Limited Amounts of Dendritic Cells Migrate into the T-Cell Area of Lymph Nodes but Have High Immune Activating Potential in Melanoma Patients

Pauline Verdijk,1 Erik H.J.G. Aarntzen,1,2 W. Joost Lesterhuis,2 A.C. Inge Boullart,1,2 Ellemieke Kok,1 Michelle M. van Rossum,3 Simon Strijk,4 Femke Eijckeler,5 Johannes J. Bonenkamp,6 Joannes F.M. Jacobs,1 Willeke Blokx,5 J. Han J.M. van Krieken,5 Irma Joosten,7 Otto C. Boerman,8 Wim J.G. Oyen,8 Gosse Adema,1 Cornelis J. A. Punt,2 Carl G. Figdor,1 and I. Jolanda M. de Vries1,2,9

Abstract

Purpose: The success of immunotherapy with dendritic cells (DC) to treat cancer is dependent on effective migration to the lymph nodes and subsequent activation of antigen-specific T cells. In this study, we investigated the fate of DC after intradermal (i.d.) or intranodal (i.n.) administration and the consequences for the immune activating potential of DC vaccines in melanoma patients.

Experimental Design: DC were i.d. or i.n. administered to 25 patients with metastatic melanoma scheduled for regionallymph node resection. To track DC in vivo with scintigraphic imaging and in lymph nodes by immunohistochemistry, cells were labeled with both [111In]-indium and superparamagnetic iron oxide.

Results: After i.d. injection, maximally 4% of the DC reached the draining lymph nodes. When correctly delivered, all DC were delivered to one or more lymph nodes after i.n. injection. Independent of the route of administration, large numbers of DC remained at the injection site, lost viability, and were cleared by infiltrating CD163+ macrophages within 48 hours. Interestingly, 87 ± 10% of the surviving DC preferentially migrated into the T-cell areas, where they induced antigen-specific T-cell responses. Even though more DC reached the T-cell areas, i.n. injection of DC induced similar antigen-specific immune responses as i.d. injection. Immune responses were already induced with <5 × 10^5 DC migrating into the T-cell areas.

Conclusions: Monocyte-derived DC have high immune activating potential irrespective of the route of vaccination. Limited numbers of DC in the draining lymph nodes are sufficient to induce antigen-specific immunologic responses.

In cancer patients, the immune system has not been able to establish an effective immune response against the tumor. Immunotherapy aims at educating the immune system to generate effective tumor-specific immune responses. Dendritic cells (DC) are specialized antigen-presenting cells that can induce de novo antitumor responses and are excellent candidates for cell-based immunotherapy. Whereas many DC-based clinical studies for the treatment of cancer have shown the feasibility and safety of DC vaccinations (reviewed in refs. 1, 2), the clinical efficacy of the therapy still needs to be improved (3). One important factor that determines the outcome of DC therapy is the delivery of the vaccine to immune-reactive sites, such as lymph nodes, and, more specifically, to the T-cell rich area, the paracortex. To exert their action, DC must closely encounter and interact with T cells. In addition, it has now been generally accepted that the site at which T-cell priming occurs significantly influences the homing characteristics of the effector cells (4–8). Therefore, the route of administration may be of crucial importance. It might be beneficial to combine different routes of administration; for example, in mice metastatic-like lung lesions were controlled by i.v. immunization and only partially by s.c. immunization (6). However, because DC migration to lymph nodes is very poor after i.v. injection (4), we study here only the intradermal (i.d.) and intranodal (i.n.) route. Currently, for treatment of solid tumors, the i.d. or s.c. administration of DC is most frequently used (91 clinical trials), followed by i.v. (47 trials) and i.n. (9 trials) injection (www.mmri.mater.org.au).

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Authors’ Affiliations: Departments of 1 Tumor Immunology, 2 Medical Oncology, 4 Dermatology, 5 Radiology, 6 Pathology, 7 Surgery, 8 Blood Transfusion and Transplantation Immunology, 5 Nuclear Medicine, and 8 Pediatric Hemato-Oncology, Nijmegen Centre for Molecular Life Science, Radboud University Nijmegen Medical Centre, Nijmegen, The Netherlands

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Requests for reprints: I. Jolanda M. de Vries, Department of Tumor Immunology, Radboud University Nijmegen Medical Centre, P.O. Box 9101, 6500 HB Nijmegen, The Netherlands. Phone: 31-24-3617600; Fax: 31-24-3540339; E-mail: j.devries@ncmls.ru.nl.

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Scintigraphic imaging of cancer patients injected with DC labeled with $^{111}$In-iodinum or $^{99m}$Tc-tecnethium showed that only after i.d. or s.c. (9–16), i.n. (9, 11, 17), and intra-lymphatic (11, 18) injection did DC migrate to the draining lymph node regions. More substantial evidence was obtained in our previous study, in which DC were labeled with both $^{111}$In and paramagnetic iron oxide particles (SPIO), which is a suitable contrast agent for magnetic resonance imaging (17).

We showed that after i.n. injection, DC were indeed present in the injected node and had migrated to nearby lymph nodes. However, these images do not show how many cells actually reach the T-cell area and provide no information on the phenotype and quality of the injected cells and whether the label is still present in the administered cells. SPIO-labeled DC can be easily visualized in resected lymph nodes and allow a detailed analysis of the fate and local immune effects of the injected DC in the targeted lymph nodes. We show that, although the majority of nonmigrating DC at the injection site are phagocytosed by macrophages, part of the SPIO-labeled cells migrate into the T-cell areas, exhibit a DC phenotype, and activate specific T cells. Because the immune responses in both groups of patients were comparable, there was no advantage of i.n. injection over i.d. administration.

**Patients, Materials, and Methods**

**Patients.** Eligibility criteria included stage III and IV melanoma according to the American Joint Committee on Cancer criteria (19), HLA-A2.1 phenotype, and melanoma expressing the melanocyte-associated antigens gp100 and tyrosinase. The study was approved by our Institutional Review Board, and written informed consent was obtained from all patients.

**Treatment schedule.** At day 7, peripheral blood mononuclear cells (PBMC) were obtained by leukapheresis for DC culturing. DC loaded with both keyhole limpet hemocyanin (KLH) and tumor-associated antigen (TAA) peptides were injected either i.n. into a lymph node of the region that was to be resected or i.d. in close vicinity of this lymph node region. On day 0, patients received either an i.n. injection of $^{111}$In-labeled DC ($7.5 \times 10^6$) mixed with SPIO-labeled DC ($7.5 \times 10^6$; total volume 100 $\mu$L) directly into a lymph node or an i.d. injection of DC labeled with both $^{111}$In and SPIO ($15 \times 10^6$; total volume 100 $\mu$L). i.n. injections were done under ultrasound guidance by an experienced radiologist. Scintigraphic imaging of the lymph node region was done 30 min after i.n. injection, and 24, 48, and/or 48 h after i.n. or i.d. injection, before dissection of the regional lymph nodes. Two patients received an i.n. injection in both the left and the right lymph node region.

Radiolabeled lymph nodes were dissected from the surgical specimen under the guidance of a $\gamma$-probe (Europrobe; Eurorad) and then fixed in Unifix (Klinipath). The lymph nodes were embedded in paraffin and were processed for histology and immunohistochemistry. Patients received three more vaccinations at days 14, 28, and 42.

**DC culture and labeling.** DC were generated from adherent PBMC by culturing in the presence of interleukin 4 (500 U/mL) and granulocyte-macrophage colony-stimulating factor (800 U/mL; both Cellgenix). On day 4, DC were loaded with the control antigen KLH (10 $\mu$g/mL; Calbiochem) and were labeled with SPIO by adding 100 $\mu$g Ferumoxide/mL (Endorem; Laboratoire Guerbet) for 3 d after the onset of DC culturing (17). On day 5, DC were matured with autologous monocyte-conditioned medium supplemented with prostaglandin E$_2$ (10 $\mu$g/mL; Pharmacia & Upjohn) and 10 ng/mL recombinant tumor necrosis factor-$\alpha$ (Cellgenix) for 48 h as described previously (20, 21).

After maturation, 60 ± 23% of the SPIO DC and 77 ± 17% of unlabeled DC expressed high levels of CD83, and <10 ± 8% of all DC expressed residual levels of CD14, analyzed by fluorescence-activated cell sorting analysis as described below. DC were pulsed with the melanoma peptides gp100:154-167, gp100:280-288, and tyrosinase:369-376 as described previously (9). Mature DC were labeled with $^{111}$In-oxine (Coviden) in 0.1 mL Tris-HCl (pH 7.0) for 15 min at room temperature as described previously (9, 20), resulting in 4 MBq activity per 7.5 $\times 10^6$ cells.

**Scintigraphic imaging.** *In vivo* and *ex vivo* planar scintigraphic images ($256 \times 256$ matrix, 174 and 247 keV $^{111}$In photon peaks with 15% energy window) of the injection depot and the corresponding lymph node basin were acquired with a $\gamma$-camera (Siemens ECAM) equipped with medium energy collimators at day 0 and day 2. Migration was quantified by region of interest analysis of the individual nodes visualized on the images and expressed as the relative fraction of $^{111}$In-labeled DC that had migrated from the injection depot to successive lymph nodes after 2 d.

**Iron staining and immunohistochemistry of histopathologic sections.** Sections (5 $\mu$m) of the resected radiolabeled lymph nodes were stained with H&E or with Prussian blue to detect SPIO-labeled cells. Slides were stained with 2% potassium hexacyanoferrate (II)-trihydrate in 0.2 mol/L HCl for 15 min and were counterstained with 0.05% nuclear fast red in 5% aluminum sulphate. Immunohistochemical stainings were done on paraffin-embedded tissue sections with the use of monoclonal antibodies. Paraffin sections were dewaxed and rehydrated. All reactions were done at room temperature unless stated otherwise. Endogenous peroxidase activity was blocked by incubation in PBS containing 3% H$_2$O$_2$ for 30 min. After rinsing with PBS, antigen retrieval consisted of microwave boiling in either 10 mmol/L sodium citrate buffer (pH 6.0) or 10 mmol/L EDTA/1 mmol/L Tris buffer (pH 8.0 or 9.0) for 10 min, depending on the primary antibody. After boiling, slides were allowed to cool down for at least 2 h. After rinsing with PBS, slides were pretreated with 20% normal horse serum for 10 min to reduce nonspecific staining. All sera and antibodies were dissolved in PBS with 1% bovine serum albumin. Subsequently, slides were incubated in a humidity chamber with the primary antibody at 4°C for 16 to 20 h. The following primary monoclonal antibodies were used: CD4, CD8 (both from Beckman Coulter), CD25, CD69, CD83, and CD163 (Novocea-stra). The avidin-biotin complex (Vector) method was used for visualization with 3,3'-diaminobenzidine hydrochloride solution. Staining of the 3,3'-diaminobenzidine substrate was intensified with a 0.5% copper sulfate solution. Slides were counterstained with hematoxylin solution or nuclear fast red. Sections were analyzed by microscopy (Zeiss Axioskop 2 plus; Zeiss or Leica DMLB microscope; Leica) with the use of ProgRes CapturePro (Jenoptik) or Leica IM500 (Leica) software.

**Humoral responses to KLH.** Antibodies against KLH were measured in the serum of vaccinated patients by ELISA (22). Microtiter plates (96 wells) were coated overnight at 4°C with KLH (25 $\mu$g/mL in PBS per well). After washing the plates, different concentrations of patient serum (range, 1 in 100 to 1 in 500,000) were added for 1 h at room temperature. After extensive washing, specific antibodies (IgG, IgG1, IgG2, and IgG4) labeled with horseradish peroxidase were allowed to bind for 1 h at room temperature. Peroxidase activity was revealed with the use of 3,3',5,5-tetramethylbenzidine as substrate and was measured in a microtiter plate reader at 450 nm.

**Proliferation assay.** Cellular responses against KLH were measured in a proliferation assay. Briefly, PBMC isolated from blood samples taken...
before each DC vaccination were plated in a 96-well tissue culture microplate with or without KLH. After 4 d of culture, 1 μCi per well of tritiated thymidine was added for 8 h, and incorporation of tritiated thymidine was measured in a β-counter.

**Isolation of lymph node DC and T cells.** Cell suspensions were made from resected radiolabeled lymph nodes from three patients. Lymph node tissue was cut into small fragments in Hanks Balanced Salt Solution medium (GIBCO) with 50 μg/ml collagenase type 1A, 10 μg/ml DNase, and 1 μg/ml trypsin inhibitor (Sigma Chemical Co.). The fragments were incubated for 30 min at 37°C. For the isolation of DC–T cell rosettes, the large fragments were left to settle down, and the supernatant was transferred to a fresh tube. SPIO-containing DC were isolated with the use of a Dynal MPC magnet. Cells were washed with RPMI supplemented with 7% human serum. The isolated SPIO DC were spun onto microscope slides and were stained with H&E. From two patients, lymph node cells were cultured for 4 to 6 wk in RPMI/7% human serum and 200 IU/ml of interleukin 2. KLH-specific T-cell proliferation was measured in a proliferation assay. Fresh autologous PBMC were loaded with KLH overnight, irradiated, and used as stimulator cells for lymph node T cells (ratio, 1:1). After 4 d of culture, 1 μCi per well of tritiated thymidine was added for 8 h, and incorporation of tritiated thymidine was measured in a β-counter.

**Delayed-type hypersensitivity.** One to 2 wk after the four DC vaccinations, a delayed-type hypersensitivity (DTH) skin test was done. Briefly, DC pulsed with peptides and DC pulsed with peptides plus KLH (2 × 10^5 DC each) were injected i.d. into the skin of the back of the patients at four different sites. The diameter (in millimeters) of induration was measured after 48 h, and punch biopsies (6 mm) were obtained under local anesthesia. Biopsies were cut in half; one part was frozen directly in liquid nitrogen for immunohistochemistry, and the other part was cut into small pieces and cultured in RPMI/7% human serum supplemented with interleukin 2 (100 U/ml). Every 7 d, half of the medium was replaced by fresh interleukin 2—containing RPMI/7% human serum. After 2 to 4 wk of culturing, the T cells were tested for antigen recognition or tested for tetramer binding. For tetramer binding, the PBMC of T cells were incubated with allopregocyanin-labeled tetrameric MHC containing the gp100:154-167, gp100:280-288, or tyrosinase:369-376 peptide (Sanquin) for 1 h, and then were washed and analyzed by flow cytometry.

**Antigen-recognition assay.** Antigen recognition was determined by the production of cytokines by DTH-derived cells in response to T2 cells pulsed with the indicated peptides or BLM (a melanoma cell line expressing HLA-A2.1 and with no endogenous expression of gp100 and tyrosinase), transfected with control antigen G250, or with gp100 or an allogenic HLA-A2.1-positive, gp100-positive, and tyrosinase-positive tumor cell line (MEL624). Cytokine production was measured in supernatants after 16 h of coculture by the cytometric bead array (Th1/Th2 Cytokine CBA; BD Pharmingen).

**Statistics.** Data were analyzed with the use of unpaired Student’s t test; P values <0.05 were considered to be statistically significant.

## Results

**Biodistribution of ^111^In/SPIO DC after i.d. and i.n. vaccination.** To study the distribution of the DC vaccine after i.d. and i.n
injection in patients with metastatic melanoma, DC were labeled with $^{111}$In and SPIO for the first vaccination and injected in (the vicinity of) a lymph node region 24 to 72 hours before scheduled lymph node resection. Delivery of the DC to skin-draining lymph nodes after i.d. ($n = 12$) and i.n. injection ($n = 13$) was monitored by scintigraphic imaging of the lymph node region (Table 1). After i.d. administration, never >4% migration was observed (Fig. 1A; mean, $1.1 \pm 1.1\%$). In contrast, the percentage of cells reaching nearby lymph nodes after i.n. administration was highly variable and ranged from 0% to 56% (Fig. 1B; mean, $10.6 \pm 17.5\%$). In five patients, distribution to more than one lymph node had already taken place within the first 30 minutes after injection, indicating that DC had spread via the lymphatic system during injection. Migration to subsequent lymph nodes increased in the next 24 to 72 hours. In six patients, DC remained localized at the injection site after i.n. administration was indeed confirmed with magnetic resonance imaging as previously described (17). In the resected lymph node basin from the sixth patient, an $^{111}$In-positive lymph node was isolated, showing that the injection was correct but that the cells had not migrated to nearby lymph nodes. Taken together, although maximally 4% of the DC reached the draining lymph nodes after i.d. injection, all DC were delivered to one or more lymph nodes after i.n. injection when correctly delivered.

Numerous $^{111}$In/SPIO DC die at the injection site and are phagocytosed by macrophages. Because the presence of $^{111}$In/SPIO DC inside lymph nodes does not necessarily imply that DC actually reach the T-cell area, we analyzed the dissected radio-labeled lymph nodes by immunohistochemistry to visualize $^{111}$In/SPIO DC. Lymph nodes were obtained from eight and nine patients that received i.d. or i.n. injections, respectively. In addition, biopsies from the i.d. injection site were obtained and analyzed in the same way. Although single $^{111}$In/SPIO DC were present in the sinuses and paracortex of draining lymph nodes, a large depot of SPIO$^+$ cells was found at the site of injection, as was expected from the scintigraphs. This was not only observed after i.d. vaccination (Fig. 1C) but also after i.n. injection (Fig. 1D). In lymph nodes from patients in which immediate redistribution of the DC after injection was observed, $^{111}$In/SPIO DC also accumulated in the sinuses of subsequent nodes (not shown).

To verify whether the SPIO$^+$ cells at the i.d. or i.n. injection site were indeed DC and not phagocytes that have taken up label, lymph node sections were stained for the DC marker CD83 and the macrophage marker CD163 (23). This revealed that in the dermis, only part of the SPIO$^+$ cells in the injection site expressed CD83 (Fig. 2A). In the CD83-negative...
areas, all SPIO\(^+\) cells were positive for the macrophage marker CD163 (Fig. 2A and B). Staining with H&E revealed that in areas with the most CD83\(^+\) SPIO DC, cells were enlarged and exhibited pale-pink nuclei, typical for necrotic cells. At the same time, the SPIO\(^+\) cells in the CD163\(^+\) area seemed viable, exhibiting regular hematoxylin staining of the nuclei (Fig. 2B), suggesting that macrophages had infiltrated the injection site and had phagocytosed dead \(^{111}\)In/SPIO DC. Comparable results were found after i.n. injection (Fig. 2C and D). Most of the SPIO\(^+\) cells at the site of injection still expressed CD83 at 24 hours (one patient), but numerous small macrophages containing no or little SPIO were present in between enlarged SPIO\(^+\) cells (Fig. 2C, top). In lymph nodes resected after 48 hours, CD83 expression was rare and was mainly found in the center of the depot, whereas macrophages were present throughout the injection site (Fig. 2C, bottom). In H&E stainings, SPIO is visible as brown granula. In immunostainings, SPIO was visualized by Prussian blue staining, and nuclei were stained with nuclear fast red.

Indeed, SPIO\(^+\) cells were predominantly found in the T-cell areas (93 ± 9\%) and sometimes in the subcapsular sinuses (Fig. 3A, left). Of all the SPIO\(^+\) cells in the paracortex, 85 ± 13\% were CD83\(^+\) DC (Fig. 3B), indicating that mainly viable \(^{111}\)In/SPIO DC and only a few SPIO\(^+\) macrophages had migrated to the draining lymph nodes and into the T-cell areas.

After i.n. injection, more SPIO\(^+\) cells were dispersed over the lymph nodes than after injection in the dermis. SPIO\(^+\) cells were present in high numbers both in the sinuses and the paracortex. We estimated the total number of SPIO\(^+\) cells in the paracortex in single lymph node sections after i.n. injection to be 10-fold to 30-fold higher than after i.d. administration. However, many SPIO\(^+\) macrophages were also present both in the injected node and in nearby lymph nodes. Analysis of the paracortex of all lymph nodes revealed that 54 ± 21\% of all SPIO\(^+\) cells in the paracortex were DC. Within each section, the ratio between \(^{111}\)In/SPIO DC and SPIO\(^+\) macrophages was variable and correlated with the distance from the depot. In close vicinity of the injection depot, 5\% of the SPIO\(^+\) cells were CD83\(^+\) \(^{111}\)In/SPIO DC, whereas at more distant sites, virtually all of the SPIO\(^+\) cells expressed CD83 (not shown). Quantitative analysis of the total number of SPIO\(^+\) cells was only possible in sections of the lymph nodes where DC had migrated to. Quantification of CD83\(^+\) and CD163\(^+\) expression revealed that only 35 ± 26\% of all SPIO\(^+\) cells were CD83\(^+\) DC (Fig. 3A, right, and C). Interestingly, 87 ± 10\% of these CD83\(^+\) \(^{111}\)In/SPIO DC were found in the T-cell area, indicating that \(^{111}\)In/SPIO DC that do survive preferentially migrate into the paracortex.

**DC in T-cell area are fully functional and activate T cells.** We next analyzed whether \(^{111}\)In/SPIO DC that had migrated into
the paracortex of the (draining) lymph nodes were still functional and able to interact with resident T cells. SPIO+ DC in the T-cell area colocalized with both CD4+ and CD8+ T cells (Fig. 4A). Interestingly, in various cells the staining for CD8 was most prominent at the interface between the T cell and the SPIO-positive cells (Fig. 4B, second panel). This explicit localized expression of CD8 molecules suggests that the T cells are actively engaged with the DC in a way that stimulated the redistribution of the CD8 molecules and likely the formation of an immunologic synapse. Activation of T cells surrounding the injected DC was further supported by the expression of the early T-cell activation marker CD69 and the activation marker CD25 (Fig. 4B). Of the SPIO+ cells in the T-cell area, 65% ± 22% were in the immediate proximity of CD69+ cells (8 lymph nodes from six patients). Similarly, 64% ± 13% of the SPIO+ cells in the paracortex were surrounded by CD25+ T cells (10 lymph nodes from six patients).

To further investigate the immune activating potential of the migrated 111In/SPIO DC, they were isolated from a lymph node cell suspension by a magnet. We found that some of these DC were still forming rosettes with T cells (Fig. 4C). In addition, we could show that T cells cultured from lymph nodes suspensions of two i.n. vaccinated patients showed KLH-specific proliferation (Fig. 4D). Thus, 111In/SPIO DC interacted with T cells and induced antigen-specific T cell responses within 48 hours after vaccination.

Systemic immune responses after i.d. and i.n. vaccination with DC are comparable despite difference in number of DC in T-cell area. To test whether the differences in 111In/SPIO-DC distribution after vaccination by different routes of administration had any effect on the strength of immune activation, we compared the immune responses elicited in patients that received multiple injections. To monitor the capacity of vaccinated DC to initiate an immune response, DC were not only pulsed with TAA peptides but also loaded with the foreign protein KLH. We observed that most patients (i.n., 11 of 13; i.d., 10 of 11) developed an IgG antibody response against KLH after DC vaccination, with serum levels increasing after each vaccination (Fig. 5A). Levels of KLH-specific IgG antibody were comparable after i.n. and i.d. vaccination. In addition, KLH-specific proliferation of CD4+ T cells was induced in all i.n. and i.d. injected patients (Fig. 5B). Thus, despite the difference in the number of DC that reach the skin-draining lymph nodes, there was no effect on the strength of the immune response against KLH after i.n. or i.d. injection.

Next we compared the capacity of i.d. and i.n. injected DC to induce specific CD8+ T-cell responses against the tumor-associated peptides used for loading. TAA responses were tested by staining T cells with HLA-A2.1 tetramers encompassing gp100:154, gp100:280, or tyrosinase:369 peptides. In one patient in each group, we could already detect tetramer-specific T cells in PBMC freshly isolated after four vaccinations. Because the frequency of peptide-specific T cell in the blood is very low and often undetectable, DTH responses were induced by injecting peptide-loaded DC i.d. We have shown previously that sampling of DTH sites is a very powerful approach to
detect vaccine-related T cells (24). No significant differences were found in the number of patients with tetramer+ and IFN-γ–producing T cells isolated from DTH reactions. TAA tetramer–positive T cells were found in 4 out of 11 patients after i.d. vaccination and in 6 out of 11 patients after i.n. vaccination (Table 1; example in Fig. 5C). The functionality of these T cells was tested by coculturing with T2 cells loaded with TAA peptides and measuring the IFN-γ production. Both after i.d. (4 of 10 patients) and after i.n. (4 of 8 patients), T cells from DTH biopsies produced IFN-γ in cocultures with peptide-loaded target cells (Table 1; example in Fig. 5D). Thus, despite the limited number of DC migrating to subsequent lymph nodes, immunologic responses were found in most patients, and no significant differences were found in the immunologic responses after i.n. and i.d. DC vaccination.

Discussion

Previously, we have shown that DC can be tracked in vivo both by scintigraphic imaging and magnetic resonance imaging (9, 17). In this study, we monitored the migration of DC into the T-cell area, the paracortex, of the lymph node at a microscopic level. Although in vivo imaging provided us with a lot of information on the distribution of the DC vaccine after injection, the limitation is that lymph nodes can only be generally qualified as positive for the cell-tracking label. Here we exploit immunohistochemistry to show that many DC remained at the site of injection where they died and were cleared by macrophages. Nevertheless, small numbers of labeled DC did reach the lymph node and migrated into the T-cell area, where they interacted with T cells and induced immune responses against KLH and TAA. Importantly, despite the fact that more DC reached the T-cell areas after i.n. injection, DC were equally well capable of inducing immune responses.

In vivo tracking of ex vivo–generated cells is an important readout for the evaluation of cellular therapy. Our results explicate that in vivo tracking of therapeutic cells primarily tracks the cell label and may need to be confirmed with immunohistochemical studies. For example, whereas quantification of migrated DC after i.d. injection proved to be quite accurate (~85% of the migrated cells were indeed DC),

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**Fig. 4.** 111In/SPIO DC interact with lymph node T cells and activate KLH-specific T cells. A, T-cell area after i.n. injection stained for CD83, CD163, CD4, and CD8 (>630). Bar, 20 μm. B, T-cell area of lymph node after i.n. injection stained for CD8 (second panel, detail of first panel), CD69 (third panel), and CD25 (last panel; original magnification >630). Bar, 20 μm. Iron oxide was visualized by Prussian blue staining. Nuclei were stained with hematoxylin (A and E, right) or nuclear fast red (D and E, left). C, rosette of a SPIO-labeled DC with T cells magnetically isolated from a lymph node of a patient after i.n. injection. Bar, 20 μm. D, KLH–specific proliferation of lymph node–derived T cells from two patients after one i.n. DC injection. T cells were cocultured with irradiated autologous PBMC loaded with (black bars) or without (white bars) KLH. Proliferation was measured in a tritiated thymidine incorporation assay.
quantification of the number of viable therapeutic cells that had actually migrated after i.n. injection was not possible because the migrated cells as well as the depot were present within the same lymph nodes. Therefore, not only viable DC but also nonviable DC and phagocytes that have taken up the tracking agent were imaged. Moreover, migration may be overestimated because therapeutic cells and phagocytes containing the tracking label cannot be distinguished by in vivo imaging. Thus, depending on the method of administration and the nature of the vaccine, immunohistochemical analysis of target organs will still be decisive for quantitative and qualitative evaluation of the efficacy of cellular therapy.

i.d. administration of DC vaccines is not only attractive for imaging purposes, it is also easier to do, and less time-consuming and less expensive than i.n. injection, which needs to be done under ultrasound guidance by an experienced radiologist. We found that after i.d. injection, never >4% of DC migrated to the draining lymph node, whereas up to 56% of the DC redistributed to distinct lymph nodes after i.n. injection. Despite these major differences in migrating DC, immunologic responses were comparable after both i.n. and i.d. DC injection. Up till now, the minimal amount of DC that is needed to induce an efficient immune response in humans is unknown. From our data, we can calculate that after i.d. injection, <5 × 10⁵ DC reach the T-cell area (in the case of a maximal migration of 4%, of which 84% accounts for viable DC after injection of 15 × 10⁶ DC). Apparently, this low number of DC is sufficient to induce de novo immune responses. It is interesting that after four vaccinations, immune responses after i.n. injection of DC were not significantly higher than after i.d.
injection even though at least 10-fold to 30-fold more SPIO+ cells reach the paracortex with each vaccination. An important difference is that after i.d. injection, only few macrophages migrate from the injection site into the lymph nodes, whereas after i.n. injection all macrophages are already inside. The presence of macrophages that have phagocytosed 111In/SPIO DC in the paracortex (~50% of SPIO+ cells after i.n. injection versus 13% after i.d. injection) may have a negative effect on the immune response; for instance, by the secretion of anti-inflammatory cytokines. CD163 was found to be exclusively expressed by anti-inflammatory (m3c2) and not by proinflammatory macrophage (m2) subsets cultured from monocytes (25). These m2c2 macrophages can induce regulatory T cells via the expression of membrane-bound tumor growth factor-β (26). Thus, the presence of large numbers of CD163/SPIO+ macrophages may dampen the immune response induced by the 111In/SPIO DC.

Although we detected anti-KLH immune responses in most patients, TAA-specific immune responses were observed less frequently. One of the reasons may be the absence of MHC class II–restricted peptides. Also, the threshold for the activation of T cells specific for TAA tyrosinase and gp100 is likely higher than for the immunogenic foreign antigen KLH. Most importantly, the binding of exogenous-loaded TAA peptides to DC is not stable, and peptides may therefore be presented by fewer cells at lower densities and for a shorter period of time (27). An alternative way of antigen delivery would be loading of DC with antigens that need to be intracellularly processed. The use of antigens requiring processing results in prolonged antigen presentation (27), thereby increasing the time span during which the DC can activate specific T cells. Furthermore, DC may amplify the response by the transfer of antigen to resident DC in addition to presenting processed antigen to T cells themselves (28, 29). Increasing antigen presentation, immune stimulatory potential, and migratory capacities of DC for immunotherapy may greatly enhance the efficacy of DC therapy (reviewed in ref. 30). At present, we are comparing peptide-loaded DC with DC loaded with mRNA encoding for TAA. Initial findings show that when DC electroporated with TAA mRNA are injected into a lymph node, TAA protein expression was detectable up until 24 hours after injection.10 Simultaneously, our research focuses on the generation of DC with high immune stimulatory capacity (31) and on maximizing migratory capacities and vaccine delivery. In addition, it will be worthwhile to investigate whether the injection of no more than 5 × 106 DC directly into a lymph node will already be sufficient to induce immunologic responses, as this will greatly reduce the need for large-scale production of DC and may make the use of DC that circulate in small numbers in the peripheral blood feasible. Optimizing both the generation of high migratory and immune stimulatory DC, and improving application and treatment strategies will be necessary to enhance the number of responding patients and advance experimental DC immune therapy into a standard treatment.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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