where T_c decreases as the layer thickness of the nonsuperconducting PrBa₂Cu₃O₇ increases (21–23).

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- Complete Nucleotide Sequence of Saccharomyces cerevisiae Chromosome VIII

M. Johnston, S. Andrews, R. Brinkman, J. Cooper, H. Ding, J. Dover, Z. Du, A. Favello, L. Fulton, S. Gattung, C. Geisel, J. Kirsten, T. Kucaba, L. Hillier, M. Jier, L. Johnston, Y. Langston, P. Latreille, E. J. Louis,* C. Macri, E. Mardis,
S. Menezes, L. Mouser, M. Nhan, L. Rifkin, L. Riles, H. St. Peter, E. Trevaskis, K. Vaughan, D. Vignati, L. Wilcox, P. Wohldman, R. Waterston, R. Wilson, M. Vaudin

The complete nucleotide sequence of *Saccharomyces cerevisiae* chromosome VIII reveals that it contains 269 predicted or known genes (300 base pairs or larger). Fifty-nine of these genes (22 percent) were previously identified. Of the 210 novel genes, 65 are predicted to encode proteins that are similar to other proteins of known or predicted function. Sixteen genes appear to be relatively recently duplicated. On average, there is one gene approximately every 2 kilobases. Although the coding density and base composition across the chromosome are not uniform, no regular pattern of variation is apparent.

 ${f T}$ o identify all of the genes that constitute a simple eukaryotic cell, an international collaborative effort is under way to determine the sequence of the Saccharomyces cerevisiae genome. This is an important goal because of the central importance of yeast as a model organism for the study of functions basic to all eukaryotic cells. The sequences of the first two yeast chromosomes to be completed (1, 2) have revealed that more than two-thirds of yeast genes have not been previously recognized and are thus novel, and the functions of more than half of these cannot be predicted, because they are not similar to proteins of known function. Here, we describe the DNA sequence of yeast chromosome VIII, which provides another 210 previously unrecognized genes and further illuminates features of yeast chromosome organization.

The sequence was determined (3) from the set of 23 partially overlapping phage λ

and cosmid clones shown in Fig. 1 that were previously mapped by Riles *et al.* (4). The order of Hind III and Eco RI sites predicted from the sequence is consistent with the physical map of these sites determined independently by Riles *et al.* (4), which confirms that the sequence was assembled correctly. We estimate the accuracy of the sequence to be better than 99.99% (5). The genes and other features of the chromosome VIII sequence are listed in Table 1.

The sequence contains 269 nonoverlapping open reading frames (ORFs) greater than 300 base pairs (bp). On the basis of the analysis of Dujon *et al.* (2, 6), approximately 7% of these are likely to be false genes. Thirteen of these ORFs (4.8%) are predicted to be interrupted by introns at the extreme 5' end of each gene. The average gene size is 482 codons; the longest ORF (YHR099w) spans 11,235 bp (3745 codons).

Fifty-nine of the genes (22%) were previously identified (that is, already present in the public databases). Another 65 of the ORFs (24%) are predicted to encode pro X. Li, T. Kawai, S. Kawai, Jpn. J. Appl. Phys. 33, L18 (1994).

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Table 1. List of genes and features of chromosome VIII. The number of the cosmid (as submitted to GenBank) and its accession number are listed above the elements included in that database entry. Column 1: Nucleotide position of the start of each designated element (ATG for ORFs, or the first nucleotide of all other elements). For the 200 LTRs of the Ty elements, the beginning of the left LTR and the end of the right LTR is listed. Column 00 2: Genes are named according to established January convention: Y designates yeast; H designates chromosome VIII; L and R designate the left or right chromosomal arm, respectively; w and c designate that the gene is encoded on the top or bottom strand, respectively; and a superscript 5 "s" denotes genes predicted to be spliced. Genes are numbered from the CEN toward each TEL (telomere). Transfer RNA names also follow www.sciencemag convention: t designates tRNA; the next letter is the one-letter code for the amino acid inserted by the tRNA (abbreviations for the amino acid residues are A, Ala; F, Phe; H, His; P, Pro; Q, Gln; S, Ser; T, Thr; and V, Val.); the letters in parentheses are the codon recognized by the tRNA; and w and c designate that the tRNA is on the top (w) or bottom (c) strand. Retrotransposon LTRs in brackets are partial elements. Column 3: Genetic from names of genes previously identified. Note that one previously identified gene does not have a locus name (YHR042w) and that two genes (HX15/YHR096c and ACT5/YHR129c) were or named during the course of this work. **Column 4**: A description of the function of the genes. A de-scription of the protein most similar to the other genes is also listed. Genes with no listing in this column have no homologs (RLASTY course in column have no homologs (BLASTX score usually less than 70). Column 5: The BLASTX (18) score for the alignment of the encoded protein to its closest homolog. Note that BLASTX scores are not listed for previously identified genes, because the two sequences are identical. BLASTX scores greater than 100 are generally considered to indicate a significant relation between two proteins; scores between 70 and 100 are considered suggestive of a relation. Column 6: Database accession number of the closest homolog. In the few cases where comparison of predicted proteins to the BLOCKS database (19) revealed potential similarities not found by BLAST, the number of the BLOCKS entry is given.

teins that are similar to genes of known or predicted function (see Table 1 for a list). Thus, the function of only 46% of the encoded proteins is known or can be predicted (in some cases, only the biological process that the protein is involved in is

Genome Sequencing Center and Department of Genetics, Washington University School of Medicine, St. Louis, MO 63110, USA.

^{*}Yeast Genetics, Institute of Molecular Medicine, John Radcliffe Hospital, Oxford OX3 9DU, England.

Pos.	Gene or element	Locus	Function or homology	BLAST score	Acc. no.	Pos.	Gene or element	Locus	Function or homology	BLAST score	Acc. no.
1	TEL		9196/U11583 C(1-3)A	repeat	ense de S	127772 129473	YHR011w YHR012w ^s		Seryl-tRNA synthetase	369	gplX75627
36	Y' element		Y' subtelomeric repeat	Tepeat		131438	YHR013c	ARD1	Arrest-defective protein		spIP073471
3310	YHL050cs		Hyp. protein in Y' repeat region (pseudogene?)	1088	spIP240891	132038		SPO13	Meiosis-specific sporulation protein		spiP23624
1540	YHL049c		Hyp. protein in Y' repeat region (pseudogene?)	1371	pirlS312141	133099	tS(TCT)c	01010	tRNA-Ser		0011 200241
5051	X element		X subtelomeric repeat		P	133665	delta				
6400	YHL048w		YKL219w	653	spIP360341	134313	tQ(CAA)w		tRNA-GIn		
7993	Ty5 LTR				na da	134545	YHR015w		Poly(A)-binding protein	627	gpID26442
0211	YHL047c		YKR106w; YCL070c; YCL071c; YCL073c	1372	splP361731	138446	YHR016cs		SH3 domain in COOH-terminus	111	gplX59932
2283	YHL046c		Pau1p;YKL224c et al.; stress-induced proteins	583	gplL251231	138685	YHR017w				
2500	YHL045w		YCR103c; YKL223w	163	spIP256091	141393	YHR018c	ARG4	Arginosuccinate lyase		spIP04076
3563	YHL044w		YCR007c	130	spIP253541	gami, Hi			8082/U10399		
4899	YHL043w		YKL219w	179	spIP360341	143549	YHR019c		Filarial antigen (nematode); Asp-tRNA-synthetase	937	gplJ032661
5665	YHL042w		YKL219w	178	spIP360341	143987	YHR020w		Multifunctional aminoacyl tRNA-synthetase	956	spIP28668
7390	YHL041w					146305	tA(GCT)c		tRNA-Ala		
0968	YHL040c		YKR106w	1456	gplZ282021	146322	sigma				
1780	YHL039w					148660	YHR021cs		40S ribosomal prot. S27; potential Zn finger	429	spIP359971
5506	YHL038c	CBP2	Cytochrome b pre-mRNA processing protein		gplK00138l	150336	YHR022c		RAS-related protein	68	gpIU02928
6177	YHL037c					151657	YHR023w	MYO1	Myosin		spIP089641
6239	YHL036w		Amino acid permease	151	gplL25068l	159183	YHR024c	MAS2	Mitochondrial processing peptidase		spIP11914
2754	YHL035c		Multidrug resistance protein (ABC transporter)	630	spIP360281	159429	YHR025w	THR1	Homoserine kinase		gpIM37692
4075	YHL034c	SSB1	Single-strand nucleic acid binding protein		spIP100801	160835	YHR026w	PPA1	Proteolipid protein of proton ATPase		spIP239681
6023	YHL033c	RPL4A	60S ribosomal protein L7A-1, same as MAK7		spIP17076l	164702	YHR027c	12213136	Stoll, or Mancherer, mark		2011 (A. 1997)
8506	YHL032c	GUT1	Glycerol kinase		spIP321901	167425	YHR028c	DAP2	Dipeptidyl aminopeptidase B		splP189621
9484	YHL031c					168552	YHR029c		Thymidylate synthase (putative)	112	gplX59273
0082	YHL030w								8179/U00062		
7966	YHL029c					170335	YHR030c	SLT2	Protein Ser-Thr kinase		gpIX59262
8761	YHL028w		Ser-Thr rich			172961	YHR031c		Pif1p (mito. DNA repair/recomb. prot.)	388	spIP07271
1109	YHL027w	RIM1	Pos. regulator of meiosis (Cys-His Zn fingers)		spIP334001	173335	YHR032w				
4023	YHL026c					175539	YHR033w		Pro1p (gamma-glutamyl kinase)	997	spIP322641
			9433/U11582			177990	YHR034c				
64848	YHL025w	SNF6	Transcription factor		spIP188881	178210	YHR035w		Sec23p (yeast protein transport protein)	90	spIP153031
56646	YHL024w		RNA binding proteins	90	splQ011301	180336	YHR036w				
\$2560	YHL023c					181968	YHR037w	PUT2	P5C dehydrogenase		gpIU00062
62752	tH(CUC)w		tRNA-His			184057	YHR038w				
64154	YHL022c	SPO11	Sporulation protein		spIP231791	186800	YHR039c		Aldehyde dehydrogenase	159	spIP17445
5855	YHL021c					187915	YHR040w		Hit1p, required for high-temperature growth	98	pirlS30869
67452	YHL020c	OPI1	Neg. regulator of phospholipid biosyn.		spIP219571	189855	YHR041c°	SRB2	Transcription factor		spIP34162
39544	YHL019c		Clathrin coat associated protein AP54	156	splQ00776l	190534	YHR042w		NADPH-cytochrome P-450 reductase		gplD13788
69704	YHL018w		Dimerization cofactor of NF1-a	85	spIP800951	193536	YHR043c				
70272	YHL017w		Probable transmembrane protein YKL039w	150	pirlS377391	194799	YHR044c				
4240	YHL016c	DUR3	Urea active transporter		spIP33413I	195542	YHR045w				
75408	YHL015w		S10P family of 40S ribosomal proteins	337	spIP23403I	198276	YHR046c		Inositol monophosphatase, QUTG protein	189	pirlS11944
77310	YHL014c		Glycogen phosphorylase; GTP-binding protein	60	spIP004891	201301	YHR047c	AAP1	Ala-Arg aminopeptidase (Zn metalloprotease)		gblL125421
78349	YHL013c					204598	YHR048w		Various drug resistance proteins	293	pirlJC1173
78931	YHL012w		UDP-glucose pyrophosphorylase	228	spIP088001	206453	YHR049w				
31611	YHL011c		Phosphoribosyl pyrophosphate synthetase	518	spIP11908I	207646	YHR050w		Smf1p (mitochrodrial membrane protein)	441	bbs1119299
3716	YHL010c					209697	YHR051w	COX6	Cytochrome c oxidase subunit VI		spIP00427
			L5018/U11581		1.01 CM	210840	YHR052w				
5055	YHL009c		bZIP DNA-binding protein	124	splP19880l	Impanyl	beelena	03557	8025/U00061		shame
5367	tV(GUU)c		tRNA-Val			212720	YHR053c	CUP1	Copper metallothionein		spIP07215
5383	[sigma]					214249	YHR054c		ORFX in CUP1 repeat region		
5534	tau					214718	YHR055c	CUP1	Copper metallothionein		splP07215
	Ty4					217681	YHR056c	01/17-1	ORFX' (extended) in CUP1 repeat region		
91755	tau					218844	YHR057c	CYP2	Peptidyl-prolyl cis-trans isomerase		spIP23285
1767	delta					219885	YHR058c				
2095	[delta]			a standard a		220109	YHR059w				
4505	YHL008c		Potential formate transporter NirC (E. coli)	62	spIP358391	220726	YHR060w				
7932	YHL007c	STE20	Protein Ser-Thr kinase, pheromone response		gblL04655l	222479	YHR061c				
8789	YHL006c					223759	YHR062c				
9214	YHL005c					225170	YHR063c			100	
Stopp.	AL AL		9780/U10555		199909200	227244	YHR064c		Hsp70 heat shock protein	432	spIP22202
9213	YHL004w	MRP4	Mitochondrial ribosomal protein		spIP329021	229164	YHR065c		RNA helicase (DEAD box)	562	spIP34580
1877	YHL003c		Hypothetical protein YKL008c	1549	spIP284961	229336	YHR066w				
2605	YHL002w		SH3 domain	151	spIP293541	230971	YHR067w				
4270	YHL001ws		Hypothetical protein YKL006w	677	splP361051	232134	YHR068w			075	ebiliozea
05579	CDEIII					234659	YHR069c		Hyp. protein upstream of abl (human)	275	gbIU07561
5000	CEN					234882	YHR070w		Of /O multic	74	an D04027
5689	CDEI			1500		237005	YHR071w		G1/S cyclin	74	spIP24867
6048	YHR001w		Hyp. prot. YKR003w; oxysterol-binding prot.	1596	splQ022011	237940	tF(TTC)1c ^e		tRNA-Phe		
1210	YHR002w		Mitochondrial carrier/Grave's disease prot.	192	gplX660351	237995	[delta]		9205/110556		
1310	YHR003c		Hypothetical protein YKL027w	344	gplZ280271	220000	VHD070	EPCT	9205/U10556		gplU04841
3087	YHR004c	GRAd	G protoin alpha subusit		en/D00500	239099	YHR072w	ENG/	Lanosterol synthase	170	spIP22059
4910	YHR005c	GPA1	G protein alpha subunit		spIP085391	242583	YHR073w		Oxysterol-binding protein	172 83	
6172	tT(ACT)c		tRNA-Thr			246194	YHR074w		Spore outgrowth factor B (B. subtilis)	83	splP08164
6745	delta		Za finanz protein (00110 track 01 d	507	00000171	249642	YHR075c				
7807	YHR006w	EDOIN	Zn finger protein (C2H2 type) Stp1p (yeast)	507	splQ009471	251102	YHR076w		Highly soldie COOH terminue		
21676	YHR007c	ERG11	Cyto. P-450 L1 (Lanosterol 14-a-demethylase)		spIP106141	255650	YHR077c		Highly acidic COOH-terminus		
2500	VLIDOGG	6000	L2825/U10400		0010004471	256361	YHR078w	IRE1	Protoin kinaso		epiDoooct
23583	YHR008c	SOD2	Superoxide dismutase		spIP004471	261571	YHR079c	IRE1	Protein kinase		spIP32361
5658	YHR009c		Difference in the LOT	101		266839	YHR080c				
6513	YHR010w ^s		Ribosomal protein L27	424	pirlS004011	267539	YHR081w				

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Pos.	Gene or element	Locus	Function or homology	BLAST score	Acc. no.	Pos.	Gene or element	Locus	Function or homology	BLAST score	Acc. no.
71549	YHR082c	0.005	Protein Ser-Thr kinase	136	gplM204871	402966	YHR154w				
2628	YHR083w					407103	YHR155w		Sip3p (Snf1p interacting protein)	363	gpIU03376
4175	YHR084w	STE12	Transcriptional activator		spIP13574I	412406	YHR156c				
6765	YHR085w					412907	YHR157w	REC104	Meiotic recombination protein		spIP33323
			9332/U00060			417179	YHR158c				
8154	YHR086w	NAM8	RNA binding protein		gpIU000601	417549	YHR159w		Aminopophidase D.P. proline disortidase		DI 00401
30821 31496	YHR087w YHR088w					420072 422286	YHR160c YHR161c		Aminopeptidase P & proline dipeptidase		BL00491
33299	YHR088w YHR089c	GAR1	snRNP required for pre-rRNA processing		spIP280071	422200	THRIDIC		9986/U00027		
84626	YHR090c	GAHT	sinner required for pre-mine processing		Spir 200071	423072	YHR162w		Rat brain 0-44 mRNA, segment 2	221	gplM13095
B6771	YHR091c		Arginyl-tRNA synthetase	472	splP118751	423630	YHR163w		That brain of 44 minute, segment 2	661	gpiwrooso
88813	YHR092c	HXT4	Hexose transporter		spIP324671	429177	YHR164c		DNA-binding prot. for G-rich single strands	147	ap L14754
89144	YHR093w					436947	YHR165c	PRP8	U5 snRNP, pre-mRNA splicing factor		splP33334
92627	YHR094c	HXT1	Hexose transporter		spIP324651	439049	YHR166c	CDC23	Cell division cycle protein		spIP16522
92945	YHR095w					439341	YHR167w				
96449	YHR096c	HXT5	Hexose transporter	576	spIP324671	440376	YHR168w		GTP-binding proteins	214	spIP20964
98611	YHR097c ^s					442179	YHR169w		RNA helicase (DEAD box)	319	spIP34580
01936	YHR098c					443826	YHR170w				
02763	YHR099w					445710	YHR171w		Molybdopterin biosynthesis protein ChIN	141	spIP12282
			8263/U00059			448332	YHR172w				
14675	YHR100c					451150	YHR173c				
15970	YHR101c [®]					451324	YHR174w	ENO2	Enolase 2 (2-phosphoglycerate dehydratase)		pirlA01148
16574	YHR102w		Protein Ser-Thr kinase	325	splQ034971	452869	YHR175w		THE CONTRACTORS AND A STREET	13326-01	
20416	YHR103w			K. ALA		454226	YHR176w		Flavin-containing monooxygenase	97	gplL10037
23411	YHR104w		Aldo-keto reductase	495	splP318671	456589	YHR177w		7		and the second
24768	YHR105w		Bact. reg. prot. (helix-turn-helix, arsR group)		BL00846	459294	YHR178w	01/5-	Zinc finger (6-Cys) protein	95	spIP08657
05000	VUDIO		This sedenis reductors			462497	YHR179w	OYE2	NADPH oxidoreductase (Old Yellow enzyme)		splQ03558
25600	YHR106w	00010	Thioredoxin reductase	457	gplZ23109l	405470	YHR180w		9186/U00028		
28038	YHR107c YHR108w	CDC12	Cell division cycle protein		spIP324681	465173					
28305 30312	YHR108w YHR109w					466528 466906	delta				
32284	YHR110w		Glycoprotein 25L; involved in protein sorting?	149	splP278691	466985	[sigma) tT(ACA)w		tRNA-Thr		
33074	YHR111w		Molybdopterin biosynthesis protein moeB	313	spiP122821	467223	YHR181w				
35665	YHR112c		Cystathionine gamma-synthase	221	spiP009351	468214	YHR182w				
336339	YHR113w		Vacuolar aminopeptidase	249	splP149041	470955	YHR183w		6-phosphogluconate dehydrogenase	800	gplM80598
38085	YHR114w		SH3 domain	100	splP278701	472739	YHR184w		o priopriogradoriato derry drogenade	000	gpinicococ
41361	YHR115c					DAN LAN			9998/U00030		
341667	YHR116w					475335	YHR185c				
342351	YHR117w		Mito. protein import receptor; TPR repeats	616	splP072131	475782	tV(GTG)c		tRNA-Val		
345624	YHR118c					480619	YHR186c				
346045	YHR119w		Trithorax protein (COOH-terminus)	232	spIP206591	480985	YHR187w				
349576	YHR120w	MSH1	DNA mismatch repair protein		spIP258461	483808	YHR188c				
852758	YHR121w					484023	YHR189w				
			9315/U10398			484840	YHR190w	ERG9	Farnesyl-diphosphate farnesyltransferase		gblX59959
353627	YHR122w					486626	YHR191c				
354817	YHR123w ^s	EPT1	Ethanolaminephosphotransferase		splP221401	486821	YHR192w				
356563	YHR124w					488231	YHR193c				
358571	tF(TTC)2cs		tRNA-Phe			488652	YHR194w				
358698 358861	[delta] YHR125w					490742 491926	YHR195w YHR196w				
359081	[delta]					491920	YHR190W YHR197w				
60183	YHR126c		Tir2p (Cold shock-induced protein)	81	spIP338901	497275	YHR198c		YHR199c gene product	160	gplU00030
360915	YHR1200 YHR127w		(concentration proton)	01	op.1 000001	497275	YHR1980 YHR1990		YHR198c gene product	160	gp1000030
362012	YHR128w	FUR1	Uracil phosphoribosyltransferase		splP185621	499074	YHR200w		Serie product	100	gp1000000
64155	YHR129c	ACT5	Actin-related protein; centractin	564	gplZ14978l	501138	YHR201c	PPX1	Exopolyphosphatase		gplL28711
65302	YHR130c		states proton, contractin	004	90121-10101	502383	YHR202w				Shirroull
67864	YHR131c		Highly acidic COOH-terminus			505525	YHR203cs	RPS7A	Ribosomal protein S7		gpIM64293
69795	YHR132c		Carboxypeptidases	279	spIP150891	506314	YHR204w		Alpha-mannosidase	81	gpIU03458
71597	YHR133c								9177/U00029		0,000,00
71749	YHR134w					509361	YHR205w	SCH9	cAMP-dependent protein kinase		gplX57629
74310	YHR135c	YCK1	Casein kinase homolog I		splP232911	512727	YHR206w		Heat shock transcription factor	239	splP10961
75100	YHR136c				and the second second	516480	YHR207c				
75709	YHR137w					517527	YHR208w		Teratocarcinoma protein	475	spIP24288
77699	YHR138c					519432	YHR209w		Hyp. yeast prot. between DMC1-BMH1	158	gblL11229
79199	YHR139c	SPS100	Sporulation-specific wall maturation prot.		splP131301	521732	YHR210c		UDP-glucose-4-epimerase (GalE, Gal10p)	304	spIP04397
80575	YHR140w					525387	YHR211w		Flo1p (flocculation prot.; FLO8 gene?)	1075	spIP32768
82751	YHR141cs	RPL4B	60S ribosomal prot. L41, same as MAK18		gpID10578I	538089	YHR212c		RAA19 gene on chr. I right arm (identical)	555	gplL28920
			9666/U10397		92 80	539146	YHR213w		Flo1p (flocculation protein)	653	splP32768
83538	YHR142w		Car The risk			541646	YHR214w				
85510	YHR143w	0001	Ser-Thr rich		- IDC	543605	delta				
88726	YHR144c	DCD1	dCMP deaminase		spIP067731	Edoood	Ty1				
88995	tP(CCA)c ^s		tRNA-Pro; probable SUF8 gene (spans most of delta element)		Andream Station Station	549631	delta	DUCIO	Acid pheephotoco	0470	aniDoro
89337 89509	YHR145c delta		(spans most of delta element)			552094 554391	YHR215w YHR216w	PHO12	Acid phosphatase	2479	spIP35842
90300	YHR146w					556098	X element		IMP dehydrogenase (PUR5?) X subtelomeric repeat	1351	gplL22608
93283	YHR140W YHR147c	MRP-I 6	Mitochondrial ribosomal protein L6		splP329041	556640	Y' element		Y' subtelomeric repeat		
93534	YHR1470		40 <i>S</i> ribosomal protein YS11 (YP28)	136	spiP057551	557037	YHR217c				
96659	YHR149c			100	opii 007001	558009	YHR218w		Hyp. protein in Y' repeat region (pseudogene?)	1871	splP24089
97251	YHR150w					560168	YHR219w		Hyp. protein in Y' repeat region (pseudogene?)	3143	pirIS28368
00848	YHR151c					562451	TEL		TG(1-3) repeat		
	YHR152w	SPO12	Sporulation protein		splP171231				in an		
01434											

known). Nearly half of the ORFs (124, or 46%) are predicted to encode proteins that are not significantly similar to sequences in the public databases. Finally, 21 genes (7.8%) are predicted to encode proteins that are similar to proteins of unknown function. Only two of these (YHR069c and YHR162w) are similar to gene products of other organisms; most of the rest (13 of 19) lie very near the ends of the chromosome, where large segments are extensively duplicated in analogous regions of other yeast chromosomes.

Eleven transfer RNA (tRNA) genes were identified, three of which are interrupted by introns. Nine of these are preceded by complete or partial copies of the long terminal repeats (LTRs) of yeast retrotransposons (six with partial or complete δ elements, one with a σ element, and two with a partial σ element and a complete δ element), which reside 14 to 566 bp upstream of the tRNA genes. Except for the two $\boldsymbol{\delta}$ sequences that are part of the Tyl element on the right arm of the chromosome, all δ elements are associated with tRNA genes, as are the three complete or partial σ elements. The close association of these retrotransposon LTRs with tRNA genes is a general feature of the yeast genome (7). Four complete or partial τ sequences, two of

A Genetic map

which are associated with a Ty4 element on the left arm and one Ty5 LTR (8) were also identified.

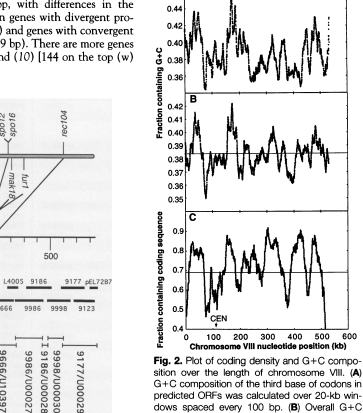
The CUP1 gene, encoding copper metallothionein, is contained in a 1998-bp repeated sequence that also includes an ORF of unknown function upstream of CUPI (YHR054c, previously called ORFX). The repeated region has been estimated to span 29.9 kb in the strain we used (4), which would encompass 15 repeats, but the number of repeats varies among yeast strains (9). We sequenced into the repeat region from each end and determined the sequence of one complete repeat. However, because the ORF upstream of CUP1 continues into unique sequence in the first copy of the repeat [the right, or centromere (CEN) distal copy], we included two copies of the repeat in the final sequence in order to include this novel ORF (YHR056c). Thus, the sequence includes two copies of the CUP1 gene (YHR053c and YHR055c).

The coding sequence comprises 69.2% of the chromosome, with one gene every 2087 bp. The average distance between genes is 629 bp, with differences in the spacing between genes with divergent promoters (731 bp) and genes with convergent terminators (479 bp). There are more genes on the top strand (10) [144 on the top (w)

cdc12

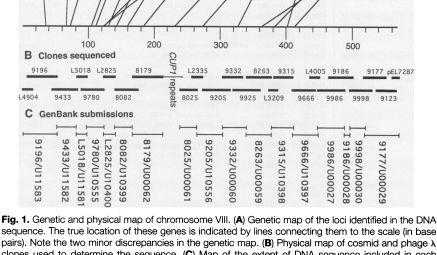
strand and 124 on the bottom (c) strand], but nearly all the excess w strand genes are accounted for by a stretch of approximately 35 kb where 17 of the 18 ORFs are arrayed on the top strand (coordinates 439341 to 474454). Disregarding this unusual cluster of genes, there are nearly equal numbers of genes on each strand. These properties of the sequence are similar to those found for the two yeast chromosomes previously sequenced (1, 2).

The base composition of the chromosome is clearly not uniform over its length (Fig. 2, A and B): there are two major G+C-rich peaks toward the left end of the chromosome and several minor peaks in the right half of the chromosome. On the basis of statistical analysis, we are confident that at least the two major G+C-rich peaks and the one major G+C-poor peak in the left ∞ half of the chromosome are significant (11). A similar degree of nonuniformity in base



0.46

G+C composition of the third base of codons in predicted ORFs was calculated over 20-kb windows spaced every 100 bp. (B) Overall G+C composition was calculated over 20-kb windows spaced every 100 bp. The horizontal line marks the average G+C composition (38.45%). (C) Coding density was calculated over 20-kb windows spaced every 100 bp. The horizontal line marks the average coding density (69.2%). For all three plots, similar results were obtained if the window size was varied between 10 and 50 kb or if the window size was the next 15 ORFs.



sequence. The true location of these genes is indicated by lines connecting them to the scale (in base pairs). Note the two minor discrepancies in the genetic map. (B) Physical map of cosmid and phage λ clones used to determine the sequence. (C) Map of the extent of DNA sequence included in each GenBank entry. The GenBank entry name and accession number are listed below each line. In addition, the entire (nonoverlapping) sequence (562,638 bp) is available via anonymous ftp (genome-ftp.stanford.edu in the /pub/yeast/genome_seq/chrVIII directory; ncbi.nlm.nih.gov in the /repository/yeast/CHVIII directory; mips.embnet.org in the /anonymous/yeast/chrviii directory).

500

600

composition was observed for chromosomes III and XI (2, 12). Although the regional variations in chromosome XI seem to occur in an almost regular pattern, those in chromosome VIII appear less regular. Thus, a regular periodicity of base composition does not appear to be a universal feature of yeast chromosomes. These base composition and gene density variations could be of functional importance (that is, having to do with processes such as replication or chromosome packaging) or could reflect the evolutionary history of the chromosome.

Similarly, the amount of protein coding sequence is not uniformly distributed over the length of chromosome VIII: there are six or seven regions of the chromosome with a coding density that is higher than average (Fig. 2C), a phenomenon also noted for chromosome XI (2). Perhaps not surprisingly, the G+C-rich regions correlate roughly, though certainly not precisely, with the regions of increased coding density, as was also noted for chromosome XI (2).

Several regions of chromosome VIII are duplicated on chromosomes I, III, or XI. The most extensive of these is an approximately 30-kb region very near the right telomere (bases 525393 to 555891) that is more than 90% identical to the similar region on the right arm of chromosome I. In addition, a smaller portion of this region of the right arms of chromosomes I and VIII is also duplicated on the left arm of chromosome I (13). This duplication, which was previously recognized (14), includes six genes whose order and orientation are preserved in the two copies. A Tyl element present in the duplicated region of chromosome VIII was probably originally present and subsequently lost from the homologous region of chromosome I, because chromosome I retains one of the LTRs of the retrotransposon at this location. A remarkable feature of this duplication is that its borders coincide almost precisely with the coding sequence (YHR211w at the left border and YHR216w at the right border). In addition, the high degree of sequence conservation between these regions of chromosomes I and VIII extends through a noncoding sequence, which suggests that this is a relatively recent duplication. Alternatively, the duplication could be more ancient, but extensive enough for the duplicated regions to pair infrequently in mitosis or meiosis and to be homogenized by gene conversion. A few other comparable duplications have been recognized on other yeast chromosomes (10, 15).

There are also several shorter duplicated segments of the subtelomeric region of the left arm of chromosome VIII at analogous positions of chromosomes III and XI. [This is in addition to the X and Y' subtelomeric

repeats, which are present at the ends of nearly all yeast chromosomes (7, 16).] These duplicated segments, which are scattered throughout the region between coordinates 5000 and 13000, vary in identity from about 54 to about 94% and are largely limited to four ORFs (YHL045 to YHL048).

Six other individual genes on chromosome VIII appear to be recently duplicated. This is clearly recognizable at the DNA level [BLASTN score cutoff of 300 (17)], in contrast to duplications of clearly older origin, which can be recognized only at the protein level. In each case, the duplicated sequences are confined to nearly the entire coding region of the duplicated gene. Four of the duplicated genes (YHL003c, YHL001w, YHR001w, and YHR003c) reside near the centromere, and three of the four homologs of these genes (YKL008c, 70% identical to YHR003c; YKL006w, 96% identical to YHL001w; and YKR003w, 72% identical to YHR001w) are also very near the centromere of chromosome XI [the other homolog is also on chromosome XI but is somewhat distant from the centromere, and the duplication is much less extensive and much less conserved (YKL027w, 57 to 63% identical to YHR003c over less than half the length of these genes)]. Two other duplicated genes (YHL047w and YHR021c) are dispersed on chromosome VIII, though homologs (YKL156w and YKL157w, respectively) are adjacent on chromosome XI. Thus, a total of 16 genes on chromosome VIII appear to be recently duplicated. In addition, another obvious case of less recent gene duplication on chromosome VIII is a cluster of three hexose transporter genes (YHR092c/HXT4, YHR094c/HXT1, and YHR096c/HXT5). The amount of redundancy recognized in the yeast genome will undoubtedly grow as the sequence of additional chromosomes becomes available.

We imagine two ways these duplications could have arisen. First, some of these genes could represent processed genes that were inserted into the genome relatively recently, a view that is consistent with the conservation of sequence only in the coding regions. However, all of these cases would appear to be created by integration of fulllength complementary DNAs, because none appear to be pseudogenes and this is unexpected in this model. In addition, one of the homologous gene pairs includes introns in both genes (which are 63% identical; their exons are 96% identical), which suggests that at least these genes were not duplicated by this mechanism. Alternatively, the clustering of four of the duplicated genes near the centromeres of their respective chromosomes compels us to consider the idea that entire genomic regions were duplicated. This centromeric duplication would appear to be ancient, because the DNA sequence has clearly diverged outside the coding regions, but the high degree of DNA sequence conservation in the coding region would appear to be at odds with this view.

Analysis of the sequence of chromosome VIII corroborates our current view of the organization of yeast chromosomes. The high coding density and close spacing of genes on chromosome VIII is similar to that of the other two yeast chromosomes sequenced, and the degree of genetic redundancy is also similar. However, the apparent organization of chromosome XI into regularly spaced intervals of G+C-rich and G+C-poor segments does not appear to hold for chromosome VIII, making the generality of this phenomenon unlikely. The most immediate and wide-ranging impact of this work is likely to be the identification of ∞ the 210 novel genes found on chromosome VIII, most of which we are unable to predict a function for at the present time. The $\underline{\mathfrak{O}}$ sophisticated genetic techniques available Jary for manipulating yeast cells provide the possibility of determining the function of many go of these genes. It seems certain that S. of these genes. It seems certain that S. cerevisiae will become even more important 5 for understanding the function of eukaryorg otic cells as the sequence of more chromofrom www.sciencemag somes is made available to the scientific community by the several groups collaborating internationally to complete the sequence of the entire yeast genome.

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- 3. The clones sequenced all originate from strain AB972, which is derived from the common laboratownloaded rv strain S288C (4). The sequence of the entire yeast DNA insert of each cosmid clone was determined. We sequenced the yeast DNA inserts in the phage λ clones after converting them into plasmids by recombination in yeast [J. Erickson and M. Johnston, Genetics 134, 151 (1993)]. Gaps that exist between two pairs of cosmid clones and between a cosmid clone and the left end of the CUP1 repeat were short enough to be recovered as polymerase chain reaction (PCR) products, using as a template the clones that span the gaps (λ 3209 and 4005 and cosmid 9181), which were then sequenced in their entirety. Finally, the sequence of the extreme right end of the chromosome, including the telomere, was determined mined from a plasmid clone generated by integration at the TG1-3 repeats of the telomere, followed by excision of the plasmid and capture of the flanking sequences (E. Louis, unpublished results). The details of the sequencing strategy have been described elsewhere [R. Wilson et al., Nature 368, 32 (1994)]. Briefly, 1- to 2-kb sheared fragments of the substrate DNA (cosmid, plasmid, or PCR product) were subcloned into M13 and sequenced on automated fluorescent DNA sequencing machines with universal primer. The sequence was assembled into contigs after 600 to 800 random subclones of each cosmid (fewer for the smaller λ clones and PCB products) had been sequenced (approximately sixto eightfold redundancy in the data). At this point, a directed sequencing strategy was used to join contigs, to sequence regions not represented on both strands, and to resolve discrepancies in the sequence. The sequence of both strands of each clone was determined (the sequence of overlapping re-

gions of cosmids was finished for only one clone), and all ambiguities in the sequence were resolved before the sequence of a clone was considered finished. The finished sequences were compared with the public sequence databases for protein and nucleic acid homologies [SWISSPROT (release 28.0), PIR (release 40.0), and GENPEPT (release 82.0)], with BLASTX (for protein similarities) and BLASTN (for nucleotide similarities) (18) and searched for tRNAs with TRNASCAN IG. Fichant and C. Burks, J. Mol. Biol. 220, 659 (1991)]. The sequence of each cosmid was also compared to the yeast sequences in GenBank, and discrepancies were examined in our sequence and corrected when possible (however, we judged that very few of these differences were due to mistakes in our sequence). The finished sequences were assembled and interactively annotated with AScDB, a version of the Caenorhabditis elegans database program ACeDB (R. Durbin and J.-T. Mieg, unpublished results) modified (by E. Sonhammer and R. Durbin and L. Hillier) for use with veast data. At this point, any potential frameshift errors were recognized, and the appropriate regions were resequenced to resolve the problems. Portions of the chromosome (usually individual cosmids) were submitted to GenBank, as shown in Fig. 1 (entry names and accession numbers are also listed in Table 1). Only a small number of overlapping bases were included in each database entry to facilitate joining of the sequences or to keep a gene intact. In addition, the entire (nonoverlapping) 562,638 bp of DNA that comprise chromosome VIII are available via anonymous file transfer protocol (ftp) (genome-ftp-.stanford.edu in the directory: /pub/yeast/genome_seq/chrVIII; ncbi.nlm.nih.gov in the directory: /repository/yeast/CHVIII). All ORFs containing at least 100 codons (including the ATG and translation termination codons) were identified. This analysis was done in batch with two scripts (ASCPREP1 and ASCPREP2; L. Hillier, unpublished results) that prepare the sequence and the database search results for entry into AScDB, which was used interactively to annotate the sequence. Genes were chosen with the help of the GENEFINDER program (P. Green and L. Hillier, unpublished results) modified (by L. Hillier, E. Sonhammer, and R. Durbin) for use with S. cerevisiae. All genes larger than 100 codons were annotated, except in the case of overlapping genes, where the longest gene or the gene that had homology to another gene was chosen. The first ATG codon in an ORF was always chosen as the beginning of the gene. Splice sites were used as necessary and when possible to construct a gene; a TACTAAC box 5 to 134 bases upstream of the 3' splice site [B. C. Rymond and M. Rosbash, in The Molecular and Cellular Biology of the Yeast Saccharomyces, E. Jones, J. Pringle, J. Broach, Eds. (Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1992), vol. 2, pp. 143–192] was demanded in each case. We sought delta (8), sigma (σ), and tau (τ) elements by comparing the sequence using BLASTN and FASTA against a representative member of each element.

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- 5. This is a conservative accuracy estimate based on our analysis of the yeast sequence as well as of the C. elegans sequence that has been determined in our sequencing center. We identified mistakes in the yeast sequence by comparing our sequence to sequences already in GenBank and by recognizing apparent frameshift errors. In 425 kb of yeast sequence checked in this way, 24 potential errors were identified (two by comparison to sequences in GenBank and 22 by recognition of apparent frameshifts)-approximately one error in 17 kb (most of these errors were corrected). An independent comparison of 17.208 bp of C. elegans sequence to an independently determined sequence already in GenBank revealed one error (L. Hillier, unpublished results), corroborating our estimate of approximately one mistake per 17 kb.
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Specific Cleavage of Model Recombination and Repair Intermediates by the Yeast Rad1-Rad10 DNA Endonuclease

A. Jane Bardwell,*† Lee Bardwell,*‡ Alan E. Tomkinson, Errol C. Friedbera§

The RAD1 and RAD10 genes of Saccharomyces cerevisiae are required for both nucleotide excision repair and certain mitotic recombination events. Here, model recombination 💬 and repair intermediates were used to show that Rad1-Rad10-mediated cleavage occurs Q at duplex-single-strand junctions. Moreover, cleavage occurs only on the strand containing the 3' single-stranded tail. Thus, both biochemical and genetic evidence indicate taining the 3' single-stranded tail. Thus, both biochemical and genetic evidence indicate a role for the Rad1-Rad10 complex in the cleavage of specific recombination intermediates. Furthermore, these data suggest that Rad1-Rad10 endonuclease incises DNA 5' to damaged bases during nucleotide excision repair.

The S. cerevisiae RAD1 and RAD10 genes are involved in both nucleotide excision repair (1) and mitotic recombination (2-9). RAD1 is the probable homolog of the human XPF (ERCC4) gene, which is defective in the cancer-prone disease xeroderma pigmentosum (10, 11); RAD10 is homologous to human ERCC1 (12). Rad1 and Rad10 proteins form a stable complex (13, 14) that catalyzes the endonucleolytic degradation of single-stranded bacteriophage DNA but is inactive on linear duplex DNA (15, 16). Here we demonstrate that rather than exhibiting a generalized single-strand DNA endonuclease activity as previously indicated (15, 16), Rad1-Rad10 protein is a

A. J. Bardwell, L. Bardwell, E. C. Friedberg, Laboratory of Molecular Pathology, The University of Texas Southwestern Medical Center at Dallas, Dallas, TX 75235, USA. A. E. Tomkinson, Institute for Biotechnology, Center for Molecular Medicine, The University of Texas Health Science Center at San Antonio, San Antonio, TX 78245, USA.

duplex-3' single-strand junction-specific endonuclease. The characterization of this structure-specific activity greatly clarifies \underline{O} the role of Rad1-Rad10 protein in recombination and DNA repair.

Single-stranded, duplex, or partial du-plex model DNA substrates were generated from synthetic oligonucleotides 18 to 50 nucleotides in length (Table 1). Rad1-Rad10 endonuclease did not degrade a single-stranded 49-nucleotide oligomer (S1 in Table 1 and Fig. 1, A and B) or a 49-base pair (bp) duplex structure (D in Table 1 and Fig. 2, A and B). However, when S1 was annealed to shorter complementary oligonucleotides, partial duplex molecules containing 3' single-stranded tails (TD1 and TD2 in Table 1) were cleaved by the enzyme (Fig. 1A), whereas substrate TD3 (Table 1) containing a 5' single-stranded tail was not (Fig. 1A). In a similar manner, substrate S3 (Table 1) was not cleaved as a singlestranded oligonucleotide (Fig. 2B), nor as a partial duplex derivative with a 5' singlestranded tail (TD4 in Table 1 and Fig. 1A). A partial duplex derivative with a 3' tail was cleaved (TD5 in Table 1 and Fig. 1A).

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Analyses with denaturing gels demon-

^{*}These authors contributed equally to this study. †Present address: Genelabs Technologies Inc., 505 Pe nobscot Drive, Redwood City, CA 94063, USA ‡Present address: Department of Molecular and Cell Biology, University of California at Berkeley, Berkeley, CA 94720, USA

[§]To whom correspondence should be addressed