Original Article

Mouse dendritic cell migration in abdominal lymph nodes by intraperitoneal administration

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Abstract: Dendritic cell (DC) based immunotherapy is a promising approach for cancer treatment and has been approved in clinical settings for decades. Clinical trials have demonstrated relatively poor therapeutic efficacy. The efficacy of DC immunotherapy is strongly influenced by their ability to migrate to the draining lymph nodes (LNs). Therefore, it is critical to deliver DCs and monitor the in vivo biodistributions of DCs after administration. The purpose of this study is to determine whether a novel injection route of DCs improves DC migration to LNs, tissues, organs and lymphatics. In the present study, a modified method was investigated to acquire DCs from mouse bone marrow. Cultured antibody labeled DCs were analyzed by flow cytometry. India ink was used to visualize mouse abdominal LNs and PKH26 was utilized to label DCs for intraperitoneal (IP) injection, results were evaluated by histology. Our results showed that large amounts of DCs with a relatively high purity were acquired. IP injection of India ink marked the abdominal LNs and PKH26 labeled DCs showed IP was an effective administration route to increase the absorption of viable DCs, and different time points after IP inject showed no significant difference of the migrated DCs. The findings indicated that large amounts of high purity DCs can be acquired through our method and IP injection accelerates DCs migration to abdominal LNs, which can be directly translated to clinical settings, especially for abdominal cancers. This study makes a foundation for future researches of DC-based immunotherapy as a treatment modality against cancer.

Keywords: Dendritic cells, lymph node, migration, abdomen, intraperitoneal administration

Introduction

Dendritic cells (DCs) can present antigens on their surface to other cells of the immune system [1, 2] and have been used in this capacity as potent therapeutic vaccines against human cancers [3]. However, immature DCs in a tumor environment may be functionally defective [4-6]. This limitation can be overcome by generating immature DCs from bone marrow cells and activating them with tumor antigens to allow maturation of these DCs with proinflammatory stimuli in vitro [7, 8]. Many experimental immune therapies are now based upon the DC immunotherapies of cancer patients with autologous DCs [6, 9-11]. DC-based immunotherapy has clinically relevant mechanisms of action with great potential for the systemic treatment of many cancers in clinical settings [12-14]. However, clinical trials have not yet demonstrated positive therapeutic efficacy and clinical response has been limited to a minority of patients [9-11, 15]. The effectiveness of immunization with DC-based immunotherapies is strongly influenced by their successful migration to secondary lymphoid organs and tissues where they orchestrate immune response [6, 9-11, 16].

The administration route and frequency of DC-based vaccine injections along with the number of injected cells are strongly relevant with the
viable DCs migration efficacy and subsequently determine the result of the treatment. Currently the standardized methodology of DC-based cancer vaccine application is not established yet [17]. Though ex vivo generated DCs loaded with specific tumor antigens have been proved to be feasible and superior, the administration route remains controversial [10]. Given that substantial population of LNs reside in the abdomen and together with spleen, consisting the most important secondary lymphoid organs [18], it’s theoretically effective that we inject the DCs intraperitoneally (IP), particularly for the abdominal tumors. However, few studies have demonstrated the efficiency of DC-based cancer vaccine employed through this route.

In this study, to determine whether a novel injection route of DCs improves DC migration to LNs, tissues, organs and lymphatics, which may give new insights into the DC-based anti-cancer regimen and provide approaches that can be directly translated to clinic and improve the outcome.

Materials and methods

The study was approved by institutional animal care and use committee (IACUC) and were strictly performed in compliance with NIH guidelines.

Animals and reagents

The mouse bone marrow derived DCs were prepared as previously described with some modifications [19]. Briefly, C57BL/6 mice (4-6 weeks age; Charles River, Wilmington, MA) were sacrificed with CO2. After immersed in 70% ethanol for 5 mins, tibias and femurs were carefully dissected. All the muscles and tissues attached to the bones were cleaned with sterilized gauze, and the bones were disinfected by immersion in 70% ethanol for 5 mins, tibias and femurs were carefully dissected. All the muscles and tissues attached to the bones were cleaned with sterilized gauze, and the bones were disinfected by immersion in 70% ethanol for 5 mins. Then the bones were flushed in half with fetal bovine serum (FBS) free RMPI 1640 (Gibco, Waltham, MA) in the hood. When the middle of the bones became visually white, the end of the bone was cut and flushed again. The bone marrow cells were collected and red blood cells were lysed. A total of 1×10^6 cells in 10 mL DC culture medium that containing 10% FBS (Gibco, Waltham, MA), 100 units/mL penicillin, 100 µg/mL streptomycin, 0.25 µg/mL amphotericin b (antibiotic-antimycotic 100x, Gibco, Waltham, MA), 10 ng/mL rm-GM-CSF and 1 ng/mL rm-IL-4 (both from Shenandoah Biotechnology, Warwick, PA) were plated in a petri dish. If a mature state was preferred, a cocktail of IFNg, TNFα and LPS (all from Shenandoah Biotechnology, Warwick, PA) was added at 7th day. After 8 days’ culture, floating and loosely attached cells were ready to be harvested and tested by flow cytometry.

FACS

The cultured mouse bone marrow derived cells were collected and washed by cold PBS, incubated for 40 mins at 4°C with 2 µg/3×10^5 cells anti-mouse PerCP-Cy5.5 CD11c, PerCP-Cy5.5 CD11b, APC CD40, APC CD86, PE CD80, PE H-2Db, FITC H-2 Kb (all from BD Bioscience, San Jose, CA), PE MHC II (Southern Biotech, Birmingham, AL) and appropriate isotype controls. Cells were identified by flow cytometry (BD LSRFortessa™ cell analyzer, San Jose, CA) and the data were analyzed by FlowJo (FlowJo LLC, Ashland, OR).

Dendritic cell labelling

The labeling of DCs was performed according to production manual. Briefly, at the 8th day of culture when the DCs were mature, 2×10^6 washed cells in 100 µL Diluent C were mixed with 100 µL Diluent C containing 0.4 µL PKH 26 dye for 5 mins (Sigma-Aldrich, St. Louis, MO). After adding 200 µL FBS and 3 times washing, the DCs were labeled with PKH26 (red) and the labeling efficiency was evaluated by fluorescent microscope (Axiovert 40 CFL, Carl Zeiss, Ontario, CA).

Abdominal LNs visualization

Six mice were IP injected with 100 µL 10% india ink (BD, Sparks, MD) and 2×10^6 PKH26 (red) labeled DCs respectively. After 30 mins, the india ink injected mice were sacrificed and abdomen were opened. India ink stained LNs and lymphatics can be visually seen. Spleen, pancreas, intestine and surrounding tissues were then collected and fixed in formalin for histological verification of india ink staining. As to PKH26 (red) labeled DCs injected mice, the interval before euthanasia is 1 hour. The collected tissues were embedded in OCT compound (Fisher HealthCare, Houston, TX) infused modes that were placed on dry ice, and freeze at -80°C after 2 mins.
Mouse DC generation and its migration in abdomen

Histology

For India ink injected mice, the organs and tissues collected after euthanization were immediately fixed in formalin solution for 72 hours, and then embedded in paraffin. Slices were cut in 4 μm thick and stained with H&E. All the slices were analyzed with optical microscope (EVOS FLC Imaging System, Life Technologies, Carlsbad, CA) from low magnification (10x) to high magnification (40x) by the same person. For PKH26 (red) labeled DCs injected mice, the samples that were placed in OCT compound and freeze at -80°C were cut in 5 μm with the microtome portion of the cryostat and picked up on slides. One group of the slides were mounted by cover glasses with ProLong Glod Antifade Reagent with DAPI (Cell Signaling Technology, Danvers, MA). Another group were stained with H&E, and all the slices were observed under optical microscope as well as florescent microscope (Axioimager Z1, Carl Zeiss, Ontario, CA).

Statistics

Total cells amount and PKH26 positive cells amount of the representative figures of spleen
Mouse DC generation and its migration in abdomen

A total amount of 1.0-1.2×10^7 cells could be acquired from one mouse bone marrow through our method, as diagrammatized in Figure 1, in contrast to flushing in half only with 0.5-0.8×10^7 cells. Representative photographs of the cultured cells with typical DC characteristics in a dynamic development were shown in Figure 2. After 8 days' culture, the bone marrow derived cells with initial concentration of 1×10^5 cells/mL would proliferate to occupy all the 10 cm petri dishes, and we were able to obtain 1-2×10^8 cells at this point.

**Results**

**BMDC amount and morphology development**

A total amount of 1.0-1.2×10^7 cells could be acquired from one mouse bone marrow through our method, as diagrammatized in Figure 1, in contrast to flushing in half only with 0.5-0.8×10^7 cells. Representative photographs of the cultured cells with typical DC characteristics in a dynamic development were shown in Figure 2. After 8 days' culture, the bone marrow derived cells with initial concentration of 1×10^5 cells/mL would proliferate to occupy all the 10 cm petri dishes, and we were able to obtain 1-2×10^8 cells at this point.

**BMDC purity and maturity**

The cultured bone marrow derived cells were collected and tested by flow cytometry. Results suggested a relatively high purity and maturity

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**Figure 3.** Flow cytometry analysis of BMDC. DCs were harvested after 8 days' culture. CD11c and CD11b were chosen as DC specialty markers, CD40, CD80, CD86, MHC II, H-2Db and H-2Kb were utilized as DC maturity markers (A, B). (C) Average percentage of the chosen markers expression. Error bars represented mean ± SD. Three independent experiments were performed.

Slides were counted by ImageJ software (NIH, Bethesda, MD). Student’s t test was used to compare the differences between different time points after IP injection of PKH26 labeled DCs. Data were presented as mean ± SD. *P* < 0.05 was considered as statistically significant.
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of DCs with substantial expression of CD11c, CD11b and other DC maturity markers compared to isotype controls (Figure 3A, 3B). Independent repeating experiments of analogous DC markers expression confirmed this method as a stable and repeatable approach to achieve extensive quantity mouse bone marrow derived DCs (Figure 3C).

Abdominal LNs Visualization by India ink

After 30 mins india ink IP injection, the mouse abdominal LNs and lymphatics could be visually distinguished (Figures 4A, 4B, 5B). The india ink stained organs and tissues were collected for histologic demonstration of the staining efficiency. H&E staining of the slices indicated significant uptake of india ink by abdominal lymph organs (Figures 4C, 4D, 5D). As spleen is the largest lymph organ of the body where large amounts of lymphocytes exist, we compared the macroscopical and histological features of spleen after india ink and equal PBS IP injection, respectively. Results showed considerable stained cells in the spleen (Figure 5).

Migration of fluorescence labeled BMDCs to abdominal LNs via IP injection

The PKH26 (red) labeling efficiency of mouse bone marrow derived mature DCs was verified under fluorescent microscope (Figure 6A, 6B). 2×10^6 labeled

Figure 4. Abdominal LNs visualization by india ink. Mice are injected 100 µL of 10% india ink intraperitoneally and sacrificed after 30 mins. The abdomen is opened and pancreas as well as intestine with visual India ink staining are collected for histology analysis (A, B). Representative pictures of pancreas and intestine slices with H&E staining are shown (C, D). Black arrows indicate LNs and lymphatics stained by india ink. Scale bar represents 40 µm.

Figure 5. Macroscopical and histological pictures of spleen after india ink injection. Spleens are collected 30 mins after IP injection of 100 µL India ink at 10% concentration or 100 µL saline solution (A, B). Corresponding slices of H&E staining are presented with black arrows indicating LNs and lymphatics staining with india ink (C, D). Scale bar represents 40 µm.
Mouse DC generation and its migration in abdomen

Figure 6. Fluorescent labeled BMDC in abdominal LNs. The cultured DCs labeled with PKH26 (red) were observed by optical and fluorescent microscope (A, B). Scale bar represents 20 µm. Spleen, pancreas and intestines along with surrounding tissues were collected for frozen sections after 1-hour IP injection of labeled DCs. H&E staining characterized the tissue structure and the labeled DCs were detected in the LNs, tissues, organs and lymphatics under fluorescent microscope (C-H). (C-E) Scale bar represents 40 µm. (F-H) Scale bar represents 20 µm. PKH26 positive DCs in spleen were quantified by ImageJ in different time points after IP injection (I). (#), no statistical difference.
Mouse DC generation and its migration in abdomen

cells were injected intraperitoneally and the mice were euthanized 1 hour later. The same organs and tissues to Figures 4 and 5 were collected for frozen sections. After the organ structure were confirmed by H&E staining (Figure 6C-E), fluorescent microscope analysis of the same slides detected the labeled DCs in the LNs, tissues, organs and lymphatics, which was in consistent with india ink injected mice results (Figure 6F-H). The results of spleen were quantified by ImageJ and no significant difference was observed between different time points after IP injection (1 h vs. 6 h, 7.26% vs. 7.73%, \( P = 0.4842 \); 1 h vs. 12 h, 7.26% vs. 7.59%, \( P = 0.7462 \); 6 h vs. 12 h, 7.73% vs. 7.59%, \( P = 0.8908 \)) (Figure 6I).

Discussion

Ex vivo isolation of DC precursors from murine bone marrow and following differentiation into DCs has been extensively studied, and it is reported to be feasible and effective [20, 21]. However, unified standard is still not reached among diverse exploration of the mouse bone marrow isolation process and cell culture recipe [22-24]. In the present study, a modified method was used to acquire large amounts of DCs from mouse bone marrow. The tips of bones were preserved at first to avoid possible contamination and cut it after first flush to fully wash out the bone marrow. Furthermore, the ratio of rm-GM-CSF and rm-IL-4 combination was investigated for the DC culture medium and found out 10 ng/mL rm-GM-CSF with 1 ng/mL rm-IL-4 to be the preference in our assays. A relatively high DC purity and maturity were founded after 8 days’ culture rather than 6 days, similar as Lutz’s conclusions [25]. These slight modifications enabled us to gain large quantities of highly pure DCs from mouse bone marrow for further research compared to other reports [20, 21].

The administration route of DC-based cancer vaccine is directly correlated with consequent antitumor immune response and immune memory [26, 27]. Currently, administration routes include subcutaneous injection and intra-LN injection [28, 29]; Low dose of DCs can migrate to LNs and organs by subcutaneous injection [28, 29]. A cluster of DCs directly delivers to LNs, which affects DC therapeutic function by intra-LN injection [28, 29]. Regarding in the anatomic basis of abdomen. It turned out the uptake of IP injected labeled DCs by abdominal LNs, tissues, organs and lymphatics was comparable to india ink IP injection, indicating this administration method may increase the amount of viable DC-based vaccines that are actively absorbed by abdominal lymphoid organs and thus improve outcomes. Our results suggested a more suitable and potential administration route for abdominal cancers.

Previous reports showed that DC-based vaccine might be more valuable and efficient when applied in the early stage of the disease. Current attempts to combine DC-based cancer vaccine with immunostimulatory cytokines, immunomodulating agents, chemotherapy, targeted therapy and immune checkpoint inhibitors have shown clinical benefits [30-33], while the optimal combination regimen remains to be established. However, large amount of highly pure DCs from mouse bone marrow will be needed and IP delivery of DCs will improve DC therapeutic efficacy due to more DCs to abdominal LNs.

In conclusion, we present a modified method to acquire considerable amount of mouse bone marrow derived DCs and testify the efficiency of IP injection of labeled DCs, making a foundation for future researches of DC-based vaccine immunotherapy as a part of comprehensive treatment against cancer, especially for abdominal tumors.

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Disclosure of conflict of interest

None.

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