Two Compound Replication Origins in *Saccharomyces cerevisiae* Contain Redundant Origin Recognition Complex Binding Sites

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While many of the proteins involved in the initiation of DNA replication are conserved between yeasts and metazoans, the structure of the replication origins themselves has appeared to be different. As typified by ARS1, replication origins in *Saccharomyces cerevisiae* are <150 bp long and have a simple modular structure, consisting of a single binding site for the origin recognition complex, the replication initiator protein, and one or more accessory sequences. DNA replication initiates from a discrete site. While the important sequences are currently less well defined, metazoan origins appear to be different. These origins are large and appear to be composed of multiple, redundant elements, and replication initiates throughout zones as large as 55 kb. In this report, we characterize two *S. cerevisiae* replication origins, ARS101 and ARS310, which differ from the paradigm. These origins contain multiple, redundant binding sites for the origin recognition complex. Each binding site must be altered to abolish origin function, while the alteration of a single binding site is sufficient to inactivate ARS1. This redundant structure may be similar to that seen in metazoan origins.

The replication of eukaryotic chromosomes initiates at multiple origins during each S phase. These DNA replication origins are best understood in the budding yeast *Saccharomyces cerevisiae*, in which they were initially recognized by their ability to promote the autonomous replication of plasmids. For this reason, they are referred to as autonomously replicating sequence (ARS) elements (29, 55). The paradigm *S. cerevisiae* replication origin is ARS1. It has a modular structure that spans about 120 bp and includes a small essential region, domain A, and three small accessory sequences, B1, B2, and B3, mutations in which reduce but do not abolish activity (40). Domain A, which encompasses the essential match to the 11-bp ARS consensus sequence (ACS), is the core of the binding site for the *S. cerevisiae* replication initiator protein, the origin recognition complex (ORC). The six-subunit ORC complex also contacts and protects DNA in the B1 element, and some mutations in B1 compromise ORC binding in vitro (3, 37, 52). The B3 element contains a binding site for the transcriptional activator-repressor Abf1p, which can be replaced by the binding sites for the transcriptional regulators Rap1p and Gal4p (40). The precise role of the B2 element has not been defined, although at least one of its functions is likely to be unwinding the DNA duplex to allow entry of the replication machinery (38, 41).

Other well-studied ARS elements, including ARS307 (49, 57), ARS305 (30), ARS121 (62), and the H4-ARS (7), seem to fit the ARS1 paradigm in that they contain a single, essential ACS flanked by a B domain. However, details of the structure of the B domain differ, and some ARS elements also contain stimulatory sequences on the other side of domain A, a region called domain C.

In this paper, we describe our characterization of two ARS elements, ARS101 and ARS310, which differ from the ARS1 paradigm in that they contain multiple ORC binding sites, all of which must be inactivated to abolish function. Following the precedent established by Hurst and Rivier (31), we will refer to these elements as compound ARS elements to distinguish them from ARS elements that have a single, essential match to the ACS. The occurrence of redundant ORC binding sites is reminiscent of the structure of *Schizosaccharomyces pombe* ARS elements, which appear to contain redundant functional elements (13, 21, 33, 44), and is similar to one model for the initiation zones detected at mammalian origins (reviewed in references 14 and 25).

MATERIALS AND METHODS

Strains. *Escherichia coli* strain DH5α (Life Technologies, Grand Rapids, N.Y.) was used for routine cloning, and strain GM2929 was used to prepare DNA lacking dam modification (46). Ura3-containing DNA was prepared from strain CJ236 (36).

*S. cerevisiae* strain 1C6 (ATCC 201543) was used for plasmid stability assays, which were performed as described previously (57). Strain YP45 (54) was used for the analyses of replication intermediates of ARS101 and its mutant derivatives. Strain CN31C, which lacks the 30-kb duplication present in strain YP45 of the region that includes ARS310 and the adjacent Ty element (43, 63; A. Dereshowitz and C. S. Newlon, unpublished data and data not shown), was used for analyses of ARS310 replication intermediates.

All transformations were performed by electroporation (2).

Plasmids. (i) ARS101. pLF34 was provided by David Kaback (Dept. of Microbiology and Molecular Genetics, UMDNJ-New Jersey Medical School, Newark, N.J.). The 1.5-kb *Kpn*I-*I* fragment of pLF34 was cloned into pBl KS (Stratagene); the 3.0-kb *Fsp*I fragment of this construct was ligated to the 4.7-kb *Fsp*I fragment of pRS326 (57) and the 2.6-kb *Fsp*I fragment of pRS306 (54) to yield yARS1KG and yARS1KG0, respectively. yARS1KG was isolated from *E. coli* strain GM2929; the 650-bp *EcoRV*-ClaI fragment was blunted with T4 polymerase (New England Biolabs [NEB]) plus deoxynucleoside triphosphates (dNTPs) (Roche Molecular Biochemicals), HindIII linkers (NEB) were attached, and the fragment was cloned into pRS326 to yield yAR1VCA, which contains the *EcoRV*-ClaI fragment in the same orientation as in yARS1KG. The *EcoRV*-ClaI fragment contains nucleotides 159457 to 161018 of the chromosome I sequence. The mutation in the 11 of 11 match to the ACS was made as described by Kunkel (36), and the mutation in the 9 of 11 match was made by fusion PCR (28). The

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11 of 11 match mutation introduces an XbaI site, while the 9 of 11 match introduction introduces an Msel site. yARS3CM (See Fig. 1E) was created by digesting the 9 of 11 match mutant, yARS5VCA-Msc, with Msel plus XhoI, blunting with T4 polymerase plus dNTPs, and recircularizing. yARS5VHA (see Fig. 1F) was created by first cloning the 310-bp HindIII-HindIII fragment of the 11 of 11 match mutant (blunted with T4 polymerase plus dNTPs) into the filled-in HindIII site of pRS326 and then digesting the resulting plasmid with XhoI and recircularizing.

Mutants constructed in the yARS5VCA backbone were transferred to the yARS5KG0 backbone for integration into the chromosome via a two-step process: First, the 530-bp PstI-KpnI fragment of the mutant was ligated to the 4.8-kb PnuII-KpnI fragment of yARS5G0. Next, the 1.5-kb NsiI fragment of the resulting plasmid was ligated to the 4.3-kb NsiI fragment of yARS5KG0. This step transferred the 150-bp PstI-NsiI fragment of the mutant into the yARS5KG0 backbone. Plasmids were linearized by digestion with Ehel to direct integration for two-step gene replacements (6), which resulted in a precise substitution of mutant sequences for wild-type (WT) sequences.

The 3.3-kb BamHI-EcoRV and the 2.6-kb BamHI-EcoRI fragments of pLF34 were cloned into pBS-KS to yield yARSleft and yARSright, respectively. These fragments were used to probe fork direction gels.

(ii) ARS310. BamHI links (NEB) were added to the 850-bp EcoRV fragment containing ARS310 (43), and the fragment was cloned into the BamHI site of pRS326. The EcoRV fragment contains nucleotides 166495 to 167340 of the chromosome III sequence. Using this backbone, mutant derivatives were made by the method of Kunkel (36), except for the 106-bp deletion, which was generated by fusion PCR (28). These alterations, indicated by lowercase letters, were introduced in the ACS match B (ATTTATCaAA), match C (TTTTACTCTtA), and match E (ATTTATGAGAT). The 2.2-kb BamHI-XbaI fragment of plasmid HB1-1 (43) was cloned into a derivative of pRS306 (54) deleted for sequences between the Kpnl and Smal sites of the polylinker to yield ARS310BX. To transfer the mutations into the chromosome, the 500-bp SpeI-HpaI fragment from the mutants was ligated to ARS310BX digested with the same enzymes. The resulting plasmids were linearized with XhoI to direct integration for two-step gene replacement (6) as was done for ARS101.

The 2.9-kb PstI-SacII fragment of K3B (43) was cloned into pRS304 (54) to yield 3100BF.

Analysis of replication intermediates. Preparation of genomic DNA and two-dimensional gel electrophoresis were performed as described previously (57). For the fork direction analysis of ARS310, 60 μg of DNA was digested with BamHI plus PstI. The reaction mix was adjusted to 1 M NaCl and subjected to benzoylated naphthoylated DEAE-cellulose (Sigma Chemical Co., St. Louis, Mo.) chromatography as described by Dijkwel et al. (18), except that all volumes were halved. Images were obtained on a Molecular Dynamics (Sunnyvale, Calif.) PhosphorImager (model PSI or 445SI). Fork direction gels were quantitated by drawing [32P]ATP (New England Nuclear) and polynucleotide kinase (NEB) and one unlabeled primer. For ARS101, the labeled primer was positioned 70 bp upstream of the closest match to the ACS, and the unlabeled primer was the T7 reverse primer. For ARS310, the labeled primer was positioned 95 bp upstream of the closest match to the ACS, and unlabeled M13 reverse primer was used. Footprinting reactions were performed as described by Richter-Klemm et al. (34). Recombinant ORC was a generous gift from Stephen Bell (MIT). Chemical sequencing reactions were performed as described by Richter-Klemm et al. (51).
To determine which of these weak ORC binding sites was responsible for ARS activity, the mutant 650-bp EcoRV-ClaI fragment was digested with Hinfl, and the two halves were cloned. Only the 310-bp Hinfl-ClaI fragment was Ars⁺ (data not shown), suggesting that the 9 of 11 match to the ACS (TTATATGTCCTA, nonconsensus positions in boldface) is responsible for the residual ARS activity in the mutant. To test this hypothesis, we altered this sequence to TTATATGgCcA (Msc mutant) in both the WT 650-bp fragment and the Xba mutant fragment. The double mutation abolished ARS activity (Fig. 1D), while the Msc mutation alone had only a modest effect on plasmid stability (Fig. 1C).

To assess the effect of these mutations on origin activity, they were used to replace the WT chromosomal copy of ARS101. While the WT origin was quite active, the double mutant was completely inactive, as indicated by the absence of bubble-shaped replication intermediates (compare Fig. 2A with 2D). As expected from their effects on plasmid stability, the mutation in the strong ORC binding site (the Xba mutant) caused a more dramatic reduction in origin activity than the mutation in the weak binding site (the Msc mutant) (compare Fig. 2B to 2C).

Modification of the 2D gel electrophoresis technique to include an in-gel digestion of replication intermediates between running the first and second dimensions allows one to separate the intermediates arising from a replication fork traversing a given fragment in one direction from those arising from a fork traversing the fragment in the opposite direction (22). In an attempt to quantitate the effects of these mutations on chromosomal replication origin activity, fork direction analyses
were performed in the regions flanking ARS101. In the case of ARS101, these analyses were complicated by two factors. First, when ARS101 was inactivated, forks initiated at flanking origins converged on this region, resulting in a termination zone. The double-Y termination intermediates can be seen in the 2D gel patterns of the mutants (Fig. 2). Second, a Ty1 element lies immediately to the right of ARS101 (Fig. 4A), necessitating the examination of fragments on the opposite side of this repetitive element. Figure 4 shows the fork direction analysis of the WT and double mutant on both sides of ARS101, and the quantitation of these and other results is presented in Table 1. In the patterns obtained from the WT ARS (Fig. 4D and E), the most intense signals are from forks moving away from the ARS, while in the double mutant (Fig. 4F and G) the patterns are reversed, with the most intense signals reflecting forks moving toward the ARS. The best estimate of the activity of WT ARS101 is provided by the analysis of the left side, because the fragment analyzed is directly adjacent to the ARS element. In this case, about 90% of the forks are coming from ARS101 in the WT, while only 20% are moving in the same direction in the double mutant, which is inactive, as judged from 2D gels. If the 20% signal seen in the double mutant contributes to the signal seen in the WT, it would suggest that ARS101 is active only about 70% of the time. The results of the analysis of the right side are consistent with this conclusion, with about two-thirds of the forks moving away from the ARS in the WT and only 8% of the forks moving away from the ARS in the mutant. Mutation of the 11 of 11 match to the ACS reduced chromosomal origin activity of ARS101 to about half of the WT level, while mutation of the 9 of 11 match had no significant effect on chromosomal origin activity in this analysis (Table 1).

**Inactivation of ARS310 requires mutation of three ACS matches.** We also examined ARS310 as part of our analysis of replication origins on chromosome III (15, 19, 42, 56, 57). ARS310 was localized to an 850-bp EcoRV fragment (43, 48). This fragment contains one 11 of 11 match to the ACS (match B, ATTTACATTTA) and three 10 of 11 matches (A, C, and E). Each match was mutated independently in a plasmid carrying the 850-bp fragment, and none of these mutations abolished ARS activity, though mutation of one of the 10 of 11 matches (match C, TTTTAC TTTT) dramatically reduced activity (Fig. 5 and data not shown). The match C mutation was paired with mutations in each of the other matches. One pair (B"C", Fig. 5F) abolished ARS activity while the remaining

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**FIG. 3.** In vitro ORC footprint of ARS101. The left panel shows the footprint of WT ARS101 labeled near the ClaI end (see Fig. 1A). The black arrow marks the position of the 11 of 11 match to the ACS, and the grey arrow marks that of the 9 of 11 match. The region of protection resulting from ORC binding is indicated by the bracket, and the solid black arrowheads mark the positions of two relatively unaffected sites within this region. Lanes labeled 0 contain no ORC protein, and the triangle indicates lanes with increasing amounts of ORC (12.5, 25, 50, and 100 ng for the WT, 100 and 200 ng for the Xba mutant). R and T, A+G and T sequencing lanes, respectively. The black box indicates a weak ORC binding site near the top of the gel. The right panel shows the footprint of the 11 of 11 match mutant. Notice the lack of protection over that match. Protection (region delineated by the bracket) is seen over the 9 of 11 match (grey arrow), and the open arrowheads mark the positions of three hypersensitive sites induced by ORC binding to the 9 of 11 match.
double mutant pairs had activities similar to that of the match C mutant alone (data not shown).

To examine the effect of the $B^{-}C^{-}$ double mutation on origin activity, a two-step gene replacement strategy was used. Two surprising results were obtained. The replacement construct, which carried the double mutation on a 2.2-kb BamHI-XbaI fragment (Fig. 5A), was weakly Ars$^{+}$ (data not shown), and the doubly mutant chromosomal origin was also weakly active, as evidenced by the presence of bubble-shaped intermediates on 2D gels (Fig. 6E). We also noted the presence of

**FIG. 4.** Fork direction analyses flanking ARS101. (A) The upper line depicts the 13.9-kb BamHI fragment containing ARS101. The positions of the SEN34 ORF and YARCTy1-1 are indicated by arrows. The position of the 650-bp EcoRV-ClaI fragment containing ARS101 is shown by the grey box below the line. Below this are the 5.6-kb BamHI-BglII fragment, used to examine fork direction to the left of ARS101, and the 4.5-kb BglII-BamHI fragment, used on the right. The arrows mark the positions of the EcoRV and EcoRI sites used for in-gel digestion prior to the second dimension. The probes used are indicated below these lines. As discussed by Friedman and Brewer (22), in-gel digestion allows one to distinguish rightward-moving replication forks from leftward-moving ones. How these replication intermediates are resolved depends on the geometry of the origin, the site of digestion, and the probe used to detect them. (B) Schematic diagram for fork direction analysis to the left of ARS101. Replication intermediates from leftward-moving (thick arrow) forks, including those emanating from ARS101, are shown as a thick line. Intermediates from rightward-moving (thin arrow) forks, i.e., those moving towards ARS101 are shown as a thin line. (C) Schematic diagram for fork direction analysis to the right of ARS101. Replication intermediates from rightward-moving (thick arrow) forks, including those emanating from ARS101, are shown as a thick line. Intermediates from leftward-moving (thin arrow) forks, those moving towards ARS101 are shown as a thin line. (D) Left side, WT. (E) Right side, WT. (F) Left side, double mutant. (G) Right side, double mutant.
TABLE 1. Quantitation of fork direction analyses of ARS101

| ARS101 derivative | Mean % of forks \( \pm \) SD |
|-------------------|-----------------|
|                   | Left of ARS101  | Right of ARS101 |
| WT                | 87 \( \pm \) 1   | 66 \( \pm \) 4   |
| 11 of 11 match mutant | 48 \( \pm \) 1   | 17 \( \pm \) 8   |
| 9 of 11 match mutant  | 95 \( \pm \) 4   | 75 \( \pm \) 7   |
| Double mutant     | 20 \( \pm \) 1   | 8 \( \pm \) 2    |

* Results are expressed as the percentage of forks coming from the direction of ARS101, i.e. from the right on the left side and from the left on the right side.

a replication fork pause site, indicated by intense spots on the Y arcs in Fig. 6E and F (arrows) that reflect accumulation of large Y-shaped replication intermediates. Replication fork pause when they encounter a tRNA gene whose direction of transcription opposes the direction of movement of the replication fork (16). The direction in which forks traverse this fragment when ARS310 is not active (see below) and the position and orientation of a glutamine tRNA gene (Fig. 6A) are consistent with the observed pause site.

The WT, B⁻, C⁻, and B⁻ C⁻ BamHI-XbaI fragments were transferred to a CEN vector, and ARS activity was quantitated. As indicated in Fig. 5B, plasmids carrying the WT ARS310 in the BamHI-XbaI fragment are considerably more stable than plasmids carrying the 850-bp EcoRV fragment. Since ARS activity was detected only in the 850-bp EcoRV fragment and not in flanking fragments (43), sequences present in the larger context must stimulate the activity of the minimal ARS310 fragment. Experiments are under way to characterize these stimulatory sequences (data not shown). These stimulatory sequences also act on the B⁻, C⁻, and B⁻ C⁻ mutants. The B⁻ mutant shows approximately the same plasmid stability as the WT in both fragment contexts (Fig. 5C), while the C⁻ mutant shows reduced plasmid stability in both contexts (Fig. 5D).

While the B⁻ C⁻ double mutant appears to be Ars⁻ in the 850-bp EcoRV fragment, extremely weak ARS activity is detected in the context of the BamHI-XbaI fragment (Fig. 5F). Presumably, in the absence of these stimulatory sequences, the ARS activity of the double mutant is reduced to the point where the primary transformants fail to give rise to colonies when streaked on selective plates. Hence, the double mutant appears to be Ars⁻ in the context of the 850-bp EcoRV fragment.

As in the case of ARS101, ORC footprinting was used to identify the sequences responsible for the residual activity of the B⁻ C⁻ mutant. In the WT fragment, ORC footprinted over the B and C matches, apparently binding with similar affinity at both sites (Fig. 7). Binding at both sites resulted in the induction of hypersensitive sites in similar positions relative to the ACS matches as seen in other ARS elements. The B and C matches to the ACS are separated by only 45 bp, yet ORC was able to bind at both sites simultaneously. Binding to these sites did not appear to be cooperative, since ORC bound to match B in the C⁻ mutant and to match C in the B⁻ mutant with affinities similar to those seen with the WT fragment.

In the B⁻ C⁻ fragment, as expected, ORC failed to bind to the mutant matches. Instead, binding was detected over another 10 of 11 match to the ACS (match E, ATTTATGTTAT), which is separated from match C by 26 bp (Fig. 7). This difference is most clearly seen in the region of the footprint delineated by the uppermost brackets. In the WT and B⁻ fragments, a pair of strong hypersensitive sites was induced at the bottom of this region and a weak pair of hypersensitive sites was induced at the top. The region between these pairs of hypersensitive sites was strongly protected. In the C⁻ and B⁻ C⁻ fragments, only the lower site of the pair of hypersensitive sites at the bottom of the region was induced. In these mutants, the hypersensitive sites at the top of the region were missing, as was the region of strong protection seen in the WT fragment. Instead, all the DNase I cleavage sites in this region decreased to the same extent. To confirm that the binding detected in the B⁻ C⁻ mutant is due to match E, the sequence was altered to create the B⁻ C⁻ E⁻ mutant. As shown in Fig. 7, ORC fails to bind to any of the mutated sites in this fragment.

The in vitro footprinting data suggested that the residual ARS and origin activity of the B⁻ C⁻ mutant was due to ORC recognition of match E. To confirm this hypothesis, the triple mutant was tested for ARS activity in the context of the 2.2-kb BamHI-XbaI fragment and found to be Ars⁻ (Fig. 5G). Similarly, replacing the WT ARS in the chromosome with the B⁻ C⁻ E⁻ mutant abolished origin activity (Fig. 5F). Thus, inactivation of ARS310 required mutation of three matches to the ACS within a 104-bp region. Despite being unaltered in the B⁻ C⁻ E⁻ mutant, the remaining 10 of 11 match (match A, TTICATGTTTA) apparently does not contribute to either ARS or chromosomal replication origin function.

To quantitate the effects of these mutations on origin activity, fork direction analyses were performed. No effects of these mutations were seen in the region to the right of ARS310, which was replicated by forks moving rightward through the
fragment (data not shown). The analysis of the region to the left of $ARS310$ is presented in Fig. 8 and Table 2. Figure 8A is a diagram of the 4.8-kb $PstI-BamHI$ fragment used for this analysis. The $BamHI$ site defining the right end of this fragment is the same as the site defining the left end of the 2.2-kb $BamHI-XbaI$ fragment shown in Fig. 5A. The position of the $SacII$ site used for in-gel digestion is indicated. The WT origin was very active, since nearly 90% of the forks came from the direction of $ARS310$, to the right of the fragment analyzed (Fig. 8C and Table 2). The mutations in matches B and C had little effect individually, but the $B^-C^-E^-$ double mutation dramatically reduced origin activity (Fig. 8D and Table 2). In the $B^-C^-E^-$ mutant, the fraction of forks coming from the right was even further reduced, to about 10% (Fig. 8E and Table 2). We conclude that the triple mutation completely inactivates the

**FIG. 6.** Chromosomal replication origin activity of $ARS310$ and its mutant derivatives. (A) Diagram of the 5.3-kb $HindIII$ fragment examined by 2D gels. The positions of the $RSG1$ and the included portion of the YCR026C ORFs are indicated by arrows. The position of a Gln tRNA gene is indicated by the arrowhead; this tRNA gene is in the correct position and orientation to cause the pause site which appears when $ARS310$ is inactivated. The grey box marks the position of the 0.85-kb $EcoRV$ fragment containing $ARS310$. The lines within this box indicate the positions of the three matches to the ACS, B, C, and E. (B) WT. (C) $B^-$ mutant. (D) $C^-$ mutant. (E) $B^-C^-$ double mutant. (F) $B^-C^-E^-$ triple mutant. Arrows in B to F point to a spot resulting from the accumulation of Y-shaped replication intermediates, indicative of a pause site.
chromosomal origin, based on the observation that similar frequencies of forks moving leftward through the fragment analyzed are detected in the 106-bp deletion mutant, in which all three matches were removed, and the triple mutant (Table 2).

**DISCUSSION**

We have described our characterization of two ARS elements, *ARS101* and *ARS310*, both of which are active as origins in their native locations. These ARS elements differ from those described previously in that multiple matches to the ACS must be altered to inactivate ARS and origin function. In previous analyses, alteration of a single match to the ACS was sufficient to abolish ARS function (7, 47, 56, 57, 61, 62). Despite the fact that *ARS1* contains multiple weak ORC binding sites (3), mutations in the exact match to the ACS, the preferred ORC binding site, abolish ARS activity (40). In contrast, in the case of *ARS101*, two matches to the ACS, separated by 8 bp, must be altered to eliminate ARS and origin activity. These two matches appear to represent independent ARS elements, as

![Diagram of ORC footprint analysis](image)

**FIG. 7.** In vitro ORC footprint of *ARS310*. The footprints of WT *ARS310* and four mutant derivatives, single knockouts of ACS matches B and C, the B’ C’ double mutant, and the B’ C’ E’ triple knockout mutant, are shown. The positions of ACS matches B, C, and E are marked by the arrows on the right side. The uppermost brackets mark the region where ORC binding to match C versus match E is readily distinguished. The middle and lower brackets mark the regions protected by ORC binding to matches C and B, respectively. Hypersensitive sites are marked by dots. Labels on individual lanes are as in Fig. 3. The amounts of ORC used for the WT fragment were 10, 20, 40, and 60 ng, and for the mutant fragments they were 20 and 60 ng.

| **TABLE 2. Fork direction analyses to the left of *ARS310*** |  |
|-----------------------------------------------------------|--|
| **ARS310 derivative** | **Mean % of forks** ± SD |
| WT | 88 ± 1 |
| B’ mutant | 83 ± 1 |
| C’ mutant | 78 ± 4 |
| B’ C’ double mutant | 24 ± 5 |
| B’ C’ E’ triple mutant | 10 ± 4 |
| Deletion mutant | 15 |

*Coming from the right, the direction of *ARS310*.
they can be cloned separately (Fig. 1). They do not contribute equally to activity, however. As one might expect, the 11 of 11 match to ACS is the preferred ORC binding site in vitro (Fig. 3), and alteration of this site has much more dramatic effects on both ARS (Fig. 1) and origin (Fig. 2) function. The contribution of the 9 of 11 match to the ACS in the WT ARS is unclear. While the 9 of 11 match mutation gave slightly reduced ARS activity, as measured by plasmid stability (Fig. 1), it had no significant effect on origin activity, as measured by 2D gel (Fig. 2) and fork direction (Table 1) analyses.

The analysis of ARS310 revealed that three matches to the ACS, one 11 of 11 match and two 10 of 11 matches, within a 104-bp region must be altered to inactivate both ARS and origin activity. Somewhat surprisingly, the 11 of 11 match does not seem to be the major contributor to activity. While the mutation of match B, the 11 of 11 match, caused little or no reduction in ARS activity, it was the mutation of match C which dramatically reduced activity (Fig. 5). However, this apparent dominance of match C was not mimicked by ORC binding, since ORC footprinted equally well over matches B and C in vitro (Fig. 7), nor by the effects of these mutations on origin activity, since the B<sup>−</sup> and C<sup>−</sup> mutants had similar small reductions in activity (Fig. 6 and Table 2). At this point, we do not understand the basis of the differential effects of the C<sup>−</sup> mutation in the plasmid and chromosomal contexts.

While these analyses do not allow us to assess the contributions of the individual ACS matches to origin function, it should be possible to do so utilizing the replication initiation point (RIP) mapping technique of Gerbi and Bielinsky (24). The chromosomal copy of ARSI shows a single initiation point located 30 bp from the ACS (4). Assuming that each match to the ACS of ARS310 specifies its own start site, it should be
possible to determine the frequency of initiation for each match in the WT origin by RIP mapping. Applying this technique to mutant ARS310 derivatives would also reveal whether each match uses its own start site or if they all use a common one.

The compound nature of ARS310 is not conserved in a closely related Saccharomyces species. We have analyzed several ARS elements from the homeologous chromosome III present in the brewing strain Saccharomyces carlsbergensis (58, 65). ARS310<sup>carl</sup> has a single essential match to the ACS which is in a short region of homology (18 of 21 bp) that includes match E in S. cerevisiae ARS310. Matches B and C are not conserved in ARS310<sup>carl</sup>. In S. cerevisiae, match E has a T-to-A transversion at position 10 of the ACS relative to its S. carlsbergensis counterpart. This change in the ACS is known to inactivate ARS307 (59), and the modification of position 10 strongly inhibits ORC binding at ARS1 (37). Therefore, it is not surprising that ORC has a lower affinity for match E than it does for matches B and C.

Having found these two unusual replication origins, we wondered how frequent such origins might be in the genome. We adopted a simple definition for a compound origin: functionally redundant ACS matches within the same intergenic region. For practical reasons, we used inter-ORF regions in our analysis. We realize that this is not a perfect definition. For example, the essential matches to the ACS for ARS604 and ARS605 reside within the BLM3 and MSH4 ORFs, respectively. In addition, the B3 element of ARS1 lies within the TRP1 ORF. Therefore, sequences important for origin function can reside within an ORF. However, single essential matches to the ACS have been defined for 22 ARS elements, and 20 of these fall within intergenic regions. The inactivation of ARS603 required the mutation of two closely spaced matches, both of which lie in an intergenic region. An analysis of these 23 ARS elements revealed four additional compound origins (Table 3). We suggest that, like the two fragments of ARS101, ARS601 and ARS602 should be considered a single compound element, as should ARS302, ARS303, and ARS320. These five ARS elements were given individual ARS designations because nonoverlapping subclones were shown to have ARS activity (43, 53, 61). However, given the close proximity of the essential ACSs (250 bp for ARS601/602 and approximately 600 bp for ARS302/303/320), it is unlikely that these clusters of ARS elements function independently. In fact, it has been reported that, both on plasmids and in the chromosome, ARS elements separated by as much as 6 kb interfere with each other, so that only one of the two ARS elements fires in any replication cycle (8, 9, 59). The sequences responsible for the activity of the HMRE ARS are less well defined than the others, but Hurst and Rivier (31) reported that three separate fragments, spanning 865 bp, have ARS activity, while Palacios DeBeer and Fox (45) extended this observation by demonstrating chromosomal replication origin activity for three fragments in HMRE.

In ARS603, the ACS matches are oriented so that each match lies within the first 16 bp of domain B of the other match, making it unlikely that nonoverlapping subclones with ARS activity could be found (53). ARS310 presents a somewhat similar situation. Matches C and E are in opposite orientations, separated by 26 bp. Therefore, this 26-bp interval is shared by the domain B regions of both matches. Matches B and C are in the same orientation, with match B only 45 bp upstream of match C. While the individual functional elements have not been defined experimentally, the domain B regions for the different ACS matches of ARS310 clearly overlap.

In summary, 6 of the 22 ARS elements examined are compound elements. This would suggest that one-quarter to one-third of the origins in the S. cerevisiae genome are compound origins. Could this frequency of compound origins arise by chance? We have analyzed the complete sequences of three regions of the yeast genome in which ARS elements have been identified systematically, chromosomes III and VI and a 131-kb region of chromosome XIV (23). This 716 kb represents about 6% of the genome. There are 33 ARS elements, of which 27 are detectably active as chromosomal replication origins, in these three regions, which also contains 359 inter-ORF regions (23, 42, 53, 64). Since less than 5% of the inter-ORF regions contain origins, the frequency of compound origins is much higher than expected. The origin-containing regions are distinguished from their counterparts neither by their size nor by the orientations of their flanking ORFs (C. S. Newlon, unpublished data).

While S. cerevisiae has proven to be an excellent model for many aspects of mammalian DNA replication, S. cerevisiae DNA replication origin structure has appeared to be different from that of other eukaryotes. ARS1, the paradigm S. cerevisiae origin, is small, about 120 bp (40), and has a single binding site for ORC, which contains the essential match to the ACS (3). In addition, replication initiates at a discrete site, as determined both by 2D gels (10) and RIP mapping (4, 5). In contrast to the highly specific replication initiation sites typical of S. cerevisiae, replication initiation events in mammalian cells appear to be distributed through large “initiation zones” (reviewed by DePamphilis [14]). The extreme case (out of more than 10 mammalian replication origins analyzed) is the well-studied dihydrofolate reductase (DHFR) replication origin of Chinese hamster ovary cells, in which 2D gel analyses detect bubble-shaped-replication intermediates throughout a 55-kb region (17, 60). Other approaches to mapping replication initiation sites, e.g., detection of the earliest-labeled fragments, nascent-strand abundance assays, and fork polarity assays, often reveal preferred initiation sites within these initiation zones. In the DHFR origin, these approaches have identified three preferred sites, oriβ, oriβ′, and oriγ (11, 26, 27, 35). While the detection of preferred initiation sites suggests the presence of multiple functional elements within initiation zones, it is also possible that a single element specifies initiation events anywhere within a broad region. Support for the presence of such an element in the DHFR origin has been provided recently by the finding that a 3.2-kb fragment at one end of the initiation zone appears to be required for all initiation activity within the

| ARS     | No. of matches | Spacing (bp) |
|---------|---------------|--------------|
| ARS101  | 2             | 8            |
| ARS310  | 3             | 82           |
| ARS603  | 2             | 5            |
| ARS601/602 | 2          | 241          |
| ARS302/303/320 | 3    | 580          |
| HMRE    | ≥3            | ≤865         |
The presence of multiple elements would clearly be analogous to the compound origins of *S. cerevisiae*, which are composed of multiple binding sites for the replication initiator protein ORC.

The analysis of *S. pombe* replication origins has provided clear examples of origins containing multiple functional elements. Individual *S. pombe* ARS elements are larger than their *S. cerevisiae* counterparts, 0.5 to 1.5 kb, and are themselves composed of multiple redundant elements. Detailed dissections have been performed on four ARS elements, *ars1* (13), *ars3001* (33), *ars3002* (20), and *ars2004* (44), and a feature common to all of them is the presence of redundant elements important for ARS function. It is not yet clear if this functional redundacy is analogous to that seen in the *S. cerevisiae* compound origins, i.e., multiple, redundant binding sites for ORC. However, two observations are particularly intriguing in this regard. First, these redundant elements are A+T rich, with a biased strand distribution, i.e., with one strand containing predominantly A and the other strand predominantly T, a pattern reminiscent of the biased strand distribution of A’s and T’s in the *S. cerevisiae* ACS. Second, Orp4p, a component of the *S. pombe* ORC homolog, binds DNA via AT hooks, a motif that recognizes A+T tracts (12), suggesting that the redundant elements might be binding sites for the replication initiator protein. On a larger scale, *S. pombe* also provides a precedent for the presence of multiple, separable ARS elements within a single replication origin. The *ura4* origin consists of a cluster of three ARS elements, *ars3002*, *ars3003*, and *ars3004*, within a 5.5-kb region (21).

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