Site-Directed Mutagenesis of the Simian Virus 40 Large T-Antigen Gene: Replication-Defective Amino Acid Substitution Mutants That Retain the Ability to Induce Morphological Transformation

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We used a heteroduplex deletion loop mutagenesis procedure for directing sodium bisulfite-induced mutations to specific sites on viral or plasmid DNA to generate a series of SV40 large T-antigen point mutants. The mutations were directed to a region of the T-antigen gene, 0.5 map units, that is thought to be important for interaction of the protein with the viral origin of DNA replication. Of the 16 mutants reported here, 10 had lost the ability to replicate their DNA, and 3 others showed a reduced level of replication compared to wild type. All of the mutants tested were capable of transforming rat cells in culture by the dense focus assay. We conclude that the sequences of the early region around 0.5 map units are critical for the replication of viral DNA but not for the transformation function of T antigen.

The large tumor antigen (T antigen) of simian virus 40 (SV40) is a regulatory protein that is expressed early after infection and functions to initiate viral DNA replication (71), activate late region transcription (1, 12, 25, 26, 45, 55), and autoregulate early region transcription (1, 26, 53, 73). In addition, T antigen is capable of inducing the morphological transformation of a number of cells types in culture (3, 6, 19, 27, 28, 38, 44, 59, 68, 70, 72). During infection of both permissive and nonpermissive cells, T antigen induces a number of changes in cellular metabolism. For example, T antigen stimulates cellular DNA synthesis (7, 15, 21, 22, 57, 65, 75) and activates the transcription of certain cellular genes (66-68).

Three biochemical activities of T antigen have been described: a DNA-binding activity (41, 42, 52, 58, 74), an ATPase activity (8, 16, 76), and the ability to bind a host protein of 53,000 daltons, p53 (30, 33). The binding of T antigen to specific sequences at the origin of viral replication is thought to be required for the autoregulation of early region transcription, the stimulation of late region transcription, and the initiation of viral DNA replication. Furthermore, systems that replicate SV40 DNA in vitro are dependent upon T antigen (2, 32). The activities that function to induce morphological transformation have not been identified.

One approach to understanding the molecular mechanism of T antigen action is to correlate the biochemical activities of this molecule with its multiple biological effects. To this end, we have attempted to separate the activities of T antigen by mutating the T-antigen gene at specific sites. Experiments with deletion mutants that synthesize fragments of the protein have demonstrated that some activities can be dissociated (9, 15, 49, 65). These studies also indicated that certain activities seemed to reside in discrete functional domains of the protein. For example, the ability to stimulate cellular DNA synthesis requires residues between amino acids 82 and 272 and is unaffected by mutations outside this region (65). Similarly, the function that activates the transcription of silent rRNA genes resides in the central portion of the protein (65). On the other hand, the abilities to initiate viral DNA replication and to induce morphological transformation of cells in culture were sensitive to mutations in several regions of the molecule and were not dissociated from each other (49). This indicates that either the same biochemical activities of T antigen that induce transformation are required for viral DNA replication or the mutational lesions alter the protein structure such that independent activities are lost coordinately.

Results from Shortle et al. (62) and Margolskee and Nathans (37) have implicated the region of SV40 T antigen around 0.5 map units (m.u.) as being involved in viral DNA replication. Mutations at the origin of DNA replication leading to a reduced ability to replicate could be suppressed by mutations in T antigen, and two second-site revertants have been found to have lesions that map to 0.5 m.u. on the SV40 genome, changing amino acids 157 and 166, respectively. Additional evidence that this region of the protein is involved with the DNA replication activity has come from the studies of Stringer (69), Gluzman and Ahrens (18), and Manos and Gluzman (35, 36). Viral genomes isolated from SV40-transformed monkey cells (18, 35, 36) or rat cells (69) have been found to carry mutations in the T antigen gene that eliminate the ability of the protein to replicate its DNA while maintaining the transformation activity. Therefore, to determine whether in vitro-constructed mutations in this region would affect the DNA replication activity, and also to map the extent of this region, it was decided to target 0.5 m.u. of SV40 for site-directed mutagenesis. With a series of mutants it should be possible to characterize which biochemical activities are involved with DNA replication and transformation. A similar approach has been taken recently by Kalderon and Smith (24). We report here the isolation and characterization of in vitro-constructed amino acid substitu-

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tion mutants that eliminate the DNA replication function of T antigen while leaving the transformation activity relatively unaffected.

MATERIALS AND METHODS

Cells, viruses, DNAs, and bacterial strains. The small plaque strain of SV40 (77b) was grown in BSC40 cells, as described previously (4, 5). REF52 and Rat-1 (79) cells were obtained from W. Topp, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y., and grown in Earle minimum essential medium supplemented with 10% fetal bovine serum (HyClone; Sterile Systems, Inc., Logan, Utah). SV40 deletion mutants are described by Pipas et al. (49) and in references within their study. The plasmid pKP38 is pBR322 from which the Ava I and Sal I sites have been mutated by repairing the four unpaired nucleotides, followed by ligation, as described by Peden and Nathans (47). MM294 and BD1528 have been described previously (43, 47).

Preparation of plasmid DNA. Small-scale preparations of plasmid DNA were made by the method of Holmes and Quigley (23). Large-scale preparations were carried out as described previously (46, 47).

Construction of nucleotide substitution mutations. The procedure used followed closely that described previously (47). DNA from the plasmid pSVB40, a recombinant of wild-type SV40 and pKP38 joined through their BamHI sites, was digested with KpnI, and DNA from pSVd1137 and pSVd1138 was digested with Clal. KpnI-digested pSVB40 (2 μg) was mixed with Clal-digested pSVd1137 (2 μg) and, separately, with Clal-digested pSVd1138 (2 μg) in 1.620 μl of H2O, denatured by addition of 1 M NaOH (180 μl; final concentration, 0.1 M) and incubation at 37°C for 10 min, and neutralized by addition of 360 μl of neutralizing solution (1 M Tris hydrochloride [pH 7.2], 1 M HCl; 2:1; see reference 29). Renaturation was allowed to proceed at 60°C for 2 h. The extent of heteroduplex formation was determined by agarose gel electrophoresis, since these molecules migrate with form II species, and varied between 20 and 40% of the linear, homoduplex molecules. This mixture was concentrated by ethanol precipitation and dissolved in 200 μl of 10 mM Tris hydrochloride (pH 8)–1 mM EDTA.

Mutagenesis with sodium bisulfite was carried out as described previously (17, 47, 60). Between 50 ng and 1 μg of the heteroduplex mixture was incubated with 2.5 M sodium bisulfite–1 mM hydroquinone at 37°C under paraform oil in the dark for 1 h, followed by dialysis by the method of Shortle and Nathans (60). This mutagenized DNA was used directly to transform BD1528 to ampicillin resistance (31), and plasmid DNA was prepared from the pooled transformants (23). Full-length molecules were enriched by digestion of the deletion mutant DNA with Ava I and Sal I, and DNA was prepared from individual MM294 transformants. Those full-length molecules (ascertained by restriction enzyme mapping; HindIII plus TaqI for the pSVd1137 series, HindII for the pSVd1138 series) were subjected to sequence analysis. Fifty to 80% of those molecules sequenced had nucleotide changes.

DNA sequence analysis. Mutants were subjected to sequence analysis by the method of Maxam and Gilbert (39) after labeling the 3' ends of restriction fragments with Micrococcus luteus DNA polymerase I and the appropriately labeled [α-32P]dNTP and unlabeled dNTP. Cleavage products were resolved on 8% polyacrylamide gels (56). Mutants in the d1137 region were labeled with [α-32P]dGTP at the Ddel site at nucleotide 4499 (78), and the fragments were separated by electrophoresis on 6% polyacrylamide gels and recovered by the method of Maxam and Gilbert (39). Mutants in the dl1138 region were labeled with [α-32P]dCTP at the HinfI site at nucleotide 4376, and the fragments were separated by electrophoresis on 2.5% agarose gels and recovered on NA45 membranes by a protocol of Lizardi supplied by Schleicher & Schuell, Inc., Keene, N.H.

Indirect immunofluorescence in monkey cells. BSC40 cell monolayers in 5-cm dishes were transfected with 1 μg of plasmid DNA from the mutants by the DEAE-dextran method (40). The next day, the cells were trypsinized, plated in chamber slides, allowed to attach, and fixed in ethanol. T antigen was detected with hamster SV40 anti-T serum (50).

Viability and complementation tests. Viability was determined at 32, 37, and 40°C by transfecting BSC40 cell monolayers with 1 ng of excised and ligated DNA per 5-cm dish by the DEAE-dextran method (40). Plaques were scored at 32°C between 3 and 4 weeks and at 37 and 40°C between 12 days and 3 weeks.

Complementation tests were done by coinfecting BSC40 cell monolayers with mixtures of 10 ng each of excised and ligated mutant DNA and dl1007 DNA per 5-cm dish. Plaques were scored after 12 to 14 days at 37°C.

Marker rescue. Marker rescue was performed by using modifications of published procedures (18, 29). SV40 fragments used for the marker rescue were either cloned into pBR322 or purified from agarose gels by electrophoresis and DEAE-cellulose chromatography (63). Wild-type viral HindIII fragments were inserted into the HindIII site of pBR322, HaeIII fragments were inserted into the BstNI site, and AluI fragments were inserted into the PvuII site of pBR322. BamHI-digested mutant DNA (30 to 50 ng), HaeIII-digested plasmid containing the SV40 HaeIII C fragment (120 ng), and either the appropriately digested plasmid containing the fragment to be tested (120 ng) or the gel-puriﬁed fragment (10 to 20 ng) were mixed and then denatured by addition of 0.1 volume of 1 M NaOH and incubation at 37°C for 10 min. After neutralization by addition of 0.2 volume of neutralizing solution (see above), heteroduplex formation was allowed to occur (in a final volume of 260 μl) by incubation at 68°C for 30 min. DEAE-dextran (260 μl) was added to 250 μg/ml, and the mixture was used to transfect two 5-cm dishes of BSC40 monolayers. After 1 to 2 h at 37°C, minimum essential medium plus 10% fetal bovine serum was added, and the cells were incubated at 37°C overnight. The following day, the medium was replaced with agar for plaque determination as described above.

DNA replication in monkey cells. Viral DNA replication of the mutants was measured in BSC40 cells as described previously (48, 49). After digestion with DpnI plus BclI, DNA was fractionated by electrophoresis on 1% agarose gels and transferred to nitrocellulose (67), and viral sequences were detected by hybridization with 32P-labeled SV40 DNA (labeled by nick translation [34, 54]). Hybridization and washing of filters were done as described previously (46).

Transformation of rat cells. Rat embry fibroblasts (REF52) and Rat-1 cells (79) were seeded into 5-cm dishes at approximately 5 × 10^5 cells per dish the day before transfection. Mutant DNA (1 μg per dish) was introduced, using the calcium phosphate procedure (20) and glycerol shock (14).

Dishes were fed every 3 days with minimal essential medium plus 10% fetal bovine serum, and foci were scored at 3 weeks after transfection.
RESULTS

Mutant construction. Base substitution mutations were generated in the T antigen gene at 0.5 m.u. by the heteroduplex deletion loop mutagenesis method described previously (47). In this method, a heteroduplex is prepared between a deletion mutant and a wild-type molecule, and the cytosine residues in the resulting single-stranded loop at the position of the deletion become targets for deamination by sodium bisulfite. Two deletion mutants, d11137 and d11138 (49), were used as templates for mutant construction. The procedure is described above and outlined in Fig. 1.

Sequence determination of mutants. Figures 2 and 3 show the nucleotide sequence changes together with the predicted amino acid substitutions of mutants constructed with d11137 and d11138 as templates, respectively. Both cytosine-to-thymine and guanine-to-adenine transitions were found, as expected, since both strands are able to participate in heteroduplex formation. Of the 11 cytosine-guanine base pairs (bp) susceptible to sodium bisulfite in the exposed 31 nucleotides of d11137, 8 were changed. More importantly, of the six amino acids that could be changed by this method, four were mutated. In the case of the d11138 series, 11 cytosine-to-guanine bp were targets and 10 were mutated, resulting in the changing of five of the six susceptible amino acids. Those mutations that did not change the amino acid sequence are not included here, since several of this class of mutant had no detectable phenotypic alterations.

Cellular localization of T antigen by indirect immunofluorescence. The ability of the mutants to produce a stable T antigen was determined by indirect immunofluorescence with a hamster polyclonal SV40 anti-T serum after transfection of BSC40 monolayer cultures with plasmid DNAs. All mutants produced nuclear fluorescence with intensities approximately that of the wild-type SV40 plasmid, pSVB3. Mutants 5017, 5018, and 5028 produced weak cytoplasmic fluorescence as well.

Viability on BSC40 cells. The ability of each mutant to plaque on BSC40 cell monolayers was measured at 32, 37, and 40°C, and the ability to complement the late deletion mutant helper, d11007, was determined at 37°C (Table 1). None of the mutants except 5024 showed any temperature-dependent growth. Three mutants were viable and produced plaques with morphology indistinguishable from that produced by wild-type SV40 (5013, 5021, and 5032), and two mutants produced small plaques (5015, 5016). Mutant 5024 showed a slight cold-sensitive phenotype, producing very small plaques 4 to 5 days later than wild type at 32°C and 1 to 2 days later at 37°C.

All defective and partially defective mutants could be complemented by d11007, demonstrating that they are defective only in early function. Complementation plaques with the nonviable point mutants were smaller than those obtained with an early deletion mutant that produces no T antigen, such as d11136 (49). This could be due to interference with wild-type T antigen function by the mutant protein.

Marker rescue of mutants. To confirm that the phenotype of the mutants was a consequence of the nucleotide changes determined by DNA sequence analysis, marker rescue experiments were carried out. The locations of the fragments used and of the deletions in d11137 and d11138 are shown in
FIG. 2. Sequences of point mutants isolated by using dl1137 as template. The nucleotide sequence of SV40 DNA together with the corresponding amino acid sequence of T antigen from amino acid residues 121 to 133 is shown across the top. Nucleotides 4454 and 4422 are the limits of the deletion of dl1137. Mutant designations are in the left column, and the nucleotide changes for each mutant are shown. Amino acid substitutions are underlined.

FIG. 3. Sequences of point mutants isolated by using dl1138 as template. The format of the figure is as described for Fig. 2.
Fig. 4. Mutations in both the 1137 and 1138 regions were rescued (Table 2) by both the HindIII B fragment (nucleotides 5171 to 4002) and the RsaI B fragment (nucleotides 4876 to 4168) but not by HaeIII-C (nucleotides 2799 to 2259; the fragment used to cyclize the heteroduplex), HaeIII-E (nucleotides 5191 to 4862), or HindIII-D (nucleotides 4002 to 3476; data not shown), thus localizing the lesions to the RsaI B fragment. The AluI C fragment (nucleotides 4643 to 4314) rescued mutants 5014, 5017, and 5018, which were isolated by using di1137 as template, and some of the mutants isolated in the di1138 region (mutants 5024, 5026, and 5027). Since the end of the AluI C fragment lies within the deletion limits of 1138, it would be anticipated that not all the mutants would be rescued. In fact, mutants 5023 and 5025, which have mutations causing amino acid changes either outside or at the last nucleotide of the AluI C fragment, are not rescued. In the case of mutants 5020, 5022, 5026, and 5028, which have mutations leading to amino acid changes both within and outside AluI-C, only 5026 is rescued by AluI-C, indicating that the substitution Ala-169 to Val is inconsequential. This was confirmed by obtaining DNA from two such rescants and determining their DNA sequence. Both had the original cytosine-to-thymine transition at nucleotide 4312 of mutant 5026; this viable mutant is number 5035 in Fig. 3.

Hinfl-C (nucleotides 4376 to 3610) failed to rescue the lesions in the 1137 series mutants but rescued all those in the 1138 series, as predicted (Table 2). Thus, the mutations in the 1137 series map to 267 bp between nucleotides 4643 and 4376, and those in the 1138 series to 208 bp between nucleotides 4376 and 4168, or, if rescued by AluI-C, to the 62 bp between nucleotides 4376 and 4314. Since over 100 nucleotides of each mutant were sequenced and no additional alterations were found, we feel confident that the nucleotide substitutions of each mutant presented in Fig. 2 and 3 are responsible for the observed phenotypes.

DNA replication in BSC40 cells. To test whether the mutants retained the ability to replicate their DNA, we used the Dpnl assay described previously (48, 49). Examples of this assay are shown in Fig. 5, in which the replication efficiencies of wild-type SV40 and mutants 5024, 5018, 5015, and 5016 are compared. In 10 out of 16 cases, no DNA replication was detectable. Those mutants that produced plaques (5013, 5015, 5016, 5021, 5024) all replicated their DNA as expected. Since mutant 5032 produced plaques of wild-type morphology, it would also be expected to be positive in this assay although it was not tested. Mutant 5024, which formed late-appearing small plaques at 37 and 32°C, replicated its DNA to approximately 40% that of wild type at both temperatures (Table 3).

Morphological transformation of nonpermissive rodent cells. The ability of the T-antigen mutants to transform nonpermissive cells in culture was tested on two rat cell lines, the REFS2 line and the Rat-1 line, using the dense focus assay of Todaro and Green (77). All mutants, whether viable or not, were able to induce foci with various efficiencies on both cell lines (Table 3). Thus, the transformation function by this assay is relatively insensitive to mutations in the region of the T-antigen gene between nucleotides 4450 and 4492 and nucleotides 4330 and 4312.

**Table 2. Marker rescue results**

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Plaques per dish produced by the following restriction fragment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HindIII-C</td>
</tr>
<tr>
<td>5013</td>
<td>&gt;100. &gt;100</td>
</tr>
<tr>
<td>5014</td>
<td>0. 0</td>
</tr>
<tr>
<td>5015</td>
<td>&gt;100. &gt;100b</td>
</tr>
<tr>
<td>5016</td>
<td>&gt;100. &gt;100b</td>
</tr>
<tr>
<td>5017</td>
<td>0. 0</td>
</tr>
<tr>
<td>5018</td>
<td>0. 0</td>
</tr>
<tr>
<td>5019</td>
<td>0. 0</td>
</tr>
<tr>
<td>5022</td>
<td>0. 0</td>
</tr>
<tr>
<td>5023</td>
<td>0. 0</td>
</tr>
<tr>
<td>5024</td>
<td>&gt;100. &gt;100b</td>
</tr>
<tr>
<td>5025</td>
<td>0. 0</td>
</tr>
<tr>
<td>5026</td>
<td>0. 0</td>
</tr>
<tr>
<td>5027</td>
<td>0. 0</td>
</tr>
<tr>
<td>5028</td>
<td>0. 0</td>
</tr>
</tbody>
</table>

* ND, Not done.

**Table 3. Summary of biological activities**

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Amino acid substitutions</th>
<th>Viability</th>
<th>Replication (%) of wild type</th>
<th>Transformation (% of wild type)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5013</td>
<td>Arg-130 → Lys</td>
<td>+*</td>
<td>100</td>
<td>14</td>
</tr>
<tr>
<td>5014</td>
<td>Pro-124 → Ser</td>
<td>−</td>
<td>0</td>
<td>110</td>
</tr>
<tr>
<td>5015</td>
<td>Pro-126 → Leu</td>
<td>+*</td>
<td>20</td>
<td>58</td>
</tr>
<tr>
<td>5016</td>
<td>Pro-126 → Ser</td>
<td>+*</td>
<td>15</td>
<td>37</td>
</tr>
<tr>
<td>5017</td>
<td>Pro-125 → Ser, Pro-126 → Ser</td>
<td>−</td>
<td>0</td>
<td>115</td>
</tr>
<tr>
<td>5018</td>
<td>Ser-123 → Phe</td>
<td>−</td>
<td>0</td>
<td>60</td>
</tr>
<tr>
<td>5020</td>
<td>Thr-164 → Ile, Ala-168 → Val</td>
<td>−</td>
<td>0</td>
<td>32</td>
</tr>
<tr>
<td>5021</td>
<td>Ala-169 → Thr</td>
<td>+*</td>
<td>100</td>
<td>57</td>
</tr>
<tr>
<td>5022</td>
<td>Thr-164 → Ile, Ala-168 → Val</td>
<td>−</td>
<td>0</td>
<td>15</td>
</tr>
<tr>
<td>5023</td>
<td>Ala-168 → Thr</td>
<td>−</td>
<td>0</td>
<td>58</td>
</tr>
<tr>
<td>5024</td>
<td>Glu-166 → Lys</td>
<td>+*</td>
<td>40</td>
<td>23</td>
</tr>
<tr>
<td>5025</td>
<td>Ala-168 → Val</td>
<td>−</td>
<td>0</td>
<td>19</td>
</tr>
<tr>
<td>5026</td>
<td>Thr-164 → Ile, Ala-169 → Val</td>
<td>−</td>
<td>0</td>
<td>60</td>
</tr>
<tr>
<td>5027</td>
<td>Thr-163 → Ile, Thr-164 → Ile</td>
<td>−</td>
<td>0</td>
<td>40</td>
</tr>
<tr>
<td>5028</td>
<td>Thr-163 → Ile, Thr-164 → Ile</td>
<td>−</td>
<td>0</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>Ala-168 → Val</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5032</td>
<td>Ala-169 → Val</td>
<td>+*</td>
<td>ND</td>
<td>108</td>
</tr>
</tbody>
</table>

* Viability was measured as the ability to form plaques on monolayers of BSC40 cells at 37°C. replication was measured in BSC40 cells at 37°C, and transformation was measured by dense focus assay on rat cell lines REFS2 and Rat-1. The numbers of foci per dish on REFS2 cells varied between 80 and 110 with the wild-type SV40-pBR322 recombinant psV/B3, but on Rat-1 cells the numbers were 20 to 30 foci per dish. Data are shown only for the REFS2 cell line. Foci were present in assays on Rat-1 cells with all of the mutants listed except 5032, which was not assayed.

# DISCUSSION

We report the isolation and preliminary characterization of 16 new SV40 mutants with amino acid substitutions in the large T antigen. The mutations were targeted to a region near 0.5 m.u. on the viral genome, using the heteroduplex deletion loop mutagenesis procedure for directing sodium bisulfite-induced transitions. Nucleotide changes were determined by DNA sequence analysis, and in those cases in which a mutant exhibited an altered phenotype, marker rescue experiments were carried out to confirm the locations of the mutations responsible for the phenotype.
Three of the mutants (5013, 5021, 5032) had normal plaque morphologies and, where tested, replicated their DNA to levels similar to that of wild-type SV40. These mutants were also capable of inducing dense foci on REF52 cells. We conclude that substitutions of Lys for Arg at position 130 and of either Val or Thr for Ala at position 169 do not alter the ability of T antigen to support productive viral infection or transformation.

Three mutants (5015, 5016, 5024) displayed a reduced ability to replicate their DNA as well as an altered plaque morphology. Mutants 5015 and 5016 change Pro-126 to Leu or to Ser, respectively. Both formed small plaques relative to wild type at the three temperatures tested. Therefore, the ability to undergo a productive infection is reduced but not abolished in these mutants.

The mutation in 5024 results in the substitution of Lys for Glu at position 166. This same residue is changed to Gln in a mutant selected as a pseudorevertant of an origin-defective mutant (37, 62). Since the revertant produces a T antigen capable of functioning with a number of mutant origins (61) that are defective with the wild-type T antigen (37), it indicates that this region of the protein is important for the interaction with the origin of replication. Mutant 5024 has a cold-sensitive phenotype, forming small plaques that appear late at 32°C. Viral DNA replication is reduced to about 40% of the level of wild-type SV40 at both 32 and 37°C. Since 5024 is capable of replicating its DNA at 32°C but forms very small plaques at this temperature, we suggest that the T antigen of 5024 is defective in an essential function for productive infection other than DNA replication, such as activation of late transcription, processing or translation of late mRNA, or encapsidation.

The remaining 10 mutants were defective for plaque formation at all three temperatures tested, and all failed to replicate their DNA to a detectable level. Two of these mutants, 5014 and 5017, contain a substitution of a Ser for Pro at position 125: mutant 5014 carries the single substitution while 5017 has an additional change of Pro-126 to Ser. Thus, while replacement of Pro-126 with either Leu (mutant 5015) or Ser (mutant 5016) leads to a partially active protein, substitution of the adjacent Pro-125 with Ser results in a T antigen unable to replicate its DNA and unable to form plaques. Similarly, a substitution of Phe for Ser at position 123 (mutant 5018) and either Thr (mutant 5023) or Val (mutant 5025) for Ala-168 leads to the DNA replication-negative phenotype. Thr-164 may be critical for the replication function of T antigen, since mutant 5026, which carries the double substitution Thr-164 to Ile and Ala-169 to Val, is defective, while mutant 5032, which has the single substitution at position 169, is viable. Two mutants (5027, 5028) carry substitutions of Ile for Thr-163. However, the importance of this change cannot be assessed, since both also have the lethal replacement Thr-164 to Ile. In summary, we obtained amino acid substitutions at nine positions. Alterations at four of these residues, Ser-123, Pro-125, Thr-164, and Ala-168, destroyed the ability of T antigen to support the replication of viral DNA. Substitutions at another two positions, Pro-126 and Glu-166, reduced the level of viral DNA replication relative to wild type. Therefore, the replication function of large T antigen is very sensitive to mutations in this region of the protein.

In contrast, the ability of SV40 large T antigen to induce cellular transformation was not affected appreciably in any of the mutants so far tested. Of the 10 mutants that showed no detectable viral DNA replication in monkey cells, all were capable of transforming the REF52 rat cell line by the dense focus assay. We conclude that at least one of the activities of T antigen necessary for DNA replication in permissive cells is not obligatory to induce the transformed phenotype in nonpermissive cells and, therefore, the two activities are separable by mutation.

This conclusion is supported by the findings of others (18, 36, 39) that SV40 genomes isolated from transformed cells frequently possess point mutations between 0.45 and 0.5 m.u. of the T-antigen gene that result in the same phenotype. Replacements of Thr for Asn-153 (18), Gln for His-203 (36),
Glu for Lys-214 (69), and Glu for Lys-224 (36) also conferred the replication-negative, transformation-competent phenotype. Furthermore, Pearson-White and Nathans (personal communication) constructed a mutant, 1153, containing an additional Ile residue inserted between amino acids 143 and 144. This mutant also possessed the same phenotype.

Recently, Kalderon and Smith (24) described an extensive series of mutants constructed in vitro by a method analogous to the one used here and reached similar conclusions. From the combined results, it can be concluded that the region of T antigen where mutations can lead to the replication negative, transformation-competent phenotype can be extended to include amino acids 106 to 224. It remains possible, of course, that mutations in other regions of the protein will also result in this phenotype. In fact, Manos and Gluzman (35) have shown recently that substitution of Arg for Lys-516 or of Ser for Pro-522 results in the same phenotype and, therefore, this part of T antigen must be involved, either directly or indirectly, in DNA replication.

While the molecular basis for the mutational separation of the transformation and replication functions of T antigen is not understood, the simplest possibility is that these mutations destroy a biochemical activity of T antigen required for viral DNA replication while leaving those activities that mediate transformation untouched. A detailed examination of the biochemical properties of the mutant proteins will be necessary to differentiate between the possible mechanisms. Of interest in this regard will be whether the mutant T antigens bind specifically to the origin of SV40 DNA replication. Experiments with other replication-negative, transformation-competent mutant T antigens have shown that some are able to bind in a McKay assay (41) while others are not (36, 51), suggesting that the inability to bind to the origin of DNA replication cannot be the only mechanism that results in the replication-negative phenotype.

Mutants have been isolated that replicate their DNA but have a reduced potential to induce morphological transformation of nonpermissive cells. Cosman and Trevethia (11) described a mutant, tsA1642, that is temperature sensitive for lytic growth, replicates its DNA, but is unable to transform mouse embryo fibroblasts. Mutant 5002, which has two amino acid substitutions in the first exon of T antigen, is unable to transform REF52 cells but replicates its DNA to about 30% of wild-type levels (K. W. C. Peden, unpublished observations). These two mutants suggest that the replication activity may not be sufficient for the transformation of nonpermissive cells.

Finally, we point out that site-directed mutagenesis provides a powerful tool for introducing changes into regions of proteins thought to be of functional significance. Such regions may be identified by virtue of their primary structure or indicated by previous genetic analysis. Thus, specific hypotheses concerning structure-function relationships can be tested. Along these lines, we are currently directing mutations to other parts of T antigen such as the amino acids known to be phosphorylated (80), the regions conserved...
among the various papovavirus T antigens (13), the residues to which the ATPase activity has been localized (8, 10), and the major hydrophobic region of the protein.

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