Antitumor effect of murine dendritic and tumor cells transduced with IL-2 gene

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Abstract: Interleukin (IL-) 2 acts on a number of types of immune cells promoting their effector functions. To replace systemic administration of recombinant form of this cytokine, various genetically modified cells have been used in different preclinical models for tumor growth inhibition. In this study, dendritic or tumor cells transduced with retroviral vector carrying IL-2 gene (JAWS II/IL-2, X63/IL-2, MC38/IL-2 cells) alone or combined with tumor antigen-stimulated dendritic cells (JAWS II/TAg) were exploited to treat colon carcinoma MC38-bearing mice. After the peritumoral injection of vaccine cells, the tumor growth delay and the increase in the number of tumor infiltrating CD4+ and CD8+ T lymphocytes were noted. A considerable increase in CD4+ cell influx into tumor tissue was observed when JAWS II/IL-2 cells or JAWS II/TAg with syngeneic MC38/IL-2 cells were applied. The increase in intensity of CD8+ cell infiltration was associated with immune reaction triggered by the same combination of applied cells or JAWS II/TAg with allogeneic X63/IL-2 cells. The effect observed in vivo was accompanied by MC38/0 cell specific cytotoxic activity of spleen cells in vitro. Thus, the application of vaccines, including IL-2-secreting cells of various origins, was able to induce different antitumor responses polarized by exogenous IL-2 and the encountered tumor antigen. (Folia Histochemica et Cytobiologica 2012, Vol. 50, No. 3, 414–419)

Keys words: MC38 murine colon carcinoma, IL-2 gene transduction, dendritic or tumor cell vaccine

Introduction

Interleukin (IL-) 2 is a pleiotropic cytokine acting on a number of types of immune cells to promote their effector functions. Due to this, IL-2 has been used as a tool in numerous clinical trials to fight growing tumors. Because of serious side effects observed after the systemic administration of IL-2, a number of alternative forms of therapy replacing the recombinant cytokines with genetically modified tumors as well as immune cells have been put forward [1–3]. Many studies have demonstrated that vaccination with IL-2-secreting tumor cells can cause considerable tumor growth inhibition, and additionally induce immunological memory [3–8]. On the other hand, results obtained in mouse models as well as initial clinical trials have demonstrated that genetically modified dendritic cells (DCs) can attend the induction of anti-tumor immune responses followed by tumor regression [8].

The aim of our study was to compare the effects of the application of IL-2-transduced cells in a model of colon carcinoma MC38. The main question was whether IL-2 transductants of various origins would be able to support the antitumor activity of tumor antigen (TAg)-stimulated DCs with the same efficiency. The cytokine carriers were represented by syngeneic tumor (MC38/IL-2) cells or allogeneic plasmacytoma (X63/IL-2) cells. Genetically modified DCs originating from the immature dendritic cell of JAWS II line represented immune cells.

The obtained results demonstrate that the vaccines including particular IL-2-secreting-cells, independently of their origin, induced similar tumor growth delay. However, the use of IL-2-secreting-cells in combination with TAg-stimulated DCs diversified their effect on the antitumor immune response.
Material and methods

Mice. Female C57BL/6 mice, 8–10 weeks old, were obtained from Mossakowski Medical Research Center, Polish Academy of Sciences, Warsaw. All animal experiments were approved by the Local Ethics Committee.

Cell lines. Parental cells of the MC38/0 murine colon carcinoma line [9] and genetically modified cells were maintained in RPMI 1640 supplemented with 5% fetal bovine serum (FBS). The C57BL/6 murine bone marrow-derived DCs of the JAWS II line (ATCC Cat. No. CRL-11904, USA), as well as cells modified with IL-2 gene carried by retroviral vector, were cultured in a culture medium containing 10% FBS and 5 ng/ml GM-CSF (Cytogen). The X63/Ag8.653 murine plasmacytoma cells transduced with IL-2 gene (Institute of Immunology and Experimental Therapy) were maintained in culture medium supplemented with 10% FBS. JAWS II cells and cells of both murine tumor lines were transduced with retroviral vector pQON carrying murine IL-2 gene (JAWS II/IL-2, MC38/IL-2, X63/IL-2) according to the previously described protocol [10]. Transductants produced cytokine at a relatively constant level and were not tumorigenic (data not shown).

Therapeutic schedules. Mice were inoculated subcutaneously (s.c.) in the right flank with MC38/0 colon carcinoma cells (1 × 10^6 cells/0.2 ml in PBS per mouse). On the 14th day after tumor inoculation, mice with palpable nodules were vaccinated peritumorally (p.t.) with dendritic JAWS II/IL-2 or tumor (X63/IL-2, MC38/IL-2) cells alone or in combination with JAWS II cells stimulated in vitro with tumor antigens (JAWS II/Tag). The vaccine consisted of ca. 1 × 10^6 cells (when combined — 0.7 × 10^6 of JAWS II/Tag cells were mixed with 0.3 × 10^6 of transductants). The cells were applied once a week, for three consecutive weeks. Tumor diameters were measured every 2–3 days. On the 7th day after the last injection (35th day of the experiment), some of the mice were sacrificed, and spleens as well as tumors were dissected.

Tumor sections and immunohistological analysis. Tumor tissue cryo-sections, 5–10 μm thick, were fixed in acetone and incubated for 2 hours with rat anti-CD4 or anti-CD8 mAb (BD Pharmingen) followed by goat anti-rat antibody conjugated with AlexaFluor488® (Molecular Probes) and counterstained with PI. The tissue sections were analyzed using a BioRad MRC 1024 scanning confocal fluorescence microscope equipped with LaserSharp software. Cryosections from three distinct parts of tumor from each individual mouse were analyzed.

Cytotoxicity assay of activated spleen cells. Spleen cells were co-cultured with mitomycin C treated MC38/0 cells for 5 days. After this in vitro stimulation, spleen cell cytotoxicity was investigated using a LIVE/DEAD Cell-Mediated Cytotoxicity Kit (Molecular Probes). The cytotoxic index was determined by the percentage of dead (PI-positive) among DiO MC38/0 target cells. The analyses were carried out using BD FACSCalibur apparatus with the CELLQuest software.

Results/Discussion

Our study analyzed the effect of the vaccines containing cells transduced with retroviral vector pQON carrying murine IL-2 gene on the induction of antitumor immunity. Transductants produced cytokine at the level range 60–120 LU/ml. They were represented by non-tumorigenic syngeneic tumor cells (MC38/IL-2 cells) and allogeneic plasmacytoma cells (X63/IL-2 cells) and JAWS II/IL-2 cells originating from an immature dendritic cell line (for detailed characteristics of these cells, see [10]). The analysis was conducted on mice bearing advanced murine colon carcinoma (MC38) treated with IL-2-producing cells and/or tumor antigen-stimulated dendritic (JAWS II/Tag) cells (Figure 1).

The obtained data revealed similar therapeutic effects of IL-2-producing tumor cell vaccines. The injections of MC38/IL-2 or X63/IL-2 cells elicited ΔTRV (difference in the median time required for the tumor to reach a volume of 1 cm³ compared to the control group of mice) amounting to 5.5 or 6.5 days, and TGI (tumor growth inhibition) amounting to 53 or 55%, respectively. The application of JAWS II/IL-2 cells resulted in stronger tumor growth delay (up to seven days for ΔTRV, and TGI of 56%; see insert in Figure 1). An explanation of their stronger effect on tumor growth delay could be based on two activities — they could act in a complex manner as a source of IL-2 and, to some extent [see 10], as antigen presenting cells. This suggests that although both tumor and dendritic vaccine cells were applied in the same numbers, their therapeutic effects depended not only on their ability to produce IL-2 but also on their various origins.

We noted that the application of JAWS II/Tag cells caused tumor growth delay for only four days compared to the control group; this is three days less than vaccine containing the same number of JAWS II/IL-2 cells. Given this, we decided to replace the 30% of JAWS II/Tag cells by JAWS II/IL-2 cells as well as by MC38/IL-2 or X63/IL-2 cells. The application of JAWS II/Tag cells along with JAWS II/IL-2 did not elicit further tumor growth delay. Thus, JAWS II/IL-2 cells which exhibited only low antitumor activity in vivo [10] were not able to balance the effect of the part of Tag-stimulated DCs and obviously were not able to augment the vaccine’s effectiveness.

We determined which type of vaccine tumor cells would be able to trigger efficient immunity against advanced tumor — the syngeneic cells presenting the same Tag as growing tumor or the allogeneic cells
expressing antigens of distinctive MHC and representing the other type of malignancy. Based on the data described by others, we expected that the allogeneic IL-2-secreting transductants ought to be strong stimulators inducing response not only against themselves but also against the growing tumor. Meanwhile, the application of vaccines consisting of JAWS II/TAg+MC38/IL-2 cells caused the highest tumor delay (ΔTRV — seven days and TGI — 61%), and after JAWS II/TAg+X63/IL-2 cell administration the weakest effect was observed (ΔTRV — four days, TGI — 40%, Figure 1). Moreover, the use of JAWS II/TAg+X63/IL-2 cells shortened the survival time of mice (data not shown). Generally, it was revealed that the application of JAWS II/IL-2 cells as well as JAWS II/TAg+MC38/IL-2 cells affected stronger tumor growth delay than other groups (illustrated by ΔTRV amounting to seven days and TGI — to 56% and 61%, respectively), and the use of JAWS II/TAg+X63/IL-2 cells triggered adversarial immune reaction. Overall, tumor growth delays elicited by particular vaccines were noted, but the differences between them were not statistically significant. Nevertheless, our observations allow us to hypothesize that vaccines containing various type of cells elicited final results which may be mediated by different components of the immune system. In many human cancers, the presence of T cells infiltrating into tumor tissue is believed to be a good prognostic marker for tumor regression [11]. The same relationship between the number of such lymphocytes and their antitumor effect can be observed...
in experimental tumors. In our model, the application of dendritic and/or tumor cell vaccines induced changes of the tumor tissue structure which were associated with the increase in the number of both CD4+ and CD8+ T cells. Generally, the influx of CD4+ cells was more intensive than that of CD8+ cells. However, the locations of these cells were various: CD4+ cells were found in connective tissue in tumor nodules, whereas CD8+ cells were preferentially located between tumor cells. The intensity of lymphocyte influx depended on the type of vaccinated cells (Figure 2, Table 1).

The highest infiltration of tumor tissue by both CD4+ and CD8+ cells was observed after the injection of JAWS II/IL-2 cells. The application of IL-2-secreting tumor cells caused a moderate influx, higher than in the control group but comparable with that elicited by JAWS II/TAg cells. On the other hand, after the application of the combined cell vaccines, a differential lymphocyte influx was noted. An inten-
Table 1. CD4+ or CD8+ cell infiltration into tumor tissues after the cellular vaccine treatment

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*Intensity of CD4 and CD8 cell influx was estimated based on 4–5 representative tumor sections and is shown by the number of crosses, i.e. “+” = few cells, and “++ + + + +” = many cells.

A significant influx of CD4+ cells was observed after treatment with JAWS II/TAg+MC38/IL-2 cells, a moderate influx after JAWS II/TAg+JAWS II/IL-2 cells, and a small influx after JAWS II/TAg+X63/IL-2 cells. In turn, the combined cell vaccines containing transduced tumor cells caused a more intensive influx of CD8+ lymphocytes than JAWS II/TAg+JAWS II/IL-2 cells.

Our results demonstrate that the most intensive influx of both CD4+ and CD8+ cells into tumor tissue was observed when IL-2-producing DCs or, alternatively, TAg-stimulated DCs along with IL-2-secreting syngeneic tumor cells, were delivered into the tumor surroundings. Moreover, when both CD4+ and CD8+ lymphocytes appeared in tumor tissue at a high level, a positive correlation between the intensity of the lymphocyte influx and the inhibition of the tumor growth could be observed. As observed by Bos [12], CD4+ cells presented in tumor tissue could help in overcoming the immune tolerance by promoting the CD8+ cell recruitment and activation of their cytolytic function together with the enhancement of the effector and memory CD8+ cell survival. The application of JAWS II/TAg+X63/IL-2 cell combination caused an influx of CD8+ cells similar to the other combined vaccines. Nevertheless, they elicited an unexpected overall effect inducing a weaker inhibition of MC38 tumor growth (the lowest TGI) and shortening the survival time of mice (data not shown). Presumably, this was a consequence of the simultaneous presence, in the vicinity of a growing tumor, of the TAg-stimulated DCs and allogeneic tumor cells. This parallel reaction directed towards two types of tumor cells probably caused an arrest of tumor growth delay. Thus, the application of allogeneic X63/IL-2 cells alone caused more effective tumor growth delay than the application of JAWS II/TAg+X63/IL-2 cell combination despite the level of CD8+ lymphocyte infiltration increasing after the latter vaccination. The application of vaccines of various origins seemed to be able to polarize the antitumor response by exogenous IL-2 and the encountered tumor antigen. This could result in a division of tumor infiltrating CD8+ or CD4+ lymphocyte, down-regulated by tumor microenvironment.

To answer the question as to whether an increase in the number of infiltrating cells would be associated with the level of lymphoid cell cytotoxicity, the splenocytes obtained from the treated mice were in vitro co-cultivated with MC38/0 cells for five days. After this time (restimulation), splenocytes were co-cultured with target MC38 cells stained with DiO (MC38/DiO). Next, MC38/0-DiO cells were stained with PI and analyzed using flow cytometry. Cytotoxic index was determined by the percentage of dead (PI-positive) cells among MC38/0-DiO cells.
Acknowledgements

This work was supported by grants from the Polish Ministry of Science and Higher Education (No N401 255334) and by the National Science Center of Poland (decision number DEC-2011/01/N/N24/01725).

References


Submitted: 15 December, 2011
Accepted after reviews: 22 February, 2012