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Enhanced transduction of CAR-negative cells by protein IX-gene deleted adenovirus 5 vectors

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A B S T R A C T
In human adenoviruses (HAdV), 240 copies of the 14.3-kDa minor capsid protein IX stabilize the capsid. Three N-terminal domains of protein IX form triskelions between hexon capsomers. The C-terminal domains of four protein IX monomers associate near the facet periphery. The precise biological role of protein IX remains enigmatic. Here we show that deletion of the protein IX gene from a HAdV-5 vector enhanced the reporter gene delivery 5 to 25-fold, specifically to Coxackie and Adenovirus Receptor (CAR)-negative cell lines. Deletion of the protein IX gene also resulted in enhanced activation of peripheral blood mononuclear cells. The mechanism for the enhanced transduction is obscure. No differences in fiber loading, integrin-dependency of transduction, or factor-X binding could be established between protein IX-containing and protein IX-deficient particles. Our data suggest that protein IX can affect the cell tropism of HAdV-5, and may function to dampen the innate immune responses against HAdV particles.© 2010 Elsevier Inc. Open access under the Elsevier OA license.

Introduction

Protein IX is a non-essential protein in the capsid of human adenoviruses (HAdV). The protein has a size of 14.3 kDa, is present at 240 copies per virion, and has three highly conserved regions present in the amino (N) terminus, the central part (alanine-rich), and the carboxy (C) terminus (leucine-rich). The location and function of protein IX in the virus capsid has been the subject of investigation and debate for many years (Vellinga et al., 2005b). Recent work by different groups has brought consensus on its location and topology in the capsid (Fabry et al., 2009; Saban et al., 2006). The N-terminus of the protein is located in between hexon cavities of the groups of nine (GON) hexons, presumably stabilizing the GONs. The C-terminus of the protein forms an alpha helix and is exposed on the capsid surface in close contact with hexon hypervariable region 4 (HVR4) (Saban et al., 2006). C-terminal domains of three protein IX molecules associate in a parallel orientation, whereas a fourth domain binds in an antiparallel orientation (Fabry et al., 2009). The role of protein IX in the capsid remains enigmatic. In vitro analysis revealed the N-terminus of protein IX to confer a thermostable phenotype on HAdV-5 capsids (Vellinga et al., 2005a). Propagation of protein IX gene deleted HAdV-5 in cell culture yields wild-type like virus titers, demonstrating that protein IX is dispensable for virus replication in vitro.

Protein IX has potential as an anchor for the attachment of different types of polypeptides to the viral capsid. Targeting of HAdV-5 to tumor cells has been achieved by genetically fusing protein IX to a single-chain T cell receptor directed against MHC class I in complex with MAGE-A1 peptides (de Vrij et al., 2008). Similarly, integrin-binding arginine–glycine–aspartate (RGD) peptides, as well as single-chain antibody fragments have been incorporated in this way (Vellinga et al., 2007, 2004). Alternatively, targeting ligands can be coupled to protein IX via the genetic inclusion of cysteine residues and subsequent chemical coupling of ligands to the reactive thiol groups (Corjon et al., 2008). Multiple polypeptides can be incorporated simultaneously (Tang et al., 2009). A triple-mosaic HAdV-5 vector was developed with a poly-lysine motif, the herpes simplex virus type 1 (HSV-1) thymidine kinase, and the monomeric red fluorescent protein fused with protein IX, thereby combining targeting, therapeutic, and imaging modalities. Recently, it was demonstrated that HAdV-5 vaccine vectors with pathogen-specific antigens fused to pIX can stimulate robust protective immune responses in animals, suggesting a new route for the development of improved HAdV-5 based recombinant vaccines (Bayer et al., 2010; Boyer et al., 2010).
Here we report on the enhanced delivery of transgenes into CAR-negative cell lines as a result of protein IX-gene deletion from a HAdV-5-based vector. Furthermore, the protein IX-deficient particles demonstrated enhanced activation of peripheral blood mononuclear cells (PBMCs), and had a different in vivo distribution after intravenous delivery in a mouse model. The exact molecular mechanism behind this ΔpIX effect remains to be delineated. Our data suggest that protein IX can affect the cell tropism of HAdV-5, and may function to dampen the innate immune responses against HAdV particles.

**Results**

**Enhanced transgene expression in CAR-negative cells with Ad5ΔE1ΔpIX**

To study the role of protein IX in the HAdV-5 transduction of cells, we compared the vectors Ad5ΔE1+pIX and Ad5ΔE1ΔpIX for luciferase transgene expression in a panel of cell lines (Fig. 1A). Cell lines with varying expression levels of CAR were included (Fig. 1B). Whereas similar expression levels were obtained with both vectors in the CAR-positive cell lines HeLa, A549, and MEL2A, the vector Ad5ΔE1ΔpIX yielded higher levels than Ad5ΔE1+pIX in the CAR-negative cell lines MZ2-MEL3.0 and VH10. Since these results suggested a specific role of the protein IX lacking vector in mediating relatively higher transduction in the absence of CAR, Ad5ΔE1+pIX and Ad5ΔE1ΔpIX were analyzed for reporter gene expression in MZ2-MEL3.0 cells versus MZ2-MEL3.0/CAR cell lines (Fig. 2B). MZ2-MEL3.0/CAR cells stably expressed CAR via transduction with a recombinant lentivirus, which was confirmed by flow cytometry and immunofluorescence staining (Fig. 2A). In MZ2-MEL3.0 cells the reporter gene expression upon infection with Ad5ΔE1ΔpIX was found to be ten-fold increased compared to infection with Ad5ΔE1+pIX, while in MZ2-MEL3.0/CAR cells the difference was a mere two-fold (Fig. 2B). The enhanced transgene expression for Ad5ΔE1ΔpIX on the CAR-negative cell line MZ2-MEL3.0 appeared to be not affected by the establishment of protein IX expression in the cells (by using the recombinant lentivirus LV-CMV-pIX-IRES-NPTII; Vellinga et al., 2006) prior to the transduction (result not shown).

As a next step, the involvement of the C-terminal region of protein IX in the observed phenomenon was investigated. This domain, which is rich in leucine amino acids and is exposed on the HAdV-5 capsid as an alpha-helical structure (Fabry et al., 2009; Saban et al., 2006), is highly conserved in human adenoviruses. The biological function of this conserved domain of protein IX is unknown. We analyzed the vector Ad5ΔE1pIXΔLEU, which lacks a major part of the C-terminal region of protein IX (amino acids 100 to 114) for reporter gene expression in MZ2-MEL3.0 and MZ2-MEL3.0/CAR. Ad5ΔE1pIXΔLEU demonstrated enhanced transduction of the CAR-negative cell line, very similar to the Ad5ΔE1ΔpIX vector (Fig. 2C).

To assess the appearance of the vector particles and to check for the absence of microaggregation, electron microscopy was performed on Ad5ΔE1+pIX and Ad5ΔE1ΔpIX vector batches. This showed identically shaped virus particles (Fig. 3A). No signs of microaggregation were observed. The Ad5ΔE1ΔpIX stock appeared to contain more small particulate matter, possibly virus debris. As previously described, pIX-deficient HAdV-5 particles have an enhanced tendency to partly dissociate into fiber- and penton base-lacking particles (Fabry et al., 2005). However, our vectors had similar capsid incorporation levels of fiber and hexon proteins, as evident from immunoblot analyses (Fig. 3B), thus ruling out differences in particle dissociation for the vector preparations.

**Transduction with Ad5ΔE1ΔpIX is integrin-dependent**

Wild-type HAdV-5 enters cells via high affinity binding of the fiber knob domain to CAR (Bergelson et al., 1997). Subsequently low affinity interaction of the penton base with cellular integrins αvβ3 and αvβ5 promotes virus internalization in clathrin-coated pits (Nemerow and Stewart, 1999; Wickham et al., 1993). To answer the question if Ad5ΔE1ΔpIX still uses integrins for cellular uptake, we analyzed Ad5ΔE1+pIX and Ad5ΔE1ΔpIX for transgene expression (GFP) in the presence or absence of bivalent cations, which are necessary for integrin-mediated uptake of wild-type HAdV-5 into cells (Wickham et al., 1993) (Fig. 4A). This experiment again displayed a stronger reporter gene expression of Ad5ΔE1ΔpIX in MZ2-MEL3.0 cells compared to Ad5ΔE1+pIX. For both vectors the transduction appeared to be totally dependent on the presence of bivalent cations, with a complete reduction to background GFP levels for the cation-negative incubation. This is consistent with integrin-mediated uptake for both vectors. More specifically, the integrin dependency of Ad5ΔE1ΔpIX was confirmed by a small but significant (approximately two-fold)}
decrease in transduction after incubation of MZ2-MEL3.0 cells with antibodies directed against αVβ3 and αVβ5 integrins (Fig. 4B).

Similar antibody-blocking (1.5-fold reduced transduction for anti-αVβ3 and anti-αVβ5) was observed for Ad5ΔE1+plIX. Incubating the cells with higher concentrations of antibodies did not result in further reductions in transduction levels (data not shown). Anti-integrin-mediated blocking of transduction was also observed on A549 cells (Fig. 4B). From these data we conclude that the vector Ad5ΔE1ΔplIX still uses integrins for cell internalization in CAR-deficient cells.

**Reduced virus spread of the replication-competent virus Ad5ΔplIX**

Our data from the comparative transduction analysis suggest an alternative interaction of HAdV-5 particles lacking protein IX with the cell surface. In parallel to cell tropism extending capsid modifications

![Image](A)

**Fig. 2.** Transduction assays on MZ2-MEL3.0 and MZ2-MEL3.0/CAR. (A) Detection of CAR expression in MZ2-MEL3.0 cells by immune-fluorescence staining with anti-CAR antibody and FITC-labeled secondary antibody. The insets represent flow cytometry histograms after staining with anti-CAR antibody and PE-labeled secondary antibody. (B) Luciferase expression in MZ2-MEL3.0 and MZ2-MEL3.0/CAR after Ad5ΔE1+plIX and Ad5ΔE1ΔplIX transduction. Error bars represent SEM (n = 3). (C) Fold enhancement of MZ2-MEL3.0 transduction with Ad5ΔE1ΔplIX and Ad5ΔE1pIXLEU as compared to the transduction with Ad5ΔE1+plIX. The fold enhancements are normalized to the vector transduction ratios on MZ2-MEL3.0/CAR (*p < 0.05, **p < 0.005 versus Ad5ΔE1+plIX). Errors bars represent SEM (n = 3).

![Image](B)

**Fig. 3.** (A) Electron microscopy on Ad5ΔE1+plIX and Ad5ΔE1ΔplIX samples with negative staining of the vector particles in phosphotungstic acid. (B) Immunoblot detection on Ad5ΔE1+plIX and Ad5ΔE1ΔplIX lysates to analyze capsid incorporation levels of protein IX, hexon, and fiber proteins.
described for other viruses (de Haan et al., 2005), it is likely that protein IX deletion from a replication-competent HAdV-5 virus would result in a modified ability to spread in monolayer cell cultures. To investigate this, we constructed the replication-competent HAdV-5 viruses Ad5+pIX and Ad5ΔpIX. Both viruses expressed GFP, allowing accurate measurement of plaque size. On A549 cells the plaque size for the Ad5ΔpIX virus (median 30 arbitrary surface units (ASU), range 20–170) appeared to be significantly smaller than the plaque size for Ad5+E1+pIX (median 100 ASU, range 30–290). A similar difference in plaque size was observed on the CAR-negative cell line VH10, with Ad5ΔpIX (median 50 ASU, range 30–150) yielding much smaller plaques compared to Ad5+E1+pIX (median 100 ASU, range 75–280).

From these analyses we conclude that protein IX-gene deletion from the genome of the replication-competent virus results in a decrease in virus spread in CAR-positive (A549) as well as CAR-negative (VH10) monolayer cell cultures.

Enhanced activation of peripheral blood mononuclear cells by Ad5ΔE1ΔpIX

Our findings on the modified transduction characteristics of protein IX-deficient HAdV-5 vectors are of relevance for: (1) fundamental adeno-virology (as the findings point towards a novel biological function of protein IX), and (2) the development of protein...
IX-modified HAdV-5 vectors for gene therapies. For both these aspects, it will be highly interesting to determine the effect of protein IX-deletion on the interaction of HAdV-5 vectors with white blood cells. Therefore, we incubated freshly isolated human peripheral blood mononuclear cells with Ad5ΔE1+pIX and Ad5ΔE1ΔpIX, and analyzed GFP expression and the expression of cellular activation markers. This revealed relatively high levels of GFP expression in the monocyte population. The percentage of GFP-positive monocytes was similar for both vectors, varying between 10% and 30% at 100 pp/cell, depending on the donor (data not shown). For both vectors the GFP expression in the T cell, B cell, and NK cell populations was very low (<1% GFP-positive cells). Although Ad5ΔE1+pIX and Ad5ΔE1ΔpIX showed identical GFP expression levels in the monocytes, the incubation with Ad5ΔE1ΔpIX resulted in a remarkably higher level of monocyte activation, as indicated by enhanced CD86 expression (Fig. 5A). The percentage of CD86 positive monocytes as well as the mean fluorescence intensity for CD86 was significantly higher for the pIX-lacking vector. This enhancement in monocyte activation was observed for monocytes derived from PBMCs of three different donors and with different virus batches. The up-regulated CD86 level observed for monocytes derived from PBMCs of three different donors and with different virus batches. The up-regulated CD86 level involved the entire Ad5ΔE1ΔpIX-incubated monocyte population, not only the GFP-positive cells (Fig. 5A). Incubation with Ad5ΔE1ΔpIX also resulted in enhanced activation of NK cells, as demonstrated by an increase in CD69 expression (Fig. 5B). Interferon-gamma (IFN-γ) ELISA of PBMC supernatants revealed higher levels of IFN-γ production after incubation with Ad5ΔE1ΔpIX at the higher input virus levels (Fig. 5C).

Enhanced liver transduction upon intravenous administration of Ad5ΔpIX

To study the functional consequences of protein IX-gene deletion on biodistribution in mice, Ad5ΔE1+pIX and Ad5ΔE1ΔpIX viruses were administered via tail vein injection. Luciferase expression in multiple organs was determined at 3 days post injection. The vector lacking protein IX yielded a more than ten-fold higher luciferase activity in the liver (Fig. 6A). These data show that the absence of protein IX in the viral capsid strongly affected the biodistribution of HAdV-5 particles.

Recent reports have described the involvement of plasma proteins, such as the blood coagulation factor X (FX), in HAdV-5 transduction of the liver (Kalyuzhnyi et al., 2008; Waddington et al., 2008). To study whether removal of protein IX influences the effects of clotting-factor binding we compared the FX-mediated transduction for Ad5ΔE1+pIX and Ad5ΔE1ΔpIX (Fig. 6B). In vitro incubation of A549 cells and HepG2 cells with FX resulted in a similar enhancement in transduction for Ad5+pIX and Ad5ΔpIX. As expected, no effect on transduction was observed after incubation with the mutant FX (FX\textsuperscript{MUT}), which lacks the domain necessary for binding to the HAdV-5 capsid (Waddington et al., 2008). From these data we conclude that the absence of protein IX does not affect the binding of coagulation factor X.

Discussion

From our data we conclude that the omission of protein IX from HAdV-5 vectors enhances viral transduction of cell lines that are low in expression of the adenovirus receptor CAR. This finding is of relevance for the development and implementation of protein IX-gene modified HAdV-5 vectors. Also, the findings enhance our knowledge on HAdV-5 biology and evolution, which especially becomes clear if stating our conclusion in a 'backwards' manner: the introduction of protein IX in HAdV-5 (making it wild-type HAdV-5) decreases viral transduction of cell lines that are low in CAR expression. Although speculative, it is very well possible that the presence of protein IX in the HAdV-5 capsid negatively interferes with non-specific cell transduction, and thereby plays a role in determining the virus tropism. Noteworthy, the extended cell tropism of the Ad5ΔE1ΔpIX vector, as presented by its enhanced transduction of CAR-negative cells, did not come with a loss in ability of CAR-mediated transduction of cells. This is clear from the comparison on M22-MEL3.0 and M22-MEL3.0/CAR. Introduction of CAR in the CAR-negative cells significantly increased transduction levels with Ad5ΔE1ΔpIX. It is conceivable that in CAR-expressing cells the protein IX-deficient particles can use either the CAR/fiber-dependent

![Fig. 6](image-url)
mechanism, or the CAR-independent pIX-dependent mechanism. Quantifying the relative contribution of each of these mechanisms to the total transduction requires tools for specifically blocking the new pathway. Such inhibitors remain to be identified.

Interestingly, the vector Ad5ΔE1pIXΔE1EU had Ad5ΔE1ΔapIX-like properties, implicating the importance of the C-terminal domain of protein IX in inhibiting transgene expression in CAR-negative cell lines. The specificity for the C-terminal domain of protein IX excludes differences in viral capsid stability as a cause for the observed phenomenon, since deletion of this domain does not result in reduced capsid stability (Vellinga et al., 2005a).

Wild-type HAdV-5 enters cells via high affinity binding of the fiber knob domain to CAR (Bergelson et al., 1997), followed by interaction of the RGD motif of the penton base with cellular integrins αvβ3 and αvβ5 promoting rapid adenovirus cell entry into clathrin-coated vesicles (Nemerow and Stewart, 1999; Wickham et al., 1993). Similar to the Ad5ΔE1−1+pIX control virus, Ad5ΔE1ΔapIX requires the presence of bivalent cations for its transgene delivery, indicating the usage of cellular integrins for cell internalization (Wickham et al., 1993). More specifically, blocking cells with anti-integrin antibodies resulted in a decrease in transduction for Ad5ΔE1ΔapIX, thereby confirming the integrin-dependency. Unfortunately, our efforts to compare the vectors for a general difference in cell-binding affinity, using Alexa488-TFP (tetrafluorophenyl) labeled vector particles, were not conclusive as a result of strong and reproducible negative effects of the labeling procedure on the pIX-deficient vector particles. The effects were not identical for protein IX-positive and protein IX-negative particles, making the results obtained with these particles in comparative binding assays unreliable. Alternative protocols for fluorescent- or radio-labeling of vector particles might be more suitable for comparing the cell-binding affinity. Labeled vector particles might also be used for analyzing differences in cell surface motility between protein IX-containing and protein IX-lacking vectors. Through largely unknown mechanisms HAdV-5 particles migrate on the cell surface and alterations in viral movement can result in modified transduction (Patterson and Russell, 1983).

The removal of the protein IX gene from a replication-competent HAdV-5 virus results in a small-plaque phenotype on CAR-positive as well as CAR-negative cell lines. This suggests the tropism modifying mutation affects the virus’s capacity to spread from cell to cell. Such small-plaque phenotypes of extended tropism mutants is not unprecedented: similar phenotypes have been described for murine corona virus mutants that acquired the capacity to bind to cell. Such small-plaque phenotypes of extended tropism mutants have important consequences for the virus particles in the blood to complement factors and immunoglobulins (mediating uptake in Kupffer cell macrophages) (Kalyuzhniy et al., 2008; Xu et al., 2008), and coagulation factors (resulting in hepatocyte transduction) (Kalyuzhniy et al., 2008; Waddington et al., 2008). The enhanced transduction of the liver with Ad5ΔE1ΔapIX observed in our mouse model seems not to be a result of more efficient binding of the vector to coagulation factor X (FX), as can be concluded from our in vitro FX-binding assay. An alternative explanation might be that the absence of protein IX extends the HAdV-5 tropism, enabling the transduction of cells in the liver that do not present CAR. Of interest, primary human hepatocytes were recently found to have CAR localized at cellular junctions that are inaccessible to the hepatic blood flow (Au et al., 2007). This localization is in contrast to the CAR molecules on hepatocellular carcinoma cells (like HepG2), being highly available for HAdV-5 binding (Au et al., 2007; Bangari et al., 2005).

Protein IX is strongly conserved in all primate adenoviruses indicating the importance of the protein. A biological role for protein IX in HAdV-5 capsid stabilization has been proposed, based on in vitro heat-stability assays (Vellinga et al., 2005a). Our findings point toward other biological functions of protein IX in (i) determining the cell tropism of HAdV-5, and (ii) negatively interfering with the innate immune response against HAdV-5. More insight into the mechanisms by which the presence of protein IX affects gene transfer and activation of immune cells may be of use for enhancing the efficiency of current (e.g. Atencio et al., 2006) and future gene therapies involving protein IX modified HAdV-5 vectors.

Materials and methods

Cell lines

All cell lines were maintained as monolayers at 37 °C in a humidified atmosphere of 5% CO₂. The human cell lines HeLa (cervical
Table 1

| Oligonucleotides used in the cloning procedures. |
|-----------------------------------------------|
| **FWD CAR_Pul** | 5′-GATGATGCGAGATGCAGGGCTCCGCGCTG-3′ |
| **REV CAR** | 5′-CCAGCCCTGCTTGATGACCATGAGCCATCGGTCTG-3′ |
| **Nhel** | |
| **FWD** | 5′-TTGAGGACGCGCGCGACACTGGCAGAATTCGATTGAGG-3′ |
| **Scal_correct** | |
| **REV** | 5′-CCATCAAACGATGTTGCTCTACGTAGGGGCGCGCTCTCGAA-3′ |
| **Scal_correct** | |
| **FWD** | 5′-GTTTTCCGCTGAAAGGCTTACTGGCTCAGGCGAGAACG-3′ |
| **Spel_correct** | |
| **REV** | 5′-GGTTTTACCGCAATGCTAGCTACTGCTG-3′ |
| **Spel_correct** | |
| **pIX_Scal** | 5′-CCGCGAAGTACAGTAACCGCCACTGCGTGTAGG-3′ |
| **pIX_Spel** | 5′-CCGACTAGTCTTAAACCAGGATCGGAGGGG-3′ |

HAdV-5 **BsrGI-MfeI** fragment containing the E1 genes (nucleotides 193–3925) was isolated from pTG3602 (kindly provided by Dr. M. Luksi, Transgene, Strasbourg, France), and cloned into the **BsrGI-MfeI** digested pTrackCMV-GFP/LUC (Vellinga et al., 2006), thereby replacing the GFP/LUC genes with the HADV-5 E1 region. By using site-directed mutagenesis (QuikChange site-directed mutagenesis kit; Stratagene) (primers 3, 4, 5, 6; Table 1) two restriction sites were introduced in the protein IX gene; a Scal site at the start codon of protein IX and a Spel site upstream of the protein IX stop codon, thereby creating pShuttle+E1+plxScalSpel. Next, the pSh+plx plasmid was constructed by Scal/Spel digestion and re-ligation of the protein IX sequence (amplified from pAd5pix (Vellinga et al., 2004) by using primers 7 and 8 (Table 1)) into the Scal/Spel linearized pShuttle+E1+plxMfeI/Spel. The Scal site was restored to the wild-type HADV-5 sequence by exchanging the Scal-overlapping MfeI/HindIII fragment with the corresponding fragment from pTG3602. The Spel site and the downstream ‘plx-remainder sequence’ were left intact, since part of this sequence forms a hairpin-loop structure situated over the polyA site of the E1B transcript, which might be essential for efficient polymerase slippage needed for polyadenylation (Sittler et al., 1995).

The pBB backbone plasmid was constructed by replacing the E3-lacking Spel–PacI fragment (nucleotides 27238–33443) of pDAdEasy-1 (He et al., 1998) with the corresponding Spel–PacI fragment of pShuttle-ΔE3-ADP-EGFP-F2 (Ono et al., 2005), thereby introducing eGFP in the E3 region under control of the viral major late promoter. The coding sequence for the E3 Adenovirus Death Protein (ADP) was retained. The kanamycin resistance gene (inserted with the pShuttle-ΔE3-ADP-EGFP-F2 fragment) was removed by ClaI digestion and re-ligation of the two largest fragments.

Recombination of pBB with pSh+plx and pShΔpix in E. coli and subsequent virus rescue in A549 cells were performed as described elsewhere (He et al., 1998). Virus was purified by a standard double cesium chloride gradient protocol, dialyzed against sucrose buffer (5% sucrose, 140 mM NaCl, 5 mM NaHPO₄, 2H₂O, 1.5 mM KH₂PO₄) and stored at −80 °C. The virus titer was determined by the PicoGreen-DNA binding assay (Murakami and McCaman, 1999) (for pfu/ml measurement), and a plaque assay on A549 cells (Fallaux et al., 1996) (for pfu/ml measurement).

For analysis of virus spread, GFP-positive plaques were photographed (Olympus Camera Digital Camera C-3030, installed on an Olympus CK40 microscope) and the plaque size was determined in arbitrary units (Olympus DP-sof v.5.0 Soft Imaging System software). The median plaque size of Ad5Δpix was normalized to the plaque size for Ad5+plx.

**Analysis of CAR presentation on the cell surface**

Flow cytometry was performed to determine the levels of CAR presentation on the cell surface. Cells in suspension (in PBS with 0.5% bovine serum albumin and 0.02% sodium azide) were incubated with mouse monoclonal anti-CAR antibody (clone Rmcb, Upstate Biotechnology, Lake Placid, NY, diluted 1:1000) for 30 min on ice, followed by incubation with phycoerythrin (PE)-conjugated rabbit-anti-mouse secondary antibody (Caltac Laboratories, Burlingame, CA, USA) for 30 min on ice. Flow cytometry data were analyzed with CellQuest software (Becton Dickinson).

Immunohistochemistry was performed on the cell line MZ2-MEL3.0/CAR. After washing with phosphate-buffered saline (PBS), the cells were fixed in acetone/methanol (1:1) for 10 min at room temperature. Staining was performed with the anti-CAR antibody (clone Rmcb, Upstate Biotechnology, Lake Placid, NY, diluted 1:500). Fluorescein isothiocyanate (FITC)-conjugated rabbit-anti-mouse antibody (Jackson ImmunoResearch, France) was used as secondary antibody.
Virus transduction assays

Luciferase expression

The transduction efficiency of CAR-positive (HeLa, A549, MEL2A, M22-MEL3.0/CAR) and CAR-negative (M22-MEL3.0, VH10) cell lines by Ad5ΔΔE1+pIX and Ad5ΔΔE1ΔpIX was compared by measuring luciferase expression. Transduction was performed in triplicate in 24-well plate wells in 500 μl DMEM/8% FBS. After a two-hour incubation the virus-containing medium was replaced with fresh medium without virus. At 24 h post transduction the cells were washed once with PBS and lysed in 100 μl LUC-lysis mix (25 mM Tris–phosphate (pH 7.8), 2 mM CDTA, 2 mM DTT, 10% glycerol and 1% Triton-X in PBS). Luciferase production was determined with the Promega Luciferase Assay by adding 25 μl luciferase assay reagent to 10 μl lysate. Light intensity measurement was performed in a Victor Wallac 2 microplate reader (PerkinElmer, Inc., Waltham, MA, USA).

Integrin blocking

Indirect blocking of integrin-mediated virus uptake was performed by incubating cells with EDTA. M22-MEL3.0 cells were harvested from semi-confluent tissue culture plates, washed three times in PBS with 5 mM EDTA, and resuspended in standard PBS or PBS supplemented with 0.9 mM CaCl2 and 0.5 mM MgCl2 (PBS++). The Ad5ΔΔE1+pIX and Ad5ΔΔE1ΔpIX stocks were adjusted to equal pp concentrations by adding sucrose buffer and were diluted 1:1 in a 5 mM EDTA solution in PBS. Virus (100 pp/cell) was added to 500,000 cells in 1 ml PBS or PBS++ and incubation was performed for 60 min at 37 °C under constant agitation. Subsequently, the cells were pelleted by centrifugation, dissolved in 5 ml medium and transferred to 24-well plate wells (500 μl per well). Cells were incubated for 24 h and analyzed for GFP expression by flow cytometry. Data were analyzed with CellQuest software (Becton Dickinson).

The anti-human CD51/61 monoclonal antibody (MAb LM609), an αVβ3 integrin antagonist, and MAb P1F6, an αVβ5 integrin antagonist (both obtained from Millipore) were used to test the inhibitory effect of anti-integrin antibodies on virus transduction. Cells grown as monolayers were pre-incubated with medium only or with medium containing integrin function-blocking MAb (10 ng/ml). After 30 min of incubation, the excess antibody was removed by gentle washing followed by virus transduction (100 pp/cell). Reporter gene expression was measured 24 h post transduction by performing a standard luciferase assay.

Virus incubation with coagulation factor X (FX)

HepG2 and A549 cells were plated in 24-well plate wells. After a PBS wash step, Ad5ΔΔE1+pIX or Ad5ΔΔE1ΔpIX (100 pp/cell) was added in serum-free medium containing 8 μg/ml Factor X (FX) (HCX-0050, Haemotologic Technologies Inc.), 8 μg/ml Gla-domainless Factor X (FXMUT) (HCX-GD, Haemotologic Technologies Inc.) or no FX/FXMUT. After 2 h the medium was replaced by normal medium. Luciferase expression was measured 24 h post transduction.

Immunoblot analysis and electron microscopy

Immunoblot analyses were performed to assess the incorporation of proteins into the capsid of Ad5ΔΔE1+pIX and Ad5ΔΔE1ΔpIX. The western blotting and detection procedures were described previously (de Vrij et al., 2008). Virus lysates were prepared by adding 5×109 virus particles directly to western sample buffer. Capsid proteins were visualized with rabbit polyclonal anti-protein IX serum (1:20000) (Caravokyri and Leppard, 1995), goat polyclonal anti-hexon (1:1000, ab19998, Abcam, Cambridge, UK), and mouse monoclonal anti-fiber (1:5000, 4D2, Abcam) (Hong and Engler, 1991). Secondary antibodies were horseradish peroxidase (HRP)-conjugated goat-anti-rabbit and rabbit-anti-mouse (Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA).

Electron microscopy was performed on Ad5ΔΔE1+pIX and Ad5ΔΔE1ΔpIX samples adsorbed into glow-discharged carbon coated copper grids and negatively stained for 30 s with 2% phosphotungstic acid (pH 7). The viruses were examined with a FEI Tecnai Spirit BioTwin transmission electron microscope operating at 120 kV. Images were recorded on a 4 k × 4 k Eagle CCD camera.

PBMC analysis

Buffy coats were obtained from healthy donors after consent (Sanquin Bloodbank, Leiden, The Netherlands) and centrifuged on a Ficoll gradient to obtain PBMC. PBMC (1 × 106) were added to a well of a 24-well plate in 0.5 ml medium (RPMI/10%FCS) and virus was added at the indicated MOI in 0.5 ml medium. After incubation at 37 °C and 5% CO2 for two days, the supernatants were isolated and frozen for interferon gamma (IFN-γ) measurement, and the cells were prepared for flow cytometry analysis. Cells were washed twice with PBS/0.02% sodium azide, fixed for 10 min in 4% paraformaldehyde, washed twice with PBS/0.5% BSA/0.02 sodium azide, and stained with antibodies. Antibodies used were anti-CD3-PerCP-Cy5.5, anti-CD4–PE, anti-CD14–APC, anti-CD14–PerCP-Cy5.5, anti-CD19–PE, anti-CD19–PerCP-Cy5.5, anti-CD69–PE and anti-CD86–PE (Becton Dickinson, Franklin Lakes, NJ, USA), anti-CD8-APC and anti-CD56-APC (Beckman Coulter, Brea, CA, USA). Activation of NK cells was evaluated as increased CD69 expression on CD3−, CD14−, CD19−, CD56+ cells. Monocyte activation was analyzed as increased CD86 expression on CD3−, CD19−, CD14+ cells. Fluorescence was measured by flow cytometry on a FACS Calibur (Becton Dickinson) and data were analyzed with CellQuest software (Becton Dickinson). IFN-γ in supernatants was measured by ELISA using the PeliPair reagent set for human IFN-γ (Sanquin, Amsterdam, NL).

Viral distribution after intravenous delivery

Ad5ΔΔE1+pIX or Ad5ΔΔE1ΔpIX (1010 pp) was injected in the tail vein of 6-week-old athymic nude mice (NMRI nu/nu; Taconic M&B A/S, Ry, Denmark), followed by sacrificing the animals and harvesting of multiple organs at 3 days post injection. Four hours before Ad5ΔΔE1+pIX and Ad5ΔΔE1ΔpIX injection, pre-dosing was performed with the empty vector HAdv-5.CMV (replication-deficient and not encoding a transgene) (5 × 1010 pp) to saturate Kupffer cell macrophages. Tissue samples from each organ were lysed in LUC–lysis mix and the luciferase expression was measured according to the Promega Luciferase Assay. The protein concentration in the lysates was determined by using the bicinchoninic acid protein assay (Pierce, Perbio Science BV, Etten-Leur, The Netherlands), enabling the calculation of luciferase expression per total protein. The experiment was performed under the Dutch Experiments on Animals Act that serves the implementation of “Guidelines on the protection of experimental animals” by the Council of Europe (1986), Directive 86/609/EC, and only after a positive recommendation by the Animal Experiments Committee.

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