In Vitro Replication of Adeno-Associated Virus DNA

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The study of eukaryotic viral DNA replication in vitro has led to the identification of cellular enzymes involved in DNA replication. Adeno-associated virus (AAV) is distinct from previously reported systems in that it is believed to replicate entirely by leading-strand DNA synthesis and requires coinfection with adenovirus to establish completely permissive replication. In previous work, we demonstrated that two of the AAV nonstructural proteins, Rep78 and -68, are site-specific endonucleases and DNA helicases that are capable of resolving covalently closed AAV termini, a key step in AAV DNA replication. We have now cloned the AAV nonstructural proteins Rep78, Rep68, and Rep52 in the baculovirus expression system. Using the baculovirus-expressed proteins, we have developed an efficient in vitro AAV DNA replication system which mimics the in vivo behavior of AAV in every respect. With no-end AAV DNA as the starting substrate, the reaction required an adenovirus-infected cell extract and the presence of either Rep78 or Rep68. Rep52, as expected, did not support DNA replication. A mutant in the AAV terminal resolution site (trs) was defective for DNA replication in the in vitro assay. Little, if any, product was formed in the absence of the adenovirus-infected HeLa cell extract. In general, uninfected HeLa extracts were less efficient in supporting AAV DNA replication than adenovirus-infected extracts. Thus, the requirement for adenovirus infection in vivo was partially duplicated in vitro. The reduced ability of uninfected HeLa extracts to support complete DNA replication was not due to a defect in terminal resolution but rather to a defect in the reinitiation reaction or in elongation. Rep78 produced a characteristic monomer-dimer pattern of replicative intermediates, but surprisingly, Rep68 produced little, if any, dimer replicative form. The reaction had a significant lag (30 min) before incorporation of $^{32}$P-deoxyribonucleoside triphosphate could be detected in DpnI-resistant monomer replicative form and was linear for at least 4 h after the lag. The rate of incorporation in the reaction was comparable to that in the simian virus 40 in vitro system. Replication of the complete AAV DNA molecule was demonstrated by the following criteria. (i) Most of the monomer and dimer product DNAs were completely resistant to digestion with DpnI. (ii) Virtually all of the starting substrate was converted to heavy-light or heavy-heavy product DNA in the presence of bromo-UTP when examined on CsCl density gradients. (iii) Both flip and flop orientations of the terminal repeat were present in the product DNA, suggesting that each of the terminal repeats underwent at least two rounds of terminal resolution. The availability of an efficient in vitro AAV DNA replication assay should help to identify cellular proteins required exclusively for leading-strand DNA synthesis.

Much of what is known about eukaryotic DNA replication has come from in vitro studies of the replication of simian virus 40 (SV40), a small DNA virus which relies largely on cellular replication enzymes for amplification (5, 18, 20, 21, 30, 33, 49, 50, 52–56). Adeno-associated virus (AAV) is a single-stranded human DNA virus whose genome size is approximately 5 kb (43; see references 1 and 31 for reviews). Like SV40, AAV appears to rely on cellular enzymes for DNA replication (1, 3, 17, 31, 45). However, unlike SV40, AAV appears to use exclusively leading-strand DNA synthesis (Fig. 1) (2, 9, 10, 24, 25, 46, 47). Thus, the development of an efficient in vitro AAV DNA replication system may be useful for identifying cellular enzymes that are used primarily for leading-strand synthesis. In addition, AAV is unusual in that coinfection with a helper virus (either herpesvirus or adenovirus (Ad)) is essential for establishing a completely permissive environment for viral amplification (1, 31). In the case of Ad, which has been studied extensively, it appears that the helper virus is used primarily for the induction of AAV or cellular genes that are required for AAV replication (1, 3, 17, 31, 45). The Ad DNA polymerase and terminal protein are not required for AAV DNA synthesis, and the Ad DNA-binding protein, although serving an essential helper function, appears to be required primarily for efficient AAV gene expression (3, 17, 45).

Two AAV genes are required for viral DNA replication. The first is the viral origin of DNA replication (9, 25, 37, 39), which consists of a 145-bp terminal repeat (TR). The second is the rep gene, which codes for four nonstructural proteins that are synthesized from a single open reading frame by the use of alternate promoters and splicing (1, 31, 43). Viruses with mutations in the two larger proteins, Rep78 and Rep68, are defective for viral DNA replication (11, 48), while mutations in the smaller Rep proteins, Rep52 and Rep40, do not affect the production of replicative-form (RF) DNA (4). Like other paroviruses, the AAV genome replicates by a self-priming strand displacement mechanism (2, 9, 10, 24, 25, 35, 46, 47). The first step in DNA replication is the conversion of the single-stranded input genome into a linear duplex molecule in which at least one of the ends of the molecule is covalently joined. This is accomplished by using the terminal AAV palindrome as the primer for synthesis of the complementary strand. The covalently joined end is then converted into an open duplex end by a process called terminal resolution or hairpin transfer (Fig. 1) (2, 42, 47). A consequence of terminal
resolution is that the terminal AAV sequence is inverted (24, 25, 35, 42). Following resolution, the ends of the molecule are believed to reform the terminal hairpins, thus providing a 3' OH primer for strand displacement synthesis (reinitiation [Fig. 1]). Elongation from the hairpin primer then generates a single-stranded genome (which is presumably packaged) and a new RF molecule which again can undergo terminal resolution. Several other types of RF molecules can theoretically be formed during AAV replication. One of these is a dimer RF which presumably is formed when strand displacement synthesis proceeds through a terminal sequence that has not yet resolved (Fig. 1). The two major forms of RF species found in vivo are monomer and dimer duplex molecules (46).

In previous work, we have purified Rep68 to apparent homogeneity from AAV-infected cells and have partially purified Rep78 and Rep52 (15, 16). In addition, we developed several in vitro assays to characterize the biochemical activities of these proteins. For a substrate, we used a linear AAV DNA molecule in which both ends were covalently joined (Fig. 1), called no-end (NE) DNA (42). We demonstrated that both Rep78 and Rep68 were capable of correctly resolving a hairpinned end when supplemented with an uninfected HeLa cell extract (15, 16, 40, 42) to provide DNA polymerase activity. In the absence of a HeLa cell extract, both Rep78 and Rep68 could bind to the AAV terminal hairpin and cut at the terminal resolution site (trs). The trs endonuclease reaction was ATP dependent and was both site specific and strand specific, resulting in a singly nicked product that contained a molecule of Rep covalently attached at the trs site (15, 16, 40). Finally, both enzymes were found to have an ATP-dependent DNA helicase activity (15, 16). In contrast, Rep52 had neither terminal resolution nor trs endonuclease activity, and we could detect no specific binding to the hairpin (14, 16).

Although these experiments provided a plausible, direct role for the two larger Rep proteins in DNA replication, they were not successful in producing net DNA synthesis in vitro. Two other laboratories have reported the replication of AAV DNA in vitro. Yalkinoglu et al. (59) reported that a circular plasmid containing a single AAV terminal repeat replicated as a circle in an in vitro assay supplemented with crude extracts from Chinese hamster or mouse cells. DNA synthesis in this system was dependent on the presence of an AAV TR and was stimulated by pretreatment of the cells with a carcinogen. This was consistent with in vivo studies which showed that transformed cells treated with carcinogens became partially permissive for AAV replication (38, 57, 58). However, the absence of a requirement for viral Rep proteins and the production of circular rather than linear intermediates suggested that this system did not accurately reflect the conditions of fully permissive AAV replication. Hong et al. (12) have also described an in vitro replication system in which the substrate was a circular plasmid containing a complete copy of AAV. This assay produced linear replicative intermediates but was relatively inefficient, producing only a limited amount of replication over an extended period of time. In this article, we describe the development of an efficient in vitro AAV DNA replication assay using NE DNA as the substrate. The reaction mimics the in vivo behavior of AAV DNA replication in virtually every respect.

MATERIALS AND METHODS

Cells, plasmids, and virus. Cells, plasmids, and virus stocks were prepared and maintained as previously described (16, 29). Mammalian cells were routinely tested for latent AAV proviruses as described elsewhere (29). SPf insect cells, wild-type baculovirus, and the baculovirus transfer plasmids described below were obtained from M. Summers (Texas A&M University) and maintained as described elsewhere (22, 23).

Substrates and chromatography materials. NE substrate DNA was prepared from pusb201 plasmid DNA as previously described (36, 42). Ribonucleoside triphosphates were purchased from Promega or Sigma. They were dissolved in water and neutralized with NaOH. Radioactive nucleotides were purchased from ICN. Creatine phosphate, creatine phosphokinase, and dithiothreitol (DTT) were purchased from Sigma and dissolved in water. Restriction and DNA modifying enzymes and a bacteriophage DNA were from New England Biolabs. Molecular weight protein standards were from Bio-Rad. DEAE cellulose and phosphocellulose were purchased from Whatman; mono-S and mono-Q fast protein liquid chromatography columns were purchased from Pharmacia.

Baculovirus expression vectors. AAV Rep52 was expressed from a recombinant baculovirus containing AAV nucleotides 964 to 4493 which constituted the BclI-to-XbaI fragment from the previously described plasmid pIM29 (28). The plasmid used to generate this recombinant was derived from pNM348, which contained the XbaI fragment of pIM29 in the polylinker XbaI site of the baculovirus transfer vector pAC610 (22, 26). pNM348 was subsequently grown in Escherichia coli dam dem, and the BclI-to-XbaI fragment was recloned into BamHI- and XbaI-digested pAC610 to generate a baculovirus transfer construct in which the first ATG start codon under the control of the polyhedron promoter initiates translation of the p19 rep gene product. This plasmid, 348Bcl (Fig. 2), was cotransfected with purified wild-type Autographa californica nuclear polyhedrosis virus DNA into SF9 insect cells, and occlusion-negative recombinant virus plaques were isolated. Expression
of full-length Rep52 was confirmed by immunoblotting with Rep-specific polyclonal and monoclonal antibodies (13, 15).

The Rep78 baculovirus construct contained AAV nucleotides 321 to 4493. Oligonucleotide directed mutagenesis (19) was used to delete sequences between the left-end XbaI site and the start codon for Rep translation in plM29 to create plasmid 29dl-320 (28). The XbaI fragment from this plasmid was ligated into the BamHI site of the baculovirus vector pVL941 (23) via XbaI linkers to make the transfer plasmid 941Xp5M (Fig. 2). Recombinant baculovirus was generated as described above except that recombients containing Rep78 sequences were isolated by probing dot blots of end point dilutions of virus with oligonucleotide 5'-ATCTCGTAAAAAC CCCGGCAT, which is specific for the region encoding the NH₂ terminus of Rep78. The recombinant virus was subsequently plaque purified.

The Rep68 baculovirus construct was the same as the Rep78 clone except that the intron (AAV nucleotides 1906 to 2228) had been precisely deleted by oligonucleotide directed mutagenesis with oligonucleotide 5'-ACTACGCCAGACAGATT GGCTCGAGGA. This deletion mutant was made in the background of plM29, sequenced, and was designated dlint (19, 26, 27). Subsequently, the BamHI fragment from 29dl-320 was inserted into the BamHI site of dlint to form plasmid dl1-320int. The XbaI fragment from dl1-320int was then inserted into the BamHI site of the baculovirus transfer vector pVL941 to make the transfer plasmid 941XR68 (Fig. 2). Recombinant baculovirus was screened by dot blotting at limiting dilution with an AAV probe and then for the production of Rep68 by immunoblotting with the anti-52/40 monoclonal antibody previously described (13, 15). In the case of the anti-52/40 mouse monoclonal antibody, an ammonium sulfate-precipitated fraction of an ascites preparation was used at a 1/10,000 dilution. All baculovirus plaques were replated three times before use.

Preparation of baculovirus extracts. One-molar salt extracts of baculovirus-infected nuclei were prepared as follows. One liter of SF9 suspension cells was grown to a density of 1.2 × 10⁸ cells per ml in Grace's media supplemented with 10% fetal calf serum, penicillin, streptomycin, and 0.1% pluronic S68 (GIBCO). The cells were infected with the appropriate baculovirus recombinant at a multiplicity of 8 and incubated at room temperature for 2 to 5 days. After harvesting by centrifugation, the cells were washed once by suspension in buffer A (20 mM HEPES [N-2-hydroxyethylpiperazine-N'-2-ethanesulfonic acid] [pH 7.5], 5 mM KCl, 0.5 mM MgCl₂, and 0.5 mM DTT) containing 10% sucrose. The cells were then suspended in 40 ml of buffer A for 30 min on ice and lysed by 10 strokes with a type A Dounce homogenizer. Nuclei were harvested by centrifugation for 10 min at 2,000 × g and resuspended in 8 ml of buffer B (50 mM HEPES [pH 7.5], 10% sucrose, 0.4 mM EDTA, 0.3 mM phenylmethylsulfonyl fluoride, 2 µg of leupeptin per ml, and 1 µg of pepstatin per ml). The suspension was adjusted to 1 M NaCl, incubated on ice for 1 h, and then centrifuged for 1 h at 100,000 × g. The protein concentration of the 1 M nuclear S100 Rep extract was approximately 6 to 7 mg/ml in a final volume of approximately 10 ml. For Rep78, the specific activity was approximately 2 × 10⁶ U (see below for definition) of replication activity per mg of protein.

Purification of baculovirus-expressed Rep68. Rep68 was purified to apparent homogeneity from a 1 M nuclear extract of baculovirus-infected cells (1 liter) by a modification of the method previously described for Rep68 expressed in HeLa cells (15). Phenylsepharose and single-stranded DNA cellulose chromatography was performed as previously described (15). The single-stranded DNA cellulose pool was loaded onto a 1-ml HR5/5 mono-Q column, washed with 10 column volumes of buffer C (25 mM Tris-HCl [pH 7.5], 0.1 mM EDTA, 1 mM DTT, 50 mM NaCl) containing 0.1 mM phenylmethylsulfonyl fluoride, 0.2 mM pepstatin A, and 15% glycerol. The column was washed with 10 column volumes of the starting buffer and eluted with 20 column volumes of a convex 50 mM-to-1 M NaCl gradient in buffer C. Rep-containing fractions were identified by immunoblotting, pooled, and loaded onto a 1-ml HR5/5 mono-Q column. Elution from the mono-Q column was done in the same way as that from the mono-S column. The peak Rep-containing fraction was dialyzed against buffer C containing 50% glycerol, frozen in liquid nitrogen, and stored at −80°C.

Preparation of uninfected and Ad-infected HeLa cell extracts. Two kinds of extracts were used in this study. The first
was a cytoplasmic S100 extract prepared as described previously by Stillman and Gluzman (44). The second was a cytoplasmic extract supplemented with a 0.2 M NaCl nuclear wash and was prepared essentially as described by Wobbe et al. (53) and modified by Ward and Berns (51). Both types of extracts are called S100 extracts in this article. HeLa suspension cells (11) were grown to a density of 5 × 10⁵ cells per ml and harvested at 40 h postinfection by centrifugation to make uninfected extracts or extracts infected with Ad type 2 or type 5 ts149 (obtained from Bruce Stillman, Cold Spring Harbor Laboratory) at a multiplicity of infection of 5. After centrifugation at 100,000 × g for 1 h, the extracts were dialyzed overnight against 20 mM Tris (pH 7.5)–50 mM NaCl–10% glycerol–0.1 mM EDTA–1 mM DTT. The final protein concentrations of the uninfected and Ad-infected HeLa extracts were approximately 17 mg/ml, and the extracts were stored at −80°C.

Protein was measured with the Bradford reagent (Bio-Rad) using pooled bovine gamma globulin as the standard.

In vitro DNA replication assay. The standard AAV DNA replication assay was similar to the assay used previously for terminal resolution (42). It contained in 30 μl 30 mM HEPES (pH 7.5); 7 mM MgCl₂; 0.5 mM DTT; 100 μM each dATP, dGTP, dCTP, and dTTP; 25 μg of [α-32P]dATP (3 μCi/ml); 4 mM ATP; 40 mM creatine phosphate; 1 μg of creatine phosphokinase; 255 μg of Ad-infected HeLa S100 extract; 0.1 μg of NE substrate DNA (0.032 pmol of AAV DNA or 300 pmol of nucleotide); and 1 to 80 U of Rep78 or Rep68 baculovirus extract (0.5 to 8 μg). A unit of Rep activity was arbitrarily defined as an amount of Rep protein that would catalyze the incorporation of 1 pmol of radioactive deoxyxynucleoside monophosphate into DpnI-resistant monomer or dimer duplex AAV DNA in 2 h at 37°C. Where indicated, the reactions were supplemented with CTP, GTP, and UTP (0.2 mM each). The reaction mixtures were incubated for a minimum of 2 h and up to 6 h, as indicated. Following incubation, the reaction mixture was adjusted to 70 μl containing 0.3% sodium dodecyl sulfate (SDS), 0.7 mg of proteinase K per ml, and 17 mM EDTA. Proteinase K digestion was at 37°C for 1 h. The products were then extracted with phenol and chloroform and precipitated with ethanol. The ethanol precipitate was dissolute in 18 μl of water and, where indicated, digested with DpnI for 2 h at 37°C. The products (or a portion of them) were separated on either 0.8 or 1% agarose gels by electrophoresis for 4 h at 6 V/cm. The radioactivities in monomer and dimer RF products were counted in dried gels by a scanning gas flow (AMBIS) counter. X-ray film was exposed for 5 min to 16 h without a screen at room temperature.

In addition to the replication assay described above, total incorporation of radioactive precursors into acid-insoluble product was measured by the DE-81 filter paper (Whatman) method. Aliquots of products were spotted onto 1-cm² DE-81 filters and immediately immersed in 0.5 M dibasic potassium phosphate (pH 8.6). After 3 min, the filters were washed four more times (3 min each) with 0.5 M phosphate solution, three times with water, and twice with ethanol. The filters were then dried and counted in a liquid scintillation counter.

RESULTS

Expression of Rep proteins in baculovirus vectors. In our previous experiments, we had purified Rep68 to apparent homogeneity and we had partially purified Rep78. Both were purified from HeLa cell extracts that had been infected with Ad and AAV and both were capable of resolving the AAV terminal repeats. However, neither of these proteins was able to promote net DNA synthesis, and although it was possible to separate the two proteins chromatographically, the yields were poor (15, 16, 42). There were several possible explanations for the failure to achieve net synthesis. The simplest was that the level of Rep protein in HeLa cell crude or partially purified extracts was too low to support DNA synthesis. To obtain extracts that contained higher levels of Rep protein expression, we cloned Rep78, Rep68, and Rep52 into baculovirus vectors and prepared 1 M NaCl extracts of infected insect cell nuclei as described in Materials and Methods. Rep78 and Rep52 clones were constructed by inserting the appropriate region of the Rep coding sequence downstream of the baculovirus polyhedrin promoter. The Rep68 clone was constructed in a similar way except that a cDNA clone was constructed by oligonucleotide mutagenesis to remove the Rep68 intron. Figure 2A illustrates the structure of the baculovirus clones and demonstrates that each of the clones expressed a Rep protein of the appropriate size by immunoblot analysis with a monoclonal antibody (13) which recognizes an epitope that is present in all four Rep proteins. In addition, all of the baculovirus clones synthesized Rep proteins that comigrated with Rep proteins synthesized in AAV-infected HeLa cells (data not shown). None of the Rep baculovirus clones produced detectable levels of the other Rep species (Fig. 2A). This confirmed the earlier report by Owens et al. (32) that none of the other Rep proteins could be detected in extracts from their independently isolated Rep68-expressing baculovirus clone. Two different preparations of Rep78 are shown in Fig. 2A. The earlier one (far left lane) contains a number of shorter Rep species due to proteolysis, but none of these migrate at the position of Rep68 or Rep52. The addition of a several protease inhibitors during extraction (see Materials and Methods) largely eliminated these breakdown products (Fig. 2A, far right lane). Rep78 and Rep52 are each found in HeLa cell extracts as two slightly differently migrating forms (14, 15, 32, 34). In contrast, and as reported previously by Owens et al. (32) for Rep78, the baculovirus-expressed Rep78 and -68 were each present as predominantly one species (Fig. 2A).

The crude 1 M nuclear extracts in Fig. 2A were used as the source of the Rep proteins in the experiments described below. In the case of Rep68, we also purified the protein by a modification of a procedure we reported previously for the purification of Rep68 from HeLa cells (15) (Fig. 2B; see Materials and Methods). The most highly purified Rep68 fraction, a mono-Q fraction, is also shown in Fig. 2A (Rep68 Q). This fraction was apparently homogeneous. When 1.8 μg of the mono-Q fraction was analyzed on a silver-stained SDS-polyacrylamide gel (Fig. 2B), very little, if any, contamination by other protein species was detected. The Rep68 mono-Q fraction had a specific activity in the DNA replication assay of approximately 10⁷ U/mg of protein and was approximately 100- to 200-fold purified relative to the crude baculovirus preparation. The total yield of pure Rep68 was approximately 200 to 300 μg from 1 liter of infected SF9 cells.

The ability of Rep78, -68, and -52 to promote in vitro AAV DNA replication. To determine whether the baculovirus-expressed Rep proteins could promote AAV DNA synthesis, we tested each of them in an assay that was similar to the one we used previously for in vitro terminal resolution (42). The substrate for the reaction was NE DNA (Fig. 1), which consists of a linear AAV DNA molecule in which both ends are covalently joined (42). In addition to radiolabeled deoxyxynucleoside triphosphates, MgCl₂, and an ATP-generating system, the reaction mixture contained one of the three Rep proteins (supplied in a crude baculovirus extract) and a crude...
Ad-infected HeLa cell extract, which could supply the cellular or Ad-encoded replication factors that might be needed for DNA synthesis. The reaction mixtures were incubated for 2 to 6 h. The products were then digested with DpnI and separated by electrophoresis on neutral agarose gels. Successful net synthesis would result in the production of DpnI-resistant monomer and dimer linear AAV RFs (Fig. 1).

We anticipated from previous genetic and biochemical studies (4, 16, 40) that Rep52 would be unable to promote DNA synthesis by itself but that either Rep78 or Rep68 (or both) might be capable of generating DpnI-resistant AAV RFs. The products of a typical replication assay are shown in Fig. 3. As expected, Rep52 did not promote the incorporation of detectable levels of radioactive nucleotide into the NE substrate (Fig. 3, lane 5). The Rep52 reaction also served as an internal control which showed that, without the appropriate Rep protein, the enzymes in the HeLa cell extract or in the baculovirus extract were not capable by themselves of promoting significant background incorporation.

In contrast, the addition of either Rep68 or Rep78 produced DpnI-resistant AAV RF molecules (Fig. 3). In the case of Rep78, both monomer duplex and dimer duplex forms were synthesized (Fig. 3, lane 2) and the pattern of RFs was similar to that seen in vivo. Surprisingly, in the case of Rep68, only the monomer RF accumulated to any significant extent (Fig. 3, lane 3). To determine whether there might be an inhibitor in the Rep68 extracts that specifically prevented the accumulation of dimer RFs, we assayed the baculovirus-expressed Rep68 mono-Q fraction described above. It also produced primarily monomer RF (Fig. 3, lane 4). Thus, the inability to accumulate dimer RF species appeared to be an intrinsic property of Rep68, and it was the first clear difference that we have noticed between the biochemical activities of Rep68 and Rep78. In addition, the experiments in which we used purified Rep68 to drive the replication reaction allowed us to calculate the ratio of Rep to DNA molecules in our assay. Titration of the mono-Q fraction (data not shown) revealed that optimal activity was achieved at a ratio of approximately 20 Rep68 molecules to 1 terminal repeat in the starting substrate. This ratio was twice that used in the experiment whose results are shown in Fig. 3 and was based on a theoretical molecular mass of Rep68 of 61,000 Da. We could not be certain, however, that all of the Rep68 molecules in the mono-Q fraction were active.

In addition to the monomer and dimer RF species generated by Rep78, several additional DpnI-resistant species were often seen, particularly a doublet that migrated just more slowly than the dimer duplex (Fig. 3, lanes 1 and 2). The larger species of this doublet had a mobility consistent with that of a tetramer RF molecule (see also Fig. 4B), but the structures of the other species in the doublet as well as the even higher-molecular-weight forms in the Rep78 lanes have not yet been determined. All of these species have also been seen in vivo. Finally, a significant amount of radioactivity was incorporated into the DpnI-sensitive product during the course of the reaction. A comparison of the Rep52 lane with the Rep78 and Rep68 lanes (Fig. 3) suggests that a substantial amount of the DpnI-sensitive incorporation was also Rep dependent. A possible explanation for this effect is that the DpnI-sensitive products represent partially replicated molecules that have incorporated radioactive nucleotides in a Rep-dependent fashion.

**Reaction kinetics.** As expected, the amount of radioactive nucleotide incorporated and the level of DpnI-resistant monomer and dimer RFs synthesized were dependent on the amount of Rep extract added to the reaction mixture (Fig. 4A). Although some incorporation into DpnI-sensitive products was seen at all levels of Rep, incorporation into the DpnI-resistant product appeared to be linear with Rep extract only after a threshold level of Rep extract had been reached (Fig. 4A). This may reflect the need to assemble a complex of Rep or Rep and cellular factors prior to full-length DNA synthesis. A time course of the DNA replication reaction also suggested the need to preassemble an active replication complex. No DpnI-resistant full-length DNA was detected until approximately 30 min after the reaction was started (Fig. 4B). If total incorporation into DNA was measured by the DE-81 filter assay (see Materials and Methods), the lag was 17 to 20 min (data not shown). (The difference in lag time between the two assays presumably reflected the time that it took to synthesize the first full-length [DpnI-resistant] product after incorporation began.) After the initial 30-min lag, however, the accumulation of monomer and dimer RFs was approximately linear for the next 4 h. This suggested that there was some rate-limiting step early in the reaction, perhaps the need to assemble a protein complex or the need to inactivate an inhibitor. The rate of incorporation of radioactive dAMP into monomer and dimer duplexes was approximately 20 to 40 pmol/h after the initial 30-min lag. This is similar to the rate of incorporation in the SV40 DNA replication systems reported previously (21, 44, 53). Given the amount of starting substrate DNA in the standard 30-μl reaction mixture (300 pmol of nucleotide; approximately 75 pmol of dAMP), the incorporation of 150 pmol of dAMP in approximately 4 h (Fig. 4B) is consistent with each starting molecule being replicated up to two times.

**Semiconservative replication.** To confirm that the DpnI-resistant DNA produced was the result of semiconservative replication, a reaction was performed without any radioactive label and with bromo-dUTP in place of dTTP. The products of the reaction were separated in a CsCl density gradient, and the species in each fraction were analyzed by Southern hybridiza-
The reaction was incubated for 4 h and treated with proteinase K. Aliquots (one-fifth) of each reaction were electrophoresed on a 1% agarose gel both before (left) and after (right) DpnI digestion. The BssHII and BstEII lanes contain λ bacteriophage DNA digested with the indicated restriction enzyme and 3’ labeled with Klenow fragment to provide size markers. The sizes of the BssHII fragments are shown on the left. The relevant BstEII fragment sizes are indicated on the right. Monomer length sub201 AAV DNA in which both ends have been resolved is expected to be 4.702 kbp long (36, 43); the dimer RF with both ends resolved is expected to be 9.279 kbp. Each unresolved end would reduce the size of an RF molecule by 0.124 kbp. The amount of [32P]dAMP incorporated into DpnI-resistant monomer and dimer RF molecules was counted and is displayed graphically below. The incorporation was adjusted to represent the amount incorporated in a 15-μl reaction mixture. (B) Time course of AAV DNA synthesis. The standard AAV DNA replication reaction mixture (30 μl) contained 3.4 μg of crude Rep78 nuclear extract and 255 μg of Ad-infected HeLa S100 extract. The reaction mixtures were incubated for the indicated time, treated with proteinase K and DpnI, and electrophoresed on a 1% agarose gel. The positions of the monomer (md) and dimer (dd) duplex RF species are indicated on the right, as are the incompletely replicated DpnI-sensitive products (DpnI). The amount of [32P]dAMP incorporated into DpnI-resistant monomer and dimer RF molecules was counted and is displayed graphically below.

![Graph](https://via.placeholder.com/150)

**FIG. 4.** (A) Titration of Rep78. The standard DNA replication assay was used, except that it was adjusted to a total volume of 15 μl containing 127 μg of Ad-infected HeLa S100 extract and (from left to right) 0.1, 0.2, 0.4, 0.9, 1.7, or 6.8 μg of Rep78 baculovirus nuclear extract. The reaction mixtures were incubated for 4 h and treated with proteinase K. Aliquots (one-fifth) of each reaction were electrophoresed on a 1% agarose gel both before (left) and after (right) DpnI digestion. The BssHII and BstEII lanes contain λ bacteriophage DNA digested with the indicated restriction enzyme and 3’ labeled with Klenow fragment to provide size markers. The sizes of the BssHII fragments are shown on the left. The relevant BstEII fragment sizes are indicated on the right. Monomer length sub201 AAV DNA in which both ends have been resolved is expected to be 4.702 kbp long (36, 43); the dimer RF with both ends resolved is expected to be 9.279 kbp. Each unresolved end would reduce the size of an RF molecule by 0.124 kbp. The amount of [32P]dAMP incorporated into DpnI-resistant monomer and dimer RF molecules was counted and is displayed graphically below. The incorporation was adjusted to represent the amount incorporated in a 15-μl reaction mixture. (B) Time course of AAV DNA synthesis. The standard AAV DNA replication reaction mixture (30 μl) contained 3.4 μg of crude Rep78 nuclear extract and 255 μg of Ad-infected HeLa S100 extract. The reaction mixtures were incubated for the indicated time, treated with proteinase K and DpnI, and electrophoresed on a 1% agarose gel. The positions of the monomer (md) and dimer (dd) duplex RF species are indicated on the right, as are the incompletely replicated DpnI-sensitive products (DpnI). The amount of [32P]dAMP incorporated into DpnI-resistant monomer and dimer RF molecules was counted and is displayed graphically below.

**FIG. 5.** Isopycnic centrifugation of in vitro-synthesized AAV DNA. DNA was synthesized under standard reaction conditions except that 100 μM 5’bromo-dUTP was used instead of dTTP in a total reaction volume of 120 μl for 6 h in the absence of radioactive deoxynucleoside triphosphates. After incubation, the reaction products were treated with proteinase K and centrifuged to equilibrium in a neutral CsCl solution containing 10 mM Tris-HCl (pH 7.5), 1 mM EDTA, and 1.750 g of CsCl per ml in a total volume of 4 ml. Fractions were collected from the bottom and electrophoresed on a 0.8% agarose gel. The presence of AAV DNA was detected by Southern blotting using the randomly labeled AAV-containing PvuII fragment from pFub201 plasmid DNA as a probe. The density of the fractions was determined by weight to confirm that the density gradient was linear (not shown). The expected positions of unreplicated, light-light (LL) DNA, once replicated heavy-light (HL) DNA, and doubly replicated heavy-heavy (HH) DNA are indicated at the top. md, monomer duplex; dd, dimer duplex.

flop orientation during the first round of resolution (36, 42). Reinitiation and strand displacement synthesis of the full-length molecule would then generate a covalently closed end that would generate the flip orientation after a second round of resolution. In our previous experiments, in which no net synthesis and only one round of terminal resolution had occurred, we detected only the flip orientation in the products of the reaction (42). Under the conditions of net synthesis used here, we expected to find both orientations in the products of the reaction. To see whether we could detect both orientations of the terminal repeats, we isolated the resolved terminal XbaI fragment from the products of the replication reaction as described previously (42) (see the legends to Fig. 6A and 7B). Digestion of the terminal XbaI fragment with SmaI revealed that both orientations were present in the products of the reaction in approximately equal amounts (Fig. 6A). This suggested that, indeed, more than one round of terminal resolution was occurring during the course of the replication reaction.

In addition, we demonstrated in our previous work that the apparent molecular weight of the in vitro-replicated SmaI terminal flip fragment was slightly larger than expected (42). This was due to the fact that, during in vitro resolution, a molecule of Rep protein was covalently attached to the 5’ end of the terminal repeat, and even after treatment of the products with proteinase K, a portion of the Rep protein still remained attached (15, 40, 42). When the SmaI flop fragment was treated with proteinase and proteinase K, its apparent molecular weight was reduced further, presumably because additional amino acids of the Rep protein were removed. To see whether the same would be true of the terminal fragments generated in the DNA replication assay, some of the SmaI terminal fragments were digested with proteinase in addition to proteinase K (Fig. 6A). Both the flip (68 bp) and the flop (46 bp) terminal fragments were shifted to a lower molecular weight after digestion with proteinase. This suggested that the bulk of the product DNA generated in the in vitro replication reaction contained a covalently attached Rep protein.

**Requirement for a functional AAV origin.** To determine
whether the in vitro AAV replication reaction was specific for a wild-type AAV origin, a mutant origin was tested in the reaction. The HpaIA mutant substrate consisted of NE DNA in which both termini had identical 8-bp linker insertions near the terminal resolution site. In previous work, we have shown that this mutant substrate is approximately 50-fold less efficient in its ability to be nicked by the Rep protein at the terminal resolution site in vitro (41), and Samulski and his colleagues (34a) have shown that it is equally defective for DNA replication in vivo. When the HpaIA mutant substrate was compared with the wild-type origin in the in vitro replication assay, there was a clear preference for the wild-type terminal repeat (Fig. 6B). Only small amounts of monomer and dimer RF species could be detected in the products of the reaction with the mutant substrate, even after prolonged exposure of the gel.

**FIG. 6.** (A) Orientation of the resolved termini. The standard DNA replication reaction mixture was adjusted to 15 μl containing 127 μg of Ad-infected HeLa S100 extract and 1.7 μg of 1 M baculovirus Rep78 nuclear extract and incubated at 37°C for 4.5 h. The products were treated with protease K, digested with XbaI and PstI, and electrophoresed on a 6% polyacrylamide gel. The resolved, 32P-labeled XbaI terminal fragment was isolated from the gel, digested with SmaI, and electrophoresed on an 8% nondenaturing polyacrylamide gel. The left lane contains products that had been treated only with proteinase K (prot K); the right lane contains fragments that had been further treated with pronase (pr) prior to electrophoresis. The diagram on the right illustrates the relative positions of SmaI sites in the flip and flop orientations of the resolved termini. Arrowheads indicate the 3′ ends of each strand. Ovals indicate the fact that, after proteolytic digestion, a portion of the Rep protein remains covalently attached to the 5′ end of the terminal repeat. Numbers indicate the sizes (in base pairs) of the expected SmaI-digested products. (The small 10-bp fragment that contains the sequence between the two SmaI sites is lost on these gels.) As shown previously by Snyder et al. (42), the variation in the sizes of the flip-terminal SmaI fragment (68 bp) and the flop-terminal fragment (46 bp) reflects the fact that the alternate proteolytic treatments leave different amounts of the Rep protein covalently attached to the fragments. X, the resolved terminal XbaI fragments that escaped SmaI digestion or were partially digested (immediately below the X fragment). (B) Comparison of wild-type and mutant origins. Duplicate reactions were performed with wild-type (wt) or HpaIA mutant (mut) NE template DNA. The mutant contains an 8-bp linker inserted at the tsr site. It is 50-fold reduced in its ability to be nicked by the Rep protein in vitro (41) and defective for DNA replication in vivo (56a). The DNA replication reaction conditions were the same as those for Fig. 4B. The products of the reaction were electrophoresed on a 0.8% agarose gel.

**FIG. 7.** Requirement for Rep protein and an Ad-infected extract. (A) NE DNA was incubated with S100 extracts from either uninfected or Ad-infected HeLa cells in the presence of no additional extract, a wild-type baculovirus (wt bac) nuclear extract, or a baculovirus extract containing Rep78 (BR78) under standard replication reaction conditions in a total reaction volume of 120 μl for 6 h at 37°C. Wild-type and Rep78 baculovirus extracts (20.4 μg) were added to the indicated reactions mixtures. Following incubation, the reaction mixtures were treated with proteinase K, and one-sixth each reaction mixture was analyzed on a 0.8% agarose gel, either with (+) or without (−) DpnI digestion. dd and md, DpnI-resistant dimer duplex and monomer duplex AAV DNAs, respectively; DpnI, the DpnI-sensitive partially replicated products. (B) Aliquots of the reaction mixtures in panel A and two additional reaction mixtures with baculovirus Rep68 extract were treated with PstI and XbaI and electrophoresed on a 6% polyacrylamide gel. Cx and Dx, internal PstI-XbaI fragments; X and Xn, resolved and unresolved terminal XbaI fragments, respectively. The marker lane contains 3′-labeled pBR322MspI fragments.

**Requirement for an Ad-infected extract.** As mentioned earlier, AAV DNA replication is unusual in that coinfection with Ad is usually necessary to establish fully permissive conditions for DNA replication. To see whether the in vitro reaction also depended on Ad infection, the abilities of uninfected and Ad-infected HeLa cell extracts to support AAV DNA replication were compared (Fig. 7 and 8). The results in Fig. 7A show that, in the presence of Rep78, uninfected HeLa extracts supported incorporation of radioactive nucleotides into product DNA but virtually all of the product DNA was DpnI sensitive. In contrast, the Ad-infected extract efficiently generated DpnI-resistant monomer and dimer RF species. As expected, both the uninfected and the Ad-infected extracts promoted little incorporation in the absence of Rep protein or in the presence of a wild-type baculovirus extract. The difference between uninfected and Ad-infected HeLa extracts was confirmed by titrating each extract in the presence of a constant amount of Rep (Fig. 8). As more Ad-infected extract was added to the reaction mixture, the yield of DpnI-resistant monomer and dimer RF products increased (Fig. 8, lanes 5 to 8). In contrast, it was difficult to detect DpnI-resistant products with the uninfected HeLa extract at any concentration that was tried (Fig. 8, lanes 1 to 4). The same difference between uninfected and Ad-infected extracts was seen when Rep68 was used in the replication assay (Fig. 8, lanes 9 and 10). The effect
of Ad infection suggested that some factor(s) that directly participated in AAV DNA replication was missing (reduced) in an uninfected HeLa extract. Other explanations, however, are also possible (see Discussion).

Finally, in previous work we had demonstrated that an uninfected HeLa cell extract was capable of supporting the terminal resolution reaction (Fig. 1) provided it was supplemented with purified Rep protein (40). To see whether this was true of the HeLa extracts used in this study, we digested the products of the in vitro replication assay with XbaI and PstI and compared the yields of the resolved terminal XbaI fragment produced in the presence of uninfected and Ad-infected extracts (Fig. 7B). Both types of extracts were found to generate approximately the same level of the resolved XbaI fragment in the presence of either Rep78 or Rep68 (Fig. 7B; compare the X fragments in lanes 4 and 8 with those in lanes 5 and 9). This confirmed our earlier results and suggested that the block in AAV DNA replication with an uninfected HeLa extract occurred at some step after terminal resolution, either in the formation of the reinitiation intermediate or in elongation (Fig. 1).

Other components required for the reaction. As already discussed, the in vitro reaction required the NE DNA substrate with wild-type terminal repeats and either Rep78 or Rep68 (Table 1). As expected from other DNA replication systems, the AAV replication reaction also required the presence of MgCl₂, ATP, and an ATP-generating system (creatine phosphate and creatine phosphokinase). Omission of the other ribonucleoside triphosphates (UTP, GTP, and CTP) did not affect the reaction (Table 1).

DISCUSSION

We have developed an efficient in vitro AAV DNA replication assay that should be useful for identifying and purifying the cellular (or helper virus) components involved in AAV DNA replication. Our assay is up to 100-fold more efficient than previously described assays, both in the utilization of the starting substrate and in the rate of incorporation (12, 59). Density shift experiments indicated that almost all of the starting substrate was replicated at least once and a portion of the substrate (less than 5%) was replicated twice during a 6-h incubation (Fig. 5). The level of labeled nucleotide incorporated suggested that two rounds of DNA replication could occur (Fig. 4). The discrepancy between these experiments was likely due to an uncertainty in the amount of starting substrate used or to inherent differences in the extracts used in the two experiments. A comparison of the DNA replication assay presented here and that of Hong et al. (12) suggests that the difference in efficiency is probably due to the use of circular AAV plasmids as the starting substrate for replication. Before AAV replication can proceed efficiently, a linear molecule must be rescued from the circular plasmid. Aside from being a potential rate-limiting step, we (6-8) and Ward and Berns (51) have shown that in vitro rescue of AAV linear DNA can occur by at least two different mechanisms which do not always produce products that are active substrates for subsequent AAV DNA replication (31a). It should also be noted that our results are not necessarily in conflict with those of the experiments reported by Yalkinoglu et al. (59). We have suggested previously that there may be two different mechanisms for AAV DNA replication, the one described here and a second one which may be used during integration of AAV proviruses (29). The latter may rely on the use of the AAV terminal repeat as a cellular origin, as suggested by Yalkinoglu et al. (59).

The reaction faithfully reflected in vivo AAV DNA replication by several criteria. First, the major replicative intermediates formed during the reaction were monomer and dimer duplex linear DNAs, and most of the products were DpnI resistant, suggesting that full-length AAV DNA replication had occurred. Second, the reaction required the presence of one of the two larger Rep proteins, Rep78 or Rep68. Third, a mutant origin that we had previously shown to be defective for nicking by Rep68 (41) was also defective as a substrate in the DNA replication assay. Fourth, the in vitro products contained both orientations of the terminal repeat (flip and flop) in approximately equal amounts, as in vivo. This indicated that at least two rounds of terminal resolution had occurred in a large fraction of the termini. Fifth, the reaction was much more efficient in the presence of an Ad-infected HeLa cell extract.

### Table 1. Components required for in vitro replication of AAV DNA

<table>
<thead>
<tr>
<th>Component omitted</th>
<th>% of complete reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>100</td>
</tr>
<tr>
<td>MgCl₂</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>ATP</td>
<td>3.3</td>
</tr>
<tr>
<td>CTP, GTP, UTP</td>
<td>100</td>
</tr>
<tr>
<td>dGTP, dCTP, dTTP</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>CP, CPK</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>NE substrate</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>Rep78</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>+Baculovirus*</td>
<td>&lt;0.5</td>
</tr>
<tr>
<td>Ad-infected HeLa S100*</td>
<td>&lt;0.5</td>
</tr>
</tbody>
</table>

*See "In vitro DNA replication assay" for the complete reaction mixture.

Additional components required for the Rep78 replication reaction were supplied by 3.4 μg of wild-type-infected baculovirus extract. The latter contained S100 and Rep78. S100 can substitute for the Rep78-containing baculovirus extract. A wild type-infected baculovirus extract was substituted for the Rep78-containing baculovirus extract.

*An uninfected HeLa S100 extract was substituted for the Ad-infected S100 extract.
than it was with an uninfected cell extract. This was also consistent with AAV replication in vivo.

The stoichiometry of the products that we observed in the in vitro reaction can be explained by the current model for AAV DNA replication as follows. The hairpinned termini in the starting NE substrate are first resolved at both ends to produce the flp orientation at both ends. One of the resolved ends is then converted to the double hairpin intermediate (illustrated in Fig. 1 as the reorientation substrate), which then primes a round of full-length strand displacement synthesis to produce a duplex molecule with one hairpinned (or covalently closed) end and a single-stranded molecule. Resolution of the hairpinned end in the duplex molecule would then produce a duplex (heavy-light) molecule that is resolved at both ends and has one flp end and one flop end. Similarly, the single-stranded product of the strand displacement reaction would be elongated by hairpin priming from the 3’ end to produce a duplex product with one end in the hairpinned configuration. Terminal resolution of this molecule would also produce a double-stranded product with one flp end and one flop end. Thus, the net result of one round of strand displacement synthesis, followed by elongation of the single-stranded product and the resolution of all hairpinned ends, would be two double-stranded hybrid (heavy-light) molecules and an equal molar ratio of flp and flop ends. This is consistent with our observations (Fig. 5 and 6A). As yet, we have not examined the product DNA to determine the distribution of flp and flop ends on individual molecules. Dimer molecules (Fig. 1) would be the result of reinitiation and strand displacement synthesis on molecules in which only one of the ends had been resolved. Such events, followed by the resolution of all dimer termini, would also produce hybrid (heavy-light) molecules with equimolar ratios of flp and flop ends. This also is consistent with our observations (Fig. 5). Finally, we note that other mechanisms could explain the equimolar ratio of flp and flop ends in the product DNA. For example, repeated resolution and partial elongation could occur at an end to generate what would appear to be an equimolar flip/flop ratio. Such a mechanism would be expected to generate small DNA products that contained only the terminal repeat sequence. Although such products were not seen during the course of these studies, we have not rigorously excluded them.

One aspect of the reaction that may not faithfully reflect in vivo AAV DNA replication was the covalent linkage of the Rep protein to the ends of product DNA. Although we have not demonstrated this conclusively, the shift in size in the terminal Smal fragments after sequential proteolytic digestion (Fig. 6) suggested that most of the product synthesized in vitro contained a covalently attached Rep protein. It is not known whether this is also the case with replicative intermediates in vivo, but there is no evidence that AAV DNA packaged in vivo contains a covalently attached protein. Thus, some processing step that would be required to remove the covalently attached protein prior to or during packaging does not occur in our in vitro DNA replication reaction. It should be noted that no AAV capsid proteins were present in our reactions and Rep and capsid proteins have been found to colocalize in AAV-infected nuclei (13).

In previous work, we have compared the biochemical properties of Rep78 and Rep68 with respect to DNA binding, its endonuclease activity, and DNA helicase activity (14–16). The inability of Rep68 to generate significant levels of dimer RF intermediates is the first qualitative difference that we have found between the two enzymes. The reason for the difference is not yet clear, but one possible explanation is that Rep68 is more efficient in resolving internal AAV terminal sequences that are present in a dimer AAV-AAV junction. This may have some physiological significance for the rescue of AAV proviruses and for normal AAV DNA replication.

The requirement for an Ad-infected extract suggests that AAV DNA replication needs one or more components that are not needed for the well-characterized SV40 DNA replication system. In principle, the AAV DNA replication mechanism involves only leading-strand DNA synthesis. Extrapolating from what is known about SV40 replication, we might expect that AAV replication would require (in addition to Rep protein) RPA, RFC, polymerase 6, and PCNA (20, 21, 30, 33, 49, 50, 52–56). All of these components are present in ample amounts in uninfected HeLa cell extracts, but these extracts do not efficiently support AAV replication in vitro. This suggests that some additional component that is either coded by Ad or coded by the cell and induced following Ad infection is required for AAV DNA replication. Previous genetic experiments have shown that neither the Ad DNA polymerase nor the Ad terminal protein is required for AAV DNA replication or virus production (17, 29a, 45). Studies of Ad DNA-binding protein mutants suggest that, although they do affect virus production, they have little, if any, effect on AAV DNA replication in vivo (3; also references therein). Thus, the missing component required for AAV DNA synthesis is likely to be a cellular enzyme. Finally, both Rep78 and Rep68 were capable of resolving the hairpinned terminal sequences in the presence of uninfected cell extracts (40) (Fig. 7). Thus, whatever the missing component is, it appears to be required either for the reinitiation step of AAV DNA replication (in which a putative double hairpin intermediate is formed) or for elongation during strand displacement synthesis (Fig. 1). It remains to be seen whether the difference between uninfected and Ad-infected extracts we observed is due to a novel cellular factor or simply to the induction of higher levels of known replication factors by Ad infection. We note also that there are several other possible explanations for the improved DNA synthesis in Ad-infected extracts. For example, uninfected extracts may contain an inhibitor of AAV DNA replication that is inactivated during Ad infections. Alternatively, Ad infection may lead to posttranslational modifications of some of the known cellular replication proteins. Ultimately, it will be necessary to reconstitute AAV DNA replication with purified proteins to determine the role of the helper virus.

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