Journal of Bacteriology

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M Kuenzler, T Balmelli, C M Egli, G Paravicini and G H Braus *J. Bacteriol.* 1993, 175(17):5548.

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Cloning, Primary Structure, and Regulation of the HIS7 Gene Encoding a Bifunctional Glutamine Amidotransferase:Cyclase from Saccharomyces cerevisiae

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Received 12 April 1993/Accepted 30 June 1993

The Saccharomyces cerevisiae HIS7 gene was cloned by its location immediately downstream of the previously isolated and characterized ARO4 gene. The two genes have the same orientation with a distance of only 416 bp between the two open reading frames. The yeast HIS7 gene represents the first isolated eukaryotic gene encoding the enzymatic activities which catalyze the fifth and sixth step in histidine biosynthesis. The open reading frame of the HIS7 gene has a length of 1,656 bp resulting in a gene product of 552 amino acids with a calculated molecular weight of 61,082. Two findings implicate a bifunctional nature of the HIS7 gene product. First, the N-terminal and C-terminal segments of the deduced HIS7 amino acid sequence show significant homology to prokaryotic monofunctional glutamine amidotransferases and cyclases, respectively, involved in histidine biosynthesis. Second, the yeast HIS7 gene is able to suppress His auxotrophy of corresponding Escherichia coli hisH and hisF mutants. HIS7 gene expression is regulated by the general control system of amino acid biosynthesis. GCN4-dependent and GCN4-independent (basal) transcription use different initiator elements in the HIS7 promoter.

Histidine is synthesized in an invariable series of 11 enzymatic reactions from ATP and phosphoribosyl-pyrophosphate (PRPP) in all histidine-autotrophic organisms studied so far (Fig. 1A). Enzymatic regulation of the unbranched pathway is achieved by feedback inhibition of the first step of the pathway by its final product, histidine, in both prokaryotic and eukaryotic microorganisms studied (9, 26). This biochemical invariability faces a considerable diversity in organization, structure, and regulation of the genes coding for the various enzymatic activities even within a biological kingdom.

In the best-studied organisms, Escherichia coli and Salmonella typhimurium, the 11 enzymatic activities are encoded by eight genes organized in a single operon [hisGD CBHAF(IE)] (Fig. 1) (11, 54). Three of the eight genes code for bifunctional enzymes [hisD, hisB, and his(IE)]. This situation, however, is not typical of prokaryotes and not even of eubacteria. In Lactococcus lactis, nine his genes exist, of which eight are clustered in an operon and one is located elsewhere on the chromosome (14). In Streptomyces coelicolor, eight genes map at three loci, one of them grouping six genes in an operon and the other two containing single genes (25, 30), whereas in Bacillus subtilis the eight genes are organized in two loci with seven genes and one gene, respectively (44). In the nitrogen-fixing eubacterium Azospirillum brasilense, at least four his genes are clustered (5, 17), and in Staphylococcus aureus, six his genes are clustered (42). In some of these organisms, not only the organization but also the structure of individual genes differs from the situation in enteric eubacteria. The two enzymatic activities encoded by the hisB gene in E. coli and S. typhimurium reside on separate genes in L. lactis, S. coelicolor, and A. brasilense. In addition, in S. coelicolor the his(IE) activities are separated as well. Physically separated his(IE) activities are also found in the methanogenic archae-bacterium Methanococcus vannielii (6).

In fungi, the genes encoding the enzymatic activities of various biosynthetic pathways are scattered throughout the genome. In S. cerevisiae, the genetic information for the histidine biosynthetic enzymes is encoded by seven genes, which are located on six different chromosomes (HIS1-7) (Fig. 1) (10). The structure of the genes differs from that of enterobacteria in that the enzymatic activities of hisD and his(IE) are combined to a multifunctional enzyme catalyzing four steps in the histidine biosynthetic pathway encoded by the HIS4 gene (16). Such a multifunctional enzyme exists as well in Neurospora crassa (29) and Candida albicans (2). In addition, as in L. lactis, A. brasilense, and S. coelicolor the hisB activities of E. coli and S. typhimurium are encoded by two independent genes in S. cerevisiae (HIS2 and HIS3) (45, 50). The only plant gene cloned so far is a cDNA from the cabbage Brassica oleracea corresponding to the hisD gene of enterobacteria and encoding a bifunctional histidinol dehydrogenase (36), indicating that the organization and structure of the genes involved in histidine biosynthesis are also variable within the eukaryotic kingdom.

Coordinate regulation of the histidine-biosynthetic genes strongly depends on the gene organization in the corresponding organism. In *E. coli* and *S. typhimurium*, where all *his* genes are clustered in a single operon, coregulation is achieved by attenuation control and positive metabolic regulation of the operon (54). In *S. cerevisiae*, the scattered genes are part of a complex regulatory network which couples the transcriptional derepression of at least 30 structural genes involved in multiple-amino-acid biosynthetic pathways under environmental conditions of amino acid starvation (23). The final step in this general control system is the binding of the transcriptional activator protein GCN4

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А	ATP + PRPP
	1 L ATP PR-transferase (EC 2.4.2.17)
	2 C PR-ATP pyrophosphohydrolase
	3 PR-AMP cyclohydrolase (EC 3.5.4.19)
	4 T PR-formimino-5-amino-1-PR-4-imidazole-
	5 J Glutamine amidotransferase
	6 🗸 Cyclase
	7 🗸 Imidazoleglycerol-phosphate dehydratase (EC 4.2.1.19)
	8 🕂 Histidinol-phosphate aminotransferase (EC 2.6.1.9)
	9 THistidinol-phosphate phosphatase (EC 3.1.3.15)
	11 Histidinol dehydrogenase (EC 1.1.1.23)
	Histidine

B E. coli: Thom 8 97 5 4 6 32 hisGDCBHAF(IE) (44.1 min)

S. cerevisiae:	🔢 <i>HIS1</i> (chr.V)
	In <i>HIS2</i> (chr.VI)
	🔽 <i>HIS3</i> (chr. XV)
	3/2/10/11 HIS4 (chr. III)
	⑧-≁≁- ₄ <i>HIS5-HIS6</i> (chr. IX)
	5/6 <i>HIS7</i> (chr. II)

FIG. 1. Gene-enzyme relationships in histidine biosynthesis in *E. coli* and *S. cerevisiae*. (A) Schematic representation of the biosynthetic steps from ATP and PRPP to histidine with enzyme designations and assigned Enzyme Commission (EC) numbers. (B) Organization of the genetic information for the histidine biosynthetic enzymes in *E. coli* and *S. cerevisiae*. Genes are represented by boxes which are shaded for genes that are cloned and sequenced. Encoded enzymatic activities are indicated by numbers referring to panel A. Multiple numbers separated by slashes symbolize multifunctional enzymes. The chromosomal locations are given as map positions for *E. coli* (min) and as chromosome numbers for *S. cerevisiae* (chr.). The *S. cerevisiae HIS5* and *HIS6* genes are located on the same arm of chromosome IX.

to the promoters of the target genes resulting in elevated transcription of these genes.

In this paper, we describe the cloning and characterization of the HIS7 gene, which codes for the enzyme catalyzing the fifth and sixth step of the histidine biosynthetic pathway in *S. cerevisiae*. The derived amino acid sequence and heterologous complementation of corresponding *E. coli* mutants suggests that the HIS7 gene product is a bifunctional enzyme with an N-terminal glutamine amidotransferase and a C-terminal cyclase domain. Furthermore, we find that transcription of the HIS7 gene is regulated by the general control system of amino acid biosynthesis and that the start sites for GCN4-dependent and GCN4-independent (basal) transcription are different.

MATERIALS AND METHODS

Strains and culture conditions. All strains used in this study are listed in Table 1. Yeast strains were all derivatives of the S. cerevisiae laboratory strain S288C ($MAT\alpha$ gal2 SUC2 mal CUP1).

Cultivation of S. cerevisiae was performed at 30° C in either YEPD complete medium (46) or MV minimal medium (33). Appropriate supplements were added to the medium in recommended amounts (46). LB complete medium and M9 minimal medium for E. coli are described by Sambrook et al.

(48). LB medium containing ampicillin (50 mg/liter) was used to select for transformants. *E. coli* was cultivated at 37°C.

Crossing of S. cerevisiae. Crossing of compatible yeast strains was performed as described previously (46). Selection for diploids after mating was done on MV minimal medium.

DNA techniques and sequencing. Enzymatic manipulation and cloning of DNA were performed as described by Sambrook et al. (48). *E. coli* MC1061 (12) was used for the propagation of plasmid DNA. DNA sequences were determined for both strands by the chain termination method (49) and the M13 subcloning technique (31). Oligonucleotide primers were purchased from Microsynth (Windisch, Switzerland). The M13 host JM101 (31) and the M13-based vectors M13mp18 and M13mp19 (56) were used for the isolation of single-stranded template DNA.

PCR. The polymerase chain reaction (PCR) technique for the amplification of cloned DNA fragments by using sequence-specific oligonucleotides was described previously (47). In this work, the technique was exploited for the production of a DNA fragment used for S1 nuclease mapping of the *HIS7* mRNA 3' end. As standard reactions using Super *Taq* polymerase (P. H. Stehelin & Cie AG, Basal, Switzerland), 30 cycles of 1 min at 94°C, 1 min at 50°C, and 1 min at 72°C were performed in a Biometra Trioblock thermocycler (Biometra, Göttingen, Germany).

Yeast genomic DNA library. The yeast genomic DNA library contains DNA of strain YPH1 (MATa ura3-52 bys2-801 ade2-101 GAL⁺ SUC⁺) partially digested with Sau3AI in a YCp50 derivative in which the yeast URA3 gene is replaced by the yeast LEU2 gene.

Construction of the $\Delta(aro4-his7)$::URA3 disruption strain. S. cerevisiae RH1447 carrying a disrupted ARO4-HIS7 locus was constructed as follows. In the course of subcloning of the ARO4 gene, a chromosomal 3.5-kb XbaI-BamHI fragment ranging from an XbaI site located approximately 0.9 kb upstream of the 5' end of the region shown in Fig. 2A to the indicated BamHI site was cloned into pGEM7Zf(+) (Promega, Madison, Wis.), yielding plasmid pME638. From this plasmid, a 2.4-kb AccI fragment comprising the complete ARO4 gene and the 5' end of the HIS7 gene (see Fig. 2A) was isolated and replaced by the chromosomal 1.1-kb URA3 fragment in the same orientation as the substituted genes, resulting in plasmid pME642. Transformation of S. cerevisiae RH1377 with the 2.2-kb XbaI-BamHI fragment from plasmid pME642 and selection for a Ura⁺ phenotype in the presence of supplementing amounts of histidine resulted in strain RH1447. The strain was examined for its His⁻ and concomitant Aro^- phenotypes in the presence of 5 mM phenylalanine and by Southern blot analysis.

Construction of strains with an integrated translational *HIS7-lacZ* **fusion.** The translational *HIS7-lacZ* fusion was constructed based on plasmid pNM482 (32). A 0.6-kb *HpaI-AccI* fragment containing the complete *ARO4-HIS7* intergenic region and the N-terminal 56 amino acids of the *HIS7* open reading frame (see Fig. 2A and 3) was inserted into pNM482 restricted with *SmaI* and *AccI*. From this plasmid the *HIS7-lacZ* fusion gene was isolated as a 4-kb *EcoRI-Csp451* fragment and inserted into vector pGEM7Zf(+) containing the 0.5-kb *HindIII-BamHI 3'* end of the yeast *ADH1* gene (7) to yield plasmid pME688. Therefore, this plasmid contains a 4.5-kb *EcoRI-BamHI HIS7-lacZ* translational fusion cassette.

For the construction of an integrative HIS7-lacZ fusion cassette, the 1.9-kb BamHI-HindIII HIS7 fragment (same fragment as BamHI fragment from pME692) was cloned as a

Species and strain or plasmid	Relevant characteristics	Reference or source
E. coli		
MC1061	Δ (lacIPOZYA)X74 galU galK StrA ^r hsdR Δ (ara-leu)	12
JM101	$\Delta(lac-pro)$ thi supE F'(traD36 proAB lacI ^q Z Δ M15)	31
W3110	Wild type	E. coli Genetic Stock Center
UTH6	λ^{-} his A 323	E. coli Genetic Stock Center
UTH860	ara-14 glnV44 galK2 λ^- rpsL 145 malT1(λ^r) xylA5 mtl-1 hisF860 λ^r	E. coli Genetic Stock Center
UTH1767	malA1(λ^{r}) xyl-5 mtl-1 rpsL145 hisH1767 λ^{-1}	E. coli Genetic Stock Center
S. cerevisiae		
RH1377	ΜΑΤα Δυτα3	ETH collection ^b
RH1447	MATα Δura3 Δ(aro4-his7)::URA3	This work
C20-2C	MATa his7 ade2 ade4 ura1 lys2 tyr1 arg4 leu1 trp5 gal	Yeast Genetic Stock Center ^c
RH1631	MATa ura3-52	ETH collection
RH1632	MATa ura3-52 gcd2-1	ETH collection
F194	MATa ura3-52 gal2 gcn4-103	22
RH1371	MATa aro3-2 Lura3 gcd2-1	ETH collection
RH1372	MATa aro3-2 Δura3	ETH collection
RH1381	MATa aro3-2 ura3-52 gcn4-101	ETH collection
RH1614	MATa aro3-2 Δura3 Δhis7::lacZ gcd2-1	This work
RH1615	MATa aro3-2 Δura3 Δhis7::lacZ	This work
RH1616	MATα aro3-2 ura3-52 Δhis7::lacZ gcn4-101	This work
Plasmids		
pME638	pGEM7Zf(+) ^d containing a 3.5-kb XbaI-BamHI ARO4-HIS7 fragment	ETH collection
pME642	pME638 with a $\Delta(aro4-his7)$::URA3 disruption	This work
pME688	pGEM7Zf(+) containing a 4.5-kb EcoRI-BamHI HIS7-lacZ cassette	This work
pME692	pGEM7Zf(+) containing a 1.9-kb BamHI-HindIII HIS7 fragment	This work
pME693	pME688 with a 1.9-kb BamHI HIS7 fragment from pME692	This work
pME694	pGEM7Zf(+) containing a 1.9-kb SphI-BamHI ARO4-HIS7 fragment	This work
pME696	pME694 with a 6.1-kb BsmI-Nsil Δhis7::lacZ fragment from pME693	This work
pME979	pGEM7Zf(+) containing a 2.4-kb EcoRV-HindIII HIS7 fragment	This work

TABLE 1. Strains and plasmids

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homologous downstream region behind the *ADH1* 3' end region in pME688, resulting in plasmid pME693. To ensure proper integrative recombination of the fusion, its homologous upstream region was subsequently enlarged to the *SphI* site by ligating a 6.1-kb *BsmI-NsiI* fragment from the latter plasmid into plasmid pME694 containing the 1.9-kb *SphI*-*Bam*HI fragment of the *ARO4-HIS7* locus (see Fig. 2A) on pGEM7Zf(+) to yield plasmid pME696.

A two-step procedure was used for the integration of the translational $\hat{H}IS7$ -lacZ fusion at the original HIS7 locus on the yeast chromosome, resulting in a $\Delta his7::lacZ$ genotype. In the first step, the 1.1-kb KpnI-AccI fragment of the chromosome was replaced by the URA3 gene in strains RH1371, RH1372, and RH1381 as described for the construction of RH1447. The resulting strains had both His⁻ and an Aro⁻ phenotypes, with the latter due to an aro3-2 mutation. Transformation of the disruption strains with plasmid pME696 restricted with XbaI and selection for an Aro⁺ phenotype in the presence of supplementing amounts of histidine and uracil resulted in strains RH1614, RH1615, and RH1616 carrying a translational HIS7-lacZ fusion instead of the original HIS7 locus and an intact ARO4 gene. Strains were examined for Ura⁻ and His⁻ phenotypes and by Southern blot analysis.

Poly(A)⁺ RNA isolation. Yeast RNA enriched for polyadenylated RNA species was isolated as described previously (20). Isolated RNA was stored in 40% (vol/vol) isopropanol-120 mM sodium acetate at -20° C.

Primer extension analysis. The primer extension method to determine RNA 5' ends was performed as described by Kassavetis and Geiduschek (27). For the mapping of the *HIS7* transcript 5' ends, 50 μ g of poly(A)⁺ RNA of each strain was hybridized against an excess of a 5'-³²P-end-labelled 51-bp primer complementary to nucleotide positions +14 to +64 relative to the *HIS7* translational start site. Annealed primers were elongated with avian myeloblastosis virus reverse transcriptase. Elongation products were separated on a 6% polyacrylamide standard sequencing gel together with a T ladder generated by using the same primer as for the primer extension reactions.

S1 nuclease mapping. The S1 nuclease protection method for mapping RNA 5' and 3' ends was performed as described by Furter et al. (20). For *HIS7* mRNA 5' end mappings, 30 μ g of poly(A)⁺ RNA of each strain was hybridized against an excess of a *KpnI-Bam*HI fragment (ranging from nucleotide positions -769 to +415) which was ³²P labelled at the 3' end of the antisense strand. The resulting hybrid molecules were digested with S1 nuclease, and the protected DNA strands were separated on a 6% polyacrylamide standard sequencing gel. As a size standard, ³²P-labelled pBR322 plasmid DNA restricted with *HpaII* was used. Mapping of the *HIS7* transcript 3' ends employed a fragment ranging from nucleotide positions +1503 to +2145 and ³²P labelled at the 5' end of the antisense strand. This fragment was generated by a standard PCR reaction using two primers, ranging from positions +1467 to +1486 (20 bp) and complementary to positions +2116 to +2145 (30 bp) respectively, followed by subsequent cleavage with *NarI* and filling in of the 5' protruding end with $[\alpha^{-3^2}P]dCTP$. Nuclease S1 digestion products were separated on a 6% polyacrylamide standard sequencing gel together with a sequence ladder generated by using the PCR primer complementary to nucleotide positions +2116 to +2145. Torula yeast RNA (30 µg) was used as a negative control.

Northern (RNA) analysis. Poly(A)⁺ RNA (30 μ g) of each strain was separated on a formaldehyde agarose gel, electroblotted onto a nylon membrane, and hybridized against DNA fragments which were ³²P labelled by using the oligolabelling technique described by Feinberg and Vogelstein (18). Probes used were made from a chromosomal 1.1-kb *HindIII URA3* fragment, a chromosomal 0.7-kb *HpaI ARO4* fragment, and a chromosomal 0.9-kb *BamHI-XbaI HIS7* fragment. The *URA3* transcript was chosen as an internal standard for the amount of RNA, as this gene is not regulated by the GCN4 protein.

β-Galactosidase assay. β-Galactosidase activities were determined by using permeabilized yeast cells and the fluorogenic substrate 4-methylumbelliferyl-β-D-galactoside (Fluka Chemie AG, Buchs, Switzerland). Yeast was cultivated in 5 ml of MV minimal medium supplemented with histidine and uracil to an optical density at 546 nm of between 1 and 4. Typically, cells from 0.5 ml of yeast culture were washed once with water and resuspended in 1 ml of reaction buffer (25 mM Tris-HCl [pH 7.5], 125 mM NaCl, 2 mM MgCl₂, 12 mM 2-mercaptoethanol). The cells were permeabilized by vortexing for 10 s after the addition of 50 μ l of CH₂Cl₂ and 0.1% (wt/vol) sodium dodecyl sulfate. Then, 40 µl of permeabilized cells was incubated with 160 µl of reaction buffer containing 0.3 mM 4-methylumbelliferyl-B-D-galactoside for 30 min at 37°C. The reaction was stopped after 30 min by adding 50 µl of 25% (wt/vol) trichloroacetic acid. The cells were spun down, and the fluorescence of the supernatant was determined in an at least 1/4 dilution in glycine/carbonate reagent (133 mM glycine, 83 mM Na₂CO₃) with a Hoefer model TKO 100 fluorometer (Hoefer Scientific Instruments, San Francisco, Calif.). The concentration of product formed during the reaction was determined based on a standard curve in a range from 0 to 40 µM 4-methylumbelliferone (MUF) in reaction buffer. Product concentrations were normalized to the reaction time and the optical density of the culture. One unit of β -galactosidase activity is defined as 1 nmol of MUF h^{-1} ml⁻¹ optical density at 546 nm⁻¹. The given values are means of at least three independent cultures. The standard error of the mean was less than 25%.

Sequence data analysis. Sequence data were analyzed with the Genetics Computer Group Sequence Analysis Software (15). Multisequence alignments were produced by the program PILEUP, and pairwise alignments and identity or similarity value calculations were done by using the program GAP.

Nucleotide sequence accession numbers. The nucleotide sequence presented in this paper has been assigned Gen-Bank/EMBL accession numbers X61107 (ARO4) and X69815 (HIS7).

RESULTS

Cloning and sequencing of the HIS7 gene. The ARO4 gene of S. cerevisiae, encoding the tyrosine-inhibitable desoxy-arabino-heptulosonate-phosphate synthase (DAHPS), was previously isolated and assigned to chromosome II (28). A disruption of ARO4 replacing a chromosomal 2.4-kb AccI

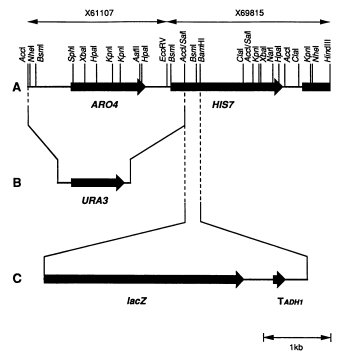


FIG. 2. Restriction map of the ARO4-HIS7 locus in different S. cerevisiae strains. Assigned GenBank/EMBL sequence accession numbers for the two genes are indicated above. (A) Wild-type situation. (B) Disrupted locus in strain RH1447. (C) Translational fusion of the HIS7 gene to the E. coli lacZ gene with the 3' end of the yeast ADH1 gene (T_{ADH1}) integrated at the original chromosomal locus in strains RH1614, RH1615, and RH1616.

fragment by the yeast URA3 gene (Fig. 2B) resulted in an additional His auxotrophy. The fact that the HIS7 gene is located on the same chromosome as the ARO4 gene (34) led us to the presumption that a concomitant disruption of the HIS7 gene could have caused the additional phenotype. To test this, we introduced the above-described gene disruption in MAT α strain RH1377 to yield RH1447 (Fig. 2B), and crossed both wild-type and disruption strain with a MATa his7 mutant (strain C20-2C) from the Yeast Genetic Stock Center. Wild-type strain RH1377, but not disruption strain RH1447, was able to complement the his7 mutation of strain C20-2C. This indicated that the HIS7 gene was located adjacent to the ARO4 gene and codisrupted with the ARO4 gene. The relative location of the genes was determined by Northern analysis. A 1.8-kb poly(A)⁺ RNA, which was not present in a disruption strain, could be detected by using the 0.5-kb EcoRV-BamHI fragment shown in Fig. 2A as a probe (data not shown). The HIS7 gene was therefore located immediately downstream of the ARO4 gene. Additional evidence for the location of HIS7 adjacent to ARO4 was obtained by cloning of the complete HIS7 gene by functional complementation of disruption strain RH1447 with a yeast genomic DNA library on a yeast centromeric plasmid. One of the transformants contained a 10.5-kb insert of yeast DNA. Subcloning localized the complementing activity on a chromosomal 6.2-kb HindIII fragment. This fragment was able to confer growth to RH1447 in the presence of 5 mM phenylalanine in the medium. Under these conditions the isoenzyme of the ARO4 gene product, the phenylalanineinhibitable, ARO3-encoded DAHPS, is fully inhibited and growth depends on an intact ARO4 gene. This indicated that

the 6.2-kb *Hin*dIII fragment contained both the *ARO4* and the *HIS7* gene.

A comparison of the restriction map of the fragment with the one of the ARO4 locus revealed a 2.4-kb EcoRV-HindIII fragment of chromosomal DNA located immediately downstream of the previously sequenced ARO4 AccI-EcoRV fragment (Fig. 2A) (28). The nucleotide sequence of the EcoRV-HindIII fragment contained a single open reading frame of 1,656 bp (Fig. 3). The HIS7 gene product is thus predicted to consist of 552 amino acids with a calculated molecular weight of 61,082.

Bifunctional nature of the HIS7 gene product. The HIS7 gene was previously assigned to both the fifth and sixth step in histidine biosynthesis in S. cerevisiae, converting phosphoribosyl-formimino-5-amino-1-phosphoribosyl-4-imidazole-carboxamide to imidazoleglycerol-phosphate (19, 26). This assignment was based on the analysis of accumulation products caused by mutational blocks in the biosynthetic pathway. In prokaryotic microorganisms, by contrast, these reactions are carried out by two monofunctional enzymes, a glutamine amidotransferase (fifth step) and a cyclase (sixth step). In E. coli and S. typhimurium these enzymes are encoded by two different genes, the hisH gene (glutamine amidotransferase) and the hisF gene (cyclase). In both organisms the two genes are part of a single histidine operon and separated by the hisA gene, which codes for phosphoribosyl-formimino-5-amino-1-phosphoribosyl-4-imidazole-carboxamide catalyzing the fourth step of the histidine biosynthetic pathway (Fig. 1).

The deduced amino acid sequence for the HIS7 gene was aligned to the prokaryotic glutamine amidotransferase and cyclase sequences currently available in the GenBank/ EMBL data base (Fig. 4). Alignments revealed significant homology of the N-terminal segment of the HIS7-derived amino acid sequence (amino acids 1 to 213) with the various *hisH*-derived amino acid sequences and of the C-terminal segment (amino acids 235 to 552) with the various *hisF*-derived amino acid sequences. Thus the primary structure of the HIS7 gene is consistent with its product being a bifunctional enzyme with a N-terminal glutamine amidotransferase and a C-terminal cyclase domain.

Many yeast genes like HIS2 (45) and HIS3 (50) have been functionally expressed in E. coli. In order to confirm the dual function of the HIS7 gene product on a functional level, we tested the HIS7 gene for its ability to complement different E. coli his mutants. Therefore we transformed E. coli K-12-derived strains W3110 (wild-type strain), UTH6 (hisA mutant), UTH860 (hisF mutant), and UTH1767 (hisH mutant) with plasmid pME979 carrying the 2.4-kb EcoRV-HindIII HIS7 fragment on vector pGEM7Zf(+) (Promega, Madison, Wis.) and the empty vector as a negative control. Selection for transformants was done on LB complete medium containing ampicillin. The transformants were tested for the His phenotype by streaking them on M9 minimal glucose agar and incubating the plates for 3 days at 37°C. The untransformed strains were plated on M9 minimal glucose agar and M9 minimal glucose agar supplemented with 20 mg of histidine per liter as a control. Plasmid pME979 containing the yeast HIS7 gene was able to suppress His auxotrophy of E. coli K-12 hisH and hisF mutants, whereas the hisA mutant could not be complemented (Fig. 5). Thus, the yeast HIS7 gene product can functionally replace both the hisHencoded glutamine amidotransferase and the hisF-encoded cyclase activity in E. coli.

In summary, both structural and functional findings reveal that the S. cerevisiae HIS7 gene encodes a bifunctional enzyme homologous to prokaryotic hisH and hisF gene products.

HIS7 regulation by the general control system of amino acid biosynthesis. In S. cerevisiae many genes involved in amino acid biosynthetic pathways are coordinately regulated by the general control system of amino acid biosynthesis. As the final step of a regulatory cascade under the environmental conditions of amino acid starvation, this system activates transcription of target genes by binding of the protein GCN4 to distinct recognition elements in the promoters of the corresponding genes (23). For five histidine-biosynthetic gene products (HIS1 to HIS5), a regulation by this system was demonstrated on the enzymatic level (26). After the isolation of the corresponding genes the regulation was confirmed on the transcriptional level for HIS1, HIS3, HIS4, and HIS5 (16, 24, 39, 51). The HIS7 gene could not be tested for a regulation by the general control system because of the lack of a convenient enzyme assay for the HIS7 gene product. The isolation of the HIS7 gene enabled us to perform an analysis of HIS7 transcription regulation. Two independent assays were used to demonstrate regulation of HIS7 transcription by the general control system of amino acid biosynthesis. Both assays made use of regulatory mutants in the general control system and a wild-type strain as a control. Strains carrying a gcd2-1 mutation express GCN4 at a constitutively high level and therefore mimic the situation of amino acid starvation (38), whereas in gcn4-103 and gcn4-101 mutants no functional GCN4 is present (22). Wildtype strains exhibit intermediate GCN4 levels (22). In the first method, HIS7 mRNA levels were determined relative to those of ARO4 and URA3 in S. cerevisiae RH1632 (gcd2-1), RH1631 (wild type), and F194 (gcn4-103) by Northern analysis (Fig. 6A). In a second approach, β -galactosidase activities of strains RH1614 (gcd2-1), RH1615 (wild type), and RH1616 (gcn4-101) carrying a translational HIS7-lacZ fusion integrated at the HIS7 locus were measured (Fig. 6B). The experiments revealed an up to sixfold derepression of the HIS7 gene in a gcd2-1 mutant compared with that in a gcn4 mutant. Thus, the HIS7 gene is regulated by GCN4.

Transcript 5' end mapping of GCN4-regulated genes under both repressing and derepressing conditions has revealed two types of transcription initiation patterns. In the HIS3 (44) and the TRP4 (20, 35) genes, the start sites for GCN4dependent transcription differ from those for GCN4-independent (basal) transcription, whereas other genes like HIS1 (24), HIS4 (16), HIS5 (39), ARO3 (41), and ARO4 (28) show the same pattern under both conditions. For the HIS7 gene the situation was analyzed by mapping the mRNA 5' ends in the general control regulatory mutants RH1371 (gcd2-1) and RH1381 (gcn4-101) and the wild-type strain RH1372 by both primer extension and S1 nuclease protection analysis (Fig. 7A). Three major HIS7 mRNA 5' ends could be mapped at positions -96, -88/89, and -60/64. The pattern of transcription start sites used was the same in all three genetic backgrounds, but the intensity of the signals at position -60/-64 relative to those at -88/89 and -96 was at least fivefold stronger in a gcd2-1 background than in a gcn4-101 background. Thus GCN4-dependent transcription uses preferentially the downstream located initiator elements at position -60/-64. The HIS7 gene belongs therefore to the same transcription initiation type as HIS3 and TRP4. The HIS7 transcript 3' ends were determined by the S1 nuclease protection method. Three ends located at positions +1723, +1739, and +1747 could be mapped in both a gcd2-1 and a gcn4-101 background (Fig. 7B). Thus the 3' untranslated region of the HIS7 gene has a length of 64 to 88 bp taking the

-476 ARO4	AatII
-376	TACCACTGCCAATTCGGTATTATTTAATTGTGTTTAGCGCTATTTACTAATTAACTAGAAACTCAATTTTTAAAGGCAAAGCTCGCTGACCTTTCACTGA
-276	poly(dA:dT)/GCRE1
-176	GCRE2 . TATA
-76 1	AAAGAAAAGAGAGGCAGCTATTGAAGTAGCAGTATCCAGTTTAGGTTTTTTAATTATTTACAAGTAAAGAAAAAGAGAATGCCGGTCGTTCACGTGATTGAC M P V V H V I D
- 25	GTTGAAAGTGGTAACCTACAGTCACTAACCAATGCAATTGAGCATTTAGGTTACGAAGTACAACTGGTGAAATCACCAAAGGATTTTAACATATCAGGCA
9	VESGNLQSLTNAIEHLGYEVQLVKSPKDFNISGT
125 43	CGTCAAGATTGATTTGCCTGGTGTCGGAAATTATGCCCATTTC <u>GTCGAC</u> AATTTATTAATAGAGGATTCGAAAAGCCGATAAGAGAATACATTGAATC S R L I L P G V G N Y A H F V D N L F N R G F E K P I R E Y I E S
225 76	TGGAAAACCAATAATGGGAATTTGCGTCGGGCTACAAGCGCTCTTTGCCGGTCCGTGGAAAGCCCTAAGAGTACGGGTCTGAACTACATTGATTTTAAG G K P I M G I C V G L Q A L F A G S V E S P K S T G L N Y I D F K BamHI
325 109	TTGTCCAGGTTCGATGATTCAGAAAAGCCAGTACCAGAAATAGGTTGGAATTCTTGCATTCCCTCGGAAAACCTATTCTTTGGATT <u>GGATCC</u> ATACAAGA L S R F D D S E K P V P E I G W N S C I P S E N L F F G L D P Y K R
425 143	GGTACTATTTCGTCCATTCTTTGCTGCCATTCTGAATTCAGAAAAGAAAAAAAA
525 176	AGAGGAATTTATTGCGGCAGTCAACAAGAATAATATATAT
625 209	TTGAAGCAACAAAGTCCTCCGATTCCAAACTATAGTGCGGAAGAAGAAGGAAG
725 243	CTTGTCTTGATGTACGTACTAATGACCAAGGTGATTTGGTGGTTACTAAAGGTGATCAATACGATGTACGTGAAAAAAAGTGATGGTAAAGGTGTTAGAAA C L D V R T N D Q G D L V V T K G D Q Y D V R E K S D G K G V R N
825 276	CCTTGGTAAGCCTGTTCAGTTGGCACAGAAATATTACCAACAGGGTGCGGATGAAGTAACATTTTTGAATATAACTTCTTTTAGAGATTGTCCTTTGAAG L G K P V Q L A Q K Y Y Q Q G A D E V T F L N I T S F R D C P L K
925 309	GATACTCCGATGCTAGAGGTTCTGAAACAAGCCGCAAAGACAGTCTTTGTTCCATTGACAGTCGGTGGGGGGGATCAAGGATATTGTCGATGTTGATGGAA D T P M L E V L K Q A A K T V F V P L T V G G G I K D I V D V D G T
1025 343	CCAAAATACCTGCTTTAGAAGTTGCAAGTCTATACTTCAGATCTGGTGCTGATAAAGTATCGATCG
1125 376	CGAGTTGGGTAACAGAGGAGATGGAACGTCACCAATAGAGACAATCTCGAAAGCATACGGTGCTCAGGCAGTTGTTATTTCTGTCGACCCTAAGAGAGATA E L G N R G D G T S P I E T I S K A Y G A Q A V V I S V D P K R V
1225 409	TATGTAAATTCACAAGCAGATACGAAGAACAAAGTCTTCGAGACAGAATATCCGGGCCCCCAATGGAGAGAAATACTGCTGGTACCAATGTACAAATCAAAG Y V N S Q A D T K N K V F E T E Y P G P N G E K Y C W Y Q C T I K G
1325 443	. Xbai GTGGAAGAGAA <u>TCTAGA</u> GACCTTGGTGTGTGGGAATTTAACAAGGGCATGTGAAGCTCTAGGTGCTGGGGAGATTTTATTGAACTGCATAGACAAGGATGG G R E S R D L G V W E L T R A C E A L G A G E I L L N C I D K D G
1425 476	NarI CTCTAATTCTGGTTATGATCTGGAATTGATAGAACATGTTAAAGATGCGGTCAAGATTCCCGTCATTGCATCCAGT <u>GGCGCC</u> GGTGTACCCGAACATTTC S N S G Y D L E L I E H V K D A V K I P V I A S S G A G V P E H F
1525 509	E E A F L K T R A D A C L G A G M F H R G E F T V N D V K E Y L L E
1625 543	Z/S. AGCACGGATTAAAGGTTAGAATGGATGAAGAGTAATGTGGTTGGA <u>TATGTA</u> TACTTTATAATCTTGACTCAGTCTATATACGCAATAATGATAGATGTTA H G L K V R M D E E
1725	AATCAGACATTTCACAACACAAGAGGATGTACAGCTTGGAGAAATTGTACCAACTTATATGGTTGTATTATTGGTGGTGTCAGTAGGGGAAGAAATAGAA
1825	CATATTTTTCCACTTTTTCATTTTTTTTTTTTTTAGCGAGGCATCGGAAATGAAAATTTTTTAAAAAATCGATGAGCTCCCACTTCTTCAACATTGACGAAAGG
1925	AAATATGCACTAAGTTGTTTTAAATCCAAGATTTGTCTCGTTTTAAGACTTACAGATAAAACAATATATTAGAAAGATTAACTATAATGGCCAGAGCATC
2025	CTCTACTARAGCCAGAAAACAGAGGCATGATCCACTTTTAAAGGATTTAGATGCAGCTCAAGGTACCTTGAAAAAAAA
2125	AACGATGCTGCAAAATCACGATGCTGCAAAATGAGGAAGATGGATACATAGACTCCAAAGCATCAAGAAAAAATTTTGCAGTTGGCCAAGGAACAACAGGATG
2225	AAATTGAAGGTGAGGAACTTGCTGAATCAGAAAGAAACAAGCAATTTGAAGCCAGATTCACCACCATGAGCTATGATGATGAAGACGAAGACGAAGACGA Hindiii
2325 TG. 3.	AGACGAAGCTT Nucleotide sequence of the HIS7 gene and deduced amino acid sequence for the encoded bifunctional glutamine amidotransferas

FIG. 3. Nucleotide sequence of the HIS7 gene and deduced amino acid sequence for the encoded bifunctional glutamine amidotransferase: cyclase. Nucleotide numbering refers to the A (+1) of the first ATG in the open reading frame. The presented sequence comprises the whole ARO4-HIS7 intergenic region and the last 19 codons of the ARO4 open reading frame (ARO4). Mapped HIS7 transcript 5' and 3' ends are indicated by solid arrowheads; ARO4 transcript 3' ends (28) are indicated by open arrowheads. Relevant restriction sites, the Zaret/Sherman consensus sequence for transcript 3' end formation (Z/S), a poly(dA-dT) stretch, two putative GCN4 recognition elements (GCRE), and a putative TATA element in the HIS7 promoter region, are indicated and underlined.

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ScHIS7		MPVWHVID	VESCINLOSILT	NATEHLGYEV	QLVKSP	34
<i>Ec</i> HisH		MNWILD	TGCANLNSVK	SALARHGY	EPKVSR	31
StHish		MNVVIID	T.CANLSSVK	SAGAPR.L.	HPGGQR	29 32
AbHisH		METMALID	YGSGNLRSPA	KALERAAAGC	HASFOVINTS	38
St cHisH	LTAAVPAGAT	GRAKKVVVFD	YGFGNVRSAE	RALARAGA	QLVKSP EPKVSR HPGGQR ETVISR HASFQVLVTS DVEITR	44
				-		
ScHIS7	KDFNISGTSR	LILPOVONYA	HFVDNLFNRG	FEKPIREYIE	SGKPIMGICV	84
StHieH	FAFTVLBADK	LEUPOVGTAQ	AAMDQVRERE	LFDLIKA	CTOPVLGICL	78 76
LlHisH	DLEEIRKADA	LILPGVGAFP	TAMNNLKKFN	LIELIOERAA	AGIPILGICL	82
Abhish	DADAVRKADR	VVIPAVGAFA	DCKRGLSEVP			68
<i>Stc</i> HisH	DYDKAMNADG	LILPGVGNYA LFLPGVGTAQ LFLPGVGTAQ LILPGVGAFP VVLPAVGAFA LLVPGVGAFA	ACMEGLKAAR	GDWIVDRRLS	GGRPVMGICV	94
ScHIS7						120
EcHisH	GMO.LLGRRS	EESNGVDLLG	IIDEDVPKMT	DFG. LPIPH	IGWNSCIP MGWNRVYPQA	125
StHisH	GMQ.LLGRRS	EETRGVDLLN	IIEQDVPKMT	DFG. LPIPH	MGWNRVYPOR	123
LlHisH	GMONLFEKGY	EIE.EROGLG	LLKGEVIPIK	TNEK. IPH	MGWNOLNLAK	128
Abhish	GMOLMAEGGR	EYGVT.ECIG	WIKGEVVKLE	PADPTLKIPH	MGWNRVYPQA MGWNRVYPQR MGWNQLNLAK MGWNELDIRR MGWNTVEAPA	117
StcHisH						
ScHIS7	SENLFFGLDP	YKRYYFVHSF	AAILNSEKKK	NLENDGWKIA	KAKYGSEEFI	179
<i>Ec</i> HisH	GNRLFQGIED	GAYFYFVHSY	AMP	VNPWTIA	QCNYGEPFTA	165
StHisH	GNRLFQGIED	GAYFYFVHSY	AMP	VNPWTIA	QCNYGEPFTA	163
LIHISH	TSPTTHYLSG	NDEVYFVHSY	QATCPD	DELIA	QCNYGEPFTA QCNYGEPFTA YTTYGEVKIP SADYGGPF.A	169
StcHisH	DSOLFAGLDA	DARFYFVHSY	AVHEWTOESH	NPLIAEPRVT	WSTHGKPFVA	190
00011011						
ScHIS7	AAVNKNNIFA	TOFHPERSGK	AGLNVIENFL	KQQSPPIPNY	SAEEKELLMN	229
EcHisH	A. WOKDNFYG	VOFHPERSGA	AGAKLLKNFL	EM		196
StHisH LlHisH	A.WOKDNFFG	OFHPERSGA	AGAQLLKNEL	EM		194
AbHish	AVVGRDNLVG	TOFHPERSOE	TGLALVANEL	TWRV		192
StcHisH	A.MENGALWA	TOFHPEKSGD	AGAQLLTNWI	ETL	SAEEKELLMN	222
<i>SC</i> HIS7 <i>EC</i> HISF	DYSNYGUTRR	ILACLOVETN	DOGDLVVIKG	DOMDVREKSD	GKGVRNLCKP	279
StHisF	MILAKE	TIPCLOVEDG			RNHEIIGDI	32
<i>Ll</i> HisF	MUTKR	ITPCLDIKNG	KWWKG	INF	GKGVRNIGKP .RNHEIIGDI .RNHEIIGDI .VGLREIGDP	32
AbHisF	MUKMR	VIPCLDVKDG	R	VNE	.VDLIDAGDP	32
0-11707						
<i>Sc</i> HIS7 <i>Ec</i> HisF	VIDIADRITIOU	GADEV TELNI	TASSOCPUT	TPMLEVLKQA	ARTVENPUTIV	329 79
StHisF	VPLAKRYADE	GADELVEYDI	TASSOGRVVD	KSWVARV	AEVIDIPHOV	79
				C FITCRA		79
<i>Ll</i> HisF	VELAKIMEEQ	CADEIIVELDI	TASPEREII	G ELIGRA	hater 2 th tutal	
LlHisF AbHisF	VELAKIMEEQ VEQARVMDRE	CADELVELDI GADELTELDI	TASHERRETT	YDVVRRT	AEQVEMPLITV	79
<i>Ll</i> HisF AbHisF	VOIAOKIYOO VPIAKRYAEE VPIAKRYADE VEIAKIYEEO VEOARVYORE					79
<i>Ll</i> HisF AbHisF ScHIS7	GGGIKDIVDV	DGTKIPALEV	ASLYFRSGAD	KVSIGTDAVY	AAEKYYELGN	79 379
LlHisF AbHisF ScHIS7 EcHisF StHisF	GGGIKDIVDV	DGTKIPALEV	ASLYFRSGAD	KVSIGTDAVY	AAEKYYELGN	79 379
LlHisF AbHisF ScHIS7 EcHisF StHisF LlHisF	GGGIKDIVDV	DGTKIPALEV	ASLYFRSGAD	KVSIGTDAVY	AAEKYYELGN	79 379
LlHisF AbhisF SchIS7 EchisF SthisF LlHisF AbhisF	GGGIKDIVDV AGGIKSLED. AGGIRSIDD. GGGIRSIDD. GGGVRTVDD.	DGTKIPALEV	ASLYFRSGAD AAKIISFGAD AAKIISFGAD FRRIIARGAD IRKLILAGAD	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSAAIE KVSINTAAIH	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE	79 379 118 118 118 118 118
LlHisF AbhisF SchIS7 EchisF SthisF LlHisF AbhisF	GGGIKDIVDV AGGIKSLED. AGGIRSIDD. GGGIRSIDD. GGGVRTVDD.	DGTKIPALEV	ASLYFRSGAD AAKIISFGAD AAKIISFGAD FRRIIARGAD IRKLILAGAD	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSAAIE KVSINTAAIH	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE	79 379 118 118 118 118 118
LlHisF AbhisF SchIS7 EchisF SthisF LlHisF AbhisF	GGGIKDIVDV AGGIKSLED. AGGIRSIDD. GGGIRSIDD. GGGVRTVDD.	DGTKIPALEV	ASLYFRSGAD AAKIISFGAD AAKIISFGAD FRRIIARGAD IRKLILAGAD	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSAAIE KVSINTAAIH	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE	79 379 118 118 118 118 118
LlHisF AbhisF SchIS7 EchisF SthisF LlHisF AbhisF	GGGIKDIVDV AGGIKSLED. AGGIRSIDD. GGGIRSIDD. GGGVRTVDD.	DGTKIPALEV	ASLYFRSGAD AAKIISFGAD AAKIISFGAD FRRIIARGAD IRKLILAGAD	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSAAIE KVSINTAAIH	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE	79 379 118 118 118 118 118
LlHisF AbhisF SchIS7 EchisF SthisF LlHisF AbhisF	GGGIKDIVDV AGGIKSLED. AGGIRSIDD. GGGIRSIDD. GGGVRTVDD.	DGTKIPALEV	ASLYFRSGAD AAKIISFGAD AAKIISFGAD FRRIIARGAD IRKLILAGAD	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSAAIE KVSINTAAIH	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE	79 379 118 118 118 118 118
LlHisF AbHisF ScHIS7 EcHisF StHisF AbHisF AbHisF ScHIS7 EcHisF LlHisF AbHisF	GGGTKDIVDV AGGTKSLED. AGGTRSIDD. GGGTRSIDD. GGGTRSIDD. RGDGTSPIET R. R. E. K.	DGTKIPALEV	ASLYFRSGAD AAKTISFGAD AAKTISFGAD FRRTIARGAD IRKITLAGAD VISUDPKRVY VVSIDTW VVSIDTW VVSIDTW VVAIDAK	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSPALA KVSINTAAIH VNSQADTKNK	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN F F Q	79 379 118 118 118 118 429 133 133 133 133
LlHisF AbHisF ScHIS7 EcHisF StHisF AbHisF AbHisF ScHIS7 EcHisF LlHisF AbHisF	GGGTKDIVDV AGGTKSLED. AGGTRSIDD. GGGTRSIDD. GGGTRSIDD. RGDGTSPIET R. R. E. K.	DGTKIPALEV	ASLYFRSGAD AAKTISFGAD AAKTISFGAD FRRTIARGAD IRKITLAGAD VISUDPKRVY VVSIDTW VVSIDTW VVSIDTW VVAIDAK	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSPALA KVSINTAAIH VNSQADTKNK	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN F F Q	79 379 118 118 118 118 429 133 133 133 133
LlHisF AbHisF ScHIS7 EcHisF StHisF AbHisF AbHisF ScHIS7 EcHisF LlHisF AbHisF	GGGTKDIVDV AGGTKSLED. AGGTRSIDD. GGGTRSIDD. GGGTRSIDD. RGDGTSPIET R. R. E. K.	DGTKIPALEV	ASLYFRSGAD AAKTISFGAD AAKTISFGAD FRRTIARGAD IRKITLAGAD VISUDPKRVY VVSIDTW VVSIDTW VVSIDTW VVAIDAK	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSPALA KVSINTAAIH VNSQADTKNK	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN F F Q	79 379 118 118 118 118 429 133 133 133 133
LlHisF AbHisF ScHIS7 EcHisF StHisF AbHisF AbHisF ScHIS7 EcHisF LlHisF AbHisF	GGGTKDIVDV AGGTKSLED. AGGTRSIDD. GGGTRSIDD. GGGTRSIDD. RGDGTSPIET R. R. E. K.	DGTKIPALEV	ASLYFRSGAD AAKTISFGAD AAKTISFGAD FRRTIARGAD IRKITLAGAD VISUDPKRVY VVSIDTW VVSIDTW VVSIDTW VVAIDAK	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSPALA KVSINTAAIH VNSQADTKNK	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN F F Q	79 379 118 118 118 118 429 133 133 133 133
LlHisF AbHisF ScHIS7 EcHisF StHisF AbHisF AbHisF ScHIS7 EcHisF LlHisF AbHisF	GGGTKDIVDV AGGTKSLED. AGGTRSIDD. GGGTRSIDD. GGGTRSIDD. RGDGTSPIET R. R. E. K.	DGTKIPALEV	ASLYFRSGAD AAKTISFGAD AAKTISFGAD FRRTIARGAD IRKITLAGAD VISUDPKRVY VVSIDTW VVSIDTW VVSIDTW VVAIDAK	KVSIGTDAVY KISINSPALA KISINSPALA KVSVNSPALA KVSINTAAIH VNSQADTKNK	AAEKYYELGN DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN F F Q	79 379 118 118 118 118 429 133 133 133 133
LlHisF AbHisF ScHIS7 EcHisF StHisF AbHisF AbHisF ScHIS7 EcHisF LlHisF AbHisF	GGTKDIVDV AGGTKSLDD. GGGTKSLDD. GGGTKSIDD. GGGTKSIDD. GGGTKSIDD. GGGTKTVDD. R. R. K. GGKYCW20CT GAETGKHVN DATGKYEVN VEPGR. BIF	DGTKIPALEV 	ASLYERSGAD AAKILISIGAD AAKILISIGAD FRIILAIGAD IRKILILAGAD VISYODERKVY VVGIDTW VVGIDTW VVGIDTW VVGIDAK VVGIDAK VVGIDALDAK DLGVWELT VTQWETLDWV AGLDLADAR AGLDLADAR	KVSICTDAVY KISINSALA KISINSALA KVSVNSAIE KVSINTAAIH VNSQADTKNK 	AAEKYYELGN DPTLITRLAD DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN 	79 379 118 118 118 118 429 133 133 133 133 133 133 133 183 183 179 178
LlHisF AbHisF ScHIS7 EcHisF StHisF AbHisF AbHisF ScHIS7 EcHisF LlHisF AbHisF	GGTKDIVDV AGGTKSLDD. GGGTKSLDD. GGGTKSIDD. GGGTKSIDD. GGGTKSIDD. GGGTKTVDD. R. R. K. GGKYCW20CT GAETGKHVN DATGKYEVN VEPGR. BIF	DGTKIPALEV 	ASLYERSGAD AAKILISIGAD AAKILISIGAD FRIILAIGAD IRKILILAGAD VISYODERKVY VVGIDTW VVGIDTW VVGIDTW VVGIDAK VVGIDAK VVGIDALDAK DLGVWELT VTQWETLDWV AGLDLADAR AGLDLADAR	KVSICTDAVY KISINSALA KISINSALA KVSVNSAIE KVSINTAAIH VNSQADTKNK 	AAEKYYELGN DPTLITRLAD DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN 	79 379 118 118 118 118 429 133 133 133 133 133 133 133 183 183 179 178
LlHisF AbHisF ScHIS7 EcHisF ScHisF ScHIS7 EcHisF ScHIS7 EcHisF AbHisF ScHIS7 EcHisF ScHIS7 ScHIS7 EcHisF	GGTKDIVDV AGGTKSLDD. GGGTKSLDD. GGGTKSIDD. GGGTKSIDD. GGGTKSIDD. GGGTKTVDD. R. R. K. GGKYCW20CT GAETGKHVN DATGKYEVN VEPGR. BIF	DGTKIPALEV 	ASLYERSGAD AAKILISIGAD AAKILISIGAD FRIILAIGAD IRKILILAGAD VISYODERKVY VVGIDTW VVGIDTW VVGIDTW VVGIDAK VVGIDAK VVGIDALDAK DLGVWELT VTQWETLDWV AGLDLADAR AGLDLADAR	KVSICTDAVY KISINSALA KISINSALA KVSVNSAIE KVSINTAAIH VNSQADTKNK 	AAEKYYELGN DPTLITRLAD DPTLITRLAD DPTLITRLAD NPELIRQAAN RPEFVQEAAE VFETEYPGPN 	79 379 118 118 118 118 429 133 133 133 133 133 133 133 183 183 179 178
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Organism	Glutamine- amidotransferase (HisH)	Cyclase (HisF)
Escherichia coli	62.4 / 39.2 %	61.9 / 36.6 %
Salmonella typhimurium	57.3 / 35.9 %	61.1 / 36.2 %
Lactococcus lactis	54.3 / 36.0 %	65.4 / 44.9 %
Azospirillum brasilense	57.1 / 36.3 %	64.3 / 40.4 %
Streptomyces coelicolor	56.3 / 33.5 %	-/-

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FIG. 4. Comparison of deduced amino acid sequences for different prokaryotic monofunctional glutamine amidotransferases (HisH) and cyclases (HisF) with the HIS7 gene product from S. cerevisiae. (A) Multisequence alignment. The various sequences (Ec. E. coli; St: S. typhimurium; Ll: L. lactis; Ab: A. brasilense; Stc: S. coelicolor) were obtained from the GenBank/EMBL data base and aligned with the deduced HIS7 amino acid sequence (top line). S. coelicolor genes are designated as suggested by Limauro et al. (30). Residues similar in all compared sequences are boxed. (B) Pairwise comparisons. Identities and similarities (%) of the N-ter-

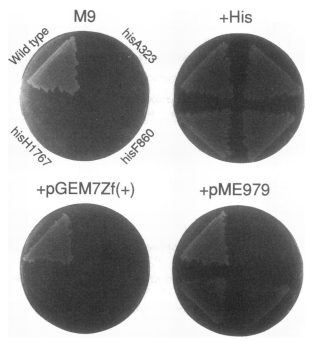


FIG. 5. Suppression of His auxotrophy in *E. coli* by the yeast *HIS7* gene. *E. coli* K-12 derivatives W3110 (wild type), UTH6 (*hisA323*), UTH860 (*hisF860*) and UTH1767 (*hisH1767*) harboring either no plasmid (M9, +His), the empty vector [pGEM7Zf(+) (Promega, Madison, Wis.)] or pGEM7Zf(+) bearing a 2.4-kb *EcoRV-Hind*III *HIS7* fragment (pME979) were streaked onto M9 minimal glucose agar [M9, +pGEM7Zf(+), +pME979] or M9 minimal glucose agar supplemented with 20 mg of histidine (+His) per liter and incubated for 3 days at 37°C. Complementation of a *hisA* mutation was tested as the *hisA* gene is located between the genes *hisH* and *hisF* in the *E. coli* histidine operon (Fig. 1).

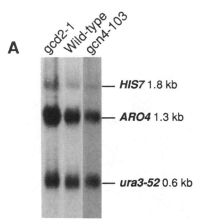
translational stop codon at position +1657 into account. According to the transcript end-mapping experiments, the *HIS7* transcripts have a length of approximately 1.8 kb, which corresponds to the length determined by Northern analysis (Fig. 6A).

DISCUSSION

In this report, we provide primary structure, complementing activity in *E. coli*, and regulation of the *S. cerevisiae HIS7* gene. The gene codes for the first eukaryotic glutamine amidotransferase and cyclase catalyzing the fifth and sixth step in the histidine biosynthetic pathway. The primary structure and function of the two enzymatic activities are shown to be conserved from *E. coli* to *S. cerevisiae*, whereas structure, organization, and regulation of the corresponding genes differ considerably in the two organisms (Fig. 1).

In *E. coli* the two enzymatic activities are encoded by two separate cistrons, hisH (glutamine amidotransferase) and hisF (cyclase), organized in a single operon. The two cistrons are separated by the hisA cistron, which codes for

minal segment of the HIS7 sequence (amino acids 1 to 213) with the various glutamine amidotransferase sequences (HisH) and of the C-terminal segment (amino acids 235 to 552) with the various cyclase sequences (HisF) were calculated on the basis of pairwise alignments.



Strain	Relevant genotype	B-Galactosidase- activity (U)
RH1614	gcd2-1	143
RH1615	Wild-type	35
RH1616	gcn4-101	25

FIG. 6. *HIS7* expression analysis in *S. cerevisiae* strains expressing different amounts of GCN4 protein. (A) Northern analysis. Poly(A)⁺ RNA of strains RH1632 (*gcd2-1*, high amount of GCN4 protein), RH1631 (wild type, intermediate amount of GCN4 protein), and F194 (*gcn4-103*, no GCN4 protein) was hybridized against DNA probes for the *HIS7*, *ARO4*, and *URA3* transcripts. Sizes of the various transcripts are indicated. The *URA3* transcript was chosen as a negative control and the *ARO4* transcript was chosen as a positive control for a regulation by the general control system of amino acid biosynthesis. (B) β -Galactosidase activity of integrated *HIS7-lacZ* fusions. Activities have been determined for strains indicated. Yeast strains harboring no *E. coli lacZ* gene did not show any detectable β -galactosidase activity (data not shown).

phosphoribosyl-formimino-5-amino-1-phosphoribosyl-4-imidazole-carboxamide isomerase (EC 5.3.1.16) catalyzing the fourth step of the pathway. In S. cerevisiae both enzymatic activities are fused on a single polypeptide chain. The HIS7 gene codes for a bifunctional enzyme with an N-terminal glutamine amidotransferase and a C-terminal cyclase domain. Besides the HIS7 enzyme, there is only one other multifunctional enzyme in the histidine biosynthetic pathway in S. cerevisiae (Fig. 1). The HIS4 gene product catalyzes four steps in the pathway and seems to be unique to fungi, as the four enzymatic activities are catalyzed by at least two distinct enzymes both in prokaryotes and plants (see introduction). The HIS7 gene product probably represents an analogous situation, as in no prokaryote studied so far are these enzymatic steps fused on a single polypeptide chain (5, 11, 14, 17, 30) and furthermore in N. crassa the steps seem to be genetically coupled as well (1). By contrast, the physical uncoupling of the seventh and ninth step of the pathway in S. cerevisiae compared with the situation in enterobacteria is not unique to fungi, as the same situation is found both in eubacteria such as L. lactis, S. coelicolor, and A. brasilense (14, 17, 30) and in the methanogenic archaebacterium M. vannielii (6).

The HIS7 gene was located on chromosome II immediately downstream of the previously isolated and characterized ARO4 gene (28). The ARO4 gene codes for the tyrosineinhibitable DAHPS catalyzing the first step in the biosynthetic pathway of the aromatic amino acids. The two genes have the same orientation with a distance of only 416 bp between the two open reading frames. The mapped ARO4 mRNA 3' ends and HIS7 mRNA 5' ends are only 121 bp apart (Fig. 3) which leaves little space for signal sequences directing termination of ARO4 transcription and initiation of HIS7 transcription. Such a close packing of two independent genes is not untypical for S. cerevisiae (40). Nevertheless, it is remarkable that both genes are involved in amino acid biosynthetic pathways and coregulated by the general control system of amino acid biosynthesis in S. cerevisiae. Both the HIS7 and the ARO4 gene (28) are derepressed severalfold under amino acid starvation conditions. To our knowledge, there is no other example of two coregulated genes adjacent to each other in S. cerevisiae.

In the HIS7 gene GCN4-dependent transcription uses only selected initiator elements used by GCN4-independent (basal) transcription, whereas in the ARO4 gene the same elements are used under both repressing and derepressing conditions (28). In this respect the HIS7 gene resembles the HIS3 gene coding for imidazoleglycerol-phosphate dehydratase (51). Of the two mapped HIS3 transcripts, only the one located more downstream is subject to GCN4 control. Constitutive and regulated HIS3 transcription differ not only by their utilization of initiator elements but also by their required upstream promoter elements. Upstream elements in the HIS3 promoter include a poly(dA-dT) stretch for constitutive transcription and two GCN4 recognition elements (GCRE) (21) for maximal induction by GCN4. In addition, two classes of TATA elements have been suggested, responsible for constitutive and GCN4-regulated HIS3 transcription, respectively. In the HIS4 gene the start site patterns for GCN4-dependent and GCN4-independent transcription are identical (16). The HIS4 promoter contains not less than five GCREs for derepression by GCN4 (3). GCN4-independent transcription of the HIS4 gene is controlled by the global activators BAS1 and BAS2 (4, 52) which have also been shown to regulate purine biosynthesis (13). In addition, BAS2 (also known as PHO2) is involved in the regulation of phosphate metabolism (53) and tryptophan biosynthesis (8). The HIS4 TATA element (37) is required for correct mRNA start site selection by GCN4-dependent transcription but not by GCN4-independent transcription (43).

In the *HIS7* promoter region various putative upstream elements are present. Two possible GCREs, a one-mismatch consensus sequence (ATGACTCAA in inverse orientation) at position -228 (GCRE1) and a two-mismatch consensus sequence (CTGACTCTT in inverse orientation) at position -142 (GCRE2), are found (Fig. 3). The latter element contains the hexanucleotide sequence TGACTC followed by a T. This motif is found in several binding sites of BAS1 to DNA (13). As a further putative upstream element for

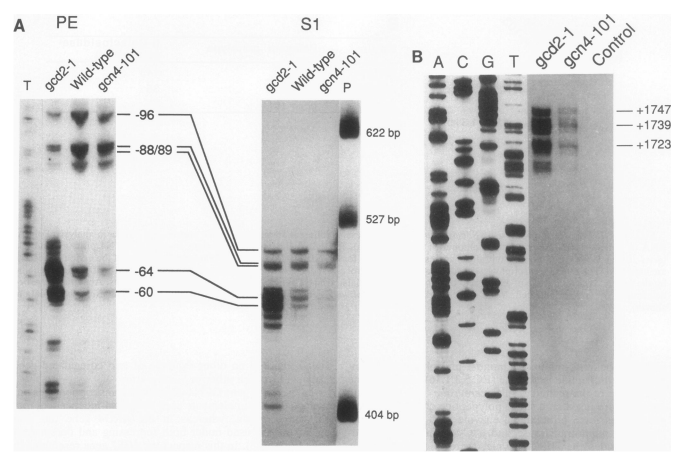


FIG. 7. Mapping of HIS7 transcript ends in S. cerevisiae strains expressing different amounts of GCN4 protein. (A) HIS7 mRNA 5' end mapping. The 5' ends of the HIS7 transcript were determined in strains RH1371 (gcd2-1), RH1372 (wild type), and RH1381 (gcn4-101) both by primer extension analysis (PE) and S1 nuclease mapping (S1). A T ladder (T) produced with the same primer as for the primer extension reactions was used as a standard for the size of the elongation products. For S1 nuclease mapping, plasmid pBR322 DNA cut with HpaII (P) served as a size marker. (B) HIS7 mRNA 3' end mapping. The 3' ends of the HIS7 transcript were determined in strain RH1371 (gcd2-1) and RH1381 (gcn4-101). The DNA fragment used for S1 nuclease mapping of the mRNA 3' ends was produced by PCR technique. As a size standard a sequence ladder generated with one of the primers used in the PCR reaction was used. Torula yeast RNA (30 μ g) was chosen as a negative control.

constitutive transcription, an 11-bp poly(dA-dT) stretch is located at position -236, immediately upstream of GCRE1. The only sequence element in the *HIS7* promoter region resembling a TATA element (55) is found as TACATAAG in normal orientation at position -122. In the 3' untranslated regions of both the *HIS7* and *ARO4* genes perfect matches to the sequence TATGTA (57) are found as possible signal sequences for transcript 3' end formation (Fig. 3).

Further experiments will concentrate on the identification and characterization of control elements in the ARO4-HIS7 intergenic region involved in GCN4-independent (basal) or GCN4-dependent HIS7 transcription and eventually in preventing transcriptional interference between the two adjacent and coregulated genes.

ACKNOWLEDGMENTS

We thank Ralf Hütter for his continued interest and support. We are especially grateful to Phil Hieter for providing the yeast genomic DNA library and Barbara Bachmann for providing the *E. coli his* mutants. We also appreciate Hans-Ueli Mösch for helpful discussions and Roney Graf and Brigitta Mehmann for critical reading of the manuscript.

This work was supported by the Swiss National Foundation (grant no. 31.29926.90) and the Swiss Federal Institute of Technology.

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