A novel modulation of structural and functional changes of mouse bone marrow derived dendritic cells (BMDCs) by interleukin-2(IL-2)

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Abbreviations: IL-2, Interlukin-2; ACP, Acidic phosphatase; BMDCs, Bone marrow derived dendritic cells; APCs, LPS; Lipopolysaccharide; TEM; Transmission electron microscopy; DAB 3,3'-diaminobenzidine; MTT 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide; FCM, Flow cytometry; MACS, Magnetic activated cell sorting; CTL, Cytotoxic lymphocyte

IL-2 is a pleiotropic cytokine produced by T cell after antigen activation of T cell and it is so called T cell growth factor. A large number of documents suggest that IL-2 plays pivotal roles in the immune response and now IL-2 is an approved drug being used for various kinds of diseases such as cancer and dermatitis. The aim of present exploration was to look at effect of IL-2 on structural, phenotypic and functional maturation of murine BMDCs. The structural and phenotypic maturation of BMDCs under influence of IL-2 were evaluated by light microscope and flow cytometry (FCM). The functional maturation of BMDCs was confirmed by cytochemistry assay, FITC-dextran, acid phosphatase (ACP) activity, bio-assay and enzyme linked immunosorbent assay (ELISA). We elucidated that IL-2 up-regulated the expression of key surface markers such as: CD80, CD83, CD86, CD40 and MHC II molecules on BMDCs, down-regulated phagocytosis activity, induced more production of IL-12 and TNF-α secreted by BMDCs. Therefore it can be concluded that IL-2 effectively enhance the maturation of BMDCs. Our results provide direct evidence to support IL-2 would be used as a potent adjuvant in preparation of DC-based vaccines, as well as an immune remedy for cancer situation.

Introduction

IL-2 is an interleukin, a type of cytokine and a soluble signaling molecule mediating in the immune system. It was first reported simultaneously by Shinpei Kasakura and Louis Lowenstein, plus Julius Gordon and Lloyd MacLean for its mitogenic activity in the culture media of mixed leukocytes. IL-2 is also a small protein that regulates white blood cells in body’s natural response to infection. As documented, IL-2 mediates a series of immunological effects by binding to IL-2 receptors on the surface of lymphocytes and the cellular responses in our body are regulated by the amount of IL-2 produced in response to antigens. It also stimulates the production of IL-2 receptors on the surface of immune cells and acts through autocrine or paracrine way. As result of this the expansion and activation of macrophages, natural killer cells, B lymphocytes etc. would happen.

The unraveled data on IL-2 has considered IL-2 as a major growth and apoptosis factor for activated T lymphocytes. IL-2 also plays necessary role in maintaining self-tolerance and it has been used for over 30 years, and remains one of the most extensively studied cytokines. Especially in recent years there were more and more reports on antitumor activity of IL-2.

Dendritic cells (DCs) were first described by Paul Langerhans (Langerhans cells) in late stage of nineteenth century. It wasn’t until 1973, however, that the term “dendritic cells” was coined as part of immune cells by Ralph M. Steinman and Zanvil A. Cohn, representing a milestone of immunology. The main function of DCs is to process and present antigen on the surface to be recognized by T cell and therefore initiate T cell response, functioning as antigen-presenting cells (APC). DCs are not only important for the induction of primary immune responses, but also crucial for the induction of immunological tolerance.

There were many articles describing relation between dendritic cell and IL-2, including transformation of monocyte to DC under influence of IL-2, combined with other cytokines, IL-2 promotes the motility of dendritic cells and their accumulation in lung and skin. IL-2 production by dendritic cells and its immune-regulatory functions, IL-2 induces development of dendritic cells from cord blood CD34+ cells. However, so far there is no systematic study from data bank on the impact of IL-2 on phenotypic and functional maturation of DCs. Therefore we performed the following research to look at phenotypic and functional effect on DC by IL-2.
Results

Confirmation of change of key surface markers on the BMDCs by FCM

Post treatment with 25μg /ml IL-2 for 48h immature BMDCs developed into maturation confirmed by upregulated expression of key surface markers such as: CD86, CD80, CD83, CD40 and MHC-II, which will work as key signal and co-signals to activate T cell. The detailed changes are shown in Figure 1.

T cell activation driven by BMDCs treated with IL-2

Usually immature BMDCs are with a powerful ability to capture and process antigen, but with a lower potential to initiate T-cell response. IL-2 induced maturation of BMDCs with increase in driving T lymphocytes response. The index of induced BMDCs to drive expansion of allogeneic lymphocytes was reflected by a corresponding value of OD492nm (A492) and yielded following numbers as shown in Figure 2.

Figure 1. Detailed changes of surface markers on the BMDCs after treatment with IL-2.

Figure 2. Assay for treated BMDCs to drive T cells.
Immunohistochemical analysis of phagocytosis of BMDCs

Post staining with DAB kit, BMDCs observed under conventional microscope exhibited the process of phagocytosing. The picture revealed a process of phagocytosing labeled horseradish peroxidase by BMDC and exhibited more phagosomes in immature BMDC as compared to less phagosomes in mature BMDC shown in Figure 3. This result was corresponding to ACP activity.

Phagocytosis study by FCM

Antigen arrested inside BMDCs also underwent FITC-dextran test. IL-2 weakened the phagocytosis of BMDCs to the labeled dextran. Reversely BMDCs without IL-2 treatment showed strong phagocytosis, as shown Figure 4.

Test of ACP activity

With the processing of BMDCs maturation ACP activity changed accordingly. The decrease of ACP activity was an indication of gradual termination of phagocytosis and the number of ACP activity could be reflected by a value of OD520nm (A520) in Figure 5.

Assay for production of IL-12 and TNF-α

The immature BMDCs stimulated by 25μg/ml IL-2 for 96h developed into maturation. Besides maturation in morphology as proved by upregulated expression of key surface markers, the BMDCs also underwent functional maturation by secreting higher levels of IL-12 and TNF-α. IL-12 and TNF-α could play important role in regulating a spectrum of innate and adaptive immune responses, as exhibited in Figures 6A,B.

Discussion

IL-2 is a natural messenger protein of human immune system, called a cytokine which activates parts of immune system and induces substantial increase in number of tumor-reactive T cells in vitro. Experimental data prove that T cells cultured with suitable dose of IL-2 can be more effective in vivo for showing specific tumor cytotoxicity. IL-2 does not kill tumor cells directly like a chemotherapy drug. Instead, IL-2 activates the expansion of T-cells and natural killer cells (NK), both of which can kill cancer cells directly. Now IL-2 has some clinical application in cancer therapy such as: as specific cellular anti-tumor immunity and cytokine induced killer (CIK) cells in cancer therapy.18–20

Early documented research results in data bank have indicated that clinical significance of IL-2 to boost and maintain the balance of both innate and adaptive cellular immunity and even now IL-2 continues to be a key solution for most of melanoma and renal cancer.21–23

In our current research we have proven that IL-2s affects on phenotypes and functions of BMDCs were present. From our results we could see that IL-2 induced the maturation of BMDCs with the following identified fact: (1) 25μg/ml of IL-2 and 48h culture were optimal condition to boost BMDCs; (2) BMDCs under induction of IL-2 showed typical maturation with reduced phagosomes inside BMDCs, which would downregulate phagocytosis of BMDCs, with simultaneously intensified antigen (Fig.3) presentation; (3)IL-2 increased expression of key surface markers of CD80(B7.1), CD40, CD83, CD86(B7.2) and MHC II on the BMDCs(Fig.1), which would mount antigen presentation of BMDCs to initiate specific T cell response(Fig.2) ; (4) IL-2 decreased ACP activity, which is consistent with decreased phagocytosis and increased BMDCS driving T lymphocytes reaction(Figs.3, 4) and (5)IL-2 upregulated secretion of IL-12 and TNF-α, which would be helpful in intensifying the signals between the DC and CD4+T cell, increasing Th1 branch responses and amplifying macrophage function(Fig.6).

Additionally we should pay attention to that higher secretion of IL-12 and TNF-α by BMDCs would not only act on themselves in an autocrine or paracrine manner, resulting in more activated BMDCs, but also will amplify the signal to direct CD4+T cells to be functioning and thus the stability between DCs and CD4+T cells would be firmed. As a positive feedback, this chain signals will enhance more production of cytokines by BMDCs and TNF-β or IFN-γ by CD4+T cells, involved in stimulating immune responses.
macrophage to secret many types of cytokines, coordinating a chain of immune responses in the body.  

It is very surprising to us that the results for IL-2 treatment are statistically different from controls only at a 1:1 ratio and the low response obtained with LPS at only 1:1 and 1:5 ratios. These might be caused by our laboratory condition, cell purity or medicines we used.

Really immune system is a diverse network with rather complication, in which all of immune cells are interacting directly or indirectly through mediation of immune molecules, forming a precise feedback. When TNF-α or IL-12 is playing role no matter whether it works alone, or is interacting with other signal molecules, it can exert impact on BMDCs in a direct way or indirect way by activating other cells such as macrophages, neutrophils and NK cells. These activated cells can then secret high levels of cytokines, acting on BMDCs. For example, activated NK or neutrophils can release more IFN-γ to induce BMDCs to maturation and, as consequence of this, BMDCs will secret more IL-12 forming an interacting loop, maintaining the balanced immunity. IL-2 can play a very important role as immune adjuvant to be used in vaccines preparation.

DCs, originally discovered by Steinman (1972), represent marked progress in immunology and in cancer therapy. Due to the most powerful antigen presentation to initiate naïve T cell response they are therefore critical to specific cellular immunity and cellular cytotoxicity.

So far continuous emerging scientific data are supporting DC activation will be one of the best solutions to cancer therapy. This meaningful exploration with enriched data contribute to the further understanding of immunomodulatory role of IL-2 in optimizing immune system or immune functions and further understanding of deepening mechanisms of IL-2 action at subcellular or molecular level.

Finally this investigation also presents a meaningful mode for IL-2 and highlights the clinical potential of IL-2 as an adjuvant used in cancer therapy, an useful adjuvant in vaccine designing.

**Conclusion**

We believe that it is the first time that we release the data on a systematic study demonstrating that IL-2, at defined concentration could enhance both phenotypic and functional maturation of BMDCs. The induced BMDCs could also supply with more IL-12, TNF-α and other key surface markers of CD40, CD80, CD86, CD83 and MHC II to work together to intensify antigen presentation by BMDCs to activate T response.

**Materials and Methods**

**Reagents**

rIL-2 was a product of PeproTech USA. The impact of a range of concentrations of IL-2 from 1.56μg/ml to 200 μg/ml on the growth of BMDCs in vitro was tested, and the concentration of 25 μg/ml was found to be optimal concentration. So 25 μg/ml was token for current work. Recombinant murine cytokines of IL-4 and GM-CSF were provided by PeproTech Inc. LPS (from Escherichia coli, serotype 055:B5) was a product of Sigma-Aldrich. The mAbs used in the study include PE-anti-CD83, FITC-conjugated anti-CD40, PE-anti-CD80, PE-anti-CD86 and PE-anti-MHC-II, which were bought either from eBioscience or BD PharMingen. ELISA kits for cytokines assays were bought from R&D System. Other chemicals and reagents, being frequently used in our laboratory were all made in Sigma-Aldrich or BD Pharmingen.
Induction of bone marrow cells to get BMDCs

The preparation of BMDCs from mice referred to a method as described previously. The bone marrow cells from the femurs and tibias of female C57BL/6 mice were removed of red cells and counted to 10^7/ml. They were grown in 24-well plates charged with 2 ml of RPMI 1640 containing standard 10% fetal bovine serum, 10ng/ml GM-CSF, 10ng/ml IL-4, 2 mM L-glutamine, 100units/ml penicillin, 10μg/ml streptomycin. After culture for 4h the adherent cells was obtained and 1μg/ml LPS was added to the culture medium on 7d of culture for overnight to get more number to fit purification with CD11c-MACS magnet beads and the process of purification was performed by referring published method and instruction manual included in MACS kit (Miltenyi Biotech.). The CD11c+ BMDCs were sorted out using anti-CD11c-coated magnetic beads and the auto- MACS system and the purity of the purified cells was confirmed by FCM with > 90% purity.

Confirmation of elevated key surface molecules by FCM

The BMDCs co-cultured in presence of 25μg /ml IL-2 for 48h were confirmed of expression of key surface molecules by FCM. The treated BMDCs were reacted together with anti-CD80, anti-CD86, anti-CD83, anti-CD40, and anti-MHC II antibodies at 4℃ for 30 min. After thorough rinsing, the cells were measured with use of FACS Calibur (Becton Dickinson, San Diego, CA) to quantitate change percentage.

T cell activation driven by BMDCs treated with IL-2

The BMDCs post treatment with 25μg /ml IL-2 were collected on 7d. The purified CD8+ T cells (purity>90%, with a similar way to purify DC.) from the splenocytes of C57BL/6 mice were sorted out with magnetic beads and subsequently the CD8+ T cells were grown with10μg/ml ConA for 48h to a number of 10^7/ml. The CD8+ T cells (5 × 10^5 /well) were incubated in 96-well culture plates with differential BMDCs at a ratio of 1:1, 1:5, 1:10, 1:50, 1:100 for 5d. The proliferation of CD8+ T cells was determined by testing the OD absorbance at 492nm (A492) using a bichromatic microplate reader. Results from triplicate wells were expressed as the mean ± SE.

ACP activity determination

One × 10^6/ml BMDCs treated with 25μg /ml IL-2 for 48h were used with the phenol-4-AAP (amino anti-pyrine) plus ACP testing kit (Jiancheng Bio-engineering Institute of the South) to test ACP activity inside the BMDCs. OD number of absorbance at 520 nm (A520) represent activity.

Immunohistochemical analysis for phagocytic process by BMDCs

The BMDCs incubated with 25μg /ml IL-2 for 48h were combined with 0.08mg/ml horseradish peroxidase, then fixed with methanol for 4h, followed by with DAB staining and finally the stained sample was observed under microscope.

Phagocytosis study by FCM

The BMDCs incubated with 25μg /ml IL-2 for 48h were collected and100μl FITC-Dextran (40,000 D) 28–30 was added to the BMDCs culture at 4℃ for 2h, subsequently at 37℃ for 1h. Finally the sample was checked with FACS Calibur (Becton Dickinson, San Diego, CA) for localization of phagocytosis process.

Assay of cytokine production of IL-12p70 and TNF-α by ELISA

The BMDCs incubated with 25μg /ml IL-2 for 96h were collected for assay of cytokine production of IL-12p70 and TNF-α by ELISA respectively per instruction manual in the ELISA kit. The absorbance at 450 nm (A450) was obtained using a bichromatic microplate reader (BIO-TEK, USA).

Statistical analysis

All results were processed using statistical program SPSS (Statistical Package for Social Sciences, Version 16.0) for Windows and variables number were presented as mean ± SE. The divergence evaluated by ANOVA indicated statistical significance while P < 0.05.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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