Tracking in atomic detail the functional specializations in viral RecA helicases that occur during evolution

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ABSTRACT

Many complex viruses package their genomes into empty protein shells and bacteriophages of the Cystoviridae family provide some of the simplest models for this. The cystoviral hexameric NTPase, P4, uses chemical energy to translocate single-stranded RNA genomic precursors into the procapsid. We previously dissected the mechanism of RNA translocation for one such phage, φ12, and have now investigated three further highly divergent, cystoviral P4 NTPases (from φ6, φ8 and φ13). High-resolution crystal structures of the set of P4s allow a structure-based phylogenetic analysis, which reveals that these proteins form a distinct subfamily of the RecA-type ATPases. Although the proteins share a common catalytic core, they have different specificities and control mechanisms, which we map onto divergent N- and C-terminal domains. Thus, the RNA loading and tight coupling of NTPase activity with RNA translocation in φ8 P4 is due to a remarkable C-terminal structure, which wraps around the outside of the molecule to insert into the central hole where RNA binds to coupled L1 and L2 loops, whereas in φ12 P4, a C-terminal residue, serine 282, forms a specific hydrogen bond to the N7 of purines ring to confer purine specificity for the φ12 enzyme.

INTRODUCTION

Viruses protect their genome by condensing it into a compartment, the virion. Many complex viruses rely on rapid encapsidation by energy-dependent transport of the nucleic acid into an empty preformed capsid (procapsid). This process requires the presence of portal complexes, which are conduits for nucleic acid molecules, and molecular motors that convert the chemical energy gained from nucleoside triphosphate (NTP) hydrolysis into mechanical movement, resulting in nucleic acid translocation.

Some viruses, including herpesvirus and tailed double-stranded DNA (dsDNA) bacteriophages, package their genome using a multi-protein packaging motor (terminase) that transiently assembles at a single vertex (1–4). These complexes are relatively elaborate, consisting of a large dodecameric portal that is an integral part of the capsid and an oligomeric transiently associated terminase, neither of which can work in the absence of the other. The ATPase-nuclease terminase subunit is responsible for recruiting the viral DNA to the procapsid. Compacting relatively stiff dsDNA into a small volume of the procapsid has a high energy cost. Single-molecule experiments have revealed that viral packaging proteins can exert forces as high as 110 pN on dsDNA, making them some of the strongest known biological motors (5).

Similarly, dsRNA bacteriophages of the Cystoviridae family (bacteriophages φ6 through to φ14, and φ2954) encapsidate single-stranded RNA (ssRNA) genomic precursors into procapsids (6). However, their packaging
machinery is less complex, consisting of a hexamer that is at the same time the physical portal and the active genome translocating motor (7,8). Although this motor shares the same function of translocating the genomic nucleic acid into the procapsid, the challenges differ between ssRNA and dsDNA. ssRNA is significantly more flexible (persistence length l_p ~ 1–2 nm) than dsDNA (l_p ~ 50 nm) (9), and the packaging densities are less than those found for dsDNA viruses (10); therefore, high forces are probably not required. However, naturally occurring ssRNAs, such as the genomic precursors, exhibit extensive local secondary-structure (11,12), and thus the packaging motor has to exhibit helicase activity.

The lipid-enveloped bacteriophages of the Cystoviridae family infect Gram-negative bacteria, mainly plant-pathogenic Pseudomonas species (13) and share similarities with the members of the Reoviridae family, including bluetongue virus and rotavirus (14). Their genome of ~14 kb consists of three dsRNA segments small (S), medium (M) and large (L), which are sequentially encapsidated as ssRNA precursors into the icosahedrally symmetric procapsid by the packaging NTPase P4 (15–23).

P4 NTPases are structural components of the procapsid, built by co-assembly of 120 copies of the major structural protein P1 with ~10 copies of the viral RNA-dependent RNA polymerase P2, 10 hexamers of P4 and 12 trimers of the assembly cofactor P7 (24) (Figure 1). In bacteriophage φ6, P4 hexamers nucleate procapsid assembly in vitro (7,25), are essential for genome packaging (21) and also have a role in transcription (21,26). Up to 12 P4 hexamers lie on the 5-fold symmetry axes of facets of the procapsid (16,24,27), creating a symmetry mismatch. Although the P4 hexamer constitutes the packaging motor, the specificity for viral RNA is mediated by RNA-binding sites on the P1 shell, which recognize three distinct packaging signals on the genomic precursors (28,29).

Previous studies have revealed the structure and mechanism of φ12 P4 (30–32). P4 is a protein of ~35 kDa, which can assemble into a hexameric ring. NTP-binding sites are located on the external perimeter of the ring at the interfaces between adjacent subunits, whereas the nucleic acid binding sites are found in the central channel (31) (Figure 1). P4 proteins are the only known RNA-specific helicases belonging to helicase Superfamily 4 (SF4) (33). SF4 encompasses mainly DNA helicases and is characterized by five conserved sequence motifs (H1, H1a, H2, H3 and H4) (34). Motifs H1, H1a and H2 are involved in nucleotide binding and hydrolysis, whereas H3 is involved in the coupling of NTP hydrolysis to nucleic acid translocation, and H4 in oligonucleotide binding. Crystal structures of φ12 P4 at different key catalytic states of the protein unveiled a power stroke mechanism by which a conformational change associated with sequential NTP hydrolysis is responsible for RNA translocation (31,35,36).

P4 NTPases show little sequence similarity; however, they are believed to share a common architecture and mechanism of action. When recombinant P4 proteins are studied in isolation, they show variation in their in vitro biochemical properties (Table 1): φ8 and φ13 P4 NTPases form stable complexes with RNA and their ATPase activities are strongly stimulated by RNA (φ8 has no detectable ATPase activity in absence of RNA), whereas φ6 and φ12 P4s bind RNA transiently and are only weakly stimulated; the isolated P4 hexamers of φ8 and φ13 have measurable helicase activities in vitro in contrast to φ6 P4, which only acquires processive helicase activity in the context of the procapsid (30); the φ12 P4 hexamer has low translocation processivity and lacks helicase activity (36); the NTPase activity of φ12 P4 is specific to purine bases (26), whereas the other P4s can also accept pyrimidine bases (8,40). These differences in biochemical properties are presumably reflected in the hexamer architecture and structural details of different domains. To gain further insights into RNA loading, interaction and translocation mechanisms and the structural evolution of these packaging enzymes, we have solved the crystal structures of three additional P4 proteins, from φ8, φ13 and from the prototype virus of the cystoviral family, φ6. We also report here the structural and/or biochemical characterization of φ12 P4 mutants to explain nucleotide specificity and RNA recognition. We compare these structures with that of wild-type φ12, whose structure has already been reported (31), creating a series of structurally related viral packaging motors.

**MATERIALS AND METHODS**

**Cloning, expression and purification**

Recombinant full-length P4 from φ8, φ13 and C-terminally truncated φ8 P4Δ281 (missing residues 281–321) and φ6 P4Δ310 (missing residues 310–331) were expressed from plasmids pSJ1b (41), pDK3 (8), pDK10 (42) and pJTJ7.3/7 (43), respectively. Point mutations were introduced into φ12 gene 4 using plasmid pPG27 (32) as a template to introduce amino acid substitutions S252Q, R272A, Q278A, S292A, Y288A and TTS202-204 by site-directed mutagenesis (QuikChange, Stratagene) following the manufacturer recommendations. The corresponding plasmids were designated as pDK33, pDK35, pDK30, pDK31, pDK29 and pDK249 respectively. The insertion of LKK instead of TTS (residues 202–204) was introduced by amplifying the N-terminal portion of the P4 gene with primers 1 and 2 (Supplementary Table S1) and the

![Figure 1. The cystovirus P4 protein, a molecular packaging motor. (A) Cartoon showing the position of the P4 hexamer (grey) on the empty cystovirus procapsid (black) while packaging ssRNA. (B) Cartoon model of the mechanism of RNA translocation by P4. The energy derived from the hydrolysis of ATP is mechanically converted to the translocation of single-stranded ssRNA.](http://nar.oxfordjournals.org/)

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C-terminus part with primers 3 and 4. PCR products were digested with NdeI/AflII (N-terminal part) and AflII/EcoRI (C-terminal part) and ligated into pT7-7 vector at NdeI-EcoRI sites. Sequencing was used to confirm the mutations.

Recombinant P4 proteins were expressed in *Escherichia coli* BL21(DE3) or B834(DE3) and purified to homogeneity as previously described (31,32,42). Briefly, *E. coli* cells were grown at 37°C in Luria-Bertani medium until OD540nm reached 0.5–0.6. Cultures were then chilled on ice and induced with 1 mM isopropyl-β-thiogalactopyranoside. Induced cells were further incubated for 12–14 h at 17–18°C, harvested by centrifugation and lysed with a French pressure cell. P4 proteins were purified by chromatography: Heparin and Q-sepharose columns (GE Healthcare) followed by size exclusion chromatography (Superdex 200, GE Healthcare).

Cloning, expression, purification and characterization of C-terminally His-tagged φ8 P4 (φ8 P4His), which exhibits full RNA-induced ATPase activity, was described previously (44).

Crystallization

Crystallization conditions of the P4 proteins have been previously described (32,42). In brief, crystals of φ6 P4Δ310 proteins were grown at 24°C from a 3.5 mg/ml protein solution in 20 mM HEPES (pH 8.0), 5 mM MgCl2, 2 mM CaCl2, 5 mM adenosine diphosphate (ADP) and 100 mM NaCl, and they appeared after 9 months in drops in which 3 ml of protein had been mixed with 3 ml of a reservoir solution consisting of 6% PEG 4000 and 90 mM sodium acetate (pH 4.5). Crystals were cryo-protected by transferring them into reservoir solution with a final glycerol concentration of 25% before freezing in a nitrogen-gas stream at −173°C.

From a 12 mg/ml protein solution, φ13 P4 crystals were grown at 20°C using 100 mM Tris–HCl (pH 7.0), 900 mM trisodium citrate and 200 mM NaCl as precipitant. Crystals were cryo-protected as φ6 P4Δ310, but using a final glycerol concentration of 20%.

The φ8 P4 crystals were grown at 24°C in 100 mM sodium acetate (pH 4.6) and 2.2 M ammonium sulphate as a precipitant. Drops consisted of 0.9 μl of protein at a concentration of 3 mg/ml, 0.9 μl of reservoir solution and 0.4 μl of 100 mM dithiothreitol (DTT). Crystals of φ8 P4Δ281 obtained from a protein solution concentrated to 5 mg/ml appeared in 100 mM Tris (pH 8.0) and 18% PEG 1000. Crystals were cryo-protected following the protocol for φ6 P4Δ310.

Crystals of φ12 P4 mutants were obtained in a solution composed of 10% PEG 1500 in 100 mM sodium acetate (pH 4.8) and 5 mM AMPcPP. Crystals of wild-type φ12 P4 with UTP were obtained with the same precipitant and 5 mM UTP.

Data collection and structure determination

Data collection was performed as previously detailed (32,42), and all data were indexed, integrated and scaled using HKL2000 (45). crystallographic statistics for the data are detailed in Supplementary Table S2.
Structures of $\phi 12$ P4 mutants $\phi 12$ P4-Q278A and $\phi 12$ P4-S292A were solved by molecular replacement using the program PHASER (46) with wild-type $\phi 12$ P4 (PDB code 1W4B) as the search model.

The structure of $\phi 13$ P4 was solved by single-wavelength anomalous dispersion as described elsewhere (47). The substructure was determined using the program SHELX (48), and phases were refined using SHARP (49). After 6-fold non-crystallographic symmetry averaging using General Averaging Program (unpublished program available from D. I. Stuart or J. M. Grimes), an interpretable electron density map was obtained into which the structure could be built.

The structure of $\phi 6$ P4 was solved by molecular replacement with the crystal structure of the $\phi 13$ P4 as a search model. The search model included one hexamer in which each chain was truncated to the conserved ATPase core of the protein. A weak molecular replacement solution comprising two truncated hexamers was found by the program AMoRe (50). The preliminary phases were greatly improved by 12-fold non-crystallographic symmetry averaging and phase extension from low resolution using General Averaging Program. The last 34 residues of the $\phi 6$ P4A310 construct were not visible in the electron density; their absence might be due to proteolysis, which would explain the long crystallisation period.

The structure of $\phi 8$ P4 was initially solved by single-wavelength anomalous dispersion from crystals of the selenomethionine labelled protein in space group $P6_22$ containing one monomer in the asymmetric unit. HKL2MAP (48) was used to identify the selenium sites, which were then fed into PHENIX AUTOSOL (51), resulting in an interpretable electron density map for the ATPase core domain. The electron density corresponding to the rest of the protein was not interpretable owing to the statistically disordered crystal reported previously (42). The hexameric P4 was formed by applying the crystallographic symmetry and used as search model for molecular replacement with the program PHASER (46) to find a solution for $\phi 8$ P4 ($R32$ space group) and $\phi 8$ P4A281 ($P2_1;2_1;2_1$ space group).

Manual building was performed with the program COOT (52) and restrained refinement (with TLS) with either AUTOBUSTER (53) or REFMAC5 (54). The final models were validated with MolProbity (55). Refinement statistics are provided in Supplementary Table S2; in summary; the resolution (Å)/R-factor(%)/R-free(%) for the structures were $\phi 6$ P4: 2.8/21.7/24.4, $\phi 8$ P4-His: 3.1/29.6/30.9, $\phi 12$ P4 UTP: 1.9/19.4/20.4, $\phi 13$ P4: 1.7/16.4/18.8.

Hydrogen-deuterium exchange mapping

Previously published hydrogen-deuterium exchange (HDX) data for $\phi 8$ P4 were used (37) and mapped onto the high-resolution structure presented in this work using average rate colouring as described (37).

ATPase activity of mutants

ATPase activity of $\phi 12$ P4-binding site mutants was assayed using the EnzChek phosphate assay kit (Invitrogen) (39).

Evolutionary analysis of structures

The coordinates of the ATPase core of P4 from $\phi 8$ (residues 104–261) were submitted to the DALI Server (56), a program that identifies and ranks proteins by structural similarity. The DALI search returned 47 proteins, which have significant structural similarity to P4. All these proteins were then truncated to their core ATPase domains, and using the program SHP superimposed onto one another, and a matrix of structural relationships was calculated (57).

RESULTS AND DISCUSSION

Overall fold

All P4 proteins form a hexameric ring with a central channel varying in size from 13 to 21 Å (30 Å for $\phi 8$ P4A281) and external diameter of ~100 Å (Figure 2). However, the hexamers have different charge distributions on their surfaces (Supplementary Figure S1) and different outline shapes: $\phi 6$ P4, $\phi 8$ P4 and $\phi 13$ P4 form hexagonal notched rings, whereas $\phi 12$ P4 has a smoother contour. The subunit interface within hexamers varies in size from ~1500 to 1900 Å$^2$, and the number of hydrogen bonds, salt bridges and hydrophobic interactions shows substantial variation (Supplementary Table S3). The interfaces within the P4 hexamers are more polar than expected for a stable oligomer. This is because rings of hexameric helicases are generally required to open to load the nucleic acid strand into the central cavity (Table 1) (58,59). The rounnder $\phi 12$ P4 subunits bury the biggest surface area and form the highest number of hydrogen bonds and salt bridges, whereas the interaction area is least for $\phi 8$ P4, which harbours fewer hydrogen bonds and only three salt bridges. The buried area does not correlate with P4 ring stability. For example, $\phi 12$ P4 has been shown to exhibit frequent ring opening unless it is bound to the procapsid (38), leading to low translocation processivity (36). On the other hand, $\phi 8$ P4 is a processive translocase and opens only during loading a new RNA strand into the central channel (37). Ring stability correlates instead with the fraction of buried polar interactions (hydrogen bonds and salt bridges) per buried area. The less stable $\phi 6$ and $\phi 12$ hexamers have 0.016 and 0.018 polar contacts per Å$^2$ respectively, whereas the more stable $\phi 8$ and $\phi 13$ exhibit values of 0.13 and 0.15, respectively.

ATPase core domain

Within the hexamer, the different P4 monomers adopt similar orientations and can be divided into three domains: an N-terminal region (110–150 residues), a central core NTPase domain of ~160 residues and a smaller C-terminal domain (~40–50 residues) (blue, grey and red, respectively, in Figure 2). Strikingly, despite low overall sequence conservation ranging from 9 to 21% amino acid sequence identity, the key structural features of the ATPase core domain (motifs H1, H1a, H2, H3 and H4) are well-conserved (Figure 3A andB). The ATPase domain is a Rossman-type nucleotide-binding domain consisting of a twisted seven-stranded β-sheet with
Figure 2. The overall fold of cystoviral P4 proteins. (A) The P4 hexamers of bacteriophages φ6, φ8, φ12 and φ13 (left to right) are viewed from the top and coloured by chain. (B) Side view of the P4 hexamers. (C) The panel shows structures of monomeric P4 in two orientations, the upper orientation of the monomer corresponding to the one depicted in cyan in (B); the lower one has undergone a rotation of 140° to show the C-terminal domains. The core domain is coloured in grey, the N-terminal domain in blue and the C-terminal domain in red. Nucleotides, if present, are depicted as sticks with carbon, oxygen, nitrogen and phosphorus atoms coloured in yellow, red, blue and orange, respectively. Dotted lines represent the disordered region of the proteins.
Figure 3. Structural conservation between P4 proteins. (A and B) Sequence and structural conservation of the helicase motifs in P4 proteins. Motifs H1, H1a and H2 are involved in nucleotide binding and hydrolysis, H3 is involved in the coupling of NTP hydrolysis to nucleic acid translocation, and H4 in oligonucleotide binding. Motifs H1, H1a, H2, H3, H4 are coloured in red, yellow, green, blue and brown, respectively; the arginine fingers are coloured purple, whereas the L1 and L2 loops are black and cyan, respectively. (A) Structure-based acid sequence alignment of the ATPase core domain of $\phi 6$, $\phi 8$, $\phi 12$ and $\phi 13$ P4. Functionally important residues that are conserved amongst the different cystoviruses are indicated by stars, whereas a sphere marks the lysine in loop L2 (K241 in $\phi 12$ P4), which is not conserved in $\phi 8$ P4. (B) Cartoon representations of $\phi 6$, $\phi 8$, $\phi 12$ and $\phi 13$ P4 structures in equivalent orientations. The arginine fingers and the nucleotides are shown in a ball-and-stick representation. The colour coding is the same as in (A). (C) Topology diagrams of the N-terminal domains of $\phi 6$, $\phi 8$, $\phi 12$ and $\phi 13$ P4. Secondary structural elements are coloured in green (strands) and yellow (helices). Topologically similar domains are shaded in pink ($\phi 6$ and $\phi 8$) and orange ($\phi 8$ and C2). The topology for C2 was derived from PDB entry 2ENP.
mixed parallel and antiparallel topology flanked by five helices. Residues previously demonstrated to be critically important in the mechano-chemically coupling of ATP hydrolysis to RNA translocation in Pφ24 P4 (35) are structurally conserved in other Pφ4s (Figure 3A and B, Table 2), except for one residue in motif H4 (residue K241 in Pφ24 P4), which has no equivalent in Pφ8 P4 (see explanation for this later in the text). It is therefore likely that all cystoviral Pφ4 NTPases use an RNA translocation mechanism similar to that described for Pφ24 P4 (31), although details may vary, especially for Pφ8 P4 where a tight coupling between ATPase activity and RNA binding is observed (Table 1).

Structural classification based on the ATPase core structures shows that cystovirus Pφ4 proteins are closely related to each other and only distantly related to other P-loop ATPases (Figure 4 and Supplementary Figure S2). They most closely resemble RecA-type ATPases (35), such as ATP synthase-like proteins (RHO, F1-ATPase, etc.), RecA-like proteins (RepA, TP4, etc.) and Rad51-like protein (Rad51, RecA, etc.). Many of these proteins are involved in nucleotide repair and recombination and have similar functional properties to Pφ4 proteins. This indicates that the cystoviral Pφ4 proteins form a distinct subfamily of RecA-type ATPases.

**N-terminal domain**

The structural conservation across Pφ4 proteins of the central ATPase core domains does not extend to the N- and C-terminal domains. Most of the N-terminal domain residues of Pφ4 from Pφ6 and Pφ8 are visible in our crystal structures (starting from amino acid residues 2 and 12, respectively), whereas Pφ13 Pφ4 lacks the first 32 residues [which are predicted to be disordered (60)]. In all Pφ4 structures, the N-terminal domain covers the apical part of the hexamer (Figure 2), and in Pφ12 Pφ4, an N-terminal domain α-helix projects from one subunit to the adjacent one, giving the hexamer a more rounded appearance. Pφ6 Pφ4 lacks such a helix and might stabilize the hexamer by strengthening subunit interfaces with nucleotides. Pφ6 Pφ4 is the only Pφ4 that needs nucleotides and divergent cations to form hexamers (7). It is also conceivable that NTP binding triggers a conformational change in the Pφ6 Pφ4 subunits allowing them to form hexamers. Interestingly, Pφ8 and Pφ13 Pφ4 also lack such a stabilizing helix; however, the first 12 and 31 residues, respectively, are not visible in the crystal structures and might play such a stabilizing role.

The N-terminal domains of cystoviral Pφ4s are highly divergent (Figures 2, 3B and C). However in Pφ6 and Pφ13, more than half of their residues can be superimposed with a root-mean-square deviation of 2.1 Å, including two parallel helices and two small anti-parallel β-sheets, creating a topologically identical sub-domain (Figure 3C). In Pφ8 and Pφ12, the N-terminal domains have higher secondary structure content but are completely unrelated to each other and to those in Pφ6 and Pφ13. In Pφ12 Pφ4, the N-terminal domain is composed of two orthogonal α-helices and three anti-parallel β-sheets (Figure 3C). The Pφ8 Pφ4 N-terminal domain is composed of two helices separated by a four-stranded antiparallel β-sheet (Figure 3C). Structural alignment searches against the PDB database returned no significant matches for any of the N-terminal domains, aside from a weak structural similarity (43 of 87 residues within 3.7 Å) of Pφ8 Pφ4 to one half of a C2 domain (domain involved in targeting proteins to cell membranes; Figure 3C). Intriguingly, Pφ8 lacks the Pφ8 nucleocapsid protein layer present in other cystoviruses so that Pφ4 proteins (together with P1 shell) interact directly with the viral lipid membrane (10).

**C-terminal domain**

The C-terminal domain of Pφ4 comprises ~40–50 amino acid residues downstream of the ATPase core (Figure 2) expected to be located at the bottom of the hexamer and to be essential for binding to the capsid protein P1 (38,61). The C-terminal domains of Pφ4 proteins diverge substantially. In Pφ6 and Pφ13, the C-termini are predicted to be disordered with little secondary structure (60), and indeed, no density for these domains could be found in our crystal structures. In contrast, the corresponding regions in Pφ8 and Pφ12 are predicted to be mostly ordered (60) with a C-terminal helix preceded by a flexible loop. In Pφ4 Pφ12, the strand following the arginine finger motifs extends back into the ATP-binding site contributing two residues (Y288 and S292), which help position the nucleotide ring (see later in the text). The density for the amino acid chain then disappears to re-emerge into a C-terminal helix stacked at the bottom of the hexamer (Figure 2). In Pφ4 Pφ8, the strand following the arginine fingers motifs does not extend as far as the ATP-binding site but instead climbs back along the side of the hexamer (partially

### Table 2. Conserved residues and their function within Pφ6, Pφ8, Pφ12 and Pφ13 Pφ4 proteins

| Amino acid          | Pφ6 | Pφ8 | Pφ12 | Pφ13 | Function                  | Walker motif |
|---------------------|-----|-----|------|------|---------------------------|--------------|
| Lysine/Threonine    | K132| K116| K136 | K159 | Phosphate binding         | H1           |
| Glutamate           | E150| E141| E160 | E176 | Catalytic base             | H1a          |
| Asparagine          | D187| D171| D189 | D213 | Coordinate Mg              | H2           |
| Lysine              | K239| K185| K241 | K265 | RNA binding                | H4           |
| Arginine            | R268| R263| R272 | R294 | Arginine finger            |              |
| Glutamine           |     | Q278|      |      | Base stacking              |              |
| Arginine            | R273| R266| R279 | R299 | Arginine finger            |              |
| Tyrosine/Phenylalanine| F275| F247| F247 | F288 | Base stacking              |              |

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Figure 4. Structure-based phylogenetic tree of ATPase enzymes. The matrix of evolutionary distances was calculated with SHP (56). The rectangle corresponds to a close-up view of the members of the RecA family. Abbreviations (in alphabetical order; Protein Data Bank accession codes are quoted in brackets): AfGspE, archaeal secretion ATPase, (2Oap); CFTR, Cystic Fibrosis Transmembrane Conductance Regulation, (1Xmi); Clamp Loader, eukaryotic clamp loader, (1Sxj); CobA, corrinoid adenosyltransferase, (1G64); CobU, adenosylcobinamide kinase/adenosylcobinamide phosphate guanylyltransferase, (1Cbu); DMC1, meiotic recombination protein, (2Zjh); DnaB, Thermus aquaticus DnaB, (2Q6t); Elp4, elongator complex protein 4, (4A8j); ESCN, prototypical T3ss ATPase EscN, (2Obl); F1-ATP Synthase-a, ATP synthase subunit-a, heart isoform, (2Jj1); F1-ATPase-β, bovine mitochondrial F1-ATPase, (1E1r); FbpC, Fe(3+) ions import ATP-binding protein FbpC, (3Fvq); FtsK, DNA translocase FtsK, (2Jut); G40P, ATPase domain of G40P, (3Bh0); Get3, ATPase Get3, (3Sja); GskA, Geobacillus kaustophilus DnaC, (2Vyf); GsDnaB, Geobacillus steinthermophilus DnaB, (2R6c); IoID, Aquifex aeolicus ABC transporter, (2Pcj); KaiC, Circadian clock protein kinase KaiC, (3Koe); MalK, maltose/maltodextrin import ATP-binding protein, (2Awn); MipZ, bacterial cell division regulator protein MipZ, (2Xit); MMAA, methylmalonic aciduria type A protein, (2Www); Msb8, Thermotoga maritima Abc transporter ATPp-binding protein, (1Vpl); MutS, DNA mismatch repair protein MutS, (1Ewq); P-gp, multidrug resistance protein Pgp-1, (4F4c); PH0273, Protein PH0273 Protein, (2Dr3); PiT, twitching motility protein PiT, (2Gs); Psy3, Platinum sensitivity protein 3, (4D1t); Rad50, DNA Double-Strand Break Repair Rad50 ATPase, (3Qku); Rad51, DNA repair protein Rad51, (1Ssp); RadA, DNA repair and recombination protein RadA, (4Dc9); RecA, Recombinase A, (1Mo4); RepA, regulatory protein RepA, (1G8s); Rho, transcription termination factor Rho, (3Jie); Rli1p, translation initiation factor, (3Jle); RNT1, regulator of nonsense transcripts 1, (2Wjy); SMC, chromosome partition protein, (4F9h); Sso2452, putative uncharacterized protein, (2W0m); T7Gp4, T7 DNA Primase/Helicase, (1Cr1); TK, thymidine kinase, (2Ja); TrwB, conjugal transfer protein TrwB, (1E9r); V1-ATPase, V-Type sodium ATPase, (3VR4); VirB4, type IV secretion pathway VirB4 components-like protein, (4Ag6); Vps4, vacuolar protein sorting-associated protein 4, (3Eih); XDP, Xpd/Rad3 related DNA helicase, (3Crv).
disordered) to re-emerge into as C-terminal helix at the top of the hexamer (Figure 2B), followed by a loop that dives into the central channel restricting its diameter by more than half (see later in the text for more discussion on the C-terminal domain).

Nucleotide binding site

The φ6 P4 was crystallised with ADP-Mg\(^{2+}\) bound in the nucleotide binding site, whereas P4 from φ8 and φ13 were crystallized in their apo form. As for φ12 P4, and other hexameric NTPases, the nucleotide binding sites in φ6 P4 are located at the interfaces between neighbouring subunits. The ADP phosphate groups are bound via the conserved Walker A (H1) motif residues (K132, S133) (Figure 5); a conserved glutamate E150 (H1a) is positioned to catalyse the nucleophilic attack on the γ-phosphate, whereas D187, a conserved aspartate in the Walker B motif (H2), co-ordinates the magnesium ion. A sensor motif detecting the presence or absence of the γ-phosphate of NTP and modulating allosteric transitions of the RNA binding loop L2 in response to ATP binding and hydrolysis was identified in P4 from φ12 (N234) (31). The equivalent residue in φ6 P4, N232, is positioned to contact the γ-phosphate of the NTP (Figure 5) and might fulfil the same role. As the mechanism of NTP binding and hydrolysis is similar, it is likely that the equivalent conserved residues in P4 from φ8 and φ13 (Figure 5 and Table 2) play analogous roles.

It has been shown that φ12 P4 possesses two essential ‘arginine fingers’ (35). We find that all P4 proteins follow this unusual pattern (Figure 5 and Table 2). Arginine fingers can contact the γ-phosphate of the triphosphate from a neighbouring subunit, and the insertion of this residue in a catalytic site is believed to stabilize the transition state, thus facilitating ATP

Figure 5. Cartoon representation of the nucleotide binding sites of φ6 (A), φ8 (B), φ12 (C) and φ13 (D) P4s. Within hexamers, adjacent monomers are coloured in yellow and grey. Nucleotides (ADP), if present, are depicted as sticks with carbon atoms coloured in green. Oxygen, nitrogen and phosphorus atoms are coloured in red, blue and orange, respectively, and the position of Mg\(^{2+}\) (φ12 P4) or Ca\(^{2+}\) (φ6 P4) is indicated with a cyan sphere.
hydrolysis. Arginine fingers in P4 proteins are all contributed from the same region (a loop between two strands in the C-terminal region) but display different conformations (Figure 5). In P4 from φ6, φ12 and φ13, the arginine fingers are pointing towards the catalytic sites, making the subunits competent and primed for hydrolysis. However, in φ8 P4, these residues are displaced >8 Å from that position and therefore cannot contribute to catalysis. This suggests that in φ8 P4, extensive conformational changes occur as a consequence of nucleotide and/or oligonucleotide binding, which render the enzyme competent for catalysis. Indeed, nucleotide binding kinetics revealed a first-order rate limiting step, which is consistent with a conformational change associated with ATP binding (39,62).

In RecA-like ATPases, bound nucleotides are stabilized by stacking of the adenine moiety between side chains, but these side chains are not conserved and are contributed from different regions. In RepA and T7 helicases, the ATP base stacks against residues belonging to the subunit carrying the catalytic site. In φ12 P4 (31), as in RepA (63), the nucleotide base is sandwiched between Y288 from the catalytic subunit and Q278 from the neighbouring subunit. In φ6 P4, a much looser stacking of the nucleotide base is observed, with only one side chain (F275) stabilizing the adenine ring (Figure 5). From our structures, we predict similar loose arrangements in P4 from φ8 and φ13 where F247 (from the same subunit) and F301 (from a neighbouring subunit) seem to be in the correct orientation to stack the nucleotide base. The difference in the arrangement of the nucleotide binding motifs is likely to explain the mechanism of base-specific hydrolysis in different P4s. Of the P4s, only φ12 is purine specific, with pyrimidines also being accepted by φ16, φ18 and φ13 (Table 1).

To understand this catalytic mechanism in detail, we performed side-directed mutagenesis of the residues in φ12 P4 involved in binding the nucleotide ring and analysed the mutants structurally and biochemically. In φ12 P4, the stacking interaction is critical for nucleotide binding, as replacement of the tyrosine with alanine (Y288A) completely abolished ATP binding and ATPase activity (Table 1) so that the apoprotein structure is found even in the presence of ATP (data not shown). However, the mutation Q278A had only a moderate effect on ATPase activity and virtually no effect on the structure of the bound ATP analogue AMPcPP when compared with the wild-type (Figure 6A and C), primarily increasing the Km as a result of reduced nucleotide affinity (Table 1). Hence, the stacking interactions primarily determine nucleotide affinity but not specificity. A specific feature in φ12 P4 is a hydrogen bond between the hydroxyl of S292 and N7 of the purine ring. The substitution S292A did not prevent ATP binding but completely abolished ATPase activity owing to misplacement of the triphosphate moiety in the active site (Figure 6D). A displacement is also seen when the AMPcPP bound wild-type structure is compared with that of UTP bound hexamer (Figure 6A and B). This confirms that pyrimidine triphosphates can bind the hexamer without being hydrolysed (36) and should act as competitive inhibitors. Indeed, we find that UTP effectively competes with ATP and inhibits hydrolysis (data not shown). Hence, purine specificity is achieved by locking the base by hydrogen bonding to the N7 site of a purine. The correct coordination of the base results in the precise alignment of the nucleotide that is essential for catalysis so that UTP is misaligned and not hydrolysed. This is probably the mechanism underpinning the dependence of helicase efficiency on the type of nucleotide. For example, T7 gp4 helicase activity is optimal in presence of dTTP (58).

### Nucleic acid binding site

It has been proposed that P4 hexamers bind nucleic acid through their central channel via two protruding loops named L1 and L2 (31) (Figure 3A and B, Supplementary Figure S3). Mutagenesis studies confirmed that these loops are essential for nucleic acid binding and translocation (30,35,37). Structurally homologous loops were reported to bind ssDNA and ssRNA, respectively, in crystals of E. coli helicase of bovine papilloma virus and Rho of E. coli (59). The L1 loops in P4 are rich in residues that contribute to flexibility (in φ12 P4 they are disordered), whereas the L2 loops are mainly composed of hydrophilic residues, amongst them a lysine, which in φ12 P4 (K241) was shown to be essential for RNA binding (35). The structures of P4 from φ6 and φ13 show ordered L1 loops, which line the central channel and contact the L2 loops (Supplementary Figure S2). The L2 loops are found with lysine residues (K239 and K265, respectively) projecting towards the centre of the channel, in the same position as K241 in φ12, suggesting a conserved mechanism for binding and translocating RNA. Although the L2 loop of φ8 P4 contains hydrophilic residues (DDENVD), it does not project a lysine side chain towards the central channel. Nevertheless, the L1 loop contains a motif (LKK) that has been shown to be crucial for RNA binding (35). The first lysine of this motif (K185) is found in the equivalent position to K241 of φ12 P4 and is also seen interacting with D220 of loop L2. We therefore postulate that K185 (loop L1) in φ8 P4 plays the same role in RNA binding as K241 (loop L2) in φ12 P4, and that the coupling of the movement of the L1 and L2 loops to ATP hydrolysis via motion of helix 6, as proposed for φ12, may be a general feature of all P4 molecules (Supplementary Figure S2). The importance of the L1 loop is further supported by mutational analysis in φ12 P4: deleting L1 loop central residues T202-T203-S204 or mutating them into the equivalent residues of φ8 P4 (LKK) completely abolishes the ATPase activity (Table 1). This demonstrates that the integrity of the L1 loop is essential for ATP hydrolysis, despite being distal to the ATP active site.

### RNA loading in φ8 P4 and the structural basis of processive translocation

The φ8 P4 ATPase activity is tightly coupled to ssRNA translocation, as it will only hydrolyse ATP in the presence of ssRNA. As noted earlier in the text, the RNA binding motif LKK in loop L1 is located in the middle of the central channel (37). Nucleic acids are likely to bind in
the channel, ensuring topological enclosure of the strand and processive translocation.

Based on transient cooperative exposure of subunit interfaces to HDX on RNA binding (residues 198–209 in Figure 7), it was suggested that RNA enters the central channel via a transient ring opening (37). The deletion of the C-terminal portion of the protein (residues 282–321) more than doubles the diameter of the central channel (from 13 to 30 Å), as the C-terminus wraps upwards from the base of the hexamer, along the inter-subunit cleft, to stick down into the central channel (Figure 8). As the C-terminal domain is (i) necessary for ATP hydrolysis (data not shown), (ii) restricts the diameter of the central channel and (iii) blocks the interface through which RNA is thought to be loaded, we postulate that the C-terminal region needs to be displaced by RNA for ring opening and subsequent ATP hydrolysis to occur. To verify this hypothesis, previous HDX experiments (37) were further analysed by mapped to the φ8 P4 structure.

The C-terminal region exhibits the fastest HDX within the protein (Figure 7). However, the distal C-terminal portion that extends into the central channel is marginally protected in the absence of RNA and becomes fully exposed only on addition of RNA, implying that this region becomes further exposed presumably by expulsion from the central channel (Figure 8B). Thus, it appears that φ8 P4 has developed a specific mechanism to regulate ATPase activity and couple it with ssRNA binding such that RNA displaces the C-terminal domain, to allow ATP hydrolysis to occur. This would explain the tight coupling observed between ATP hydrolysis and translocation.

**Figure 6.** Cartoon representation of the nucleotide binding site of φ12 P4. (A) Wild-type φ12 P4 bound to non-hydrolysable ATP analogue AMPcPP (PDB: 1W48) or (B) UTP. (C) Q278A mutant bound to AMPcPP. (D) S292A mutant bound to AMPcPP. Within hexamers, adjacent monomers are coloured in yellow and grey. AMPcPP bound to wild-type φ12 P4 is depicted in sticks, and the carbon atoms are coloured cyan (A), whereas carbon atoms in the UTP bound to φ12 P4 (B) and AMPcPP bound to the P4 mutants Q278A and S292A (C and D) are coloured in green. Oxygen, nitrogen and phosphorus atoms are coloured in red, blue and orange, respectively. (B–D) The position of the AMPcPP bound to wild-type P4 is represented in transparent for comparison.
CONCLUSION

The current study broadens our understanding of the mechanism used by dsRNA bacterial viruses to package RNA genome during assembly. Interestingly, P4 proteins are only remotely related to packaging ATPases of dsDNA viruses such as gp17 from bacteriophage T4 (64) or pUL15 from Herpes Simplex virus 1 (65), which have more complicated portal complexes. Recently, however, it has been suggested that the ATPase of the phi29 DNA packaging motor is a member of the hexameric AAA+ superfamily (66), indicating that the mechanism of nucleic acid packaging might be similar.

A structure-based phylogeny (Figure 4) suggests that the RecA-like proteins may be the closest cellular relatives of the P4, with φ12 being the most similar to the cellular proteins, φ8 being rather divergent and φ6 and φ13 rather similar to each other and intermediate in terms of divergence from the cellular proteins. These structural variations map onto the various functional specializations of the molecules so that although the motors have a common catalytic mechanism, they have developed somewhat different specificity and control mechanisms. We identify a specific hydrogen bond (serine 292 and N7 of the purine ring) responsible for the purine specificity of φ12 P4 catalysed NTP hydrolysis reaction and find that an extraordinary insertion of the C-terminal peptide into the central channel of the hexamer explains the tight coupling of ATPase activity and RNA translocation in φ8. Furthermore, the φ8 P4 structure revealed a novel

![Figure 7](http://nar.oxfordjournals.org/)

Mapping of HDX data on the φ8 P4 structure. HDX rates are coloured from slow-exchange (blue) to fast-exchange rates (red). Previously measured HDX rates (53) for φ8 P4 in the presence/absence of AMP, ADP, ATP and RNA (as indicated) were mapped onto the φ8 P4 monomer structure. The central box shows on the left, the orientation of all the monomers of the figure within the hexamer, and on the right, the same monomer in which the N- and C-terminal domains are coloured in blue and red, respectively.
mechanism of power transduction to the RNA in which RNA is engaged with the L1 loop, which, in turn, is coupled to the L2 loop. Comparison between the P4 structures suggest that coupling between the two loops may be a general mechanistic feature of P4 and perhaps other SF4 helicases. Overall, the P4 machine represents a remarkable test bed where, by virtue of high mutational rates over long periods of time, nature has been able to devise a range of functional variations on the basic theme of regulated RNA translocation, resulting in an array of systems where although the molecular engine remains largely similar, the ignition and transmission systems have diverged markedly.

ACCESSION NUMBERS
Coordinates and structure factors of ADP-bound $\phi 6$ P4$\Delta 310$, $\phi 8$ P4, $\phi 8$ P4$\Delta 281$, UTP-bound $\phi 12$ P4, AMPcPP-bound $\phi 12$ P4-Q278A and AMPcPP-bound $\phi 12$ P4-S292A and $\phi 13$ P4 have been deposited in the Protein Data Bank under accession codes 4BLO, 4BWY, 4BLQ, 4BLR, 4BLS, 4BLT and 4BLP, respectively.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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