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Surveying *Saccharomyces* Genomes to Identify Functional Elements by Comparative DNA Sequence Analysis

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Comparative sequence analysis has facilitated the discovery of protein coding genes and important functional sequences within proteins, but has been less useful for identifying functional sequence elements in nonprotein-coding DNA because the relatively rapid rate of change of nonprotein-coding sequences and the relative simplicity of non-coding regulatory sequence elements necessitates the comparison of sequences of relatively closely related species. We tested the use of comparative DNA sequence analysis to aid identification of promoter regulatory elements, nonprotein-coding RNA genes, and small protein-coding genes by surveying random DNA sequences of several *Saccharomyces* yeast species, with the goal of learning which species are best suited for comparisons with *S. cerevisiae*. We also determined the DNA sequence of a few specific promoters and RNA genes of several *Saccharomyces* species to determine the degree of conservation of known functional elements within the genome. Our results lead us to conclude that comparative DNA sequence analysis will enable identification of functionally conserved elements within the yeast genome, and suggest a path for obtaining this information.

Identifying functional elements in DNA sequence is a significant challenge. It is difficult enough to predict correctly protein-coding genes; an even greater challenge is to identify functional sequences that do not code for protein, such as sequences regulating gene expression, sequences governing chromosome replication, structure and stability, and sequences of nonprotein-coding RNAs. This task is complicated by the diverse nature of these sequence elements. For example, sequences that regulate gene expression are usually short, often independent of orientation, and can reside at varying distances from their target gene. Genes encoding RNAs can be difficult to identify because they contain few hallmarks and because their folded structure, rather than their primary sequence, dictates their function.

Because functional sequences are maintained in evolution, they often can be recognized by their conservation among different organisms (Tagle et al. 1988; Hardison et al. 1997). This approach has most often been used to identify functional sequences in proteins, because they are relatively large and evolve relatively slowly, causing their functional elements to remain conserved in diverse organisms. In a few cases potential nonprotein-coding functional sequence elements have been identified by comparing DNA sequences of relatively diverged species (usually from different genera) (Venkatesh et al. 1997; Gellner and Brenner 1999; Wentworth et al. 1999; Gelfand et al. 2000; Kent and Zahler 2000; McGuire et al. 2000). The comparative approach for finding functional nonprotein-coding sequences will likely be more

productive with DNA sequences of relatively closely related organisms. We sought to gain experience with this by comparing the DNA sequences of yeasts of the *Saccharomyces* genus because (1) the complete *Saccharomyces cerevisiae* genome sequence is available for comparison (Goffeau et al. 1996); (2) the genomes of *Saccharomyces* species are relatively small, with relatively compact noncoding sequences (~30% of the genome); (3) their phylogeny is well characterized, with many related species at various evolutionary distances; and (4) it is possible to test the functionality of sequences with relatively simple and incisive experiments.

The *Saccharomyces* genus is composed of three subgroups (Barnett 1992). The sensu stricto species are physiologically similar to *S. cerevisiae*, are capable of forming stable diploids with each other, and have a very similar karyotype. The sensu lato and petite-negative *Saccharomyces* species are quite diverged from *S. cerevisiae*. They have significantly different physiological characteristics, seldom form diploids with *S. cerevisiae*, and have a smaller number of chromosomes (Petersen et al. 1999). To identify which *Saccharomyces* species are optimally diverged from *S. cerevisiae* for comparative DNA sequence analysis, and to evaluate the number of sequences that must be compared to obtain useful information, we determined the sequence of randomly generated genomic clones of seven *Saccharomyces* species. In addition, we determined the DNA sequences of the regulatory regions of a few specific genes and of a few genes encoding nonprotein-coding RNAs from several *Saccharomyces* species to determine the degree of conservation of known functional elements in the genome. Our results suggest that comparative DNA sequence analysis will enable identification of many functional gene regulatory elements and other nonprotein-coding sequences, and suggest a path for obtaining this information.

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RESULTS

Sequence Summary of *Saccharomyces* Species

We obtained sequence from ~1000 genomic DNA clones from each of four species of the sensu stricto group (*Saccharomyces bayanus*, *Saccharomyces paradoxus*, *Saccharomyces cariocanus*, and *Saccharomyces mikatae*) (Naumov et al. 2000) and from >2000 clones from each of two species of the sensu lato group (*Saccharomyces castellii* and *Saccharomyces unisporus*) and the petite-negative species *Saccharomyces kluyveri* (Table 1). Over 4.3 Mb of unassembled sequence was obtained, a sample sufficient for genomic exploration of the *Saccharomyces* genus.

Comparisons of the nucleotide sequences (using BLASTN, Table 1) and inferred protein sequences (using BLASTX, Table 2) to *S. cerevisiae* sequence are instructive for determining the relative genetic distance of the species from *S. cerevisiae* and for identifying which species are likely to yield the most data in sequence comparisons. Most (94.5%–98.2%) of the sensu stricto sequences align to *S. cerevisiae* coding or noncoding sequence using BLASTN. As expected, the sequences are most similar to *S. cerevisiae* DNA sequence in protein-coding genes, but the sequences of *S. paradoxus* and *S. cariocanus* are also highly conserved within nonprotein-coding regions. *S. mikatae* and *S. bayanus* have lower similarity and yield significantly fewer alignments to nonprotein-coding regions of the genome using the same BLASTN parameters (Table 1).

The sensu lato and petite-negative species seem to lie a similar evolutionary distance from *S. cerevisiae*, because the average identities of their protein-coding DNA (Table 1) and predicted proteins (Table 2) to *S. cerevisiae* sequences are similar for each sensu lato species. These species are more diverged from *S. cerevisiae* than we expected from ribosomal RNA comparisons. Many presumably orthologous DNA sequences (identified by aligning the protein-coding sequences using BLASTX) do not align to *S. cerevisiae* sequence with BLASTN, and many of those that do align are near the lower limits of detection by BLASTN.

Although most of the proteins of the sensu stricto group

Table 1. Summary of BLASTN Comparisons of *Saccharomyces* Species' DNA Sequences to *S. cerevisiae* DNA Sequence

Species	# of sequences	% identity CODING	% identity NON-CODING	% of sequences aligning CODING/NON
<i>S. paradoxus</i>	728	88.3	81.6	77.9/36.3
<i>S. cariocanus</i>	867	88.0	80.5	77.5/39.1
<i>S. mikatae</i>	1136	83.6	74.8	79.9/27.9
<i>S. bayanus</i>	851	79.7	73.6	65.7/19.5
<i>S. castellii</i>	2290	70.3	62.5*	48.9/1.7*
<i>S. kluyveri</i>	2145	70.2	72.8*	38.6/1.8*
<i>S. unisporus</i>	2357	69.2	67.6*	40.4/2.2*

Coding sequences are all ORFs known or predicted to encode protein. The remaining yeast genomic sequences (minus the sequences encoding RNA genes, transposable elements, and CEN sequences) were considered non-coding. BLASTN comparisons of sensu stricto yeast species used MATCH/MISMATCH parameters M = 5, N = 5 with the default gap penalties. This change was made to more accurately reflect the similarity of the sequences and equals a PAM distance of ~30 (States et al. 1991). Using these strict MATCH/MISMATCH parameters, only significant alignments were obtained. Default parameters were used for sensu lato and petite-negative comparisons to *S. cerevisiae* using a wordlength (W) of 5. The dust filter was used to mask low complexity nucleotide sequences. Alignments with a *P*-value <5.0e-06 were counted as significant.

*Many of the alignments are to repetitive sequences (ribosomal rDNA repeat alignments were not counted).

Table 2. Summary of BLASTX Comparisons between *Saccharomyces* Species' Protein Sequences and *S. cerevisiae* Protein Sequence

Species	# of sequences	% of sequences aligning	% sequence identity
<i>S. paradoxus</i>	728	78.3	91.2
<i>S. cariocanus</i>	867	78.8	90.3
<i>S. mikatae</i>	1136	81.8	87.4
<i>S. bayanus</i>	851	83.2	83.2
<i>S. castellii</i>	2290	71.0	60.6
<i>S. kluyveri</i>	2145	63.7	59.8
<i>S. unisporus</i>	2357	55.0	59.6

BLASTX comparisons of protein coding sequences included only the highest scoring pair in most cases. Additional HSPs were included only if the *S. cerevisiae* encoded proteins were adjacent on a chromosome and could be spanned by a sequence read (123 of the sensu stricto sequences and 104 of the sensu lato and petite-negative sequences aligned to adjacent proteins). Sensu stricto comparisons were performed with a PAM matrix of 40 whereas sensu lato and petite-negative comparisons used the default BLOSUM62 matrix. Several additional alignments could be generated using alternative matrices (such as PAM120) and altered GAP penalties, but essentially equal numbers of alignments were generated using less stringent matrices since these parameters often miss short alignments near the ends of the proteins. The seg and xnu filters were used to mask low complexity protein coding regions. Essentially all of the alignments are significant when the filters are used.

are >80% identical to their *S. cerevisiae* homologs, 29 alignments show <50% identity (Fig. 1A). These low similarity alignments may indicate rapidly evolving proteins or genes that duplicated in other *Saccharomyces* species (but not in *S. cerevisiae*), allowing one copy to diverge from its *S. cerevisiae* ortholog. A large number of proteins of the sensu lato species (>30%) are <50% identical to their *S. cerevisiae* homologs (Fig. 1B), emphasizing the substantial divergence of these species from *S. cerevisiae*.

We noted many discrepancies in open reading frame (ORF) boundaries in the different sensu stricto species (data not shown). Some of these are likely caused by sequencing errors (most likely in our sequence, but potentially in the *S. cerevisiae* sequence); some may possibly be attributable to real DNA sequence differences between these closely related species. ORF length polymorphisms are even more common in the sensu lato alignments, which is not surprising considering the relatively greater divergence of these species from *S. cerevisiae*.

Species-Specific Sequences

A small percentage of the sequences of sensu stricto species have no significant similarity to any *S. cerevisiae* sequence at the nucleotide or amino acid levels, regardless of the BLAST parameters that were used. Several of these sequences are pre-

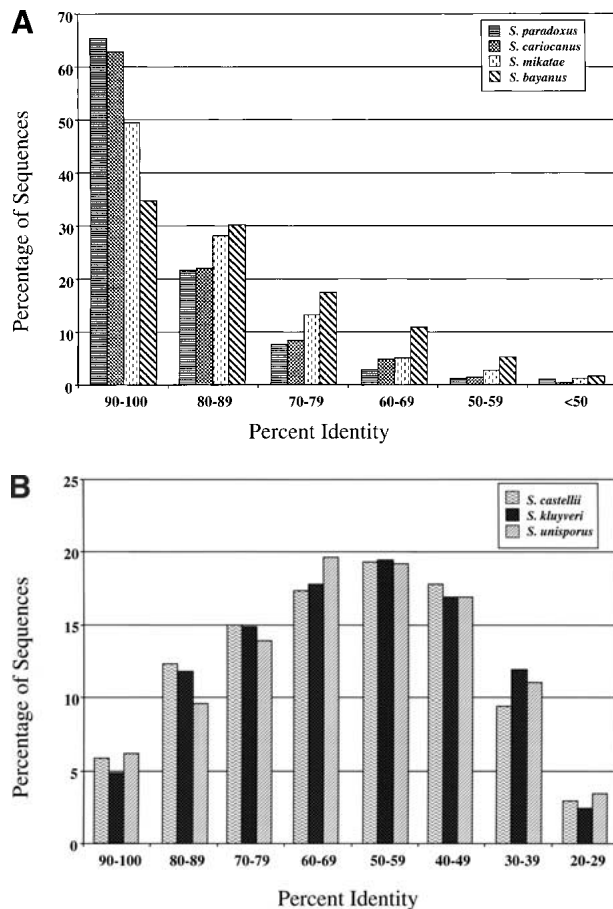


Figure 1 Summary of protein sequence identities between *Saccharomyces* sequences and their *S. cerevisiae* homologs. (A) Sensu stricto BLASTX comparisons to *S. cerevisiae* using a PAM40 matrix. (B) Sensu lato and petite-negative comparisons using the BLOSUM62 matrix.

dicted to encode proteins similar to those found in other species. For example, we identified a Ty5 transposon protein that is unique to *S. paradoxus*, and a gene in *S. bayanus* and *S. cariocanus* predicted to encode a protein related to an amidase of *S. pombe*. Many sensu lato and petite-negative sequences are predicted to encode proteins similar to those in species other than *S. cerevisiae* (Table 3). Similar findings have been made in *Kluyveromyces lactis* (Ozier-Kalogeropoulos et al. 1998) and other hemiascomycete species (Souciet et al. 2000).

To verify that these DNA sequences are specific to certain *Saccharomyces* species, we designed oligonucleotide primers to the amidase-encoding gene identified in *S. cariocanus*, and to two other unique sequences confined to *S. mikatae* and *S. paradoxus* that do not appear to encode protein. The sequences were amplified by PCR from genomic DNA of the corresponding species and produced products of the expected size. (Three other species-specific sequences produced PCR products larger than expected.) The amidase PCR product from *S. cariocanus* hybridized to a specific DNA fragment of *S. cariocanus* and *S. paradoxus* DNA in Southern blots (data not shown). The *S. mikatae*-specific sequence hybridized to genomic DNA from *S. mikatae*, but not to genomic DNA from the other sensu stricto species. Because we determined the DNA sequence of only about 1/32nd of

the four sensu stricto species, it is likely that several more *Saccharomyces* sequences not present in *S. cerevisiae* remain to be identified.

Identification of Small Protein-Coding Genes

Some of the intergenic regions in the *S. cerevisiae* genome likely encode small proteins (<100 amino acids) that have not been annotated. Comparisons of *S. cerevisiae* intergenic sequences to those of the other *Saccharomyces* species using TBLASTX revealed many potential protein-coding sequences. Such comparisons of the sensu stricto sequences are not very informative because their relatively high degree of similarity to *S. cerevisiae* sequence produces many spurious alignments, but the sequences of the sensu lato and petite-negative species are well suited for identification of small ORFs (smORFs, <100 codons) because their significant divergence from *S. cerevisiae* sequence yields relatively few TBLASTX alignments outside of protein-coding sequences. Most of the high-scoring TBLASTX alignments to *S. cerevisiae* sequence are extensions or fusions of known ORFs, but we identified 11 alignments of smORFs that potentially encode a protein (TBLASTX *P* value <1.0e-05; Table 4). Two of the smORFs were predicted previously by searching the *S. cerevisiae* genome for transcripts that originate from large intergenic regions of the genome (Olivas et al. 1997); two smORFs are similar to small ORFs in other yeast species (Table 4).

Identifying Functional Non-Protein-Coding Sequences

Our primary goal is to use sequence conservation to identify functional nonprotein-coding sequences. However, much of the similarity of the nonprotein-coding sequences of the sensu stricto species is likely caused simply by an inadequate amount of evolutionary time for the accumulation of sequence changes. Which species are sufficiently diverged so that functional nonprotein-coding sequences are apparent? How many different sequences need to be compared to reveal regions of sequence similarity that are functionally significant? We began to address these questions by searching for nonprotein-coding RNA genes and promoter regulatory sequences in non protein-coding portions of the *S. cerevisiae* genome.

Identification of Non-Protein-Coding RNAs

Many highly conserved sequences in intergenic regions were identified that are unlikely to encode proteins because no ORF was apparent in the *S. cerevisiae* genome sequence. In the few cases where we obtained sequences from more than one species that are similar to the same *S. cerevisiae* sequence, the conserved sequence elements readily stand out from surrounding sequences. Because there are only a few regions of the genome for which we have sequences from multiple species, we determined the DNA sequence of a few genes encoding nonprotein-coding RNAs from many different *Saccharomyces* species to compare how quickly they diverge across the genus, and to help us decide which *Saccharomyces* species are best suited for identifying these genes by comparative sequence analyses. We amplified the sequences by PCR using oligonucleotide primers chosen from *S. cerevisiae* flanking sequences that seemed likely to be conserved. We determined the sequence of *SNR39*, encoding a C/D

Table 3. *Saccharomyces* Genes not Present in *S. cerevisiae*

Species	Predicted protein	Closest homolog	P-value	Accession no.
<i>S. bayanus</i>	putative amidase	<i>S. pombe</i>	2.3e-24	pir T39112
<i>S. cariocanus</i>	putative amidase	<i>S. pombe</i>	8.9e-46	
<i>S. paradoxus</i>	Ty-5 associated protein	<i>S. paradoxus</i>	8.6e-40	gb U19263.1
<i>S. castellii</i>	Squalene synthase	<i>C. glabrata</i>	2.6e-07	dbj BAB12207.1
	NAD(P)H dehydrogenase	<i>P. aeruginosa</i>	3.1e-40	gb AAG04613.1
	hypothetical protein	<i>C. albicans</i>	6.2e-35	emb CAA21971.1
	putative p150	<i>H. sapiens</i>	5.1e-25	gb AAA88038.1
	unnamed protein	<i>H. sapiens</i>	5.3e-09	dbj BAA91193.1
	hyphally regulated protein	<i>C. albicans</i>	4.7e-08	pir S58135
<i>S. kluyveri</i>	hypothetical protein	<i>X. fastidiosa</i>	1.8e-08	gb AAF83652.1
	hypothetical protein	<i>S. pombe</i>	5.9e-30	pir T42242
	RNA binding protein, pumilio-family	<i>S. pombe</i>	1.3e-09	pir T41065
	probable dipeptidase	<i>S. pombe</i>	3.0e-26	pir T41665
	WSC4 homologue	<i>K. lactis</i>	6.5e-11	emb CAB50897.1
	hypothetical protein jhp1324	<i>H. pylori</i>	6.9e-15	pir G71821
	coiled-coil protein	<i>S. Pombe</i>	9.1e-13	pir T40559
	sphingomyelin phosphodiesterase	<i>M. musculus</i>	1.2e-17	pir S27393
	neuronal thread protein AD7c-NT	<i>H. sapiens</i>	1.6e-14	gb AAC08737.1
	ser/arg-rich protein specific kinase	<i>M. musculus</i>	3.9e-06	pir C5930
	hypothetical protein SPAC630.13c	<i>S. pombe</i>	2.9e-19	pir T38991
	purine nucleoside permease	<i>C. albicans</i>	1.8e-41	gb AAC64324.1
	beta-glucosidase	<i>K. lactis</i>	2.1e-59	emb CAA29353.1
	stearyl-CoA desaturase	<i>C. albicans</i>	2.1e-06	pir T18228
	iron/ascorbate dependent oxidoreductase	<i>C. jejuni</i>	7.6e-08	emb CAB73453.1
	N-carbamoyl-amidohydrolase	<i>P. aeruginosa</i>	1.5e-07	gb AAG03833.1
	Nef attachable protein	<i>H. sapiens</i>	2.9e-08	dbj BAA95214.1
	hypothetical protein B0252.2	<i>C. elegans</i>	1.5e14	pir T15291
	unnamed protein product	<i>H. sapiens</i>	7.2e-20	dbj BAA91131.1
	CLIP-170	<i>R. norvegicus</i>	6.5e-06	emb CAB92974.1
	putative transaminase	<i>L. lactis</i>	1.5e-19	emb CAB97148.1
	splicing factor u2af 35 kd subunit	<i>S. pombe</i>	1.6e-18	pir T39243
	cytoskeleton assembly control protein	<i>C. albicans</i>	3.7e-09	gb AAC62773.1
	Rbt2p (repressed by TUP1)	<i>C. albicans</i>	2.0e-08	gb AAG09788.1
	uric acid-xanthine permease	<i>E. nidulans</i>	1.7e-10	emb CAA50681.1
<i>S. unisporus</i>	hypothetical protein SPAC1687.07	<i>S. pombe</i>	3.4e-11	pir T37750
	homolog of co-factor B Alp11p	<i>S. pombe</i>	1.5e-10	pir T38860
	polyprotein	<i>O. sativa</i>	2.1e-32	gb AAD27547.1

Alignments having a *P* value <1.0e-05 are shown. Sequences that had high scoring alignments to known genes were not included if they also had significant similarity to *S. cerevisiae* rDNA sequence.

box snoRNA required for methylation of the 25s ribosomal subunit (Kiss-Laszlo et al. 1996) and *SNR44*, encoding an H/ACA box snoRNA involved in pseudouridylation of ribosomal RNA subunits (Ganot et al. 1997), both of which are located within introns of well-conserved ribosomal protein genes.

All eight *SNR39* sequences align well using CLUSTALW, with 58% of the 93 nucleotides of *SNR39* being conserved among eight *Saccharomyces* species. The C and D boxes as well as the guide sequence are conserved perfectly throughout the genus (Fig. 2). The gene is even readily distinguishable from surrounding intron sequence in most two-way alignments with *S. cerevisiae* sequence, although it is indistinguishable in an alignment of *S. cerevisiae* sequence to *S. paradoxus* (too similar), and is difficult to discern in an alignment to *S. exiguus* sequence (too diverged).

The *SNR44* gene (encoding an H/ACA snoRNA) sequences also align using CLUSTALW, but are less well conserved than *SNR39* (Fig. 3). The alignment reveals four highly conserved blocks, ranging from 5 to 15 nucleotides in length, which, surprisingly, do not include the ACA sequence (AAA in

the sensu stricto species) or the H-box (Fig. 3), known functional elements in this snoRNA

Identification of Gene Regulatory Sequences

We expect potential gene regulatory sequences to be manifested as short blocks of sequence similarity in intergenic regions of the genome. These are difficult to recognize in the random DNA sequences of the *Saccharomyces* species for two reasons. On one hand, the sequences of the sensu stricto species are usually so similar to *S. cerevisiae* sequence that few isolated runs of identical nucleotides are found. In rare cases where we can align DNA sequence of two or more species, the background sequence similarity is reduced and potential regulatory elements begin to stand out (e.g., Fig. 4), but these alignments never extend over the entire promoter. On the other hand, the DNA sequences of the sensu lato species are generally too different from *S. cerevisiae* sequence to align with local alignment algorithms (such as BLASTN or BEST-FIT). Some sensu lato sequences can be anchored to their presumed orthologs in *S. cerevisiae* gene using BLASTX, then

Table 4. Small *S. cerevisiae* ORFs (smORFs) with Homologs in Other *Saccharomyces* species

Location	Length	Species	% ID	Similar proteins
Chr2 363049-362771	92#	<i>S. unisporus</i>	25/36 (69%)	
Chr2 419123-418869*	85	<i>S. kluyveri</i>	52/77 (67%)	
Chr2 684936-685216+	94	<i>S. castellii</i>	51/91 (56%)	
Chr3 100839-101343	142#	<i>S. unisporus</i>	67/124 (54%)	<i>S. pombe</i>
Chr4 603805-603590+	72	<i>S. kluyveri</i>	35/81 (43%)	<i>C. elegans</i>
Chr4 691007-691204	66	<i>S. kluyveri</i>	34/69 (49%)	
Chr5 261045-260935*	37	<i>S. kluyveri</i>	25/45 (55%)	
Chr5 551117-550863	85	<i>S. unisporus</i>	42/66 (63%)	
Chr7 836659-836384	92	<i>S. castellii</i>	23/61 (37%)	
Chr10 159545-159324	74	<i>S. kluyveri</i>	40/50 (80%)	
Chr10 316571-316377+	65	<i>S. castellii</i>	38/63 (60%)	
		<i>S. kluyveri</i>	20/25 (80%)	

Location refers to SGD nucleotide coordinates. smORFs with an asterisk correspond to smORFs identified by Olivas and Parker (1997). Amino acid lengths followed by a pound sign indicate that a frame shift or an RNA splice is required to create the reading frame in the current annotation of the *S. cerevisiae* sequence. Species encoding sequences similar to the smORFs are listed (*S. pombe* accession no. T37750, protein information resource; *C. elegans* accession no. AAF59588.1, GenBank. A (+) refers to smORFs that are conserved in other hemiascomycetes yeast species (Blandin et al. 2000).

extended into the promoter region. Of the 4296 sensu lato sequences that align to *S. cerevisiae* proteins with BLASTX, 866 extend >30 nucleotides into the promoter region. We were able to align only 398 of them to their *S. cerevisiae* ortholog (using BLASTN, with a word length of five and only 111 of the alignments extend >100 bp into the promoter region. Many potential regulatory elements are apparent in these alignments. For instance, we can clearly see conservation of DNA-binding sites for the Alpha1 and Mcm1 proteins in the *MFA2* and *STE3* promoters of *S. castellii* (Fig. 5A,B). A consensus Hap2 binding site (ACCAATNA; Svetlov and Cooper 1995) is included in one of the three conserved runs of sequence present in the promoter of *ATP14* (Fig. 5C), a gene that is plausibly regulated by Hap2. Many other alignments contain short runs of conserved sequence that are potential gene regulatory elements.

To assess *Saccharomyces* species more broadly for their suitability for identifying regulatory sequences in gene pro-

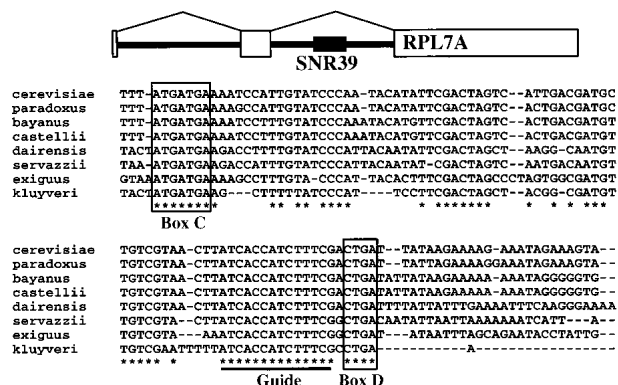


Figure 2 A CLUSTALW alignment of *SNR39* sequences (encoding a snoRNA) from eight different *Saccharomyces* species. Box C and Box D are known functional elements; the "guide" is the sequence complementary to rRNA sequence adjacent to the methylation site. The structure of the *RPL7A* transcript including the intronic *SNR39* gene is shown above the sequence alignment.

moters by comparative DNA sequence analysis, we amplified by PCR and determined the sequences of the promoters of three well-characterized genes from many *Saccharomyces* species (see Methods for details). Although the sequences of the promoter region between *GAL1* and *GAL10* of the sensu stricto species align well using CLUSTALW, the sensu lato species' sequences do not align. We searched for Gal4 and Mig1 protein-binding sites by a simple pattern search [FINDPATTERNS] and, as expected, they are apparent in the *GAL1-10* promoter of all 10 species (Fig. 6). The spacing of these binding sites is well conserved in the sensu stricto species, with only a few sites missing from some of the species. The spacing and number of Gal4 and Mig1 binding sites are not conserved in the sensu lato species. Of these,

only the *S. castellii* sequence aligned with the *S. cerevisiae* sequence using local alignment algorithms (BLASTN or BESTFIT).

We obtained from each species the DNA sequence of at least one of the two nearly identical copies of the divergently transcribed *HHT* and *HHF* genes (*HHT1-HHF1* and *HHT2-HHF2*), encoding histones H3 and H4, respectively. In some cases we obtained the sequence of both copies (though they are so diverged from the *S. cerevisiae* sequence in the sensu lato species that it was usually difficult to distinguish between the two copies using BLASTN alignments). Although conserved blocks of sequence begin to emerge from 3-way CLUSTALW alignments of sequences of the sensu stricto species (data not shown), addition of a sensu lato sequence makes these elements clearly stand out (Fig. 7A). Both (TATA) boxes are conserved, as are sequences within and near CCA boxes 1 and 2 that were defined previously as regulatory elements of this promoter (Freeman et al. 1992). One of the conserved sequences (enclosed in dashed lines in Fig. 7A) is present only in one of the *HHT-HHF* copies in each species, so the two pairs of genes may be regulated differently. Other potential CCA boxes (CCA box 3 and 4) that were not recognized previously seem apparent. Using the pattern recognition program AlignAce (Roth et al. 1998), we identified many variations of the CCA box within the histone promoters (60 different motifs in the 13 promoter sequences) some of which were not identified by full-length sequence alignments (Fig. 7B); the logo consensus (Schneider and Stephens 1990) CCA box sequence is shown in Figure 7C. Each copy of the divergent promoter contains three to six of the relaxed CCA box sequence motifs.

We succeeded in amplifying the *GAL4* promoter from only three *Saccharomyces* species (presumably because the flanking protein-coding sequences are not well conserved). One of these sequences (from *S. paradoxus*) is too similar to the *S. cerevisiae* sequence to be useful, but alignments of the other two sequences (one from the sensu stricto species *S. bayanus*, the other from the petite-negative species *S. kluyveri*) seem very informative (Fig. 8). The promoter elements that were previously defined genetically (Griggs and Johnston

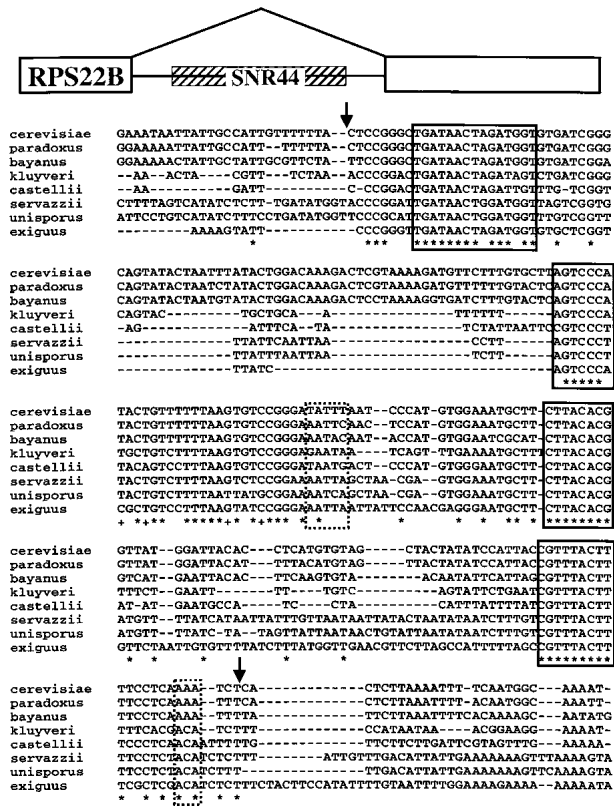


Figure 3 A CLUSTALW alignment of *SNR44* sequences encoding an H/ACA box snoRNA. The positions of the start and end of *S. cerevisiae SNR44* are marked with arrows. Highly conserved sequences are boxed. The H and ACA boxes are enclosed in dashed lines. The plus symbols in the consensus line indicates that seven of eight nucleotides in the alignment are conserved and were added to highlight conservation of sequence upstream of the H box. The structure of the *RPS22B* transcript including the intronic *SNR44* gene is shown above the sequence alignment.

1993) clearly stand out as conserved in a CLUSTALW alignment of these sequences with the *S. cerevisiae* sequence.

DISCUSSION

We have investigated the feasibility of using comparative DNA sequence analysis to identify functional sequences in the genome of *S. cerevisiae*, with the goal of identifying regulatory sequences and sequences specifying nonprotein-coding RNAs. We are confident that promoter regulatory sequences,



Figure 4 A CLUSTALW alignment of *SEC35* promoter sequences of sensu stricto species reveals conserved elements that could function as regulatory elements. The conserved sequences are boxed; the *S. cerevisiae* ATG start codon and putative start codons for the other species are underlined.

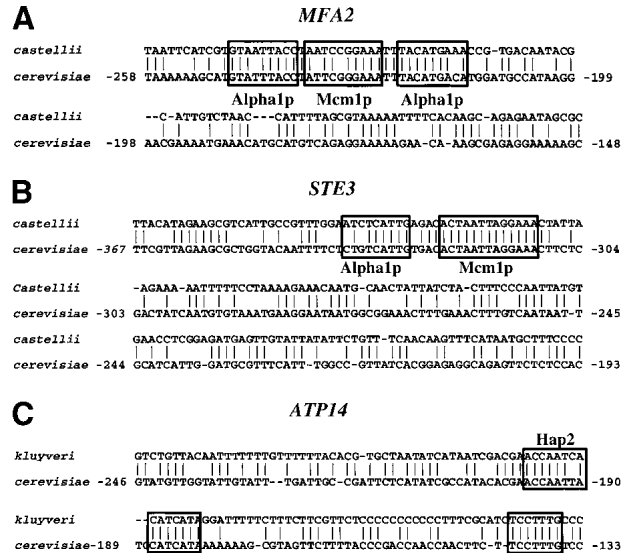


Figure 5 Known functional elements are conserved in BLASTN alignments between *S. cerevisiae* promoters and orthologous sequences of sensu lato species. (A) An alignment of *MFA2* promoter sequences from *S. castellii* and *S. cerevisiae* with known binding sites for Alpha1 and Mcm1 boxed. (B) An alignment of *STE3* promoter sequences from *S. castellii* and *S. cerevisiae* with known binding sites for Alpha1 and Mcm1 boxed. (C) An alignment of the *ATP14* promoter sequences from *S. kluyveri* and *S. cerevisiae* showing three conserved runs of sequence, one of which includes a consensus Hap2 (ACCAATNA) binding site. Numbers refer to the position of the *S. cerevisiae* sequence relative to the ATG start codon.

RNA genes, and smORFs could be identified by comparisons of orthologous *Saccharomyces* sequences.

Analysis of Protein-Coding Genes

Before discussing our analysis of non-protein-coding sequence, a few observations regarding protein-coding genes

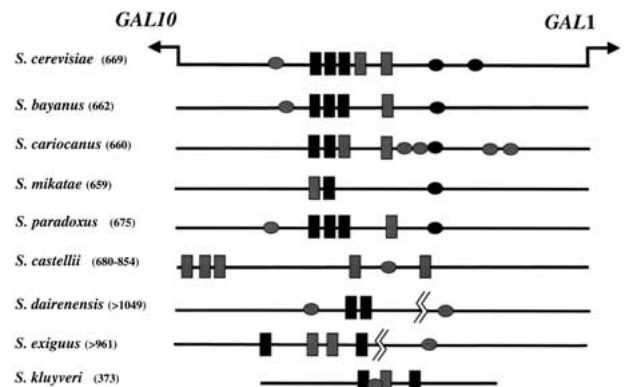


Figure 6 The location of Gal4 and Mig1 regulatory elements in the *GAL1-GAL10* promoter sequences from many *Saccharomyces* species. The length of the promoter (where known) is shown in parentheses. Gal4 binding sites are shown as black boxes with gray boxes indicating putative binding sites that are altered by one nucleotide from the consensus site (CGG_nCCG). Potential Mig1 binding sites are shown as ellipses. Black ellipses refer to experimentally verified Mig1 sites (in *S. cerevisiae*) or Mig1 sites that clearly align to *S. cerevisiae* sites in pairwise alignments. Gray ellipses refer to putative Mig1 sites (CCC followed by AT-rich sequence).

a conserved hexamer is expected to occur by chance about once in every eight promoters on average. We realize that these “back of the envelope” calculations must be interpreted cautiously because it is unlikely that even a probable optimal multiple sequence alignment will correspond precisely to the biological events that generated the sequences being aligned (Thorne and Churchill 1995), but in any case, four-way alignments seem to be the minimum required for efficient recognition of conserved regulatory sequences.

Candidate Species for Comparative Analysis

Which species' sequences will yield the most information? Two considerations need to be balanced in making this decision. The sequences need to be similar enough so that most of them can be aligned with their *S. cerevisiae* ortholog, a requirement that favors the sensu stricto species, as more of their sequence reads can be aligned to *S. cerevisiae* sequence (Tables 1, 2). Conversely, if the sequences are too similar the functional elements do not stand out, a consideration that favors the sensu lato species. It seems clear that at least one of each group of species need to be compared.

Two of the sensu stricto species for which we obtained DNA sequence — *S. mikatae* and *S. bayanus* — seem like good candidates for large-scale sequence comparisons. *S. cariocanus* and *S. paradoxus* seem too similar to *S. cerevisiae* for this purpose (Tables 1, 2). Another good candidate for comparative sequence analysis is *S. kudriavzevii*, a recently described species (Naumov et al. 2000), that seems to lie between *S. bayanus* and *S. mikatae* in its evolutionary distance from *S. cerevisiae* (Fischer et al. 2000).

There are few differentiating characteristics for selecting one sensu lato species over another, and sequence information from any of these species will likely be useful. The species that we studied are a similar evolutionary distance from *S. cerevisiae*. *S. unisporus* is topologically much closer to *S. cerevisiae* than *S. kluyveri* in most phylogenies, but some show a much longer branch length for *S. unisporus* (James et al. 1997; Kurtzman and Robert 1991; Oda et al. 1997), consistent with our data. Two species, *S. servazzii* and *S. dairenensis* branch closely with species that we studied (*S. unisporus* and *S. castellii*, respectively). It is unlikely that any known sensu lato species are significantly closer to *S. cerevisiae* than the sensu lato species we studied (Petersen et al. 1999). We favor *S. kluyveri* and *S. castellii* over *S. unisporus* for comparative sequence analysis for a few reasons. First, we obtained a few more BLASTX alignments for these species (Table 2), and *S. castellii* has a significantly greater number of BLASTN alignments (Table 1) that will help in locating promoter sequences. Second, we identified more snoRNA sequences by BLASTN comparisons from *S. kluyveri* and *S. castellii* than from *S. unisporus* (data not shown). Finally, these two species are estimated to have the smallest genomes within the *Saccharomyces* genus (Vaughan-Martini et al. 1993; Petersen et al. 1999) thus reducing the amount of sequence that would need to be determined.

METHODS

Sequencing Methods

The DNA sequence of random genomic clones from different *Saccharomyces* species was determined by the Washington University Genome Sequencing Center (WUGSC). A brief description of each step follows. Detailed protocols are provided in Mardis (1997), and at <http://genome.wustl.edu/gsc/Protocols>.

Table 5. Yeast Strains Used in this Study

Strains	Name	Other designation	Obtained from
<i>S. bayanus</i> *	623-6C		E. Louis ^a
<i>S. cariocanus</i> *	50791		E. Louis
<i>S. mikatae</i> *	1815		E. Louis
<i>S. paradoxus</i> *	N17		E. Louis
<i>S. bayanus</i>	NRRL Y-12624	(CBS 380)	C. Kurtzman ^b
<i>S. castellii</i> *	NRRL Y-12630	(CBS 4309)	C. Kurtzman
<i>S. dairenensis</i>	NRRL Y-12639	(CBS 421)	C. Kurtzman
<i>S. exiguus</i>	NRRL Y-12640	(CBS 379)	C. Kurtzman
<i>S. kluyveri</i> *	NRRL Y-12651	(CBS 3082)	C. Kurtzman
<i>S. paradoxus</i>	NRRL Y-17217	(CBS 432)	C. Kurtzman
<i>S. servazzii</i>	NRRL Y-12661	(CBS 4311)	C. Kurtzman
<i>S. unisporus</i> *	NRRL Y-1556	(CBS 398)	C. Kurtzman

An asterisk indicates species there were used for sequencing random genomic DNA.

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^bUSDA, Peoria, IL

Preparation of Genomic DNA

Derivatives of each yeast species lacking mitochondrial DNA were obtained by plating colonies on YPD agar media containing ethidium bromide (Slonimski et al. 1968) (except in the case of *S. kluyveri* which requires mitochondrial activity for viability). Genomic DNA was extracted from cells (50–100 mL cultures grown overnight to stationary phase in YPD). Cells were harvested and resuspended in 2 cell volumes of cold buffer (50 mM EDTA, 20 mM Tris at pH 8 containing 1 mg/mL ethidium bromide). Glass beads (0.5 mm, acid washed) were added to the cells and the suspension was vortexed for 5 min at 4°C. Sarkosyl (1% final concentration) was added and mixed by inversion. Cell debris was removed by centrifugation (2000 g for 10 min) and cesium chloride was added to the supernatant (1 g/mL). The mixture was centrifuged again (5000 g for 10 min) to remove additional debris. The supernatant was centrifuged at 100,000 G for 6 h in a Beckman TL-100 rotor.

Preparation of Clone Libraries

After quantification, the DNA was sheared by sonication, the ends repaired with mung bean nuclease, and fragments of 1–2 kb were selected by electrophoresis through a 0.8% agarose gel. The DNA fragments were excised and extracted from the gel, ligated to the plasmid (pBC) vector, and introduced by electroporation into competent *Escherichia coli* DH10B cells. DNA sequence of a representative sample of the resulting plasmid subclones was determined to assess library quality.

DNA Sequencing

Plating and sequencing of plasmid library subclones, as well as sample loading, data collection and processing was done as described by Mardis and Wilson, (1997) (description of methods also available at <http://genome.wustl.edu/gsc/Protocols/protocols.shtml>). *E. coli* colonies carrying plasmid subclones were picked and their plasmid DNA prepared by a simple method that lyses cells with microwaves (Marra et al. 1999). DNA sequencing used “big dye terminator” chemistry in cycle sequencing reactions. Cycle sequencing reactions were ethanol precipitated, resuspended in loading buffer and loaded onto an ABI 3700 capillary sequencing machine. Statistics were monitored constantly, and representative traces were inspected for each run. The data was processed by XGASP (Wendl et al. 1998), a script that directs vector clipping, quality analysis, data transfer, and initial assembly. The traces

were processed with PLAN, which performs PHRED basecalling (Ewing and Green 1998; Ewing et al. 1998).

BLAST Sequence Comparisons

Sequence alignments were initially generated using WU-BLAST 2.0 (<http://blast.wustl.edu>). *S. cerevisiae* sequence databases were obtained from SGD (<http://genome-www.stanford.edu/Saccharomyces>). The databases included *S. cerevisiae* genomic DNA (all_sacchdb.dna), *S. cerevisiae* protein-coding sequences and genes (orf.trans.fasta and orf_coding.fasta) and a library of intergenic DNA (NotFeature.fasta). This latter database consists of *S. cerevisiae* DNA from which was removed all genes (encoding both proteins and RNA), LTRs, and transposons. Another library was created by fusing the sequences of a 5' UTR library (consisting of the 500 bp upstream of the ATG start codon) to the genes sequences in orf_coding.fasta. BLAST output was parsed and processed with ad hoc PERL scripts. Percent identities of BLAST alignments were taken from the highest scoring alignment, to avoid biasing the calculations with low scoring and/or multiple alternative alignments of the same sequence segment.

TBLASTX comparisons used default parameters. Sensu lato and petite-negative sequences were compared to *S. cerevisiae* "not-feature" DNA sequences (those not known or predicted to encode proteins or functional RNAs) to identify potential protein-coding sequences. High scoring alignments were manually selected from all TBLASTX alignments. *S. cerevisiae* sequences with interesting alignments were visualized using AceDB (Eeckman and Durbin 1995) to look for potential new ORFs and extensions of known ORFs.

Multiple sequence alignments were created with CLUSTALW (Thompson et al. 1994). Some alignments were edited manually to investigate alternative equal scoring alignments. Local and global sequence alignments were generated using BESTFIT and GAP (GCG Corp.) with default parameters. FINDPATTERNS (GCG Corp.) was used to identify specific motifs within sequences.

Identification of Random Sequences Encoding Proteins Absent from *S. cerevisiae*

Random genomic sequences were compared with BLASTX to a nonredundant protein database similar to that maintained at NCBI. Because the database contains *S. cerevisiae* sequences, most of the identified homologs were from *S. cerevisiae*. In the sensu stricto species only five DNA sequences had significant hits to proteins not found in *S. cerevisiae*. The situation was more complex for the sensu lato and petite-negative sequences. In addition to finding genes not encoded in *S. cerevisiae*, we also found many sequences that had more similarity to sequences of species other than *S. cerevisiae*. Often the proteins with higher similarity are from closely related fungi, but occasionally they are from a distantly related organism, with the *S. cerevisiae* homolog showing much weaker similarity.

PCR Amplification and Southern Blotting of Species-Specific Sequences

Sequences were amplified by a standard PCR using the oligonucleotide primers listed below with genomic templates prepared as described by Hoffman and Winston (1987). PCR products were randomly labeled with ³²P (Ausubel et al. 1998) and used as probes for Southern blots of HindIII-digested genomic DNA from different species fractionated on a 1% agarose gel and transferred to a charged nylon filter.

S. cariocanus amidase
OM2186 AAACAACACAGCACCAGC
OM2187 TATCCTCAGACGCAGTCG

S. cariocanus unique sequences
OM2196 ATGGTGTGCGTTGTTATC
OM2197 TCAACATGCTGCATTTCG
OM2198 GCTAGTAGTTCGCGTGGTG
OM2199 CATCCCGACTCTGTCTTG

S. paradoxus unique sequence
OM2191 GGTACTTGGAGTTCTGTG
OM2192 TGGTGCTTTGGCAACAAG

S. mikatae unique sequence
OM2194 CACGTCCACATTAACCTG
OM2195 GATGTGGACATCATGCTTG

S. bayanus unique sequence
OM2206 CAATAACACGGATGCTCAAC
OM2207 TGTCGGAGGTCACAGGAG

Cloning of Specific Genomic Regions

Yeast genomic DNA was prepared as described by Hoffman and Winston (1987). The sequences of oligonucleotide primers were chosen based on conserved protein-coding regions that flank the sites of interest. Nested primers were used in a second reaction to amplify the *GAL1-10* and the *GAL4* promoters from some species. Initial PCRs were diluted 1/250 in TE and 1 μL was used as template in the second PCR. PCR products were cloned into a TA cloning vector (pGEMT-EASY [Promega] or pCRII [Invitrogen]). In some cases the PCR products were ligated directly into the vector; in other cases the PCR product was first purified by preparative agarose gel electrophoresis and extracted with QIAGEN gel purification columns. Plasmids were isolated using QIAGEN miniprep columns, checked by restriction digest for inserts of the correct size, and the DNA sequence of the insert was determined on an ABI 377 automated sequencer using universal or T7/SP6 primers and Big-Dye Terminator reactions (ABI). Sequence reads were processed with PHRED (Ewing and Green 1998; Ewing et al. 1998), PHRAP, and CONSED (Gordon et al. 1998). Sequences from *S. bayanus* and *S. paradoxus* were obtained from strains NRRL Y-12624 and NRRL Y-17217, respectively.

PCR Primers (asterisks indicate nested primers):

26s rDNA
OM1833 GCATATCAATAAGCGGAGGAAAAG
OM1834 GGTCCTGTTTCAAGACGG

GAL1-10
OM1975 AGCCGTGAGTTCAAACATCACC
OM1866* CCATGTATCCAGCACCACC
OM1867 AGTCACAATAATCAATATGTTCCACC

GAL4
OM1941 TTCCCACCAATAACATCATTTGACTGGAA
OM1920 TTCCACTTCTGTGAGATGTGCCCTAGTCAGCGG
OM1921* GACTCGAACAAAATCATTATTCTAGATATGAG
OM2208* GATAAACCAACATGTCATGGAGGC

S. paradoxus: OM1920-OM1941

S. bayanus: OM1920-OM1941 / OM1920-OM1921

S. kluyveri: OM1920-OM2208

HHT-HHF
OM1837 CACCAGTGGACTTTCTTGCTGTTTG
OM1838 CCACCTTACCTAGACCTTACCACCTTACC

SNR39
OM1959 GAAAAAATCTTGACCCCAAGTCTCAGTTGAAGA
OM1961 CTTGTTAATACCCTTGATTCTGACAACGAA

SNR44

OM1835 TCATCAAGTTTTTACAAGTTATGCAAAAAGC

OM1836 CGACAATCTTACCAGATCTGTGGTC

Estimation of Hexamer Frequency in Multiple Sequence Alignments

Random DNA sequences were generated in the computer with Seq-Gen (Rambaut and Grassly 1997). One thousand groups of four sequences (500 nucleotides long) were generated. After the first sequence was generated, two sequences having 70% identity and one having 40% identity to the first sequence were generated. The separate groups were aligned with CLUSTALW and the number of aligned n-mers ($n = 1-10$) were computed using an ad hoc PERL script.

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