Fusion of Glioblastoma Tumor Antigens to Herpes Simplex Virus-1 Glycoprotein D Enhances Secondary Adaptive Immune Responses in a DNA Vaccine Strategy

Rios WM1, De Molfetta JB2, Brandão IT1, Masson AP1, Peripato R2, Silva ID2, Rodrigues RP3, Arnoldi F1, De Souza PRM3, Diniz MDO4, Ferreira LCDS4 and Silva CL1

1The Center for Tuberculosis Research, Department of Biochemistry and Immunology, School of Medicine of Ribeirao Preto, University of Sao Paulo, Brazil
2Farmacore Biotechnology, Brazil
3University Federal of Sergipe, Brazil
4Department of Microbiology, Biomedical Sciences Institute, University of Sao Paulo, Brazil

Received date: Oct 27, 2015; Accepted date: Dec 04, 2015; Published date: Dec 08, 2015

Copyright: © 2015 Rios WM, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Abstract

Glioblastoma multiforme (GBM) is a recurrent and fatal cancer. EGFRvIII, MAGE-3 and GLEA-2 are antigens that are found in this highly heterogeneous tumor and that are absent in normal tissue. Usually, conventional GMB treatments do not prevent recurrence, reinforcing the need for new therapeutic strategies. Vaccination can be an alternative GMB therapy capable to induce long-lasting and specific immune responses to tumors antigens but requires the activation of strong cellular responses. Fusion of tumor antigens with microbial-derived proteins is a rather simple approach that can enhance the immunogenicity of vaccines, particularly, DNA vaccines. In this study, we constructed DNA vaccines encoding GBM tumor antigens fused to glycoprotein D from herpes simplex virus-1 and evaluated their immunogenicity in C57BL/6 mice. The tumor antigens were correctly expressed by the DNA vaccines and induced cell mediated immune responses under experimental conditions. The vaccines encoding antigen genetically fused with gD induced higher cellular immune responses, associated with IFN-γ and IL-10 production, than vaccines encoding non-fused GMB antigens. Therefore, DNA vaccinations induced a Th1-biased immune response. We concluded that the strategy could be an effective immunotherapeutic approach for GBM.

Keywords: Glioblastoma antigens; Immunotherapy; DNA vaccine; Glycoprotein D; IFN-γ; IL-10

Abbreviations
gD: glycoprotein D; EGFRvIII: Variant of the type I epidermal growth factor receptor; MAGE: Melanoma antigen gene; GLEA: Glioma-expressed antigen; TMZ: temozolomide; BBB, blood-brain barrier; KLH, keyhole limpet hemocyanin; CT: Cancer testis; HVEM: Herpes virus entry mediator; BTLA: B and T-lymphocyte attenuator; MDSCs: Myeloid-derived suppressor cells.

Introduction

Glioblastoma multiforme (GBM) (WHO grade IV astrocytoma) is the most common and malignant primary cancer in the central nervous system (CNS) [1-3]. Despite the conventional and aggressive treatment, including maximal surgical resection followed by temozolomide (TMZ) and radiation, the prognosis for patients with GBM remains poor, with a median survival of less than 15 months [4,5]. One great challenge in developing GBM therapies is the complexity of the GBM microenvironment [6]. The typical characteristics of this disease include uncontrolled cellular proliferation, diffuse infiltration, necrotic tendency, significant angiogenesis, apoptosis resistance and tumor heterogeneity [7]. Furthermore, patients with GBM display profound immunosuppression [8], which seems, partially, triggered by the cancer cells, as reported by Wei and colleagues [9]. Therefore, effective therapies must not only be directly cytotoxic to a molecularly diverse tumor cell population but also must overcome the pro-tumorigenic properties of the GBM microenvironment. Thus, studies have actively focused on testing new therapeutic approaches, including immunotherapy. Active immunotherapy, represented by therapeutic anti-cancer vaccines, is a primarily attractive approach to fight different forms of cancer since it provides the advantage of cellular specificity and generation of long-term immune surveillance to recurrent cancer cells [2]. Furthermore, the change in the concept that the CNS is immunologically privileged has generated enthusiasm for a potential role for immunotherapy in GBM. The primary line of active immunological defense for the brain is composed of specialized resident cells called microglia [10]. These cells are activated when they migrate toward inflammatory areas, where they obtain phagocytic properties and produce cytokines and chemokines, allowing the recruitment of other immune cells [10]. Additionally, macrophages and dendritic cells (DCs) can act as powerful antigen-presenting cells (APCs) in the CNS [11,12]. Some studies have shown that antigens originating within the CNS are drained in the cerebrospinal fluid to nasal and cervical lymph nodes where adaptive immune reactions are initiated [13,14]. Moreover, subpopulations of activated T cells expressing a particular phenotype 4/7 integrin overexpression exhibit tropism for the brain and cross the blood-brain barrier (BBB) [15]. Taken together, these findings represent the evolution of the knowledge regarding the interactions between CNS immune cells and GBM, stimulating the development of immunotherapeutic strategies to improve the survival of GBM patients.
between the CNS and the immune system and strengthen the idea that immunotherapy can be an effective weapon in GBM treatment.

Selection of good antigens remains a considerable challenge both for passive and active immunotherapies for cancer. Some antigens associated with GBM have been described. EGFRvIII, the most common and well-characterized variant of the type I epidermal growth factor receptor (EGFR), was first identified in primary human GBM [16,17].

This variant form is constitutively active, and its expression confers tumorigenic properties to the cell [18,19]. Clinical trials have been performed with a 14-mer-derived peptide EGFRvIII, PEPvIII [16], conjugated to KLH (keyhole limpet hemocyanin), delivered like peptide-based vaccines or pulsed with DCs [20-23]. The results showed that this strategy could induce antigen-specific cellular and humoral immune responses and increase patient median survival [22,23]. The MAGE-3 (melanoma antigen gene) antigen has also been described in GBM. Chi et al. showed that MAGE-3 mRNA was present in 33% of the GBM cases but it is absent in normal tissue [24]. This antigen, which is a member of the MAGE family, is characterized as a cancer-testis (CT) antigen that is expressed in normal testis and several tumors [25-27]. Other researchers have identified MAGE-3 peptides that can be recognized by CD4+ T cells of individuals with or without cancer [28,29], indicating that this antigen may be the target of an immune response. The GLEA-2 (glioma-expressed antigen 2) antigen, identified in a GBM cDNA library [30], is immunogenic and considered a putative GBM antigen. Fischer and collaborators showed that 43% of the studied GBM patients presented antibodies against this antigen [30]. Furthermore, patients undergoing radiotherapy presented increased frequencies of anti-GLEA-2 antibody [31].

Although all these characteristics indicate that these antigens can be potential targets for GBM immunotherapy, the immunogenicity of cancer antigens is usually limited. Therefore, these antigens must be combined with other substances to achieve the desired response. The genetic fusion of glycoprotein D (gD) from herpes simplex virus-1 (HSV-1) to tumor antigens encoded by DNA vaccines has been used as an alternative to increase the immunogenicity of proteins, generating increased frequencies of anti-GLEA-2 antibody [31].

Materials and methods

Construction of DNA vaccines

The coding sequences were provided from human proteins named EGFRvIII, MAGE and GLEA. The sequencing data were submitted to the GenBank databases under accession numbers NM_005228 (EGFRvIII), NM_005362.3 (MAGE), and AF258787 (GLEA). These sequences without their stop codons were combined to the gD nucleotide sequence in silico. The fusion was made between two unique sites (PvuI and Apal) that are present in gD [45]. The gDEGRvIII, gDMAGE, gDGLEA fragments were constructed by Epoch Biolabs (Missouri, TX) and cloned into the pBluescript II SK vector (pBSK). The pBSKgDEGRvIII, pBSKgDMAGE and pBSKgDGLEA plasmids were cleaved with NheI and XbaI restriction enzymes (Fermentas, Thermo Scientific, Cat. #FD0974, #FD0684), whose sites were included at the beginning and end, respectively, of all fragments. The cleaved gDEGRvIII, gDMAGE, gDGLEA fragments were cloned into the pVAX (Invitrogen, Cat. V260-20) vector. Cloning was confirmed by restriction enzyme digestion, PCR and DNA sequencing. The pBSKgDEGRvIII, pBSKgDMAGE and pBSKgDGLEA plasmids were also used as templates for EGFRvIII, MAGE and GLEA amplification. Reactions were performed using the following primers (IDT, Integrated DNA Technologies):

Forward

- EGFRpVAX_S
  - 5’-TACTAGCTAGCACATGGCTGAGGAAAAGAAAGTATATTATGTTGACA-3’
- MAGEpVAX_S
  - 5’-TACTAGCTAGCACATGGCTGAGGAAAAGAAAGTATATTATGTTGACA-3’
- GLEApVAX_S
  - 5’-TACTAGCTAGCACATGGCTGAGGAAAAGAAAGTATATTATGTTGACA-3’

Reverse

- EGFRpVAX_R
  - 5’-ATTTCTAGATCAGCGTGCCCTTCTGAGCGAG-3’
- MAGEpVAX_R
  - 5’-ATTTCTAGATCAGCGTGCCCTTCTGAGCGAG-3’
- GLEApVAX_R
  - 5’-ATTTCTAGATCAGCGTGCCCTTCTGAGCGAG-3’

In the forward primer sequences, the bold letters represent the Kozak sequence, and the Nhel restriction site is underlined. In the reverse primers, the bold letters represent stop codons, and the XbaI restriction site is underlined. The conditions for PCR were as follows: a 100 µL PCR mixture composed of 100-150 ng of template, 0.2 mM each dNTP (Invitrogen, Cat. 10297-018), 0.1 mM each primer and 5 units of High Fidelity PCR enzyme mix (Fermentas, Thermo Scientific, Cat. #K0192). The following amplification program was used: 94°C for 5 min, 24 cycles of 94°C for 1 min, 48°C for 1 min, and 72°C for 1 min (GLEA, 2 min), with a final step at 72°C for 10 min. The amplified EGFRvIII, MAGE and GLEA fragments were cleaved with Nhel and XbaI restriction enzymes and cloned into pVAX without gD. Cloning was confirmed by restriction enzyme digestion, PCR and DNA sequencing. Large-scale purification of the DNA vaccines pVAXgDEGRvIII, pVAXgDMAGE, pVAXgDGLEA, pVAXEGFRvIII, pVAXMAGE and
pVAXGLEA was conducted by ion-exchange chromatography using an Endofree Plasmid Giga Kit (Qiagen, Cat.12391) according to the manufacturer's instructions. DNA vaccines were quantified by spectrophotometry (Nanodrop 1000®, Thermo Scientific, Wilmington, DE 19810 USA) and analyzed for the endotoxin presence using a QCL-1000 LAL Kit (Cambrex, Cat. 50-647U) according to the manufacturer's instructions.

Construction of expression vectors

The pBSKDEGFRvIII, pBSKDMAGE and pBSKDGLEA plasmids were used as templates for EGFRvIII, MAGE and GLEA amplification. Reactions were performed using the following primers:

**Forward**
- EGFRepET28a_S
- TACTAGCTAGCAGCTGCCTCTTGAGCAGAG-3'
- MAGEpEpET28a_S
- TACTACTAGCAGCATGCCCTCTTGACGAG-3'
- GLEApET28a_S 5’-TACTAGCTAGCATGAAACATCATCCACC-3'

**Reverse**
- EGFRepET28a_R 5’-GATCTCGAGTCAGCAGTGGGGGCCGTC-3'
- MAGEpET28a_R 5’-GATCTCGAGTCAGCTGCTCCCTCCTC-3'
- GLEApET28a_R 5’T-GATCTCGAGTCAGCTGCTCCCTCCTGAC-3'

In the forward primer sequences, the underlined letters represent the Nhel restriction site. In the reverse primer sequences, the bold letters represent stop codons, and the Xhol restriction site is underlined. The conditions for PCR were as follows: a 100 μL PCR mixture composed of 100-150 ng of template, 0.2 mM each dNTP, 0.1 mM each primer and 5 units of High Fidelity PCR enzyme mix. The amplification program was the same as described above. The amplified EGFRvIII, MAGE and GLEA fragments were cleaved with Nhel and Xhol (Fermentas, Thermo Scientific, Cat. #ER0695) restriction enzymes and cloned into the pET28a vector (pET) (Novagen, Cat. 69864-3). Cloning was confirmed by restriction enzyme digestion, PCR and DNA sequencing.

Expression and purification of the recombinant proteins

The EGFRvIII, MAGE and GLEA fragments were cloned into the pET, which encodes the N-terminal 6XHis-tag. The E. coli BL21 and Rosetta strains were transformed with the pETEGFRvIII, pETMAGE, pETGLEA plasmids. For the production of the recombinant proteins EGFRvIII, MAGE and GLEA, the transformed bacteria were grown at 37°C in LB broth (USB, Cat. 75852) supplemented with 50 µg/mL kanamycin (Gibco, Cat. 905037IP) to an optical density of 0.5-0.6 at 600 nm (OD600=0.5-0.6), and protein expression was induced with 0.5 mM IPTG (isopropyl-β-D-galactoside) (Invitrogen, Cat. 15259-019) at 30°C overnight (MAGE was induced with 0.5 mM IPTG at 37°C for approximately 4 h, and GLEA was induced with 0.2 mM IPTG at 37°C for approximately 4 h). The cell pellets were collected, analyzed by 10% SDS-PAGE with Coomassie Brilliant Blue and visualized using the peroxidase substrate (46), with Coomassie Brilliant Blue R-250 (USB, Cat. 32826) and used for protein purification.

Recombinant proteins were purified by affinity chromatography on Ni-NTA (GE Healthcare, Cat. 17-5318-02) resin using 6XHis-tag according to the manufacturer's protocol and analyzed by 10% SDS-PAGE with Coomassie Brilliant Blue. The EGFRvIII and GLEA recombinant proteins were purified under denaturing conditions (8 M urea); MAGE, under native conditions. The purified protein fractions were dialyzed against PBS (phosphate-buffered saline), concentrated using a GE Healthcare Vivaspin (Cat. 28-9323-61) and quantified using a Coomassie PlusTM Protein assay kit (Pierce, Thermo Scientific, Cat. #1856210) [47]. When the EGFRvIII and GLEA proteins precipitated after dialysis, they were centrifuged, and the supernatants were used in the assays.

Production of antibodies against the recombinant proteins

Female BALB/c mice at 6-8 weeks of age from the Animal Breeding Center of the School of Medicine of Ribeirao Preto University of Sao Paulo were used in this protocol. All procedures involving the handling and killing of animals were performed according to approved protocols and in accordance with recommendations for the proper use and care of laboratory animals at the School of Medicine of Ribeirao Preto-University of Sao Paulo (number 082/2009). Three mice were immunized subcutaneously in the back with 50 µg of recombinant protein (MAGE or GLEA) or precipitated protein (EGFRvIII) emulsified with Freund's incomplete adjuvant (Sigma, Cat F5605) at a ratio of 1:1 (oilyaqueous phase), which was administered 3 times in 12 day intervals. Then, the mice were sacrificed and bled by cardiac puncture 20 days after the last immunization. Sera were obtained from the whole blood samples, pooled and used in the DNA vaccine characterization.

DNA vaccine characterization

Human embryonic kidney 293 cells were cultivated in 24-well culture plates (Falcon, Cat. 353047) at a density of 2 x 10^5 cells per well in 1 ml of DMEN medium (Gibco, Cat. 12100-046) supplemented with 10% fetal bovine serum (FBS) (Gibco, Cat. 12657-029) and 1% penicillin/streptomycin (Sigma, Cat. A5955) and incubated for 24 h at 37°C in an atmosphere of 5% CO₂. Next, the cells were transfected with 500 ng of DNA vaccine and CaCl₂ according to the protocol described by Kingston et al. and incubated again [48]. Negative control cells were transfected with the pVAX vector. After 24 h, the transfected cells were harvested, washed with PBS, lysed with RIPA buffer and evaluated by western blot. Cell samples were analyzed by 10% SDS-PAGE, transferred onto a nitrocellulose membrane (Life Technologies, Cat. 1465MB). The membrane was blocked with PBS supplemented with 0.05% Tween 20 (Yetec, Cat. 1280) (PBS-T) and 2% bovine serum albumin (InLab, Cat. 1870) for 2 h at room temperature and then reacted with anti-gD (1:20000, kindly provided by Luis Carlos de Souza Ferreira) or sera (item Production of antibodies against the recombinant proteins) containing anti-recombinant protein antibody (1:1000) (anti-EGFRvIII, MAGE or GLEA) overnight at 4°C. After the membrane was washed three times with PBS-T, it was incubated with anti-mouse IgGHRP (1:5000) (Invitrogen, Molecular Probes, Cat. F21453) for 1 h at room temperature. The immunoreactive protein bands were visualized using the peroxidase substrate 3,3'-diaminobenzidine (DAB) (Vector, Peroxidase Substrate Kit SK-4100).

Mouse immunization

Groups of 8 to 12 female C57BL/6 mice at 6-8 weeks of age from the same animal breeding center were immunized with the DNA vaccines (pVAXDEGFRvIII, pVAXDMAGE, pVAXDGLEA, pVAXEGFRvIII, pVAXMAGE and pVAXGLEA) 3 times intramuscularly in 12 days intervals. The control group was
immunized with the pVAX vector. Each dose of 100 μg of DNA was diluted in 25% saccharose, divided into two 50 μl aliquots and injected into the tibialis anterior muscle of each hind limb.

**Assessment of humoral and cellular immune responses**

Twenty days after the last immunization, the mice were sacrificed and bled, and the spleens were removed. Sera were obtained from the whole blood samples, and cell suspensions were obtained from the macerated spleen samples. Cells were treated for 2 min with 2 ml ammonium chloride potassium buffer to lyse red blood cells, washed with PBS and suspended in RPMI 1640 medium (Sigma, Cat. R6504) supplemented with 10% FBS, 1% penicillin/streptomycin, 30 μg/ml polymyxin B (Sigma, Cat. P4932), 10 μg/ml gentamicin (Gibco, Cat. 15710-064) and 50 μM 2-mercaptoethanol (Sigma, Cat. M7522). Then, 5 × 10^6 cells per well were cultured in 1 ml of medium in the presence of 20 μg/ml EGFvIII, MAGE or GLEA recombinant proteins for 48 h at 37°C in an atmosphere of 5% CO₂. Positive and negative controls were treated with 40 μg/ml concanavalin A (ConA) (Sigma, Cat. C2631) and medium only, respectively.

Sera from vaccinated mice were tested for the presence of IgG1 and IgG2a antibodies against the recombinant proteins by ELISA. Briefly, 96-microwell plates (Maxisorb, Nunc, Cat. 442404) were coated with 100 μl of coating buffer (15 mM Na₂CO₃ and 35 mM NaHCO₃, pH 9.6) containing 5 μg/ml recombinant protein and incubated overnight at 4°C. After the plates were washed with PBS-T, they were blocked with 200 μl of blocking solution (0.1 M NaH₂PO₄, pH 9.0) containing capture antibody [IFN-γ (R4-6A2) (Cat. 551216) and IL-5 (TRFK5) (Cat. 554397) at 0.5 μg/ml; IL-10 (SXC-1) (Cat. 554423) at 1 μg/ml; IL-5 (TRFK5) (Cat. 554397) at 0.5 μg/ml; IL-10 (SXC-1) (Cat. 554423) at 1 μg/ml] (BD Biosciences) and incubated overnight at 4°C. After the plates were washed, 100 μl of serum samples diluted (1:10) in blocking solution was added to the plates, incubated for 2 h at 37°C and washed. Next, 100 μl of biotinylated anti-IgG1 (A85-1) or anti-IgG2a (R19-15) (BD Bioscience) diluted 1:1000 in blocking buffer (15 mM Na₂CO₃ and 35 mM NaHCO₃, pH 9.6) containing 5 μg/ml recombinant protein and incubated overnight at 4°C. After the plates were washed, 100 μl of PBS-T containing 10% FBS (block solution) for 1 h at 37°C. After the plates were washed again, 100 μl of biotinylated secondary antibody [IFN-γ (R4-6A2) (Cat. 551216) and IL-5 (TRFK5) (Cat. 554397) at 0.5 μg/ml; IL-10 (SXC-1) (Cat. 554423) at 1 μg/ml] (BD Biosciences) and incubated overnight at 4°C. After the plates were washed, 100 μl of streptavidin-biotin-horseradish peroxidase (1:1000 dilution; Vector, Vectastain ABC-kit) for 30 min at room temperature. Detection was performed with 100 μl of TMB substrate (BD Bioscience, Cat. 555214). After 30 min at room temperature, the enzyme reaction was stopped by 50 μl of 16% sulfuric acid, and the absorbance was measured at 490 and 570 nm.

Cytokine production in the supernatants of cell cultures was evaluated by ELISA. Briefly, 96-microwell plates were coated with 100 μl of coating buffer (0.1 M Na₂HPO₄, pH 9.0) containing capture antibody [IFN-γ (R4-6A2) (Cat. 551216) and IL-5 (TRFK5) (Cat. 554397)] at 1 μg/ml; IL-10 (JES5-2A5) (Cat. 551215) at 2 μg/ml (BD Biosciences) and incubated overnight at 4°C. After the plates were washed, 100 μl of PBS-T containing 10% FBS (block solution) for 1 h at 37°C. After the plates were washed again, 100 μl of serum samples was added to the plates, incubated overnight for 1-2 h at room temperature. After the plates were washed again, 100 μl of supernatant samples was added to the plates, incubated overnight at 4°C and washed. Next, 100 μl of biotinylated secondary antibody [IFN-γ (XGM1.2) (Cat. 554410) and IL-5 (TRFK4) (Cat. 554497)] at 0.5 μg/ml; IL-10 (SXC-1) (Cat. 554423)] at 1 μg/ml] (BD Biosciences) and incubated overnight at 4°C. After the plates were washed, 100 μl of streptavidin-biotin-horseradish peroxidase (1:1000 dilution; Vector, Vectastain ABC-kit) for 30 min at room temperature. Detection was performed with 100 μl of TMB substrate (BD Bioscience, Cat. 555214). After 30 min at room temperature, the enzyme reaction was stopped by 50 μl of 16% sulfuric acid, and the absorbance was measured at 490 and 570 nm.

**Statistical analysis**

Comparisons were performed using ANOVA with Tukey’s test a posteriori and GraphPad Prism 4.02 version software (GraphPad Software, San Diego, CA, USA). Statistical significance was attributed to p values <0.05.

**Results**

**Construction of DNA vaccines**

The coding sequences of EGFRvIII, MAGE-3 (MAGE) and GLEA-2 (GLEA) proteins without their stop codons were fused to the gD nucleotide sequence in silico. The fusion to gD resulted in the deletion of the gD central portion with 214 amino acids. Because the EGFRvIII sequence is extremely large, only the initial 900 base pairs (bp) were utilized in this assembly. gDEGFRvIII, gDMage, gDGlea fragments were constructed byEpoch Biolabs and cloned into the pBluescript II SK vector (pBSK). The pBSKgDEGFRvIII, pBSkgDmage and pBSkgDGlea plasmids were cleaved to remove the gD-antigen fragments and were used as templates for EGFRvIII, MAGE and GLEA amplification. All fragments were cloned into the pVAX1 (pVAX) vector. Because pVAX is a transcription plasmid, all the fragments already had the Kozak sequence (ACCATG) [49]. Restriction analyses (Figure 1) and sequencing (data not shown) confirmed that the DNA vaccines were correctly constructed. As shown in Figure 1, pVAXgDEGFRvIII, pVAXgDmage, pVAXgDGlea, pVAXEGFRvIII, pVAXMAGE and pVAXGLEA constructs presented sizes that matched the vector associated with the insert: 4,357 bp, 4,399 bp, 5,086 bp, 3,820 bp, 3,859 bp and 4,546 bp, respectively.

**Figure 1:** Digestion of DNA vaccines. Large-scale purification of the DNA vaccines pVAXEGFRvIII (1), pVAXMAGE (2), pVAXGLEA (3), pVAXEgDGFRvIII (4), pVAXDmage (5) and pVAXGLEA (6) was conducted using an Endofree Plasmid Giga Kit. DNA vaccines were quantified by spectrophotometry and digested with the NheI restriction enzyme, which was used in the cloning. L: 1 Kb Plus DNA Ladder (Invitrogen).
Expression and purification of the recombinant proteins

EGFRvIII, MAGE and GLEA were also amplified from pBSKgDEGFRvIII, pBSKgDMAGE and pBSKgDGLEA plasmids for cloning into the E. coli expression vector pET. The recombinant proteins were expressed and analyzed by 10% SDS-PAGE (Figure 2a). The recombinant EGFRvIII, MAGE and GLEA proteins were separated by SDS-PAGE and were 36 kDa, 43 kDa and 72 kDa, respectively. The MAGE and GLEA proteins showed larger molecular masses than expected (37 and 64 kDa, respectively). The recombinant EGFRvIII, MAGE and GLEA proteins were purified by affinity chromatography and analyzed by 10% SDS-PAGE (Figure 2b). The recombinant proteins were used for mouse immunization, antibody production and immunogenicity assays.

DNA vaccines characterization

To verify whether the DNA vaccines could express the expected proteins, human embryonic kidney 293 (HEK293) cells were transiently transfected with the constructs. One day later, cell extracts were obtained, and the proteins were detected by western blot with the appropriate antibodies (Figure 3). The extract from cells transfected with the empty pVAX vector failed to bind to gD, EGFRvIII, MAGE and GLEA-specific antibodies. Anti-gD antibody stained extracts from cells transfected with vaccines expressing the gDEGFRvIII, gDMAGE and gDGLEA fusion proteins (Figure 3a). Cells transfected with pVAXgDEGFRvIII and pVAXEGFRvIII were stained by anti-EGFRvIII antibody specific to the corresponding tumor antigen (Figure 3b). Extracts from cells transfected with pVAXgDMAGE and pVAXMAGE were stained by anti-MAGE antibody (Figure 3c), and extracts from cells transfected with pVAXDGLEA and pVAXGLEA were stained by anti-GLEA antibody (Figure 3d). Collectively, these results demonstrate that the proteins encoded by the DNA vaccines were correctly expressed by the cellular machinery.

Assessment of humoral and cellular immune responses

C57BL/6 mice received three doses of the DNA vaccines (pVAXgDEGFRvIII, pVAXgDMAGE, pVAXDGLEA, pVAXEGFRvIII, pVAXMAGE or pVAXGLEA) at a 12 days interval regimen. Twenty days after the last immunization, blood and spleen samples were collected. Sera were obtained from the whole blood samples and were tested for the presence of IgG1 and IgG2a antibodies against the recombinant proteins EGFRvIII, MAGE and GLEA by
The production of specific IgG2a antibodies was inconsistent for the pVAXgDEGFRvIII (Figure 4a) and pVAXgDMAGE (Figure 4b) groups. Only the pVAXgDGLEA group produced a high level of specific IgG2a antibodies (Figure 4c). Specific IgG1 antibodies were not detected in any group (data not shown).

Figure 4: Humoral immune response induced by DNA vaccines. Sera from vaccinated mice with three doses of 100 µg of DNA vaccines were collected 20 days after the last immunization and tested for the presence of IgG2a antibodies against recombinant proteins by ELISA. a) Anti-EGFRvIII detection. The data presented are from one experiment (6 mice per group). b) Anti-MAGE detection. Representative data presented are from one experiment (6 mice per group). c) Anti-GLEA detection. The data presented are from two experiments (12 mice per group).

* p<0.05, one-way ANOVA and Tukey's posttest.

To characterize antigen-specific cell-mediated responses, cells were obtained from macerated spleen samples and were stimulated ex vivo with the recombinant proteins or with concanavalin A (ConA) as a positive control. The negative control was performed with unstimulated cells. Cytokine production by the stimulated cells was quantified by ELISA using culture supernatants. IFN-γ (Figure 5) and IL-10 (Figure 6) production was significantly higher in the pVAXgDEGFRvIII (Figure 5a and 6a), pVAXgDMAGE (Figure 5b and 6b) and pVAXgDGLEA (Figure 5c and 6c) groups compared with the other groups. IL-5 production was not detected in any group (data not shown).

Figure 5: Assessment of cellular immune responses induced by DNA vaccines by measuring IFN-γ production. Mice were vaccinated with three doses of 100 µg of DNA vaccines. Spleen cells were collected 20 days after the last immunization and stimulated ex vivo for 48 h. IFN-γ production levels in the culture supernatants were quantified by ELISA. a) Culture stimulated with EGFRvIII protein. The data presented are from two experiments (12 mice per group). b) Culture stimulated with MAGE protein. The data presented are from two experiments (12 mice per group). c) Culture stimulated with GLEA protein. The data presented are from one experiment (8 mice per group). The differences in the cellular responses were determined by one-way ANOVA and Tukey's posttest. ** p<0.01, *** p<0.001. The negative control was performed without any stimulus, and if detected, cytokine production was discounted for data tabulation of other stimuli. The positive control was performed using ConA as the polyclonal stimulus.

Discussion

Despite multimodal treatment with maximal surgical resection followed by temozolomide and radiation, GBM remains a recurrent and fatal cancer [4,5]. Thus, immunotherapy combined with conventional therapy constitutes an alternative intervention, providing the advantage of inducing a long-term, specific immune response that could prevent GBM recurrence. An effective vaccine for cancer should elicit not only a humoral response but also a cellular response. However, vaccines containing antigens alone do not induce efficient cellular responses. Thus, antigens need to be combined with an efficient adjuvant or delivery system that can be used in humans. Therefore, we developed DNA vaccines encoding GBM antigens fused
to gD from HSV-1. Our results demonstrated that our DNA vaccines were constructed correctly and that the proteins were expressed in transfected HEK293 cells.

**Figure 6:** Assessment of cellular immune responses induced by DNA vaccines by measuring IL-10 production. Mice were vaccinated with three doses of 100 µg of DNA vaccines. Spleen cells were collected 20 days after the last immunization and stimulated ex vivo for 48 h. IL-10 production levels in the culture supernatants were quantified by ELISA. a) Culture stimulated with EGRFvIII protein. The data presented are from two experiments (12 mice per group). b) Culture stimulated with MAGE protein. The data presented are from two experiments (12 mice per group). c) Culture stimulated with GLEA protein. The data presented are from two experiments (12 mice per group). The differences in the cellular responses were determined by one-way ANOVA and Tukey’s posttest. *p<0.05, **p<0.01. The negative control was performed without any stimulus, and if detected, cytokine production was discounted for data tabulation of other stimuli. The positive control was performed using ConA as the polyclonal stimulus.

Previous observations have indicated that gD can enhance the immunogenicity of antigens present in DNA vaccines when these antigens are fused to gD [32-34,50]. Antigens and secondary signals are required for the activation of adaptive immune responses. Secondary signals are delivered by co-stimulatory molecules, and gD can perform this function when interacting with cell surface receptors, thus activating type I IFN and IL-12 production by DCs [43,44]. Lasaro et al. also suggested that a secondary signal could originate from the inhibition of the HVEM-BTLA/CD160 inhibitory pathway by gD, which interacts with HVEM in the same site to which other molecules bind [32,51]. BTLA is an inhibitory receptor present in T cells that inhibits IL-2 secretion when activated. BTLA-deficient T cells display increased proliferation in specific T lymphocytes stimulated by antigen-DGs [52]. CD160 presents BTLA-like features, thus inhibiting T CD4+ activation [42]. Therefore, blocking the HVEM-BTLA/CD160 interaction can enhance the T cell response.

In this study, DNA vaccines codifying only tumor antigens showed low immunogenicity. Mice immunized with pVAXgDEGFRvIII, pVAXMAGE and pVAXgGLEA did not present relevant secondary humoral or cellular immune responses. Many studies have shown that DNA vaccines can stimulate both humoral and cellular immune responses [53-55]. However, most of the proteins produced by DNA immunization are not able of priming the immune response by themselves, thus, is usually required to do the enhancement of the potency of genetic vaccination [56-58]. Satisfying this condition, the pVAXgDEGFRvIII, pVAXgMAGE and pVAXgDGLEA vaccines encoding the tumor antigens genetically fused to the gD protein were able to induce cellular immune responses, despite low induced humoral responses. The immune effects induced by these vaccines are suggestive that the structural integrity of the gD-HVEM binding domains is preserved. The induction of adaptive immune response (humoral and cellular) was evaluated after stimulation with the recombinant proteins EGRFvIII, MAGE and GLEA, which were expressed in E. coli and purified by affinity chromatography. The MAGE and GLEA recombinant proteins showed larger molecular masses than expected, as demonstrated previously by Kocher et al. [59]. The authors showed that the MAGE protein, whether recombinant or native in melanoma cells, presented larger molecular masses than expected [59]. Because the MAGE protein does not show post-translational modifications, the authors suggested that the size could be changed due to its amino acid composition [59]. The prevalence of amino acids rich in acidic residues can alter the electrophoresis pattern of the protein. The amino acid composition may also explain the increased size of the GLEA protein, which also does not undergo post-translational modifications when expressed in *E. coli*.

Analysis of the cellular immune responses showed that the IFN-γ production in the supernatants of spleen cells from mice vaccinated with pVAXgDEGFRvIII, pVAXgMAGE and pVAXgDGLEA were significantly higher than those levels verified in the animals of other immunization groups. This result indicates that tumor antigens associated with gD drive the T cell response toward a Th1-type immune response profile. Immunomodulatory effects of gD were also demonstrated in other studies with DNA vaccines codifying E5, E6 and E7 tumor antigens fused to gD [32-34]. Mice immunized with these DNA vaccines presented marked increases in specific CD8+ T cells [32-34]. In another study by the same group, trivalent DNA vaccines that simultaneously encoded HIV (p24), HSV (gD) and HPV (E7) antigens stimulated functional and protective p24-, gD- and E7-specific CD8+ T cell responses in vaccinated mice, thus promoting efficient anti-virus (HIV/HSV-1) and anti-tumor (HPV) effects [60].

For all vaccination strategies, cell-mediated immune response and IFN-γ production have been considered essential for the generation of a long-lasting specific immune response against tumor cells. IFN-γ participates in immunosurveillance that protects against
spontaneously arising, transplantable, chemically induced tumors [61]. The effects of IFN-γ on host cells include activation of the Th1-cell lineage, enabling macrophage activation and CTL maturation [62,63], and inhibition of regulatory T cell generation and/or activation [64]. In contrast, IFN-γ also presents a direct anti-tumor mechanism that promotes apoptosis through its effects on the expression of caspases, FAS and TRAIL [65-67], thus inhibiting cellular proliferation and angiogenesis [68,69].

Spleen cells from mice vaccinated with pVAXgDEGFrVL11, pVAXgDMAGE and pVAXgDGLEA stimulated with recombinant proteins produced significantly higher levels of IL-10 than mice immunized with the other constructs. The functions of some cytokines produced in the tumor environment may have different effects when compared with situations in which the same cytokine is produced outside the tumor environment [70]. This situation has been confirmed with IL-10; its role in tumor immunology is contradictory. IL-10 was initially classified as a Th2-derived cytokine [71]; however, it is produced by a range of CD4+ T cell subsets and by macrophages, DCs, B cells, eosinophils and mast cells [72]. Generally, IL-10 is considered an immunosuppressive molecule [72,73], and in glioma cancer, it is produced by cancer cells to escape the immune response [74].

However, some studies have shown the importance of IL-10 in controlling tumor cells. T cells primed by autologous DCs loaded with GBM tumor cell lysates acquired a Th1 profile and the capacity to produce IL-10 [75]. This production correlated with antitumor T cell activity [75]. In a vaccine strategy to protect against tumor challenge, Shin and collaborators showed that IL-10 administration promotes the proliferation and maintenance of T CD8+--activated antigen-specific cells [76]. These cells produced IFN-γ and displayed an increased cytotoxic capacity, which supports tumor reduction [76]. Other researchers demonstrated that IL-10-deficient mice developed a greater number of skin tumors and less granzymes than wild-type animals. Furthermore, the administration of IL-10 promoted tumor rejection in several tumor types even when the tumor was well established. These authors clearly showed that the anti-tumor activity of IL-10 is dependent on IFN-γ induction in CD8+ T cells. The anti-tumor CD8+ T cells present higher IL-10 receptor expression, which explains its activation via this cytokine [77]. These results indicate that IL-10 may be a helpful cytokine in immunization strategies because it improves the effectiveness of anti-tumor CD8+ T cells.

Other studies have revealed that IL-10/- mice develop many colon polyps with accelerated growth and lung metastasis when compared to control animals [78]. These animals, despite their IL-10 deficiency, have a greater number of regulatory T cells (Tregs) and myeloid-derived suppressor cells (MDSCs), which increase immunosuppression and inhibit the anti-tumor immune response [78]. Instead, the myeloid cells of these animals expressed higher levels of IL-1a and IL-1β, which are pro-angiogenic and pro-tumorigenic [78]. The inhibitory signaling of these cytokines decreases angiogenesis and Tregs and increases CD8+IFN-γ+ T cells resulting in tumor reduction [78]. Thus, tumor development in IL-10/- animals results from a linked mechanism of immunosuppression and inflammation that impairs anti-tumor responses.

In contrast, a new theory suggests that Th1 cells are regulated via the co-induction of IL-10 in addition to IFN-γ in the same cells [79,80]. These cells that produce IL-10 develop from IFN-γ+ Th1 cells when appropriate signals are received and represent the ‘endpoint’ of an effective immune response (one component of the life cycle of a Th1 cell) [81]. Jankovic and collaborators suggested an explanation to this theory, namely, that IFN-γ is important cytokine for combatting pathogens and that IL-10 protects against tissue injury that can be triggered by immune responses [82]. Thus, Th1 IFN-γ+IL-10+ cells promote and control immune responses by effector and regulatory cytokine induction, following recognition of the same antigen [81]. The two roles of IL-10, i.e., favoring or controlling the immune response, are important for destroying tumors and for controlling immune responses that should not cause damage to normal tissues.

In summary, this study indicates the potential value of DNA vaccination using the expression of GBM antigens genetically fused with the HSV-1 gD. DNA vaccination with chimeric tumor antigens stimulated cell-mediated immunity, revealed by the specific IFN-γ and IL-10 responses, which is considered essential for combating tumors. Collectively, these results provide evidences for the potential application of an active immunotherapy method to be used with conventional therapy for GBM treatment.

Acknowledgments
This work was supported by the Fundação de Amparo à Pesquisa do Estado de São Paulo (FAPESP, Grant number 2009/51280-8), Coordenação de Aperfeicoamento de Pessoal de Nível Superior (CAPES), Conselho nacional de Pesquisa (CNPq) and Financiadora de Estudos e projetos (FINEP).

Conflicts of Interest
No other funding or conflicts of interest to declare.

References
1. Kleihues P, Louis DN, Scheithauer BW, Ortega-Pelayo MR, Reifenberger G, et al. (2002) The WHO classification of tumors of the nervous system. J Neuropathol Exp Neurol 61: 215-225.
2. Jackson C, Ruzek J, Phallen J, Belcaid Z, Lim M (2011) Challenges in immunotherapy presented by the glioblastoma multiforme microenvironment. Clin Dev Immunol 2011: 732413.
3. Wilson TA, Karajannis MA, Harter DH (2014) Glioblastoma multiforme: State of the art and future therapeutics. Surg Neurol Int 5: 64.
4. Stupp R, ME Hegi, WP Mason, MJ van den Bent, MJ Taphoorn, et al. (2009) Effects of radiotherapy with concomitant and adjuvant temozolomide versus radiotherapy alone on survival in glioblastoma in a randomised phase III study: 5-year analysis of the EORTC-NCIC trial. Lancet Oncol 10: 459-466.
5. Stupp R, Mason WP, van den Bent MJ, Weller M, Fisher B, et al. (2005) Radiotherapy plus concomitant and adjuvant temozolomide for glioblastoma. N Engl J Med 352: 987-996.
6. Charles NA, Holland EC, Gilbertson R, Glass K, Kettenmann H (2011) The brain tumor microenvironment. Glia 59: 1169-1180.
7. Fornari FB, T Fenton, RM Bachoo, A Mukasa, JM Stommel, et al. (2007) Malignant astrocytic glioma: genetics, biology, and paths to treatment. Genes Dev 21: 2683-2710.
8. Kesari S (2011) Understanding glioblastoma tumor biology: the potential to improve current diagnosis and treatments. Semin Oncol 40: S4-10.9.
9. Wei J, Barr J, Kong LY, Wang Y, Wu A, et al. (2010) Glioma-associated cancer-initiating cells induce immunosuppression. Clin Cancer Res 16: 461-473.
10. Tambuyzer BR1, Ponsarta P, Nouwen EJ (2009) Microglia: gatekeepers of central nervous system immunity. J Leukoc Biol 85: 352-370.
11. Hussain SF, AB Heinberger (2005) Immunotherapy for human glioma: innovative approaches and recent results. Expert Rev Anticancer Ther 5: 777-790.
Citation: Rios WM, De Molfetta JB, Brandão IT, Masson AP, Peripato R, et al. (2015) Fusion of Glioblastoma Tumor Antigens to Herpes Simplex Virus-1 Glycoprotein D Enhances Secondary Adaptive Immune Responses in a DNA Vaccine Strategy. J Vaccines Vaccin 6: 302. doi: 10.4172/2157-7560.1000302

12. Serot JM, Bo Foliuguet, MC Béné, GC Faure (1997) Ultrastructural and immunohistochemical evidence for dendritic-like cells within human choroid plexus epithelium. Neuroreport 8: 1995-1998.

13. Goldmann J, K Kwizdinski, C Brandt, J Mahlo, D Richter, et al. (2006) T cells traffic from brain to cervical lymph nodes via the cribriform plate and the nasal mucosa. J Leukoc Biol 80: 797-801.

14. Weller RO, Engelhardt B, Phillips MJ (1996) Lymphocyte targeting of the central nervous system: a review of afferent and effector CNS-immune pathways. Brain Pathol 6: 275-288.

15. Calzascia T, F Masson, W Di Berardino-Bessone, E Contassot, R Wilmotte, et al. (2005) Homing phenotypes of tumor-specific CD8 T cells are predetermined at the tumor site by crosspresenting APCs. Immunity 22: 175-184.

16. Humphrey PA, AJ Wong, B Vogelstein, MR Zalutsky, GN Fuller, et al. (1990) Anti-synthetic peptide antibody reacting at the fusion junction of deletionmutant epidermal growth factor receptors in human glioblastoma. Proc Natl Acad Sci U S A 87: 4207-4211.

17. Wong AJ, JM Ruppert, SH Bigner, CH Grzeschik, PA Humphrey, et al. (1992) Structural alterations of the epidermal growth factor receptor gene in human gliomas. Proc Natl Acad Sci U S A 89: 2965-2969.

18. Batra SK, S Castelino-Prabhu, C Wikstrand, X Zhu, PA Humphrey, et al. (1995) Epidermal growth factor ligandindependent, unregulated, cell-transforming potential of a naturally occurring human mutant EGFRvIII gene. Cell Growth Differ 6: 1251-1259.

19. Nishikawa R, XD Ji, RC Harmon, CS Lazar, GN Gill, et al. (1994) A mutant epidermal growth factor receptor common in human glioma confers enhanced tumorigenicity. Proc Natl Acad Sci U S A 91: 7727-7731.

20. Heimberger A, S Hussain, K Alapde, R Sawaya, G Archer, et al. (2006) Tumor-specific peptide vaccination in newly-diagnosed patients with GBM. J Clin Oncol Abstrat 2529.

21. Sampson JH, G E Archer, DA Mitchell, AB Heimberger, DB Bigner (2008) Tumor-specific immunotherapy targeting the EGFRvIII mutation in patients with malignant glioma. Semin Immunol 20: 267-275.

22. Sampson JH, Archer GE, Mitchell DA, Heimberger AB, Herndon JE, et al. (2009) An epidermal growth factor receptor variant III-targeted vaccine is safe and immunogenic in patients with glioblastoma multiforme. Mol Cancer Ther 8: 2773-2779.

23. Schmitting RJ, Archer GE, Mitchell DA, Heimberger A, Pegram C, et al. (2008) Detection of humoral response in patients with glioblastoma receiving EGFRvIII-KLH vaccines. J Immunol Methods 339: 74-81.

24. Chi DD, Merchant RE, Rand R, Conrad AJ, Garrison D, et al. (1997) Molecular detection of tumor-associated antigens shared by human cutaneous melanomas and gliomas. Am J Pathol 150: 2143-2152.

25. Chomier P, De Backer O, Bertrand M, De Plaen E, Boon T, et al. (2001). Targeting of antigen to the herpesvirus entry mediator augments primary adaptive immune responses. Nat Med 14: 205-212.

26. Diniz MO, Lasaro MO, RE, Ferreira LC (2010) Immun response and therapeutic antitumor effects of an experimental DNA vaccine encoding human papillomavirus type 16 oncoproteins genetically fused to herpesvirus glycoprotein D. Clin Vaccine Immunol 17: 1576-1583.

27. Heimberger A, S Hussain, K Aldape, R Sawaya, G Archer, et al. (2006) Expression of herpes simplex virus type 1 glycoprotein D inhibit virus penetration. J Virol 61: 3356-3364.

28. Spear PG (2004) Herpes simplex virus: receptors and ligands for cell entry. Cell Microbiol 6: 401-410.

29. Montgomery RI, Warner MS, BM J, Speir PG (1996) Herpes simplex virus-1 entry into cells mediated by a novel member of the TNF/NF receptor family. Cell 87: 427-436.

30. Mauri DN, Ebner R, Montgomery RI, Kochel KD, Cheung TC, et al. (1998) LIGHT, a new member of the TNF superfamily, and lymphotoxin alpha are ligands for herpesvirus entry mediator. Immunity 6: 21-30.

31. Gonzalez LC, Loyet KM, Caulemine-Fenaux J, Chauhan V, Wrankel B, et al. (2003) A coreceptor interaction between the CD28 and TNF receptor family members B and T lymphocyte attenuator and herpesvirus entry mediator. Proc Natl Acad Sci U S A 102: 1116-1121.

32. Cai G, Anumantha B, Brown JA, Greenfeld EA, Zhu B, et al. (2008) CD160 inhibits activation of human CD4+ T cells through interaction with herpesvirus entry mediator. Nat Immunol 9: 176-185.

33. Medici MA, Sciorrenti MT, Perri D, Amici C, Avitabile E, et al. (2003) Protection by herpes simplex virus glycoprotein D against Fas-mediated apoptotic role of nuclear factor kappaB. J Biol Chem 278: 36059-36067.

34. Pollara G, Jones M, Handley ME, Rajpopat M, Kwan A, et al. (2004) Herpes simplex virus type 1-induced activation of myeloid dendritic cells: the roles of virus cell interaction and paracrine type I IFN secretion. J Immunol 173: 4108-4119.

35. Cohen GH, Wilcox WC, Soda LA, Long D, Levin JZ, et al. (1988) Expression of herpes simplex virus type 1 glycoprotein D deletion mutants in mammalian cells. J Virol 62: 1932-1940.

36. Laemmli UK (1970) Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature 227: 680-685.

37. Bradford MM (1976) A rapid and sensitive method for the quantitation of microgram quantities of protein utilizing the principle of protein-dye binding. Anal Biochem 72: 248-254.

38. Kingston RE, Chen CA, Okayama H (2001) Calcium phosphate transfection. Curr Protoc Immunol Chapter 10: Unit 10.13.

39. Kozak M (1990) Downstream secondary structure facilitates recognition of initiator codons by eukaryotic ribosomes. Proc Natl Acad Sci USA 87: 8301-8305.

40. Lásaro MO, Alves AM, Botoso V, Durigon EL, Ferreira LC (2003) Antibody-inducing properties of a prototype bivalent herpes simplex virus/enterotoxigenic Escherichia coli DNA vaccine. FEMS Immunol Med Microbiol 35: 25-31.
Citation: Rios WM, De Molfetta JB, Brandão IT, Masson AP, Piripato R, et al. (2015) Fusion of Glioblastoma Tumor Antigens to Herpes Simplex Virus-1 Glycoprotein D Enhances Secondary Adaptive Immune Responses in a DNA Vaccine Strategy. J Vaccines Vaccin 6: 302. doi: 2157-7560.1000302

51. Lasaro MO, M Sazanovich, W Giles-Davis, P Mrass, RM Bunte, et al. (2011) Active immunotherapy combined with blockade of a co-inhibitory pathway achieves regression of large tumor masses in cancer-prone mice. Mol Ther 19: 1727-1736.

52. Watanabe N, M Gavrieli, JR Sedy, J Yang, F Fallarino, (2003) BTLA is a lymphocyte inhibitory receptor with similarities to CTLA-4 and PD-1. Nat Immunol 4: 670-679.

53. Conry RM, DT Curiel, TV Strong, SE Moore, KO Allen, et al. (2002) Safety and immunogenicity of a DNA vaccine encoding carcinoembryonic antigen and hepatitis B surface antigen in colorectal cancer patients. Clin Cancer Res 8: 2782-2787.

54. Tagawa ST, P Lee, J Snively, W Boswell, S Ounpraseuth, et al. (2003) Phase I study of intranodal delivery of a plasmid DNA vaccine for patients with Stage IV melanoma. Cancer 98: 144-154.

55. Liu MA (2003) DNA vaccines: a review. J Intern Med 253: 402-410.

56. Pavlenko M, AK Roos, A Lundqvist, A Palmborg, AM Miller, et al. (2002) Naked DNA and adenalovirual immunizations for immunotherapy of prostate cancer: a phase II clinical trial. Eur Urol 38: 208-217.

57. Low L, A Mander, K McCann, D Dearnaley, T Tjelle, I et al. (2002) DNA vaccination with electroporation induces increased antibody responses in patients with prostate cancer. Hum Gene Ther 20: 1269-1278.

58. Kocher T, E Schultz-Thater, F Gadot, C Schaefer, G Casorati, et al. (1995) Identification and intracellular location of MAGE-3 gene product. Cancer Res 55: 2236-2239.

59. Mincheff M, S Tchakarov, S Zoubak, D Loukinov, C Botev, et al. (2000) Interleukin-10 elicits the maintenance of antitumor CD8(+) T-cell effector function in situ. Blood 98: 2143-2151.

60. Meng RD, WS El-Deiry (2001) p53-independent upregulation of KILLER/DR5 TRAIL receptor expression by glucocorticoids and interferon gamma. Exp Cell Res 262: 154-169.

61. Chin YE, Kitagawa M, Su WC, You ZH, Iwamoto Y, et al. (1996) Cell growth arrest and induction of cyclin-dependent kinase inhibitor p21 WAF1/CIP1 mediated by STAT1. Science 272: 719-722.

62. Bromberg JF, Horvhath CM, Wen Z, Schreiber RD, Darnell JE (1996) Transcriptionally active Stat1 is required for the antiproliferative effects of both interferon alpha and interferon gamma. Proc Natl Acad Sci U S A 93: 7673-7678.

63. Meng RD, Darcy PK, Smyth MJ (2011) Stable IL-10: a new therapeutic that promotes tumor immunity. Cancer Cell 20: 691-693.

64. Th1-directed effector cells after stimulation with tumor lysate-loaded dendritic cells. J Neurooncol 84: 131-140.

65. Fuji S, Shimizu K, Shimizu T, Lotze MT (2001) Interleukin-10 promotes the maintenance of antitumor CD8(+) T-cell effector function in situ. Blood 98: 2143-2151.

66. Mumm JB, Emmerich J, Zhang X, Chan I, Wu L, et al. (2011) IL-10 elicits IFN-γ-dependent tumor immune surveillance. Cancer Cell 20: 781-796.

67. O'Garra A, Vieira P (2007) T(H)1 cells control themselves by producing interleukin-10. Nat Rev Immunol 7: 425-428.

68. Fujii S, Shimizu K, Shimizu T, Lotze MT (2001) Interleukin-10 promotes the maintenance of antitumor CD8(+) T-cell effector function in situ. Blood 98: 2143-2151.

69. De Vleeschouwer S, Spencer Lopes I, Ceuppens JL, Van Goew Wol (2007) Persistent IL-10 production is required for glioma growth suppressive activity by Th1-directed effector cells after stimulation with tumor lysate-loaded dendritic cells. J Neurooncol 84: 131-140.

70. Cope A, Le Friec G, Cardone J, Kemper C (2011) Interleukin-10 ablation promotes tumor development, growth, and metastasis. Cancer Res 72: 420-429.

71. Trinchieri G (2007) Interleukin-10 production by effector T cells: Th1 cells show self control. J Exp Med 204: 239-243.

72. Cope A, Le Friec G, Cardone J, Kemper C (2011) The Th1 life cycle: molecular control of IFN-γ to IL-10 switching. Trends Immunol 32: 278-286.

73. Jankovic D, Kullberg MC, Feng CG, Goldsmdz RD, Collazo CM, et al. (2007) Conventional T-bet(+)Foxp3(-) Th1 cells are the major source of host-protective regulatory IL-10 during intracellular protozoan infection. J Exp Med 204: 273-283.