Neutralized Adenovirus-Immune Complexes Can Mediate Effective Gene Transfer via an Fc Receptor-Dependent Infection Pathway

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Neutralization of adenovirus (Ad) by anti-Ad neutralizing antibodies in serum involves formation of Ad-immune complexes that prevent the virus from interacting with target cells. We hypothesized that Ad-immune complexes likely contain viable Ad vectors which, although no longer capable of gaining access to receptors on target cells, may be able to express transgenes in cells bearing Fc receptors for immunoglobulins, i.e., that antibody-based “neutralization” of Ad vectors may be circumvented by the Fc receptor pathway. To test this hypothesis, we expressed the Fcγ receptor II (FcγRII) in A549 lung epithelial cells or human dermal fibroblasts and evaluated gene transfer in the presence of human neutralizing anti-Ad serum. FcγRII-expressing cells bound and internalized copious amounts of Ad, with a distinct population of internalized Ad trafficking to the nucleus. The dose-response curves for inhibition of gene transfer revealed that FcγR-expressing cells required a more-than-10-fold higher concentration of anti-Ad serum to achieve 50% inhibition of Ad-encoded β-galactosidase expression compared with non-FcγR-expressing cells. The discrepancy between neutralization of Ad during infection of FcγR-expressing cells and neutralization of Ad during infection of non-FcγR-expressing cells occurred with either heat-inactivated or non-heat-inactivated sera, blocked by addition of purified Fc domain protein, and did not require the cytoplasmic domain of FcγR, suggesting that immune complex internalization proceeded via endocytosis rather than phagocytosis. FcγR-mediated infection by Ad-immune complexes did not require expression of the coxsackie virus-Ad receptor (CAR) since similar data were obtained when CAR-deficient human dermal fibroblasts were engineered to express FcγR. However, interaction of the Ad penton base with cell surface integrins contributed to the difference in neutralization between FcγR-expressing and non-FcγR-expressing cells. The data indicate that complexes formed from Ad and anti-Ad neutralizing antibodies, while compromised with respect to infection of non-FcγR-expressing target cells, maintain the potential to transfer genes to FcγR-expressing cells, with consequent expression of the transgene. The formation of Ad-immune complexes that can target viable virus to antigen-presenting cells may account for the success of Ad-based vaccines administered in the presence of low levels of neutralizing anti-Ad antibody.

One ubiquitous challenge regarding the use of viral vectors for gene transfer relates to the ability of immune-competent hosts to develop neutralizing humoral immunity against the viral capsid. In the case of adenovirus (Ad) gene transfer vectors based on subgroup C viruses, approximately one-half of the general patient population has a detectable neutralizing antibody titer (3, 10, 11, 19, 21, 21, 53, 61, 65, 68). The problem is exacerbated when one considers the impact of repeated administration of Ad vectors. Following each successive administration, the neutralizing antibody titer tends to increase and the efficacy of gene transfer decreases dramatically (10, 19, 33, 41, 42, 86).

Neutralization of Ad by antibodies has typically been described with one of two possible outcomes: intracellular neutralization or extracellular neutralization. Intracellular neutralization refers to an Ad-immune complex that enters the cell but fails to accomplish gene delivery to the nucleus. With purified antibodies against individual capsid proteins, several groups have demonstrated intracellular neutralization with accumulation of Ad inside organelles in the cytoplasm (12, 52, 55, 79–81). Extracellular neutralization, in which formation of Ad-immune complexes prevents Ad from interacting with target cells, is the predominant form of neutralization for unfractionated, anti-Ad sera (52, 74). However, it is not clear whether extracellularly neutralized Ad is necessarily neutralized with respect to intracellular trafficking. In other words, there may exist viable Ad capsids in extracellular Ad-immune complexes that could traffic to the nucleus and express viral genes if given the opportunity to interact with cells.

The relevance of this question lies in the fact that antigen-presenting cells express receptors for immune complexes and can internalize immune complexes via Fc receptors. The Fc receptor family includes both high-affinity and low-affinity receptors for the Fc portion of immunoglobulins (17). The low-affinity receptors (FcγRII and FcγRIII) clear immune complexes from tissue and serum and enhance the immune response to foreign antigens contained in the antibody-antigen complex (30). In the event that viable viral capsids gain entry to an antigen-presenting cell via Fc receptor interaction, there exists a potential viral gene expression with the antigen-presenting cell (14, 44), a mechanism that can lead to particularly strong immune responses to virus-encoded antigens.

With the knowledge that formation of Ad-immune complexes prevents Ad access to target cells, we hypothesized that Ad-immune complexes may contain viable virus that is capable of intracellular trafficking and infection if the Ad-immune complex is forced to interact with target cells via Fc receptors.
To address this hypothesis, Fcγ receptor IIA (FcγRII) was expressed in a nonhematopoietic target cell line. This strategy was based on the fact that this isoform of FcγRII was previously shown to contain an immunoreceptor tyrosine-based activation domain and thus was capable of directing phagocytosis of immune complexes (46). In addition, the use of nonhematopoietic and nonmyeloid cell lines avoided potential confusion from endogenous expression of FcγRII, a strategy previously validated in the literature (29, 30). Following formation of Ad-immune complexes with neutralizing anti-Ad serum, the intracellular trafficking of Ad capsids and Ad-mediated gene transfer were assessed over a range of ratios of Ad to neutralizing anti-Ad serum to determine the concentration of anti-Ad serum that gave 50% inhibition of gene transfer (IC50).

MATERIALS AND METHODS

Ad vectors. All of the Ads used in this study were E1–E3– replication-deficient, recombinant Ad gene transfer vectors with an expression cassette inserted into the E1 position. The expression cassette included the cytomegalovirus promoter-enhancer, a simian virus 40 polyadenylation signal, and a transgene. The study included vectors encoding a β-galactosidase transgene (Adβgal) (22), an FcγRII (CD122) transgenic (AdFcγRII) (8), a vector with a form of the Fc receptor that lacked the cytoplasmic domain (AdtalesisFcγR) (8), and a vector that was used as a control for vector infection into which no cDNA had been inserted (AdNull) (22). In addition to these vectors, one tropism-modified vector was employed. The vector was identical to Adβgal except that the nucleic acid sequence encoding the penton base protein was altered to modify the three amino acids (RGD) that confer integrin binding (AdRGDβgal) (76). All vectors were propagated, purified, and stored as described previously (56, 57). All vectors had particle-to-PFU ratios of less than 100. Virus concentration was determined in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum, 50 U/ml penicillin, and 50 μg/ml streptomycin at 37°C in 5% CO2. All cell culture reagents were obtained from Life Technologies, Inc. (Gaithersburg, MD).

Cell culture. A549 lung epithelial carcinoma cells (CCL-185; American Type Culture Collection, Manassas, VA) express the high-affinity coxsackievirus Ad receptor (CAR) (23) and were used as a model for CAR-sufficient cells. Human dermal fibroblasts are deficient in CAR expression (23). A549 cells were maintained in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum, 50 U/ml penicillin, and 50 μg/ml streptomycin at 37°C in 5% CO2. Fibroblasts were maintained in RPMI 1640 cell culture medium supplemented with 10% fetal bovine serum, 50 U/ml penicillin, and 50 μg/ml streptomycin at 37°C in 5% CO2. All cell culture reagents were obtained from Life Technologies, Inc. (Gaithersburg, MD).

Anti-Ad neutralizing sera, titer determination, and heat inactivation. Human neutralizing sera were obtained in the course of gene therapy clinical trials that have been previously described in detail (19). Sera were previously assayed for neutralizing anti-Ad titer by propagation of wild-type Ad5 infection in a monolayer culture of A549 cells and were stored at −80°C until use (19). Neutralizing titers of individual sera are described in the text according to the inverse of the lowest dilution that gave a 90% inhibition of adenovirus serotype 5 infection. Except where noted otherwise, the neutralizing titer of the anti-Ad serum was 2,560 (i.e., a 2,560-fold dilution was the lowest dilution to give 90% inhibition of adenovirus infection). Upon thawing, sera were heat inactivated at 55°C for 60 min prior to use, except where noted otherwise. This research was performed in compliance with the Health Information Privacy and Protection Act. This report does not contain information on the identities of patients from whom samples were collected.

Preinfection of A549 cells or human dermal fibroblasts with AdFcγRII. FcγR expression was obtained by preinfecting cells with AdFcγRII. For A549 cells, the preinfection conditions were 1,000 particles per cell, 60 min, 37°C. For human dermal fibroblasts, the preinfection conditions were 10,000 particles per cell, 60 min, 37°C. The use of a higher concentration of Ad to infect human dermal fibroblasts was previously shown to be required because of the absence of CAR (19, 39). Preinfected cells were prepared in parallel with uninfected cells (naive), as well as cells preinfected with AdNull as a control for vector infection. Where noted, preinfection was performed in the same manner with AdtalesisFcγR. After 3 h, preinfected A549 cells or fibroblasts were replated at a density of 104 cells per well on a 96-well plate for analysis of gene transfer or at a density of 105 cells per well of a coverslip bottom dish for microscopic analysis (19, 39).

Evaluation of Ad binding and FcγR expression. Ad-immune complexes were formed as previously described, by preparing serial dilutions of neutralizing sera (0.01, 0.03, 0.1, 0.3, 1, 3, and 10%) in binding buffer (modified Eagle’s medium without phenol red [Life Technologies], 1% bovine serum albumin [BSA; Sigma Chemical Co., St. Louis, MO], and 10 mM HEPES, pH 7.3 [BioFluids, Rockville, MD]) (19, 39). After preparation of the serial dilutions of serum, sufficient Ad was added to the mixture to yield a concentration of 1011–1013 Ad particles/ml. The mixture was then incubated at 37°C for 5 min. Prior to infection, the cells were washed three times with binding buffer. A 30-μl volume of the Ad-serum mixture was added to 104 cells in the well of a coverslip bottom dish. The infection proceeded for 10 min at 37°C. The unbound virus was then washed away (three times with binding buffer) and returned to complete cell culture medium. Cells were incubated for at least six well washes before fixation, two-phosphate-buffered saline (PBS), and fixed with paraformaldehyde (4% in PBS; 20 min, 22°C; Electron Microscopy Sciences, Inc., Fort Washington, PA). After fixation, cells were prepared for indirect immunofluorescence by blocking with 5% goat serum (Calbiochem, La Jolla, CA) and 1% BSA in PBS plus 0.1% Triton X-100 (Sigma Chemical Co., St. Louis, MO) for 20 min. Primary antibody (fluorescein isothiocyanate-conjugated anti-CD32; Pharmingen/BD Biosciences, San Diego, CA) or an irrelevant primary antibody (fluorescein isothiocyanate-conjugated anti-CD64; Pharmingen/BD Biosciences, San Diego, CA) was added to the cells at a 1:100 dilution in block for 60 min. Cells were washed with 1% BSA in PBS, and nuclei were stained with 4′,6-diamidino-2-phenylindole (DAPI; Molecular Probes, Eugene, OR) at a concentration of 1 μg/ml in PBS. Cells were imaged with a Nikon Microphot SA microscope with a 60× N.A. 1.4 PlanApo differential interference contrast objective and DAPI, fluorescein, and Cy3 filters; a Princeton Instruments cooled charge-coupled device camera; and MetaMorph image analysis software (Universal Imaging, Inc., Downingtown, PA) as previously described (19, 39).

Evaluation of Ad-mediated β-galactosidase gene expression. Monolayers of naïve cells, AdNull-infected cells, or AdFcγR-infected cells were infected with Adβgal in a manner identical to the method described for infection with Cy3Ad above, with the exceptions that Adβgal was substituted for Cy3Ad and that infection occurred in wells of a 96-well plate instead of a coverslip bottom dish. Each well was preinfected with at least six well washes. After infection, cells were incubated for 24 h and β-galactosidase transgene expression was evaluated in cell lysates by quantitative chemiluminescence detection (Tropix, Bedford, MA) and normalized to the protein concentration of the cell lysate with the bichiniconinic acid reagent (Bio-Rad, Hercules, CA). In A549 cells, the β-galactosidase activity measured in cells infected in the absence of neutralizing serum was, on average, approximately 3 × 105 relative light units/mg or 2,000-fold above the background. In fibroblasts, the β-galactosidase activity measured in the absence of neutralizing serum was 2 × 105 relative light units/mg or 100-fold above the background. To determine the IC50s for each condition, data were normalized to the control value for the data set (infection in the absence of serum). Normalized data from independent experiments were averaged to generate a dose-response curve for inhibition of Ad-mediated gene transfer by the serum. The data were used to generate a summary dose-response curve and were fitted with a spline (SigmaPlot for Windows; SPSS, Inc., Chicago, IL). The point at which the curve intersected a normalized transgene expression level of 0.5 (corresponding to 50% inhibition relative to the control) was taken as the IC50 for the experiment.

Competition with purified Fc domain. To test the dependence of Ad-immune complex-mediated gene transfer on interaction of Fc domains with Fc receptors, complexes of Adβgal with 0.1% neutralizing serum (titer of 2,560) were used to infect AdNull- or AdFcγR-infected A549 cells. Infections were performed in the absence or presence of the purified Fc domain at 1 mg/ml (Calbiochem). Infection conditions and analysis of β-galactosidase activity were performed as described above.

Viability. Because of reports of cytotoxic effects of Ad-immune complexes in the literature (34–37), the viability of cells was checked 1 h after the 10-min treatment with Ad-immune complexes created with dilutions of the serum with a titer of 2,560. Cell viability was assessed with two fluorophores, one that stains live cells (calcine AM) and one that stains dead cells (ethidium homodimer 1; Invitrogen Cytotoxicity Kit: Molecular Probes). Counts of live and dead cells were performed on fluorescence micrographs with automated image analysis software (MetaMorph; Universal Imaging).

Statistics. All data are reported as the mean ± the standard error. Student’s t test was used to compare conditions, with P values of <0.05 indicative of a significant difference.
RESULTS

FcγR-dependent cell association of Ad-immune complexes.

Immune complexes were formed by combining Cy3Ad with a range of concentrations of neutralizing serum (0.01% to 10%). The immune complexes were applied to A549 cells previously infected with either AdNull or AdFcγR. In AdNull-infected cells, cell-associated Ad decreased as the concentration of neutralizing serum increased, similar to data previously reported for naive A549 cells (74). At a concentration of 0.1% serum, cell-associated Cy3Ad was noticeably decreased and virtually...
all Cy3Ad was prevented from reaching the cells at a serum concentration of 1.0% (Fig. 1A). In contrast, A549 cells expressing FcγR maintained the ability to bind Ad-immune complexes at serum concentrations as high as 10% (Fig. 1B). Fluorescence intensity of cell-associated Cy3Ad indicated that the amount of viral capsid taken up by FcγR/H9253R-expressing cells exceeded the amount of Cy3Ad internalized by a CAR-dependent mechanism in naive A549 cells. One striking observation concerned the ability of Cy3Ad to traffic to the nucleus. The nuclear perimeter in Fc receptor-bearing cells was demarcated by Cy3Ad even when Cy3Ad was mixed with as much as 1.0% neutralizing serum, indicating that some of the capsids internalized as Ad-immune complexes were capable of trafficking in a fashion similar to that of naive Cy3Ad. When AdNull-infected cells were compared to AdFcγR-infected cells, both in the presence of 1% neutralizing serum, it was clear that much of the capsid associated with FcγR-expressing cells was neutralized with respect to AdNull-infected cells. When FcγR-bearing A549 cells were exposed to Cy3Ad-immune complexes in the presence of neutralizing serum, the amount of Cy3Ad found in the cytoplasm rather than at the nucleus increased as the concentration of neutralizing serum increased (Fig. 1B). The intense, punctate concentrations of Cy3Ad in the cytoplasm indicated that some of virus was unable to traffic to the nucleus.

To understand the significance of the subcellular localization of Ad following infection of FcγR-bearing cells with Ad-immune complexes, localization of the Ad capsid was compared to localization of the FcγRII protein. The amount of immune complexes associated with A549 cells corresponded to the level of FcγR expression in the cells (Fig. 2A). FcγRII-positive staining was found at the plasma membrane, as well as in a punctate pattern, typical of intracellular, endocytic vesicles that internalize and recycle the FcγR (Fig. 2B). The distribution of Cy3Ad included perinuclear accumulation, as well as a punctate pattern that colocalized with the punctate portion of FcγRII (Fig. 2B). Colocalization of Cy3Ad and FcγRII suggested that a portion of Cy3Ad failed to escape from FcγR-containing organelles. These data showed that anti-Ad neutralizing serum induced primarily extracellular neutralization at lower concentrations (e.g., 1% in the case of this serum) and induced complete intracellular neutralization only at higher concentrations (e.g., 10% in the case of this serum).

FcγR-dependent Ad-mediated gene expression. To confirm that trafficking of Cy3Ad corresponded to successful gene transfer, A549 cells were exposed to Ad-immune complexes that were formed with an Ad vector carrying the β-galactosidase transgene (Adβgal). In earlier in vitro studies using a different protocol for the formation of Ad-immune complexes, significant cytotoxicity was attributed to antibody-aggregated Ad (34–37). Differential cell viability could complicate the interpretation of the gene expression results. In the present studies, cell cultures were assessed for cell viability following infection with Ad-immune complexes. A549 cells maintained >90% viability regardless of the ratio of neutralizing serum to Ad particles (not shown).

The kinetics of neutralization were compared in naive A549 cells and A549 cells preinfected with either AdNull or Ad FcγR. Neutralization of Adβgal in AdNull-treated A549 cells was comparable to the kinetics of neutralization in naive A549 cells (Fig. 3). Ad-mediated gene expression was reduced by 50% when Ad was exposed to a low concentration of neutralizing serum (<0.1%). In contrast, AdFcγR-treated cells exhibited elevated levels of gene transfer to A549 cells and did not reach 50% neutralization until exposed to >1% serum (Fig. 3). Comparison of IC₅₀ measured in naive A549 cells and A549 cells preinfected with either AdNull or Ad FcγR showed that cells expressing the Fc receptor required a 37-fold higher concentration of this neutralizing serum in order to achieve 50% inhibition of gene transfer compared with control AdNull-infected cells (Fig. 3). The precise shift in the IC₅₀ is likely to be a unique property of a given serum and based on the precise combination of anti-Ad epitopes and anti-Ad isotypes in the serum.

To test whether the shift in IC₅₀ was generally related to the neutralizing titer of the anti-Ad serum, two other sera with lower and higher neutralizing titers were evaluated. As predicted, the IC₅₀ measured in naive A549 cells corresponded to
the anti-Ad titer of the serum. With the same dilution series of sera with a low or high anti-Ad neutralizing titer (undiluted titer of either 20 or 49,000), the observed IC\textsubscript{50} shifted correspondingly, indicating that the inhibition curve was governed by the anti-Ad antibodies rather than another serum component (Fig. 4). For each serum, the IC\textsubscript{50} for Fc\textsubscript{R} expressing cells was significantly higher than for AdNull-infected or naive cells. In fact, 50% inhibition did not occur in the serum with the lowest titer (undiluted titer, 20), even when the serum was present at the highest concentration (10%). The IC\textsubscript{50} for one serum (undiluted titer, 2,560), either with or without prior heat inactivation, were similar, suggesting that complement was not involved in the interaction of Ad-immune complexes with Fc\textsubscript{R}-expressing cells (Fig. 4).

Characterization of the role of the Fc receptor in gene transfer following uptake of Ad-immune complexes. To confirm the importance of the Fc-Fc receptor interaction in the rescue of Ad-immune complexes, the uptake of Ad-immune complexes was assayed in the absence or presence of the purified Fc domain, a competitive inhibitor of antibody-Fc receptor binding. AdNull-infected A549 cells supported a minimal level of transgene expression in the presence of 0.1% serum, regardless of the presence or absence of the purified Fc domain (Fig. 5). In contrast, the high level of gene transfer in Fc receptor-bearing A549 cells was reduced by more than 75% in the presence of the purified Fc domain, indicating the importance of the Fc-Fc receptor interaction in uptake of Ad-immune complexes.

Prior studies of Fc\textsubscript{R} have demonstrated that the cytoplasmic tail is required for phagocytosis of large ligands (29). In contrast, the cytoplasmic tail of endocytic receptors is normally

FIG. 3. Fc\textsubscript{R}-mediated gene transfer by neutralized Ad-immune complexes. An Ad vector expressing the β-galactosidase transgene (Ad\textsubscript{βgal}) was mixed with various concentrations of human serum (0.01% to 10%) containing anti-Ad neutralizing antibodies (titer, 2,560). After 5 min, the Ad\textsubscript{βgal}-serum mixture was applied to monolayers of naive A549 cells or A549 cells previously infected with either AdNull or AdFc\textsubscript{R}. β-Galactosidase activity was assayed after 24 h and normalized to cellular protein content (relative light units per milligram of protein). Data were normalized to Ad\textsubscript{βgal} infection in the absence of serum for each target cell type (dashed line at y = 1.0). The IC\textsubscript{50} was determined as the concentration of serum that resulted in a 50% decrease in β-galactosidase gene expression (dotted line at y = 0.5). Comparable dose-response curves were obtained in the presence of neutralizing serum during infection of either naive or AdNull-infected A549 cells. In contrast, addition of neutralizing serum during infection of AdFc\textsubscript{R}-infected A549 cells shifted the dose-response inhibition curve, resulting in a higher IC\textsubscript{50}.

FIG. 4. Effect of Fc receptor expression on IC\textsubscript{50} of anti-Ad sera. Ad-immune complexes were formed and applied to naive, AdNull-infected, or AdFc\textsubscript{R}-infected A549 cells with dilutions of sera with various titers, including titers of 20, 2,560, and 49,000. In addition, anti-Ad serum (titer, 2,560) was used to form Ad-immune complexes without prior heat inactivation of the serum. IC\textsubscript{50}s were determined as described in the legend to Fig. 3.
required for optimal internalization but is not necessary to accomplish endocytosis of small ligands (32, 50). To test whether the cytoplasmic domain is required, A549 cells were preinfected with Ad vectors carrying either full-length FcγR or a truncated version of the FcγR gene that lacked the cytoplasmic domain (AdtaillessFγR). When challenged with a range of doses of neutralizing serum, A549 cells expressing the full-length Fc receptor or the tailless Fc receptor had identical kinetics of inhibition, indicating that the cytoplasmic tail of the Fc receptor is not required for binding and entry of Ad-immune complexes into cells (Fig. 6).

Receptor dependence of FcγR-mediated rescue of neutralized Ad. A549 cells express both the Ad high-affinity receptor for the Ad fiber protein (CAR) and αv integrins, which serve as the lower-affinity receptor for the Ad penton base protein (23, 62). Since capsid-receptor interactions play a critical role during Ad escape to the cytosol (48, 49, 63, 77), the fiber-CAR interaction and the penton base-integrin interaction were investigated.

To evaluate the effect of loss of fiber-CAR interaction, CAR-deficient human dermal fibroblasts previously treated with AdNull or Ad FcyR were evaluated for uptake of Ad-immune complexes. Each condition was analyzed for viral trafficking with Cy3Ad-immune complexes and for gene expression with Adβgal-immune complexes. As expected, very little Cy3Ad bound to AdNull-infected fibroblasts, regardless of the presence of serum, since the fibroblasts lacked a high-affinity receptor for either the Ad capsid or immunoglobulins in the Ad-immune complexes. Ad FcyR-infected fibroblasts bound a small amount of Cy3Ad in the absence of serum, but in the presence of 0.1% neutralizing anti-Ad serum, copious amounts of Cy3Ad were associated with FcγR-bearing fibroblasts (Fig. 7). A smaller proportion of internalized Ad was observed at the nuclear envelope compared to A549 cells. This difference likely reflected the propensity of Ad to traffic to the microtubule organizing center, as well as the nucleus, in fibroblasts (5, 66).

To examine gene transfer via uptake of Ad-immune complexes in the absence of CAR, naive fibroblasts or fibroblasts previously infected with AdNull or Ad FcyR were infected with immune complexes formed with an Ad vector that carried the β-galactosidase transgene (Adβgal) (Fig. 8A). The most striking aspect of the neutralization curve for Fc receptor-bearing fibroblasts was the dramatic increase in gene transfer at low concentrations of serum. The presence of 0.3% neutralizing serum led to an increase in gene transfer to Fc receptor-bearing fibroblasts of 64-fold compared to gene transfer in the absence of serum. The dramatic increase in gene transfer reflected the introduction of a high-affinity immunoglobulin G-FcγR interaction that enabled Ad-immune complexes to bind to cells. The high level of gene transfer to Fc receptor-bearing fibroblasts was accompanied by a 6.5-fold increase in the IC50 of the neutralizing serum (Fig. 8B). The IC50 for AdNull-infected fibroblasts was 0.2% serum, while the IC50 for Ad FcyR-infected fibroblasts was 1.3% serum.

To evaluate the requirement for the penton base-integrin interaction during FcγR-dependent infection of A549 cells, Ad-immune complexes were created with either wild-type Ad capsid or an Ad capsid that had been genetically altered to remove the RGD integrin-binding motif in the penton base (AdΔRGDβgal) (76). A comparison of the kinetics of gene transfer following formation of Adβgal- or AdΔRGDβgal-immune complexes demonstrated striking differences in gene
transfer in the presence or absence of integrin interaction (Fig. 9A). In A549 cells expressing FcγR, different IC₅₀s were determined for the same serum, depending on the Ad vector that had been incorporated into the immune complex. Strikingly, when Ad-immune complexes were formed with AdΔRGDβgal, the IC₅₀ of the serum for inhibiting gene transfer to FcγR-expressing A549 cells (0.09%) was comparable to the IC₅₀ of immune complexes formed with Adβgal on non-FcγR expressing cells (0.03 to 0.06%) (Fig. 9B, black bars). These data differ from the result obtained when immune complexes were formed with Adβgal (Fig. 9B, white bars), suggesting that the penton base-integrin interaction is important for Ad-immune complexes to deliver viable Ad capsids to FcγR-expressing cells.

**DISCUSSION**

The experiments described herein were designed to answer the following question: is neutralized Ad in Ad-immune complexes no longer infectious? This question was based on the observation that increasing concentrations of neutralizing serum led primarily to extracellular neutralization, i.e., abrogation of viral interaction with cells (74). This observation left open the possibility that viable Ad capsids, as well as nonviable Ad capsids, were caught in the Ad-immune complexes but that the viable Ad capsids no longer had access to the target cell surface. To answer this question, an experimental system was designed to force Ad-immune complexes to associate with target cells. Target cells were genetically modified to express FcγR, a receptor for the Fc domain of immunoglobulins. While it is a formal possibility that some nonspecific effect of the exogenous membrane of a membrane protein accounts for the observations described here, the fact that the enhanced infection of Fc receptor-expressing target cells was abrogated by an excess of the purified Fc domain argues that it was, in

![FIG. 7. Association of Ad-immune complexes with CAR-deficient human dermal fibroblasts modified to express FcγR. Cy3Ad-immune complexes were prepared and applied to human dermal fibroblasts as described in the legend to Fig. 1. The target cells used included human dermal fibroblasts previously infected with either AdNull or AdFcγR. After a 10-min infection, unbound virus was removed by washing. Cells were incubated for an additional 60 min to allow Cy3Ad trafficking within the cell. After fixation, cell nuclei were stained with a fluorophore (DAPI) and the position of virus within the cell was evaluated by fluorescence microscopy. Note that 0.1% neutralizing serum was sufficient to prevent Ad association with AdNull-infected fibroblasts. However, with the same concentration of neutralizing serum, a high level of cell-associated Cy3-Ad was observed. Bar = 10 μm.](http://jvi.asm.org/)

![FIG. 8. FcγR-mediated gene transfer by neutralized Ad-immune complexes in CAR-deficient fibroblasts. To determine the contribution of the native high-affinity interaction of Ad with CAR during immune complex uptake, immune complexes were exposed to naive (CAR-deficient) fibroblasts or fibroblasts infected with AdNull or AdFcγR. Immune complex formation and evaluation of gene transfer were performed as described in the legend to Fig. 3. (A) Ad-mediated gene transfer as a function of the anti-Ad serum concentration in naive, AdNull-infected, and AdFcγR-infected A549 cells. (B) IC₅₀ for all of the conditions tested.](http://jvi.asm.org/)
fact, the Fc-Fc receptor interaction that led to immune complex uptake. The Fc-Fc receptor interaction is physiologically relevant in that this receptor is expressed on tissue macrophages and dendritic cells and leads to uptake and antigen presentation of opsonized materials in vivo. Historically, viral neutralization has been characterized on the basis of quantitative and qualitative analyses of the neutralizing serum. This study demonstrates that viral neutralization is also a function of the characteristics of target cells, potentially leading to new interpretations of viral neutralization in the complexity of an in vivo setting.

The data clearly demonstrated that uptake of Ad-immune complexes via the Fc receptor not only led to elevated viral gene expression but also effectively changed the neutralization properties (IC50) of the serum. It is important to note that the extent to which a given serum will show a change in its IC50 is likely to be determined by a number of factors, including the isotypes of antibodies against the virus and the number and identity of epitopes contributing to the antiserum. Unlike prior studies of viral neutralization by serum in vitro which employed extended incubations of virus with neutralizing serum prior to infecting cells, the present study intended to model the interactions that might occur upon administration of a gene transfer vector to a patient with preexisting anti-Ad immunity. With the knowledge that viable Ad capsids are quickly cleared from serum (2, 70, 83), an acute in vitro model of immune complex formation and target cell infection was designed. To mimic acute exposure of Ad to immune sera, followed by rapid access to target cells, the experimental protocol called for mixing of Ad and anti-Ad serum for only 5 min prior to infection of a monolayer of target cells for 10 min. The concentration of virus used for the infections (1011 particles per ml) is comparable to the dose of virus administered in clinical trials for local or systemic administration (19).

In the present study, no immune complex-induced cytotoxicity was observed. Kjellen and Ankerst used a protocol in which 2 h was allowed for Ad-immune complex formation with wild-type Ad and 2 to 6 h was allowed for immune complex exposure to cells (34–37). These investigators reported cytotoxicities as high as 100%, depending on the ratio of serum to Ad and the time of exposure of complexes to target cells. In contrast, the protocol employed herein limited the time of formation of Ad-immune complexes and the duration of exposure of complexes to target cells as described above. As a result, cell viability was >90% for all conditions. The observation that gene transfer occurred independent of the presence of the cytoplasmic tail of the Fc receptor underscored the likelihood that endocytosis rather than phagocytosis was sufficient to internalize the small Ad-immune complexes formed by this protocol. There exists a potential for formation of cytotoxic Ad-immune complexes in vivo, and importantly, Ad-immune complexes formed with wild-type, replication-competent Ad have been proposed as causative agents in the canine eye disorder anterior uveitis and in canine and human nephrotoxicity (1, 84, 85, 87). However, cytotoxicity due to the formation of Ad-immune complexes from replication-deficient gene transfer vectors has not been reported, and under the conditions used in this study, significant cytotoxicity was not observed.

A number of avenues of research have focused on the ability of antibodies to modulate viral infection. The effect of antibodies reported here confirms previous reports of antibody-enhanced infection in a number of viruses, including human immunodeficiency virus (HIV), coxsackie virus B4, and rhinovirus (6, 24, 25, 69). Unlike the observations reported here, antibody-enhanced infection by viruses that are tropic for FcγR-bearing cells did not constitute a change in the tropism of the virus. The observations presented here show that binding of neutralizing antibodies to Ad may shift the tropism of
Ad away from native target cells (CAR-expressing epithelial cells) toward a new target cell (FcγR-bearing antigen-presenting cells). This report is also distinct from studies in which nonneutralizing antibodies have been used to retarget Ad to FcγR-bearing cells (14, 44, 45), since modification of tropism was observed following treatment with whole, unfraccionated neutralizing serum rather than selected monoclonal antibodies. Finally, this study is distinct from studies in which human sera have been analyzed to determine the relative contribution of neutralizing antibodies against each of the major capsid proteins (12, 52, 55, 59, 60, 67, 68, 73, 75, 79) since those studies examined the effects of subsets of antibodies rather than whole, unfraccionated neutralizing serum on the infectivity of Ad. While this report does not test the hypothesis that Fc receptor-mediated uptake of Ad-immune complexes could be a means of propagating a wild-type infection in a host organism, it is interesting that persistent subgroup C Ad infections have been reported in lymphoid tissue of humans and mice (13, 15, 18, 31, 38, 51, 54, 58, 64, 72), where a high concentration of Fc receptor-bearing cells resides. Also of interest, follicular dendritic cells, T lymphocytes, and B lymphocytes are all capable of expressing Fc receptors (40, 71) and persistent Ad infections have been noted in patients with neutralizing anti-Ad antibodies (64). In contrast, these cell types have low or undetectable levels of CAR expression (26–28, 78, 82). Among the three lymphoid cell types, CD4+ CD8− CD3+ T lymphocytes appear to be the major harbor for Ad in lymphoid tissue and support low levels of viral replication, although a small but detectable population of Ad was detected in a CD4+ CD8− CD3+ population that included dendritic cells (16, 43). Garnett et al. (16) noted that the level of Ad DNA decreased with age in the study population, correlating with the age-related loss of Fc receptor expression on follicular dendritic cells (4). The latent infection by Ad in lymphoid tissue may then result from a complex interaction of viruses, anti-Ad antibodies, Fc receptor-expressing cells, and low rates of viral replication.

The observation that a virus might be capable of gene expression in FcγR-bearing cells when it has been effectively neutralized with respect to infection in other cell types may have important implications for Ad-mediated therapeutic gene transfer and, in particular, Ad-based vaccines. The observations in the present report suggest a pathway for antigen presentation in which a host with preexisting anti-Ad immunity is exposed to Ad. After the formation of Ad-immune complexes, FcγR-bearing antigen-presenting cells may phagocytose the complexes, leading to presentation of viral capsid proteins via major histocompatibility complex (MHC) class I. However, some small fraction of the virus may escape from the phagosome and productively infect the antigen-presenting cell, leading to presentation of viral proteins via MHC class I. The implication of cross-priming both in pathogenic viral infection and in the clinical setting of viral gene transfer deserves additional study. The idea that the class I and class II pathways could be activated in a host with preexisting anti-Ad neutralizing antibody titers further predicts that an Ad-based vaccine could be effective in a patient with an anti-Ad neutralizing antibody titer. Some prior reports support this hypothesis. Although expression of marker genes cannot be detected following systemic administration of Ad vectors in the presence of a neutralizing anti-Ad antibody titer (9, 19, 33, 41, 42, 86), neutralizing antibodies and cytotoxic T-lymphocyte responses that can confer protective immunity have been developed following delivery of gene transfer vectors under similar conditions (7, 20, 60, 68). In fact, in a recent analysis of data relating to the use of Ad vectors to deliver an HIV vaccine in a human clinical trial, the trial sponsor chose to stratify the data, with one group of patients carrying preexisting anti-Ad antibody titers of <200 while a second tier had anti-Ad antibody titers of ≥200. The group with the detectable but low anti-Ad antibody titer developed an immune response to the Ad-encoded HIV antigen (N. Chirmule, personal communication). Even though neutralizing antibody titers are likely to prevent therapeutic gene transfer to a target cell, gene expression in antigen-presenting cells mediated by Ad-immune complexes might provide sufficient stimulation to boost an immune response. The observation that Ad-immune complexes can be used to transfer genes to dendritic cells in vitro (45) also supports the possibility that this phenomenon could contribute to immune recognition of Ad-encoded gene products in vivo.

The analysis of Ad receptors that contribute to viral gene expression from Ad-immune complexes yielded important insights with implications for Ad-mediated gene therapy. FcγR-mediated infection by Ad-immune complexes formed with the native Ad capsid versus an AdRGD capsid in A549 cells demonstrated that the RGD sequence in the penton base was essential to observe the shift in IC₅₀. If uptake of Ad-immune complexes by antigen-presenting cells in vivo leads to cross-priming through viral gene expression, then use of an AdRGD capsid may limit MHC class I presentation in antigen-presenting cells and therefore might blunt the cell-mediated immune response against vector-encoded genes in patients with preexisting anti-Ad humoral immunity.

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