Antitumor Effect of Embryonic Stem Cells in a Non-Small Cell Lung Cancer Model: Antitumor Factors and Immune Responses

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Abstract

Research in recent years has revealed that embryonic stem cells (ESCs) could generate obvious antitumor effects in both vitro and vivo. In vitro, ESCs could secrete soluble factors that are capable of blocking cancer cells proliferation, moreover, embryonic microenvironments could effectively inhibit tumorigenesis and metastasis; while in vivo, administration of ESCs in tumor-bearing mice could generate significant antitumor effects by indirectly activating the antitumor immune system. In this study, non-small cell lung cancer cells (Lewis Lung Carcinoma cells, LLCs) and ESCs were co-injected together into mice, after that subcutaneous tumor growth was monitored, cellular and humoral immune responses were detected, and different control groups were set to compare the results in different conditions. Our results suggested that compared to be injected alone, ESCs co-injected with cancer cells could inhibit cancer cell growth more efficiently in vivo, with more CD8+ lymphocytes generated in both peripheral circulation and spleen, and with higher serum anticancer cytokine level (interleukin (IL)-2 and interferon (IFN)-γ). We conclude that the boosted antitumor effects induced by ESCs and cancer cells co-injection may be both the effects of antitumor factors secreted by ESCs and immune responses induced by ESCs in vivo.

Key words: embryonic stem cells; immunotherapy; antitumor immunity; antitumor factors.

1. Introduction

Since Lobstein and Recamier put forward the hypothesis that neoplasm originates from embryo cells, investigators made great efforts in the comparative study between neoplasm formation and embryonic development in the last few decades [1]. They found that cancer cells and stem cells have some common biological properties, including unlimited proliferation, the ability of migration and invasion, and they have some similar gene products and signal transduction pathways [2-5]. Moreover, cancer cells and ESCs share some cell surface markers and antigens that not presented by adult tissues [6-8]. On the other hand, ESCs were applied to generate antitumor effects in both vitro and vivo. For example, ESCs could produce soluble factors arresting or slowing population growth of cancer cell lines [9], and embryonic microenvironments have the potential to reprogram cancer cells into a less invasive pheno-
type and help preventing tumorigenesis and metastasis [10, 11]. In vivo studies, mice vaccinated with ESCs could induce specific antitumor responses effectively protecting them from tumor formation and development [12]; even in mice with minor and heavy tumor load, administration of ESCs could generate effective antitumor immunity, which obviously improved the immune dysfunction and inhibit tumor growth [13].

So far, it is not hard to conclude that ESCs have double hit against cancer: direct inhibition by antitumor factors and indirect suppression by activating the body immune system. We hypothesized that these two aspects can give a combined effect when ESCs can get direct contact with cancer cells in vivo. In this study, mice were co-injected with ESCs and cancer cells simultaneously, and two different control groups were set. As expected, the results showed that ESCs co-injected with cancer cells could induce stronger anticancer immune responses in vivo, with more CD8+ lymphocytes generated and higher anticancer cytokine level (IL-2 and IFN-γ) compared to injected alone.

2. Results and Discussion

2.1 Inhibition of tumor growth by administration of ESCs

Lewis Lung Carcinoma cell is a well-established carcinoma cell line in mice [14], and 1×10⁶ cells were enough to induce subcutaneous tumor masses and eventual death. The tumor formation time and tumor growth of all groups of mice were closely monitored after LLCs administration (Fig.1A, C), and the tumor weight was recorded at the endpoint of experiment (Fig.1B). The results suggested that mice with ESCs injection (LLC+ESC and LLC+contra ESC group) generated obvious antitumor responses and rejected tumor masses from formation and development effectively compared to the LLC+MEF (Mouse Embryonic Fibroblast cells) group (p<0.05); moreover, mice in LLC+ESC group obtained stronger antitumor effects in inhibition of tumor masses compared with mice in LLC+contra ESC group (p<0.05), which means more significant antitumor effect could be induced when ESCs got direct contact with tumor cells in vivo.

Fig 1. (A) Tumor size of all groups was measured and tumor volumes were calculated after LLCs inoculation. The results indicated statistical difference in tumor size between mice injected with ESCs (LLC+contra ESC and LLC+ESC) and MEFs (LLC+MEF) from day 5 on (p<0.05), and between groups in which ESCs could get direct contact with cancer cells (LLC+ESC) or not (LLC+contra ESC) (p<0.05 from day 9 on). (B) All mice were sacrificed under anesthesia and tumor weight was detected at the end of the experiment. The results suggested significant difference in tumor weight between ESCs (LLC+contra ESC and LLC+ESC) and MEFs (LLC+MEF) groups, and between the LLC+ESC and LLC+contra ESC group (p<0.05). (C) Tumor formation time of all groups. The results suggested the tumor formation time was statistically different between the experimental (LLC+ESC) and control groups (LLC+MEF and LLC+contra ESC) (p<0.05). Error bars denote SD.
2.2 Increased proliferation of lymphocytes and secretion of cytokines

To explore the specific changes in immune system, we examined the proliferation of lymphocytes in the peripheral circulation and spleen of all mice induced by ESCs administration by flow cytometry and IHC in our study (Fig. 2 and 3). We found statistically significant proliferation of both CD4+ and CD8+ lymphocytes in both blood and spleen of mice injected by ESCs (LLC+ESC and LLC+contra ESC group) compared to the LLC+MEF group \( (p < 0.05) \). And compared to the LLC+contra ESC group, mice in LLC+ESC group generated more CD8+ lymphocytes in both peripheral circulation and spleen \( (p < 0.05) \).

To further study the changes in cellular immune responses induced by ESCs administration, we detected the level of immune important cytokines in serum of all mice. The concentration of IL-4, IL-2 and IFN-\( \gamma \) was detected by ELISA kits (Fig. 4), and the results suggested that mice in LLC+ESC and LLC+contra ESC group had significantly increased level of IL-2 and IFN-\( \gamma \) \( (p < 0.05) \), but no obvious change in the serum level of IL-4. Furthermore, compared with the LLC+contra ESC group, mice in LLC+ESC group had a statistically higher level of IFN-\( \gamma \) in serum \( (p < 0.05) \).

2.3 Increased cell apoptosis in tumor tissues

When the mice were sacrificed at the endpoint of the study, we detected the cell apoptosis in the tumor tissues by TUNEL kits (Fig. 5). We found more cell apoptosis in the LLC+ESC and LLC+contra ESC group compared to the LLC+MEF group \( (p < 0.05) \). Moreover, the apoptosis level of tumor cells in mice from LLC+ESC group was statistically different from mice in LLC+contra ESC group \( (p < 0.05) \).
There was statistical difference in the serum level of IL-2 and IFN-γ between mice injected with ESCs (LLC+contra ESC and LLC+ESC) and MEFs (LLC+MEF) ($p<0.05$), but no obvious change in the serum level of IL-4. Furthermore, compared with the LLC+contra ESC group, mice in LLC+ESC group had a statistically higher level of IFN-γ in serum ($p<0.05$).

The mean optical density for TUNEL cell apoptosis was $0.041\pm0.0034$ for LLC+MEF group, $0.073\pm0.0040$ for LLC+contra ESC group and $0.092\pm0.0047$ for LLC+ESC group. The results suggested statistical difference between mice injected with ESCs (LLC+contra ESC and LLC+ESC) and MEFs (LLC+MEF) ($p<0.05$), moreover, compared with the LLC+contra ESC group, mice in LLC+ESC group had a statistically higher level of cell apoptosis ($p<0.05$).

2.4 Discussion

Lung cancer is known to be the most commonly diagnosed cancer as well as the leading cause of cancer related death around the world [15]. The five-year survival rate remains less than 15% despite of improvement in surgery, radiotherapy and chemotherapy. There has been a strong requirement in new effective and better tolerated treatment strategies. Lack of appropriate lymphocytes activation for tumor rejection was thought to be an important reason why immune system failed to eliminate a tumor mass [16, 17]. The mechanisms for cancer cells to escape from immune recognition may be the alteration of proteins involved in antigen processing and presentation, loss of major histocompatibility complex class I expression and presence of immunosuppressive factors. And finally, insufficient activation of antitumor immunity lead to poor protection from tumors [18-20].

Since the first observation that embryonic materials could be used as vaccines to prevent tumor formation and development in animal models, there has been accumulative evidence supporting the fact that embryonic/fetal materials could generate effective antitumor responses in vivo [21]. Now it has been well proved that ESCs and their derivatives could generate effective cellular immune response and active CD4+ and CD8+ lymphocytes [22-24]. Because of the antigenic similarities between ESCs and tumor cells, immune responses against ESCs could cross-react with tumor cells, so antitumor immunity in vivo was generated [12, 25].

In our study, we found a strong and steady activation of antitumor cellular immune response by ESCs administration, we detected obvious proliferation of lymphocytes (in both peripheral circulation and spleen) and especially a dramatic increase of CD8+ lymphocytes (cytotoxic T lymphocytes) which was believed to consist the main part of antitumor immunity and play an crucial role in tumor rejection [26-28]. Moreover, increase in the serum level of Th1 cytokines (IL-2 and IFN-γ) secreted by lymphocytes
was also detected, these cytokines played an important role in regulating the antitumor immune responses and differentiation of cytotoxic T lymphocytes [29-31]. So we concluded that the induced immune responses were not only an increase in quantity of lymphocytes but also an enhancement in cellular immune function, which help to rebuild the host’s immunity against cancer [13]. And these antitumor immune responses lead to obvious retardation in the formation and development of tumor masses, including the delayed tumor formation time, inhibited tumor growth in both size and weight, and more cell apoptosis in tumor sections.

However, the more attractive part in this study was that, compared with the LLC+contra ESC group, mice in LLC+ESC group seemed to generate more effective antitumor responses when ESCs could get direct contact with cancer cells in vivo. In these mice, we found more significant inhibition in formation and growth of tumor masses, more effectively activation of lymphocytes proliferation and cytokine secretion, and significant difference in tumor cell apoptosis without obvious clinical signs of autoimmune disease. There are several potential mechanisms that could account for the boosted antitumor response in the LLC+ESC group. We believed that it was a complex result of both direct inhibition effects by ESCs and induced immune responses. As recent studies indicated that stem cells could secrete antitumor factors and inhibit cancer cells proliferation [9, 32, 33], and embryonic microenvironments have an essential role in preventing carcinogenesis by providing signals to inhibit malignant cells proliferation and promote differentiation [10, 11, 34]. On the other hand, the immunogenicity of ESCs played an important part in tumor rejection, which induced cellular immune responses, recruited cytotoxic T lymphocytes and lead to tumor cells apoptosis as specifically described above.

However, it is still difficult to illustrate a specific mechanism responsible for the rejection of tumors, as it is known to be a complicated one involving lymphocytes and other immune effector cells, soluble factors such as nitric oxide, chemokine ligand and cytokines [12, 35]. And some issues are still necessary to be explored for further experiments, such as more cell lines and multiple experiment are required to make a general statement, whether the immune responses are tumor specific, what kind of effector cells are involved in the co-injection site of ESCs and cancer cells, etc. But our study may introduce a new dimension on the antitumor application of ESCs and therapeutic strategy for cancer.

3. Materials and methods

3.1 Mice and cell lines

Wild type female C57 BL/6 mice were obtained from the Institute of Laboratory Animal Science, Chinese Academy of Medical Science, 4-6 weeks old on arrival and raised under special pathogen-free conditions for 2 weeks before use. The C57 BL/6 Mouse Embryonic Stem Cells were obtained from Cyagen Bioscience Inc., cultured on plates pre-coated with gelatin solution and γ-ray irradiated C57 BL/6 Mouse Embryonic Fibroblast cells (MEFs) (Cyagen Bioscience Inc.) as feeder cells, and grew in Mouse Embryonic Stem Cell Growth Medium (Cyagen Bioscience Inc.). The ESCs were in passage 8-10 approximately and pre-irradiated (15Gy/3 fractions) when used as cellular therapy. The Lewis Lung Carcinoma cells were purchased from the Cell Bank of Chinese Academy of Medical Science, cultured in Dulbecco’s Modified Eagle’s Medium (DMEM, Hyclone) supplemented with 10% Fetal Bovine Serum (FBS, Hyclone), and were in passage 10-15 when inoculated.

3.2 Reagent and antibodies

Most reagents used in cell culture were purchased from Hyclone Biochemical Products Co., Ltd unless specially indicated. The antibodies and reagents used in flow cytometry were obtained from ebioscience (San Diego, CA, USA), and antibodies used in immunohistochemistry were from BioWorld Technology Ltd (Barrie, ON, Canada) and Biosynthesis Biotechnology Co., Ltd (Beijing, China). The ELISA kits were obtained from Jingmei Biotech Co., Ltd (Beijing, China) and one step TUNEL apoptosis assay kits were purchased from Beyotime Biotechnology (Haimen, China).

3.3 Experiment protocols

Thirty female C57 BL/6 mice were randomly divided into three groups: mice in experimental group (LLC+ESC) (n=10) received subcutaneous administration of a mixture of 1×106 LLCs and 1×106 ESCs in the left axilla; mice in control group (LLC+MEF) (n=10) received subcutaneous injection of a mixture of 1×106 LLCs and 1×106 MEFs in left axilla; and mice in another control group (LLC+contra ESC) (n=10) received subcutaneous administration of 1×106 LLCs in left axilla and 1×106 ESCs in right axilla. Tumor sizes were measured by digital calipers in longitudinal (L) and transverse (W) diameters every other day since tumor cells inoculation and tumor volumes were calculated according to the formula \( V = \frac{L \times W^2}{2} \) [36]. The endpoint for this study was set to be three weeks after tumor cells administration, and then all mice were sacrificed under anesthesia and

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tumors were exercised and weighed. During the experiment, all mice were monitored for general health indicators: general behavior, feeding and defecation, body weight, neuromuscular tone, appearance of fur, etc. All animal experiments were carried out using a protocol approved by the Ethics Committee of Provincial Hospital affiliated to Shandong University, Shandong University.

3.4 Flow cytometry

Three days after the cell injection, citrate-anticoagulated blood was acquired from venous sinus of the eye orbits in mice from all groups as described [37]. Sampled blood was stained with antibodies for CD3, CD4 and CD8 positive lymphocytes and analyzed on an EPICS XL flow cytometer (Beckman Coulter Corp.), and the results were processed by the WinMDI software (Scripps Institute).

3.5 ELISA

Fresh blood was obtained from all mice at the same time and in the same way as previously described, and serum was collected. Quantification of IL-4, IL-2 and IFN-γ was performed using commercially available enzyme-linked immunosorbent assay kits in accordance with the instructions and analyzed on a micro-plate reader (Wellscan MK3, Labsystems Dragon).

3.6 Histology and immunohistochemistry (IHC)

All mice were sacrificed under anesthesia at the endpoint of the study, and the primary tumors and spleens were excised and fixed in 4% formaldehyde solution. Tissue samples were embedded in paraffin wax and serially sectioned. Tissue sections were deparaffinized and rehydrated through a graded ethanol series, followed by incubation with 3% hydrogen peroxide for 15 min. After 15-min rinses in three changes of Phosphate Buffered Saline (PBS), sections were blocked with normal goat serum to suppress nonspecific background staining, and incubated at 4°C overnight with diluted (1: 100) CD4 and CD8 antibodies. Then the sections were incubated with biotinylated goat anti-rabbit IgG for 30 min at room temperature and processed according to the manufacturer’s protocol, and finally examined under a Nikon E200 light microscope (Nikon, Tokyo, Japan) equipped with a digital camera system.

3.7 TUNEL

The tumor sections were deparaffinized and rehydrated through a graded ethanol series, incubated with protease K for 25 min at room temperature for antigen retrieval, and then operated according to the protocol of the TUNEL apoptosis detection kit. The FITC-labeled positive cells were imaged under a fluorescent microscope using 488-nm excitation and 530-nm emission [38]. Quantification of the positive expression of both immunohistochemistry and TUNEL was performed using the mean optical density method as described [13, 39].

3.8 Statistical analysis

All data were expressed as the mean ± standard error. Statistical significance between groups was determined by ANOVA test and Student-Newman-Keuls multiple comparisons test after confirming the equality of variance with the Levene test using SPSS 17.0 statistical software (SPSS Inc, Chicago, IL). For all tests, a p value less than 0.05 was considered to be statistically significant.

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Competing Interests

The authors have declared that no competing interest exists.

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