

The Extended “Two-Barrel” Polymerases Superfamily: Structure, Function and Evolution

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Abstract

DNA and RNA polymerases (DNAP and RNAP) play central roles in genome replication, maintenance and repair, as well as in the expression of genes through their transcription. Multisubunit RNAPs carry out transcription and are represented, without exception, in all cellular life forms as well as in nucleo-cytoplasmic DNA viruses. Since their discovery, multisubunit RNAPs have been the focus of intense structural and functional studies revealing that they all share a well-conserved active-site region called the two-barrel catalytic core. The two-barrel core hosts the polymerase active site, which is located at the interface between two double-psi β -barrel domains that contribute distinct amino acid residues to the active site in an asymmetrical fashion. Recently, sequencing and structural studies have added a surprising variety of DNA and RNA to the two-barrel superfamily, including the archaeal replicative DNAP (PoID), which extends the family to DNA-dependent DNAPs involved in replication. While all these polymerases share a minimal core that must have been present in their common ancestor, the two-barrel polymerase superfamily now encompasses a remarkable diversity of enzymes, including DNA-dependent RNAPs, RNA-dependent RNAPs, and DNA-dependent DNAPs, which participate in critical biological processes such as DNA transcription, DNA replication, and gene silencing. The present review will discuss both common features and differences among the extended two-barrel polymerase superfamily, focusing on the newly discovered members. Comparing their structures provides insights into the molecular mechanisms evolved by the contemporary two-barrel polymerases to accomplish their different biological functions.

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Introduction

Polymerases are essential for life being responsible for preserving genetic information by replicating and repairing nucleic acid molecules as well as for the expression of genes through transcription [1]. While all nucleotide polymerases likely use a two-metal ion catalytic general mechanism [2–4], nucleotide polymerization has been invented several times during evolution. Most nucleotide polymerases can be divided into three major groups [5,6]. The first group contains nucleotide polymerases that structurally resemble the *Escherichia coli* Pol-I Klenow-fold [7]. Their overall fold is characterized by thumb, finger, and palm subdomains and is shared by most eukaryotic, archaeal, and viral replicative DNA polymerases (DNAPs), the bacteriophage single-

subunit RNA polymerase (RNAP), the mitochondrial RNAP, and reverse transcriptases [8,9]. The second group, often referred to as Pol β -like polymerases, includes the bacterial replicative DNAPs and several eukaryotic DNAPs involved in DNA repair [10]. While they share a similar three-dimensional arrangement of catalytic aspartates in the active site with the first group, they show a completely different topology of the palm subdomain. The third group includes multisubunit RNA polymerases (msRNAPs), which carry out transcription and is represented, without exception, in all cellular life forms as well as in nucleo-cytoplasmic DNA viruses (NCLDV). Since their discovery about 60 years ago, cellular msRNAPs have been the focus of intense structural and functional studies revealing that they all share a well-conserved active-site region, consisting of two

Table 1. Subunit composition and activity of two-barrel polymerases

Organisms			Two-barrel core subunits		Other core subunits	Biological function
			DPBB-1 (DPBB-B)	DPBB-2 (DPBB-A)		
DNA-dependent RNAP	Eukaryotes	RNAPI	A135	A190	AC19, AC40, RPB4–10, RPB12	Transcription of ribosomal RNA
		RNAPII	RPB2	RPB1	RPB3–12	Transcription of pre-messenger RNA
		RNAPIII	C128	C160	AC19, AC40, RPB5–6, RPB8, RPB10, RPB12, C11, C17, C25	Transcription of small RNAs
		RNAPIV (Plant)	NRPD/E2	NRPD1	RPB3, RPB5, RPB8, RPB10–12, NRPD/E4, NRPD7, NRPD9b	Production of small interfering RNAs
		RNAPV (Plant)	NRPD/E2	NRPE1	RPB3, NRPE5, RPB8, RPB9–12, NRPD/E4, NRPE7	Transcription of the whole gene repertoire
	Archaea	<i>M. jannaschii</i>	RPO2(A)	RPO1(A)	RPO1(A) [*] , RPO3(D), RPO4(F), RPO5(H), RPO6(K), RPO7(E), RPO8(G), RPO10(N), RPO12(P)	Transcription of the whole gene repertoire
	Bacteria	(e.g., <i>E. coli</i>) Epsilonproteobacteria	β-Subunit Fusion β/β'-subunit	β'-Subunit	α-Subunit, ω-subunit α-Subunit, ω-subunit	
	Chloroplasts	Plastid-encoded prokaryotic-type RNAP	RPOB	RPOC ₁	RPOA, RPOC ₂ , PAP1–10	Major RNAP activity in mature chloroplasts
	Viruses	NCLDVs (e.g., poxviruses)	RPO2	RPO1	RPO7, RPO22, RPO30	Transcription (cytoplasmic phase of the virus cycle)
		φKZ giant phage <i>Baculoviruses</i>	GP71–73 LEF-8	GP55 LEF-9	GP68, GP74, GP123 LEF-4, P47	Late gene expression
<i>Firmicutes phages</i>		YonO			Late gene expression	
RNA-dependent RNAP	Yeast	pGKL2	ORF6	ORF7	pGKL1 (and 2?) gene expression	
	Plasmids	(e.g., <i>K. lactis</i>)			PTG silencing	
	Eukaryotes	<i>N. crassa</i> <i>N. crassa</i> <i>N. crassa</i>	QDE-1 SAD-1 RRP-3		Meiotic silencing PTG silencing?	
DNA-dependent DNAP	Archaea	PolD	DP2	DP1	DNA replication	

Alternative names that are common in the literature are shown in brackets.

double-psi β-barrel (DPBB) subdomains [11,12]. However, sequencing and structural studies have added a surprising variety of nucleotide polymerases to the two-barrel superfamily [13–15] (Table 1). Recently, the two-barrel polymerase superfamily was substantially diversified with the structure elucidation of PolD, an archaeal replicative DNAP, which extends the family to DNA-dependent DNAPs involved in replication [16,17].

All two-barrel polymerases share a common catalytic core, which is located at the interface between two DPBB subdomains that contribute distinct amino acid residues to the active site in an asymmetrical fashion (Table 1). The first DPBB (DPBB-1 and DPBB-B in msRNAPs, respectively) harbors two canonical lysine residues involved in substrate binding, while the second DPBB (DPBB-2 and DPBB-A in msRNAPs, respectively) contributes

invariant aspartic residues that coordinate a catalytic Mg²⁺. The present review will discuss both common features and differences among members of the extended two-barrel polymerase superfamily, focusing on the newly discovered members. While all these polymerases share a minimal core that must have been present in their common ancestor, the two-barrel polymerase superfamily now encompasses a remarkable diversity of enzymes, including DNA-dependent RNAPs, RNA-dependent RNAPs [18,19], and DNA-dependent DNAPs, which participate to critical biological processes such as DNA transcription, DNA replication, and gene silencing. Comparing their structures provides insights into the remarkable variety of molecular mechanisms, which were evolved by the contemporary two-barrel polymerases in order to achieve their different biological functions.

msRNAPs: The Remarkably Conserved Cellular Transcriptases

In all forms of cellular life, the genomes are transcribed by a highly conserved family of msRNAPs, which was discovered about 60 years ago and has since then been the focus of intense structural and functional studies [11]. Bacteria and archaea have one single type of msRNAP to transcribe their gene repertoire, whereas eukaryotes have three RNAPs, responsible mainly for the synthesis of ribosomal RNA (RNAPI), pre-messenger RNA (RNAPII), and small RNAs including transfer RNAs (RNAPIII), respectively (Table 1). Plastids encode two types of RNAPs: two different single-subunit, nuclear-encoded phage-type RNAPs, and one plastid-encoded multi-subunit RNAPs, whose subunits are homologous to the bacterial RNAP [20]. Over the past two decades, x-ray crystallographic and cryo-electron microscopy (cryo-EM) structures have been determined for msRNAPs from the three domains of life, revealing that these enzymes operate by closely related molecular mechanisms [12].

All cellular msRNAPs share an exceptionally conserved large active-site region, composed of five core subunits, which resemble a crab claw, the jaws of which interact with the downstream duplex DNA template [21–27] (Fig. 1). The two large subunits form the core of the enzyme and host the polymerase catalytic center (Rpb1 and Rpb2 in Eukaryotic RNAPII, Rpo1 and Rpo2 in archaeal RNAP, β and β' in bacterial RNAP), which have been pinpointed by extensive biochemical and genetic analyses [29–31] (Table 1). These two catalytic subunits are anchored and stabilized by three smaller “assembly” subunits (the Rpb3/Rpb11 heterodimer and Rpb6 in Eukaryotic RNAPII; the Rpo3/Rpo11 heterodimer and Rpo6 in archaeal RNAP; the α/α homodimer and ω subunits in bacterial RNAP) [11]. In addition, the activity of msRNAPs is modulated by several transcription factors during the three distinct phases of the transcription cycle: initiation, elongation, and termination. The active site of msRNAPs is located at the interface between the two core subunits and can be dissected within several functionally critical regions [21,32] (Fig. 1).

In the transcribing RNAP, the downstream DNA traverses the major DNA-binding channel, lining the “floor,” until it encounters the active center of the RNAP “wall.” The DNA-binding channel forms a positively charged cleft, lined from one side by a mobile clamp, and from the other side by the lobe and protrusion domains. The DNA–RNA hybrid rises up from the active-site perpendicular to the downstream duplex DNA. The hybrid and clamp

domains secure about one helical turn of the DNA–RNA hybrid, whose strands are separated by the RNAP lid. The catalytic core harbors the active site, including the Mg^{2+} -bound two-barrel catalytic core, the bridge, the DNA–RNA hybrid-binding helix, and trigger helix, which are essential for the nucleotide translocation cycle [33–37]. The two-barrel catalytic center is organized around two catalytic Mg^{2+} ions, each Mg^{2+} being harbored by one of the two barrels. The DPBB-A of the largest RNAP catalytic subunit carries three invariant aspartic residues in the highly conserved NADFDGD motif, which coordinates a catalytic Mg^{2+} ion. A second Mg^{2+} is required for catalysis but comes with the substrate. The DPBB-B of the second catalytic subunit possesses two invariant lysine residues, whose side chains face the last two nucleotides of the RNA primer in the elongating bacterial and eukaryotic RNAPs [38]. The bridge helix and the two-barrel catalytic center line a perforation in the floor of the cleft, which widens toward the exterior, creating an inverted funnel, named the pore, or secondary channel. The pore allows substrates and cleavage factors to access the active site, and allows extrusion of the transcript during backtracking. The outer rim of the funnel is lined by the funnel region.

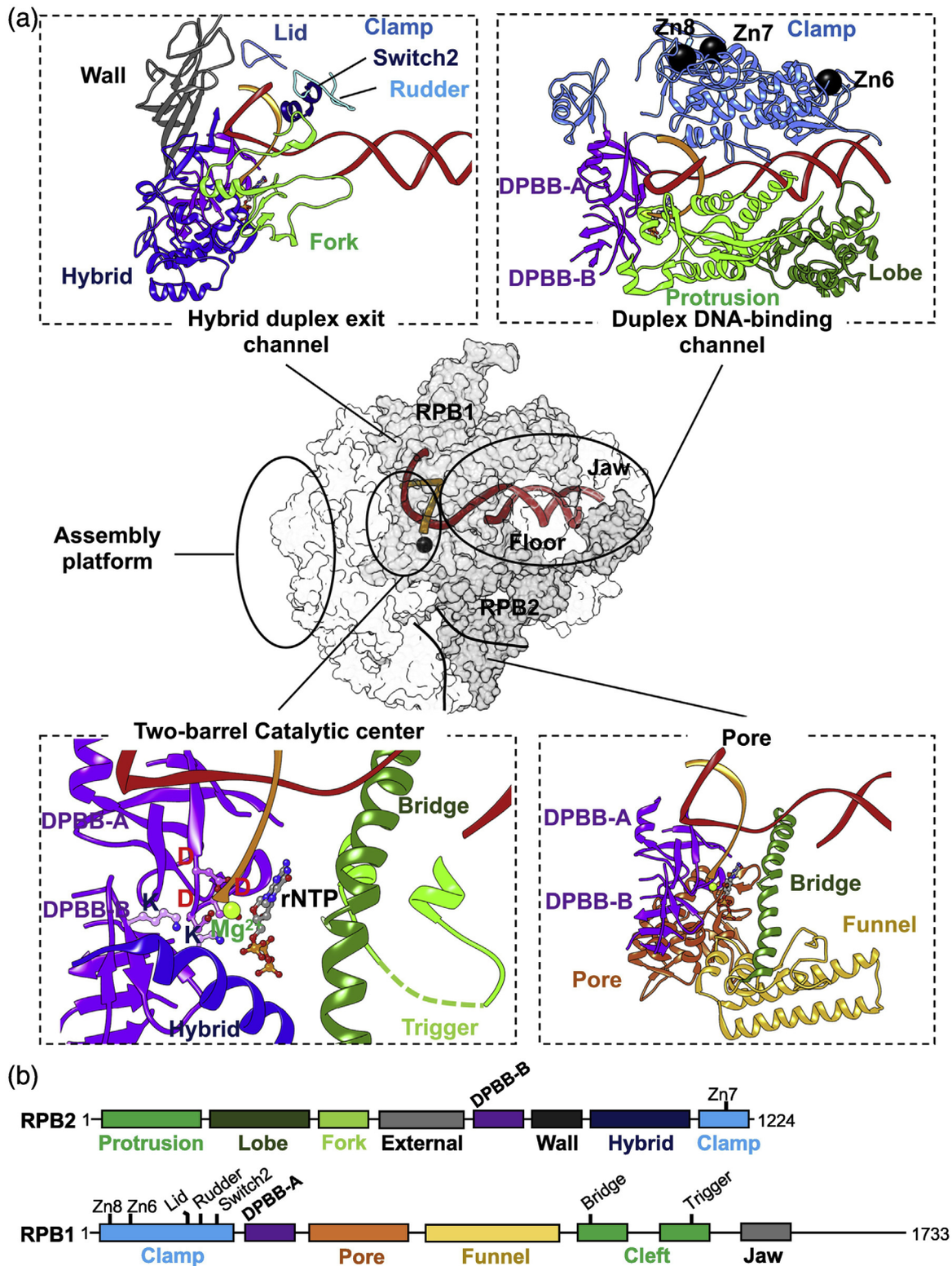
Non-Canonical msRNAPs from DNA Viruses

In addition to being present in all forms of cellular life, msRNAPs are encoded by several lineages of NCLDV, which are one of the largest viral division that includes viruses with large DNA genomes that infect diverse eukaryotes [39]. NCLDVs encode their own msRNAP, which presumably derives from the eukaryotic RNAPI form [40] and operates during the cytoplasmic phase of their infectious cycle, when the viral DNA is not accessible to the host nuclear transcription system. Their multi-subunit structure systematically includes the two largest subunits, Rpb1 and Rpb2, as well as several “assembly” subunits, which play a critical role in the organization of the active-site region (Table 1) [41]. Interestingly, several DNA viruses encode non-canonical msRNAPs that do not include any classical “assembly” subunits.

For example, the well-characterized insect Baculovirus encodes a non-canonical msRNAP, which is responsible for transcribing a relatively small number of viral late genes, including the polyhedrin gene, which is used to overexpress foreign proteins in the baculovirus expression systems [42–44]. These msRNAPs are made of four subunits: p47, LEF4, LEF8, and LEF9 (LEF stands for late expression factor), all with unknown structures (Table 1) [45].

These subunits are unrelated to canonical RNAPs, except for a limited similarity of its LEF-9 and LEF-8 subunits to Rpb1 and Rpb2, respectively [46–48].

An updated sequence analysis revealed that this region of homology not only includes the two barrels, but encompasses also the switch 2 domain and the



funnel domain [13]. However, the two subunits (LEF4 and p47) have no sequence similarity with any classical “assembly” subunit or with any known msRNAP subunits [46,49,50].

Recent sequencing and biochemical studies have revealed that giant phages encode non-canonical msRNAPs [51]. While the transcription of DNA phages usually depends on a phage-encoded Klenow-like monomeric RNAP or on the host transcription system, the sequencing of the giant phages from *Pseudomonas aeruginosa* (Φ EL, Φ KZ180) and *Pseudomonas chlororaphis* (201 Φ 2) have revealed two sets of proteins distantly related to the β' and β subunits of bacterial msRNAPs [51,52]. Biochemical characterization of the non-canonical msRNAP encoded by the Φ KZ giant phage has revealed that it specifically initiates transcription of late phage promoters in a rifamycin-resistant manner [53]. The Φ KZ phage-encoded msRNAP includes five subunits: four subunits are homologous to the cellular msRNAPs; the fifth subunit is a protein of unknown function (Table 1). Like the msRNAP from Baculovirus, the Φ KZ msRNAP lacks identifiable “assembly” and promoter specificity subunits characteristic for cellular msRNAPs.

Non-canonical msRNAPs from insect viruses (baculoviral and nudiviral lineages) and giant phages may respectively originate from eukaryotic (insect viruses) and bacterial (phages) RNAP ancestor genes, which underwent a fast and highly divergent evolution obliterating anything that was not strictly needed for DNA transcription [13]. Strikingly, none of these enzymes harbor a subunit that may play a role in RNAP assembly. The reduced repertoire of the different viral msRNAPs supports the idea that primordial msRNAPs were mainly composed of the two catalytic core subunits. This hypothesis is further supported by the fact that a variety of two-barrel polymerases carry the entire active site on a single polypeptide chain, suggesting that they are closer to the ancestral protein than their msRNAPs counterparts.

Single-chain RNA- and DNA-dependent RNAPs

Eukaryotic cells use small RNAs (sRNAs) guides to limit proliferation of viruses and transposable

elements, maintain proper chromosomal structure, and control gene expression [54–56]. In Fungi, plants, and several animal systems, sRNA production often depends on template-dependent RNA synthesis catalyzed by single-chain RNA-dependent RNAPs [57]. RNA-dependent RNAPs have been initially described in plants as a cellular RNAP activity induced upon viral infection that synthesized antisense RNA in a primer-dependent or independent manner [58–60]. Subsequent genetic studies showed that mutations in RNA-dependent RNAP genes impaired gene silencing in a variety of systems [58,61]. Within the same organism, they often coexist under structurally and functionally diverse forms such as the three paralogous genes present in *Neurospora crassa*, one of the best-studied models for gene silencing [61,62]. *N. crassa* thus encodes three distinct RNA-dependent RNAPs: SAD-1, that is involved in meiotic silencing [63]; RRP-3 and quelling defective phenotype (QDE-1), which participate in post-transcriptional gene silencing [61,62]. Isolation and biochemical characterization of recombinant QDE-1 orthologs from related fungi shows that their mode of action differs considerably. While *N. crassa* QDE-1 efficiently produces full-length copies of short 9- to 21-nucleotide copies scattered throughout the input ssRNA templates and is capable to extend complementary primers [64], *Thielavia terrestris* QDE-1 produces predominantly short RNA copies via primer independent initiation [65].

Crystal structures of large C-terminal fragments of QDE-1 from both *N. crassa* and *T. terrestris* have been determined by x-ray crystallography [19,65]. QDE-1 forms a functional homodimer, which was confirmed by gel filtration, sedimentation assays [19], and electron microscopy [65]. Consistently, crystal structures of the QDE-1 dimers show a pyramidal shape with the two-barrel catalytic core located at the base of each subunit (Fig. 2A). Each subunit includes a catalytic domain that houses the two-barrel catalytic core and the catalytic loop. The catalytic loop hosts three canonical aspartate residues that coordinate a Mg^{2+} ion, which has been shown to be required for enzymatic activity (Fig. 2B) [64]. While they are arranged sequentially on a single polypeptide chain, the DPBBs subdomains in QDE-1 and msRNAPs are structurally very similar and almost identically disposed. The two-barrel catalytic core forms a cleft that is surrounded by

Fig. 1. Structure of the msRNAP catalytic subunits. (A) The central panel is a simplified diagram that illustrates the overall architecture of msRNAPs. Important structural and functional features discussed in the main text are highlighted. Top and bottom boxes show enlarged cartoon representations of functionally important regions of the RNAP: the hybrid duplex exit channel, the DNA-binding channel, the two-barrel catalytic center, and the pore. Structural information is derived from *S. cerevisiae* RNAPII [Protein Data Bank (PDB) ID: 2e2i] [28]. Individual domains are named according to Ref. (21). (B) Distribution of the conserved domains and motifs on the RPB1 and RPB2 subunits. Domains are colored as in panel A.

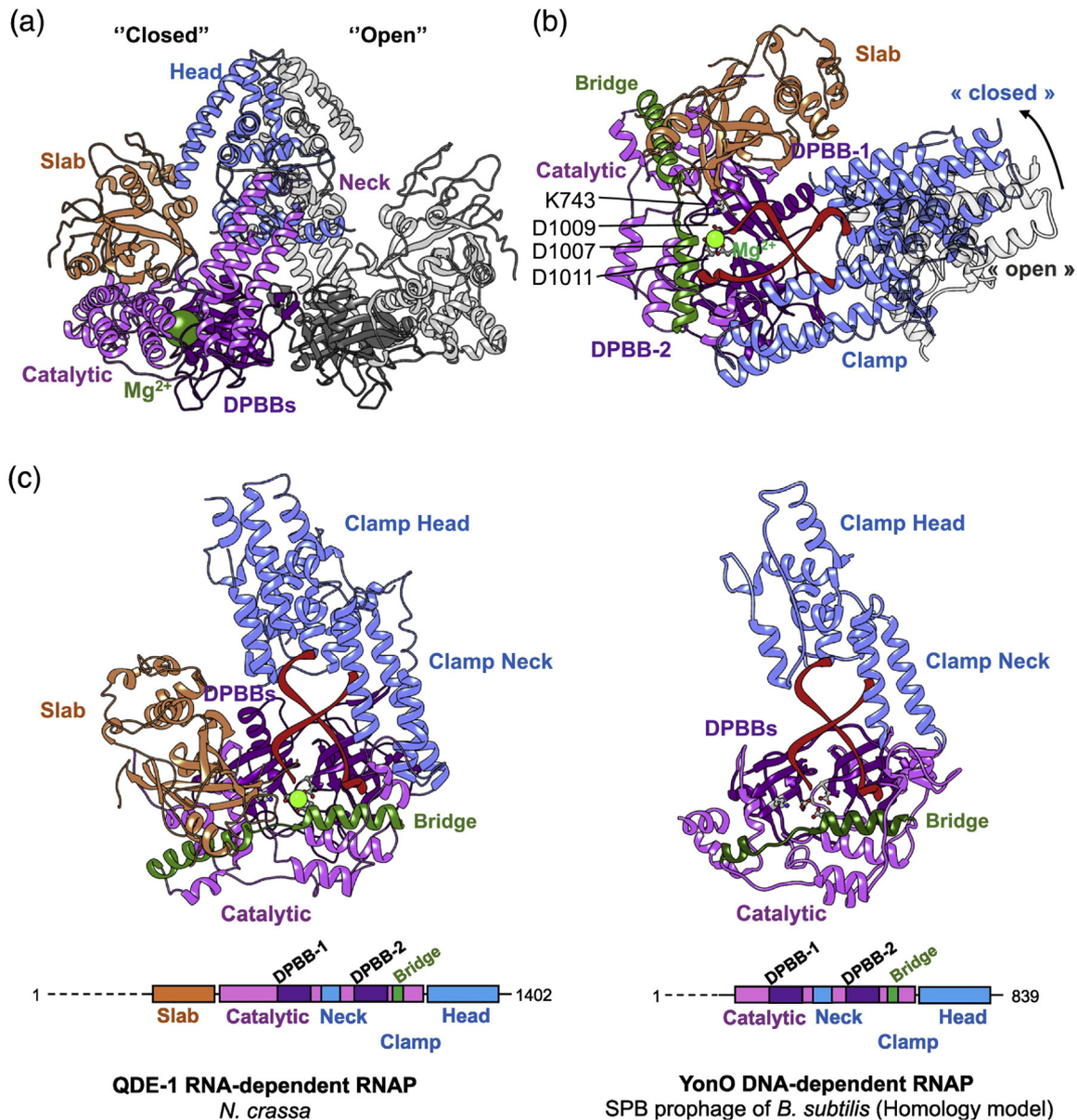


Fig. 2. Structures of the single-chain two-barrel RNAPs. (A) Ribbon diagrams of the *N. crassa* QDE-1 homodimer (PDB ID: 2J7N) [19] showing the subunit A colored according to domains and subunit B colored gray. Mg^{2+} ions are marked as spheres. (B) Cartoon representation of the QDE-1 subunit in its “closed” conformation colored according to domains. For comparison, the head domain in its “open” conformation is colored in gray. The direction of the movement of the head domain is indicated as a gray arrow. The RNA duplex is modeled based on a superposition with an elongation complex of *S. cerevisiae* RNAPII [28]. (C) Comparison of the structure of a *N. crassa* QDE-1 monomer (left panel) with a homology model of the YonO protein (right panel), originating from the prophage SP β . The homology model that covers about 70% of the YonO sequence was predicted using Phyre 2 [66], based on the crystal structure of *N. crassa* QDE-1. Distribution of the domains on the QDE-1 and YonO polymerases is shown in the bottom panels. The RNA/DNA hybrid duplex is modeled based on a superposition with an elongation complex of *S. cerevisiae* RNAPII [28].

the head and slab domains, which form an extensive groove that is well adapted for accommodating the template and the nascent RNA product. In addition, QDE-1 also contains the flap subdomain peripheral to the active-site cleft and the neck domain that

connects the catalytic domain to the head domain. The QDE-1 structure shows a highly positively charged channel formed between the slab and head of each subunit, which leads to the active site and has been proposed to accommodate the

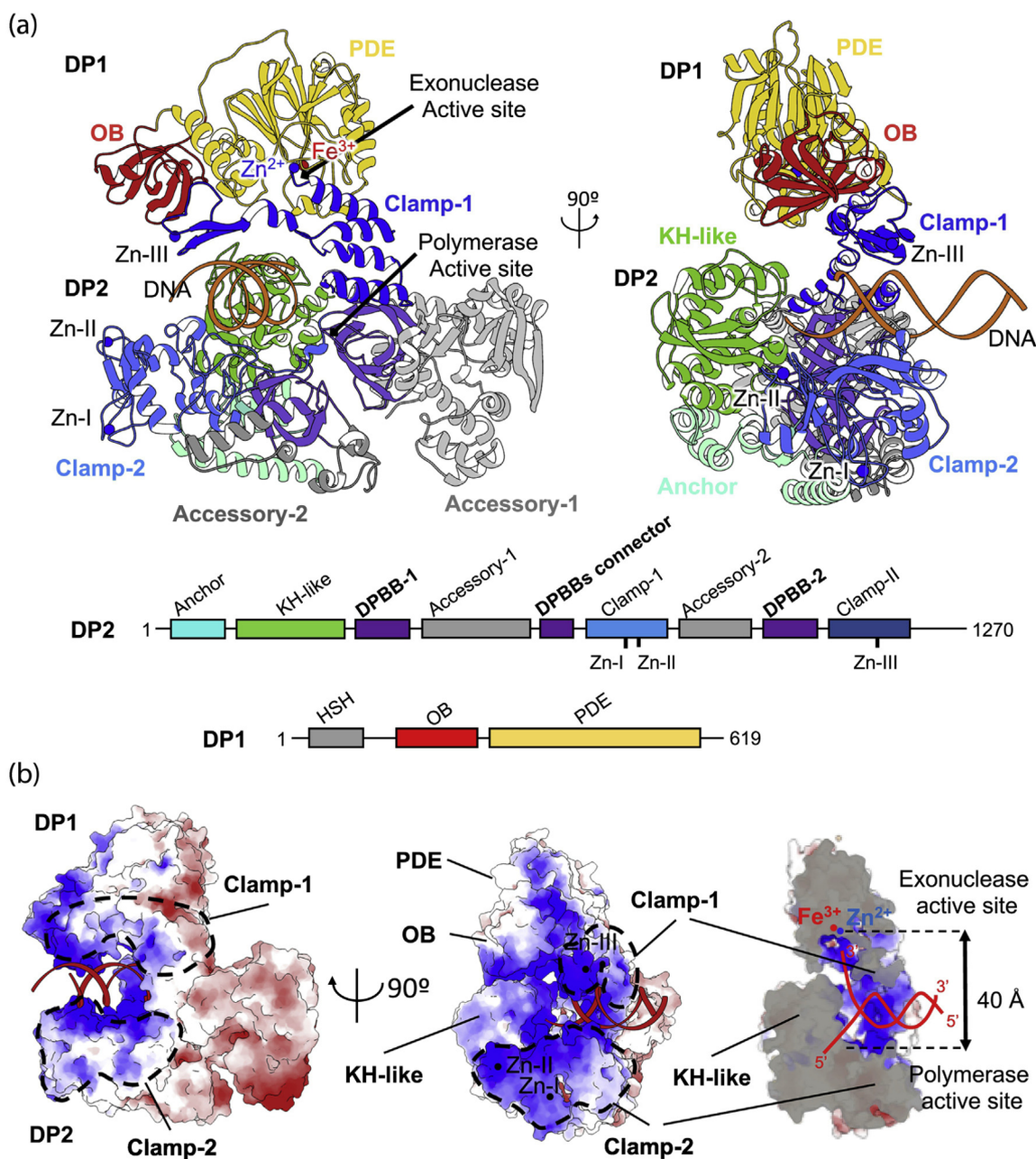
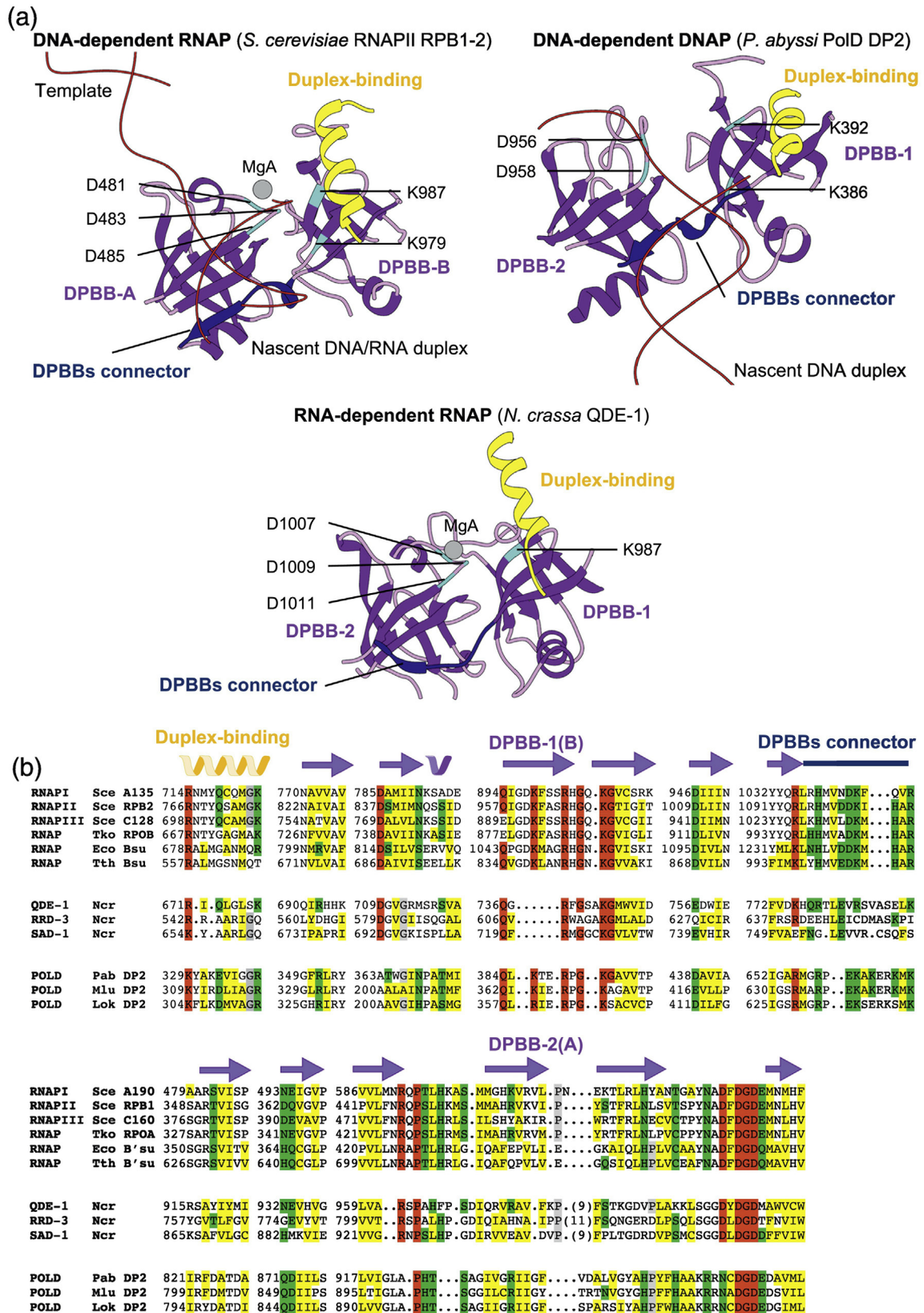


Fig. 3. Overall structure and surface architecture of PolD: the two-barrel DNAP. (A) Ribbon diagrams of the DNA-bound PolD structure highlighting the domains and domain-like regions that compose the DP1 and DP2 subunits. Zn^{2+} and Fe^{3+} ions are shown as spheres (PDB ID: 6hms). The bottom panels show the distribution of the conserved domains and motifs on the DP1 and DP2 subunits of PolD. Domains are colored as above. (B) Surface charge distribution of the PolD–DNA-binding site. The surface of PolD is colored according to the electrostatic surface potential, with negative, neutral, and positive charges shown in red, white, and blue, respectively. The electrostatic potential was calculated as discussed in Ref. (17) using a PolD model containing all side-chain atoms. The clamp-1 and clamp-2 domains are highlighted by dotted lines. The DNA backbone is colored red. The right panel shows a cutaway view of PolD showing a putative path for the DNA being digested by the proofreading exonuclease domain.

nascent dsRNA product [19]. A second charged tunnel may be a route for the entry of nucleic acid triphosphates. QDE-1 shares with its DNA-dependent msRNAPs counterparts the two-barrel catalytic

core [18], the bridge helix, and the proposed nucleic acid triphosphate tunnel that matches with the one proposed in yeast RNAPII [38]. The mobile head domain may be equivalent to the clamp domain in



multi-subunit RNAPs by closing down on the slab domain to stabilize the RNA product during polymerization. Interestingly, the proposed dsRNA product-binding channels are not identical in the two subunits and adopt either an open or a closed conformation. This is mainly due to the disposition of the head domains that is different in the two subunits. In the subunit showing a closed conformation, the head domain is clamped down on the active-site cleft, while in the subunit showing an open conformation, rotation of the head domain provides space for an RNA duplex [19] (Fig. 2B). The structure suggests that only one catalytic site is active, represented by the closed conformation, with the inactive subunit being held open. The dimeric QDE-1 polymerase has been proposed to work as a “two-stroke” engine [19]. In this mechanism, substrate binding to one active site would prime the other, thereby enabling an efficient re-initiation, leading to the effective production of appropriate-length dsRNA triggers.

Iyer and colleagues [18] identified a similarity between the single-chain RNA-dependent RNAPs and the YonO protein of *Bacillus subtilis*, originating from an inserted prophage SP β . Yono-like open-reading frames have since then been found in several other firmicute phages [13,67]. Recently, YonO was shown to be a highly processive DNA-dependent RNAP that specifically transcribes the late genes of the SP β prophage [68]. Strikingly, the homology of YonO to msRNAPs is restricted to only few amino acids, and most conserved domains of msRNAPs are absent in YonO. Instead, as noted previously [18], YonO shows a much stronger similarity to its single-chain two-barrel RNA-dependent RNAPs counterparts. Interestingly, a putative 3D model covering ~70% of the YonO sequence could be predicted using Phyre 2 [66], based on the crystal structure of *N. crassa* QDE-1 (Fig. 2C). This similarity encompasses most of the catalytic and clamp domains of QDE-1. Interestingly, QDE1 regions contributing to the interactions between the two monomers corresponds to insertions that appear

to be absent in YonO, thereby suggesting that homodimerization is a specificity of RNA-dependent RNAPs. However, biochemical and structural data are required to confirm this model and decipher the molecular specificities of single-chain YonO-like DNA-dependent RNAPs over single-chain RNA-dependent RNAPs.

PoID: The Archaeal DNA-Dependent Two-Barrel Replicative DNAP

In all forms of cellular life, DNAPs play central role in genome replication, maintenance, and repair. Over the years, all DNAPs have been grouped in different families, using sequence alignments: PolA, PolB, PolC, PolD, PolX, PolY, and reverse transcriptases [8,9,69]. The main replicative DNAPs from Eukarya are found in family B, Bacteria in family C, and Archaea in families B and D [6]. PolD is a heterodimeric replicative DNAP composed of a large catalytic subunit (DP2) and a smaller subunit with 3′–5′ proofreading exonuclease activity (DP1) [70–72]. PolD has been shown to be essential for cell viability [73–75] and is widely distributed among Archaea, being present in all four major superphyla: Euryarchaeota (including the methanogenic human symbionts); DPANN (Diapherotrites, Parvarchaeota, Aenigmarchaeota, Nanoarchaeota, Nanohaloarchaeota); the emerging Asgard superphylum; and TACK (Thaumarchaeota, Aigarchaeota, Crenarchaeota, Korarchaeota), only absent from Crenarchaeota [76]. Based on biochemical evidence, it has been proposed that PolD may act soon after initiation by the primase [77,78]. Resolving the individual crystal structures of the DP1 and DP2 catalytic cores from the archaeon *Pyrococcus abyssi* revealed that PolD is an atypical DNAP that has all functional properties of a replicative DNAP but with the catalytic core of a two-barrel RNAP [16]. Recently, the cryo-EM structure of the DNA-bound heterodimeric DP1–DP2 PolD complex was reported [17]. This structure sheds light on DNA-binding domains

Fig. 4. The minimal catalytic core shared by all two-barrel polymerases: structure and sequence conservation. (A) Cartoon representations showing the shared structural elements of the two-barrel catalytic core between *P. abyssi* PolD (PDB ID: 6hms) [17], *S. cerevisiae* RNAPII (PDB ID: 2e2i) [28], and *N. crassa* QDE1 (PDB ID: 2j7n) [19]. The two-barrel core is colored according to the conserved structural elements: the two DPBBs (purple), the duplex-binding helix (yellow), and the DPBBs-connector (blue). The invariant aspartic and lysine residues harbored by the two DPBBs are colored in cyan. Mg²⁺ ions are marked as spheres (gray). (B) Structure-based alignment of the shared structural elements within the two-barrel catalytic core of PolD, msRNAPs and single-chain RNA-dependent RNAPs. The shared secondary structure elements are shown above the alignment. Yellow boxes highlight hydrophobic residues, green boxes highlight polar residues, gray boxes highlight conserved glycine or proline positions, and red boxes highlight highly conserved catalytic motifs. The following structures were used for the structure-based alignment: *S. cerevisiae* RNAPI [89] (PDB ID: 5m5x), *S. cerevisiae* RNAPII [28] (PDB ID: 2e2i), *S. cerevisiae* RNAPIII [90] (PDB ID: 6f41), *Thermococcus kodakarensis* RNAP [91] (PDB ID: 4qiw), *E. coli* RNAP [92] (PDB ID: 6asx), *Thermus thermophilus* RNAP [93] (PDB ID: 2o5i), *N. crassa* QDE-1 [19] (PDB ID: 2j7o), and *P. abyssi* PolD [17] (PDB ID: 6hmf).

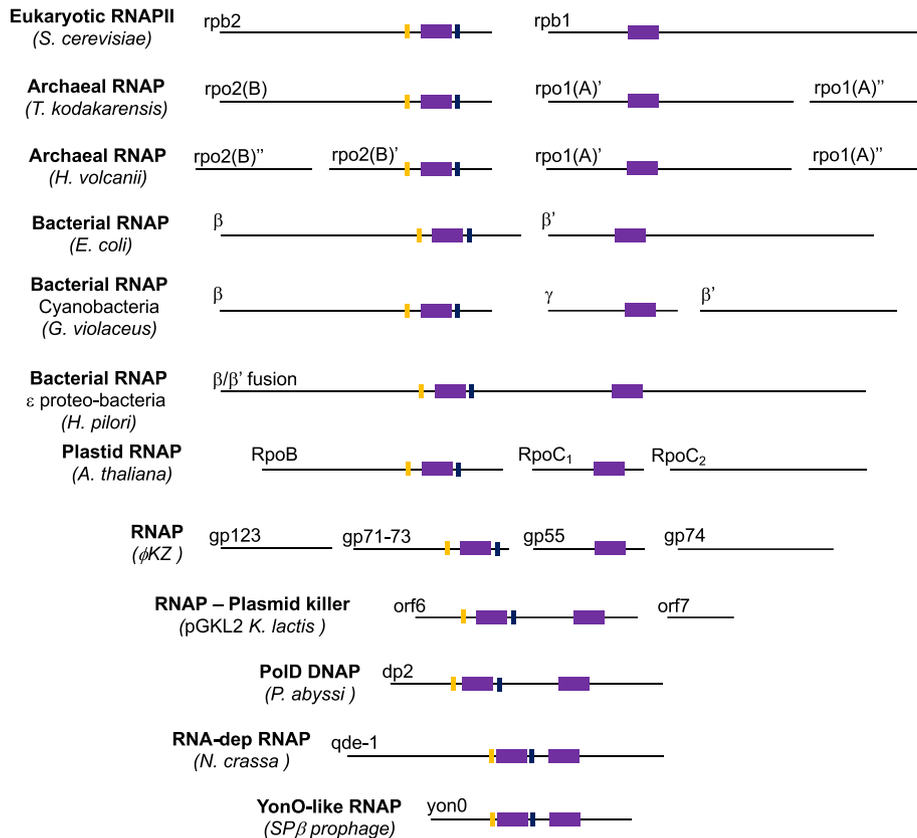


Fig. 5. Distribution of the conserved structural elements forming the catalytic core in the extended two-barrel polymerase superfamily. The conserved structural elements are colored as in Fig. 4: the two DPBBs (purple), the duplex-binding helix (yellow), and the DPBBs-connector (blue). The structural elements composing the two-barrel minimal core are clustered and share a similar organization in the primary sequence of all single-chain two-barrel polymerases. In msRNAPs, regions coding for the two halves of the two-barrel minimal core are, respectively, clustered at the C-terminal end of one core subunit and at the N-terminal end of the second core subunit.

evolved by PoID to perform DNA replication and extends the repertoire of protein domains known to be involved in DNA replication (Fig. 3A).

All replicative DNAPs have evolved protein domains named palm, fingers, and thumb domains arranged to form the DNA-binding cleft. The palm domain carries the catalytic residues, the fingers domain drapes over the nascent base pair, and the thumb domain holds the DNA duplex during replication and contributes to processivity [79,80]. PoID has evolved a specific claw-shaped active site, at the center of which is the two-barrel catalytic core and at the edge of which are three zinc-binding modules named the Zn-I, Zn-II, and Zn-III domains. The DNA substrate is cradled between a bipartite clamp domain, named 1 and 2, emanating from DPBB-1 and DPBB-2, respectively (Fig. 3B). Clamp-1 and clamp-2 domains contribute a central cleft with a diameter of 30 Å, which is located upstream of the DP2 polymerase catalytic center. The Zn-III domain plays a critical role in replication by interacting with the minor groove of the nascent duplex [81]. The Zn-I and Zn-II domains contribute to stabilizing clamp-

2, which is otherwise composed mainly of loops. The clamp domain is barricaded from one side by a K-homology (KH)-like domain located in the N-terminal region of DP2, which is ideally located to orient the DNA template in the active site. This domain shares structural homology with the archaeal/bacterial type-II KH domains, which are ancestral single-stranded nucleic acid-binding folds [82]. KH domains had been predicted in the N-terminal region of bacterial PoIC [83], and the cryo-EM structure of PoID shows in a structural context, a KH domain associated with a replicative DNAP. It is noteworthy that KH domains are found in transcription factors, including in the NusA elongation factor, which harbors a type-I KH domain [84,85]. Two accessory domains, named – 1 and – 2, emanating from insertions in DPBB-1 and DPBB-2 subdomains, respectively, play a structural role by scaffolding together the essential two-barrel catalytic cores and the clamp-1 and clamp-2 domains. The KH-like domain is connected to the anchor domain, which firmly attaches it to the clamp-2 domain and orients it in the active site (Fig. 3A).

The exonuclease and polymerase active sites of PoID are hosted by two distinct subunits [72,86] (Fig. 3). DP1 shows an oligonucleotide binding domain that is inserted within the N-terminal region of a large Mre11-like nuclease phosphodiesterase domain [87] whose active-site entry faces the 3' end of the nascent DNA strand. The DP1 nuclease and DP2 polymerase active sites sandwich the 3' end of the nascent DNA strand and are about 40 Å distant from each other, a feature shared with other replicative DNAPs [80]. Although the cryo-EM PoID structure shows no contacts between DP1 and the bound DNA, the exonuclease active site is suitably located to catch the 3' end of the nascent DNA strand. The phosphate moiety of the 3'-terminal nucleotide of the primer lies at 25 Å away from the exonuclease active site of DP1, which could be accounted for by a 4-nt-long single-stranded DNA, a value shared with other DNAPs with proofreading activity [80]. It is noteworthy that msRNAPs have evolved a proofreading mechanism called backtracking, which involves a nucleolytic reaction that occurs in the two-barrel catalytic core, at the same site as polymerization [88]. While PoID also hosts a polymerase two-barrel core, it evolved another proofreading mechanisms, which recruited a phosphodiesterase domain domain. In that sense, PoID differs from its two-barrel counterparts but resembles other replicative DNAPs, which contain a distinct nuclease-specific active site.

The Minimal Catalytic Core Shared by All Two-Barrel Polymerases

The two-barrel polymerase superfamily encompasses DNA-dependent RNAPs, RNA-dependent RNAPs [13,18,19], and DNA-dependent DNAPs [16]. Comparing their structures enables us to delineate the minimal core shared by all these distantly related DNAPs or RNAPs and discuss the molecular basis for their respective substrate specificities (Fig. 4). All these polymerases share a two-barrel catalytic core with several invariant amino acids located in the immediate vicinity of their catalytic Mg^{2+} cations. Superimposing these two-barrel polymerases reveals that the relative disposition of their DPBB subdomains is highly conserved. Interestingly, perturbing the disposition of the two barrels in the individual DP2 crystal structure of PoID severely impairs the activity of PoID [16,17]. All two-barrel polymerases share an α -helix that is connected to the N-terminal end of DPBB-1 (DPBB-B, in msRNAPs) and shows a conserved orientation with respect to the two-barrel catalytic core and the nascent duplex in both PoID and RNAPII DNA-bound structures (Fig. 4A). This helix, arbitrarily named duplex-binding helix, contains two canonical basic residues and shares a similar orientation with

respect to the two-barrel catalytic core and the nascent duplex in both families of enzymes [28]. Finally, PoID and two-barrel RNAPs share a loop, arbitrarily named DPBBs-connector, that emanates from the C-terminal end of DPBB-1 (DPBB-B, in msRNAPs) and is connected to the adjacent DPBB through secondary structure interactions. The DPBBs-connector likely participates in maintaining the correct relative disposition of the two DPBBs: in the individual crystal structure of DP2, the DPBBs-connector is found to be disordered and the canonical disposition of the two DPBBs is lost. It is noteworthy that the active sites of both PoID and msRNAPs orient the DNA template entry and the nascent duplex exit in a same relative axis with respect to the two-barrel catalytic core.

A structure-based alignment of the two-barrel polymerases sequences over the structurally homologous catalytic core reveals that their common architecture is underpinned by the conservation of hydrophobic, polar, and charged residues (Fig. 4B). In particular, several positively charged residues occupy conserved positions in this shared minimal core. These residues are located in several conserved motifs of the two DPBBs subdomains, the duplex-binding helix and the DPBBs-connector. They contribute to form a circular clamp that circumvents the nascent duplex in both RNAPII and PoID DNA-bound structures. In particular, DPBB-B in msRNAPs and DPBB-1 in PoID carry two extremely conserved positively charged residues (RPB2-K979 and RPB2-K987 in *Saccharomyces cerevisiae* RNAPII; DP2-K386 and DP2-K392 in *P. abyssi* PoID). In both DNA-bound structures, the positively charged side chains of these residues are found to contact the phosphate moieties of the 3' end of the nascent RNA or DNA. Genetic studies in *S. cerevisiae* [94] and in *E. coli* [95] revealed that even precise inter-atomic distance between the two residues is highly critical for RNAP activity, given the lethality of lysine to arginine mutations.

Comparison of two-barrel polymerases also reveals structural determinants that are specific to DNAPs versus RNAPs. The catalytic site of all RNA- or DNA-dependent two-barrel RNAPs is characterized by two catalytic Mg^{2+} that are coordinated by four invariant aspartic residues (479DFDGD483 in RPB1 and D837 in RPB2 in *S. cerevisiae* RNAPII; 1007DYDGD1011 and D709 in *N. crassa* QDE-1) [38,96]. Only two aspartic residues are strictly conserved in PoID: 956DGD958, respectively, corresponding to 481DGD483 in *S. cerevisiae* RPB1 and 1009DGD1011 in *N. crassa* QDE-1, possibly explaining why no density accounting for the presence of Mg^{2+} has been observed so far in the active site of PoID, neither in the DNA-bound PoID cryo-EM structure [17] or in the DP2 individual crystal structure [16]. Phosphates of the incoming nucleotide may thus be required in order to bind the

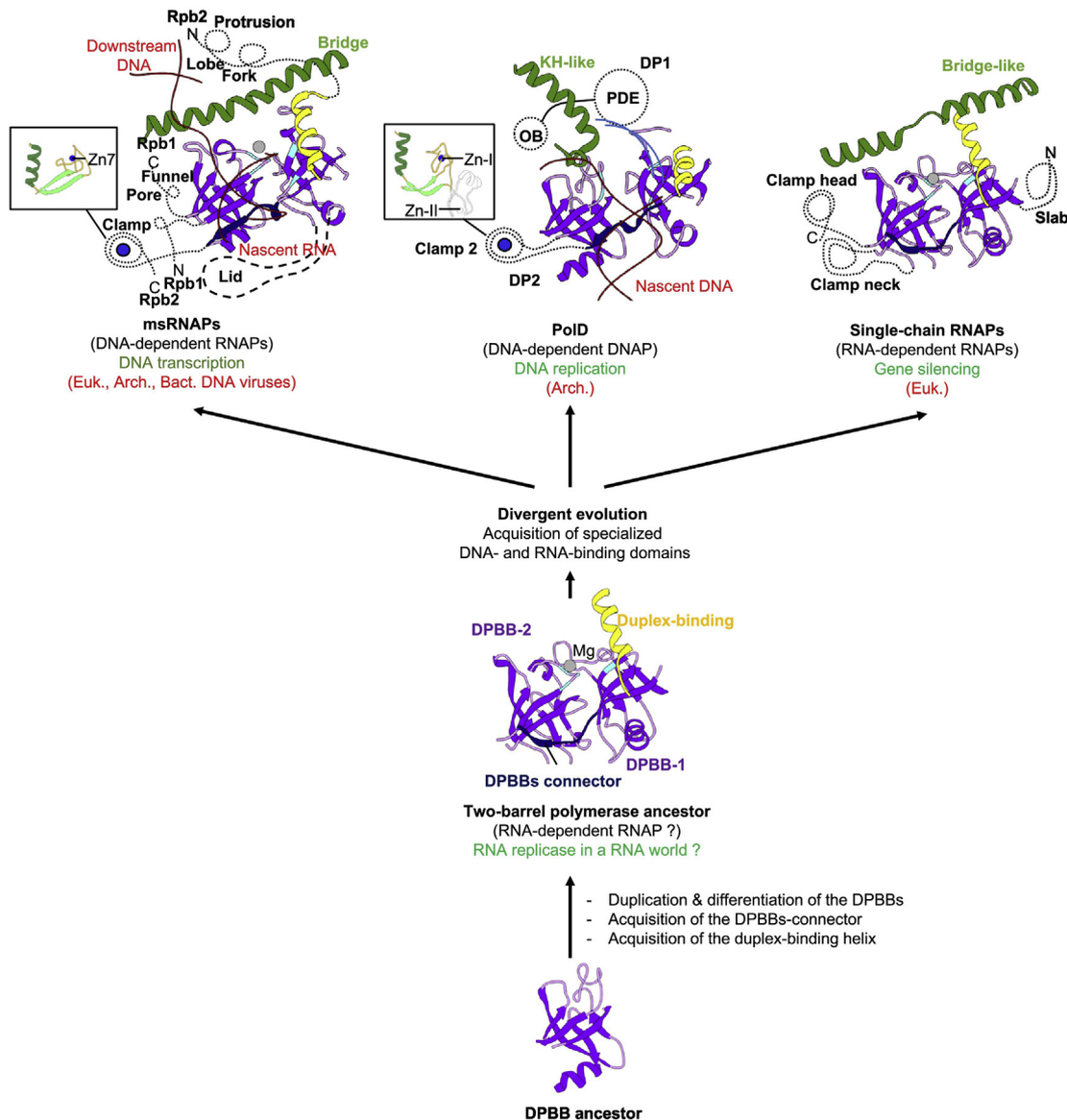


Fig. 6. A hypothetical scheme of evolution of the two-barrel catalytic core.

two catalytic Mg^{2+} ions in the active site, as observed for other DNAPs [97].

A Conserved Arrangement of the Genes Encoding the Structural Elements of the Two-Barrel Minimal Core

Interestingly, the structural elements composing the two-barrel minimal core are clustered and share a similar organization in the primary sequence of all single-chain two-barrel polymerases: DNA-dependent DNAPs (PoID), RNA-dependent RNAPs (QDE-1, SAD-1 and RPR-3), and DNA-dependent RNAPs (YonO-like polymerases) (Fig. 5). In contrast to

single-chain two-barrel polymerases, the gene organization of msRNAPs reveals a much higher level of complexity. In most cellular msRNAPs, the two-barrel minimal core is split within two distinct subunits: RPB1 and RPB2 in eukaryotic RNAPII, RPO1, and RPO2 in archaea, β and β' in bacteria (Table 1). However, it is striking that the regions coding for the two halves of the two-barrel minimal core are, respectively, clustered at the C-terminal end of one core subunit and at the N-terminal end of the second core subunit, thereby suggesting that although msRNAPs share a conserved ancestor with their single-chain counterparts, diversification and complexification may have resulted in a scission of their minimal catalytic core into two separate

genes (Fig. 5). Splitting the two-barrel catalytic core onto separate subunits may thus have facilitated the radical evolution of the common single-chain two-barrel polymerase ancestor into the complex msRNAPs [14]. This hypothesis is supported by the fact that epsilon proteo-bacteria harbor a fused single catalytic subunit encompassing both β and β' subunits [98]. Furthermore, the yeast *Kluyveromyces lactis* killer DNA plasmids encodes a non-canonical two-barrel RNAP that resembles a fusion of portions of the two RPB1 and RPB2 core subunits [99–101]. It is puzzling that the splits occurred in the middle of the two-barrel minimal core, leaving two distinct subunits, each carrying one half of the active site. Splitting the active site in two distinct subunits ensures that only correctly-assembled msRNAPs are active and may be important for regulating the activity of these enzymes.

Evolution of Two-Barrel Polymerases

All two-barrel polymerases evolved from a common ancestor, probably at a very early stage of evolution. One might hypothesize that their common ancestor has been responsible for transmission of RNA genomic information in a RNA-based world. The capacity of PoID to use RNA-primed DNA and occasionally incorporate ribonucleotides may be a property inherited from its common ancestor to RNAPs [77,102]. Similarly, the potentially ancient RNA-dependent activity of msRNAPs is illustrated by the fact that they can use RNA templates in special cases [103,104]. Two-barrel polymerases most likely evolved from a single DPBB ancestor (Fig. 6). Indeed, DPBB is an ancient fold that is found as single copy in the active site of several metabolic enzymes in all domains of life [18]. Gene duplication led to a molecule with two DPBBs on a single polypeptide chain, which differentiated and recruited both DPBBs-connector and duplex-binding structural elements to produce a molecule capable of efficient RNA polymerization. This primordial two-barrel minimal core may have served as a RNA-binding protein cofactor to a self-replicating ribozyme in an ancient RNA world, before taking over the polymerization catalytic activity.

While the structures of two-barrel polymerases have undergone significant diversifications from their common ancestor, these distantly related polymerases show intriguing similarities. All two-barrel polymerases thus evolved a nucleic acid binding clamp domain, which occupies the same relative location with respect to the two-barrel active site. In PoID and msRNAPs, clamp domains host several loops that participate in binding the nascent DNA–DNA or RNA–DNA duplex and several zinc-binding domains (Zn-I and Zn-II domains in PoID; Zn-7, Zn-8, and Zn-6 in *S. cerevisiae* RNAP-II), which

stabilize the loop-rich clamp domains. Whereas these clamps do not share sequence motifs, Zn-I of PoID and Zn-7 of RNAP-II share the same topology, which is conserved in all eukaryotic/archaeal msRNAPs. Consistently, when the structures of PoID and the elongation complex of *S. cerevisiae* RNAPII are superimposed on their two-barrel catalytic cores, the nascent RNA/DNA hybrid in RNAPII and the nascent DNA duplex in PoID show a very similar orientation with respect to their two-barrel catalytic core [17].

The two-barrel catalytic core is found to be associated with a wide range of structurally distinct DNA-binding domains, which contribute to the wide range of substrate specificities (DNA-dependent or RNA-dependent), activities (RNAPs and DNAPs), and functions (DNA transcription, DNA replication, and gene silencing) exhibited by contemporaneous two-barrel polymerases. The two-barrel minimal core is remarkably versatile and evolved toward a wide range of biological functions in modern DNA world by accretion of distinct sets of additional domains. Comparing their structures illustrates how this minimal catalytic core has recruited different domains in order to perform distinct biological functions. As an example, msRNAPs that are involved in DNA transcription have evolved the lid domain (absent in PoID and QDE-1), which is inserted in DPBB-B, to prevent the formation of an extended upstream RNA–DNA hybrid. In PoID, two ancestral single-stranded DNA binding domains, oligonucleotide binding and KH-like, have been recruited for DNA replication to guide the DNA template into the active sites and a phosphodiesterase domain to perform proofreading. Although conserved in all two-barrel RNAPs [34], the bridge helix is notably absent in PoID. Instead, the DNA template seems to be guided in the active site of PoID by a KH-like domain [17]. Both the bridge helix in RNAPs and the KH-like domain of PoID display a canonic motif of residues that are colocalized with respect to the DNA substrate, suggesting that they may play a similar role in guiding the DNA substrate into the active site.

Since their discovery, cellular msRNAPs have been the focus of intense structural and functional studies revealing, with an unprecedented level of details, how these molecular machines perform DNA transcription. However, many questions remain unanswered regarding the molecular specificities of their non-canonical two-barrel RNA and DNAPs counterparts. Further structural and functional studies are thus required to decipher the molecular mechanisms evolved by these polymerases for polymerization, template binding, nucleotide selection, and proofreading. In particular, no structure of the non-canonical viral msRNAPs has been reported yet and their elucidation may provide valuable models for comparative analysis of the mechanism and evolution of the extended two-barrel polymerases superfamily.

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cryo-EM, cryo-electron microscopy; msRNAPs, multisubunit RNA polymerases; NCLDVs, nucleo-cytoplasmic DNA viruses; LEF, late expression factor; DNAP, DNA polymerase; DPBB, double-psi β -barrel; KH, K-homology; PDB, Protein Data Bank; QDE-1, quelling defective phenotype; RNAP, RNA polymerase.

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