Genetic Disorders 2020: Gene Therapy for Cancer Treatment: Past, Present and Future- Deanna Cross, PhD, Center for Human Genetics, Marshfield Clinic Research Foundation, 1000 North Oak Avenue, Marshfield, WI 54449 USA

Abstract

The broad field of gene therapy promises a number of innovative treatments that are likely to become important in preventing deaths from cancer. In this review, we discuss the history, highlights and future of three different gene therapy treatment approaches: immunotherapy, oncolytic virotherapy and gene transfer. Immunotherapy uses genetically modified cells and viral particles to stimulate the immune system to destroy cancer cells. Recent clinical trials of second and third generation vaccines have shown encouraging results with a wide range of cancers, including lung cancer, pancreatic cancer, prostate cancer and malignant melanoma. Oncolytic virotherapy, which uses viral particles that replicate within the cancer cell to cause cell death, is an emerging treatment modality that shows great promise, particularly with metastatic cancers. Initial phase I trials for several vectors have generated excitement over the potential power of this technique. Gene transfer is a new treatment modality that introduces new genes into a cancerous cell or the surrounding tissue to cause cell death or slow the growth of the cancer. This treatment technique is very flexible, and a wide range of genes and vectors are being used in clinical trials with successful outcomes. As these therapies mature, they may be used alone or in combination with current treatments to help make cancer a manageable disease.

Key Words: Cancer gene therapy, Gene transfer, Immunotherapy, Oncolytic virotherapy

Introduction:

New therapy options need to be developed if the National Cancer Institute’s bold plan 1 of eliminating cancer death and suffering by 2015 is to be achieved. Five-year survival rates for pancreatic (4%), lung (15%), liver (7%) and glioblastoma (5%), a common form of brain cancer, remain abysmally low.2 Even prostate and breast cancers, which are highly amenable to treatment with 5-year survival rates better than 80%, still respond poorly to treatment at later stages and together result in more than 60,000 deaths a year.3 Current treatments often have far reaching negative side effects. The systemic toxicity of chemotherapy regimens, while not as severe as they once were, were, still often result in acute and delayed nausea, mouth ulcerations and mild cognitive impairments.4 In addition, long-term side effects from chemotherapy can include an increased risk of developing other types of cancers.5 Less serious, but potentially just as debilitating, side effects can also occur. Treatment for metastatic prostate cancer, while prolonging life, often causes hot flashes, impotence, incontinence and an increased risk of bone fractures.6 Therefore, entirely new
treatment methods are called for in order to alleviate the death and suffering caused by cancers.

The emerging field of cancer gene therapy offers a number of exciting potential treatments. The term gene therapy encompasses a wide range of treatment types that all use genetic material to modify cells (either *in vitro* or *in vivo*) to help effect a cure. Numerous *in vitro* and preclinical animal models, testing a wide variety of gene therapy agents, have shown remarkable efficacy. In lung cancer models, for example, survival benefits have been demonstrated using gene therapy to create cancer vaccines, target viruses to cancer cells for lysis and death, decrease the blood supply to the tumor, and introduce genes into the cancer cells that cause death or restore normal cellular phenotype. Preclinical gene therapy tests have also been performed on gliomas, pancreatic cancer and liver cancer, as well as many other cancers.

As with any new type of therapy, there are serious safety concerns. Initial enthusiasm for gene therapy as a treatment modality was curtailed by the death of a patient participating in a dose escalation gene therapy trial in 1999. While this was a trial to use gene therapy to correct a metabolic disease (ornithine transcarbamylase deficiency) and not a cancer trial, all gene therapy trials were reevaluated for safety. Since that time, newer and safer gene therapy delivery agents have been created and thousands of cancer patients globally have participated in gene therapy trials with remarkably few treatment side effects. However, when compared with the side effects of conventional chemotherapeutic treatments, these side effects are minimal.

This review focuses on the gene therapy trials that have progressed beyond the preclinical stage and are now in clinical trials in the United States. In order to explain these treatments, we have broken the field of cancer gene therapy treatments into three broad categories: immunotherapy, oncolytic virotherapy and gene transfer. Each section includes a brief history of the gene therapy category, a brief discussion of the techniques being used, a discussion of the state of current clinical trials and the future directions for the therapy.

**Immunotherapy**

**History**

Immunotherapy, or the concept of boosting the immune system to target and destroy cancer cells, has been a goal of cancer treatment for over 100 years. However, limited success has been achieved with traditional immunotherapy, as cancer cells tend to evolve mechanisms that evade immune detection. A wide array of gene therapy techniques are being used to overcome this limitation.

Currently gene therapy is being used to create recombinant cancer vaccines. Unlike vaccines for infectious agents, these vaccines are not meant to prevent disease, but to cure or contain it by training the patient’s immune system to recognize the cancer cells by presenting it with highly antigenic and immunostimulatory cellular debris. Initially cancer cells are harvested from the patient (autologous cells) or from
established cancer cell lines (allogeneic) and then are grown \textit{in vitro}. These cells are then engineered to be more recognizable to the immune system by the addition of one or more genes, which are often cytokine genes that produce pro-inflammatory immune immune stimulating molecules, or highly antigenic protein genes. These altered cells are grown \textit{in vitro} and killed, and the cellular contents are incorporated into a vaccine (figure 1A\textsuperscript{20}). Immunotherapy is also being attempted through the delivery of immunostimulatory genes, mainly cytokines, to the tumor \textit{in vivo}. The method of introducing a gene to the tumor varies and is discussed in more detail in the gene transfer section of this review. Once in the cancer cell, these genes will produce proteins that unmask the cells from immune evasion and encourage the development of antitumor antibodies (figure 1B\textsuperscript{21}).

Figure 1.


Another unique immunotherapy strategy facilitated by gene therapy is to directly alter the patient’s immune system in order to sensitize it to the cancer cells. One approach uses mononuclear circulating blood cells or bone marrow gathered from the patient. A tumor antigen, or other stimulatory gene, is then added to the selected cell type. These altered cells are now primed to cause an immune reaction to the cancer cells leading to cancer eradication (figure 1C\textsuperscript{22}). Alternatively, the gene can be added \textit{in vivo} using a targeted delivery system, such as an altered viral particle.\textsuperscript{23}

Initial trials using first generation vaccines have produced mixed results, highlighting both the potential for this therapy and the areas that still need to be perfected before these engineered cancer vaccines become part of standard cancer treatment. Early preclinical cancer vaccine models demonstrated positive results. Murine models using murine colon adenocarcinoma cells expressing human carcinoembryonic antigen (CEA) (CEA) demonstrated tumor reduction and long-lasting immune response when immunized with a vaccinia virus engineered to express CEA.\textsuperscript{24} However, when this type of vaccine was used in patients with breast cancer, no clinical response was
observed.\textsuperscript{25} This early experimental treatment highlights one of the limitations of using self-antigens for a vaccine. Although vaccines used for infectious agents generally lead to antigen-specific T-cell precursors in the range of 10\%, with self-antigens of cancer this response is often <1\%.\textsuperscript{26} Even when a successful immune response is mounted in clinical trials, it can be difficult to sustain. In a prostate cancer vaccine trial, a patient who achieved normal prostate specific antigen (PSA) levels for the year of the trial, developed rising PSA levels after vaccination was stopped. His PSA levels were stabilized again only with reinitiation of the vaccine therapy.\textsuperscript{27} These results, while not entirely positive, have given scientists a better understanding of the immune reaction to cancer and have led to the development of the next generation of cancer vaccines.

**Current Clinical Trials**

The next generation of vaccines is already in clinical trials for several cancer types. Table 1 provides a list of the more advanced clinical trials in this field, including phase, the type of cell used and the gene used to create a better immune response. These trials were picked to illustrate the fact that there are wide ranges of trials in different stages of efficacy testing using a variety of vectors for many cancer types.

<table>
<thead>
<tr>
<th>Cancer</th>
<th>Stimulating genes</th>
<th>ClinicalTrials.gov identifier</th>
<th>Description</th>
<th>Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prostate</td>
<td>CD-80, cytokine co-stimulatory molecule; CEA, carcinoembryonic antigen; GM-CSF, granulocyte-macrophage colony stimulating factor; IL-2, interleukin-2; MUC-1, mucin-1</td>
<td>NCT00105053</td>
<td>Mouse protein-sugars are expressed on allogeneic prostate cells to induce a hyperacute rejection response</td>
<td>II</td>
</tr>
<tr>
<td>Pancreatic</td>
<td>CEA and MUC-1</td>
<td>NCT00088660</td>
<td>Replication incompetent vaccinia and fowlpox viruses engineered to produce CEA and MUC-1 given</td>
<td>III</td>
</tr>
<tr>
<td>Tumor Type</td>
<td>Gene(s)</td>
<td>NCT Number</td>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>---------</td>
<td>------------</td>
<td>-------------</td>
<td></td>
</tr>
<tr>
<td>Prostate</td>
<td>GM-CSF</td>
<td>NCT00122005</td>
<td>Allogeneic prostate cells expressing the GM-CSF gene are used to induce immune response following chemotherapy and peripheral blood mononuclear cells infusion.</td>
<td></td>
</tr>
<tr>
<td>Lymphoma</td>
<td>GM-CSF and CD40L</td>
<td>NCT00101101</td>
<td>Autologous tumor cells are combined with allogeneic cells that express GM-CSF and CD40L and incorporated into a vaccine with low doses of IL-2.</td>
<td></td>
</tr>
<tr>
<td>Melanoma</td>
<td>IL-2</td>
<td>NCT00059163</td>
<td>Autologous tumor cells engineered to express IL-2 are incorporated into a vaccine.</td>
<td></td>
</tr>
<tr>
<td>Kidney</td>
<td>CD-80</td>
<td>NCT00040170</td>
<td>A modified replication incompetent adenovirus containing the</td>
<td></td>
</tr>
</tbody>
</table>
tumor antigen CD-80 is injected subcutaneously along with the cytokine IL-2 to produce an immune response to the prostate cancer.

Vaccines using engineered cells are showing great promise for the treatment of many cancers that respond poorly to conventional therapy. Vaccine therapy for non-small cell lung cancer is an example of an autologous vaccine therapy that has had good results in clinical trials. Recent clinical trials with GVAX, a vaccine made from autologous tumor cells modified to express granulocyte-macrophage colony-stimulating factor (GM-CSF) have led to further clinical testing. The initial phase I/phase II trial resulted in 3 of 33 subjects experiencing complete remission and an additional 7 who achieved stable disease for an average of 7 months. A further phase II study comparing GVAX alone to GVAX combined with cyclophosphamide in advanced stage patients demonstrated a clinical effect with 14 of the 53 participants experiencing stable disease and 1 patient experiencing stable disease for over 2 years. Median overall survival was between 5.4 months and 9.5 months with longer survival times seen on the cyclophosphamide arm of the study. The vaccine is now being tested in at least two larger phase II trials and phase III testing is planned. Unlike past trials, these trials have shown a demonstrable, but somewhat modest, effect on patient survival and, if phase III studies continue to show this impact, have the potential to become part of a treatment regimen. In addition to lung cancer, GVAX is also being tested in other cancers. An allogeneic GVAX vaccine using a combination of prostate cell lines that are engineered to express GM-CSF is being tested as a treatment for prostate cancer. This vaccine has been shown to increase the PSA doubling time of patients with progressive prostate cancer and increase time to disease progression by several months. Currently, several large phase III trials are underway to determine if there is an increase in life expectancy as well.

Other clinical trials are demonstrating the potential of unmasking the tumor from immune invasion using immunostimulatory genes inserted directly into the tumor tissue. For example, MDA-7 (IL-24), a cytokine that induces cancer cell death, is currently in clinical trials for its ability to cause a systemic immune reaction in malignant melanoma patients. Melanomas have long been observed to elicit an immune response from the patient and many attempts have been made over the years to bolster this reaction to effect a cure. After packaging in a replication incompetent adenovirus, the MDA-7 gene is injected intratumorally and induces apoptosis. A clinical trial demonstrated that this treatment lead to complete response and partial response in 2 of 28 patients. In 22 other patients, systemic immune activation was
observed, as well as local apoptosis. The clinical response observed in these trials has led to a phase II study to determine if this response can induce apoptosis in distant metastasis via a systemic immune reaction when the vector is injected intratumorally.

Current clinical trials seeking to directly stimulate the immune system for cancer destruction also show promising results. One example of this type of immunotherapy is the current clinical trial using the TRICOM vaccines. These vaccines incorporate a cancer antigen into a modified virus, either vaccinia or fowlpox, that also contain three immunostimulatory genes: B-lymphocyte activation antigen B7-1 (B7-1), intercellular adhesion molecule 1 (ICAM-1) and lymphocyte function-associated antigen 3 (LFA-3). The PANVAC-VF vaccine is a vaccinia virus modified to deliver deliver mucin-1 (Muc-1) and CEA, in addition to the immunostimulatory genes. The vaccine is injected subcutaneously and followed by boosting vaccines of a fowlpox virus modified in the same manner as the vaccinia virus. This vaccine strategy recently completed a phase III trial in pancreatic cancer. In addition Prostvac, a vaccine vaccine that uses the fowlpox virus engineered to express Muc-1 (a gene highly expressed in tumors) to induce an immune response, is exhibiting promising results. Phase I data revealed a 3- to 4-fold increase in PSA doubling time when patients were given the vaccine. Currently, large phase II studies are underway.

**Future Directions**

While current clinical trials are progressing much better than earlier ones, there are still some areas that could be improved. For example, many of the most promising vaccines rely on autologous cells for vaccine production. These vaccines do give the patient a truly personalized vaccine; however, they may also present a long-term problem because of the expense and effort needed to create it. Few hospitals contain a facility for vaccine production and substantial time and expertise are required to grow the cells and create a custom vaccine. One way around this obstacle is the creation of allogeneic alternative vaccines, though efforts to create an effective allogeneic alternative to GVAX have not been as successful in trials as the autologous GVAX vaccine. However, other allogeneic strategies, such as GM.CD40L, have been more successful. GM.CD40L is a vaccine composed of autologous tumor cells mixed with allogeneic tumor cells that have been engineered to produce both GM-CSF and CD40L. GM.CD40L is currently in a phase II trial for treatment of malignant melanoma. Combining these genes may lead to a stronger immune response than either gene used alone. In addition, in a pancreatic cancer trial, a vaccine of allogeneic pancreatic cancer cells engineered to produce GM-CSF combined with surgery has shown impressive phase II results with 76% survival at 2 years compared to the historic average of <50% at 2 years.

As with any cancer monotherapy, combination therapy using vaccines may be more effective than vaccine therapy alone. Cancer vaccines that have presented only modest immune response may find usefulness as an adjuvant therapy for use after surgery or chemotherapy to eliminate any remaining cancer cells. With the current round of ongoing clinical trials, the potential of gene therapy cancer vaccines is close to being fulfilled. The initial phases of vaccine development are being completed and it is likely that there soon will be effective cancer treatments that incorporate vaccines into the therapy regimen.

**Oncolytic Agents**
**History**

Another growing area of gene therapy treatment for cancer is the use of oncolytic vectors for cancer destruction. Like immunotherapy, this is a concept that has been around for almost a century and, like immunotherapy, it is undergoing a renaissance due to gene therapy. Oncolytic gene therapy vectors are generally viruses that have been genetically engineered to target and destroy cancer cells while remaining innocuous to the rest of the body. Oncolytic vectors are designed to infect cancer cells and induce cell death through the propagation of the virus, expression of cytotoxic proteins and cell lysis (figure 2↓). A number of different viruses have been used for this purpose, including vaccinia, adenovirus, herpes simplex virus type I, reovirus and Newcastle disease virus. These viruses have been chosen, in many cases, for their natural ability to target cancers, as well as the ease at which they can be manipulated genetically.

![Figure 2. Schematic diagram of oncolytic virotherapy.](image)

Initial trials of oncolytic therapies have highlighted both its incredible power, as well as unique obstacles to treatment implementation. Mammalian models of oncolytic gene therapy have worked remarkably well. In murine models, both colon and bladder cancer have shown survival benefits and reduced metastasis using oncolytic viral agents. In a canine model, using an oncolytic virus designed to destroy osteosarcoma, survival was prolonged even in immunocompetent dogs with syngenic osteosarcoma. However, there are several unique stumbling blocks for oncolytic virotherapy in humans. Most people have antibodies to the common viruses used for therapy development which often leads to an immune response that clears the viral agent before it has had time to infect cells. In addition, the use of replication competent viral particles often calls for increased safety precautions, making clinical trials more expensive and cumbersome. In a trial using a modified vaccinia virus to treat breast and prostate cancer, patients were required to be isolated in a specialized hospital facility for a week to ensure that the virus had completely cleared before being allowed back into the general population. Because of these limitations, there have been relatively few trials with oncolytic therapy. However, new vectors are being being created and past experience is being incorporated into current trials to enhance results so that they mimic those in animal studies.

**Current Clinical Trials**

Even in this early stage, oncolytic viral therapy has demonstrated some success. Both adenovirus and herpes virus agents have ongoing clinical trials for intractable cancers. The most notable adenoviral therapy is the ONYX-015 viral therapy. ONYX-015 is an adenovirus that has been engineered to lack the viral E1B protein. Without this protein, the virus is unable to replicate in cells with a normal p53 pathway. In addition,
the E1B protein is essential for RNA export during viral replication. Cancer cells often have deficiencies in the p53 pathway due to mutations and thus, allow ONYX-015 to replicate and lyse the cells. Cancer cells also exhibit altered RNA export mechanisms that allow for the export of viral RNA even in the absence of the E1B protein. ONYX-015 has been tested in phase I and II trials on squamous cell carcinoma of the head and neck that resulted in tumor regression which correlated to the p53 status of the tumor. Tumors with an inactive pathway demonstrated a better response. Phase II trials of ONYX-015, in combination with chemotherapy, demonstrated even better tumor response and have led to a phase III study. In addition to squamous cell carcinoma, ONYX-015 is currently being tested as a preventative treatment for precancerous oral tissue, the theory being that even in the precancerous state, there are p53 pathway inactivating mutations that will allow the oncolytic adenovirus to replicate and eliminate the cells before they become cancerous.

The second type of oncolytic virotherapy undergoing clinical trials uses herpes simplex simplex virus type 1 (HSV-1). Two vectors, G207 and NV1020, are currently in phase I and phase II trials for treatment of intractable cancers. Mutations in several genes of these herpes viruses ensure that they replicate efficiently only in cancerous cells. G207 is mutated so that it has attenuated neurovirulence and cannot replicate in nondividing cells. NV1020, a derivative originally used for vaccine studies, has multiple mutations, including a deletion in the thymidine kinase region and a deletion across the long and short components of the genome, and an insertion of the thymidine kinase gene under the control of the α4 promoter. These viral vectors have two distinct cell killing mechanisms. The lytic portion of the life cycle directly kills cells and the thymidine kinase that is expressed from the viral genes sensitizes cells to ganciclovir. These viral therapy vectors have been used with great success in vitro and in model animals against a wide number of solid cancers. Clinical trials trials using these vectors include a phase I trial of G207 for treatment of malignant glioma and a phase I/II trial of NV1020 for treatment of colorectal cancer metastases to the liver. In addition, NV1020 has also been tested for treatment of glioblastoma.

**Future Directions**

Because oncolytic virotherapy is not yet a mature technology, there is plenty of room for improved treatment vectors. In order for virotherapy to be successful, viral particle production rates in the infected cancer cells must outstrip the growth rate of the uninfected cancer cells. This may be difficult to achieve with large established tumors and may mean that virotherapy must be combined with an existing therapy, such as surgery, to decrease the number of cancer cells in the initial treatment. In addition, the most effective treatment delivery method is yet to be determined. In preliminary studies, systemic injection required 1000x the viral load necessary to achieve results than injection intratumorally.

However, once these factors are overcome, there are many benefits to oncolytic therapy. The selective nature of the virotherapy ensures that healthy tissue will be minimally impacted. In addition, when combined with cytotoxic gene expression, this therapy can affect not only rapidly dividing cells, but those in the surrounding tissue making the microenvironment less favorable for cancer growth. The combination of the
the powerful killing nature of these vectors combined with the selectivity makes them an exciting avenue for lowering the number of cancer deaths.

**Gene Transfer**

**History**

One of the most exciting treatments to emerge from the concept of gene therapy is that of gene transfer or insertion. This is a radically new treatment paradigm involving the introduction of a foreign gene into the cancer cell or surrounding tissue. Genes with a number of different functions have been proposed for this type of therapy, including suicide genes (genes that cause cellular death when expressed), antiangiogenesis genes and cellular stasis genes (figure 3). A number of different viral vectors have been used in clinical trials to deliver these genes, but most commonly have used a replication incompetent adenovirus. Nonviral methods, including naked DNA transfer and oligodendroner DNA coatings, as well as electroporation are also viable modes of gene delivery.56 The type of delivery vehicle chosen depends on the desired specificity of the gene transfer therapy, as well as the length of time the gene must be expressed in order to be effective. For instance, a replication incompetent adenoviral vector containing the herpes simplex virus thymidine kinase (HSVtk) gene needs only transient expression to accomplish cell death and is generally delivered via an adenoviral vector.57 However, antiangiogenesis antiangiogenesis genes, such as sFLT-1 and statin-AE, need continuous expression for therapeutic effect and have been delivered using plasmids that contain a transposon to insert the gene into the cellular DNA.58

![Figure 3.](image)

Initial attempts to implement gene transfer therapy have highlighted its promise, as well as some delivery difficulties. Delivery of the therapeutic gene to the target cells has to be effective enough to elicit a response and has been difficult to achieve with many of the current technologies. In addition, extra precautions must be taken to ensure the therapeutic gene does not integrate into unwanted cell types, such as reproductive tissues. Earlier gene transfer trials suffered from gene silencing so that even if the gene was effectively introduced into the cell, it was not expressed or was expressed only for a limited length of time.59 Despite these hurdles, solid tumors such as prostate, lung and pancreatic tumors have been treated successfully in animal models using a variety of genes and transfer methods.60–62 Special precautions must be taken if DNA is inserted into the cell chromosome. The insertional site must be in an area of the genome that does not promote cancer. Retrotransposons, such as sleeping beauty (an artificially constructed retrotransposon that is used to insert genes into vertebrate chromosomes) often insert into actively transcribed genes causing potential problems for cellular function.63 Preclinical models using gene insertion
techniques, such as murine models for glioma, showed significantly greater survival when they administered antiangiogenic genes via a retrotransposon system injected intracranially.58

**Current Clinical Trials**

Because gene transfer technology encompasses such a diverse set of therapeutic options, it is impossible to describe examples for every treatment. However, a partial list of treatments in significant current clinical trials with a brief description of each is presented in table 2. Below, we highlight several of the late stage clinical trials, as well as some exciting innovative approaches that further highlight the promise of gene transfer therapy.

### Table 2.

**Selected recent gene transfer clinical trials.**

<table>
<thead>
<tr>
<th>Cancer</th>
<th>Transferred genes</th>
<th>ClinicalTrials.gov identifier</th>
<th>Description</th>
<th>Phase</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HSVtk</strong>, herpes simplex virus thymidine kinase; TNF-α, tumor necrosis factor alpha</td>
<td></td>
<td></td>
<td>A cytocidal cyclin G1 construct accumulates preferentially in the tumor cells to block the action of cyclin G1 and initiate cell death</td>
<td>I</td>
</tr>
<tr>
<td>Pancreatic</td>
<td>Rexin-G</td>
<td>NCT00121745</td>
<td>The HSVtk gene is introduced into glioblastoma cells via a mouse retrovirus. Glioblastoma cells with the HSVtk gene are then sensitive to the drug ganciclovir which is administered</td>
<td>I</td>
</tr>
<tr>
<td>Glioblastoma</td>
<td>HSVtk</td>
<td>NCT00001328</td>
<td></td>
<td>I</td>
</tr>
</tbody>
</table>
### Table

<table>
<thead>
<tr>
<th>Disease</th>
<th>Gene</th>
<th>Trial ID</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Head and neck</td>
<td>p53</td>
<td>NCT00041613</td>
<td>Transfer of the p53 gene via a replication incompetent adenovirus to tumor cells to inhibit cell growth and induce apoptosis</td>
</tr>
<tr>
<td>Melanoma</td>
<td>MDA-7</td>
<td>NCT00116363</td>
<td>MDA-7 a novel tumor suppressor molecule is introduced into the melanoma cells and overexpression inhibits cellular proliferation and induces apoptosis</td>
</tr>
<tr>
<td>Pancreatic</td>
<td>TNF-α</td>
<td>NCT00051467</td>
<td>The TNF-α gene under the control of a radiation inducible promoter is introduced into tumor cells and in combination with the radiation therapy induces cell death</td>
</tr>
</tbody>
</table>

TNFerade is one such treatment option that is currently in late stage II trials. This agent is a replication incompetent adenoviral vector that delivers the tumor necrosis factor-α (TNF-α) gene under the transcriptional control of a radiation inducible promoter. TNF-α is a cytokine with potent anticancer properties and high systemic toxicity, and TNF-α gene therapy provides a way to target this molecule to only the cancer cells through the use of intratumoral injections and a promoter that is activated by radiation therapy. Once TNFerade is injected, the patient then receives radiation therapy to the tumor to activate the gene. The gene then produces the TNF-α molecule which in combination with the radiation therapy promotes cell death in the affected cancer cells and surrounding cells. A phase I study of patients with soft tissue sarcoma using TNFerade demonstrated an 85% response rate including 2 complete responses. In another large phase I study of patients with histologically confirmed...
advanced cancer, 43% of the patients demonstrated an objective response with 5 of 30 exhibiting complete response to the treatment. Larger studies are being conducted using TNFerade for treatment in pancreatic, esophageal, rectal cancer and melanoma.

Another exciting gene therapy treatment agent is Rexin-G, the first injectable gene therapy agent to achieve orphan drug status from the Food and Drug Administration for treatment of pancreatic cancer. This gene therapy agent contains a gene designed to interfere with the cyclin G1 gene and is delivered via a retroviral vector. The gene integrates into the cancer cell’s DNA to disrupt the cyclin G1 gene and causes cell death or growth arrest. In a phase I trial, 3 of 3 patients experienced tumor growth arrest with 2 patients experiencing stable disease. These results have led to larger phase I and II trials.

A gene transfer technology that shows great promise is the replication incompetent adenovirus delivering the HSVtk gene to a tumor followed by ganciclovir treatment. Ganciclovir is not toxic unless metabolized by the HSVtk gene, and therefore only the cancer cells that are treated with the gene and the surrounding cells will be affected by treatment. In a large phase I study involving glioblastoma patients, the HSVtk-engineered viral treatment increased median survival from 39 weeks to 70.6 weeks and was the first glioblastoma gene therapy trial to show any measurable improvement in survival.

Several agents that use a replication incompetent adenoviral vector to deliver the p53 gene to cancer cells are also currently in phase II and III trials. The p53 gene is an important cell cycle regulator that has been extensively studied and is mutated in 50% to 70% of human tumors. Mutations in this gene are often linked to aggressiveness. It has been shown that restoration of a functional p53 gene in cancer cells results in tumor cell stasis and often apoptosis. Using this information, INGN 201, an adenoviral vector containing p53 for gene transfer, is in current phase III testing for squamous cell carcinoma of the head and neck, and has completed phase I studies on prostate, ovarian, glioma and bladder cancer.

**Future Directions**

Gene transfer, while a radical new type of treatment, is also the only gene therapy product to obtain regulatory approval in any global market, as demonstrated by China’s 2003 approval of Gendicine for clinical use. Gendicine is a modified adenovirus that delivers the p53 gene to cancer cells and is approved for the treatment of head and neck squamous cell carcinoma. Since approval, thousands of patients have been treated in China; some with repeated injections. As yet, large-scale efficacy trial results have not been published; the results of which are eagerly awaited.

Gene transfer technology allows an incredible diversity of treatment possibilities. This diversity can be used to complement traditional therapies, as well as provide radically new frontiers for treatment. Gene transfer therapy can rely on the current information known about the genetics of cancer formation, bringing a more sophisticated and personalized approach to therapy. Current gene transfer trials have demonstrated statistically significant survival improvements for cancers such as glioblastoma and pancreatic cancer, as discussed previously. These studies have provided very
encouraging signs that current research is on the right path. New delivery methods and more sophisticated gene expression cassettes will create better therapeutic alternatives to make the goal of cancer treatment and eradication achievable.

Conclusions

The field of cancer gene therapy is rapidly maturing and will no doubt be part of the future of cancer therapeutics. Several very exciting cancer vaccine treatments are in late stage trials, thanks to the advent of genetic engineering. In addition, gene transfer technology for cancer treatment holds great promise for increasing the effectiveness of current chemotherapeutic treatment regimens. Significant advances have been made in the field of oncolytic virotherapy, and trials are in progress that incorporate this technique for precancerous, as well as cancerous treatment. Many of the past obstacles to treatment are being actively overcome and current second and third generation therapeutics are being tested. While not all the current trials will lead to a viable therapeutic agent, there is great hope that these advances will help relegate cancer to a manageable chronic disease without severe suffering and death.

References


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