Targeting breast cancer stem cells by dendritic cell vaccination in humanized mice with breast tumor: preliminary results

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Abstract

Breast cancer (BC) is one of the leading cancers in women. Recent progress has enabled BC to be cured with high efficiency. However, late detection or metastatic disease often renders the disease untreatable. Additionally, relapse is the main cause of death in BC patients. Breast cancer stem cells (BCSCs) are considered to cause the development of BC and are thought to be responsible for metastasis and relapse. This study aimed to target BCSCs using dendritic cells (DCs) to treat tumor-bearing humanized mice models.

Materials and methods: NOD/SCID mice were used to produce the humanized mice by transplantation of human hematopoietic stem cells. Human BCSCs were injected into the mammary fat pad to produce BC humanized mice. Both hematopoietic stem cells and DCs were isolated from the human umbilical cord blood, and immature DCs were produced from cultured mononuclear cells. DCs were matured by BCSC-derived antigen incubation for 48 hours. Mature DCs were vaccinated to BC humanized mice with a dose of 10⁶ cells/mice, and the survival percentage was monitored in both treated and untreated groups.

Results: The results showed that DC vaccination could target BCSCs and reduce the tumor size and prolong survival.

Conclusion: These results suggested that targeting BCSCs with DCs is a promising therapy for BC.

Keywords: breast cancer, breast cancer stem cells, targeting cancer therapy, humanized mice, targeting cancer stem cells

Introduction

Breast cancer (BC) is the second leading cause of cancer-related death in women.¹ Despite advances in treatment methods, such as surgery, chemotherapy, radiation therapy, and biological therapy, the percentage of death in BC patients remains high. Although targeted therapies using antibodies, such as pertuzumab and trastuzumab, have significantly improved the treatment of BC in recent years,²⁻⁴ some investigations reported that 30%–70% of BC patients relapse after 5 years.⁵

In recent years, dendritic cell (DC) vaccination has emerged as a promising therapy for cancer treatment. DCs are professional antigen-presenting cells in the human body that originate from bone marrow precursors.⁶ In an immature state, DCs exhibit high endocytic activity and low T-cell activation. Upon contact with an antigen, they become mature and can strongly activate the T-cells via cell–cell contact or by producing a pool of cytokines.⁷ These cells highly express costimulators, major histocompatibility complex molecules (CD80, CD86), and CD40. Through interaction between CD40 (on DCs) and CD40 ligand (on T-cells), DCs can proliferate and present the
antigens to T-cells. In BC patients, DCs reportedly exhibit abnormalities that prevent them from efficiently presenting the tumor antigens to T-cells. In fact, it has been shown that DCs in cancer patients exhibit reduced antigen uptake, reduced antigen processing, low expression of costimulators, weak migration, and decreased interleukin-12 (IL-12) production. It was previously demonstrated that DCs in BC are dysfunctional and show weaker migration to lymph nodes, lower expression of human leukocyte antigen (HLA) and CD86, and lower ability to induce IL-12 secretion in vitro compared with those in healthy patients. To address these problems, DC therapy was used to produce a large number of functional DCs ex vivo. Specifically, both hematopoietic stem cells (HSCs) and monocytes were collected and induced to DCs using a cocktail of granulocyte-macrophage colony-stimulating factor (GM-CSF) and IL-4. Subsequently, the immature DCs were loaded with antigens in the form of DNA, RNA, proteins, peptides, or cell lysates to produce the mature DCs for further applications.

DC therapies have been used in both preclinical and clinical trials for various cancers, such as prostate cancer, multiple myeloma, renal cell carcinoma, pancreatic cancer, leukemia, melanoma, colorectal cancer, glioma, and BC. In almost all cases, DC vaccination was demonstrated to be a safe and effective method for treating metastatic patients. Importantly, some DC vaccinations have been approved by governmental regulatory agencies as official methods to treat cancers. For example, Sipuleucel-T has been approved by the United States Food and Drug Administration to treat human prostate cancer, and VacciCell has been approved by the Japanese Food and Drug Administration. To date, there are >289 clinical studies of DC-based cancer vaccines that are registered and under investigation (https://www.ClinicalTrials.gov). More importantly, among the 289 cases, six are in Phase III, and two are in Phase IV.

Recently, it has been shown that DC vaccination can improve BC treatment. For example, Brossart et al showed that DC vaccination with HLA-A2-restricted HER2 or MUC peptide-pulsed DCs induced immunologic responses in patients. However, the clinical efficacy of DC vaccination was not recorded in this study. In a separate study, Avigan et al fused DCs with BC cells and recorded immunological and antitumor responses. More recently, Qi et al adopted a novel inducing method for DCs by using tumor lysate, but the results were limited, with only a partial response. DC vaccine, in combination with IL-2 or IL-12, also recorded specific immunity against introduced antigen. In order to improve the outcome, some investigators combined DCs with cytokine killer cells and found significant improvement in the progresssion-free survival and overall survival of patients. Thus, to date, DC therapy has had limitations in the improvement of the clinical outcome.

Breast cancer stem cells (BCSCs) were discovered over 10 years ago, by Al-Hajj et al, and have been shown to be the cause of breast tumor development and the drivers of therapeutic resistance in BC. New therapies aimed at targeting BCSCs have shown an increase in patient outcome. Therefore, we hypothesized that the existence of BCSCs in tumors may be responsible for the low efficacy of DC therapy. In this study, we aimed to evaluate the preclinical trial efficacy of a DC vaccination from DCs primed from BCSC lysate using BC humanized mice models.

**Materials and methods**

**Animals, BCSCs, and umbilical cord blood**

NOD/SCID mice were bought from Jackson Laboratory (Charles River). Mice manipulations were approved by the Institutional Animal Care and Use Committee of Stem Cell Research and Application Laboratory, University of Science, Vietnam National University, Ho Chi Minh. All mice were housed in individual ventilated cages and were carefully monitored daily as The Institutional Animal Care and Use Committee guidelines (followed by Guide for the Care and Use of Laboratory Animals, Eighth Edition, National Institute of Health, US, published by The National Academies Press, Washington, DC, USA).

BCSCs were used from the previously published study. BCSCs were thawed and allowed to proliferate in suitable conditions. BCSCs were cultured in mammosphere medium without fetal bovine serum supplement (ie, Dulbecco’s Modified Eagle’s Medium/F12 supplemented with 1% (v/v) prostate-specific antigen, 2% (v/v) B-27 supplement, 20 ng/mL epidermal growth factor and basic fibroblast growth factor, 4 ng/mL heparin, and 10 μg/mL insulin) in 10% O2, 5% CO2, as published previously. BCSCs were validated by flow cytometry using the surface markers with phenotype CD44+CD24- before being used in the experiments. Briefly, BCSCs were stained with both anti-CD44 monoclonal antibody conjugated with antigen-presenting cells and anti-CD24 monoclonal antibody conjugated with fluorescein isothiocyanate (FITC) (BD Biosciences, San Jose, CA, USA). Stained cells were analyzed in FASCalibur machine with CellQuest Pro at 10,000 events.

Umbilical cord blood (UCB) was collected as described previously. Briefly, UCB was collected from the umbilical cord vein with informed consent from the mother. The collection was performed in accordance with the ethical standards of the local ethics committee (Van Hanh General Hospital, Ho Chi Minh City, Vietnam).
Isolation of HSCs from UCB
In this study, HSCs were used as unpurified, mononuclear cells (MNCs) that were isolated from UCB. To isolate MNCs, each UCB unit was diluted into a ratio of 1:1 with phosphate-buffered saline (PBS), and 10 mL of diluted blood was loaded onto 25 mL Ficoll Hypaque solution (1.077 g/mL; Code 10771; Sigma-Aldrich Co., St Louis, MO, USA) in a 50 mL tube. After centrifuging at 2,500 rpm for 5 minutes, MNCs were derived from the interphase layer and washed twice with PBS.

To determine the dose of HSCs for transplantation, the obtained MNCs were used to enumerate the HSCs. The number of HSCs was determined using an Enumeration Pro-Count Kit (BD Biosciences) following the manufacturer’s guidelines.

Humanized mice
NOD/SCID mice were intraperitoneally injected with busulfan (25 mg/kg) prepared in dimethyl sulfoxide. After 48 hours, total white blood cells (WBCs) and body weight were measured for each mouse. Only mice with WBCs <1,000 cells/mL were used to graft HSCs. Mice were anesthetized using ketamine (100 µL/mouse). Mouse tails were relaxed in warm water (37°C) for 3–5 minutes. Tails were then cleaned with 70% alcohol. HSCs (1×10⁶) in MNCs were transfused into the tail vein with a total volume of 0.5 mL at a rate of 10 mL/h using an electronic pump.

The presence of human cells was determined by measuring the percentage of human leukocytes (CD45 cells) in the peripheral blood. HSC localized in the bone marrow was determined by the existence of human HSCs (CD34 cells) in the bone marrow. HSC differentiation was determined based on the presence of CD3, CD4, CD8, CD1a, and CD56 in the peripheral blood. To determine the presence of CD1a, CD3, CD4, CD8, CD34, CD56, and CD1a, peripheral blood was collected at the tail vein with anticoagulation agent (ACD-A; BD Biosciences). Blood samples were stained with antibodies for CD1a, CD3, CD4, CD8, CD34, CD56, and CD45 conjugated with FITC for 20 minutes at room temperature. Then, blood was lysed with Pharm Lyse Lysing buffer (BD Biosciences). Lysed blood was diluted with 100 µL sheath fluid for analysis in a FACSCalibur machine. The human immunoglobulin G (IgG) was detected and measured in peripheral blood using enzyme-linked immunosorbent assay (ELISA) technique.

BC humanized mice
Humanized mice were injected with 1×10⁶ BCSCs with GFPs into the fat pad to produce BC humanized mice. Mice were monitored for tumor growth and invasion. BCSCs were transduced with GFP as described in a previously published study.⁴¹

Immature DC production
MNCs were cultured in RPMI 1640 supplemented with 10% fetal bovine serum, 20 ng/mL GM-CSF, 50 ng/mL IL-4, 1% l-glutamine, and 1% penicillin/streptomycin (Sigma-Aldrich Co.). At day 3 of culturing, 75% of the medium was removed, and fresh medium was added. On day 6, DC maturation was induced by supplementing the complete medium with 10,000 ng/mL BCSC-derived lysate and tumor necrosis factor-α (100 ng/mL). At day 3, cells were harvested and used in subsequent experiments.

To evaluate the phenotype of mature DCs, the differentiated cells were stained with specific surface markers CD40 (CD40-FITC), CD80 (CD80-PE-Cy7), CD86 (CD86-PE), and HLA-DR (HLA-DR-FITC) in staining buffer (PBS pH 7.4, 0.5% bovine serum albumin, 0.02% azide). The stained cells were analyzed by BD FACSCalibur flow cytometer.

T-lymphocyte proliferation stimulated by DCs and measurement of IL-12 were evaluated as described in a previously published study.⁴² There were five experimental groups with different ratios of DCs:lymphocytes (0.25:100, 0.5:100, 1:100, 2:100, and 8:100) and three control groups with DCs + phytohemagglutinin (PHA), PHA alone, or PHA + lymphocytes. The T-lymphocyte concentration was measured by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide assay kit according to the manufacturer’s instructions (Sigma-Aldrich Co.). Optical density values were read at a wavelength of 490 nm with the reference wavelength of 620 nm. The stimulation ability of DCs was calculated based on A-values. A-values were offset from optical density values measured for control samples (lymphocyte + PHA) and experimental groups. To measure IL-12 concentration, mature DCs were incubated in the fresh culture medium in a 24-well plate for 24 hours. Then, supernatants were collected and frozen at −80°C until analysis. IL-12 concentration in the supernatants was measured by ELISA kits (IL-12 High Sensitivity Human ELISA Kit; Abcam, Cambridge, UK), and the results were analyzed with the DTX880 Multimode Detector (Beckman Coulter, Inc., Brea, CA, USA).

BCSC antigen production
BCSCs grown to 70%–80% confluence were detached with 0.25% trypsin/ethylenediaminetetraacetic acid and washed two times with Dulbecco’s PBS. BCSC lysates were generated from 5×10⁶ tumor cells by three rapid freeze-thaw
cycles (7 minutes in liquid nitrogen and 7 minutes in the water bath). The tumor cell lysates were then spun at 700 rpm at 4°C for 10 minutes to remove cellular debris, and the supernatant was collected. All lysates were stored at -80°C until use.

**DC vaccination and efficiency monitoring plan**

On day 3, tumor-bearing mice were treated with intravenous injections of mature DCs in 100 µL of PBS. Tumor-bearing mice were sorted into two groups: 1) the treated group (ten mice), which was injected with $1 \times 10^6$ primed mature DCs (only one time), and 2) the untreated control (ten mice), which was injected with 100 µL of PBS.

From day 0 until day 30 of treatment, tumor size was measured using calipers in two dimensions, and size was calculated using the following formula: $a \times b^2/2$, where “a” is the tumor length and “b” is the diameter. The survival percentages of the mice were recorded in both treated and untreated groups during the 120 days of treatment.

**Statistical methods**

The results were expressed as the mean ± SD. One-way analysis of variance and two-tailed tests were utilized for all statistical analyses performed with GraphPad Prism software, Version 4.0 (GraphPad Software, Inc., La Jolla, CA, USA). $P$-values <0.05 were considered statistically significant.

**Results**

**Humanized mice models**

In order to generate humanized mice, we treated NOD/SCID mice with 25 mg/kg busulfan in order to destroy the murine bone marrow. WBCs strongly decreased from $1,788\pm655$ cells/µL to $763\pm571$ cells/µL, while in control mice, WBCs did not significantly change ($1,659\pm571$ cells/µL to $1,518\pm207$ cells/µL). Busulfan also strongly decreased CD4 cells from $11.26\pm2.45\%$ to $1.83\pm0.70\%$. However, the percentage of CD8 cells nonsignificantly reduced compared with that before busulfan injection.

Ten NOD/SCID mice in the experimental group were injected with HSC via the tail vein, and ten NOD/SCID mice in the control group were injected with PBS. HSCs were counted based on the expression of CD45 as markers (Figure 2E–J). The results showed that after 2 weeks, the human CD45 cell percentage changed between the ten mice with large amplitude. The highest percentage of human CD45 leukocyte in peripheral blood achieved 37% in WBCs isolated from peripheral blood at day 28 after HSC transplantation.

After 28 days of transplantation, transplanted mice were used to evaluate the existence of CD34 cells in bone marrow (Figure 2C and D). After staining with anti-human CD34-FITC, the results showed that 100% of mice had HSC homing to the bone marrow. Compared with the total of HSCs in bone marrow, the results showed that the highest human HSCs in bone marrow achieved 38.58% (n=3). In addition, the existence of different kinds of human leukocytes was investigated, using CD3, CD4, CD8, CD19, CD45, and CD56 as markers (Figure 2E–J). The results showed that after 35 days of transplantation, the leukocytes appeared in the peripheral blood, and the percentages of which rapidly increased in 42 days, 49 days, and 55 days.

The function of human B-cells was also evaluated using human IgG in peripheral blood. The human IgG concentrations at 42 days, 49 days, and 55 days were $78\pm9.6$ µg/mL, $89\pm13.1$ µg/mL, and $110\pm15.6$ µg/mL, respectively. However, the HSC transplantation significantly reduced survival in mice. Compared with the control group, after 120 days, there were 20% of mice alive, while only 6.67% of mice alive in the experimental group. HSC transplantation also caused mice to die earlier than in the control group. In fact, 75 days after transplantation, only ~26.67% of HSC-transplanted
mice were still alive, while 73.33% of control mice were still alive.

BC humanized mice

At day 30 after HSC transplantation, humanized mice were injected with $1 \times 10^6$ BCSCs into the fat pad. The results showed that 100% of humanized mice formed tumors, and invasion appeared at day 60 after BCSC transplantation (Figure 3A and B). More importantly, our results showed that BCSCs invade into the bones of 100% of the mice (Figure 3C). The results presented in Figure 3D–E clearly showed that the BCSC transplantation also caused mice to die earlier than in the control group.

UCB-derived DCs

To match HLA between DCs and leukocytes in humanized mice, both DCs and HSCs were isolated from human UCB. MNCs isolated from human UCB were induced by GM-CSF, IL-4, and BCSC-derived antigens for 10 days. The results are presented in Figure 4 and show that the DCs exhibited important characteristics of functional DCs. Specifically, they expressed some costimulator markers, such as CD40, CD80, CD86, and HLA-DR. After phagotogizing with BCSC antigen, they strongly expressed CD40, CD80, CD86, and HLA-DR (Figure 4E–H). More importantly, they successfully stimulated the allogenic T-lymphocytes (Figure 4I) and produced IL-12 (Figure 4J).
DC vaccination lengthened the survival of BC humanized mice

Changes in tumor size

In our previous study, we found that the tumors in NOD/SCID metastasized after 30 days of treatment. Therefore, in this study, we only monitored the tumor size after treatment with DCs in the first month (30 days) and from day 30 to 75 days of treatment; the efficiency of treatment was evaluated by the survival percentage of mice.

In the first month, tumors clearly responded with DC treatment. Before treatment, the tumor size was about 0.0954±0.0046 cm³ when injected with 1×10⁶ BCSCs into the fat pad. These tumors strongly increased in size in control mice, while they slowly increased their size in the treated groups. In fact, at day 30, the tumor size in the control mice increased from 0.044±0.003 cm³ to 0.745±0.114 cm³, while in treated mice, the tumor size slightly increased from 0.043±0.007 cm³ to 0.196±0.091 cm³. Taken together, we found that the tumor size increased eight times in control mice in the first month, while the tumor size only doubled in treated mice.

Changes in mice body weight in control and treated groups

The results showed that there was a slight change in the weight of the treated mice compared with that of control mice. Specifically, in control mice, the weight was maintained around 25 g during the first 30 days of treatment, while in treated mice, the weight of the mice slightly increased in week 1 after vaccinations and the mice maintained the weight during the first month. In the next month, weights of the mice gradually decreased, particularly in mice that eventually died.
In almost all cases, the deceased mice weighed between 15 g and 20 g in body weight.

**BCSC-primed DCs vaccination changed the survival time of treated mice**

All mice were monitored during the 120 days from the first day of the experiment. Figure 5 showed that the BCSC-primed DCs changed the percentage of survival time of treated mice ($P<0.05$). In fact, after 87 days of experiment, 70% of treated mice prolonged their survival, while only 40% of mice survived in the control group. There was 10% (one of ten mice) of mice in the treated group still alive after 120 days of monitoring, while 100% of mice in the control group died.

**Discussion**

BC is one of the most common cancers in women and is the second leading cause of death in women. Despite recent advances in the treatment of this disease, the percentage of relapse remains high. In particular, the existing therapies are unable to cure patients with metastases. Therefore, novel therapies are needed to overcome the current paradigm. DC vaccination has been suggested as a suitable strategy for the prevention of relapse in combination with radiation and chemotherapies. Therefore, DC vaccination has been used in the treatment of some diseases, included BC.

However, a recent analysis about DC vaccination in BC treatment showed that the efficacy of this therapy is much lower compared with the same in other cancers, such as prostate cancers or melanoma cancer. We proposed that the treatment efficacy depended on antigens used to induce DCs. This study aimed to investigate the treatment efficacy of BC using BCSC-derived antigen-primed DCs in the BC humanized mice models.

In the first experiment, the humanized mice models were produced by human HSC transplantation to replace the mouse’s blood system. HSCs were collected from human UCB. NOD/SCID mice were transplanted with HSCs according to a previously reported dose. In this study, we showed that busulfan could efficiently destroy the murine bone marrow. Some previous studies also succeeded to produce humanized models with busulfan as a bone marrow-destroying agent.

Although radiation can absolutely kill the murine HSCs, this method can cause a high percentage of death in...
weight of the mice increased after mice receiving human HSCs, suggesting that the human HSCs could maintain a healthy hematopoietic system in the mouse. This observation was confirmed by the existence of human CD45 cells in the peripheral blood, and this cell population gradually increased after 2 weeks of HSC transplantation. More importantly, in the WBCs in the peripheral blood, there were 50% of cells expressing CD45, which was slightly higher than was reported in a previous study.48 According to Hayakawa et al, after receiving 2×10^6 HSCs for 5–8 weeks, ~60% of human CD45 cells were detected in the peripheral blood of the mice.

The gold standard in HSC transplantation is homing of HSC to bone marrow to recover the blood system. The mice. According to Hayakawa et al,49 a radiation dose of 91.7 cGy/min significantly reduced mice survival. Hayakawa et al used busulfan to replace the radiation and showed higher efficacy in humanized mice creation. In our study, at a dose of 25 mg/kg of busulfan, mice had a significant reduction in body weight, as well as a reduction in WBCs. The results showed that busulfan exhibited its effects on the blood system. In fact, busulfan is an alkylation agent that interferes with the cell division, especially HSCs in murine bone marrow. By flow cytometry analysis, the results also showed that busulfan efficiently reduced the CD4 T-cells compared with CD8 T-cells.

Human HSC transplantation to busulfan-treated NOD/SCID mice rescued the murine bone marrow. In fact, body weight of the mice increased after mice receiving human HSCs, suggesting that the human HSCs could maintain a healthy hematopoietic system in the mouse. This observation was confirmed by the existence of human CD45 cells in the peripheral blood, and this cell population gradually increased after 2 weeks of HSC transplantation. More importantly, in the WBCs in the peripheral blood, there were >50% of cells expressing CD45, which was slightly higher than was reported in a previous study.48 According to Hayakawa et al, after receiving 2×10^6 HSCs for 5–8 weeks, ~60% of human CD45 cells were detected in the peripheral blood of the mice.

The gold standard in HSC transplantation is homing of HSC to bone marrow to recover the blood system. The
results showed that >50% of CD34+ cells in bone marrow were human CD34+ cells. The existence of human CD34+ cells in bone marrow helped peripheral blood CD45+ cells increase with time. More importantly, human HSCs in the bone marrow successfully differentiated into leukocytes, including T-cells (CD3, CD4, CD8, CD45), B-cells (CD19), and natural killer cells (CD56). In peripheral blood, human IgG was also detected by ELISA, demonstrating that human HSCs in the bone marrow successfully differentiated into leukocytes, including T-cells (CD3, CD4, CD8, CD45), B-cells (CD19), and natural killer cells (CD56). In peripheral blood, human IgG was also detected by ELISA, demonstrating that human HSCs in the bone marrow successfully differentiated into leukocytes, including T-cells (CD3, CD4, CD8, CD45), B-cells (CD19), and natural killer cells (CD56).

Subsequently, BCSCs with GFP as a reporter were injected into the fat pad to produce the BC humanized models. In line with the characteristics of BCSCs, the injected cells formed tumors in humanized mice. Based on GFP signal, after 30 days, dead mice revealed that the breast tumors had metastasized into the bone.

The results showed that BCSC-derived antigen-primed DC vaccination significantly inhibited the tumor growth and mice survival. The first evidence of the effects of the DC vaccination was tumor regression, compared with control mice. While the tumor size significantly increased in the control group, tumors in the treated mice maintained or slightly increased in size. The beneficial effect of DC vaccination was also recorded via body weight. In treated mice, their body weight increased. The most important results of BCSC-derived antigen-primed DC vaccination was that the BC mice survival was significantly prolonged after vaccination.

The main limitation of this study was noncomparison of treatment efficacy between BCSC-derived antigen-primed DCs and BC cell-derived antigen-primed DCs in breast tumor-bearing humanized mice models. However, these results showed that targeting BCSCs exhibited a good effect on inhibiting tumor progression and prolonged survival of breast tumor-bearing humanized mice models.

These results indicated that DCs successfully presented BCSC-derived antigens to other WBCs, such as T-cells and B-cells. These activated WBCs attacked tumor cells or BCSCs and inhibited the tumor progression, as well as tumor invasion. In a recent study, Kai et al used LBH589 and salinomycin to target BCSCs in mice model. They showed that in xenograft mouse models treated with LBH589 and salinomycin, the drug combination effectively and synergistically inhibited tumor growth of ALDH1-positive cells. Using another strategy to target BCSCs, Marcato et al successfully inhibited the BCSCs by oncolytic reovirus. Oncolytic reovirus has the potential to induce tumor regression in BC. Targeting BCSCs by Ad.mda-7 also showed that Ad.mda-7 inhibited tumor growth associated with a decrease in proliferation and angiogenesis in nude mice models.

**Conclusion**

Targeting BCSCs is an important strategy to treat BC. DC therapy is a promising alternative approach to targeting BCSCs. This study showed that DC-primed BCSC-derived antigens efficiently inhibited the tumor progression and prolonged the survival of BC mice. Moreover, this study also showed that the BC humanized mice models were feasible models to study and evaluate the DC therapy. Although further studies need to be performed to determine the mechanism and side effects of BCSC-targeting DC-based therapy, these results suggest that BCSC-derived antigen-primed DC therapy is a promising therapy for BC treatment.

**Acknowledgments**

This study was funded by Ministry of Science and Technology under grant number DTDL.2011-T/30.

**Author contribution**

PVP performed HSC isolation, developed the humanized NOD/SCID and DC vaccination treatment. HTL and BTV
carried out humanized mice. VQP and PML prepared the DCs. NLCP and NVT prepared the BC cells and BCSC-derived antigens. HTLN and STN analyzed the flow cytometry and monitored the treatment efficacy. TLN and NKP took care of the NOD/SCID mice, performed DC vaccination. All authors contributed toward data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

Disclosure
The authors report no conflicts of interest in this work.

References