Use of Macrophages to Target Therapeutic Adenovirus to Human Prostate Tumors

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Abstract

New therapies are required to target hypoxic areas of tumors as these sites are highly resistant to conventional cancer therapies. Monocytes continuously extravasate from the bloodstream into tumors where they differentiate into macrophages and accumulate in hypoxic areas, thereby opening up the possibility of using these cells as vehicles to deliver gene therapy to these otherwise inaccessible sites. We describe a new cell-based method that selectively targets an oncolytic adenovirus to hypoxic areas of prostate tumors. In this approach, macrophages were cotransduced with a hypoxia-regulated E1A/B construct and an E1A-dependent oncolytic adenovirus, whose proliferation is restricted to prostate tumor cells using prostate-specific promoter elements from the TARP, PSA, and PMSA genes. When such cotransduced cells reach an area of extreme hypoxia, the E1A/B proteins are expressed, thereby activating replication of the adenovirus. The virus is subsequently released by the host macrophage and infects neighboring tumor cells. Following systemic injection into mice bearing subcutaneous or orthotopic prostate tumors, cotransduced macrophages migrated into hypoxic tumor areas, upregulated E1A protein, and released multiple copies of adenovirus. The virus then infected neighboring cells but only proliferated and was cytotoxic in prostate tumor cells, resulting in the marked inhibition of tumor growth and reduction of pulmonary metastases. This novel delivery system employs 3 levels of tumor specificity: the natural “homing” of macrophages to hypoxic tumor areas, hypoxia-induced proliferation of the therapeutic adenovirus in host macrophages, and targeted replication of oncolytic virus in prostate tumor cells. Cancer Res; 71(5): 1805–15. © 2011 AACR.

Introduction

In recent years, 2 crucial components of anticancer gene therapies have emerged—the need to both develop effective methods for delivering the “therapeutic” gene to the tumor site and restrict the expression of genes to the tumor site alone (to avoid extratumoral gene expression and concomitant side effects). Transcriptional control, exploiting promoter elements from the prostate-specific antigen (PSA) gene, have been used to confine therapeutic gene expression to prostate epithelial cells (1). The presence of multiple areas of extremely low oxygen tension (hypoxia) in tumors, including prostate carcinomas (1), has also been exploited to target therapeutic gene expression. In this case, the therapeutic gene is placed under the control of hypoxia-regulated promoter elements so it is only expressed in hypoxic cells (2–4).

An array of viral, synthetic, and cell-based vectors have been developed to transport such therapeutic gene constructs from the bloodstream into tumors but most have shown poor practical efficacy (5). Only large doses (>10^12 particles) of virus injected directly into tumors resulted in widespread infection of tumor cells. Intravenous injection is considerably less efficient, as most virus fails to penetrate the tumor and is ultimately sequestered by the liver (6, 7). New strategies are urgently required to deliver therapeutic genes to sites deep within primary and metastatic tumors.

Recently, central inflammatory cells: macrophages—and their precursors in the bloodstream, monocytes—have attracted considerable attention as gene delivery vehicles as they are continually recruited into tumors and migrate into avascular, hypoxic/necrotic areas (8–10). There, they upregulate expression of hypoxia-activated transcription factors...
(HIFs 1α and 2α), which then bind to hypoxic response elements (HRE) within the promoters of multiple tumor-promoting and adaptive genes to activate their expression (8, 9). This prompted us to propose that macrophages could be exploited to deliver HIF-regulated therapeutic genes to otherwise inaccessible areas in tumors. Initially, we showed that human macrophages adenovirally infected with a HIF-regulated reporter or a therapeutic gene and then cocultured with human tumor spheroids in vitro, migrated into the hypoxic centre of these structures and expressed the exogenous gene (11). In this study, we show for the first time that macrophages can be engineered to silently transport to, and then produce high titres of therapeutic adenovirus specifically in hypoxic areas of experimental prostate tumors. An extra level of (prostate) tumor targeting was achieved by placing either the exogenous gene in the virus and/or the replication of the virus itself under the control of promoter sequences from such prostate-specific genes (12).

Materials and Methods

Immunolabeling of hypoxia and macrophages in human prostate tumors

Consenting patients with localized prostate adenocarcinoma received 0.5 gm/m² pimonidazole (PIMO) i.v. 16–24 hours before radical prostatectomy (Mount Vernon Hospital, London). Serial sections (4 μmol/L) were immunostained with CD68 (1:100, Dako) and anti-PIMO (Hypoxprobe Store) and visualized with DAB or Vector Red (Vector Laboratories), respectively. To determine the distribution of tumor-associated macrophage (TAM) relative to blood vessels, 30 cases of adenocarcinoma of the prostate were selected at random from the 1997–1999 archives (Royal Hallamshire Hospital, Sheffield, UK). Sections were double-immunostained with Factor VIII (FVIII; Dako, UK) and CD68 and visualized using Vector Red. Sections were scanned at low power (x100–x100) to identify areas of macrophage and vascular density—the “hot-spots.” The mean densities of macrophages (M/μl) and blood vessels (MVD) within macrophage and vascular hotspots were determined using the Chalkley point array method (7).

Cell lines

HEK 293, A549 (human lung carcinoma), and LNCaP (human prostate carcinoma) were purchased from the ECACC. HEK 293s are purchased yearly to obtain low passage cells for viral transduction. Tumor cell lines are routinely tested for authenticity by microsatellite genotyping at the ECACC.

Isolation of human monocytes and generation of monocyte-derived macrophages

Macrophages were prepared from mononuclear cells isolated fromuffy coats (Blood Transfusion Service; ref. 9).

Plasmid construction and adenoviruses

The techniques for constructing HRE-regulated E1A/B plasmids and adenoviruses (CMV-AdV5-GFP (driven by a CMV promoter) and PSA-AdV5-GFP (driven by the prostate-specific promoter, PSA), are outlined in the supplemental methods. Optimal transduction of HEK 293 cells with adenovirus (CMV-AdV5-GFP) was achieved with an MOI of 5 PFU/cell as measured by flow cytometry for GFP expression (Supplementary Fig. 1).

Cotransduction of primary monocyte-derived macrophages

To prevent undesirable viral recombination events, the HRE-regulated E1A/B gene constructs were transferred into macrophages by plasmid transfection rather than coinfection with a second viral vector. Monocyte-derived macrophages (MDM) that had been cultured for 3 days were infected with an MOI of 100 PFU/cell. For cotransduction, macrophages (2 × 10⁶) were incubated overnight with the virus and then transduced with 5 μg pcDNA3.1(+)–HRE-E1A/B (HRE-E1A/B) construct or the pcDNA3.1(+)–No-HRE-E1A/B (No-HRE-E1A/B) using the Amaxa Macrophage Nucleofection Kit (Amaxa Biosystems).

Protein detection by 1-D SDS-PAGE

SDS page was carried out on MDMs lysed in SDS buffer all techniques and antibodies used were described by us recently (13).

Complementation system

Clariﬁed supernatants from cotransduced MDM cultured in normoxic or hypoxic conditions were applied to 293 cells at increasing dilutions. Plaque formation and GFP development was monitored daily in an end-point dilution assay. Viral particles (vp) were calculated by measuring the optical density (OD) at 260 nm after particle lysis in PBS/0.5% SDS for 15 minutes at 37°C. Titres were calculated using OD260 × 1.1 × 10¹³ vp/mL (14, 15).

Infiltration of primary human macrophages into tumor spheroids in vitro

Tumor spheroids were generated using LNCaP and A549 cells by seeding 5 × 10⁶ cells into each well of a well of a 2% agarose-coated 96-well tissue cultureplate. After 7–10 days, each well contained a 700–800 μm spheroid, to which 3 × 10⁶ cotransduced macrophages were added. These experiments were then repeated in the presence of antienoviral antibodies obtained from pooled human serum (as outlined in detail in the supplemental methods).

Macrophage trafficking to hypoxic prostate tumor xenografts

Male CD1 athymic mice (aged 6–8 weeks; Harlan laboratories) were injected subcutaneously with 2 × 10⁶ human prostate cancer PC3 cells. Mice were injected with 3 × 10⁶ either; cotransduced (CMV-AdV5-GFP or PSA-AdV5-GFP and HRE-E1A/B plasmid) macrophages, singly transduced (CMV-AdV5-GFP) macrophages, untransduced macrophages, or PBS alone through the tail vein. After 48 hours, mice were injected with 60 mg/kg PIMO for 1 hour before sacrifice. For flow cytometry tumor, liver, lung, kidney, and spleen were made into cell suspensions and incubated with phycoerythrin (PE)-conjugated anti-CD14 (Serotec Ltd.). For microscopic studies, 7 μmol/L frozen sections were stained with human anti-CD68,

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rabbit anti-PIMO, and rabbit anti-GFP (1:100), PE-anti-mouse 
(1:250; Dako), Alexa-647-anti-rabbit (1:250), and Alexa-488-
anti-rabbit (1:250) secondary antibodies, were incubated 
tumor sections for 30 minutes and then DAPI (3nmol/L) 
stained. Prolong Gold Anti-fade used to preserve fluorescence 
(all reagents from Molecular Probes Inc). Images were cap-
tured using a Zeiss LSM 510 confocal microscope (magnifica-
tion x400).

For the therapeutic studies, the replication competent 
adenovirus Ad[I/PPT-E1A] replaced the CMV-AdV5-GFP 
and we used the LNCaP xenograft model (both subcutaneous 
and orthotopic) as Ad[I/PPT-E1A] has been shown to be 
highly lytic in this cell line (16). Macrophage trafficking studies 
conducted in the presence of human serum are detailed in the 
Supplementary Methods.

(i) Subcutaneous model

LNCaPs were mixed 1:1 with matrigel (BD Biosciences) and 2 \times 10^6 injected subcutaneously into the hind flank region. When the tumors reached 4 mm in diameter, mice were injected via tail vein with 100 \muL PBS containing either 3 million singly transduced human MDMs (infected with the oncolytic virus Ad[I/PPT-E1A] at MOI 100), or cotransduced macrophages (Ad[I/PPT-E1A] at MOI 100 and HRE-E1A/B), 5 \times 10^10 Ad[I/PPT-E1A] only or PBS alone. Tissues including 
tumors, kidney, liver, lungs, and spleen, were paraffin 
 wax embedded. Human and mouse macrophage popu-
lations were identified in sections by staining with 
 antibodies to human CD68, and murine F4/80 (AbD 
Serotec) all sections were H&E stained.

(ii) Orthotopic model

One million cells were mixed 1:1 in Matrigel and 
injected into the dorsolateral prostate. Tumor take was 
monitored by ultrasound using Vevo 770 High resolu-
tion imaging platform (Visual Sonics) and mice were 
treated as above. Thirty days after treatment, animals 
were sacrificed and tumors/organisms were collected, 
sectioned, and stained by H&E. Whole tumor sections 
were scanned on an Aperio slide scanner (Aperio slide 
scanner) to facilitate tumor diameter measurements.

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Macrophages are present in hypoxic areas of human prostate carcinoma. Sections were dual stained for CD68/FVIII to identify the areas of greatest intratumoral macrophage and vascular density—the "hotspots." Mean MVD and mean M/D were assessed microscopically using a 25-point Chalkley array eyepiece graticule (x250 magnification; field area 0.189mm²). Representative bright-field images of CD68 positive macrophages (brown; arrows) costained (red) for pimonidazole. A, a nonhypoxic tumor area. B, a hypoxic (PIMO positive) area. Bar, 50 \mu m. C, M/D were highest in areas of 
 poor vascularity (P < 0.001; i), whereas MVD was significantly (P < 0.001) higher in areas of low macrophage counts (ii).
MTS assay
Singly transduced (Ad[I/PPT-E1A]) or cotransduced macrophages (Ad[I/PPT-E1A] and HRE-E1A/B) were incubated for 16 hours at 37°C in humidified Heto multigas incubators (0.5% oxygen). Medium was collected and centrifuged at 10,000 g for 5 minutes to remove cell debris, and then applied to LNCaPs at different dilutions (Neat, $10^2$, $10^3$, $10^4$, $10^5$). LNCaPs were also transduced with Ad[I/PPT-E1A] (50 pfu/cell) as a positive control. Cell viability was analyzed on day 6 after transduction using the MTS Cell Titer 96 Aqueous One Solution Cell Proliferation Assay (Promega).

In vitrō tubule formation assay
Conditioned medium was collected from LNCaP spheroids or spheroids infiltrated with untreated MDMs or Ad[I/PPT-E1A] cotransduced MDMs after 48 hours of culture. Serum starved human umbilical vein endothelial cells (HUVEC; 8000 cell/well) were resuspended in conditioned medium and seeded onto growth factor-reduced Matrigel (BD Biosciences)-coated wells. After 12 hours, tubule formation was measured using ImageJ software (National Institute of Health).

Statistical analysis
Statistical comparison was done using the GraphPad prism program (GraphPad software). Nonparametric statistical tests were used to analyze the data and the logrank test was used to compare tumor survival curves using Kaplan–Meier. A value of $P = 0.05$ was the limit of statistical significance.

Results
TAMs accumulate in hypoxic areas of human prostate carcinoma
TAMs can be detected in both well-oxygenated (Fig. 1A) and hypoxic (Fig. 1B) areas of human prostate tumors. Poorly vascularised (hypoxic) areas contained higher numbers of TAMs (Fig. 1Ci) than highly vascularised ones (Fig. 1Cii).
E1A complementation and replication of adenovirus in macrophages under hypoxic conditions

The E1A complementation system in cotransduced MDMs in the presence of tumor hypoxia is illustrated in Fig. 2A. To test this, MDMs were infected with the adenovirus vector CMV-Ad5-GFP at increasing MOIs (0–100). The efficiency of infection (i.e., GFP expression with this vector) was routinely found to be 75 ± 9.5% (n = 5) and reached its peak after 72 hours of infection with minimal cytotoxicity compared with uninfected macrophages (Supplementary Fig. 1A and B).

Cotransduced macrophages, exposed to hypoxia (0.1% O2) for 16 hours, upregulated HIF-1α, HIF-2α, and (hypoxia regulated) E1A proteins compared with normoxic conditions (Fig. 2B). Furthermore, when supernatants from cotransduced MDMs were titrated in HEK 293 cells, only supernatants from hypoxic MDMs (Fig. 2Ci) resulted in production of detectable infectious virus. Average yields of 5001 ± 1495 PFU/infected macrophage (n = 5) were confirmed in the plaque assays, by both limiting dilution (Fig. 2Ci) and flow cytometry even when supernatants were diluted to 1 in 1 × 10^6 (Fig. 2D). No infectious adenovirus was detected under normoxia (Fig. 2Cii and D) or in cells transfected with the No-HRE-E1A/B construct (Fig. 2C i, ii and D) indicating that the hypoxic trigger for viral replication was tightly regulated.

Hypoxic activation of adenovirus production by macrophages infiltrating prostate tumor spheroids in vitro

Inner areas of tumor spheroids are severely hypoxic due to limited diffusion of oxygen from the surrounding culture fluid (11, 17). Lung (A549) and prostate (LNCaP) tumor spheroids were used to assess the prostate specificity of the macrophage-based viral delivery system. The inner, perinecrotic regions of both sets of spheroids were confirmed as hypoxic (Fig. 3A). MDMs cotransduced with the HRE-E1A/B plasmid and a CMV-AdV5-GFP were seen to infiltrate both LNCaP and A549 spheroids equally (Fig. 3B). Cotransduced MDMs released sufficient adenovirus to cause widespread CMV-driven GFP expression throughout A549 and LNCaP spheroids (normoxic and hypoxic areas) within 72 hours of infiltration (Fig. 3C).

When MDMs cotransduced with an adenovirus where GFP expression was regulated by the PSA promoter (PSA-AdV5-GFP) and HRE-E1A/B infiltrated LNCaP and A549 spheroids widespread expression of PSA-driven GFP was only seen LNCaPs (Fig. 3D and E).

Macrophage delivery of adenovirus to hypoxic areas of human prostate tumors

(i) Delivery of a GFP-expressing adenovirus to prostate tumor xenografts in vivo.

Initially, we confirmed the relevance of our prostate tumor xenograft models to human prostate tumors by showing a similar level of hypoxia in prostate xenografts and primary human prostate carcinomas (Supplementary Fig. 2S).

We then cotransduced MDMs with CMV-Ad5-GFP and HRE-E1A/B and administered these via tail vein injection to mice bearing subcutaneous human prostate (PC3) tumors. Control mice received PBS alone, untransduced MDMs, or singly transduced MDMs (no HRE-E1A/B). Triple immunofluorescence microscopy on frozen human tumors labeled with antibodies against PIMO (white), human CD68 (red), and GFP (green) revealed that human MDMs had migrated into hypoxic/perinecrotic areas of the tumor (Fig. 4A). No human CD68+ cells were detected in tumors from mice injected with PBS alone and no GFP was detected in mice injected with untransduced MDMs (data not shown). In mice...
receiving singly transduced MDMs, GFP expression was largely restricted to CD68+ human macrophages (Fig. 4A, top). This contrasted with the abundance of GFP+ cells in hypoxic epithelial cells and macrophages in tumors treated with cotransduced MDMs (Fig. 4A, bottom). Therefore, hypoxia present in prostate tumor xenografts triggered viral replication within MDMs and viral spread from MDMs resulted in the infection of neighboring tumor cells.

In contrast to the hypoxic tumors, the frequency of human MDMs in the liver and lung was low (<1 cell/field/view in a total of 10 fields) and no MDMs were detected in the kidney or spleen (data not shown). These immunohistochemical data were supported by flow cytometry. Similar levels of human MDM infiltration into tumors (2%–5%) were seen in all mice injected with MDMs (Fig. 4B). This equates to approximately 442,000 ± 25,000 human MDMs per LNCaP xenografts (i.e., 2.21% of the 3 × 10⁶ MDMs injected), as determined using the regression equation described by Meyskins et al. (1984; ref. 18) [No. of cells/tumor = 2.40 (tumor diameter)².378/(cell diameter)².804]. In contrast, mice injected with PBS showed negligible levels of GFP (1.5% ± 0.4%); whereas in mice injected with singly transduced MDMs, this increased to 5.6% ± 1.4% of cells. However, in tumors from mice injected with cotransduced MDMs GFP expression was significantly higher (21% ± 5.4 of cells; Fig. 4B, right). These data were reproduced when mice bearing LNCaP xenografts were injected with macrophages cotransduced with Ad5-PSA-GFP and HRE-E1AB. The presence of human macrophages and adenovirus in these tumors was confirmed using antibodies to human CD68 and E1A (data not shown).

(ii) Use of macrophages to target oncolytic adenovirus, Ad[I/PPT-E1A] to prostate tumors

(a) Immunophenotype of MDMs cotransduced to express the Ad[I/PPT-E1A].

The oncolytic adenovirus, Ad[I/PPT-E1A], specifically replicates in prostate tumor cells (12, 16), as E1A gene expression is controlled by a recombinant regulatory sequence designated PPT, consisting of a PSA enhancer, PSMA enhancer, and a T-cell receptor γ-chain alternate reading frame protein promoter (12). However, the therapeutic efficacy of Ad[I/PPT-E1A] has only been seen following intratumoral administration into mice bearing prostate tumors (12, 16, 19). Systemic delivery of such adenoviruses is limited by liver sequestration, complement, pre-immune IgM, and neutralizing antibodies (20).
The impact of cotransduction on the phenotype of macrophages was investigated before injection in vivo. Flow cytometric analysis of cotransduced MDMs revealed no significant alteration in the expression of CD11b, CD14, CD68, or classic anti-inflammatory markers, such as CD36, MHC-class II, and mannose receptor when compared with untreated MDMs or singly transduced MDMs (data not shown). However, an increase in cell surface expression of toll-like receptors (TLR) 2 and 4 was evident (Supplementary Fig. 3A). TLRs mediate adenoviral regulation and transgene expression in human peripheral blood mononuclear cells following infection, activating their expression of proinflammatory cytokines (21, 22). This was confirmed when cotransduced MDMs were found to release elevated levels of various proinflammatory cytokines (Supplementary Fig. 3B).

(b) Effect of MDMs cotransduced with HRE-E1A/B and Ad[I/PPT-E1A] on tumor spheroids.

Infiltration of LNCaP spheroids by cotransduced MDMs triggered sufficient adenoviral production to cause widespread cell death after 48 hours (Fig. 5A and B). The ability of hypoxic, cotransduced MDMs to kill monolayers of LNCaPs was confirmed using the MTS assay (Fig. 5C). No cell death was evident for monolayers of A549 cells infected in parallel MTS assays (data not shown).

As macrophages in tumors can be proangiogenic (9), we investigated whether this might be the case with our cotransduced MDMs, by taking media conditioned by LNCaP spheroids alone or following infiltration by cotransduced MDMs and adding this to HUVEC. Cotransduced MDMs failed to induce tubule formation (Supplementary Fig. 4A and B). Interestingly, the presence of cotransduced MDMs significantly (P = 0.035) reduced rather than increased the release of the proangiogenic cytokine, VEGF, by LNCaP spheroids. It also increased release of the proinflammatory cytokines TNFα, IL-1β, IL-6, and IL-8 (Supplementary Fig. 4C) while leaving IL-10, IL-12p70, or GMCSF levels in the media unaffected (data not shown).

Figure 5. MDMs cotransduced to express Ad[I/PPT-E1A] efficiently lyse tumor cells in LNCaP spheroids in vitro. Human prostate tumor (LNCaP) spheroids were cocultured with 3 x 10⁴ MDMs 24 hours after being cotransduced with Ad[I/PPT-E1A] and HRE-E1A/B. Parallel sets of spheroids were also infiltrated with singly transduced MDMs or Ad[I/PPT-E1A] (50 pfu/cell). A, representative dot plots of enzymatically dispersed spheroids. B, light microscopy images of whole, individual spheroids. C, loss of LNCaP tumor cell viability was also confirmed using the MTS assay. The data are averages for triplicate samples. Bar (B), 200 μm.
Effect of MDMs cotransduced with HRE-E1A/B and Ad[I/IPPT-E1A] on subcutaneous and orthotopic LNCaP xenografts in vivo.

Mice bearing subcutaneous LNCaP xenografts were injected with 100 μL PBS containing either 3 × 10^6 singly transduced MDMs, cotransduced MDMs, or 5 × 10^10 Ad[I/IPPT-E1A] particles. As early as day 5 after treatment initiation, a single injection of cotransduced MDMs resulted in 50% reduction in tumor size, and by day 15, 2 mice were tumor-free. Even after 50 days, and a single treatment, the 3 remaining mice in the cotransduced macrophage group had tumors less than 20% of their original volume (Fig. 6A), whereas a single injection of Ad[I/IPPT-E1A] alone initially delayed tumor growth for up to 30 days, but tumors then regrew and were 50% greater than their starting volume. Inhibition of tumor growth between these 2 groups was significant (P = 0.001). By contrast, mice that received PBS or singly transduced MDMs had to be killed early due to large tumors.

Survival data are summarized in Fig. 6A (right). Tumor-bearing mice that received cotransduced MDMs survived longer than the other treatment groups. Fig. 6B shows representative images of tumors from each group of mice. Tumors from mice receiving cotransduced MDMs were necrotic and contained few viable cells (Fig. 6B). Interestingly, the tumors from mice treated with Ad[I/IPPT-E1A] or cotransduced MDM were considerably paler and less vascularized (Fig. 6B). In the case of Ad[I/IPPT-E1A] injected mice,

Figure 6. Systemic delivery of MDMs cotransduced with HRE-E1A/B plasmid and Ad[I/IPPT-E1A] markedly slows the growth of both subcutaneous and orthotopic LNCaP xenografts. A and B, subcutaneous LNCaP tumor model. A, only injections of cotransduced MDMs and Ad[I/IPPT-E1A] significantly reduced tumor volume compared with PBS treatment (P = 0.003 and P = 0.01, respectively) and singly transduced MDM (P = 0.0001 and P = 0.0012, respectively). Survival data for all 4 groups were plotted as a Kaplan–Meier survival curve (right). B, photographs of the tumors following treatment (top). Representative (x 20) images for necrosis (middle) and murine macrophage infiltration (F480; bottom) are presented. C, orthotopic LNCaP tumor model. Representative images of H&E tumor sections and analysis of tumor volumes (bottom). Data shown are means ± SEMs.
this could have resulted from the reduction in blood vessels on the under surface of tumor and/or the marked leukocyte infiltration that reportedly occurs with oncolytic adenoviruses (20). Indeed, tumors in the Ad[I/PPT-E1A] injected group had significantly more murine F4/80+ cells (64±13%) compared with the other groups (Fig. 6B). By contrast, mice receiving cotransduced MDMs had low F4/80+ cell infiltration—most likely because these tumors were very small and mainly necrotic/acellular. This implies that the anti-angiogenic effect of cotransduced MDMs seen in vitro also occurred in LNCaP tumors in vivo.

Tumor growth in mice bearing orthotopic LNCaPs was significantly inhibited when mice were injected with Ad[I/PPT-E1A] alone or cotransduced macrophages (Fig. 6C), the latter having a more marked effect. Significantly more F4/80+ TAMs (71%±17) were detected in mice injected with Ad[I/PPT-E1A] than those injected with PBS alone, singly transduced MDMs or cotransduced MDMs (data not shown). In both tumor models considerably more necrosis was detected in tumors from mice injected with cotransduced MDMs (Fig. 7A and B). Notably, although lung metastases were visible in mice in control groups (Fig. 7A and B ii, iii), these were absent in the mice treated with cotransduced MDMs.

(iii) Macrophages protect adenovirus from adenoviral neutralizing antibodies in human sera.

In humans, preexisting humoral immunity to adenovirus can interfere with the systemic delivery of adenovirally based therapies. We hypothesized that our macrophage delivery system would protect adenoviruses against these preexisting antianimal neutralizing antibodies (NAb). Indeed, the presence of high titer NAb in cultures of cotransduced MDMs (CMV-AdV5-GFP and HRE-E1A/B) did not prevent...
viruses in various tumor models have shown a heterogeneous viral copies by each hypoxic macrophage). virus (due to the amplification and release of thousands of macrophages resulted in widespread dissemination of the grafts with relatively small numbers of cotransduced human Interestingly, infiltration of tumor spheroids or tumor xenografts and in clinical trials but these had to be injected directly into primary tumors and at very high viral titres of adenovirus. Moreover, this approach fails to target metastatic tumors growing at distant sites (23–26).

We then used macrophages to systemically deliver an oncolytic adenovirus in which HIFs regulate gene expression have been used to target hypoxic tumor cells in human tumor xenografts and in clinical trials but these had to be injected directly into primary tumors and at very high viral titres of adenovirus. Moreover, this approach fails to target metastatic tumors growing at distant sites (23–26).

Our finding that macrophages accumulate in hypoxic areas of human prostate tumors and human prostate tumor xenografts prompted us to develop a means of using these cells to deliver therapeutic adenovirus to these sites and via a systemic route permitting the targeting of primary tumors and their metastases. As hypoxia also exists in diseased tissues other than tumors (27) and mild hypoxia can exist in healthy tissues (28), we added a further degree of tumor targeting by placing the exogenous gene (e.g., GFP) in the virus or the further replication of the virus itself when released by macrophages—under the control of prostate-specific promoters.

We found hypoxia-induced replication and release of adenovirus by infiltrating cotransduced macrophages resulted in the widespread infection of tumor cells and expression of the PSA-driven GFP reporter gene construct in the adenovirus. Interestingly, infiltration of tumor spheroids or tumor xenografts with relatively small numbers of cotransduced human macrophages resulted in widespread dissemination of the virus (due to the amplification and release of thousands of viral copies by each hypoxic macrophage).

We then used macrophages to systemically deliver an oncolytic adenovirus (Ad[I/PPT-E1A]). This virus previously has shown therapeutic potential but only when injected directly into prostate tumors (16, 19). Studies using different oncolytic viruses in various tumor models have shown a heterogeneous and incomplete dissemination of virus (29–31) as adenoviruses interact with human blood cells following systemic delivery. This could be responsible for reducing the therapeutic effect (32). The compromised vascular supply within necrotic areas, the distorted functional properties of tumor vessels, and the elevated tumor interstitial fluid pressure also contribute to an unequal viral distribution within tumors (30).

Mice bearing both subcutaneous and orthotopic tumors exhibited remarkable tumor regression following a single injection of cotransduced MDMs and showed no signs of regrowth. Viral delivery by macrophages therefore resulted in a lasting antitumor effect with negligible metastatic frequency. Moreover, we required at least 1,000-fold fewer adenoviral particles in this system, compared with animals administered with virus alone. We have therefore exceeded previous benchmarks by greatly reducing the amount of virus tumor-bearing mice receive, circumventing the need for systemic intratumoral administration. We are aware however, that our in vivo studies were carried out in immunocompromised mice. This choice of animal model was because human Ad replication is known to be negligible in rodent cells so syngeneic murine tumor models could not be used (33, 34). Although host immunity against such cotransduced, “therapeutic” autologous macrophages may be activated in a fully immunocompetent host, the fact that the adenovirus is in a latent phase during trafficking of macrophages to hypoxic tumor sites, suggests that expression of viral antigens is likely to be extremely low. The oncolytic adenovirus used in this study has been tested extensively against numerous human tissues as part of a clinical assessment of its safety in possible clinical trials. Even at high multiplicities of infection this virus showed no ability to replicate in primary cells from such healthy human tissues as endothelium, urothelium, lungs, liver etc. (Maitland NJ and coworkers unpublished observations). Moreover, neutralizing antibodies from human sera administered to human prostate tumor-bearing mice, had no impact on the activity of this oncolytic virus when delivered systemically via our macrophage system suggesting that patients with preexisting adenovirus immunity would be suitable candidates for i.v. administration of such viruses (35). The application of a cell-mediated viral delivery system like the one described in this study may also prevent adenovirus neutralization from taking place by offering protection to the virus from humoral immunity (36, 37).

In summary, we describe a novel system whereby the natural ability of macrophages to harbor and support viral replication was used as “silent carriers” to trigger oncolytic viral production in hypoxic areas of tumors, resulting in intratumoral spread and a lasting therapeutic effect.

Discussion

Adenoviruses in which HIFs regulate gene expression have been used to target hypoxic tumor cells in human tumor xenografts and in clinical trials but these had to be injected directly into primary tumors and at very high viral titres of adenovirus. Moreover, this approach fails to target metastatic tumors growing at distant sites (23–26).

Our finding that macrophages accumulate in hypoxic areas of human prostate tumors and human prostate tumor xenografts prompted us to develop a means of using these cells to deliver therapeutic adenovirus to these sites and via a systemic route permitting the targeting of primary tumors and their metastases. As hypoxia also exists in diseased tissues other than tumors (27) and mild hypoxia can exist in healthy tissues (28), we added a further degree of tumor targeting by placing the exogenous gene (e.g., GFP) in the virus or the further replication of the virus itself when released by macrophages—under the control of prostate-specific promoters.

We found hypoxia-induced replication and release of adenovirus by infiltrating cotransduced macrophages resulted in the widespread infection of tumor cells and expression of the PSA-driven GFP reporter gene construct in the adenovirus. Interestingly, infiltration of tumor spheroids or tumor xenografts with relatively small numbers of cotransduced human macrophages resulted in widespread dissemination of the virus (due to the amplification and release of thousands of viral copies by each hypoxic macrophage).

We then used macrophages to systemically deliver an oncolytic adenovirus (Ad[I/PPT-E1A]). This virus previously has shown therapeutic potential but only when injected directly into prostate tumors (16, 19). Studies using different oncolytic viruses in various tumor models have shown a heterogeneous and incomplete dissemination of virus (29–31) as adenoviruses interact with human blood cells following systemic delivery. This could be responsible for reducing the therapeutic effect (32). The compromised vascular supply within necrotic areas, the distorted functional properties of tumor vessels, and the elevated tumor interstitial fluid pressure also contribute to an unequal viral distribution within tumors (30).

Mice bearing both subcutaneous and orthotopic tumors exhibited remarkable tumor regression following a single injection of cotransduced MDMs and showed no signs of regrowth. Viral delivery by macrophages therefore resulted in a lasting antitumor effect with negligible metastatic frequency. Moreover, we required at least 1,000-fold fewer adenoviral particles in this system, compared with animals administered with virus alone. We have therefore exceeded previous benchmarks by greatly reducing the amount of virus tumor-bearing mice receive, circumventing the need for systemic intratumoral administration. We are aware however, that our in vivo studies were carried out in immunocompromised mice. This choice of animal model was because human Ad replication is known to be negligible in rodent cells so syngeneic murine tumor models could not be used (33, 34). Although host immunity against such cotransduced, “therapeutic” autologous macrophages may be activated in a fully immunocompetent host, the fact that the adenovirus is in a latent phase during trafficking of macrophages to hypoxic tumor sites, suggests that expression of viral antigens is likely to be extremely low. The oncolytic adenovirus used in this study has been tested extensively against numerous human tissues as part of a clinical assessment of its safety in possible clinical trials. Even at high multiplicities of infection this virus showed no ability to replicate in primary cells from such healthy human tissues as endothelium, urothelium, lungs, liver etc. (Maitland NJ and coworkers unpublished observations). Moreover, neutralizing antibodies from human sera administered to human prostate tumor-bearing mice, had no impact on the activity of this oncolytic virus when delivered systemically via our macrophage system suggesting that patients with preexisting adenovirus immunity would be suitable candidates for i.v. administration of such viruses (35). The application of a cell-mediated viral delivery system like the one described in this study may also prevent adenovirus neutralization from taking place by offering protection to the virus from humoral immunity (36, 37).

In summary, we describe a novel system whereby the natural ability of macrophages to harbor and support viral replication was used as “silent carriers” to trigger oncolytic viral production in hypoxic areas of tumors, resulting in intratumoral spread and a lasting therapeutic effect.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

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