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The Structure of the Nucleoprotein Binding Domain of lyssavirus Phosphoprotein reveals a structural relationship between the N-RNA binding domains of Rhabdoviridae and Paramyxoviridae

Delmas Olivier¹, Rene Assenberg²,³, Jonathan M Grimes² and Hervé Bourhy¹*.

¹Institut Pasteur, Unit Lyssavirus Dynamics and Host Adaptation, WHO Collaborating Centre for Reference and Research on Rabies, 75724 Paris CEDEX 15, France

²Division of Structural Biology and Oxford Protein Production Facility, Wellcome Trust Centre for Human Genetics, University of Oxford, Roosevelt Drive, Oxford OX3 7BN, United Kingdom

³Current address: Center for Proteomic Chemistry/Expertise Platform Proteases, Novartis Institutes for BioMedical Research, Fabrikstrasse 16, CH-4002 Basel, Switzerland.

* Corresponding author:

Hervé Bourhy. E-mail: hbourhy@pasteur.fr, Phone: +33 1 45 68 87 85, Fax: +33 1 40 61 30 20

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Abstract

The phosphoprotein P of non-segmented negative-sense RNA viruses is an essential component of the replication and transcription complex and acts as a co-factor for the viral RNA-dependent RNA polymerase. P recruits the viral polymerase to the nucleoprotein-bound viral RNA (N-RNA) via an interaction between its C-terminal domain and the N-RNA complex. We have obtained the structure of the C-terminal domain of P of Mokola virus (MOKV), a lyssavirus that belongs to the Rhabdoviridae family and mapped at the amino acid level the crucial positions involved in interaction with N and in the formation of the viral replication complex. Comparison of the N-RNA binding domains of P solved to date suggests that the N-RNA binding domains are structurally conserved among paramyxoviruses and rhabdoviruses in spite of low sequence conservation. We also review the numerous other functions of this domain and more generally of the phosphoprotein.

Key words: Lyssavirus, Phosphoprotein, RNA-dependant RNA polymerase, nucleoprotein.
Introduction

Viruses belonging to the Mononegavirales order are enveloped viruses comprising four distinct families, the *Filoviridae*, *Bornaviridae*, *Rhabdoviridae* and *Paramyxoviridae*. The Mononegavirales infect numerous mammal hosts, and exhibit high genetic diversity and rapid evolutionary rates due to the lack of proofreading activity of the replication complex, therefore constituting a large variety of potential and actual human pathogens. Among them the *Paramyxoviridae* and *Rhabdoviridae* will be the main focus of this report as they include several highly pathogenic viruses like measles virus (a paramyxovirus) and rabies virus (a rhabdovirus), respectively. Measles virus kills more than 200,000 children in developing countries in which vaccination provides a significant decrease of paediatric mortality\(^1\). The measles virus has also been associated with severe respiratory presentations and with rare but extremely serious neurological complications. The agents of rabies, leading invariably to fatal viral encephalitis, all belong to the lyssavirus genus. They are responsible for an estimated 55,000 deaths per year worldwide\(^2\). Seven different species are recognized by the ICTV based on genome sequence homology. New viruses have been recently described, although not yet classified, that could represent new lyssavirus genotypes\(^3-5\).

The genome of Mononegavirales consists of a negative sense non-segmented RNA, which is encapsidated by the nucleoprotein N and encodes at least 5 viral proteins common to all viruses of this family, in the order from 3' to 5' : the nucleoprotein (N), the phosphoprotein (P), the matrix (M) and the large protein (L). The latter contains approximately 2000 amino acids and is the central protein in the replication and transcription of the viral genome, and carries RNA dependent RNA polymerase, RNA polyadenylation, RNA guanylyltransferase and RNA methyltransferase activities. After the first steps of attachment, penetration and uncoating of the virus particle, the nucleocapsid (the N-RNA complex) and all the components necessary for the early transcription are released in the cytoplasm of the infected
cell where the transcription and then the replication steps of the virus life cycle occur. During these processes the genome (and anti-genome) remains bound to the N polymer at all times and recently structures have been solved for N-RNA complexes from Rabies, VSV and respiratory syncytial virus (RSV) \(^6\)\(^8\). Importantly, during transcription the viral mRNAs are not encapsidated by N, probably due to the fact that leader and trailer sequences are absent from mRNA conversely to genomic and antigenomic RNA. Thus, the viral transcripts are, like their cellular counterparts, capped and poly-adenylated at their 5’ and 3’ ends respectively. One important unresolved question remains how the viral polymerase switches between replication and transcription, which both utilise the same genomic RNA strand. However, in the case of transcription the polymerase recognises specific RNA signals that demarcate the start and end points of the viral genes to produce a gradient of mRNA transcripts, whilst during replication these are ignored. The Mononegavirales have to deal with a limited number of proteins (5-10) to achieve all the necessary functions for their life cycle. As such the P exhibits a modular organization with each of its domains is involved in several aspects of the P "cellular functions" (Fig.1) The P protein is an essential cofactor of the viral RdRP although the precise role of P in facilitating polymerase activity remains to be established. It is thought that one function of P is to deliver, during replication, free N (referred to as “N-degree” or N\(^0\)) to the newly synthesised viral RNA. In addition, it is thought that P acts as a bridge between the L polymerase and the N-RNA template to gain access to the genomic RNA. Consistent with this idea is that P has two distinct N-binding sites, one specific for N\(^0\) (located at the N-terminus of P) and one for N-RNA (located in the C-terminal domain of P [PCDT]). The general domain organization of P is conserved across the Mononegavirales, although the details differ. In between the two N-binding domains, P contains a multimerisation domain, which in rhabdoviruses mediates P dimerisation \(^9\) whilst in paramyxoviruses it mediates tetramerisation \(^10\). The multimerisation domain is flanked by
disordered regions of variable length, probably explaining the difficulty of obtaining full length P crystals, as well as its susceptibility to in vitro proteolysis (see below). Interestingly, in paramyxoviruses, and in rhabdoviruses, the P gene is edited during transcription by L, yielding additional proteins termed C and V in paramyxovirus and shorter P in lyssavirus termed P2, P3, P4 and P5 \(^{11,12}\). These P products appear to function primarily to evade the host immune response.

In lyssavirus, the PCDT modulates the sub-cellular localization of P during infection. The PCTD have been shown to contain a nuclear localization signal (NLS) \(^{13}\) and a nuclear export signal (NES) \(^{14}\). The NLS involves the amino acids that also contribute to the interaction with N. The NES involves amino acids positions 227 to 232. These two signals contribute to the cellular localization of P and phosphorylation by protein kinase C may be involved in the switch mechanism to balance the import and export of P to and from the nucleus \(^{14}\). The association of P with microtubules also regulates its nuclear trafficking. The CTD of P contains motifs that participate in P association with microtubules but these associations are dependent on oligomerization of P \(^{15}\). In the nucleus, the PCTD also interacts with interferon-induced promyelocytic leukaemia (PML) bodies \(^{11}\). The exact role of this interaction remains to be fully understood but may be involved in virus pathogenesis.

The PCTD also plays a role to limit the capacity of defense of the hosts. It has been shown that P prevents interferon regulatory factor 3 (IRF3) phosphorylation thus inhibiting the key factor involved in IFN alpha/beta response in infected cells \(^{16}\). P is also able to target and interact with STAT1 and STAT 2 \(^{17-19}\). This interaction in combination with microtubule association prevents STAT 1 and STAT 2 nuclear localization \(^{15}\). The region of P involved in this interaction is also located within the PCTD between positions 267 and 297 \(^{18}\).

We have recently obtained the structure of the phosphoprotein C-terminal domain (PCTD) of one species of the lyssavirus genus, the Mokola virus (MOKV) and mapped at the amino acid
level the crucial positions involved in interaction with N involved in the viral replication complex formation $^{20}$. In this paper we will focus on the structural aspects of the C-terminal domain in relation to virus replication.
Results

Structure determination of the C-terminal domain of Mokola virus P

Full length MOKV P was recombinantly expressed in E. coli, purified and subjected to crystallization experiments. The structure was solved by molecular replacement, using the structure of the RABV P N-RNA binding domain as a model (PDB 1VYI), which revealed that a proteolytic fragment had been crystallized. Mass spectrometry analysis of the protein immediately before the crystallization trials showed the protein to be intact at that point confirming that truncation had occurred during crystallisation. The structure of MOKV P obtained was of the C-terminal domain of the protein, formed by six tightly folded α-helices, a 3_10-helix and a small β-sheet consisting of two β-strands (Fig. 2), with a morphology resembling a sliced pear, with a flat and curved face.

Analysis of the crystal structure of MOKV P revealed that one of the contacts points in crystal packing involves the C-terminal 4 residues (E300, E301, A302, E303) of one monomer packing against a predominantly positively charged region consisting of residues located on β1, β2, α4 and α6 (K212, K213, Y214, K215, I223, L225, R261 and Y295) in another monomer. K212, K213, K215, R261 and Y295 engage in both main- and side chain polar contacts with the acidic tail, with residues Y214, K215, I223, L225 and R261 contributing additional hydrophobic interactions.

This positively charged region on P has been implicated previously in the P-N interaction. To confirm and map precisely the relative contribution of each of these amino acids, we tested by yeast two hybrid the interaction of N with single point mutants of the P C-terminal domain. Mutations of Y214, K215 and R261 resulted in the most significant growth defects of the yeast strain whilst for K212 and L225 higher stringency conditions were necessary to reveal an effect on N binding. The degree of association and the reduction in accessible
surface area upon binding the acidic tail by each of the residues shows a good overall correspondence with the yeast two hybrid results.

One of the major roles of P is as part of the replication complex together with N and L. The P is thought to bridge between N (associated with the viral genome) and the L polymerase. This viral replication complex is active in the Negri bodies. Negri bodies are viral cytoplasmic inclusions that contain N, P and L and viral genome whose formation is dependent on Toll like receptor 3. The PCTD interacts with N alone or with N associated with RNA and the amino acids K212, Y214, K215, L225 and 261 have been implicated in this interaction. Whether these N-P interactions are involved in Negri bodies formation remains to be explored.

PCTD motif and structure conservation among Lyssaviruses

Lyssaviruses taxonomy is defined on the basis of genetic similarity of the nucleoprotein sequence. Seven species of lyssavirus are recognized by the ICTV (International Committee on Taxonomy of Viruses). Recently, four new lyssaviruses have been described from bats originated from Eurasia (Aravan virus (ARAV), Khujand virus (KHUV), Irkut virus (IRKV)) or from Caucasian region (West Caucasian Bat Lyssavirus (WCBV)). A virus related to LBV has also been described. Each of these new viruses could represent new species.

The lyssavirus P protein, including those from newly described viruses show a percentage of sequence identity ranging from 42 % (minimum intergenotype identity) to 81% (minimum intragenotype identity). The PCTD is known to be the most conserved part of the protein. The structure of MOKV PCDT is very similar to the PCDT solved previously of RABV P, (another species of lyssavirus) (PDB ID 1VYI) (Fig. 2B) where 101 amino acids can be aligned with a root mean square deviation (rmsd) of 0.66 Å, consistent with the high degree of sequence identity (68%). There are only minor changes in the length of helixes at the N and C
termini of the domain, with a shorter $\alpha_1$ helix but an extended $\alpha_6$ helix in MOKV compared to RABV.

All the amino acids positions which were implicated in P-N association are conserved in all lyssaviruses further reinforcing the probable role of this charged interaction surface of the P in the virus life cycle (Fig 3).

**A conserved organization of P proteins in the families of Rhabdoviridae and Paramyxoviridae**

The structures of functionally equivalent domains of the P proteins from several other members of the order Mononegavirales have been solved. The structures of the N-RNA binding domain of Sendai virus (SeV) and Measles virus (MeV) P protein consist of an anti-parallel triple helix bundle and, as for the Rhabdoviridae, the N-RNA binding domains among paramyxoviruses are also structurally conserved in spite of low sequence conservation. Careful analysis of the structures reveals a structural relationship between the N-RNA binding domains of the P proteins of the Rhabdoviridae and the Paramyxoviridae, where the 3 helix bundle of SeV and MeV P aligns with helices $\alpha_3$, $\alpha_4$ and $\alpha_6$ of MOKV (and RABV) (Fig. 4). Structural superposition using SHP shows that 43 and 40 residues of MOKV P (out of 107) can be aligned with SeV P and MeV P, with rmsds of 3.3 Å and 3.2 Å respectively over the aligned residues, despite 14% and 0% sequence identity. The similarity between VSV P and the paramyxovirus P proteins is much poorer due to the shorter C-terminal helix in VSV P ($\alpha_4$ in VSV, $\alpha_6$ in MOKP) and the absence of a helix equivalent to MOKV $\alpha_3$ (Fig 4), although the alignment does suggest that VSV P $\alpha_3$ is conserved ($\alpha_4$ in MOKV P).

The structure based sequence alignment reveals that the highest degree of sequence similarity resides in the region spanning 7 residues in helix $\alpha_6$ in MOKV P (equivalent to helix $\alpha_3$ in
SeV and MeV P) (Fig. 4). Although the alignment indicates that MeV P shows a much poorer sequence conservation in this area, SeV also resembles lyssavirus P in the character of the P-N interactions which are primarily electrostatic (whereas in MeV P with N the interactions are primarily hydrophobic).

This alignment is in agreement with functional data where the structurally equivalent helix in the rhabdo- and paramyxoviruses (helix $\alpha_3$) to MOKV P $\alpha_6$ is involved in binding N. For MeV P a co-crystal structure was solved with the N-RNA binding domain of P bound to a MeV N segment (spanning one $\alpha$-helix) as shown in Fig. 4. The latter is bound to the topologically equivalent site in MOKV P (the positive cluster) as shown in the superposition of MeV P bound to N with MOKV P in Fig. 4. Clearly however the precise details of the interactions between P and N differ. For instance, none of the MOKV P residues identified in this study as important for the interaction with MOKV N, appear to be conserved in the VSV P structure, suggesting that VSV P interacts differently with N than the lyssavirus P. In addition, biochemical data also indicate that P-N and P-L interactions are likely to be different. This has been confirmed by the structure of VSV N-RNA complexed with the C-terminal domain of VSV P which reveals that those residues of VSV-P that are involved in binding lie close to but do not map directly onto the residues that we predict are involved in MOK N-P binding. This lack of agreement is probably due primarily to differences in the size of the P proteins (the CTD of MOK P is 50% larger than the equivalent CTD of VSV).

Thus, although clearly the interactions and mode of binding has diverged between the different members of the Mononegavirales, it seems likely that the P proteins of Rhabdo- and Paramyxoviridae originated from a common ancestor in spite of the high degree of amino acid divergence.

A recent analysis of structure and sequence conservation of P of two rhabdoviruses and one paramyxovirus combined the results from multiple prediction methods as well as two-hybrid
and biochemical approaches to locate the boundaries between disordered regions and structured domains and to characterize their potential functions. The proposed modular organization predicted for RABV P confirmed both our and earlier findings that the putative structured domains correspond to autonomous folding units, and that CTD of P is central to the formations of N-P complexes.

**Conclusion**

The phosphoprotein of the rhabdoviruses has a modular multidomain architecture and is central to the formation of an active RNP. The C-terminal domain of lyssavirus P has been shown to be multifunctional, and structural and biochemical analysis has implicated binding surfaces involved in N-P complex formation. P is found both in the cytoplasm and nucleus of infected cells and this compartmentalization defines to some degree the function of P. Thus P in the cytoplasm either forms part of the active viral polymerase complex or sequesters free N, and it has been shown that P oligomerisation is required for polymerase function. This oligomerisation may be required for the correct presentation of binding surfaces of P to N or L. It is unclear what the oligomerisation state of P is in the nucleus, but its role in inhibiting the IFN response is dependent on its nuclear import and retention.

What role if any structural switching of P has on modulating these different functions remains unclear but further steps in understanding the regulation of these functions will be to obtain the structure of P in complex with its partners; as recently obtained for VSV PCTD in association with N-RNA rings. What is clear from our structure and the analysis of those amino acids crucial to N-P association is that they are different from those observed in the VSV N-P complex structure. Whether this is due to differences in the mode of association or to structural changes to P when in complex with N-RNA remains an open question, but this lack of structural correlation is probably due partly to differences in the size of the P proteins.
Our structural analysis of the N-RNA binding domains of the P proteins from the Rhabdo- and Paramyxoviridae reveals a low but detectable degree of structural similarity as well as the conservation of the binding surfaces that suggest that these P proteins originated from a common ancestor in spite of the high degree of amino acid divergence.

Obtaining the structure of complexes of P with other proteins involved in the replication complex will help in our understanding of how so many functions could be retained and regulated by this viral protein. It will also allow us to understand the viral RNA replication / transcription complex and model how the viral polymerase accesses the viral RNA genome that is totally covered by the N protein. Understanding this process would help in the design of specific antiviral compounds that could target functional interaction surface on viral proteins and could help in preventing rabies and more widely underpin the treatment of viruses belonging to the Order Mononegavirales.

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Figure Legend

Fig1: Schematic representation of lyssavirus phosphoprotein and its functions in virus pathogenesis/host interplay and virus replication. Abbreviations: PML promyelocytic leukaemia proteins NLS, nuclear localization signal; NES, nuclear export signal, NTD aminoterminal domain, CED central domain, CTD carboxy terminal domain.

Fig2: Structure of MOKV and RABV PCTD. Side-by-side comparison of rhabdovirus P structures: (A) MOKV P; (B) RABV P (PDB accession no. 1VYI); (C) VSV P (PDB accession no. 2K47). The structures are rainbow ramped from blue (N terminus) to red (C terminus).

Fig3: PCTD alignment of all lyssavirus genotypes. Arrowheads show conserved aminoacids involved in nucleoprotein binding. Homology plot of each position is indicated in the bottom.

Fig4: Comparison of the structures of Rhabdovirus and Paramyxovirus N-RNA binding domains of P. (a) Side-by-side comparison of the superposed conserved elements of MOKV P (orange), VSV P (yellow) (PDB accession no. 2K47), and SeV (magenta) (PDB accession no. 1R4G), based on SHP alignment of the Ca’s (40) (b) Side-by-side comparison of MOKV P (orange) and MeV P bound to MeV N (cyan, PDB Id 1T6O), (based on the superposition from SHP) highlighting the overlap between the N-binding site in the two P proteins.