Adenovirus entry into cells, a quantitative fluorescence microscopy approach

Urs F. Greber, Michel Nakano & Maarit Suomalainen
Institute of Zoology
University of Zurich
Winterthurerstrasse 190
CH-8057 Zurich
Switzerland

Fax: (41) 1 635 6822
Tel: (41) 1 635 4841
ufgreber@zool.unizh.ch

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1. INTRODUCTION

Adenoviruses carry their DNA genome into post-mitotic nuclei of a variety of human cells, either within an organism (in vivo) or outside an organism in culture (ex vivo) (1). Recombinant adenoviruses are developed in many laboratories as gene delivery vehicles to treat hereditary and acquired human disorders of somatic cells (2; 3). Diseased lungs of cystic fibrosis patients have been pioneered for treatment with recombinant adenovirus vectors (4). Preliminary results are promising, but demonstrate that the disease has not yet been cured by the emerging gene transfer technology (5). One reason for limited success was that the transgenes were not expressed adequately in the diseased tissues, either due to low efficiency of virus delivery to the target cell or inefficient DNA import into the nucleus. In this chapter, we describe a quantitative method to determine transport of fluorescently labeled wild type adenovirus 2 to the nucleus of a model cell line, HeLa cells. This protocol should be directly applicable to recombinant adenoviruses in a variety of cell lines, including peripheral blood cells, fibroblasts, polarized epithelial cells and differentiated neurons.

1.1. Adenovirus structure

Adenoviruses contain at least 11 different polypeptides (6). A hierarchically organized network of interactions between capsid proteins encloses a linear double-stranded DNA genome of about 34 kb,
which is linked to the capsid via the internal protein VI (7). Individual building blocks of the major capsid protein hexon join with minor elements, such as the facet-stabilizing protein IIIa and the hexon-cementing protein IX, as well as the vertex proteins penton base and protruding fiber. Inside the nucleus of an infected cell, adenovirus DNA is packaged into the capsid together with condensing proteins V, VII and μ and cysteine protease L3/p23.

1.2. Internalization and penetration

Adenovirus type 2 uses a stepwise cell entry program (8). Its fibers attach to an epithelial cell surface receptor of the immunoglobulin gene family (9-11). In the case of hematopoietic cells, an alpha_beta integrin has also been reported as a high affinity receptor attaching to the virus penton base protein (12). In epithelial cells, fibers are shed from the particle and virus enters by receptor-mediated endocytosis via fibronectin-binding αvβ5 integrins binding to penton base (13; 14). After 10 min of internalization, most virus particles are found within endosomes. Penetration of individual virions across the endosomal membrane is facilitated by slightly acidic pH and appears to require binding of penton base to the αvβ5 integrin (15; 16). When added to cells in high amounts virions can also get into the cytoplasm in the absence of acidified endosomes (14). In the cytoplasm, the internal capsid protein VI is degraded by the reactivated viral protease and protein IX is dissociated from the capsid. Protease is reactivated by two signals, penton base binding to integrin receptors at the plasma membrane and the reducing cytoplasmic milieu. These triggers prepare the capsid for disassembly near the nuclear membrane (17).

1.3. Nuclear transport

All the proteins needed for virus assembly must enter the nucleus after synthesis in the cytoplasm. They either directly or indirectly contain nuclear targeting information, such as nuclear localization sequences (NLSs) (for review of nuclear import of proteins, see 18). It is possible that NLSs present on capsid protein(s) are involved in nuclear targeting of incoming virus particles. In addition, mechanical forces, generated by cytoplasmic motor proteins, may be required for virus localization to the nucleus. In any case, structurally altered adenoviruses reach the nuclear membrane and attach to the cytoplasmic side of the nuclear pore complex (14). Nuclear import of the DNA and associated proteins, such as protein VII, then occurs after DNA is dissociated from the capsid (17).
2. MATERIALS

Commonly used chemicals were from Sigma or Fluka. All stock solutions were made up in double distilled water and filtered through 0.2 µm polyether sulfon membrane filters (Semadeni, Switzerland). Working solutions were diluted into double distilled water before use.

2.1. Cell growth and virus purification

2.1.1. Cells and Chemicals

Cells for virus growth are available from American Type Culture Collection (ATCC), e.g., HeLa cells (cervical epithelioid carcinoma), HeLa S3 cells (cervical epithelioid carcinoma), KB cells (nasopharyngal carcinoma), 293 cells (transformed embryonal kidney cells) and A549 cells (human lung carcinoma).

100 mm dishes (Costar, cell culture grade, 78.5 cm²). Dulbecco’s modified Eagle’s medium (Gibco-BRL, DMEM) containing 7% Clone III serum (Hyclone), 1% non-essential amino acids, 1% glutamine and 100 units/ml streptomycin, 0.1 mg/ml penicillin (all from Gibco-BRL). Bovine serum albumin (BSA, Sigma, A-9418: 10% (w/v) filtered through 0.2 µm membrane). DMEM containing 0.2% BSA. 10% sodium dodecyl sulfate (SDS, molecular biology grade).

1 x phosphate buffered saline (PBS), prepared from a premixed salt stock (Gibco-BRL). 1 M Tris-HCl, pH 8.1. 5 M NaCl. 1 M MgCl₂. 22.4% (w/w) cesium chloride (CsCl, ultrapure grade): dissolve 60 g in 0.05 M Tris-HCl, pH 8 yielding a total of 267.8 g. 42.2% (w/w) CsCl: dissolve 72.5 g in 0.05 M Tris-HCl, pH 8 yielding a total of 171.8 g. Immediately before use, prepare CsCl step gradient for SW 55 rotor: 1.5 ml of 22.4% solution over 1.5 ml of 42.2% solution. Freon (1,1,2-Trichloro-1,2,2-
Trifluoroethane, sds, France), saturated with 1/3 vol of 0.01 M Tris-HCL, pH 8.0. 0.2 M Phenyl-methyl-sulfonyl fluoride (PMSF): 34 mg/ml in ethanol. This solution is stable at -20° C for a few days. Glycerol (autoclaved). 2% (w/v) uranyl-acetate. 2% (w/v) silicotungstic acid, pH 7 adjusted with 1 N KOH.

2.1.2. Hardware

Cell incubator, equilibrated to 5% humidified CO₂ atmosphere at 37° C. Rocking plate fitting into incubator. Ultracentrifuge, SW 55 (or SW 50.1) rotor and corresponding ultra clear thermoplastic tubes (Beckman). Table top centrifuge (Heraeus). Low speed centrifuge holding 50 and 15 ml screw capped plastic tubes. Pasteur pipettes, autoclaved. Vacuum dialysis apparatus with fitting 75 kD collodion membrane bag (Schleicher & Schuell). 0.45 µm sterile filters (Millex-GS filter, Millipore). 50 ml and 15 ml screw capped plastic centrifuge tubes. Spectrophotometer including quartz cuvette (e.g., Pharmacia Ultrospec 2000) for measurements at 260 and 280 nm. Transmission electron microscope (e.g., Zeiss EM 902A at acceleration voltage of 80’000) and palladium film and carbon-coated EM grids (SYNAP TEK G-2793, Plano, Germany). Sodium dodecyl sulfate polyacrylamide gel electrophoresis system (SDS-PAGE, Hoefer).

2.2. Plaque assay

A549 (human lung carcinoma) cells grown in 30 mm plastic dishes (Costar) as described in sections 2.1. and 3.1.. 5 x concentrated DMEM (made up from powder and stored at 4°C for less than 2 months protected from light, Gibco-BRL). BSA (Sigma, A-9418). 100 ml glass flask, autoclaved, 500 ml of ddH₂O, autoclaved. 1 x phosphate buffered saline (PBS), prepared from a premixed salt stock (Gibco-BRL). 37% formaldehyde: for working solution, dilute 10 x into 100 ml 1 x PBS. To prepare 100 ml of 1 x maintenance medium, mix 20 ml of 5 x DMEM, 10 ml Tryptose Phosphate Broth (Gibco-BRL),
9 ml sodium bicarbonate (2.25 g/100 ml, Gibco-BRL), 2 ml Clone III serum (Hyclone), 2 ml 1 M MgCl₂, 1 ml penicillin/streptomycin (10'000 units/10 mg per ml, Gibco-BRL), 2 ml L-glutamine (100 x, Gibco-BRL) and ddH₂O to 80 ml. This solution can be filtered through a 0.2 µm membrane and kept light protected in the cold for about 4 weeks. To prepare 100 ml of overlay agar medium, microwave 0.625 g agarose (sea plaque, low gelling, FMC) in a screw capped glass flask containing 20 ml of ddH₂O, until agar is completely dissolved (no sign of Schlieren). Equilibrate in 37°C water bath for about 1-2 hours and add to 80 ml of prewarmed maintenance medium. Mix well avoiding air bubbles and add immediately over cells. Cristal Violet: to obtain a 50 x stock solution, dissolve 0.25 g of cristal violet (Sigma) in 5 ml of ethanol. For working solution dilute into ddH₂O.

2.3. Fluorescent virus entry

2.3.1. Chemicals and cells

Micro BCA assay for protein determination (Pierce). CsCl purified adenovirus. Virus dialysis buffer consisting of 0.1 M sodium bicarbonate, 0.05 M sodium chloride, 1 mM MgCl₂ pH 8.2. Collodion membrane (75 kD cut off) and dialysis apparatus (Schleicher & Schuell). Texas-red-X (dissolved at 5 mg/ml in water-free DMSO, Molecular Probes). Freshly made 1.5 M hydroxylamine pH 8.5.

Cells grown on glass coverslips in DMEM-clone III serum placed in 24 well dish (Costar) in a humidified 37°C CO₂ incubator. Virus binding medium: RPMI-0.2% BSA containing 15 mM Hepes, pH is adjusted to 7.4 with concentrated NaOH. Virus internalization medium: DMEM-0.2% BSA. PBS. 3.3% para-formaldehyde made up in PBS according to standard procedures (19). 0.5 M ammonium chloride (NH₄Cl). PBS containing 2% 1,4,-Diazabicyclo[2,2,2]octane (DABCO). Nailpolish.
2.3.2. Hardware

Round glass coverslips (12 mm diameter, Assistent, Winiger, Switzerland) are washed in 1 N hydrochloric acid for 10 min (HCl), followed by three changes in ddH$_2$O, methanol and again ddH$_2$O. They are placed between 2 filter papers in a glass petri dish, autoclaved and dried in an oven at 60° C. Aluminium plate. Ice bucket. Wipping rocker kept at 4° C. Reichert-Jung Polyvar microscope (Merck, Switzerland) equipped with a 40 x oil immersion objective (numerical aperature 1.0), Nomarski differential interference optics (DIC) and a texas red filter set (excitation filter 530 to 585 nm, emission filter LP 615) linked to a Charge Coupled Device (CCD) video camera (Hamamatsu C5405, Hamamatsu Photonics, Germany). Macintosh computer on-line to the camera system loaded with Argus-20 imaging acquisition program (Hamamatsu Photonics, Germany) and Photoshop Version 3.05 (Adobe). NIH image analysis software (version 1.6): World Wide Web at http://rsb.info.nih.gov/nih-image/.
3. METHODS

3.1. Adenovirus purification

The following protocol was modified according to (20).

3.1.1. Procedure

1. Grow KB cells on ten 100 mm tissue culture dishes in DMEM containing 7% Clone III serum to
90% confluency corresponding to about $8 \times 10^7$ cells.

2. Infect cells with 1 to 5 plaque forming units per cell in 3 ml of DMEM-BSA on a rocking platform
at 37°C in a CO$_2$ incubator for 90 min.

3. Remove inoculum, add 7 ml of DMEM-7% Clone III serum and incubate in CO$_2$ incubator at 37°C until cytopathic effect (CPE) occurs about 3 days later. Make sure CPE is complete and cells are
detached from the dish.

4. Collect cells with a short Pasteur pipette and transfer to 50 ml screw capped centrifuge tube. Pellet
cells at 600 x g for 10 min. Wash pellet in 10 ml of PBS and transfer suspension to 15 ml centrifuge
tube. Pellet cells.

5. Resuspend cells in 0.01 M Tris-HCl, pH 8.1, 0.5 mM PMSF at $2 \times 10^7$ cells/ml and freeze-thaw 3
times in liquid nitrogen and a 37°C water bath. At this point, the preparation can be kept frozen at -70°C for several months.
6. Extract cells with an equal volume of Freon by gently vortexing and shaking by hand for 1 to 2 min. The suspension should become viscous and homogenous. Separate organic and aqueous phase by centrifugation at 1000 x g for 5 min at 4° C and collect upper aqueous phase.

6 a. To prepare stock virus, filter-sterilize aqueous phase through 0.45 µm disk filter and shock freeze aliquots in liquid nitrogen. Store at -70° C.

7. Layer upper aqueous phase on top of a CsCl step gradient and spin for 2 hours at 32’000 rpm in SW55 rotor at 4° C.

8. Collect virus band by carefully removing liquid from the top using a Pasteur pipette. Dilute virus (approximately 400 µl) with 0.01 M Tris-HCl, pH 8.1 to 2 ml and overlay over a second CsCl step gradient as described in step 7. Spin isopycnically for 18 hours at 32’000 rpm at 4° C.

9. Collect virus band as described in step 8.

10. Vacuum-concentrate virus in collodion membrane by dialysis against 0.01 M Tris-HCl, pH 8.1, 0.15 M NaCl, 1 mM MgCl₂. Observe approximately every 5 min and concentrate to 1 to 2 mg/ml (about 100 to 200 µl). Disconnect vacuum and continue dialysis for another 2 to 3 hours on ice. Change buffers at least twice. Use virus immediately or add glycerol to 10% and shock-freeze aliquots in liquid nitrogen. Store at -70° C up to several months.
3.1.2. Quality control

3.1.2.1. Optical measurements

The amount of optical particle units is determined by the absorbance of dissociated virus at 260 nm.

1. Dissolve a small aliquot of virus in 0.5% SDS, 0.02 M Tris-HCl, pH 7.4 at 37° C for 15 min with occasional shaking.

2. Place disrupted virus into quartz cuvette and measure absorbances at 260 nm (OD260) and 280 nm.

3. Determine virus concentration by multiplying OD260 value with the dilution factor and dividing by the extinction coefficient ($\varepsilon = 9 \times 10^{-13}$ OD ml cm virus$^{-1}$) as determined for adenovirus type 5 (21). Optically pure virus should have an absorbance ratio 260/280 of 1.33 to 1.45 (22). 1 OD260 unit equals $1.1 \times 10^{12}$ optical virus particles, based on the measurement that 13% of the virus dry weight is accounted for by DNA (23). 1 OD260 is equivalent to about 1.2 mg/ml protein. This value roughly corresponds to the amount of protein (0.8 mg/ml), determined in our Micro BCA protein assay (Pierce).

3.1.2.2. Plaque assay

Plaque assays determine the biological activity of a virus preparation. They are the most stringent criterium for virus functionality. Results are expressed as plaque forming units (PFU).

1. Seed A549 cells 48 h prior to infection into 30 mm plastic dishes and grow to about 95% confluency. Set up enough dishes to perform test in duplicates including 2 dishes as non-infected controls.

2. Prepare 10-fold serial dilutions of virus into 2 ml of DMEM-BSA, typically ranging from $10^{-5}$ to $10^{-9}$.

3. Remove medium from the cells, wash cells once with DMEM-BSA and add 0.5 ml of diluted virus. Incubate on a gently rocking plate in a humidified CO$_2$ incubator @ 37° C for 90 min.

4. Suck off inoculum and add 2 ml of overlay agar medium. Let agar solidify in the cold for 10 min and place dishes into humidified CO$_2$ incubator @ 37° C for 2 to 3 days.

5. Overlay with an additional 2 ml of overlay agar medium and incubate for another 3 to 4 days @ 37° C as above. Examine plaque formation by eye against a dark surface beginning at day 5 after infection.

6. When no more new plaques grow (e.g., at day 7), determine the number of plaques by staining cells with crystal violet. Carefully remove agar with a spatula (avoid scratching the cells). Fix cells in 3.7% formaldehyde-PBS for 10 min and stain cells with crystal violet for 5 min. Suck off solution and count plaques.

3.1.2.3. Electron microscopy and negative staining

To determine if isolated adenovirus is free of other virus particles, such as adeno-associated virus (AAV), the following procedure can be applied. All steps are carried out at room temperature.

1. Spot 5 µl of a 1:10 dilution (into 0.01 M Tris-HCl, pH 8) of stock virus onto a piece of parafilm foil.

2. Place a carbon coated plastified EM grid (glow discharged) upside down on virus droplet and absorb virus to the grid for 2 min.
3. Pick up grid with a pair of fine tweezers and remove excess liquid with a piece of Whatman filter paper.

4. Place grid upside-down on 5 µl of 2% uranyl-acetate and incubate for 20 sec. Alternatively, stain virus with 2% silicotungstic acid (made up in ddH2O, pH adjusted with 1 N KOH to 7) for 10 sec.

5. Blot off excess stain with filter paper and dry specimen for 1 hour.

6. Observe sample in the transmission electron microscope at 80 kV and 80’000 to 200’000 fold magnification.

**3.1.2.4. SDS-PAGE**

Virus homogeneity can also be tested by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) followed by Coomassie blue staining, e.g., as described (8; 24). Pure adenovirus is characterized by a predominant hexon band, proteins V and VII and fainter bands representing penton base, fiber, IIIa, VI, and IX (Fig. 1, lane 1).

**3.2. Quantitative subcellular analysis of incoming fluorescent virus**

**3.2.1. Texas-red and FITC labeling of adenovirus**

1. Dialyse purified adenovirus against 0.1 M sodium bicarbonate, 0.05 M sodium chloride, 1 mM MgCl₂ pH 8.2 using a 75 kD cut off collodion membrane as described in section 3.1.1., step 10.
2. Determine protein concentration using the Micro BCA assay according to the manufacturer’s conditions (Pierce).

3. To 0.4 ml of 0.8 mg/ml adenovirus add dropwise 8 µl of 0.5 mg/ml texas red (diluted from 5 mg/ml stock with dialysis buffer immediately before use). Incubate for 1 hour in the dark on a rocker at room temperature.

4. To block excess texas red dye, add 40 µl of freshly made 1.5 M hydroxylamine pH 8.5 and incubate for another hour as described in step 3.

5. Repurify virus on CsCl step gradient as described in section 3.1.1., steps 7 and 10.

6. Determine protein concentration using Micro BCA assay (Pierce) and freeze virus in aliquots of 0.4 mg/ml.

3.2.2. Cold-synchronized virus entry into cells

1. Two days before infection, seed HeLa cells (or any other cell line) on round glass coverslips in DMEM growth medium placed in a 24 well dish. At the day of infection cells should be 60 to 90% confluent.

2. Transfer 24 well dish to an aluminum plate coated with a wet kim wipe and kept on ice. Wash cells once with cold binding medium (RPMI-0.2% BSA, 15 mM Hepes-NaOH, pH 7.4).

3. Dilute stock texas-red labeled virus (typically 0.2 µl of 0.26 mg/ml per coverslip, approximately
1'000 virus particles per cell) into cold binding medium and add 0.2 ml of this dilution to each coverslip.

4. Bind virus to the cell surface in the cold on a gently rocking platform for 90 min.

5. Wash off unbound virus with cold binding medium and briefly add 0.5 ml of warm internalization medium (DMEM-0.2% BSA) to each coverslip. Remove medium and add fresh 0.5 ml of warm DMEM-0.2% BSA. Incubate for a given time in a 37°C CO₂ incubator.

6. Wash cells quickly in PBS at room temperature and fix immediately in 3.3% para-formaldehyde for 15 min at room temperature.

7. Quench unreacted para-formaldehyde with 50 mM NH₄Cl in PBS for 5 min at room temperature and wash briefly with PBS.

8. Mount slide in PBS containing DABCO and seal edges of coverslip with nailpolish.

9. For best results analyze within 24 hours.

3.2.3. Quantification of particles

1. Record a digital picture of a group of 2 to 4 cells with a CCD camera in DIC mode using a 40 x objective across the middle of the cells (examples see Fig. 2 a, c, e).

2. Transfer image as a TIFF file on-line to the hard disk of a personal computer.
3. Record the corresponding fluorescence picture using the texas red or fluorescein filter set at maximal
intensity settings integrating a total of 50 frames (examples see Fig. 2 b, d, f).

4. Repeat step 2..

5. Open the recorded TIFF file of the DIC image in the NIH image program.

6. Outline the cells and nuclei in plain white color using the “pencil” tool.

7. Set threshold such that only the white outlines are visible. Make image binary. Select all.

8. Load Macro “image_registration”.

9. Open the corresponding fluorescence image and enhance contrast using the built in automatic
software function under the “Process” menu.

10. Under “Special”, “define image to register” and the DIC outline image to “register”.

11. Use the “Blend” and “Divide” commands to merge the images.

12. Define the “Density Slice” such that all the fluorescent particles become colored and are selected.
Save image.

13. Outline the area of interest (nucleus or whole cell) using the “free hand” mode and determine the
number of fluorescent particles by the “Analyze Particles” function.
14. Import data into a statistics program (e.g., Microsoft’s Excel or Adelbeck’s KaleidaGraph) for further analysis and data presentation (example see Fig. 3).

15. Process TIFF files and arrange with Adobe Photoshop software program for printing, e.g., on a Fuji Pictography 3000 sublimation printer.

As can be seen in Figure 2, virus bound to the cell surface of HeLa cells in the cold is more or less evenly distributed over the cell. The ratio of virus particles over the nucleus compared to the cytoplasm (Nuc/Cyt) is about 0.29 (Fig. 3). After 10 min of virus internalization at 37° C, a qualitatively similar result is obtained, with a nuclear to cytoplasmic ratio of fluorescent viruses approximating about 0.32. After 60 min of internalization, a different picture emerges. The majority of viruses has now moved towards the nucleus. The nuclear to cytoplasmic distribution coefficient at 60 min of warming cells is about 1.28 (Fig. 3).

It is important to note that by conventional fluorescence microscopy it is not possible to determine how close to the nuclear membrane the virus particles in fact are. Confocal microscopy and thin section electron microscopy have been applied to resolve this question. The results will presented elsewhere demonstrating that the fluorescently-labeled viruses enter the cells and localize to nuclear pore complexes with similar efficiencies as unlabeled control virus (Greber et al. submitted). No aggregated viruses inside or outside the cells are detected by electron microscopy suggesting that each fluorescent dot represents a single virus particle. Thus, the method described here truely measures virus transport from the cell surface or endosomes across the cytoplasm towards the nucleus.
4. NOTES

1. Typical yields of virus vary from cell to cell and of course depend on the number of cells on the dish. We found that ten 100 mm dishes of KB cells (approximately $4 \times 10^7$ cells) yield about 1 mg virus.

2. Freon helps dissociating aggregated virions during virus extraction from the cells. To prevent virus aggregation after cell lysis, it is important not to overload the CsCl gradients. One SW50 tube should not contain more than 0.5 mg of virus. For long term storage, a concentration of 2 mg/ml virus should not be exceeded.

3. Check pH of the silicotungstic acid stock solution each time before use (pH will drop with time).

4. If the adenovirus preparation is contaminated with AAV, at least one strong band of AAV protein 3 (VP-3) shows up below the adenovirus IIIa/fiber bands in SDS-PAGE (data not shown, see also 25). AAV proteins 1 and 2 (VP-1 and VP-2) are less abundant than VP-3 and stain weakly in Coomassie blue.

5. Texas-red labeled virus prepared under these conditions contains maximally 2.4 molecules of texas red per hexon monomer. Texas red labeled virus has the same specific infectivity (per protein) as unlabeled virus as determined by plaque assay. This virus preparation remains active for months.

6. Fluorescein isothiocyanate (FITC) labeled adenovirus can be prepared using a similar protocol. Virus is dialyzed against 0.01 M sodium bicarbonate containing 0.14 M NaCl, 1 mM MgCl$_2$, for 2 hours and protein concentration is determined using Micro BCA assay (Pierce). The pH of the virus solution is adjusted to 9 by adding 1/5 of the volume 0.5 M sodium carbonate buffer, pH 9.0. Labeling is initiated by adding 16 µl of 0.5 mg/ml FITC solution (diluted immediately before use into 0.1 M sodium carbonate buffer, pH 9 from a 5 mg/ml stock solution made in DMSO) to 125 µl of 1.8 mg/ml virus suspension. Incubate for one hour in the dark on a rocker at room temperature. Treat with hydroxylamine and repurify virus on CsCl gradient as described above. Analysis of FITC labeled virus by SDS-PAGE and fluoroimaging indicated that 47% of the label is
incorporated into hexon, 13% into penton base, 18% into protein IIIa and fiber, 3% into protein VI, 7% into protein VII and 12% into protein IX (Fig. 1, lane 2).

7. PBS containing 0.1% sodium azide can be used instead of PBS to minimize bacterial contamination.

8. DABCO-containing mounting media only lasts a few weeks and should be made fresh on a regular basis. An alternative mounting medium is based on NPGT (N-Propyl-gallate-Glycerol-Tris). For 25 ml, mix 17.5 ml glycerol (87%), 7.5 ml 0.1 M Tris-HCl pH 9.5, 2.5 g N-propyl-gallate in a 50 ml screw cap plastic tube. Place tube into water bath sonicator (e.g., Branson Model 1210) and sonicate for 15 min at 37° C. Transfer solution into vacuum flask and degas extensively under house vacuum. Store aliquots at -20° C. NPGT medium is recommended for texas red- or rhodamine-labeled probes, but appears to quench fluorescein-labeled probes.

9. A third mounting medium consists of 80% glycerol, 20 mM Tris-HCl, pH 8.8, 0.5% Paraphenylene-diamine (Sigma). This medium has a slightly brownish color and lasts several months if stored at -20° C. It works equally well with both, Texas red and FITC probes.

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5. FIGURES

**Fig. 1:** SDS 10-15% polyacrylamide gel electrophoresis (PAGE) of FITC-labeled adenovirus type 2. 8 µg of FITC-labeled virus was denatured in SDS-sample buffer at 95°C for 3 min and fractionated by SDS-PAGE. FITC-containing protein bands were immediately analyzed on a fluoroimager Model 575 (Molecular Dynamics) at 488 nm excitation wave length using the 530 DF30 band pass emission filter (Fluorogram). The relative fluorescence intensities of each protein band were determined by the ImageQuant software (Molecular Dynamics). Results are expressed as % of total FITC labeling (n.d. not detected). The gel was then stained with 0.1% Coomassie brilliant blue in 20% methanol, 5% acetic acid and digitized by a UTA-IE scanner (UMAX Data System Inc, Taiwan, R.o.C.). The result is shown on the right side (Coomassie).
Fig. 2: Entry of texas-red labeled adenovirus into HeLa cells. Virus was bound to the cell surface in the cold for 90 min (a,b) and internalized for 10 min (c,d) or 60 min at 37°C (e,f). Nomarski DIC optics was used to visualize the cells (Nom). Virus was detected by fluorescence microscopy using the Texas red filter set (Ad-tx red). The focus of the microscope objective was set to the middle of the cells. The apparent depth of field was estimated to be in the order of several µm (see also 26). Outlines of cells and nuclei in Figures b, d, f were generated with the NIH image software program.
Fig. 3: Summarized data of cell entry of texas red-labeled adenovirus particles. Images, as presented in Fig. 2, were analyzed for subcellular localization of fluorescent virus particles. Results are expressed as the mean ratio of nuclear to cytoplasmic viruses (Nuc/Cyt Ad). Standard deviations are indicated by error bars including the number of cells analyzed for each experiment (n).
6. REFERENCES


