

A DNA binding protein specific for an origin of replication of herpes simplex virus type 1

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ABSTRACT We have identified a protein that binds specifically to an origin of replication (*ori_S*) of the herpes simplex virus type 1 genome. The *ori_S* binding protein, detectable only in nuclear extracts of infected cells, shows the same time course of appearance as the herpesvirus-induced DNA polymerase and the DNA binding protein ICP8. The partially purified *ori_S* binding protein generates a DNase I "footprint" that spans 18- of the 90-base-pair minimal *ori_S* sequence. The *ori_S* binding protein may, therefore, be analogous to other origin-specific binding proteins that are required for the initiation of viral and chromosomal DNA replication.

The initiation of DNA replication occurs at well-defined sequences termed origins that act to regulate initiation as well as to facilitate assembly of the multi-enzyme complex required for semiconservative DNA replication (1). To understand the molecular mechanisms that underlie the initiation of DNA replication in eukaryotic cells, we have chosen to study herpes simplex virus type 1 (HSV-1). The herpesvirus family, whose members are highly cell-associated, appears to have genomic signals for lytic replication as well as for maintenance during latency. It is likely that they may encode functions that regulate the decision between these alternatives (2).

The 150-kilobase genome of HSV-1 contains elements, *ori_S* and *ori_L*, that can serve as origins of replication when isolated from the viral genome and are believed to be responsible for the initiation of DNA replication during lytic growth of the virus (Fig. 1) (2). The genome has two copies of *ori_S*, which are present within the reiterated *c* sequence of the S component, and one copy of *ori_L*, which maps in the unique sequences (*U_L*) of the L component (Fig. 1) (3-6). Plasmids containing cloned *ori_S* or *ori_L* sequences can replicate in mammalian cells superinfected with HSV-1 (4, 5, 7, 8). Analysis of these sequences has demonstrated that a 90-base-pair (bp) region of the *ori_S* sequence is sufficient to function as a minimal origin *in vivo* and that the nucleotide sequences of *ori_S* and *ori_L* show substantial homology in this region (5, 8, 9).

Our approach has been to identify viral proteins that specifically recognize the *ori_S* sequence and could, as a consequence, function in the initiation of HSV-1 DNA replication. Here we report on the identification and partial purification of a virus-induced DNA binding protein that is specific for the *ori_S* sequence.

MATERIALS AND METHODS

Cells and Viruses. RA305 (10), a thymidine kinase-deficient mutant of HSV-1 [F], was used to infect roller bottle cultures of Vero cells using a multiplicity of infection of 5-10 plaque-forming units per cell.

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Construction of *ori_S*-Containing Plasmids. The plasmid pON103 was constructed by inserting a 294-bp *Ava* I fragment spanning *ori_S* from pRB370 (11) into the plasmid vector pMT11 (12). This fragment, whose nucleotide sequence is identical to the comparable region of HSV-1 [17] (13), was positive for origin function by cotransfection with intact HSV-1 [F] into Vero cells (data not shown) as described (6). A 360-bp *Eco*RI-*Hind*III fragment from pON103 containing the *ori_S* sequence was introduced into M13 mp18 (Fig. 1) (14). The M13 mp18 *ori_S* phage were grown under standard conditions (15), and closed-circular duplex DNA was prepared by the alkaline lysis method (16).

Labeling of *ori_S*-Containing Fragment. The *Eco*RI-*Hind*III *ori_S*-containing fragment was labeled using the large fragment of DNA polymerase I and [α -³²P]dTTP (800 Ci/mmol; 1 Ci = 37 GBq) or [α -³²P]dCTP (3000 Ci/mmol) (16).

Construction of *ori_S* Deletion Mutant. The oligonucleotide CGCGAAGCGTGCCTGGCGC synthesized by the phosphoramidite method (17, 18) using a 380 A DNA synthesizer (Applied Biosystems, Foster City, CA) was phosphorylated at the 5' end by treatment with T4 polynucleotide kinase and [γ -³²P]ATP and purified by Sep-Pak chromatography (19). A 100-fold excess of the labeled oligonucleotide was annealed to M13 mp18 *ori_S* single-stranded DNA in 10 μ l of 0.01 M Tris-HCl (pH 8.0)/0.01 M MgCl₂/0.05 M NaCl (buffer A). An equal volume of 1 mM ATP/0.25 mM each of the four dNTPs/10 mM dithiothreitol in buffer A containing 5 units of the large fragment of DNA polymerase I and 1 unit of T4 DNA ligase was added. Incubation was for 12 hr at 12°C. The product closed-circular duplex DNA was isolated by electrophoresis in a 0.8% SeaPlaque agarose gel. It was cut out of the gel (which was melted), diluted with 2 vol of 20 mM Tris-HCl (pH 7.5), 1 mM EDTA, and used to transform competent *Escherichia coli* MC1061 cells (20). The M13 phage were grown in *E. coli* TG1 (21), and the single-stranded DNA was isolated by standard procedures. Dideoxy sequencing (22) of the mutant phage obtained by this procedure confirmed that the entire 45-bp inverted repeat present in the *ori_S* sequence had been deleted; it was designated M13 mp18 Δ *ori_S* (Fig. 1).

Partial Purification of *ori_S* Binding Protein from HSV-1-Infected Cells. HSV-1-infected cells from 50 roller bottles (20-30 g of cells), grown as described above, were harvested 18 hr after infection by shaking vigorously or by scraping with a rubber policeman and then collected by centrifugation. The cells were washed in 250 ml of cold 20 mM Hepes adjusted to pH 7.6 with NaOH/150 mM NaCl/0.5 mM dithiothreitol and subsequently lysed in 200 ml of 20 mM Hepes adjusted to pH 7.6 with NaOH/0.5 mM dithiothreitol/10 mM NaHSO₃, pH 7.0/0.5 mM phenylmethylsulfonyl fluoride/leupeptin at 2 μ g/ml (buffer B) using a Dounce homogenizer with a tight fitting pestle. Nuclei were collected by low-speed centrifugation and washed with buffer B. The nuclear pellet was finally suspended in 40 ml of buffer B containing 10%

Abbreviations: bp, base pairs; HSV-1, herpes simplex virus type 1; RF, closed-circular duplex DNA.

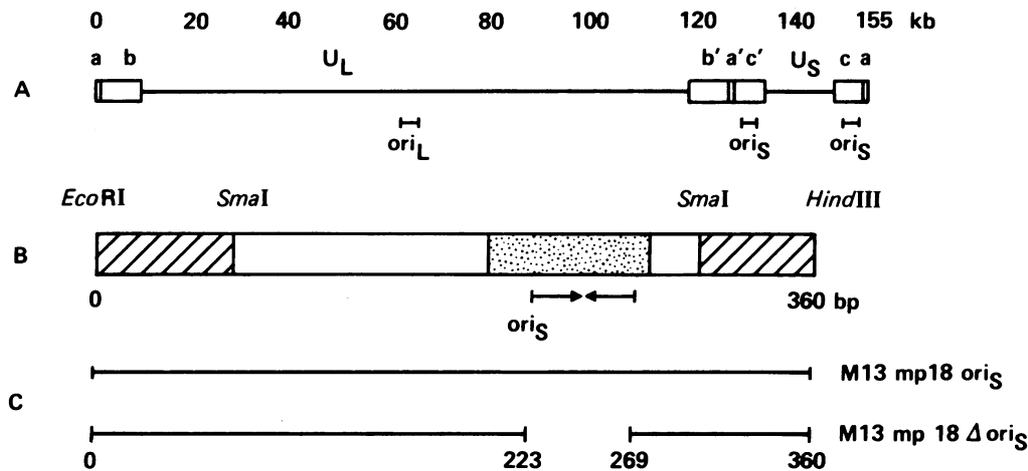


FIG. 1. Diagram of the HSV-1 genome, *ori_S* region, and M13 constructions. (A) The HSV-1 genome showing the location of the putative origins of replication. (B) Cloned fragment containing the *ori_S* sequence. The hatched area indicates poly-linker sequences bordering the HSV-1 *Sma*I-*Sma*I fragment (235 bp). The minimal origin is shown as a stippled box, and the arrows indicate the 45-bp inverted repeat. (C) Structure of the M13 mp18 derivatives used in this study. M13 mp18Δ*ori_S* has a 45-bp deletion of the entire inverted repeat.

(wt/vol) sucrose and stored at -80°C . Four such nuclear preparations were thawed and pooled by swirling in a large flask with an equal volume of 3.4 M NaCl in buffer B. The highly viscous extract was centrifuged at 27,000 rpm in a Beckman SW28 rotor for 2 hr at 4°C . Solid $(\text{NH}_4)_2\text{SO}_4$ (50 g/100 ml) was added to the pooled supernatants. The precipitate was collected by centrifugation for 1 hr at $20,000 \times g$ at 4°C and dissolved in a minimal volume of buffer B containing 10% (wt/vol) glycerol, 0.5 mM EDTA and dialyzed for 12 hr against the same buffer. The precipitate remaining after dialysis was removed by centrifugation, and the supernatant was adjusted to approximately 0.05 M NaCl by dilution with the dialysis buffer. This fraction (100–250 mg of protein in approximately 100 ml) was then applied to a phosphocellulose column (2.5×15 cm) equilibrated with 20 mM Hepes adjusted to pH 7.6 with NaOH/0.5 mM dithiothreitol/0.5 mM EDTA/0.5 mM phenylmethylsulfonyl fluoride/10% (wt/vol) glycerol (buffer C). The column was eluted with a linear gradient from 0.1 to 0.7 M NaCl in a total volume of 1 liter of buffer C. Fractions (5 ml) were collected and analyzed by NaDodSO₄/polyacrylamide gel electrophoresis (23). Assays for DNA polymerase and *ori_S* binding protein are described below. Fractions containing the *ori_S* binding protein were pooled and dialyzed against buffer C. The precipitate formed during dialysis that contained the *ori_S* binding protein was collected by centrifugation and dissolved in 2 ml of 0.5 M NaCl in buffer C. This fraction, which could be stored at -80°C for at least 2 months without significant loss of activity, was used for the experiments described below unless otherwise indicated.

Assays. ICP8 was identified as the predominant band with an apparent molecular weight of 130,000 by NaDodSO₄/polyacrylamide gel electrophoresis (24). It was quantitated by scanning Coomassie blue stained gels with a GS300 scanning densitometer (Hoefer, San Francisco).

The HSV-1 DNA polymerase was assayed in a reaction mixture (25 μl) containing 20 mM Tris-HCl (pH 7.5); 0.1 mM EDTA; bovine serum albumin at 0.04 mg/ml; 4% (wt/vol) glycerol; 3 mM MgCl₂; 5 mM dithiothreitol; 0.06 mM each of dATP, dGTP, and dCTP; 0.005 mM [³H]dTTP (40 Ci/mmol); 150 mM $(\text{NH}_4)_2\text{SO}_4$; and activated calf thymus DNA at 0.09 mg/ml (25). Incubation was for 10 min at 37°C , and the incorporation of labeled nucleotide into trichloroacetic acid-precipitable material was monitored as described (26).

The *ori_S* binding protein was assayed by nitrocellulose filter binding. The reaction mixture (25 μl) contained 50 mM Hepes adjusted to pH 7.5 with NaOH, 5 mM MgCl₂, 0.1 mM

EDTA, 0.5 mM dithiothreitol, 10% (wt/vol) glycerol, 100 mM NaCl, and 20 fmol of the ³²P-labeled *Eco*RI-*Hind*III *ori_S*-containing fragment. Sonicated calf thymus DNA (4 mg/ml, average size 600–900 bp) was added as a competitor. After incubation for 10 min at 25°C , the reaction mixture was diluted with 1 ml of 50 mM Hepes adjusted to pH 7.5 with NaOH/5 mM MgCl₂/0.1 mM EDTA/0.5 mM dithiothreitol/10% (wt/vol) glycerol (buffer D), and immediately filtered through 0.45- μm Millipore filters type HA. The filters were dried, and their radioactivity was determined using a toluene-based scintillation fluid. A background, which ranged from 2000 to 6000 cpm due to nonspecific retention of the labeled fragment, was subtracted.

DNase I "Footprinting." The *ori_S* binding protein was incubated with the ³²P-labeled *Eco*RI-*Hind*III fragment under conditions identical to those used in the standard binding assay except that the concentration of competitor DNA was reduced to 0.04 mg/ml. Following incubation for 10 min at 25°C , DNase I was added to a final concentration of 4 $\mu\text{g}/\text{ml}$. Digestion was interrupted after 1 min at 25°C by the addition of 1 ml of buffer D. DNA-protein complexes were isolated on Millipore filters as described above. The DNA fragments retained were eluted as described by Fuller *et al.* (27), extracted with phenol/chloroform, 1:1 (vol/vol), and analyzed on 6%, 0.2-mm sequencing gels (28). The positions of the DNase I cuts were identified by means of Maxam-Gilbert G and G+A reactions (29) as well as with standard restriction fragments.

RESULTS

Identification of *ori_S* Binding Protein. We have used a nitrocellulose filter assay to detect proteins that bind specifically to the restriction fragment containing the *ori_S* sequence. To eliminate the competitive effect of other DNA binding proteins a large excess of sonicated calf thymus DNA was added. As shown in Fig. 2, nuclear extracts from cells infected with HSV-1 contained an activity that promoted retention of the labeled *ori_S* fragment on nitrocellulose filters even in the presence of a 10,000-fold excess of calf thymus DNA. To reduce further the background due to nonspecific DNA binding proteins, we partially purified the *ori_S* binding activity from nuclear lysates of cells 18 hr after infection by phosphocellulose chromatography. As shown in Fig. 3, ICP8, the major DNA binding protein, eluted at about 0.15 M NaCl, and the virus-encoded DNA polymerase was eluted at 0.3 M

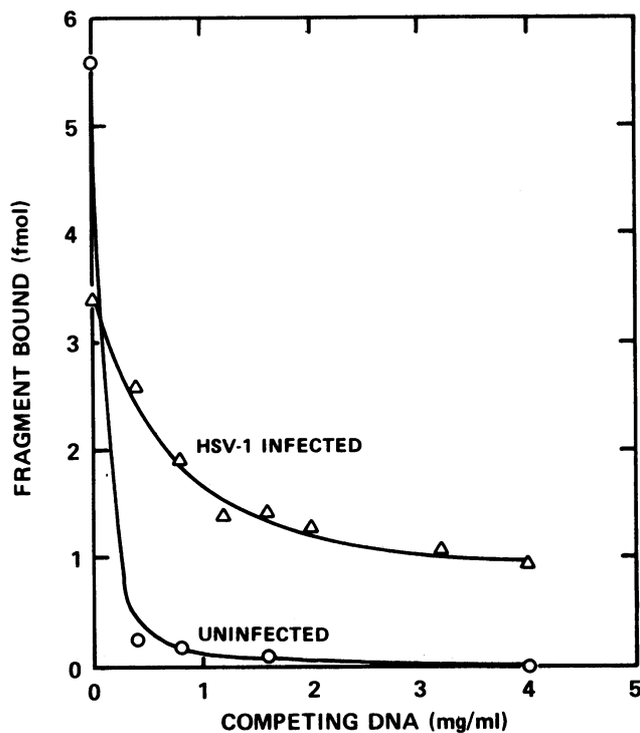


FIG. 2. Retention of the *ori_S* fragment on nitrocellulose filters. Nuclear extracts were prepared from uninfected cells (○) and from cells 12 hr after HSV-1 infection (△). Equivalent amounts of extract (5 μ l, 2 mg/ml) were used in the filter binding assay.

NaCl. The *ori_S* binding protein eluted at 0.5 M NaCl. The *ori_S* binding fractions were pooled and purified further.

Specificity of *ori_S* Binding Protein. The *ori_S* binding protein was specific for the 360-bp *EcoRI-HindIII* fragment containing *ori_S*. As shown in Fig. 4, M13 mp18 RF DNA lacking this sequence failed to compete for binding. Furthermore, M13 mp18 Δ *ori_S* RF DNA, in which the 45-bp dyad within the origin was deleted, showed little if any competition for binding with the wild-type *ori_S* sequence, indicating that critical binding sites are located within the dyad.

The specific sequence with which the *ori_S* binding protein interacts was determined by means of the DNase I "footprinting" technique (30). A complex was formed under conditions identical to those used in the binding assay except that the 360-bp fragment was end-labeled at its *HindIII* site with [α -³²P]dCMP. Following brief digestion with DNase I

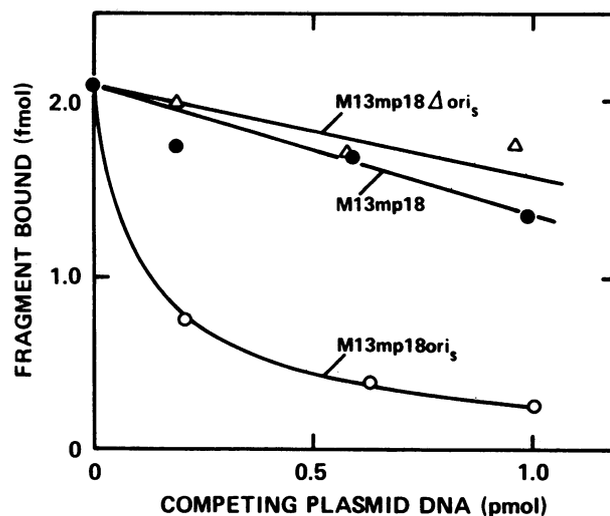


FIG. 4. Failure of *ori_S* deletion mutant to compete for binding to *ori_S* sequence. Phosphocellulose peak fraction (5 μ g) (Fig. 3) was used as the source of *ori_S* binding protein in the nitrocellulose filter binding assay. M13 mp18 RF DNA (●), M13 mp18*ori_S* RF DNA (○), and M13 mp18 Δ *ori_S* RF DNA (△) were added as indicated.

the complex was isolated and subjected to electrophoresis on a 6% sequencing gel. As shown in Figs. 5 and 6, a stretch of 18 bp was protected from DNase I digestion. The protected region starting at adenosine-219 and ending at cytidine-236 includes a portion of one of the inverted repeats and a short sequence immediately adjacent to its border. There was no evidence of protection of the remaining inverted repeat nor of any other sequence in this region.

Time Course of Appearance of *ori_S* Binding Protein. The time course of appearance of the *ori_S* binding protein following HSV-1 infection of Vero cells is shown in Fig. 7. *ori_S* binding activity was absent from nuclear extracts made from uninfected cells. Activity was, however, detectable at 8 hr after infection and reached a maximum at 18 hr. As shown in Fig. 7, the appearance of the *ori_S* binding protein coincided with two other proteins associated with HSV-1 DNA replication, the HSV-1 DNA polymerase and ICP8.

DISCUSSION

We have identified and partially purified a protein that specifically binds an origin of replication, *ori_S*, of the HSV-1 genome. This *ori_S* binding protein is detectable only in

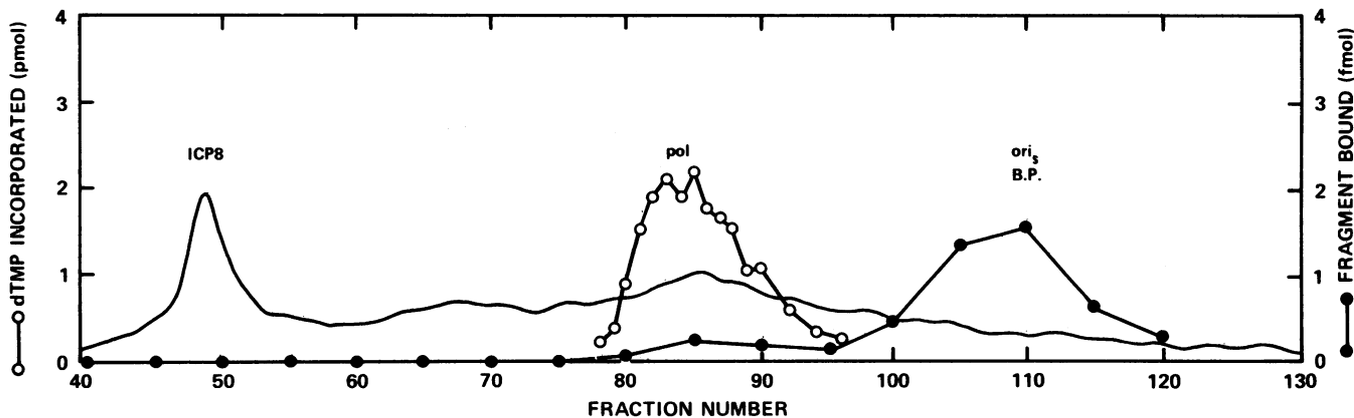


FIG. 3. Separation of ICP8, HSV-1 DNA polymerase, and *ori_S* binding protein by phosphocellulose chromatography. The solid line indicates the UV profile recorded continuously by an absorbance monitor. Solid line only, ICP8; open circles, HSV-1 DNA polymerase; and closed circles, *ori_S* binding protein (*ori_S* B.P.).

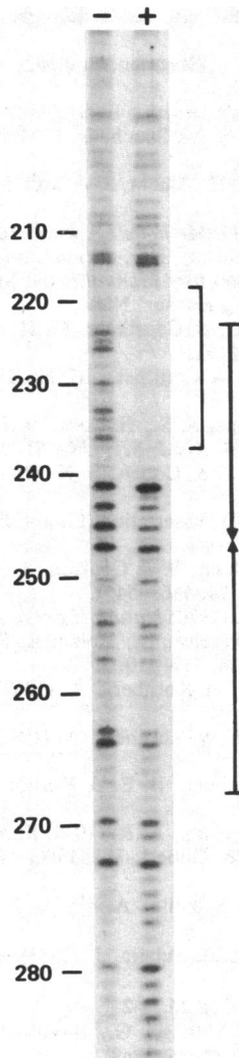


FIG. 5. "Footprint" of the *ori_S* binding protein on *ori_S*-containing fragment. DNase I digest of the complex between the *ori_S* binding protein and the *EcoRI-HindIII* *ori_S*-containing fragment was carried out. -, Control without protein; +, "footprint" obtained with 10 μ g of *ori_S* binding protein.

nuclear extracts from infected cells and accumulates during infection coincident with the HSV-1-induced DNA polymerase and the DNA binding protein ICP8. It may, therefore, be analogous to the sequence-specific binding proteins that are essential for the origin-dependent initiation of DNA synthesis in a variety of eukaryotic and prokaryotic cells (27, 31-34). The direct involvement of the *ori_S* binding protein in the initiation of DNA replication remains, however, to be established.

The partially purified *ori_S* binding protein produces a DNase I "footprint" covering 18 bp of the *ori_S* sequence.

HSV - 1 MINIMAL ORIGIN (*ori_S*)



FIG. 6. Sequence of the minimal *ori_S*. The position of the inverted repeat as well as the protected region is indicated.

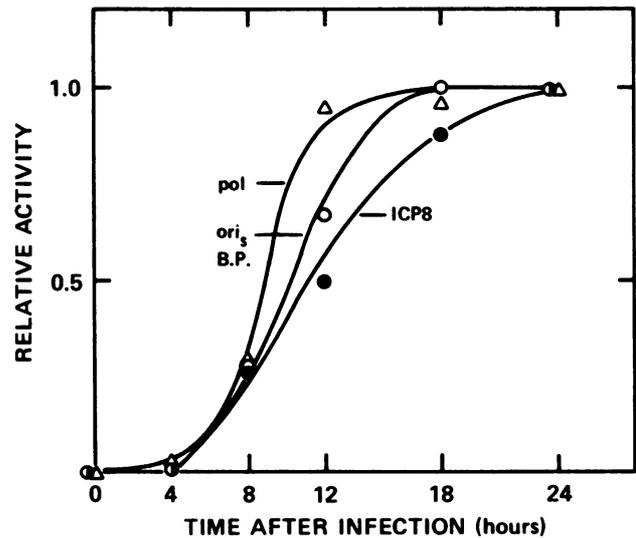


FIG. 7. Time course of appearance of *ori_S* binding protein following HSV-1 infection. Nuclei were prepared from Vero cells at various times after infection. The amount of ICP8 (●) was determined by densitometric scanning of Coomassie blue-stained NaDodSO₄/polyacrylamide gels. HSV-1 DNA polymerase (pol) (Δ) was assayed in the unfractionated nuclear extract. *ori_S* binding protein (*ori_S* BP) (○) was assayed after chromatography on a 0.2-ml phosphocellulose column using a step elution between 0.15 and 0.7 M NaCl in buffer C. Activities were normalized to the total activity found in nuclear extracts of cells harvested 24 hr after infection.

Fourteen of these are present in one part of the 45-bp palindrome that is a characteristic feature of the *ori_S* sequence (5). We have not observed protection of the other arm of the palindrome indicating that there is only a single binding site in the *ori_S* sequence. Since the two potential binding sites within *ori_S* differ at only three positions, it will be informative to determine whether point mutations introduced by site-directed mutagenesis can convert the active to an inactive site, or alternatively, produce two active binding sites within *ori_S*. Stow (35) has shown that an *ori_S*-containing plasmid with a 23-bp deletion in the center of the dyad is unable to replicate in HSV-1-infected cells. The fact that this plasmid lacks four of the nucleotides that were found to be protected in our "footprinting" experiment further supports a role for the *ori_S* binding protein in HSV-1 replication.

The minimal sequence required for replication of plasmid DNA in cells superinfected with HSV-1 (i.e., *ori_S*) consists of 90 nucleotides (5). It is noteworthy that one of the non-functional plasmids (S13) used in defining the minimal origin lacks two nucleotides of the sequence protected by the *ori_S* binding protein but includes the entire region of dyad symmetry (5). Inasmuch as this plasmid is unable to replicate, sequences immediately bordering the palindrome may be essential because of their interaction with the *ori_S* binding protein. The finding that sequences immediately outside the 45-bp region of dyad symmetry of *ori_S* are required for an interaction with the *ori_S* binding protein is also consistent

with our observation that a synthetic 20-bp duplex DNA fragment containing one-half of the dyad cannot compete with the 360-bp *ori_S*-containing restriction fragment in the nitrocellulose filter-binding assay (P.E., M.E.O'D., and I.R.L., unpublished observations).

The precise recognition site for the *ori_S* binding protein is still a matter of speculation. However, it is worth noting that an element of 2-fold symmetry exists within the protected sequence; as shown in Fig. 6, part of the sequence CGAAGCG occurs as an inverted repeat within this region.

In addition to *ori_S*, the HSV-1 genome contains another origin of replication, *ori_L*. The nucleotide sequence of *ori_L* is homologous to *ori_S*, and contains two equivalent sites within the 177-bp region of dyad symmetry that differ from the protected region of *ori_S* at only one position. These sites should endow *ori_L* with a 2-fold symmetry of binding that may be related to the bidirectional initiation of DNA replication (8, 9).

Finally, a comparison of the origins of replication in the HSV-1 and HSV-2 genomes reveals that the potential binding sites for the *ori_S* binding protein are completely conserved (36, 37).

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