Degradation of Fibrillar Collagen in a Human Melanoma Xenograft Improves the Efficacy of an Oncolytic Herpes Simplex Virus Vector

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Abstract

Oncolytic viral therapy provides a promising approach to treat certain human malignancies. These vectors improve on replication-deficient vectors by increasing the viral load within tumors through preferential viral replication within tumor cells. However, the inability to efficiently propagate throughout the entire tumor and infect cells distant from the injection site has limited the capacity of oncolytic viruses to achieve consistent therapeutic responses. Here we show that the spread of the oncolytic herpes simplex virus (HSV) vector MGH2 within the human melanoma Mu89 is limited by the fibrillar collagen in the extracellular matrix. This limitation seems to be size specific as nanoparticles of equivalent size to the virus distribute within tumors to the same extent whereas smaller particles distribute more widely. Due to limited viral penetration, tumor cells in inaccessible regions continue to grow, remaining out of the range of viral infection, and tumor eradication cannot be achieved. Matrix modification with bacterial collagenase coinjection results in a significant improvement in the initial range of viral distribution within the tumor. This results in an extended range of infected tumor cells and improved virus propagation, ultimately leading to enhanced therapeutic outcome. Thus, fibrillar collagen can be a formidable barrier to viral distribution and matrix-modifying treatments can significantly enhance the therapeutic response. (Cancer Res 2006; 66(5): 2509-13)

Introduction

Oncolytic vectors, mutant viruses that replicate preferentially in tumor cells, have shown promise in various preclinical tumor models (1). Oncolytic viral therapy employs a novel method of tumor destruction mediated by viral replication and selective lysis of cancer cells (2). Creation of more virus by infected tumor cells and the resultant infectious spread improve performance over more passive forms of therapeutic delivery (3, 4). Early-phase human trials of G207, an oncolytic herpes simplex virus (HSV) vector, for treatment of recurrent malignant glioblastomas have shown both safety and efficacy (5). However, the inability to efficiently propagate and infect cells distant from the injection site limits the capacity of oncolytic viruses to achieve consistent therapeutic responses (6). In this study, we show that fibrillar collagen, a major barrier to macromolecular transport in the tumor interstitium (7–9), also limits HSV vector distribution within a melanoma. Direct degradation of the fibrillar collagen network improves viral distribution, leading to improved oncolytic viral therapy.

Materials and Methods

Viral vectors. The HSV-1 recombinant viruses used in this study were the replication defective mutant Gal4 (ICP4−, lacZ+; from Dr. Neal DeLuca, University of Pittsburgh, Pittsburgh, PA; ref. 10) and MGH2 (ICP6−, lacZ+, eGFP+; from Drs. E. Antonio Chiocca and Yoshi Saeki, The Ohio State University, Columbus, OH). MGH2 is a replication conditional virus attenuated by two nonessential viral gene deletions, ICP6 and lacZ (11). Virus replication is impaired in nontumor cells but not in tumor cells. Gal4 and MGH2 stocks were propagated in E5 and E26 cells (from Dr. Neal DeLuca; ref. 12), respectively, which supply the HSV-1 ICP4 protein (E5) or HSV ICP4 and ICP27 proteins (E26) in trans. To obtain GFP-labeled HSV particles, E5 and E26 cells were transfected with a plasmid encoding the fusion protein VP16-GFP (pVP16-GFP; ref. 13) and infected with Gal4 and MGH2, respectively. After purification and concentration, the titer of each virus preparation was quantified by counting lacZ-positive cells for Gal4 and GFP-positive cells for MGH2.

Dorsal skinfold window preparation. Human melanoma Mu89 cells were grown in dorsal skinfold chambers in severe combined immunodeficient (SCID) mice as previously described (9). All animal experiments were done with the approval of the Institutional Animal Care and Use Committee.

Injection and imaging of labeled vectors and tracers. For dorsal chamber tumor studies, HSV vectors labeled with VP16-GFP were mixed with either 0.2 μg/μL bacterial collagenase (Sigma, C0773, St Louis, MO) or PBS to a final titer of 108 transducing units/μL. For microsphere experiments, quantum dot–encoded silica microspheres were synthesized according to a previously developed procedure (14). For all injections, 1 μL of solution was infused into the tumor at constant pressure (~1 μL/10 minutes) using a glass micropipette connected to a syringe pump. Images were obtained using a custom-built multiphoton laser scanning microscope (9) using a 20× 0.5 numerical aperture objective lens. Excitation was at 880 nm, with simultaneous detection of second harmonic generation (9) via a 453DF30 emission filter and GFP via a high-pass 475 dichroic and a 525DF100 emission filter. Cascade blue–conjugated dextran was visualized by exciting at 780 nm and imaging the same region (for Cascade blue–conjugated dextran and GFP), followed by image registration. Microspheres (containing 642-nm maximum emission quantum dots) were imaged with a 610DF70 emission filter. Three-dimensional image stacks containing 20 images of 5-μm thickness were obtained wherever fluorescence intensity peaked.
from the injected particles was detected. A maximum intensity z-projection of each colored stack generated a two-dimensional image. Images of consecutive adjacent regions in the x and y directions were combined into a montage, generating a single image of the entire injection site.

**Image analysis.** The pixel intensities of collagen (red pixels) and injected particles (viral vectors, green pixels; dextran, blue pixels) were spatially compared along lines drawn perpendicular to the periphery of virus containing regions. Analysis was done for one injection in each case, in five distinct image stacks, and at different depths, for a total of 20 lines measured within each tumor. The mean pixel intensities were plotted as a function of the relative distance from the observed interface with fibrillar collagen. All lines were registered such that the largest change in second-harmonic generation intensity was maintained at the origin of the graph. For quantification of viral vector distribution following injection, the entire area of viral distribution was outlined on the images. The border of the viral focus was determined as the location at which the intensity dropped to 10% of the mean intensity at the center of injection. The calculation of the area was done with imaging software (ImageJ, U.S. NIH, Bethesda, MD).6

**Flank tumor growth delay.** Mu89 cells were implanted s.c. in the flank of SCID mice and allowed to reach 100 mm³ average volume. Mice were then randomized into four groups (six to seven animals per group) and given 10 µL intratumoral injections of either PBS, 1.0 µg collagenase, 10⁶ transducing units of MGH2, or a mixture of 10⁶ transducing units of MGH2 and 1.0 µg collagenase. A second injection was done 2 days later. Tumor volume was measured every 2 to 3 days and calculated as volume = \( \pi AB^2 / 6 \), where A and B are maximum and minimum diameters, respectively. Mice died from natural progression of disease or were euthanized when (a) tumor mass exceeded a size of 2,000 mm³ or (b) premorbid behavior was noted. For tumors failing to reach 10 times the initial volume due to morbidity, the time to last measurement substituted as a conservative approximation of growth.

**Statistical analysis.** Data are expressed as mean ± SE. Statistical significance between groups was determined by unpaired Student’s t test. Statistical analysis was done with the StatView 4.51 software (SAS Institute, Inc., Cary, NC). Differences were considered statistically significant at \( P < 0.05 \).

**Results**

**Distribution of virion particles is hindered by collagen-rich regions.** To quantify virus distribution, 1 µL containing 10⁶ viral transducing units of VP16-GFP labeled nonreplicative HSV-1 virions (Gal4) was directly injected into Mu89 human melanomas grown in dorsal skin windows in SCID mice. In vivo, multiphoton imaging of viral particles with simultaneous second-harmonic generation imaging of fibrillar collagen was done ~30 minutes following injection. Viral particles distributed primarily within collagen-free areas of the tumor with limited penetration into

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similar to those of HSV vectors (Fig. 1A). Microspheres were excluded from collagen-rich tumor areas directly injected into Mu89 tumors. As with HSV particles, these spread more uniformly from the injection site with collagenase. Collagenase increased the area of viral distribution by nearly 3-fold (Fig. 1B). Collagenase-treated tumors recover from any induced hemorrhage by day 11 (C, bottom). No significant treatment response was observed in any of the tumors injected with MGH2 alone or with bacterial collagenase alone (Fig. 2A and C, bottom).

In contrast, when the same amount of oncolytic virus was coinjected with collagenase (0.2 μg/μL), the initial viral distribution was greater relative to virus alone (data not shown), translating into an improved area of tumor cell infection (Fig. 2B, middle). Therapeutic response was observed in all four collagenase cotreated tumors with a significant decrease in tumor size in two cases (Fig. 2B, bottom).

We then tested if the coinjection of collagenase and MGH2 would increase the therapeutic efficacy of MGH2 over longer time intervals. When Mu89 tumors growing in the flank of SCID mice reached 100 mm³, they were injected intratumorally with either 1 μg collagenase, 10⁶ transducing units of MGH2, or both collagenase and MGH2, followed by similar injections 2 days later. Control tumors were injected with PBS alone. The time for the tumor to reach 10 times the initial volume (mean ± SE) was compared for each group (Fig. 3). Both collagenase treatment alone (19 ± 1 days) and MGH2 injection alone (27 ± 3 days) had no significant effect on tumor growth compared with PBS control (24 ± 3 days; P > 0.05, both cases). With MGH2 treatment alone, one tumor showed marked regression but recurred after 10 days. Coinjection of MGH2 with collagenase (50 ± 8 days) significantly delayed the growth of tumors compared with all other treatment groups (P < 0.05 for all.
cases). Two of seven tumors failed to grow to 200 mm$^3$, even 60 days after treatment; a third experienced apparently complete regression although it recurred 20 days later.

**Improved efficacy is due to initial improved distribution of viral particles.** To investigate the mechanism of improved efficacy, tumors were treated as before with MGH2, either alone or with collagenase, and analyzed 2 days after the second injection. Tissue sections were stained for structural virion proteins, counterstained for nuclei (4',6-diamidino-2-phenylindole), and imaged for GFP expression using confocal microscopy to determine viral distribution. As expected, HSV antigen was present within and surrounding cells expressing GFP. In MGH2-treated tumors, virion particles and infected cells distributed in a localized fashion reminiscent of the needle track (Supplementary Fig. S1A). In contrast, for MGH2 and collagenase treatment, a diffuse distribution of infected cells was observed throughout the entire tumor section, spanning an area of up to 3 × 7 mm (Supplementary Fig. S1B). Later time points showed continued viral spread but only within collagen-free areas.

**Discussion**

The development of strategies to improve both the initial vector distribution within tumors and the ability of these vectors to propagate through the entire tumor mass is critical to the success of oncolytic viral therapy (1). Previous reports show that protease pretreatment can increase the therapeutic efficacy of a non-replicative viral vector (15). However, the mechanism of improved efficacy was undefined due to the use of nonspecific digestive enzymes, normally used to dissociate tissues, which degrade multiple extracellular matrix components. Thus, we wished to determine whether fibrillar collagen, previously shown to be a major barrier to macromolecular diffusion in tumors (7, 8), is the matrix component which limits viral distribution in certain tumors.

Our results show the important role that fibrillar collagen can play in regulating the initial distribution of viral vectors in certain fibrous tumors. Whereas smaller tracers (2 × 10$^6$ MW dextran, $R_{Ht} \sim 20$ nm, as well as immunoglobulin G, $R_{Ht} \sim 5$ nm; data not shown) distributed relatively uniformly within the tumor, the vast majority of HSV virions (150 nm in diameter) were located only in collagen-poor areas. Furthermore, silica microspheres similar in size to the viral particles, but lacking their ability to bind to cell and matrix proteins, were excluded from collagen (Supplementary Fig. S2). This suggests that the effective pore size cutoff of the collagen network is smaller than the size of these viral particles. This may have a significant effect because many tumors in humans show extensive stromal infiltration with extracellular matrix and collagen deposition (16).

Fibrillar collagen is also important in the propagation of oncolytic viral vectors through the tumor. Oncolytic vectors are thought to overcome some of the delivery issues faced by nonreplicating viral vectors through their ability to propagate on site in tumors (thereby amplifying the input dose) and spread from tumor cell to tumor cell. We found that the collagen network, in addition to restricting the initial distribution, limited the maximal spread of MGH2 infection within the tumor. Tumor cell infection remained confined to a small area and the tumor continued to grow. Coinjection of MGH2 with collagenase resulted in a broad, uniform distribution of viral particles and infected cells, with substantial tumor regression and enhanced efficacy. This dispersed distribution of virus can improve therapeutic outcome in several ways: (a) the broad initial virion distribution improves the chance that viral vectors can penetrate all regions of the tumor; (b) the occurrence of multiple infections of the same tumor cell decreases whereas the number of distinct tumor cell infections increases; and (c) once the virus replicates and lyses the cell it has infected, it has access to a greater number of previously uninfected neighboring cells. All together, these processes can lead to increased oncolytic activity as shown schematically in Fig. 4.

Researchers have developed other methods to try to overcome the limited distribution of oncolytic vectors in tumors (17). Some use multiple injections, either on successive days or with fractionation of the initial dose at multiple sites (18). However, in the absence of extracellular matrix modification, viral distribution at each individual injection site would still be limited by collagen. Indeed, a phase II trial with an oncolytic adenoviral vector showed limited improvement in efficacy even with daily injections that included fractionation (19).

We noted that intratumoral hemorrhages occurred in many tumors treated with collagenase. Whereas bleeding from collagenase treatment alone did not affect tumor growth, this phenomenon shows the complex interactions between the extracellular matrix and cells within the tumor, including both tumor and host endothelial cells. It is possible that collagenase treatment of tumors may increase the risk of metastasis although no increase in metastasis was observed in our preliminary study (Supplementary Table S1) or a previous study in which tumors were treated with direct injections of proteases (collagenase/dispaire, trypsin; ref. 15). The development of this matrix-modulating technique for clinical applications may require the use of specific matrix proteases, such as matrix metalloproteinase 8, which degrades collagen and has been shown to decrease metastasis (20).

In conclusion, we determined that even with the on-site generation of viral particles provided by the replication-competent nature of an oncolytic HSV vector, fibrillar collagen still prevents viral spread throughout the tumor in a melanoma model. Collagen

![Figure 3. Effect of collagenase on MGH2-induced tumor growth delay. Tumors were grown s.c. in the hind flank of SCID mice. When tumors reached ~100 mm$^3$, animals were divided into four groups (n = 6-7) and treated twice (day 0 and day 2) with 10 μL of PBS (green), collagenase (0.1 μg/μL; black), MGH2 (10$^6$ transducing units) in PBS (blue), or MGH2 (10$^6$ transducing units) and collagenase (0.1 μg/μL) in PBS (red). Tumor volumes were measured every 2 to 3 days. Points, mean time to reach a given volume for each group; bars, SE. The time to reach 10 times the initial volume was compared. There was no significant difference between PBS (24 ± 3 days) and either collagenase treatment alone (19 ± 1 days) or MGH2 alone (27 ± 3 days; P > 0.05 for both cases). However, MGH2 and collagenase cotreatment (50 ± 8 days) induced a significant tumor growth delay relative to all other groups (P < 0.05 for all cases).](https://cancerres.aacrjournals.org/content/canres/66/5/2512/F2.large.jpg)
network disruption increases initial vector distribution and subsequent propagation through the tumor mass, significantly improving therapeutic outcome. This result has implications for other viral particles, gene delivery strategies, and nanotechnology-based delivery systems, as all face the problem of insufficient delivery to the target cells. Furthermore, any method decreasing tumor collagen content may have similar effects. These findings suggest ways to increase the potency of gene therapy in certain types of cancer and other diseases.

References

Figure 4. A representative model of improvement in oncolytic viral distribution and tumor cell infection by collagenase treatment. Following direct intratumor injection, viral spread (red area) is limited by fibrillar collagen (red lines) and results in a cluster of infected cells (light green). The collagen network also restricts the distribution of subsequent viral progeny and tumor cell infection beyond the initial injection site is not achieved. In contrast, coinjection of virus with collagenase results in a more diffuse distribution of viral particles and a greater number of initially infected cells (light green). Viral particles released by these cells have greater access to neighboring uninfected cells. This process results in more widespread secondary infection (dark green) and ultimately greater therapeutic efficacy.

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