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The yeast tcm1 gene, which codes for ribosomal protein L3, has been isolated by using recombinant DNA and genetic complementation. The DNA fragment carrying this gene has been subcloned and we have determined its DNA sequence. The 20 amino acid residues at the amino terminus as inferred from the nucleotide sequence agreed exactly with the amino acid sequence data. The amino acid composition of the encoded protein agreed with that determined for purified ribosomal protein L3. Codon usage in the tcm1 gene was strongly biased in the direction found for several other abundant Saccharomyces cerevisiae proteins. The tcm1 gene has no introns, which appears to be atypical of ribosomal protein structural genes.

Eucaryotic ribosomes consist of a complex of four RNA species and more than 70 proteins (39). The biosynthesis of these gene products in *Saccharomyces cerevisiae* is regulated in a remarkably coordinated manner (16, 21, 38), yet with one exception (12), no genetic linkage among them has yet been demonstrated (5, 12, 40). The mechanism for this coordinate regulation is not obvious. However, at least in the case of one ribosomal protein (r-protein) gene, it has been suggested that this regulation might be affected by autogenous feedback at the translational level (30). This has beem amply demonstrated in *Escherichia coli* (15, 23, 28).

Among those genes which affect protein synthesis in S. cerevisiae, there are several which have an easily identifiable genetic phenotype. These include three drug-resistant genes: tricodermin resistance (tcml; 18), cyclohexamide resistance (cyh2; 34), and cryptopleurine resistance (cry1; 17). The first two of these code for rproteins L3 and L29, respectively (13, 14). In addition, there are at least three genetic loci which cause omnipotent suppression, apparently as a result of RNA misreading by the ribosome: sup35 and sup45 (19) and SUP46 (25). It is quite possible that these also code for r-proteins. These six genes offer a twofold advantage. First, the phenotype can be used as a possible selection method to clone the structural gene. This approach has been used by Fried and Warner (13, 14) in their isolation of the genes responsible for tricodermin resistance (tcm1) and cycloheximide resistance (cyh2). Those studies established that the tcm1 and cyh2 genes code for rproteins L3 and L29, respectively (13, 14). Second, one might be able to make use of a selectable phenotype in the isolation of further mutants, for example, those which lie in sequences of regulatory importance.

We have chosen to work with the tcml gene. This codes for the largest r-protein, L3, of S. cerevisiae (13), and its biosynthesis appears to be quite strictly regulated (30). We present the nucleotide sequence of this gene and its environs as a basis for detailed genetic studies of its biosynthetic regulation.

## MATERIALS AND METHODS

Microbial strains and plasmids. The tricoderminresistant strain CLP1 (a leu2 tcml; obtained from J. Davies [18]) of S. cerevisiae was used as the source of DNA for constructing the clone library. S. cerevisiae strain LL20 (a leu2-3 leu2-112 his3-11 his3-15 can1; obtained from G. R. Fink) was the recipient for direct isolation of the tcml gene. E. coli strain JF1754 (hsdR lac gal metB leuB hisB) was used as the bacterial recipient. The cloning vehicle was pJZ1 (constructed by J. Zeissler; Fig. 1); this is a typical yeast shuttle vector expressing Leu2<sup>+</sup>, ampicillin resistance (Ap<sup>r</sup>), and tetracycline resistance (Tc<sup>r</sup>) (35). Growth and transformation of S. cerevisiae and E. coli were as previously described (27). In the isolation of the tcml gene, the S. cerevisiae genomic clone library carried on pJZ1 in E. coli JF1754 was transformed into S. cervisiae strain LL20, selecting for Leu2<sup>+</sup> on minimal medium plus histidine after embedding the spheroplasts in 3% agar. After 18 h at 23°C to allow expression of tricodermin resistance, the thin slab of agar containing the transformants was transferred to the surface of selection agar containing minimal medium

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FIG. 1. Method of isolation of the S. cerevisiae tcml gene. 2 µm, Yeast 2-µm circle. B, BamHI; H, HindIII, and E, EcoRI, sites for restriction endonuclease cleavage.

plus histidine and 5  $\mu$ g of tricodermin per ml. This is a convenient way to transfer spheroplasts from one growth condition to another. The drug concentration used was sufficient for complete growth inhibition of sensitive LL20 yeast cells (data not shown). After 2 days of incubation at 30°C, Leu<sup>+</sup> and tricoderminresistant (Tcm<sup>r</sup>) colonies were picked, purified, and retested. The proportion of Leu<sup>+</sup> colonies which were also Tcm<sup>r</sup> was 1 per 1,600.

DNA manipulations and nucleotide sequence analysis. Enzymes and reagents were purchased from Boehringer-Mannheim Canada, Bethesda Research Laboratories, New England Nuclear, and New England Biolabs. Enzyme reactions were carried out according to the procedures supplied by the manufacturers. DNA manipulations were by standard methods (24). DNA sequencing was carried out according to the method of Maxam and Gilbert (26). Fine-structure mapping of restriction sites was carried out by the method of Smith and Birnstiel (33) with DNA fragments labeled at the 3' end with the Klenow fragment of DNA polymerase I and  $\alpha$ -<sup>32</sup>P-labeled deoxyribonucleotide triphosphates.

# **RESULTS AND DISCUSSION**

Cloning the tcm1 gene. The scheme for cloning the *tcml* gene (which is a mutant allele) made use of the fact that drug resistance is semidominant over sensitivity at low concentrations of tricodermin (2 to 5  $\mu$ g/ml) which completely inhibit sensitive strains (unpublished data). Thus, a partial diploid cell in which tricodermin resistance is carried on a medium-copy replicating plasmid but which bears a chromosomal drug-sensitive allele might be expected to show elevated drug resistance. A clone library in E. coli from DNA fragments of a total BamHI digest of DNA from the tricodermin-resistant S. cerevisiae strain CLP1 was prepared with pJZ1 as vector (Fig. 1). Transformants of strain LL20 were initially selected for the Leu<sup>+</sup> phenotype. After an expression time of 18 h at 23°C, the thin top agar slab containing the transformants was transferred to a selection agar containing minimal medium plus histidine plus 5  $\mu$ g of tricodermin per ml to select directly for Leu<sup>+</sup> Tcm<sup>r</sup> transformants.

Yeast plasmid DNA was prepared from several Leu<sup>+</sup> Tcm<sup>r</sup> transformants and used to transform *E. coli* strain JF1754 to ampicillin resistance. Plasmid pLS2-II was isolated from one such *E. coli* transformant; this carried a 13.5kilobase *Bam*HI yeast DNA insert containing the *tcm1* gene. This plasmid DNA retransformed *S. cerevisiae* strain LL20 to both Tcm<sup>r</sup> and Leu<sup>+</sup> at a high frequency.

Restriction map and nucleotide sequence of the *tcm1* gene. The restriction map of the insert carried on pLS2-II was determined by conventional means (24) (Fig. 2a). This restriction map

agrees closely with the map obtained by Fried and Warner (13) for the BamHI fragment carrying the tcml gene. It has been shown by Fried and Warner (13) that the *tcml* gene is present on the 3.4-kilobase BamHI-AvaI region (heavy line in Fig. 2a) of the 13.5-kilobase BamHI fragment. This BamHI-AvaI fragment contains the information necessary to transform sensitive yeast cells to tricodermin resistance (13; our unpublished data). To facilitate further analysis, the BamHI-Aval fragment was subcloned onto pBR325 (4), producing a plasmid called pLS5T. With pLS5T, a more detailed restriction map of the BamHI-Aval region was determined (Fig. 2b). Figure 2c shows the tactics used in determining the nucleotide sequence of the region



FIG. 2. (a) Restriction endonuclease map of DNA insert carrying the *tcm1* gene. (b) Detailed restriction endonuclease map of the *BamHI-AvaI* portion of the DNA shown in (a), which carries the *tcm1* gene. (c) Tactics for nucleotide sequence determination of the *tcm1* gene. Arrowheads indicate the direction and extent of sequence determination by the method of Maxam and Gilbert (26). The predicted coding region for r-protein L3 (from Fig. 3) is shown by the boxed region between ATG and TAA. The heavy arrow indicates the inferred direction of transcription. bp, Base pairs.

NUCLEOTIDE SEQUENCE OF THE YEAST tcml GENE Vol. 155, 1983 -210 - 230 - 220 -200 - İ 90 -İ80 -i70 - İ 50 - 240 -160 -140 - 130 -120 -i10 -100 - 60 - 60 -80 - ŻO - 60 -40-- 30 - 20 - io 20 50 ò io żо 40 oArgLysArgAlaAlaSerIleArgAlaArgValLysAlaPheProLysAspAspArgSerLysProValAlaLeuThrSerPheLeuGlyTyrLysAla AAGAAAGAGAGCTGCCTCCATCAGAGCTAGAGTTAAGGCTTTTCCAAAGGATGACAGATCCAAGCCAGTTGCTCTAACTTCCTTGGGTTACAAGGCT 60 żo 80 90 Ìoo İ10 İ50 i20 i30 i40 G1yMetThrThrIleValArgAspLeuAspArgProG1ySerLysPheHisLysArgG1uValValG1uAlaValThrValValAspThrProProValV GGTATGACCACCATTGTCAGAGATTTGGACAGACCAGGTTCTAAGTTCCACAAGCGTGAAGTTGTCGAAGCTGTCACCGTTGTGACACTCCACCAGTTG 160 i70 180 190 200 210 220 230 240 250 a1Va1G1yVa1G1yTyrVa1G1uThrProArgG1yLeuArgSerLeuThrThrVa1TrpA1aG1uHisLeuSerAspG1uVa1LysArgArgPh TCGTTGTTGGTGTTGCGGTTACGTCGAAACCCCCAAGAGGTTTGAGATCTTTGACCACCGTCTGGGCTGAACATTTGTCTGACGAAGTCAAGAGAAGATT 260 . 270 . 280 . 290 300 310 320 330 340 350 360 370 380 400 410 420 **Å**30 390 440 450 470 500 520 530 540 460 480 490 510 550 1yG1ySerI1eSerG1uLysVa1AspTrpA1aArgG1uHisPheG1uLysThrVa1A1aVa1AspSerVa1TheG1uG1nAsnG1uMetI1eAspA1aI1 GTGGTTCCATCTCTGAAAAGGTTGACTGGGCTCGTGAACAATTCGAAAAGACTGTTGCTGTCGACAGCGTTTTTGAACAAAACGAAATGATTGACGCTAT 570 560 **580** 590 600 610 620 630 640 650 660 670 680 200 ÷10 720 730 240 250 κon IleGlyAlaCysHisProAlaHisValMetTrpSerValAlaArgAlaGlyGlnArgGlyTyrHisSerArgThrSerIleAsnHisLysIleTyrArgV ATTGGTGCTTGCCATCCAGCCCACGTTATGTGGAGTGTGCCAGAGCTGGTCAAAGAGGTTACCATTCCAGAACCTCCATTAACCACAAGATTTACAGAG . 760 770 . 780 . 790 . 800 810 830 \$20 Å₄∩ 850 alGlyLysGlyAspAspGluAlaAsnGlyAlaThrSerPheAspArgThrLysLysThrIleThrProMetGlyGlyPheValHisTyrGlyGluIleLy TCGGTAAGGGTGATGATGATGAAGCTAACGGTGCTACCAGCTTCGACAGAACCAAGAAGACTATTACCCCCAATGGGTGGTTTCGTCCACTACGGTGAAATTAA 860 940 950 870 880 890 900 910 920 **9**30 sAsnAspPheIleMetValLysGlyCysIleProGlyAsnArgLysArgIleValThrLeuArgLysSerLeuTyrThrAsnThrSerArgLysAlaLeu GAACGACTTCATCATGGTTAAAGGTTGTATCCCAGGTAACAGAAAGAGAATTGTTACTTTGGAAAAGTCTTTGTACACCAACACTTCTAGAAAGGCTTTG . 80 960 970 900 1000 1010 i020 1030 i040 1050 GluGluValSerLeuLysTrpIleAspThrAlaSerLysPheGlyLysGlyArgPheGlnThrProAlaGluLysHisAlaPheMetGlyThrLeuLysL GAAGAAGTCAGCTTGAAGTGGATTGACACTGCTTCTAAGTTCGGTAAGGGTAGGATTCCAAACCCCAGCTGAAAAGCATGCTTTCATGGGTACTTTGAAGA 1070 . 1080 1090 . 1100 . 1110 . 1120 . 1130 i140 . 1150 1060 ysAspleuter Aggactigtaagaagtittgttagaaaataaatcattitttaattgagcattcttattcctattttattaatagtittatgtattgttagctacatac 1190 i200 1210 i220 1230 1240 1250 . 1160 İ170 i180

AACAGTTTAAATCAAATTTTCTTTTTCCC

1260 1270 1280

FIG. 3. Nucleotide sequence of the S. cerevisiae tcml gene and flanking regions.

believed to encode the *tcm1* gene (13); over 95% of the sequence was determined for both DNA strands. The sequence itself is shown in Fig. 3.

Features of the nucleotide sequence of the tcm1 gene. The amino acid sequence inferred from the nucleotide sequence yields a protein of 43.5 kilodaltons, a size similar to that reported by Fried and Warner (13) for r-protein L3. The predicted amino acid composition of this protein agrees with that determined by Itoh et al. (20) for the purified yeast r-protein L3. Furthermore, the sequence of 20 amino acid residues at the amino

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		L3	Ŷ			L3	Y			L3	Y			L3	Y
Phe	T TT TTC	2 13	5 63	Ser	TCT TCC	10 7	79 73	Tyr	TAT TAC	0 12	2 73	Cys	TGT TGC	2	20 2
Leu	TTA TTG	1 18	14 148		TCA TCG	0	1	och amb	TAA TAG	1 0		umb Trp	TGA TGG	0 6	26
Leu	CTT CTC CTA CTG	0 0 2 0	0 1 6 0	Pro	CCT CCC CCA CCG	0 0 14 0	6 1 71 0	His Gln	CAT CAC CAA CAG	7 10 7 0	3 56 44 0	Arg	CGT CGC CGA CGG	3 0 0 0	2 0 0 0
Ile Met	ATT ATC ATA ATG	11 7 0 7	58 66 1 39	Thr	ACT ACC ACA ACG	12 15 0 0	49 66 0 0	Asn Lys	AAT AAC AAA AAG	0 8 3 40	3 86 21 151	Ser Arg	AGT AGC AGA AGG	1 3 30 0	1 1 63 1
Val	GTT GTC GTA GTG	19 16 0 0	112 93 3 1	Ala	GCT GCC GCA GCG	28 4 1 0	182 62 3 4	Asp Glu	GAT GAC GAA GAG	5 11 20 0	37 100 67 3	Gly	GGT GGC GGA GGG	31 0 0 0	164 11 1 1

FIG. 4. Codon usage for the S. cerevisiae tcml gene and several highly expressed S. cerevisiae genes. The values represent the actual number of times each codon is present in the structural gene. The numbers under the columns headed L3 are for the tcml gene, and those under the columns headed Y are summed for two glyceraldehyde-3-phosphate dehydrogenase genes, two enolase genes, alcohol dehydrogenase isozyme I, and alcohol dehydrogenase isozyme II. The data for these six are from Ammerer et al. (1).

terminus as determined by direct analysis of purified r-protein L3 (E. Otaka, personal communication) agrees exactly with that inferred from the nucleotide sequence. Finally, there is no evidence for a noncoding intervening sequence in the tcml gene. Our sequencing results substantiate the results of Fried and Warner (13), who showed on the basis of translation of hybrid-selected mRNA in vitro that the tcml gene codes for r-protein L3. Schwindinger and Warner (personal communication) have shown by S1 mapping that the *tcm1* gene is a rarity among r-protein genes examined thus far in that it does not have an intervening sequence (6, 12); this supports the conclusion drawn from our sequence data.

The data in Fig. 4 show the frequency of codon usage for the S. cerevisiae tcml gene. A summation of the total codon usage for six other yeast genes, two for glyceraldehyde-3-phosphate dehydrogenase, two for enolase, alcohol dehydrogenase isozyme I, and alcohol dehydrogenase isozyme II, is also shown (1, 3). These results clearly show that there is a pronounced bias in the choice of amino acid codons in the tcml gene, similar to the bias in codon usage seen for a number of highly expressed yeast genes (1, 3). These data also show that rprotein L3 is very rich in the basic amino acids lysine and arginine, as is to be expected for an rprotein.

Other features of the DNA sequence include the following. (i) A TATA box, in this case the sequence TATATAAAA, is located 113 nucleotides upstream from the translation start of the tcm1 gene. This presumably represents an essential feature of the promoter in eucaryotic cells (8). Although we have not determined the transcription start site, by analogy with several other yeast genes (2, 10, 11, 36) one might predict this to lie about 40 to 70 nucleotides on the 3' side of the TATA box. If this is so, then the untranslated leader of the mRNA would be about 40 to 70 nucleotides long. It is this stretch of RNA that might be implicated in autogenous translational regulation or r-protein gene expression in S. cerevisiae (30). The 5'-leader region is very adenine-thymine rich and does not appear to allow for a great deal of secondary structure. This is in contrast to several E. coli r-protein leaders with which it has been suggested that the secondary structure is important for binding the autogenous regulator (7, 28, 29).

(ii) Zaret and Sherman (41) observed that the 3' untranslated regions of many S. cerevisiae genes show sequence similarities in a region that is postulated to be involved in transcription termination. Based on a comparison of the 3' untranslated regions from 15 yeast genes, Zaret and Sherman (41) have proposed the following general sequence as a possible signal for transcription termination: . . . (T rich) . . . TAG . . . TAGT . . . . (AT rich) . . . TTT. The *tcm1* gene sequence shows similarities to this general sequence in the region from +1,217 to +1,262(Fig. 3); in particular, there is a TAG sequence at +1,228 and a TATGT sequence at +1,234. The 3'-flanking region of the tcml gene is also very adenine-thymine rich. There are several AATAA sequences, which for other genes have been suggested as possible polyadenylic acid addition sites (31).

(iii) A comparison of the nucleotide sequence of *tcml* in the region of the ATG start codon (CTACAACAATCAATCATGTCTCAC) with the corresponding region of 11 other *S. cerevisiae* genes (1) shows several similarities. (a) There is a scarcity of G residues in the 15 nucleotides of the plus-strand sequence just preceding the ATG. Instead, this region is rich in sequences such as CA, CAA, and CAAA. (b) There appears to be an invariant A residue at position -3. (c) There is a prevalent T residue at position +6.

(iv) Perhaps most striking, the tcm1 gene has no intron, in contrast to most other yeast rprotein structural genes so far studied (6, 13, 22, 32). Thus, one can conclude that coordinate regulation of r-protein biosynthesis does not depend directly on intron processing. Despite this, r-protein L3 biosynthesis appears to be affected by the *rna2* mutation (16, 37), which is involved with the processing of introns from precursor mRNA (9, 32). This raises the question of whether there is a mechanism of r-protein regulation in which the expression of non-introncontaining r-protein genes is dependent on the expression of other r-protein genes that do contain introns.

In total, five r-protein genes from S. cerevisiae have now been sequenced (22; M. Rosbash, J. Warner, and J. Woolford, personal communication). A detailed structural comparison of all five genes will be presented in a joint publication (manuscript in preparation).

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