Research article

Evolutionary connection between the catalytic subunits of DNA-dependent RNA polymerases and eukaryotic RNA-dependent RNA polymerases and the origin of RNA polymerases Lakshminarayan M Iyer, Eugene V Koonin and L Aravind*

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Abstract

Background: The eukaryotic RNA-dependent RNA polymerase (RDRP) is involved in the amplification of regulatory microRNAs during post-transcriptional gene silencing. This enzyme is highly conserved in most eukaryotes but is missing in archaea and bacteria. No evolutionary relationship between RDRP and other polymerases has been reported so far, hence the origin of this eukaryote-specific polymerase remains a mystery.

Results: Using extensive sequence profile searches, we identified bacteriophage homologs of the eukaryotic RDRP. The comparison of the eukaryotic RDRP and their homologs from bacteriophages led to the delineation of the conserved portion of these enzymes, which is predicted to harbor the catalytic site. Further, detailed sequence comparison, aided by examination of the crystal structure of the DNA-dependent RNA polymerase (DDRP), showed that the RDRP and the β' subunit of DDRP (and its orthologs in archaea and eukaryotes) contain a conserved double-psi β -barrel (DPBB) domain. This DPBB domain contains the signature motif DbDGD (b is a bulky residue), which is conserved in all RDRPs and DDRPs and contributes to catalysis via a coordinated divalent cation. Apart from the DPBB domain, no similarity was detected between RDRP and DDRP, which leaves open two scenarios for the origin of RDRP: i) RDRP evolved at the onset of the evolution of eukaryotes via a duplication of the DDRP β ' subunit followed by dramatic divergence that obliterated the sequence similarity outside the core catalytic domain and ii) the primordial RDRP, which consisted primarily of the DPBB domain, evolved from a common ancestor with the DDRP at a very early stage of evolution, during the RNA world era. The latter hypothesis implies that RDRP had been subsequently eliminated from cellular life forms and might have been reintroduced into the eukaryotic genomes through a bacteriophage. Sequence and structure analysis of the DDRP led to further insights into the evolution of RNA polymerases. In addition to the β ' subunit, β subunit of DDRP also contains a DPBB domain, which is, however, distorted by large inserts and does not harbor a counterpart of the DbDGD motif. The DPBB domains of the two DDRP subunits together form the catalytic cleft, with the domain from the β' subunit supplying the metal-coordinating DbDGD motif and the one from the β subunit providing two lysine residues involved in catalysis. Given that the two DPBB domains of DDRP contribute completely different sets of active residues to the catalytic center, it is hypothesized that the ultimate ancestor of RNA polymerases functioned as a homodimer of a generic, RNA-binding DPBB domain. This ancestral protein probably did not have catalytic activity and served as a

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cofactor for a ribozyme RNA polymerase. Subsequent evolution of DDRP and RDRP involved accretion of distinct sets of additional domains. In the DDRPs, these included a RNA-binding Zn-ribbon, an AT-hook-like module and a sandwich-barrel hybrid motif (SBHM) domain. Further, lineage-specific accretion of SBHM domains and other, DDRP-specific domains is observed in bacterial DDRPs. In contrast, the orthologs of the β ' subunit in archaea and eukaryotes contains a four-stranded $\alpha + \beta$ domain that is shared with the α -subunit of bacterial DDRP, eukaryotic DDRP subunit RBP11, translation factor eIF1 and type II topoisomerases. The additional domains of the RDRPs remain to be characterized.

Conclusions: Eukaryotic RNA-dependent RNA polymerases share the catalytic double-psi β -barrel domain, containing a signature metal-coordinating motif, with the universally conserved β ' subunit of DNA-dependent RNA polymerases. Beyond this core catalytic domain, the two classes of RNA polymerases do not have common domains, suggesting early divergence from a common ancestor, with subsequent independent domain accretion. The β -subunit of DDRP contains another, highly diverged DPBB domain. The presence of two distinct DPBB domains in two subunits of DDRP is compatible with the hypothesis that the ultimate ancestor of RNA polymerases was a RNA-binding DPBB domain that had no catalytic activity but rather functioned as a homodimeric cofactor for a ribozyme polymerase.

Background

Polymerization of ribonucleoside triphosphates (NTPs), which is central to a variety of crucial biological processes, including transcription, primer synthesis during DNA replication, addition of polyA tails to mRNA, addition of CCA to tRNA, uridylation in RNA editing, replication of RNA viruses, amplification of RNA in eukaryotic posttranscriptional gene silencing (PTGS), and oligoadenylate synthesis during interferon signaling, is catalyzed by a broad variety of distinct enzymes [1-4]. These polymerases belong to two major mechanistic categories, those that copy a nucleic acid template and those that are independent of a template. The DNA-dependent RNA polymerases (DDRPs) involved in the transcription of cellular and DNA viral genes, primases, and RNA-dependent RNA polymerases (RDRPs) of RNA viruses and cellular PTGS systems are template-dependent polymerases. Templateindependent RNA polymerases (nucleotidyltransferases) include CCA-adding enzymes, polyA polymerases, uridylyl transferases and oligoA-synthetases. Some of the RNA polymerases function as single-subunit proteins, whereas others form large complexes that consist of multiple, distinct subunits; however, in all known cases, the basic catalytic activity maps to a single polypeptide containing a characteristic metal-chelating active site [5-7]. Typically, active sites of polymerases contain acidic or polar residues, which coordinate divalent metal cations, most often Mg²⁺. The metal cations direct a 5' nucleoside triphosphate to form a phosphoester bond with the 3' hydroxyl of the preceding nucleotide (the 3'-terminal nucleotide of a growing polynucleotide chain), with the elimination of pyrophosphate [6].

Despite the basic biochemical similarity among all RNA polymerases, sequence and structure comparisons indi-

cate that these enzymes belong to at least five evolutionarily unrelated folds. The RDRPs of RNA viruses define one major lineage of nucleic acid polymerases, which additionally includes reverse transcriptases, archaeo-eukaryotic DNA polymerases, and nucleotide cyclases [8–13]. The DNA-dependent RNA polymerase of certain bacteriophages, such as T7, and the archaeo-eukaryotic primase (also detected in some bacteria) are divergent derivatives of the same fold [11,14]. The core catalytic domain of all these enzymes, the so-called "palm" domain, has an RNArecognition motif (RRM)-like fold with strategically placed metal-coordinating residues, which form the active site [11,15,16]. In contrast, bacterial DnaG-type primases (also present in archaea and some eukaryotes) contain a polymerase domain of the Rossmann-like TOPRIM fold, which is shared with topoisomerases and OLD-family nucleases [17-19]. The recently solved structures of the DDRPs from yeast and the thermophilic bacterium Thermus thermophilus indicate that the β ' subunit (according to the subunit nomenclature of Escherichia coli DDRP, which we hereinafter employ to designate all orthologs of the respective E. coli subunits) of these enzymes defines another distinct catalytic scaffold, which is unrelated to any of the above template-dependent RNA polymerases [20-24]. Additionally, the structural and evolutionary affinities of two other template-dependent RNA polymerases, namely RDRPs involved in PTGS [25-27] and primases of herpesviruses [28], remain obscure. In contrast to the template-dependent RNA polymerases, which have several distinct scaffolds of the catalytic domains, all template-independent RNA polymerases have the same fold of the principal catalytic domain and belong to the pol β superfamily of nucleotidyl transferases [29,30].

Thus, RNA polymerase activity apparently has been independently "invented" on several occasions. Nevertheless, DDRP is one of the most conserved enzymes, which is represented, without exception, in all cellular life forms [31,32]. The DDRP complex from most organisms consists of 5 to 15 polypeptides. Of these, four subunits, which correspond to the bacterial α , β , β' and ω , are universally present in all cellular DDRPs and constitute the conserved DDRP core [7]. Orthologs of the β and β ' subunits are also encoded in the genomes of several families of large eukaryotic DNA viruses [33-35]. Biochemical studies have shown that the catalytic site of DDRP resides in the β ' subunit and contains three invariant aspartates that coordinate a Mg²⁺ cation [22-24,36]. Recent structural analyses demonstrated that the core of DDRP is assembled around the β and β' subunits, which interact with each other to form a positively charged nucleic-acid-binding cleft. The α subunit further stabilizes this cleft, whereas the ω subunit interacts with the β ' subunit [7]. Given their ubiquitous presence in all cellular life forms and high level of sequence conservation, it seems evident that the core subunits of this complex RNA synthesis machine were already present in the last universal common ancestor (LUCA) of the extant cellular life forms and performed functions mechanistically similar to those of modern DDRP.

In contrast to the ubiquitous DDRP, cellular RDRPs that are involved in PTGS so far have been detected only in eukaryotes [1,26]. The PTGS phenomenon covers a variety of complementarity-dependent silencing pathways, such as RNA interference (RNAi) in animals and slime mold, co-suppression (silencing of transgene and the corresponding endogenous genes) and virus gene resistance in plants, and quelling in fungi, all of which share a common mechanism of RNA turnover [28,37-40]. Essentially, double-stranded (ds) RNA, which is formed in these processes, triggers the activation of a sequence-specific RNA degradation system, which targets homologous RNAs [41,42]. The dsRNA is broken down by the Dicer enzyme into 21-25 nucleotide (nt) fragments called small interfering RNA or siRNAs [43-45]. The siRNAs subsequently associate with a complex of proteins called RISC and target homologous RNA by serving as guides for multiple rounds of RNA cleavage [45,46].

The RDRP is a component of the PTGS system that has been initially described in plants as a cellular RNA polymerase activity induced upon viral infection that synthesized antisense RNA in a primer-dependent or independent manner [26,47–50]. Subsequent genetic studies showed that mutations in RDRP genes impaired PTGS in a variety of systems [25,38,51,52]. Experimental studies in *C. elegans* and in plants suggested that the RDRP is involved in production and amplification of dsRNA using the target RNA as template and siRNAs as primers or guides and resulting in amplification of the RNA silencing response [53–56].

The RDRP is present in one or more copies in a wide range of eukaryotes, from early-branching parabasalids, such as *Giardia*, to multicellular forms, including fungi, plants and animals [57,58]. This broad representation in eukaryotes, including organisms that are considered primitive, suggests that RDRP might have been encoded in the genome of the common ancestor of all modern eukaryotes. However, this gene has been subsequently lost in several eukaryotic lineages, such as the yeast *Saccharomyces cerevisiae*, insects and vertebrates. Although RDRPs display some diversity in domain architecture and the level of sequence conservation, they form a tight, well-conserved family of large proteins with no detectable prokaryotic or viral homologs.

Here, we investigate the evolutionary history of RDRPs and DDRPs in an attempt to unveil the origin of the RDRP. We identify the first homologs of the cellular RDRP outside the eukaryotic clade, in bacteriophages, and show that RDRPs and DDRPs share a homologous catalytic core, which comprises a six-stranded double-psi- β -barrel (DPBB) domain. Evidence is presented that the ultimate ancestor of RNA polymerases might have been a RNAbinding DPBB domain, which functioned as a cofactor for a polymerase ribozyme, and that subsequent evolution of DDRPs and RDRPs proceeded via accretion of various simple modules, such as the sandwich-barrel hybrid motif domain, around this conserved core.

Results and discussion Bacteriophage homologs of the eukaryotic RDRPs and prediction of their active site

To investigate the evolutionary affinities of the RDRPs, we carefully examined the results of BLAST searches of the non-redundant (NR) protein sequence database (National Center for Biotechnology Information, NIH, Bethesda) for various RDRP sequences. These searches identified a large region of 700-800 residues as the conserved module shared by all RDRPs. This region was further examined by searches of NR using the PSI-BLAST program, which was iterated to convergence with a profile inclusion threshold of E = 0.01. These searches, e.g. the search initiated with the core RDRP module from *Petunia* (residues 37–775), readily retrieved RDRPs from plants, C. elegans, Dictyostelium discoideum, several fungi, and Giardia lamblia with statistically highly significant expectation (E) values. Interestingly, the second iteration of this search retrieved the YonO protein from the Bacillus subtilis phage Spβc2 with a significant E-value (7 \times 10⁻³). YonO was also detected with significant E-values in searches initiated with other eukaryotic RDRP modules, e.g., that of Schizosaccha*romyces pombe*. Reciprocal searches started with the YonO protein sequence retrieved closely related proteins encoded by prophages in the genome of *Clostridium acetobutylicum* (CAC1139) and *Clostridium perfringens* (CPE1103). Hereinafter, we refer to these bacteriophage proteins as the YonO-like RDRP homologs (YRHs).

The Gibbs sampling alignment procedure detected 9 statistically significant motifs that are conserved across the entire set of RdRps and YRHs, with a probability of occurrence by chance in this set of protein sequences estimated as <10⁻¹⁸. A multiple alignment of the RDRPs and YRHs was constructed using the T-Coffee program and adjusted using the alignments reported by PSI-BLAST and secondary structure prediction. Secondary structure prediction, which was produced using the multiple alignment as the query, included 16 α -helices and 20 β -strands, which are conserved in RDRPs and YRHs (Fig. 1). Conserved sequence motifs are distributed throughout the aligned region; two motifs with the highest density of conserved residues are located between strands 8 and 11 and strands 18 and 20 (Fig. 1). Twelve residues are conserved in all sequences from the two families, of which nine are charged and one is polar (serine) (Fig. 1). The conserved charged and polar residues conceivably might be involved in catalysis and substrate-binding. The most notable signature shared by the RDRPs and the YRH proteins is the DbDGD (b is a bulky residue) motif located between strands 19 and 20 (Fig. 1). Closely spaced acidic residues that coordinate divalent cations are characteristic of the active sites of most nucleic acid polymerases, in spite of the fact that they belong to several unrelated structural folds [9,13,14,17,28,29,59]. Given that the DbDGD motif is the only set of closely spaced acidic residues shared by the RDRPs and the YRHs, it is likely to form part of the nucleotidyltransferase active site of these enzymes. Thus, these comparisons identify previously undetected prokaryotic homologs of the eukaryotic RDRPs, the YRH proteins, which probably also have RNA-dependent RNA polymerase activity.

Structural and evolutionary relationship between the catalytic domains of the RDRPs and the DDRPs

Interestingly, the PSI-BLAST searches started with the sequences of the RDRP module of the RDRPs and the YRH proteins consistently retrieved the β ' subunits of DDRPs, albeit at statistically not significant E-values. The highscoring segment pairs (HSPs) detected in these searches aligned the highly conserved region of the RDRPs between the predicted strands 18 and 20, including the Db-DGD motif, with the portion of the DDRP β ' subunit sequence, which contains the metal-chelating active site, with a similar conserved motif, DxDGD. Additionally, a search of the NR database with a position-specific scoring matrix (PSSM) that included all unique RDRP and YRH sequences detected, with borderline E-values, a large protein from Corynebacterium glutamicum (Cgl1702), which also contained a DXDGD motif. Reciprocal searches showed that this protein contained two regions of statistically significant similarity to the DDRPs. The Nterminal region of similarity corresponded to the conserved portion of the β subunits and the C-terminal region corresponded to the core of the β ' subunit and included the DXDGD motif (see discussion below). Despite the low statistical significance of the similarity between RDRPs and DDRPs, the presence of the same signature motif in the experimentally identified or predicted active sites of the two classes of enzymes prompted us to perform a detailed comparison of the catalytic domains of these two classes of RNA polymerases.

Among nucleic acid polymerases, a DxDxD motif is conserved only in the RDRPs, the DDRP β ' subunits and the euryarchaea-specific DNA polymerase subunit II. However, in the latter polymerase, the motif has the signature DGDED, as opposed to the DxDGD pattern shared by the RDRPs and DDRPs. To investigate the distribution of this motif in the entire protein database, we conducted pattern searches using the GREF program of the SEALS package, with queries corresponding to the residue conservation profile in the predicted active sites of the RDRPs. All these queries contained the DxDGD motif with additional flanking conserved residues. The only proteins with evolutionarily conserved DxDGD motifs that were detected in these searches were the RDRPs, the β ' subunit of DDRPs, the integrin calcium-binding module, and the EF-hand domain (data not shown). Among these domains, the motif was embedded in the context of conserved (predicted) β-strands only in the RDRPs and DDRPs. Moreover, in searches conducted with extended queries, such as Db-DGDxhxh or DGDxhxh patterns (h is a hydrophobic residue), the RDRPs and the β' subunits of DDRPs were identified as the only protein families in which these motifs were conserved. We also ran database searches using the PHI-BLAST program that combines a regular BLAST search with a pattern search [60]. In these searches, the active site motif was used as the pattern query, and various RDRP sequences as the sequence queries. All these searches retrieved the YonO proteins with significant E-values and some of the DDRP β' subunits with borderline E-values. For example, searches seeded with the RDRP (RrpB gene product) sequence from *Dictyostelium* (gi:14475571) and the pattern [GAS]-D- [FLYMQN]-D- [G]-D-X-[ACLIVMFY]-X- [ACLIVMFY] detected the rice DDRP catalytic subunit with an E-value of 0.048.

Examination of the recently solved structures of the DDRP showed that, although β ' subunit is a large protein, the active site containing the DxDGD motif mapped to a small, compact β -barrel domain. A search of the PDB database



Figure I

Multiple sequenced alignment of the conserved RDRP module from eukaryotic RNA-dependent RNA polymerases and their bacteriophage homologs. The sequences are denoted by gene names, abbreviated species names and Gene Identification (GI) numbers from the GenBank database. Species name abbreviations: At: *Arabidopsis thaliana*, Cac: *Clostridium acetobutylicum*, Ce: *Caenorhabditis elegans*, Cpe: *Clostridium perfringens*, Ddi: *Dictyostelium discoideum*, Diam: *Diaporthe ambigua*, Diper: *Diaporthe perjuncta*, Giin: *Giardia intestinalis*, Lyes: *Lycopersicon esculentum*, Nc: *Neurospora crassa*, Nt: *Nicotiana tabacum*, Os: *Oryza sativa*, Pethy: *Petunia hybrida*, Phsp: *Phomopsis* sp., Sp: *Schizosaccharomyces pombe*, Spbc2: Bacteriophage Spβc2. The positions of the first and the last residue of the aligned region in the corresponding protein are indicated before and after each sequence, respectively. The numbers between aligned blocks represent poorly conserved inserts that are not shown. The coloring is based on the 95% consensus shown underneath the alignment; h indicates hydrophobic residues (ACFILMVWY), a indicates aromatic residues (FYW), I indicates aliphatic residues (ILVA), p indicates polar residues (STED-KRNQHC), c indicates charged residues (DEKR), bi indicates bulky residue (ILFYWMKREQ), **s** indicates small residues (AGS-VCDN). The predicted secondary structure elements are shown below the alignment; H indicates α -helix and E indicate extended conformation (β -strand). The predicted helices are marked H1-16 and the predicted strands S1-20. The signature motif DbDGD, which is predicted to form part of the catalytic site, is shown in reverse shading. The three sequences at the bottom are those of bacteriophage homologs of the RDRPs (the YRH proteins); the remaining are RDRP sequences.

using the DALI program [61], with this β -barrel from *Thermus thermophilus* β' subunit submitted as a query (pdb: 1iw7, chain D region: 625–749), retrieved the N-terminal domain of the CDC48-like AAA ATPases, formate dehydrogenases, aspartate decarboxylases and Barwin. The reported structural alignments span ~70 residues with

significant Z-scores (8.8–6.6) and root mean square deviations (RMSD) between the C- α backbones in the range of 1.8–2.5 Å, which is characteristic of homologous domains. The region of structural similarity shared by the catalytic domain of the β ' subunit of the DDRPs and the above proteins corresponds to a distinct globular fold known as the double-psi beta-barrel (DPBB) [62–64]. The SCOP database included the RNA polymerase subunits in the multidomain protein class but mentions the presence of a DPBB domain in each of the β and β ' subunits ([65]; see below).

The region of sequence similarity between the RDRPs and the β ' subunits of DDRPs, that was detected in the searches described above, almost precisely mapped to the DPBBfold catalytic domain of DDRP. The multiple-alignmentbased secondary structure prediction for the RDRPs and the YRHs indicated that, like the similar region of the β' subunit, this region was enriched in β -strands (Fig. 1). Multiple alignment constructed on the basis of sequence alignments produced in the PSI-BLAST and PHI-BLAST searches and superposition of the predicted secondary structure elements of RDRP and the experimentally determined structure of DDRP revealed considerable concordance between the RDRP and DDRP families. In addition to the DbDGD motif, which is located in the loop between strands 5 and 6, several residues are conserved between these polymerase families (Fig. 2). These include hydrophobic residues in the β -strands, a conserved proline after strand 2, a small (typically, glycine) and polar (typically, aspartate) residues before strand 3, small residues that typically mark the boundaries of secondary structure elements and several charged and polar residues (Fig. 2). A comparison of the RDRP and DDRP sequences using the Gibbs sampling method identified two conserved blocks, which were highly statistically significant (E $< 10^{-16}$) within the analyzed sequence set. One of these blocks centered at the DbDGD motif, whereas the other one encompassed strands 3 and 4, which contain several partially conserved hydrophobic and polar positions (Fig. 2).

Importantly, approximately 10 conserved residues that are detected in the structure-based alignments of the DPBB domains are also conserved in the RDRPs (Fig. 3). These include a doublet comprised of a small and a polar residues preceding each of strands 3 and 6, a small residue after strand 2 (proline in the DDRP β ' subunit and RDRP), bulky and polar residues in strand 3, a small residue immediately downstream of strand 5, a bulky residue in the loop between strands 5 and 6, and two hydrophobic residues in strand 6 (Fig. 3). The predicted DPBB domains of the RDRPs differ from most of the DPBB domains of DDRP β' subunits in that the former lack a long and variable, in both length and sequence, insert between strands 2 and 3 that is characteristic of the latter (Fig. 2). However, the DPBB from the RNA polymerases of yeast killer plasmids and the divergent predicted corynebacterial polymerase Cgl1702 also lack this insert (Fig. 2). Taken together, these observations suggest that, although RDRPs and the β ' subunits of DDRPs show limited sequence similarity, these two RNA polymerase families share a homologous catalytic domain, which consists of a double-psi-βbarrel containing the metal-coordinating DbDGD motif.

The DPBB domains and early evolution of the catalytic domain of RNA polymerases

The SCOP database, while classifying the DDRP subunits in the multidomain category, recognizes two DPBBs in the DDRP complex. One of these corresponds to the catalytic, metal-coordinating domain, containing the DbDGD motif, in the β' subunit (residues: 626-750 of pdb id:1iw7 chain D), whereas the other one corresponds to the conserved core domain of the β subunit (residues:673–994 of pdb id: 1iw7 chain C). The DPBB domains of the β and β' subunits interact to form the catalytic cleft of the RNA polymerases, with two conserved lysines in the β subunit protruding into the cleft and interacting with the substrate (Fig. 4). However, the DPBB domain in the β subunit is distorted by two large inserts (Fig. 4 and see below) and shows no detectable sequence similarity to the β' subunit DPBB. Because of these inserts and the resulting distortion, the DPBB of the β subunit does not show similarity to any domains in structure database searches. Nevertheless, a structural alignment based on visual inspection confirms the presence of all the bona fide features of the DPBB domain in the β subunit (Fig. 3). The Cgl1702 protein from Corynebacterium, which we identified as one of the most divergent members of the DDRP clade, showed similarity to the other DDRPs only in two regions, which corresponded to the DPBBs from the β and the β ' subunit. This observation, taken together with the spatial proximity of the two DPBBs in the catalytic cleft of the DDRP (Fig. 4), indicates that these domains comprise the ancestral conserved core of the RNA polymerases. The arrangement of the two DPBBs in the β and β' subunits suggests that, in the primordial ancestor of the DDRPs, the two DPBBs formed a "head-to-tail" homodimer that bound RNA at the domain interface.

Although the cores of the two DDRP subunits are homologous and are likely to have evolved from a common ancestor (see discussion below), they have completely different sets of conserved residues and contribute distinct moieties to the catalytic cleft (Figs. 3, 4). Thus, these specific active residues apparently emerged after the divergence of the β and β ' subunit DPBBs, whereas the ancestral DPBB homodimer probably did not have catalytic activity and merely bound a ribozyme that originally catalyzed the RNA polymerization. Subsequently, the two subunits diverged from each other, with one (β ') acquiring a divalent cation-binding site in the insert between the two terminal strands of the DPBB, whereas the DPBB in the other (β) subunit acquired two basic residues that interacted with the nucleic acid. At this stage, which might be considered the emergence of modern-type RNA

		S1	S2	S3	S4	S5	S6		
Secondary Str.		EEEEEEEE	.EEEEE.	EEEE	EEEEEEE	EEE	EEEEEE		
RrpB_Ddi_14475571	1121	RKYEK <mark>V</mark> IE G L	- <mark>V</mark> MVIKNP	CTHPGD-VRYLKAVDNLRLRH	-LRNVL <mark>V</mark> FSTKG	DVPNFKE <mark>I</mark>	SGSDLDGD-RYFFCY	D 1191	Eukaryotic
RrpA Ddi 14475569	1297	RKYEK <mark>V</mark> IEGL	- <mark>V</mark> MVIKNP	CTHPGD-VRYLKAVDNIRLRH	-LRNVL <mark>V</mark> FSTKG	DVPNFKE <mark>I</mark>	SGSDLDGD-RYFFCY	D 1367	RDRP
At2q19910 At 15225263	606	ILHSGOISGD	- <mark>V</mark> LVYRNP	GLHFGD-IHVLKATYVKALED	3 NAKFAVFFPOKG	PRSLGDE	AGGDFDGD-MYFISR	N 678	
P0419B01.14 Os 14209603	855	ILDYGOVSGD	- <mark>V</mark> LVYKYP	GLHPGD-IHVLKATYSSDIEK	3 NSKHAILFPTTG	ORSLADE	ANSDFDGD-IYWVSL	N 927	
RdRp Giin 15419599	599	GLNISCVEGR	-VVAKNP	CFYRGD-VRIYHAVNIFERLR	23 MYRG VI<mark>C</mark>FPKYY	EGRPMTDC	SGSDLDGD-IYWVSW	D 694	
F26A3 3 Ce 7499876	2497	VHGSETVTGT		CIVEGD-VRIFEAVDIPELHH				2567	
F26A3 8 p Ce 17508703	888	VHGSETVTGT	VLLTKNP	CIVEGD-VRIFEAVDIPELHH		PRPHPDE		958	
M01G12 12 p Ce 17508365	869	VLCSETVTCT	VILTEND	GIVE CD VRIFEAVDI PELHH		DPDHDDF		939	
FG01 Ce 17506499	915	NAAROVI.TCT	VLLTKNP	CIVAGD-VRIFEAVDI PELHH				985	
EGO1_CC_1/500155	1000	CMDDI	VI TTEND		LUDWUEDOUG			1160	
PDD 1 Dimor 19656909	1090	TTNVENTOT		ALUDOD WRVENUDRIDU				1000	
RDP_1_Diper_18050898	939			ALHPGD-VKVVEAVDVPELKH				1019	
RDR_1_D1am_15422180	940			SLHPGD-VRVVEAVDALELRH	-LKNVVVFPRTG			0 1010	
Rarp_Prisp_1/22/102	559	TINYE <mark>V</mark> IIGI	CVVGRNP	SLHPGD-VRVVEAVDVPELRH			SGG_M_GDYFVYW	0 629	
SADI_NC_13641451	910	NGGYR <mark>V</mark> IT G T	-CVVGRNP	SLHPGD-IRVVEAVDVPALRH	-LRDVV <mark>V</mark> FPL TG	DRDVPSMC	SGGDLDGD-DFFVIW	D 980	
RdRP_Sp_19114207	840	KLKTR <mark>V</mark> IVGL	- <mark>C</mark> IVARNP	SLHPGD-VRVCKAVRCDELMH	-LKNVI <mark>V</mark> FPTTG	DRSIPAM <mark>C</mark>	SGGDLDGD-EYTVIW	D 910	
B1074C08.21_Os_21328146	828	DKNTE <mark>V</mark> ILGT	- <mark>V</mark> VIAKNP	CLHPGD-VRILEAVDVPELHH	-LVDCL <mark>V</mark> FPQKG	ERPHANE <mark>A</mark>	SGSDLDGD-LYFVTW	D 898	
SGS2_At_15229153	804	KTDLE <mark>V</mark> VK G Y	- <mark>V</mark> AIAKNP	CLHPGD-VRILEAVDVPQLHH	-MYDCL <mark>I</mark> FPQKG	DRPHTNE <mark>A</mark>	SGSDLDGD-LYFVAW	D 874	
SDE1_At_8248473	804	KKDLE <mark>V</mark> VK G Y	- <mark>V</mark> AIAKNP	CLHPGD-VRILEAVDVPQLHH	-MYDCL <mark>I</mark> FPQKG	DRPHTNE <mark>A</mark>	SGSDLDGD-LYFVAW	D 874	
RdRp_Nt_7489197	744	ANCDF <mark>I</mark> LKGN	- <mark>V</mark> VVAKNP	CLHPGD- <mark>IRV</mark> LRAVD <mark>V</mark> PA <mark>L</mark> HH	-MVDCV <mark>V</mark> FPQKG	K <mark>R</mark> PHPNE <mark>C</mark>	SGS <mark>DLD</mark> G <mark>D-IY</mark> FVCW	D 814	
RdRp_Lyes_7489070	742	TNSNF <mark>I</mark> LKGN	- <mark>V</mark> VVAKNP	CLHPGD- <mark>IRV</mark> LKAVN <mark>V</mark> RA <mark>L</mark> HH	-MVDCV <mark>V</mark> FPQKG	K <mark>R</mark> PHPNE <mark>C</mark>	SGS <mark>DLD</mark> G <mark>D-IY</mark> FVCW	D 812	
At1g14790_At_15223906	738	PGRRF <mark>I</mark> ITGP	- <mark>V</mark> VVAKNP	CLHPGD- <mark>VRV</mark> LQAVN <mark>V</mark> PA <mark>L</mark> NH	-MVDCV <mark>V</mark> FPQKG	L <mark>R</mark> PHPNE <mark>C</mark>	SGS <mark>DLD</mark> G <mark>D-IY</mark> FVCW	D 808	
At4g11130_At_15237051	769	DEETS <mark>V</mark> VIGK	- <mark>v</mark> vvtkn p	CLHPGD-IRVLDAIYEVHFEE	1 GYLDCI <mark>I</mark> FPQKG	ERPHPNE <mark>C</mark>	SGG <mark>DLD</mark> G <mark>D-QF</mark> FVSW	D 841	
QDE1_Nc_4803727	948	ESFTLLSDCD	- <mark>V</mark> LVARSP	AHFPSD-IQRVRAVFKPELHS	-LKDVI <mark>I</mark> FSTKG	DVPLAKK <mark>I</mark>	SGG <mark>DYD</mark> G <mark>D-MA</mark> WVCW	D 1018	
RdRp_Pethy_4138343	435	ANYNF <mark>I</mark> LKGN	- <mark>v</mark> vvaknp	CLHPGD-IRVLKAVDVPALHH	-MVDCV <mark>V</mark> FPQKG	KRPHPNE <mark>C</mark>	SGSDLDGD-IYFVCW	D 505,	/
CPE1103_Cpe_18310085	478	LAEKE <mark>F</mark> YI A GWE	-GEKVTI S 4	ACYQEI-QKTILTDKLDKYCK	DYTSEI <mark>I</mark> FFNQ-	KDNTHML <mark>M</mark>	ISGADEDGD-GCKCFK	N 554	YRH
yon0 Spbc2 9630188	473	LSGSD <mark>V</mark> HC S LYD	-EGYIDIL 1	SPHLFR-EHGVRWNKKNEEYE	1 WFI TP G <mark>V</mark> YTSI-	HDPISKL <mark>I</mark>	O-F <mark>DND</mark> GD-KALIIS	D 546	
CAC1139 Cac 15894423	452	FSDGE <mark>V</mark> YT P -LFK	-DSEEIVG 2	NPHVCS-GNVLLVKNKYDWRL	5 FTENIV <mark>A</mark> VNLI-	DFPLODT <mark>l</mark>	OGSDODSD-TVLLTN	N 531,	/
14.7D mbbb 01720240	606		UDGGLD 47		ON FORM	TOLUDIU			DDDD
DD0011 Dm 15905036	642	SGRSVIVGPQLK-	LUCCOUP 47	EVITEDVINULNDADT				P 750	URDP
DR0911_D1_15805950	250	SGRSVIVGPULK-		EVIEDAVVELINAPILARDGI				P /0/	
Rpoc_Ec_15804578	350	SGRSVIIVGPILK	HQCGLP 44	EVIREHPVLLNRAPTLHRLGI	QAFEPVLIEGKA		AAYNA FOGO-QMAVHV	P 4/1	
rpoc_Atu_1/935845	352	SGRSVIVIGPELK	HQCGLP 44	EVIREHPVLENRAPTEREGI	QAFEPMLVEGKA		TAFNA FOG -QMAVHV	P 4/3	
mir02//_MIO_134/0543	1720	SGRSVIVIGPELK	HQCGLP 44	EVIREHPVLLNRAPTLHRLGI	QAFEPILIEGKA		TAFNA FOG - MAVHV	P 4/3	
P1198_Hp_15645812	1/39	SGRSVIVVGPNLK	MDECGLP 44	EITEGYPVLLNRAPTLHKQSI	QAPHPKLIDGKA	IQLHPLV <mark>C</mark>	SAFNADFDGD-QMAVHV	P 1860	
rpoC_Uu_13357746	408	SGRSVIVVGPELK	MYEVGIP 53	KVIKQRP VIL NRAPTLHRLGI	QAF EP K M VDG KA	IRLHPLVI	"TAFNADFDGD-QMAVHI	P 538	
HrpoC_Bs_16077176	339	SGRSV <mark>I</mark> VVGPHLK	-MYQCGLP 44	SVIKEHP VLL NRAPTLHRLGI	QAF EP T <mark>L</mark> VEG RA	IRLHPLV <mark>C</mark>	TAYNADFDGD-QMAVHV	P 460	
rpoC_Mpn_13508254	411	SGRSV <mark>I</mark> VVGPELK	MYEVGIP 53	DVIKDRP VLL NRAPTLHRLGI	QAF EP R <mark>I</mark> VDG KA	IRLHPLVI	TAFNADFDGD-QMAVHV	P 541	
TM0459_Tm_15643225	641	SGRAV <mark>I</mark> VVGPNLK	HQCGIP 46	EVIKGSV <mark>VLL</mark> NRAPT <mark>L</mark> HR <mark>M</mark> SI	QAF EP K <mark>L</mark> VEG NA	IQLHPVV <mark>C</mark>	PPFNADFDGD-QMAVHV	P 764	
rpoC1_Ssp_16330858	357	SGRSV <mark>I</mark> VVGPNLK	- <mark>I</mark> YQCGLP 44	EVITGHP VML NRAPT <mark>L</mark> HR <mark>L</mark> GI	QAF EP I <mark>L</mark> VEG RA	IQLHPLV <mark>C</mark>	PAFNADFDGD-QMAVHV	P 478	
rpoC_Ct_15605035	353	SGRSV <mark>I</mark> IVGPELK	- <mark>f</mark> nqcglp 44	EIIK G HP <mark>VLL</mark> NRA PTLHRLGI	QAF EP V <mark>L</mark> IEGK A	IRVHPLV <mark>C</mark>	AAFNADFDGD-QMAVHV	P 474	
TP0242_Tp_15639234	339	SGRSV <mark>I</mark> VVGPELK	- <mark>l</mark> wqcglp 44	EVVKEHP VML NRAPT <mark>L</mark> HR <mark>L</mark> GI	QAF EP V <mark>L</mark> VEGK A	IRLHPLV <mark>C</mark>	KPF NADFDGD-QM AVHV	P 460	
rpoC_Aae_15606950	480	SGRSV <mark>I</mark> VVGPELQ	- <mark>m</mark> hqcglp 44	EVVKEHP <mark>VLL</mark> NRA PTL HRPSI	QAF EP V <mark>L</mark> VEGK A	IQLHPLV <mark>C</mark>	PPFNADFDGD-QMAVHV	P 601	
rpoC_Mtu_15607808	425	SGRSV <mark>I</mark> VVGPQLK	- <mark>l</mark> hqcglp 44	EVIAEHP <mark>VLL</mark> NRA PTL HR <mark>L</mark> GI	QAF EP M <mark>L</mark> VEGK A	IQLHPLV <mark>C</mark>	EAFNADFDGD-QMAVHL	P 546	
rpoA1_Sso_15897171	321	SSRTV <mark>I</mark> SPDPNIS	- <mark>I</mark> DEVGVP 69	HLIDGDI <mark>VLF</mark> NRQPS <mark>L</mark> HR <mark>I</mark> SM	MAHRVR <mark>V</mark> LKGLT	F <mark>R</mark> LNLLV <mark>C</mark>	PPY NA<mark>DFD</mark>G<mark>D-EM</mark>NLHV	P 467	
APE1853_Ap_14601674	332	SARTV <mark>I</mark> SPDPYIS	- <mark>I</mark> NEVGVP 70	HLINGDI <mark>VLF</mark> NRQPS <mark>L</mark> HR <mark>M</mark> SI	MGHIVR <mark>V</mark> MPGKT	F <mark>R</mark> LNLLV <mark>C</mark>	PPY NA<mark>DFD</mark>G<mark>D-EM</mark>NLHV	P 479	
rpoA1_Mac-C2A_20090127	321	SARTV <mark>I</mark> SPDPNLS	- <mark>I</mark> NEVGVP 68	QLKDGDT <mark>VLF</mark> NRQPS <mark>L</mark> HK <mark>M</mark> SI	MAHSVK <mark>V</mark> LPNK T	FRLNPAV <mark>C</mark>	PPYNA <mark>DFD</mark> G <mark>D-EM</mark> NMH <mark>A</mark>	L 466	
rpoA1_Hsp_16554525	393	SARTV <mark>I</mark> SPDPTLS	- <mark>L</mark> NEVGVP 68	HLIDGDI <mark>IIF</mark> NRQPS <mark>L</mark> HR <mark>M</mark> SI	MAHEVV <mark>V</mark> MPYKT	FRLNTVV <mark>C</mark>	PPY NA<mark>DFD</mark>G<mark>D-EM</mark>NMH<mark>A</mark>	L 538	
RPB1_Sc_6320061	348	SARTV <mark>I</mark> SGDPNLE	- <mark>L</mark> DQVGVP 67	HIMDNDP <mark>VLF</mark> NRQPS <mark>L</mark> HK <mark>M</mark> SM	MAHRVK <mark>V</mark> IPY ST	FRLNLSVI	SPY NA<mark>DFD</mark>G<mark>D-EM</mark>NLH<mark>V</mark>	P 492	
RNAPII_Hs_7434727	362	SARTVITPDPNLS	- <mark>I</mark> DQVGVP 67	HMCDGDIVIFNRQPTLHKMSM	MGHRVR <mark>I</mark> LPWST	FRLNLSVI	TPY NA<mark>DFD</mark>G<mark>D-EM</mark>NLHL	P 506	
J6R_VV_9791022	285	TARSV <mark>I</mark> GPSTSIT	- <mark>V</mark> NEVGMP 64	AVQEYTS <mark>IIF</mark> GRQPS <mark>L</mark> HR <mark>Y</mark> NV	IAS <mark>S</mark> IR <mark>A</mark> TEGD T	IKISPGI <mark>A</mark>	NSQNADFDGD-EEWMIL	E 426	
RPB1 At 15233338	360	SARTV <mark>I</mark> TPDPTIN	- <mark>I</mark> DELGVP 70	HLODGDF <mark>VLF</mark> NROPSLHKMSI	MGHRIR <mark>I</mark> MPYST	FRLNLSVT	SPYNADFDGD-EMNMHV	P 507	
	479	AARSV <mark>I</mark> SP D PNIE	TNEIGVP 81	HIKNRDV <mark>VLM</mark> NRQPTLHKASM	mghkvr <mark>v</mark> lpnek	TLRLHYANT	GAY NADFDGD-EM NMHF	P 638	
rpc-1 Ce 21913089	360	SGRTV <mark>I</mark> SPDPNLR	IDOVGVP 69	HLDDNDVVLFNROPSLHKISI	MSHRAK <mark>V</mark> MPGRT	FRFNECAC	TPYNADFDGD-EMNLHL	P 506	
rpoC1 At 7525024	379	SGRSVIVVGPSUS	LHRCGLP 44	EVMOGHPVLLNRAPTLHRLGT	OSFOPILVEGRT	ICLHPLVC	KGFNADFDGD-OMAVHV	P 500	
RPC1 Sc 6324690	376	SGRTVISPDPNUS-	IDEVAVP 69	HLEDGDVVLFNROPSI HRLST	LSHYAKIRPWRT	FRINECVC	TPYNADEDGD-EMNLHV	P 522	
CG17209 Dm 7293125	363	SGRTVISPDPNIM	INOVGVP 69	HURDGDTVLFNROPSIHKMST	MCHRAKVOPORT	FRENECAC	TPYNADEDGD-EMNLHL	P 509	
Call702 Cal 19552977	1521	SATAVWTGDPRIS	VDEVAMN 7	GVLDNGYAMVWRDPVT RDGGV	RYLRVVINDI.H	3VAVNPVS	KSFDGDFDGD-SVGLVG	N 1606	
ORF6 Klla 133456	786	CIRSUTVPDPTID	TOTIKIP 2	ANIGCEYGLINROPSUMUST	KI WKI KOGSNKT		OSENADED CD-EMNTYC	T 865	
Lef-9 BNV 9630869	107	TTNYL TLPHFTLWD	NEVIELN 78	RDVDNVRAWTTRHPNTSOT	OVSDVHTNFSST	DWNVKVGLGT	PGANTDCDGDKKTTTET	P 268	/
consensus/85%	107	h s	h P	c lbh c ssh ph	ns lh ng	n h	suDbDGD bb b b	- 200,	
001100110000/0000				······································	· · · bo · · · · bo	· · · P · · · · · ·		-	

Multiple alignment of the double-psi β -barrel (DPBB) domains from the β ' subunit of DNA-dependent RNA polymerases with the predicted DPBB domains of RNA-dependent RNA polymerases. The conventions for naming sequences and coloring conserved residues are as described in the legend to Figure 1. The shared secondary structure elements are shown above the alignment with E denote the β -strand (extended) conformation. Non-conserved regions are depicted as numbers with inserts. The species abbreviations are as in figure Fig. 1. The species abbreviations that are not listed in the legend to Figure 1 are: Thth: Thermus thermophilus, Dr: Deinococcus radiodurans, Ec: Escherischia coli, Atu: Agrobacterium tumefaciens, MIo: Mesorhizobium loti, Hp: Helicobacter pylori, Uu:Ureaplasma ureolyticum, Bs: Bacillus subtilis, Tm: Thermotoga maritima, Ssp: Synechocystis species, Aae: Aquifex aeolicus, Ct Chlamydia trachomatis, Mpn Mycoplasma pneumoniae, Mtu: Mycobacterium tuberculosis, Tp: Treponema pallidum, Sso: Sulfolbus solfataricus, Ap: Aeropyrum pernix, Mac: Methanosarcina acetivorans, Hsp: Halobacterium species, Sc: Saccharomyces cerevisiae, Hs: Homo sapiens, VV: Vaccinia virus, BNV: Bombyx mori Nuclear Polyhedrosis virus, Cgl: Corynebacterium glutamicum, Klla: Kluyveromyces lactis.

polymerases, the catalytic activity probably was taken over by the protein. The basic DPBB fold consists of six β -strands and two variable regions, which are located after strand 2 and strand 5 and, at least in some case, adopt a helical conformation [62–64]. The domain can be further split into two

		S1		S2		S3		S4		S5			S6		
Secondary Str.		EEEEEEEE		.EEEEE.		EEEE		. EEEEEEE		EEE			EEEEE.		
RrpA_Ddi_14475569	1297	RKYEKVIEGL		VMVIKNP	4	GD-VRYLKAV	б	H-LRNVLVFSTKG		DVPNF	4	GSDLDGD-	R <mark>Y</mark> F <mark>F</mark> CYD	1367	\RdRp
EG01_Ce_17506499	915	NAARQVLTGT		VLLTKNP	4	GD-VRIFEAV	б	H-MCD V VVFPQHG		PRPHP	4	GSDLDGD-	E <mark>Y</mark> S <mark>I</mark> IWD	985	
SAD1_Nc_13641451	910	NGGYRVITGT		CVVGRNP	4	GD-IRVVEAV	6	H-LRD V VVFPLTG		DRDVP	4	GGDLDGD-	D <mark>f</mark> f <mark>v</mark> iwd	980	
RdRP_Sp_19114207	840	KLKTRVIVGL		CIVARNP	4	GD-VRVCKAV	6	H-LKNVIVFPTTG		DRSIP	4	GGDLDGD-	E <mark>Y</mark> T <mark>V</mark> IWD	910	ĺ
SDE1_At_8248473	804	KKDLEVVKGY		VAIAKNP	4	GD-VR <mark>I</mark> LEAV	6	H-MYD C LIFPQKG		DRPHT	4	GSDLDGD-	L <mark>Y</mark> F <mark>V</mark> AWD	874	/
1iw7D_Thth_21730340	626	SGRSVIVVGPQLK		LHQCG <mark>L</mark> P	51	GKVVL<mark>L</mark>NR AP	6	IQAFQ P VLVEGQS		I Q <mark>L</mark> HP	6	NADFDGD-	Q <mark>M</mark> A <mark>V</mark> HVP	750	\DRDPbetaprime
RpoC_Ec_15804578	350	SGRSVITVGPYLR		LHQCG <mark>L</mark> P	51	EHPVL <mark>L</mark> NRAP	б	IQAFEPVLIEGKA		i q <mark>l</mark> hp	6	NADFDGD-	Q <mark>M</mark> A <mark>V</mark> HVP	471	
RpoA1_Sso_15897171	321	SSRTVISPDPNIS		IDEVG <mark>V</mark> P	73	GDIVLFNRQP	б	MMAHRVLKGLT		F R <mark>L</mark> NL	6	NADFDGD-	E <mark>M</mark> N <mark>L</mark> HVP	467	
RpoA1_Hsp_16554525	393	SARTVISPDPTLS		lnevg <mark>v</mark> p	72	GDII IF NRQP	6	IMAHE V VVMPYKT		FRLNT	6	NADFDGD-	E <mark>M</mark> NMHAL	538	
RPB1_Sc_6320061	348	SARTVISGDPNLE		ldqvg <mark>v</mark> p	71	NDPVL <mark>F</mark> NRQP	6	$\mathtt{MMAHR} \mathbf{V} \mathtt{KVIPYST}$		FR <mark>L</mark> NL	6	NADFDGD	E <mark>M</mark> N <mark>L</mark> HVP	492.	/
DR0912_Dr_15805937	732	TIAIMPFDGFNFE	3	CINED <mark>L</mark> V	140	GD-KVANRHG		N K GVVSKIVRPED	2	IV <mark>F</mark> NP	107	GEPISGPV	V <mark>V</mark> G <mark>I</mark> MY <mark>V</mark>	1054	\DRDPbeta
RpoB_Ec_15834164	801	RVAFMPWNGYNFE	3	lvser <mark>v</mark> v	239	GD-KMAGRHG		NKGVISKINPIED	2	i v l np	117	GEQFERPV	T <mark>V</mark> G <mark>Y</mark> MY	1232	
aq_1939_Aae_15606949	885	LVAFMPWRGYNFE	3	VISER <mark>L</mark> V	259	GD-KMAGRHG		NKGVISVVLPVED	2	i v l np	123	GEPFDFEV	T <mark>V</mark> G <mark>Y</mark> MH	1342	ĺ
RpoB_Hp_15645812	811	RVAFMPWNGYNFE	3	VVSEC <mark>I</mark> T	257	GD-KMAGRHG		NKGIVSNIVPVAD	2	I V L NP	121	GEKMRERV	'N <mark>V</mark> G <mark>Y</mark> MY <mark>M</mark>	1264	
RpoB_Ana_17229086	687	VVAYMPWEGYNYE	3	LISER <mark>L</mark> V	139	GD-KMAGRHG		NKGIISRILPIED	2	I V L NP	78	GEAFDRPV	T <mark>I</mark> G <mark>V</mark> AY <mark>M</mark>	979	
TP0241_Tp_15639233	769	LVGFVPWNGYNYE	3	LISHR <mark>V</mark> V	139	GD-KMAGRHG		NKGIVARILPEED	2	VCLNP	63	GDYFQNPV	F <mark>V</mark> G <mark>V</mark> IY <mark>F</mark>	1046	
RpoB_Bs_16077175	760	MVGFMTWDGYNYE	3	IMSER <mark>L</mark> V	139	GD-KMAGRHG		N K GVI S KILPEED	2	IMLNP	63	GEPFDNRV	'S <mark>V</mark> G <mark>I</mark> MY <mark>M</mark>	1037	
PH1546_Ph_14591329	720	VVAVLAYHGYNME	3	IINKASI	130	GD-KFASRHG		QKGVIGLIVPQED	2	LIVNP	63	GKRFEADI	F <mark>I</mark> G <mark>I</mark> IY <mark>Y</mark>	988	
VNG2665G_Hsp_15791389	210	VVAVMSYEGFNIE	3	VMNKGSV	131	GD-KFASRHG		QKGVVGHLAPQED	2	LVLNP	63	GEKIEAEI	F <mark>V</mark> G <mark>T</mark> IF <mark>Y</mark>	479	
MJ1041_Mj_15669230	212	VVAIMSYEGYNME	3	vfnks <mark>a</mark> i	130	GD-KFASRHG		QKGVMGLTVPQED	2	I I I NP	63	GKKFEVEI	Y <mark>I</mark> G <mark>I</mark> AY <mark>Y</mark>	480	
RpoB1_Af_11499471	207	VVAVLSYEGYNIE	3	IMNKGSV	130	GD-KFASRHG		QKGVVGLLVPEED	2	L I I NP	63	GRRYLVDI	F <mark>V</mark> G <mark>V</mark> IY <mark>Y</mark>	475	
APE1856_Ape_14601675	760	VVAIMSYTGYNIE	3	IFNKGSI	131	GD-KFASRHG		QKGVIGMIFPRYD	2	VI <mark>L</mark> NP	63	G QI I E A PI	T <mark>I</mark> G <mark>V</mark> AY <mark>Y</mark>	1029	
RpoB1_Sso_15897172	248	ILAVMSFTGYNME	3	IMNRSSV	129	GD-KFASRHG		QKGVIGMLIPQVD	2	I I L NP	63	GQKIKSRI	Y <mark>f</mark> g <mark>v</mark> vy <mark>y</mark>	515	
RPB2_Sc_6324725	824	IVAIACYSGYNQE	3	IMNQSSI	130	GD-KFASRHG		QKGTIGVTYRHED	2	L I I NP	63	GKKLMAQV	'F <mark>F</mark> GPTY <mark>Y</mark>	1092	
liw7C_Thth_21730339	673	LVAIMPFDGYNFE	3	VISEE <mark>l</mark> L	145	GD-KLANRHG		NKGVVAKILPVED	2	VI <mark>L</mark> NP	106	GEPIEGPI	V <mark>V</mark> GQMF <mark>I</mark>	994.	/
1cr5A_Sce_6730184	5	TRHLKVSNCPNNS	3	anvaa <mark>v</mark> s	6	NIYI <mark>II</mark> DN		$\texttt{LFVFT}{\textbf{T}}\texttt{RHSNDIP}$	3	IG <mark>F</mark> NG	6	GWSL-NQ-	D <mark>VQ</mark> AKAF	81)	\FDHC-like
lqcsA_Crgr_5107651	7	GRSMQAARCPTDE	3	SNCAV <mark>V</mark> S	6	GQHVIVRTS-	3	KYIFTLRTHPSVV	3	v a f sl	6	GLSI-GQ-	E <mark>IEV</mark> AL <mark>Y</mark>	87	
1e32A_Mm_14488635	24	NRLIVDEAINED-		nsvvs <mark>ls</mark>	11	GDTVLLKGK-	3	EAVCIVLSDDTCS	3	I R <mark>M</mark> NR	6	RVRL-GD-	V <mark>I</mark> S <mark>I</mark> QP <mark>C</mark>	105	
1kqfA_Ec_20150976	887	QFPYVGTTYRLTE	17	EQFVE <mark>I</mark> S	11	GDRVT <mark>V</mark> SSK-	2	FIRAVAVVTRRLK	11	V G <mark>I</mark> PI	28	PEYK-AF-	L <mark>V</mark> N <mark>I</mark> EK <mark>A</mark>	1015	
1g8jA_Alfae_12084495	693	KYRFWLNNGRNNE	19	MAYIE <mark>MN</mark>	11	GDIVEVYND-	2	STFAMVYPVAEIK	3	TF <mark>M</mark> LF	21	PYYK-GT-	wgd <mark>i</mark> rk <mark>v</mark>	807	
1dmr_Rhca_2981779	676	KYPLHIAASHPFN	19	hepcl <mark>m</mark> h	11	GDVVR <mark>V</mark> HND-	2	QILTG V KVTDAVM	3	i q i ye	36	NCGQ-TV-	L <mark>a</mark> evek <mark>y</mark>	806	
2napA_Dede_6137474	608	EYPLYLTSMRVID	19	IAFVE <mark>IN</mark>	11	GDSVI <mark>V</mark> ETR-	2	AMELPARVSDVCR	3	i a <mark>v</mark> pf	21	PEYK-IC-	A <mark>A</mark> RVRKA	723	
law8_Ec_400729	10	LHRVKVTHADLHY		EGSCA <mark>I</mark> D	11	NEAID <mark>I</mark> WNVT	3	RFSTY A IAAERGS	3	I S <mark>V</mark> NG	5	CASV-GD-	I <mark>V</mark> I I ASF	90	
lcz4A_Thac_6435754	6	GIILRVAEANSTD	2	GMSRVRL	12	GDVVEIEK	2	KTVGRVYRARPE-	5	i vrid	7	GASI-GD-	K <mark>v</mark> kvrkv	90.	/
2eng_Huin_1942898	1	ADGRSTRYWDCC-	57	LGFAAT <mark>S</mark>	10	CACY <mark>EL</mark> TFTS	6	$\texttt{KMVVQ}{\textbf{S}}\texttt{TSTGGDL}$	4	FD <mark>L</mark> NI	49	FKNAD N P-	S <mark>F</mark> SFRQ <mark>V</mark>	187	\Barwin-like
1bw4_Hovu_442694	6	VRATYHYYRPAQN	28	lgwta <mark>f</mark> C	11	GKCLRVTNPA	3	QITARIVDQCA	3	LDLDW	12	GYQQGHL-	N <mark>V</mark> N <mark>Y</mark> QF <mark>V</mark>	121	/
consensus/80%				hs		spblpp		s		h.h		sb.sp.	.h.hh		
				*		** ** *		*				* * **	* *		

A structure-based multiple alignment of the DPBB domains from the β and β ' subunits of DDRPs with a selection of other structurally characterized DPBB domains and the predicted DPBB domains of RDRP. The alignments were generated by structural superposition of representatives of the DPBB domains followed by addition of sequence neighbors. The predicted DPBB domain of the RDRPs was included on the basis of the alignment with the DDRP β ' subunit's DPBB domain. The functionally important lysines in the β subunits and the metal-coordinating aspartates in the β ' subunits are boxed. The conserved positions shared by the RDRPs with the structurally characterized DPBBs are indicated below the alignment by asterisks. The other conventions and abbreviations are as in the legends to Fig. 1 and Fig. 2.

symmetrical structural units, each with three core β strands. These two units are combined into a barrel with a complex topology, with strand 5 wedged between strands 1 and 2 and strand 2 wedged between strands 4 and 5, yielding two 'psi' loop structures (Fig. 4). The psi-loops between strands 1 and 2 and between strands 4 and 5 often harbor the active site or substrate-interaction residues of the respective DPBBs. Sequence comparisons and identification of specific conserved motifs showed that the DPBB fold includes five major superfamilies. These superfamilies are typified by the formate dehydrogenase C-terminal (FDHC) domain, Barwin endoglucanase, aspartyl protease, and the DPBB domains from the β and β' subunits of DDRP. Two of these superfamilies show a restricted phyletic distribution. Barwin-like endoglucanases are present only in plants and fungi, whereas aspartyl proteases are found in eukaryotes, retroviruses and a minority of bacteria. The latter superfamily also has a circular permu-

tation, which suggests secondary derivation. Thus, none of these superfamilies are likely to have been present in the last universal common ancestor (LUCA) of extant life forms. In contrast, the DPBBs detected in DDRP subunits are universally present in all modern life forms and hence are traceable to LUCA. The FDHC superfamily includes several distinct families. The N-terminal domain of the CDC48-like AAA ATPases is present largely in eukaryotes and archaea and the N-terminal domain of thymidine phosphorylases is archaea-specific. In contrast, the FDHClike DPBB domain associated with oxidoreductases, such as formate dehydrogenase and molybdopterin-containing dehydrogenases, is widespread in archaea and all major lineages of free-living bacteria. Thus, at least one DPBB domain of the FHDC superfamily probably was present in LUCA.



Structure of the catalytic cleft of DDRP formed by interacting DPBB domains of the β and β' subunits. The metal-coordinating DbDGD motif of the β' subunit and the functionally important lysines projecting into the catalytic cleft of the β subunit are shown in ball and stick representation. The two double psi-barrels are juxtaposed in an asymmetric head to tail configuration.

The above analysis shows that LUCA probably encoded three distinct DPBB domains. Although the DPBB domain in the RNA polymerase β subunit is distorted by inserts and shows extreme sequence divergence, its role in the formation of the catalytic cleft and the spatial arrangement with respect to the more canonical DPBB of the β' subunit (Fig. 4) are compatible with the hypothesis that the two DPBB domains of DDRP shared a common ancestor to the exclusion of the other ancient, FDHC-like DPBB. A structure-based sequence alignment shows that the DPBB domains of the RNA polymerases share several conserved residues with other DPBB domains. These include 7 hydrophobic residues, two sets of symmetrically positioned small (mostly glycine) and polar (mostly aspartate) residues preceding strands 3 and 6, respectively, and at least three other small residues, two polar residues and two bulky residues (Fig. 3). Of these, the small residues preceding strands 3 and 6 contribute a positive ϕ angle, which results in turning of the C- α backbone to stabilize the β -barrel [62]. These shared features notwithstanding, the ancient DPBBs have no conserved residues in their catalytic or substrate-binding sites. This is best compatible with the notion that the common ancestor of the DPBB domains was a generic binding domain devoid of high specificity or catalytic activity (see also discussion above).

The same three-stranded structural units that interlock to give rise to DPBB also form another type of β -barrel via terminal dimerization. This is the <u>E</u>longation factor-<u>I</u>somerase (EI)-barrel found in ribosomal proteins, such as L3, GTPase translation factors, ferredoxin reductase, and L-fucose isomerase [62,66]. It seems likely that the EI-barrel and the DPBB evolved from ancient three-stranded ancestral units, which were stabilized via the formation of different barrel structures through homo-dimerization.

Several versions of the DPBB, e.g., those in RNA polymerases, and of the EIB, such as those in translation elongation factors and ribosomal protein L3, are parts of ribonucleoprotein complexes. Other ancient forms, especially those that occur as accessory domains in enzymes, appear to bind small molecules. Hence, the primordial three-strand unit might have formed barrel-shaped dimers that non-specifically associated with RNA or with small molecules. These generic barrel domains subsequently diverged to occupy different functional niches in ribonucleoproteins and metabolic enzymes. These considerations, together with the inferred relationships between the different versions of the DPBB domain, lead to the scenario of early evolution outlined in Figure 5. Only the last bifurcation of the DPBB domains in the proposed tree yielded the distinct forms present in the β and β' subunits of DDRP, which apparently acquired catalytic activity. Substantial divergence of the β -barrels seems to have occurred at a stage of evolution when the actual RNA polymerase activity resided in a ribozyme whose functioning was facilitated by a non-specific, RNA-binding protein cofactor.

Ancient conserved domains and evolution of DNA-dependent RNA polymerases

While all bacteria, archaea and eukaryotes encode orthologs of at least four distinct subunits of DDRP, the divergent RNA polymerases of baculoviruses and yeast killer plasmids and the Corynebacterium Cgl1702 protein only contain counterparts to the β and β' subunits. The minimal RNA polymerase seems to be represented by the predicted catalytic core of Cgl1702, which appears to consist entirely of the β -type and β '-type DPBBs, in this case juxtaposed in the same polypeptide. These observations are not particularly surprising given that β and β' subunits form the catalytic cleft of the DDRP (Fig. 3), whereas the other subunits occupy peripheral positions in the complex. In an attempt to gain further insight into the early evolution of the catalytic core of RNA polymerases, we sought to identify additional globular domains in β and β' subunits.

Using the experimentally determined structure as a guide, we split the β and β' subunits of the *Thermus thermophilus* DDRP and the largest two subunits of the yeast DDRP into individual globular domains and investigated them by visual inspection of the topology, structural searches of the PDB database using the DALI program and iterative searches of the NR database using PSI-BLAST. An obvious ancient conserved domain thus detected was a rubredoxin-like Zn-ribbon. The Zn-ribbons are widespread, small domains that are comprised of two β-hairpins bounded by consecutive extended regions stabilized by metal-coordinating residues in the hairpin loops [67-70]. Zn-ribbons are present in a variety of nucleic-acid-binding proteins, including several ribosomal proteins, translation factors, aminoacyl-tRNA synthetases, RNA polymerase cofactors in archaea and eukaryotes, and several transcription factors. These domains are also present in other contexts where they function as structural scaffolds and as regulators of redox reactions. The β ' subunit of bacterial DDRPs contains a single Zn-ribbon at the N-terminus (region 56–81 of *Thermus* β ' subunit; Fig. 6). The orthologous largest subunits of the archaeal and eukaryotic DDRPs have two Zn-ribbons in the N-terminal portion of the protein (regions 59-88 and 103-173 in the yeast protein). The distal ribbon is distorted by a large insert between the two core halves of the domain and a substitution of asparagine for one of the metal-coordinating cysteines.

Although the sequences of the Zn-ribbons of bacterial and archaeo-eukaryotic DDRP subunits have diverged



A hypothetical scheme of evolution of two types of 6-stranded β -barrels from 3-stranded units. The scheme was derived as the most parsimonious explanation for the phyletic patterns and structural peculiarities of each lineage of 6-stranded barrels. The emergence of particular properties or characters characteristic of a given clade is indicated by horizontal bars.

considerably, their similar location in the orthologous polypeptides from different kingdoms suggests that β' subunit of LUCA already contained a Zn-ribbon. Based on the crystal structure, it has been proposed that the region containing the Zn ribbon may interact with the promoter during transcription initiation in bacteria [23]. Hence, it is plausible that a nucleic-acid-binding Zn-ribbon had been recruited to the ancestral polymerase core, which consisted of the DPBB domains, and contributed to non-specific interactions with the template. Consistent with this, the Zn-ribbon had been subsequently reused in several basal

archaeo-eukaryotic transcription factors, such as TFIIB, TFIIE and TFIIS [67,68,71–73].

A potential additional nucleic-acid-binding moiety that is conserved in all β '-subunits is the AT-hook like module (Fig. 6). The AT-hook is a simple, small protein module that consists of a flap-like structure with a positively charged surface that is inserted into the minor groove of DNA [74]. The two AT-hook-like modules (residues 583– 603 of 1IW7 chain D) in the DDRP β ' subunit are inserted within the α -helix that is located immediately upstream of the DPBB domain (Fig. 6). Previously, AT-hooks have



Domain architectures of the β **and** β ' **subunits of DDRP.** Domain designations: DPBB, double-psi β -barrel, SBHM, sandwich-barrel hybrid motif, ZnR, Zn-ribbon, ATL, AT-hook like, BBM, β , β '-(specific) module. β G is a domain containing a minimal version of the β grasp fold. Other designations: A, archaea, B, bacteria, E, eukaryota, Pr, Proteobacteria, Aqae, Aquifex, Spi, spirochetes, Chl, Chlamydia, Tth, *Thermus thermophilus*, Dra, *Deinococcus radiodurans*, Tma, *Thermotoga maritima*; Af, *Archaeoglobus fulgidus*, Hsp, *Halobacterium* sp., Mj, *Methanocaldococcus jannaschii*, Mth, *Methanothermobacter thermoautotrophicus*. Other globular regions that are conserved between the cellular DDRPs are shown by blue rectangles and non-conserved regions are shown as gray lines. Globular regions conserved in the archaeo-eukaryotic lineage are shown by red rectangles, whereas those conserved in all bacteria are shown by yellow rectangles. The proteins are not shown to scale. Splits indicated by arrows and slashes are instances when different portions of the respective subunits are encoded in separate genes. been detected primarily in eukaryotic chromatin-associated proteins, particularly transcription factors, where they function as accessory DNA-binding domains [74]. The ubiquity of the AT-hook-like modules in the β ' subunits of DDRPs suggests that they were recruited for nucleic acid recognition in LUCA if not earlier.

Analysis of a β -strand-rich insert between strands 2 and 3 of the β subunit DPBB (Figs. 4, 6) led to the identification of another ancient conserved domain that could be traced back to ancestral RNA polymerases. Visual examination of the structure of this insert domain revealed a topology with elements of both a sandwich and a barrel, and two "waist"-like structures that are characteristic of the sandwich-barrel hybrid motif (SBHM) fold [75,76] (see also the SCOP database [65]). The SBHM fold is present in biotin/lipoate carrier domains and their homologs found in a variety of enzymes and transporters [76]. A DALI search with this insert domain from the β subunit of *Thermus* DDRP showed similarity to the SBHMs from various biotin/lipoate-binding enzymes, such as acetyl-CoA carboxylase (1bdo) or dihydrolipoamide acetyltransferase (1iyu) with Z scores of \sim 6 and RMSD \sim 2.5 Å in the aligned regions spanning approximately 70 residues. An alignment of the SBHM domains from the DDRP β subunit with the previously identified SBHMs showed conservation of most residues characteristic of the SBHM superfamily (Fig. 7). The defining structural feature of this domain is seen in the loop between strands 5 and 6 (Figs. 7,8). This loop is bounded by small residues at each end (mostly valine at the N-terminus and glycine at the C-terminus) and has a distinct signature, with small (mostly glycine), charged (mostly aspartate) and hydrophobic residues occurring in succession, in the middle of the loop (Figs. 7,8). These observations reveal an evolutionary relationship between the SBHM domain of DDRP and the SB-HMs found in other, functionally distinct proteins.

The core SBHM domain consists of a repeat of two threestranded units with the characteristic "waist"-like structure occurring between the second and third strands of each repeat (Fig. 8). In the complete SBHM, strand 1 packs with strand 6 and the loops bounded by the two "waist"like regions adopt an extended conformation resulting in the formation of a barrel. In the crystal structure of DDRP, the SBHM domain inserted into the β -subunit DPBB is in contact with the outward-projecting insert between strands 2 and 3 of the β '-subunit DPBB. Furthermore, the SBHM also contributes to the stabilization of the catalytic site by forming an interface between the two DPBBs (Fig. 4). The universal presence of the SBHM inserted in the DPBB domain of the DDRP β -subunit suggests that this domain was already present in the common ancestor of all extant DDRPs. The wide spread in different phylogenetic lineages of two other SBHM domains, namely those found in dihydrolipoamide acetyltransferase and the glycine-cleaving enzyme, is indicative of their presence in LUCA [76]. The SBHM of the DDRPs is specifically related to the biotin/lipoate-binding SBHMs. The two share a conserved basic residue that is present immediately prior to strand 4. In all biotin/lipoate-binding SBHMs, this position is occupied by a lysine, which covalently binds organic radical ligands (Fig. 7), but there is no evidence for such a role in the DDRP β subunits. These observations suggest that the ancestral SBHM interacted with various ligands with a low specificity. From such a precursor, which existed prior to LUCA, the SBHM domain apparently diversified into two forms, one that covalently bound organic radicals, and another that specialized in non-covalent interactions with proteins or nucleic acids. This latter form was recruited into the RNA polymerase catalytic core where it provided additional surfaces for the interaction between the DPBB domains and formed part of the interface between the catalytic domains. Thus, it appears likely that the ancestral DDRP evolved from the coalescence of at least 3 distinct domains: i) the DPBB that probably originally bound RNA and, subsequently, segregated into the cores of the β ' subunit, where it acquired the metal-coordinating active site, and of the β subunit, with the two lysines projecting into the catalytic cleft (Fig. 4), ii) the Zn-ribbon, which was probably involved in interactions with the template, and iii) the SBHM domain that stabilized the interactions between the DPBB domains forming the catalytic cleft (Fig. 6). Additionally, two AThook like modules, which probably had an accessory DNA-binding function, were also inserted into a conserved α -helix directly N-terminal of the DPBB domain in the β' subunit.

The remaining conserved regions in the β and β' subunits of the DDRPs do not show detectable relationship with any other ancient conserved domains. Examination of these structures shows that they are principally composed of large α -helical hairpins that form coiled coil structures or stretches with successive shorter α -helices. Some of the other conserved regions consist of β -meanders with unusually long β -strands. These regions probably emerged, respectively, through duplication and divergence of a simple α -helical unit or through stabilization of long loops by selection for hydrophobic residues, resulting in the formation of the core units of the β -meanders.

Proliferation of the Sandwich-Barrel Hybrid Motif and lineage-specific innovations of the large subunits of bacterial DDRP

Further visual examination and structural alignments revealed the presence of one additional SBHM domain in the β subunit (1iw7 chain C: region 592–659) and several repeats of this domain in the β ' subunit (1iw7 chain D, regions: 163–195, 248–308, 311–368, 369–419 and 1270–

	S1 S2	S3	S4	S5	S	6	
Sec. Structure	EEEELLLLLLLEEEEE	EEEEEE.	EEEEEE	EEE.LL	LLLELLLE	EEEE	
lci3 PhLa 11386726	214 NAVYNASAAGV <mark>I</mark> TA	IAKADDGSAEVKIRTE	DGTTIVDKIP	AGPELI <mark>V</mark> SE	GEEWAAGA	A <mark>L</mark> TNNP 276	Cytochrome F like SBHM
1hcz Brra 1942179	168 NTVYNATAGGIISK	TLRKEKGGYETTTVDA	SNEROVIDITP	RGLELL <mark>V</mark> SE	GESTKLDC	PLTSNP 237	1
CutE Cap 1190E1		TAMARAACTE	AC PUIDTID	ACDEUTICE		ATTINING 270	
Cycr_539_110051	205 MAGRADA IGI <mark>1</mark> IN	INVINIARGIDITISTE	AG 1011F	AGFLVI	GVA LAAG	ADIMME 270	'!
1gpr_Bs_442961	49 EGIVVSPVRGK <mark>I</mark> LN	VFPTKHAIGLQSD-	GGRE <mark>I</mark> LIHFGIDTVS	3LKGEGFTSF <mark>V</mark> SE	GDRQEPGQ	K <mark>l</mark> levd 116	5
BglF_Ec_131486	513 VGEVRSPVAGR <mark>I</mark> AS	LFATLHAIGIESD-	DGVE <mark>I</mark> LIHVGIDTVF	(LDGKFF <mark>S</mark> AH <mark>V</mark> NV	GDKVNTGE	R <mark>L</mark> ISFD 580	
CRR EC 131514	57 GNKMVAPVDGTIGK	TEETNHAESIESD-	SGVELEVHEGIDTVE	ELK GEGEKRTAEE	GOR VKVG	TVTEFD 124	LÍ
N1-D D- 460715				NID THE WORLD			
N1pD_EC_462715	289 GQAIIAIADGRVVY	AGNALRGIGNLIIIKHN	DDILSAIAH	NDTMLVRE	QQEVRAGQ	AIMG 350	1
LppB_Haeso_1170823	255 GQAVNAAAAGR <mark>V</mark> VY	AGDALRGYGNLIIIKHN	DSYLSAYAH	NESIL <mark>V</mark> KD	QQEQKAGQ	Q <mark>I</mark> AKMG 316	o/
liw7D Thth 21730340	163		YG-KOETYPL-PP	GVDAL <mark>V</mark> KD	GEEVVKGC	ELAPGV 195	RpoC SBHM
liw7D Thth 21730340	248 PYLFRAEEEGVVEL	KELEEGA F LVLRRE	DE-PVATYFL-PV	GMTPL <mark>V</mark> VH	GEIVEKGC	PLAEAK 30F	3 Í -
liw7D Thth 21730340	311 LEMPROVEAAOVEA	EERGETVYLTLELE	WT-EPKDYRV-OP		GARVEAG	KTV 368	d .
liw7D Thth 21730340	360 AATDDEEE WINEA	ECULUI MEDACTI	INT DELEG	IVDEED	DUE VOTO D	DUADOD 410	
11W/D_111C11_21/30340	309 AAIDPEREVIAEA	EGVVHLHEPASIL	VVKAR	VIPFED	DVEVSIGL	RVAPGD 415	
11W/D_Thth_21/30340	1270 AKAVISEIDGVVRI	EETEEKLSVFVES	EGFSKEYKLPK	EARLL <mark>V</mark> KD	GDYVEAGQ	PLITRGA 1329	·
RpoC_Dr_6458629	1287 TQAVVADRDGVIRI	EEEEERYLVRIEA	2 EQYSSKT <mark>A</mark> TKVPR	VLRMT <mark>V</mark> KD	GERQEAGQ	PITRGA 1350	1
RpoC_Ec_15804578	1152 EPAILAEISGIVSF	GKETKGKRRLVITP	D-GSDPYEEMIPK	WRQLN <mark>V</mark> FE	GERE	VISDGP 1214	ŧ į
RpoC_Mlo_13470543	1145 DHAIIAEIDGT <mark>I</mark> RF	GRDYKNKRRIIIEP	DSTLEPVEYLIPK	GKPFH <mark>L</mark> QD	GDVEEGE	Y <mark>I</mark> LDGN 1208	3
RpoC Hp 15645812	2663 DVAILSEVDGI <mark>V</mark> SF	GKPIRNKEHIIVTS	L D-GRSMD <mark>Y</mark> FVDK	GKQIL <mark>V</mark> HA	DEFE	AMTDGV 2724	L
RpoC Ci 11261227	1295 NAAVIAEIDGV <mark>V</mark> RF	DKPLRSKERIIIOA	L D-GTSAE <mark>Y</mark> LIDK	SKHIO <mark>V</mark> RD	GEFE	K <mark>L</mark> TDGV 1356	i i
Rpod Ct 15605035	1155 DAADTAKTDGVVDF	KGTOKNKRTLWVRD	T-GMEERHUTSU	TKHLT	GDSVIKGC	OLTDGL 1215	1
Rpod Tp 15639234	1157 NAAVLAOTSQUVSE	KGLEKCKPIWAVPD	-CKEVKULUSM	SPOLL VPD	GDTVEAGE	PLCDCC 1211	
Brod Tm 15643335	1397 CENTIONU DOPUND	TANDESCRIPTINE	A CNILLA VUDY	DAVUDUEV	COX NIDC E	TTTCCA 1440	
Rpoc_111_15043225	1387 SEATLCEVDGFVRD	IAIDESGRIVIIIE	A-GNIHAIRVPR	KARVRVER	GQR	1013GA 1445	
RpoC_Mpn_13508254	1063 ERCVISEVKGVVKS	ITTTQNAQEVLIES	S-VDERTYSIPF	SAQLR <mark>V</mark> KV	GDAVELGS	KITEGS 1123	
RpoC_Bs_16077176	967 GQATITEIDGT <mark>V</mark> VE	INEVRDKQQEIVVQ	L A-VETRS <mark>Y</mark> TAPY	NSRLK <mark>V</mark> AE	GDKQ	VLTGGS 1028	1
RpoC_Mtu_15607808	1042 GKAPIADVTGR <mark>V</mark> RL	EDGERFYKITIVPD	LGGEE <mark>V</mark> VYDKI	SKRQRLR <mark>V</mark> FK 1	1 GDHQEVGQ	Q <mark>L</mark> MEGS 1115	5
RpoD_Pmar_23123296	1029 DSSILCKKSGV <mark>V</mark> QI	KEGTDEESVSLSV	B D-DSISE <mark>Y</mark> QLLM	GQNIM <mark>V</mark> SD	GQQE	LLTDGP 1093	
RpoD Ssp 16329956	997 EACVLARAPGVCQV	EYLEDESVDIKV	B D-GTVSE <mark>Y</mark> PLLP	GONAM <mark>V</mark> TD	GORH	A <mark>L</mark> TDGY 1058	3
NusG Tm 128916	140 NEEYICELDGKIVE	TERMKKVVVOTP-	D-GEODVYYTP	UD V FD	RDRK	EVKOGE 194	LÍ
rpoC Ec 15804578	956 GSTKLSNVKSVVNS	SGKLVITSENTELKLIDE	G-RTKES	GAVLAKGD	GEO	TVANWD 1027	
rpoC Ec 15804578	1058 SIMUDEAFRTACC		T-DMDAOVELDC		GVOTSSGT	TTADID 1125	
1000_10_1004970	1000 SHVIDDA EKINGG	KOLKFALKIVDAQ (T DHEAQUE DEG			TORRET 1125	
rpoc_Aqae_15606950	1284 NPAILSEIDGIVKI	IEDADEVIIFNPR	I-GEIANISIKK	DELILVRH	GQFVKKGQ	ALIEIK 1343	
rpoc_Aqae_15606950	1340 TETKVAEIDGQ <mark>V</mark> RI	KGRGFK V IVYNPE	T-GLQRE <mark>Y</mark> FVPK	GKFLL <mark>V</mark> KE	GDFE	Q <mark>L</mark> TDGT 1399	97
liw7C_Thth_21730339	592 LAALYAEEDGE <mark>V</mark> VK	VDG-TRIAVRYE-	-DGRRLVEH-PLRRYARS	SNQGTAFDQRPR <mark>V</mark> RV	GQRE	LLADGP 659	NRpoB SBHM
RpoB Atu 17935846	723 GAAIAARRGGV <mark>V</mark> DQ	VDA-TRIVIRATE	5 KS-GGVDIY-RLOKFORS	SNONTCVNORPL <mark>V</mark> SV	GDAISKGE	IIADGP 796	5
RpoB Bs 16077175	671 GAAVICKHPGIVER	VEA-KNVWVRRYE	7 KG-NNLDKY-SLLKFVRS	SNOGTCYNORPIVSV	GDEE	ILADGP 746	i i
RpoB_Mtu 15607807	633 GDVVVARESGVTEE	VSA-DVTTVMHD-	NG-TTRRTY-RMRKEARS	NHGTCANOCPT	GDRVEAGC	VTADGP 700	ni
Doop Eq. 15934164	713 CUTAVAND COVIDE	VDA CDTUTKUNE	7 ACTOTY NUTRYTE	NONTGINOMDOUGI	GED VEDG D	VINDOR 700	
RDOB_EC_15054104	694 CULUKAKO DOTUAN	VCC FRINCCAARC	CE FORTUN DI KYOP	NODTOYUODDINUU	GDP WOVG F	ALADGE 787	
RDOB_1D_15039233	664 GVLVKAKQDGIVAI	VBS-ERIVVCSAAAS	GE-EQEV <mark>V</mark> I-PLBRIQR		GDR	ADADGP 755	-
RpoB_BD_15594/34	665 GVVVKAKRSGEVIL	ATS-SKIVVKPFE	3AKDLDEY-HIVKYER:	INQDICFNQSVL <mark>V</mark> KE	GQKVERGE	ILADGP 736	
RpoB_Aqae_15606949	794 HAVVVAKRGGV <mark>V</mark> EE	VDS-SKTIIRVNP 10)IGID <mark>I</mark> Y-ELRKFQR1	INQKTCVNQRPI <mark>V</mark> RK	GEKQ	I <mark>I</mark> ADGH 871	- [
RpoB_Hp_15645812	727 WGAIKANRAGV <mark>V</mark> EK	IDS-KNIYILGES 4	1YID <mark>A</mark> Y-SLQKNLR7	INQNTSFNQVPI <mark>V</mark> KV	GDKQGAGQ	I <mark>I</mark> ADGP 797	7
RpoB_Cj_15791842	728 WEAVKAKRGGV <mark>V</mark> EK	VDN-KSIFILGED 4	I -FIDHY-TMEKNLR1	INQNTNYIQHPI <mark>V</mark> KK	GDIQKAGQ	I <mark>I</mark> ADGP 798	3
RpoB Nm 11261156	740 ATAIVARRGGV <mark>V</mark> EY	VDA-NRVVVRVHD	7VGVD <mark>I</mark> Y-NLVKFTRS	SNOSTNINORPA <mark>V</mark> KA	GDVEQRGE	LVADGA 814	L
RpoB Mlo 13470542	723 GAAIGARRGGI <mark>V</mark> DO	VDA-TRIVIRATE 6	5SGVDIY-RLMKFORS	SNONTCINORPL <mark>V</mark> RM	GDRVNKG	I <mark>I</mark> ADGP 796	i i
RpoB Car 16124757	720 GAVVIAKRTGVVEO	TDG-TRTVIRATE (SGVDTY-RMSKEORS	NOSTCINOPPL	GDRTVAGT	TADGP 797	d .
BroB Ct 15605036	659 CATTWACE DOWNEY	UDC VETUINKEN	I KDDV OI KKEID	NECTOINOTDICEU		VINDOR 725	
RpoB_CC_15005050	538 GALLVAVEDGVVEI	VDS-TELVVARRIN		NOGICINQIPLESV	GDVVINGL	VLADGE 121	
RDOB_350_10329937	SOF GROUNDER	VDA-TELRVQPH3 (5KGEIVI-PIQKIQK2	SNODICENQRPEVIA	GED	VLADGS 043	
Rbog_pt_12802631	642 GISVVSDVNGRVSY	VDA-RALQVTLSE 8	SAGVRTF-ELIRFTRS	SNQGTNLDQHPI <mark>V</mark> SV	GDEQRVGQ	VIADGP /18	5
RpoB_Mge_12045200	812 GLTMSSPCSGV <mark>V</mark> SY	VDN-SKTIITSDS	–KKETV–NLVKFERS	SNQNTCYNHKPI <mark>V</mark> EI	GQRE	IIVDGP 879	97
livu Azvi 1942194	1 SETTRUPDI GODGEVIE	LLVKTGDLTEVEOGEVULES-	A - A SMEVPSPKA	GUVKSVSVK	GDKTKEG	ATTELE 77	NB/LBD-like SBHM
Acor Fg 16129109	2 ALEIKUPDIC ADEVETTE	TI WAYCONNENE COL TIMEC	DE ACMENDODOD	CTUKETKUCU	CDK TOTC A	TMIED 7/	
ACEF_EC_10128108	Z AIEIKVPDIGADEVELIE	ILVRVGDRVEAEQSEIIVEG-	DR-ASME PSPQA	GIVKEIK SV	GDKIQIGA	CINIFD /4	
IdczA_Prir_8569597	8 EGEIPAPLAGTVSK	ILVKEGDTVKAGQTVLVLEA-	MR-METEINAPTD	GKVEKVL <mark>V</mark> KE	RDAQQGGQ	GLIKIG //	
IDdo_EC_182/931	4 GHIVRSPMVGT <mark>F</mark> YR /	AFIEVGQKVNVGDT E CIVEA-	Ma-MMNQ1EADKS	GIVKAIL <mark>V</mark> ES	GODR	5 <mark>7</mark> AATE 80	1
1htp_Pisa_999900	22 VATIGITDHAQDHLGEVV <mark>F</mark> VE	-LPEPGVSVTKGKGFGAVES-	VK-ATSDVNSPIS	GEVIEVNTGL	1 GKPGL-INSSPYEDGW	M <mark>I</mark> KIKP 105	5
GcvH Ec 121078	24 TYTVGITEHAQELLGDMVFVD	-LPEVGATVSAGDDCAVAES-	VK-AASDIYAPVS	GEIVAVNDAL	1 DSPEL-VNSEPYAGGW	IFKIKA 107	7
GovH Bs 12229788	22 KARIGITHFAOSELGDIVEVE	-LPEVGAETKADEPEGSVES-	VK-TVSELYAPIN	GTVVEVNEDI	1 DSPEE-VNESPYEKAW	MIVVEP 10F	si
GCSH Hs 4758424	66 IGTVGISNFAOFALGDVVYCS	-LPEVGTKLNKODEFGALES-	VX-AASELYSPLS	GEVTEINEAL	1 ENPGL-VNKSCYEDGW	LIKMTL 149	
CONTRACTOR 23821627		AKOKCEKUNCDENT EGIEC	WICK PDI DIE	CEVEDUNEEU	1 KNDGI INEBDYD CW	TWEETEN 117	
GCVH5_350_23821027	25 VVSIGVIDLGQIMAGKIPQVI	-ARQRGERVINGRSVEFSIES-	MA-WIGKERLPIE	GEVEDVINEEV	1 RAPSI-INERPID-3W	IVAIRV III	·
PAB0559_Pab_/441699	39 IVLVGITDIAQKELGDIA <mark>I</mark> VE	-LPEVGKEVKKGEV E CEVES-	VAPVS-AVSEVIAPVS	GEVIEVNEEL	I DSPER-INEDPIG-AW	IANINP 121	-/
liw7C_Thth_21730339	708 YEIEARDTKLGPER <mark>I</mark> TR 16	GVVRIGAEVKPGDILVGRTSF 23	3 V <mark>K</mark> DTS <mark>L</mark> RVPPGEG	GIVVRTVRLR	1 GDPGVE <mark>L</mark> KPGVRE	V <mark>V</mark> RVYV 827	/\DRDP-like SBHM
RpoB_Dr_15805937	767 DEIEARDTKLGPEK <mark>I</mark> TR 16	GIVRVGAEVKPGDILVGKTSF 23	3 V <mark>K</mark> DTS <mark>L</mark> RVQSGQG	GIVVKTVRFR	1 GDEGVDLKPGVRE	M <mark>V</mark> RVY V 886	5
RpoB_Ec_15834164	836 LACVSRDTKLGPEE <mark>I</mark> TA 16	GIVYIGAEVTGGDILVGKVTP 2	3 VKDSSLRVPNGVS	GTVIDVQ <mark>V</mark> FT 10	3 GDDLAPGVLK	1 <mark>V</mark> KVYL 1054	4
RpoB Aae 15606949	920 LEVEARETKVGEFETTR 16	GIVRVGTYVKPGDILVGKVTP 2	VKDASLRCPPGVE	GIVIDVOVFT 12	3 RRDLPPGVIT	LVKVFI 1158	ai
RpoB Hp 15612186	846 KEVDARELKHGVEERTA 16	GIVKVGTYVSAGMILVGKTSP 2	VVNKSLYCPPSLE	GTVIDVKVFT 12	1 DDTKPNGVTK	KVKLYT 108:	e i
PpoB Con 16753076	786 FELTADOTKICKEPTTD 16	GIIRIGAEVERGDIWCKITED 2	WE-DASETURD	GUUMDUKWEC 13	A GDADLDHGVI P	OVKUVV 1034	
Rpob_Cpii_10/330/0	700 FELIARDIRUGREELIK 10	GIIRIGAEVAPODIEVORITP 2	DNCT DUDNCTY	CDURDUD FS 12	CDE CDECAN	VICVIV 1020	
RDOD_AD3_1/229086	722 IEIEARQIKLGPEELTR 16	GIIRIGAWVEAGDIEVGKVTP 2	DISTRUCTION	GRVVDVRHFT	3 GDEBPPGANM	vyr.viv 840	
KDOR_R8_100//1/2	/95 YESEARDTKLGPEE <mark>I</mark> TR 16	GIIRIGAEVKDGDLLVGKVTP 2	vRDTSLRVPHGGG	GIIHDVK <mark>V</mark> FN	S GDEQ	PAKANT A13	2
rpoB1_Mj_15669230	247 YDACERRYPGGQMDRF 22	GIVAVESHVKGGDVIVGKTSP 16	RRDSSVVRHGEE	GYIDKVI <mark>L</mark>	TETREGNR	L <mark>V</mark> KVKV 356	
APE1856_Ap_14601675	795 YTGIENEYPGGERDRI 23	GIVAPETEVEGGEILIGRVSP 16	5 R <mark>R</mark> DTS <mark>I</mark> PMRLGEK	GVVDMVV <mark>I</mark>	STTK	L <mark>V</mark> KVRV 905	5
RPB2p_Sc_6324725	859 YMDIEKRQGMKALETF 22	GLIAPGVRVSGEDIIIGKTTP 16	5 K <mark>R</mark> DASTPLRSTES	GIVDQVL <mark>L</mark>	TTNGDGSK	F <mark>V</mark> KVRV 968	87
consensus/80%	ph	pb		p <mark>1</mark>	spls	.ls	

A structure-based multiple alignment of the sandwich-barrel hybrid motif (SBHM) domains from β and β ' subunits of DDRPs and other proteins. The alignments were generated by structural superposition of representatives of the SBHM domains followed by addition of sequences neighbors of the representative structures. The conserved basic residue that forms a covalent linkage with the organic radical in biotin/lipoate-binding domain-type SBHM is shown in reverse shading. The individual sequence families of SBHMs are indicated to the right of the alignment. The waist-like loops are marked as 'L' in the secondary structure shown above the alignment. Species abbreviations are as in Fig. 1 and Fig. 2 Additional species abbreviations not given above are: Ana, Anabaena sp., Azvi, Azotobacter vinelandii, Bb, Borrelia burgdorferi, Brra, Brassica rapa, Ccr,Caulobacter crescentus, Cj, Campylobacter jejuni, Cpn, Chlamydophila pneumoniae, Haeso, Haemophilus somnus, Mge, Mycoplasma genitalium, Mj, Methanocaldococcus jannaschii, Nm, Neisseria meningitidis, Pab, Pyrococcus abyssi, PhLa, Phormidium laminosum, Pisa, Pisum sativum, Pmar, Prochlorococcus marinus, Prfr, Propionibacterium freudenreichii

1329) of the bacterial DDRPs (Fig. 6). A PSI-BLAST search started with the N-terminal SBHM from the *Aquifex aeolicus* DDRP β ' subunit detected several SBHMs from proteins other than the β ' subunit, such as acetylornithine

deacetylase (E = 2×10^{-7} , iteration 2), the C-terminal region of bacterial cytochrome F (E = 10^{-4} , iteration 2), biotin carboxyl carrier domain of biotin transcarboxylases (E = 6×10^{-4} , iteration 2) and the phosphotransferase sys-



An evolutionary scenario for the SBHM domains from the DDRP subunits and other proteins. The scenario for the derivation of various versions of the SHBM fold from the simple, ancestral 3-stranded unit was inferred from phyletic patterns, structural features and the internal duplication. The conserved basic residue present in the biotin/lipoate-binding domain-type SBHMs and the β subunit of the DDRPs is shows as a ball-and-stick model. The emergence of different lineage-specific specializations is indicated to the side of each clade.

tem enzyme II (E = 10^{-3} , iteration 2). This search also retrieved the bacteria-specific C-terminal SBHM domain of the β subunit (e.g. *Thermotoga*, E ~ 10^{-3}) and, interestingly, also a SBHM in the N-terminal region of the NusG protein from *Thermotoga maritima* (E = 2×10^{-7} , iteration 2). Reverse searches with the C-terminal region of cytochrome F (gi: 11467387, residues 150–321) retrieved the bacterial β ' sequences in the second and subsequent iterations with statistically significant E-values. None of these additional SBHMs were detectable in the archaeal or eukaryotic DDRPs, either in sequence searches or in direct comparisons of the crystal structures. Thus there appears to have been a lineage-specific proliferation and dissemination of the SBHM domains in bacteria. Bacterial β and β' subunits show major variability in the distribution of the SBHM domains (Fig. 6). All bacterial β subunits share one additional SBHM, which is located Nterminal of the DPBB domain. Similarly, an extra SBHM, located C-terminal of the DPBB, is conserved in bacterial β ' subunits. In some cyanobacteria, the portion of the β ' subunit containing the SBHM occurs as a separate polypeptide. In addition to the SBHMs that are conserved among all bacterial lineages, a cassette of up to four SB-HMs are present N-terminal of the DPBB domain in the β' subunit of Thermus, Deinococcus and Thermotoga (Fig. 6). Similarly, the β ' subunit of proteobacteria, Aquifex, spirochaetes, cyanobacteria and chlamydiae contains an additional set of SBHMs C-terminal of the DPBB domain, preceding the C-terminal SBHM that is shared by all bacteria (Fig. 6). The bacterial-specific SBHM domains present in the β and β' subunits are typically located on the periphery of the holoenzyme, which makes them accessible for interactions with other proteins. Two of these SBHMs (regions 163–195 and 369–419 in 1iw7 chain D) are involved in the interaction with the σ 70 subunit, which is critical for transcription initiation in bacteria [23]. Thus, it appears likely that, originally, the SBHM domain of the β subunit participated in generic protein-protein interactions that allowed the ancestral DDRP to recruit accessory subunits. Subsequently, in the bacterial lineage, proliferation of SBHMs provided for several other, specific interactions involved in initiation and elongation.

Several of the SBHMs that are specific to the bacterial DDRPs show degradation of the "waist"-like loop of the first internal repeat. This is clearly a derived state because the SBHM originally evolved through the duplication of a single ancestral unit that contained an intact "waist"-like loop between strands 2 and 3. The form of the SBHM with a shortened loop in the first internal repeat is additionally seen in several bacterial-specific families of SBHMs. These include the SBHM of the phosphotransferase system enzyme II (glucose permease or PTS-EII) found in several lineages of free-living bacteria, the one at the C-termini of cytochrome F from cyanobacteria and chloroplasts, and the NlpD-like cell wall peptidases that are present in most bacteria. In addition to the specific structural relationship between some of the SBHMs of the DDRPs and those present in these bacterial proteins, they also show detectable sequence similarity. For example, a PSI-BLAST search initiated with the SBHM from NlpD retrieved the derived SBHMs of the bacterial RNA polymerase subunits (e.g. *Streptococcus* β subunit, E = 10⁻⁴, iteration 2) well before the classical SBHMs (eg. Mesorhizobium dihydrolipoamide acetyltransferase, $E = 10^{-3}$, iteration 4). The β and β ' subunits of the bacterial DDRP are the only proteins that contain both versions of the SBHM (Fig. 7). Hence, it appears likely that the form of the SBHM with the degraded loop in the first internal repeat initially evolved in the bacterial DDRP subunits. Subsequently, these domains appear to have been recruited for various interactions in other bacterial proteins.

Examination of the structure and multiple sequence alignments of bacterial β and β' subunits led to the identification of two additional, distinct domains, which so far are not detectable in any proteins other than the DDRP and were accordingly designated Beta-Beta' Module 1 (BBM1, region 119-165, 1iw7 chain D) and Beta-Beta' Module 2 (BBM2, region 1109-1190, 1iw7 chain D) (Fig. 6). Both these domains share an unusual, common structural core that we term the BBM-core (BBMC) of about 40-50 residues with an RMSD of 2Å over the aligned region. The BBMC consists of N- and C-terminal helices that bound a central region with 4 unusual, extended regions that fold into a curved hairpin structure (Fig. 9A). The BBM1 domain is inserted after the Zn-ribbon in all bacterial B' subunits and, additionally, into the SBHM domain of the β subunit in proteobacteria, Aquifex and chlamydiae (Fig. 6). Notably, this is the same set of bacteria that contain extra SBHM domains inserted in the C-terminal region of the β ' subunit (Fig. 6 and see above). The BBM2 domain shows a similar pattern of insertion: the C-terminal portion of all bacterial β ' subunit contains one copy, Thermotoga has two extra copies distal of the pan-bacterial copy. Proteobacteria, Aquifex, spirochetes and chlamydiae have a BBM2 inserted into the N-terminal region of the β subunit (Fig. 6). The functions of these previously unnoticed modules and the nature of the apparent congruence in the evolution of the two RDRP subunits in a subset of bacteria remain to be elucidated.

The RNA polymerase α subunit-core-related domain and other elaborations of the archaeo-eukaryotic RNA polymerase catalytic subunits

The bacterial DDRPs have two α -subunits that form a homodimer at the "base" of the β - β ' dimer and stabilize the catalytic-binding cleft. In the archaeo-eukaryotic lineage, there is a duplication of the α -subunit, resulting in two paralogous subunits, RBP3 and RBP11. In the eukaryotic DDRP complex, these subunits occupy a spatial position very similar to that of the α -subunit homodimer of the bacterial DDRPs. All these proteins share a common $\alpha + \beta$ core domain, with two α -helices, which form the dimer interface. In the SCOP database, the α -subunit core domains have been classified together with several other domains, including the ligand-binding domain of the bacterial arginine repressor, the dimerization cofactor of the transcription factor HNF1 (DCOH), and a domain specific to the bacterial aspartyl-tRNA synthetases. To investigate this domain further, we conducted structural similarity searches of the PDB database using the DALI program, with the α -subunits of the bacterial and



Structures of other conserved modules of DDRPs. (A) The core BBMC module of the BBM1 and BBM2 domains **(B)** Different versions of the α -subunit-core related (ASCR) domain ASCR domains from the bacterial DDRP α subunit, RBP11 (the eukaryotic ortholog of α), the β ' subunit of the archaeo-eukaryotic DDRPs, topoisomerase II/gyrase globular domain 3, and the eukaryotic translation initiation factor 1 (Sui1) are shown. Inserts or regions of poor X-ray diffraction are shown with dotted lines. The ASCR domains from the archaeo-eukaryotic β ' subunits are most similar to those from the α -subunits.

eukaryotic DDRPs used as the queries. These searches showed that the core domain of the α -subunits were most similar (Z scores 5.4–7, with RMSDs of 2.2–3 Å over the aligned C- α residues) to the arginine repressor ligandbinding domain [77] and several domains that were not previously recognized as being related (Fig. 9B). These newly detected domains included the SUI1 domain, that is found in eIF1 and related proteins [78], the third globular domain of DNA gyrase/Topoisomerase II and, most interestingly, a distinct globular domain in the yeast DDRP β ' subunit (PDB: 1k83; region 1143–1271). Reciprocal DALI searches with these domains readily recovered the α -subunits of DDRP as the best hits. For example, the domain from the yeast β ' subunit hit the α -subunits of bacterial DDRPs with Z-scores of 5–7 and RMSD of 2.8–3 Å. Hereinafter we refer to these domains as the α -subunit core-related (ASCR) domains. The ASCR fold has a β 2- α - β 2- α topology and forms a two-layered structure with a 4stranded β -sheet and characteristic strand order of 1-2-4-3 (Fig. 9B). The ASCR domains of bacterial α -subunits and eukaryotic RBP3 share a common insert domain between helix-1 and strand 3, which apparently has been lost in the RBP11 subunit of the eukaryotic DDRPs.

The ASCR domain in the yeast β' subunit is inserted into another α + β domain. Visual examination of the topology and structural searches of the PDB database using the DALI program showed that this domain it adopts an RNA Recognition Motif (RRM)-like structure with a β - α - β 2- α - β topology forming a 4-stranded β-sheet. These two domains comprise a conserved module that is specifically shared by the archaeo-eukaryotic DDRPs to the exclusion of the bacterial β' subunits (Fig. 6). This two-domain module is inserted precisely in the same region as the Cterminal SBHMs seen in the bacterial lineage (Fig. 6). The closest relatives of the ASCR domain detected in the β ' subunit are those present in the α -subunits (Fig. 9B). This observation suggests that the primary event in the divergence of the archaeo-eukaryotic β' subunit was the insertion of an RRM-like domain into the ancestral core. This was followed by the insertion of the ASCR domain, which probably evolved as a result of a duplication of the RBP3/ 11 subunits, into this RRM-like domain. Both the spatial arrangement of the ASCR domain in the β ' subunits and its extreme sequence divergence with respect to the ASCR domains of the α -subunit suggests that it has acquired a new function. The RRM-like domain and the ASCR domain are positioned in the exterior entrance to the catalytic cleft and could interact with cofactors or with nucleic acid. There is no evidence that the ASCR domain in the DDRP a-subunits binds nucleic acids and studies on SUI1 protein and the arginine repressor ligand-binding domain implicate the ASCR domain in protein-protein interactions and small molecule binding [77,79]. However, most of the ASCR domains are present in proteins that interact with nucleic acids (see above), raising the possibility of a role in non-specific nucleic acid binding.

Mirroring the insertion of the ASCR domain in the β ' subunit, a distinct $\alpha + \beta$ globular domain, N-terminal to the DPBB domain (region 578–632 of 1K83, Chain B), is inserted in the β subunits of all archaeal and eukaryotic DDRPs (Fig. 6). This domain contains a 4-stranded sheet with a $\beta 2-\alpha-\beta 2$ topology, with the strands in a 2-1-4-3 order. Visual examination of the topology and searches of the PDB database using the DALI program showed that this insert domain has the β -grasp fold [80,81]. For example, in the DALI search, the *Staphylococcus* immunoglobulin-binding protein (IGBP; PDB code 2igd) was detected with a *Z* score of 5.2 and a RMSD of 2.7 over ~50 residues. This insert domain in the archaeo-eukaryotic β subunits resembles the version of the β -grasp domain that is present in the bacterial translation initiation factor IF3 and the IGBPs, in contrast to the distinct version found in ubiquitins, ferredoxins and ThiS. Specifically, the β -subunit insert domain shares a 4-stranded core with the former group and, like these proteins, lacks a long insert between the two terminal strands (data not shown). Given that this domain lies in the periphery of the holoenzyme, it is likely to participate in interactions with cofactors, which is compatible with the protein-protein interaction function identified in several proteins with the β -grasp fold [80,81].

Thus, a common set of ancient conserved modules appears to have been repeatedly reused in the evolutionary diversification of the RNA polymerase subunits from their ancestral cores. The lineage-specific domain insertions in bacteria and in archaea-eukaryotes correlate with the fundamental differences between the basal transcription machinery of these two lineages [82,83]. This suggests congruent evolution of the RNA polymerase catalytic subunits and the basal transcription factors [68].

Origin of the RDRPs

The evidence of an evolutionary relationship between the core catalytic domains of the RDRPs and the DDRPs has implications for the origin of the former group of proteins. Given that DDRP is universal, whereas RDRP is eukaryote-specific, a plausible hypothesis might be that RDRP evolved at the onset of eukaryotic evolution through extensive, rapid sequence divergence following a duplication of the β' subunit of the ancestral DDRP. However, despite extensive comparisons using PSSM and careful visual examination, we failed to detect any residual sequence similarity between DDRP and RDRP, outside the core DPBB domain containing the metal-binding active site. In particular, no traces of the Zn-ribbon and the AT-hook-like modules, which are conserved in all β ' subunits (Fig. 6), were detected. All RDRP sequences contain two closely spaced lysines, which are conserved in the YRH proteins (Fig. 1) and might functionally correspond to the conserved lysines of the β -type DPBB. These residues are embedded in a secondary structure context that is enriched in predicted β -strands (Fig. 1); hence, in principle, this region might correspond to the β -type DPBB of the DDRPs. However, the respective region of the RDRP sequences shows no detectable similarity to the to this DPBB domain, suggesting that the presence of two functionally important lysines in both RDRP and DDRP might be convergent. In principle, radical divergence could have eroded all similarity between RDRP and DDRP beyond the DPBB domain. An alternative, perhaps more likely explanation is that, while the core catalytic domains of the two polymerases evolved from the same ancestral DPBB, they subsequently accreted distinct sets of peripheral modules around this core. The bacteriophage YRH proteins show no specific relationship with the RDRPs from any one eukaryotic lineage, arguing against a recent lateral transfer from eukaryotes to the phages. Taken together, all these observations point to a scenario for the origin of the RDRP, under which the RDRPs and the YRHs diverged from the DDRPs at a stage of evolution that succeeded the divergence of the DPBB domains of the β and β ' subunits (Fig. 5) but preceded the emergence of eukaryotes and even LUCA. One class of substrates of the extant eukaryotic RDRPs are small RNAs, such as microR-NAs and stRNAs [84-90], and analogous small RNAs have been identified in bacteria [91-94]. Given that small RNAs are likely to have been abundant in the hypothetical ancient RNA world, RDRPs might have been involved in the replication of such RNAs since an early stage of evolution. Should that be the case, the YRH proteins could be late surviving derivatives of these ancient enzymes, which apparently have been shunned from cellular life forms during the evolution of prokaryotes. Eukaryotes might have acquired the YRH gene from a bacteriophage at an early stage of their evolution and retained the RDRPs thanks to the selective advantage conferred by the new level of gene expression regulation mediated, in part, by these enzymes.

The eukaryotic RDRP is part of a large ensemble of functionally linked proteins, which include the Dicer helicasenuclease, PIWI family proteins, the Hen-2/Corymbosalike RNA methylase, AlkB-related RNA demethylases, and Lin28-like RNA-binding proteins [1,45,46]. Since no closely related homologs of any of these proteins are encoded in the genomes of the phages that encode YRH proteins, the latter are unlikely to function in a biological context similar to that of the eukaryotic RDRPs. Nevertheless, the extended similarity between the YRH and RDRP sequences and the presence of potential nucleic-acidbinding modules, such as a C-terminal Zn-ribbon in the SpβC2 YonO protein, suggest that the YRH proteins also have a RNA polymerase activity. Possible functions of the YRH proteins include a regulatory activity via amplification of antisense RNAs or perhaps a more conventional role as a DNA-dependent RNA polymerase of the respective phages.

General discussion and conclusions

Here, we identified bacteriophage homologs of the eukaryotic RDRP, which is involved in post-transcriptional gene silencing. We also present evidence that RDRP and the ubiquitous β' subunit of DDRP contain a homologous, metal-coordinating, catalytic domain. This domain adopts the DPBB fold, in which the signature DbDGD motif, shared by RDRP and DDRP, is ensconced in the insert between the ultimate and penultimate β -strands. A second DPBB domain is present in the β subunit of the

DDRPs. This version of the DPBB domain lacks the metalcoordinating motif but contributes two lysine residues, which are critical for substrate interactions in the catalytic cleft. The DPBB domain in the β subunit is distorted by large inserts and shares no detectable sequence similarity with the DPBB from the β ' subunit. However, the DPBBs from the two DDRP subunits are spatially juxtaposed in the DDRP crystal, with the catalytic cleft located between them. This arrangement suggests that the primordial RNA polymerase was a head to tail homodimer of a DPBB domain; a duplication of this primordial DPBB domain probably gave rise to the two DPBBs of modern DDRPs. The presence of completely different conserved residues and the asymmetric head to tail arrangement of the monomers in the pair suggest that the ancestral DPBB dimer was not specialized, had no catalytic activity, and merely bound RNA. Originally, these DPBB domains might have functioned as protein cofactors that stabilized a ribozyme RNA polymerase and eventually displaced the ribozyme as they acquired key residues required for protein-based polymerase activity. This evolutionary scenario complements a similar model proposed for the evolution of the other major polymerase class, the palm-domain-containing RNA and DNA polymerases, whose primordial core apparently consisted of a RNA-binding domain of the RRM fold [11].

The β and β' subunits of DDRP additionally contain a Znribbon and a SBHM domain respectively. The former is a widespread nucleic-acid-binding domain, whereas the latter functions in a variety of biochemical contexts as a small-molecule-binding and protein-protein interaction domain. Analysis of the conserved protein superfamilies within these folds and their phyletic distribution patterns suggests that diverse versions of these domains with distinct functions were already present in LUCA. Thus, several duplication-divergence events apparently preceded the emergence of the forms of these domains that are seen in the catalytic core of DDRPs (e.g., Fig. 5). The ancestral versions of these domains would not have the adaptations for specific roles possessed by their descendant forms, such as those in extant DDRP or RDRP, and probably functioned as generic RNA-binding and protein-protein interaction domains, with the specificity conferred by catalytic RNA molecules. Hence, considerable diversity of protein domains appears to have emerged prior to the "crystallization" [95] of a transcription machinery similar to the one that operates in modern cells and probably had been already in place in LUCA [96]. Similar conclusions regarding the early stages of protein domain evolution have been reached previously as a result of evolutionary analysis of proteins involved in translation, such as aminoacyl-tRNA synthetases and GTPases: substantial diversification of protein domains had occurred prior to the

"crystallization" of elaborate, modern-type translation machinery [66,97].

Although the major events in the evolution of polymerases occurred at very early stages of evolution and LUCA already had an advanced DDRP resembling the extant forms, subsequent lineage-specific elaborations of considerable magnitude have occurred. The most notable of these is the proliferation and dissemination of SBHM and the BBM domains in the bacterial β and β ' subunits. In the archaeo-eukaryotic lineage, another domain, apparently derived through duplication of the dimerization domain of the DDRP α -subunit, was inserted into the C-terminal region of the β ' subunit. Additionally, the Zn-ribbon underwent duplication and elaboration in the β ' subunit and was also recruited in various transcription factors. Similarly in the archaeo-eukaryotic β subunit, a minimal β -grasp domain was added to the conserved core.

While we provide evidence that the catalytic cores of the RDRPs, their bacteriophage homologs and the DDRPs evolved from a common ancestor, the provenance of the rest of the conserved RDRP module and, accordingly, the evolutionary scenario for RDRP remain less clear. No similarity to DDRPs was detected in the RDRP sequences beyond the core DPBB domain. One possibility is that the RDRP module has emerged at the onset of the evolution of eukaryotes through extreme sequence divergence following a duplication of the DDRP β ' subunit. An alternative scenario holds that the RDRPs are ancient enzymes that diverged from the evolutionary precursor of DDRP at a very early, pre-LUCA stage of evolution, albeit after the duplication that led to the differentiation of the β and β' versions of the DPBB domain. In fact, it appears likely that the ancestral protein RNA polymerase, at the time of the differentiation of the β -type and β '-type DPBB domains, functioned as an RDRP because DNA probably had not evolved by that stage [96]. The switch of the ancestral enzyme to the DDRP activity and the emergence of the evolutionary precursor of RDRP might be tentatively linked to one another and associated with the advent of DNA. This view of the evolutionary history of RDRPs is compatible with the presence of RDRP homologs in phages and with their role in replicating microRNAs (potential relics of the RNA world) during PTGS in eukaryotes. Subsequently, this enzyme apparently has been expunged from the cellular RNA synthesis systems and survived only in some parasitic elements, such as bacteriophages, through which it might have been reintroduced into the genome of an ancestral eukaryote. This scenario shows parallels to the probable evolutionary history of another major, unrelated class of polymerases, the RDRPs and reverse transcriptases containing the palm domain. Among extant biological entities, these polymerases are encoded largely by RNA viruses and retroid viruses, but a reverse transcriptase had been recruited by eukaryotes as the catalytic subunit of the telomerase [98,99]. There seems to be a distinct possibility that both classes of RDRPs are vestiges of the ancient RNA world that have been largely displaced from cellular biochemical machinery at an early stage of evolution, because of the deleterious effects of RNA replication in DNA-based organisms, while surviving in selfish genetic elements.

Material and Methods

The non-redundant (NR) database of protein sequences (National Center for Biotechnology Information, NIH, Bethesda) was searched using the BLASTP program [100]. Position-specific scoring matrix (PSSM) searches were conducted using the PSI-BLAST program, typically with a PSSM inclusion expectation (E) value threshold of 0.01, and were iterated until convergence [100,101]. Prior to PSI-BLAST searches, query sequences were evaluated for compositional bias using the SEG program [102]. If no such bias was detected, searches were run with the composition-based statistics turned off, in order to maximize sensitivity [103]. Multiple alignments of protein sequence were constructed using the T_Coffee program [104], followed by manual correction based on the PSI-BLAST results. Identification and statistical evaluation of conserved motifs in multiple protein sequences were performed using the Gibbs sampling method as implemented in the MACAW program [105,106]. Pattern searches were conducted using the GREF program of the SEALS package [107], and pattern-initiated BLAST searches were carried out using the PHI-BLAST program [60]?. Protein structure databases were searched for similar structures using the DALI program [108]. Protein secondary structure was predicted using the PHD program implemented on the PredictProtein server with a multiple alignment submitted as the [109,110]. Protein structures were visualized and manipulated using the Swiss-PDB viewer program [111] and the ribbon diagrams were made using MOLSCRIPT [112].

Authors' contributions

All three authors (LMI, EVK, LA) contributed to making the discoveries reported here. Author 1 and 3 (LMI, LA) prepared all the figures and the manuscript. All authors read and approved the final manuscript.

References

- Anantharaman V, Koonin EV and Aravind L Comparative genomics and evolution of proteins involved in RNA metabolism. Nucleic Acids Res 2002, 30:1427-1464
- 2. Nelson DL and Cox MM InLehninger Principles of Biochemistry. Worth Publishers Inc 2000,
- 3. Goodman MF and Tippin B **The expanding polymerase universe.** *Nat Rev Mol Cell Biol* 2000, **1**:101-109
- Burgers PM, Koonin EV, Bruford E, Blanco L, Burtis KC, Christman MF, Copeland WC, Friedberg EC, Hanaoka F and Hinkle DC Eukaryotic DNA polymerases: proposal for a revised nomenclature. *J Biol Chem* 2001, 276:43487-43490

- 5. Cheetham GM and Steitz TA Insights into transcription: structure and function of single-subunit DNA-dependent RNA polymerases. *Curr Opin Struct Biol* 2000, 10:117-123
- Steitz TA A mechanism for all polymerases. Nature 1998, 391:231-232
- Cramer P Multisubunit RNA polymerases. Curr Opin Struct Biol 2002, 12:89-97
- Kamer G and Argos P Primary structural comparison of RNAdependent polymerases from plant, animal and bacterial viruses. Nucleic Acids Res 1984, 12:7269-7282
- Delarue M, Poch O, Tordo N, Moras D and Argos P An attempt to unify the structure of polymerases. Protein Eng 1990, 3:461-467
- Poch O, Sauvaget I, Delarue M and Tordo N Identification of four conserved motifs among the RNA-dependent polymerase encoding elements. *Embo J* 1989, 8:3867-3874
- Aravind L, Mazumder R, Vasudevan S and Koonin EV Trends in protein evolution inferred from sequence and structure analysis. *Curr Opin Struct Biol* 2002, 12:392-399
- 12. Pei J and Grishin NV GGDEF domain is homologous to adenylyl cyclase. Proteins 2001, 42:210-216
- Makarova KS, Aravind L, Grishin NV, Rogozin IB and Koonin EV A DNA repair system specific for thermophilic Archaea and bacteria predicted by genomic context analysis. Nucleic Acids Res 2002, 30:482-496
- Koonin EV, Wolf YI, Kondrashov AS and Aravind L Bacterial homologs of the small subunit of eukaryotic DNA primase. J Mol Microbiol Biotechnol 2000, 2:509-512
- 15. Artymiuk PJ, Poirrette AR, Rice DW and Willett P **A polymerase I** palm in adenylyl cyclase? *Nature* 1997, **388:**33-34
- Murzin AG How far divergent evolution goes in proteins. Curr Opin Struct Biol 1998, 8:380-387
- Aravind L, Leipe DD and Koonin EV Toprim a conserved catalytic domain in type IA and II topoisomerases, DnaG-type primases, OLD family nucleases and RecR proteins. Nucleic Acids Res 1998, 26:4205-4213
- Keck JL, Roche DD, Lynch AS and Berger JM Structure of the RNA polymerase domain of E. coli primase. Science 2000, 287:2482-2486
- Podobnik M, McInerney P, O'Donnell M and Kuriyan J A TOPRIM domain in the crystal structure of the catalytic core of Escherichia coli primase confirms a structural link to DNA topoisomerases. J Mol Biol 2000, 300:353-362
- 20. Bushnell DA, Cramer P and Kornberg RD Structural basis of transcription: alpha-amanitin-RNA polymerase II cocrystal at 2.8 A resolution. Proc Natl Acad Sci U S A 2002, 99:1218-1222
- 21. Gnatt AL, Cramer P, Fu J, Bushnell DA and Kornberg RD Structural basis of transcription: an RNA polymerase II elongation complex at 3.3 A resolution. Science 2001, 292:1876-1882
- 22. Cramer P, Bushnell DA and Kornberg RD **Structural basis of tran**scription: RNA polymerase II at 2.8 angstrom resolution. *Science* 2001, **292**:1863-1876
- 23. Vassylyev DG, Sekine S, Laptenko O, Lee J, Vassylyeva MN, Borukhov S and Yokoyama S Crystal structure of a bacterial RNA polymerase holoenzyme at 2.6 A resolution. *Nature* 2002, 417:712-719
- 24. Zhang G, Campbell EA, Minakhin L, Richter C, Severinov K and Darst SA Crystal structure of Thermus aquaticus core RNA polymerase at 3.3 A resolution. *Cell* 1999, **98**:811-824
- Mourrain P, Beclin C, Elmayan T, Feuerbach F, Godon C, Morel JB, Jouette D, Lacombe AM, Nikic S and Picault N Arabidopsis SGS2 and SGS3 genes are required for posttranscriptional gene silencing and natural virus resistance. *Cell* 2000, 101:533-542
- Schiebel W, Pelissier T, Riedel L, Thalmeir S, Schiebel R, Kempe D, Lottspeich F, Sanger HL and Wassenegger M Isolation of an RNAdirected RNA polymerase-specific cDNA clone from tomato. Plant Cell 1998, 10:2087-2101
- 27. Vaistij FE, Jones L and Baulcombe DC **Spreading of RNA targeting** and **DNA methylation in RNA silencing requires transcrip**tion of the target gene and a putative **RNA-**dependent **RNA** polymerase. *Plant Cell* 2002, **14:**857-867
- Dracheva S, Koonin EV and Crute JJ Identification of the primase active site of the herpes simplex virus type I helicase-primase. J Biol Chem 1995, 270:14148-14153
- Holm L and Sander C DNA polymerase beta belongs to an ancient nucleotidyltransferase superfamily. Trends Biochem Sci 1995, 20:345-347

- Aravind L and Koonin EV DNA polymerase beta-like nucleotidyltransferase superfamily: identification of three new families, classification and evolutionary history. Nucleic Acids Res 1999, 27:1609-1618
- Sweetser D, Nonet M and Young RA Prokaryotic and eukaryotic RNA polymerases have homologous core subunits. Proc Natl Acad Sci U S A 1987, 84:1192-1196
- 32. Ebright RH RNA polymerase: structural similarities between bacterial RNA polymerase and eukaryotic RNA polymerase II. J Mol Biol 2000, 304:687-698
- Passarelli AL, Todd JW and Miller LK A baculovirus gene involved in late gene expression predicts a large polypeptide with a conserved motif of RNA polymerases. J Virol 1994, 68:4673-4678
- 34. Passarelli AL and Miller LK Identification of genes encoding late expression factors located between 56.0 and 65.4 map units of the Autographa californica nuclear polyhedrosis virus genome. Virology 1993, 197:704-714
- Iyer LM, Aravind L and Koonin EV Common origin of four diverse families of large eukaryotic DNA viruses. J Virol 2001, 75:11720-11734
- Dieci G, Hermann-Le Denmat S, Lukhtanov E, Thuriaux P, Werner M and Sentenac A A universally conserved region of the largest subunit participates in the active site of RNA polymerase III. *Embo* J 1995, 14:3766-3776
- 37. Iyer LM, Kumpatla SP, Chandrasekharan MB and Hall TC **Transgene** silencing in monocots. *Plant Mol Biol* 2000, **43:**323-346
- Cogoni C and Macino G Post-transcriptional gene silencing across kingdoms. Curr Opin Genet Dev 2000, 10:638-643
- 39. Grishok A and Mello CC RNAi (Nematodes: Caenorhabditis elegans). Adv Genet 2002, 46:339-360
- 40. Fire A RNA-triggered gene silencing. Trends Genet 1999, 15:358-363
- Montgomery MK and Fire A Double-stranded RNA as a mediator in sequence-specific genetic silencing and co-suppression. Trends Genet 1998, 14:255-258
- Parrish S, Fleenor J, Xu S, Mello C and Fire A Functional anatomy of a dsRNA trigger: differential requirement for the two trigger strands in RNA interference. *Mol Cell* 2000, 6:1077-1087
 Zamore PD, Tuschl T, Sharp PA and Bartel DP RNAi: double-
- Žamore PD, Tuschl T, Sharp PA and Bartel DP RNAi: doublestranded RNA directs the ATP-dependent cleavage of mRNA at 21 to 23 nucleotide intervals. Cell 2000, 101:25-33
- Hamilton AJ and Baulcombe DC A species of small antisense RNA in posttranscriptional gene silencing in plants. Science 1999, 286:950-952
- Hammond SM, Bernstein E, Beach D and Hannon GJ An RNA-directed nuclease mediates post-transcriptional gene silencing in Drosophila cells. Nature 2000, 404:293-296
- 46. Hutvagner G and Zamore PD A microRNA in a multiple-turnover RNAi enzyme complex. Science 2002, 297:2056-2060
- Astier-Manifacier S and Cornuet P [RNA-dependent RNApolymerase in Brassica chinensis L]. C R Acad Sci Hebd Seances Acad Sci D 1970, 270:2587-2590
- Astier-Manifacier S and Cornuet P RNA-dependent RNA polymerase in Chinese cabbage. Biochim Biophys Acta 1971, 232:484-493
- Ikegami M and Fraenkel-Conrat H Characterization of the RNAdependent RNA polymerase of tobacco leaves. J Biol Chem 1979, 254:149-154
- Schiebel W, Haas B, Marinkovic S, Klanner A and Sanger HL RNAdirected RNA polymerase from tomato leaves. I. Purification and physical properties. J Biol Chem 1993, 268:11851-11857
- 51. Cogoni C and Macino G Gene silencing in Neurospora crassa requires a protein homologous to RNA-dependent RNA polymerase. Nature 1999, 399:166-169
- 52. Pickford AS, Catalanotto C, Cogoni C and Macino G Quelling in Neurospora crassa. Adv Genet 2002, 46:277-303
- Litiere K, van Eldik GJ, Jacobs JJ, Van Montagu M and Cornelissen M Posttranscriptional gene silencing of gn1 in tobacco triggers accumulation of truncated gn1-derived RNA species. Rna 1999, 5:1364-1373
- Martens H, Novotny J, Oberstrass J, Steck TL, Postlethwait P and Nellen W RNAi in Dictyostelium: the role of RNA-directed RNA polymerases and double-stranded RNase. *Mol Biol Cell* 2002, 13:445-453

- Sijen T, Fleenor J, Simmer F, Thijssen KL, Parrish S, Timmons L, Plasterk RH and Fire A On the role of RNA amplification in dsRNAtriggered gene silencing. *Cell* 2001, 107:465-476
- Meins F Jr RNA degradation and models for post-transcriptional gene-silencing. Plant Mol Biol 2000, 43:261-273
- 57. Nishikura K A short primer on RNAi: RNA-directed RNA polymerase acts as a key catalyst. *Cell* 2001, **107**:415-418
- Ahlquist P RNA-dependent RNA polymerases, viruses, and RNA silencing. Science 2002, 296:1270-1273
- Argos P A sequence motif in many polymerases. Nucleic Acids Res 1988, 16:9909-9916
- Zhang Z, Schaffer AA, Miller W, Madden TL, Lipman DJ, Koonin EV and Altschul SF Protein sequence similarity searches using patterns as seeds. Nucleic Acids Res 1998, 26:3986-3990
- Holm L and Sander C Touring protein fold space with Dali/ FSSP. Nucleic Acids Res 1998, 26:316-319
- Castillo RM, Mizuguchi K, Dhanaraj V, Albert A, Blundell TL and Murzin AG A six-stranded double-psi beta barrel is shared by several protein superfamilies. *Structure Fold Des* 1999, 7:227-236
- 63. Mizuguchi K, Dhanaraj V, Blundell TL and Murzin AG N-ethylmaleimide-sensitive fusion protein (NSF) and CDC48 confirmed as members of the double-psi beta-barrel aspartate decarboxylase/formate dehydrogenase family. Structure Fold Des 1999, 7:R215-216
- 64. Coles M, Diercks T, Liermann J, Groger A, Rockel B, Baumeister W, Koretke KK, Lupas A, Peters J and Kessler H The solution structure of VAT-N reveals a 'missing link' in the evolution of complex enzymes from a simple betaalphabetabeta element. Curr Biol 1999, 9:1158-1168
- Lo Conte L, Brenner SE, Hubbard TJ, Chothia C and Murzin AG SCOP database in 2002: refinements accommodate structural genomics. Nucleic Acids Res 2002, 30:264-267
- Leipe DD, Wolf YI, Koonin EV and Aravind L Classification and evolution of P-loop GTPases and related ATPases. J Mol Biol 2002, 317:41-72
- Wang B, Jones DN, Kaine BP and Weiss MA High-resolution structure of an archaeal zinc ribbon defines a general architectural motif in eukaryotic RNA polymerases. Structure 1998, 6:555-569
- Aravind L and Koonin EV DNA-binding proteins and evolution of transcription regulation in the archaea. Nucleic Acids Res 1999, 27:4658-4670
- 69. Hard T, Rak A, Allard P, Kloo L and Garber M The solution structure of ribosomal protein L36 from Thermus thermophilus reveals a zinc-ribbon-like fold. J Mol Biol 2000, 296:169-180
- Chen HT, Legault P, Glushka J, Omichinski JG and Scott RA Structure of a (Cys3His) zinc ribbon, a ubiquitous motif in archaeal and eucaryal transcription. Protein Sci 2000, 9:1743-1752
- Ferri ML, Peyroche G, Siaut M, Lefebvre O, Carles C, Conesa C and Sentenac A A novel subunit of yeast RNA polymerase III interacts with the TFIIB-related domain of TFIIIB70. Mol Cell Biol 2000, 20:488-495
- 72. Bell SD and Jackson SP The role of transcription factor **B** in transcription initiation and promoter clearance in the archaeon Sulfolobus acidocaldarius. J Biol Chem 2000, 275:12934-12940
- 73. Hahn S and Roberts S The zinc ribbon domains of the general transcription factors TFIIB and Brf: conserved functional surfaces but different roles in transcription initiation. Genes Dev 2000, 14:719-730
- 74. Aravind L and Landsman D AT-hook motifs identified in a wide variety of DNA-binding proteins. Nucleic Acids Res 1998, 26:4413-4421
- Neuwald AF, Liu JS, Lipman DJ and Lawrence CE Extracting protein alignment models from the sequence database. Nucleic Acids Res 1997, 25:1665-1677
- Anantharaman V, Koonin EV and Aravind L Regulatory potential, phyletic distribution and evolution of ancient, intracellular small-molecule-binding domains. J Mol Biol 2001, 307:1271-1292
- Van Duyne GD, Ghosh G, Maas WK and Sigler PB Structure of the oligomerization and L-arginine binding domain of the arginine repressor of Escherichia coli. J Mol Biol 1996, 256:377-391
- Aravind L and Koonin EV Novel predicted RNA-binding domains associated with the translation machinery. J Mol Evol 1999, 48:291-302

- Fletcher CM, Pestova TV, Hellen CU and Wagner G Structure and interactions of the translation initiation factor eIF1. Embo J 1999, 18:2631-2637
- Kraulis PJ Similarity of protein G and ubiquitin. Science 1991, 254:581-582
- 81. Murzin AG Familiar strangers. Nature 1992, 360:635
- 82. Baumann P, Qureshi SA and Jackson SP Transcription: new in-
- sights from studies on Archaea. Trends Genet 1995, 11:279-283
 83. Bell SD and Jackson SP Mechanism and regulation of transcrip-
- tion in archaea. Curr Opin Microbiol 2001, 4:208-213
 84. Lagos-Quintana M, Rauhut R, Lendeckel W and Tuschl T Identification of novel genes coding for small expressed RNAs. Science
- 2001, **294:**853-858 85. Dennis C **The brave new world of RNA.** *Nature* 2002, **418:**122-124
- Reinhart BJ, Weinstein EG, Rhoades MW, Bartel B and Bartel DP MicroRNAs in plants. Genes Dev 2002, 16:1616-1626
- Pasquinelli AE MicroRNAs: deviants no longer. Trends Genet 2002, 18:171-173
- Grosshans H and Slack FJ Micro-RNAs: small is plentiful. J Cell Biol 2002, 156:17-21
- Llave C, Kasschau KD, Rector MA and Carrington JC Endogenous and silencing-associated small RNAs in plants. Plant Cell 2002, 14:1605-1619
- Llave C, Xie Z, Kasschau KD and Carrington JC Cleavage of Scarecrow-like mRNA targets directed by a class of Arabidopsis miRNA. Science 2002, 297:2053-2056
- Wassarman KM, Repoila F, Rosenow C, Storz G and Gottesman S Identification of novel small RNAs using comparative genomics and microarrays. Genes Dev 2001, 15:1637-1651
- 92. Davids W, Amiri H and Andersson SG Small RNAs in Rickettsia: are they functional? Trends Genet 2002, 18:331-334
- 93. Eddy SR Non-coding RNA genes and the modern RNA world. Nat Rev Genet 2001, 2:919-929
- 94. Rivas E, Klein RJ, Jones TA and Eddy SR Computational identification of noncoding RNAs in E. coli by comparative genomics. *Curr Biol* 2001, 11:1369-1373
- Woese C The universal ancestor. Proc Natl Acad Sci U S A 1998, 95:6854-6859
- 96. Leipe DD, Aravind L and Koonin EV Did DNA replication evolve twice independently? *Nucleic Acids* Res 1999, 27:3389-3401
- Aravind L, Anantharaman V and Koonin EV Monophyly of class I aminoacyl tRNA synthetase, USPA, ETFP, photolyase, and PP-ATPase nucleotide-binding domains: implications for protein evolution in the RNA. Proteins 2002, 48:1-14
- Lingner J, Hughes TR, Shevchenko A, Mann M, Lundblad V and Cech TR Reverse transcriptase motifs in the catalytic subunit of telomerase. Science 1997, 276:561-567
- Nakamura TM, Morin GB, Chapman KB, Weinrich SL, Andrews WH, Lingner J, Harley CB and Cech TR Telomerase catalytic subunit homologs from fission yeast and human. Science 1997, 277:955-959
- 100. Altschul SF, Madden TL, Schaffer AA, Zhang J, Zhang Z, Miller W and Lipman DJ Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. Nucleic Acids Res 1997, 25:3389-3402
- 101. Aravind L and Koonin EV Gleaning non-trivial structural, functional and evolutionary information about proteins by iterative database searches. J Mol Biol 1999, 287:1023-1040
- 102. Wootton JC Non-globular domains in protein sequences: automated segmentation using complexity measures. Comput Chem 1994, 18:269-285
- 103. Schaffer AA, Aravind L, Madden TL, Shavirin S, Spouge JL, Wolf YI, Koonin EV and Altschul SF Improving the accuracy of PSI-BLAST protein database searches with composition-based statistics and other refinements. Nucleic Acids Res 2001, 29:2994-3005
- 104. Notredame C, Higgins DG and Heringa J T-Coffee: A novel method for fast and accurate multiple sequence alignment. J Mol Biol 2000, 302:205-217
- 105. Neuwald AF, Liu JS and Lawrence CE Gibbs motif sampling: detection of bacterial outer membrane protein repeats. Protein Sci 1995, 4:1618-1632
- 106. Schuler GD, Altschul SF and Lipman DJ A workbench for multiple alignment construction and analysis. Proteins 1991, 9:180-190

- 107. Walker DR and Koonin EV SEALS: a system for easy analysis of lots of sequences. Proc Int Conf Intell Syst Mol Biol 1997, 5:333-339
- Holm L and Sander C Dali/FSSP classification of three-dimensional protein folds. Nucleic Acids Res 1997, 25:231-234
- 109. Rost B and Sander C Prediction of protein secondary structure at better than 70% accuracy. J Mol Biol 1993, 232:584-599
 110. Rost B, Sander C and Schneider R PHD – an automatic mail serv-
- 110. Rost B, Sander C and Schneider R PHD an automatic mail server for protein secondary structure prediction. Comput Appl Biosci 1994, 10:53-60
- III. Guex N and Peitsch MC SWISS-MODEL and the Swiss-Pdb-Viewer: an environment for comparative protein modeling. Electrophoresis 1997, 18:2714-2723
- 112. Kraulis PJ MOLSCRIPT: A Program to Produce Both Detailed and Schematic Plots of Protein Structures. Journal of App Crystallography 1991, 24:946-950

