Differential Effects of DNA Double-Strand Break Repair Pathways on Single-Strand and Self-Complementary Adeno-Associated Virus Vector Genomes

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The linear DNA genomes of recombinant adeno-associated virus (rAAV) gene delivery vectors are acted upon by multiple DNA repair and recombination pathways upon release into the host nucleus, resulting in circularization, concatemer formation, or chromosomal integration. We have compared the fates of single-strand rAAV (ssAAV) and self-complementary AAV (scAAV) genomes in cell lines deficient in each of three signaling factors, ATM, ATR, and DNA-PKcs, orchestrating major DNA double-strand break (DSB) repair pathways. In cells deficient in ATM, transduction as scored by green fluorescent protein (GFP) expression is increased relative to that in wild-type (wt) cells by 2.6-fold for ssAAV and 6.6-fold for scAAV vectors, arguing against a mechanism related to second-strand synthesis. The augmented transduction is not reflected in Southern blots of nuclear vector DNA, suggesting that interactions with ATM lead to silencing in normal cells. The additional functional genomes in ATM−/− cells remain linear, and the number of circularized genomes is not affected by the mutation, consistent with compartmentalization of genomes into different DNA repair pathways. A similar effect is observed in ATR-deficient cells but is specific for ssAAV vector. Conversely, a large decrease in transduction is observed in cells deficient in DNA-PKcs, which is involved in DSB repair by nonhomologous end joining rather than homologous recombination. The mutations also have differential effects on chromosomal integration of ssAAV versus scAAV vector genomes. Integration of ssAAV was specifically reduced in ATM−/− cells, while scAAV integration was more profoundly inhibited in DNA-PKcs−/− cells. Taken together, the results suggest that productive rAAV genome circularization is mediated primarily by nonhomologous end joining.

Recombinant adeno-associated virus (rAAV) vectors are an important tool for gene transfer, with a growing presence in gene therapy clinical applications. The single-stranded DNA genome of conventional rAAV vectors (ssAAV) contains only the desired transgene and the 145-bp palindromic inverted terminal repeat (ITR) sequences, which function as the viral origins of replication, packaging sequences, and priming sites for synthesis of the cDNA strand. A derivative rAAV vector, termed self-complementary AAV (scAAV), is packaged as a single-stranded inverted repeat, which can fold into double-stranded DNA (dsDNA) without the requirement for DNA synthesis, leading to more efficient transduction (20, 21).

Once released into the nucleus, the free DNA ends of rAAV genomes are targets for DNA repair pathways, leading to circularization or concatemerization and, infrequently, chromosomal DNA integration. At low multiplicities of infection (MOIs), monomeric circular episomes are the predominant stable form of the rAAV genome, and at higher MOIs, increased opportunities for end-to-end joining lead to the formation of concatemeric episomes, which also circularize. These recombination events appear to follow the conversion of the ssDNA genome to dsDNA (41). We have found similar processes acting on scAAV vectors, demonstrating that ssDNA is not a prerequisite for the formation of circles and concatemers (5).

Our primary interest in rAAV circularization is how it relates to the persistence of rAAV vector genomes in the nucleus and to chromosomal integration, with its potential for genotoxicity and oncogenesis (9, 23, 24, 29). It is currently not known whether the same DNA recombination processes that mediate circularization also lead to integration. However, the most likely starting substrate for either fate is a linear rAAV genome that has been converted to dsDNA, with an ITR in either a covalently closed hairpin or an open duplex configuration at each end. Either structure is subject to recombination, though the efficiencies and host factor requirements differ (5). The genome ends interact with each other to form circles and concatemers or with preexisting chromosomal DNA double-strand breaks (DSBs), leading to integration (22).

Circularization of ssAAV is highly dependent on the nonhomologous end-joining (NHEJ) pathway for DSB repair in some animal tissues but not others (12, 16a, 25, 34, 35). While NHEJ plays an important role in processing ssAAV genomes, it also appears to have negative effects on transduction efficiency. A deficiency in Ku86, an essential core component of NHEJ-mediated repair, leads to higher levels of transduction from ssAAV vectors in cultured cells (42). The NHEJ pathway also participates in AAV Rep protein-mediated site-specific

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integration into the chromosome 19 AAV-S1 site, though with differential effects on scAAV and ssAAV vector genomes (8).

The conversion of the ssAAV genome to dsDNA may be intricately linked to its subsequent recombination to circular or concatameric forms. Transfection from ssAAV vectors is significantly increased by treatments that induce DNA damage repair pathways (1). Transduction is also increased by the adenovirus (Ad) E4ORF6 gene product, which disrupts an important DSB repair function through the segregation of cellular Mre11 (10, 13, 15, 31). Mre11 is a component of the MRN complex (Mre11, Rad50, and Nbs1), which promotes DSB repair by both homologous recombination (HR) and the NHEJ pathway (28). The MRN complex colocalizes with ssAAV genomes through interaction either with the ssDNA or with the ITR hairpin (3, 31). Consistent with the positive effects of Ad E4ORF6 on ssAAV transduction, MRN interactions with the rAAV genome inhibit transduction, possibly at the step of conversion to dsDNA. This is also consistent with reports that mutations in ATM (ataxia telangiectasia mutated) increase transduction from ssAAV vectors (30, 42). ATM is the primary signaling kinase responding to DSB in association with the MRN complex, and it is involved in the repair of DSB caused by ionizing radiation. These studies cumulatively suggest that interactions with the ATM-mediated pathway of DSB repair block conversion of the ssAAV genome to dsDNA.

Because scAAV vectors do not require complementary strand synthesis, they can be used to specifically test the contributions of DNA repair factors to recombination independent of dsDNA conversion. We measured the ratio of circularized to linear genomes in cultured cells (NBS1), but not both, other DNA repair factors had significant effects in either muscle-injected into the muscles of ATM-deficient mice. Mutations in ATM, which resulted in an apparent 70 to 80% decrease in the ratio of circularized to linear genomes. However, the greatest inhibition came from mutations in the DSB repair pathways that can operate on rAAV vector (ssAAV-GFP, ssAAV-GFP-CD, scAAV-GFP, and scAAV-GFP-CD). The ssAAV and scAAV vectors suggested that the damage response was specific not to the ssDNA or the hairpin ITR but to a cis-acting sequence within the wt AAV P5 promoter region (16). While rAAV vectors lacking the P5 promoter element do not provoke a pan-nuclear DNA damage response and G2/M cell cycle arrest, it remains likely that the ssDNA associates with RPA in the context of dsDNA conversion, making it a potential substrate for recombination through the ATR pathway. In contrast, the scAAV genome folds rapidly into double-stranded DNA, without the requirement for DNA synthesis, and therefore lacks significant regions of ssDNA. This raises the possibility that ssAAV and scAAV vector genomes could undergo differential processing to reach their stable isofoms.

In order to determine how the various DNA repair pathways interact with ssAAV and scAAV vector genomes to promote circularization or chromosomal integration, we used reporter vector-based assays with cells deficient in three phosphatidylinositol 3-kinase (PI3)-like kinase activities that orchestrate DNA repair (ATM, ATR, and DNA-PK\textsubscript{cs}).

**MATERIALS AND METHODS**

**Cell lines, maintenance, and treatments.** Cell lines overexpressing wt ATR (GW33) or kinase-dead ATR (ATR\textsuperscript{KD} (GK41)) under the control of a tetracycline (Tet)-inducible promoter were a kind gift from Paul Nghiem (26). wt ATR and ATR\textsubscript{KD} cells were grown in alpha minimal essential medium (MEM) (GIBCO) supplemented with 10% fetal bovine serum (FBS), 200 µg/ml G418, and 200 µg/ml hygromycin B. For vector transduction assays, overexpression of the transgene was induced with 1 µg/ml doxycycline for 48 h prior to viral infection and maintained until green fluorescent protein (GFP) expression was evaluated at 24 h postinfection.

Simian virus 40 (SV40)-transformed fibroblast cell lines from a normal individual (wt cells, GM00637 J) and from an ataxia telangiectasia (AT) homozygous patient (ATM<sup>-/-</sup> cells, GM05849 E) were purchased from Coriell Institute (Camden, NJ). wt and ATM<sup>-/-</sup> cells were grown in Eagle MEM with Earle’s balanced salt solution (BSS) (Lonza BioWhittaker) supplemented with 10% FBS and 1% GlutaMAX (GIBCO). To inhibit ATM or DNA-PK\textsubscript{cs} activity in the transformed fibroblasts, cultures were treated with 10 µM KU55933 (Toxic), respectively, for 24 h before infection, and the treatment was maintained for the duration of the experiment.

Human malignant glioblastoma cells lines expressing normal levels of DNA-PK (DNA-PK<sup>+</sup> cells, M059K) or lacking DNA-PK activity (DNA-PK<sup>−</sup> cells, M059J) were purchased from ATCC (Manassas, VA). DNA-PK<sup>−</sup> and DNA-PK<sup>+</sup> cells were grown in Dulbecco modified Eagle medium (DMEM)–balanced salt solution (BSS) (Lonza BioWhittaker) supplemented with 10% FBS and 1% GlutaMAX (GIBCO). To inhibit ATM or DNA-PK<sub>cs</sub> activity (G418)–mediated rounds of infection and quantification by flow cytometry.

Infection and quantification by flow cytometry. For transduction and circularization assays with the inducible wt ATR and ATR<sup>KD</sup> cell lines (GW33 and GK41), cells were seeded on 12-well plates on day 1 at 12.5% confluence and treated with doxycycline (1 µg/ml). At day 3, the cells were infected with each rAAV vector (ssAAV-GFP, ssAAV-GFP-CD, scAAV-GFP, and scAAV-GFP-CD) and analyzed by flow cytometry.
with a multiplicity of infection (MOI) of 0.5 HeLa C12 infectious unit per cell. Medium with virus was replaced with fresh medium at 2 h postinfection. Doxycycline treatment was maintained until cells were harvested. wt, ATM^−/−, DNA-PK^−/−, and DNA-PK^−/− cells were seeded on 12-well plates on day 1 at 25% confluence and infected on day 2 at an MOI of 1. All cell lines were harvested at 24 h postinfection, washed with ice-cold phosphate-buffered saline, and fixed with 4% paraformaldehyde in phosphate-buffered saline on ice. GFP expression was detected by flow cytometry (Coulter XL cytometer).

For integration assays, wt ATR and ATR^−/− cells were seeded and treated with doxycycline as described above. At day 3, cells were infected at an MOI of 260 for 2 h. Medium with virus was removed and replaced with fresh medium with doxycycline until day 2 postinfection, when doxycycline treatment was removed to maintain cell viability. wt, ATM^−/−, DNA-PK^−/−, and DNA-PK^−/− cell lines were seeded on 24-well plates on day 1 at 25% confluence and infected at day 2 at an MOI of 1,400 for 2 h. At day 2 postinfection, cells were transferred to 12-well plates. All cells lines were passed at 1:5 or 1:10 every 4 to 7 days. At each passage, cell samples were collected and fixed as described above for flow cytometry analysis. Cells were passed continuously until the percentage of GFP-positive cells reached a plateau.

Southern blotting. wt and ATM^−/− cell lines were seeded on 10-cm plates on day 1 at 30% confluence and infected at day 2 with scAAV-GFP vector at an MOI of 250 (determined on HeLa C12 cells). Medium with virus was replaced with fresh medium at 5 h postinfection. At day 3, cells were harvested and cytoplasmic (C) and nuclear (N) fractions were isolated as described previously (37). Nuclear and cytoplasmic fractions were resuspended in 10 mM Tris–10 mM EDTA (pH 8.0) plus 1% SDS and incubated with 1 μl RNase (10 mg/ml) for 0.5 h at 37°C, followed by 2 h incubation with protease K (20 μg/ml) at 37°C. After the addition of NaCl (1 M), samples were refrigerated at 4°C overnight. Following centrifugation at 16,000 × g, supernatants were phenol- chloroform extracted and ethanol precipitated in the presence of 10 μg of yeast RNA carrier. All samples were digested with Sphi I and Bsa6I (noncutters in AAV vector) to digest episomal SV40 DNA in the SV40-transformed fibroblasts. Nuclear fractions were divided for further digestion with restriction enzymes (Xhol, MluI, and Plasmid Safe PS-DNase). Samples were electrophoresed on 0.8% neutral agarose gels, blotted onto a nylon membrane, and probed with 32P-labeled probe specific for the cytomegalovirus (CMV) promoter and the GFP protein-coding region. Vector DNA excised from plasmids was subjected to Southern blotting.

Comparison of ssAAV and scAAV transduction and genome circularization in ATM-deficient cells. We had previously determined that ATM played a role in the circularization of scAAV vector genomes but that ATR did not. To determine whether ATM had differential effects on ssAAV and scAAV genomes, we compared transduction and circularization in a transformed ATM^−/− fibroblast cell line (GM05849E) and transformed normal fibroblasts (GM00637J).

Vector transduction and circularization were determined by quantifying expression from vectors containing either an intact GFP gene or a split GFP gene cassette. The circularization-dependent split GFP (GFP-CD) has the left and right halves situated at the ends of the genome and fused to a split intron such that it can make the gene product only when two vector genome ends have been joined together by DNA recombination. The multiplicity of infection (MOI) was kept low (<1) in these experiments to ensure that expression from GFP-CD would be the result of circularization, rather than end-to-end joining of two vector genomes within the same nucleus, and that each GFP-positive cell was the result of a single recombination event involving a single, productively infecting rAAV genome.

In ATM^−/− cells, the overall levels of transduction from the ssAAV and scAAV were increased by 2.6- and 6.6-fold, respectively, compared to those in normal fibroblasts, as evidenced by the increase in GFP positive cells from the intact GFP vector (Fig. 1a [open bars]; Table 1). This was consistent with a previously described effect on ssAAV vectors in ATM^−/− cells, which had been attributed to increased second-strand synthesis (30). However, transduction from the scAAV vector was increased to an even greater extent, suggesting that the previously observed effect on ssAAV may not have been entirely due to increased second-strand synthesis, since this would not have affected transduction from the already-double-stranded scAAV vector.

While transduction levels increased dramatically in ATM^−/− cells, the numbers of circularized genomes did not change significantly for either the ssAAV or scAAV vector, suggesting that ATM is not directly involved in productive circularization (Fig. 1a, hatched bars). These results further suggest that previous interpretations of increased transduction, or decreased circularization relative to transduction, in the absence of ATM may instead be viewed as loss of functional rAAV genomes in normal cells through interaction with ATM-mediated recom
The level of vector DNA.

of ATM on the numbers of rAAV genomes in the nucleus, we form.

degrad linear dsDNA or ssDNA but not supercoiled or fourth fraction was treated with plasmid-safe DNase, which create linear, circular, and total vector DNA (Fig. 2a). Thepared to digests with singly or multiply cut enzymes to delin-

tcrease at the molecular level, in either total vector DNA in the

cells compared to the wt, we observed no corresponding in-

While our transduction-based assay showed a >5-fold in-
crease exclusively from linear scAAV genomes in ATM^−/− cells compared to the wt, we observed no corresponding in-
crease at the molecular level, in either total vector DNA in the
nucleus, linear DNA, or circular forms (Fig. 2c, compare lanes 1 to 4 with lanes 9 to 12 for scAAV-GFP vector and lanes 5 to 8 with lanes 13 to 16 for scAAV-GFP-CD vector). Uncut unit-length linear genomes ran at approximately 2,432 bp (lanes 1, 5, 9, and 13), with uncut circular genomes just below (lanes 4, 8, 12, and 16). Single-cut linear genomes dropped to 1,713 and 1,631 bp, respectively, for scAAV-GFP and scAAV-GFP-CD (lanes 2, 6, 10, and 14). Single-cut circular genomes were converted to unit-length linear DNA, running at 2,432 bp (head-to-head and tail-to-tail concatemers were not detected in this experiment). Linear and circular genomes were quantified from the single-cut vector DNA in lanes 2, 6, 10, and 14. There was no evidence of a 5-fold increase in the linear form between the wt and ATM^−/− cells. The ratio of linear to circular genomes was approximately 2.6:1 in wt cells and 3.6:1 in the ATM^−/− cells, which was consistent with the expression levels in ATM^−/− cells but not in wt cells, where the majority of the expressed genomes were circular. Again, no differences between wt and ATM^−/− cells were observed in these products. Digestion with the double-cut enzyme converts all vector DNA to single bands of 2,123 and 1,218 bp for scAAV-GFP and scAAV-GFP-CD, respectively (lanes 3, 7, 11, and 15). While it was possible that some of the linear genomes detected were derived from vector DNA that had remained encapsi-
dated within the nucleus, the circular DNA could be formed only by interaction with cellular factors, and the ratio of cir-
cular to linear DNA was consistent with the excess transduc-
tion by linear vector genomes in ATM^−/− cells as determined by GFP expression.

Similarly, Southern hybridization of DNA from ssAAV-infected cells showed no significant differences between ATM^−/− and wt cells in the ratios of circular to linear genomes (data not shown). Because the amounts and ratios of circular and linear genomes did not change between wt and ATM^−/− cells, even though there were clearly more linear genomes being expressed in ATM^−/− cells, this suggested that the differential GFP expression may be a consequence of silencing of ATM-interacting vector genomes in normal cells.

An alternate possibility was that the high vector dose (MOI, 250) required for Southern blot detection saturated the effects of the ATM mutation such that there was no increase in transduction under these conditions. While we could not assess increased transduction simply by GFP expression at this dose because all of the cells are multiply transduced, we were able to test this notion in a competition experiment, where the doses of the GFP reporter vectors were kept constant at an MOI of 1 and increasing amounts of competing scAAV vector were added (Fig. 3). The competitor contained an irrelevant transgene and a liver-specific promoter, which was not active in these cells, to preclude transcriptional quenching effects. We observed competition for GFP transduction in wt cells beginning at a competitor MOI of 60. At MOIs of 120 and 240, GFP expression was reduced by half with each doubling of compet-
itor, indicating that we had reached saturation for the number of genomes that can be productively delivered to the nucleus. The previously observed increases in transduction by linear

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<th>Mutation</th>
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<tr>
<td></td>
<td>Transduction^a</td>
<td>Circularization^a</td>
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<td>DNA-PKcs</td>
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^a Positive values indicate an increased percentage of GFP-positive cells relative to the wt. Negative values indicate a decrease relative to the wt. A value of 1.0 indicates no change.

^b Fold difference in intact GFP vector transduction between wt and mutant cell lines (data are from Fig. 1, 4, and 5).

^c Fold difference in circularized genomes between wt and mutant cell lines (data are from Fig. 1, 4, and 5).

^d Fold difference in integrated genomes between wt and mutant cell lines (data are from Fig. 8).
vector genomes in ATM−/− cells relative to wt cells was clearly
evident at all concentrations of competitor vector, up to the
dose (MOI, 250) used in the Southern blot experiment. Thus,
if there had been an increase in linear vector DNA in the
nucleus matching the increased GFP expression from linear
vector in ATM−/− cells, it should have been detected in the
Southern blot.
Interestingly, although the increased transduction in
ATM−/− cells was evident at all vector concentrations, com-
petition for transduction in ATM−/− cells began earlier and
exhibited a higher slope than transduction in wt cells or trans-
duction with genomes that would ultimately be circularized.
This suggests that the pathway that leads to interaction with
ATM is more readily saturated than the pathway that leads to
productive circularization.

**Transduction and circularization in ATR-deficient cells.**
The PI3-like kinase ATR is activated by DNA damage involv-
ing significant regions of ssDNA, including UV lesions, signif-
icantly resected DSBs, and stalled replication forks. We there-
fore reasoned that the ssAAV was more likely to be a substrate
for ATR-mediated recombination than scAAV, because the
latter folds quickly into dsDNA after uncoating. The recombi-
nation efficiencies of the two kinds of vectors were compared in
isogenic cell lines that overexpressed either a kinase-dead
dominant negative ATR (ATR KD) or the wt ATR from a
Tet-inducible promoter. Doxycycline induction in the ATR KD
The cell line, GK41, leads to effective inhibition of ATR activity, while induction of wt ATR in GW33 cells has no effect (26). Doxycycline-induced overexpression of control wt ATR did not affect transduction or circularization of either type of vector (Fig. 4, wt ATR and wt ATR + DOX). As we had previously observed, overexpression of the dominant-negative ATRKD had no significant effect on the circularization of scAAV vectors, and the overall transduction efficiency was unchanged (Fig. 4, ATRKD and ATRKD + DOX, gray bars). In contrast, induction of ATRKD led to a significant (30%) increase in transduction from the ssAAV vector, similar to the effects of ATM on both vectors. Again, the number of circularized ssAAV genomes did not reflect the increase in active genomes available (Fig. 4, ATRKD and ATRKD + DOX, white bars). We concluded that ATR does not recognize the scAAV vector but does recognize and interact with the ssAAV vector. Similar to the effects of ATM, this interaction leads to a loss of functional genomes.

**Transduction and circularization in DNA-PKCS-deficient cells.** In order to determine whether the NHEJ pathway had differential effects on ssAAV and scAAV vectors, we also compared transduction and circularization in two cell lines with or without DNA-PKCS activity (M059K and M0597J, respectively) (Fig. 5). In contrast to the case for deficiencies in the HR pathways (ATM and ATR), transduction levels were greatly decreased for both ssAAV and scAAV intact GFP vectors in DNA-PKCS-deficient cells. Also in contrast to the

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**FIG. 3.** Transduction response to ATM−/− mutation with increasing vector dose. wt and ATM−/− cells were infected at an MOI of 1 with each of the four reporter vectors plus the indicated increasing amounts of competitor vector containing an inactive (liver-specific) promoter and irrelevant (mouse PD-L1) transgene. Expression of GFP was assayed at 24 h postinfection. Error bars indicate standard deviations.

**FIG. 4.** Transduction and circularization with ssAAV and scAAV in ATR-deficient cells. Cells containing doxycycline-inducible wt (wt ATR) or dominant-negative (ATR KD) ATR transgenes were infected with the four reporter vectors used for Fig. 1 at an MOI of 0.5 for 2 h. Doxycycline (DOX) treatment was initiated 48 h before infection and maintained through the experiment. GFP expression was quantified by flow cytometry at 24 h postinfection. The indicated (*) differences between wt and ATR-deficient cells were significant (P < 0.0010). Error bars indicate standard deviations.

**FIG. 5.** ssAAV and scAAV circularization assay with DNA-PKCS−/− deficient cells. Cells were infected with the four reporter vectors (see Fig. 1) at an MOI of 1 for 2 h. GFP expression was quantified by flow cytometry at 24 h postinfection. Means decreases for transduction in DNA-PK+ and DNA-PK− cells for each virus are all significantly different (P < 0.0001 for ssAAV-GFP, ssAAV-GFP-CD, and scAAV-GFP-CD; P = 0.0002 for scAAV-GFP). Error bars indicate standard deviations.
case for cells deficient in HR pathways, most of the GFP-expressing genomes were circularized in the DNA-PK CS/H11002/H11002 cell line.

Southern blots from infected M059K and M059J cells were inconclusive as to whether the decreased transduction in the DNA-PKCS cell line was reflected in the amount of vector DNA in the nucleus, largely due to a larger amount of vector taken up in the cytoplasm of the mutant cells (data not shown). This led to various levels of potential contamination of the nuclear fraction with cytoplasmic vector. However, it should be noted that while the DNA-PKCS mutant cells took up more vector in the cytoplasm, they displayed less transduction than the wt cells.

Because the M059K and M059J cell lines were not isogenic, we tested the effects of NU7441, a specific inhibitor of DNA-PKCS activity, on transduction and circularization in normal and ATM/H11002/H11002 fibroblasts to ensure that the observed differences in transduction were not due to unrelated differences in the infectivities of the two cell lines (data not shown). This led to various levels of potential contamination of the nuclear fraction with cytoplasmic vector. However, it should be noted that while the DNA-PKCS mutant cells took up more vector in the cytoplasm, they displayed less transduction than the wt cells.

Because the M059K and M059J cell lines were not isogenic, we tested the effects of NU7441, a specific inhibitor of DNA-PKCS activity, on transduction and circularization in normal and ATM/H11002/H11002 fibroblasts to ensure that the observed differences in transduction were not due to unrelated differences in the infectivities of the two cell lines (Fig. 6) (18). In each case, NU7441 treatment reduced transduction from both ssAAV and scAAV vectors, as had been observed in the mutant cell lines, although the change was not as great. We also tested the DNA-PKCS inhibitor in ATM/H11002/H11002 cells to determine how the two DSB repair pathways interacted (Fig. 6a to d). In ATM/H11002/H11002 cells, the DNA-PKCS inhibitor caused a decrease in transduction relative to that in untreated ATM/H11002/H11002 cells, similar to its effect in normal cells. Further, the inhibitor-treated ATM/H11002/H11002 cells showed the characteristic increase in transduction relative to inhibitor-treated wt cells. Again, the excess transduction was coming from vector genomes that remained linear. This suggested that that there was little overlap between the effects of ATM and DNA-PKCS on rAAV vector genomes; i.e., there appears to be a commitment to one or the other pathway early in infection, and the two pathways do not appear to complement or compete with each other, at least through the next 24 h.

Chromosomal integration in DNA repair-deficient cells. Having determined the effects of deficiencies in each of the three DSB signaling PI3-like kinases on circularization, we tested whether there were corollary effects on rAAV integration. For each cell line, integration was assayed by infection at a high MOI (approximately 260 to 1,400 HeLa cell infectious units per cell), followed by continuous passage until a stable percentage of GFP-expressing cells was reached, which represented the percentage of cells with integrated vector (Fig. 7).

The increased GFP transduction with both scAAV and ssAAV linear vectors in ATM/H11002/H11002 cells was not reflected in a corresponding increase in integrated vector (Fig. 8a). Indeed, the integration of ssAAV was significantly decreased compared to that in wt cells, while scAAV vector integration was not significantly changed. This decrease was surprising because the numbers of circularized and linear genomes, as determined by Southern blotting, had remained unchanged for both vectors in the presence and absence of ATM, suggesting that circularization and integration were not mediated through the same processes.

To test the effects of ATR on integration, either the wt ATR or ATRKD was induced with doxycycline at 48 h prior to infection and maintained for an additional 48 h postinfection. Doxycycline was then discontinued, because ATR activity is essential for long-term cell viability. Induction of wt ATR had no effect on the rates of integration of either vector, while
induction of ATR\textsubscript{KD} lead to a modest increase in integration for both the ssAAV and scAAV vectors (Fig. 8b). This suggested that ATR did not participate directly in the integration of either rAAV vector, despite its effects on the ssAAV genome at the transduction level.

Finally, the effect of DNA-PK\textsubscript{CS} deficiency on integration was tested in the M059K and M059J cells. Integration was reduced by 8.5-fold and 10.5-fold for the ssAAV and scAAV vectors, respectively, in the DNA-PK\textsubscript{CS}-deficient cells (Fig. 8c). While this generally followed the decreased transduction observed for both vectors in these cells, the effect on transduction with ssAAV vector had been greater than that with scAAV, converse to the effects on integration.

**DISCUSSION**

We have examined the roles of three principal factors orchestrating DSB repair in the circularization and integration of rAAV genomes. Our results suggest that there are qualitative and quantitative differences between ssAAV and scAAV vectors in terms of their interactions with different host DNA repair and recombination pathways. A striking and unexpected observation from these experiments is that deficiencies in DNA repair factors lead to significant changes in transduction efficiencies independent of dsDNA conversion and circularization. In the absence of ATM, transduction from the ssAAV and scAAV vectors increased by 2.6- and 6.6-fold, respectively (Table 1). In contrast, in the absence of DNA-PK, transduction is decreased 8.0- and 2.5-fold for the ssAAV and scAAV vectors, respectively.

Under the conditions used in these experiments, that is, in rapidly dividing cell cultures, rAAV genomes are acted upon by either HR or NHEJ pathways. Interactions with HR appear to lead to significant loss of functional genomes, as evidenced by the large increase in transduction with intact GFP vectors in ATM-deficient cells. This effect is greater for scAAV than ssAAV, arguing against a mechanism based strictly on conversion to dsDNA genomes, as had been previously suggested (30, 42). In the absence of ATM, the rescued genomes remain linear, seemingly trapped in this form and inaccessible to the NHEJ pathway, at least over the next 24 h. This suggests that the interaction choice between the major DNA repair pathways is fixed early after infection and is not easily altered. We also noted in our competition assay that the pathway that leads to interaction with ATM appears to be more easily saturated than the pathway that leads to productive circularization. Whether this suggests that the fate of the genome is determined through chance interactions with DSB repair factors that reside in separate compartments or that the different pathways are available at different stages of the cell cycle has not been determined.

A similar, though smaller, effect on transduction, specific for ssAAV genomes, is observed in ATR-deficient cells, suggesting a general negative outcome for interactions with HR pathways. Alternatively, it remains possible that ATR blocks conversion of ssAAV to dsDNA, since it does not affect scAAV genomes. However, it is clear that the fate of the majority of ssAAV vector genomes is determined by ATM, as evidenced by the greater increase in transduction in ATM-deficient cells than in ATR-deficient cells. This suggests that most interactions with HR repair pathways occur after dsDNA conversion, even in ATR-competent cells.

It is unlikely that the loss of GFP expression in HR-competent cells was due to excessive loss of sequence near the DNA ends, because the intact ssGFP vector used in these experiments contained 900 to 1,000 bp of stuffer sequences at each end, which would buffer the GFP cassette against imprecise end joining. Further, most circular rAAV junctions show end-to-end joining within the ITR sequences (11). Further, our Southern blot results are not consistent with extensive degradation of vector DNA, suggesting that the loss of GFP signal is due to silencing of gene expression in genomes interacting with the HR pathway. The possibility of a silencing mechanism is consistent with observations that ATM-mediated phosphorylation of histone H2AX is associated with reduced transcription in regions undergoing DSB repair (33). Further, parovirus and rAAV vector genomes have been demonstrated to associate with chromatin, and rAAV vector infection induces a slight H2AX phosphorylation (2, 7, 19, 27). Future experiments employing immunoprecipitation of rAAV with specific chromatin components may address these issues.

We had previously reported that ATM and related HR factors were important for scAAV circularization, based on the ratio of circularized to noncircularized genomes in wt versus mutant cells and on similar observations with vectors injected into muscle of mutant mice in vivo (4). While we still observe...
the decreased ratio of circularized to noncircularized genomes, our previous interpretation did not take into account the present observation that deficiency in HR factors leads to a net gain in functional genomes which do not circularize. Because the presence or absence of either ATM or ATR did not substantially alter the number of circular genomes, we now conclude that NHEJ is likely to be the principal pathway for rAAV genome circularization.

Converse to the effects of HR-mediated DNA repair pathways, interactions with the NHEJ pathway lead to circularization without loss of functional genomes, as evidenced by the large decrease in transduction efficiency in the absence of DNA-PK\(_{\text{CS}}\) activity. However, while the majority of functional genomes are lost in these cells, most of those remaining are circularized. It is not clear what mechanism is circularizing these genomes in the absence of DNA-PK\(_{\text{CS}}\) activity, but it does not appear to be ATM, since similar numbers of genomes are circularized in ATM\(^{-/-}\) cells in the presence of a DNA-PK\(_{\text{CS}}\) inhibitor as in normal cells with the inhibitor. This may be a consequence of nonclassical NHEJ, which is DNA-PK independent (39). Taken together, these results suggest that interaction of rAAV genomes with HR DNA repair pathways is costly in terms of functional genomes, while circularization through NHEJ is more sparing. This has important implications for gene therapy, because quiescent cells rely predominantly on the NHEJ pathway for DNA repair and make up the majority of relevant target cell types for rAAV applications.

Interpretation of the effects of the various PI3-like kinases on integration is hampered somewhat by the effects of the mutations on the number of functional genomes available in the nucleus and their circularization status, leaving few fixed points of reference. However, our results suggest that there is not a simple relationship between rAAV vector genome circularization and integration, as might have been expected if they were competing parallel processes mediated by the same factors. In ATM-deficient cells, the excess of functional, noncircularized genomes, with free ends available for recombination with chromosomal DSBs, does not lead to increased integration. However, if transcriptional silencing is indeed the mechanism responsible for the loss of functional genomes in normal cells, it remains possible that silencing of episomal vector DNA is relieved over time, and these genomes can subsequently be scored in the integration assay.

There are clearly differences between ssAAV and scAAV vector integration in terms of their responses to ATM or DNA-PK\(_{\text{CS}}\) deficiency. There is a significant decrease in the integration of ssAAV in ATM-deficient cells, while scAAV vector integration remains unchanged, despite the fact that ATM deficiency has a greater effect on the number of functional linear scAAV genomes in the nucleus. This suggests that

![FIG. 8. Integration of ssAAV-GFP and scAAV-GFP vectors in ATR\(^{-}\), ATM\(^{-}\), and DNA-PK\(_{\text{CS}}\)-deficient cells. (a) wt and ATM\(^{-}\) cells were infected at an MOI of 1,400 for 2 h. Cells were passed and GFP expression was quantified by flow cytometry until the percentage of GFP-positive cells reached a plateau at 22 days postinfection. The indicated (*) differences between wt and ATM-deficient cells were significant (\(P < 0.0038\)). (b) Doxycycline-treated and untreated wt ATR and ATR\(_{\text{KD}}\) cells were infected for 2 h at an MOI of 260. Treatment with DOX was maintained until 2 days postinfection. Cells were passed and GFP expression was quantified by flow cytometry until the percentage of GFP-positive cells reached a plateau at 25 days postinfection. Means for percent GFP-positive cells for ssAAV-GFP and scAAV-GFP vectors were not significantly different for wt ATR and wt ATR + DOX cells or for ATR\(_{\text{KD}}\) and ATR\(_{\text{KD}} + \text{DOX}\) cells. (c) DNA-PK\(^{-}\) and DNA-PK\(^{-}\) cells were infected at an MOI of 1,400 for 2 h. Cells were passed and GFP expression was quantified by flow cytometry until percentage of GFP-positive cells reached a plateau at 48 days postinfection. Means for percent GFP-positive cells for ssAAV-GFP and scAAV-GFP vectors are significantly different (\(P = 0.0018\) and \(P = 0.0088\), respectively). Error bars indicate standard deviations.]
ATM-mediated HR contributes productively to ssAAV integration but not scAAV integration. In DNA-PKcs-deficient cells, there are decreases in both the number of functional and the number of integrations. For ssAAV, the decrease in integration is similar to the decrease in functional genomes (8.5-fold and 8.0-fold, respectively). In contrast, the decrease in integrated scAAV genomes is greater than the decrease in transduction (10.5-fold and 2.5-fold, respectively). While it is not clear why there is a disproportionately large decrease in scAAV integration, it is consistent with the observations of Daya et al., who noted an overall decrease in integration but not scAAV integration. In DNA-PK CS-deficient ATM-mediated HR contributes productively to ssAAV integration and DNA-PKcs-deficient scAAV integration. J. Mol. Biol. 372:979–986.


