

Eukaryotic Multi-subunit DNA dependent RNA Polymerases: An Insight into their Active Sites and Catalytic Mechanism

Abstract

Aim: To analyze the most complex multi-subunit (MSU) DNA dependent RNA polymerases (RNAPs) of eukaryotic organisms and find out conserved motifs, metal binding sites and catalytic regions and propose a plausible mechanism of action for these complex eukaryotic MSU RNAPs, using yeast (*Saccharomyces cerevisiae*) RNAP II, as a model enzyme.

Study Design: Bioinformatics, Biochemical, Site-directed mutagenesis and X-ray crystallographic data were analyzed.

Place and Duration of Study: School of Biotechnology, Madurai Kamaraj University, Madurai, India, between 2007- 2013.

Methodology: Bioinformatics, Biochemical, Site-directed mutagenesis (SDM) and X-ray crystallographic data of the enzyme were analyzed. The advanced version of Clustal Omega was used for protein sequence analysis of the MSU DNA dependent RNAPs from various eukaryotic sources. Along with the conserved motifs identified by the bioinformatics analysis, the data already available by biochemical and SDM experiments and X-ray crystallographic analysis of these enzymes were used to confirm the possible amino acids involved in the active sites and catalysis.

Results: Multiple sequence alignment (MSA) of RNAPs from different eukaryotic organisms showed a large number of highly conserved motifs among them. Possible catalytic regions in the catalytic subunits of the yeast Rpb2 (= β in eubacteria) and Rpb1 (= β' in eubacteria) consist of an absolutely conserved amino acid R, in contrast to a K that was reported for DNA polymerases and single subunit (SSU) RNAPs. However, the invariant 'gatekeeper/DNA template binding' YG pair that was reported in all SSU RNAPs, prokaryotic MSU RNAPs and DNA polymerases is also highly conserved in eukaryotic Rpb2 initiation subunits, but unusually a KG pair is found in higher eukaryotes including the human RNAPs. Like the eubacterial initiation subunits of MSU RNAPs, the eukaryotic initiation subunits, viz. Rpb2, exhibit very similar active site and catalytic regions but slightly different distance conservations between the template binding YG/KG pair and the catalytic R. In the eukaryotic initiation subunits, the proposed catalytic R is placed at the -9th position from the YG/KG pair and an invariant R is placed at -5 which are implicated to play a role in nucleoside triphosphate (NTP) selection as reported for SSU RNAPs (viral family) and DNA polymerases.

Similarly, the eukaryotic elongation subunits (Rpb1) are also found to be very much homologous to the elongation subunits (β') of prokaryotes. Interestingly, the catalytic regions are highly conserved, and the metal binding sites are absolutely conserved as in prokaryotic MSU RNAPs. In eukaryotes, the template binding YG pair is replaced with an FG pair. Another interesting observation is, similar to the prokaryotic β' subunits, in the eukaryotic Rpb1 elongation subunits also, the proposed catalytic R is placed double the distance, i.e., -18 amino acids downstream from the FG pair unlike in the SSU RNAPs and DNA polymerases where the distance is only -8 amino acids downstream from the YG pair. Thus, the completely conserved FG pair, catalytic R with an invariant R, at -6th position are proposed to play a crucial role in template binding, NTP selection and polymerization reactions in the elongation subunits of eukaryotic MSU RNAPs. Moreover, the Zn binding motif with the three completely conserved Cs is also highly conserved in the eukaryotic elongation subunits. Another important difference is that the catalytic region is placed very close to the N-terminal region in eukaryotes.

Conclusions: Unlike reported for the DNA polymerases and SSU RNA polymerases, the of eukaryotic MSU RNAPs use an R as the catalytic amino acid and exhibit a different distance conservation in the initiation and elongation subunits. An invariant Zn²⁺ binding motif found in the Rpb1 elongation subunits is

50 proposed to participate in proof-reading function. Differences in the active sites of bacterial and human
51 RNA polymerases may pave the way for the design of new and effective drugs for many bacterial
52 infections, including the multidrug resistant strains which are a global crisis at present.

53 **Keywords:** *Multi-subunit DNA dependent RNA polymerases, Eukaryotic RNA polymerases, RNA*
54 *polymerase II, Saccharomyces cerevisiae, Conserved motifs, Polymerase active site, Polymerization*
55 *mechanism, Transcription slippage diseases, Drug design.*

56

57 1. INTRODUCTION

58 RNAPs (EC 2.7.7.6) are key enzymes which play a vital role in the flow of genetic information
59 from DNA to proteins and the proteins are final players in all cellular processes. Therefore, they are found
60 in all living organisms and play a crucial role in copying DNA sequences into RNA molecules which are
61 subsequently translated into proteins. Thus, transcription forms the first step and a key control point in
62 gene regulation and expression. Errors in the transcription process can potentially lead to aberrant gene
63 products and ultimately lead to various diseases including cancer. One major class of transcription error,
64 known as transcriptional slippages, are implicated in the development of a wide variety of diseases, such
65 as colon cancer, non-familial Alzheimer's, Down's syndrome, etc. [1]. Recently the *in vitro* transcribed
66 mRNAs have come into focus as a potential new class of drugs known as 'mRNA therapeutics' to
67 deliver genetic information through mRNAs to correct the malfunction(s) [2] and also potential vaccines
68 against cancer treatment [3]. In fact, many anti-cancer drugs act by inhibiting the transcription step
69 itself. Therefore, understanding the mechanism and regulation of RNAPs have been a key goal of
70 molecular biologists since its discovery. Interestingly, the sequences, overall 3D framework and functions
71 of these DNA dependent RNAPs are universally conserved in viruses, bacteria, plants and animals with
72 small but significant differences in their active sites and catalytic amino acids. For the discussion about
73 the discovery and dynamics of the polymerization reactions [refer to 4]. Instead of a single type of RNAPs
74 in prokaryotes, Robert Roeder and William Rutter discovered in 1969 the existence of three different
75 RNAPs in eukaryotic cells that were responsible for transcription of all types of RNAs in the eukaryotic
76 cells [5].

77 Though RNAPs are found in all organisms, their number and composition vary across taxa,
78 possibly due to evolutionary consequences, changes in the genome structure and the complexity of the
79 transcription process. However, the MSU RNAPs display a conserved core structure across all major
80 domains of life, viz. viruses, bacteria, archaea and eucarya. For example, viruses contain mainly two
81 different RNAPs, viz. DNA dependent RNAPs and RNA dependent RNAPs. Both eubacteria and
82 archaeobacteria contain a single type of MSU RNAPs, while eukaryotes contain at least five distinct types
83 of MSU RNAPs (I-V). While the eubacterial enzymes are composed of 5 different subunits, the eukaryotic
84 enzymes are made up of up to 12-16 different subunits. Despite such differences, there are striking
85 similarities among the transcriptional mechanisms by various types of RNAPs across the three major
86 domains of life [6 and references therein].

87 Transcription and transcriptional regulation are very important in eukaryotes as it underlies all
88 aspects of cellular metabolism including oncogenesis (cancer) and morphogenesis (development). The
89 major type of RNA polymerase, i.e., the RNA polymerase II, which involves in the transcription of genes,
90 is a large (550 kDa) complex made up of 12 subunits. Unlike in prokaryotes, gene expression, and
91 therefore, RNAP II activity is controlled by several proteins known as general transcription factors. In
92 other words, as there are no Sigma like factors in eukaryotic cells for promoter recognition and to initiate
93 transcription, the eukaryotic RNAPs interact with a variety of protein factors known as general
94 transcription factors. Large volumes of genomic and protein sequence data are available for most of the
95 prokaryotic and eukaryotic MSU RNAPs. Understanding the biological principles buried in these
96 sequencing data is a significant challenge for scientists. Therefore, the purpose of the study is to analyze
97 the available protein sequences of the MSU RNAPs from eukaryotes to find out the highly conserved
98 motifs among them and to understand the structure- function relationships and mechanism of action of
99 these key enzymes, along with the X-ray crystallographic and experimental data available on these
100 enzymes from various sources. A comparative analysis of the bacterial and human RNAPs for their
101 transcription mechanism will pave way to design new and effective drugs for many bacterial infections,
102 including the antibiotic resistance, especially the multi-drug resistance, which has become a global crisis
103 [7, 8 and references therein]. MSA has been proven to be very useful for assigning a function to a given

104 sequence, by comparing the existing experimental data with the protein sequence data and thus, bridging
105 the gap between the protein sequence and experimental data.

106 RNAPs catalyze the chemical reaction that synthesizes an RNA strand from a DNA template with
107 all the 4 NTPs and a metal ion, usually a Mg^{2+} ion. RNA synthesis in eukaryotes involves four steps, viz.
108 formation of pre-initiation complex, initiation, elongation and termination. The nucleotides are added one
109 at a time to the growing 3' end. The newly formed RNA copies serve as blueprints for the synthesis of
110 proteins during the next step of translation. The basic transcription unit is the distance between the sites
111 of transcription start site (TSS) and transcription termination site (TTS), and may have one or more genes
112 between them (e.g., mono or poly-cistronic mRNAs, poly-cistronic mRNAs are uncommon in eukaryotes).

113 2. Types of MSU DNA dependent RNAPs of prokaryotes and eukaryotes

114 There are at least 4 different types of MSU RNAPs in living cells, viz.

- 115 1) MSU DNA dependent RNAPs of eubacteria
- 116 2) MSU DNA dependent RNAPs of archaeobacteria
- 117 3) MSU DNA dependent RNAPs of chloroplasts (Plastid encoded)
- 118 4) MSU DNA dependent RNAPs of eukaryotes

119 2. 1 Types of MSU DNA dependent RNAPs in Eukaryotes

120 As mentioned elsewhere, bacteria and archaea have only a single RNAP to transcribe all of its
121 genes whereas eukaryotic cells employ different RNAPs to transcribe different types of genes, viz.
122 rRNAs, tRNAs and mRNAs (Table 1).

123 There are at least 7 major types of RNAPs in eukaryotes. The nuclear genome utilizes three
124 major types of RNAPs which are localized in the nucleus. For example, the RNAP I is localized in the
125 nucleolus and primarily involves in the synthesis of the rRNAs and the other two RNAPs, viz. RNAPs II
126 and III are localized in the nucleoplasm and mainly involved in the synthesis of mRNAs and tRNAs,
127 respectively (Table 1). Plants are unique among eukaryotes in having five nuclear MSU RNAPs. Two
128 plant-specific RNAPs, polymerases IV and V are 12-subunit enzymes that are evolved as specialized
129 forms of Pol II. (Pols IV and V are nonessential for viability but play important roles in RNA-mediated
130 gene silencing pathways that tame transposons and defend against invading viruses [9]).

131 Table 1 Types and functions of eukaryotic RNAPs

Type of RNA Polymerase	Type of RNA(s) synthesized
1. RNA Polymerase I (Nucleolus)	rRNA genes (5.8S, 18S, 28S rRNAs from 45S pre-rRNA)
2. RNA Polymerase II (Nucleoplasm)	mRNAs, snRNAs, microRNAs
3. RNA Polymerase III (Nucleoplasm)	tRNAs & 5S rRNA, scRNAs, U6 SnRNA
4. RNA Polymerase IV (Plant specific)	siRNAs in plants
5. RNA Polymerase V (Plant specific)	Plant specific RNAs involved in siRNA directed heterochromatin formation in plants.
6. MSU type (Chromosome encoded)	Eubacterial and Chloroplasts
7. SSU type (Plasmid & Nucleus encoded)	Mitochondrial and Chloroplasts

132

133 2.2 Basic structure and composition of the eukaryotic RNAPs

134 All the 5 eukaryotic enzymes (RNAPs I-V) are MSU enzymes. The 12-subunit RNAP II is the
135 enzyme largely responsible for transcription of protein-encoding genes and thus, forms the central
136 component of the eukaryotic transcription machinery. Similarities between the eukaryotic and prokaryotic
137 MSU enzymes, suggest a common lineage in the evolutionary tree. For example, counterparts for all the
138 five core eubacterial RNAP subunits ($\alpha_2\beta\beta'\omega$) are found in the eukaryotic RNAPs I, II and III [10] (Table
139 2). The five orthologs include the two largest catalytic subunits Rpb1 and Rpb2, which correspond to the
140 bacterial β' and β subunits, respectively [11]. Furthermore, Rpb3 and Rpb11 correspond to the two copies

141 of the bacterial α subunit, and the Rpb6 subunit corresponds to the bacterial ω subunit [12]. In addition to,
 142 the two large subunits of RNAP II share some sequence homology and antigenic determinants with the
 143 corresponding subunits of RNAPs I and III [13] (Refer to also Mix and Match analysis Figs. 5 and 6).
 144 Rpb3 is involved in RNAP II assembly. The subunits Rpb4 and Rpb7 form a heterodimer and associate
 145 reversibly with the '10-subunit core polymerase' and involves in the initiation process which is otherwise
 146 defective in initiation. Rpb5, an evolutionarily highly conserved, universal eukaryotic RNAP subunit,
 147 shared by all three enzymes, facilitates communication between the RNAP core with a variety of basal
 148 and gene-specific transcription factors [14]. Rpb7 is essential for the functioning of the RNAP-II as
 149 deletion is also found to be lethal. A cleft is formed by jaws; the upper jaw is formed by regions of Rpb1,
 150 Rpb2 and Rpb9 whereas the lower jaw is formed by Rpb1 and Rpb5. The jaws are thought to grab the
 151 incoming DNA template [15] (Fig. 1).

152 Consistent with the increased complexity of the eukaryotic genome, all three major RNAPs of the
 153 eukaryotic transcriptional machinery have several additional subunits that do not have bacterial
 154 counterparts. Table 2 shows the subunit compositions of the three major types of nuclear MSU RNAPs.
 155 Apart from the five orthologs, as discussed above, the nuclear RNAPs share four common subunits, while
 156 the remaining subunits are RNAP dependent. Thus, the three major classes of eukaryotic RNAPs: I, II
 157 and III, in addition to comprising of two large catalytic subunits are also made up of 10-14 smaller
 158 subunits (Table 2).

159 Table 2 Composition of the three major eukaryotic nuclear RNAPs

Features	Pol I	Pol II	Pol III
Subunits unique to each RNAP ($\alpha_2\beta\beta'\omega$ - like)	5	5*	5
Common subunits to all 3 RNAPs	4	4	4
Additional unique subunits to each RNAPs	5	3	7
Total No. of subunits	14	12	16
Products	pre-rRNAS (45S RNA \rightarrow 28S, 5.8S, 18S).	pre-mRNAs 5 snRNAs [^] Sno RNAs, microRNAs.	pre-tRNAs 5S & 7S RNAs [§] , U6-snRNA.
Sensitivity to α -Amanitin	Nil	High (1 μ g/ml)	Moderate (10 μ g /ml)
Sensitivity to Actinomycin-D#	0.05 μ g/ml	0.5 μ g/ml	5.0 μ g/ml

160 **NB:**

161 *The largest subunit, Rpb1, has unique Carboxy Terminal Domain (CTD); # [16]

162 Subunit nomenclatures RNAPs I, II & III: RP A1-A14; RP B1-B12; RP C1-C16

163 [^] U1-U5 of ~200 bases; involves in the formation of spliceosomes

164 [§] 7S RNA from the signal recognition particle (SRP) which is involved in the transport of proteins into the
 165 endoplasmic reticulum.

166

167 2.3 Characteristics of the C-terminal domain (CTD) in the largest subunit Rpb1

168 Though exact functions of all the eukaryotic RNAPs are more or less completely understood, yet
 169 much data are not available on the individual subunits except for the larger subunits. An interesting
 170 feature of the RNAP II, a 12-subunit complex, is its unique CTD in the largest subunit Rpb1. The Rpb1
 171 elongation subunit consists of a tandem repeat of a conserved heptapeptide repeat sequences (-
 172 **YS²PTS⁵PS⁷**-). These heptapeptide repeats are completely conserved in sequence and function from
 173 yeast to human with varying lengths from 25 to 52. (The CTD domain does not exist in RNAPs I and III
 174 suggesting its special importance to mRNA processing). The Ser residues are phosphorylated and
 175 dephosphorylated during the active transcription process. Therefore, the RNAP-II exists in two forms, i.e.,
 176 unphosphorylated and phosphorylated forms (II and II^P), respectively [17]. In fact, the transition between
 177 the two forms facilitates different functions during transcription. (The phosphorylation of CTD is catalyzed
 178 by TFII-H, one of the six general transcription factors associated with RNAP II. TFII-H plays a dual role:
 179 one is to unwind the DNA at the transcription start site and the other is to phosphorylate the heptapeptide
 180 repeats. (TFII-H is a large protein complex that contains among others the Cdk7/Cyclin-H kinase complex
 181 for phosphorylation of CTD and an ATP dependent DNA helicase to unwind the DNA and open up the

182 transcription bubble at TSS). It also involves in transcription-coupled DNA mismatch repair. (Mutations in
183 the human XPD kinase cause *Xeroderma pigmentosum* and *Trichothiodystrophy* [18].

184 The elongation of initiation is accomplished by the phosphorylation of Ser⁵ of the heptapeptide, (-
185 **YS²PTS⁵PS-**) by the TFII-H. The Ser⁵ phosphorylation recruits enzymes to cap the 5' end of the newly
186 synthesized mRNA and the 3' processing factors to poly(A) sites. Once the second Ser is phosphorylated,
187 i.e., Ser², elongation is activated. West and Corden have shown substitution of Ala or Glu for Ser in
188 positions 2 or 5 is lethal. In addition, changing Tyr in position 1 to Phe is also lethal [19]. In order to
189 terminate elongation, dephosphorylation is accomplished by phosphatases. Once the domain is
190 completely dephosphorylated, the RNAP II is "recycled" and catalyzes the same process with another
191 initiation site. Thus, the CTD acts as a platform for various transcription factors as it binds or dissociates
192 them, depending upon their requirements during the transcription process. Thus, the phosphorylation and
193 dephosphorylation of the CTD is an important regulatory mechanism exhibited only by the RNAP II [20].
194 Ser7 phosphorylation is required for the transcription of SnRNAs and a mutation of Ser7 to Ala causes a
195 specific defect in snRNA expression [21].

196 The number of heptapeptide repeats in the CTD increases with genomic complexity; for example,
197 17 in *Plasmodium*, 26 in yeast, 32 in *Caenorhabditis elegans*, 45 in *Drosophila*, and 52 in mammals). The
198 CTD deletion experiments have shown the CTD's central role in coupling transcription to all three of the
199 main mRNA processing events [22]. For example, deletion of most of the CTD can result in inefficient
200 capping, splicing, and polyadenylation *in vivo*. SDM experiments of the yeast RNAP II has found that at
201 least 10 repeats are needed for the viability of the process [23, 24].

202 **2.4 Salient features of the yeast (*S. cerevisiae*) MSU RNAPs**

203 To-date one of the most well-studied eukaryotic RNAPs is from the baker's yeast, *S. cerevisiae*.
204 In fact, our present understanding of the eukaryotic transcription system is mainly based on the yeast
205 system. (For solving the structure and molecular basis of transcription by the yeast RNAP II, Roger
206 Kornberg was awarded Nobel Prize in chemistry in 2006). Like other eukaryotic cells, the yeast cells also
207 contain 3 different polymerases as discussed above. All the three yeast polymerases have five core
208 subunits that exhibit good homology to the β , β' , α and ω subunits of *E. coli* RNAP. RNAPs I and III
209 contain the same two non-identical α -like subunits, whereas polymerase II has two copies of a different α -
210 like subunit. All three yeast polymerases share four other common subunits as mentioned elsewhere. In
211 addition, each RNAP contains three to seven unique smaller subunits as shown in Table 3.

212 The yeast RNAP II, which involves in the production of mRNAs, is extensively characterized and
213 a great deal of information is available. The yeast RNAP II is composed of 12 subunits and the largest
214 subunit (Rpb1) contains the essential CTD, containing 26 heptapeptide repeats (YSPTSPS). Thus, the
215 Rpb1, Rpb2, Rpb3 and Rpb11 subunits are related to the subunits of the eubacterial RNA polymerase
216 core enzyme, whereas the 5 subunits, viz. Rpb5, Rpb6, Rpb8, Rpb10 and Rpb12 are shared among
217 yeast RNA polymerases I, II and III [25] (Table 3).

218 The yeast RNAP II core enzyme is composed of 12 subunits (Fig. 1). The 7 subunits, viz. Rpb1-4,
219 Rpb7, Rpb9 and Rpb11 are unique to RNAP II while the 5 subunits, viz. Rpb5, Rpb6, Rpb8, Rpb10 and
220 Rpb12 are shared between the three RNAPs (Table 3). The $\Delta rpb4$ phenotypes can be suppressed by
221 overexpression of Rpb7, and the high level of Rpb7 allows its interaction with RNAP II in the absence of
222 Rpb4, suggesting that Rpb7 is the critical component of the Rpb4-Rpb7 complex and the role of Rpb4 is
223 to stabilize the complex [26]. MSA analysis of the Rpb7 from various eukaryotic sources have shown 3
224 template binding pairs, possibly decide the right orientation of the template DNA by three-point
225 attachment in addition to two long stretches of conserved motifs (data not shown). Rpb2, a protein of
226 138,750 Daltons, exists as a single copy in the haploid yeast genome and disruption of the gene is lethal
227 to the yeast cell [11].

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234 Table 3. The composition of the *S. cerevisiae* RNAP-II

235	Subunit	Size (~kDa)	Subunit	Size (~kDa)
236	Rpb1 (β')	192E	Rpb7**	19E
237	Rpb2 (β)	139E	Rpb8*	17E/S
238	Rpb3 ($\sim\alpha$)	35E	Rpb9	14NE
239	Rpb4	25NE	Rpb10*	8.8E/S
240	Rpb5*	25E/S	Rpb11	14E
241	Rpb6* (ω)	18E/S	Rpb12*	7.7E/S

242 Adapted from [25]; E, Essential; NE, Nonessential; S, Shared.

243 *The 5 Rpb subunits, viz. 5, 6, 8, 10 and 12 are common for all 3 RNAPs, viz. RNAP I, II & III

244 The Rpb6 promotes RNAP assembly and contains 9 C2H2 zinc fingers [27].

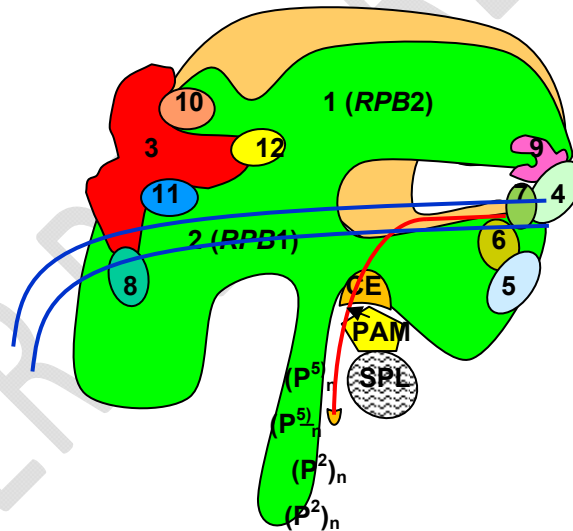
245 ** Rpb7 is unique to RNAP II. (Rpbs7 and 4 form a dimer and initiate transcription, not required for
246 elongation)

247 Rpb3 is not absolutely equivalent to α subunit of *E. coli*.

248 Subunits Rpb1, Rpb2, Rpb3 and Rpb7 are absolutely required for activity.

249 Rpb1 is known to bind strongly to Rpb5.

250 Rpb12 interacts with Rpb3.



265 **Fig. 1** A schematic diagram of the 12-subunit yeast MSU RNAP-II with the CTD and processing
266 components

267 CE Capping Enzyme; PAM, Poly-Adenylation Machinery; SPL, Spliceosome;

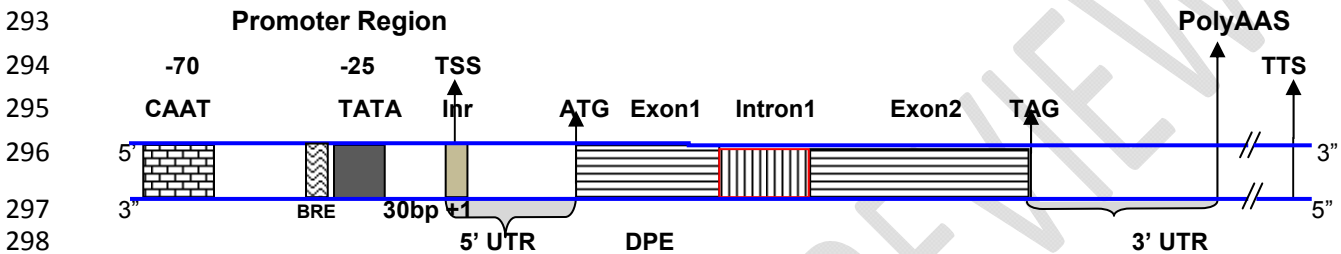
268 $(P^5)_n$, phosphorylation at Ser5 positions (during initiation); $(P^2)_n$ phosphorylations at Ser2 positions (during
269 elongation).

270
271 During the production of the primary transcript by RNAP-II, the phosphorylation state (P) of the
272 CTD changes to allow the transcribing polymerase to associate with the capping, splicing,
273 polyadenylation and mRNA export machinery [22]. X-ray crystallographic results also provide evidence
274 for RNA exit in the vicinity of the carboxyl-terminal repeat domain, coupling synthesis to RNA processing
275 by enzymes bound to this domain [28]. These associations are essential for normal processing of pre-
276 mRNAs to generate mature mRNAs and to export them to the cytoplasm and also for normal termination
277 of transcription by RNAP II. The subunit Rpb1 (*RPO21*) is the largest and catalytic component of RNAP II
278 and similar to the eubacterial β' . Bacterial RNA polymerase subunit ω and eukaryotic RNA polymerase
279 subunit Rpb6 (shared by RNAPs I, II, and III) are the sequence, structural, and functional homologs and
280 promote RNA polymerase assembly [12].

281 In this communication, the yeast RNAP II is used as the model system and the structural studies
 282 of yeast RNAP II are directly relevant to RNAP II enzymes in higher organisms, as the yeast RNAP
 283 II subunits are very well conserved in sequence and function as discussed elsewhere. These analyses
 284 not only bridge the data obtained by protein sequence analysis and the experimental results obtained on
 285 the yeast RNAP II but also will pave for more SDM experiments and genetic analysis to dissect the
 286 transcription mechanism in eukaryotes in much detail. The following brief discussion on the basic
 287 structure of the transcription unit, transcription cycle and the participation of the RNAP II subunits will be
 288 useful to further understand the analysis and conclusions.

289 3. Basic Structure of the Transcription Units in Eukaryotes

290 The transcription units in eukaryotes are slightly different from the one described in prokaryotes
 291 [4] as most of the eukaryotic genes are interrupted with intervening sequences called introns. A typical
 292 transcription unit in eukaryotes is shown in Fig. 2.



299 **Fig. 2** Schematic diagram of a typical transcription unit in eukaryotes

300 **NB:**

301 **Blue lines**, DNA strands; **TATA**, TATA box (Hogness box) (some promoters are TATAless), **CAAT**, CAAT box; The
 302 starting nucleotide (nt) is always an **A** in eukaryotes; **UTR**, UnTranslated Region); **PolyAAS**, Poly-A Addition Site
 303 (AATAAA); **TSS**, Transcription Start Site; **Inr**, Initiation region -pyTCACA-; **DPE**, Downstream Promoter Element (is
 304 seen ~ +30 bp in many genes, AGACA); **TTS**, Transcription Termination Site.

305 [Some transcription units have an InR start site, TATA-like site, Down Stream Promoter Elements (DPE), Upstream
 306 activator elements/Enhancer elements, Repressor and even Insulator and Silencer sequences. The poly-A cleavage
 307 site is about 30-35 or even 100 nucleotides downstream of the STOP codon (TAG). This sequence at the mRNAs' 3'
 308 end is called poly-A signal. The sequence present in the terminal region of the gene is TTATTT (poly-A signal) is
 309 usually followed by poly-U of 20 to 35 nts downstream from the poly-A signal.

310 5' UTR is the portion of an mRNA from the 5' end to the first codon; the 3' UTR is the portion of an mRNA from the
 311 last codon to the poly-A site. The UTRs play crucial roles in mRNA stability, transport, translation efficiency, etc].

312 4. Transcription Processes in Eukaryotes

313 The basic transcription unit in eukaryotes is the distance between the sites of transcription start
 314 site (TSS) and transcription termination site (TTS), and the coding region is usually interrupted by one to
 315 many introns (Fig. 2)

316 The RNA synthesis in eukaryotes is carried out in the nucleus and involves four distinctive steps,
 317 viz. formation of pre-initiation complex (PIC), initiation, elongation and termination. It is interesting to note
 318 that the eukaryotic RNAPs do not possess any initiator subunits similar to the σ factors, as reported in
 319 prokaryotes. However, unlike in prokaryotes where the initiation starts with the specific binding of the
 320 single σ subunit, in eukaryotes, at least about half a dozen protein factors involve in the initiation of
 321 transcription and are collectively known as General Transcription factors (GTFs). These GTFs assemble
 322 at the promoter region to form the PIC. Once the PIC is formed and the RNAP is positioned at TSS, the
 323 initiation subunit Rpb2 initiates RNA synthesis by making short RNA primers of ~10 nts, from which the
 324 elongation subunit Rpb1 extends and terminates at TTS. In eukaryotes, there is no well-defined
 325 transcriptional terminator region as in prokaryotes and therefore, the transcription progresses well beyond
 326 1000 or more nts downstream from the stop codon. Unlike RNAP I and III, RNAP II lacks any specific
 327 termination signals.

328 In my earlier communication, SSU RNAPs and MSU RNAPs (prokaryotic and prokaryotic type,
 329 i.e., plastid-encoded) RNAPs were analyzed in detail [29, 4, 30]. In this communication, the most complex

330 MSU RNAPs of eukaryotic origin are analyzed for their conserved motifs, active sites, metal binding
 331 regions and based on these findings, a plausible mechanism of action is proposed for these MSU
 332 eukaryotic enzymes using the yeast MSU RNAP II as the model enzyme.

333 5. MATERIALS AND METHODS

334 A large number of MSU RNAPs from eubacteria and eukaryotes have been isolated, purified,
 335 characterized, cloned and sequenced [17, 31, 6 and references therein]. Complete nucleic acid and
 336 protein sequence data are available for these enzymes from different eukaryotic sources. Thus, these
 337 data have become valuable tools in analyzing and understanding the structure-function relationships of
 338 these most complex enzymes which play a vital role gene expression. This communication presents a
 339 consensus model for initiation and elongation processes and also a plausible mechanism of action for
 340 these enzymes.

341 The *S. cerevisiae* DNA-dependent MSU RNAP II is used as the model system for delineating the
 342 polymerization mechanism. Biochemically and genetically as this is the most well-studied enzyme, a large
 343 amount of data on biochemical, SDM and X-ray analyses of its subunits make this enzyme a convenient
 344 model for investigating the physicochemical aspects of transcription in eukaryotes. For MSA of various
 345 eukaryotic RNAP II, the sequences were retrieved from SWISS-PROT and PUBMED sites and analyzed
 346 using Clustal Omega, an accurate, fast and widely accepted algorithm, available on their website.

347 6. RESULTS AND DISCUSSION

348 6.1 MSA of eukaryotic MSU RNA polymerases from different sources

349 The eukaryotic RNAP II is composed of various subunits as described elsewhere (Table 2). Only
 350 the two catalytic subunits, viz. Rpb2 and Rpb1 which involve in the initiation and elongation processes are
 351 analyzed to find out the conserved and active site regions among them. Figures 3 and 4 show the MSA of
 352 the two catalytic subunits of MSU RNAP II from various eukaryotic sources. To reduce the length of the
 353 article, only the relevant and highly conserved regions are shown in the Figures. The possible catalytic,
 354 template and substrate binding motifs are highlighted in yellow and the metal binding regions are
 355 highlighted in green/orange. The subunit sequences of standard organisms are highlighted in
 356 yellow/magenta and used for numbering.

357 6.1.1 MSA of Rpb2 subunits of eukaryotic MSU RNAPs (RNAP II)

358 Figure 3 shows the MSA analysis and conserved motifs of the Rpb2 initiation subunits in
 359 eukaryotic MSU RNAPs II. There are large numbers of conserved motifs are observed and some are
 360 found to be long stretches (highlighted). The yeast (*S. cerevisiae*) Rpb2 subunit (1224 amino acids) is
 361 used as the standard for numbering and highlighted in magenta. The catalytic region is found ~400 amino
 362 acids from the N-terminal. The invariant template binding YG is replaced by a KG pair in higher
 363 eukaryotes including human. The invariant catalytic R is placed at -9 from the template binding pair. This
 364 distance conservation is in close agreement with Pal and Luse findings that the transcription slippage
 365 abruptly stopped once the Rpb2 makes about 9 nts [32]. Interestingly, the transition from abortive to
 366 productive elongation cycle occurred once the RNAP register +10 nts [33]. The absolutely conserved R,
 367 which is implicated in NTP selection in SSU and MSU RNAPs and DNA polymerases, is placed -5
 368 position. In fact, in all the eubacterial β subunits the catalytic R is placed at -7th position from the YG pair
 369 and completely conserved R is placed at -8th position downstream from the catalytic R. However, catalytic
 370 R is placed at -8th position from the YG pair the completely conserved R was at -4th position in SSU
 371 RNAPs and DNA polymerases [29,34]. This strongly suggests that the DNA polymerases, SSU and MSU
 372 RNAPs use the same set of amino acids for template, substrate binding and catalysis establishing a
 373 structure-function relationship among the DNA and RNA polymerases. The immediate upstream amino
 374 acid from catalytic K in DNA polymerases is usually a G or A [34], but in SSU viral RNA polymerases it is
 375 a Q [29] and in MSU eubacterial β subunits, it is a D in all [4] and in eukaryotic Rpb2 it is S/T, suggesting
 376 a possible role in the substrate binding and catalysis. Another catalytic like region is located in ~400
 377 amino acids from the N-terminal but with a YG/FG pair and a catalytic R at -9 and long highly conserved
 378 stretches on both the sides. The Zn binding motif with 3 completely conserved Cs is found in the C-
 379 terminal region (highlighted in orange) which is followed by a long stretch of conservation till the C-
 380 terminal end. There are many highly conserved YG/FG/IG/LG pairs and a WG pair in the Rpb2 subunits.

381 At least three of them may provide by three-point attachment for stereo specific binding on to the template
382 DNA. However, the DNA polymerases show only one YG pair. Interestingly, the RNA dependent RNAPs
383 do not have the YG pair [35]. A long stretch of the C-terminal region is highly conserved in all Rpb2
384 subunits. Interestingly the higher eukaryotic Rpb2 subunits show a completely conserved motif till the end
385 of the C-terminal whereas the lower eukaryotic Rpb2 subunits also show a completely conserved stretch
386 but not covering all the amino acids till the C- terminal with a consensus sequence "PYAxKLLFQELMxM".
387 A long stretch of highly conserved end suggests a possible role in the initiation process.

388

UNDER PEER REVIEW

473	sp	P30876	RPB2_HUMAN	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
474	tr	G3V8Y5	G3V8Y5_RAT	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
475	tr	A0A250Y753	A0A250Y753_CASCN	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
476	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
477	tr	A0A286XI09	A0A286XI09_CAVPO	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
478	tr	I3M351	I3M351 ICTTR	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
479	tr	G7P5R6	G7P5R6_MACFA	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
480	tr	H2QPI8	H2QPI8_PANTR	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
481	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
482	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
483	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	155
484	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	155
485	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
486	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
487	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
488	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
489	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	155
490	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
491	tr	A0A096NEY4	A0A096NEY4_PAPAN	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	162
492	tr	C9J2Y9	C9J2Y9_HUMAN	NLTYSALYVVDITKTVIKEG-----EEQLQTHQKTFIGKIPIMLRSTYCL	155
493	tr	G8BY61	G8BY61_TETPH	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	178
494	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	176
495	tr	J7RV95	J7RV95_KAZNA	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	176
496	tr	H2AVJ8	H2AVJ8_KAZAF	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	176
497	sp	Q6FLD5	RPB2_CANGA	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	178
498	sp	P08518	RPB2_YEAST	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	180
499	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	180
500	tr	A0A0L8RB33	A0A0L8RB33_SACEU	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	180
501	tr	GOVJ71	GOVJ71_NAUCC	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	180
502	tr	G8ZM49	G8ZM49_TORDC	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	178
503	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	180
504	tr	A0A0N7IS35	A0A0N7IS35_9SACH	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	178
505	tr	A0A212MG88	A0A212MG88_ZYGBA	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	179
506	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	179
507	tr	S6ESB4	S6ESB4_ZYGB2	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	179
508	tr	B6K5Q5	B6K5Q5_SCHJY	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	165
509	sp	Q02061	RPB2_SCHPO	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	165
510	tr	S9R8U4	S9R8U4_SCHOY	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	165
511	tr	S9W8C6	S9W8C6_SCHCR	NLTYSGLFVDVVKRRTYEADVDPGRELKYNLIAEESEEDSDSGKVFIGRLPIMLRSENCY	165
512	****:	****:	****:	****:	****:
513	****:	****:	****:	****:	****:
514	****:	****:	****:	****:	****:
515	****:	****:	****:	****:	****:

516	sp	P30876	RPB2_HUMAN	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
517	tr	G3V8Y5	G3V8Y5_RAT	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
518	tr	A0A250Y753	A0A250Y753_CASCN	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
519	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
520	tr	A0A286XI09	A0A286XI09_CAVPO	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
521	tr	I3M351	I3M351 ICTTR	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
522	tr	G7P5R6	G7P5R6_MACFA	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
523	tr	H2QPI8	H2QPI8_PANTR	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
524	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
525	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	214
526	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
527	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	214
528	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
529	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
530	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
531	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
532	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	214
533	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
534	tr	A0A096NEY4	A0A096NEY4_PAPAN	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	221
535	tr	C9J2Y9	C9J2Y9_HUMAN	LNGLTDRLDCLNECPDLDGGYFIINGSEKVLIAQEKMATNTVYVFAKKD-SKYAYTGEC	214
536	tr	G8BY61	G8BY61_TETPH	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	238
537	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	236
538	tr	J7RV95	J7RV95_KAZNA	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	236
539	tr	H2AVJ8	H2AVJ8_KAZAF	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	236
540	sp	Q6FLD5	RPB2_CANGA	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	238
541	sp	P08518	RPB2_YEAST	LSEATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	240
542	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	LSEATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	240
543	tr	A0A0L8RB33	A0A0L8RB33_SACEU	LSEATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	240
544	tr	GOVJ71	GOVJ71_NAUCC	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	240
545	tr	G8ZM49	G8ZM49_TORDC	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	238
546	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	240
547	tr	A0A0N7IS35	A0A0N7IS35_9SACH	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	238
548	tr	A0A212MG88	A0A212MG88_ZYGBA	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	239
549	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	239
550	tr	S6ESB4	S6ESB4_ZYGB2	LSDATESDLYLKECPFDMGGYFIINGSEKVLIAQERSAGNIVQVFKKAAPSPISHVAEI	239
551	tr	B6K5Q5	B6K5Q5_SCHJY	LNGVSDSELYDLNECPYDGGYFIINGSEKVI IAQERSAANIVQVFRKAAPSP IAVVAEI	225
552	sp	Q02061	RPB2_SCHPO	LNGVSDSELYDLNECPYDGGYFIINGSEKVI IAQERSAANIVQVFRKAAPSP IAVVAEI	225
553	tr	S9R8U4	S9R8U4_SCHOY	LNGVSDSELYDLNECPYDGGYFIINGSEKVI IAQERSAANIVQVFRKAAPSP IAVVAEI	225
554	tr	S9W8C6	S9W8C6_SCHCR	LNGVSDSELYDLNECPYDGGYFIINGSEKVI IAQERSAANIVQVFRKAAPSP IAVVAEI	225
555	****:	****:	****:	****:	****:
556	****:	****:	****:	****:	****:

557	sp	P30876	RPB2_HUMAN	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
558	tr	G3V8Y5	G3V8Y5_RAT	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
559	tr	A0A250Y753	A0A250Y753_CASCN	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
560	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
561	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
562	tr	I3M351	I3M351 ICTTR	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
563	tr	G7P5R6	G7P5R6_MACFA	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
564	tr	H2QPI8	H2QPI8_PANTR	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
565	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
566	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
567	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	274
568	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	274
569	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
570	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
571	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
572	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
573	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	274
574	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
575	tr	A0A096NEY4	A0A096NEY4_PAPAN	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	281
576	tr	C9J2Y9	C9J2Y9_HUMAN	RSCLNSSRPTSTIIVWSMLARGGGGAKKSAIGQRI	VATLPHYKQEVPIIIVFRALGFVSD	274
577	tr	G8BY61	G8BY61_TETPH	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	292	
578	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	RSALFKGSRFISTLQVKLYGRES-----STRTIKATLPHYKQDIPIVIIFRALGIIPD	290	
579	tr	J7RV95	J7RV95_KAZNA	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	290	
580	tr	H2AVJ8	H2AVJ8_KAZAF	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	290	
581	tr	Q6FLD5	RPB2_CANGA	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	292	
582	sp	P08518	RPB2_YEAST	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	294	
583	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	294	
584	tr	A0A0L8RB33	A0A0L8RB33_SACEU	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	294	
585	tr	GOVJ71	GOVJ71_NAUCC	RSALFKGSRFISTLQVKLYGRES-----SDARTINATLPHYKQDIPIVIIFRALGIIPD	294	
586	tr	G8ZM49	G8ZM49_TORDC	RSALFKGSRFISTLQVKLYGRES-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	292	
587	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	294	
588	tr	A0A0N7IS35	A0A0N7IS35_9SACH	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	292	
589	tr	A0A212MG88	A0A212MG88_ZYGBA	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	293	
590	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	293	
591	tr	S6ESB4	S6ESB4_ZYGB2	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	293	
592	tr	B6K5Q5	B6K5Q5_SCHJY	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
593	tr	Q02061	RPB2_SCHPO	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
594	sp	S9R8U4	S9R8U4_SCHOY	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
595	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
596	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
597	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
598	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
599	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
600	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
601	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
602	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
603	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
604	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
605	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
606	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
607	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
608	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
609	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
610	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
611	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
612	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
613	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
614	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
615	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
616	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
617	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
618	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
619	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
620	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
621	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
622	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
623	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
624	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
625	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
626	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
627	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
628	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
629	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
630	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
631	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
632	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
633	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
634	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
635	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
636	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
637	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
638	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
639	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	
640	tr	S9W8C6	S9W8C6_SCHCR	RSALFKGSRFISTLQVKLFAREG-----SSRTIKATLPHYKQDIPIVIIFRALGIIPD	280	

725	sp	P30876	RPB2_HUMAN	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
726	tr	G3V8Y5	G3V8Y5_RAT	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
727	tr	A0A250Y753	A0A250Y753_CASCN	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
728	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
729	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
730	tr	I3M351	I3M351 ICTTR	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
731	tr	G7P5R6	G7P5R6_MACFA	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
732	tr	H2QP18	H2QP18_PANTR	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
733	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
734	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
735	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	514
736	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	514
737	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
738	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
739	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
740	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
741	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	514
742	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
743	tr	A0A096NEY4	A0A096NEY4_PAPAN	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	521
744	tr	C9J2Y9	C9J2Y9_HUMAN	ARAGVSVQLNR	IFASTLSHLRR	LNSPIGRDGK	LAKPRLHNT	LWGMVCPAET	PPEGHA	VG	514
745	tr	G8BY61	G8BY61_TETPH	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	532	
746	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	530	
747	tr	J7RV95	J7RV95_KAZNA	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	530	
748	tr	H2AVJ8	H2AVJ8_KAZAF	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	530	
749	sp	Q6FLD5	RPB2_CANGA	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	532	
750	sp	P08518	RPB2_YEAST	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	534	
751	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	534	
752	tr	A0A0L8RB33	A0A0L8RB33_SACEU	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	534	
753	tr	GOVJ71	GOVJ71_NAUCC	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	534	
754	tr	G8ZM49	G8ZM49_TORDC	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	532	
755	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	534	
756	tr	A0A0N7IS35	A0A0N7IS35_9SACH	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	532	
757	tr	A0A212MG88	A0A212MG88_ZYGBA	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	533	
758	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	533	
759	tr	S6ESB4	S6ESB4_ZYGB2	SRAGVSVQLNR	YFYSTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGLVCPAET	PPEGQACG	533	
760	tr	B6K5Q5	B6K5Q5_SCHJY	NRVGVSVQLNR	YFASTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGMVCPAET	PPEGQACG	520	
761	sp	Q02061	RPB2_SCHPO	NRVGVSVQLNR	YFASTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGMVCPAET	PPEGQACG	520	
762	tr	S9R8U4	S9R8U4_SCHOY	NRVGVSVQLNR	YFASTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGMVCPAET	PPEGQACG	520	
763	tr	S9W8C6	S9W8C6_SCHCR	NRVGVSVQLNR	YFASTLSHLRR	TNP	IGRDGKLAKPRLHNT	HWGMVCPAET	PPEGQACG	520	
764				*	*****	*::*****	*:*****	**::*****	**::*****	*	

767	sp	P30876	RPB2_HUMAN	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
768	tr	G3V8Y5	G3V8Y5_RAT	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
769	tr	A0A250Y753	A0A250Y753_CASCN	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
770	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
771	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
772	tr	I3M351	I3M351 ICTTR	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
773	tr	G7P5R6	G7P5R6_MACFA	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
774	tr	H2QP18	H2QP18_PANTR	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
775	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
776	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
777	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	574
778	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	574
779	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
780	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
781	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
782	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
783	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	574
784	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
785	tr	A0A096NEY4	A0A096NEY4_PAPAN	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	581
786	tr	C9J2Y9	C9J2Y9_HUMAN	LVKNLALMAY	ISVGSQ	PSPILEFLEEWS	MENLEEI	SPAAIADATK	IFVNGC	VWGIHKD	PE	574
787	tr	G8BY61	G8BY61_TETPH	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	592
788	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	590
789	tr	J7RV95	J7RV95_KAZNA	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	590
790	tr	H2AVJ8	H2AVJ8_KAZAF	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	590
791	sp	Q6FLD5	RPB2_CANGA	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	592
792	sp	P08518	RPB2_YEAST	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	594
793	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	594
794	tr	A0A0L8RB33	A0A0L8RB33_SACEU	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	594
795	tr	GOVJ71	GOVJ71_NAUCC	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	594
796	tr	G8ZM49	G8ZM49_TORDC	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	592
797	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	594
798	tr	A0A0N7IS35	A0A0N7IS35_9SACH	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	592
799	tr	A0A212MG88	A0A212MG88_ZYGBA	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	593
800	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	593
801	tr	S6ESB4	S6ESB4_ZYGB2	LVKNLSLMSC	ISVGTDP	MPPIITFLSEWG	MEPLEDY	VPHQSE	DATRVFVNGV	WHGVHRN	PA	593
802	tr	B6K5Q5	B6K5Q5_SCHJY	LVKNLSLMSC	ISVSVGS	SPAPIIEFLEEWG	MESLEDY	NPSASE	NATKVFVNGV	WLGVHRD	PV	580
803	sp	Q02061	RPB2_SCHPO	LVKNLSLMSC	ISVSVGS	SPAPIIEFLEEWG	MESLEDY	NPSASE	NATKVFVNGV	WLGVHRD	PA	580
804	tr	S9R8U4	S9R8U4_SCHOY	LVKNLSLMSC	ISVSVGS	SPAPIIEFLEEWG	MESLEDY	NPSASE	NATKVFVNGV	WLGVHRD	PA	580
805	tr	S9W8C6	S9W8C6_SCHCR	LVKNLSLMSC	ISVSVGS	SPAPIIEFLEEWG	MESLEDY	NPSASE	NATKVFVNGV	WLGVHRD	PA	580
806				*****	***	***	**	***	**	***	**	

1145	sp	P30876	RPB2_HUMAN	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1146	tr	G3V8Y5	G3V8Y5_RAT	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1147	tr	A0A250Y753	A0A250Y753_CASCN	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1148	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1149	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1150	tr	I3M351	I3M351 ICTTR	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1151	tr	G7P5R6	G7P5R6_MACFA	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1152	tr	H2QPI8	H2QPI8_PANTR	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1153	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1154	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1155	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1075
1156	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1075
1157	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1158	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1159	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1160	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1161	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1075
1162	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1163	tr	A0A096NEY4	A0A096NEY4_PAPAN	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1082
1164	tr	C9J2Y9	C9J2Y9_HUMAN	RGNEVLVNGFTGRKITSQIFIGPTYYQLKHMVDDKIHSRARGPIQLNRQPMEGRSRDG	1075
1165	tr	G8BY61	G8BY61_TETPH	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1126
1166	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1123
1167	tr	J7RV95	J7RV95_KAZNA	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1122
1168	tr	H2AVJ8	H2AVJ8_KAZAF	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1124
1169	tr	Q6FLD5	RPB2_CANGA	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1125
1170	sp	P08518	RPB2_YEAST	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1126
1171	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1126
1172	tr	A0A0L8RB33	A0A0L8RB33_SACEU	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1126
1173	tr	GOVJ71	GOVJ71_NAUCC	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1126
1174	tr	G8ZM49	G8ZM49_TORDC	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1124
1175	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1126
1176	tr	A0A0N7IS35	A0A0N7IS35_9SACH	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1124
1177	tr	A0A212MG88	A0A212MG88_ZYGBA	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1125
1178	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1125
1179	tr	S6ESB4	S6ESB4_ZYGB2	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1125
1180	tr	B6K5Q5	B6K5Q5_SCHJY	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1181	tr	Q02061	RPB2_SCHPO	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1182	tr	S9R8U4	S9R8U4_SCHOY	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1183	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1184	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1185	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1186	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1187	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1188	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1189	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1190	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1191	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1192	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1193	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1194	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1195	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1196	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1197	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1198	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1199	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1200	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1201	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1202	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1203	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1204	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1205	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1206	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1207	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1208	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1209	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1210	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1211	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1212	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1213	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1214	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1215	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1216	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1217	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1218	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1219	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1220	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1221	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1222	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1223	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1224	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1225	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1226	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1227	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115
1228	tr	S9W8C6	S9W8C6_SCHCR	RGFEVMYNGHTGKKLMAQIFFGPTYYQLRHMVDDKIHSRARGPMQVLRQPVEGRSDG	1115

1187	tr	P30876	RPB2_HUMAN	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1188	tr	G3V8Y5	G3V8Y5_RAT	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1189	tr	A0A250Y753	A0A250Y753_CASCN	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1190	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1191	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1192	tr	I3M351	I3M351 ICTTR	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1193	tr	G7P5R6	G7P5R6_MACFA	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1194	tr	H2QPI8	H2QPI8_PANTR	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1195	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1196	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1197	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1134
1198	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1134
1199	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1200	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1201	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1202	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1203	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1134
1204	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1205	tr	A0A096NEY4	A0A096NEY4_PAPAN	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1141
1206	tr	C9J2Y9	C9J2Y9_HUMAN	GLRFGEMERDCQIAHGAAQFLRERLFEASDPYQVHVCNLCGMA-IANTRTHTYEGRGCR	1134
1207	tr	G8BY61	G8BY61_TETPH	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMTVI AKLNHNQFECKGCD	1186
1208	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1183
1209	tr	J7RV95	J7RV95_KAZNA	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1182
1210	tr	H2AVJ8	H2AVJ8_KAZAF	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1184
1211	tr	Q6FLD5	RPB2_CANGA	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1185
1212	tr	P08518	RPB2_YEAST	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMTVI AKLNHNQFECKGCD	1186
1213	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMTVI AKLNHNQFECKGCD	1186
1214	tr	A0A0L8RB33	A0A0L8RB33_SACEU	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMTVI AKLNHNQFECKGCD	1186
1215	tr	GOVJ71	GOVJ71_NAUCC	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1186
1216	tr	G8ZM49	G8ZM49_TORDC	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1184
1217	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1186
1218	tr	A0A0N7IS35	A0A0N7IS35_9SACH	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1184
1219	tr	A0A212MG88	A0A212MG88_ZYGBA	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1185
1220	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1185
1221	tr	S6ESB4	S6ESB4_ZYGB2	GLRFGEMERDCMIAHGAAFLKERLMEASDAFRVHICGICGLMSVIAKLSHNQFECKGCD	1185
1222	tr	B6K5Q5	B6K5Q5_SCHJY	GLRFGEMERDCQISHGCSVLRERLFDSDAYRVVICDLGLIA-IASPKKNSYECRSQ	1174
1223	tr	Q02061	RPB2_SCHPO	GLRFGEMERDCQISHGCSVLRERLFDSDAYRVVICDLGLIA-IASPKKNSYECRSQ	1174
1224	tr	S9R8U4	S9R8U4_SCHOY	GLRFGEMERDCQISHGCSVLRERLFDSDAYRVVICDLGLIA-IASPKKNSYECRSQ	1174
1225	tr	S9W8C6	S9W8C6_SCHCR	GLRFGEMERDCQISHGCSVLRERLFDSDAYRVVICDLGLIA-IASPKKNSYECRSQ	1174
1226	tr	S9W8C6	S9W8C6_SCHCR	GLRFGEMERDCQISHGCSVLRERLFDSDAYRVVICDLGLIA-IASPKKNSYECRSQ	1174
1227	tr	S9W8C6	S9W8C6_SCHCR	GLRFGEMERDCQISHGCSVLRERLFDSDAYRVVICDLGLIA-IASPKKNSYECRSQ	1174
1228	tr	S9W8C6	S9W8C6_SCHCR	GLRFGEMERDCQISHGCSVLRERLFDSDAYRVVICDLGLIA-IASPKKNSYECRSQ	1174

1229			
1230			
1231	sp	P30876 RPB2_HUMAN	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1232	tr	G3V8Y5 G3V8Y5_RAT	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1233	tr	A0A250Y753 A0A250Y753_CASCN	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1234	tr	A0A1U7R4C7 A0A1U7R4C7_MESAU	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1235	tr	A0A286XIQ9 A0A286XIQ9_CAVPO	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1236	tr	I3M351 I3M351 ICTTR	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1237	tr	G7P5R6 G7P5R6_MACFA	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1238	tr	H2QPI8 H2QPI8_PANTR	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1239	tr	A0A1U7V0T5 A0A1U7V0T5_TARSY	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1240	tr	A0A1S2ZSL2 A0A1S2ZSL2_ERIEU	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1241	tr	A0A0D9QYL1 A0A0D9QYL1_CHLSB	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1242	tr	A0A2K5ZNR7 A0A2K5ZNR7_MANLE	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1243	tr	A0A2I2ZIU3 A0A2I2ZIU3_GORGO	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1244	tr	A0A1D5QGA5 A0A1D5QGA5_MACMU	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1245	tr	A0A2J8S2N1 A0A2J8S2N1_PONAB	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1246	tr	A0A2K5K5J5 A0A2K5K5J5_COLAP	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1247	tr	A0A2J8PEW7 A0A2J8PEW7_PANTR	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1248	tr	A0A2K5CY83 A0A2K5CY83_AOTNA	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1249	tr	A0A096NEY4 A0A096NEY4_PAPAN	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1250	tr	C9J2Y9 C9J2Y9_HUMAN	NKTQISLVRMPYACKLLFQELMSMSIAPRMMSV----
1251			
1252	tr	G8BY61 G8BY61_TETPH	NKIDYQIRIPYAAKLLFQELMAMNITPRLYTDRSKNF
1253	tr	A0A1X7QYA1 A0A1X7QYA1_9SACH	NKIDYQIQIPYAAKLLFQELMAMNITPRLYTDRSRDF
1254	tr	J7RV95 J7RV95_KAZNA	NKIDYQIRIPYAAKLLFQELMAMNITPRLYTDRSRDF
1255	tr	H2AVJ8 H2AVJ8_KAZAF	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSRDF
1256	sp	Q6FLD5 RPB2_CANGA	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSRDF
1257	sp	P08518 RPB2_YEAST	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSRDF
1258	tr	A0A0L8VHA5 A0A0L8VHA5_9SACH	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSRDF
1259	tr	A0A0L8RB33 A0A0L8RB33_SACEU	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSRDF
1260	tr	G0VJ71 G0VJ71_NAUCC	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSRDF
1261	tr	G8ZM49 G8ZM49_TORDC	NKIDYQINIPYAAKLLFQELMAMNITPRLYTDRSKDF
1262	tr	A0A1Q3A090 A0A1Q3A090_ZYGRO	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSKDF
1263	tr	A0A0N7IS35 A0A0N7IS35_9SACH	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSKDF
1264	tr	A0A212MG88 A0A212MG88_ZYGBA	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSKDF
1265	tr	A0A1S7HHE1 A0A1S7HHE1_9SACH	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSKDF
1266	tr	S6ESB4 S6ESB4_ZYGB2	NKIDYQIHIPYAAKLLFQELMAMNITPRLYTDRSKDF
1267	tr	B6K5Q5 B6K5Q5_SCHJY	NRTRFSQIYIPYAAKLLFQELMSMNIAPRLFTKNHKN--
1268	sp	Q02061 RPB2_SCHPO	NRTRFSQVYIPYAAKLLFQELMSMNIAPRLFTKNHKN--
1269	tr	S9R8U4 S9R8U4_SCHOY	NRTRFSQIYIPYAAKLLFQELMSMNIAPRLFTKNHKN--
1270	tr	S9W8C6 S9W8C6_SCHCR	NRTRFSQVYIPYAAKLLFQELMSMNIAPRLFTKNHKN--
1271			***.*****.*.*.***:

1272 **Fig. 3** MSA of the Rpb2, initiation subunits, of eukaryotic RNAPs II

1272	P30876 RPB2_HUMAN, <i>Homo sapiens</i>	G3V8Y5_RAT, <i>Rattus norvegicus</i>
1273	A0A250Y753_CASCN, <i>Castor Canadensis</i>	A0A1U7R4C7_MESAU, <i>Mesocricetus auratus</i>
1274	A0A286XIQ9_CAVPO, <i>Caviaporcellus</i>	tr G7P5R6_MACFA, <i>Macaca fascicularis</i>
1275	I3M351 ICTTR, <i>Ictidomys tridecemlineatus</i>	H2QPI8_PANTR, <i>Pan troglodytes</i>
1276	A0A1U7V0T5_TARSY, <i>Tarsiussyrichtha</i>	A0A1S2ZSL2_ERIEU, <i>Erinaceus europaeus</i>
1277	A0A0D9QYL1_CHLSB, <i>Chlorocebus sabaeus</i>	A0A2K5K5J5_COLAP, <i>Colobus angolensis palliatus</i>
1278	A0A2I2ZIU3_GORGO, <i>Gorilla gorilla gorilla</i>	A0A1D5QGA5_MACMU, <i>Macaca mulatta</i>
1279	A0A2J8PEW7_PANTR, <i>Pan troglodytes</i>	A0A2K5CY83_AOTNA, <i>Aotus nancymaeae</i>
1280	A0A2J8S2N1_PONAB, <i>Pongo abelii</i>	A0A2K5ZNR7_MANLE, <i>Mandrillus leucophaeus</i>
1281	A0A096NEY4_PAPAN, <i>Papio Anubis</i>	C9J2Y9_HUMAN, <i>Homo sapiens</i>
1282	G8BY61_TETPH, <i>Tetrapisporaphaffii</i>	tr A0A1X7QYA1_9SACH, <i>Kazachstania saulgeensis</i>
1283	J7RV95_KAZNA, <i>Kazachstaniana ganishii</i>	H2AVJ8_KAZAF, <i>Kazachstania africana</i>
1284	Q6FLD5_CANGA, <i>Candida glabrata</i>	P08518_Yeast, <i>Saccharomyces cerevisiae</i>
1285	A0A0L8VHA5_9SACH, <i>Saccharomyces boulardii</i>	A0A0L8RB33_SACEU, <i>Saccharomyces eubayanus</i>
1286	G0VJ71_NAUCC, <i>Naumovozyma castellii</i>	G8ZM49_TORDC, <i>Torulasporeadel brueckii</i>
1287	A0A1Q3A090_ZYGRO, <i>Zygosaccharomyces rouxii</i>	
1288	A0A0N7IS35_9SACH, <i>Zygosaccharomyces kombuchaensis</i>	
1289	A0A212MG88_ZYGBA, <i>Zygosaccharomyces bailii</i>	
1290	A0A1S7HHE1_9SACH, <i>Zygosaccharomyces parabailii</i>	
1291	S6ESB4_ZYGB2, <i>Zygosaccharomyces bailii</i> (strain CLIB 213)	
1292	B6K5Q5_SCHJY, <i>Schizosaccharomyces japonicas</i>	
1293	Q02061_SCHPO, <i>Schizosaccharomyces pombe</i>	
1294	S9R8U4_SCHOY, <i>Schizosaccharomyces octosporus</i>	
1295	S9W8C6_SCHCR, <i>Schizosaccharomyces cryophilus</i>	
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1300 Figure 4 shows the MSA analysis and conserved motifs in the Rpb1 elongation subunits in
 1301 eukaryotic MSU RNAPs II. The yeast subunit with 1733 amino acids is used as standard and highlighted
 1302 in magenta. Like the Rpb2 subunits, there are large numbers of conserved motifs and some are found to
 1303 be long stretches (highlighted). However, the Rpb1 subunits are more conserved than the Rpb2
 1304 subunits. The 'template binding' pair is invariably an FG rather than a YG pair as reported in other MSU
 1305 RNAPs. However, the catalytic R is completely conserved, including distance conservation in all the
 1306 eukaryotic Rpb1 subunits and the upstream neighbour is mostly S/T but in lower eukaryotes like yeasts, it
 1307 is N. Unlike in prokaryotic elongation subunit, the catalytic region is placed very close to the N-terminal (~
 1308 100 amino acids). It is interesting to note that the distance between the catalytic R and the FG pair is 18
 1309 amino acids, i.e., placed exactly double the distance as compared to the Rpb2 initiation subunits. This
 1310 distance closely agrees with the transcription bubble which extends ~ 20 bp from the TATA box [32]. The
 1311 template binding and catalytic conservations strongly suggest that the DNA, SSU and MSU RNA
 1312 polymerases use the same set of amino acids for template, substrate binding and catalysis establishing a
 1313 structure-function relationship among the DNA and RNAPs. Interestingly the catalytic region harbouring
 1314 the Zn binding motif is found very close to the N-terminal region. This is in sharp contrast to the
 1315 equivalent region in the β' subunit in all eubacteria (Table 1). The Zn binding motif was originally
 1316 identified by X-ray crystallographic analysis in the equivalent β' subunit of the thermophilic bacterium,
 1317 *Thermus aquaticus* [36]. The Zn binding 3 invariant Cs were located in the catalytic region and suggested
 1318 in possible proof-reading activity during elongation. A completely conserved R found upstream from the
 1319 catalytic R is located at -6th position which is implicated in NTP selection. It is interesting to note a
 1320 completely conserved R found upstream from the catalytic R is missing in eubacteria (Table 1) [29, 34].
 1321 The immediate downstream amino acid from catalytic K in DNA polymerases is usually a G or A [29], but
 1322 in viral RNA polymerases it is a K or R, in MSU β' subunits, it is a D and in all β' subunits it is an S and it
 1323 is an S/T in eukaryotes suggesting a possible role in NTP selection. In SSU RNAPs, an invariant hydroxyl
 1324 amino acid located very close to the YG pair is shown to involve in NTP selection by SDM experiments
 1325 [37]. In addition to the template binding FG pair, there are 1 YG pair, 4 LG pairs and 3 I/VLG pairs in this
 1326 subunit. A C-terminal conservation - **SPDSDEEN**- (where the penultimate E is replaced with
 1327 functionally equivalent D in some species) is seen in all higher forms of life and such sequence was
 1328 conspicuously absent in the lower eukaryotes like yeasts and also not found in the initiation subunits,
 1329 Rpb2. The -DXD- is a metal binding motif commonly found in glycosyl transferases and interestingly
 1330 similar motif -DXDXT- in metal-dependent phosphatases (DxDXE in C-terminals of Rpb1 elongation
 1331 subunits) where it plays a role for Ser phosphate removal from the CTD at the end of the cycle [38].
 1332 Interestingly, this motif invariably ends in N preceded by D/E in all the cases in Rpb1 (Fig 4). As expected
 1333 the CTD is filled with the "heptapeptide repeats" (YSPTSPS) in all species and its role is discussed
 1334 elsewhere (Not shown in the Figure) and results provide evidence for RNA exit in the vicinity of the
 1335 carboxyl-terminal repeat domain, coupling synthesis to RNA processing by enzymes bound to this
 1336 domain [28].

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1410 tr |A0A1U8DYN0|A0A1U8DYN0_ALLSI LVDSNNPKIKD-ILGSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 175
1411 tr |A0A1L8H4P4|A0A1L8H4P4_XENLA LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDI 174
1412 tr |H9GLG5|H9GLG5_ANOCA LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDIT 175
1413 tr |H2R1J6|H2R1J6_PANTR LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1414 tr |G1MCZ1|G1MCZ1_ALLME LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1415 tr |O08847|O08847_MOUSE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1416 tr |S7PWZ6|S7PWZ6_MYOBR LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1417 tr |D4A5A6|D4A5A6_RAT LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1418 sp |P08775|RPB1_MOUSE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1419 sp |P24928|RPB1_HUMAN LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1420 tr |A0A1S3EWL2|A0A1S3EWL2_DIPOR LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1421 sp |P11414|RPB1_CRIGR LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1422 tr |O35559|O35559_CRIGR LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1423 tr |A0A2I3M9H2|A0A2I3M9H2_PAPAN LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1424 tr |F7HB40|F7HB40_MACMU LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1425 tr |A0A2K6RYW9|A0A2K6RYW9_SAIBB LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEGDEDLT 176
1426 tr |W5N8Z6|W5N8Z6_LEPOC LVDSNNPKIKD-ILGSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 175
1427 tr |I3JRW6|I3JRW6_ORENI LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 174
1428 tr |A0A0R4IMS9|A0A0R4IMS9_DANRE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 173
1429 tr |A0A1A7X327|A0A1A7X327_9TELE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 174
1430 tr |A0A1A8UKD7|A0A1A8UKD7_NOTFU LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 174
1431 tr |A0A1A8ER05|A0A1A8ER05_9TELE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 174
1432 tr |A0A1A8DQ60|A0A1A8DQ60_9TELE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 174
1433 tr |A0A1A8NSR8|A0A1A8NSR8_9TELE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 174
1434 tr |A0A1W4YLM7|A0A1W4YLM7_9TELE LVDSNNPKIKD-ILAKSKGQPKRRLTHVYDLCKGKNI CRGGEEMDNKFGVEQTEEEDIT 174
1435 tr |A0A1M8A6L7|A0A1M8A6L7_MALS4 KADPISDPVFKSLQSTRANRRKRFQVWEYCSKITITCEADEGKDE-----DELGDTTQ 169
1436 sp |P04050|RPB1_YEAST LLDHNEELMRQAAL---AIKDSKKRFAAIWTLLTKMVCCTDVPSED-----DPT- 159
1437 tr |A0A1B2J8C6|A0A1B2J8C6_PICPA LLDETNPMTMAQAI---RIRDPKKRFNAVWQLCKTKMVCCTADAPVDE-----YSEQ 160
1438 tr |F2QW17|F2QW17_KOMPC LLDENPAMAQAI---RIRDPKKRFNAVWQLCKTKMVCCTADAPVDE-----YSEQ 160
1439 tr |A3GID7|A3GID7_PICST LLDENPAMAQAI---KIRDPKKRFNAVWQLCKTKMVCCTDIEEG-----ATE- 160
1440 tr |A0A1D8PUA6|A0A1D8PUA6_CANAL LLDENPAMAQAI---KIRDPKKRFNAVWQLCKTKMVCCTDNEDE-----MTD- 160
1441 tr |G8BEH9|G8BEH9_CANPC LLDENPLMAQAI---KIRDPKKRFNAVWQLCKTKMVCCTATSEE-----MND- 160

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1445 tr |A0A1U8DYN0|A0A1U8DYN0_ALLSI KEKGHGCCRYPRIIRRVGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIA 228
1446 tr |A0A1L8H4P4|A0A1L8H4P4_XENLA KEKGHGCCRYPRIIRRTGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIT 227
1447 tr |H9GLG5|H9GLG5_ANOCA KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 228
1448 tr |H2R1J6|H2R1J6_PANTR KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1449 tr |G1MCZ1|G1MCZ1_ALLME KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1450 tr |O08847|O08847_MOUSE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1451 tr |S7PWZ6|S7PWZ6_MYOBR KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1452 tr |D4A5A6|D4A5A6_RAT KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1453 sp |P08775|RPB1_MOUSE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1454 sp |P24928|RPB1_HUMAN KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1455 tr |A0A1S3EWL2|A0A1S3EWL2_DIPOR KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1456 sp |P11414|RPB1_CRIGR KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1457 tr |O35559|O35559_CRIGR KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1458 tr |A0A2I3M9H2|A0A2I3M9H2_PAPAN KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1459 tr |F7HB40|F7HB40_MACMU KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1460 tr |A0A2K6RYW9|A0A2K6RYW9_SAIBB KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 229
1461 tr |W5N8Z6|W5N8Z6_LEPOC KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 228
1462 tr |I3JRW6|I3JRW6_ORENI KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 227
1463 tr |A0A0R4IMS9|A0A0R4IMS9_DANRE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIA 226
1464 tr |A0A1A7X327|A0A1A7X327_9TELE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 227
1465 tr |A0A1A8UKD7|A0A1A8UKD7_NOTFU KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 227
1466 tr |A0A1A8ER05|A0A1A8ER05_9TELE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 227
1467 tr |A0A1A8DQ60|A0A1A8DQ60_9TELE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 227
1468 tr |A0A1A8NSR8|A0A1A8NSR8_9TELE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 227
1469 tr |A0A1W4YLM7|A0A1W4YLM7_9TELE KEKGHGCCRYPRIIRRSGLELYAEWKHVNE-----DSQEKKILLSPERVHEIFKRIS 227
1470 tr |A0A1M8A6L7|A0A1M8A6L7_MALS4 QKIGHGCCRFPQPAIRKEALKLFVSWVKQSKDEDEDSSGMAQSEKRPPLPASEVHTILKKIT 229
1471 sp |P04050|RPB1_YEAST QLVSRGGCCNTQPTIRKDGKLVGSKKDRATGD---ADEPELRLVSTEEILNIFKHIS 215
1472 tr |A0A1B2J8C6|A0A1B2J8C6_PICPA KVVSRGGCCNTQPVVVRKDGKLVGSKKDRATGD---DAQPERKLLTPGEILLNVFKHIS 216
1473 tr |F2QW17|F2QW17_KOMPC KVVSRGGCCNTQPVVVRKDGKLVGSKKDRATGD---DAQPERKLLTPGEILLNVFKHIS 216
1474 tr |A3GID7|A3GID7_PICST -TTTRGGCCHTQPTIIRRDGLKLVGSKKDRATGD---NEQPERRLLTPSEILNLVKHIS 215
1475 tr |A0A1D8PUA6|A0A1D8PUA6_CANAL -QPSRGGCCHTQPTIIRRDGLKLVGSKKDRATGD---NDQPERRLLTPSEILNLVKHIS 215
1476 tr |G8BEH9|G8BEH9_CANPC -HNVRRGGCCHTQPTIIRRDGLKLVGSKKDRATGD---NDQPERRLLTPSEILNLVKHIS 215

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1494	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	DDECLVLGMDPKFARPEWMI	CTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	288	
1495	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	DEECFLGMDPKFARPEWLI	ITVLPVPE	LCVPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1496	tr	H9GLG5	H9GLG5_ANOCA	DEECFLGMDPKFARPEWMI	CTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	288	
1497	tr	H2R1J6	H2R1J6_PANTR	DEECFLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1498	tr	G1MCZ1	G1MCZ1_AILME	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1499	tr	O08847	O08847_MOUSE	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1500	tr	S7PWZ6	S7PWZ6_MYOBR	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1501	tr	D4A5A6	D4A5A6_RAT	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1502	sp	P08775	RPB1_MOUSE	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1503	sp	P24928	RPB1_HUMAN	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1504	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1505	tr	P11414	RPB1_CRIGR	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1506	tr	O35559	O35559_CRIGR	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1507	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1508	tr	F7HB40	F7HB40_MACMU	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1509	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	DEECFVLGMEPRYARPEWMI	IVTVLPVPE	LSVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	289	
1510	tr	W5N8Z6	W5N8Z6_LEPOC	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	288	
1511	tr	I3JRW6	I3JRW6_ORENI	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1512	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	286	
1513	tr	A0A1A7X327	A0A1A7X327_9TELE	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1514	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1515	tr	A0A1A8ER05	A0A1A8ER05_9TELE	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1516	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1517	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1518	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	DEEDIILGMDPKFARPEWMI	IVTVLPVPE	LAVRPAVVMQGSARN	ODDLTHK	LADIVK	INNQ	287	
1519	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	PEDVVTILGSLMDEPAQDPDMV	LVTVLPVPE	PQVRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	289	
1520	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	PKDFTS	GGNEVSRPEWMLIT	QLPVPE	PPVPRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	275
1521	sp	P04050	RPB1_YEAST	PKDFTS	GGNEVSRPEWMLIT	QLPVPE	PPVPRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	275
1522	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	PEDCFRLG	NEDYARPEWMLIT	TVLPVPE	PQVRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	276
1523	tr	F2QW17	F2QW17_KOMP	PEDCFRLG	NEDYARPEWMLIT	TVLPVPE	PQVRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	276
1524	tr	A3GID7	A3GID7_PICST	SLDCLR	GGNEVSRPEWMLIT	TVLPVPE	PPVPRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	275
1525	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	PEDCYR	GGNEVSRPEWMLIT	TVLPVPE	PPVPRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	275
1526	tr	G8BEH9	G8BEH9_CANPC	PEDCYR	GGNEVSRPEWMLIT	TVLPVPE	PPVPRPSIAFNDTARGE	DDLTHK	LADIVK	IKANIN	275
1527				:	..	:.*:	
1528				:	
1529				:	

1530	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		348
1531	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1532	tr	H9GLG5	H9GLG5_ANOCA	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		348
1533	tr	H2R1J6	H2R1J6_PANTR	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1534	tr	G1MCZ1	G1MCZ1_AILME	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1535	tr	O08847	O08847_MOUSE	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1536	tr	S7PWZ6	S7PWZ6_MYOBR	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1537	tr	D4A5A6	D4A5A6_RAT	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1538	sp	P08775	RPB1_MOUSE	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1539	sp	P24928	RPB1_HUMAN	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1540	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1541	sp	P11414	RPB1_CRIGR	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1542	tr	O35559	O35559_CRIGR	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1543	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1544	tr	F7HB40	F7HB40_MACMU	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1545	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		349
1546	tr	W5N8Z6	W5N8Z6_LEPOC	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		348
1547	tr	I3JRW6	I3JRW6_ORENI	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1548	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	LKRNESGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		346
1549	tr	A0A1A7X327	A0A1A7X327_9TELE	LKRNESGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1550	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	LKRNESGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1551	tr	A0A1A8ER05	A0A1A8ER05_9TELE	LKRNESGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1552	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	LKRNESGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1553	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	LKRNESGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1554	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	LRNEQNGAAAHVIAEDV	KLLQFHV	ATMVDNELPGLPRAMQK	SGRPLKSLK	RLKGKEGR		347
1555	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	LRMEQEGAPAHILNDFADLL	QYHCATYMDNDIAGLP	QSLQSGR	VPKAIR	RLKGKEGR		349
1556	sp	P04050	RPB1_YEAST	LETLEHNGAPHAIIEAE	SILLQFHV	ATYMDNDIAGQP	QALQK	SGRPVKSIR	RLKGKEGR	335
1557	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	VQKLEMDGSPQHI	INEVEQLLQFHV	ATYMDNDIAGQP	QALQK	SGRPVKSIR	RLKGKEGR	336
1558	tr	F2QW17	F2QW17_KOMP	VQKLEMDGSPQHI	INEVEQLLQFHV	ATYMDNDIAGQP	QALQK	SGRPVKSIR	RLKGKEGR	336
1559	tr	A3GID7	A3GID7_PICST	VQKLEMDGSPQHI	INEVEQLLQFHV	ATYMDNDIAGQP	QALQK	SGRPVKSIR	RLKGKEGR	335
1560	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	VQKLEMDGSPQHI	INEVEQLLQFHV	ATYMDNDIAGQP	QALQK	SGRPVKSIR	RLKGKEGR	335
1561	tr	G8BEH9	G8BEH9_CANPC	VQKLEMDGSPQHI	INEVEQLLQFHV	ATYMDNDIAGQP	QALQK	SGRPVKSIR	RLKGKEGR	335
1562			
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1733	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	QVFSLIIPGHINCVRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	648
1734	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGLVMGILCKKSLG	754	647
1735	tr	H9GLG5	H9GLG5_ANOCA	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	648
1736	tr	H2R1J6	H2R1J6_PANTR	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1737	tr	G1MCZ1	G1MCZ1_ALLME	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1738	tr	O08847	O08847_MOUSE	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1739	tr	S7PWZ6	S7PWZ6_MYOBR	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1740	tr	D4A5A6	D4A5A6_RAT	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1741	sp	P08775	RBP1_MOUSE	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1742	sp	P24928	RBP1_HUMAN	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1743	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1744	sp	P11414	RBP1_CRIGR	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1745	tr	O35559	O35559_CRIGR	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1746	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1747	tr	F7HB40	F7HB40_MACMU	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1748	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	QIFSLIIPGHINCIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VVENGELIMGILCKKSLG	754	649
1749	tr	W5N8Z6	W5N8Z6_LEPOC	QVFSLIIPGHINAIRTHSTHPDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	648
1750	tr	I3JRW6	I3JRW6_ORENI	QIFSLIIPGHINVIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	647
1751	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	QIFSLIIPGHINAIRTHSTHPDDEDSGPYKNI	SPGDTKVI	VENGELIMGILCKKSLG	754	646
1752	tr	A0A1A7X327	A0A1A7X327_9TELE	QVFSLIIPGHINVIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	647
1753	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	QVFSLIIPGHINVIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	647
1754	tr	A0A1A8ER05	A0A1A8ER05_9TELE	QVFSLIIPGHINVIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	647
1755	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	QVFSLIIPGHINVIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	647
1756	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	QVFSLIIPGHINVIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	647
1757	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	QVFSLIIPGHINAIRTHSTHPDDEDSGPYKHI	SPGDTKVI	VENGELIMGILCKKSLG	754	647
1758	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	QLLSMAIPKGINVFLG-----	DAKAAANNFLKDDGVH	ENGEIMYGVINKKVVGSSA	754	640
1759	sp	P04050	RBP1_YEAST	QILSVIAIPSGIHLQRF-----	DEGT-TLLSPKDNGLM	IIDGQIIFGVVDDKKTGATG	754	626
1760	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	QLLSIAIPSGIHLQRT-----	DGGN-SLLSPKDNGLM	IVDGNVMFVVDKKTGSGG	754	627
1761	tr	F2QW17	F2QW17_KOMPC	QLLSMAIPSGIHLQRT-----	DGGN-SLLSPKDNGLM	IVDGKVMFVVDKKTGSSG	754	627
1762	tr	A3GID7	A3GID7_PICST	QLLSMAIPKGIHLQRF-----	DGGK-DLLSPKDTGML	IVDGEIMFVVDKKTGATG	754	626
1763	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	QLLSLAIPKGIHLQRF-----	DGGR-DLLSPKDTGML	IVDGEIMFVVDKKTGATG	754	626
1764	tr	G8BEH9	G8BEH9_CANPC	QMLSMIAIPKGIHLQRF-----	DGGR-DLLSPKDTGML	IVDGEIMFVVDKKTGATG	754	626
1765				***: ** *	*	* : : : : ** *		
1766								
1767	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	GSLVHISYLEMGHDVTRLXXXXXXXPQTPLALHFLVXKRGRGHTI		GIGDSIADAKTYQ	754	708
1768	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADAKTYQ	698
1769	tr	H9GLG5	H9GLG5_ANOCA	GSLVHISYLEMGHDVTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADAKTYQ	699
1770	tr	H2R1J6	H2R1J6_PANTR	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1771	tr	G1MCZ1	G1MCZ1_ALLME	GSLVHISYLEMGHDVTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1772	tr	O08847	O08847_MOUSE	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1773	tr	S7PWZ6	S7PWZ6_MYOBR	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1774	tr	D4A5A6	D4A5A6_RAT	GSLVHISYLEMGHDVTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1775	sp	P08775	RBP1_MOUSE	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1776	sp	P24928	RBP1_HUMAN	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1777	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1778	sp	P11414	RBP1_CRIGR	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1779	tr	O35559	O35559_CRIGR	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1780	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1781	tr	F7HB40	F7HB40_MACMU	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1782	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	GSLVHISYLEMGHDTRLFYFSNIQTVIN-----	N--	WLLIEGHTI	GIGDSIADSKTYQ	700
1783	tr	W5N8Z6	W5N8Z6_LEPOC	GSLVHISYLEMGHDVTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	699
1784	tr	I3JRW6	I3JRW6_ORENI	GSLVHISYLEMGHDTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	698
1785	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	GSLVHISYLEMGHDTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYQ	697
1786	tr	A0A1A7X327	A0A1A7X327_9TELE	GSLVHISYLEMGHDTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	698
1787	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	GSLVHISYLEMGHDTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	698
1788	tr	A0A1A8ER05	A0A1A8ER05_9TELE	GSLVHISYLEMGHDTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	698
1789	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	GSLVHISYLEMGHDTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	698
1790	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	GSLVHISYLEMGHDTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	698
1791	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	GSLVHISYLEMGHDVTRLFYFSNIQTVVNN-----		WLLIEGHSI	GIGDSIADAKTYL	698
1792	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	GGLIHIIFRERGPVVCDFSGVQRVLFN-----		WLLHNGFSI	GIGDVTADKATTA	691
1793	sp	P04050	RBP1_YEAST	GGLIHVVTREKGPQVCAKLFNGIQKVVNF-----		WLLHNGFSI	GIGDVTADKATTA	677
1794	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	GGLIHTVMREKGPKICAEFLFGNIQKVVNY-----		WLLHNGFSI	GIGDAIADASTMK	678
1795	tr	F2QW17	F2QW17_KOMPC	GGLIHTVMREKGPKICAEFLFGNIQKVVNY-----		WLLHNGFSI	GIGDAIADASTMK	678
1796	tr	A3GID7	A3GID7_PICST	GGLIHTVMREKGPQVCAQLFSSIQKVTNY-----		WLLHNGFSI	GIGDVTADKATTA	677
1797	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	GGLIHTVMREKGPQVCAQLFSSIQKVVNY-----		WLLHNGFSI	GIGDVTADKATTA	677
1798	tr	G8BEH9	G8BEH9_CANPC	GGLIHTVMREKGPQVCAQLFSSIQKVVNF-----		WLLHNGFSI	GIGDVTADKATTA	677
1799				*. *: *		*. *: *		
1800								
1801								

1802	tr	A0A1U8DYN0 A0A1U8DYN0_ALLSI	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	768
1803	tr	A0A1L8H4P4 A0A1L8H4P4_XENLA	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1804	tr	H9GLG5 H9GLG5_ANOCA	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	759
1805	tr	H2R1J6 H2R1J6_PANTR	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1806	tr	G1MCZ1 G1MCZ1_ALLME	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1807	tr	O08847 O08847_MOUSE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1808	tr	S7PWZ6 S7PWZ6_MYOBR	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1809	tr	D4A5A6 D4A5A6_RAT	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1810	sp	P08775 RPB1_MOUSE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1811	sp	P24928 RPB1_HUMAN	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1812	tr	A0A1S3EWL2 A0A1S3EWL2_DIPOR	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1813	sp	P11414 RPB1_CRIGR	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1814	tr	O35559 O35559_CRIGR	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1815	tr	A0A2I3M9H2 A0A2I3M9H2_PAPAN	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1816	tr	F7HB40 F7HB40_MACMU	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1817	tr	A0A2K6RYW9 A0A2K6RYW9_SAIIBB	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	760
1818	tr	W5N8Z6 W5N8Z6_LEPOC	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	759
1819	tr	I3JRW6 I3JRW6_ORENI	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1820	tr	A0A0R4IMS9 A0A0R4IMS9_DANRE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	757
1821	tr	A0A1A7X327 A0A1A7X327_9TELE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1822	tr	A0A1A8UKD7 A0A1A8UKD7_NOTFU	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1823	tr	A0A1A8ER05 A0A1A8ER05_9TELE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1824	tr	A0A1A8DQ60 A0A1A8DQ60_9TELE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1825	tr	A0A1A8NSR8 A0A1A8NSR8_9TELE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1826	tr	A0A1W4YLM7 A0A1W4YLM7_9TELE	DIQNTIKKAKQDVIEVIEKAHNNELEPTPGNTRLRQTFENQVNRILNDARDKTGSSAQKSL	758
1827	tr	A0A1M8A6L7 A0A1M8A6L7_MALS4	NIQTITAKAKAVMDLIIQAARNDLWKADPGMTRLESFEAHVNRILNKRARDVGSBAEQNL	751
1828	sp	P04050 RPB1_YEAST	EITHTAISSAKEQVQEI IYKQHNLELELPGMTRLESFEAGEVSRTLNDARDTAGRSAEMNL	737
1829	tr	A0A1B2J8C6 A0A1B2J8C6_PICPA	EITHTAISSAKEQVQEI IYKQHNLELELPGMTRLESFEAGEVSRTLNDARDTAGRSAEMNL	738
1830	tr	F2QW17 F2QW17_KOMPC	DITSTSEAKIKVQEI ILDAQSNKLEPEPGMTRLESFEHNSRVLNQRDITAGRSAEMNL	737
1831	tr	A3GID7 A3GID7_PICST	DVNKTIQEAQKQVQEI ILDAQHNKLEPEPGMTRLESFEHNSRVLNQRDITAGRSAEMSL	737
1832	tr	A0A1D8PUA6 A0A1D8PUA6_CANAL	DITKTIQEAQKQVQEI ILDAQHNKLDPEPGMTRLESFEHNSRVLNQRDITAGRSAEMSL	737
1833	tr	G8BEH9 G8BEH9_CANPC	:: * : * . * : * : : * * : * : * : * : * * : * : * : * : *	
1835				
1836				
1837	tr	A0A1U8DYN0 A0A1U8DYN0_ALLSI	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	828
1838	tr	A0A1L8H4P4 A0A1L8H4P4_XENLA	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1839	tr	H9GLG5 H9GLG5_ANOCA	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	819
1840	tr	H2R1J6 H2R1J6_PANTR	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1841	tr	G1MCZ1 G1MCZ1_ALLME	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1842	tr	O08847 O08847_MOUSE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1843	tr	S7PWZ6 S7PWZ6_MYOBR	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1844	tr	D4A5A6 D4A5A6_RAT	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1845	sp	P08775 RPB1_MOUSE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1846	sp	P24928 RPB1_HUMAN	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1847	tr	A0A1S3EWL2 A0A1S3EWL2_DIPOR	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1848	sp	P11414 RPB1_CRIGR	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1849	tr	O35559 O35559_CRIGR	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1850	tr	A0A2I3M9H2 A0A2I3M9H2_PAPAN	SEYNNFKSLVFPHT-GKVFVCIRVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	819
1851	tr	F7HB40 F7HB40_MACMU	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1852	tr	A0A2K6RYW9 A0A2K6RYW9_SAIIBB	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	820
1853	tr	W5N8Z6 W5N8Z6_LEPOC	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	819
1854	tr	I3JRW6 I3JRW6_ORENI	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1855	tr	A0A0R4IMS9 A0A0R4IMS9_DANRE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	817
1856	tr	A0A1A7X327 A0A1A7X327_9TELE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1857	tr	A0A1A8UKD7 A0A1A8UKD7_NOTFU	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1858	tr	A0A1A8ER05 A0A1A8ER05_9TELE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1859	tr	A0A1A8DQ60 A0A1A8DQ60_9TELE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1860	tr	A0A1A8NSR8 A0A1A8NSR8_9TELE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1861	tr	A0A1W4YLM7 A0A1W4YLM7_9TELE	SEYNNFKSMVVSAGKSGSKINISQVIAVVGQQIVVEGKRIPFGFKHRTLPHFIKDDYIPESR	818
1862	tr	A0A1M8A6L7 A0A1M8A6L7_MALS4	PDWNNVKQMVVAGSKGSFINISQMSACVGGQIVVEGKRIPFGFRHRTLPHFIKDDYIPESR	811
1863	sp	P04050 RPB1_YEAST	KDLNNVKQMVVAGSKGSFINIAQMSACVGGQIVVEGKRIPFGFVDRRTLPHFIKDDYIPESK	797
1864	tr	A0A1B2J8C6 A0A1B2J8C6_PICPA	KDLNNVKQMVVAGSKGSFINIAQMSACVGGQIVVEGKRIPFGFADRSRLPHFIKDDYIPESK	798
1865	tr	F2QW17 F2QW17_KOMPC	KDLNNVKQMVVAGSKGSFINIAQMSACVGGQIVVEGKRIPFGFADRSRLPHFIKDDYIPESK	798
1866	tr	A3GID7 A3GID7_PICST	KDLNNVKQMVVAGSKGSFINISQMSACVGGQIVVEGKRIPFGFSRRTLPHFIKDDYIPESK	797
1867	tr	A0A1D8PUA6 A0A1D8PUA6_CANAL	KDLNNVKQMVVAGSKGSFINISQMSACVGGQIVVEGKRIPFGFADRSRLPHFIKDDYIPESK	797
1868	tr	G8BEH9 G8BEH9_CANPC	KDLNNVKQMVVAGSKGSFINISQMSACVGGQIVVEGKRIPFGFADRSRLPHFIKDDYIPESK	797
1869			:: * : * . * : * : : * * : * : * : * : * * : * : * : * : *	
1870				
1871				

1872	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	888	
1873	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	878	
1874	tr	H9GLG5	H9GLG5_ANOCA	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	879	
1875	tr	H2R1J6	H2R1J6_PANTR	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1876	tr	G1MCZ1	G1MCZ1_ALLME	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1877	tr	O08847	O08847_MOUSE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1878	tr	S7PWZ6	S7PWZ6_MYOBR	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1879	tr	D4A5A6	D4A5A6_RAT	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1880	sp	P08775	RPB1_MOUSE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1881	sp	P24928	RPB1_HUMAN	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1882	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1883	sp	P11414	RPB1_CRIGR	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1884	tr	O35559	O35559_CRIGR	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1885	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	879	
1886	tr	F7HB40	F7HB40_MACMU	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1887	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	880	
1888	tr	W5N8Z6	W5N8Z6_LEPOC	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	879	
1889	tr	I3JRW6	I3JRW6_ORENI	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDGTVR	878	
1890	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	877	
1891	tr	A0A1A7X3Z7	A0A1A7X3Z7_9TELE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	878	
1892	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	878	
1893	tr	A0A1A8ER05	A0A1A8ER05_9TELE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	878	
1894	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	878	
1895	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	878	
1896	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	GFVENSYLAGLTPTEFFPHAMG	GREGLIDTAVKTAETGYIQRRLIKSMESV	MVKYDATVR	878	
1897	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	GFVENSYLRGLTAPQEFFPHAMG	GREGLIDTAVKTAETGYIQRRLVKALEDV	TMVHYDGTTR	857	
1898	sp	P04050	RPB1_YEAST	GFVENSYLRGLTAPQEFFPHAMG	GREGLIDTAVKTAETGYIQRRLVKALEDV	TMVHYDGTTR	871	
1899	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	GFVENSYLRGLTAPQEFFPHAMG	GREGLIDTAVKTAETGYIQRRLVKALEDV	TMVHYDGTTR	858	
1900	tr	F2QW17	F2QW17_KOMPC	GFVENSYLRGLTAPQEFFPHAMG	GREGLIDTAVKTAETGYIQRRLVKALEDV	TMVHYDGTTR	858	
1901	tr	A3GID7	A3GID7_PICST	GFVENSYLRGLTAPQEFFPHAMG	GREGLIDTAVKTAETGYIQRRLVKALEDV	TMVHYDGTTR	857	
1902	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	GFVENSYLRGLTAPQEFFPHAMG	GREGLIDTAVKTAETGYIQRRLVKALEDV	TMVHYDGTTR	857	
1903	tr	G8BEH9	G8BEH9_CANPC	GFVENSYLRGLTAPQEFFPHAMG	GREGLIDTAVKTAETGYIQRRLVKALEDV	TMVHYDGTTR	857	
1904				*****	*****	*****	*****	
1905								
1906								
1907	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVMK	948
1908	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1909	tr	H9GLG5	H9GLG5_ANOCA	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVMK	939
1910	tr	H2R1J6	H2R1J6_PANTR	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1911	tr	G1MCZ1	G1MCZ1_ALLME	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1912	tr	O08847	O08847_MOUSE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1913	tr	S7PWZ6	S7PWZ6_MYOBR	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1914	tr	D4A5A6	D4A5A6_RAT	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1915	sp	P08775	RPB1_MOUSE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1916	sp	P24928	RPB1_HUMAN	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1917	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1918	sp	P11414	RPB1_CRIGR	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1919	tr	O35559	O35559_CRIGR	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1920	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	939
1921	tr	F7HB40	F7HB40_MACMU	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1922	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDLVK	940
1923	tr	W5N8Z6	W5N8Z6_LEPOC	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	939
1924	tr	I3JRW6	I3JRW6_ORENI	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1925	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	937
1926	tr	A0A1A7X3Z7	A0A1A7X3Z7_9TELE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1927	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1928	tr	A0A1A8ER05	A0A1A8ER05_9TELE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1929	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1930	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1931	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1932	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	NSINQVVQLRYGEDGLAGE	SVEFQNLATLTKPSNKA	FEKKFKFDYANERALRR	TQEDVVK	938
1933	sp	P04050	RPB1_YEAST	NSLNQVIFQIYGEDGIDG	AMVERQKLI	THGLNDKEFRFRKVDLSHGGF	--KGTLRAGL	929
1934	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	NSLGDIIQPIYGEDGIDG	TQVEKQSVDT	IPGSDKAFHKRYVVDLMDKENS	IKADVIEYAA	918
1935	tr	F2QW17	F2QW17_KOMPC	NSLGDIIQPIYGEDGIDG	TQVEKQSVDT	IPGSDKAFHKRYVVDLMDKENS	IKADVIEYAA	918
1936	tr	A3GID7	A3GID7_PICST	NSLGDIIQPIYGEDGIDG	TQVEKQSVDT	IPGSDKAFHKRYVVDLMDKENS	IKADVIEYAA	917
1937	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	NSLGDIIQPIYGEDGIDG	TQVEKQSVDT	IPGSDKAFHKRYVVDLMDKENS	IKADVIEYAA	917
1938	tr	G8BEH9	G8BEH9_CANPC	NSLGDIIQPIYGEDGIDG	TQVEKQSVDT	IPGSDKAFHKRYVVDLMDKENS	IKADVIEYAA	917
1939				**	*****	*****	*****	
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2335	tr	A0A1U8DYN0 A0A1U8DYN0_ALLSI	----	1926
2336	tr	A0A1L8H4P4 A0A1L8H4P4_XENLA	SPDSDSEEDN----	1968
2337	tr	H9GLG5 H9GLG5_ANOCA	SPDSDSEEN----	1969
2338	tr	H2R1J6 H2R1J6_PANTR	SPDSDSEEN----	1960
2339	tr	G1MCZ1 G1MCZ1_ALLME	SPDSDSEEN----	1973
2340	tr	O08847 O08847_MOUSE	-----SEEN----	1966
2341	tr	S7PWZ6 S7PWZ6_MYOBR	SPDSDSEEN----	1970
2342	tr	D4A5A6 D4A5A6_RAT	SPDSDSEEN----	1970
2343	sp	P08775 RPB1_MOUSE	SPDSDSEEN----	1970
2344	sp	P24928 RPB1_HUMAN	SPDSDSEEN----	1970
2345	tr	A0A1S3EWL2 A0A1S3EWL2_DIPOR	SPDSDSEEN----	1970
2346	sp	P11414 RPB1_CRIGR	SPDSDSEEN----	1970
2347	tr	O35559 O35559_CRIGR	SPDSDSEEN----	1970
2348	tr	A0A2I3M9H2 A0A2I3M9H2_PAPAN	SPDSDSEEN----	1931
2349	tr	F7HB40 F7HB40_MACMU	SPDSDSEEN----	1932
2350	tr	A0A2K6RYW9 A0A2K6RYW9_SAIBB	SPDSDSEEN----	1932
2351	tr	W5N8Z6 W5N8Z6_LEPOC	SPDSDSEEDN----	1959
2352	tr	I3JRW6 I3JRW6_ORENI	SPDSDSEENN----	1966
2353	tr	A0A0R4IMS9 A0A0R4IMS9_DANRE	SPDSDSEENN----	1965
2354	tr	A0A1A7X327 A0A1A7X327_9TELE	SPDSDSESEENN	1969
2355	tr	A0A1A8UKD7 A0A1A8UKD7_NOTFU	SPDSDSESEENN	1969
2356	tr	A0A1A8ER05 A0A1A8ER05_9TELE	SPDSDSESEENN	1969
2357	tr	A0A1A8DQ60 A0A1A8DQ60_9TELE	SPDSDSESEENN	1962
2358	tr	A0A1A8NSR8 A0A1A8NSR8_9TELE	SPDSDSESEENN	1962
2359	tr	A0A1W4YLM7 A0A1W4YLM7_9TELE	SPDSDSDDN----	1968
2360	tr	A0A1M8A6L7 A0A1M8A6L7_MALS4	-----	1803
2361	sp	P04050 RPB1_YEAST	-----	1733
2362	tr	A0A1B2J8C6 A0A1B2J8C6_PICPA	-----	1743
2363	tr	F2QW17 F2QW17_KOMPC	-----	1743
2364	tr	A3GID7 A3GID7_PICST	-----	1739
2365	tr	A0A1D8PUA6 A0A1D8PUA6_CANAL	-----	1728
2366	tr	G8BEH9 G8BEH9_CANPC	-----	1746

Fig. 4 MSA of the Rpb1 the elongation subunits of eukaryotic RNAP II

2370	A0A1U8DYN0_ALLSI	<i>Alligator sinensis</i>	A0A1L8H4P4_XENLA	<i>Xenopus laevis</i>
2371	H9GLG5_ANOCA	<i>Anolis carolinensis</i>	H2R1J6_PANTR	<i>Pan troglodytes</i>
2372	G1MCZ1_AILME	<i>Ailuropoda melanoleuca</i>	O08847_MOUSE	<i>Mus musculus</i>
2373	S7PWZ6_MYOBR	<i>Myotis brandtii</i>	D4A5A6_RAT	<i>Rattus norvegicus</i>
2374	P08775_MOUSE	<i>Mus musculus</i>	P24928 RPB1_HUMAN	<i>Homo sapiens</i>
2375	A0A1S3EWL2_DIPOR	<i>Dipodomys ordii</i>	P11414_CRIGR	<i>Cricetulus griseus</i>
2376	O35559_CRIGR	<i>Cricetulus griseus</i>	A0A2I3M9H2_PAPAN	<i>Papio Anubis</i>
2377	F7HB40 _MACMU	<i>Macaca mulatta</i>	A0A2K6RYW9_SAIBB	<i>Saimiri boliviensis</i>
2378	W5N8Z6_LEPOC	<i>Lepisosteus oculatus</i>	I3JRW6_ORENI	<i>Oreochromis niloticus</i>
2379	A0A0R4IMS9_DANRE	<i>Danio rerio</i>	A0A1A7X327_9TELE	<i>Aphyosemion striatum</i>
2380	A0A1A8UKD7_NOTFU	<i>Nothobranchius furzeri</i>	A0A1A8ER05_9TELE	<i>Nothobranchius korthausae</i>
2381	A0A1A8DQ60_9TELE	<i>Nothobranchius kadleci</i>	A0A1A8NSR8_9TELE	<i>Nothobranchius rachovii</i>
2382	A0A1W4YLM7_9TELE	<i>Scleropages formosus</i>	A0A1M8A6L7_MALS4	<i>Malassezia sympodialis</i>
2383	P04050 RPB1_YEAST	<i>Saccharomyces cerevisiae</i>	A0A1B2J8C6_PICPA	<i>Komagataella pastoris</i>
2384	F2QW17_KOMPC	<i>Komagataella phaffii</i>	A3GID7_PICST	<i>Scheffersomyces stipitii</i>
2385	A0A1D8PUA6_CANAL	<i>Candida albicans</i>	G8BEH9_CANPC	<i>Candida parapsilosis</i>

2386 7. Mix and Match analysis of eubacterial and eukaryotic initiation and elongation 2387 subunits

2388 For this analysis a small number of initiation and elongation subunits of eubacterial and
2389 eukaryotic MSU RNAPs were subjected to MSA with ClustalW Omega programme and presented in Figs.
2390 5 and 6. For eubacteria, *E. coli* MSU RNAP and for eukaryote, *S. cerevisiae* MSU RNAP II sequences
2391 were used as standards. Only highly conserved regions are shown in the figures.

2392 7.1 Mix and Match analysis of the *E. coli* (β) and *S. cerevisiae* (Rpb2) initiation subunits

2393 Figure 5 shows the mix and match analysis of the initiation subunits from pro- and eukaryotic
2394 MSU RNAPs. The eukaryotic sequences are shown in red. The active site regions are highlighted in
2395 yellow the representative sequences are highlighted in yellow. There are a good number of sequences
2396 aligning in both. Significant among them are
2397 FIVINGS/TEK/RVL/II/VA/SQ- (~200), IETPE (~500) preceding this sequence a WG in eukaryotic and YG
2398 in prokaryotic sequences, -ASI/LIPF-, the metal ion binding regions -GYNQ/FEDS- (~800), -
2399 LDD/ED/SGL/I- (~850); -GDKF/MAS/GRHGXKG- (~1000), HLVDK/MHAR (~1050), RFGEME (~1100).
2400 The catalytic regions are matching but among themselves only and there is no complete consensus
2401 among them, i.e., they are located at different regions. However, the metal binding motifs are aligning in

2402 both the initiation subunits suggesting the metal binding regions are preserved as a domain and did not
2403 diverge much during evolution. The prokaryotic active site region is placed around 540-555/1342 amino
2404 acids whereas eukaryotic active site region is placed around 851-866/1224 amino acids. The notable
2405 difference in the eukaryotic template binding pair sis KG in lower eukaryotes like human and in lower
2406 eukaryotes it uses YG like yeasts. The eukaryotes follow two very similar ending sequences at the C-
2407 terminal end as –ACKLLFQELMSMSIAPRMSV- (~1150) and –
2408 AKLLFQELMAMNITPRLYT- (~1200) (deviating amino acids are shown in red) and the prokaryotes
2409 follow an altogether different consensus end sequence as –SFNVLLKEIRSL- (~1280). Thus, the
2410 eukaryotic and prokaryotic ending sequences are different and their significance is not clear now.

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UNDER PEER REVIEW

2414 CLUSTAL O(1.2.4): Mix and Match analysis of eukaryotic Rpb2 and eubacterial beta initiation subunits 5

2415	sp	P30876	RPB2_HUMAN	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2416	tr	G3V8Y5	G3V8Y5_RAT	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2417	tr	A0A250Y753	A0A250Y753_CASCN	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2418	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2419	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2420	tr	I3M351	I3M351_ICTTR	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2421	tr	G7P5R6	G7P5R6_MACFA	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2422	tr	H2QP18	H2QP18_PANTR	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2423	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2424	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2425	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	212
2426	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	212
2427	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2428	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2429	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2430	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2431	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	212
2432	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2433	tr	A0A096NEY4	A0A096NEY4_PAPAN	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	219
2434	tr	C9J2Y9	C9J2Y9_HUMPH	LLNGLTDRDLCELNECPLDPGGYFI	INGSEKVLIAQEKMATNTVYVF-AKDD-SKYAYTG	212
2435	tr	G8BY61	G8BY61_TETPN	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	236
2436	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	234
2437	tr	J7RV95	J7RV95_KAZNA	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	234
2438	tr	H2AVJ8	H2AVJ8_KAZAF	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	234
2439	sp	Q6FLD5	RPB2_CANGA	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	236
2440	sp	P08518	RPB2_YEAST	YLSEATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	238
2441	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	YLSEATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	238
2442	tr	A0A0L8RB33	A0A0L8RB33_SACEU	YLSEATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	238
2443	tr	GOVJ71	GOVJ71_NAUCC	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	238
2444	tr	G8ZM49	G8ZM49_TORDC	YLSDATELDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	236
2445	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	238
2446	tr	A0A0N7IS35	A0A0N7IS35_9SACH	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	236
2447	tr	A0A212MG88	A0A212MG88_ZYGBA	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	237
2448	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	237
2449	tr	S6ESB4	S6ESB4_ZYGB2	YLSDATESDLYKLKCEPFDMGGYFI	INGSEKVLIAQERSAGNIVQVF-KKAAPSPISHVA	237
2450	tr	B6K5Q5	B6K5Q5_SCHJY	ILNGVSDSELYDLNECPYDQGGYFI	INGSEKVLIAQERSAANIVQVF-RKAAPSPYAYVA	223
2451	sp	Q02061	RPB2_SCHPO	ILNGVSDSELYDLNECPYDQGGYFI	INGSEKVLIAQERSAANIVQVF-RKAAPSPYAYVA	223
2452	tr	S9R8U4	S9R8U4_SCHOY	ILNGVSDAELYDLNECPYDQGGYFI	INGSEKVLIAQERSAANIVQVF-RKAAPSPYAYVA	223
2453	tr	S9W8C6	S9W8C6_SCHCR	ILNGVSDAELYDLNECPYDQGGYFI	INGSEKVLIAQERSAANIVQVF-RKAAPSPYAYVA	223
2454	sp	Q8RQE9	RPOB_THET8	-----DGSFI	INGADRVIVSQIHRSPGVYFTDPDPA--PGRYIA	160
2455	ASR51304.1			-----NGTFV	INGTERVIVSQMHRSPGVLFPHDRGKTHSSGKLYF	184
2456	OXR47929.1			-----TGSFV	INGTERVIVSQLHRSPGVVFFEHDRGKTHSSGKLLF	176
2457	WP_093971860.1			-----TGSFV	INGTERVIVSQLHRSPGVVFFEHDRGKTHSSGKLLF	176
2458	sp	Q2NWR6	RPOB_SODGM	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2459	sp	B4EYU9	RPOB_PROMH	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2460	sp	A7FNI3	RPOB_YERP3	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2461	sp	Q1C1U1	RPOB_YERPA	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2462	sp	B2K113	RPOB_YERPB	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2463	sp	A8G8E7	RPOB_SERP5	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2464	sp	Q6DANO	RPOB_PECAS	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2465	sp	C6DHR5	RPOB_PECBP	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2466	sp	Q7N9A4	RPOB_PHOLL	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2467	sp	C5BHE3	RPOB_EDWI9	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2468	sp	A7MQQ9	RPOB_CROS8	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2469	sp	B5XYF5	RPOB_KLEP3	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2470	sp	P0A8V2	RPOB_ECOLI	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2471	sp	C5A0S7	RPOB_ECOBW	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2472	sp	Q31U10	RPOB_SHIBS	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2473	sp	Q32AF9	RPOB_SHIDS	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2474	sp	A8AKT9	RPOB_CITK8	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2475	sp	B5RFK1	RPOB_SALG2	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2476	sp	B5BJQ3	RPOB_SALPK	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2477	sp	B4T0Y9	RPOB_SALNS	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2478	sp	P06173	RPOB_SALTY	-----NGTFV	INGTERVIVSQLHRSPGVVFFSDSKGKTHSSGKVLV	172
2479						
2480						
2481	sp	P30876	RPB2_HUMAN	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2482	tr	G3V8Y5	G3V8Y5_RAT	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2483	tr	A0A250Y753	A0A250Y753_CASCN	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2484	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2485	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2486	tr	I3M351	I3M351_ICTTR	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2487	tr	G7P5R6	G7P5R6_MACFA	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2488	tr	H2QP18	H2QP18_PANTR	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2489	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2490	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2491	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	494
2492	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	494
2493	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2494	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501
2495	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	YSLATGNWGDQKKAHQARAGVSVQLNRLTF	ASTLSHLRRLNSP---IGRDGHLAKP	501

2496	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	YSLATGNWGDQKKAHQARAGVSVQLNRLTFASLTLSHLRLNSP----	IGRDKLAKP	501
2497	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	YSLATGNWGDQKKAHQARAGVSVQLNRLTFASLTLSHLRLNSP----	IGRDKLAKP	494
2498	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	YSLATGNWGDQKKAHQARAGVSVQLNRLTFASLTLSHLRLNSP----	IGRDKLAKP	501
2499	tr	A0A096NEY4	A0A096NEY4_PAPAN	YSLATGNWGDQKKAHQARAGVSVQLNRLTFASLTLSHLRLNSP----	IGRDKLAKP	501
2500	tr	C9J2Y9	C9J2Y9_HUMAN	YSLATGNWGDQKKAHQARAGVSVQLNRLTFASLTLSHLRLNSP----	IGRDKLAKP	494
2501	tr	G8BY61	G8BY61_TETPH	YALATGNWGEQKAMTSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	512
2502	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	510
2503	tr	J7RV95	J7RV95_KAZNA	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	510
2504	tr	H2AVJ8	H2AVJ8_KAZAF	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	510
2505	sp	Q6FLD5	RPB2_CANGA	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	512
2506	sp	P08518	RPB2_YEAST	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	514
2507	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	514
2508	tr	A0A0L8RB33	A0A0L8RB33_SACEU	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	514
2509	tr	GOVJ71	GOVJ71_NAUCC	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	514
2510	tr	G8ZM49	G8ZM49_TORDC	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	512
2511	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	514
2512	tr	A0A0N7IS35	A0A0N7IS35_9SACH	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	512
2513	tr	A0A212MG88	A0A212MG88_ZYGBA	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	513
2514	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	513
2515	tr	S6ESB4	S6ESB4_ZYGB2	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	513
2516	tr	B6K5Q5	B6K5Q5_SCHJY	YSLATGNWGDQKRGLANRVGVSQVLNRYTFASLTLSHLRRNTNP----	IGRDKLAKP	500
2517	sp	Q02061	RPB2_SCHPO	YALATGNWGEQKAMSSRAGVSVQLNRYTYSSTLSHLRRNTNP----	IGRDKLAKP	500
2518	tr	S9R8U4	S9R8U4_SCHOY	YSLATGNWGDQKRSMNVRVGSQVLNRYTFASLTLSHLRRNTNP----	IGRDKLAKP	500
2519	tr	S9W8C6	S9W8C6_SCHCR	YSLATGNWGDQKRSMNVRVGSQVLNRYTFASLTLSHLRRNTNP----	IGRDKLAKP	500x8
2520	sp	Q8RQE9	RPB2_THET8	-----	EFFRSQSLQSFQKDETNPLSLRHKRI	430x8
2521	ASR51304.1			-----	EFFGSSQSLQSFMDQTNPLAEVTHKRRV	564
2522	OXK47929.1			-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	556
2523	WP_093971860.1			-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	556
2524	sp	Q2NWR6	RPB2_SODGM	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2525	sp	B4EYU9	RPB2_PROMH	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2526	sp	A7FNI3	RPB2_YERP3	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2527	sp	Q1C1U1	RPB2_YERPA	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2528	sp	B2K113	RPB2_YERPB	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2529	sp	A8G8E7	RPB2_SERP5	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2530	sp	Q6DANO	RPB2_PECAS	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2531	sp	C6DHR5	RPB2_PECCEP	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2532	sp	Q7N9A4	RPB2_PHOLL	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2533	sp	C5BHE3	RPB2_EDWI9	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2534	sp	A7MQQ9	RPB2_CROS8	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2535	sp	B5XYF5	RPB2_KLEP3	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2536	sp	POA8V2	RPB2_ECOLI	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2537	sp	C5A0S7	RPB2_ECOWB	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2538	sp	Q31U10	RPB2_SHIBS	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2539	sp	Q32AF9	RPB2_SHIDS	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2540	sp	A8AKT9	RPB2_CITK8	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2541	sp	B5RFK1	RPB2_SALG2	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2542	sp	B5BJQ3	RPB2_SALPK	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2543	sp	B4T0Y9	RPB2_SALNS	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2544	sp	P06173	RPB2_SALTY	-----	EFFGSSQSLQSFMDQTNPLSEITHKRRV	550
2545				*** . . . : : * * * . : : . * . *		
2546						
2547	sp	P30876	RPB2_HUMAN	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2548	tr	G3V8Y5	G3V8Y5_RAT	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2549	tr	A0A250Y753	A0A250Y753_CASCN	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2550	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2551	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2552	tr	I3M351	I3M351 ICTTR	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2553	tr	G7P5R6	G7P5R6_MACFA	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2554	tr	H2QPI8	H2QPI8_PANTR	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2555	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2556	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2557	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		554
2558	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		554
2559	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2560	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2561	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2562	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2563	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		554
2564	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2565	tr	A0A096NEY4	A0A096NEY4_PAPAN	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		561
2566	tr	C9J2Y9	C9J2Y9_HUMAN	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		554
2567	tr	G8BY61	G8BY61_TETPH	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		572
2568	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		570
2569	tr	J7RV95	J7RV95_KAZNA	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		570
2570	tr	H2AVJ8	H2AVJ8_KAZAF	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		570
2571	sp	Q6FLD5	RPB2_CANGA	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		572
2572	sp	P08518	RPB2_YEAST	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		574
2573	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		574
2574	tr	A0A0L8RB33	A0A0L8RB33_SACEU	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		574
2575	tr	GOVJ71	GOVJ71_NAUCC	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		574
2576	tr	G8ZM49	G8ZM49_TORDC	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		572
2577	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		574
2578	tr	A0A0N7IS35	A0A0N7IS35_9SACH	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		572
2579	tr	A0A212MG88	A0A212MG88_ZYGBA	HNTLWGLVCPAETPEGHAVGLVKNLALMAYISVGSQSPPILEFLEEWSMENLEEISPAAI		573

2664	sp	Q7N9A4	RPOB_PHOLL	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2665	sp	C5BHE3	RPOB_EDW19	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2666	sp	A7MQQ9	RPOB_CROS8	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2667	sp	B5XYF5	RPOB_KLEP3	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2668	sp	POA8V2	RPOB_ECOLI	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2669	sp	C5A0S7	RPOB_ECOWB	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2670	sp	Q31U10	RPOB_SHIBS	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2671	sp	Q32AF9	RPOB_SHIDS	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2672	sp	A8AKT9	RPOB_CITK8	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2673	sp	B5RFK1	RPOB_SALG2	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2674	sp	B5BJQ3	RPOB_SALPK	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2675	sp	B4T0Y9	RPOB_SALNS	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2676	sp	P06173	RPOB_SALTY	VVSVGASLIPFLEHDDANRALMGANMQRQAVPTLRADKPLVGTGMERAVAVDSGVTSAVK	719
2677				... : . : ** : : : * : * : * : : . : . :	
2678					
2679	sp	P30876	RBP2_HUMAN	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2680	tr	G3V8Y5	G3V8Y5_RAT	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2681	tr	A0A250Y753	A0A250Y753_CASCN	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2682	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2683	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2684	tr	I3M351	I3M351 ICTTR	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2685	tr	G7P5R6	G7P5R6_MACFA	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2686	tr	H2QP18	H2QP18_PANTR	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2687	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2688	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2689	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	810
2690	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	810
2691	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2692	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2693	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2694	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2695	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2696	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	810
2697	tr	A0A096NEY4	A0A096NEY4_PAPAN	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	817
2698	tr	C9J2Y9	C9J2Y9_HUMAN	-----SMEYLRFRELPAGINSIVAIASYTPTGQEDWIMNRSVAVDRGFFPSVFFRYSYKEQ	810
2699	tr	G8BY61	G8BY61_TETPH	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	862
2700	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	-----SMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	859
2701	tr	J7RV95	J7RV95_KAZNA	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	858
2702	tr	H2AVJ8	H2AVJ8_KAZAF	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	860
2703	sp	Q6FLD5	RBP2_CANGA	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	861
2704	sp	P08518	RBP2_YEAST	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	862
2705	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	862
2706	tr	A0A0L8RB33	A0A0L8RB33_SACEU	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	862
2707	tr	GOVJ71	GOVJ71_NAUCC	-----SMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	862
2708	tr	G8ZM49	G8ZM49_TORDC	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	860
2709	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	862
2710	tr	A0A0N7IS35	A0A0N7IS35_9SACH	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	860
2711	tr	A0A212MG88	A0A212MG88_ZYGBA	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	861
2712	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	861
2713	tr	S6ESB4	S6ESB4_ZYGB2	-----AMEYKLFRELPAGONAIVAIACYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	861
2714	tr	B6K5Q5	B6K5Q5_SCHJY	-----SMEYKLFRELPAGONAIVAILCYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	851
2715	sp	Q02061	RBP2_SCHPO	-----SMEYKLFRELPAGONAIVAILCYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	851
2716	tr	S9R8U4	S9R8U4_SCHOY	-----SMEYKLFRELPAGONAIVAILCYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	851
2717	tr	S9W8C6	S9W8C6_SCHCR	-----SMEYKLFRELPAGONAIVAILCYSPTGQEDWIMNQSSIDRGLFRLSFFRYSYMDQ	851x6
2718	sp	Q8RQE9	RPOB_THET8	GDLLADGPAENGFALGQNVLAIMPFDGYNFEDSLVISELLKRDFTYTSIHIEREYIE	711
2719		ASR51304.1		GDIIADGPSTDLGELALGQNVLAIMPFDGYNFEDSLVISELLKRDFTYTSIHIEREYIE	852
2720		OXR47929.1		GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	845
2721		WP_093971860.1		GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	845
2722	sp	Q2NWR6	RPOB_SODGM	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2723	sp	B4EYU9	RPOB_PROMH	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2724	sp	A7FNI3	RPOB_YERP3	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2725	sp	Q1C1U1	RPOB_YERPA	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2726	sp	B2K113	RPOB_YERPB	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2727	sp	A8G8E7	RPOB_SERP5	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2728	sp	Q6DAN0	RPOB_PECAS	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2729	sp	C6DHR5	RPOB_PECPC	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2730	sp	Q7N9A4	RPOB_PHOLL	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2731	sp	C5BHE3	RPOB_EDW19	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2732	sp	A7MQQ9	RPOB_CROS8	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2733	sp	B5XYF5	RPOB_KLEP3	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2734	sp	POA8V2	RPOB_ECOLI	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2735	sp	C5A0S7	RPOB_ECOWB	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2736	sp	Q31U10	RPOB_SHIBS	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2737	sp	Q32AF9	RPOB_SHIDS	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2738	sp	A8AKT9	RPOB_CITK8	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2739	sp	B5RFK1	RPOB_SALG2	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2740	sp	B5BJQ3	RPOB_SALPK	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2741	sp	B4T0Y9	RPOB_SALNS	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2742	sp	P06173	RPOB_SALTY	GDVLADGASTDLGELALGQNMVAFMPWNGYNFEDSLVISELLKRDFTYTSIHIEREYIE	839
2743				* * * : * : * * : : : * * * : : . : . : . :	
2744					
2745					
2746					
2747					

2832	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	IGDKFASRHGQKGTFCGIQYRQEDMPFTCEGITPDIINPHAIPSRMTIGHLECLQGVKS	983
2833	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	IGDKFASRHGQKGTFCGIQYRQEDMPFTCEGITPDIINPHAIPSRMTIGHLECLQGVKS	990
2834	tr	A0A096NEY4	A0A096NEY4_PAPAN	IGDKFASRHGQKGTFCGIQYRQEDMPFTCEGITPDIINPHAIPSRMTIGHLECLQGVKS	990
2835	tr	C9J2Y9	C9J2Y9_HUMAN	IGDKFASRHGQKGTFCGIQYRQEDMPFTCEGITPDIINPHAIPSRMTIGHLECLQGVKS	983
2836	tr	G8BY61	G8BY61_TETPH	IGDKFASRHGQKGTIGITYGREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1035
2837	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	IGDKFASRHGQKGTIGITYRREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1032
2838	tr	J7RV95	J7RV95_KAZNA	IGDKFASRHGQKGTIGITYSREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1031
2839	tr	H2AVJ8	H2AVJ8_KAZAF	IGDKFASRHGQKGTIGITYRREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1033
2840	sp	Q6FLD5	RBP2_CANGA	IGDKFASRHGQKGTIGITYRREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1034
2841	sp	P08518	RBP2_YEAST	IGDKFASRHGQKGTIGITYRREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1035
2842	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	IGDKFASRHGQKGTIGITYRREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1035
2843	tr	A0A0L8RB33	A0A0L8RB33_SACEU	IGDKFASRHGQKGTIGITYRREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1035
2844	tr	GOVJ71	GOVJ71_NAUCC	IGDKFASRHGQKGTIGITYRREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1035
2845	tr	G8ZM49	G8ZM49_TORDC	IGDKFASRHGQKGTIGITYGREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1033
2846	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	IGDKFASRHGQKGTIGITYGREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1035
2847	tr	A0A0N7IS35	A0A0N7IS35_9SACH	IGDKFASRHGQKGTIGITYSREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1033
2848	tr	A0A212MG88	A0A212MG88_ZYGBA	IGDKFASRHGQKGTIGITYSREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1034
2849	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	IGDKFASRHGQKGTIGITYSREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1034
2850	tr	S6ESB4	S6ESB4_ZYGB2	IGDKFASRHGQKGTIGITYSREDMPTAEGIVPDLIINPHAIPSRMTVAHLECLLSKVA	1034
2851	tr	B6K5Q5	B6K5Q5_SCHJY	IGDKFASRHGQKGTIGMTRYREDMPTSAQGVVDIIINPHAIPSRMTVAHLECLLSKVS	1024
2852	sp	Q02061	RBP2_SCHPO	IGDKFASRHGQKGTIGMTRYREDMPTSAQGVVDIIINPHAIPSRMTVAHLECLLSKVS	1024
2853	tr	S9R8U4	S9R8U4_SCHOY	IGDKFASRHGQKGTIGMTRYREDMPTSAQGVVDIIINPHAIPSRMTVAHLECLLSKVS	1024
2854	tr	S9W8C6	S9W8C6_SCHCR	IGDKFASRHGQKGTIGMTRYREDMPTSAQGVVDIIINPHAIPSRMTVAHLECLLSKVS	1024
2855	sp	Q8RQE9	RPOB_THET8	VGDKLANRHGKGVVAKILPVEDMPLPDGTPVDVILNPLGVPSRMNIGQIILETHLGLAG	894
2856	ASR51304.1			PGDKMAGRHNKGVIISRIIPVEDMPLFEDGTHVDFVLNPLGVPSRMNIGQIILETHLGLAA	1147
2857	OXR47929.1			PGDKMAGRHNKGVIISRIIPVEDMPLHMDGTPADIVLNPGLVPSRMNIGQIILETHLGLAA	1140
2858	WP_093971860.1			PGDKMAGRHNKGVIISRIIPVEDMPLHMDGTPADIVLNPGLVPSRMNIGQIILETHLGLAA	1140
2859	sp	Q2NWR6	RPOB_SODGM	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2860	sp	B4EYU9	RPOB_PROMH	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2861	sp	A7FN13	RPOB_YERP3	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2862	sp	Q1C1U1	RPOB_YERPA	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2863	sp	B2K113	RPOB_YERPB	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2864	sp	A8G8E7	RPOB_SERP5	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2865	sp	Q6DAN0	RPOB_PECAS	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2866	sp	C6DHR5	RPOB_PECPC	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2867	sp	Q7N9A4	RPOB_PHOLL	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2868	sp	C5BHE3	RPOB_EDWI9	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2869	sp	A7MQQ9	RPOB_CROS8	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2870	sp	B5XYF5	RPOB_KLEP3	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2871	sp	P0A8V2	RPOB_ECOLI	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2872	sp	C5A0S7	RPOB_ECOWB	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2873	sp	Q31U10	RPOB_SIBS3	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2874	sp	Q32AF9	RPOB_SHIDS	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2875	sp	A8AKT9	RPOB_CITK8	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2876	sp	B5RFK1	RPOB_SALG2	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2877	sp	B5BJQ3	RPOB_SALPK	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2878	sp	B4T0Y9	RPOB_SALNS	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2879	sp	P06173	RPOB_SALTY	PGDKMAGRHNKGVIISRIIPVEDMPLYDENGTPVDIVLNPGLVPSRMNIGQIILETHLGLAA	1121
2880				***.*.***.*	
2881				****.*	
2882				*.:.:*	
2883	sp	P30876	RBP2_HUMAN	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2884	tr	G3V8Y5	G3V8Y5_RAT	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2885	tr	A0A250Y753	A0A250Y753_CASCN	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2886	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2887	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2888	tr	I3M351	I3M351_ICTTR	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2889	tr	G7P5R6	G7P5R6_MACFA	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2890	tr	H2QPI8	H2QPI8_PANTR	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2891	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2892	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2893	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1087
2894	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1087
2895	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2896	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2897	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2898	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2899	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1087
2900	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2901	tr	A0A096NEY4	A0A096NEY4_PAPAN	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1094
2902	tr	C9J2Y9	C9J2Y9_HUMAN	RKITSQIFIGPTYYQRLKHMVDDKIHSRARGPIQILNRPQMEGRSRDGGIRFGEMERDCQ	1087
2903	tr	G8BY61	G8BY61_TETPH	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1138
2904	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1135
2905	tr	J7RV95	J7RV95_KAZNA	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1134
2906	tr	H2AVJ8	H2AVJ8_KAZAF	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1136
2907	sp	Q6FLD5	RBP2_CANGA	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1137
2908	sp	P08518	RBP2_YEAST	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1138
2909	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1138
2910	tr	A0A0L8RB33	A0A0L8RB33_SACEU	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1138
2911	tr	GOVJ71	GOVJ71_NAUCC	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1138
2912	tr	G8ZM49	G8ZM49_TORDC	KKLMAQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1136
2913	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	KKLMSQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1138
2914	tr	A0A0N7IS35	A0A0N7IS35_9SACH	KKLMSQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1136
2915	tr	A0A212MG88	A0A212MG88_ZYGBA	KKLMSQIFFGPTYYQRLRHMVDDKIHSRARGPMQVLTROPVEGRSRDGGIRFGEMERDCM	1137

2916	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	KKLMSQIFFGPTYQRLRHHVDDKIHARARGPMQVLT	TRQPVV	GRSRDGGI	RF	GEMERDCM	1137		
2917	tr	S6ESB4	S6ESB4_ZYGB2	KKLMSQIFFGPTYQRLRHHVDDKIHARARGPMQVLT	TRQPVV	GRSRDGGI	RF	GEMERDCM	1137		
2918	tr	B6K5Q5	B6K5Q5_SCHJY	RKLVAQVFLGPTYQRLKHLVDDKIHARARGPVQIL	TRQPVV	GRSRDGGI	RF	GEMERDCQ	1127		
2919	sp	Q02061	RBP2_SCHPO	RKLVAQVFLGPTYQRLKHLVDDKIHARARGPVQIL	TRQPVV	GRSRDGGI	RF	GEMERDCQ	1127		
2920	tr	S9R8U4	S9R8U4_SCHOY	RKLVSQVFLGPTYQRLKHLVDDKIHARARGPVQIL	TRQPVV	GRSRDGGI	RF	GEMERDCQ	1127		
2921	tr	S9W8C6	S9W8C6_SCHCR	RKLVSQVFLGPTYQRLKHLVDDKIHARARGPVQIL	TRQPVV	GRSRDGGI	RF	GEMERDCQ	1127		
2922	sp	Q8RQE9	RPOB_THET8	EPIEGPIVVQGMFIMKLYHIVDDKMHARSTG	YPYSL	IT	QQPL	GGKAQFGG	RF	GEMEVWAL	1040
2923		ASR51304.1		DKFDRKVTVGYIYMLKLHHLVDDKIHARST	IGPYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1310
2924		OXR47929.1		EQFERPVTVGMYMLKLHHLVDDKMHARST	IGPYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1306
2925		WP_093971860.1		EQFERPVTVGMYMLKLHHLVDDKMHARST	IGPYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1306
2926	sp	Q2NWR6	RPOB_SODGM	EQFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2927	sp	B4EYU9	RPOB_PROMH	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2928	sp	A7FNI3	RPOB_YERP3	EQFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2929	sp	Q1C1U1	RPOB_YERPA	EQFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2930	sp	B2K113	RPOB_YERPB	EQFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2931	sp	A8G8E7	RPOB_SERP5	EQFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2932	sp	Q6DANO	RPOB_PECCAS	EQFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2933	sp	C6DHR5	RPOB_PECCP	EKFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2934	sp	Q7N9A4	RPOB_PHOLL	EQFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2935	sp	C5BHE3	RPOB_EDWI9	ERFERQVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2936	sp	A7MQQ9	RPOB_CROS8	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2937	sp	B5XYF5	RPOB_KLEP3	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2938	sp	POA8V2	RPOB_ECOLI	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2939	sp	C5A0S7	RPOB_ECOWB	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2940	sp	Q31U10	RPOB_SHIBS	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2941	sp	Q32AF9	RPOB_SHIDS	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2942	sp	A8AKT9	RPOB_CITK8	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2943	sp	B5RFK1	RPOB_SALG2	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2944	sp	B5BJQ3	RPOB_SALPK	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2945	sp	B4T0Y9	RPOB_SALNS	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2946	sp	P06173	RPOB_SALTY	EQFERPVTVGMYMLKLHHLVDDKMHARST	IGSYSL	VT	QQPL	GGKAQFGG	RF	GEMEVWAL	1278
2947				:	:	:	:	:	:	:	:
2948				:	:	:	:	:	:	:	:
2949	sp	P30876	RBP2_HUMAN	ACKLLFQELMSMSIAPRMMSV							1174
2950	tr	G3V8Y5	G3V8Y5_RAT	ACKLLFQELMSMSIAPRMMSV							1174
2951	tr	A0A250Y753	A0A250Y753_CASCN	ACKLLFQELMSMSIAPRMMSV							1174
2952	tr	A0A1U7R4C7	A0A1U7R4C7_MESAU	ACKLLFQELMSMSIAPRMMSV							1174
2953	tr	A0A286XIQ9	A0A286XIQ9_CAVPO	ACKLLFQELMSMSIAPRMMSV							1174
2954	tr	I3M351	I3M351_ICTTR	ACKLLFQELMSMSIAPRMMSV							1174
2955	tr	G7P5R6	G7P5R6_MACFA	ACKLLFQELMSMSIAPRMMSV							1174
2956	tr	H2QP18	H2QP18_PANTR	ACKLLFQELMSMSIAPRMMSV							1174
2957	tr	A0A1U7V0T5	A0A1U7V0T5_TARSY	ACKLLFQELMSMSIAPRMMSV							1174
2958	tr	A0A1S2ZSL2	A0A1S2ZSL2_ERIEU	ACKLLFQELMSMSIAPRMMSV							1174
2959	tr	A0A0D9QYL1	A0A0D9QYL1_CHLSB	ACKLLFQELMSMSIAPRMMSV							1167
2960	tr	A0A2K5ZNR7	A0A2K5ZNR7_MANLE	ACKLLFQELMSMSIAPRMMSV							1167
2961	tr	A0A2I2ZIU3	A0A2I2ZIU3_GORGO	ACKLLFQELMSMSIAPRMMSV							1174
2962	tr	A0A1D5QGA5	A0A1D5QGA5_MACMU	ACKLLFQELMSMSIAPRMMSV							1174
2963	tr	A0A2J8S2N1	A0A2J8S2N1_PONAB	ACKLLFQELMSMSIAPRMMSV							1174
2964	tr	A0A2K5K5J5	A0A2K5K5J5_COLAP	ACKLLFQELMSMSIAPRMMSV							1174
2965	tr	A0A2J8PEW7	A0A2J8PEW7_PANTR	ACKLLFQELMSMSIAPRMMSV							1167
2966	tr	A0A2K5CY83	A0A2K5CY83_AOTNA	ACKLLFQELMSMSIAPRMMSV							1174
2967	tr	A0A096NEY4	A0A096NEY4_PAPAN	ACKLLFQELMSMSIAPRMMSV							1174
2968	tr	C9J2Y9	C9J2Y9_HUMAN	ACKLLFQELMSMSIAPRMMSV							1167
2969	tr	G8BY61	G8BY61_TETPH	AAKLLFQELMAMNITPRLYTDRSKNF							1224
2970	tr	A0A1X7QYA1	A0A1X7QYA1_9SACH	AAKLLFQELMAMNITPRLYTDRSRDF							1221
2971	tr	J7RV95	J7RV95_KAZNA	AAKLLFQELMAMNITPRLYTDRSRDF							1220
2972	tr	H2AVJ8	H2AVJ8_KAZAF	AAKLLFQELMAMNITPRLYTDRSRDF							1222
2973	sp	Q6FLD5	RBP2_CANGA	AAKLLFQELMAMNITPRLYTDRSRDF							1223
2974	sp	P08518	RBP2_YEAST	AAKLLFQELMAMNITPRLYTDRSRDF							1224
2975	tr	A0A0L8VHA5	A0A0L8VHA5_9SACH	AAKLLFQELMAMNITPRLYTDRSRDF							1224
2976	tr	A0A0L8RB33	A0A0L8RB33_SACEU	AAKLLFQELMAMNITPRLYTDRSRDF							1224
2977	tr	G0VJ71	G0VJ71_NAUCC	AAKLLFQELMAMNITPRLYTDRSRDF							1224
2978	tr	G8ZM49	G8ZM49_TORDC	AAKLLFQELMAMNITPRLYTDRSKDF							1222
2979	tr	A0A1Q3A090	A0A1Q3A090_ZYGRO	AAKLLFQELMAMNITPRLYTDRSKDF							1224
2980	tr	A0A0N7IS35	A0A0N7IS35_9SACH	AAKLLFQELMAMNITPRLYTDRSKDF							1222
2981	tr	A0A212MG88	A0A212MG88_ZYGBA	AAKLLFQELMAMNITPRLYTDRSKDF							1223
2982	tr	A0A1S7HHE1	A0A1S7HHE1_9SACH	AAKLLFQELMAMNITPRLYTDRSKDF							1223
2983	tr	S6ESB4	S6ESB4_ZYGB2	AAKLLFQELMAMNITPRLYTDRSKDF							1223
2984	tr	B6K5Q5	B6K5Q5_SCHJY	AAKLLFQELMSMNIAPRLFTHSHH							1210
2985	sp	Q02061	RBP2_SCHPO	AAKLLFQELMSMNIAPRLFTHNHK							1210
2986	tr	S9R8U4	S9R8U4_SCHOY	AAKLLFQELMSMNIAPRLFTHNHI							1211
2987	tr	S9W8C6	S9W8C6_SCHCR	AAKLLFQELMSMNIAPRLFTHNHN							1211

2990	sp Q8RQE9 RPOB_THET8	SFNVLVKELQALALD	VQTLDEKDN--PVDIFEGLASKR	1119
2991	ASR51304.1	SFNVLVKEMRSLGLN	VELNSIDALPDPDEIAEAAE---	1388
2992	OXR47929.1	SFNVLVKEIRSLSLD	MDLERN-----	1370
2993	WP_093971860.1	SFNVLVKEIRSLSLD	MDLERN-----	1370
2994	sp Q2NWR6 RPOB_SODGM	SFNVLLKEIRSLGINI	ELEED-----	1342
2995	sp B4EYU9 RPOB_PROMH	SFNVLLKEIRSLGINI	ELEDE-----	1342
2996	sp A7FNI3 RPOB_YERP3	SFNVLLKEIRSLGINI	ELEEE-----	1342
2997	sp Q1C1U1 RPOB_YERPA	SFNVLLKEIRSLGINI	ELEEE-----	1342
2998	sp B2K113 RPOB_YERPB	SFNVLLKEIRSLGINI	ELEEE-----	1342
2999	sp A8G8E7 RPOB_SERP5	SFNVLLKEIRSLGINI	ELEGE-----	1342
3000	sp Q6DAN0 RPOB_PECAS	SFNVLLKEIRSLGINI	ELEEK-----	1342
3001	sp C6DHR5 RPOB_PECCP	SFNVLLKEIRSLGINI	ELEEE-----	1342
3002	sp Q7N9A4 RPOB_PHOLL	SFNVLLKEIRSLGINI	ELEGE-----	1342
3003	sp C5BHE3 RPOB_EDWI9	SFNVLLKEIRSLGINI	ELEDE-----	1342
3004	sp A7MQQ9 RPOB_CROS8	SFNVLLKEIRSLGINI	ELEDE-----	1342
3005	sp B5XYF5 RPOB_KLEP3	SFNVLLKEIRSLGINI	ELEDE-----	1342
3006	sp P0A8V2 RPOB_ECOLI	SFNVLLKEIRSLGINI	ELEDE-----	1342
3007	sp C5A0S7 RPOB_ECOBW	SFNVLLKEIRSLGINI	ELEDE-----	1342
3008	sp Q31U10 RPOB_SHIBS	SFNVLLKEIRSLGINI	ELEDE-----	1342
3009	sp Q32AF9 RPOB_SHIDS	SFNVLLKEIRSLGINI	ELEDE-----	1342
3010	sp A8AKT9 RPOB_CITK8	SFNVLLKEIRSLGINI	ELEDE-----	1342
3011	sp B5RFR1 RPOB_SALG2	SFNVLLKEIRSLGINI	ELEDE-----	1342
3012	sp B5BJQ3 RPOB_SALPK	SFNVLLKEIRSLGINI	ELEDE-----	1342
3013	sp B4T0Y9 RPOB_SALNS	SFNVLLKEIRSLGINI	ELEDE-----	1342
3014	sp P06173 RPOB_SALTY	SFNVLLKEIRSLGINI	ELEDE-----	1342
3015		: : * : * : : : :		
3016				

Fig. 5 Mix and Match analysis of the initiation subunits β of *E. coli* and Rpb2 of yeast RNAP-II
For legends follow Figures 3-5.

Figure 6 shows the mix and match analysis of the elongation subunits from pro- and eukaryotes MSU RNAPs. The active site regions are highlighted in yellow and the representative sequences are highlighted in yellow. In eukaryotic elongation subunits, the catalytic region is brought to the N-terminal region whereas it is found very close to the C-terminal region. There are 4 Cs at the N-terminal region in prokaryotic elongation subunits (out of which 2 are aligning in both) suggesting a possible additional Zn binding region. There are a very few motifs aligning in both. Significant among them are -GHIELA-, -NLM/LGKRVDF/YS-, possible metal binding region -DFDGDE/QM-, -DTAV/LKTAE/NT/SGYI/L-, -L/V/IAAQSIGEPA/GTQM/LTL/MXTFH-. The metal binding region -NADFDGD- is aligning in both and placed in the middle of the enzyme. The C-terminal ends in eukaryotes usually end in -PDDSD~~E~~/~~E~~/DN- (except in yeasts and Alligator) and -GSD~~N~~/~~E~~- in prokaryotes, suggesting a possible metal binding site (DxD) at the ends. It is interesting to note that the prokaryotic active site is placed at the C-terminal region (~900 amino acids) whereas the eukaryotic active site is placed at the N-terminal region (~90 amino acids) with their built-in Zn binding motifs.

3034 CLUSTAL O(1.2.4):MSA of eubacterial β' and eukaryotic Rpb1 subunits 6

3035	tr	A0A1U8DYN0 A0A1U8DYN0_ALLSI	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	99	
3036	tr	A0A1L8H4P4 A0A1L8H4P4_XENLA	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3037	tr	H9GLG5 H9GLG5_ANOCA	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	99	
3038	tr	H2R1J6 H2R1J6_PANTR	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3039	sp	P08775 RPB1_MOUSE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3040	tr	G1MCZ1 G1MCZ1_ALLME	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3041	tr	O08847 O08847_MOUSE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3042	tr	S7PWZ6 S7PWZ6_MYOBR	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3043	tr	D4A5A6 D4A5A6_RAT	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3044	tr	A0A1S3EWL2 A0A1S3EWL2_DIPOR	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3045	sp	P11414 RPB1_CRIGR	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3046	tr	O35559 O35559_CRIGR	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3047	tr	A0A2I3M9H2 A0A2I3M9H2_PAPAN	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3048	tr	F7HB40 F7HB40_MACMU	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3049	tr	A0A2K6RYW9 A0A2K6RYW9_SAIBB	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3050	tr	W5N8Z6 W5N8Z6_LEPOC	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	100	
3051	tr	I3JRW6 I3JRW6_ORENI	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3052	tr	A0A0R4IMS9 A0A0R4IMS9_DANRE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3053	tr	A0A1A7X327 A0A1A7X327_9TELE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3054	tr	A0A1A8UKD7 A0A1A8UKD7_NOTFU	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3055	tr	A0A1A8ER05 A0A1A8ER05_9TELE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3056	tr	A0A1A8DQ60 A0A1A8DQ60_9TELE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3057	tr	A0A1A8NSR8 A0A1A8NSR8_9TELE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3058	tr	A0A1W4YLM7 A0A1W4YLM7_9TELE	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3059	tr	A0A1M8A6L7 A0A1M8A6L7_MALS4	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3060	tr	A0A1M8A6L7 A0A1M8A6L7_MALS4	DP	ROGVIERTGR	-----	COTCA	-GNM--TE	CGHFGHIELAKPVFHVGF	98	
3061	sp	P04050 RPB1_YEAST	DP	RLGSDIRNFK	-----	COTCE	-GM--NE	CGHFGHIELAKPVFHVGF	96	
3062	tr	A0A1B2J8C6 A0A1B2J8C6_PICPA	DP	RLGSDIRNFK	-----	COTCE	-GM--AE	CGHFGHIELAKPVFHVGF	96	
3063	tr	F2QW17 F2QW17_KOMPC	DP	RLGSDIRNFK	-----	COTCE	-GM--AE	CGHFGHIELAKPVFHVGF	96	
3064	tr	A3GID7 A3GID7_PICST	DP	RLGSDIRNFK	-----	COTCE	-DM--AE	CGHFGHIELAKPVFHVGF	97	
3065	tr	A0A1D8PUA6 A0A1D8PUA6_CANAL	DP	RLGSDIRNFK	-----	COTCE	-DM--AE	CGHFGHIELAKPVFHVGF	97	
3066	tr	G8BEH9 G8BEH9_CANPC	DP	RLGSDIRNFK	-----	COTCE	-DM--AE	CGHFGHIELAKPVFHVGF	976,19	
3067	tr	AEQ34223.1	DER	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3068	tr	ASR51305.1	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3069	tr	OKR47930.1	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3070	sp	A7MQQ8 RPOC_CROS8	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3071	sp	Q32AG0 RPOC_SHIDS	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3072	sp	Q0SY12 RPOC_SHIF8	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3073	sp	B2TWH4 RPOC_SHIB3	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3074	sp	P0A8T7 RPOC_ECOLI	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3075	sp	Q3YU26 RPOC_SHISS	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3076	sp	B1XBZ0 RPOC_ECODH	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3077	sp	A8A787 RPOC_ECOHS	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3078	tr	A0A237JUP3 A0A237JUP3_SHISO	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3079	tr	A0A0F1RFB2 A0A0F1RFB2_ENTAS	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3080	tr	A0A1B3EWG0 A0A1B3EWG0_ENTCL	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3081	tr	A0A0F0XM62 A0A0F0XM62_9ENTR	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3082	sp	Q5PK92 RPOC_SALPA	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3083	sp	A9MHE9 RPOC_SALAR	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3084	tr	A0A232XM43 A0A232XM43_SALMU	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3085	tr	B5RFK0 B5RFK0_SALG2	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3086	sp	P0A2R5 RPOC_SALTI	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3087	sp	Q57H68 RPOC_SALCH	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3088	sp	P0A2R4 RPOC_SALTY	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3089	sp	A6TGP1 RPOC_KLEP7	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3090	tr	A0A0J2K6S7 A0A0J2K6S7_9ENTR	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3091	tr	A0A0G3RZQ0 A0A0G3RZQ0_KLEOX	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3092	tr	A0A212HDS5 A0A212HDS5_9ENTR	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3093	tr	A0A1R0FP41 A0A1R0FP41_CITBR	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3094	tr	A0A078LHA5 A0A078LHA5_CITKO	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3095	sp	ABAKT8 RPOC_CITK8	CAR	IFGPKDYEL	CG	SKYKRLKHRGVI	CEK	DVEVTQTKVRRERM	GHIELASPTAH	117
3096			:	:	:	:	:	:	:	
3097			:	:	:	:	:	:	:	
3098			:	:	:	:	:	:	:	
3099	tr	A0A1U8DYN0 A0A1U8DYN0_ALLSI	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	373
3100	tr	A0A1L8H4P4 A0A1L8H4P4_XENLA	AT	MVDNE	IPLPRAMQKSGRPLKSVKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	372
3101	tr	H9GLG5 H9GLG5_ANOCA	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	373
3102	tr	H2R1J6 H2R1J6_PANTR	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3103	sp	P08775 RPB1_MOUSE	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3104	tr	G1MCZ1 G1MCZ1_ALLME	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3105	tr	O08847 O08847_MOUSE	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3106	tr	S7PWZ6 S7PWZ6_MYOBR	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3107	tr	D4A5A6 D4A5A6_RAT	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3108	tr	A0A1S3EWL2 A0A1S3EWL2_DIPOR	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3109	sp	P11414 RPB1_CRIGR	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3110	tr	O35559 O35559_CRIGR	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3111	tr	A0A2I3M9H2 A0A2I3M9H2_PAPAN	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3112	tr	F7HB40 F7HB40_MACMU	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3113	tr	A0A2K6RYW9 A0A2K6RYW9_SAIBB	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	374
3114	tr	W5N8Z6 W5N8Z6_LEPOC	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	373
3115	tr	I3JRW6 I3JRW6_ORENI	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	372
3116	tr	A0A0R4IMS9 A0A0R4IMS9_DANRE	AT	MVDN	LPLPRAMQKSGRPLKSLKQRLKGGK	EGRV	RGNLMGKR	VDFSARTVIT	PPDNLS	371

3201	sp	B1XBZ0	RPOC_ECODH	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3202	sp	A8A787	RPOC_ECOHS	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3203	tr	A0A237JUP3	A0A237JUP3_SHISO	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3204	tr	A0A0F1RBF2	A0A0F1RBF2_ENTAS	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3205	tr	A0A1B3EWG0	A0A1B3EWG0_ENTCL	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3206	tr	A0A0F0XM62	A0A0F0XM62_9ENTR	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3207	sp	Q5PK92	RPOC_SALPA	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3208	sp	A9MHE9	RPOC_SALAR	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3209	tr	A0A232XM43	A0A232XM43_SALMU	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3210	tr	B5RFK0	B5RFK0_SALG2	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3211	sp	POA2R5	RPOC_SALTI	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3212	sp	Q57H68	RPOC_SALCH	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3213	sp	POA2R4	RPOC_SALTY	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3214	sp	A6TGP1	RPOC_KLEP7	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3215	tr	A0A0J2K6S7	A0A0J2K6S7_9ENTR	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3216	tr	A0A0G3RZQ0	A0A0G3RZQ0_KLEOX	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3217	tr	A0A212HDS5	A0A212HDS5_9ENTR	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3218	tr	A0A1R0FP41	A0A1R0FP41_CITBR	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3219	tr	A0A078LHA5	A0A078LHA5_CITKO	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3220	sp	A8AKT8	RPOC_CITK8	EE----	AVVVDILDEVI	REHPV	LLNRAPTLHRL	GIQAFEPV	LIEGKAIQLHPLVCAAYNA	459
3221				:	:	:	:	:	:	:
3222				:	:	:	:	:	:	:
3223	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		553
3224	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3225	tr	H9GLG5	H9GLG5_ANOCA	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		553
3226	tr	H2R1J6	H2R1J6_PANTR	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3227	sp	P08775	RPB1_MOUSE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3228	tr	G1MCZ1	G1MCZ1_AILME	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3229	tr	O08847	O08847_MOUSE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3230	tr	S7PWZ6	S7PWZ6_MYOB	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3231	tr	D4A5A6	D4A5A6_RAT	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3232	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3233	sp	P11414	RPB1_CRIGR	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3234	tr	O35559	O35559_CRIGR	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3235	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3236	tr	F7HB40	F7HB40_MACMU	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3237	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		554
3238	tr	W5N8Z6	W5N8Z6_LEPOC	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		553
3239	tr	I3JRW6	I3JRW6_ORENI	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3240	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		551
3241	tr	A0A1A7X327	A0A1A7X327_9TELE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3242	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3243	tr	A0A1A8ER05	A0A1A8ER05_9TELE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3244	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3245	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3246	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	DFDGD	EMNLHLPQ	SLETRAEIQELAMVPRMIVTPQSNRPVMGIVQD	TLTAVRKF	TKRDVF		552
3247	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	DFDGD	EMNLHVPQ	SEEARAEIAI	IAWVPRQIVSPQANKPVMGIVQD	TLTAVRKF	TKRDVF	553
3248	sp	P04050	RPB1_YEAST	DFDGD	EMNLHVPQ	SEETRAELSQLCAVPLQIVSPQSNKPCMGIVQD	TLTAVRKF	TKRDVF		540
3249	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	DFDGD	EMNLHVPQ	SEETRAELSQLCAVPLQIVSPQSNKPCMGIVQD	TLTAVRKF	TKRDVF		541
3250	tr	F2QW17	F2QW17_KOMPC	DFDGD	EMNLHVPQ	SEETRAELSQLCAVPLQIVSPQSNKPCMGIVQD	TLTAVRKF	TKRDVF		541
3251	tr	A3GID7	A3GID7_PICST	DFDGD	EMNLHVPQ	SEETRAELSQLCAVPLQIVSPQSNKPCMGIVQD	TLTAVRKF	TKRDVF		540
3252	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	DFDGD	EMNLHVPQ	SEETRAELSQLCAVPLQIVSPQSNKPCMGIVQD	TLTAVRKF	TKRDVF		540
3253	tr	G8BEH9	G8BEH9_CANPC	DFDGD	EMNLHVPQ	SEETRAELSQLCAVPLQIVSPQSNKPCMGIVQD	TLTAVRKF	TKRDVF		540
3254		ABG34223.1		DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			797
3255		ASR51305.1		DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-E			519
3256		OKR47930.1		DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3257	sp	A7MQQ8	RPOC_CROS8	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3258	sp	Q32AG0	RPOC_SHIDS	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3259	sp	Q0SY12	RPOC_SHIF8	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3260	sp	B2TWH4	RPOC_SHIB3	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3261	sp	POA8T7	RPOC_ECOLI	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3262	sp	Q3YU26	RPOC_SHISS	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3263	sp	B1XBZ0	RPOC_ECODH	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3264	sp	A8A787	RPOC_ECOHS	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3265	tr	A0A237JUP3	A0A237JUP3_SHISO	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3266	tr	A0A0F1RBF2	A0A0F1RBF2_ENTAS	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3267	tr	A0A1B3EWG0	A0A1B3EWG0_ENTCL	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3268	tr	A0A0F0XM62	A0A0F0XM62_9ENTR	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3269	sp	Q5PK92	RPOC_SALPA	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3270	sp	A9MHE9	RPOC_SALAR	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3271	tr	A0A232XM43	A0A232XM43_SALMU	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3272	tr	B5RFK0	B5RFK0_SALG2	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3273	sp	POA2R5	RPOC_SALTI	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3274	sp	Q57H68	RPOC_SALCH	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3275	sp	POA2R4	RPOC_SALTY	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3276	sp	A6TGP1	RPOC_KLEP7	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3277	tr	A0A0J2K6S7	A0A0J2K6S7_9ENTR	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3278	tr	A0A0G3RZQ0	A0A0G3RZQ0_KLEOX	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3279	tr	A0A212HDS5	A0A212HDS5_9ENTR	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3280	tr	A0A1R0FP41	A0A1R0FP41_CITBR	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3281	tr	A0A078LHA5	A0A078LHA5_CITKO	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-K			518
3282	sp	A8AKT8	RPOC_CITK8	DFDGD	EMNVHVP	LSSFAQAEARIQMLSAHNLLSPASGEPLAKPSRDI	ILGLYITQVR-I			518
3283				:	:	:	:	:	:	:
3284				:	:	:	:	:	:	:

3285	tr	AOA1U8DYN0 A0A1U8DYN0_ALLSI	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	911
3286	tr	AOA1L8H4P4 A0A1L8H4P4_XENLA	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3287	tr	H9GLG5 H9GLG5_ANOCA	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	902
3288	tr	H2R1J6 H2R1J6_PANTR	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3289	tr	P08775 RPB1_MOUSE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3290	tr	G1MCZ1 G1MCZ1_AILME	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3291	tr	O08847 O08847_MOUSE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3292	tr	S7PWZ6 S7PWZ6_MYOB	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3293	tr	D4A5A6 D4A5A6_RAT	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3294	tr	AOA1S3EWL2 AOA1S3EWL2_DIPOR	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3295	sp	P11414 RPB1_CRIGR	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3296	tr	O35559 O35559_CRIGR	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3297	tr	AOA2I3M9H2 AOA2I3M9H2_PAPAN	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	902
3298	tr	F7HB40 F7HB40_MACMU	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3299	tr	AOA2K6RYW9 AOA2K6RYW9_SAIBB	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	903
3300	tr	W5N8Z6 W5N8Z6_LEP	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	902
3301	tr	I3JRW6 I3JRW6_ORENI	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3302	tr	AOA0R4IMS9 AOA0R4IMS9_DANRE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	900
3303	tr	AOA1A7X327 AOA1A7X327_9TELE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3304	tr	AOA1A8UKD7 AOA1A8UKD7_NOTFU	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3305	tr	AOA1A8ER05 AOA1A8ER05_9TELE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3306	tr	AOA1A8DQ60 AOA1A8DQ60_9TELE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3307	tr	AOA1A8NSR8 AOA1A8NSR8_9TELE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3308	tr	AOA1W4YLM7 AOA1W4YLM7_9TELE	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3309	tr	AOA1M8A6L7 AOA1M8A6L7_MALS4	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	901
3310	tr	AOA1M8A6L7 AOA1M8A6L7_MALS4	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	894
3311	sp	P04050 RPB1_YEAST	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	880
3312	tr	AOA1B2J8C6 AOA1B2J8C6_PICPA	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	881
3313	tr	F2QW17 F2QW17_KOMP	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	881
3314	tr	A3GID7 A3GID7_PICST	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	880
3315	tr	AOA1D8PUA6 AOA1D8PUA6_CANAL	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	880
3316	tr	G8BEH9 G8BEH9_CANPC	REGLIDTAVKTAETGYI	ORRLIKSMESVMVKYDATVRNSINQVVQLRYGEDGLAGESVEF	880
3317	tr	AEG34223.1	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3318	tr	ASR51305.1	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	827	
3319	tr	OXR47930.1	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3320	sp	A7MQQ8 RPOC_CROS8	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3321	sp	Q32AG0 RPOC_SHIDS	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3322	sp	Q0SY12 RPOC_SHIF8	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3323	sp	B2TWH4 RPOC_SHIB3	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3324	sp	P0A8T7 RPOC_ECOLI	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3325	sp	Q3YU26 RPOC_SHISS	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3326	sp	B1XBZ0 RPOC_ECODH	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3327	sp	A8A787 RPOC_ECOHS	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3328	tr	AOA237JUP3 AOA237JUP3_SHISO	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3329	tr	AOA0F1RBF2 AOA0F1RBF2_ENTAS	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3330	tr	AOA1B3EWG0 AOA1B3EWG0_ENTCL	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3331	tr	AOA0F0XM62 AOA0F0XM62_9ENTR	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3332	tr	Q5PK92 RPOC_SALPA	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3333	sp	A9MHE9 RPOC_SALAR	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3334	tr	AOA232XM43 AOA232XM43_SALMU	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3335	tr	B5RFK0 B5RFK0_SALG2	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3336	sp	POA2R5 RPOC_SALTI	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3337	sp	Q57H68 RPOC_SALCH	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3338	sp	POA2R4 RPOC_SALTY	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3339	sp	A6TGP1 RPOC_KLEP7	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3340	tr	AOA0J2K6S7 AOA0J2K6S7_9ENTR	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3341	tr	AOA0G3RZQ0 AOA0G3RZQ0_KLEOX	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3342	tr	AOA212HDS5 AOA212HDS5_9ENTR	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3343	tr	AOA1R0FP41 AOA1R0FP41_CITBR	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3344	tr	AOA078LHA5 AOA078LHA5_CITKO	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3345	sp	ABAKT8 RPOC_CITK8	RKGLADTALKTANSGLYLRRLVDVADLVVTEDDCGTHEGIMMTPVIEGGDVK	832	
3346			***		
3347					
3348	tr	AOA1U8DYN0 A0A1U8DYN0_ALLSI	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1090	
3349	tr	AOA1L8H4P4 A0A1L8H4P4_XENLA	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1080	
3350	tr	H9GLG5 H9GLG5_ANOCA	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1081	
3351	tr	H2R1J6 H2R1J6_PANTR	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3352	sp	P08775 RPB1_MOUSE	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3353	tr	G1MCZ1 G1MCZ1_AILME	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3354	tr	O08847 O08847_MOUSE	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3355	tr	S7PWZ6 S7PWZ6_MYOB	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3356	tr	D4A5A6 D4A5A6_RAT	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3357	tr	AOA1S3EWL2 AOA1S3EWL2_DIPOR	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3358	sp	P11414 RPB1_CRIGR	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3359	tr	O35559 O35559_CRIGR	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3360	tr	AOA2I3M9H2 AOA2I3M9H2_PAPAN	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1081	
3361	tr	F7HB40 F7HB40_MACMU	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3362	tr	AOA2K6RYW9 AOA2K6RYW9_SAIBB	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1082	
3363	tr	W5N8Z6 W5N8Z6_LEP	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1081	
3364	tr	I3JRW6 I3JRW6_ORENI	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1080	
3365	tr	AOA0R4IMS9 AOA0R4IMS9_DANRE	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1079	
3366	tr	AOA1A7X327 AOA1A7X327_9TELE	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1080	
3367	tr	AOA1A8UKD7 AOA1A8UKD7_NOTFU	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1080	
3368	tr	AOA1A8ER05 AOA1A8ER05_9TELE	VNGDDPLSRQAQENATLLFNHLRSTLCSRRMAEEFRLSQAQAFDILLGIEISKFNQIAH	1080	

3369	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	VNGDDPLSRQAQENATLLFNHLRSTLCSRMTTEEFRLSMEAFDWLLGEIETKFNQSVIH	1080
3370	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	VNGDDPLSRQAQENATLLFNHLRSTLCSRMTTEEFRLSMEAFDWLLGEIETKFNQSVIH	1080
3371	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	VNGDDPLSRQAQENATLLFNHLRSTLCSRMTTEEFRLSMEAFDWLLGEIETKFNQSVIH	1080
3372	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	IRGNDPISRSMQENATLLFKLHMRSLCTKQVIEVHHLREAWEWILGIEGQFARSVAQ	1071
3373	sp	P04050	RPB1_YEAST	LRGKNEIQNAQRDAVTLFCCLLRSLRATRRVLQEVRLTKQAFDWLWSLNLIAEQFLRSVVH	1059
3374	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	LRGENELIKEAQQNATSLFQCLVRARLATRRILEEFRLNRDAFEWVWLGTEAQQFQSLVH	1061
3375	tr	F2QW17	F2QW17_KOMP	LRGENELIKEAQQNATSLFQCLVRARLATRRILEEFRLNRDAFEWVWLGTEAQQFQSLVH	1061
3376	tr	A3GID7	A3GID7_PICST	VRGDTLKVKEAQNATLLFQCLVRSRLASRRVIEEFKLNRSSEFEWVWGEIETQFQKSVIH	1060
3377	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	VRGDTPLVKEAQNATLLFQCLLRSLRSLARRVIEEFKLNRSSEFEWVWGEIETQFQKSVIH	1060
3378	tr	G8BEH9	G8BEH9_CANPC	VRGKTKLAKEAQNATLLFQCLVRSRLAARRVIEEFKLNRSSEFEWVWGEIETQFQKSVIH	1060
3379	sp	AEG34223.1		-GEIQ-----EVPVRSPLTCCRYGVCCKQ-----YGDLS-----MARPVN	1216
3380	sp	ASR51305.1		-LGLQ-----AARISPLICEATMGVCKKQ-----YGRDLA-----RGTPVN	905
3381	sp	OXR47930.1		-LGVD-----EVKIRTLTCEITRRGLCAHQ-----YGRDLG-----RGLSVN	910
3382	sp	A7MQQ8	RPOC_CROS8	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3383	sp	Q32AG0	RPOC_SHIDS	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3384	sp	Q0SY12	RPOC_SHIF8	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3385	sp	B2TWH4	RPOC_SHIB3	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3386	sp	FOA8T7	RPOC_ECOLI	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3387	sp	Q3YU26	RPOC_SHISS	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3388	sp	B1XBZ0	RPOC_ECODH	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3389	sp	A8A787	RPOC_ECOHS	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3390	tr	A0A237JUP3	A0A237JUP3_SHISO	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3391	tr	A0A0F1RBF2	A0A0F1RBF2_ENTAS	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3392	tr	A0A1B3EWG0	A0A1B3EWG0_ENTCL	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3393	tr	A0A0F0XM62	A0A0F0XM62_9ENTR	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3394	sp	Q5PK92	RPOC_SALPA	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3395	sp	A9MHE9	RPOC_SALAR	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3396	tr	A0A232XM43	A0A232XM43_SALMU	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3397	tr	B5RFK0	B5RFK0_SALG2	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3398	sp	POA2R5	RPOC_SALTI	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3399	sp	Q57H68	RPOC_SALCH	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3400	sp	POA2R4	RPOC_SALTY	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3401	sp	A6TGP1	RPOC_KLEP7	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3402	tr	A0A0J2K6S7	A0A0J2K6S7_9ENTR	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3403	tr	A0A0G3RZQ0	A0A0G3RZQ0_KLEOX	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3404	tr	A0A212HDS5	A0A212HDS5_9ENTR	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3405	tr	A0A1R0FP41	A0A1R0FP41_CITBR	-NSVD-----SVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3406	tr	A0A078LHA5	A0A078LHA5_CITKO	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910
3407	sp	ABAKT8	RPOC_CITK8	-NSVD-----AVKVRSVVSCIDDFGVCAHQ-----YGRDLA-----RHHIIN	910x16
3409					
3410	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1142
3411	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1132
3412	tr	H9GLG5	H9GLG5_ANOCA	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1133
3413	tr	H2R1J6	H2R1J6_PANTR	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3414	sp	P08775	RPB1_MOUSE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3415	tr	G1MCZ1	G1MCZ1_ALLME	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3416	tr	O08847	O08847_MOUSE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3417	tr	S7PWZ6	S7PWZ6_MYOBR	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3418	tr	D4A5A6	D4A5A6_RAT	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3419	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3420	sp	P11414	RPB1_CRIGR	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3421	tr	O35559	O35559_CRIGR	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3422	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1133
3423	tr	F7HB40	F7HB40_MACMU	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3424	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKKP	1134
3425	tr	W5N8Z6	W5N8Z6_LEPOC	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1133
3426	tr	I3JRW6	I3JRW6_ORENI	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1132
3427	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1131
3428	tr	A0A1A7X327	A0A1A7X327_9TELE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1132
3429	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1132
3430	tr	A0A1A8ER05	A0A1A8ER05_9TELE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1132
3431	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1132
3432	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1132
3433	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	PGEMVGALAAQSLGEPATQMTLNTFHYAGVSAKNVTLGVPR-----LKEILNISKRP	1132
3434	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	PGEMCGTLAAQSLGEPATQMTLNTFHYAGVSSKNVTLGVPR-----LKEILNCAENI	1123
3435	sp	P04050	RPB1_YEAST	PGEMVGVLAAQSLGEPATQMTLNTFHYAGVASKKVTSGVPR-----LKEILNVAKNM	1111
3436	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	PGEMVGVLAAQSLGEPATQMTLNTFHYAGVSSKNVTLGVPR-----LKEILNVAKNI	1113
3437	tr	F2QW17	F2QW17_KOMP	PGEMVGVLAAQSLGEPATQMTLNTFHYAGVSSKNVTLGVPR-----LKEILNVAKNI	1113
3438	tr	A3GID7	A3GID7_PICST	PGEMVGVLAAQSLGEPATQMTLNTFHYAGVSSKNVTLGVPR-----LKEILNVAKNI	1112
3439	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	PGEMVGVLAAQSLGEPATQMTLNTFHYAGVSSKNVTLGVPR-----LKEILNVAKNI	1112
3440	tr	G8BEH9	G8BEH9_CANPC	PGEMVGVLAAQSLGEPATQMTLNTFHYAGVSSKNVTLGVPR-----LKEILNVAKNI	1112
3441	sp	AEG34223.1		IGEAIVGLAAQSLGEPGTQMTLNTFHYGGVAG-----LKEILNVAKNI	1248
3442	sp	ASR51305.1		IGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAQLNE-TSNLEAVADGTLQYRDI-PTIVNK	963
3443	sp	OXR47930.1		IGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAMASSVETKAAAGTVSFGVSMRVVINA	970
3444	sp	A7MQQ8	RPOC_CROS8	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3445	sp	Q32AG0	RPOC_SHIDS	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3446	sp	Q0SY12	RPOC_SHIF8	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3447	sp	B2TWH4	RPOC_SHIB3	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3448	sp	FOA8T7	RPOC_ECOLI	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3449	sp	Q3YU26	RPOC_SHISS	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3450	sp	B1XBZ0	RPOC_ECODH	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3451	sp	A8A787	RPOC_ECOHS	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3452	tr	A0A237JUP3	A0A237JUP3_SHISO	KGEAIVGLAAQSLGEPGTQMTLNTFHYGGAAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969

3453	tr	A0A0F1RBF2	A0A0F1RBF2_ENTAS	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNA-KSVVNS	969
3454	tr	A0A1B3EWG0	A0A1B3EWG0_ENTCL	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNA-KSVVNS	969
3455	tr	A0A0F0XM62	A0A0F0XM62_9ENTR	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNA-KSVVNS	969
3456	sp	Q5PK92	RPOC_SALPA	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3457	sp	A9MHE9	RPOC_SALAR	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3458	tr	A0A232XM43	A0A232XM43_SALMU	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3459	tr	B5RFK0	B5RFK0_SALG2	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3460	sp	P0A2R5	RPOC_SALTI	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3461	sp	Q57H68	RPOC_SALCH	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3462	sp	P0A2R4	RPOC_SALTY	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNV-KSVVNS	969
3463	sp	A6TGP1	RPOC_KLEP7	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIKLSNA-KSVVNS	969
3464	tr	A0A0J2K6S7	A0A0J2K6S7_9ENTR	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIRLSNA-KSVVNS	969
3465	tr	A0A0G3RZQ0	A0A0G3RZQ0_KLEOX	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIRLSNA-KSVVNS	969
3466	tr	A0A212HDS5	A0A212HDS5_9ENTR	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIRLSNA-KSVVNS	969
3467	tr	A0A1R0FP41	A0A1R0FP41_CITBR	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIRLSNA-KSVVNS	969
3468	tr	A0A078LHA5	A0A078LHA5_CITKO	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIRLSNA-KSVVNS	969
3469	sp	A8AKT8	RPOC_CITK8	KGEAIGVIAAQSIGEPGTQLTMRPFHIGGAASRAAAESSIQVKNKGSIRLSNA-KSVVNS	969
3470				** * :****:*** ** : * * * * *	
3471					
3472	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1589
3473	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	--WSPS---IGSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1579
3474	tr	H9GLG5	H9GLG5_ANOCA	--WSPS---VSGMTPGAAGFSPSAASDA-\$GLSPGYSPAWSPTPGSPGSPGSSPY---	1580
3475	tr	H2R1J6	H2R1J6_PANTR	GLWSPALA-----LTYA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1571
3476	sp	P08775	RPB1_MOUSE	--WSPSV---GSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1581
3477	tr	G1MCZ1	G1MCZ1_AILME	--WSPSVDITGSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1584
3478	tr	O08847	O08847_MOUSE	--WSPSV---GSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1581
3479	tr	S7PWZ6	S7PWZ6_MYOBR	--WSPSV---GSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1581
3480	tr	D4A5A6	D4A5A6_RAT	--WSPSV---GSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1581
3481	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	--WSPSV---GSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1581
3482	sp	P11414	RPB1_CRIGR	--WSPSV---GSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1581
3483	tr	O35559	O35559_CRIGR	--WSPSV---GSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1581
3484	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	-----GMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1542
3485	tr	F7HB40	F7HB40_MACMU	-----GMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1543
3486	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	-----GMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSSPY---	1543
3487	tr	W5N8Z6	W5N8Z6_LEPOC	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1580
3488	tr	I3JRW6	I3JRW6_ORENI	--WSPS---VSGMTPGGAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1579
3489	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1578
3490	tr	A0A1A7X327	A0A1A7X327_9TELE	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1579
3491	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1579
3492	tr	A0A1A8ER05	A0A1A8ER05_9TELE	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1579
3493	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1579
3494	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1579
3495	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	--WSPS---VSGMTPGAAGFSPSAASDA-\$GFSPGYSPAWSPTPGSPGSPGSPASPY---	1579
3496	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	---DDF---R--VQQQAMFSPPLVQVGGDE---GGY---SDYLSAGQSP---	1542
3497	sp	P04050	RPB1_YEAST	---N-A---DLDVDLMLFSPPLVDSGNSNDAMAGGF---TAYGGADYGEA-TSPF---	1523
3498	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	---DEF---NHDDVADVMFSPMAETGSGDDRSGL---TEYAGIQSPYQP---	1527
3499	tr	F2QW17	F2QW17_KOMPC	---DEF---NHDDVADVMFSPMAETGSGDDRSGL---TEYAGIQSPYQP---	1527
3500	tr	A3GID7	A3GID7_PICST	---DD---KIQFEEGAGFSPHIAHQVQ-DVSGGL---TSYGGQPTSPSATSFPFSYG	1529
3501	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	---DE---NIDIDAGAGFSPHIAQMNENIGGL---TSYGGQPTSPAATSFPFSYG	1527
3502	tr	G8BEH9	G8BEH9_CANPC	---DD---RIQVDESAGFSPVHQAPTAEGMNGGL---TSYGGQPTSPSATSFPFSYS	1526
3503		ABG34223.1		GKQA-----	1524
3504		ASR51305.1		-----	1403
3505		OKR47930.1		-----	1416
3506	sp	A7MQQ8	RPOC_CROS8	GSDNE-----	1407
3507	sp	Q32AG0	RPOC_SHIDS	GSDNE-----	1407
3508	sp	Q0SY12	RPOC_SHIF8	GSDNE-----	1407
3509	sp	B2TWH4	RPOC_SHIB3	GSDNE-----	1407
3510	sp	P0A8T7	RPOC_ECOLI	GSDNE-----	1407
3511	sp	Q3YUZ6	RPOC_SHISS	GSDNE-----	1407
3512	sp	B1XBZ0	RPOC_ECODH	GSDNE-----	1407
3513	sp	A8A787	RPOC_ECOHS	GSDNE-----	1407
3514	tr	A0A237JUP3	A0A237JUP3_SHISO	GSDNE-----	1407
3515	tr	A0A0F1RBF2	A0A0F1RBF2_ENTAS	GSDNE-----	1407
3516	tr	A0A1B3EWG0	A0A1B3EWG0_ENTCL	GSDNE-----	1407
3517	tr	A0A0F0XM62	A0A0F0XM62_9ENTR	GSDNE-----	1407
3518	sp	Q5PK92	RPOC_SALPA	GSDNE-----	1407
3519	sp	A9MHE9	RPOC_SALAR	GSDNE-----	1407
3520	tr	A0A232XM43	A0A232XM43_SALMU	GSDNE-----	1407
3521	tr	B5RFK0	B5RFK0_SALG2	GSDNE-----	1407
3522	sp	P0A2R5	RPOC_SALTI	GSDNE-----	1407
3523	sp	Q57H68	RPOC_SALCH	GSDNE-----	1407
3524	sp	P0A2R4	RPOC_SALTY	GSDNE-----	1407
3525	sp	A6TGP1	RPOC_KLEP7	GSDNE-----	1407
3526	tr	A0A0J2K6S7	A0A0J2K6S7_9ENTR	GSDNE-----	1407
3527	tr	A0A0G3RZQ0	A0A0G3RZQ0_KLEOX	GSDNE-----	1407
3528	tr	A0A212HDS5	A0A212HDS5_9ENTR	GSDNE-----	1407
3529	tr	A0A1R0FP41	A0A1R0FP41_CITBR	GSDNE-----	1407
3530	tr	A0A078LHA5	A0A078LHA5_CITKO	GSDNE-----	1407
3531	sp	A8AKT8	RPOC_CITK8	GSDNE-----	1407
3532					

3533	tr	A0A1U8DYN0	A0A1U8DYN0_ALLSI	-----	1926
3534	tr	A0A1L8H4P4	A0A1L8H4P4_XENLA	PDPSDEDN	1968
3535	tr	H9GLG5	H9GLG5_ANOCA	PDPSDEEN	1969
3536	tr	H2R1J6	H2R1J6_PANTR	PDPSDEEN	1960
3537	sp	P08775	RBP1_MOUSE	PDPSDEEN	1970
3538	tr	G1MCZ1	G1MCZ1_AILME	PDPSDEEN	1973
3539	tr	O08847	O08847_MOUSE	---DREN	1966
3540	tr	S7PWZ6	S7PWZ6_MYOBR	PDPSDEEN	1970
3541	tr	D4A5A6	D4A5A6_RAT	PEPSDEEN	1970
3542	tr	A0A1S3EWL2	A0A1S3EWL2_DIPOR	PDPSDEEN	1970
3543	sp	P11414	RBP1_CRIGR	PDPSDEEN	1970
3544	tr	O35559	O35559_CRIGR	PDPSDEEN	1970
3545	tr	A0A2I3M9H2	A0A2I3M9H2_PAPAN	PDPSDEEN	1931
3546	tr	F7HB40	F7HB40_MACMU	PDPSDEEN	1932
3547	tr	A0A2K6RYW9	A0A2K6RYW9_SAIBB	PDPSDEEN	1932
3548	tr	W5N8Z6	W5N8Z6_LEPOC	PDPSDEEN	1959
3549	tr	I3JRW6	I3JRW6_ORENI	PDPSDEEN	1966
3550	tr	A0A0R4IMS9	A0A0R4IMS9_DANRE	PDPSDEEN	1965
3551	tr	A0A1A7X327	A0A1A7X327_9TELE	PDPSDEEN	1969
3552	tr	A0A1A8UKD7	A0A1A8UKD7_NOTFU	PDPSDEEN	1969
3553	tr	A0A1A8ER05	A0A1A8ER05_9TELE	PDPSDEEN	1969
3554	tr	A0A1A8DQ60	A0A1A8DQ60_9TELE	PDPSDEEN	1962
3555	tr	A0A1A8NSR8	A0A1A8NSR8_9TELE	PDPSDEEN	1962
3556	tr	A0A1W4YLM7	A0A1W4YLM7_9TELE	PDPSDDDN	1968
3557	tr	A0A1M8A6L7	A0A1M8A6L7_MALS4	-----	1803
3558	sp	P04050	RBP1_YEAST	-----	1733
3559	tr	A0A1B2J8C6	A0A1B2J8C6_PICPA	-----	1743
3560	tr	F2QW17	F2QW17_KOMPC	-----	1743
3561	tr	A3GID7	A3GID7_PICST	-----	1739
3562	tr	A0A1D8PUA6	A0A1D8PUA6_CANAL	-----	1728
3563	tr	G8BEH9	G8BEH9_CANPC	-----	1746

Fig. 6 Mix and Match analysis of the elongation subunits β' of *E. coli* and Rpb1 of yeast RNAP-II.

For legends refer to Figs. 3 and 4

8. Active site analysis of the *S. cerevisiae* MSU RNAP- II

8.1. Catalytic region

It has been found that almost all DNA polymerases and SSU RNAPs use an invariant K for catalysis, i.e., in the initial proton transfer reactions [34,29]. However, in all MSU RNAPs analyzed, no K was found at the expected distance from the template binding YG/FG pair but an equivalent invariant R (Table 4). However, a detailed analysis has shown that all prokaryotic DNA polymerases II also use an invariant R in catalysis with similar distance conservations [34, 4] instead of a usual K; interestingly, an enzyme also possesses primase activity and along with associated 3'→5' exonuclease activity.

Table 4 Amino acids around the catalytic amino acid K/R and the YG/FG pair in DNA polymerases, SSU and MSU RNAPs.

Polymerase Type	Catalytic Region
Viral SSU RNA pol (T7)	-TR ⁴ VTKR ¹ SVMTLAY ⁸ GS-
Mitochondrial SSU RNA pol (Yeast)	-TR ⁴ KVVKQ ¹ TVMTNVY ⁸ GV--
Chloroplast SSU pol (ARATH)	-DR ⁴ KLVKQ ¹ TVMTSVY ⁸ GV-
<i>E. coli</i> DNA pol I (SSU)	-QR ⁴ RSKA ¹ INFGLIY ⁸ GM-
Initiation subunits of MSU RNAPs	
<i>E. coli</i> MSU RNAP β subunit	⁵³⁹ TR ⁸ ERAGFEV RD ¹ VHPTHY ⁷ G ⁸ RV ⁵⁵⁸ -
<i>S. cerevisiae</i> MSU RNAP II Rpb2 subunit	⁸⁵¹ FR ⁵ SLFFRS ¹ YMDQEKKY ⁹ GMSI ⁸⁷⁰ -
Human MSU RNAP II Rpb2 subunit	⁸⁰⁶ FR ⁵ SVFYRS ¹ YKEQESK ⁹ GFDQ ⁸²⁵ -
Elongation subunits of MSU RNAPs	
<i>E. coli</i> MSU RNAP β' subunit	⁸³³ NSV ⁶ DAVKVRS ¹ VVSC ⁵ DTDFGVC ¹² AHC ¹⁵ Y ¹⁶ G ¹⁷ RDL ⁸⁶¹ -
<i>S. cerevisiae</i> MSU RNAP II Rpb1 subunit	⁵⁵ DPR ⁶ LGSIDRN ¹ LKC ⁴ QTC ⁷ QEGMNEC ¹⁴ PGHF ¹⁸ G ¹⁹ HI ⁸⁴ -
Huan MSU RNAP II Rpb1 subunit	⁵⁹ DPR ⁶ QGVIERT ¹ GRC ⁴ QTC ⁷ AGNMTEC ¹⁴ PGHF ¹⁸ G ¹⁹ HI ⁸⁸ -

NB: MSU RNA polymerases which use R in the catalytic site is shown in bold.

The invariant R, at -6/-7 is not found in β' subunits of eubacteria

3579 The eukaryotic elongation subunits an FG instead of a YG

3580 Table 4 shows the invariant template binding YG pair with its catalytic R in the initiation
 3581 (β and Rpb2) and in the elongation subunits (β' and Rpb1) of eubacterial and eukaryotic MSU RNAPs,
 3582 respectively (Figs. 3-7). The catalytic R in both in the initiation and elongation subunits of the MSU
 3583 RNAPs will serve as a proton abstractor for initiating the catalysis as explained by Palanivelu [4]. Table 4
 3584 shows the invariant YG/FG pair and its catalytic R in the initiation and elongation subunits of eubacterial
 3585 and eukaryotic MSU RNAPs.

3586 It is interesting to note that the YG pair appears to be specific for polymerases using DNA as the
 3587 template (including the prokaryotic and eukaryotic MSU RNAPs, data not shown) as it is not reported in
 3588 RNA dependent RNAPs where they use RNA as the template [35]. In fact, Kotsyuk et al [37] have shown
 3589 that the DNA dependent T7 RNAP requires both the strands for activity and uses two YG pairs and there
 3590 was no activity when single-stranded DNA was used as the substrate. However, the eukaryotic initiation
 3591 subunit Rpb2 uses a functional equivalent FG and the initiation subunit from higher eukaryotes including
 3592 human uses a completely different pair, KG but followed by an F. The elongation subunits in all
 3593 eukaryotes use a functionally equivalent FG pair. However, it is interesting to note the catalytic amino
 3594 acid is R and is completely conserved in all MSU RNAPs II. Three invariant Cs between catalytic R and
 3595 YG/FG pair in the elongation subunits (β' and Rpb1) are highly conserved in both pro- and eukaryotes but
 3596 the three Cs are placed at a different distances as 5+12+15 and 4+7+14 with an 11 amino acids span in
 3597 both the cases (Table 4). These completely conserved Cs are implicated in Zn binding in the bacterial
 3598 elongation subunits by crystallographic analysis [36].

3599 8.2 Mechanism of NTP and dNTP discrimination in eukaryotic MSU RNA polymerases

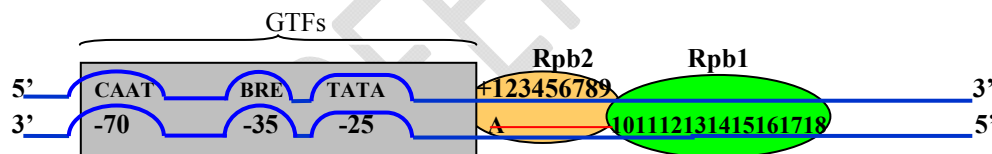
3600 NTP and dNTP discrimination in nucleic acid polymerases is usually achieved by base pairing, base
 3601 stacking, hydrogen bonding and also by specific interaction(s) of completely conserved amino acid(s) in
 3602 the nucleotide interacting domains. A completely conserved R at -5 in the Rpb2 and -6 in Rpb1 is known
 3603 to participate in the nucleotide discrimination and select only NTPs in the polymerization site (Table 4).
 3604 Similar absolute conservation is seen in SSU RNAPs from viruses, mitochondria and chloroplasts and
 3605 DNA polymerases too. A conspicuous absence of an invariant R at the expected distance at -6 in the
 3606 elongation subunit β' of eubacterial MSU RNAPs (Table 4) raises the question of how they discriminate
 3607 the NTPs from dNTPs. Modelling of the substrate NTP bound to the *T. thermophilus* RNAP active site
 3608 suggests that N⁴⁵⁸ (numbering from *E. coli* RNA polymerase) within a highly conserved sequence motif
 3609 ⁴⁵⁸NADFDGD⁴⁶⁴ that includes the catalytic Asp triad (D460, D462, D464) could mediate specific
 3610 recognition of the O2 ribose atom [39, 36]. Functional *in vitro* analysis demonstrated that the substitutions
 3611 of the corresponding β' N⁴⁵⁸ residue led to the loss of discrimination between NTP and dNTP substrates
 3612 as well as to defects in RNA chain extension [39]. It is interesting to note that substitution of the
 3613 corresponding amino acid in yeast Rpb1 (N479Y) is lethal in the same sequence motif ⁴⁷⁸YNADFDGD in
 3614 eukaryotes [40] The absolutely conserved N in both the cases could discriminate the NTP from dNTP by
 3615 recognizing the 2'-OH of the ribose and suggested that the crucial N could interact with both the 2'-OH as
 3616 well 3'-OH [40].

3617 The highly conserved S/T adjacent (N in *S. cerevisiae*) to the catalytic R in the initiation and
 3618 elongation subunits likely recognizes the 2'-OH in the NTPs and possibly makes a hydrogen bond and
 3619 discriminate dNTPs, which lack a 2'-OH (the yeast elongation subunit, Rpb1 could also use the invariant
 3620 N⁴⁷⁹ to make the necessary H bond with the 2'-OH as well as 3'-OH as shown in elongation subunit β' of
 3621 eubacterial MSU RNAPs [4]. Kaplan et al [41] have shown by SDM experiments that the completely
 3622 conserved His1085 both in prokaryotes (β') and eukaryotes Rpb1 might involve in NTP selection and
 3623 substitution of Ala or Phe resulted in inviability. This H1085 in the trigger loop (amino acids from 1060 to
 3624 1101) was shown to make a contact with the β phosphate either through H-bond or salt bridge and the
 3625 L1081 was located at the 3' end of the RNA [42]. Mix and match analysis also shows and that particular
 3626 His is completely conserved in both prokaryotes and eukaryotes in highly conserved stretch (Fig. 6).
 3627 Another amino acid Q1078 in the same block is also shown to be essential and replacing Q1078 in Rpb1
 3628 to either N or S is lethal in yeast [41], consistent with a key role of these residues in NTP/dNTP
 3629 discrimination.

3630 9. The initiation and elongation subunits work in tandem during transcription cycles in 3631 eukaryotic MSU RNAP II

3632 A close similarity is observed between the prokaryotic and eukaryotic transcription cycles. For
 3633 example, it has been shown in prokaryotes (*E. coli*) that the initiation of transcription by the β subunit is
 3634 not smooth and it makes many aborted transcripts of sizes 2-7 nts before the elongation step is taken
 3635 over by the β' subunit [43]. It was found that RNA/DNA hybrids of less than 8-bp display markedly less
 3636 stability than those that are 8 bp or longer. In fact, the *E. coli* MSU RNAPs use these short RNA/DNA
 3637 hybrids of 8 bp or longer for the ternary elongation complex (TEC) formation and further processivity [44].
 3638 Furthermore, Zaychikov et al [45] have shown that an ~ 17 bp region of the DNA called the 'transcription
 3639 bubble' was melted to expose the template strand for transcription in *E. coli*. MSA analysis agreed with
 3640 these findings with the 7 amino acid gap between the catalytic R and the template binding YG pair in the
 3641 initiation subunit β and 17 amino acid gap ('transcription bubble') between the catalytic R and the
 3642 template binding YG pair in the elongation subunit β' (Table 5) [4].

3643 A similar observation of abortive initiations in eukaryotic Rpb2 was also reported by Pal and Luse
 3644 [32]. In addition to, as in prokaryotes, in eukaryotes also a 9 bp DNA-RNA stable hybrid is formed which
 3645 extends from the active centre at nearly right angles to the entering DNA during the elongation cycle [16,
 3646 33]. Fiedler and Timmers [33] results further support the MSA findings where the transition from abortive
 3647 to productive elongation cycle occurred once the RNAP register +10 nts (Table 5). These observations
 3648 are further supported by Luse who have shown that the promoter clearance was complete with ~ 10 nts
 3649 [46]. Pal et al on analyzing the 'transcription bubble' have found that regardless of promoter spacing, the
 3650 upstream edge of the 'transcription bubble' formed 20 bp from TATA which is in close agreement with
 3651 MSA analysis data [47]. A similar finding was reported by Giardina and Lis [48]. Furthermore, Holstege et
 3652 al analyzed the transcription initiation by the yeast RNAP II in a highly efficient *in vitro* transcription
 3653 system composed of essentially homogeneous protein preparations and found that the downstream part
 3654 of the 'transcription bubble' expands in a continuous motion, but the initially opened region on the non-
 3655 template strand reclosed abruptly when transcription reached 11, which was accompanied by a switch
 3656 from abortive initiations to productive mRNA synthesis (elongation) [49]. Similar findings were also
 3657 reported by Barnes et al by analyzing the crystal structure of the transcribing RNAP II complex. The
 3658 transcription bubble was unwound ~ 18 -25 bases and transcripts of 10 nts or more resulted in promoter
 3659 escape with stabilization of a mature bubble [50].



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 3665 **Fig. 7** A consensus model showing pre-initiation, initiation and elongation events by GTFs, Rpb2 and Rpb1
 3666 subunits of MSU RNAP- II of *S. cerevisiae*

3667 NB: Thick lines indicate DNA strands and the thin line indicates the mRNA transcript.

3668 A is shown as the first nucleotide at the +1 TSS

3669 GTFs, General Transcription Factors (TFII-D, -A, -B, -F, -E, -H)

3670 BRE, TFII-B Recognition Element (G/C-G/C-G/A-C-G-C-C)

3671 In contrast to prokaryotes, in eukaryotes, it has been found that an A is inserted (and not a usual G, as
 3672 in prokaryotes) in the initiation site as it invariably uses a G for capping enzyme which is also
 3673 associated with the RNAP II.

3674 MSA analysis agrees with these findings with the 9 amino acid gap between the catalytic R and the
 3675 template binding YG pair in the initiation subunit Rpb2 and 19 amino acid gap ('transcription bubble')
 3676 between the catalytic R and the template binding FG pair in the elongation subunit Rpb1. Furthermore,
 3677 Gnatt et al [16] found that the contacts to the downstream and upstream parts of the hybrid are made by
 3678 Rpb1 and Rpb2, respectively, which further supports the tandem arrangement of the initiation and
 3679 elongation subunits as proposed in this model.

3680 Figure 7 shows a consensus model of the yeast MSU RNAP II subunits, from promoter recognition
 3681 to initiation and elongation events during the transcription process. This has been confirmed that the
 3682 RNAP II undergoes abortive initiations until it reached a position beyond +9 (i.e., the distance between
 3683 the template binding FG pair and the catalytic R), at which stage the RNAP II was released from its
 3684 promoter contacts and an elongation complex (TEC) is formed and a 20 bp 'transcription bubble' formed
 3685 from the TATA box [47]. A modular structure is proposed for the initiation and elongation subunits'
 3686 function in eukaryotes also as suggested by in prokaryotes [51].

3687 **10. Metal binding sites**

3688 The metal binding sites *S. cerevisiae* MSU RNAP II is arrived at from the data obtained by the
 3689 MSA (this work) and X-ray crystallographic analyses [36] and SDM experiments [39] on eubacterial MSU
 3690 RNAPs. The eukaryotic Rpb2 and Rpb1 subunits show many possible metal binding sites (highlighted in
 3691 green). A Mg^{2+} ion binding site is found in the mRNA initiation subunit, Rpb2, from all eukaryotes.
 3692 Similarly, a Mg^{2+} and a Zn^{2+} binding motifs are also found in the elongation subunits, Rpb1, of all
 3693 eukaryotic RNAPs II. Crystallographic analysis of the *T. aquaticus* RNAP by Zhang et al have shown that
 3694 the Mg atom is chelated at an absolutely conserved **-NADFDGD-** motif in the β' elongation subunit and
 3695 surprisingly the same invariant motif is found by MSA in all eukaryotic elongation subunits as well [36].
 3696 Moreover, this is one of the regions that align in both the pro- and eukaryotes by mix and match analysis
 3697 (Fig. 6). Interestingly, substitution of these Ds by A (D→A) gave rise to a dominant lethal phenotype and
 3698 showed no detectable enzyme activity [45]. In addition to that, a Zn binding motif is also recognized in the
 3699 eubacterial and eukaryotic elongation subunits with three conserved Cs. However, their distance
 3700 arrangements between the three Cs are found to be different (Table 5). Another distinguishing feature
 3701 between the eukaryotic and eubacterial elongation subunits is the catalytic and Zn binding regions, i.e., in
 3702 *E. coli* β' subunit, the amino acids 888, 895 and 898 are shown to be involved in Zn binding (Table 5) but
 3703 in eukaryotic elongation subunits it is placed at the very beginning of the N-terminal and covers the amino
 3704 acids region 57-82 (Table 5). The Zn binding motif that is located in Rpb1 subunits may play a similar role
 3705 in the Zn mediated proof-reading function as proposed for DNA polymerases and MSU RNAPs from
 3706 eubacteria [4, 29].

3707 Table 5. Metal binding sites in prokaryotic and eukaryotic MSU RNAPs

Subunit (Organism)	Metal binding site	Method and Reference
β eubacteria (<i>E. coli</i>)	⁻⁶⁷¹ LEHDDA/ ⁻⁸⁰⁹ GYNFEDS* - (Mg^{2+})	MSA (This communication)
β' eubacteria (<i>E. coli</i>)	⁻⁴⁵⁸ YNADFDGDQM - (Mg^{2+}) & ⁻⁸⁸³ RS ¹ VVSC ⁵ DTDFGVC ¹² AHC ¹⁵ Y ¹⁶ GR ⁹⁰¹ -(Zn^{2+})*	X-ray crystallographic data [36]
Rpb2 Eukaryote (<i>S. cerevisiae</i>)	⁻⁸⁹³ LDDDG ⁸⁹⁷ - ⁻⁸³² GYNQED*S ⁸³⁸ -(Mg^{2+})	MSA (This communication)
Rpb1 Eukaryote (<i>S. cerevisiae</i>)	⁻⁴⁷⁸ YNAD*FD*GDEM ⁴⁸⁷ - (Mg^{2+}) & ⁻⁵⁶ PR ⁶ LGSIDRN ¹ LKC ⁴ QTC ⁷ QEGMNEC ¹⁴ PGHF ¹⁸ GH ⁸³ -(Zn^{2+})	MSA (This communication)

3708 *Possible metal binding sites arrived at by MSA and SDM*

3709 NB: The β' and Rpb1 elongation subunits of eubacteria and eukaryotes contain both the Mg^{2+} and Zn^{2+}
 3710 binding sites. In both the cases, the Zn^{2+} binding site is built in the catalytic region with the 3 invariant Cs
 3711 which coordinates the Zn atom.

3712 **11. Mechanism of Action of the MSU RNAP II of *S. cerevisiae***

3713 A minimal number of steps involved in the catalytic cycle of RNAPs consist of NTP selection,
 3714 Watson-Crick base pairing with the complementary nucleotide to the template, catalysis, pyrophosphate
 3715 release and translocation. As the catalytic regions are found to be similar in both the initiation and
 3716 elongation subunits, the polymerization mechanism could be also similar in both the initiation and
 3717 elongation reactions. Figs. 8 and 9 describe the reactions involved in the initiation and elongation cycles
 3718 during transcription in *S. cerevisiae*. (all participating amino acids are not shown in the figures).

3719 **11.1 Mechanism of initiation by Rpb2 subunit of the MSU RNAP II from *S. cerevisiae***

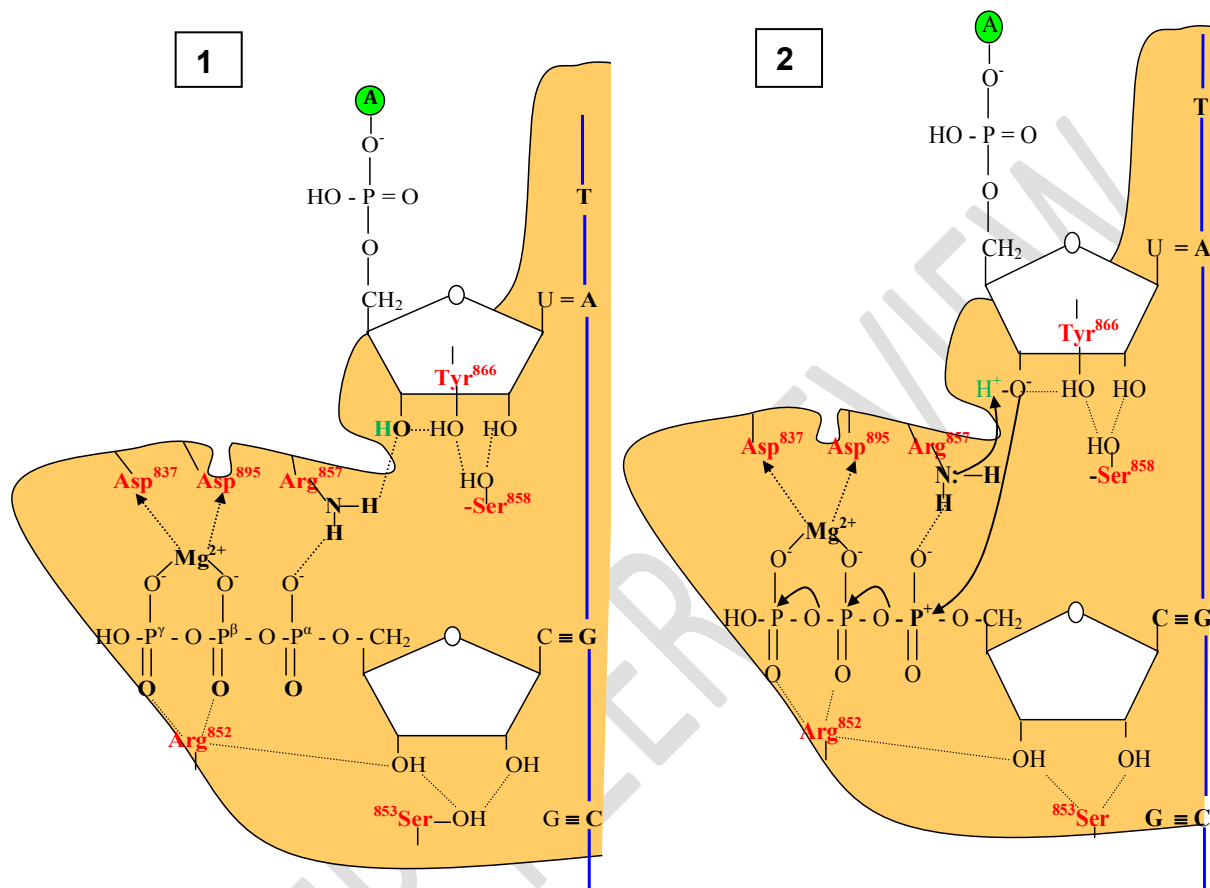
3720 **Step 1. Enzyme and the NTP at the Entry Site:** Template binding pair (Tyr-Gly) and nucleotide
 3721 discrimination by the invariant amino acids, Watson-Crick base pairing of the incoming NTP with the
 3722 template DNA. The catalytic site amino acid Arg is positioned for proton abstraction. ATP is the
 3723 initiating nucleotide.

3724 **Step 2. Proton abstraction and nucleophilic attack of the α -phosphate of NTP:** Electronic
 3725 transition at the active site for proton abstraction by the active site Arg and an electrophilic and
 3726 nucleophilic attack of the incoming NTPs on the 3'-OH.

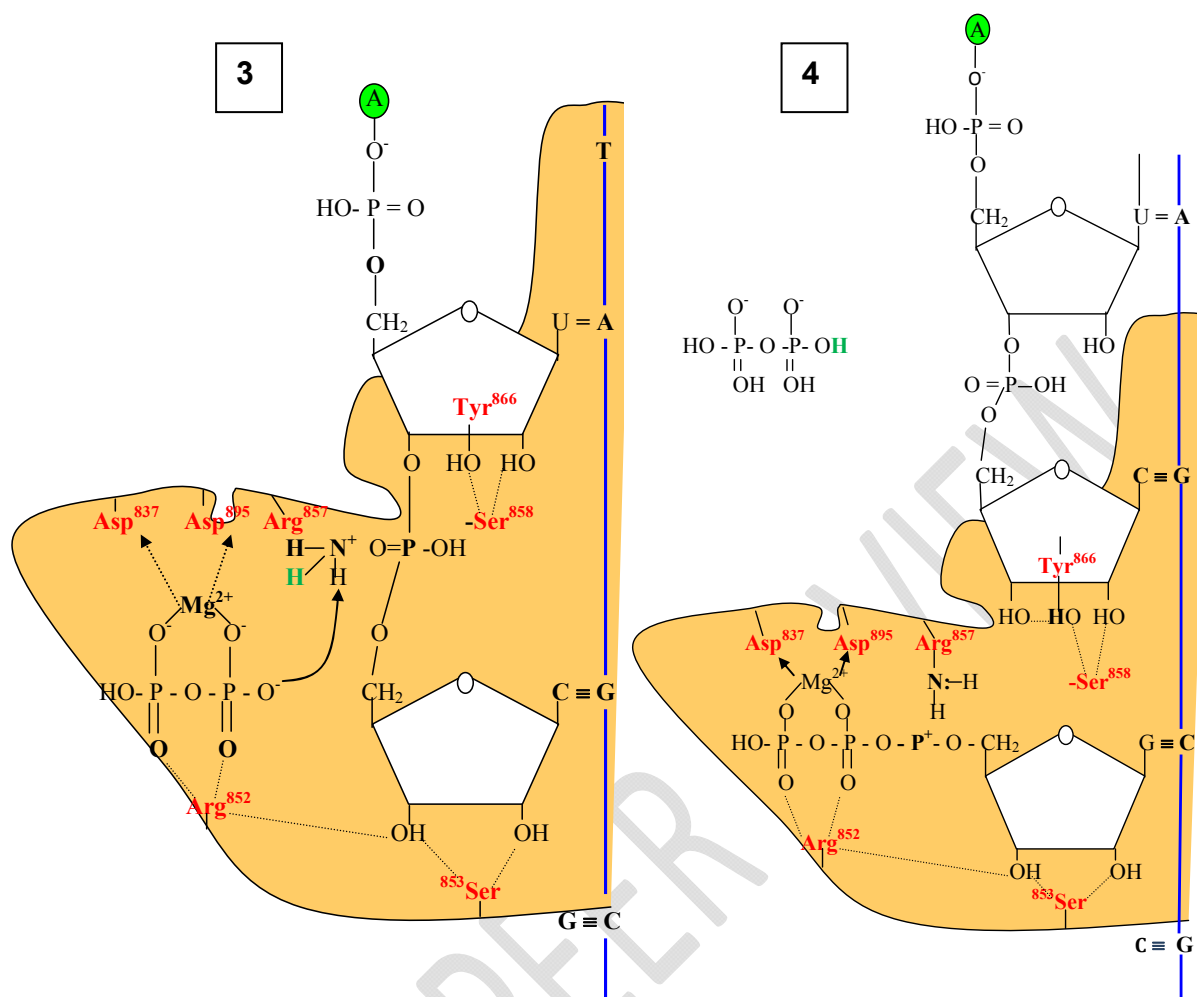
3727 **Step 3. Phosphodiester bond formation:** Proton abstraction by the active site amino acid Arg with
 3728 simultaneous formation of 3'→5' phosphodiester bond with the incoming NTPs.

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Step 4. Inorganic pyrophosphate formation and translocation of the enzyme to next nucleotide: Proton transfer from the active site amino acid Arg, formation of inorganic pyrophosphate, active site restoration and translocation of the enzyme to the next nucleotide (Fig. 8).



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Fig. 8 Steps (1-4) proposed mechanism of the polymerization reactions of transcription by the yeast initiation subunit Rpb2, of the MSU RANAP II

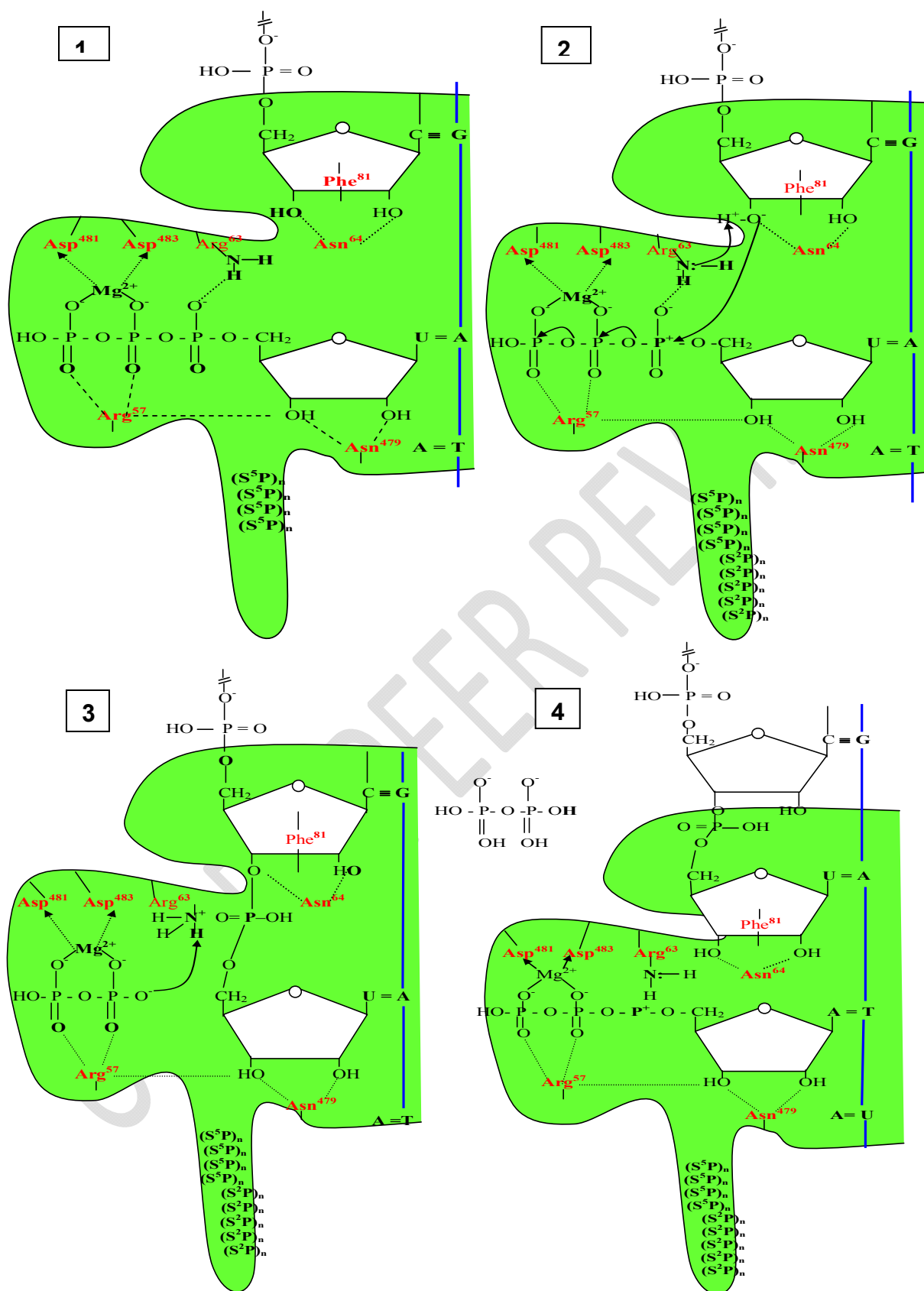
3743 11.2 Mechanism of elongation by Rpb1 subunit of the MSU RNAP II from *S. cerevisiae*

3744 **Step1: Enzyme and the NTP at the Entry Site:** Template binding by the template binding amino
3745 acids (Phe-Gly) and nucleotide discrimination by the invariant amino acids, Watson-Crick base
3746 pairing of the incoming NTP with the template DNA. The catalytic site amino acid Arg is positioned for
3747 proton abstraction.

3748 **Step 2. Proton abstraction and nucleophilic attack of α -phosphate of NTP:** Electronic transition
3749 at the active site for proton abstraction by the active site Arg and an electrophilic and nucleophilic
3750 attack of the incoming NTPs on the 3'-OH.

3751 **Step 3. Phosphodiester bond formation:** Proton abstraction by the active site amino acid Arg with
3752 simultaneous formation of 3'→5' phosphodiester bond with the incoming NTPs.

3753 **Step 4. Inorganic pyrophosphate formation and translocation of the enzyme to next
3754 nucleotide:** Proton transfer from the active site amino acid Arg, formation of inorganic
3755 pyrophosphate, active site restoration and translocation of the enzyme to the next nucleotide (Fig. 9).
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Fig. 9 Steps (1-4) proposed for the polymerization reactions of elongation of transcription by the yeast elongation subunit, Rpb1, of the MSU RNAP II.

3762 12. Proof-reading mechanism and Transcription termination in *S. cerevisiae* MSU RNAP II

3763 Maintaining high fidelity during transcription is essential for the accurate transfer of genetic
 3764 information from DNA to RNA as the first step in gene expression. The proof-reading mechanism is well
 3765 established in DNA polymerases [34]. However, it is poorly understood in RNAPs. Transcription accuracy
 3766 is relatively high, as RNAPs generally misincorporate one wrong nucleotide/~100000 bases. As RNAPs
 3767 are also Zn metalloenzymes the Zn-mediated deletion of the misincorporated NTP could be a possible
 3768 mechanism as explained for DNA polymerases and eubacterial MSU RNAPs as both use DNA template
 3769 to produce a complementary DNA/RNA strand [34, 4, 29, 30]. There is no distinct proof-reading domain
 3770 or separate proof-reading subunit as reported in DNA polymerases. However, in eubacterial and
 3771 eukaryotic elongation subunits of the MSU RNAPs a highly conserved Zn binding motif built into the
 3772 catalytic region itself might involve in the proof-reading function [4]. Presence of two metal ions at the
 3773 active site is supported by a 2.8 Å difference Fourier map, one persistently bound and the other possibly
 3774 exchangeable during RNA synthesis [28]. Unlike the DNA polymerases, the RNAPs could move forward
 3775 (when correct NTP is bound) and backtrack (when wrong NTP is bound and secondary structures) [42].
 3776 When the enzyme stalls due to secondary structures and/or misincorporation, the proof-reading begins
 3777 with fraying of the misincorporated nucleotide away from the DNA template and subsequent backtracking
 3778 of RNAP by one position followed by nucleolytic cleavage of a dinucleotide that contains the
 3779 misincorporated nucleotide [52]. Sydow and Cramer [52] have also proposed that the exo- or
 3780 endonucleolytic cleavage occurs at the same active site that is used for polymerization, which is further
 3781 confirmed by X-ray crystallographic analysis of the *E. coli* and *T. thermophilus* MSU RNAPs suggesting a
 3782 possible Zn mediated cleavage [36]. The complete conservation of the active sites and Zn binding motif in
 3783 all prokaryotic and eukaryotic elongation subunits, found by MSA analysis also further proves this point.
 3784 The proof-reading mechanism may not be that much important during initiation events by the Rpb1
 3785 subunits, as they transcribe only very small regions, i.e., only ~ 10 nts and again such transcripts are well
 3786 within the 5' untranslated regions of the RNA; so any error it makes may not be deleterious. Furthermore,
 3787 a Zn binding motif is not found in the active site region of the initiation subunits. However, proof-
 3788 reading is very important for the elongation subunit, viz., Rpb1 to produce an error-free transcripts, as they
 3789 are going to be translated into active proteins. Therefore, when the enzyme stalls at the wrong nucleotide, it
 3790 could backtrack and make an endonucleolytic cleavage or simply remove the wrong nucleotide by
 3791 exonuclease action [53, 34]. Unlike in DNA-dependent RNA polymerases, where the polymerization and
 3792 proofreading functions are separated, in RNAPs, synthesis and degradation are performed by the same
 3793 active centre [54]. In support of this, Liu et al have shown that following the initiation, RNAP II alone was
 3794 capable of RNA transcript elongation and of proofreading [55]. Thus, the fidelity in the transcription is
 3795 ensured by two mechanisms in MSU RNAPs, viz. by the selection of correct NTPs at the entry site and an
 3796 efficient Zn mediated proof-reading mechanism.

3797 The enzyme could use the same strategy for transcription terminations (intrinsic types) at TTSS.
 3798 That is, when the enzyme stalls due to the stem-loop structures commonly found at the TTSS, it could
 3799 backtrack and make an endo- or exonucleolytic cleavage and release the RNA transcript, using the same
 3800 active site by Zn mediated reaction.

3801 13. Other conserved regions and amino acid(s) in the MSU RNAP II

3802 The above analysis to find out the conserved motifs, active sites and invariant amino acids, with
 3803 respect to the substrate binding and catalytic region, form only a small region of the MSU RNAPs. Apart
 3804 from these, there are a large number of single amino acid conservations like Ys, Ws, Cs, Ps and Gs,
 3805 diads, triads and long stretches of conserved amino acids in all these polymerases (Figs. 4 – 6). The
 3806 highly conserved stretches of amino acids may represent the different conserved structural motifs such as
 3807 loops, rudders, jaws, bridge helix, exit/entry points for DNA/RNA, etc. Thus, they could form small
 3808 functional domains. A good number of completely conserved Ps in these polymerases is implicated in
 3809 making the necessary bents on the enzyme's structure during unique folding, substrate and template
 3810 binding. The completely conserved Cs other than the catalytic region may play a role in the formation of
 3811 disulphide bridges to make the enzyme more compact and stable.

3812 14. CONCLUSIONS

3813 Unlike reported for the DNA polymerases and SSU RNA polymerases, the MSU enzymes of
 3814 eukaryotic MSU RNAPs are similar to the prokaryotic ones and use an R as the catalytic amino acid.
 3815 However, they maintain a different distance conservation in the initiation and elongation subunits. An
 3816 invariant Zn²⁺ binding motif found in the Rpb1 elongation subunits, is proposed to participate in the proof-

3817 reading function. Therefore, it is clear from this communication that even though the transcription
 3818 machinery of eukaryotes is much more complex than that of prokaryotes, the general principals of
 3819 transcription and its regulation are found to be highly preserved. Hence, it is proposed that the MSU
 3820 RNAP II of eukaryotes may also follow very similar polymerization and proof-reading mechanisms like
 3821 their counterparts in eubacteria. MSA data and the available experimental data show that both the
 3822 eubacterial and eukaryotic MSU RNAPs would have evolved from a common ancestor.

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