Gutted adenoviral vector growth using E1/E2b/E3-deleted helper viruses

Catherine Barjot1†
Dennis Hartigan-O'Connor1,2,3
Giovanni Salvatori1‡
Jeannine M. Scott1,3
Jeffrey S. Chamberlain1,2,3*

1Department of Human Genetics, University of Michigan, Ann Arbor, MI, USA
2Program in Cellular and Molecular Biology, University of Michigan, Ann Arbor, MI, USA
3Department of Neurology, University of Washington School of Medicine, HSB Room K243, Box 357720, Seattle, WA 98195-7720, USA

*Correspondence to: Dr Jeffrey S. Chamberlain, University of Washington School of Medicine, Department of Neurology, Box 357720, 1959 N.E. Pacific Street, Seattle, WA 98195-7720, USA.
E-mail: jsc5@u.washington.edu
†Current address: UMR INRA 703, ENVN – Atlanpole La Chantrerie, BP 40706, 44307 Nantes Cedex 3, France.
‡Current address: Department of Immunology, Sigma-Tau, 00040 Pomezia, Italy.

Abstract

Background Helper-dependent, or gutted, adenoviruses (Ad) lack viral coding sequences, resulting in reduced immunotoxicity compared with conventional Ad vectors. Gutted Ad growth requires a conventional Ad to supply replication and packaging functions in trans. Methods that allow high-titer growth of gutted vectors while reducing helper contamination, and which use safer helper viruses, will facilitate the use of gutted Ad vectors in vivo.

Methods Replication-defective helper viruses were generated that are deleted for Ad E1, E2b and E3 genes, but which contain loxP sites flanking the packaging signal. Complementing Ad packaging cell lines (C7-cre cells) were also generated by transfecting 293 cells with the Ad E2b genes encoding DNA polymerase and pre-terminal protein, and with a cre-recombinase plasmid.

Results We show that C7-cre cells allow efficient production of gutted Ad using ΔE1+ΔE2b+ΔE3 helper viruses whose growth can be limited by cre-loxP-mediated excision of the packaging signal. Gutted Ad vectors carrying ~28 kb cassettes expressing full-length dystrophin were prepared at high titers, similar to those obtained with E2b+ helpers, with a resulting helper contamination of <1%.

Conclusions These new packaging cell lines and helper viruses offer several significant advantages for gutted Ad vector production. They allow gutted virus amplification using a reduced number of passages, which should reduce the chances of selecting rearranged products. Furthermore, the residual helper contamination in gutted vector preparations should be less able to elicit immunological reactions upon delivery to tissues, since E2b-deleted vectors display a profound reduction in viral gene expression.

Keywords adenovirus; gutted adenovirus; helper virus; Duchenne muscular dystrophy; dystrophin; gene therapy

Introduction

Gene therapy applications of adenoviral vectors have been extensively explored. Despite great progress, shortcomings in vector production, immunogenicity and cloning capacity have imposed significant limitations on their use in vivo [1]. The development of helper-dependent (hdAd) or “gutted” adenoviral vectors, whereby the sequences and regulatory elements of a therapeutic transgene are flanked by the minimal adenovirus sequences required for replication and packaging, holds great promise to overcome...
Materials and methods

C7-cré cell lines

C7 cells [18] were co-transfected with 10 μg of pOG231 (expressing cré recombinase; kindly provided by Dr. S. O’Gorman; [19]), and 1 μg of pcDNA3 (for neomycin resistance; Invitrogen). Cré-recombinase expression from pOG231 is driven by the CMV promoter and a synthetic intron. Prior to transfection, the plasmids were linearized with PvuII and NotI, respectively. C7 cells were transfected using the calcium phosphate-DNA precipitation method and stable transfectants were selected for and subcloned by colony isolation in medium supplemented with 1 μg/ml G418. All reagents for cell culture were purchased from GibcoBRL. Isolated cell lines were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum in the presence of 5 U/ml each penicillin and streptomycin, plus 0.5 μg/ml G418 to maintain cre expression and 100 μg/ml hygromycin B for Ad Pol and pTP expression.

Cre-recombinase activity in the subcloned lines was initially compared following transient transfection with pGKintβgal. This plasmid contains a polyadenylation site flanked by loxP sites located between the PGK promoter and the β-galactosidase (β-gal) cDNA. In the presence of cre recombinase, the internal polyadenylation sequence is excised, allowing expression of β-gal. The relative levels of cre-recombinase activity in each cell line were estimated by monitoring β-gal expression after transfection with pGKintβgal [20]. Clones displaying the highest β-gal expression were further subcloned and expanded for additional analyses.

Construction of helper viruses

The human placental alkaline phosphatase (hpAP) cDNA and SV40 large T intron and poly(A) site were subcloned from pRSVhAPT40 (a gift from Dr. Gary Nabel) downstream of the edcsyne-inducible promoter in pIND (Invitrogen). The promoter-cDNA cassette was then excised and inserted into pABS.4 (Microbix) along with a 1.8 kb EcoRI stuffer fragment isolated from human dystrophin intron 45 (J. S. Chamberlain, unpublished). The resulting PacaI fragment (which included a kanamycin-resistance gene derived from pABS.4) was inserted into the E3 deletion site of pBHG10 (Microbix), then the resistance gene derived from pABS.4 was inserted into pABS.4 (Microbix). Cre-recombinase expression from pOG231 is driven by the CMV promoter and a synthetic intron. Prior to transfection, the plasmids were linearized with PvuII and NotI, respectively. C7 cells were transfected using the calcium phosphate-DNA precipitation method and stable transfectants were selected for and subcloned by colony isolation in medium supplemented with 1 μg/ml G418. All reagents for cell culture were purchased from GibcoBRL. Isolated cell lines were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum in the presence of 5 U/ml each penicillin and streptomycin, plus 0.5 μg/ml G418 to maintain cre expression and 100 μg/ml hygromycin B for Ad Pol and pTP expression.

Cre-recombinase activity in the subcloned lines was initially compared following transient transfection with pGKintβgal. This plasmid contains a polyadenylation site flanked by loxP sites located between the PGK promoter and the β-galactosidase (β-gal) cDNA. In the presence of cre recombinase, the internal polyadenylation sequence is excised, allowing expression of β-gal. The relative levels of cre-recombinase activity in each cell line were estimated by monitoring β-gal expression after transfection with pGKintβgal [20]. Clones displaying the highest β-gal expression were further subcloned and expanded for additional analyses.
LoxP sites were added on either side of the packaging signal by ligating synthetic double-stranded oligonucleotides containing loxP sites into the AdIII (0.5 mu) and BglIII (1 mu) sites of plasmid pAdBglIII to form pAdloxPBglIII. 10 µg of this ‘flxed’ left-end shuttle plasmid were linearized by digestion with Nhel and co-transfected into C7 cells with 1 µg of Clal-digested viral DNA isolated from adenoviral strains dl7001 or dl7001ΔPol to create viruses AdloxP(+)/Pol7 and AdloxPΔPol7. DNA (1 µg) from each of these two viruses was then digested with EcoRI and AseI and separately co-transfected with 1 µg of BclI-digested AdL60 genomic DNA into C7 cells to generate the two helper viruses Ad(+)/loxP(+)/PolAP and Ad(+)/loxPΔPolAP (Figure 1A). Sixteen hours post-transfection, the cells were passaged and replated with fresh, untransfected C7 cells and distributed into 24-well plates as described previously [13]. Cells that displayed evidence of virus-induced cytopathic effect (CPE) were: MCK forward, 5′-CGCAACGAGAAGCTATGTCCAA-3′; MCK reverse, 5′-GCTTGTAATCCTGCTCTTCCTT-3′; L2 probe, 5′-VIC-CAGGTCATCGCGCCGGAGATCTA-TAMRA-3′; L2 forward, 5′-CGCAACGAGAAGCTATGTCCAA-3′; L2 reverse, 5′-GCTTGTAATCCTGCTCTTCCTT-3′; probe, 5′-VIC-CAGGTCATCGCGCCGGAGATCTA- TAMRA-3′. Prior to PCR amplification, virion capsids were destroyed by digestion with proteinase K at 37°C for 1 h, followed by inactivation at 95°C for 20 min. Serial dilutions of the digested samples were used as template in the PCRs and were compared with standards of known quantity. All samples were amplified and fluorescence of the reporter dyes recorded using the Applied Biosystems 7700 sequence detection system.

Quantitative PCR

Purified virus stocks were quantified by real-time PCR, using primers and a probe for the mouse muscle creatine kinase (MCK) gene promoter for the gutted virus, and for the Ad L2 gene for the helper. Primer sequences were: MCK forward, 5′-CGCAACGAGAAGCTATGTCCAA-3′; MCK reverse, 5′-GCTTGTAATCCTGCTCTTCCTT-3′; MCK probe, 5′-VIC-CAGGTCATCGCGCCGGAGATCTA- TAMRA-3′; L2 forward, 5′-CGCAACGAGAAGCTATGTCCAA-3′; L2 reverse, 5′-GCTTGTAATCCTGCTCTTCCTT-3′; L2 probe, 5′-VIC-CAGGTCATCGCGCCGGAGATCTA- TAMRA-3′. Prior to PCR amplification, virion capsids were destroyed by digestion with proteinase K at 37°C for 1 h, followed by inactivation at 95°C for 20 min. Serial dilutions of the digested samples were used as template in the PCRs and were compared with standards of known quantity. All samples were amplified and fluorescence of the reporter dyes recorded using the Applied Biosystems 7700 sequence detection system.

Colorimetric assays to determine viral transducing unit titer

Infected cell lysates were diluted in 500 µl complete medium containing 5 µM ponasterone A (Invitrogen), an analogue of ecdysone, and used to infect monolayers of Ecr-293 cells (Invitrogen) in a poly-L-lysine-coated 24-well tissue culture plate (Biocoat). After 16 h, the medium was removed, the cells were washed twice with phosphate-buffered saline (PBS) and fixed with 0.5% glutaraldehyde in PBS, and stained overnight for either alkaline phosphatase or β-gal expression, as described [3]. The number of cells that stained positively for β-gal or alkaline phosphatase activity was used to calculate viral titers as transducing units per ml (tu/ml).

Figure 1. Structure of helper and gutted viruses. (A) Structure of helper adenoviruses indicating features that are relevant to this study: the viral DNA packaging signal (Ψ) is flanked by loxP sites ([+]/loxP), the human placental alkaline phosphatase (AP) reporter gene is driven by an ecdysone-inducible promoter, and the E1 and E3 regions are deleted. Dyst-intron refers to a 1.8-kb fragment from human dystrophin intron 45 that was used as a stuffer. Also shown is the location of the viral DNA polymerase gene ([+]/loxPΔPol7). DNA ([+]/loxPΔPolAP (see text); the same vector lacking the Pol gene is referred to as Ad([+]/loxPΔPolAP. The total size of the helper viruses is ∼36 kb. (B) GE Dys is a 28-kb gutted adenovirus containing a 21-kb expression cassette composed of the full-length 13.9-kb murine dystrophin cDNA regulated by the 6.5-kb mouse muscle creatine kinase (MCK) promoter/enhancer, an ecdysone-inducible β-gal reporter gene, and one loxP site between the left ITR and the viral DNA packaging signal (G. Salvatori, M. Hauser, J. S. Chamberlain et al., manuscript in preparation).
Southern analysis of cre-recombinase activity on viral DNA substrates

C7 and C7-cre cells were infected in parallel with equal volumes of cell lysate or banded virus from Ad(+)/loxP(+)PolAP infected C7 cells. After 48 h, total viral DNA was extracted by the method of Hirt [21], digested with HindIII, and separated on a 0.8% agarose gel. The resolved fragments were transferred to a nylon membrane and hybridized with a 32P-dCTP by random-priming. Digestion of wild-type viral DNA generates a fragment of 3371 bp which contains the viral DNA packaging signal. Cre-recombinase-mediated excision of the sequences between the loxP sites, which includes the packaging signal, results in a band of 3100 bp. Hybridized bands were visualized using a phosphorimagery and quantified with Imagequant software (Molecular Dynamics).

Growth of gutted virus

To initiate production of the gutted adenovirus GEβDys (Figure 1B; G. Salvatori, M. Hauser and J. S. Chamberlain, in preparation), C7-cre cells were plated at 2.5 × 10⁶ cells/ml in a 60-mm dish and cultured to 70% confluency. The cells were then transfected with 10 µg of pGEβDys (the GEβDys viral genome contained in a plasmid backbone) in the presence of 0.1 mM chloroquine. After 4 h at 37 °C, the cells were subjected to a 30-s shock with 15% glycerol in 1× HEPES-buffered saline, pH 7.05. The cells were immediately washed twice with warm PBS and then incubated at 37 °C in tissue culture medium. At 18 h post-transfection, the cells were infected with either Ad(+)/loxP(+)/PolAP helper virus at a multiplicity of infection (moi) of 1 or Ad(+)/loxPΔPolAP helper virus at a moi of 5. The cells were harvested when they displayed complete CPE. (usually 10–15 days later for C7-cre8.2 cells, and 5–7 days for C7-cre8 cells) and gutted and helper virus titers were determined using the β-gal or alkaline phosphatase assays described above. Cell lysates containing infectious gutted virus were then used in subsequent passages for virus amplification and to determine optimal conditions for gutted Ad vector growth. The amount of lysate necessary to infect more fresh cells was calculated according to the titers of the gutted and helper viruses. For the first passage, one-third of the lysate was typically used to infect a fresh 60-mm dish of C7-cre cells to keep the moi of the input gutted virus as high as possible. For subsequent passages, an input gutted virus moi of 1–2 tu/cell was used. For each passage, purified helper virus was added to the lysate to achieve a final input helper virus moi of 5, except for the final passage, for which helper virus was added at a moi of 3. After each round of amplification, the cells and medium were harvested when the cells displayed complete CPE, usually 5–7 days after inoculation (for C7-cre8.2 cells; 3–5 days for C7-cre8 cells).

Purification of gutted virus

After three or four serial passages, the yield from a typical gutted virus preparation approached 10¹¹ tu in a crude cell lysate. When complete CPE was observed in the final passage, the cells were collected from the plates, pelleted at 150 g for 10 min at 4 °C and resuspended in two cell pellet volumes of 10 mM Tris-HCl, pH 8.0, 1 mM MgCl₂. Cell membranes were disrupted by three rounds of freeze/thaw followed by three rounds of sonication, for 1 min each, using a model 550 sonic dismembrator (Fisher Scientific) at level 5 with a microprobe. The lysate was then clarified by centrifugation at 1500 g for 10 min at 4 °C. The supernatant was stored on ice while the resulting pellet of cell debris was resuspended in 10 ml of 10 mM Tris-HCl, pH 8.0, 1 mM MgCl₂, and subjected to three additional rounds of sonication. The supernatants from the clarified lysates were then pooled and treated with 50 µg/ml each of DNaseI and RNaseA to degrade unpackaged viral and cellular nucleic acids. Both gutted and helper viruses were co-purified on a CsCl step gradient of 25% 1.4 g/ml CsCl and 50% 1.3 g/ml CsCl. The remaining 25% included the prepared virus solution brought to 1.1 g/ml with CsCl before layering onto the gradient. The gradients were centrifuged at 53 000 g for 4–16 h at 4 °C in a Beckman SW-28 rotor. The gutted and helper viruses were not separated on this gradient but were subsequently resolved on a self-forming gradient of 1.34 g/ml CsCl centrifuged for 12 h at 320 000 g for 12 h followed by 73 000 g at 4 °C in a Beckman MTV65 rotor. The viral band was then pulled from the gradient using an 18-gauge needle, and in the case of gutted virus, further purified from helper virus through one or two additional self-forming gradients. Purified virus was dialysed against 10 mM HEPES buffer, pH 7.6, containing 5% sucrose. Viral titers were determined using the colorimetric and real-time PCR assays as described above.

Results

Generation of loxP-containing, polymerase-deleted helper viruses

The C7 cell line is a derivative of the adenovirus packaging cell line 293 that expresses the E2b-encoded polymerase and pre-terminal proteins of AdS [14,18,22]. As a result of the additional adenoviral proteins, C7 cells produce adenovirus by efficiently rescuing supercoiled or linear viral genome templates [15]. C7 cells also allow growth of polymerase-deleted adenoviral vectors [13]. For enhanced production of helper-dependent or “gutted” adenoviral vectors, we further modified the C7 cells by making them capable of site-specific excision of the DNA packaging signal in first-generation or helper viruses, thereby increasing the yield and simplifying purification of gutted viral vectors. For this site-specific excision, we chose the cre-loxP system [6].
Figure 1A illustrates the basic features of the various helper viruses we generated. AdL60 is an E1/E3-deleted virus carrying an edcysone-inducible alkaline phosphatase expression cassette. We also generated four viruses that contain loxP sites flanking the packaging signal in an E1/E3-deleted backbone [(+)-loxP], and which either contain or lack the Ad DNA polymerase gene [(+)-Pol or APol] and the inducible alkaline phosphatase reporter cassette (AP). Most of the studies described below utilized either AdL60, Ad(+)-loxP(+)-PolAP or Ad(+)-loxPΔPolAP (Figure 1A; see ‘Materials and methods’).

We next sought to determine whether our insertion of loxP sites adjacent to the viral DNA packaging signal would have any detrimental effects on the replication of the virus. We compared the viral output from C7 cells infected with a virus that either contained, or lacked, loxP sites. Figure 2 shows that the production of the (+)-loxP virus is nearly identical to the (−)-loxP virus. We concluded that the insertion sites for the loxP sequences did not impair viral replication in C7 cells.

Isolation of C7-cre cells

C7 cells were stably transfected with a cre-recombinase expression construct and subcloned as described in ‘Materials and methods’. The initial transfectants were screened for their ability to excise a loxP-flanked fragment of DNA present in pGKint such that it blocks expression of the β-gal reporter gene. Proper excision of the internal fragment enabled expression of the β-gal cDNA, leading to the production of quantifiable β-gal. C7-cre subclones that produced appreciable β-gal activity by this assay were expanded for additional analysis (data not shown).

The ability of these subclones to excise the loxP-flanked viral packaging signal was evaluated in two ways: Southern analysis of viral DNA from infected C7-cre cells, and comparison of relative viral titers produced in C7-cre vs. C7 cells. For DNA analysis, the cells were infected with Ad(+)-loxP(+)-PolAP for 48 h followed by total DNA extraction, digestion and Southern analysis. We observed that the virus produced by the C7 cells yielded DNA with a left-end fragment of the predicted full-length size (3371 bp) that includes the packaging signal (Figure 3A). In contrast, the C7-cre clones produced viral DNA with a shorter left-end fragment (3100 bp), indicating that the packaging signal had been excised. This experiment also revealed that the individual C7-cre subclones varied in their ability to excise packaging signals. For example, quantitative analysis of the relative intensity of the bands indicated that clones #8 and #25 were more efficient (>95%) than clone #15, which excised the packaging signal from only ~35% of the viral genomes. Clone #8 was further subcloned, and after initial testing with the pGKintβgal expression system, four new sublines were compared with C7 and C7-cre8 cells by infection with purified virus and Southern analysis. All four of the selected sublines excised the packaging signal with >99% efficiency (Figure 3B).

The presence of active cre recombinase in the C7-cre cell lines should lead to a reduction in the yields of packaged (+)-loxP viruses. To examine viral yield we performed parallel infections of C7 and the C7-cre cell lines with CsCl-purified Ad(+)-loxP(+)-PolAP virus. For these studies all cell lines used had undergone a similar number of passages since their transfection with the Ad Pol and pTP constructs. The cells were infected at a moi of 3 after reaching 90% confluency in a 60-mm dish. After 48 h, cell lysates were prepared and viral yields determined using a colorimetric assay for alkaline phosphatase. The results showed a dramatic reduction in viral yields in the cre-recombinase-expressing cells. Cell line #25 produced the most virus (4.1%), while line #8.2 produced the least (0.4%), compared with the yield obtained in C7 cells (Figure 3C). These results correlated well with the levels of excision seen in the Southern analysis studies, indicating that the cre-expressing cells efficiently reduce the amount of virus produced when that virus contains a packaging signal flanked by loxP sites.

To verify that this reduction in viral yield was due to the cre-mediated excision of the packaging signal and not to a general reduction in their ability to produce virus, we compared C7 and C7-cre cells for the ability to produce virus independent of the cre-loxP system. The cells were infected at a moi of 3 with AdL60, which is identical to the Ad(+)-loxP(+)-PolAP virus except that it lacks loxP sites (Figure 1A); 48 h post-infection, the amount of virus produced by C7-cre cells was not different from the amount produced by C7 cells (Figure 3D).
Growth of gutted Ad vectors using C7-cre cells

The C7-cre cell line #8 (C7-cre8) was subsequently used to optimize a procedure for amplification of gutted adenoviruses. To simplify titering, we used the gutted adenoviral vector GEβDys, which contains a β-gal cDNA under the control of the edcsyone-inducible promoter, as well as a full-length dystrophin expression cassette (Figure 1B; G. Salvatori, M. Hauser and J. S. Chamberlain, in preparation). Expansion of this virus can therefore be monitored using the EcR-293 colorimetric assay system similar to that for the alkaline phosphatase reporter gene in our helper viruses. Gutted and helper virus production can be assessed in parallel following each round of amplification. Furthermore, GEβDys contains a loxP site between the L-ITR and the packaging signal. This loxP site ensures that the packaging signal of the helper virus retains two loxP sites in the event that homologous recombination occurs between the sequences of the gutted and helper virus packaging signals [7].

To determine the optimal moi of both helper and gutted viruses that would maximize the yield of gutted virus in the C7-cre cells, we evaluated viral yield after infecting 60-mm dishes with varying amounts of each virus, but always using a helper to gutted transforming unit ratio (H/G) of either 5 or 1. The gutted virus GEβDys was derived from a CsCl-purified stock from which the integrity and helper contamination were checked by Southern and quantitative PCR analyses, respectively; the helper virus was added from a CsCl-purified stock to achieve the desired moi. Figure 4 shows the viral yields from these infections and demonstrates that maximum yields of gutted virus were produced when the input helper virus moi was 5 times greater than that of the gutted virus (H5/G1, H25/G5, or H50/G10). An exception was noted when the total input virus moi was 300. In this case, the yields of helper and gutted viruses were equal suggesting that the cre-recombinase levels were insufficient to process efficiently such large amounts of input virus. It is interesting to note that, although the overall yields were lower with a H/G ratio of 1.0 then with a ratio of 5.0, equal input ratios of the gutted and helper viruses achieved the best relative yield of the two vectors, except at the highest input moi tested (50). These data again suggested that the total viral load had exceeded the ability of the C7-cre cells to excise efficiently the viral packaging signal. Based on these data, we selected an input moi ratio of 5 (H/G) for subsequent experiments.

To examine growth of the gutted virus during serial passages, the gutted plasmid pGEβDys was transfected into C7-cre8 cells and subsequently infected with the helper virus Ad(+)/loxP(+)/PolAP (see ‘Materials and
The gutted virus GEβDys was recovered, or rescued, from this plasmid transfection (termed passage zero or P0) as a cell lysate and subsequently expanded to 100 x 150 mm tissue culture dishes during three passages (Figure 5A). The amount of gutted virus increased with each of the first two passages (P1 and P2), then reached a plateau between $10^7$ and $10^8$ tu/ml, while the amount of helper virus rescued after each passage remained between $10^6$ and $10^7$ tu/ml. Following P3, the (+)loxP helper virus titers were 3–4% that of the gutted virus. Purification of the gutted virus through two continuous CsCl gradients reduced the helper contamination to less than 1% (the lowest we have achieved is 0.2% as determined by real-time PCR). It is possible that the total amount of gutted virus could be increased even beyond this yield since a lower input moi of gutted virus (moi 1 vs. 5) results in similar yields in the lysate, thus allowing one to infect more dishes starting with a given stock of virus or lysate (Figure 4). Similar results were observed for the C7-cre8.2 subline (data not shown).

Figure 5B shows the theoretical yield if all of the infected lysate generated in the first two passages were used at the maximum dilution to infect additional plates at an optimal moi. The lysate produced in P2 typically yields enough gutted virus to inoculate 200 x 150 mm dishes of C7-cre cells (at a gutted moi of 2) with a potential yield of ~$2 \times 10^{11}$ β-gal transforming units (tu). In summary, the use of the C7-cre cell lines with a helper virus carrying loxP sites on each side of its packaging signal not only reduces the helper virus contamination obtained with C7 cells, but also reduces to only 3–4 the number of passages necessary to reach the virus production plateau. This limit has been reported previously to be approximately $10^7$ tu/ml of gutted virus with Cre-293 cells [6], whereas the C7-cre8 lines routinely support plateau growth at >$10^8$ tu/ml (Figures 5A and 6A).

**Growth of gutted Ad vectors with an E1-, E2b- and E3-deleted helper virus**

The gutted Ad system described above allows vectors to be produced with helper virus contamination levels of <1%, as assayed by quantitative, real-time PCR. Nonetheless, if very high doses of gutted Ad vectors were to be delivered in vivo, the residual first-generation Ad helper could lead to undesired immunological consequences. We have shown previously that the C7 cell line is able to support efficient, high-titer growth of replication-defective E2b-deleted Ad vectors that display a profound reduction in...
viral gene expression both in vitro and in vivo [13]. E2b-deleted helper viruses also display increased persistence and reduced cellular toxicity in vivo [17]. We therefore explored whether Ad vectors with deletions of the E2b gene encoding DNA polymerase could support high-titer production of gutted Ad vectors in C7-cre cells. For these studies we constructed a helper virus that contains loxP sites flanking the packaging signal and which lacks the polymerase gene, Ad(+)/loxPΔPolAP (Figure 1A). This helper virus was used to expand GEβDys in C7-cre cells. Using a helper virus moi of 5, the amount of gutted virus rescued from the transfection/infection with this ΔPol helper virus was ∼5 × 10^4 tu/ml (Figure 6A). This yield was 10-fold lower than the titer of gutted virus produced with the helper virus that contained the Pol gene, suggesting that the ΔPol helper was less able to support gutted Ad growth, possibly because it displayed less robust replication. Indeed, these results are consistent with burst assay experiments that showed a 55% reduction in titer of ΔPol helper virus compared with the (+)Pol helper virus when grown in C7 cells (data not shown).

Despite the slight growth disadvantage of the ΔPol helper virus, titers of nearly 10^8 tu/ml of gutted virus were achieved after three serial passages, similar to yields obtained with the (+)Pol helper (Figure 6A). In addition, the theoretical yield following expansion of the entire lysate would be the same, >10^11 tu (Figure 6B).

### Discussion

Gutted adenoviral vector technology is a promising approach to reducing the immunological barriers that have limited the use of conventional Ad vectors for in vivo gene transfer. The lack of viral genes combined with a large cloning capacity enables gutted vectors to deliver transgene products in tightly regulated or tissue-specific patterns, reducing the propensity to activate immune effector cells [5,23–29]. Nonetheless, the need to use a helper virus to grow a gutted vector complicates the use of this system by introducing low levels of viral gene contamination. In addition, the requirement for serial passaging to obtain high-titer stocks of vector raises the risk of vector rearrangement during growth. The modified helper viruses and packaging cell lines we describe here provide several important improvements to gutted vector technology that should help reduce these concerns by enabling robust growth of vectors in C7 cells using E2b-deleted helper viruses whose growth can be controlled by cre recombinase.

Since C7 cells stably express the E2b genes Pol and pTP, they should allow the use of a helper virus deleted for either, or both, genes. We have shown that a helper virus deleted for the Ad DNA polymerase gene can support high-titer gutted Ad vector growth (Figure 6). Although slightly less efficient than a polymerase-positive helper virus, the ΔPol helper virus was still able to generate at least 10^11 tu (∼10^13 particles) of gutted virus in only three serial passages. Titers of 10^7 tu of gutted vector/ml in crude cellular lysates were generated in only two passages, in contrast to the previous need for six to seven passages reported by us in 293 cells, and by Parks et al. using Cre-293 cells and a polymerase positive helper virus [3,6]. We feel the use of a ΔPol helper can offer several significant advantages. Polymerase-deleted Ad vectors have been shown to be replication-defective in the absence of a trans-complementing cell line [13]. This replication defect results in a 4–5 log reduction in viral late gene expression, and enables a significant prolongation of vector retention in vivo as well as reduced toxicity when delivered to the liver [13,17]. In addition, multiply deleted Ad vectors have been shown to display a significantly lowered propensity to recombine with viral genes in packaging cell lines to generate replication-competent genomes [30–33]. This latter risk can be further reduced by inserting sequences into the E3 region.
to make the helper genome too large to be packaged if it acquires E1 sequences from the packaging cells [6]. The ΔPol helper we describe in Figure 1 also incorporated this feature.

The levels of Pol and pTP expressed by C7-cre cells also support efficient conversion of plasmid substrates into replicating Ad genomes [15], and high-titer viral yields can be obtained starting with both linear and plasmid-embedded versions of the helper vector [5]. The resulting improvements in gutted virus titer following initial transfection/infections (P0) enables infection of packaging cells with the optimal input moi of gutted vector (between 1 and 5) as early as P1 or P2. Since cre-recombinase expression prevents helper virus propagation, an input moi of 1–5 for the gutted vector is critical for maximal yields as it ensures that most of the cells infected with helper virus are co-infected with gutted vector. The ability to infect cells with optimal titers early in the preparations results in high-titer gutted vector yields with fewer serial passages, not only allowing faster production of gutted virus, but also reducing the number of rounds of vector growth, thereby reducing the opportunities for a vector to rearrange or recombine. In theory, it may be possible to reduce the number of passages required for gutted virus growth even further. We have recently shown that conversion of gutted vector DNA into replicating viral DNA can be enhanced by performing the transfection using gutted DNA that has been covalently modified to contain terminal protein at the ends of the genome [16]. This form of the substrate is in a conformation that more closely models the optimal template necessary for initiation of viral DNA replication.

Regardless of whether the helper contains or lacks the polymerase gene, it appears difficult to achieve gutted virus titers higher than 10^8 tu/ml of lysate. In our experience, the titer of the gutted virus increases only slightly as the input gutted moi is increased from 1 to 5. The use of an input gutted moi higher than 5 does not increase the gutted titer and can lead to reduced yields, even with the use of a higher helper moi. In contrast to the growth of a first- or second-generation adenovirus, gutted virus amplification requires a fine balance between the gutted and the helper ratios. We suspect that too much gutted virus inhibits replication of the helper virus by competing with cellular factors necessary for viral DNA replication. Increasing the amount of helper virus in parallel with the input gutted moi also does not lead to increased yields since high helper loads lead to a cellular cytopathic effect before efficient replication of the gutted vector can occur.

Using the cre-loxP system, the main concern with gutted vector growth is the generation of a packagable helper virus that would contaminate the final gutted virus preparation. One event that can give rise to packagable helper virus is a recombinant that leads to replacement of the packaging signal of the helper by the packaging signal of a gutted vector that has no loxP sites. This possibility has been shown to be reduced considerably by either placing a loxP site between the left ITR and the packaging signal of the gutted virus [7] or by limiting contiguous homology between the two packaging signals by replacing the eight ambiguous nucleotides within each A-element with sequences taken from a different A-element [8]. In the gutted virus shown in Figure 1, we added a single loxP site, although there is no reason that the packaging signal of our ΔPol helper viruses could not also be modified to reduce potential recombination.

The system we have described here thus incorporates novel and established technologies into reagents for gutted virus production that should allow transfer of a variety of genes and regulatory elements into cells in vivo. We have used this system to produce gutted adenovirus vectors able to transduce skeletal muscle of dystrophin-deficient mdx mice with full-length (426 kDa) dystrophin resulting in physiological improvement (C. DelloRusso, C. Barjot, J. M. Scott, J. S. Chamberlain et al., submitted). The large cloning capacity of these vectors, combined with the additional safety and utility afforded by the C7-cre system and E2b-deleted helper viruses, may further increase the potential for gene therapy of a variety of diseases.

Acknowledgements

We thank Nuria Morral, Christiana DelloRusso, Robert Crawford, and Michael Blankenship for helpful discussions. Supported by grants from the National Institutes of Health (AI15434) and the Muscular Dystrophy Association (USA) (to JSC). CB was supported in part by a post-doctoral fellowship from the Association Française Contre les Myopathies.

References


17. Hu H, Serra D, Amalfitano A. Persistence of an [E1-,


