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Effect of Electrically Mediated Intratumor and Intramuscular Delivery of a Plasmid Encoding IFN α on Visible B16 Mouse Melanomas

Interferon α may be used as a single agent therapy for metastatic malignant melanoma or as an adjuvant to chemotherapy. Delivery of interferon α by gene therapy offers an alternative to recombinant protein therapy. Electrically mediated delivery enhances plasmid expression in a number of tissues, for instance skin, liver, muscle and tumors including melanomas. Here we compare the effect of delivery of a plasmid encoding mouse interferon α on growth of visible B16 mouse melanomas following electrically mediated delivery to muscle or directly to the tumor. Intratumoral delivery of interferon α plasmid not only slows melanoma growth, but induces complete, long term, regression. This effect was not observed after intramuscular delivery.

Introduction

A variety of clinical trials have investigated interferon α (IFN α) as a single agent therapy or as an adjuvant to chemotherapy for metastatic malignant melanoma (1, 2). IFN α treatment increases patient survival and durability of cure. In the B16 mouse melanoma model, long term in vitro exposure of cells to IFN α increases mouse survival after injection and delays tumor growth (3). When B16 cells are irradiated and used as a vaccine, a significant level of protection is elicited which prevents tumor establishment (4). IFN α producing clones are less tumorigenic, although mice immunized with irradiated cells are only slightly protected (5).

Delivery of IFN α by gene therapy offers an alternative to recombinant protein therapy. In vivo electroporation, which has been used in phase II clinical trials for the delivery of chemotherapeutic agents to cancer tumors (6, 7) also enhances plasmid delivery and expression (8, 9). The electrically mediated enhancement of plasmid expression has been demonstrated in skin (10-12), liver (13), and muscle (14-19). Intramuscular electroporation delivery of a plasmid encoding IFN α has been used to affect growth of squamous cell carcinomas in mice (20).

Electrically mediated enhancement of plasmid expression following direct intratumor injection of reporter plasmid has been demonstrated in rat brain tumors (21), mouse melanomas (22), and rat liver tumors (23). Therapeutic responses to experimental melanomas have also been described following electrically mediated delivery of a dominant negative Stat3 variant in mouse melanomas (24) or combinations of cytokines such as interleukins 2 and 12 (25) or interleukins 12 and 18 (26). The combination of electrochemotherapy and cytokine plasmid delivery by electroporation into melanomas prevents tumor recurrence and induces long-term antitumor immunity in mice (27).

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Simple muscle injection of a plasmid encoding IFN α is effective in reducing the onset of growth of B16 melanomas and in increasing mouse survival (28). Here we directly compare the effect on growth of visible B16 mouse melanomas following electrically mediated delivery to muscle or directly to the tumor. Direct intratumoral delivery of plasmid not only slows melanoma growth, but induces complete, long term, regression. This effect was not observed after intramuscular delivery.

**Materials and Methods**

**Plasmids**

Plasmids VR1255, encoding the luciferase gene driven by the CMV promoter, and VR4111, encoding mouse interferon α (28, Vical, San Diego, CA), were prepared using an endotoxin-free method (Qiagen, Valencia, CA), suspended in sterile injectable saline, and endotoxin levels confirmed to be less than 0.1 EU/µg (QCL-1000, Biowhittaker, Walkersville, MD).

**Cell culture and tumor induction**

B16.F10 (ATCC CRL-6475) mouse melanoma cells were maintained in McCoy’s 5A medium supplemented with 10% fetal bovine serum and 90 µg/ml gentamycin at 37 C and 5% CO₂. A suspension containing 10⁶ trypsinized B16 cells (>90% viability) in 50 µl sterile injectable saline was injected subcutaneously in the left flank of 7 week old female C57Bl/6 mice (Jackson Laboratory, Bar Harbor, ME) using a 0.5 inch 30 gauge needle. Tumors were allowed to grow 8 days to a diameter of approximately 4 mm before treatment, equivalent to a volume of approximately 34 mm³. Mice were challenged by subcutaneous injection of 5 x 10⁵ cells on the opposite flank.

**Plasmid delivery**

Mice were anesthetized during all treatments using a mixture of 2% isoflurane and 98% O₂.

**Tumors**. Tumors were injected with 50 µl 2 µg/µl plasmid DNA unless otherwise noted, then pulsed immediately with six rotating 100 µs pulses at a nominal field strength of 1500V/cm and a frequency of 1 Hz with a 7.5mm diameter 6 needle electrode (29).

**Muscle**. Left and right gastrocnemius muscles were alternately injected weekly with 50 µl 2 µg/µl VR4111 plasmid. If treated with electroporation, three 20 ms pulses per tissue area were applied immediately at a nominal field strength of 100V/cm with a custom applicator containing 4 needle electrodes in a 2.5 x 5 nm rectangle (30). All pulses were delivered using a T820 Electrode porator (BTX, San Diego, CA).

**Tumor monitoring**

Tumors were measured twice weekly using a digital caliper. Tumor volume was calculated by the standard formula \(v=ab^2/6\), where \(a\) is the longest diameter, and \(b\) is the next longest diameter perpendicular to \(a\). In the case of continued tumor growth or tumor recurrence, the animal was considered incurable and humanely euthanized when the tumor volume reached 1000 mm³. Each individual tumor volume was normalized to its tumor volume on day 0, the first day of treatment.

**ELISAs**

For determination of IFN α levels after intramuscular delivery, blood was removed from animals via the tail vein, allowed to clot at 4º C, then the serum removed. For determination of IFN α levels after intratumor delivery, tumors were induced and treated as described, then mice were humanely euthanized at set times after treatment as indicated. Tumors were removed, weighed, homogenized in PBS containing protease inhibitors (P8340, Sigma-Aldrich, St. Louis, MO), and centrifuged at 1000 x g for 5 minutes. Serum and tumor supernatants were assayed for IFN α production by ELISA (PBL Biomedical Laboratories, New Brunswick, New Jersey).

![Figure 1: Effect of intramuscular delivery of plasmid encoding interferon α (VR4111) on growth of large, visible B16 mouse melanomas. Tumors were induced, 100 µg plasmid VR4111 was delivered intramuscularly with or without electroporation, and tumor growth monitored as described in methods. Lines represent mean and standard error of the mean. ○, no treatment, n = 6; □, IM VR4111 injection only, n = 7; ■, IM VR4111 with pulses as described in methods, n = 7.](image-url)
Previously, intramuscular injection of 100 µg VR4111 at 1 week intervals has been shown to successfully slow the onset of B16 melanomas (28). Here, B16 cells were injected and tumors allowed to grow 8 days to an average diameter of 4mm (day 0). Animals received three weekly treatments of 100 µg plasmid in alternating gastrocnemius muscles with or without electric pulses as described in methods. Tumors in the treated groups were compared to tumors from animals that received no treatment and continued to grow. In the groups that received injection of plasmid only or injection of plasmid followed by electric pulses, resulting in much higher levels of systemic IFN α expression, no significant effect was seen on tumor growth. However, a significant effect on survival was observed (p<0.005) in the group receiving intramuscular IFN α plasmid and electroporation when compared to untreated animals or to animals receiving plasmid injection alone. This study differs from the study by Horton et. al. (28) in that treatment began 8 days after injection of 10^6 cells rather than 3 days after injection of 10^4 cells, so the tumors were much larger when treatment began. This may explain the ineffectiveness of intramuscular treatment in this case.

Direct tumor delivery of VR4111 was more effective than intramuscular delivery in inducing B16 tumor regression (Figure 2a). Tumors were induced and allowed to grow as above, then treated with electrically mediated delivery of 100 µg control vector (VR1255). This treatment insignificantly slowed tumor growth. Electrically mediated delivery of VR4111 slowed tumor growth and induced long term complete regressions in a dose dependent manner. After delivery of 100 µg plasmid, 5 of 7 tumors regressed completely, after 50 µg, 3 of 7 regressed completely, and after delivery of 25 µg plasmid, 2 of 7 regressed completely (Figure 2b). These regressions lasted 75 days until the animals were challenged subcutaneously. Five of the 10 tumor free animals were resistant to subcutaneous challenge, while tumors grew in naive control animals, indicating that some antitumor immunity was generated.

Different electroporation parameters and electrodes were used for intratumor or intramuscular delivery. Interestingly, these different delivery methods and sites resulted in similar peak serum levels of IFN α, although the time course of expression differed (Figure 3). Serum expression peaked at day 1 following intratumor delivery and decreased immediately, while serum expression after intramuscular delivery peaked more slowly at day 4 and decreased more gradually. Since the tumors were large and B16 melanoma is particularly aggressive, the faster increase in serum IFN α levels observed after intratumor delivery may be more effective in inhibiting tumor growth.

Since intratumor delivery was effective therapeutically, tumor levels of IFN α after intratumor delivery were determined (Figure 4). No IFN α expression was noted after injection alone of VR4111 or after electrically mediated delivery of a plasmid encoding luciferase (VR1255, data not shown). Intratumor IFN α expression increased significantly after electrically mediated delivery of VR4111, peaking at 369 pg/mg (369 ng/g) tumor at day 1 while serum levels peaked approximately 575 fold lower at 643 pg/ml serum. This high level of intratumor expression may be responsible for the significant antitumor effect.
Intramuscular delivery of plasmids encoding IFN α significantly affects the growth of squamous cell carcinomas (20) and B16 melanomas (28). The data here indicate that, while intramuscular delivery of VR4111 does result in serum expression of IFN α and significantly increases survival, delivery by this method has little effect on the growth of visible experimental B16 melanomas. When tumors are accessible, delivery of VR4111 directly to the tumor appears to be a much more effective treatment method, inducing tumor regression and some level of resistance to challenge with B16 cells.

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References and Footnotes


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