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1. Introduction

1.1 Treatment limitations of patients with breast cancer metastatic to the brain
Although technical advances have resulted in marked improvement in the ability to
diagnose and treat primary breast cancer, brain metastases constitute a common and
increasing occurrence associated with considerable morbidity and mortality [Lin et al., 2008;
Smedby et al., 2009]. The present standard treatment modalities including surgical resection,
cranial irradiation and systemic chemotherapy each have serious adverse side effects and
limits in efficacy. The few long-term survivors are inevitably left with cognitive deficits and
other disabilities [Heimans & Taphoorn, 2002]. The existence of blood-brain and blood-
tumor barriers impedes drug delivery to the tumor. Finally the low therapeutic index
between tumor sensitivity and toxicity to normal brain severely limits the ability to
systemically deliver therapeutic doses of drugs or administer focal radiation therapy to the
tumor. New treatment strategies are urgently needed.

1.2 Transfer of genomic DNA from one cell type to another alters both the genotype
and the phenotype of cells that take up the exogenous DNA
Classic studies indicate that transfection of genomic DNA from one cell type to another
results in integration of the transferred DNA and stable alteration of the genotype of the
recipient cells. The transferred genes are replicated as the cells divide and are expressed. In
one study the genome of adenine phosphoribosyltransferase-deficient mouse cells was
modified to express the missing enzyme by transfer of DNA from mouse cells whose
genome included the gene for the missing enzyme [Wigler et al., 1979]. Analogous findings
were observed for membrane-associated determinants. In another study genomic DNA from
human cells was transferred into polio virus-receptor-negative mouse cells and the
transfected cells expressed the missing receptor [Mendersohn et al., 1986]. Others
[Barraclough et al., 1998; Chen et al., 1997] used this approach to identify genes involved in
metastasis. Another approach involved the generation of stable transfecants of mouse
fibroblasts [Hsu et al., 1984; Kavathas & Herzenberg, 1983]. The transfected cells expressed
human membrane T cell antigens, HLA determinants, and B2-microglobulin. The expression
of the transferred human genes by the transfected cells was stable and long-term (more than
six months). The proportion of the transfected mouse cells that expressed the human gene of
interest was surprisingly large--in the range of 1/500. The importance of these findings for
development of DNA-based tumor vaccines is that the transfer of genomic DNA into cells
resulted in the expression of genes specifying missing enzymes, genes controlling cell
proliferation and metastasis, and genes specifying membrane associated determinants. An analogous approach can be used to prepare a vaccine for use in patients with malignant gliomas. Genes specifying tumor associated antigens (TAAs) that fail to provoke anti-tumor immunity can become highly immunogenic antigenic determinants if they are expressed by highly immunogenic cells.

1.3 Multiple mutant/dysregulated genes in cancer cells specify TAAs

A major rationale for the use of DNA-transfer to prepare vaccines for use in cancer therapy is that the vaccine expresses an array of multiple altered genes which define the malignant phenotype. Genetic instability in cancer cells is responsible for the formation of TAAs. TAAs such as β-catenin [Robbins et al., 1996], gp100, Melan A/Mart-1 and tyrosinase in melanoma [de Vries et al., 1997] are differentiation antigens whose expression is dysregulated in cancer cells. Mutant genes also specify TAAs [van der Bruggen et al., 1991]. For example Boon found that a point mutation in a gene in P815 murine mastocytoma cells specified a tumor-rejection antigen [Boon et al., 1994]. Thus, the malignant cell-population is characterized by the presence of numerous TAAs, some of which are unique and others are differentially expressed by cancer cells but all are strong potential targets of immune-mediated attack.

1.4 DNA from the patient’s neoplasm is the ideal source of tumor antigens for immunotherapy

Since the total number of different TAAs within the population of malignant cells is large and diverse, successful therapy will depend upon the use of a vaccine that is capable of inducing immunity to the broad array of tumor antigens that characterizes the patient’s cancer. Therapy based on the induction of immunity to a single antigen, or peptide, is less likely to be successful. Multi-epitope vaccines are expected to be more efficacious than single-epitope vaccines [Stevenson et al., 2004]. This is especially the case for malignant astrocytomas, where clinically relevant TAAs, i.e., immunity to TAAs that leads to tumor rejection, have not been identified.

1.5 Characteristics of the modified cell line used as the recipient of tumor DNA

Among other advantages of this approach, the cells chosen as DNA-recipients can be selected for their ability to enhance the immune response. The expression of both syngeneic and allogeneic MHC-determinants by the DNA recipient cells is important in order to obtain an optimum anti-tumor response [deZoeten et al., 2002]. The syngeneic determinants provide a restriction element for direct presentation of TAAs to CTLs of the host. Allogeneic antigens served as potent immune adjuvants. Numerous investigators found that the immunogenic properties of cancer cells could be enhanced if the cells were modified to express allogeneic MHC-determinants [Fearon et al., 1988; Gattoni-Celli et al., 1988; Hammerling et al., 1986; Hui et al., 1989; Nabel et al., 1996; Ostrand-Rosenberg et al., 1990]. The modified cells, which ordinarily proliferate in syngeneic immunocompetent recipients, were recognized as “foreign” and were rejected. In the mouse, immunization with tumor cells altered by the introduction and expression of allogeneic class I genes led to immune-mediated rejection of the malignant cells and the induction of protective anti-tumor immunity. However, the introduction of genes specifying allogeneic determinants into cells from a primary neoplasm is technically challenging and not always successful. In contrast, transfer of DNA from the tumor into highly immunogenic syngeneic/allogeneic cells is consistently and reliably achieved.
1.6 Important advantages of preparing a vaccine by transfer of DNA from the patient’s neoplasm into nonmalignant fibroblasts

A vaccine prepared by transfer of DNA from the patient’s neoplasm into highly immunogenic, nonmalignant human fibroblasts has a number of important advantages. A major advantage is that the cells used as recipients of the DNA can be selected for special properties, which will enhance the anti-tumor immune response. Since the recipient cells are capable of prolonged proliferation in vitro, and the transferred DNA is replicated as the cells divide, only a small quantity of DNA from the neoplasm is required to generate the vaccine. In addition, the number of transfected fibroblasts can be expanded as needed to obtain sufficient quantities for repeated immunizations of the cancer patient. The fibroblasts used as DNA-recipients will also express allogeneic class I determinants which is a desirable feature since this leads to an augmented immune response. In addition a cell line derived from the patient’s primary neoplasm does not have to be established, which is the case if genes specifying cytokines, allogeneic MHC-determinants, co-stimulatory molecules or other immune-augmenting properties are to be introduced into the autologous tumor cells. The establishment of tumor cell lines, especially cell lines derived from astrocytomas, is technically difficult, often not feasible and may not be representative of the tumor cell population as a whole. Furthermore hybrid cell vaccines prepared by fusion of tumor cells with antigen presenting cells pose similar concerns [Gong et al., 1997; Liang & Cohen, 1976, 1977]. Immunization with tumor cells modified to secrete immune-augmenting cytokines such as IL-2 and GM-CSF has been investigated and shown to result in the development of generalized MHC-restricted anti-tumor immune responses in animal models. However tumor cells are also a source of immunosuppressive factors, which inhibit the anti-tumor activity of the effector cells [Strand & Galle, 1998; Whiteside & Rabinwich, 1998]. The DNA-based vaccines are successful because a full complement of genes is transferred to the recipient cells which results in a robust signal for the development of anti-tumor immune responses.

1.7 Advantages of DNA-based vaccines relative to other types of vaccines

A number of different vaccination strategies are currently being evaluated [Ashley et al., 1997; Condon et al., 1996; Gilboa et al., 1998; Nair et al., 1998; Nestle et al., 1998; Tighe et al., 1998]. The approaches to vaccination with TAAs include those based on: a) defined antigens or antigenic peptides, b) tumor cell lysates or lysate fractions, and c) whole irradiated tumor cells or apoptotic tumor cell bodies. Clinical trials involving vaccines prepared using TAAs or TAA-derived epitopes presented by APCs or fed to dendritic cells (DCs) have shown some promising results. However, defined antigens have to be identified and purified, a tremendous effort requiring an “antigen discovery” approach. The quantity of purified antigen must be increased, to enable multiple immunizations of the cancer patient. While new TAAs are being discovered, the question of which TAA to be used in the vaccine is uncertain and extensively debated. The heterogeneity of antigen expression in the tumor cell population is likely to be a concern. Some tumor cells may not express the antigen chosen for therapy. In one study for example, it was found that expression of known tumor antigens such as gp100 and tyrosinase was variable in different melanoma lesions in the same patient [de Vries et al., 1997]. Not all the malignant cells in the patient’s neoplasm expressed these determinants. Since the tumor cell population is heterogeneous, tumor cells that fail to express the defined antigen chosen for therapy are likely to escape destruction by the activated immune system.
The major advantage of vaccines prepared by transfer of tumor DNA into nonmalignant fibroblasts is that TAAs do not have to be purified or produced in large quantities. In comparison with protein vaccines, DNA-based vaccines provide prolonged expression and direct presentation of tumor antigens which results in robust and long-lasting activation of the immune system. From a practical point of view, these vaccines are easy and relatively inexpensive to prepare. Unlike other strategies, vaccines can be prepared from only a limited quantity of tumor-derived DNA, which can be obtained from small surgical specimens (vaccines can routinely be prepared from 50 μg of DNA). Furthermore, the recipient fibroblasts can be selected to meet the requirement for rapid expansion in culture and MHC restriction. The DNA-based vaccines offer a number of important advantages, which greatly encourage their further development for cancer immunotherapy.

1.8 Disadvantages of transfer of tumor-derived DNA transfer into fibroblasts for expression of TAAs

While vaccination based on transfer of tumor-derived DNA into highly immunogenic cells has a number of advantages, there are concerns as well. Since the proportion of total DNA that specifies TAAs is likely to be small, it is possible that a large number of the transfected cells may not express TAAs or may express TAAs at low levels. This concern is minimized, however, by preclinical data which indicate that the proportion of the cells that take-up tumor DNA and express TAAs is sufficient to induce an effective anti-tumor immune response and to significantly increase survival [Lichtor et al., 2008]. Another concern related to therapy with DNA-based vaccines is that genes specifying normal “self” antigens are likely to be expressed by the DNA-transfected cells, creating a danger that autoimmune disease might develop, although this has not been observed thus far. Inbred mice immunized with the DNA-based vaccine or tumor-bearing mice injected with therapeutic DNA-based vaccines failed to exhibit adverse effects [Lichtor et al., 2006]. Of course, protocols that depend upon the use of tumor cell-extracts, peptide eluates of tumor cells, fusion cells, cDNAs or RNAs derived from tumor cells are subject to the same concern. In DNA-based vaccines, genes encoding determinants expressed by non-neoplastic cells are likely to be present in the largest proportion relative to genes specifying TAAs. While the use of purified tumor antigen in the form of cDNA or polynucleotide vaccines specific for known TAAs eliminates this concern, those types of vaccines are dependent on the selection of the most “relevant” vaccinating epitope, as discussed above. It is also conceivable that a cellular vaccine, including one using nonmalignant fibroblasts might grow in the patient, forming a tumor. Conceivably, a transforming oncogene or a defective tumor suppressor gene might be transferred to a normal cell, provoking a neoplasm although this has not been observed. Overall, the disadvantages of DNA-based vaccines are few and are certainly no more difficult to overcome than those associated with other types of experimental tumor-vaccines.

1.9 Defects in TAA presentation by tumor cells

Defects in presentation of TAAs by tumor cells have been described in both murine as well as human tumors [Ohlen et al., 1990; Restifo et al., 1991]. They can result in tumor cell “escape” from host immunity. One mechanism is the loss of MHC determinants, which results in the impaired ability of the tumors to present TAAs. Loss of MHC antigen expression in several murine tumors is correlated with an increase in the malignant properties of the cells [Cohen & Kim, 1994]. Melanomas that recurred in mice treated with a
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vaccine prepared by transfer of DNA from murine melanoma cells into mouse fibroblasts were deficient in expression of MHC class I determinants [Kim & Cohen, 1994]. Primary and especially metastatic cells may have global or selective down-regulation of class I or class II HLA antigens, due to mutations in β2 microglobulin or TAP genes and thus they may fail to present TAAs in an immunogenic form to immune cells. Even if the host generates tumor-specific CTLs, the effector cells may not be able to eliminate the tumor. In addition to a failure to express HLA antigens, tumors may not express co-stimulatory molecules resulting in an inadequate immune response to TAAs by the host. Immunization with a DNA-based vaccine can overcome certain of these tumor “escape” mechanisms.

Significance

The most compelling reason for the vaccination strategy involving DNA-based cellular vaccines is the current lack of effective therapy for patients with malignant brain tumors such as breast cancer metastatic to the brain. This is verified by the dismal survival statistics, which have remained essentially unchanged for 30 years. Immunization with a vaccine that induces strong anti-tumor responses is an attractive addition or possibly even an alternative to conventional therapies. The DNA-based vaccines described in this review have shown remarkable therapeutic efficiency and survival benefits in some initial murine preclinical studies.

2. Experimental section

2.1 Treatment of intracerebral breast cancer in C3H mice by immunization with syngeneic/allogeneic fibroblast transfected with DNA from breast cancer cells

Whether results obtained by transfer of DNA from a tumor cell line into mouse fibroblasts can be applied to tumors that develop spontaneously is uncertain. Conclusions based on a model system involving tumor cell lines may not apply to neoplasms that arise spontaneously in patients. The appearance of spontaneous breast neoplasms in C3H mice provides an opportunity to investigate this question. DNA isolated from a breast neoplasm that arose spontaneously in the mammary gland of a C3H (H-2k) mouse in our animal colony (SB5b breast carcinoma cell line) was transferred into mouse fibroblasts (H-2k). To increase their immunogenic properties and to ensure rejection, the fibroblasts were modified to express H-2Kb determinants beforehand. H-2Kb determinants are allogeneic in C3H mice. The results indicated that C3H mice with intracerebral breast cancer treated solely by immunization with fibroblasts transfected with DNA from the same spontaneous breast neoplasm survived significantly longer (p<0.005) than mice in various control groups [Lichtor et al., 2005].

2.2 T cell mediated toxicity toward intracerebral breast cancer in mice immunized with syngeneic/allogeneic transfected fibroblasts modified to secrete IL-2, GM-CSF or IL-18

An MTS cytotoxicity assay was used to detect the presence of T cells reactive with breast cancer cells in mice injected i.c. with the mixture of SB5b cells and the modified, DNA-transfected fibroblasts. MTS is a tetrazolium compound which is bioreduced by viable cells into a formazan product that can be detected at 490 nm. The T cells obtained from the spleens of the injected mice were analyzed two weeks after the i.c. injection of the cell mixture. The results indicated that the cytotoxic response of greatest magnitude was in mice...
injected i.c. with the mixture of SB5b cells and transfected fibroblasts modified to secrete IL-2 or GM-CSF [Lichtor et al., 2005]. Lesser cytotoxic effects were present in mice injected i.c. with SB5b cells and transfected fibroblasts modified to secrete IL-18.

2.3 The proportion of T cells responsive to tumor cells in mice bearing an intracerebral tumor immunized intracerebrally with syngeneic/allogeneic transfected fibroblasts modified to secrete IL-2, IL-18 or IL-2 + IL-18

An ELISPOT-IFN-γ assay was used to determine the proportion of splenic T cells reactive with SB-5b cells in mice immunized with transfected fibroblasts modified to secrete IL-2, IL-18 or both IL-2 and IL-18. The animals were injected i.c. with a mixture of $1.0 \times 10^4$ SB-5b breast carcinoma cells and $1.0 \times 10^6$ treatment cells consisting of LMKbIL-2/SB5b, LMKbIL-18/SB5b, or a mixture of LMKbIL-2/SB5b and LMKbIL-18/SB5b cells. The animals were sacrificed at two weeks and an ELISPOT assay was done using the spleen cells to detect IFN-γ secretion in the presence of SB-5b tumor cells and antibodies against various T-cell subsets. The results indicate that the cellular anti-breast carcinoma immune response was mediated by CD4+, CD8+ and NK/LAK cells [Lichtor et al., 2005]. Although IL-18 secreting cells did not produce a significant anti-tumor immune response as detected with the ELISPOT assay, the combination of IL-2 with IL-18 secreting cells did result in an enhancement of the anti-tumor responses in comparison to animals that were treated with IL-2 secreting cells alone.

2.4 Increased numbers of responding T-cells were detected in the spleens and cervical lymph nodes of naïve mice or mice with i.c. breast cancer injected into the brain with cells from the immune$^{\text{high}}$ pool

An enrichment strategy for the vaccine was developed based on the hypothesis that if aliquots of a transfected cell population were divided into smaller populations, some populations by chance would contain more highly immunogenic cells than others. The populations with higher numbers of immunogenic cells could be identified by their stronger immunogenic response against SB5b cells in C3H/He mice. Two subpools that stimulated immunity to the greatest (immune$^{\text{high}}$ pool) and least (immune$^{\text{low}}$ pool) extents after three rounds of enrichment were selected for further study. To determine if systemic anti-tumor immunity was generated in tumor-free mice injected i.c. with cells from the immune$^{\text{high}}$ pool, cervical lymph node and spleen cells from the injected mice were analyzed by ELISPOT IFN-γ assays for responding T cells. Naïve C3H/He mice received 2 i.c. injections at weekly intervals of $1.0 \times 10^6$ cells from the immune$^{\text{high}}$ pool. One week after the second injection, mononuclear cells from the spleens and cervical lymph nodes of the immunized mice were analyzed for the presence of T cells responsive to the breast cancer cells. As controls, an equivalent number of cells from the non-selected master pool or cells from the immune$^{\text{low}}$ pool were substituted for cells from the immune$^{\text{high}}$ pool. As additional controls, the same protocol was followed except that the mice were injected i.c. with equivalent numbers of SB5b cells, with LMKb cells or with media. Mice injected with SB5b tumor cells received only one injection. The results from the cervical lymph nodes (figure 1) indicated that the highest number of responding cells was in mice injected i.c. with cells from the immune$^{\text{high}}$ pool ($p < 0.005$ vs. cells from mice in any of the other groups). Similar results were found in studies using the spleen cells from these animals [Lichtor et al., 2008].
Fig. 1. Development of anti-tumor immunity in cervical lymph nodes from naïve mice injected intracerebrally with an enriched cellular vaccine. ELISPOT IFN-γ assays for responding T cells in the cervical lymph nodes of mice injected i.c. with cells from the immuno<sup>high</sup> pool. Naïve C3H/He mice received intracerebral injections through a small burr hole two times at weekly intervals with 1.0 X 10<sup>6</sup> cells from the immuno<sup>high</sup> pool of transfected cells. One week after the second injection, mononuclear cells from the cervical lymph nodes of the injected mice were analyzed by ELISPOT IFN-γ assays for responding T cells. As controls, cells from the non-enriched master pool (LMIL-2K<sub>b</sub>/SB5b) or cells from the immuno<sup>low</sup> pool were substituted for cells from the immuno<sup>high</sup> pool. As additional controls, the same protocol was followed except that the mice were injected i.c. with media or with equivalent numbers of either SB5b or LMK<sub>b</sub> cells, or the mice were not injected. The animals injected i.c. with SB5b cells alone were injected only once. In some instances, SB5b cells (stimulated) were added to the cervical lymph node cell suspensions 16 hrs before the ELISPOT IFN-γ assays were performed (the ratio of spleen cells: SB5b cells = 10:1). In this assay the number of IFN-γ spots/10<sup>6</sup> cervical lymph node cells is measured. Error bars represent one standard deviation. *p* < 0.005 for the difference in the number of spots in the group injected with high pool LMIL-2K<sub>b</sub>/SB5b cells co-incubated with SB5b cells versus any of the other groups.

ELISPOT IFN-γ assays were also used to determine the number of responding T cells in the spleens of mice with i.c. breast cancer injected into the tumor bed with cells from the immuno<sup>high</sup> pool [Lichtor et al., 2008]. A micro cannula was placed into the right frontal lobe of C3H/He mice. SB5b cells (1.0 X 10<sup>4</sup> in 10 µl) were introduced into the brain through the cannula. On days two and nine following, the animals were injected through the cannula into the tumor bed with 1.0 X 10<sup>6</sup> cells from the immuno<sup>high</sup> pool. As controls, the same procedure was followed except that the cells from the non-enriched master pool or cells from the immuno<sup>low</sup> pool were substituted for cells from the immuno<sup>high</sup> pool. As additional controls, the tumor bearing mice were injected into the tumor bed with equivalent numbers...
of non DNA-transfected LMK\(^b\) cells or the mice were injected with SB5b cells alone. The results indicate that the highest number of responding T cells were in the spleens of tumor-bearing mice injected i.c. with cells from the immuno\(^{high}\) pool \((p < 0.05\) versus the number of responding spleen cells in mice injected with cells from the master pool and \(p < 0.005\) versus the number of spots obtained from any of the other groups).

2.5 T-reg cells are relatively deficient in the spleens of mice with i.c. breast cancer injected into the tumor bed with cells from the immuno\(^{high}\) pool

T-reg cells (regulatory T cells) are potent inhibitors of natural antitumor immunity. The success of immunotherapeutic protocols may depend upon the relative numbers of T-reg cells and cytotoxic T lymphocytes in tumor-bearing animals and patients. Quantitative RT-PCR for Foxp3, a transcription factor characteristic of T-reg cells, was used to determine the relative proportions of T-reg cells in the spleens and brains of mice with i.c. breast cancer injected into the tumor bed with cells from the immuno\(^{high}\) pool of transfected cells. Naïve C3H/He mice were injected i.c. with 5.0 X 10\(^4\) SB5b cells along with 1.0 X 10\(^6\) cells from the immuno\(^{high}\) pool of transfected cells. One week later, the animals received a second i.c. injection of cells from the immuno\(^{high}\) pool through the same Burr hole alone. As controls, the same procedure was followed except that the mice were injected with equivalent numbers of SB5b cells and cells from the non-enriched master pool or the immuno\(^{low}\) pool. The results indicate that CD4\(^+\)/CD25\(^+\)/Foxp3\(^+\) T-reg cells were relatively deficient in the spleens but not in the brains of animals injected with cells from the immuno\(^{high}\) pool [Lichtor et al., 2008]. An analysis by FACS of the spleens of the injected animals revealed a relative deficiency of CD4\(^+\)/CD25\(^+\) T cells and a corresponding increase in the relative numbers of CD8\(^+\) cells in the spleens of mice injected i.c. with cells from the immuno\(^{high}\) pool.

3. Discussion

Despite standard therapeutic approaches, the survival of patients with primary or metastatic tumors to the brain has not improved significantly in more than thirty years. There is an urgent need for new and more effective forms of treatment. Immunotherapy, designed to stimulate immunity to the autologous tumor, is under active investigation for a number of different histologic types of cancer. The enhanced immunotherapeutic properties of a vaccine prepared by transfer of a cDNA expression library derived from breast cancer cells into a mouse fibroblast cell line appears to have great potential in treatment of intracerebral tumors. As the transferred cDNA integrates spontaneously into the genome of the recipient cells, replicates as the cells divide and is expressed, the vaccine could be prepared from small amounts of tumor tissue, enabling treatment at an early stage of the disease, when tumor tissue is available in only limited amounts and the tumor is most susceptible to immune-based therapy. However, like other cellular tumor vaccines, only a small proportion of the transfected cell population was expected to have incorporated cDNA fragments that specified tumor antigens. A novel enrichment strategy has also been developed to increase the proportion of immunotherapeutic cells in the vaccine. A number of different strategies have been attempted to develop vaccines that generate enhanced anti-tumor immune responses in mice and patients with intracerebral neoplasms involving the central nervous system. Vaccines have been prepared by “feeding” antigen presenting (dendritic) cells apoptotic bodies from tumor cells or tumor cell lysates. Introduction of tumor cell-derived RNA into dendritic cells is another approach which has
been developed. Immunization with dendritic cells “fed” derivatives of tumor cells or transfected with tumor-RNA can result in the induction of immune responses against the broad array of tumor antigens expressed by the population of malignant cells including tumors of neuroectodermal origin [O et al, 2002; Rosenberg et al., 2004]. In patients, immunization with autologous dendritic cells transfected with mRNA from malignant glioma elicited tumor-specific CD8+ cytotoxic T-lymphocyte (CTL) responses against the patient’s malignant cells [Kobayashi et al., 2003]. Although results of dendritic cell immunotherapy have demonstrated promise in animal models, clinical trials have been disappointing thus far [Rosenberg et al., 2004].

Other tumor vaccination strategies have been used including modification of neoplastic cells to generate anti-tumor immune responses [Colombo et al., 1991; Gansbacher et al., 1990; Columbek et al., 1991; Mullen et al., 1992]. Immunization with tumor cells modified to secrete immune-augmenting cytokines such as IL-2 and GM-CSF has resulted in the development of generalized MHC-restricted anti-tumor immune responses in animal models [Cavallo et al., 1993; Connor et al., 1993; Dranoff et al., 1993; Marincola et al., 1994; Ohlen et al., 1990, Tahara et al., 1994]. Selective tumor regression was observed in experimental animals and patients receiving immunotherapy alone, in support of the potential of this type of treatment for patients with malignant disease [Valmori et al., 2000]. The effects of cytokine expression by central nervous system tumors (CNS) were examined initially using glioma cells that were engineered to secrete IL-4 [Yu et al., 1993]. In these studies it was demonstrated that IL-4 transduced glioma cells resulted in the development of anti-tumor immune responses. Delivery of an IFN-γ expression plasmid by cationic liposomes to the CNS tumor site was also found to induce significant anti-CNS tumor immunity in pre-clinical models [Liu et al., 2002]. Use of a high-titer adenoviral vector encoding IL-12 is another strategy that was reported to induce anti-tumor responses in a glioma model [Liu et al., 2002]. Epidermal growth factor variant III is a common alteration of the epidermal growth factor receptor found in human tumors, and a peptide vaccine has now proceeded to phase 1 and 2 clinical trials in patients bearing a malignant glioma with the ability of inducing potent T- and B-cell immunity and prolongation of survival [Sampson et al., 2008; Li et al., 2010].

In addition to immunotherapy, other gene therapy strategies have been attempted in the treatment of brain tumors. For example genetically modified, conditionally-replicating Herpes Simplex Virus Type 1 vectors with anti-tumor activity have been used with some encouraging results in Phase I and Phase II clinical trials [Cassady & Parker, 2010]. RNA interference therapy has also been shown to exhibit some therapeutic effects in pre-clinical and clinical studies in patients with gliomas [Guo et al., 2010]. High grade gliomas have been shown to express an IL-13 receptor that differs from the natural IL-13 receptor, and some pre-clinical and clinical trials have been undertaken using an adenoviral vector encoding a mutated human IL-13 fused to pseudomonas exotoxin that specifically binds to the IL-13 receptor [Candolfi et al., 2010]. Antisense oligodeoxynucleotides which target mRNA encoding TGF-beta2, a cytokine secreted by brain tumors with immunosuppressive activity, have been used in a series of Phase I and II clinical trials along with an international Phase III study with some success [Hau et al., 2009]. Finally one of the problems using large viruses in treating brain tumors is that the viruses have a limited capacity to infiltrate into the brain, but recently it has been shown that neural stem cells infected with replicating adenovirus can be used to enhance intratumoral distribution of the oncolytic vectors into a malignant glioma in comparison with virus injection alone [Tyler et al., 2009].
Previous studies indicated that transfection of genomic DNA from the malignant cells into a fibroblast cell line resulted in stable integration and expression of the transferred DNA [Cohen, 2001; Lichtor et al., 2005; Lichtor et al., 2006; Lichtor et al., 2008]. Both the genotype and the phenotype of the cells that took up the exogenous DNA were altered as portions of the transferred DNA were expressed. Immunization of tumor-bearing mice with the DNA-based vaccine resulted in the induction of cell mediated immunity directed toward the type of tumor from which the DNA was obtained, and prolongation of survival, consistent with the expression of an array of TAA by the transfected cells. This was the case for mice with melanoma, squamous cell carcinoma and in mice with breast cancer [Chopra et al., 2006; deZoeten et al., 2002; Lichtor et al., 2008]. Multiple undefined genes specifying TAA that characterize the malignant cell population were expressed by cells that took up DNA from the tumor. The number of vaccine cells could be expanded as required for multiple immunizations. In addition, the recipient cells can also be modified before DNA-transfer to increase their immunogenic properties, as for example, by the introduction of genes specifying immune-augmenting cytokines or allogeneic MHC-determinants, which act as strong immune adjuvants.

4. Conclusions

To be successful, every remaining tumor cell in the patient must be eliminated. It is unlikely that a single form of therapy is capable of achieving this goal. However immunotherapy in combination with surgery, radiation therapy and chemotherapy will likely find a place as a new and important means of treatment for patients with brain tumors. A major advantage of DNA-based vaccines is that they do not require protein purification or its production and yet they are able to elicit robust and long-lasting activation of the immune response, which results in tumor rejection. From a practical point of view, these vaccines are easy to prepare and they are relatively inexpensive. Only a limited quantity of tumor-derived DNA is required, which can be obtained from small surgical specimens. The enrichment strategy enables the generation of highly immunogenic pools of transfected cells with enhanced immunotherapeutic properties. Thus DNA-based vaccines offer a number of advantages, which greatly encourage their further development for cancer immunotherapy in general and specifically for treatment of breast cancer metastatic to the brain.

5. References


Candolfi, M., Xiong, W., Yagiz, K., Liu, C., Muhammad, A.K., Puntel, M., Foulad, D., Zadmehr, A., Ahlzadeh, G.E., Kroeger, K.M., Tesarfreund, M., Lee, S., Debinski, W.,
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