of triplicate cultures.

**Immunization of mice.** To measure protective immunity, BALB/c mice were immunized s.c. with 0.5 × 10^6 DCs, necrotic tumor cell-loaded DCs, or necrotic tumor cells in the right flank. Control mice were immunized with the same volume of PBS. Seven days after immunization, mice were inoculated s.c. with 2 × 10^5 viable tumor cells. To measure therapeutic effect, 2 × 10^5 viable tumor cells were injected s.c. in the right flank. Vaccination was performed intratumorally or contra-laterally in the left flank twice on day 3 when tumors were noticeably developed, and on day 10. The size of tumors was measured in two perpendicular dimensions with a vernier caliper twice a week.

**Statistical analysis.** Data are expressed as mean±SD. Statistical analysis was performed using the Student t-test, two-sided. Differences were considered statistically significant with *P* < 0.05.

**Results**

**Uptake of necrotic tumor cells by DC.** We first examined whether DC could efficiently phagocytose necrotic tumor cells induced by a freeze-thawing by using flow cytometric analysis of fluorescence-labeled cells. After necrosis induction, all tumor cells were positively stained with trypan blue, and fragmented into small debris during 24 h incubation. When PKH-67-labeled DCs obtained from the bone marrow of syngeneic BALB/c mice were co-cultured with PKH-26-labeled necrotic tumor cells at the ratio of 1:1 at 37 °C, phagocytosis increased as the incubation time was prolonged (Fig. 1A). When the incubation was performed on ice for 24 h to discriminate uptake of necrotic tumor cells from non-specific binding, the phagocytosis percentage was the same as the one from spontaneous uptake (data not shown). The uptake of necrotic cells appeared to occur relatively fast since a significant enhancement of double positive cells (4.8% vs. 17.4%) was already observed after the incubation for 12 h. Optimum phagocytosis was consistently observed after 24 h (~30% of DCs), and the relative percentage of phagocytosis was scarcely changed at higher ratios of DC and tumor cell numbers (data not shown). Therefore, the co-culturing condition at the ratio of 1:1 for 24 h was used for subsequent experiments. This finding was further confirmed by confocal microscopic image analysis, showing that necrotic tumor cells were phagocytosed into the cytoplasm of DCs (Fig. 1B). Tumor cell fragments could be identified in DCs that had phagocytosed dye-labeled necrotic tumor cells (Fig. 1B, middle). Characteristic morphology of DCs with surface microvilli and long fillopodia was observed by scanning or transmission electron microscopy (Fig. 1C and 1D, left). After the physical contact with necrotic tumor cells in the tissue culture, microvilli on DCs displayed interaction with tumor debris at the initial phase of phagocytosis (Fig. 1C, middle). During the phagocytic processes, uptake of cell debris by microvilli and several phagocytic cups on DC surfaces were found (Fig. 1C, right). In the cytoplasmic area of DCs, engulfment of necrotic cells that showed a collapsed nucleus or cell fragments was shown by transmission electron microscopy (Fig. 1D, middle and right). These results indicate that DCs are able to phagocytose necrotic tumor cells through the recognition of specific molecules on necrotic cells. **Uptake of necrotic tumor cells induces phenotypical change of DCs.** Next, we investigated IL-12 production of DCs pulsed with necrotic tumor cells. Tumor cells were subjected to necrosis, and supernatant and/or pellets were separated by a centrifugation to test their respective effects on IL-12 production by pulsed DCs. IL-12 p70 production was induced by an addition of pellets of necrotic tumors that are absent from the supernatant (data not shown). However, in the presence of supernatant it was markedly increased by necrotic tumor cells in a time-dependent manner, indicating that maximal IL-12 production by DCs
required not only a soluble form of some tumor components, but also the uptake of necrotic cell pellets (Fig. 2A, left). This effect was not due to the contamination of endotoxin because tumor cells were confirmed to be endotoxin-free by the Limulus amoecyte lysate assay (BioWhittaker) and the treatment of polymyxin-B in the same culture did not affect IL-12 production. In addition, phagocytosis of necrotic tumor cells including their supernatant was not different from that of the necrotic cell pellets (data not shown). Therefore, necrotic tumor cells including their supernatant were used for further experiments. The next question was whether the induction of IL-12 production is associated with tumor-related components. We prepared necrotic cells of syngeneic splenocytes and added them to the DC culture to compare with necrotic tumor cells. In necrotic tumor cell-pulsed DCs, IL-12 production was repeatedly observed, whereas after uptake of necrotic splenocytes DCs were not able to produce IL-12 (Fig. 2A, right). RT-PCR analysis revealed that IL-12 p40 mRNA could be induced in DCs pulsed with necrotic
tumor cells, but not with necrotic normal splenocytes (Fig. 2A, right, inset).

IL-12 production is a phenotype typical of mature or activated DCs, with high levels of MHC class I and class II expression as well as other markers, including B7.1 and B7.2. Therefore, we investigated whether pulsing with necrotic tumor cells would affect the expression of surface antigens involved in antigen presentation and T cell stimulation. As shown in Fig. 2B, FACS analysis indicated that DCs have...
relatively high levels of B7.1, B7.2 and CD40 expression. When these cells were incubated with necrotic tumor cells, we consistently observed a slight up-regulation of antigen expression. A moderate increase in the level of MHC molecules was also induced (data not shown). These data suggest that necrotic tumor cells are able to mature DCs with respect to IL-12 production as well as increased surface expression of costimulatory molecules.

Chemokine receptor expression on DCs has been shown previously to correlate with cell’s maturation status and in vivo homing ability (16). The function of DCs to migrate first to the site of inflammation, and then to the draining lymph nodes can be explained in terms of a switch in the expression of chemokine receptors. To assess the impact of exposure to necrotic tumor cells on chemokine receptor gene expression by DCs, we performed RT-PCR to evaluate the transcription of mRNA encoding the CCR1 and CCR7. RT-PCR analysis using mRNA extracted from DCs revealed that expression of CCR1 was significantly reduced by co-culturing with necrotic CT-NP tumor cells compared with that in untreated DCs, whereas CCR7 mRNA expression

Figure 3. Peritoneal macrophages and RAW cells efficiently phagocytose necrotic tumor cells, but they are not active in producing IL-12. A. Peritoneal macrophages or RAW cells were co-cultured with necrotic tumor cells for 12 h at the ratio of 1 : 1. Thereafter, culture supernatants were collected, and examined for IL-12 p70 production by ELISA. Macrophages and RAW cells stimulated with LPS (1 μg/ml) were used as positive control. Data represent one of two experiments, each of which produced similar results (*P<0.05; **P<0.01; Student’s t-test). B. PKH-26-labeled (FL2) and PKH-67-labeled (FL1) cells were co-cultured at the ratio of 1 : 1 for 0, 12, 24, and 32 h. Cells were gated on the FL1 positive cells where FL2 positive cell population represents necrotic cells phagocytosed by macrophages or RAW cells.
was slightly increased (Fig. 2C). In fact, migration assay showed that tumor-pulsed DC had an enhanced ability to migrate responding to CCR7 ligand, MIP-3β (Fig. 2D). Thus, in terms of CCR1 expression and responsiveness to CCR7 ligand at least, pulsing with necrotic tumor cells induced a mature phenotype of DCs. These changes were also evident in DCs exposed to parent CT-26 cells, indicating that expression of NP protein in cells does not have any differential effect on CCR gene expression by DCs.

**Pulsing with necrotic tumor cells does not induce IL-12 production in macrophages.** We next questioned whether pulsing with tumor cells could induce enhanced IL-12 production in macrophages. When other antigen-presenting cells including peritoneal macrophages and RAW 264.7 cell line were tested, we observed markedly increased production of IL-12 by LPS stimulation, but not any significant increase of IL-12 production by co-cultures with necrotic tumor cells (Fig. 3A). In order to examine the possibility that the low level of IL-12 production was due to the defect of phagocytic activity, PKH-67-labeled antigen-presenting cells were mixed with PKH-26-labeled necrotic cells and then analyzed by flow cytometry. As shown in Fig. 3B, the phagocytosis of necrotic tumor cells by primary macrophages and the macrophage cell line occurred more efficiently than the one by DCs. Thus, we concluded that IL-12 production in antigen-presenting cells does not correlate with phagocytic activity and the increased IL-12 production after contact with necrotic tumor cells is a characteristic feature of DCs.

**Induction of NK cell activity by vaccination with necrotic tumor cell-loaded DCs.** It was recently reported by our group and others that murine DCs are capable of modulating innate immunity by stimulating NK cells through cell-to-cell contact (13,17). Thus we examined NK cell stimulation by tumor-pulsed DCs in two different experiments; determining IFN-γ production after co-culturing with purified splenic NK cells in vitro and NK cell activity after DC vaccination in vivo. As shown in Fig. 4A, NK cells purified with DX-5 beads or DCs alone were not able to produce IFN-γ, whereas co-culturing them exhibited significant production of IFN-γ. However, DCs pulsed with necrotic tumor cells induced a much larger IFN-γ secretion than non-treated DCs. IFN-γ production was not observed in the culture supernatant from tumor cell-pulsed DCs, suggesting that at least DCs pulsed with necrotic tumor cells are not producers of IFN-γ. In addition, although we did not detect IL-12 after the washing of tumor-pulsed DC, the co-culture of DCs with NK cells induced IL-12 production that was significantly inhibited by a transwell culture (Fig. 4B). IFN-γ secretion was confirmed again in co-culture of DCs and NK cells (Fig. 4C), which was blocked by a transwell culture or neutralizing anti-IL-12 antibody. These results show that NK cells can be stimulated to produce IFN-γ by tumor-pulsed DCs via cell-to-cell contact as well as soluble factor such as IL-12. Our results, however, could not rule out the possibility that after contact with NK cells, DCs might be converted to IFN-γ producers.

In order to examine the NK cell activity of
splenocytes in vivo, 51Cr release assay was performed 2 or 5 days after DC vaccination using NK-sensitive YAC-1 cells as targets. As expected, DCs pulsed with necrotic tumor cells markedly induced NK cell-mediated cytolysis after a short-term restimulation of vaccinated splenocytes with IL-2 in vitro (Fig. 5A). NK cell activity was not due to contamination of cytotoxic T cells because target cell killing was not influenced by a depletion of CD8-positive T cells using specific mAb and complement (Fig. 5B). Similar to NK cell activity, vaccination with tumor-pulsed DCs influenced INF-γ mRNA expression in splenocytes, which was still elevated 5 days after vaccination when compared to vaccination with DC alone (Fig. 5C).
However, consistent with IFN-γ production, vaccination with DC alone exhibited to a certain extent enhanced NK cell activity though it was significantly lower compared to the vaccination with tumor-pulsed DCs. Together, these results suggest that necrotic tumor cell-loaded DCs stimulate NK cells to produce IFN-γ as well as induce NK cell activation to efficiently kill target cells.

**Induction of tumor-specific CTL by vaccination with necrotic tumor cell-loaded DCs.** We further examined the induction of CTL activity against tumor cells after vaccination with necrotic tumor cell-loaded DCs. The spleen cells harvested from vaccinated mice were restimulated *in vitro* with DCs pulsed with necrotic tumor cells and their cytolytic activities were determined by ⁵¹Cr release assay. As shown in Fig. 5D, vaccination with necrotic tumor cells alone did not induce cytolytic activity against CT-NP target tumor cells. However, vaccination with necrotic tumor cell-loaded DCs induced markedly enhanced cytotoxic activities that were almost completely inhibited by an *in vitro* treatment with CD8-specific mAb and complement, suggesting cytotoxic T lymphocyte-mediated target killing (data not shown). It was notable that similar to NK activity induction, moderate cytolytic activity was also observed after vaccination with DCs alone. When cytotoxic activities were tested against different target cells, the cytosis of syngeneic Renca cells was marginal, whereas cytotoxic activities against parent CT-26 tumor cells were not significantly different from those against CT-NP cells (Fig. 5D, right). These results indicate that vaccination of tumor cell-pulsed DCs is able to induce efficient tumor-specific cytotoxic T cell responses.

**Therapeutic effect of vaccination with necrotic tumor cell-loaded DCs.** To investigate the *in vivo* efficacy of immunization with necrotic tumor cell-loaded DCs, both protective and therapeutic vaccinations were performed. When necrotic tumor cell-loaded DCs as a prophylactic vaccine were injected s.c. 7 days before tumor challenge at the same injection site, we consistently observed tumor protection. In addition, even after rechallenging both CT-NP and parent CT-26 cells, tumor formation was not observed and this protective immunity lasted for at least 4 months of observation (data not shown). To evaluate the therapeutic efficacy of necrotic tumor cell-loaded DCs, viable 2×10⁵ tumor cells were inoculated in the right flank. After 3 days, when tumors were measurable, mice were immunized intratumorally with PBS buffer, necrotic tumor cells, DCs or tumor cell-loaded DCs. A second immunization was done 7 days after the first one. As shown in Fig. 6, vaccination with necrotic tumor cell-loaded DCs significantly increased the number of tumor-free mice and reduced tumor size compared with any other vaccinations when tumor size was determined on day 30 after first immunization. Importantly, a substantial reduction of tumor size was also observed in mice treated with DCs alone. Thus, we questioned whether DCs administered by intratumoral injection might behave like tumor cell-pulsed DCs *in vitro*. To test this, therapeutic vaccination was done in the opposite flank to block encounter of DCs and tumor cells. While a significant therapeutic effect was repeatedly obtained by the contra-lateral subcutaneous injection of necrotic tumor cell-loaded DCs, tumors with a similar growth rate to the control tumors were formed in mice injected contra-laterally with DCs alone (Fig. 6, inset). These results indicate that intratumoral injection of DC caused DC to uptake tumor cells *in vivo* similar to tumor-pulsed DCs and resulted in the partial inhibition of tumor growth. Furthermore, when viable tumor cells were rechallenged on the opposite left flank in completely cured mice, tumors did not form over a 30- to 40-day period, indicating that the therapeutic vaccination induced long-lasting immune memory responses against tumor cells (data not shown).
Discussion

Although a number of strategies for cancer therapy using DC-based vaccine have previously been reported, the most efficient method of vaccine preparation suitable for individual tumor types still remains to be explored. Especially, the establishment of key parameters on the immune response by DC vaccine, loaded \textit{ex vivo} with well-characterized tumor antigens, their peptides or tumor materials, is of crucial importance and necessary for the precise prediction and evaluation of vaccine efficacy. In this study, we investigated the uptake of necrotic tumor cells by dendritic cells (DCs) by means of flow cytometric, confocal and electron microscopic analysis. Although the relative ratio of tumor uptake in DCs was lower than that in primary macrophages or the macrophage cell line, they were capable of picking up necrotic tumor cells effectively. Most notably, while macrophages could not be activated to produce heterodimeric cytokine IL-12, DCs pulsed with necrotic tumor cells produced markedly increased amount of IL-12 or IL-12 p40 transcript. It seems highly unlikely that the enhanced IL-12 production in DCs is due to infection of tumor cell lines as recently demonstrated (18), because \textit{Mycoplasma} contamination of them was routinely checked and, more importantly, even after a treatment of them with cyporin, IL-12 secretion in necrotic tumor-pulsed DCs was detected to a similar extent (data not shown).

IL-12 is a central component of the cellular immune response as a surrogate marker for type 1 T cell response and used as an indicator of DC activation (19). Increased IL-12 production by DCs has been suggested as a crucial step in the process of CTL priming (20,21). IL-12 induces IFN-\(\gamma\) secretion by activated T cells, NK cells and NKT cells and is required for optimal adaptive and innate responses against tumors (22,23). Indeed, IL-12 and IL-18 have been reported to promote NK cell cytotoxicity through up-regulation of NK cell-derived IFN-\(\gamma\) production (24,25). Our observation that DCs pulsed with necrotic tumor cells produce a significant amount of IL-12, and stimulate NK cell cytolytic activity and IFN-\(\gamma\) secretion is consistent with this concept. However, IL-12 production may not be enough to result in NK cell cytotoxicity and IFN-\(\gamma\) production, because DCs showing low levels of IL-12 production also have the ability to induce increased production of IFN-\(\gamma\) in NK cells and the IFN-\(\gamma\) production is not induced by culture supernatants of tumor-pulsed DCs. In fact, tumor cell-pulsed DCs prepared for immunization \textit{ex vivo} appeared to be a weak IL-12 producer after washing of DCs for injection as shown in Fig. 4B. Thus antigen-pulsed DC and NK interaction is more important for IL-12 production and subsequent NK cell activation than necrotic tumor cell pulsing itself, and supposed to be relevant in the control and the amplification of the antitumor immune response. Moreover, immunization with DC alone could induce NK cell cytotoxicity, albeit to a lesser extent. It is important that while contralateral injection of tumor-pulsed DCs induces a significant therapeutic effect, DCs alone do not show any tumor suppressive effect, indicating that although DCs alone can, at least in part, activate NK cells, they are not sufficient for the induction of systemic antitumor immune responses.

It was recently reported that murine or human DCs were capable of modulating the innate immunity by stimulating NK cells through cell-to-cell contact (26,28). However, the requirement of DC-derived IL-12 during the development of cellular immunity is somewhat controversial since DC-derived IL-12 is not required for the generation of protective immunity against melanoma and treatment of DCs with anti-IL-12 antibody does not block the enhancement of NK cell-mediated cytolysis by DCs (27,28). Grufman et al. indeed demonstrated that it is indeed possible to generate good CTL responses by immunizing B6 mice with IL-12-deficient DC (23). They suggested that either DC-derived IL-12 may still be required, but the inoculated DC may be engulfed and presented by host APC or MHC class I-peptide complexes may be transferred from the inoculated cells to host APC, which are able to produce IL-12. Alternatively, neighboring cells may provide IL-12 required for optimal responses. In the present study, we demonstrated that intratumoral injection of DCs alone had a minor but substantial effect on the inhibition of tumor formation. Thus, the results showing that the ability of DCs to produce IL-12 correlates with NK cytotoxicity, IFN-\(\gamma\) production and tumor-specific T cell cytotoxicity are consistent with this notion. Recent work from our group has shown that NK cell depletion \textit{in vivo} correlates closely with the weakness of DC vaccine-induced protective immunity and tumor rejection (13). In another therapeutic setting using the B16 model, immunization with the DC vaccine could inhibit lung metastasis of tumor cells, whereas the depletion of NK cells before DC vaccination abrogated the therapeutic effect (manuscript in preparation). Therefore, one might expect that DCs might acquire nominal tumor antigens and stimulate tumor-specific CTLs by phagocytosis of dying tumor cells at the tumor site, presenting them in both MHC class I and class II molecules (29-32). Taking this into consideration, we speculate that IL-12 production in tumor-pulsed DCs or DCs administered into tumor sites after contact
with NK cells may enhance NK cell and tumor-specific T cell activity, thereby increasing the in vivo effect of DC immunization. Moreover, NK cell production of IFN-γ via the contact with DCs might be a critical factor in the development of adaptive T cell immunity and memory after the NK cell-mediated innate immune responses to tumors.

Recent studies have demonstrated that the migratory capability of DCs is dictated by the change in responsiveness of DCs to various chemokines during their development and maturation (16,33). We investigated the expression of chemokine receptors on DCs and found that CCR1 mRNA expression is markedly downregulated by the pulsing with necrotic tumor cells. Although CCR7 appeared to be slightly down-regulated by the pulsing with necrotic tumor cell-pulsed DCs. The down-regulation of the receptor for the inflammatory chemokines including RANTES and MIP-1q and the high level of receptors for chemokines such as MIP-3β and SLC that are expressed in secondary lymphoid organs may help DCs leave the sites of inflammation and migrate to regional lymph nodes (34). These results suggest that necrotic cell-loaded DCs can activate naive T cells and efficiently migrate into draining lymph nodes through the exchanges in the CCR expression during immune responses.

Interestingly, Tong et al. have shown recently that intratumoral injection of DCs combined with systemic chemotherapy leads to complete tumor regression in the treated mice and induces long-term antitumor immunity (32). They suggest that local DC-based therapy may be synergistic with chemotherapy in enhancing antitumor host responses. However, because the direct comparison of the antitumor effect between tumor-pulsed DCs and DCs in combination with systemic chemotherapy has not been done, the relative antitumor efficacy of them remains to be obscure, although the combination of chemotherapy with DC vaccination is meaningful for clinical settings. In our study, DCs pulsed with necrotic tumor cells that had been subjected to one cycle of freeze thawing induced tumor specific immune responses and regression of established tumors. It is thus expected that concurrent treatment with drugs inducing tumor cell killing is able to increase the therapeutic effect of tumor-pulsed DC vaccines. Moreover, the systemic addition of IL-12 may be helpful for potentiating antitumor immune responses induced by DC vaccine as demonstrated in a recently published report (35).

In summary, we have demonstrate in the present study that DCs pulsed with necrotic tumor cells, most importantly after the interaction with NK cells, are converted to IL-12 producers and IL-12 is required in responses induced by antigen-pulsed DCs, including NK cell and tumor-specific T cell activation. Furthermore, our results suggest that necrotic cell-loaded DCs could activate naïve T cells and efficiently migrate into draining lymph nodes through the exchanges in the CCR expression during immune responses. Repeated application of tumor cell-loaded DCs resulted in eradication or strong inhibition of pre-established tumors followed by long-lasting immune memory. Our data thus indicate that the production of IL-12, changes of chemokine receptor expression and NK as well as CTL activation may serve as major parameters in assessing the effect of tumor cell-pulsed DC vaccine.

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