### The primary structure of the Saccharomyces cerevisiae gene for 3-phosphoglycerate kinase

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#### ABSTRACT

The DNA sequence of the gene for the yeast glycolytic enzyme, 3-phosphoglycerate kinase (PGK), has been obtained by sequencing part of a 3.1 kbp HindIII fragment obtained from the yeast genome. gene sequence corresponds to a reading frame of 1251 bp coding for 416 amino<br>acids with no intervening DNA sequences. The amino acid sequence is acids with no intervening DNA sequences. approximately 65 percent homologous with human and horse PGK protein sequences and is in general agreement with the published protein sequence for yeast PGK. As for other highly expressed structural genes in yeast, the coding sequence is highly codon biased with 95 percent of the amino acids coded for by a select 25 codons (out of 61 possible). Besides structural DNA sequence, 291 bp of 5'-flanking sequence and 286 bp of 3'-flanking sequence were determined. Transcription starts 36 nucleotides upstream from the translational start and stops 86-93 nucleotides downstream from the translational stop. These results suggest a non-polyadenylated mRNA length of 1373 to 1380 nucleotides, which is consistent with the observed length of<br>1500 nucleotides for polyadenylated PGK mRNA. A sequence TATATATAAA is 1500 nucleotides for polyadenylated PGK mRNA. found at 145 nucleotides upstream from the translational start. This sequence resembles the TATAAA box that is possibly associated with RNA polymerase II binding.

#### INTRODUCTION

Glycolytic genes from yeast are of great interest due to their combined high expression of 65 percent or more of the soluble protein in yeast (1). Two of these enzymes, 3-phosphoglycerate kinase (PGK) and glyceraldehyde-3-phosphate dehydrogenase, individually constitute 4 to 10 percent of the soluble protein depending on qrowth conditions (1, 2). The high level of glyceraldehyde-3-phosphate dehydrogenase may result in part from the presence of three copies of the gene (3); while PGK is present as only one copy per haploid cell (4, 5). Furthermore, high expression appears to correlate with the high levels of mRNA from these genes (6).

The gene for PGK has been previously cloned from the yeast genome on a plasmid by an immunological screening technique (5). Early characterization of this plasmid suggested that it contained the PGK structural gene on a 3.1

kbp HindIII fragment. A DNA insertion at the EcoRI site (see Fig. 1) in this fragment was shown to affect gene expression adversely (5). We have further characterized the 3.1 kbp HindIII fragment by sequencing 1828 bp from a PvuI site to a HindIII site. This DNA fragment contains the EcoRI site within the structural gene as well as 5'- and 3'-flanking sequences for the PGK structural gene.

Recent studies of yeast gene expression have focused on comparisons of the DNA sequences of  $5'-$  and  $3'-$ flanking regions of isolated genes. All genes apparently have a TATAAA-like sequence at various distances upstream (5') from the translational start (3, 7-10). This sequence is thought to be associated with RNA polymerase II positioning and binding in vivo (11, 12) and is usually about 30 nucleotides upstream from the translational start in most multicellular eukaryotes (13). However, in the lower eukaryote yeast this distance is quite variable (11). Yeast genes also demonstrate variable numbers of transcription starts for the same gene with as many as seven for iso-l-cytochrome c (11) and two for alcohol dehydrogenase <sup>I</sup> (ADH-1) (14). In one gene system, this variation has been shown to be involved with the regulation of gene expression. The SUC2 gene produces a 1.8 kb transcript for constitutive levels of an intracellular invertase; while a larger 1.9 kb transcript, which includes the secretion pre-sequence of invertase, is derepressed to give the extracellular form of this enzyme (16).

The untranslated 3'-flanking regions of yeast genes are thought to be associated with at least two functions: the termination of transcription and polyadenylation of the 3' end of the transcript. A deletion of 38 bp in the 3'-flanking sequence of the CYCl gene (17) results in 5-10 percent of the wild type level of iso-l-cytochrome c and only 10 percent of the normal level of mRNA. However, the mRNA in this mutant is terminating in other regions 3' to the structural gene. By comparison of the sequence defined by this deletion with other  $3'-f$ lanking gene sequences, Zaret and Sherman (17) have suggested a consensus sequence for transcription termination and polyadenylation for most yeast genes. Bennetzen and Hall (14) have suggested another consensus sequence thought to be involved with these same functions.

We now present the sequence for the PGK structural gene and flanking regions and compare it with those of other yeast genes to possibly identify the factors involved in high expression observed for glycolytic genes. We also determine the presence and relevance of some consensus sequences thought to be involved with gene expression in yeast. Furthermore, we compare the yeast PGK amino acid sequence with amino acid sequence determined for PGKs from higher eukaryotes.

## MATERIALS AND METHODS

## **Materials**

Restriction endonucleases, T4 DNA ligase, and polynucleotide kinase were purchased from BRL or New England Biolabs and used essentially as recommended by manufacturer. E. coli DNA polymerase <sup>I</sup> (large fragment) was from Boehringer and Si nuclease from Miles. ATP and deoxyribonucleotide triphosphates were obtained from PL Biochemicals; while <sup>32</sup>P-labeled nucleotides were obtained from Amersham Radiochemicals. Glass beads (0.45-0.50 mm) were purchased from B. Braun Melsungen AG. The oligonucleotide primer (5'-ATTTGTTGTAAA-3') was synthesized by conventional means (18). Strains, plasmids, and growth conditions

E. coli K-12 strain 294 (endA thr $ThsrThsm$ ) (19) was used for bacterial transformations. S. cerevisiae strains  $20B-12$  ( $\alpha$  trpl pep4-3) (20) and GM3C-2 (a leu2-3 leu2-112 trpl-1 his4-519 cycl-1 cyp3-1) (11) were used for yeast transformations. Five plasmids were used, some of which have been previously described: pBl (5), YEp13 (21), YRp7' (22, 23), PGK-YEp13, and pFRM31. Plasmid pBl contains the 3.1 kbp PGK HindIII fragment, isolated by immunological screening (5), in the HindIII site of pBR322. Plasmid YEp13 contains the yeast  $2<sub>u</sub>$  origin of replication and the yeast LEU2 gene for complementation of the double leu2 mutation in GM3C-2. YRp7' contains the 1.4 kbp TRP1 EcoRI fragment (23) in pBR322 in the opposite orientation as compared to YRp7 (22). This fragment contains the arsl origin of replication and the TRP1 gene for complementation of the trpl mutation in either 20B-12 or GM3C-2. PGK-YEp13 contains the 3.1 kbp PGK HindIII fragment from pBl substituted for the small HindIII fragment in YEp13 (orientation of PGK is with transcription toward adjacent DNA from yeast  $2\mu$  plasmid). Plasmid pFRM31 was constructed by inserting the 3.1 kbp PGK HindIII fragment into the pBR322 HindIII site of pFRD7 (which is YRp7' with the EcoRI sites removed by filling in EcoRI ends using large fragment of E. coli DNA polymerase I).

LB medium for E. coli growth was as described by Miller (24) with the addition of 20 ug/ml ampicillin (Sigma) for selection of plasmid transformants. Yeast were grown on the following media: YEPD (nonselective) contained 1 percent yeast extract, 2 percent peptone and 2 percent glucose with or without 3 percent Difco agar. YNB+CAA (used for  $Trp^+$  selection) contained 6.7 grams of Difco yeast nitrogen base (without amino acids) (YNB), 10 mg of adenine, 10 mg of uracil, 5 grams Difco casamino acids (CAA), 20 grams glucose and with or without 30 grams agar per liter. YNB-leu (leucine absent for Leu selection) contained the same components as YNB+CAA; however, with 20 ml of -leu drop-out mix replacing the CAA (-leu drop-out mix contained these amino acids per  $100$  ml of  $H<sub>2</sub>0$ : 0.2g arg, 0.1g his, 0.6g ile, 0.4g lys, 0.1g met, 0.6g phe, 0.5g thr, and 0.4g trp). The transformed yeast were always grown in media for selective maintenance of the plasmid.

# Plasmid DNA preparation and transformations

Purification of plasmid DNAs from E. coli (25) and transformation of E. coli (26) were done in accordance with previously described procedures. E. coli plasmid identification was as described by Birmboim and Doly (27). Transformation of yeast was done as previously described (28).

# DNA sequence determination

DNA sequencing of the phosphoglycerate kinase gene and flanking regions was done both by the chemical degradation method (29), using 5'  $32p-1$ abelled restriction fragments, and by the dideoxy-method (30). For the latter approach the restriction fragments containing Sau3A ends (Fig. 1), were subcloned into the BamHI site of the single stranded phage M13 mp8 (31). The dideoxy-chain termination procedure was used as described using a synthetic phage-specific primer.

# mRNA analysis

Preparation of yeast total RNA was done as previously described (32) growing the yeast under selective conditions for plasmid retention to an A<sub>660</sub> of 1.0. Total yeast RNA was then denatured and electrophoresed on a 1.0 percent agarose gel containing 6 percent formaldehyde and 1X MOPS buffer (20 mM MOPS, 5 mM NaOAc, <sup>1</sup> mM EDTA at pH 7.0) (33). The gel was stained with ethidium bromide using the yeast or E. coli ribosomal RNAs (34) and HindIII cut phage  $\lambda$  DNA (35) as size standards. The gel was transferred to nitrocellulose and hybridized (36) with the EcoRI-BglII PGK-termination DNA fragment (from pB1) that was  $32P-1$ abelled using the calf thymus primer method (37).

The initiation of transcription of the PGK gene was determined using cDNA extension (38) from a specific  $5'$   $32P-1$ abelled DNA primer,  $5'$ ATTTGTTGTAAA 3', complementary to nucleotides -10 to -21 (Fig. 2). Polyadenylated mRNA was prepared from yeast pFRM31/20B-12 (39). The extension product was sized on a 15 percent polyacrylamide-7M urea gel

alongside a sequencing ladder.

The polyadenylation site was mapped with the Si mapping procedure (40), using the conditions of Mantei et al. (41). The BglII-HindIII PGK-terminator fragment from pBl (5), 3' labelled at the BglII site using the Klenow fragment of E. coli DNA polymerase I and  $a^{32}P$ -dCTP, was hybridized to 10 ug mRNA from pFRM31/20B-12 followed by S1 nuclease digestion. The reaction products were run on a 6 percent polyacrylamide-7M urea gel. The  $32$ P-labelled HpaII fragments of pBR322 (42) were used as size markers.

#### RESULTS

# Primary structure of the yeast PGK gene and protein

The 3.1 kb HindIII fragment, carrying the PGK gene, was isolated from plasmid pBl (5). The recognition sites for several restriction endonucleases were mapped using standard techniques. The resulting preliminary restriction map served as a basis for the DNA sequence determination. Both the Maxam-Gilbert and dideoxy method were used to determine the sequence of the PvuI-HindIII segment by the strategy shown in Fig. 1. The arrows illustrate direction and scope of the sequencing as well as the method employed. Most regions were determined by sequence analysis in both directions (both strands) or using both methods.

The DNA sequence of the PvuI-HindIII fragment (1828 bp) is shown in Fig. 2. The largest open reading frame is 1248 nucleotides long, thus coding for <sup>a</sup> protein of 416 amino acids, with a calculated molecular weight of 44,746 daltons for the unmodified polypeptide. This value agrees well with the previous estimations of 45,000 to 50,000 daltons (43). In analogy with other yeast and higher eukaryotic proteins, it is very likely that the NH<sub>2</sub>-terminal methionine is removed. The adjacent serine is then probably



Figure 1. DNA sequencing strategy for the PGK gene. The 3.1 kbp HindIII fragment from plasmid pBl (5) was sequenced from the PvuI site to the HindIII site (1828 bp) using two sequencing techniques for strands, distances, and directions as indicated by the arrows. The structural gene is shown as the bar region. Restriction sites are to scale except from HindIII to PvuI on the left which was not sequenced.

modified since automated Edman-degradation (44) of the native protein from yeast did not result in the release of terminal amino acids (43). This was similarly observed for the human (45) and horse (46) PGKs.

Recently, Dobson et al. (47) have reported the partial amino acid sequence (residues 270-400) of the CN2 cyanogen bromide fragment (amino acids 270 to 419) from yeast PGK. Their sequence corresponds well with the amino acids 268 to 398 of our deduced polypeptide sequence, as shown by the underlined residues in Fig. 2. However, assuming the removal of the terminal methionine, there is a difference of three amino acids in the numbering. Their numbering corresponds to the analogous residues in the human PGK protein (45). Our protein sequence, as deduced from the DNA sequence, differs in 10 out of the 130 residues, with their directly determined protein sequence. A single base change in the codons can account for seven of these differences (differences designated by amino acids in parentheses in Fig. 2). They can therefore be explained by some variation depending on the yeast strain. Although such differences have been seen for other yeast genes, it seems unlikely that strain variations would account for the three additional amino acid differences, which correspond to codon changes of two or three nucleotides.

## PGK gene codon bias

Recently, Bennetzen and Hall (48) have tabulated the codon usage of seven yeast genes and noted that seven codons were never used (CUC, leu; CUG, leu; CCG, pro; CGU, arg; CGC, arg; CGA, arg; CGG, arg) and that highly expressed genes use fewer codons than genes that are expressed at much lower levels. Furthermore, for the two highly expressed genes, alcohol dehydrogenase <sup>I</sup> (ADH-1) and glyceraldehyde-3-phosphate dehydrogenase, 96 percent of the amino acids are coded for by 25 of the 61 possible coding triplets. The ADH-1 gene uses only 33 of 61 possible codons and 30 of these are used more than once (14). The same preference is observed in the yeast enolase genes (49).

Fig. 3 shows that a similar codon bias is present in the yeast PGK gene. Although more different codons (38 of 61) are used than in the genes for ADH-1, glyceraldehyde-3-phosphate dehydrogenase, and enolase; a very strong preference for 25 select codons, accounting for 95 percent of the codon capacity, is apparent. Therefore PGK codon usage is similarly restricted as for the other highly expressed genes, in contrast with the less biased genes which demonstrate low level expression in yeast. As shown by asterisks in Fig. 2, non-preferred codons are randomly spaced throughout the sequence. A similarly random spacing of the less favored codons is observed in the ADH-1 gene (14). These less preferred codons in the PGK gene probably do not grossly affect or restrict higher expression levels,

-280 -260 -240 -220 -200 -180 GATCGTACTGTTACTCTC TC TCTTTCAAACAGAATTGTC CGAATC6GTGTGACAACAACAGCCTGTTCTCACACACTCTTTTCTTCTAACCAAGGGGGTGGTTTAGTTTAGTAGAACCTC -160 -140 -120 -100 -80 -60 GTGAAACTTACATTTACATATATATAAACTTGCATAAATTGGTCAATGCAAGAAATACATATTTGGTCTTTTCTAATTCGTAGTTTTTCAAGTTCTTAGATGCTTTCTTTTTCTC TTTT 20 -40 -20<br>met ser leu ser leu ser val gin asp leu asp leu lys asp | 1<br>TTACAGATCATCAAGGAAGTAATCTACTTTTTTACAACAAATATAAAACA ATG TCT TTA TCT TCA AAG TTG TCT GTC CAA GAT TTG GAC TTG AAG 20<br>Iys arg val eile arg val asp phe asn val pro leu asp gly lys lys ile thr ser asn gin arg ile val ala ala leu pro thr<br>AAG CGT GTC TTC ATC AGA GTT GAC TTC AAC GTC CCA TTG GAC GGT AAG AAG ATC ACT TCT AAC CAA AGA ATT GTT GC 50<br>ile lys tyr val leu glu his his pro arg tyr val val leu ala ser his leu gly arg pro asn gly glu arg asn glu lys tyr ser<br>ATC AAG TAC GTT TTG GAA CAC CAC CCA AGA TAC GTT GTC TTG GCT TCT CAC TTG GGT AGA CCA AAC GGT GAA AG BO<br>leu ala pro glu val glu leu gln ser leu leu gly ly asp val thr phe leu asn asp cys val gly pro glu val glu ala ala<br>TTG GCT CCA GTT GCT AAG GAA TTG CAA TCA TTG GT AAG GAT GTC ACC TTC TTG AAC GAC TGT GTC GGT CCA GAA GTT G 110 110<br>- val lys ala er ala pro qly ser val ile leu leu glu arg tyr his ile glu glu glu gly ser arg lys val asp gly gln<br>- GTC AAG GCT TCT GCC CCA GGT TCC GTT ATT TTG TTG GAA AAC TTG CGT TAC CAC ATC GAA GAA GAA GGT TCC AGA 140 – 150<br>Iys val lys phe arg his glu leu ser ser leu ala asp val tyr ile asn asp ala phe gly thr<br>AAG GTC AAG GCT TCC AAG GAA GAT GTT CAA AAG TTC AGA CAC GAA TTG AGC TCT TTG GCT GAT GTT TAC ATC AAC GAT GCC TTC GGT ACC 170 – 170<br>ala his argala is ser ser met val gly phe asp leu agn argala ala gly phe leu leu glu lys glu leu lys tyr phe gly<br>GCT CAC AGA GCT CAC TCT ATG GTC GGT TTC GAC TTG CCA CAA CGT GCC GGT TTC TTG TTG GAA AAG GAA TTG AA ZOD<br>Iys ala leu glu asn pro thr arg pro phe leu ala ile ileu gly gly ala lys val ala asp lys ile gln leu ile asp asn leu leu<br>AAG GCT TTG GAG AAC CCA ACC AGA CCA TTC TTG GCC ATC TTA GGT GGT GCC AAG GTT GCT GAC AAG ATT CAA T 230 290<br>Asp lys valley in the physics of the Same the Pass lys valley all east the gluine gly asp ser the phe<br>GAC AAG GTC GAC TCT ATC ATC ATT GGT GGT GGT AGG THE ACC TTC AAG AAG GTT TTG GAA AAC ACT GAA ATC GGT GAC TCC ATC 260)<br>asp lys ala glu ile val pro lys leu met glu lys ala lys ala lys alw Yal gl<mark>u val val eu pro val asp phe ile ile</mark><br>GAC AAG GCT GGT GCT GAA ATC GTT CCA AAG TTG ATTG GAA AAG GCC AAG GCC AAG GGC GTC GTC TTG CCA GTC GAC TT 230 – 290<br>ala asp ala phe ser ala asp ala asn thr lys thr val thr asp lys glu gly ile pro ala gly trp gln gly leu asp asn gly pro<br>GCT GAT GCT TTC TCT GCT GAT GCC AAC ACC AAG ACT GTC ACT GAC AAG GAA GGT ATT CCA GCT GGC TAG (slu)320<br>ser arg lys ale thr val ale lys ale lys thr ile val tre asn gly pro pro gly val phe glu phe glu lys phe<br>FCT AGA AAG TTE GAA AAG TTE GAA AAG GCT AAG ACC ATT GIC TGG AAC GGT CCA CCA GGT GTT TTC GAA TTC GAA AAG TTC<br>3 ala ala gly thr lys ala leu leu asp glu val val lys ser ser ala ala gly asn thr val ile ile gly gly gly asp thr ala thr<br>GCT GCT GGT ACT AAG GCT TTG TTA GAC GAA GTT GTC AAG AGC TCT GCT GGT AAC ACC GTC ATC ATT GGT GGT GGT GA 380 - 400<br><mark>1914 and a lysile ser his value in the gly gly ala ser leu glu leu glu gly gly gly ala ser leu glu leu glu gly lys<br>1917 GCT AAG AAG TAC GGT GTC ACT GAC A<u>AG ATC T</u>CC CAT GTC TCT ACT GGT GGT GGT GCT TCT TTG GAA </mark> 410  $41\overline{6}$   $40$   $40$   $60$ pro gly val ala phe leu ser glu lys lys OC CCA GOT GTT GCT TTC TTA TCC 0A8 AA4 AAA TAA ATTGAATTGAATTGAAATCGATAGATCAATTTTTTTCTTTTCTCTTTCCCCATCCTTTACGCTAAAATAATAGTT 80 100 120 140 160 180 TATTTTATTTTTGATTTTTTTATTTATATACGTATATATAGACTATTATTTACTTTTAATAGATTATTAAGATTTTTATTAAAAAAAAAtTTCGT CC CTCTTTTTAATGCCTTTTAT 200 220 240 260 280 GCAGTTTTTTTTTCCCATTCGATATTTCTATGTTCGGGTTTCAGCGTATTTTAAGTTTAATAACTCGAAAATTCTGCGTTTCGAAAAAGCTT

Translated Mol. Weight - 44746

Fi gure 2. DNA and protein sequences of the yeast PGK gene. The sense strand of DNA sequence from PvuI to HindIII is shown from <sup>5</sup>' to 3'. The translated amino acid sequence is shown with amino acids underlined from 268 through 398 which agree with published protein sequence. Amino acids in parentheses designate differences in published protein sequence (47). Asterisks designate the use of nonpreferred codons for highly expressed yeast genes. The transcription start is shown by the arrow and the polyadenylation region by the bar region at the end of the gene.

since insertion of the gene on a yeast multicopy plasmid leads to very elevated PGK expression levels (see below).

However, the codon usage in the PGK gene is not consistent with one of the codon bias rules of Bennetzen and Hall (48), since one of the seven codons that are never used in the genes which they compared is used in PGK. This exception is the CGU codon for arginine which is used three times (positions 18, 122 and 182; Fig. 2). The CGU codon for arginine is also used in the yeast enolase genes (49). Another one of these seven codons, the CCG codon for proline, has been observed in the yeast TRP5 gene (50). Homology with the human and equine PGKs

The protein sequence for two other eukaryotic PGKs, the human (45) and horse (46) enzymes, has been elucidated. However, no corresponding DNA sequence is available. The horse and human PGK have a remarkable homology with only 14 amino acid differences, which probably resulted from single base changes in the gene. Another difference in the human PGK was the insertion of an extra lysine between amino acids 38 and 39 of the horse sequence. This results in a length of 416 amino acids for horse PGK versus a length of 417 amino acids for human PGK.

The yeast PGK protein sequence, which contains 415 amino acids (assuming the removal of the  $NH_2$ -terminal methionine), is compared with the human and equine PGKs in Fig. 4. The sequences are aligned for maximal homology.



Codon Usage:

Figure 3. Codon usage for the PGK gene. Amino acids for each codon are abbreviated with the numbers indicating frequency of use in the gene.  $OC$ , abbreviated with the numbers indicating frequency of use in the gene. AM, and OP refer to the translational stops ochre, amber, and opal, respectively.

One difference is the presence of an additional amino acid at the COOH-terminus of yeast PGK. In support of this sequence, Markland et al. (43) have reported that the COOH-terminal amino acid for yeast PGK is lysine

Comparative Protein Homology For Yeast, Human, and Equine PGK <sup>Y</sup> ser leu ser ser lys leu ser val gln asp leu asp leu lys asp lys Y 206 leu ala ile leu qly qly ala lys val ala asp lys ile gln leu ile I ser leu ser asn lys leu thr leu asp lys leu asp val lys gly lys H 209 leu Ica lIe leu qly ly sla lys eal aIa asD lys ile gIn leu ile 208 leu :la ile leu qly qly ala lys eal sic ase lys lie gln leue Ie <sup>E</sup> <sup>1</sup> ser leu ser asn lys leu thr leu asp lys leu asn val lys gly lys  $\epsilon$ <sup>Y</sup> 17 arg val phe ile arg eal asp phe asn val pro leu asp gly lys lys 222 asp asn leu leu asp lys val asp ser ile ile ile gly gly qly met H 17 arg val val met arg vel asp phe asn vel pro set lys esn asn gln H 225 asn asn et leu ase lys val asn qlu et ile ile gly qly gly met <sup>E</sup> 224 asn asn met leu :sp lys eal asn gelu et ile le glly giy gly cet t 17 arg val val met arg val asp phe asn val pro met lys asn asn gin 238 ala phe thr phe lys lys val leu qlu asn thr qlu ile gly asp ser Y 33 ile thr ser asn gln arg ile val ala ala leu pro thr ile lys <sup>H</sup> <sup>33</sup> ile thr asn asn gin arg lye ie lys aclaala val pro ser ile lys H 241 ala phe thr phe leu lys val leu asn asn et glu ile qly thr ser E 33 ile thr asn asn gin arg ile lys ala ala val pro ser ile lys E 240 ala phe thr phe leu lys val leu asn asn met qlu ile gly thr ser 48 tyr val leu glu his his pro arg tyr val val leu ala ser his leu Y 254 ile phe asp lys ala qly ala glu ile val pro lys leu met glu lys H 257 leu phe asp glu glu gly ala lys ile val lys asp leu met ser lys H49 phe cys leu asp asp gly ala lys ser val val leu met ser his leu <sup>E</sup> 256 leu phe ase 9lu glu gly la lys ile val lys asn leu cet ser lys t 48 phe cys leu asp asp gly ala lys ser val val leu met ser his leu Y 64 gly arg pro asn gly glu arg asn glu lys tyr ser leu ala pro Y 270 ala lys ala lys qly val qlu val val leu pro val asp phe ile ile <sup>H</sup> 65 gly crg pro asp gly val pro \*et proe asplys tyr ser leu glu pro <sup>E</sup> <sup>64</sup> gly srg pro asp vel gly pro met proe a lys tyr ser leu gin pro H 273 alaqlu lys asp qly val lys ile thr leu pro val asp phe val thr<br>E 272 alaqlu lys asn qly val lys ile thr leu pro val asp phe val thr 79 val ala lys glu leu gln ser leu leu gly lys asp val thr phe leu <sup>Y</sup> 286 ala asp ala phe ser ala asp ala asn thr lys thr val thr asp lys H81 val ala val glu leu lys ser leu leu gly lys asp val leu phe leu H 289 ala asp lys phe asp qlu asn ala lys thr qly qlu ala thr val ali <sup>E</sup> 80 val ala vel glu leu lys ser leu leu gly lys asp val leu phe leu <sup>E</sup> 288 ala asp lys phe asp qlu his ala lys thr qly qln ila thr eel sic <sup>Y</sup> 95 asn asp cys vel gly pro glu val glu ala ala val lys ala ser ala <sup>Y</sup> 302 qlu qly ile pro asa gly tre qln qly leu asep sn gly pro qlu ser H 305 ser qlyile pro ala qlytro met qlyleu asp cys qlypro glu ser<br>E 304 ser qlyile pro ala qlytrp met qlyleu asp cys qlythr qlu ser<br>E 304 ser qlyile pro ala qlytrp met qlyleu asp cys qlythr qlu ser H97 lys asp cys val gly pro glu val glu lys ala cys ala asp pro ala E96 lys asp cys val gly pro glu val glu lys ala cys ala asp pro ala <sup>Y</sup> 111 pro gly ser val ile leu leu glu asn leu arg tyr his ile glu glu 318 arq lys leu phe ala ala thr val ala lys ala lys thr ile val trp H 113 als gly ser val ile leu leu glu asn leu arg phe his val glu glu <sup>H</sup> 321 ser lys lys tyr ala glu ala val thr arg ala lys gln ile val tre <sup>E</sup> 112 ali gly ser vel ile leu leu glu asn leu arg phe his eal glu glu E 320 ser lys lys tyr ala qlu ala val ala arq ala lys gln ile val tro r 126 glu gly ser arg lys val asp gly gin lys val lys ali ser lys <sup>Y</sup> 334 asn qly pro oro qly val phe qlu phe qlu lys phe sla ala gly thr 129 glu gly lyesgly lys asp asi ser gly asn lys val lys asi glu pro 128 glug lye gly Iys asp sic see gly ssn lys cal lys iac glu pro E H 337 ase qly pro val gly val phe qlu trp qlu ala phe sla erq qly thr <sup>E</sup> 336 asn qly pro val qly eal phe qlu tre qlu sic phe ias arq gly thr 142 glu asp val gln lys phe arg his glu leu see ser leu aia asp val × Y 350 lys ala leu leu ase qlu val val lys ser ser &la la qly asn thr 145 ala lys ile glu ala phe arg ala ser leu ser lys leu gly asp val H 353 lys ala leu met asp qlu val val lys ala thr ser arg gly cys ile<br>E 352 lys ala leu met asp qlu val lys ala thr ser arg gly cys ile<br>E 352 lys ala leu met asp qlu val lys ala thr ser arg gly cys ile n 145 alalys ile glu alaphe arg alaser leuser lys leu gly asp val<br>El 44 alalys ile glu thr phe arg alaser leuser lys leu gly asp val 158 tyr ile asn asp ala phe gly thr ala his arg ala his ser ser met Y 366 wal lie ile qly qly qly ase thr ala thr vel ale lys lys tyr qly H 161 tyr eal asn asp ali phe gly thr ala his arg aia his ser ser met <sup>H</sup> 369 thr ile ile qly qly qly se thr aia thr cys cys sla lys trp asn E 160 tyr val asn asp ala phe gly thrailahis argalahis ser ser met E 368 thr ile ile qly qly qly asp thr ala thr cys cys ala lys trp asn 174 val gly phe asp leu pro gin arg ala ala gly phe leu leu glu lys Y 382 val thr asp lys ile ser his val ser thr gly gly gly ala ser leu <sup>H</sup> <sup>385</sup> thr qln asp lys val ser his val ser thr gly qly glyla ser leu <sup>E</sup> <sup>384</sup> thr qlu asp lys vel ser his el ser thr qly gly elyla ser leeu <sup>H</sup> 177 eal gly val asn leu pro gIn lys als gly gly phe leu set lys lys <sup>E</sup> 176 eel gly eel csn leu pro gIn lys asi gly gly phe leu met lys lys <sup>Y</sup> 190 glu leu lys tyr phe gly lys ala leu glu asn pro thr arg pro phe <sup>Y</sup> 398 qlu leu leu qlu qly lys qlu leu pro qly val sic phe leu ser glu <sup>H</sup> 193 glu leu asn tyr phe asa lys sia leu glu ser pro glu srg pro phe H 401 qlu leu leu qluqly lys val leu pro qly val asp ala leu ser asn<br>E 400 qlu leu leu qluqly lys val leu pro qly val asp ala leu ser asn<br>E 400 qlu leu qluqly lys val leu pro qly val asp ala leu ser asn E 192 glu leu asn tyr phe ala lys ala leu glu ser pro glu arg pro phe 414 lys lys H 417 ile <sup>E</sup> 416 val

Fi gure 4. Comparative protein homologies for yeast, human, and equine PGKs. Y, H, and E refer to yeast, human, and equine (horse), respectively. Sequences are lined up for greatest homology with squares showing the insertion of an amino acid in one sequence with respect to another. However, the insertions with respect to yeast sequence occur within regions 69 to 72 and 128 to 132 of yeast sequence; thus, the amino acid insertion shown is arbitrary. Asterisks indicate homology.

(see Fig. 2). Other major changes are the insertion of an amino acid in the human and horse sequence in the region of positions 69 to 72 and 128 to 132 of the yeast PGK gene. An additional amino acid insertion in human PGK with respect to yeast PGK occurs between residues 38 and 39 of the yeast sequence. Interestingly, this is identical to the lysine insertion in human PGK with respect to horse PGK. The yeast PGK has about 65 percent homology with both the human and horse sequence. Although this homology is spread all over the sequence, it is clear that some regions, e.g. residues 160 to 175 and 203 to 221, are strongly conserved. This might implicate an important role of such segments for the enzymatic function.

# The PGK mRNA

Polyadenylated or total RNA from S. cerevisiae GM3C-2 containing either YEp13 or PGK-YEp13 was sized on a formaldehyde-agarose gel and "Northern" hybridization (36) was performed using a PGK-gene specific probe. YEp13 (21) is a high copy number,  $2<sub>u</sub>$  origin based plasmid; while PGK-YEp13 contains the PGK HindIII fragment between the two HindIII sites of YEp13. It should further be noted that these haploid yeast contain one copy of this PGK HindIII fragment in chromosome III (4) which is also producing mRNA. As shown in Fig. 5 a single mRNA band of about 1500 bases long was observed. The size of the PGK mRNA from PGK-YEpl3/GM3C-2 is identical to the chromosomal PGK rnRNA, isolated from YEpl3/GM3C-2. The intensity of the former PGK mRNA is obviously much stronger (10-20X), since the gene and the flanking regions are situated on a high copy number plasmid. These levels



2 Figure 5. PGK gene mRNA identification. Lane 1 is total RNA from yeast strain GM3C-2 containing plasmid PGK-YEp13. Lane 2 is total RNA from yeast strain bases GM3C-2 with plasmid YEp13 (no PGK gene present on the second plasmid). The mRNAs with PGK sequence were -3360 visualized by hybridization with ---labeled <u>EcoRi/B</u><br>fragment from the PGK gene (see Fig. 1) followed by autoradiography. Size standards are designated (see Methods) .

of mRNA correspond well with PGK-YEp13 expression of PGK at about 20 percent of the total protein as compared to <sup>1</sup> percent of the total protein by the chromosomal copy of the gene in the other yeast strain (data not shown). These results strongly suggest that all control signals needed for transcription of the PGK gene are contained within the 3.1 kb HindIII fragment, situated on PGK-YEp13.

#### Transcriptional initiation

The start of transcription of the PGK gene was determined by specifically primed cDNA synthesis on the mRNA from yeast pFRM31/20B-12. Plasmid pFRM31 contains the 3.1 kbp PGK HindIII fragment in the HindIII site of the pBR322 portion of pFRD7 (see Methods), which contains the yeast arsl chromosomal origin of replication and the yeast TRP1 gene (22, 23). The extension started from an oligonucleotide complementary to position -10 to -21 (Fig. 2) and gave a distinct product of 26 base pairs long (Fig. 6). The initiation of transcription can therefore be located at -36 (Fig. 2). It is possible that the observed 5' end is the result of post-transcriptional processing (15) and that the transcription start is further upstream. No evidence for such 5' terminal mRNA processing in yeast is available. This initiation of transcription takes place at an AG-duplet, which is also observed for the two transcripts of yeast ADH-1 (14). This seems, however,



Fi gure 6. Determination of the 5 transcription initiation of the<br>transcription initiation point of the PGK gene. A discrete product was obtained from specifically primed cDNA synthesis. This extension product, shown with an arrow in lane 5, is sized on a 15 percent polyacrylamide-7M urea gel alongside an unrelated sequencing ladder, obtained by the chemical degradation method (lanes 1-4). The length of the sequencing reaction products is shown on the left.

not to be the case in other genes (11,50). Only one discrete cDNA extension product is observed, implicating a single transcription start. This is in clear contrast to an apparent multiplicity of initiation points for several other yeast genes such as TRP5 (50), ADH-1 (14) and CYCl (11). The mechanism behind this phenomenon is unclear.

### Transcription termination and polyadenylation site

The 286 bp region following the stop codon of the yeast PGK gene has also been sequenced (Fig. 2). We used the  $SI$ -mapping procedure (40, 41) to determine the polyadenylation site of the mRNA. The BglII-HindIII PGK-terminator fragment,  $3'$   $3^2$ P-labelled at the BglII site served as hybridization probe to mRNA from pFRM31/20B-12. As shown in Fiq. 7, there is a heterogeneity in the length of the DNA fragments, protected from Sl-nuclease activity, ranging from 180 to 187 nucleotides. This implies that the polyadenylation site of the PGK mRNA is located at 86 to 93 nucleotides behind the stop codon (Fig. 2). The heterogeneity in the length of the protected DNA fragments migit be due to an overdigestion by Sl-nuclease or may result from a slight heterogeneity in the polyadenylation site. Multiple polyadenylation sites have been seen for several other eukaryotic mRNAs, including the yeast ADH-1 mRNA (14) and the yeast actin mRNA (52). Again steady state levels of mRNA were used so it cannot be concluded that this defines the actual transcription termination site for PGK mRNA. Transcription may proceed further followed by processing before polyadenyl ation.



M 1 2 Figure 7. S1-nuclease mapping of the polyadenylation site of the PGK mRNA. The RNA-DNA W https://with the contract of the PGK mRNA. The RNA-<br>hybrids were treated with 50 units SI nuclease<br>hybrids were treated with 50 units SI nuclease bases for 30 min (lane 1) or 100 units Si nuclease for 1 hr (lane 2) and then sized on a 6 percent polyacrylamide-7M urea gel. The lane M shows the <sup>32p-labelled</sup> HpaII fragments of pBR322 with  $190 - w$  their sizes indicated on the left.

## DISCUSSION

### PGK coding region and protein sequence

The PGK coding sequence is 1248 bp corresponding to 416 amino acids. Like other highly expressed genes from yeast, this gene is highly codon biased using only 25 codons out of 61 for 95 percent of the amino acids. Some less biased codons are also used; however, this does not appear to limit expression, since the PGK gene expresses up to 20 percent of the total protein when present on a high copy number plasmid. The amino acid sequence is identical with 120 out of 130 previously published residues (47).

The entire amino acid sequence of yeast PGK, as determined from DNA sequence, is 65 percent homologous with horse and human PGKs. Previous x-ray crystallographic data demonstrate two major domains in both the horse (46) and yeast (53, 54) enzymes. Furthermore, such data suggest that the mechanism of PGK enzyme activity is the contact between these two domains to bring the domains binding the two substrates, ATP and 3-phosphoglycerate, together (46). For horse PGK, ATP binds to the carboxyl domain (residues 190 to 403) and 3-phosphoglycerate binds to the amino terminal domain (residues 1-185) while the rest of the carboxyl end interacts with the amino terminal domain. Two links between these domains are residues 186-189 and residues 404-406. Yeast PGK appears to be very similar. Our protein sequence shows the greatest homology from residues 133 through 413 encompassing the two links between domains. The most severe differences are codon insertions in human and horse PGK sequences from residues <sup>1</sup> through 128 with respect to the yeast sequence. Twenty-four of 25 amino acids which are thought to be involved in substrate binding in the horse enzyme (46) are identical in the yeast enzyme. Homology between the yeast and horse enzymes both coincides and falls in between  $\alpha$ -helical and  $\beta$ -sheet structures in horse PGK (46); suggesting regions between these structural components are as important to enzyme function as the defined structural components themselves.

# PGK gene 5'-flanking sequence

The Goldberg-Hogness box or TATAAA-consensus sequence is generally found 25 to 30 nucleotides upstream from the RNA start in higher eukaryotes or eukaryotic viruses and seems to be important in the positioning of the transcription start (for review see refs. 55 and 56). A TATAAA-sequence is here found 145 bp in front of the PGK-gene translational start (Fig. 2) (109 basepairs in front of the mRNA start at -36). However, although a TATAAAsequence is present in front of the transcribed sequence of yeast genes, the

distance from the transcriptional initiation is longer and more variable than in higher eukaryotes (11,14,50,52). No unambiguous evidence for the functional importance of this sequence in yeast is as yet available.

In between the TATAAA-sequence and the transcription starts of the ADH-1 gene is a very pyrimidine-rich cluster (14) and it has been proposed that this might serve as <sup>a</sup> signal for high level transcription (9). A similar C-T rich stretch is present in the <sup>5</sup>' flanking region of the PGK gene at -70 to -52. This feature cannot, however, be clearly distinguished in front of the yeast genes for the other glycolytic enzymes, enolase (49) and glyceraldehyde-3-phosphate dehydrogenase (3).

Another potentially important sequence is located at -48 to -28. As noticed by Dobson et al. (47), who recently reported the <sup>5</sup>' flanking sequence of the PGK gene (largely corresponding to our sequence) the above-mentioned region has a similarity with the corresponding sequence for the enolase (49) and glyceraldehyde-3-phosphate dehydrogenase (3) genes. Since there is, however, no clear homology with the ADH-1 gene (14), it is doubtful if this is a typical feature of genes for the glycolytic enzymes in yeast.

The sequence 5'-PuCACACA-3' precedes by 4 to 15 residues the initiation codon of several yeast genes like CYCl (8), histone H2B1 (57), glyceraldehyde-3-phosphate dehydrogenase (3), enolase (49), TRP1 (23) and TRP5 (50). Although there is no evidence for a specific function, it has been suggested that this sequence might play <sup>a</sup> role in the initiation of translation (50). A variation of such a sequence is present in front of the initiation codon of the ADH-1 gene as 5'-PuCAATCAA-3' (-15 to -22) (14). A similar, but not identical, sequence (5'-PuCAACAA-3') is present 10 residues prior to the PGK gene (Fig. 2). However, the expression of human leukocyte interferon D cDNA in a hybrid yeast expression system demonstrates that up to 33 nucleotides of 5'-sequence adjacent (from yeast alcohol dehydrogenase <sup>I</sup> gene) to the ATG can be deleted without eliminating expression; suggesting that this sequence containing similarities among different yeast genes is not critical to expression (66).

A current widely accepted view of eukaryotic translation initiation is that the 40S ribosomal subunit binds at or near the <sup>5</sup>' end of the mRNA and moves along the mRNA until it encounters the first AUG (58-60). The efficiency of ribosome binding might be enhanced by <sup>a</sup> complementarity with the 3' end of the 18S ribosomal RNA. Such a complementarity has been noticed for several eukaryotic mRNAs (61, 62). Similarly, the sequence 5' TMTTATC 3' at -34 to -27, just behind the transcription initiation site of the yeast PGK gene, has a clear complementarity to the 3' end of the yeast 18S ribosomal RNA <sup>3</sup>' AUUACUAG 5' (63). A similar complementarity has been observed for several other yeast mRNAs, as for TRP5, by Zalkin and Yanofsky (50).

The efficiency of the ribosome binding to the mRNA is likely to be influenced by the sequence immediately preceding the ATG-codon as proposed by Kozak (60). From in vitro ribosome binding studies (60) and from in vivo studies with various sequences in front of the start codon (R.D., manuscript in preparation), it can be concluded that the position of an A at -3 is very favorable for translation initiation. This feature is observed in the yeast PGK mRNA and similarly in almost all other known yeast mRNAs (64). One exception to this apparent rule for yeast is a revertant of iso-l-cytochrome c which has a T at -3 and expresses the same level of enzyme as the wild type gene with an A at -3 (65). Furthermore, the relatively high expression of leukocyte interferon D, by a constructed hybrid expression system with a T residue at -3, also suggests that the absence of A at -3 might not grossly affect translation (66).

# PGK gene 3'-flanking sequence

On the basis of the determined transcription start and of the location of the polyadenylation site (86 to 93 nucleotides from the translation stop), it can be concluded that the size of the PGK transcript is about 1380 bases. This is compatible with the observed length of 1500 nucleotides of the polyadenylated PGK mRNA, since polyadenylation adds about 50-100 bases to the length (67). The mRNA size, considered in combination with the size of the PGK coding sequence and protein, suggests that there are no introns within the PGK coding region which is consistent with other yeast glycolytic genes (3,14,49).

The polyadenylation site of the mRNAs of higher eukaryotes is preceded 20 to 26 nucleotides by <sup>a</sup> <sup>5</sup>' AAUAAA <sup>3</sup>' (68). Although some yeast mRNAs (7,17,52,69) also contain this sequence, most others (17) clearly lack this signal sequence. Bennetzen and Hall (48) recently proposed that a related consensus sequence 5' TAAATAA<sub>G</sub> 3' might play a role in the transcription termination in yeast. This sequence or a variant of it precedes the polyadenylation site by 28 to 34 nucleotides. However the comparison of the 3' terminal sequences of yeast mRNAs by these authors shows that a relatively wide variation is present, which makes its significance less convincing. A possibly related sequence in PGK is present at 64 to 71 nucleotides behind the stop codon.

Al ternatively, Zaret and Sherman (17) proposed a sequence 5' TAG...TA(T)GT...TTT <sup>3</sup>' as being important for transcription termination. This sequence or a variant precedes the polyadenylation site of most yeast mRNAs at variable distance. At 58 to 88 nucleotides behind the stop codon, a related sequence can be recognized.

Much attention is being focused on the regulation of transcription and expression in yeast. As illustrated above with the determined structure of the yeast PGK gene and mRNA, some homologies and relations in the sequences upstream and downstream of the coding regions can be observed. However, the expected functional importance of these sequences is mainly based on structural comparisons and speculations. It is obvious that much more research is needed, which will undoubtedly lead to more insight into the structural requirements for these processes in yeast.

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