SUM1-1, a Dominant Suppressor of SIR Mutations in Saccharomyces cerevisiae, Increases Transcriptional Silencing at Telomeres and HM Mating-Type Loci and Decreases Chromosome Stability

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Transcriptional silencing in the yeast Saccharomyces cerevisiae occurs at HML and HMR mating-type loci and telomeres and requires the products of the silent information regulator (SIR) genes. Recent evidence suggests that the silencer- and telomere-binding protein Rap1p initiates silencing by recruiting a complex of Sir proteins to the chromosome, where they act in some way to modify chromatin structure or accessibility. A single allele of the SUM1 gene (SUM1-1) which restores silencing at HM loci in strains mutant for any of the four SIR genes was identified a number of years ago. However, conflicting genetic results and the lack of other alleles of SUM1 made it difficult to surmise the wild-type function of SUM1 or the manner in which the SUM1-1 mutation restores silencing in sir mutant strains. Here we report the cloning and characterization of the SUM1 gene and the SUM1-1 mutant allele. Our results indicate that SUM1-1 is an unusual altered-function mutation that can bypass the need for SIR function in HM silencing and increase repression at telomeres. A sum1 deletion mutation has only minor effects on silencing in SIR strains and does not restore silencing in sir mutants. In addition to its effect on transcriptional silencing, the SUM1-1 mutation (but not a sum1 deletion) increases the rate of chromosome loss and cell death. We suggest several speculative models for the action of SUM1-1 in silencing based on these and other data.

Cytological studies have long suggested that eukaryotic genomes are organized into two distinct types of functional domains that can influence states of gene expression (reviewed in reference 40). In general, lightly staining euchromatic regions contain transcriptionally active or potentially active genes and are replicated early during S phase. In contrast, heterochromatic regions, where chromatin appears more condensed, are typically transcriptionally inactive and late replicating. The repressive effect of heterochromatin has been known and studied genetically for many years: transposition of euchromatic genes to regions next to heterochromatin can result in variable but heritable repression of the euchromatic gene, a phenomenon known as position-effect variegation (reviewed in reference 25). Position-effect variegation not only provides a means to study the nature of heterochromatin but may also reveal ways in which stable transcriptional states are normally established in euchromatic genes (28, 49).

A well-characterized example of position effect, in which the expression of a gene depends on its location in the chromosome, occurs in the yeast *Saccharomyces cerevisiae* at the silent mating-type loci *HMR* and *HML* (reviewed in reference 38). Mating type in this yeast is determined by the information present at the *MAT* locus, near the centromere of chromosome III. The *MAT***a** and *MAT* α alleles encode transcription factors that control the expression of cell-type specific genes, thereby imparting the **a** and α mating phenotypes of haploid cells and the nonmating, sporulation-proficient phenotype of **a**/ α diploids. Yeast cells typically have additional copies of **a** and α information stored at loci called *HMR* and *HML*, respectively. These loci, found near the right and left telomeres of chromosome

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Repression of the HM loci requires a number of trans-acting factors (including the four SIR genes) and cis-acting sequences that flank these regions, called silencers (1, 6, 17). The HMR-E silencer is found to the left of the HMR locus and is both necessary and sufficient for repression (6). HMR-E consists of three partially redundant regulatory sites (A, E, and B), which are binding sites for the origin recognition complex (4), Rap1p (9, 62, 63), and Abf1p (15). Genetic studies have demonstrated a direct role for both the origin recognition complex and Rap1p in silencing at HMR (3, 18, 34, 46, 68). Position-effect repression of genes at HM loci probably results from a modification of chromatin structure, since it requires the highly conserved N-terminal tails of histones H3 and H4 (29, 45, 51, 71). Repression by the HMR silencers is not specific to matingtype genes and can also affect RNA polymerase III transcription (7, 43, 61).

Telomeres in yeast cells are also subject to a position effect similar in many respects to that observed at HM loci. Genes placed near a telomere can be transcriptionally repressed (21) by a mechanism that requires RAP1, three of the four SIRgenes (SIR2, SIR3, and SIR4) and the N-terminal tails of histones H3 and H4 (2, 36, 47, 71). The SIR1 gene, which is required for stable silencing at HM loci, appears to play no role in the transcriptional repression of telomere-linked genes (2). Perhaps for this reason, transcriptional silencing at telomeres is normally unstable, resembling position-effect variegation in multicellular eukaryotes such as *Drosophila melanogaster*.

The role of SIR genes in silencing has been investigated by

the isolation and characterization of extragenic suppressors of *sir* mutations (31, 60). One such suppressor gene is *SUM1*, a single allele of which (*SUM1-1*) was isolated as a suppressor of *mar1-1* (*MAR1* is allelic to *SIR2* [31]). *SUM1-1* restores mating to *sir2* strains by restoring transcriptional repression at *HML* and *HMR* (41). *SUM1-1* is unique among suppressors of *SIR* mutations because it is able to suppress mutations in all of the *SIR* genes (37) and thus appears to bypass the requirement for *SIR* gene function in silencing. Significantly, though, *SUM1-1* still requires some *cis* and *trans* elements normally involved in silencing, since it will not bring about repression in strains carrying a deletion of the histone H4 N terminus or a deletion of the *HMR-E* silencer (37).

Initial studies indicated that the SUM1-1 allele is recessive (31). However, subsequent work using a different strain background showed that the SUM1-1 allele can be dominant to wild-type SUM1 (37). These two contrary results have led to different interpretations of the nature of the mutant phenotype and the function of the wild-type protein. Where the mutation was found to be recessive, it was proposed that SUM1-1 is a loss-of-function allele in a gene encoding an activator required for transcription at HM loci but not at MAT. In this model, the SIR genes were proposed to encode negative regulators of SUM1. One problem with this model is that it fails to explain why the HM loci should require a special activator (SUM1) not needed for transcription from the MAT locus. Furthermore, though loss-of-function mutations might be expected to arise frequently, SUM1-1 was the only allele isolated from a heavily mutagenized culture (31). On the other hand, where SUM1-1 appeared to be dominant to wild type, it was proposed to be a gain-of-function or altered-function mutation (37). In this case, it was imagined that SUM1 might encode a component of the repressed chromatin structure at the HM loci or an assembly factor involved in its formation. The SUM1-1 mutation might then alter or increase the protein's function such that SIR gene products would no longer be required to assemble repressed chromatin at HM loci.

We report here the cloning and characterization of the SUM1-1 and SUM1 alleles. Our results show that SUM1 is involved in both HM locus silencing and telomeric position effect. However, we rule out the model that SUM1 is an activator required for expression of HM loci, since a sum1 null mutation allows full expression from the silent mating-type loci in combination with sir mutations. Instead, our results are consistent with the proposition that the SUM1-1 allele is a partially dominant altered-function mutation that bypasses the need for SIR function at HM loci and increases repression at telomeres in SIR wild-type strains. Interestingly, the SUM1-1 mutation, but not a sum1 deletion, increases the rate of cell death and chromosome loss. As expected for a protein directly involved in transcriptional silencing, the SUM1 gene product is localized to the nucleus. Our results suggest possible molecular models for the function of SUM1 and SUM1-1 in both HM repression and telomeric silencing.

MATERIALS AND METHODS

Construction of genomic libraries from *SUM1-1* **strains.** Total genomic DNA from yeast strains JRY2465 and JRY2466 was partially digested with *Sau*3AI. DNA was separated on agarose gels, and fragments within the range of 10 to 16 kb were recovered by electroelution. YCp50 (a *URA3 CENIV* vector [56]) was cleaved with *Bam*HI, which is within the tetracycline resistance gene, and dephosphorylated with calf intestinal phosphatase. This vector DNA was then mixed at a roughly 2:1 ratio (by weight) with the size-fractionated genomic DNA, and the mixture was ligated at a total DNA concentration of either 12 or 25 ng/µl. Ligation mixtures were used to transform *Escherichia coli* DH5 α (22) by electroporation. Transformants were collected by scraping cells from the surface of the plates, using LB broth, and pooled. Four libraries were generated, each containing more than 20,000 independent transformants. More than 97% of the

plasmids contain inserted yeast DNA, as judged by tetracycline sensitivity of $DH5\alpha$ transformants.

Yeast strains, media, and genetic methods. The genotypes of the yeast strains described in this paper are listed in Table 1. All yeast genetic manipulations were performed as described previously (57). Yeast transformations were performed as described previously (54), using the tester strains YDS31 (*MATa*) and YDS32 (*MATa*), unless otherwise indicated. Quantitative mating assays were performed as described previously (66). Mating efficiency was calculated as the number of diploid cells (prototrophs) formed divided by the number of viable cells added to the tester strain. The reported efficiencies represent the mean of three independent assays per strain.

Yeast spot assays for tryptophan or uracil prototrophy or 5-fluoro-orotic acid (FOA) resistance (5) were done as follows. Overnight cell cultures were five times serially diluted by a factor of 10. Each dilution (5 μ l) was transferred to either control or test plates, and the cells were allowed to grow at 30°C for 2 days before the plates were photographed. Colony-forming ability was assayed by micromanipulating individual cells from overnight liquid cultures onto YEPD agar plates, which were then incubated for 5 days at 30°C.

Plasmids. Marking of the wild-type SUM1 allele for linkage analysis was done with plasmid DM268, in which a 1.2-kb PvuII-SpeI fragment from the SUM1-1 allele was cloned into pRS405, a LEU2-containing integrating plasmid (64). The plasmid was linearized by cleavage within the insert (SmaI) and used for yeast transformation. Note that through this integration process, the C terminus of the predicted SUM1 open reading frame (ORF) was truncated at codon 906. The sum1::URA3 mutation was constructed by inserting a HindIII fragment containing the URA3 gene into the SpeI sites of an AvrII fragment containing the complete SUM1 gene. This results in the removal of the entire predicted SUM1 coding region. The sum1::URA3 allele was subcloned into pBluescript II, creating plasmid DM264, and was released by digestion with both HindIII and XbaI before yeast transformation. sum1::LEU2 (DM286) contains the NsiI-Bg/II fragment of the SUM1 gene in pRS405, an integrating vector. The plasmid was used for yeast transformation after cleavage within the SUM1 insert at a unique NruI site. The integration creates a disruption by the LEU2-containing vector with a partial duplication (NsiI-BglII fragment) of the SUM1 coding sequence. (The resulting two SUM1 gene fragments consist of [i] the start codon to codon 581 and [ii] codon 340 to the end of the gene.)

The *SUM1* gene was tagged with an epitope from the influenza virus hemagglutinin (HA) protein for immunofluorescence studies. A 4.9-kb *Avr*II fragment containing the wild-type *SUM1* gene was cloned into the multicopy vector pRS425 (13), creating plasmid DM383, and a *Not*I site was generated just 5' to the termination codon of *SUM1* by PCR mutagenesis. The *NotI* fragment from plasmid GTEP, encoding an HA triple-epitope tag (72), was inserted to create DM651. The HA-tagged *SUM1* allele was tested for complementation in MC33, as well as expression of the tagged protein on immunoblots, before being examined by immunofluorescence microscopy.

Construction of isogenic *SUM1-1* **strains.** To replace the *SUM1* gene with the mutant *SUM1-1* allele, we first replaced the wild-type gene with the *sum1::URA3* deletion/insertion by one-step gene disruption (58). Subsequently, an *AvrII* fragment containing the *SUM1-1* allele was cotransformed with the 2µm-*LEU2* plasmid pRS425 (13) into the *sum1::URA3* strains. Leu⁺ transformants were replica plated to FOA plates to select for those cells in which the *SUM1-1* allele had replaced the *sum1::URA3* mutation. The putative *SUM1-1* strains were further characterized by Southern blotting analysis with multiple restriction digests to confirm that the *SUM1-1* fragment had replaced the normal *SUM1* locus. Strains MC54, MC57, MC105, and MC113 were constructed accordingly. Other *SUM1-1* strains were derived from crosses with these four parental *SUM1-1* strains. The *SUM1-1* genotype of segregants derived from a *SUM1-1* heterozygous diploid was scored by the following criteria: slow growth, loss of a *BgIII* restriction site present in the wild-type allele, and the ability to suppress mating defects of *MATa sir2* strains.

Chromosome stability assays. The rates of mitotic chromosome loss at chromosomes III and V were assayed in diploid strains (MC105 × MC113 and GA224 \times MCY2675), using fluctuation analysis, as described previously (52). Briefly, individual colonies of independent diploids were grown at 30°C on YPD medium to an average colony size of about 107 cells. Colonies were removed, resuspended in water, and plated on YPD plates for viable cells per colony. A portion of the cell suspension ($\sim 5 \times 10^5$ cells) was mated with approximately 10^7 cells of either mating-type tester strain in YPD broth for 4 h and then plated onto minimal medium. The mating-proficient cells included both \mathbf{a} - and α -mating cells. The same amount of cell suspension was also plated on synthetic complete medium with canavanine in place of arginine and later replica plated to complete medium lacking threonine. Seven colonies of each strain were assaved, and median chromosome loss frequencies for both chromosome III and chromosome V (number of maters and number of Can^r Thr⁻ cells in the total cell population) were determined. The chromosome loss rates (number of events per cell per generation) were calculated according to the following formula: rate = $(0.4343 \times \text{median frequency})/\log N - \log N_0$, where N is the number of cells present in the colony and N_0 (the number of initial cells) = 1 (16). For each genotype, three independent diploids were assayed by fluctuation analysis, and the average loss rate is reported.

TABLE	1.	Strains used	
INDLL	1.	Strams used	

IPY2515 MATe adeJ bits fast 2002 2025;12:HIS3 SUM1-1 ura3.52 J. Rine IPY2465 MATe adeJ 2016 bits 2000 lang bit 1 big-2001 sin2:HIS3 SUM1-1 ura3.52 J. Rine IRY2466 MATe adeJ 2016 bits 2000 lang bit 1 big-2001 sin2:HIS3 SUM1-1 ura3.52 J. Rine IRY2136 MATe adeJ 2016 bits 2000 lang bit 1 big-2001 sin2:HIS3 SUM1-1 ura3.52 J. Rine W305-1B Same adeJ 2016 bits 2000 lang bit 1 big-2001 sin2:HIS3 SUM1-1 ura3.52 J. Rine W305-1B Same ade W305-14. decempt MATe R. Rothtstein W305-1W W303-14 LEU2 R. Rothtstein W1089-1 W303-18 bits 1. EEU2 R. Rothtstein W1089-1 W303-18 bits 1. EEU2 R. Rothtstein W1089-1 JRY2515 sum1.2/LW3 J. Rine WC23 W303-18 bits 0mil.2/LW3 J. Rine WC34 W303-18 bits 0mil.2/LW3 J. Rine WC35 V15546 J. Rine J. Rine WC34 W303-18 bits 0mil.2/LW3 J. Rine J. Rine WC34 W303-18 bits 0mil.2/LW3 J. Rine J. Rine WC34 W303-18 bits 0mil.2/LW34 J. Rine J. Rine V15555 sum1.2/LW43 LU26-2 J. Rine	Strain	Genotype	Source
IRV2466 M47a ade2-101 kis 2200 laz bjr-1 bjr2-00 siz::H153 SUM1-1 ura3-52 J. Rine IRV2466 M47a ade2-101 kis 2200 laz bjr1-1 bjr2-00 siz::H153 SUM1-1 ura3-52 J. Rine IRV3181 M47a ade2-101 kis 2200 laz bjr1-1 bjr2-00 siz::H153 SUM1-1 ura3-52 J. Rine W303-1A M47a ade2-100 kis 3.112 bjr2-101 ura3-1 GAL* R. Rothstein W304-1A M47a ade2-1 cont.1 bjr3-115 kis 3.112 bjr2-101 ura3-1 GAL* R. Rothstein W305-1A Same as W303-1A except M47a R. Rothstein W305-1 W303-1A × W303-1B R. Rothstein W108-1 W303-1A × W303-1B R. Rothstein W108-1 W303-1A w303-1B R. Rothstein W108-1 W303-1B burn:Stal J. Rine W12935	JRY2515	MATα ade2 his3 leu2 sir2::HIS3 ura3	J. Rine
RY2466 MAT a adc2-101 his 2200 lar2 [pi-1 [ps2-00] siz::HIS3 SUM1-1 ara3-52 J. Rine RY2138 MAT a adc2-101 his 2200 lar2 [pi-1 [ps2-00] siz::HIS3 SUM1-1 ara3-52 J. Rine W303-1B Sam adc2-101 his 2200 lar2 [pi-1 [ps2-00] siz::HIS3 SUM1-1 ara3-52 R. Rothstein W303-1B Sam adc2-101 his 2200 lar2 [pi-1 ara3-12 [pi-1 ara3-12] R. Rothstein W303-1 W303-14 LEU2 R. Rothstein W108-1 W303-14 km12/LEU2 R. Rothstein W108-1 J. Rine R. Rothstein W108-1 J. Rine J. Rine W108-1 MC13 RY395 simi_2/LEU2 J. Rine W108-1 J. Rine J. Rine J. Rine W108-1 J. Rine J. Rine J. Rine W103-1 MC21 RY395 simi_2/LEA3 J. Rine W103-1 MC31 Simi_2/LEA3 J. Rine W123-1 MC31	JRY2465	MATα ade2-101 his3 Δ200 leu2 lvs1-1 lvs2-801 sir2::HIS3 SUM1-1 ura3-52	J. Rine
IRY3138 MATa ade2: 101 his3 2000 laz2 by:1 by:2-001 siz::H153 2001-1 unx3-52 J. Rine W303-1A MATa ade2: 1001 his3.1 J1 Strept-11 unx3-1 GAL' R. Rothstein W303-1B Same as W303-1A except MATa R. Rothstein W303-1A W303-1A Except MATa R. Rothstein W303-1A W303-1A R. Rothstein W303-1A W303-1A R. Rothstein W303-1A W303-1A R. Rothstein W303-1A W303-1B R. Rothstein W303 W304-1A W303-1B W303 W304-1A R. Rothstein W303 W304-1A R. Rothstein W303 W304-1B Rescaland W303 W304-1B Rescaland W303 YLS586 sum1:2UR43 Rescaland W303 YLS404 sum1:2UR43 Rescaland W407 W304-1A ar2:2H133 sum1:2UR43 Rescaland W407 W304-1A ar2:2H133 sum1:2UR43 Rescaland W407 W304-1A ar2:2H133 sum1:2UR43 Rescaland W4031 M2:2H133 sum1:2UR43 Rescaland	JRY2466	MATα ade2-101 his3 Δ200 leu2 lys1-1 lys2-801 sir2::HIS3 SUM1-1 ura3-52	J. Rine
W305.1A M4Ta udc2-1 cont.100 his?-11/5 km2.3.112 trp1-1 ura3-1 GAL* R. Rothstein W305.1B Same as W303.1A k 2002.11 R. Rothstein W305.1 W305.1A k 2002.12 J. Rine W305.1 W305.1B km12.02.03 J. Rine W305.1 W305.1B km2.02.04 J. Rine W305.1 W305.1B km2.02.04.01 J. Rine W305.1 W305.1B k km2.02.04.01 M. Rothstein W305.1 W305.1B k km2.02.04.01 M. Rothstein W305.1 W305.1B k km2.02.04.01 M. Rothstein W305.1 W305.1B k km2.02.04.01 A. Lastig W305.1 W305.1B k km2.02.04.01.01 A. Lastig W305.1 W305.1B k km2.02.24.07.01.04.01.01.01 A. Lastig W305.1 W305.1B k km2.02.24.07.01.01.01.01.01.01.01.01.01.01.01.01.01.	JRY3138	MATa ade2-101 his3 Δ200 leu2 lys1-1 lys2-801 sir2::HIS3 SUM1-1 ura3-52	J. Rine
W3031B Same as W303-1A except MATa R. Rothstein W303 W303-1A × W303-1B R. Rothstein W1089-1 W303-1A <i>LEU2</i> R. Rothstein W1089-1 W303-1A <i>LEU2</i> R. Rothstein MC25 JR Y2515 sun1-2:LEU2 R. Rothstein MC28 W303-1B sun2:LEU2 J. Rine MC28 W303-1B intra:SAI J. Rine MC31 JR Y3935 sun1-2:M43 J. Rine MC35 W303-1B intra:SAI J. Rine MC35 Y.LS404 sun1-2:M43 MC47 MC34 W303-1B intra:SAI sun1-2:M43 MC47 MC35 Y.LS404 sun1-2:M43 MC44 MC30-1A siz:HIX3 sun1-1:M243 MC54 W303-1A siz:HIX3 sun1-1:M243 MC54 W303-1A siz:HIX3 sun1-1 MC54 W303-1A siz:HIX3 sun1-1 MC51 W303-1A Siz:HIX3 sun1-1 MC54 W303-1A siz:HIX3 sun1-1 MC54 W303-1A Siz:HIX3 sun1-1 MC54 W303-1A suz:HIX3 sun1-1 MC51 W303-1A Siz:HIX3 sun1-1 MC54 W303-1A Siz:HIX3 sun1-1 MC54 W303-1A Siz:HIX3 sun1-	W303-1A	MATa ade2-1 can1-100 his3-11,15 leu2-3,112 trp1-1 ura3-1 GAL ⁺	R. Rothstein
W303 W303-1A X W303-1B R. Rothstein W1080-1 W303-1A LLU2" R. Rothstein MC25 JR Y2515 suml 3::LEU2 R. Rothstein MC27 W303-1A suml:UR43 J. Rine MC38 W303-1B burn:Sull J. Rine MC31 JR Y3935 W303-1B burn:Sull J. Rine MC33 Y15866 W303-1B burn:A::DD2 J. Rine MC33 Y15868 W303-1B burn:A::DD2 J. Rine MC34 W303-1B burn:A::DD2 J. Rine J. Rine MC35 Y15404 suml::DR43 J. Rine J. Rine MC34 W303-1A siz::HIS3 SUM1-1 J. Suml::DR43 YDS714 W303-1B siz::HIS3 SUM1-1 YDS714 W303-1B siz::HIS3 YDS74 W303-1B UR43/4DD2-TeVIII. SUM1-1 A. Lustig MC61 W303-1A UR43/4DD2-TeVIII. SUM1-1 MC61 W303-1A UR43/4DD2-TeVIII. SUM1-1 MC61 W303-1B UR43/4DD2-TeVIII. SUM1-1 MC61 W303-1B UR43/4DD2-TeVIII. SUM1-1 MC61 W303-1B UR43/4DD2-TeVIII. SuM1-1 MC61 W303-1B UR43/4DD2-TeVIIII. SuM1-1 MC61 </td <td>W303-1B</td> <td>Same as W303-1A except $MAT\alpha$</td> <td>R. Rothstein</td>	W303-1B	Same as W303-1A except $MAT\alpha$	R. Rothstein
W1089-1 W303-LA LEU2" R. Rothstein MC25 JK Y2515 sum1-SUR43 J MC31 W 303-LB sum1-UR43 J MC31 JK Y3255 sum1-UR43 J MC31 JK Y3255 sum1-UR43 J MC33 Y LS868 sum1-UR43 J MC34 W 303-LB Imm2b::LD42 J MC35 Y LS868 sum1-UR43 J MC36 W 303-LB Imm2b::LD42 J MC47 W 303-LB Siz:MI35 sum1-UR43 J MC47 W 303-LB Siz:MI35 sum1-UR43 J MC47 W 303-LB Siz:MI35 sum1-UR43 J MC54 W 303-LB Siz:MI35 sum1-UR43 J MC54 W 303-LB Siz:MI35 SUM1-I J MC51 W 303-LB Ziz:MI35 SUM1-I J MC52 W 303-LB Ziz:MI3 SUM1-I J MC51 W 303-LB Ziz:MI35 SUM1-I J MC52 W 303-LB Ziz:MI35 SUM1-I J MC54 W 303-LB Ziz:MI15 SUM1-I J MC54 W 303-LB Ziz:MI15 SUM1-I J MC54	W303	W303-1A \times W303-1B	R. Rothstein
MC25 JRY251S suml.2:LEU2 MC27 W305.1A suml.:URA3 MC28 W305.1B suml.:URA3 MC31 JRY3935 MC33 VLS586 MC33 VLS586 MC33 VLS586 MC33 VLS586 MC33 VLS586 suml.:URA3 MC34 W303.1B sinzt.:URA3 MC35 VLS404 suml.:URA3 MC47 W305.1B sizz.:HISS suml.:URA3 MC44 W305.1B sizz.:HISS suml.:URA3 MC44 W305.1B sizz.:HISS suml.:URA3 MC44 W305.1B sizz.:HISS suml.:URA3 MC47 W305.1B sizz.:HISS suml.:URA3 MC47 W305.1B sizz.:HISS suml.:URA3 MC47 W305.1B sizz.:HISS suml.:URA3 MC47 W305.1B sizz.:HISS suml.:URA3 MC51 W305.1B sizz.:HISS suml.:URA3 MC52 W305.1B URA3.:ADE2.74/ULL SUMI-1 MC53 W305.1B URA3.:DE2.74/ULL SUMI-1 MC54 VD563 suml.:LEU2 VDV66 W305.1B URA3.:TEVIL VDV66 W305.1B URA3.:TEVIL VDV66 W305.1B URA3.:TEVIL VDV66 W305.1B URA3.:TEVIL	W1089-1	W303-1A <i>LEU</i> 2 ⁺	R. Rothstein
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MC28 W305-B5 num?::SAI J. Rine MC31 IRY3935 sun1::URA3 J. Rine MC31 IRY3935 sun1::URA3 J. Sine MC33 VLS586 sun1::URA3 J. Sine MC33 VLS586 sun1::URA3 J. Sine MC35 VLS404 wolo-1B funzAwit.2DE2 J. Sine MC37 W303-B5 in:2:URA3 J. Sine:URA3 MC47 W303-B5 in:2:URA3 J. Sine:URA3 MC44 W303-B5 in:2:URA3 J. Sine:URA3 MC54 W303-B5 in:2:URA3 J. Sine:URA3 MC54 W303-B5 in:2:URA3 J. Sine:URA3 MC57 W303-B5 in:2:URA3/ADE2-7eUTIL A. Lustig MC51 W303-B5 in:2:URA3/ADE2-7eUTIL J. Sine:URA3/ADE2-7eUTIL MC51 W303-B5 in:2:ASIADE2-7eUTIL J. Sine:2:URA3 J. Sine:URA3/ADE2-7eUTIL MC51 W303-B5 in:2:URA3/ADE2-7eUTIL J. Sine:URA3/ADE2-7eUTIL J.	MC27	W303-1A sum1::URA3	
JRY395 W305.18 hmr.:SAI J. Rine MC31 JRY3955 sumi::URA3 J. Rine MC38 W1586 W305.18 hmr.Abr.:24DE2 MC33 YL586 W1586 sumi::URA3 MC44 W303.18 hmr.Abr.:24DE2 MC35 MC35 YL5404 sumi::URA3 MC47 WC34 W303.18 str.:URA3 MC47 MC49 W303.18 str.:URA3 MC47 MC57 W303.18 str.:URA3 MC48 MC54 W303.18 str.:URA3 MC49 MC57 W303.18 str.:URA3 MC49 MC51 W303.18 str.:URA3 MC49 MC51 W303.18 str.:URA3ADE2.7 clr111. ML1 YDS714 W303.18 str.:URA3ADE2.7 clr111. ML1 MC51 W303.18 URA3ADE2.7 clr111. SUM1.1 ML51 MC53 W303.18 URA3ADE2.7 clr111. SUM1.1 MC51 MC54 W303.18 URA3ADE2.7 clr111. SUM1.1 MC51 MC64 W303.18 URA3ADE2.7 clr111. sUM1.1 MC51 MC64 W303.18 URA3ADE2.7 clr111. str.::::::::::::::::::::::::::::::::::::	MC28	W303-1B <i>sum1::URA3</i>	
$\begin{array}{llllllllllllllllllllllllllllllllllll$	JRY3935	W303-1B <i>hmr::ss</i> Δ <i>I</i>	J. Rine
YLS86 W303-1B hmr2B:20E2 W303-1B hmr2A:20E2 W303-1B hmr2A:20E2 W303-1A siz:2HK3 sum1:URA3 MC49 W303-1B siz:UK3 sum1:URA3 MC49 W303-1B siz:UK3 sum1:URA3 MC57 W303-1B siz:UK3 SUM1-1 W577 W303-1B siz:UK3 SUM1-1 YDS712 W303-1A siz:2HK3 SUM1-1 YDS712 W303-1A siz:2HK3 W303-1B URA3/ADE2-Tel/UL W571 W303-1B URA3/ADE2-Tel/UL W571 W303-1B URA3/ADE2-Tel/UL W571 W303-1A URA3/ADE2-Tel/UL W572 W303-1A URA3/ADE2-Tel/UL W573 W303-1A URA3/ADE2-Tel/UL W574 W303-1B URA3/ADE2-Tel/UL W575 W303-1A URA3-Tel/UL W575 W303-1A URA3-Tel/UL W576 W303-1A URA3-Tel/UL W575 W303-1A URA3-Tel/UL W575 W303-1A URA3-Tel/UL W575 W303-1A URA3-Tel/UL W578 W303-1A SUM1-1 W579 W303-1A SUM1-1	MC31	JRY3935 sum1::URA3	
$\begin{array}{llllllllllllllllllllllllllllllllllll$	YLS586	W303-1B $hmr\Delta B:ADE2$	
YLS404 W303-1B <i>hmr2A::ADC2</i> MC35 YLS404 <i>sun1::UR43</i> MC47 W303-1B <i>siz::HIS3 sun1::UR43</i> MC49 W303-1B <i>siz::HIS3 sun1::UR43</i> MC57 W303-1B <i>siz::HIS3 sun1::UR43</i> MC57 W303-1B <i>siz::HIS3 sun1-1</i> YDS712 W303-1B <i>siz::HIS3 SUN1-1</i> YDS712 W303-1B <i>siz::HIS3 SUN1-1</i> MC51 W303-1B <i>UR43/ADC2:TelVIL</i> MC51 W303-1B <i>UR43/ADC2:TelVIL</i> MC51 W303-1B <i>UR43/ADC2:TelVIL</i> MC51 W303-1B <i>UR43/ADC2:TelVIL</i> MC52 W303-1B <i>UR43/ADC2:TelVIL</i> MC51 W303-1B <i>UR43/ADC2:TelVIL</i> MC561 W303-1B <i>UR43/ADC2:TelVIL</i> MC561 W303-1B <i>UR43/ADC2:TelVIL</i> MC561 W303-1B <i>UR43/ADC2:TelVIL</i> MC561 W303-1B <i>UR43/ADC2:TelVIL</i> MC561 W303-1B <i>UR43/ADC2:TelVIL</i> MC563 W303-1B <i>UR43/ADC2:TelVIL</i> MC564 W303-1B <i>UR43/ADC2:TelVIL</i> MC565 W105-500 M1-1 MC568 W303-1B <i>UR43/TelVIL</i> MC566 W303-1B <i>UR43/TelVIL</i> MC577 W303-1B <i>UR43/TelVIL</i> MC58 W303-1A <i>UR43/TelVIL</i> MC58 W303-1A <i>UR43/TelVIL</i> MC58 W303-1A <i>UR43/TelVIL</i> MC59 W303-1A <i>UR43/TelVIL</i> MC59 W303-1A <i>UR43/TelVIL</i> MC59 W303-1A <i>UR43/TelVIL</i> MC59 W303-1A <i>UR43/TelVIL</i> MC59 W303-1A <i>UR43/TelVIL</i> MC50 W303-1B <i>UR43/TElVI</i> MC50	MC33	YLS586 sum1::URA3	
$\begin{array}{llllllllllllllllllllllllllllllllllll$	YLS404	W303-1B $hmr\Delta A$:: $ADE2$	
MC47 W303-14 <i>siz:HI33 sum1:URA3</i> MC49 W303-18 <i>siz:HI33 sum1:URA3</i> MC54 W303-18 <i>siz:HI33 SUM1-1</i> MC57 W303-18 <i>siz:HI33 SUM1-1</i> YDS712 W303-14 <i>siz:HI33 SUM1-1</i> YDS712 W303-14 <i>siz:HI33 SUM1-1</i> YDS714 W303-18 <i>siz:HI33 SUM1-1</i> MC51 W303-1A <i>URA3/ADE2-TaV11L SUM1-1</i> MC52 W303-1B <i>URA3/ADE2-TaV11L SUM1-1</i> MC53 W303-1A <i>URA3/ADE2-TaV11L SUM1-1</i> MC50 W303-1B <i>URA3/ADE2-TaV11L siz:HI33 SUM1-1</i> MC51 W303-1B <i>URA3/ADE2-TaV11L siz:HI33 SUM1-1</i> MC60 W303-1B <i>URA3/ADE2-TaV11L siz:HI33 SUM1-1</i> MC66 YDV66 W303-1A <i>URA3-TaV11L IntrA:TRP1</i> MC66 YDV66 Sum1:LEU2 YDV67 W303-1A <i>URA3-TaV11L IntrA:TRP1</i> MC88 W303-1A <i>URA3-TaV11L IntrA:TRP1</i> MC88 W303-1A <i>URA3-TaV11L IntrA:TRP1</i> MC88 W303-1A <i>URA3 SUM1-1</i> MC90 W303-1A <i>URA3 SUM1-1</i> MC90 W303-1A <i>URA3 SUM1-1</i> MC90 W303-1A <i>URA3 SUM1-1</i> MC92 W303-1A <i>URA3 SUM1-1</i> MC94 W303-1B <i>BVM3 Fill Pitter Pitt</i>	MC35	YLS404 sum1::URA3	
MC49 W303-18 <i>wiz::HIS3 SUM1-1</i> MC54 W303-1A <i>wiz::HIS3 SUM1-1</i> MC57 W303-18 <i>wiz::HIS3 SUM1-1</i> YDS712 W303-18 <i>wiz::HIS3 SUM1-1</i> YDS714 W303-18 <i>wiz::HIS3 SUM1-1</i> AL275-2a W303-18 <i>wiz::HIS3 SUM1-1</i> MC51 W303-18 <i>Wiz::HIS3 SUM1-1</i> MC52 W303-14 <i>URA3/ADE2-TelVILL SUM1-1</i> MC53 W303-1A <i>URA3/ADE2-TelVILL SUM1-1</i> MC54 W303-1B <i>URA3/ADE2-TelVILL SUM1-1</i> MC60 W303-1B <i>URA3/ADE2-TelVILL SUM1-1</i> MC61 W303-1B <i>URA3/ADE2-TelVILL SUM1-1</i> MC64 YDS631 sum1::LEU2 YDV66 W303-1A <i>URA3/ADE2-TelVILL sum2-HIS3 SUM1-1</i> MC64 YDS631 sum1::LEU2 YDV66 W303-1A <i>URA3/ADE2-TelVILL sum2-HIS3 SUM1-1</i> MC66 YDV65 sum1::LEU2 YDV67 W303-1A <i>URA3 SUM1-1</i> MC68 W303-1A <i>SUM1-1</i> MC78 W303-1A <i>URA3 SUM1-1</i> MC89 W303-1A <i>URA3 SUM1-1</i> MC80 W303-1A <i>URA3 SUM1-1</i> MC79 W303-1B <i>SUM1-1</i> MC71 W303-1B <i>SUM1-1</i> MC72 W303-1B <i>SUM1-1</i>	MC47	W303-1A sir2::HIS3 sum1::URA3	
MCS4 W303-IA <i>siz2:HIS3 SUMI-1</i> MCS7 W303-IB <i>siz2:HIS3 SUMI-1</i> YDS712 W303-IA <i>siz2:HIS3 SUMI-1</i> YDS714 W303-IB <i>UR43/ADE2-TeIVIL</i> MCS1 W303-IB <i>UR43/ADE2-TeIVIL</i> MCS1 W303-IB <i>UR43/ADE2-TeIVIL</i> MCS2 W303-IB <i>UR43/ADE2-TeIVIL</i> MCS2 W303-IB <i>UR43/ADE2-TeIVIL</i> MCS3 W303-IB <i>UR43/ADE2-TeIVIL</i> MCS1 W303-IB <i>UR43/ADE2-TeIVIL</i> MCS6 W303-IB <i>UR43/ADE2-TEIVIL</i> MCS8 W303-IB <i>SUMI-1</i> MCS0 W303-IA <i>UR43/ADE1/L</i> MCS9 W303-IB <i>SUMI-1</i> MCS0 W303-IA <i>UR43/SUMI-1</i> MCS0 W303-IA <i>UR43/SUMI-1</i> MCS0 W303-IA <i>UR43/SUMI-1</i> MCS0 W303-IA <i>UR43/SUMI-1</i> MCS0 W303-IA <i>UR43/SUMI-1</i> MCS9 W303-IB <i>SUMI-1</i> MCS9 W303-IB <i>SUMI-1</i> MCS9 W303-IB <i>SUMI-1</i> MCS9 W303-IB <i>SUMI-1</i> MCS9 W303-IB <i>SUMI-1</i> MCS6 W303-IB <i>SUMI-1</i> MCS9 W303-IB <i>SUMI-1</i> MCS9 W303-IB <i>SUMI-1</i> MCS6 W303-IB <i>SUMI-1</i> MCS6 W303-IB <i>SUMI-1</i> MCS6 M303-IB <i>SUMI-1</i> MCS7 W303-IB <i>SUMI-1</i> MCS7 W303-IB <i>SUMI-1</i> MCS7 M37 B <i>SUMI-1</i> MCS7 M33 B <i>SUMI-1</i> MCS7 M34 B <i>SUMI-1</i> MC57 M34 B <i>SUMI-1</i> MC5	MC49	W303-1B sir2::HIS3 sum1::URA3	
MCS7 W303-1B <i>siz2:HIS3</i> YDS712 W303-1A <i>siz2:HIS3</i> YDS714 W303-1B <i>siz2:HIS3</i> YDS714 W303-1B <i>siz2:HIS3</i> YDS714 W303-1B <i>uRa3AD622-TelVIIL SUM1-1</i> MCS1 W303-1A <i>URA3AD622-TelVIIL SUM1-1</i> MCS2 W303-1B <i>URA3AD622-TelVIIL SUM1-1</i> MCS3 W303-1A <i>URA3AD622-TelVIIL SUM1-1</i> MCS6 W303-1B <i>URA3AD622-TelVIIL siz2:HIS3</i> SUM1-1 MC61 W303-1B <i>URA3AD622-TelVIIL siz2:HIS3</i> SUM1-1 MC64 YDS631 sum1::LEU2 YDV66 W303-1A <i>URA3-TAP2-TelVIIL siz2:HIS3</i> SUM1-1 MC64 YDS631 sum1::LEU2 YDV66 W303-1A <i>URA3-TAPUIL hnrAA</i> ::TRP1 MC66 Y003 TA <i>URA3-TAPUIL hnrAA</i> ::TRP1 MC66 YDV65 suM1-1 MC68 YDV65 suM1-1 MC68 YDW5 sum1::LEU2 MC80 W303 (haploid, MAT allele not determined) <i>siz2:HIS3</i> SUM1-1 <i>URA3-TelVIIL</i> MC88 W303-1B SUM1-1 MC91 W303-1B <i>URA3</i> SUM1-1 MC92 W303-1A <i>URA3</i> SUM1-1 MC94 W303-1B <i>SUM1-1</i> MC95 W303-1B <i>SUM1-1</i> MC95 W303-1B <i>SUM1-1</i> MC94 W303-1B <i>SUM1-1</i> MC95 W303-1B <i>SU</i>	MC54	W303-1A <i>sir2::HIS3 SUM1-1</i>	
YDS712 W303-IA siz-2:HIS3 YDS714 W303-IB iS:2:HIS3 AL275-Za W303-IB UR43/ADE2-TeIVIL MCS1 W303-IA UR43/ADE2-TeIVIL SUMI-1 MCS2 W303-IA UR43/ADE2-TeIVIL SUMI-1 MCS3 W303-IA UR43/ADE2-TeIVIL SUMI-1 MCS1 W303-IA UR43/ADE2-TeIVIL siz::HIS3 SUMI-1 MC60 W303-IB UR43/ADE2-TeIVIL siz::HIS3 SUMI-1 MC61 W303-IB UR43/ADE2-TeIVIL siz::HIS3 SUMI-1 MC64 YDS631 W303-IA UR43-TeIVIL IntrAd::TRP1 MC66 YDV66 sum1::LEU2 YDV67 W303-IA UR43-TeIVIL IntrAd::TRP1 MC88 YD067 um1::LEU2 YDV67 W303-IA SUMI-1 MC88 Y303-IB SUMI-1 MC88 W303-IA SUMI-1 MC89 W303-IA SUMI-1 MC91 W303-IB SUMI-1 MC92 W303-IB SUMI-1 MC93 W303-IB SUMI-1 MC94 W303-IB SUMI-1 MC95 W303-IB SUMI-1 MC96 W303-IB SUMI-1 MC97 W303-IB SUMI-1 MC98 W303-IB SUMI-1 MC99 W303-IB SUMI-1	MC57	W303-1B <i>sir2</i> :: <i>HIS3</i> SUM1-1	
YDS/14 W303-1B $sr12:HIS3$ A. Lustig MCS1 W303-1A $UR43/ADE2$ -TeVIIL $SUM1-1$ A. Lustig MCS1 W303-1A $UR43/ADE2$ -TeVIIL $SUM1-1$ A. Lustig MCS3 W303-1A $UR43/ADE2$ -TeVIIL $SUM1-1$ A. Lustig MCS3 W303-1A $UR43/ADE2$ -TeVIIL $Sum1-1$ A. Lustig MC60 W303-1B $UR43/ADE2$ -TeVIIL $sin2::HIS3 SUM1-1$ YDS63 MC61 W303-1B $UR43/ADE2$ -TeVIIL $sin2::HIS3 SUM1-1$ YDS63 WC64 YDS63 Isum1::LEU2 YDV66 W303-1A $UR43/ADE2$ -TeVIIL $sin2::HIS3 SUM1-1$ YDV66 W303-1A $UR43/ADE2$ -TeVIIL $sin2::HIS3 SUM1-1$ YDV67 W303-1B $UR43/ADE2$ -TeVIIL $sin2::HIS3 SUM1-1$ WC66 YDV67 sum1::LEU2 YDV67 W303-1B $UR43/ADE2$ -TeVIIL $sin2::HIS3 SUM1-1 UR43-TeVIIL MC88 W303-1A SUM1-1 MC88 W303-1A SUM1-1 MC88 W303-1A SUM1-1 MC89 W303-1A SUM1-1 MC90 W303-1B Imr3.SUM1-1 MC91 W303-1B Imr3.SUM1-1 MC92 W303-1B Imr3.SUM1-1 MC93 W303-1B SUM1-1 Imr3.TRP1 isr2::HIS3 Tap1-12::UR43 MC97 MC96 W303-1B SUM1-1 Imr3.TRP1 isr2::HIS3 Tap1-12::UR43 MC98 W303-1B SUM1-1 Imr3.TRP1 isr2::HIS3 Tap1-12:$	YDS712	W303-1A <i>sir2::HIS3</i>	
ALL2/5-2a W303-1B UKA3/ADE2-TeVILL A. Lustig MCS1 W303-1A UKA3/ADE2-TeVILL SUMI-1 MCS3 MCS2 W303-1B UKA3/ADE2-TeVILL SUMI-1 MCS6 MCS0 W303-1B UKA3/ADE2-TeVILL SUMI-1 MCS6 MCS1 W303-1B UKA3/ADE2-TeVILL SUMI-1 MCS6 MCS0 W303-1B UKA3/ADE2-TeVILL size::HIS3 SUMI MCS6 MCS4 YDS631 W303-1B UKA3/ADE2-TeVILL size::HIS3 SUMI-1 MCS6 YDV66 W303-1A UKA3-TeVILL MCS6 YDV66 sum1:::EU2 YDV67 VDV67 W303-1A UKA3-TeVILL hmr $\Delta4$::TRP1 MCS8 W303-1A SUMI-1 MCS8 W303-1A SUMI-1 MCS8 W303-1A SUMI-1 MCS9 W303-1B SUMI-1 MC90 W30-1A SUMI-1 MC91 W303-1B SUMI-1 hmr $\Delta4$::TRP1 isp1-12::URA3 MC92 W303-1B SUMI-1 hmr $\Delta4$::TRP1 isp1-12::URA3 MC96 W303-1B SUMI-1 hmr $\Delta4$::TRP1 isp1-12::URA3 MC98 W303-1B SUMI-1 hmr $\Delta4$::TRP1 isp1-12::URA3 MC99 W303-1B SUMI-1 hmr $\Delta4$::TRP1 isp1-12::URA3 MC90 W30-1B SUMI-1 hmr $\Delta4$::TRP1 isp1-12::URA3 MC100 W30-1B SUMI-1 hmr $\Delta4$::TR	YDS714	W303-1B <i>sir2::H1S3</i>	
$\begin{array}{llllllllllllllllllllllllllllllllllll$	AJL2/5-2a	W303-1B URA3/ADE2-TEIVILL	A. Lustig
MC52 w 303-1B URA3/ADE2-TelVILL SUM1-1 MC53 w 303-1B URA3/ADE2-TelVILL SUM1-1 MC60 W 303-1B URA3/ADE2-TelVILL siz2:HIS3 SUM1 MC61 W 303-1B URA3/ADE2-TelVILL siz2:HIS3 SUM1-1 YD8631 W 303-1B URA3-TelVILL siz2:HIS3 SUM1-1 WC64 YD8661 sum1::LEU2 YDV66 W 303-1A URA3-TelVILL hmr\DA::TRP1 MC66 YDV67 sum1::LEU2 YDV67 W 303 (haploid, MAT allele not determined) siz2::HIS3 SUM1-1 URA3-TelVILL MC80 W 303 (haploid, MAT allele not determined) siz2::HIS3 SUM1-1 URA3-TelVILL MC80 W 303 (haploid, MAT allele not determined) siz2::HIS3 SUM1-1 URA3-TelVILL MC80 W 303-1A URA3 SUM1-1 MC90 W 303-1A URA3 SUM1-1 MC91 W 303-1B URA1: TRP1 rap1-12::URA3 MC93 W 303-1B B UM1-1 hmr\DA::TRP1 rap1-12::URA3 MC94 W 303-1B SUM1-1 hmr\DA::TRP1 rap1-12::URA3 MC95 W 303-1B SUM1-1 hmr\DA::TRP1 rap1-12::URA3 MC96 W 303-1B SUM1-1 hmr\DA::TRP1 rap1-12::URA3 MC97 W 303-1B SUM1-1 hmr\DA::TRP1 rap1-12::URA3 MC98 W 303-1B SUM1-1 hmr\DA::TRP1 rap1-12::URA3 MC100 W 303-1B SUM1-1 hmr\DA::TRP1 rap1-12::URA3 MC101	MC51	W303-1A UKA3/ADE2-TelVIIL SUMI-1	
	MC52	W303-1B URA3/ADE2-IEIVIIL SUMI-1	
	MC53	W303-1A UKA3/ADE2-TelVIIL SUMI-1	
MC01 W305-1B UK13/ADE2-1eV11L MC14 YDS631 W303-1B UK13-TeV11L hmr2A::TRP1 MC64 YDS631 sum1::LEU2 MC66 YDV66 W303-1A UK13-TeV11L MC66 YDV67 W303-1B UK13-TeV11L MC68 YDV67 W303-1B UK13-TeV11L MC80 W303-1A SUM1-1 MC88 W303-1A SUM1-1 MC90 W303-1B UK1-1 MC91 W303-1B UK1-1 MC92 W303-1B UK1-1 MC93 W303-1B UK1-1 MC94 W303-1B UK1-1 MC95 W303-1B UK1-1 MC96 W303-1B SUM1-1 MC97 W303-1B SUM1-1 MC98 W303-1B SUM1-1 MC98 W303-1B SUM1-1 MC98 W303-1B SUM1-1 MC99 W303-1B SUM1-1 MC100 W303-1B SUM1-1 MC100 W303-1B SUM1-1 MC113 MC72	MC60	W303-1B UKA5/ADE2-TelVIIL \$#2::HIS3 SUM1	
1D3011 w30316 Orbor 1012 WG64 YD813 sum1::LEU2 YDV66 W303-1A URA3-TelVIIL hmrA4::TRP1 MC66 YDV65 sum1::LEU2 YDV67 W303-1B URA3-TelVIIL hmrA4::TRP1 MC68 YDV67 sum1::LEU2 MC80 W303 (haploid, MAT allele not determined) sir2::HIS3 SUM1-1 URA3-TelVIIL MC88 W303-1A SUM1-1 MC89 W303-1A SUM1-1 MC90 W303-1A URA3 SUM1-1 MC91 W303-1B B URA3 SUM1-1 MC92 W303-1B HmrA1::TRP1 rap1-12::URA3 MC96 W303-1B B MmrA1::TRP1 rap1-12::URA3 MC96 W303-1B SUM1-1 hmrA4::TRP1 rap1-12::URA3 MC97 W303-1B SUM1-1 hmrA4::TRP1 rap1-12::URA3 MC98 W303-1B SUM1-1 hmrA4::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hmrA4::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 hmrA4::TRP1 rap1-12::URA3 MC103 GA224 SUM1-1 MC104 M4Ta his3 Leu2-3,112 trp1 ura3 MC105 GA224 SUM1-1 MC120 GA224 SUM1-1 MC121 M4Ta his3 Leu2-3,112 ura3-52 MC122 MYC2675 Sum1::URA3 MC133 MC7655 SUM1-1	NDS621	W303-1D UKAS/ADE2-10VIIL \$#2::HIS5 SUM1-1 W303-1D UDA3 TalVIII	
$\begin{array}{llllllllllllllllllllllllllllllllllll$	MC64	VDS-1D UKA5-1EW11L VDS631 sum1-11 EU12	
1D Yoo W3051 KURSTELU2 YDV67 W3051B UR43-TelVIIL hmr $\Delta 4::TRP1$ MC66 YDV67 sum1::LEU2 MC80 W303 (haploid, MAT allele not determined) sir2::HIS3 SUM1-1 UR43-TelVIIL MC80 W303 (haploid, MAT allele not determined) sir2::HIS3 SUM1-1 UR43-TelVIIL MC88 W303-1A SUM1-1 MC89 W303-1B SUM1-1 MC90 W303-1A UR43 SUM1-1 MC91 W303-1B HIS3 SUM1-1 MC92 W303-1A HIS3 SUM1-1 MC93 W303-1B HIS3 SUM1-1 MC94 W303-1B HIS3 SUM1-1 MC95 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ sir2::HIS3 rap1-12::UR43 MC96 W303-1B SUM1-1 hmr $\Delta A::TRP1$ rap1-12::UR43 MC97 W303-1B SUM1-1 hmr $\Delta A::TRP1$ rap1-12::UR43 MC98 W303-1B SUM1-1 hmr $\Delta A::TRP1$ rap1-12::UR43 MC100 W303-1B SUM1-1 hmr $\Delta A::TRP1$ rap1-12::UR43 MC100 W303-1B SUM1-1 hmr $\Delta A::TRP1$ rap1-12::UR43 MC1010 GA224 SUM1-1 MC102 GA224 SUM1-1 MC103 GA224 SUM1-1 MC104 W305-1B SUM1-1 MC105 GA224 SUM1-1 MC106 GA324 SUM1-1 MC107 </td <td>VDV66</td> <td>W303 1 A UR A3 TolVIII hmrA 4. TPP1</td> <td></td>	VDV66	W303 1 A UR A3 TolVIII hmrA 4. TPP1	
Intervention Intervention WDV67 W303-1B URA3-TelVIIL hmr $\Delta A::TRP1$ MC68 YDV67 sum1::LEU2 MC80 W303 (haploid, MAT allele not determined) sir2::HIS3 SUM1-1 URA3-TelVIIL MC88 W303-1A SUM1-1 MC90 W303-1A SUM1-1 MC91 W303-1B URA3 SUM1-1 MC92 W303-1B HIS3 SUM1-1 MC93 W303-1B HIS3 SUM1-1 MC94 W303-1B HIS3 SUM1-1 MC95 W303-1B HIS3 SUM1-1 MC96 W303-1B HIS3 SUM1-1 MC97 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ rap1-12::URA3 MC96 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ sir2::HIS3 rap1-12::URA3 MC97 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ sir2::HIS3 MC98 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ rap1-12::URA3 MC99 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ rap1-12::URA3 MC100 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ rap1-12::URA3 MC100 W303-1B SUM1-1 hmr $\Delta 4::TRP1$ rap1-12::URA3 MC100 W303-1B SUM1-1 MC100 W303-1B SUM1-1 MC120 GA224 sum1::URA3 MC121 M4Ta his3 MC925 Sum1::URA3 K. Carlson MC122 <	MC66	VDV6 sum1:1 FU2	
NO66 YDV67 sum::LEU2 MC80 W303 (haploid, MAT allele not determined) sir2::HIS3 SUMI-1 URA3-TelVIIL MC80 W303-1A SUMI-1 MC89 W303-1A SUMI-1 MC90 W303-1B URA3 SUMI-1 MC91 W303-1B URA3 SUMI-1 MC92 W303-1B URA3 SUMI-1 MC93 W303-1B HIS3 SUMI-1 MC94 W303-1B URA3 SUMI-1 MC95 W303-1B HIS3 SUMI-1 MC96 W303-1B SUMI-1 MC97 W303-1B SUMI-1 hmrA4::TRP1 sir2::HIS3 rap1-12::URA3 MC96 W303-1B SUMI-1 hmrA4::TRP1 sir2::HIS3 MC97 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC98 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC99 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC100 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC100 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC1010 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC102 GA224 SUMI-1 MC102 GA224 SUMI-1 MC102 GA224 SUMI-1 MC120 GA224 SUMI-1 MC121 MY2675 SUMI-1 MC122 MY2675 SUMI-1	YDV67	W303-1B $IIRA3$ -TelVIII. hmr Λ 4TRP1	
Initial constraint of the start of the	MC68	YDV67 sum1: 1 F1/2	
MC88 W303-1A SUMI-1 MC89 W303-1B SUMI-1 MC90 W303-1B SUMI-1 MC91 W303-1B UR43 SUMI-1 MC92 W303-1A HIS3 SUMI-1 MC93 W303-1B UR43 SUMI-1 MC94 W303-1B UR43 SUMI-1 MC95 W303-1B HIS3 SUMI-1 MC96 W303-1B SUMI-1 hmrA4::TRP1 sir2::HIS3 rap1-12::URA3 MC96 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC97 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC98 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC99 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::URA3 MC100 GA224 SUMI-1 MC100 GA224 SUMI-1 MC100 GA224 SUMI-1 MC120 GA224 SUMI-1 MC120 GA224 SUMI-1 MC131 MCY2675 Sum1:URA3 MC122 MYC2675 Sum1:URA3 YDS31 MATa his1 K. Nasmyth	MC80	W303 (haploid, MAT allele not determined) sir2::HIS3 SUM1-1 URA3-TelVIIL	
MC89 W303-1B SUMI-1 MC90 W303-1A UR43 SUMI-1 MC91 W303-1B UR43 SUMI-1 MC92 W303-1A HIS3 SUMI-1 MC93 W303-1B HIS3 SUMI-1 MC94 W303-1B HIS3 SUMI-1 MC95 W303-1B binrtA::TRP1 rap1-12::UR43 MC96 W303-1B SUMI-1 hmrA4::TRP1 sir2::HIS3 rap1-12::UR43 MC97 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::UR43 MC98 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::UR43 MC99 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::UR43 MC99 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::UR43 MC99 W303-1B SUMI-1 hmrA4::TRP1 rap1-12::UR43 MC100 GA224 SUMI-1 MC120 GA224 SUMI-1 MC121 GA224 SUMI-1 MC122 MYC2675 Sum1::UR43 MC131 MC1265 SUM1-1 MC132 M4Ta his1 K< Nasmyth	MC88	W303-1A SUM1-1	
MC90 W303-1A URA3 SUM1-1 MC91 W303-1B URA3 SUM1-1 MC92 W303-1A HIS3 SUM1-1 MC93 W303-1B HIS5 SUM1-1 YLS506 W303-1B hmrΔ4::TRP1 rap1-12::URA3 MC96 W303-1B SUM1-1 hmrΔ4::TRP1 sir2::HIS3 rap1-12::URA3 MC97 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC98 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 mm24::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 mm24::TRP1 rap1-12::URA3 MC1010 GA224 sum1: MC4::TRP1 rap1-12::URA3 MC1010 GA224 sum1::URA3 S. Gasser MC120 GA224 sum1::URA3 K. Carlson MC121 MYC2675 SUM1-1 K. Nasmyth MC122 MYC2675 sum1::URA3 K. Nasmyth	MC89	W303-1B SUM1-1	
MC91 W303-1B UR43 SUM1-1 MC92 W303-1A HIS3 SUM1-1 MC93 W303-1B HIS3 SUM1-1 MC96 W303-1B hmr ΔA ::TRP1 rap1-12::UR43 MC96 W303-1B SUM1-1 hmr ΔA ::TRP1 sir2::HIS3 rap1-12::UR43 MC97 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::UR43 MC98 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::UR43 MC99 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::UR43 MC99 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::UR43 MC100 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::UR43 MC120 GA224 SUM1-1 MC1210 GA224 sum1::UR43 M. Carlson MC122 MY2675 sum1::UR43 M. Carlson MC122 MY2675 sum1::URA3 K. Nasmyth MC123 MATa his1 K. Nasmyth MC124 W303-1B Sir3:HIS4 K. Nasmyth MC130 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig MC131 W303-1A SUM1-	MC90	W303-1A URA3 SUM1-1	
MC92 W303-1A HIS3 SUM1-1 MC93 W303-1B HIS3 SUM1-1 MC96 W303-1B bill hint2A::TRP1 rap1-12::URA3 MC96 W303-1B SUM1-1 hint2A::TRP1 sip2::HIS3 rap1-12::URA3 MC97 W303-1B SUM1-1 hint2A::TRP1 sip2::HIS3 MC98 W303-1B SUM1-1 hint2A::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hint2A::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hint2A::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 MC120 GA224 SUM1-1 MC120 GA224 sum1::URA3 MC121 MCY2675 SuM1-1 MC122 MYC2675 sum1::URA3 YDS31 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC130 W303-1B sir3::HIS4 K. Nasmyth MC131 W303-1A SUM1-1 rap1-17: ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-12::HIS3 hint2A::TRP1 URA3/ADE2-TelVIIL A. Lustig </td <td>MC91</td> <td>W303-1B URA3 SUM1-1</td> <td></td>	MC91	W303-1B URA3 SUM1-1	
MC93 W303-1B HIS3 SUM1-1 YLS506 W303-1B hmrΔ4::TRP1 rap1-12::URA3 MC96 W303-1B SUM1-1 hmrΔ4::TRP1 sir2::HIS3 rap1-12::URA3 MC97 W303-1B SUM1-1 hmrΔ4::TRP1 sir2::HIS3 MC98 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 GA224 MATα can1-100 hom3 his3 leu2-3,112 trp1 ura3 S. Gasser MC105 GA224 sUM1-1 MCY2675 SUM1-1 MC120 GA224 sUM1-1 MCY2675 SUM1-1 MC121 MY2675 SUM1-1 MC Carlson MC113 MCY2675 sum1::URA3 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC125 W303-1B sir3::HIS4 K. Nasmyth MC130 W303-1B suf1-1 rap1-17::HIS3 hmrΔ4::TRP1 URA3/ADE2-TelVIIL A. Lustig MC131 W303-1B SUM1-1 rap1-17::HIS3 hmrΔ4::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL	MC92	W303-1A HIS3 SUM1-1	
YLS506 W303-1B $hmr\Delta A::TRP1 rap1-12::URA3$ MC96 W303-1B $SUM1-1 hmr\Delta A::TRP1 sir2::HIS3 rap1-12::URA3$ MC97 W303-1B $SUM1-1 hmr\Delta A::TRP1 sir2::HIS3$ MC98 W303-1B $SUM1-1 hmr\Delta A::TRP1 rap1-12::URA3$ MC99 W303-1B $SUM1-1 hmr\Delta A::TRP1 rap1-12::URA3$ MC100 GA224 $SUM1-1$ MC120 GA224 $SUM1-1$ MCY2675 MATa his3 $\Delta 200 leu2-3,112 ura3-52$ M. Carlson MC113 MCY2675 $SuM1-1$ K. Nasmyth MC122 MYC2675 $sum1::URA3$ K. Nasmyth MC124 W303-1B $sir4:HIS4$ K. Nasmyth MC125 W303-1B $sir4:HIS4$ K. Nasmyth MC126 W303-1A $sum1::LEU2 rap1-17 ADE2-TelVIIL$ A. Lustig MC131 W303-1A $SUM1-1 rap1-12::HIS3 hmr\DeltaA::TRP1 URA3/ADE2-TelVIIL$ A. Lustig MC132 W303-1A $SUM1-1 rap1-17::HIS3 hmr\DeltaA::TRP1 URA3/ADE2-TelVIIL A. Lustig $	MC93	W303-1B HIS3 SUM1-1	
MC96 W303-1B SUM1-1 hmrΔ4::TRP1 sir2::HIS3 rap1-12::URA3 MC97 W303-1B SUM1-1 hmrΔ4::TRP1 sir2::HIS3 MC98 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 GA224 MATα can1-100 hom3 his3 leu2-3,112 trp1 ura3 S. Gasser MC105 GA224 SUM1-1 MC120 GA224 sum1::URA3 MCY2675 MATa his3 Δ200 leu2-3,112 ura3-52 M. Carlson MC113 MCY2675 SUM1-1 M. Carlson MC122 MYC2675 Sum1::URA3 K. Nasmyth YDS31 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC130 W303-1B sum1::LEU2 rap1-17 ADE2-TelVIIL A. Lustig MC131 W303-1A sum1::LEU2 rap1-17 va3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-12::HIS3 hmrΔ4::TRP1 urA3/ADE2-TelVIIL A. Lustig	YLS506	W303-1B $hmr\Delta 4$::TRP1 rap1-12::URA3	
MC97 W303-1B SUM1-1 hmr ΔA ::TRP1 sir2::HIS3 MC98 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 hmr ΔA ::TRP1 rap1-12::URA3 GA224 MATa can1-100 hom3 his3 leu2-3,112 trp1 ura3 S. Gasser MC105 GA224 SUM1-1 Mc120 GA224 sum1::URA3 M. Carlson MCY2675 MATa his3 $\Delta 200$ leu2-3,112 ura3-52 M. Carlson M. Carlson MC113 MCY2675 SUM1-1 K. Nasmyth MC124 W303-1B sir1:URA3 YDS31 MATa his1 K. Nasmyth K. Nasmyth MC125 W303-1B sir1:HIS4 K. Nasmyth MC130 W303-1B sir3::HIS3 Mc124 W303-1B sir3::HIS4 MC131 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-72::HIS3 hmr $\Delta 4$::TRP1 URA3/ADE2-TelVIIL A. Lustig	MC96	W303-1B SUM1-1 hmrΔA::TRP1 sir2::HIS3 rap1-12::UR43	
MC98 W303-1B SUM1-1 hmr $\Delta4$::TRP1 rap1-12::URA3 MC99 W303-1B SUM1-1 hmr $\Delta4$::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 hmr $\Delta4$::TRP1 rap1-12::URA3 GA224 MAT α can1-100 hom3 his3 leu2-3,112 trp1 ura3 S. Gasser MC105 GA224 SUM1-1 S. Gasser MC105 GA224 sum1::URA3 S. Gasser MC102 GA224 sum1::URA3 MC carlson MC120 GA224 sum1::URA3 M. Carlson MC113 MCY2675 SUM1-1 M. Carlson MC122 MYC2675 sum1::URA3 K. Nasmyth MC123 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC130 W303-1B sir3::HIS4 A. Lustig MC131 W303-1B SUM1-1 rap1-12::HIS3 hmr $\Delta4$::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ura3/ADE2-TelVIIL A. Lustig	MC97	W303-1B SUM1-1 hmrΔA::TRP1 sir2::HIS3	
MC99 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 MC100 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 GA224 MATα can1-100 hom3 his3 leu2-3,112 trp1 ura3 S. Gasser MC105 GA224 SUM1-1 S. Gasser MC100 GA224 sum1::URA3 S. Gasser MC100 GA224 sum1::URA3 MC Carlson MC120 GA224 sum1::URA3 M. Carlson MC113 MCY2675 SUM1-1 M. Carlson MC122 MYC2675 sum1::URA3 K. Nasmyth MC123 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC130 W303-1B sir4::HIS3 Marta his1 K. Nasmyth MC131 W303-1B sir4::HIS3 hmrΔ4::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17: MDE2-TelVIIL A. Lustig MC133 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	MC98	W303-1B SUM1-1 hmrΔA::TRP1 rap1-12::URA3	
MC100 W303-1B SUM1-1 hmrΔ4::TRP1 rap1-12::URA3 GA224 MATα can1-100 hom3 his3 leu2-3,112 trp1 ura3 S. Gasser MC105 GA224 SUM1-1 S. Gasser MC100 GA224 sum1::URA3 MC MC120 GA224 sum1::URA3 MC MC121 MATa his3 $\Delta 200$ leu2-3,112 ura3-52 M. Carlson MC113 MCY2675 SUM1-1 M. Carlson MC122 MYC2675 sum1::URA3 K. Nasmyth MC123 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC125 W303-1B sir4::HIS3 Mc130 W303-1B sir4::HIS3 MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17: ADE2-TelVIIL A. Lustig	MC99	W303-1B SUM1-1 hmrΔA::TRP1 rap1-12::URA3	
GA224 MATα can1-100 hom3 his3 leu2-3,112 trp1 ura3 S. Gasser MC105 GA224 SUM1-1 MC120 GA224 sum1::URA3 MC120 GA224 sum1::URA3 MC120 GA224 sum1::URA3 MC Yaf55 MATa his3 Δ200 leu2-3,112 ura3-52 M. Carlson MC113 MCY2675 SUM1-1 MCY2675 sum1::URA3 K. Nasmyth MC122 MYC2675 sum1::URA3 K. Nasmyth YDS31 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC125 W303-1B sir4::HIS3 Mc124 W303-1B sir4::HIS3 MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL A. Lustig MC131 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL Mc133 M23-1A SUM1-1 rap1-17 ADE2-TelVIIL	MC100	W303-1B <i>SUM1-1 hmr</i> ∆ <i>A</i> :: <i>TRP1 rap1-12</i> :: <i>URA3</i>	
$\begin{array}{llllllllllllllllllllllllllllllllllll$	GA224	MAT α can1-100 hom3 his3 leu2-3,112 trp1 ura3	S. Gasser
MC120 GA224 sum1::URA3 MCY2675 MATa his3 $\Delta 200$ leu2-3,112 ura3-52 M. Carlson MC113 MCY2675 SUM1-1 MCY2675 sum1::URA3 M. Carlson MC122 MYC2675 sum1::URA3 K. Nasmyth MC123 MATa his1 K. Nasmyth YDS31 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC125 W303-1B sir4::HIS3 K. Nasmyth MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL ALustig MC131 W303-1B SUM1-1 rap1-12::HIS3 hmr ΔA ::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	MC105	GA224 SUM1-1	
MCY2675 MATa his3 $\Delta 200 \ leu2-3,112 \ ura3-52$ M. Carlson MC113 MCY2675 SUM1-1 MC2675 Sum1::URA3 MC122 MYC2675 sum1::URA3 K. Nasmyth YDS31 MATa his1 K. Nasmyth YDS32 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC125 W303-1B sir4::HIS3 MC130 MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL A. Lustig MC131 W303-1B SUM1-1 rap1-17::HIS3 hmr ΔA ::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	MC120	GA224 sum1::URA3	
MC113 MCY2675 SUM1-1 MC122 MYC2675 sum1::URA3 YDS31 MATa his1 K. Nasmyth YDS32 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC135 W303-1B sir4::HIS3 K. Nasmyth MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL ALL MC131 W303-1B SUM1-1 rap1-12::HIS3 hmr∆4::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig MC133 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	MCY2675	MATa his $3 \Delta 200 \ leu 2-3,112 \ ura 3-52$	M. Carlson
MC122 MYC2675 sum1::URA3 YDS31 MATa his1 K. Nasmyth YDS32 MATa his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC125 W303-1B sir4::HIS3 K. Nasmyth MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL AIL440-1c MC131 W303-1A SUM1-1 rap1-12::HIS3 hmr ΔA ::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	MC113	MCY2675 SUM1-1	
Y DS 31 $MATa hisl$ K. NasmythYDS 32 $MATca hisl$ K. NasmythMC124W303-1B sir3::HIS4K. NasmythMC125W303-1B sir4::HIS3MC130MC130W303-1A sum1::LEU2 rap1-17 ADE2-TelVIILMC131W303-1B SUM1-1 rap1-12::HIS3 hmr ΔA ::TRP1 URA3/ADE2-TelVIILAJL440-1cW303-1A SUM1-1 rap1-17 ura3/ADE2-TelVIILMC132W303-1A SUM1-1 rap1-17 ADE2-TelVIILMC133W303-1A SUM1-1 rap1-17 ADE2-TelVIIL	MC122	MYC2675 sum1::URA3	** ** *
YDS32 MA1 α his1 K. Nasmyth MC124 W303-1B sir3::HIS4 K. Nasmyth MC125 W303-1B sir4::HIS3 K. Nasmyth MC130 W303-1B sir4::HIS3 hmr Δ 4::TRP1 URA3/ADE2-TelVIIL K. Nasmyth MC131 W303-1B SUM1-1 rap1-12::HIS3 hmr Δ 4::TRP1 URA3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig MC133 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	YDS31	MATa hist	K. Nasmyth
MC124 W303-1B str3::H154 MC125 W303-1B sir4::H153 MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL MC131 W303-1B SUM1-1 rap1-12::H153 hmr ΔA ::TRP1 URA3/ADE2-TelVIIL AJL440-1c W303-1A SUM1-1 rap1-17 ura3/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	YDS32	$MAI \alpha hisl$	K. Nasmyth
MC125 W303-1B sur4::HIS3 MC130 W303-1A sum1::LEU2 rap1-17 ADE2-TelVIIL MC131 W303-1B SUM1-1 rap1-12::HIS3 hmrΔA::TRP1 URA3/ADE2-TelVIIL AJL440-1c W303-1A HIS3 rap1-17 ura3/ADE2-TelVIIL MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL MC133 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL	MCI24	W303-1B str3::HIS4	
MC 150 W 50/5-1A sum1::LE/2 rap1-1/ ADE2-telVIIL MC131 W 30/3-1B SUM1-1 rap1-12::HIS3 hmr\[A::TRP1 URA3/ADE2-TelVIIL AJL440-1c W 30/3-1A HIS3 rap1-17 ura3/ADE2-TelVIIL MC132 W 30/3-1A SUM1-1 rap1-17 ADE2-TelVIIL MC133 W 30/3-1A SUM1-1 rap1-17 ADE2-TelVIIL	MC125	W 505-1B SIT4::H155	
MC151 W 305-1B SUM1-1 rap1-12::H1S5 hmr∆A::IRP1 UKA3/ADE2-TelVIIL AJL440-1c W 303-1A HIS3 rap1-17 ura3/ADE2-TelVIIL A. Lustig MC132 W 303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig WC133 W 303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	MC130	W 505-1A sum1::LEU2 rap1-1/ ADE2-TelVIIL	
AJL440-1C W305-1A H155 rap1-1/ Wa31/ADE2-TelVIIL A. Lustig MC132 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig MC133 W303-1A SUM1-1 rap1-17 ADE2-TelVIIL A. Lustig	MC131	W $305-1B$ SUM1-1 rap1-12::HIS3 hmr ΔA ::TRP1 URA3/ADE2-TelVIIL W $202.1A$ HIS2 reg 1.17 rm $2/ADE2$ TelVIIL	A T
INIC152 W 505-1A 50/M1-1 rap1-1/ ADE2-1eW IIL MC133 W303-1A \$1/M1-1 rap1-17 ADE2-Tel/VIII humA A::TPD1	AJL440-10 MC122	W 505-1A HISS rap1-1/ Uras/ADE2-1eWIIL W202 1A SUM1 1 rap1 17 ADE2 T-IV/IIL	A. Lustig
	MC132	W 505-1A SUM1-1 RUP1-1/ ADE2-TERVILL W202 1A SUM1 1 rap1 17 ADE2 TalVIII IwawA A. TDD1	

Indirect immunofluorescence of yeast spheroplasts. Indirect immunofluorescence was performed as described previously (39), with the following modifications. Cells with either the HA-tagged or wild-type *SUM1* gene on a 2μ m plasmid (DM383 or DM651) were grown overnight to high density in selective medium.

The culture were first fixed for 20 min under growth conditions by adding 1/4 culture volume of 0.5 M KP_i (pH 6.5) and 18.5% formaldehyde. The fixed cells were converted to spheroplasts by treatment with Zymolyase (0.125 mg of Zymolyase per ml in wash solution) for 2.5 to 4 h at 30°C. The final pellet was



FIG. 1. (A) Restriction map of the SUM1 locus and the eight subclones used to localize the SUM1-1 allele. (B) Nucleotide sequence of the SUM1 gene. The predicted amino acid sequence of Sum1p is shown in the one-letter code below the nucleotide sequence.

washed with and resuspended in 2 volumes of NS⁺ azide (NS [20 mM Tris-HCl {pH 7.6}, 0.25 M sucrose, 1 mM EDTA, 1 mM MgCl₂, 0.1 mM CaCl₂, 0.1 mM ZnCl₂], 1 mM phenylmethylsulfonyl fluoride, 7.2 mM β -mercaptoethanol, 0.02% sodium azide). The slides were incubated in mouse monoclonal anti-HA antibody 12CA5 (6.2 mg/ml; Berkeley Antibody Company, Berkeley, Calif.) diluted 1:500 in PBT for 2 h at room temperature. The slides were washed 10 times with PBT and then incubated with rhodamine-conjugated goat anti-mouse immuno-globulin G (Boehringer Mannheim) according to the manufacturer's instructions. Slides were viewed with a 100×, 1.32-numerical aperture oil immersion objective on a Leitz Dialux fluorescence microscope (Leitz, Rockleigh, N.J.) equipped with a 3.1× projection lens (Diagnostics Instruments, Sterling Heights, Mich.) and a Star-1 cooled charge-coupled device camera (Photometrics, Tucson, Ariz.). Images were processed by the NIH Image program (version 1.55) on a Macintosh Quadra 800 (Apple Computer Inc., Cupertino, Calif.).

Nucleotide sequence accession number. The nucleotide sequence reported in this paper is entered in GenBank with accession number U34832.

RESULTS

Cloning of the *SUM1-1* **allele.** *SUM1-1* behaves as a dominant mutation in the strain background described in a previous report (37). We decided to use this property as a basis for cloning the *SUM1-1* gene, reasoning that the introduction of this mutant allele into a *MAT* α *SUM1 sir2* strain would restore mating by suppressing the *sir2* mutation. To obtain the DNA encoding *SUM1-1*, we constructed recombinant plasmid libraries that contained genomic DNA from *SUM1-1* mutant strains. Genomic libraries, with inserts ranging from 10 to 16 kbp, were made by using DNA from yeast strains JRY2465 and JRY2466 in the centromeric vector YCp50 (see Materials and Methods). Four libraries were generated, each containing more than 20,000 independent recombinants.

Recombinant plasmid DNA prepared from each library was used to transform yeast strain JRY2515 ($MAT\alpha$ SUM1 sir2:: HIS3 ura3 leu2) to uracil prototrophy. Transformants were then mated with strain W1089-1A (MATa SUM1 SIR2 ura3 LEU2) by replica plating directly from transformation plates onto YEPD plates containing a lawn of the tester strain. Mating-competent transformants, which potentially contained a plasmid with the *SUM1-1* allele, were identified by making a second replica from the mating plate onto plates lacking both uracil and leucine. Approximately 25,000 transformants were screened for each library, and 12 that acquired the ability to mate with the tester strain were identified. Plasmids were rescued from these cells and used to retransform strain JRY2515, in order to retest their ability to confer mating competence. Two independent clones tested positive in this rescreening and were found to contain overlapping inserts. Analysis of subclones from a region in common between these two clones revealed that a 4.9-kb *Avr*II fragment could confer mating competence in the *sir2* strain JRY2515 (Fig. 1A). Northern (RNA) blot analysis had previously shown that this fragment encodes an RNA of approximately 3.7 kb (19, 27).

To determine whether the cloned DNA is derived from the SUM1 locus, we performed a linkage analysis. A PvuII-SpeI fragment of the cloned DNA was inserted into pRS405, a LEU2-containing integrating plasmid, and integrated into the chromosome of a sir2 SUM1 strain (JRY2515) by cleavage within the insert at a SmaI site (see Materials and Methods for further details). Integration of the cloned DNA at its homologous chromosomal locus was confirmed by Southern blotting analysis (data not shown). The resulting strain, MC25, still mated very poorly, indicating that the PvuII-SpeI fragment was not sufficient to confer the SUM1-1 phenotype. (This result was expected since the fragment contains only the C-terminal part of the ORF contained on the minimal active fragment.) MC25 was made mating proficient by introduction of a SIR2-containing plasmid and was then crossed to strain JRY3138 (MATa sir2::HIS3 SUM1-1 leu2). After loss of the SIR2 plasmid, the resulting diploid was induced to undergo meiosis. The haploid segregants from this cross should all contain the sir2::HIS3 mutation. Because SUM1-1 suppresses mating defects in $MAT\alpha$ sir2 strains better than in MATa sir2 strains (37), the segregants

-120	TAA	NAAT:	TAG	TCGC?	TAGT	GGTA	TAT	tg a g.	ATCA	AACG	****	GŤŤŤ	CATA	CATA	ATTA	ACAA	AATT	CGTT	TGTT	GCGG	GATG	ICTG	AGAA	CACC	ACAG	cccc	гтст	GATA	ACAT	AACCA	-1
1	ATG	AAC	AGA	GAC	TTC	CGT	CTT	GGA	CCC	AAA	GAT	GAC	GTA	GAT	ACG	TTA	CGA	CTT	ACC	AGT	GCT	CAA	AAT	CAA	GCT	AAT	AGT	TTG	AGG	AAG	90
1	M	N	R	D	F	R	L	G	P	K	D	D	V	D	T	L	R	L	T	S	A	Q	N	Q	A	N	S	L	R	K	30
91	CTA	GAT	ACA	GAT	GCG	AAC	GCT	ала	GCT	TTG	CCA	тсс	ATC	ACA	GAC	ATA	CCA	GTC	AGT	GAT	GAT	тст	GAT	ATA	AAA	CGA	CAA	GTT	GGC	TCT	180
31	L	D	T	D	A	N	A	K	A	L	P	s	I	T	D	I	P	V	S	D	D	s	D	I	K	R	Q	V	G	S	60
181	GGT	TTT	GGT	TCA	AAT	CCG	CTT	CAT	ATA	ала	GAT	TCC	GAG	GCC	TŤŤ	CCA	CAT	TCA	TCT	ATT	GAA	GCC	CTA	ала	GAG	GGT	ATG	GAC	AAA	GTC	270
61	G	F	G	S	N	P	L	H	I	К	D	S	E	A	F	P	H	S	S	I	E	A	L	К	E	G	M	D	K	V	90
271	ACA	ала	CAA	TGC	AAC	GAT	11G	AAG	ACG	GCT	ŤŤG	CTT	tcc	AAA	GAC	ACT	TCC	стс	ACT	GAT	тст	GTG	CAG	GAT	TTG	TTT	AAC	TCA	TTA	ааа	360
91	T	K	Q	C	N	D	L	K	T	A	L	L	s	K	D	T	S	L	T	D	s	V	Q	D	L	F	N	S	L	К	120
361	G77	TTA	TCA	сас	AAC	с аа	TCA	GŤŤ	TTG	GAA	AAT	AAG	CTG	GAT	GAT	GTC	ATG	AAG	AAT	CAA	GTA	AAT	ACT	GAC	ата	ttg	GTC	AAT	AAT	ŤŤG	450
121	V	L	S	Н	N	Q	S	V	L	E	N	K	L	D	D	V	M	K	N	Q	V	N	T	D	І	L	V	N	N	L	150
451	AAT	GAA	CGA	TTG	AAC	ала	TTA	TCG	ACA	ATG	TTG	с ал	AAT	ACT	ŤCG	AAA	GTA	AAT	CAC	TCT	AAC	CTT	CTC	ATA	GAA	AAT	TCA	тсс	AAC	AAT	540
151	N	B	R	L	N	К	L	S	T	M	L	0	N	T	S	K	V	N	H	S	N	L	L	I	E	N	S	s	N	N	180
541	ACC	AGT	TCA	CAG	сат	AAT	ACC	тст	тст	tct	CGA	AGG	GGA	CCT	GGC	AGG	CCA	AGA	AAA	GAT	GCC	TCG	ACC	tcc	аса	ATG	AAT	AAG	TTA	GTG	630
181	T	S	S	Q	Н	N	T	s	s	S	R	R	G	P	G	R	P	R	K	D	A	S	T	s	Т	M	N	K	L	V	210
631	TCA	AAC	GCA	GCC	TCG	GTC	AAT	CTC	AAA	AGT	GCA	тсс	AAT	CAG	GGT	GCT	сст	TTC	TCA	CCA	GTA	AAT	ATT	ACT	TTA	CCG	ACT	GCC	GTA	GTA	720
211	S	N	A	A	S	V	N	L	K	S	A	s	N	Q	G	A	Р	F	S	P	V	N	I	T	L	P	T	A	V	V	240
721	с лл	ACG	tct	AAA	TCC	ааа	AGG	TAT	TTT	GTG	GAA	CCA	TCA	ACG	AAA	с аа	GAG	TCG	CTT	TTA	TTA	тст	GCC	сст	TCA	TCA	TCA	AGA	GAT	GAT	810
241	0	T	s	K	S	К	R	Y	F	V	E	P	S	T	K	0	E	S	L	L	L	s	A	Р	S	S	S	R	D	D	270
811	GCT	GAT	ATG	tcc	TTG	ACT	тсс	GTA	CCG	CAA	AGA	ACG	AAC	AAT	GAA	AAT	GGT	AAA	GAG	CGA	CCA	тсс	ACT	GCT	AAT	тст	AGC	TCC	ATC	ACA	900
271	A	D	M	s	L	T	s	V	P	Q	R	T	N	N	E	N	G	K	E	R	P	s	T	A	N	s	S	S	I	T	300
901	CCA	ACA	CCT	GTT	ACG	CCG	AAC	AAC	ŤTA	ATT	CAA	ATC	AAA	AGA	AAA	AGA	GGC	AGG	CCC	CCA	AAA	AAG	AGA	ACA	GTG	GAA	ACA	ATG	ATA	TCC	990
301	P	T	P	V	T	P	N	N	L	I	Q	I	K	R	K	R	G	R	P	P	K	K	R	T	V	E	T	M	I	S	330
991	AAT	TCC	ACG	G AT	ACG	ATA	GAT	AAG	TCA	GAT	GCA	tct	AAT	CGG	ATT	AAA	AAC	GAG	ATT	CCA	ATA	AAT	TCC	TTG	CTT	CCA	TCT	tcc	AAA	TTT	1080
331	N	S	T	D	Ť	I	D	K	S	D	A	S	N	R	I	K	N	E	I	P	I	N	S	L	L	P	S	s	K	F	360
1081	с а т	с лл	ATA	CCA	TCA	тст	CCA	TCC	AAT	CCT	GTG	TCA	с аа	CCG	GCT	CCG	GTT	CGG	ACT	TCA	AGG	TCA	GCC	ACA	CAA	GAA	ATA	GAC	ATT	AAA	1170
361	Н	0	I	P	S	s	P	S	N	P	V	S	Q	P	A	P	V	R	T	S	R	S	A	T	Q	E	I	D	I	K	390
1171	AGT	TTA	GAA	CTG	GGC	тст	CTA	ATT	TCA	ACT	AAT	GGT	GAT	CCA	AAC	GCA	GAG	GAT	TCG	AAT	ACT	ACT	GAT	ACT	GTT	CAT	AAT	AAC	GTG	GAA	1260
391	S	L	E	L	G	S	L	I	S	T	N	G	D	P	N	A	E	D	S	N	T	T	D	T	V	H	N	N	V	E	420
1261	GGA	AAG	GTA	AAT	GTT	GAA	GAA	AAT	AAA	ACC	GAG	AAG	GAG	ааа	ATA	ATA	ACC	ATC	AAA	TCA	tcc	AGC	GAA	AAT	AGT	GGT	AAT	AAT	ACG	ACC	1350
421	G	K	V	N	V	E	E	N	K	T	E	K	E	К	I	I	T	I	K	S	s	S	E	N	S	G	N	N	T	T	450
1351	AAT	AAT	AAT	AAT	ACT	GAC	AAC	GTC	ATT	AAA	TTT	TCA	GCT	AAT	TCA	GAT	ATC	AAT	AGT	GAT	ATT	CGC	CGA	TTA	ATG	GTT	AAC	GAT	CAG	TTT	1440
451	N	N	N	N	T	D	N	V	I	K	F	S	A	N	S	D	I	N	S	D	I	R	R	L	M	V	N	D	Q	F	480
1441	TCA	TTA	AGT	TAT	GAC	GCC	AGC	GGT	AAT	ATT	ACG	GTC	AAA	ŤTA	CCA	ccc	GTT	тсс	TCT	CCA	GCA	GCA	GCA	ACA	GC T	GCC	GCT	GCT	GCG	GTT	1530
491	S	L	S	Y	D	A	S	G	N	I	T	V	K	L	P	P	V	s	S	P	A	A	A	T	A	A	A	A		V	510
1531	ACG	TCA	GAG	ATG	AAT	AGA	CAA	CAA	AGA	GAA	TTA	GAT	AAA	AGA	CGT	GAT	TCA	AGG	GAG	AAA	ATG	CTT	GTT	AAT	ATG	AAA	TAT	AAC	GAT	CGC	1620
511	T	S	E	M	N	R	Q	Q	R	E	L	D	K	R	R	D	S	R	E	K	M	L	V	N	M	K	Y	N	D	R	540
1621	GAT	AAA	GCA	AAG	TCA	TTT	ATG	GAG	TCT	AAT	AAG	AAA	CTC	TTA	AAA	GCA	ATG	ала	GAA	GAA	GAG	AGG	AGG	AAA	AGA	ATG	ACT	TCG	ATA	ATA	1710
541	D	K	A	K	S	F	M	E	S	N	K	K	L	L	K	A	M	К	E	E	E	R	R	K	R	M	T	S	I	I	570
1711	CAT	GAT	AAT	сас	TTG	AAC	TTA	AAT	TTG	AAT	GAG	ATC	тсс	ACC	CGT	TCA	AAG	ATA	AAA	AGT	GCA	GAA	AAA	CCA	ACA	ACT	AAA	GGT	TCT	TCA	1800
571	H	D	N	Н	L	N	L	N	L	N	E	I	s	T	R	S	K	I	K	S	A	E	K	P	T	T	K	G	S	S	600
1801	ATG	TCG	CCA	AAA	CCA	AGA	tcg	GCC	TCC	ATC	AGC	GGC	ATT	TCA	GAC	CAC	CAA	CAG	GAA	GGA	TAT	САА	CCA	TTA	GAG	CAA	GAA	ала	CTC	GTC	1890
601	M	S	P	K	P	R	S	A	S	I	S	G	I	S	D	H	Q	Q	E	G	Y	Q	P	L	E	Q	E	К	L	V	630
1891	GAT	TTA	GAC	AAT	GAG	GGC	TCA	AAC	GCA	AAC	AGC	GAC	TCT	стс	AAG	ATG	GGT	CTA	ACC	ATA	TCC	GCT	GCC	GAT	ACG	GTT	сас	AAA	GTT	GGA	1980
631	D	I	D	N	E	G	S	N	A	N	S	D	S	L	K	M	G	L	T	I	S	A	A	D	T	V	н	K	V	G	660
1981	ATA	CAG	tcc	ATG	CTA	AAT	TCT	GGG	GAA	GAG	GCA	ATT	ACC	AAG	GAG	AAT	GCA	GAA	TAT	GAA	AGA	AAG	ACC	CCG	GGA	GAT	GAA	GAA	ACT	ACC	2070
661	I	Q	s	M	L	N	S	G	E	E	A	I	T	K	E	N	A	E	Y	E	R	K	T	P	G	D	E	E	T	T	690
2071	ACG	TTC	GTT	CCA	TTA	GAA	AAC	TCA	САА	CCT	TCG	GAC	ACG	ATA	AGG	AAA	AGA	AC A	GCA	GGC	GAT	GAC	GGT	GCA	TTG	GAT	CAA	ACA	GAG	AAT	2160
691	T	F	V	P	L	E	N	S	Q	P	S	D	T	I	R	K	R	T	A	G	D	D	G	A	L	D	Q	T	E	N	720
2161	ACG	AGC	ATA	TCA	CCA	AAG	AAG	AGA	CGC	ACA	GAA	GAT	CAT	ACA	ала	GGT	GAA	GAA	GAT	GAG	GGA	GAA	AGG	GGT	GTT	GGT	AAT	AGC	GGA	ACG	2250
721	T	S	I	S	P	K	K	R	R	T	E	D	H	T	К	G	E	E	D	E	G	E	R	G	V	G	N	S	G	T	750
2251	CTG	GCC	ACC	GTG	GAG	AAT	GTC	TCA	GGA	GAC	ATC	TCC	GCA	GAT	TTA	TCC	ала	GGT	ACT	TCA	TCT	ATC	CAT	AAC	GAT	ACT	GAG	TCT	GCT	AAC	2340
751	L	A	T	V	E	N	V	S	G	D	I	S	A	D	L	S	К	G	T	S	S	I	H	N	D	T	E	S	A	N	780
2341	GAT	AGT	AGC	AAT	GGC	AAC	GGA	AAC	CTC	GGT	ŤŤG	GGT	ACA	GAG	TCA	CGC	AAT	ACA	TTG	TTA	ACG	GCA	ACT	CCC	ATT	GAA	TTA	ATC	TGC	CGG	2430
781	D	S	S	N	G	N	G	N	L	G	L	G	T	E	S	R	N	T	L	L	T	A	T	P	I	E	L	I	C	R	810
2431	GAA	GCG	TTC	TTC	TAC	CGG	AGA	GAT	ATT	CCG	GAC	GTT	CCC	ATT	ACC	ACA	GGA	GCG	TAC	CTG	GAA	TTT	XXX	TTC	AAG	GCG	AAA	GAA	GAG	GAA	2520
811	E	A	F	F	Y	R	R	D	I	P	D	V	P	I	T	T	G	A	Y	L	E	F	K	F	K	A	K	E	E	E	840
2521	TTG	ATC	AAC	TCT	AGC	ATC	AAT	GAG	GAG	GAT	TAT	GCT	GCT	ала	TCA	AAG	CAT	GAA	AAA	ATG	AAT	GCG	CAT	TTC	TTC	AAG	ССТ	GAT	ATT	CAA	2610
841	L	I	N	S	S	I	N	E	E	D	Y	A	A	K	S	K	H	E	K	M	N	A	H	F	F	K	Р	D	I	Q	870
2611	GAA	GAG	ACG	GAA	CTT	GCC	TTT	GAA	ATT	ŤŤG	AGC	AAG	ACA	ACG	CTG	ACA	GAG	AAA	TAC	GTT	AAC	AGT	TTG	GAG	TAC	TTC	TTG	ATG	GAA	TTC	2700
871	E	E	T	E	L	A	F	E	I	L	S	K	T	T	L	T	E	K	Y	V	N	S	L	E	Y	F	L	M	E	F	900
2701	AGG	TGG	GAG	AAT	AAA	CTA	GTT	GGT	CTA	GGC	ŤТА	AAA	CTT	CGG	GAA	TCC	AAA	AGA	ACC	TGG	CAA	AGG	AGA	AAG	GCG	TTA	TTT	GCC	CTT	TTC	2790
901	R	W	E	N	K	L	V	G	L	G	L	K	L	R	E	S	K	R	T	W	Q	R	R	K	A	L	F	A	L	F	930
2791	GAA	TTT	TGG	AGA	GAC	CAA	TCA	AGA	GAC	AAA	AGA	AGA	TTC	CAC	AAC	TAC	ACC	ATC	CTG	CAT	GCG	GTG	AAA	GAG	ATG	GAA	AAC	TAC	AGA	ATA	2880
931	E	F	W	R	D	Q	S	R	D	K	R	R	F	H	N	Y	T	I	L	H	A	V	K	E	M	E	N	Y	R	I	960
2881	TŤŤ	ATT	AAT	CGA	TCC	GTT	TCA	TGG	TTC	TAC	AAC	CAT	ATT	ACC	CTG	CTA	AAA	ATG	ATC	CTC	TAC	GAC	CTC	TGC	GAC	AAT	стс	ACC	ACT	CAA	2970
961	F	I	N	R	S	V	S	W	F	Y	N	H	I	T	L	L	K	M	I	L	Y	D	L	C	D	N	V	T	T	Q	990
2971	TGG	AGA	G A G	TGG	ATG	TTT	CCC	CAT	AAC	GAA	ACA	CTG	CCG	GCA	ŤŤG	GGT	CAG	GAC	GGC	ATT	AAC	GAA	GAC	AAT	CTG	AAT	GAA	ACC	ATC	GAC	3060
991	W	R	E	W	M	F	P	H	N	E	T	L	P	A	L	G	Q	D	G	I	N	E	D	N	L	N	E	T	I	D	1020
3061	AAC	ATG	TTA .	ATT	TTT	GAC	TTC	CTT	GAT	GAC	GGT	TCA	GAA	AAC	AAC	CAG	GTC	AAA	TAT	TCC	AGA	ATC	ATA	CCG	CCA	GAT	ATC	CGT	TAA	GCŤ	3150
1021	N	M	L	I	F	D	F	L	D	D	G	S	E	N	N	Q	V	K	Y	S	R	I	I	P	P	D	I	R	*		1049
3151	GGTC	CGTA	CGTT	GGGG	CAGT	TTCG	AGAA	TAGA	TAAA	AAAA	TGAA	AAAT	алал	АЛАА	ATA		AACA	CAAA	ATA/	ATGG#	CCGT	таст	ATGC	CCAA	CCAA	GTAA	сстс	CCAA	стта	TŤŤG	3270
3271	271 GAACATTACATATTAGGGCTTAACATACATACATACATATTAGGGCTTAACATTACATTACACCTTATCG 3339																														

-240 TATECTTTTAATACCATTGTGTTTGAAATAAATGGCTAAAAAGAATTAATACACAGGCATATTTTATCAAAAGGGCCAACAAGGGACCAAGGGACTTGTTTAAAGTAGCCAAAGGAACTAAGTATCA -121

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ACTAGTGGCTTTAATTTTTTTTTTTTTTTTTCATC -241

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should display detectable mating in patch mating assays only when they are $MAT\alpha$ SUM1-1. We observed that all of the mating-proficient spore colonies from 24 tetrads dissected were phenotypically α mating and Leu⁻. This result indicates that the cloned DNA is closely linked to the SUM1 locus.

Cloning and DNA sequencing of the wild-type SUM1 gene. Several approaches were taken to clone the wild-type copy of the SUM1 gene. The cloned SUM1-1 allele was used to probe a yeast chromosome blot and nitrocellulose filters containing an ordered set of DNA clones representing most of the yeast genome (obtained from L. Riles and M. Olsen, Washington University, St. Louis, Mo.). λ clone 6898 hybridized to the probe, placing SUM1 on chromosome IV-R between GCN2 and PEP7 (53). To obtain a full-length clone of the SUM1 gene, two yeast genomic libraries in plasmid vectors (kindly provided by M. Rose and M. Carlson) (11, 56) were then screened by colony hybridization (59). The SUM1 sequence (Fig. 1B) is derived from phage λ clone 6898 and genomic clones from each of the two plasmid libraries. The putative SUM1 gene encodes an ORF of 1,048 codons which predicts a polypeptide of approximately 115 kDa. Analysis of the Gen-Bank (release 88.0) and PIR (release 44.0) databases revealed no significant homologies to known protein sequences. Furthermore, no similarities to known sequence motifs were found (GCG sequence analysis software package; Genetics Computer Group, Inc., Madison, Wis.).

SUM1 is essential neither for normal growth nor for transcriptional silencing. To determine whether the product of the SUM1 gene is important for cell growth or for silencing, a disruption of the SUM1 gene was constructed in vitro and integrated into the chromosome by the one-step gene replacement method (58). The sum1::URA3 allele (DM264; see Materials and Methods) was introduced into a homozygous diploid strain (W303), and a Ura⁺ transformant heterozygous for the *sum1* gene disruption was identified by Southern blotting analysis. This SUM1/sum1::URA3 diploid was then sporulated, and the phenotypes of the haploid segregants were analyzed. In all 13 complete tetrads examined, the Ura⁺ phenotype segregated 2:2. There was no obvious growth difference observed between the Ura+ and Ura- spore colonies on rich medium. Because the sum1::URA3 mutation removes all of the predicted SUM1 ORF and is therefore presumably a null allele, these data indicate that SUM1 is not an essential gene, nor is it necessary for normal growth on rich medium. Furthermore, mating defects were not observed in the Ura⁺ segregants. Thus, a *sum1* null mutation does not seem to affect gene repression at the silent mating-type loci in a wild-type SIR background.

To determine whether telomeric repression is affected by the *SUM1* disruption, a *sum1::LEU2* mutation was constructed (see Materials and Methods) and introduced into strains that contained a *URA3* reporter gene immediately adjacent to a telomere created at the *ADH4* locus of chromosome VII-L (21) (strains YDS631, YDV66, and YDV67 [Table 1]). The level of telomeric repression in the *sum1::LEU2* mutant strains (MC64, MC66, and MC68) was monitored by growth on complete medium, complete medium minus uracil, and FOA medium, which kills cells that are expressing the *URA3* gene product (5). In all strains tested, expression of the telomeric *URA3* reporter was unaffected by the *sum1::LEU2* mutation (data not shown).

We also introduced the *sum1::URA3* disruption into strains containing the *sir2::HIS3* mutation, whereby the *SUM1-1* mutant allele is able to restore mating. The resulting strains (MC47 and MC49) were still defective in mating (data not shown), indicating that a *sum1* null mutation cannot suppress

the mating defects of a *sir2* mutation in the W303 background. This result strongly suggests that *SUM1-1* is not a loss-of-function mutation (see below).

sum1 mutation can slightly weaken repression at HMR loci with a mutated silencer. It is possible that the lack of a silencing phenotype for the sum1::URA3 disruption is due to the complexity of the HMR-E silencer element. As stated in the introduction, the three HMR-E silencer regulatory elements (A, E, and B) are partially redundant: any two of these three binding sites are sufficient for silencing, though the Rap1p binding site (E) is required for full repression (7, 30). To test for possible synthetic effects of the sum1::URA3 disruption allele, we began with strains YLS404 and YLS586, which contain the ADE2 reporter gene placed adjacent to an HMR-E silencer with mutation in the A and B silencer elements, respectively ($hmr\Delta A$ or $hmr\Delta B$ silencer) (69). The ADE2 gene provides a colony color marker for the transcriptional state at *HMR*: phenotypically Ade^{2–} cells accumulate a pigment and yield red colonies, whereas $Ade2^+$ cells are white (55). In these strains, the ADE2 reporter gene is normally slightly derepressed and yields primarily red colonies with few white sectors. Introduction of the sum1::URA3 allele into these strains (to produce strains MC33 and MC35) caused an increase in the number of white sectors in the colonies, indicating a further decrease in repression (Fig. 2).

Other tests of sum1 mutations, however, did not reveal effects on the activity of weakened silencers (data not shown). For example, introduction of the sum1::URA3 mutation into a strain containing a synthetic silencer in place of HMR-E and a deletion of the HMR-I silencer (JRY3935 [44]) did not result in a measurable loss of mating efficiency (strain MC31). (The synthetic silencer differs from wild-type HMR-E in that all three silencer regulatory elements are required for complete repression.) In addition, we observed no effect of the sum1:: LEU2 allele on repression of a $hmr\Delta A$::TRP1 reporter gene (comparing strains YDV66 and MC66), as judged by growth in the absence of tryptophan. We conclude from these results that deletion of SUM1 has a subtle effect on the HMR-E silencer that can be seen only when the functional redundancy of the silencer is eliminated and when a sensitive reporter gene (such as ADE2) is present at HMR.

We also considered the possibility that the relatively minor effect of *SUM1* deletion on *HM* silencing is due to functional redundancy of Sum1p itself. To begin to determine whether *SUM1* homologs exist, yeast genomic DNA was cleaved with a variety of restriction enzymes and probed with *SUM1* DNA by hybridization at low stringency on Southern blots. This approach failed to reveal any sequences homologous to *SUM1*.

Characterization of the SUM1-1 mutation and construction of an isogenic set of SUM1-1 strains. We noticed several restriction fragment length polymorphisms between the SUM1-1 mutant and wild-type alleles, as well as multiple point mutations and deletions within the 3' untranslated region. To localize the mutation(s) responsible for conferring the SUM1-1 phenotype (suppression of the mating defect of a $MAT\alpha$ sir2 mutant), a series of restriction fragment exchanges between the SUM1-1 allele and the wild-type SUM1 gene was made (Fig. 3A). The recombined alleles were transformed into strain JRY2515 (MAT α sir2) and tested for the ability to restore mating. We noted that a C-terminal restriction fragment (SphI to BsiWI, nucleotides 2847 to 3155 [Fig. 3A]), which encodes the terminal 100 amino acids of Sum1p and contains only 8 nucleotides beyond the predicted stop codon, conferred the SUM1-1 phenotype in the context of otherwise wild-type sequences (Fig. 3B). This fragment was subjected to DNA sequencing and found to have a single missense mutation at

hmr(\(\(\Delta\)B)::ADE2 SUM1

hmr(\[\]*B*)::*ADE2 sum1*::*URA3*



FIG. 2. A sum1 null mutation weakens repression of hmrΔB::ADE2. Representative colonies of isogenic SUM1 and sum1::URA3 strains are shown.

codon 974 in the predicted *SUM1* ORF that would result in a threonine-to-isoleucine change. This mutation maps to a locally hydrophobic part of the predicted protein and would presumably increase the hydrophobic character of this region. The *SUM1-1* allele that contains the point mutation at the codon 974 in the context of otherwise wild-type sequences (DM449) confers a slightly (approximately 2.5-fold) higher mating efficiency to the *sir2* mutant (JRY2515) than does the original *SUM1-1* allele. Our data indicate, therefore, that this mutation in *SUM1-1* is both necessary and sufficient to restore mating in a *MAT* α *sir2::HIS3* mutant.

We proceeded to introduce the SUM1-1 allele by gene replacement into W303-derived strains in order to study the phenotype caused by the mutation in an isogenic background (see Materials and Methods for details). In these experiments, a 4.9-kb AvrII fragment containing the entire SUM1-1 coding sequence was used for gene transplacement experiments. As shown in Table 2, the SUM1-1 mutation improved mating by a $MAT\alpha$ sir2::HIS3 strain by more than 10,000-fold. In contrast, the mating efficiency of a MATa SUM1-1 sir2::HIS3 strain was improved by 10-fold relative to that of the MATa SUM1 sir2 strain. These data are consistent with results from a previous report (37) which showed that SUM1-1 strongly suppresses the nonmating phenotype of *sir2::HIS3* mutations in $MAT\alpha$ strains but restores only weak mating in MATa strains. We confirmed that the SUM1-1 allele acts to restore transcriptional repression at HMR and HML by measuring the steady-state levels of al and αl mRNAs in these strains by Northern blotting (Fig. 4). Consistent with the large improvement in mating observed, we did not detect al transcripts in the MAT α SUM1-1 sir2:: *HIS3* strain (MC57). We also found that $\alpha 1$ transcripts were fully repressed in the MATa SUM1-1 sir2::HIS3 strain (MC54), despite the fact that SUM1-1 restored only weak mating in this strain. This severe drop in $\alpha 1$ mRNA may be in part due to residual $\mathbf{a}_{1-\alpha}$ repression. Finally, we found that the SUM1-1 plasmid also suppressed mating defects in MATa sir3::HIS3 and MATa sir4::HIS3 strains (MC124 and MC125 [data not shown]), as expected (37).

The *SUM1-1* allele is dominant to the wild-type gene in the W303 strain background, since the *SUM1-1* plasmid could suppress mating defects when transformed to a *MAT* α *SUM1 sir2* strain (YDS714 [data not shown]). However, *SUM1-1* is not a

hypermorph, since *SUM1* on a 2µm plasmid could not rescue mating defects when it was transformed into a *MAT* α *sir2*:: *HIS3* strain (JRY2515 [data not shown]). Furthermore, we found that additional copies of the wild-type *SUM1* gene in a *sir2 SUM1-1* cell (either **a** or α) actually results in a slight (fivefold) decrease in mating efficiency. We also noticed that the *SUM1-1* plasmid suppressed the mating defects of *sir2*::*HIS3* mutation at least 100-fold better in a *sum1*::*URA3* strain than in a *SUM1* strain. These findings argue against the idea that the *SUM1-1* allele is an increased function (hypermorphic) mutation and support the notion that *SUM1-1* is an altered-function mutation.

SUM1-1 restores HM silencing in $rap1^s$ and $rap1^t$ mutant strains. Because SUM1-1 has been shown to suppress the silencing defects of multiple silencer site mutations at HMR-E (37), we were interested in determining whether it could also restore repression in strains with a mutated silencer-binding protein. Alleles of RAP1, called rap1s, which are defective in silencing at $hmr\Delta A$ loci but are apparently completely unaffected in essential RAP1 functions have been identified (68). We therefore asked whether SUM1-1 could suppress a rap1-12 mutation, the most severely defective *rap1s* allele. The *rap1-12* mutation results in complete derepression of $hmr\Delta A$::TRP1. A SUM1-1 strain (MC57) was crossed to a strain containing rap1-12 and the $hmr\Delta A$::TRP1 reporter gene (YLS506). The diploid was sporulated, and haploid segregants of relevant genotypes were tested for growth in the presence and absence of tryptophan. We found that SUM1-1 fully restored repression of $hmr\Delta A$::TRP1 in a rap1-12 strain (Fig. 5). By contrast, SUM1-1 appeared only to slightly restore repression of $hmr\Delta A$::TRP1 in a sir2 mutant.

Mutations that result in truncation of the Rap1p C terminus $(rap1^t)$ have been shown to cause complete derepression of telomeres and a more modest silencing defect at *HML*. We therefore examined the effect of *SUM1-1* on the *HML* locus and on the artificial chromosome VII-L telomere in the strains harboring a severe $rap1^t$ mutation (rap1-17). For this purpose, the strain MC131 (*SUM1-1 rap1-12::HIS3 hmr*\Delta4::*TRP1 URA3-ADE2-TelVII-L*) was mated to the strain MC130 (*sum1::LEU2 rap1-17 ADE2-TelVII-L*). The diploid was sporulated, and tetrads were analyzed. The *MAT***a** rap1-17 mutant (AJL440-1c) has a mating efficiency of 2.8×10^{-2} relative to that of the



FIG. 3. (A) Schematic representation of *SUM1* alleles constructed by exchanging restriction fragments between the *SUM1-1* mutant allele and the wild-type *SUM1* gene. The stippled boxes represent the sequences derived from the *SUM1-1* allele. The plasmid designations and results of patch mating assays (see panel B) are shown to the right. Two independent constructs were tested for each recombinant allele. (B) Patch mating assays for the *sir2* strain JRY2515 transformed with plasmids containing either wild-type *SUM1*, the *SUM1-1* mutant allele (DM255), or the recombinant alleles shown in panel A.

wild-type MATa (W303-1A) cells. SUM1-1 restored wild-type levels of HML silencing in the rap1-17 mutant (MC132 or MC133), as judged by the restoration of efficient mating in MATa segregants. In contrast, SUM1-1 appeared to have little or no effect on the telomere repression defect of rap1-17. Three SUM1-1 rap1-17 URA3-ADE2-TelVII-L segregants were examined, and only one showed a slight improvement in FOA resistance (~10 fold), which has not been examined further (data not shown).

SUM1-1 increases telomeric silencing. As described above, disruption of SUM1 did not appear to have any effect on telomeric repression, nor did SUM1-1 appear to restore telomere position effect in the *rap1-17* mutant. To test whether SUM1-1 can restore telomeric repression in *sir* mutants, a SUM1-1 MAT α sir2::HIS3 strain (MC57) was crossed to a MATa strain that contained the URA3 telomeric reporter (YDV66). In analyzing haploid segregants from this cross, we found that SUM1-1 could not restore telomeric URA3 repression in *sir*2 mutant segregants (e.g., MC80 [data not shown]).

In a separate experiment, a *SUM1-1 MAT***a** *sir2::HIS3* strain (MC54) was crossed to a *MAT* α *URA3-ADE2-TelVII-L* reporter strain (AJL275-2a). Again, *SUM1-1* did not restore repression of either the *ADE2* or *URA3* reporter in *sir2* mutant segregants. Surprisingly, however, we observed that *SUM1-1*

TABLE 2. Mating efficiencies of isogenic SUM1 and SUM1-1 strains

Strain	Relevant genotype	Relative mating efficiency				
W303-1B	MATα SIR2 SUM1	1				
YDS714	W303-1B sir2::HIS3 SUM1	$\leq 3.1 \times 10^{-5}$				
MC57	W303-1B sir2::HIS3 SUM1-1	0.37				
MC89	W303-1B SIR2 SUM1-1	5.3				
W303-1A	MATa SIR2 SUM1	1				
YDS712	W303-1A sir2::HIS3 SUM1	$\leq 2.2 \times 10^{-5}$				
MC54	W303-1A sir2::HIS3 SUM1-1	$1.9 imes 10^{-4}$				
MC88	W303-1A SIR2 SUM1-1	5.4				



FIG. 4. Northern blot analysis of $\mathbf{a}I$, αI , and actin transcripts in isogenic *SUM1* and *SUM1-1* strains. Total RNA was prepared from the indicated strains, size fractionated on a 1% formaldehyde agarose gel, transferred to a Hybond-N filter, and hybridized with either $\mathbf{a}I$ or αI and actin (control) probes (see Materials and Methods). The relevant genotypes of the strains used are indicated above the autoradiograph. – represents *SUM1-1* (in the case of *SUM1*) and *sir2*:: *HIS3* (in the case of *SIR2*).

caused an increase of telomeric repression in *SIR*⁺ segregants. Normally, the telomeric *ADE2* reporter in a W303 (*SUM1*) strain background is only slightly repressed, giving rise to mostly white colonies. However, in the *SUM1-1* segregants from the cross with AJL275-2a, the telomeric *ADE2* gene was further repressed, creating white/red sectored colonies (Fig. 6A). In addition, a *URA3-ADE2-TelVII-L* strain with the *SUM1-1* mutation grew at least 10-fold better on FOA plates than an otherwise isogenic *SUM1* strain, indicating an increase of repression of the telomeric *URA3* reporter in these *SUM1-1* strains (Fig. 6B). The increased telomeric *ADE2* repression could also be observed if *SUM1-1* on a *CEN* plasmid was transformed into the *SUM1 URA3-ADE2-TelVII-L* reporter strain (AJL275-2a), although the effect was weaker in the transformants than in an isogenic *SUM1-1* strain (data not shown).

The SUM1-1 mutation increases the rates of cell death and chromosome loss. A SUM1-1 mutant strain grows more slowly than isogenic wild-type or sum1 mutants. For example, wildtype haploid cells (W303-1A and W303-1B) have a doubling time of ~130 min, whereas isogenic SUM1-1 mutants (MC88 and MC89) have a doubling time of ~ 160 min. This slowgrowth phenotype of SUM1-1 mutants can be rescued when copies of the wild-type SUM1 gene are introduced into these cells. We also noted that although heterozygous SUM1-1/ SUM1 diploids show some increase in telomeric repression (as judged by a telomeric ADE2 reporter), their growth rates appear comparable to those of isogenic wild-type diploids and are clearly much higher than those of SUM1-1 homozygous diploids (data not shown). Hence, the slow-growth phenotype of the SUM1-1 mutation, like the silencing phenotype, appears to be partially dominant in the W303 strain background.

We measured cell viability in cultures of SUM1-1 mutants,

since a decrease in cell viability could result in a slow-growth phenotype. The viability was assayed by colony-forming ability in three pairs of isogenic haploid strains (YDS3 and MC89, MC60 and MC61, and YDS712 and MC57). Cell viability of the mutant strain was reduced to about 80% of that of the wild-type cells (Table 3).

We also noted that homozygous SUM1-1 diploids produce colonies with a much larger size variation than wild-type diploids. Because this phenotype is often associated with mutations that increase chromosome loss, we decided to measure chromosome stability in SUM1-1 homozygous diploids. We monitored the loss rate of chromosomes III and V in the SUM1-1/SUM1-1 diploid (MC105 \times MC113), the sum1/sum1 diploid (MC120 \times 122), and the homozygous wild-type diploid $(GA224 \times MCY2675)$. Mating assays were used to determine the rate of chromosome III loss, since an \mathbf{a}/α nonmating diploid cell will acquire the ability to mate if it loses one (or both) of its two copies of chromosome III. This diploid strain also contains a marked chromosome V, with can1 and hom3 markers on opposite chromosome arms. A cell that loses the wildtype chromosome V will grow on plates containing canavanine but not on complete media lacking threonine (23). As shown in Table 4, the SUM1-1/SUM1-1 diploid loses both chromosome III and chromosome V at a rate (number of events per cell division per generation) approximately fivefold higher than in the isogenic SUM1/SUM1 and sum1/sum1 strains.

Cellular localization of Sum1p. The results presented above suggest that Sum1p mediates interactions that influence chromosome behavior as well as position effects. To gain further insights into SUM1 function, we have localized Sum1p by indirect immunofluorescence (Fig. 7), using a version of the protein containing an influenza virus HA antigen tag at its C terminus (see Materials and Methods for details). The HAtagged SUM1 allele complemented the partial silencing defect of a sum1 null mutation in the strain MC33 (data not shown). Nuclear extracts from the cells harboring the tagged allele were analyzed by Western blotting (immunoblotting) using the mouse monoclonal anti-HA antibody 12CA5 as a probe. A protein band of ~125 kDa, specific to the tagged SUM1 gene, was detected, indicating that intact HA-tagged Sum1p was being made (data not shown). Antibody 12CA5 also recognized a nonspecific band of ~46 kDa in yeast cells. In indirect immunofluorescence of yeast spheroplasts derived from cells lacking the HA tag, we observed weak, slightly punctate staining that appeared to be cytoplasmic, presumably as a result of the 46-kDa cross-reacting protein detected on Western blots. Despite this homogeneous nonspecific staining, we observed an obvious nuclear staining in ~ 25 to 40% of spheroplasts derived from cells which contained the HA-tagged SUM1 gene. We therefore conclude that Sum1p is localized to the nucleus, consistent with the effect of SUM1-1 on both transcriptional silencing and chromosome stability. At present, we do not know why only about one-third of the cells appear to contain



FIG. 5. Effect of the SUM1-1 mutation on an $hmr\Delta 4$::TRP1 reporter gene in a rap1-12 or sir2 mutant background, as judged by the ability of cells to form colonies in the absence of tryptophan. SC, synthetic complete medium.

B SC SUM1, URA3/ADE2-TELVILI

FIG. 6. The SUM1-1 allele increases telomeric repression. (A) Colonies of SUM1 and SUM1-1 strains containing a chromosome VII-L telomeric ADE2/UR43 reporter are shown. (B) Growth of a SUM1-1 ADE2/UR43-TelVIIL strain in synthetic complete (SC) and FOA media compared with that of an isogenic SUM1 strain.

nuclear Sum1p. This could result from variable permeabilization of the spheroplasts, and hence differences in antibody accessibility, or from differences between cells in Sum1p levels or nuclear localization. We have also constructed an equivalent HA-tagged version of the *SUM1-1* allele. Unfortunately, this modified *SUM1-1* gene does not confer the *SUM1-1* silencing phenotype, perhaps because the epitope is located near the Sum1-1p mutation, and we have not determined its cellular localization.

URA3/ADE2-TEL VIIL, SUM1

DISCUSSION

Previous genetic analyses of the SUM1-1 allele have shown that it has the unique ability to suppress the silencing defects of mutations in a number of *cis*- and *trans*-acting silencer factors, including all four of the SIR genes (37). These results have suggested that wild-type Sum1p may play a critical role in silencing. However, lack of a *sum1* null allele and the variable dominance or recessiveness of the SUM1-1 mutation in differ-

 TABLE 3. Colony-forming abilities of SUM1-1 isogenic haploid cells on YEPD medium

Star in	Nc	o. of cells
Strain	Total	No growth
YDS712 (SUM1)	50	0
MC54 (SUM1-1)	50	9
YDS3 (SUM1)	50	3
MC89 (SUM1-1)	50	11
MC60 (SUM1)	50	0
MC61 (SUM1-1)	50	9

ent strain backgrounds have prevented a clear understanding of the function of either Sum1p or the *SUM1-1* mutant gene product. As a result, two very different models for Sum1p function have been proposed, one in which the protein is an activator required for the expression of silent mating-type genes which is itself repressed by Sir proteins (31) and another in which Sum1p is directly associated with an altered chromatin structure at *HM* loci (37).

URA3/ADE2-TEL VIIL, SUM1-1

Here we have reported the cloning and initial characterization of both the *SUM1* gene and the *SUM1-1* allele. The availability of these cloned genes has allowed us to examine the null phenotype of *SUM1* and also to study the effects of the *SUM1-1* mutation in a set of isogenic strains. Our results clearly indicate that a *sum1* null mutation does not suppress the effect of *sir* mutations at *HM* loci. Therefore, full expression of *HM* loci apparently occurs in the absence of Sum1p, ruling out the possibility that the protein is an activator required for transcription at the silent mating-type loci. Instead, we found that

 TABLE 4. Chromosome stability in homozygous SUM1, SUM1-1, and sum1 diploids

Genotype	Loss rate (no. of events [10 ⁻⁶]/cell division/generation)									
	Chromosome V	Chromosome III								
<u>SUM1 CAN1 HOM3</u> SUM1 can1 hom3	0.46 ± 0.29	0.86 ± 0.14								
<u>SUM1-1</u> <u>CAN1 HOM3</u> SUM1-1 can1 hom3	2.67 ± 0.49	4.37 ± 1.92								
sum1 CAN1 HOM3 sum1 can1 hom3	0.57 ± 0.11	2.80 ± 0.78								



FIG. 7. Indirect immunofluorescence and 4',6-diamidino-2-phenylindole (DAPI) staining of cells containing either HA-tagged SUM1 or the wild-type SUM1 gene. See text for details.

a sum1 null mutation has a slight derepressing effect on HM loci in SIR^+ strains when the HMR-E silencer is weakened by mutation of either the A or the B site. This subtle phenotype of sum1 null alleles explains why loss-of-function mutations in this gene were never isolated in previous genetic screens for silencing-defective mutants. Taken together, these results suggest that *SUM1-1* is either a gain-of-function (hypermorphic) mutation or an altered-function mutation. This conclusion is also consistent with previous observations (37) and results reported here, all of which indicate that the SUM1-1 allele is dominant to the wild type. However, our observations that elevated gene dosage of the wild-type SUM1 gene diminishes the mutant phenotype (suppression of sir2 mating defects) in SUM1-1 mutants and that the SUM1-1 plasmid confers a stronger phenotype in cells devoid of Sum1p than in the wild-type cells lead us to rule out the idea that SUM1-1 is a hypermorph.

By virtue of having cloned the *SUM1-1* allele, we have been able to examine the effect of this mutation in an isogenic set of strains. This analysis is particularly important given the previously reported strain-to-strain variation in the strength of SUM1-1 suppression (37) and the observation that SUM1-1 appears to be recessive to SUM1 in some strain backgrounds (31). We found a fairly uniform, dominant SUM1-1 phenotype in isogenic strains of the W303 background. The previously reported variation in the SUM1-1 phenotype may have been the result of natural strain background differences. Alternatively, the original SUM1-1 isolate, which was derived from a heavily mutagenized culture (31), may contain other mutations that modify the effect of SUM1-1. We noted that SUM1-1 only slightly restores repression of a $hmr\Delta A$::*TRP1* reporter in a *sir2* mutant, as measured by the ability to grow in the absence of tryptophan. We do not think that this observation is contradictory to the previous observation (37) that SUM1-1 restores

repression (at least 10-fold better) of *hmr::TRP1* in *sir3::LEU2* strains. The intact *HMR-E* silencer and the more leaky *sir3:: LEU2* mutation (12a) might account for the greater suppression observed in this previous study.

With respect to the differential effect of SUM1-1 on the two HM loci, our results are consistent with previous observations that SUM1-1 more effectively restores silencing at HMRa than it does at $HML\alpha$ (31, 37, 41). At present, there seem to be at least two possible explanations for this observation. The first follows from the finding that silencing at HMR is generally more resistant than HML to mutations in genes (other than SIR genes) which have partial effects on silencing. For example, *nat1* or *ard1* mutations have no effect at a wild-type HMR locus, whereas they result in partial derepression of HML (48). The same is true of a number of different mutations in the histone H4 (HHF2) N-terminal tail (29). This difference in the strength of silencing at HMR compared with HML, which may be due to the redundancy of the HMR-E silencer (7, 30) or the specific effect of the neighboring chromosome III-R telomere (70), could explain the apparent differential effect of SUM1-1. Alternatively, weak expression of α information in *MAT***a** cells may reduce mating more than weak expression of a information in $MAT\alpha$ cells. This could explain the observation that **a** information is better repressed than α information (as measured by mating efficiency) regardless of where the genes are located (HML or HMR) (31).

We also found that SUM1-1 suppresses two different types of mutations in the silencer-binding protein Rap1 $(rap1^t \text{ and } rap1^s)$. $rap1^t$ mutants are presumably defective in recruiting Sir3p and Sir4p to the silencers (47), whereas the $rap1^s$ mutants have been proposed to create a deficiency in Sir4p (and perhaps other factors) available at the *HMR* silencer, as a result of competition by telomeres (10). We imagine that SUM1-1 sup-

presses these mutations by overcoming the requirement for Sir3p and Sir4p in silencing, thus bypassing the Sir protein recruitment function of Rap1p.

Because many of the same genes involved in HM locus silencing are also involved in the variegated silencing phenomenon observed at telomeres, we also tested the effects of sum1 and SUM1-1 mutations on telomeric repression. Although we found no effect of a sum1 null mutation on telomeric silencing, we observed that SUM1-1 increases telomeric repression of two different telomeric reporters genes (URA3 and ADE2) in SIR strains. However, SUM1-1 does not suppress the telomeric silencing defect caused by a sir2 mutation. The first observation demonstrates that the effect of SUM1-1 is not restricted to the silent mating-type loci and suggests that SUM1-1 should be viewed as a general regulator of position effects in S. cerevisiae. The failure of SUM1-1 to suppress sir defects at telomeres may be viewed within the context of several observations which indicate that telomeric silencing is inherently weaker than repression at HM loci. To begin with, telomeric silencing is normally unstable, whereas HM silencing is not (2, 21). In addition, telomeric silencing is more sensitive to histone H4 and histone H3 mutations, and mutations in the NAT1 and ARD1 genes, than are HM loci (2, 71). Finally, a sir3 suppressor mutation which partially restores HML silencing in a hhf2 mutant strain (K16G) fails to restore telomeric repression (2). Taken together, these observations are consistent with the idea that the effect of SIR mutations (SIR2 to SIR4) on telomeric silencing may simply be too severe for SUM1-1 to counteract.

In thinking about how SUM1-1 suppression might work, it is important to consider current models for silencing at HM loci and telomeres. Recent studies indicate that a complex of Sir proteins (containing at least Sir3p and Sir4p) can interact directly with the silencer- and telomere-binding protein Rap1p (47). Sir3p and Sir4p, in turn, are capable of binding in vitro with the N-terminal tails of histones H4 and H3 (24), which genetic studies have shown are involved in both HM locus and telomeric silencing (29, 45, 51, 71). These findings have led to a model in which silencing results from the recruitment of a Sir complex to silencers or telomeres and the subsequent assembly of a Sir-nucleosome complex along the chromatin fiber. Silent chromatin appears to be in an altered, more protected structure than nonsilent chromatin, as determined from its decreased accessibility to methylases (20, 36, 65), the HO-encoded endonuclease (32, 42, 67), restriction enzymes (42), and thiol-reactive reagents (12). However, a growing number of cell biological studies of S. cerevisiae suggests that nuclear localization, or more specifically attachment to the nuclear envelope, may also play an important role in silencing. Indirect immunofluorescence studies using anti-Rap1p antibodies suggest that telomeres are clustered in yeast cells and localized at or near the nuclear periphery (33, 50). Strikingly, this localization and clustering of telomeric Rap1p is lost in sir3 or sir4 mutant cells (50), in which telomeric silencing is also abolished.

An intriguing and perhaps informative phenotype of *SUM1-1* mutants is increased chromosome loss. We note that many genes involved in silencing also cause chromosome instability when they are mutated or deregulated. For example, $rap1^t$ alleles (rap1-17) display telomere elongation and elevated chromosome instability (35). Overexpression of the *RAP1* C terminus or of *SIR2* causes increased chromosome instability and cell death, similar to that which we have observed in *SUM1-1* mutants (8, 14), albeit to different degrees. In light of these results and the current working models for silencing, we suggest two models for Sum1p and Sum1-1p function, both of

which might explain the chromosome instability and decreased cell viability phenotypes of SUM1-1 mutants. In the first model, Sum1p is involved in the localization of telomeres and HM loci to the nuclear periphery. The altered localization function provided by Sum1-1p is sufficient to allow silencing to occur in the absence of SIR gene function. An increase in chromosome loss might be a consequence of this enhanced nuclear envelope attachment function. Alternatively, Sum1p may be a normal (but nonessential) component of heterochromatin in S. cerevisiae which is used for both gene silencing and chromosome condensation during mitosis. In this scenario, the mutant Sum1-1p may allow HM loci (and perhaps other sites not normally subject to silencing) to form a stably repressed chromatin structure in the absence of Sir proteins. This inappropriate formation of heterochromatin could either directly or indirectly lead to a decrease in chromosome stability. It is important to note, however, that we cannot rule out the possibility that the chromosome loss and decreased viability phenotypes of SUM1-1 are unrelated to its effect on silencing.

In summary, by cloning and characterizing SUM1 and the SUM1-1 mutant allele, we have obtained clear evidence that SUM1-1 is a dominant altered-function mutation that can either restore or improve silencing at HM loci and telomeres in a number of different genetic backgrounds (e.g., *sir*, *rap1^t* or *rap1^s* mutant or wild type). The nuclear localization of Sum1p, the effect of SUM1-1 on chromosome stability and cell viability, and the ability of the mutation to bypass the requirement for *SIR* gene function in *HM* locus silencing all point to a role for *SUM1* in chromosome function. Continued study of *SUM1* and the *SUM1-1* allele should provide new experimental approaches to address the precise function(s) of this intriguing gene.

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