Adenoviral-mediated imaging of gene transfer using a somatostatin receptor-cytosine deaminase fusion protein

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Suicide gene therapy is a process by which cells are administered a gene that encodes a protein capable of converting a nontoxic prodrug into an active toxin. Cytosine deaminase (CD) has been widely investigated as a means of suicide gene therapy owing to the enzyme’s ability to convert the prodrug 5-fluorocytosine (5-FC) into the toxic compound 5-fluorouracil (5-FU). However, the extent of gene transfer is a limiting factor in predicting therapeutic outcome. The ability to monitor gene transfer, non-invasively, would strengthen the efficiency of therapy. In this regard, we have constructed and evaluated a replication-deficient adenovirus (Ad) containing the human somatostatin receptor subtype 2 (SSTR2) fused with a C-terminal yeast CD gene for the non-invasive monitoring of gene transfer and therapy. The resulting Ad (AdSSTR2-yCD) was evaluated in vitro in breast cancer cells to determine the function of the fusion protein. These studies demonstrated that both the SSTR2 and yCD were functional in binding assays, conversion assays and cytotoxicity assays. In vivo studies similarly demonstrated the functionality using conversion assays, biodistribution studies and small animal positron-emission tomography (PET) imaging studies. In conclusion, the fusion protein has been validated as useful for the non-invasive imaging of yCD expression and will be evaluated in the future for monitoring yCD-based therapy.

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INTRODUCTION

In the past two decades, gene therapy has been developed as a promising approach to combat a variety of diseases. Over this time period, more than 1700 clinical gene therapy trials were conducted, with 65% addressing cancer (The Journal of Gene Medicine website, www.wiley.com/legacy/wileychi/genmed/clinical). Thus far, adenoviral vectors have been used in 24% of clinical trials, followed by retroviral vectors (21%) and naked/plasmid DNA (19%). Adenoviral vector-based approaches for cancer gene therapy have included: mutation compensation, genetic immunopotentiation and molecular chemotherapy. For mutation compensation, the molecular lesions in the cancer cell that are responsible for malignant transformation are corrected through gene transfer, whereas for genetic immunopotentiation, active immunization against tumor associated antigens is attempted. Molecular chemotherapy (also termed gene-directed enzyme prodrug therapy or suicide gene therapy) is an approach for generating higher tumor concentrations of a cytotoxic drug than can normally be achieved through standard chemotherapy. There are many genes that can be used for this approach, but two of the most common for this strategy are those encoding the herpes simplex virus type-1 thymidine kinase (HSV1-TK) and cytosine deaminase (CD). CD is an enzyme that can convert 5-fluorocytosine (5-FC) into the chemotherapy agent, 5-fluorouracil (5-FU). In this study, we chose to work with the CD system because, unlike with HSV1-TK, 5-FU is able to diffuse across the cell membrane into neighboring cells without the need for gap junctions, which results in a more powerful bystander effect.

Adenoviral-mediated delivery of CD has demonstrated success in animal models for treating several types of cancer including glioblastoma multiforme, colon, prostate and breast either alone or in combination with radiation therapy. In this regard, both the bacterial CD (bCD) and yeast CD (yCD) enzymes have been utilized. The yCD has been shown to have a K_m that is 22-fold lower than bCD for 5-FC, but is also more thermolabile than bCD, which may impact gene therapy strategies. Subsequent engineering of both bCD and yCD have increased their efficacy by increasing their efficiency for 5-FC conversion and their thermostability, respectively. One issue that has limited the efficacy of this approach is the inability to determine the efficiency of CD gene transfer using non-invasive methods. In this regard, we have previously utilized the human somatostatin receptor subtype 2 (SSTR2) for imaging of gene transfer using gamma camera imaging and positron-emission tomographic (PET) imaging. Somatostatin receptors are members of the G protein-coupled receptor family that have seven transmembrane domains consisting of three extracellular loops, termed E1, E2 and E3. SSTR2 is expressed on brain, spleen and kidney tissues and overexpressed in a variety of tumor types, leading to the development of tumor imaging agents. Therefore, we chose to develop a novel fusion protein consisting of yCD and SSTR2 that can be used to monitor the location, magnitude and change in magnitude over time of CD using PET imaging.

In this study, we describe the construction and characterization of a novel yCD fusion protein that consists of yCD linked to the C-terminus of the human somatostatin receptor subtype 2 (SSTR2).
to create SSTR2-yCD. This initial study uses a non-replicative adenoviral vector as a model gene delivery vehicle to evaluate SSTR2-yCD in vitro and in vivo. An adenovirus containing the SSTR2-yCD was produced and evaluated for SSTR2 function using a radioligand binding assay and for yCD function using cytotoxicity and 5-FC conversion assays in infected breast cancer cells. In vivo assays in human breast cancer xenografts injected with AdSSTR2-yCD showed conversion of 5-FC to 5-FU as well as specific uptake of a PET ligand in biodistribution and small animal PET imaging. These results demonstrated that both the SSTR2 and yCD components of the fusion protein are functional and that this imaging approach should be useful for improving the efficacy of yCD-based therapy.

MATERIALS AND METHODS

Cell lines and construction of AdSSTR2-yCD

Human breast adenocarcinoma MCF-7 and T-47D cells were obtained from the American Type Culture Collection (ATCC, Manassas, VA, USA), and both cell lines were maintained in DMEM (Life Technologies, Grand Island, NY, USA) plus 10% FBS (Hyclone, Hanford, VA, USA) and 1% heat-inactivated fetal bovine serum (Sigma Aldrich, St Louis, MO, USA). The yCD-coding sequence was fused to the C-terminal end of human SSTR2 via a seven amino acid linker in an overlap extension PCR. The resulting product was cloned into pShuttle-CMV (Agilent Technologies, Santa Clara, CA, USA) via Xhol and EcoRV sites for which were incorporated into the construct via two primers for the PCR, and the resulting plasmid was pS-SSTR2-yCD. The clone was sequence-verified to show the absence of mutations. Both the SSTR2 and yCD portions of the fusion were then validated in functional assays in transiently transfected MCF-7 and T-47D cells. Following the validation, pS-SSTR2-yCD was used to produce recombinant adenoviral plasmid, which was then transfected into AD-293 cells (ATCC) for the production of crude SSTR2-yCD adenovirus, which was sent to Q-Biogene (Montreal, Quebec, Canada) for the production of purified AdSSTR2-yCD. AdSSTR2 was produced as previously described.21

Competitive binding assay

The $B_{\text{max}}$ values of AdSSTR2- and AdSSTR2-yCD-infected T-47D and MCF-7 cells were determined by using a competitive binding assay with $^{125}$I-Tyr-1-Somatostatin-14 (PerkinElmer, Boston, MA, USA). AdSSTR2 or AdSSTR2-yCD was administered to cells at a concentration of 10 or 10 plaque-forming units (pfu) per cell, and membrane preparations were made as previously described.22 Protein concentrations were determined using the Pierce Non-Reducting Agent Compatible Kit. For each reaction, 4 μl of 0.1 M NH$_4$OAc (pH = 8.0) and 37 MBq of $^{64}$CuCl$_2$ in 0.1 N HCl was diluted in binding buffer (10 μM HEPES, 1 mM EDTA, 5 mM MgCl$_2$, 0.1% bovine serum albumin, 5 μg ml$^{-1}$ aprogin, 200 mg ml$^{-1}$ bacitracin, 10 mg ml$^{-1}$ leupeptin, 10 mg ml$^{-1}$ pepstatin) to obtain a concentration of 25 μg per 100 μl. A 96-well Multiscreen Durapore filtration plate (Millipore, Bedford, MA, USA) pretreated with 0.1% polyethyleneimine via vacuum manifold aspiration was then washed with 300 μl of wash buffer (10 μM HEPES, 1 mM EDTA, 5 mM MgCl$_2$, 0.1% bovine serum albumin) before adding 100 μl of each membrane preparation in triplicate per concentration of blocking reagent. The wells were then washed three times with wash buffer. Various concentrations of $^{125}$I-Tyr-1-Somatostatin-14 (Bachem, Torrance, CA, USA) blocking reagent, ranging from 0.01 to 55 nM, were then added to the wells in triplicate in a volume of 10 μl for both AdSSTR2 and AdSSTR2-yCD membrane preparation. $^{125}$I-Tyr-1-Somatostatin-14 (Bachem, Torrance, CA, USA) binding reagent was incubated on ice for 15 min followed by centrifugation at 1400 r.p.m. for 5 min at 4°C. The supernatant was collected, and protein concentrations were determined using the Pierce Reducing Agent Compatible Kit. For each reaction, 4 μg of respective whole-cell extract were combined with 3 μg of $^{125}$I-Somatostatin-14 (PerkinElmer) to determine the amount of 5-FU conversion in infected breast cancer cells. The reactions were incubated in a 37°C water bath, and at various time points, an aliquot was removed, and the reaction stopped by adding 1 M acetic acid and a 4 mg ml$^{-1}$ 5-FC/5-FU mixture. The samples were spotted onto thin-layer chromatography plates (20 × 20 cm thin-layer chromatography plastic sheets, Silica gel 60 F$_{254}$ (EMD Chemicals, Darmstadt, Germany), and the plates were run in a butanol, acetic acid, and water solvent. The spots were then visualized under UV light (365 nm) using a UV light transilluminator, and the 5-FU and SSTR2-yCD product was identified using a handheld UV transilluminator at 254 nm, and strips of the thin-layer chromatography plates with each corresponding compound were placed into separate scintillation tubes with 5 ml ECLume scintillation fluid (MP Biomedicals, Irvine, CA, USA). The samples were then analyzed using a TriCarb Scintillation Counter (PerkinElmer) to determine the amount of 5-FC and 5-FU at each time point. The data were plotted as pmol of 5-FU formed over time, and CD activity was determined by dividing the slope of the curve by the amount of protein added.

In vitro 5-FC cytotoxicity assay

$IC_{50}$ (50% inhibitory concentration) values for AdSSTR2-yCD-infected MCF-7 cells were determined by using in vitro cytotoxicity assays. AdSSTR2 or AdSSTR2-yCD were administered to cells at 10 or 100 pfu per cell, and the cells were then plated into 96-well plates at a concentration of 5 × 10$^4$ cells per well. After the cells had 1 day to adhere, the media was aspirated and replaced with various concentrations of 5-FU in cell culture media ranging from 0 to 1000 μg ml$^{-1}$. After incubating at 37°C for 5 days, the media was aspirated, the wells were washed with phosphate-buffered saline, and 100 μl of 2% crystal violet in 70% ethanol was added to each well. The plates were incubated at room temperature for 3 h before removing the stain and washing the wells with H$_2$O. Once the wells were dry, the optical density was measured at 540 nm on a Molecular Devices SpectraMax M100 Microplate Reader (Molecular Devices, Sunnyvale, CA, USA). Percent survival was determined by dividing the absorbance value of the wells with 5-FU by the absorbance of the wells without 5-FC. The data were entered into GraphPad Prism 4 to generate sigmoidal dose–response curves, and $IC_{50}$ values were calculated from the curves.

In vivo yCD conversion in tumor xenografts

All animal studies were performed in accordance with the guidelines for the care and use of research animals by the Washington University Animal Studies Committee. Six days prior to cell implantation, a 60-day-release β-Estradiol pellet (Innovative Research of America, Sarasota, FL, USA) was implanted into the left flank of each 4-week-old female Fox Chase C.B.17-SCID mouse (Charles River Lab, Wilmington, MA, USA). MCF-7 cells mixed 1:1 with Matrigel Basement Membrane Matrix (Becton Dickinson, Palo Alto, CA, USA) were implanted at 1x10$^7$ cells per tumor in the right flank of each mouse. The tumors were allowed to grow for 5 weeks and were then injected with saline, AdSSTR2 at 3 × 10$^4$ pfu per tumor, or AdSSTR2-yCD at 1 × 10$^4$ or 3 × 10$^4$ pfu per tumor. Tumors were dissected 48 h later and were rinsed, weighed, and minced on ice. The tumors (250–450 mg) were then resuspended in 1 ml complete extraction buffer plus a Complete Protease Inhibitor tablet (Roche, Indianapolis, IN, USA) and sonicated. After incubating on ice for 30 min, the extracts were centrifuged at 10,000 r.p.m. for 20 min at 4°C. The supernatant was then assayed for protein concentration, and yCD conversion assays were performed as described above.

Radiolabeling of CB-TE2A-Y3-TATE with $^{64}$Cu

The peptides Y3-TATE and CB-TE2A-Y3-TATE were prepared by standard literature protocols. In vivo, the complexation of $^{64}$Cu to CB-TE2A-Y3-TATE was achieved by reacting 1 μg (6.3 × 10$^{-9}$ pmol) of CB-TE2A-Y3-TATE, 118 μl of 0.1 M NH$_4$OAc (pH = 8.0) and 37 MBq of $^{64}$CuCl$_2$ in 0.1 N HCl for...
Biodistribution

β-Estradiol pellets and MCF-7 tumors were implanted and allowed to grow as above. The mice were anesthetized and intratumoral injections of AdSSTR2 (n = 3) or AdSSTR2-yCD (n = 3) (each at $3 \times 10^{10}$ pfu per tumor) were performed. Two days following the viral injections, tail vein injections of $^{64}$Cu-CB-TE2A-Y3-TATE (185 kBq (S/μCi; 6 ng)) were performed. Two other groups of animals served as negative controls, with mice receiving an intratumoral saline injection (n = 3) and another group of animals receiving an intratumoral saline injection followed by a co-injection of $^{64}$Cu-CB-TE2A-Y3-TATE and 200 μg of Y3-TATE to serve as a block (n = 4). The mice were killed 4 h after injection, and blood, liver, spleen, pancreas, kidneys, muscle, tumor, bone and tail were removed. Tissue samples were weighed and placed in a gamma counter for determination of radioligand content. The percent injected dose per gram (% ID g$^{-1}$) was calculated based on a decay-corrected standard dose.

MicroPET/computed tomography (CT) imaging studies

Mice were implanted with β-estradiol pellets on the rear flank and MCF-7 tumors on the axillary thorax and allowed to grow as described above. The mice (n = 3) were injected intratumorally with $3 \times 10^{10}$ pfu of AdSSTR2-yCD or with saline, followed by intravenous injection of $^{64}$Cu-CB-TE2A-Y3-TATE 2 days later (4.1 MBq (110 μCi); 125 ng). One group of saline-injected mice received a co-injection of $^{64}$Cu-CB-TE2A-Y3-TATE and 200 μg of Y3-TATE to serve as a block. Four hours after injection, the mice were anesthetized with 1–2% isoflurane, positioned supine and imaged on microPET FOCUS 220 or Inveon PET small animal scanners (Siemens Medical Solutions, Malvern, PA, USA). The PET acquisition times were 10 min and CT images were obtained using a MicroCAT II System (ImTek, Inc., Knoxville, TN, USA).

The images were reconstructed with an Ordered-Subset Expectation Maximization algorithm which included corrections for scatter and attenuation. Regions of interest were drawn to encompass the entire tumor to determine the maximum activity concentration (nCi/cc) in the tumor. To calculate the standardized uptake values, the nCi/cc was divided by the nCi injected (decay corrected to the scan start time) and multiplied by the mouse weight.

Statistical analysis

All data are presented as the mean ± s.e.m. The Student’s two-tailed t-test was used to determine statistical significance at the 95% confidence level, with $P < 0.05$ being considered significantly different.

RESULTS

Construction of AdSSTR2-yCD

Crude AdSSTR2-yCD was validated for SSTR2 function in a single-point binding assay, and the yCD portion of the fusion was validated in a single time point CD conversion assay. Following these validations, the crude virus was used to produce cesium chloride-purified AdSSTR2-yCD (Q-Biogene) at a titer of $3.25 \times 10^{11}$ pfu ml$^{-1}$. The AdSSTR2 had a titer of $3.16 \times 10^{11}$ pfu ml$^{-1}$.

Competitive binding assay

To demonstrate that the SSTR2 portion of the fusion protein was functional, a competitive binding assay was performed using cells infected with AdSSTR2-yCD at 10 and 100 pfu per cell. A representative curve of MCF-7 cells infected at 10 and 100 pfu per cell is shown in Figure 1. The MCF-7 and T-47D cells both showed high affinity binding of $^{125}$I-Tyr$^{11}$-SST-14 with $K_d$ of $\sim 50–200$ pM at both 10 and 100 pfu per cell (Table 1). In addition, the expression levels were evaluated and MCF-7 cells had $B_{max}$ values of 764 ± 132 and 1829 ± 293 fmol mg$^{-1}$ after infection at 10 and 100 pfu per cell, respectively, whereas T-47D cells had $B_{max}$ values of 40 ± 6 and 217 ± 25 fmol mg$^{-1}$, respectively. When compared with the previously utilized AdSSTR2, there were no significant differences in the $K_d$ or in the $B_{max}$ values at 10 and 100 pfu per cell. A binding assay on uninfected MCF-7 and T-47D cells did not show endogenous SSTR2 expression (data not shown). However, others have shown low levels of SSTR2 on T-47D cells, whereas there are conflicting reports about SSTR2 expression on MCF-7 cells.

In vitro yCD conversion assay

The yCD conversion activity was determined by measuring the conversion of $^3$H-5-FC to $^3$H-5-FU in MCF-7 and T-47D cells infected with AdSSTR2-yCD at 10 and 100 pfu per cell. Table 2 shows the conversion results, in terms of pmol min$^{-1}$ mg$^{-1}$.$^1$ The MCF-7 cells infected at 100 pfu per cell showed the highest rate of conversion at 386 ± 27 compared with 214 ± 58 for T-47D cells infected at 100 pfu per cell. By comparison, when the cells were infected at 10 pfu per cell, the conversion rates were lower, with MCF-7 cells showing 104 ± 17 pmol min$^{-1}$ mg$^{-1}$ and T-47D cells showing 48 ± 10 pmol min$^{-1}$ mg$^{-1}$.

Table 1. Binding of $^{125}$I-Tyr$^{11}$-SST-14 to MCF-7 and T-47D cell membranes infected with AdSSTR2-yCD or AdSSTR2 at 10 or 100 pfu per cell

|          | 10 pfu per cell | 100 pfu per cell |
|----------|-----------------|------------------|
|          | $K_d$ (pM) | $B_{max}$ (fmol mg$^{-1}$) | $K_d$ (pM) | $B_{max}$ (fmol mg$^{-1}$) |
| MCF-7    | 87 ± 23     | 764 ± 132        | 200 ± 30   | 1829 ± 293                 |
| T-47D    | 86 ± 47     | 40 ± 6           | 134 ± 51   | 217 ± 25                   |

|          | 10 pfu per cell | 100 pfu per cell |
|----------|-----------------|------------------|
|          | $K_d$ (pM) | $B_{max}$ (fmol mg$^{-1}$) | $K_d$ (pM) | $B_{max}$ (fmol mg$^{-1}$) |
| MCF-7    | 80 ± 13     | 751 ± 107        | 190 ± 32   | 1595 ± 157                |
| T-47D    | 50 ± 20     | 32 ± 4           | 108 ± 53   | 242 ± 51                  |

Abbreviations: Ad, adenovirus; pfu, plaque-forming unit; SSTR2, somatostatin receptor subtype 2; yCD, yeast cytosine deaminase. The binding affinity ($K_d$) is expressed in pM and the maximum expression ($B_{max}$) is expressed in fmol mg$^{-1}$. © 2015 Nature America, Inc. Cancer Gene Therapy (2015), 215 – 221
infection at 10 pfu per cell. The IC₅₀ was three experiments, each performed in triplicate. We next wanted to confirm that an intratumoral injection of AdSSTR2-yCD would result in functional yCD in mice bearing MCF-7 tumor xenografts. These results show that intratumoral injection of 1 × 10⁸ pfu of AdSSTR2-yCD resulted in a ¹³¹I-5-FC conversion rate of 193 ± 41 pmol min⁻¹ mg⁻¹ compared with 698 ± 81 pmol min⁻¹ mg⁻¹ when 3 × 10⁹ pfu were injected (Table 2). In contrast, there was no detectable conversion when AdSSTR2 or saline were injected.

**Table 2.** CD conversion activity (pmol min⁻¹ mg⁻¹) of AdSSTR2-yCD in MCF-7 and T-47D cells infected at 10 or 100 pfu per cell (in vitro) or in MCF-7 tumors directly injected with either 1 × 10⁸ or 3 × 10⁹ pfu (in vivo)

|          | In vitro | In vivo |
|----------|----------|---------|
|          | 10 pfu per cell | 100 pfu per cell | 1 × 10⁸ pfu | 3 × 10⁹ pfu |
| MCF-7    | 62 ± 11  | 386 ± 27 | 193 ± 41 | 698 ± 81   |
| T-47D    | 11 ± 1   | 214 ± 58 | NP      | NP         |

Abbreviation: pfu, plaque-forming unit. Conversion activity was not performed (NP) in T-47D tumors in vivo.

**Figure 2.** Cell anti-proliferation data from MCF-7 cells infected with AdSSTR2-yCD at 10 or 100 pfu per cell. The data were generated using GraphPad Prism 4 from triplicate absorbance measurements for each concentration of ligand in which viable cells were fixed and stained with 2% crystal violet in 70% ethanol. The fixed cells were then washed, dried and the absorbance was measured. The graph is a representative IC₅₀ curve, with each point being a triplicate measurement. The IC₅₀ (µg ml⁻¹) values are the mean ± s.e.m. of three experiments, each performed in triplicate. Infected at 10 pfu per cell, the conversion rate was lower at 62 ± 11 and 11 ± 1 for MCF-7 and T-47D cells, respectively.

**In vitro 5-FC cytotoxicity assay**
To determine the 5-FC sensitivity of MCF-7 cells to AdSSTR2-yCD-mediated suicide gene expression, the cells were infected at 10 and 100 pfu per cell and treated with various concentrations of 5-FC, and the relative cell viability was determined using the crystal violet staining assay. As the previous assays demonstrated that the MCF-7 cells were more susceptible to infection with AdSSTR2-yCD than the T-47D cells, only the MCF-7 cells were evaluated for this and future assays. A representative cytotoxicity curve is shown in Figure 2. This shows that the IC₅₀ at 100 pfu per cell was 4.2 ± 1.4 µg ml⁻¹ compared with 27.2 ± 10.7 µg ml⁻¹ after infection at 10 pfu per cell. The IC₅₀ was > 1000 µg ml⁻¹ for cells infected with AdSSTR2 at either 10 or 100 pfu per cell (data not shown). Infection with AdSSTR2-yCD or AdSSTR2 at 10 and 100 pfu per cell (without addition of 5-FC) decreased cell viability by about 29% and 26%, respectively, when compared with uninfected cells (data not shown).

**In vivo yCD conversion in tumor xenografts**
We next wanted to confirm that an intratumoral injection of AdSSTR2-yCD would result in functional yCD in mice bearing MCF-7 tumor xenografts. These results show that intratumoral injection of 1 × 10⁸ pfu of AdSSTR2-yCD resulted in a ¹³¹I-5-FC uptake after AdSSTR2-yCD injection was significantly greater than that of the control mice (1.04 ± 0.21% ID g⁻¹), but were significantly greater (P < 0.01) than the control + block mice (0.23 ± 0.03% ID g⁻¹). There were no significant differences between the two control groups except for the pancreas being higher in the control vs control + block. The liver and spleen uptake of ⁶⁵Cu-CB-TE2A-Y3-TATE after intratumoral injection of AdSSTR2 was significantly greater (P < 0.01) than all of the other groups, whereas the liver uptake was significantly greater (P < 0.01) after injection of AdSSTR2-yCD compared with control, but did not reach significance when compared with control + block. Conversely, the spleen uptake after AdSSTR2-yCD injection was significantly greater compared with control + block (P < 0.01), but not when compared with control.

**MicroPET/CT imaging studies**
To test whether AdSSTR2-yCD could be used to image in vivo gene transfer, AdSSTR2-yCD was injected intratumorally into MCF-7 tumor xenografts followed by intravenous injection of ⁶⁵Cu-CB-TE2A-Y3-TATE. Small animal PET/CT images of ⁶⁵Cu-CB-TE2A-Y3-TATE at 4 h are shown in Figure 4. Coronal (Figure 4a and b) and transaxial (Figure 4c and d) views of two mice (one receiving AdSSTR2-yCD (Figure 4a and c) and the other receiving saline...
DISCUSSION

Gene-directed enzyme prodrug therapy using the CD/5-FC system has been investigated for some time. It has been utilized as a stand-alone therapeutic approach and has been combined with other gene therapy, chemotherapy and radiotherapy strategies. In addition, the efficacy of this gene-directed enzyme prodrug therapy approach has been enhanced by creating uracil phosphoribosyltransferase fusion genes with CD, thymidine kinase fusion genes with CD or mutant bCD enzymes to increase the catalytic conversion of 5-FC to 5-FU. However, one of the major limitations of adenoviral-based gene therapy in general, and specifically CD-based molecular chemotherapy, is the inability to determine the level of in vivo gene transfer non-invasively. Tissue biopsies to determine levels of gene transfer are not ideal because they are invasive, not easily repeated and are subject to sample variability that does not give a global picture of gene transfer. Therefore, the development of a CD construct that can be used to monitor the location, magnitude and change in magnitude over time of gene transfer non-invasively would address this limitation.

Previously, several approaches have been investigated to non-invasively determine the expression of CD. One approach for monitoring CD enzyme activity has been through the use of 19F magnetic resonance spectroscopy. Another approach has been used to demonstrate the conversion of 5-FC to 5-FU in animal models and to quantify their concentrations as well as their resulting metabolites. However, the sensitivity of magnetic resonance spectroscopy is inherently limited and sufficient metabolite concentrations must be sustained during image acquisition to achieve adequate resolution. It has been suggested that 5-FC is not an effective probe for magnetic resonance spectroscopy of CD because the metabolism of 5-FC by CD does not result in persistently elevated levels of fluorinated metabolites.

Another approach has been through the use of nuclear imaging methods. This method has generally used a fusion protein consisting of HSV1-TK and CD with the main purpose of combining these prodrug strategies for increasing therapeutic efficacy. However, HSV1-TK can also be used as a reporter gene that can be imaged with radioactive probes. These probes have consisted of uracil nucleoside derivatives radiolabeled with iodine isotopes for PET and single photon emission computed tomographic imaging or acycloguanosine derivatives radiolabeled with 18F for PET imaging. These probes are transported into cells, phosphorylated by cells expressing HSV1-TK (or fusion protein), and the phosphorylated product is trapped, leading to a signal when compared with surrounding tissue not containing HSV1-TK. Hackman et al. used 125I-F1A12U to image rats with tumor cells stably expressing the CD/TK fusion gene by PET. Freytag et al. used 18F-FHBG to image CD/TK expression by PET in dogs that had been injected in the
pancreas with a replicative Ad vector. However, the correlation between HSV1-TK expression and the accumulation of radiolabeled probes when HSV1-TK is expressed at high levels is lacking.38 Thus, the HSV1-TK reporter may not be appropriate for quantifying the expression level of CD in gene therapy protocols and another reporter may be necessary.

Our previous studies used the radiolabeled somatostatin analogs [111In]-DTPA-octreotide (Octreoscan), [99mTc]-P829 (NeoTect), [99mTc]-P2045 or [99mTc]-Demotide to demonstrate localization in a tumor injected directly with an adenosinovirus (AdSSTR2) encoding the SSTR2 gene driven by a cytomegalovirus promoter. Importantly, we demonstrated that there is a strong correlation between the expression of SSTR2 and the tumor uptake of [99mTc]-P2045 after intratumoral injection of various amounts of AdSSTR2.42 Therefore, the focus of the present study was to construct and evaluate a novel fusion gene consisting of SSTR2 and CD.

In the present study, we constructed an adenosinovector encoding for a fusion protein between SSTR2 and yCD and evaluated this construct (AdSSTR2-yCD) in human breast cancer cells. The in vitro binding studies demonstrated that SSTR2 was still functional as it bound somatostatin at an affinity of ~85–200 pM in both MCF-7 and T-47D cells. This was similar to when the wild-type receptor was utilized in these cells (~50–190 pM). As expected, an increase in receptor concentration (Bmax) was observed when infecting at 100 pfu per cell compared with 10 pfu per cell. In addition, these studies showed that MCF-7 cells were more easily infected with adenovirus compared with the T-47D cells as determined by the higher expression observed in the MCF-7 cells. The conversion and cytotoxicity assays demonstrated the functionality of the yCD in the fusion protein. Similar to the binding assays, the conversion of [3H]-5-FU to [3H]-5-FU was more efficient in MCF-7 cells compared with T-47D cells as well as at 100 pfu per cell compared with 10 pfu per cell. The conversion at 100 pfu per cell (~200–400 pmol min⁻¹ mg⁻¹) was greater than when pancreatic tumor cells were infected at 25 pfu per cell with bCD (~0.2–0.7 pmol min⁻¹ mg⁻¹) and similar to a catalytically enhanced bCD (~100–250 pmol min⁻¹ mg⁻¹).12 This catalytically enhanced bCD demonstrated much higher conversion (~1000–2800 pmol min⁻¹ mg⁻¹) when evaluated in human glioma cell lines.5 Owing to the higher expression and conversion in MCF-7 cells compared with T-47D cells, we only evaluated cytotoxicity in the MCF-7 cells. Interestingly, the IC50 after infection at 100 pfu per cell (4.2 µg ml⁻¹) was similar to many of the glioma and pancreatic cells infected with the catalytically enhanced bCD, although this construct showed greater [3H]-5-FU to [3H]-5-FU conversion.5,12 This difference is likely due to the inherent sensitivity of the individual cell lines to 5-FU.

In vivo evaluation of AdSSTR2-yCD was then conducted in mice bearing MCF-7 tumor xenografts using conversion assays, biodistribution and small animal PET imaging. Enzyme activity was confirmed after intratumoral injection of AdSSTR2-yCD by removal of the tumor 2 days after injection and observing conversion of [3H]-5-FU to [3H]-5-FU. As expected, more conversion was observed after injection of a higher dose of virus. Previous studies in glioma and pancreatic cancer xenografts using the catalytically enhanced bCD also demonstrated in vivo enzyme activity, but 20–fold less virus was used in these studies.5,12 Biodistribution studies demonstrated a 1.5-fold increase in tumor uptake of [64Cu-CB-TE2A-Y3-TATE after injection of AdSSTR2-yCD compared with saline-injected control tumors. This is similar to the 1.7-fold increase observed after injection of AdSSTR2. Interestingly, there was an increase in tumor uptake of [64Cu-CB-TE2A-Y3-TATE in saline-injected control mice when compared with mice receiving saline plus an excess of Y3-TATE upon injection of [64Cu-CB-TE2A-Y3-TATE. This indicates SSTR2-receptor-specific binding in mice that received saline alone and differs from the in vitro results that do not show SSTR2-specific binding in MCF-7 cell without AdSSTR2-yCD or AdSSTR2 infection. This may be due to the upregulation of SSTR2 in the tumor xenografts owing to the exposure to estrogen from the implanted pellets, similar to other studies demonstrating increase in SSTR2 expression after exposure to estrogen.43,44 There was a 6.7-fold increase in the tumor uptake of [64Cu-CB-TE2A-Y3-TATE after AdSSTR2-yCD injection compared with the uptake after saline injection plus the blocking agent. Similar to other studies, there was increased liver and spleen accumulation of [64Cu-CB-TE2A-Y3-TATE after injection of AdSSTR2-yCD compared with controls.13,45 This is likely due to adenosinovector infection of these organs even though the adenosinovirus was injected intratumorally.46,47 MicroPET imaging clearly shows accumulation of [64Cu-CB-TE2A-Y3-TATE in tumors injected with AdSSTR2-yCD (Figure 4a and c). It is interesting to note the heterogeneity of the radioactive uptake as it is likely that there is preferential uptake of [64Cu-CB-TE2A-Y3-TATE along the injection site of AdSSTR2-yCD, which is similar to previous studies.17,48 The standardized uptake value analysis shows that there is a 9.6-fold increase in tumor uptake of [64Cu-CB-TE2A-Y3-TATE after AdSSTR2-yCD infection compared with the uptake after saline injection plus the blocking agent. This is similar to the 6.7-fold increase observed in the biodistribution studies. It is not clear why the upregulation of SSTR2 was not observed in this study as there was no difference in the standardized uptake value between saline tumors and mice that received saline plus the blocking agent. These studies demonstrate that [64Cu-CB-TE2A-Y3-TATE can be used to non-invasively image the expression of SSTR2-yCD.

In conclusion, we have shown that we can construct an adenosinovirus containing the SSTR2-yCD fusion protein that is fully functional. In vitro studies demonstrated that the fusion protein can bind a radioactive SSTR2 ligand, convert [3H]-5-F to [3H]-5-FU and be cytotoxic to cells after 5-F treatment. The activity of the fusion protein was also demonstrated in vivo, and importantly, its expression was capable of being imaged non-invasively using PET. Although the use of this model, non-replicative adenosinovector was sufficient to demonstrate the functionality of the SSTR2-yCD, improved gene delivery vehicles may be needed to optimize its therapeutic utility.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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